Fundamentals of Geophysics, second edition

This second edition of Fundamentals of Geophysics has been completely revised and updated, and is the ideal geophysics textbook for undergraduate students of geoscience with only an introductory level of knowledge in physics and mathematics.

Presenting a comprehensive overview of the fundamental principles of each major branch of geophysics (gravity, seismology, geochronology, thermodynamics, geoelectricity and geomagnetism), this text also considers geophysics within the wider context of plate tectonics, geodynamics and planetary science. Basic principles are explained with the aid of numerous figures and important geophysical results are illustrated with examples from the scientific literature. Step-by-step mathematical treatments are given where necessary, allowing students to follow the derivations easily. Text-boxes highlight topics of interest for more advanced students.

Each chapter contains a short historical summary and ends with a reading list that directs students to a range of simpler, alternative or more advanced resources. This new edition also includes review questions to help evaluate the readers’ understanding of the topics covered and quantitative exercises at the end of each chapter. Solutions to the exercises are available to instructors.

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Fundamentals of Geophysics

Second edition

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CAMBRIDGE UNIVERSITY PRESS
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Preface to the second edition

In the ten years that have passed since the publication of the first edition of this textbook exciting advances have taken place in every discipline of geophysics. Computer-based improvements in technology have led the way, allowing more sophistication in the acquisition and processing of geophysical data. Advances in mass spectrometry have made it possible to analyze minute samples of matter in exquisite detail and have contributed to an improved understanding of the origin of our planet and the evolution of the solar system. Space research has led to better knowledge of the other planets in the solar system, and has revealed distant objects in orbit around the Sun, at least one of which may be a tenth planet. Satellite-based technology has provided more refined measurement of the gravity and magnetic fields of the Earth, and has enabled direct observation from space of minute surface changes related to volcanic and tectonic events. The structure, composition and dynamic behavior of the deep interior of the Earth have become better understood owing to refinements in seismic tomography. Fast computers and sophisticated algorithms have allowed scientists to construct plausible models of slow geodynamic behavior in the Earth’s mantle and core, and to elucidate the processes giving rise to the Earth’s magnetic field. The application of advanced computer analysis in high-resolution seismic reflection and ground-penetrating radar investigations has made it possible to describe subtle features of environmental interest in near-surface structures. Rock magnetic techniques applied to sediments have helped us to understand slow natural processes as well as more rapid anthropological changes that affect our environment, and to evaluate climates in the distant geological past. Climatic history in the more recent past can now be deduced from the analysis of temperature in boreholes.

Although the many advances in geophysical research depend strongly on the aid of computer science, the fundamental principles of geophysical methods remain the same; they constitute the foundation on which progress is based. In revising this textbook, I have heeded the advice of teachers who have used it and who recommended that I change as little as possible and only as much as necessary (to paraphrase medical advice on the use of medication). The reviews of the first edition, the feedback from numerous students and teachers, and the advice of friends and colleagues helped me greatly in deciding what to do.

The structure of the book has been changed slightly compared to the first edition. The final chapter on geodynamics has been removed and its contents integrated into the earlier chapters, where they fit better. Text-boxes have been introduced to handle material that merited further explanation, or more extensive treatment than seemed appropriate for the body of the text. Two appendices have been added to handle more adequately the three-dimensional wave equation and the cooling of a half-space, respectively. At the end of each chapter is a list of review questions that should help students to evaluate their knowledge of what they have read. Each chapter is also accompanied by a set of exercises. They are intended to provide practice in handling some of the numerical aspects of the topics discussed.
in the chapter. They should help the student to become more familiar with geophysical techniques and to develop a better understanding of the fundamental principles.

The first edition was mostly free of errata, in large measure because of the patient, accurate and meticulous proofreading by my wife Marcia, whom I sincerely thank. Some mistakes still occurred, mostly in the more than 350 equations, and were spotted and communicated to me by colleagues and students in time to be corrected in the second printing of the first edition. Regarding the students, this did not improve (or harm) their grades, but I was impressed and pleased that they were reading the book so carefully. Among the colleagues, I especially thank Bob Carmichael for painstakingly listing many corrections and Ray Brown for posing important questions. Constructive criticisms and useful suggestions for additions and changes to the individual revised chapters in this edition were made by Mark Bukowinski, Clark Wilson, Doug Christensen, Jim Dewey, Henry Pollack, Ladislaus Rybach, Chris Heinrich, Hans-Ruedi Maurer and Mike Fuller. I am very grateful to these colleagues for the time they expended and their unselfish efforts to help me. If errors persist in this edition, it is not their fault but due to my negligence.

The publisher of this textbook, Cambridge University Press, is a not-for-profit charitable institution. One of their activities is to promote academic literature in the “third world.” With my agreement, they decided to publish a separate low-cost version of the first edition, for sale only in developing countries. This version accounted for about one-third of the sales of the first edition. As a result, earth science students in developing countries could be helped in their studies of geophysics; several sent me appreciative messages, which I treasure.

The bulk of this edition has been written following my retirement two years ago, after 30 years as professor of geophysics at ETH Zürich. My new emeritus status should have provided lots of time for the project, but somehow it took longer than I expected. My wife Marcia exhibited her usual forbearance and understanding for my obsession. I thank her for her support, encouragement and practical suggestions, which have been as important for this as for the first edition. This edition is dedicated to her, as well as to my late parents.

William Lowrie
Zürich
August, 2006
Acknowledgements

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1 The Earth as a planet

1.1 THE SOLAR SYSTEM

1.1.1 The discovery and description of the planets

To appreciate how impressive the night sky must have been to early man it is necessary today to go to a place remote from the distracting lights and pollution of urban centers. Viewed from the wilderness the firmaments appear to the naked eye as a canopy of shining points, fixed in space relative to each other. Early observers noted that the star pattern appeared to move regularly and used this as a basis for determining the timing of events. More than 3000 years ago, in about the thirteenth century BC, the year and month were combined in a working calendar by the Chinese, and about 350 BC the Chinese astronomer Shih Shen prepared a catalog of the positions of 800 stars. The ancient Greeks observed that several celestial bodies moved back and forth against this fixed background and called them the *planetes*, meaning “wanderers.” In addition to the Sun and Moon, the naked eye could discern the planets Mercury, Venus, Mars, Jupiter and Saturn.

Geometrical ideas were introduced into astronomy by the Greek philosopher Thales in the sixth century BC. This advance enabled the Greeks to develop astronomy to its highest point in the ancient world. Aristotle (384–322 BC) summarized the Greek work performed prior to his time and proposed a model of the universe with the Earth at its center. This *geocentric* model became imbedded in religious conviction and remained in authority until late into the Middle Ages. It did not go undisputed; Aristarchus of Samos (c.310–c.230 BC) determined the sizes and distances of the Sun and Moon relative to the Earth and proposed a *heliocentric* (sun-centered) cosmology. The methods of trigonometry developed by Hipparchus (190–120 BC) enabled the determination of astronomical distances by observation of the angular positions of celestial bodies. Ptolemy, a Greco-Egyptian astronomer in the second century AD, applied these methods to the known planets and was able to predict their motions with remarkable accuracy considering the primitiveness of available instrumentation.

Until the invention of the telescope in the early seventeenth century the main instrument used by astronomers for determining the positions and distances of heavenly bodies was the *astrolabe*. This device consisted of a disk of wood or metal with the circumference marked off in degrees. At its center was pivoted a movable pointer called the *alidade*. Angular distances could be determined by sighting on a body with the alidade and reading off its elevation from the graduated scale. The inventor of the astrolabe is not known, but it is often ascribed to Hipparchus (190–120 BC). It remained an important tool for navigators until the invention of the sextant in the eighteenth century.

The angular observations were converted into distances by applying the method of parallax. This is simply illustrated by the following example. Consider the planet P as viewed from the Earth at different positions in the latter’s orbit around the Sun (Fig. 1.1). For simplicity, treat planet P as a stationary object (i.e., disregard the planet’s orbital motion). The angle between a sighting on the planet and on a fixed star will appear to change because of the Earth’s orbital motion around the Sun. Let the measured extreme angles be $\theta_1$ and $\theta_2$ and the...
distance of the Earth from the Sun be \( s \); the distance between the extreme positions \( E \) and \( E' \) of the orbit is then \( 2s \). The distances \( p_1 \) and \( p_2 \) of the planet from the Earth are computed in terms of the Earth–Sun distance by applying the trigonometric law of sines:

\[
p_1 = 2s \frac{\sin(90^\circ - \theta_1)}{\sin(\theta_1 + \theta_2)} \quad \text{and} \quad p_2 = 2s \frac{\cos \theta_1}{\sin(\theta_1 + \theta_2)}
\]

Further trigonometric calculations give the distances of the planets from the Sun. The principle of parallax was also used to determine relative distances in the Aristotelian geocentric system, according to which the fixed stars, Sun, Moon and planets are considered to be in motion about the Earth.

In 1543, the year of his death, the Polish astronomer Nicolas Copernicus published a revolutionary work in which he asserted that the Earth was not the center of the universe. According to his model the Earth rotated about its own axis, and it and the other planets revolved about the Sun. Copernicus calculated the sidereal period of each planet about the Sun; this is the time required for a planet to make one revolution and return to the same angular position relative to a fixed star. He also determined the radii of their orbits about the Sun in terms of the Earth–Sun distance. The mean radius of the Earth’s orbit about the Sun is called an astronomical unit; it equals 149,597,871 km. Accurate values of these parameters were calculated from observations compiled during an interval of 20 years by the Danish astronomer Tycho Brahe (1546–1601). On his death the records passed to his assistant, Johannes Kepler (1571–1630). Kepler succeeded in fitting the observations into a heliocentric model for the system of known planets. The three laws in which Kepler summarized his deductions were later to prove vital to Isaac Newton for verifying the law of Universal Gravitation. It is remarkable that the database used by Kepler was founded on observations that were unaided by the telescope, which was not invented until early in the seventeenth century.

1.1.2 Kepler’s laws of planetary motion

Kepler took many years to fit the observations of Tycho Brahe into three laws of planetary motion. The first and second laws (Fig. 1.2) were published in 1609 and the third law appeared in 1619. The laws may be formulated as follows:

(1) the orbit of each planet is an ellipse with the Sun at one focus;
(2) the orbital radius of a planet sweeps out equal areas in equal intervals of time;
(3) the ratio of the square of a planet’s period \( (T^2) \) to the cube of the semi-major axis of its orbit \( (a^3) \) is a constant for all the planets, including the Earth.

Kepler’s three laws are purely empirical, derived from accurate observations. In fact they are expressions of more fundamental physical laws. The elliptical shapes of planetary orbits (Box 1.1) described by the first law are a consequence of the conservation of energy of a planet orbiting the Sun under the effect of a central attraction that varies as the inverse square of distance. The second law describing the rate of motion of the planet around its orbit follows directly from the conservation of angular momentum of the planet. The third law results from the balance between the force of gravitation attracting the planet towards the Sun and the centrifugal force away from the Sun due to its orbital speed. The third law is easily proved for circular orbits (see Section 2.3.2.3).

Kepler’s laws were developed for the solar system but are applicable to any closed planetary system. They govern the motion of any natural or artificial satellite about a parent body. Kepler’s third law relates the period \( (T) \) and the semi-major axis \( (a) \) of the orbit of the satellite to the mass \( (M) \) of the parent body through the equation

\[
GM = \frac{4\pi^2}{T^2}a^3
\]

where \( G \) is the gravitational constant. This relationship was extremely important for determining the masses of those planets that have natural satellites. It can now be applied to determine the masses of planets using the orbits of artificial satellites.

Special terms are used in describing elliptical orbits. The nearest and furthest points of a planetary orbit around the Sun are called perihelion and aphelion, respectively. The terms periapsis and apoapsis refer to the corresponding nearest and furthest points of the orbit of the Moon or a satellite about the Earth.

1.1.3 Characteristics of the planets

Galileo Galilei (1564–1642) is often regarded as a founder of modern science. He made fundamental discoveries in astronomy and physics, including the formulation of the laws of motion. He was one of the first scientists to use the telescope to acquire more detailed information about
The orbit of a planet or comet in the solar system is an ellipse with the Sun at one of its focal points. This condition arises from the conservation of energy in a force field obeying an inverse square law. The total energy \( E \) of an orbiting mass is the sum of its kinetic energy \( K \) and potential energy \( U \). For an object with mass \( m \) and velocity \( v \) in orbit at distance \( r \) from the Sun (mass \( S \))

\[
\frac{1}{2} mv^2 - \frac{G mS}{r} = E = \text{constant} \tag{1}
\]

If the kinetic energy is greater than the potential energy of the gravitational attraction to the Sun \( (E > 0) \), the object will escape from the solar system. Its path is a hyperbola. The same case results if \( E < 0 \), but the path is a parabola. If \( E < 0 \), the gravitational attraction binds the object to the Sun; the path is an ellipse with the Sun at one focal point (Fig. B1.1.1). An ellipse is defined as the locus of all points in a plane whose distances \( s_1 \) and \( s_2 \) from two fixed points \( F_1 \) and \( F_2 \) in the plane have a constant sum, defined as \( 2a \):

\[
s_1 + s_2 = 2a \tag{2}
\]

The distance \( 2a \) is the length of the major axis of the ellipse; the minor axis perpendicular to it has length \( 2b \), which is related to the major axis by the eccentricity of the ellipse, \( e \):

\[
e = \sqrt{1 - \frac{b^2}{a^2}} \tag{3}
\]

The equation of a point on the ellipse with Cartesian coordinates \( (x, y) \) defined relative to the center of the figure is

\[
\frac{x^2}{a^2} + \frac{y^2}{b^2} = 1 \tag{4}
\]

The elliptical orbit of the Earth around the Sun defines the ecliptic plane. The angle between the orbital plane and the ecliptic is called the inclination of the orbit, and for most planets except Mercury (inclination \( 7^\circ \)) and Pluto (inclination \( 17^\circ \)) this is a small angle. A line perpendicular to the ecliptic defines the North and South ecliptic poles. If the fingers of one’s right hand are wrapped around Earth’s orbit in the direction of motion, the thumb points to the North ecliptic pole, which is in the constellation Draco (“the dragon”). Viewed from above this pole, all planets move around the Sun in a counterclockwise (prograde) sense.

The rotation axis of the Earth is tilted away from the perpendicular to the ecliptic forming the angle of obliquity (Fig. B1.1.2), which is currently \( 23.5^\circ \). The equatorial plane is tilted at the same angle to the ecliptic, which it intersects along the line of equinoxes. During the annual motion of the Earth around the Sun, this line twice points to the Sun: on March 20, defining the vernal (spring) equinox, and on September 23, defining the autumnal equinox. On these dates day and night have equal length everywhere on Earth. The summer and winter solstices occur on June 21 and December 22, respectively, when the apparent motion of the Sun appears to reach its highest and lowest points in the sky.

In 1686 Newton applied his theory of Universal Gravitation to observations of the orbit of Callisto and the planets. In 1610 Galileo discovered the four largest satellites of Jupiter (called Io, Europa, Ganymede and Callisto), and observed that (like the Moon) the planet Venus exhibited different phases of illumination, from full disk to partial crescent. This was persuasive evidence in favor of the Copernican view of the solar system.
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The great planets and Pluto are gaseous. For these planets the surface on which the pressure is 1 atmosphere is taken as the effective radius. In the definition of polar flattening, $a$ and $c$ are respectively the semi-major and semi-minor axes of the spheroidal shape.

### Table 1.1 Dimensions and rotational characteristics of the planets (data sources: Beatty et al., 1999; McCarthy and Petit, 2004; National Space Science Data Center, 2004)

<table>
<thead>
<tr>
<th>Planet</th>
<th>Mass relative to Earth</th>
<th>Mean density [kg m$^{-3}$]</th>
<th>Equatorial radius [km]</th>
<th>Sidereal rotation period [days]</th>
<th>Polar flattening $f = (a - c)/a$</th>
<th>Obliquity of rotation axis [°]</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Terrestrial planets and the Moon</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mercury</td>
<td>0.0553</td>
<td>5.427</td>
<td>2.440</td>
<td>58.81</td>
<td>0.0</td>
<td>0.1</td>
</tr>
<tr>
<td>Venus</td>
<td>0.815</td>
<td>5.243</td>
<td>6.052</td>
<td>243.7</td>
<td>0.0</td>
<td>177.4</td>
</tr>
<tr>
<td>Earth</td>
<td>1.000</td>
<td>5.515</td>
<td>6.378</td>
<td>0.9973</td>
<td>0.003353</td>
<td>23.45</td>
</tr>
<tr>
<td>Moon</td>
<td>0.0123</td>
<td>3.347</td>
<td>1.738</td>
<td>27.32</td>
<td>0.0012</td>
<td>6.68</td>
</tr>
<tr>
<td>Mars</td>
<td>0.1074</td>
<td>3.933</td>
<td>3.397</td>
<td>1.0275</td>
<td>0.00648</td>
<td>25.19</td>
</tr>
<tr>
<td><strong>Great planets and Pluto</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Jupiter</td>
<td>317.8</td>
<td>1.326</td>
<td>71.492</td>
<td>0.414</td>
<td>0.0649</td>
<td>3.12</td>
</tr>
<tr>
<td>Saturn</td>
<td>95.2</td>
<td>687</td>
<td>60.268</td>
<td>0.444</td>
<td>0.098</td>
<td>26.73</td>
</tr>
<tr>
<td>Uranus</td>
<td>14.4</td>
<td>1.270</td>
<td>25.559</td>
<td>0.720</td>
<td>0.023</td>
<td>97.86</td>
</tr>
<tr>
<td>Neptune</td>
<td>17.15</td>
<td>1.638</td>
<td>24.766</td>
<td>0.671</td>
<td>0.017</td>
<td>29.6</td>
</tr>
<tr>
<td>Pluto</td>
<td>0.0021</td>
<td>1.750</td>
<td>1.195</td>
<td>6.405</td>
<td>—</td>
<td>122.5</td>
</tr>
</tbody>
</table>

The masses of the planets can be determined by applying Kepler’s third law to the observed orbits of natural and artificial satellites and to the tracks of passing spacecraft. Estimation of the sizes and shapes of the planets depends on data from several sources. Early astronomers used occultations of the stars by the planets; an occultation is the eclipse of one celestial body by another, such as when a planet passes between the Earth and a star. The duration of an occultation depends on the diameter of the planet, its distance from the Earth and its orbital speed.

The dimensions of the planets (Table 1.1) have been determined with improved precision in modern times by the availability of data from spacecraft, especially from radar-ranging and Doppler tracking (see Box 1.2). Radar-ranging involves measuring the distance between an orbiting (or passing) spacecraft and the planet’s surface from the two-way travel-time of a pulse of electromagnetic waves in the radar frequency range. The separation can be measured with a precision of a few centimeters. If the radar signal is reflected from a planet that is moving away from the spacecraft the frequency of the reflection is lower than that of the transmitted signal; the opposite effect is observed when the planet and spacecraft approach each other. The Doppler frequency shift yields the relative velocity of the spacecraft and planet. Together, these radar methods allow accurate determination of the path of the spacecraft, which is affected by the mass of the planet and the shape of its gravitational equipotential surfaces (see Section 2.2.3).

The rate of rotation of a planet about its own axis can be determined by observing the motion of features on its surface. Where this is not possible (e.g., the surface of Uranus is featureless) other techniques must be employed. In the case of Uranus the rotational period of 17.2 hr was determined from periodic radio emissions produced by electrical charges trapped in its magnetic field; they were detected by the Voyager 2 spacecraft when it flew by the planet in 1986. All planets revolve around the Sun in the same sense, which is counterclockwise when viewed from above the plane of the Earth’s orbit (called the ecliptic plane). Except for Pluto, the orbital plane of each planet is inclined to the ecliptic at a small angle (Table 1.2). Most of the planets rotate about their rotation axis in the same sense as their orbital motion about the Sun, which is termed prograde. Venus rotates in the opposite, retrograde sense. The angle between a rotation axis and the ecliptic plane is called the
1.1 THE SOLAR SYSTEM

### Box 1.2: Radar and the Doppler effect

The name *radar* derives from the acronym for *Radio Detection And Ranging*, a defensive system developed during World War II for the location of enemy aircraft. An electromagnetic signal in the microwave frequency range (see Fig. 4.59), consisting of a continuous wave or a series of short pulses, is transmitted toward a target, from which a fraction of the incident energy is reflected to a receiver. The laws of optics for visible light apply equally to radar waves, which are subject to reflection, refraction and diffraction. Visible light has short wavelengths (400–700 nm) and is scattered by the atmosphere, especially by clouds. Radar signals have longer wavelengths (~1 cm to 30 cm) and can pass through clouds and the atmosphere of a planet with little dispersion. The radar signal is transmitted in a narrow beam of known azimuth, so that the returning echo allows exact location of the direction to the target. The signal travels at the speed of light so the distance, or range, to the target may be determined from the time difference at the source between the transmitted and reflected signals.

The transmitted and reflected radar signals lose energy in transit due to atmospheric absorption, but more importantly, the amplitude of the reflected signal is further affected by the nature of the reflecting surface. Each part of the target’s surface illuminated by the radar beam contributes to the reflected signal. If the surface is inclined obliquely to the incoming beam, little energy will reflect back to the source. The reflectivity and roughness of the reflecting surface determine how much of the incident energy is absorbed or scattered. The intensity of the reflected signal can thus be used to characterize the type and orientation of the reflecting surface, e.g., whether it is bare or forested, flat or mountainous.

The *Doppler effect*, first described in 1842 by an Austrian physicist, Christian Doppler, explains how the relative motion between source and detector influences the observed frequency of light and sound waves. For example, suppose a stationary radar source emits a signal consisting of \( n_0 \) pulses per second. The frequency of pulses reflected from a stationary target at distance \( d \) is also \( n_0 \), and the two-way travel-time of each pulse is equal to \( 2(d/c) \), where \( c \) is the speed of light. If the target is moving toward the radar source, its velocity \( v \) shortens the distance between the radar source and the target by \( (vt/2) \), where \( t \) is the new two-way travel-time:

\[
t = 2 \left( \frac{d - (vt/2)}{c} \right) = t_0 - \frac{vt}{c} \tag{1}
\]

\[
t = t_0/(1 + \frac{v}{c}) \tag{2}
\]

The travel-time of each reflected pulse is shortened, so the number of reflected pulses \( n \) received per second is correspondingly higher than the number emitted:

\[
n = n_0 (1 + \frac{v}{c}) \tag{3}
\]

The opposite situation arises if the target is moving away from the source: the frequency of the reflected signal is lower than that emitted. Similar principles apply if the radar source is mounted on a moving platform, such as an aircraft or satellite. The Doppler change in signal frequency in each case allows remote measurement of the relative velocity between an object and a radar transmitter.

In another important application, the Doppler effect provides evidence that the universe is expanding. The observed frequency of light from a star (i.e., its color) depends on the velocity of its motion relative to an observer on Earth. The color of the star shifts toward the red end of the spectrum (lower frequency) if the star is receding from Earth and toward the blue end (higher frequency) if it is approaching Earth. The color of light from many distant galaxies has a “red shift,” implying that these galaxies are receding from the Earth.

**obliquity** of the axis. The rotation axes of Uranus and Pluto lie close to their orbital planes; they are tilted away from the pole to the orbital plane at angles greater than 90°, so that, strictly speaking, their rotations are also retrograde.

The relative sizes of the planets are shown in Fig. 1.3. They form three categories on the basis of their physical properties (Table 1.1). The terrestrial planets (Mercury, Venus, Earth and Mars) resemble the Earth in size and density. They have a solid, rocky composition and they rotate about their own axes at the same rate or slower than the Earth. The great, or *Jovian*, planets (Jupiter, Saturn, Uranus and Neptune) are much larger than the Earth and have much lower densities. Their compositions are largely gaseous and they rotate more rapidly than the Earth. Pluto’s large orbit is highly elliptical and more steeply inclined to the ecliptic than that of any other planet. Its physical properties are different from both the great planets and the terrestrial planets. These nine bodies are called the *major planets*. There are other large objects in orbit around the Sun, called *minor planets*, which do not fulfil the criteria common to the definition of the major planets. The discovery of large objects in the solar system beyond the orbit of Neptune has stimulated debate among astronomers about what these criteria should be, and whether Pluto should indeed be considered a planet.

#### 1.1.3.1 Bode’s law

In 1772 the German astronomer Johann Bode devised an empirical formula to express the approximate distances of
the planets from the Sun. A series of numbers is created in the following way: the first number is zero, the second is 0.3, and the rest are obtained by doubling the previous number. This gives the sequence 0, 0.3, 0.6, 1.2, 2.4, 4.8, 9.6, 19.2, 38.4, 76.8, etc. Each number is then augmented by 0.4 to give the sequence: 0.4, 0.7, 1.0, 1.6, 2.8, 5.2, 10.0, 19.6, 38.8, 77.2, etc. This series can be expressed mathematically as follows:

\[ d_n = 0.4 + 0.3 \times 2^{n-2} \]  

for \( n \geq 2 \)  

\[ d_1 = 0.4 \]  

This expression gives the distance \( d_n \) in astronomical units (AU) of the \( n \)th planet from the Sun. It is usually known as Bode’s law but, as the same relationship had been suggested earlier by J. D. Titius of Wittenberg, it is sometimes called Titius–Bode’s law. Examination of Fig. 1.4 and comparison with Table 1.2 show that this relationship holds remarkably well, except for Neptune and Pluto. A possible interpretation of the discrepancies is that the orbits of these planets are no longer their original orbits.

Bode’s law predicts a fifth planet at 2.8 AU from the Sun, between the orbits of Mars and Jupiter. In the last years of the eighteenth century astronomers searched intensively for this missing planet. In 1801 a small planetoid, Ceres, was found at a distance of 2.77 AU from the Sun. Subsequently, it was found that numerous small planetoids occupied a broad band of solar orbits centered about 2.9 AU, now called the asteroid belt. Pallas was found in 1802, Juno in 1804, and Vesta, the only asteroid that can be seen with the naked eye, was found in 1807. By 1890 more than 300 asteroids had been identified. In 1891 astronomers began to record their paths on photographic plates. Thousands of asteroids occupying a broad belt
between Mars and Jupiter, at distances of 2.15–3.3 AU from the Sun, have since been tracked and cataloged.

Bode’s law is not a true law in the scientific sense. It should be regarded as an intriguing empirical relationship. Some astronomers hold that the regularity of the planetary distances from the Sun cannot be mere chance but must be a manifestation of physical laws. However, this may be wishful thinking. No combination of physical laws has yet been assembled that accounts for Bode’s law.

1.1.3.2 The terrestrial planets and the Moon

Mercury is the closest planet to the Sun. This proximity and its small size make it difficult to study telescopically. Its orbit has a large eccentricity (0.206). At perihelion the planet comes within 46.0 million km (0.313 AU) of the Sun, but at aphelion the distance is 69.8 million km (0.47 AU). Until 1965 the rotational period was thought to be the same as the period of revolution (88 days), so that it would keep the same face to the Sun, in the same way that the Moon does to the Earth. However, in 1965 Doppler radar measurements showed that this is not the case. In 1974 and 1975 images from the close passage of Mariner 10, the only spacecraft to have visited the planet, gave a period of rotation of 58.8 days, and Doppler tracking gave a radius of 2439 km.

The spin and orbital motions of Mercury are both prograde and are coupled in the ratio 3:2. The spin period is 58.79 Earth days, almost exactly 2/3 of its orbital period of 87.97 Earth days. For an observer on the planet this has the unusual consequence that a Mercury day lasts longer than a Mercury year! During one orbital revolution about the Sun (one Mercury year) an observer on the surface rotates about the spin axis 1.5 times and thus advances by an extra half turn. If the Mercury year started at sunrise, it would end at sunset, so the observer on Mercury would spend the entire 88 Earth days exposed to solar heating, which causes the surface temperature to exceed 700 K. During the following Mercury year, the axial rotation advances by a further half-turn, during which the observer is on the night side of the planet for 88 days, and the temperature sinks below 100 K. After 2 solar orbits and 3 axial rotations, the observer is back at the starting point. The range of temperatures on the surface of Mercury is the most extreme in the solar system.

Although the mass of Mercury is only about 5.5% that of the Earth, its mean density of 5427 kg m⁻³ is comparable to that of the Earth (5515 kg m⁻³) and is the second highest in the solar system. This suggests that, like Earth, Mercury’s interior is dominated by a large iron core, whose radius is estimated to be about 1800–1900 km. It is enclosed in an outer shell 500–600 thick, equivalent to Earth’s mantle and crust. The core may be partly molten. Mercury has a weak planetary magnetic field.

Venus is the brightest object in the sky after the Sun and Moon. Its orbit brings it closer to Earth than any other planet, which made it an early object of study by telescope. Its occultation with the Sun was observed telescopically as early as 1639. Estimates of its radius based on occultations indicated about 6120 km. Galileo observed that the apparent size of Venus changed with its position in orbit and, like the Moon, the appearance of Venus showed different phases from crescent-shaped to full. This was important evidence in favor of the Copernican heliocentric model of the solar system, which had not yet replaced the Aristotelian geocentric model.

Venus has the most nearly circular orbit of any planet, with an eccentricity of only 0.007 and mean radius of 0.72 AU (Table 1.2). Its orbital period is 224.7 Earth days, and the period of rotation about its own axis is 243.7 Earth days, longer than the Venusian year. Its spin axis is tilted at 177° to the pole to the ecliptic, thus making its spin retrograde. The combination of these motions results in the length of a Venusian day (the time between successive sunrises on the planet) being equal to about 117 Earth days.

Venus is very similar in size and probable composition to the Earth. During a near-crescent phase the planet is ringed by a faint glow indicating the presence of an atmosphere. This has been confirmed by several spacecraft that have visited the planet since the first visit by Mariner 2 in 1962. The atmosphere consists mainly of carbon dioxide and is very dense; the surface atmospheric pressure is 92 times that on Earth. Thick cloud cover results in a strong greenhouse effect that produces stable temperatures up to 740 K, slightly higher than the maximum day-time values on Mercury, making Venus the hottest of the planets. The thick clouds obscure any view of the surface, which has however been surveyed with radar. The Magellan spacecraft, which was placed in a nearly polar orbit around the planet in 1990, carried a radar-imaging system with an optimum resolution of 100 meters, and a radar altimeter system to measure the topography and some properties of the planet’s surface.

Venus is unique among the planets in rotating in a retrograde sense about an axis that is almost normal to the ecliptic (Table 1.1). Like Mercury, it has a high Earth-like density (5243 kg m⁻³). On the basis of its density together with gravity estimates from Magellan’s orbit, it is thought that the interior of Venus may be similar to that of Earth, with a rocky mantle surrounding an iron core about 3000 km in radius, that is possibly at least partly molten. However, in contrast to the Earth, Venus has no detectable magnetic field.

The Earth moves around the Sun in a slightly elliptical orbit. The parameters of the orbital motion are important, because they define astronomical units of distance and time. The Earth’s rotation about its own axis from one solar zenith to the next one defines the solar day (see Section 4.1.1.2). The length of time taken for it to complete one orbital revolution about the Sun defines the solar year, which is equal to 365.242 solar days. The eccentricity of the orbit is presently 0.01671 but it varies between a
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minimum of 0.001 and a maximum of 0.060 with a period of about 100,000 yr due to the influence of the other planets. The mean radius of the orbit (149,597,871 km) is called an astronomical unit (AU). Distances within the solar system are usually expressed as multiples of this unit. The distances to extra-galactic celestial bodies are expressed as multiples of a light-year (the distance travelled by light in one year). The Sun’s light takes about 8 min 20 s to reach the Earth. Owing to the difficulty of determining the gravitational constant the mass of the Earth (M) is not known with high precision, but is estimated to be $5.9737 \times 10^{24}$ kg. In contrast, the product $GM$ is known accurately; it is equal to $3.986004418 \times 10^{14}$ m$^3$ s$^{-2}$. The rotation axis of the Earth is presently inclined at $23.439^\circ$ to the pole of the ecliptic. However, the effects of other planets also cause the angle of obliquity to vary between a minimum of $21.9^\circ$ and a maximum of $24.5^\circ$, with a period of about 41,000 yr.

The Moon is Earth’s only natural satellite. The distance of the Moon from the Earth was first estimated with the method of parallax. Instead of observing the Moon from different positions of the Earth’s orbit, as shown in Fig. 1.1, the Moon’s position relative to a fixed star was observed at times 12 hours apart, close to moonrise and moonset, when the Earth had rotated through half a revolution. The baseline for the measurement is then the Earth’s diameter. The distance of the Moon from the Earth was found to be about 60 times the Earth’s radius.

The Moon rotates about its axis in the same sense as its orbital revolution about the Earth. Tidal friction resulting from the Earth’s attraction has slowed down the Moon’s rotation, so that it now has the same mean period as its revolution, 27.32 days. As a result, the Moon always presents the same face to the Earth. In fact, slightly more than half (about 59%) of the lunar surface can be viewed from the Earth. Several factors contribute to this. First, the plane of the Moon’s orbit around the Earth is inclined at $5^\circ$ to the ecliptic while the Moon’s equator is inclined at $1^\circ32^\prime$ to the ecliptic. The inclination of the Moon’s equator varies by up to $6^\circ41^\prime$ to the plane of its orbit. This is called the libration of latitude. It allows Earth-based astronomers to see $6^\circ41^\prime$ beyond each of Moon’s poles. Secondly, the Moon moves with variable velocity around its elliptical orbit, while its own rotation is constant. Near perigee the Moon’s orbital velocity is fastest (in accordance with Kepler’s second law) and the rate of revolution exceeds slightly the constant rate of lunar rotation. Similarly, near apogee the Moon’s orbital velocity is slowest and the rate of revolution is slightly less than the rate of rotation. The rotational differences are called the Moon’s libration of longitude. Their effect is to expose zones of longitude beyond the average edges of the Moon. Finally, the Earth’s diameter is quite large compared to the Moon’s distance from Earth. During Earth’s rotation the Moon is viewed from different angles that allow about one additional degree of longitude to be seen at the Moon’s edge.

The distance to the Moon and its rotational rate are well known from laser-ranging using reflectors placed on the Moon by astronauts. The accuracy of laser-ranging is about 2–3 cm. The Moon has a slightly elliptical orbit about the Earth, with eccentricity 0.0549 and mean radius 384,100 km. The Moon’s own radius of 1738 km makes it much larger relative to its parent body than the natural satellites of the other planets except for Pluto’s moon, Charon. Its low density of $3347 \text{ kg m}^{-3}$ may be due to the absence of an iron core. The internal composition and dynamics of the Moon have been inferred from instruments placed on the surface and rocks recovered from the Apollo and Luna manned missions. Below a crust that is on average 68 km thick the Moon has a mantle and a small core about 340 km in radius. In contrast to the Earth, the interior is not active, and so the Moon does not have a global magnetic field.

Mars, popularly called the red planet because of its hue when viewed from Earth, has been known since prehistoric times and was also an object of early telescopic study. In 1666 Gian Domenico Cassini determined the rotational period at just over 24 hr; radio-tracking from two Viking spacecraft that landed on Mars in 1976, more than three centuries later, gave a period of 24.623 hr. The orbit of Mars is quite elliptical (eccentricity 0.0934). The large difference between perihelion and aphelion causes large temperature variations on the planet. The average surface temperature is about 218 K, but temperatures range from 140 K at the poles in winter to 300 K on the day side in summer. Mars has two natural satellites, Phobos and Deimos. Observations of their orbits gave early estimates of the mass of the planet. Its size was established quite early telescopically from occultations. Its shape is known very precisely from spacecraft observations. The polar flattening is about double that of the Earth. The rotation rates of Earth and Mars are almost the same, but the lower mean density of Mars results in smaller gravitational forces, so at any radial distance the relative importance of the centrifugal acceleration is larger on Mars than on Earth.

In 2004 the Mars Exploration Rover vehicles Spirit and Opportunity landed on Mars, and transmitted photographs and geological information to Earth. Three spacecraft (Mars Global Surveyor, Mars Odyssey, and Mars Express) were placed in orbit to carry out surveys of the planet. These and earlier orbiting spacecraft and Martian landers have revealed details of the planet that cannot be determined with a distant telescope (including the Earth-orbiting Hubble telescope). Much of the Martian surface is very old and cratered, but there are also much younger rift valleys, ridges, hills and plains. The topography is varied and dramatic, with mountains that rise to 24 km, a 4000 km long canyon system, and impact craters up to 2000 km across and 6 km deep.

The internal structure of Mars can be inferred from the results of these missions. Mars has a relatively low mean density ($3933 \text{ kg m}^{-3}$) compared to the other terrestrial planets. Its mass is only about a tenth that of Earth.
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(Table 1.1), so the pressures in the planet are lower and the interior is less densely compressed. Mars has an internal structure similar to that of the Earth. A thin crust, 35 km thick in the northern hemisphere and 80 km thick in the southern hemisphere, surrounds a rocky mantle whose rigidity decreases with depth as the internal temperature increases. The planet has a dense core 1500–1800 km in radius, thought to be composed of iron with a relatively large fraction of sulfur. Minute perturbations of the orbit of Mars Global Surveyor, caused by deformations of Mars due to solar tides, have provided more detailed information about the internal structure. They indicate that, like the Earth, Mars probably has a solid inner core and a fluid outer core that is, however, too small to generate a global magnetic field.

The Asteroids occur in many sizes, ranging from several hundred kilometers in diameter, down to bodies that are too small to discern from Earth. There are 26 asteroids larger than 200 km in diameter, but there are probably more than a million with diameters around 1 km. Some asteroids have been photographed by spacecraft in fly-by missions: in 1997 the NEAR-Shoemaker spacecraft orbited and landed on the asteroid Eros. Hubble Space Telescope imagery has revealed details of Ceres (diameter 950 km), Pallas (diameter 830 km) and Vesta (diameter 525 km), which suggest that it may be more appropriate to call these three bodies protoplanets (i.e., still in the process of accretion from planetesimals) rather than asteroids. All three are differentiated and have a layered internal structure like a planet, although the compositions of the internal layers are different. Ceres has an oblate spheroidal shape and a silicate core, Vesta’s shape is more irregular and it has an iron core.

Asteroids are classified by type, reflecting their composition (stony carbonaceous or metallic nickel–iron), and by the location of their orbits. Main belt asteroids have near-circular orbits with radii 2–4 AU between Mars and Jupiter. The Centaur asteroids have strongly elliptical orbits that take them to the outer solar system. The Aten and Apollo asteroids follow elliptical Earth-crossing orbits. The collision of one of these asteroids with the Earth would have a cataclysmic outcome. A 1 km diameter asteroid would create a 10 km diameter crater and release as much energy as the simultaneous detonation of most or all of the nuclear weapons in the world’s arsenals. In 1980 Louis and Walter Alvarez and their colleagues showed on the basis of an anomalous concentration of extra-terrestrial iridium at the Cretaceous–Tertiary boundary at Gubbio, Italy, that a 10 km diameter asteroid had probably collided with Earth, causing directly or indirectly the mass extinctions of many species, including the demise of the dinosaurs. There are 240 known Apollo bodies; however, there may be as many as 2000 that are 1 km in diameter and many thousands more measuring tens or hundreds of meters.

Scientific opinion is divided on what the asteroid belt represents. One idea is that it may represent fragments of an earlier planet that was broken up in some disaster. Alternatively, it may consist of material that was never able to consolidate into a planet, perhaps due to the powerful gravitational influence of Jupiter.

1.1.3.3 The great planets

The great planets are largely gaseous, consisting mostly of hydrogen and helium, with traces of methane, water and solid matter. Their compositions are inferred indirectly from spectroscopic evidence, because space probes have not penetrated their atmospheres to any great depth. In contrast to the rocky terrestrial planets and the Moon, the radius of a great planet does not correspond to a solid surface, but is taken to be the level that corresponds to a pressure of one bar, which is approximately Earth’s atmospheric pressure at sea-level.

Each of the great planets is encircled by a set of concentric rings, made up of numerous particles. The rings around Saturn, discovered by Galileo in 1610, are the most spectacular. For more than three centuries they appeared to be a feature unique to Saturn, but in 1977 discrete rings were also detected around Uranus. In 1979 the Voyager 1 spacecraft detected faint rings around Jupiter, and in 1989 the Voyager 2 spacecraft confirmed that Neptune also has a ring system.

Jupiter has been studied from ground-based observatories for centuries, and more recently with the Hubble Space Telescope, but our detailed knowledge of the planet comes primarily from unmanned space probes that sent photographs and scientific data back to Earth. Between 1972 and 1977 the planet was visited by the Pioneer 10 and 11, Voyager 1 and 2, and Ulysses spacecraft. The spacecraft Galileo orbited Jupiter for eight years, from 1995 to 2003, and sent an instrumental probe into the atmosphere. It penetrated to a depth of 140 km before being crushed by the atmospheric pressure.

Jupiter is by far the largest of all the planets. Its mass (19×10^26 kg) is 318 times that of the Earth (Table 1.1) and 2.5 times the mass of all the other planets added together (7.7×10^26 kg). Despite its enormous size the planet has a very low density of only 1326 kg m⁻³, from which it can be inferred that its composition is dominated by hydrogen and helium. Jupiter has at least 63 satellites, of which the four largest – Io, Europa, Ganymede and Callisto – were discovered in 1610 by Galileo. The orbital motions of Io, Europa and Ganymede are synchronous, with periods locked in the ratio 1:2:4. In a few hundred million years, Callisto will also become synchronous with a period 8 times that of Io. Ganymede is the largest satellite in the solar system; with a radius of 2631 km it is slightly larger than the planet Mercury. Some of the outermost satellites are less than 30 km in radius, revolve in retrograde orbits and may be captured asteroids. Jupiter has a system of rings, which are like those of Saturn but are fainter and smaller, and were first detected during analysis of data from...
Voyager 1. Subsequently, they were investigated in detail during the Galileo mission.

Jupiter is thought to have a small, hot, rocky core. This is surrounded by concentric layers of hydrogen, first in a liquid-metallic state (which means that its atoms, although not bonded to each other, are so tightly packed that the electrons can move easily from atom to atom), then non-metallic liquid, and finally gaseous. The planet's atmosphere consists of approximately 86% hydrogen and 14% helium, with traces of methane, water and ammonia. The liquid-metallic hydrogen layer is a good conductor of electrical currents. These are the source of a powerful magnetic field that is many times stronger than the Earth's and enormous in extent. It stretches for several million kilometers toward the Sun and for several hundred million kilometers away from it. The magnetic field traps charged particles from the Sun, forming a zone of intense radiation outside Jupiter's atmosphere that would be fatal to a human being exposed to it. The motions of the electric charges cause radio emissions. These are modulated by the rotation of the planet and are used to estimate the period of rotation, which is about 9.9 hr.

Jupiter's moon Europa is the focus of great interest because of the possible existence of water below its icy crust, which is smooth and reflects sunlight brightly. The Voyager spacecraft took high-resolution images of the moon's surface, and gravity and magnetic data were acquired during close passages of the Galileo spacecraft. Europa has a radius of 1565 km, so is only slightly smaller than Earth's Moon, and is inferred to have an iron–nickel core within a rocky mantle, and an outer shell of water below a thick surface ice layer.

Saturn is the second largest planet in the solar system. Its equatorial radius is 60,268 km and its mean density is merely 687 kg m$^{-3}$ (the lowest in the solar system and less than that of water). Thin concentric rings in its equatorial plane give the planet a striking appearance. The obliquity of its rotation axis to the ecliptic is 26.7°, similar to that of the Earth (Table 1.1). Consequently, as Saturn moves along its orbit the rings appear at different angles to an observer on Earth. Galileo studied the planet by telescope in 1610 but the early instrument could not resolve details and he was unable to interpret his observations as a ring system. The rings were explained by Christian Huygens in 1655 using a more powerful telescope. In 1675, Domenico Cassini observed that Saturn's rings consisted of numerous small rings with gaps between them. The rings are composed of particles of ice, rock and debris, ranging in size from dust particles up to a few cubic meters, which are in orbit around the planet. The origin of the rings is unknown; one theory is that they are the remains of an earlier moon that disintegrated, either due to an extra-planetary impact or as a result of being torn apart by bodily tides caused by Saturn's gravity.

In addition to its ring system Saturn has more than 30 moons, the largest of which, Titan, has a radius of 2575 km and is the only moon in the solar system with a dense atmosphere. Observations of the orbit of Titan allowed the first estimate of the mass of Saturn to be made in 1831. Saturn was visited by the Pioneer 11 spacecraft in 1979 and later by Voyager 1 and Voyager 2. In 2004 the spacecraft Cassini entered orbit around Saturn, and launched an instrumental probe, Huygens, that landed on Titan in January 2005. Data from the probe were obtained during the descent by parachute through Titan's atmosphere and after landing, and relayed to Earth by the orbiting Cassini spacecraft.

Saturn's period of rotation has been deduced from modulated radio emissions associated with its magnetic field. The equatorial zone has a period of 10 hr 14 min, while higher latitudes have a period of about 10 hr 39 min. The shape of the planet is known from occultations of radio signals from the Voyager spacecrafts. The rapid rotation and fluid condition result in Saturn having the greatest degree of polar flattening of any planet, amounting to almost 10%. Its mean density of 687 kg m$^{-3}$ is the lowest of all the planets, implying that Saturn, like Jupiter, is made up mainly of hydrogen and helium and contains few heavy elements. The planet probably also has a similar layered structure, with rocky core overlain successively by layers of liquid-metallic hydrogen and molecular hydrogen. However, the gravitational field of Jupiter compresses hydrogen to a metallic state, which has a high density. This gives Jupiter a higher mean density than Saturn. Saturn has a planetary magnetic field that is weaker than Jupiter’s but probably originates in the same way.

Uranus is so remote from the Earth that Earth-bound telescopic observation reveals no surface features. Until the fly-past of Voyager 2 in 1986 much had to be surmised indirectly and was inaccurate. Voyager 2 provided detailed information about the size, mass and surface of the planet and its satellites, and of the structure of the planet's ring system. The planet's radius is 25,559 km and its mean density is 1270 kg m$^{-3}$. The rotational period, 17.24 hr, was inferred from periodic radio emissions detected by Voyager which are believed to arise from charged particles trapped in the magnetic field and thus rotating with the planet. The rotation results in a polar flattening of 2.3%. Prior to Voyager, there were five known moons. Voyager discovered a further 10 small moons, and a further 12 more distant from the planet have been discovered subsequently, bringing the total of Uranus' known moons to 27. The composition and internal structure of Uranus are probably different from those of Jupiter and Saturn. The higher mean density of Uranus suggests that it contains proportionately less hydrogen and more rock and ice. The rotation period is too long for a layered structure with melted ices of methane, ammonia and water around a molten rocky core. It agrees better with a model in which heavier materials are less concentrated in a central core, and the rock, ices and gases are more uniformly distributed.
1.1 THE SOLAR SYSTEM

Several paradoxes remain associated with Uranus. The axis of rotation is tilted at an angle of 98° to the pole to the planet’s orbit, and thus lies close to the ecliptic plane. The reason for the extreme tilt, compared to the other planets, is unknown. The planet has a prograde rotation about this axis. However, if the other end of the rotation axis, inclined at an angle of 82°, is taken as reference, the planet’s spin can be regarded as retrograde. Both interpretations are equivalent. The anomalous axial orientation means that during the 84 years of an orbit round the Sun the polar regions as well as the equator experience extreme solar insolation. The magnetic field of Uranus is also anomalous: it is inclined at a large angle to the rotation axis and its center is displaced axially from the center of the planet.

Neptune is the outermost of the gaseous giant planets. It can only be seen from Earth with a good telescope. By the early nineteenth century, the motion of Uranus had become well enough charted that inconsistencies were evident. French and English astronomers independently predicted the existence of an eighth planet, and the predictions led to the discovery of Neptune in 1846. The planet had been noticed by Galileo in 1612, but due to its slow motion he mistook it for a fixed star. The period of Neptune’s orbital rotation is almost 165 yr, so the planet has not yet completed a full orbit since its discovery. As a result, and because of its extreme distance from Earth, the dimensions of the planet and its orbit were not well known until 1989, when Voyager 2 became the first – and, so far, the only – spacecraft to visit Neptune.

Neptune’s orbit is nearly circular and lies close to the ecliptic. The rotation axis has an earth-like obliquity of 29.6° and its axial rotation has a period of 16.11 hr, which causes a polar flattening of 1.7%. The planet has a radius of 24,766 km and a mean density of 1638 kg m–3. The internal structure of Neptune is probably like that of Uranus: a small rocky core (about the size of planet Earth) is surrounded by a non-layered mixture of rock, water, ammonia and methane. The atmosphere is predominantly of hydrogen, helium and methane, which absorbs red light and gives the planet its blue color.

The Voyager 2 mission revealed that Neptune has 13 moons and a faint ring system. The largest of the moons, Triton, has a diameter about 40% of Earth’s and its density (2060 kg m–3) is higher than that of most other large moons in the solar system. Its orbit is steeply inclined at 157° to Neptune’s equator, making it the only large natural satellite in the solar system that rotates about its planet in retrograde sense. The moon’s physical characteristics, which resemble the planet Pluto, and its retrograde orbital motion suggest that Triton was captured from elsewhere in the outer solar system.

1.1.3.4 Pluto and the outer solar system

Pluto is the smallest planet in the solar system, about two-thirds the diameter of Earth’s Moon, and has many unusual characteristics. Its orbit has the largest inclination to the ecliptic (17.1°) of any major planet and it is highly eccentric (0.249), with aphelion at 49.3 AU and perihelion at 29.7 AU. This brings Pluto inside Neptune’s orbit for 20 years of its 248-year orbital period; the paths of Pluto and Neptune do not intersect. The orbital period is resonant with that of Neptune in the ratio 3:2 (i.e., Pluto’s period is exactly 1.5 times Neptune’s). These features preclude any collision between the planets.

Pluto is so far from Earth that it appears only as a speck of light to Earth-based telescopes and its surface features can be resolved only broadly with the Hubble Space Telescope. It is the only planet that has not been visited by a spacecraft. It was discovered fortuitously in 1930 after a systematic search for a more distant planet to explain presumed discrepancies in the orbit of Neptune which, however, were later found to be due to inaccurate estimates of Neptune’s mass. The mass and diameter of Pluto were uncertain for some decades until in 1978 a moon, Charon, was found to be orbiting Pluto at a mean distance of 19,600 km. Pluto’s mass is only 0.21% that of the Earth. Charon’s mass is about 10–15% of Pluto’s, making it the largest moon in the solar system relative to its primary planet. The radii of Pluto and Charon are estimated from observations with the Hubble Space Telescope to be 1137 km and 586 km, respectively, with a relative error of about 1%. The mass and diameter of Pluto give an estimated density about 2000 kg m–3 from which it is inferred that Pluto’s composition may be a mixture of about 70% rock and 30% ice, like that of Triton, Neptune’s moon. Charon’s estimated density is lower, about 1300 kg m–3, which suggests that there may be less rock in its composition.

Pluto’s rotation axis is inclined at about 122° to its orbital plane, so the planet’s axial rotation is retrograde, and has a period of 6.387 days. Charon also orbits Pluto in a retrograde sense. As a result of tidal forces, Charon’s orbital period is synchronous with both its own axial rotation and Pluto’s. Thus, the planet and moon constantly present the same face to each other. Because of the rotational synchronism and the large relative mass of Charon, some consider Pluto–Charon to be a double planet. However, this is unlikely because their different densities suggest that the bodies originated independently. Observations with the Hubble Space Telescope in 2005 revealed the presence of two other small moons – provisionally named 2005 P1 and 2005 P2 – in orbit around Pluto in the same sense as Charon, but at a larger distance of about 44,000 km. All three moons have the same color spectrum, which differs from Pluto’s and suggests that the moons were captured in a single collision with another large body. However, the origins of Pluto, Charon and the smaller moons are as yet unknown, and are a matter of scientific conjecture.

Since the early 1990s thousands of new – mostly small – objects beyond the orbit of Neptune have been identified.
The Earth as a planet

The trans-Neptunian objects (Box 1.3) are mostly small, but some are comparable in size to Pluto. The new discoveries have fuelled discussion as to whether Pluto’s status in the solar system should be considered that of a planet. Curiously, there is no exact definition of what constitutes a planet. Generally, to be a planet an object must (1) be in orbit around a star (Sun), (2) be large enough so that its own gravitation results in a spherical or spheroidal shape, and (3) not be so large as to initiate nuclear fusion. Conditions (1) and (3) are met by all objects orbiting the Sun. But a spherical or spheroidal shape proves to be an inadequate criterion. If a threshold diameter of 400–500 km is set, there are orbiting bodies of irregular shape that exceed the critical size, as well as spherical bodies that are smaller.

1.1.3.5 Angular momentum

An important characteristic that constrains models of the origin of the solar system is the difference between the distributions of mass and angular momentum. To determine the angular momentum of a rotating body it is necessary to know its moment of inertia. For a particle of mass \( m \) the moment of inertia \( (I) \) about an axis at distance \( r \) is defined as:

\[
I = mr^2
\]

(1.4)

The angular momentum \( (h) \) is defined as the product of its moment of inertia \( (I) \) about an axis and its rate of rotation \( (\Omega) \) about that axis:

\[
h = I\Omega
\]

(1.5)

Each planet revolves in a nearly circular orbit around the Sun and at the same time rotates about its own axis. Thus there are two contributions to its angular momentum (Table 1.3). The angular momentum of a planet’s revolution about the Sun is obtained quite simply. The solar system is so immense that the physical size of each planet is tiny compared to the size of its orbit. The moment of inertia of a planet about the Sun is computed by inserting the mass of the planet and its orbital radius (Table 1.3) in Eq. (1.4); the orbital angular momentum of the planet follows by combining the computed moment of inertia with the rate of orbital revolution as in Eq. (1.5). To determine the moment of inertia of a solid body about an axis that passes through it (e.g., the rotational axis of a planet) is more complicated. Equation (1.4) must be computed and summed for all particles in the planet. If the planet is represented by a sphere of mass \( M \) and mean radius \( R \), the moment of inertia \( C \) about the axis of rotation is given by

\[
C = kMR^2
\]

(1.6)

where the constant \( k \) is determined by the density distribution within the planet. For example, if the density is uniform inside the sphere, the value of \( k \) is exactly 2/5, or 0.4; for a hollow sphere it is 2/3. If density increases with depth in the planet, e.g., if it has a dense core, the value of \( k \) is less than 0.4; for the Earth, \( k = 0.3308 \). For some planets the variation of density with depth is not well known, but for most planets there is enough information to calculate the moment of inertia about the axis of rotation; combined with the rate of rotation as in Eq. (1.5), this gives the rotational angular momentum.

The angular momentum of a planet’s revolution about the Sun is much greater (on average about 60,000 times) than the angular momentum of its rotation about its own axis (Table 1.3). Whereas more than 99.9% of the total mass of the solar system is concentrated in the Sun, more than 99% of the angular momentum is carried by the orbital motion of the planets, especially the four great planets. Of these Jupiter is a special case: it accounts for over 70% of the mass and more than 60% of the angular momentum of the planets.

1.1.4 The origin of the solar system

There have been numerous theories for the origin of the solar system. Age determinations on meteorites indicate that the solar system originated about \((4.5–4.6) \times 10^9\) years ago. A successful theory of how it originated must account satisfactorily for the observed characteristics of the planets. The most important of these properties are the following.

1. Except for Pluto, the planetary orbits lie in or close to the same plane, which contains the Sun and the orbit of the Earth (the ecliptic plane).
2. The planets revolve about the Sun in the same sense, which is counterclockwise when viewed from above the ecliptic plane. This sense of rotation is defined as prograde.
3. The rotations of the planets about their own axes are also mostly prograde. The exceptions are Venus, which has a retrograde rotation; Uranus, whose axis of rotation lies nearly in the plane of its orbit; and Pluto, whose rotation axis and orbital plane are oblique to the ecliptic.
4. Each planet is roughly twice as far from the Sun as its closest neighbor (Bode’s law).
5. The compositions of the planets make up two distinct groups: the terrestrial planets lying close to the Sun are small and have high densities, whereas the great planets far from the Sun are large and have low densities.
6. The Sun has almost 99.9% of the mass of the solar system, but the planets account for more than 99% of the angular momentum.

The first theory based on scientific observation was the nebular hypothesis introduced by the German philosopher Immanuel Kant in 1755 and formulated by the French astronomer Pierre Simon de Laplace in 1796. According to this hypothesis the planets and their satellites were
A trans-Neptunian object (TNO) is any object in orbit around the Sun at a greater average distance than Neptune. They include Pluto and its moon Charon, as well as numerous other bodies. The objects are grouped in three classes according to the size of their orbit: the Kuiper belt, Scattered Disk, and Oort Cloud. Their composition is similar to that of comets, i.e., mainly ice, although some have densities high enough to suggest other rock-like components.

The **Kuiper belt** extends beyond the mean radius of Neptune’s orbit at 30 AU to a distance of about 50 AU (Fig. B1.3). This disk-shaped region, close to the ecliptic plane, contains thousands of objects in orbit around the Sun. According to some estimates there are more than 35,000 Kuiper Belt objects larger than 100 km in diameter, so they are much larger and more numerous than the asteroids. Some have orbital periods that are in resonance with the orbit of Neptune, and this has given rise to some curious appellations for them. Objects like Pluto with orbital periods in 3:2 resonance with Neptune are called plutinos, those further out in the belt with periods in 2:1 resonance are called twotinos, and objects in intermediate orbits are called cubewanos. The Kuiper belt objects are all largely icy in composition, and some of them are quite large. For example, Quaoar, in an orbit with semi-major axis 43.5 AU, has a diameter of 1260 km and so is about the same size as Pluto’s moon, Charon.

Objects in orbit at mean distances greater than 50 AU are called **scattered disk objects**. A large trans-Neptunian object – provisionally labelled 2003UB313 – was identified in 2003 and confirmed in 2005 during a long-term search for distant moving objects in the solar system. On the basis of its reflectivity this object may be larger than Pluto. If Pluto retains its status as a planet, this object may become the tenth planet in the solar system. It has an orbital period of 557 yr, a highly elliptical orbit inclined at 44° to the ecliptic, and is currently near to aphelion. Its present heliocentric distance of 97 AU makes it the most distant known object in the solar system.

In 2004 another trans-Neptunian object, Sedna, was discovered at a distance of 90 AU (Fig. B1.3). It is presently closer to the Sun than 2003UB313, but its extremely elliptical orbit (eccentricity 0.855, inclination 12°) takes Sedna further into the outer reaches of the solar system than any known object. Its orbital period is 12,500 yrs and its aphelion lies at about 975 AU. The object is visible to astronomers only as a tiny speck so apart from its orbit not much is known about it. It is considered to be the only known object that may have originated in the Oort Cloud.

**The Oort cloud** is thought to be the source of most new comets that enter the inner solar system. It is visualized as a spherical cloud of icy objects at an enormous distance – between 50,000 and 100,000 AU (roughly one light year) – from the Sun. The Oort cloud has never been observed, but its existence has been confirmed from work on cometary orbits. It plays a central role in models of the origin of comets.
<table>
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<tr>
<th>Planet</th>
<th>Mean orbital radius ( r ) [10^9 m]</th>
<th>Mean orbital angular momentum ( M_\omega ) [10^39 kg m^2 s^-1]</th>
<th>Orbital angular momentum of inertia ( I_\omega ) [10^40 kg m^2 s^-1]</th>
<th>Normalized moment of inertia ( I/MR^2 )</th>
<th>Normalized rotational angular momentum ( I_\Omega ) [10^10 kg m^2 s^-1]</th>
<th>Axial rotation rate ( \Omega ) ( [10^{-6} \text{ rad s}^{-1}] )</th>
<th>Rotational angular momentum ( \Omega_0 ) [10^10 rad s^-1]</th>
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### Table 1.3: Distributions of orbital and rotational angular momentum in the solar system (data sources: Yoder, 1995; Beatty et al., 1999; McCarthy and Petit, 2004; National Space Science Data Center, 2004)

Venus, Uranus and Pluto have retrograde axial positions.
formed at the same time as the Sun. Space was filled by a rotating cloud (nebula) of hot primordial gas and dust that, as it cooled, began to contract. To conserve the angular momentum of the system, its rotation speeded up; a familiar analogy is the way a pirouetting skater spins more rapidly when he draws in his outstretched arms. Centrifugal force would have caused concentric rings of matter to be thrown off, which then condensed into planets. A serious objection to this hypothesis is that the mass of material in each ring would be too small to provide the gravitational attraction needed to cause the ring to condense into a planet. Moreover, as the nebula contracted, the largest part of the angular momentum would remain associated with the main mass that condensed to form the Sun, which disagrees with the observed distribution of angular momentum in the solar system.

Several alternative models were postulated subsequently, but have also fallen into disfavor. For example, the collision hypothesis assumed that the Sun was formed before the planets. The gravitational attraction of a closely passing star or the blast of a nearby supernova explosion drew out a filament of solar material that condensed to form the planets. However, a major objection to this scenario is that the solar material would have been so hot that it would dissipate explosively into space rather than condense slowly to form the planets.

Modern interpretations of the origin of the solar system are based on modifications of the nebular hypothesis. As the cloud of gas and dust contracted, its rate of rotation speeded up, flattening the cloud into a lens-shaped disk. When the core of the contracting cloud became dense enough, gravitation caused it to collapse upon itself to form a proto-Sun in which thermonuclear fusion was initiated. Hydrogen nuclei combined under the intense pressure to form helium nuclei, releasing huge amounts of energy. The material in the spinning disk was initially very hot and gaseous but, as it cooled, solid material condensed out of it as small grains. The grains coalesced as rocky or icy clumps called planetesimals. Asteroid-like planetesimals with a silicate, or rocky, composition formed near the Sun, while comet-like planetesimals with an icy composition formed far from the Sun’s heat. In turn, helped by gravitational attraction, the planetesimals accreted to form the planets. Matter with a high boiling point (e.g., metals and silicates) could condense near to the Sun, forming the terrestrial planets. Volatile materials (e.g., water, methane) would vaporize and be driven into space by the stream of particles and radiation from the Sun. During the condensation of the large cold planets in the frigid distant realms of the solar system, the volatile materials were retained. The gravitational attractions of Jupiter and Saturn may have been strong enough to retain the composition of the original nebula.

It is important to keep in mind that this scenario is merely a hypothesis – a plausible but not unique explanation of how the solar system formed. It attributes the variable compositions of the planets to accretion at different distances from the Sun. The model can be embellished in many details to account for the characteristics of individual planets. However, the scenario is unsatisfactory because it is mostly qualitative. For example, it does not adequately explain the division of angular momentum. Physicists, astronomers, space scientists and mathematicians are constantly trying new methods of investigation and searching for additional clues that will improve the hypothesis of how the solar system formed.
wrinkles on a desiccating apple. Horizontal tectonic displacements were known, but were considered to be a by-product of more important vertical motions. The realization that large overthrusts played an important role in the formation of nappe structures in the Alps implied amounts of horizontal shortening that were difficult to accommodate in the contraction hypothesis. A new school of thought emerged in which mountain-building was depicted as a consequence of horizontal displacements.

A key observation in this context was the congruity between the opposing coasts of the South Atlantic, especially the similar shapes of the coastlines of Brazil and Africa. As early as 1620, despite the inaccuracy and incompleteness of early seventeenth century maps, Francis Bacon drew attention to the parallelism of the Atlantic-bordering coastlines. In 1858 Antonio Snider constructed a map showing relative movements of the circum-Atlantic continents, although he did not maintain the shapes of the coastlines. In the late nineteenth century the Austrian geologist Eduard Suess coined the name Gondwanaland for a proposed great southern continent that existed during late Paleozoic times. It embodied Africa, Antarctica, Arabia, Australia, India and South America, and lay predominantly in the southern hemisphere. The Gondwana continents are now individual entities and some (e.g., India, Arabia) no longer lie in the southern hemisphere, but they are often still called the “southern continents.” In the Paleozoic, the “northern continents” of North America (including Greenland), Europe and most of Asia also formed a single continent, called Laurasia. Laurasia and Gondwanaland split apart in the Early Mesozoic. The Alpine–Himalayan mountain belt was thought to have developed from a system of geosynclines that formed in the intervening sea, which Suess called the Tethys ocean to distinguish it from the present Mediterranean Sea. Implicit in these reconstructions is the idea that the continents subsequently reached their present positions by slow horizontal displacements across the surface of the globe.

1.2.2 Continental drift

The “displacement hypothesis” of continental movements matured early in the twentieth century. In 1908 F. B. Taylor related the world’s major fold-belts to convergence of the continents as they moved away from the poles, and in 1911 H. B. Baker reassembled the Atlantic-bordering continents together with Australia and Antarctica into a single continent; regrettably he omitted Asia and the Pacific. However, the most vigorous proponent of the displacement hypothesis was Alfred Wegener, a German meteorologist and geologist. In 1912 Wegener suggested that all of the continents were together in the Late Paleozoic, so that the land area of the Earth formed a single landmass (Fig. 1.5). He coined the name Pangaea (Greek for “all Earth”) for this supercontinent, which he envisioned was surrounded by a single ocean (Panthalassa). Wegener referred to the large-scale horizontal displacement of crustal blocks having continental dimensions as Kontinentalverschiebung. The anglicized form, continental drift, implies additionally that displacements of the blocks take place slowly over long time intervals.

1.2.2.1 Pangaea

As a meteorologist Wegener was especially interested in paleoclimatology. For the first half of the twentieth century the best evidence for the continental drift hypothesis and the earlier existence of Pangaea consisted of geological indicators of earlier paleoclimates. In particular, Wegener observed a much better alignment of regions of Permo-Carboniferous glaciation in the southern hemisphere when the continents were in the reconstructed positions for Gondwanaland instead of their present positions. His reconstruction of Pangaea brought Carboniferous coal deposits into alignment and suggested that the positions of the continents relative to the Paleozoic equator were quite different from their modern ones. Together with W. Köppen, a fellow German meteorologist, he assembled paleoclimatic data that showed the distributions of coal deposits (evidence of moist temperate zones), salt, gypsum and desert sandstones (evidence of dry climate) for several geological
eras (Carboniferous, Permian, Eocene, Quaternary). When plotted on Wegener’s reconstruction maps, the paleoclimatic data for each era formed climatic belts just like today; namely, an equatorial tropical rain belt, two adjacent dry belts, two temperate rain belts, and two polar ice caps (Fig. 1.5a).

Wegener’s continental drift hypothesis was bolstered in 1937 by the studies of a South African geologist, Alexander du Toit, who noted sedimentological, paleontological, paleoclimatic, and tectonic similarities between western Africa and eastern South America. These favored the Gondwanaland reconstruction rather than the present configuration of continents during the Late Paleozoic and Early Mesozoic.

Some of Wegener’s theories were largely conjectural. On the one hand, he reasoned correctly that the ocean basins are not permanent. Yet he envisioned the sub-crustal material as capable of viscous yield over long periods of time, enabling the continents to drift through the ocean crust like ships through water. This model met with profound scepticism among geologists. He believed, in the face of strong opposition from physicists, that the Earth’s geographic axis had moved with time, instead of the crust moving relative to the fixed poles. His timing of the opening of the Atlantic (Fig. 1.5b, c) was faulty, requiring a large part of the separation of South America from Africa to take place since the Early Pleistocene (i.e., in the last two million years or so). Moreover, he was unable to offer a satisfactory driving mechanism for continental drift. His detractors used the disprovable speculations to discredit his better-documented arguments in favor of continental drift.

1.2.2.2 Computer-assisted reconstructions

Wegener pointed out that it was not possible to fit the continents together using their present coastlines, which are influenced by recent sedimentary deposits at the mouths of major rivers as well as the effects of coastal erosion. The large areas of continental shelf must also be taken into account, so Wegener matched the continents at about the edges of the continental shelves, where the continental slopes plunge into the oceanic basins. The matching was visual and inexact by modern standards, but more precise methods only became available in the 1960s with the development of powerful computers.

In 1965 E. C. Bullard, J. E. Everett and A. G. Smith used a computer to match the relative positions of the continents bounding the Atlantic ocean (Fig. 1.6). They digitized the continental outlines at approximately 50 km intervals for different depth contours on the continental slopes, and selected the fit of the 500 fathom (900 m) depth contour as optimum. The traces of opposite continental margins were matched by an iterative procedure. One trace was rotated relative to the other (about a pole of relative rotation) until the differences between the traces were minimized; the procedure was then repeated with different rotation poles until the best fit was obtained. The optimum fit is not perfect, but has some overlaps and gaps. Nevertheless, the analysis gives an excellent geometric fit of the opposing coastlines of the Atlantic.

A few years later A. G. Smith and A. Hallam used the same computer-assisted technique to match the coastlines of the southern continents, also at the 500 fathom depth contour (Fig. 1.7). They obtained an optimum geometric reconstruction of Gondwanaland similar to the visual match suggested by du Toit in 1937; it probably represents the geometry of Gondwanaland that existed in the Late Paleozoic and Early Mesozoic. It is not the only possible good geometric fit, but it also satisfies other geological evidence. At various times in the Jurassic and Cretaceous, extensional plate margins formed within Gondwanaland, causing it to subdivide to form the present “southern continents.” The dispersal to their present positions took place largely in the Late Cretaceous and Tertiary.

Pangaea existed only in the Late Paleozoic and Early Mesozoic. Geological and geophysical evidence argues in favor of the existence of its northern and southern constituents – Laurasia and Gondwanaland – as separate entities in the Early Paleozoic and Precambrian. An important source of data bearing on continental reconstructions in ancient times and the drift of the continents is provided by paleomagnetism, which is the record of the Earth’s ancient magnetic field. Paleomagnetism is described in Section 5.6 and summarized below.
1.2.2.3 Paleomagnetism and continental drift

In the late nineteenth century geologists discovered that rocks can carry a stable record of the geomagnetic field direction at the time of their formation. From the magnetization direction it is possible to calculate the position of the magnetic pole at that time; this is called the virtual geomagnetic pole (VGP) position. Averaged over a time interval longer than a few tens of thousands of years, the mean VGP position coincides with the geographic pole, as if the axis of the mean geomagnetic dipole field were aligned with the Earth’s rotation axis. This correspondence can be proved for the present geomagnetic field, and a fundamental assumption of paleomagnetism – called the “axial dipole hypothesis” – is that it has always been valid. The hypothesis can be verified for rocks and sediments up to a few million years old, but its validity has to be assumed for earlier geological epochs. However, the self-consistency of paleomagnetic data and their compatibility with continental reconstructions argue that the axial dipole hypothesis is also applicable to the Earth’s ancient magnetic field.

For a particular continent, rocks of different ages give different mean VGP positions. The appearance that the pole has shifted with time is called apparent polar wander (APW). By connecting mean VGP positions of different ages for sites on the same continent a line is obtained, called the apparent polar wander path of the continent. Each continent yields a different APW path, which consequently cannot be the record of movement of the pole. Rather, each APW path represents the movement of the continent relative to the pole. By comparing APW paths the movements of the continents relative to each other can be reconstructed. The APW paths provide strong supporting evidence for continental drift.

Paleomagnetism developed as a geological discipline in the 1950s and 1960s. The first results indicating large-scale continental movement were greeted with some scepticism. In 1956 S. K. Runcorn demonstrated that the paleomagnetic data from Permian and Triassic rocks in North America and Great Britain agreed better if the Atlantic ocean were closed, i.e., as in the Laurasia configuration. In 1957 E. Irving showed that Mesozoic paleomagnetic data from the “southern continents” were more concordant with du Toit’s Gondwanaland reconstruction than with the present arrangement of the continents. Since these pioneering studies numerous paleomagnetic investigations have established APW paths for the different continents. The quality of the paleomagnetic record is good for most geological epochs since the Devonian.

The record for older geological periods is less reliable for several reasons. In the Early Paleozoic the data become fewer and the APW paths become less well defined. In addition, the oldest parts of the paleomagnetic record are clouded by the increasing possibility of false directions due to undetected secondary magnetization. This happens when thermal or tectonic events alter the original magnetization, so that its direction no longer corresponds to that at the time of rock formation. Remagnetization can affect rocks of any age, but it is recognized more readily and constitutes a less serious problem in younger rocks.

Problems afflicting Precambrian paleomagnetism are even more serious than in the Early Paleozoic. APW paths have been derived for the Precambrian, especially for North America, but only in broad outline. In part this is because it is difficult to date Precambrian rocks precisely enough to determine the fine details of an APW path. It is often not possible to establish which is the north or south pole. In addition, the range of time encompassed by the Precambrian – more than 3.5 Ga – is about six times longer than the 570 Ma length of the Phanerozoic, and the probability of remagnetization events is correspondingly higher.
In spite of some uncertainties, Early Paleozoic paleomagnetism permits reassembly of the supercontinents Gondwanaland and Laurasia and traces their movements before they collided in the Carboniferous to form Pangaea. Geological and paleomagnetic evidence concur that, in the Cambrian period, Gondwanaland very likely existed as a supercontinent in essentially the du Toit configuration. It coexisted in the Early Paleozoic with three other cratonic centers: Laurentia (North America and Greenland), Baltica (northern Europe) and Siberia. Laurentia and Baltica were separated by the Iapetus ocean (Fig. 1.8a), which began to close in the Ordovician (about 450 Ma ago). Paleomagnetic data indicate that Laurentia and Baltica fused together around Late Silurian time to form the supercontinent Laurussia; at that time the Siberian block remained a separate entity. The Laurentia–Baltica collision is expressed in the Taconic and Caledonian orogenies in North America and northern Europe. The gap between Gondwanaland and Laurussia in the Middle Silurian (Fig. 1.8b) closed about 410 Ma ago. Paleomagnetic data indicate that Laurussia separated from Gondwanaland in the Late Devonian, but the two supercontinents began to collide again in the Early Carboniferous (about 350 Ma ago), causing the Hercynian orogeny. By the Late Carboniferous (300 Ma ago) Pangaea was almost complete, except for Siberia, which was probably appended in the Permian.

The general configuration of Pangaea from the Late Carboniferous to the Early Jurassic is supported by paleomagnetic results from the Atlantic-bordering continents. However, the paleomagnetic data suggest that the purely geometric “Bullard-fit” is only appropriate for the later part of Pangaea’s existence. The results for earlier times from the individual continents agree better for slightly different reconstructions (see Section 5.6.4.4). This suggests that some internal rearrangement of the component parts of Pangaea may have occurred. Also, the computer-assisted geometric assembly of Gondwanaland, similar to that proposed by du Toit, is not the only possible reconstruction, although paleomagnetic results confirm that it is probably the optimum one. Other models involve different relative placements of West Gondwanaland (i.e., South America and Africa) and East Gondwanaland (i.e., Antarctica, Australia and India), and imply that they may have moved relative to each other. The paleomagnetic data do not contradict the alternative models, but are not precise enough to discriminate definitively between them.

The consistency of paleomagnetic results leaves little room for doubt that the continents have changed position relative to each other throughout geological time. This lends justification to the concept of continental drift, but it does not account for the mechanism by which it has taken place. Another aspect of the paleomagnetic record – the history of magnetic field polarity rather than the APW paths – has played a key role in deducing the mechanism. The explanation requires an understanding of the
Earth’s internal structure, the distribution of seismicity and the importance of the ocean basins.

1.2.3 Earth structure

Early in the twentieth century it became evident from the study of seismic waves that the interior of the Earth has a radially layered structure, like that of an onion (Fig. 1.9). The boundaries between the layers are marked by abrupt changes in seismic velocity or velocity gradient. Each layer is characterized by a specific set of physical properties determined by the composition, pressure and temperature in the layer. The four main layers are the crust, mantle and the outer and inner cores. Their properties are described in detail in Section 3.7 and summarized briefly here.

At depths of a few tens of kilometers under continents and less than ten kilometers beneath the oceans seismic velocities increase sharply. This seismic discontinuity, discovered in 1909 by A. Mohorovicic, represents the boundary between the crust and mantle. R. D. Oldham noted in 1906 that the travel-times of seismic compressional waves that traversed the body of the Earth were greater than expected; the delay was attributed to a fluid outer core. Support for this idea came in 1914, when B. Gutenberg described a shadow zone for seismic waves at epicentral distances greater than about 105°. In 1936 I. Lehmann observed the weak arrivals of compressional waves in the gap between 105° and 143°. They are interpreted as evidence for a solid inner core.

1.2.3.1 Lithospheric plates

The radially layered model of the Earth’s interior assumes spherical symmetry. This is not valid for the crust and upper mantle. These outer layers of the Earth show important lateral variations. The crust and uppermost mantle down to a depth of about 70–100 km under deep ocean basins and 100–150 km under continents is rigid, forming a hard outer shell called the lithosphere. Beneath the lithosphere lies the asthenosphere, a layer in which seismic velocities often decrease, suggesting lower rigidity. It is about 150 km thick, although its upper and lower boundaries are not sharply defined. This weaker layer is thought to be partially molten; it may be able to flow over long periods of time like a viscous liquid or plastic solid, in a way that depends on temperature and composition. The asthenosphere plays an important role in plate tectonics, because it makes possible the relative motions of the overlying lithospheric plates.

The brittle condition of the lithosphere causes it to fracture when strongly stressed. The rupture produces an earthquake, which is the violent release of elastic energy
due to sudden displacement on a fault plane. Earthquakes are not distributed evenly over the surface of the globe, but occur predominantly in well-defined narrow seismic zones that are often associated with volcanic activity (Fig. 1.10). These are: (a) the circum-Pacific “ring of fire”; (b) a sinuous belt running from the Azores through North Africa and the Alpine–Dinaride–Himalayan mountain chain as far as S.E. Asia; and (c) the world-circling system of oceanic ridges and rises. The seismic zones subdivide the lithosphere laterally into tectonic plates (Fig. 1.11). A plate may be as broad as 10,000 km (e.g., the Pacific plate) or as small as a few 1000 km (e.g., the Philippines plate). There are twelve major plates (Antarctica, Africa, Eurasia, India, Australia, Arabia, Philippines, North America, South America, Pacific, Nazca, and Cocos) and several minor plates (e.g., Scotia, Caribbean, Juan de Fuca). The positions of the boundaries between the North American and South American plates and between the North American and Eurasian plates are uncertain. The boundary between the Indian and Australian plates is not sharply defined, but may be a broad region of diffuse deformation.

A comprehensive model of current plate motions (called NUVEL-1), based on magnetic anomaly patterns and first-motion directions in earthquakes, shows rates of separation at plate boundaries that range from about 20 mm yr\(^{-1}\) in the North Atlantic to about 160 mm yr\(^{-1}\) on the East Pacific Rise (Fig. 1.11). The model also gives rates of closure ranging from about 10 mm yr\(^{-1}\) between Africa and Eurasia to about 80 mm yr\(^{-1}\) between the Nazca plate and South America.

1.2.4 Types of plate margin

An important factor in the evolution of modern plate tectonic theory was the development of oceanography in the years following World War II, when technology designed for warfare was turned to peaceful purposes. The bathymetry of the oceans was charted extensively by echo-sounding...
and within a few years several striking features became evident. Deep trenches, more than twice the depth of the ocean basins, were discovered close to island arcs and some continental margins; the Marianas Trench is more than 11 km deep. A prominent submarine mountain chain—called an oceanic ridge—was found in each ocean. The oceanic ridges rise to as much as 3000 m above the adjacent basins and form a continuous system, more than 60,000 km in length, that girdles the globe. Unlike continental mountain belts, which are usually less than several hundred kilometers across, the oceanic ridges are 2000–4000 km in width. The ridge system is offset at intervals by long horizontal faults forming fracture zones. These three features—trenches, ridges and fracture zones—originate from different plate tectonic processes.

The lithospheric plates are very thin in comparison to their breadth (compare Fig. 1.9 and Fig. 1.11). Most earthquakes occur at plate margins, and are associated with interactions between plates. Apart from rare intraplate earthquakes, which can be as large and disastrous as the earthquakes at plate boundaries, the plate interiors are aseismic. This suggests that the plates behave rigidly. Analysis of earthquakes allows the direction of displacement to be determined and permits interpretation of the relative motions between plates.

There are three types of plate margin, distinguished by different tectonic processes (Fig. 1.12). The world-wide pattern of earthquakes shows that the plates are presently moving apart at oceanic ridges. Magnetic evidence, discussed below, confirms that the separation has been going on for millions of years. New lithosphere is being formed at these spreading centers, so the ridges can be regarded as constructive plate margins. The seismic zones related to deep-sea trenches, island arcs and mountain belts mark places where lithospheric plates are converging. One plate is forced under another there in a so-called subduction zone. Because it is thin in relation to its breadth, the lower plate bends sharply before descending to depths of several hundred kilometers, where it is absorbed. The subduction zone marks a destructive plate margin.

Constructive and destructive plate margins may consist of many segments linked by horizontal faults. A crucial step in the development of plate tectonic theory was made in 1965 by a Canadian geologist, J. Tuzo Wilson, who recognized that these faults are not conventional transcurrent faults. They belong to a new class of faults, which Wilson called transform faults. The relative motion on a transform fault is opposite to what might be inferred from the offsets of bordering ridge segments. At the point where a transform fault meets an oceanic ridge it transforms the spreading on the ridge to horizontal shear on the fault. Likewise, where such a fault meets a destructive plate margin it transforms subduction to horizontal shear.

The transform faults form a conservative plate margin, where lithosphere is neither created nor destroyed; the boundary separates plates that move past each other horizontally. This interpretation was documented in 1967 by L. Sykes, an American seismologist. He showed that earthquake activity on an oceanic ridge system was confined almost entirely to the transform fault between ridge crests, where the neighboring plates rub past each other. Most importantly, Sykes found that the mechanisms of earthquakes on the transform faults agreed with the predicted sense of strike-slip motion.

Transform faults play a key role in determining plate motions. Spreading and subduction are often assumed to be perpendicular to the strike of a ridge or trench, as is the case for ridge X in Fig. 1.12. This is not necessarily the case. Oblique motion with a component along strike is possible at each of these margins, as on ridge Y. However, because lithosphere is neither created nor destroyed at a conservative margin, the relative motion between adjacent plates must be parallel to the strike of a shared transform fault. Pioneering independent studies by D. P. McKenzie and R. L. Parker (1967) and W. J. Morgan (1968) showed how transform faults could be used to locate the Euler pole of rotation for two plates (see Section 1.2.9). Using this method, X. Le Pichon in 1968 determined the present relative motions of the major tectonic plates. In addition, he derived the history of plate motions in the geological past by incorporating newly available magnetic results from the ocean basins.

1.2.5 Sea-floor spreading

One of the principal stumbling blocks of continental drift was the inability to explain the mechanism by which drift took place. Wegener had invoked forces related to gravity and the Earth’s rotation, which were demonstrably much

![Fig. 1.12 Schematic model illustrating the three types of plate margin. Lightly hachured areas symbolize spreading ridges (constructive margins); darker shaded areas denote subduction zones (destructive margins); dark lines mark transform faults (conservative margins). The figure is drawn relative to the pole of relative motion between plates A and B. Small arrows denote relative motion on transform faults; large arrows show directions of plate motion, which can be oblique to the strike of ridge segments or subduction zones. Arrows in the asthenosphere suggest return flow from destructive to constructive margins.](Image 328x592 to 544x728)
too weak to drive the continents through the resistant basaltic crust. A. Holmes proposed a model in 1944 that closely resembles the accepted plate tectonic model (Holmes, 1965). He noted that it would be necessary to remove basaltic rocks continuously out of the path of an advancing continent, and suggested that this took place at the ocean deeps where heavy eclogite “roots” would sink into the mantle and melt. Convection currents in the upper mantle would return the basaltic magma to the continents as plateau basalts, and to the oceans through innumerable fissures. Holmes saw generation of new oceanic crust as a process that was dispersed throughout an ocean basin. At the time of his proposal the existence of the system of oceanic ridges and rises was not yet known.

The important role of oceanic ridges was first recognized by H. Hess in 1962. He suggested that new oceanic crust is generated from upwelling hot mantle material at the ridges. Convection currents in the upper mantle would rise to the surface at the ridges and then spread out laterally. The continents would ride on the spreading mantle material, carried along passively by the convection currents. In 1961 R. Dietz coined the expression “sea-floor spreading” for the ridge process. This results in the generation of linedate marine magnetic anomalies at the ridges, which record the history of geomagnetic polarity reversals. Study of these magnetic effects led to the verification of sea-floor spreading.

1.2.5.1 The Vine–Matthews–Morley hypothesis

Paleomagnetic studies in the late 1950s and early 1960s of radiometrically dated continental lavas showed that the geomagnetic field has changed polarity at irregular time intervals. For tens of thousands to millions of years the polarity might be normal (as at present), then unaccountably the poles reverse within a few thousand years, so that the north magnetic pole is near the south geographic pole and the north magnetic pole is near the south geographic pole. This state may again persist for a long interval, before the polarity again switches. The ages of the reversals in the last 5 million years have been obtained radiometrically, giving an irregular but dated polarity sequence.

A magnetic anomaly is a departure from the theoretical magnetic field at a given location. If the field is stronger than expected, the anomaly is positive; if it is weaker than expected, the anomaly is negative. In the late 1950s magnetic surveys over the oceans revealed remarkable striped patterns of alternately positive and negative magnetic anomalies over large areas of oceanic crust (Fig. 1.13), for which conventional methods of interpretation gave no satisfactory account. In 1963 the English geophysicists F. J. Vine and D. H. Matthews and, independently, the Canadian geologist L. W. Morley, formulated a landmark hypothesis that explains the origin of the oceanic magnetic anomaly patterns (see also Section 5.7.3).

Observations on dredged samples had shown that basalts in the uppermost oceanic crust carry a strong remnant magnetization (i.e., they are permanently magnetized, like a magnet). The Vine–Matthews–Morley hypothesis integrates this result with the newly acquired knowledge of geomagnetic polarity reversals and the Hess–Dietz concept of sea-floor spreading (Fig. 1.14). The basaltic lava is extruded in a molten state. When it solidifies and its temperature cools below the Curie temperature of its magnetic minerals, the basalt becomes strongly magnetized in the direction of the Earth’s magnetic field at that time.
Along an active spreading ridge, long thin strips of magnetized basaltic crust form symmetrically on opposite sides of the spreading center, each carrying the magnetic imprint of the field in which it formed. Sea-floor spreading can persist for many millions of years at an oceanic ridge. During this time the magnetic field changes polarity many times, forming strips of oceanic crust that are magnetized alternately parallel and opposite to the present field, giving the observed patterns of positive and negative anomalies. Thus, the basaltic layer acts like a magnetic tape recorder, preserving a record of the changing geomagnetic field polarity.

### 1.2.5.2 Rates of sea-floor spreading

The width of a magnetic lineation (or stripe) depends on two factors: the speed with which the oceanic crust moves away from a spreading center, and the length of time that geomagnetic polarity is constantly normal or reversed. The distance between the edges of magnetized crustal stripes can be measured from magnetic surveys at the ocean surface, while the ages of the reversals can be obtained by correlating the oceanic magnetic record with the radiometrically dated reversal sequence determined in subaerial lavas for about the last 4 Ma. When the distance of a given polarity reversal from the spreading axis is plotted against the age of the reversal, a nearly linear relationship is obtained (Fig. 1.15). The slope of the best fitting straight line gives the average half-rate of spreading at the ridge. These are of the order of 10 mm yr$^{-1}$ in the North Atlantic ocean and 40–60 mm yr$^{-1}$ in the Pacific ocean. The calculation applies to the rate of motion of crust on one side of the ridge only. In most cases spreading has been symmetric on each side of the ridge (i.e., the opposite sides are moving away from the ridge at equal speeds), so the full rate of separation at a ridge axis is double the calculated half-rate of spreading (Fig. 1.11).

The rates of current plate motion determined from axial anomaly patterns (Fig. 1.11) are average values over several million years. Modern geodetic methods allow these rates to be tested directly (see Section 2.4.6). Satellite laser-ranging (SLR) and very long baseline interferometry (VLBI) allow exceptionally accurate measurement of changes in the distance between two stations on Earth. Controlled over several years, the distances between pairs of stations on opposite sides of the Atlantic ocean are...
increasing slowly at a mean rate of $17 \text{ mm yr}^{-1}$ (Fig. 1.16). This figure is close to the long-term value of about $20 \text{ mm yr}^{-1}$ interpreted from model NUVEL-1 of current plate motions (Fig. 1.11).

Knowing the spreading rates at ocean ridges makes it possible to date the ocean floor. The direct correlation between polarity sequences measured in continental lavas and derived from oceanic anomalies is only possible for the last 4 Ma or so. Close to the axial zone, where linear spreading rates are observed (Fig. 1.15), simple extrapolation gives the ages of older anomalies, converting the striped pattern into an age map (Fig. 1.13). Detailed magnetic surveying of much of the world’s oceans has revealed a continuous sequence of anomalies since the Late Cretaceous, preceded by an interval in which no reversals occurred; this Quiet Interval was itself preceded by a Mesozoic reversal sequence. Magnetostratigraphy in sedimentary rocks (Section 5.7.4) has enabled the identification, correlation and dating of key anomalies. The polarity sequence of the oceanic anomalies has been converted to a magnetic polarity timescale in which each polarity reversal is accorded an age (e.g., as in Fig. 5.78). In turn, this allows the pattern of magnetic anomalies in the ocean basins to be converted to a map of the age of the ocean basins (Fig. 5.82). The oldest areas of the oceans lie close to northwest Africa and eastern North America, as well as in the northwest Pacific. These areas formed during the early stages of the breakup of Pangaea. They are of Early Jurassic age. The ages of the ocean basins have been confirmed by drilling through the sediment layers that cover the ocean floor and into the underlying basalt layer. Beginning in the late 1960s and extending until the present, this immensely expensive undertaking has been carried out in the Deep Sea Drilling Project (DSDP) and its successor the Ocean Drilling Project (ODP). These multinational projects, under the leadership of the United States, are prime examples of open scientific cooperation on an international scale.

1.2.6 Plate margins

It is important to keep in mind that the tectonic plates are not crustal units. They involve the entire thickness of the lithosphere, of which the crust is only the outer skin. Oceanic lithosphere is thin close to a ridge axis, but thickens with distance from the ridge, reaching a value of
80–100 km; the oceanic crust makes up only the top 5–10 km. Continental lithosphere may be up to 150 km thick, of which only the top 30–60 km is continental crust. Driven by mechanisms that are not completely understood, the lithospheric plates move relative to each other across the surface of the globe. This knowledge supplies the “missing link” in Wegener’s continental drift hypothesis, removing one of the most serious objections to it. It is not necessary for the continents to plow through the rigid ocean basins; they are transported passively on top of the moving plates, as logs float on a stream. Continental drift is thus a consequence of plate motions.

The plate tectonic model involves the formation of new lithosphere at a ridge and its destruction at a subduction zone (Fig. 1.17). Since the mean density of oceanic lithosphere exceeds that of continental lithosphere, oceanic lithosphere can be subducted under continental or oceanic lithosphere, whereas continental lithosphere cannot under-ride oceanic lithosphere. Just as logs pile up where a stream dives under a surface obstacle, a continent that is transported into a subduction zone collides with the deep-sea trench, island arc or adjacent continent. Such a collision results in an orogenic belt. In a continent–continent collision, neither plate can easily subduct, so relative plate motion may come to a halt. Alternatively, subduction may start at a new location behind one of the continents, leaving a mountain chain as evidence of the suture zone between the original colliding continents. The Alpine–Himalayan and Appalachian mountain chains are thought to have formed by this mechanism, the former in Tertiary times, the latter in several stages during the Paleozoic. Plate tectonic theory is supported convincingly by an abundance of geophysical, petrological and geological evidence from the three types of plate margin. A brief summary of the main geophysical observations at these plate margins is given in the following sections. Later chapters give more detailed treatments of the gravity (Section 2.6.4), seismicity (Sections 3.5.3 and 3.5.4), geothermal (Section 4.2.5) and magnetic (Section 5.7.3) evidence.

### 1.2.6.1 Constructive margins

Although the ridges and rises are generally not centrally located in the ocean basins, they are often referred to as mid-ocean ridges. The type of oceanic basalt that is produced at an oceanic spreading center is even called a mid-ocean ridge basalt (MORB for short). Topographically, slow-spreading ridges have a distinct axial rift valley, which, for reasons that are not understood, is missing on faster-spreading ridges. Partially molten upper mantle rocks (generally assumed to be peridotites) from the asthenosphere rise under the ridges. The decrease in pressure due to the changing depth causes further melting and the formation of basaltic magma. Their chemical compositions and the concentrations of long-lived radioactive isotopes suggest that MORB lavas are derived by fractionation (i.e., separation of components, perhaps by precipitation or crystallization) from the upwelling peridotitic mush. Differentiation is thought to take place at about the depth of the lower crustal gabbroic layer beneath the ridge in a small, narrow magma chamber. Some of the fluid magma extrudes near the central rift or ridge axis and flows as lava across the ocean floor; part is intruded as dikes and sills into the thin oceanic crust. The Vine–Matthews–Morley hypothesis for the origin of oceanic magnetic anomalies requires fairly sharp boundaries between alternately magnetized blocks of oceanic crust. This implies that the zone of dike injection is narrow and close to the ridge axis.

The distribution of earthquakes defines a narrow band of seismic activity close to the crest of an oceanic ridge. These earthquakes occur at shallow depths of a few kilometers and are mostly small; magnitudes of 6 or greater are rare. The seismic energy released at ridges is an insignificant part of the world-wide annual release. Analyses show that the earthquakes are associated with normal faulting, implying extension away from the ridge axis (see Section 3.5.4).

Heat flow in the oceans is highest at the ocean ridges and decreases systematically with distance away from the
1.2 THE DYNAMIC EARTH

ridge. The thermal data conform to the model of sea-floor spreading. High axial values are caused by the formation of new lithosphere from the hot uprising magma at the ridge axis. The associated volcanism on the floor of the axial rift zones has been observed directly from deep-diving submersibles. With time, the lithosphere spreads away from the ridge and gradually cools, so that the heat outflow diminishes with increasing age or distance from the ridge.

Oceanic crust is thin, so the high-density mantle rocks occur at shallower depths than under the continents. This causes a general increase of the Earth’s gravity field over the oceans, giving positive gravity anomalies. However, over the ridge systems gravity decreases toward the axis so that a “negative” anomaly is superposed on the normally positive oceanic gravity anomaly. The effect is due to the local density structure under the ridge. It has been interpreted in terms of anomalous mantle material with density slightly less than normal. The density is low because of the different mantle composition under the ridges and its high temperature.

The interpretation of magnetic anomalies formed by sea-floor spreading at constructive margins has already been discussed. The results provide direct estimates of the mean rates of plate motions over geological time intervals.

1.2.6.2 Destructive margins

Subduction zones are found where a plate plunges beneath its neighbor to great depths, until pressure and temperature cause its consumption. This usually happens within a few hundred kilometers, but seismic tomography (Section 3.7.6) has shown that some descending slabs may sink to great depths, even to the core–mantle boundary. Density determines that the descending plate at a subduction zone is an oceanic one. The surface manifestation depends on the type of overriding plate. When this is another oceanic plate, the subduction zone is marked by a volcanic island arc and, parallel to it, a deep trench. The island arc lies near the edge of the overriding plate and is a few hundred kilometers apart. Several examples are seen around the west and northwest margins of the Pacific plate (Fig. 1.11). Melting of the downgoing slab produces magma that rises to feed the volcanoes. The intrusion of magma behind an island arc produces a back-arc basin on the inner, concave side of the arc. These basins are common in the Western Pacific. If the arc is close to a continent, the off-arc magmatism may create a marginal sea, such as the Sea of Japan. Back-arc basins and marginal seas are floored by oceanic crust.

A fine example of where the overriding plate is a continental one is seen along the west coast of South America.

Compression between the Nazca and South American plates has generated the Andes, an arcuate-folded mountain belt near the edge of the continental plate. Active volcanoes along the mountain chain emit a type of lava, called andesite, which has a higher silica content than oceanic basalt. It does not originate from the asthenosphere-type of magma. A current theory is that it may form by melting of the subducting slab and overriding plate at great depths. If some siliceous sediments from the deep-sea trench are carried down with the descending slab, they might enhance the silica content of the melt, producing a magma with andesite-type composition.

The seismicity at a subduction zone provides the key to the processes active there. Where one plate is thrust over the other, the shear causes hazardous earthquakes at shallow depths. Below this region, earthquakes are systematically distributed within the subducting plate. They form an inclined Wadati–Benioff seismic zone, which may extend for several hundred kilometers into the mantle. The deepest earthquakes have been registered down to about 700 km.

Studies of the focal mechanisms (Section 3.5.4) show that at shallow depths the downgoing plate is in a state of down-dip extension (Fig. 1.18a). Subducting lithosphere is colder and denser than the underlying asthenosphere. This gives it negative buoyancy, which causes it to sink, pulling the plate downward. At greater depths the mantle is more rigid than the asthenosphere, and its strength resists penetration (Fig. 1.18b). While the upper part is sinking, the bottom part is being partly supported by the deeper layers; this results in down-dip compression in the lower part of the descending slab and down-dip extension in the upper part. A gap in the depth distribution of seismicity may arise where the deviatoric stress changes from extensional to compressional. In a very deep subduction zone the increase in resistance with depth causes down-dip compression throughout the descending slab (Fig. 1.18c). In some cases part of the slab may break off and
sink to great depths, where the earthquakes have compressional-type mechanisms (Fig. 1.18d); a gap in seismicity exists between the two parts of the slab.

Heat flow at a destructive plate margin reflects to some extent the spreading history of the plate. The plate reaches its maximum age, and so has cooled the furthest, by the time it reaches a subduction zone. The heat flow values over deep ocean basins are uniformly low, but the values measured in deep-sea trenches are the lowest found in the oceans. In contrast, volcanic arcs and back-arc basins often have anomalously high heat flow due to the injection of fresh magma.

Gravity anomalies across subduction zones have several distinctive features. Seaward of the trench the lithosphere flexes upward slightly before it begins its descent, causing a weak positive anomaly; the presence of water or low-density sediments in a deep-sea trench gives rise to a strong negative gravity anomaly; and over the descending slab a positive anomaly is observed, due in part to the mineralogical conversion of subducted oceanic crust to higher-density eclogite.

Subduction zones have no particular magnetic signature. Close to an active or passive continental margin the contrast between the magnetic properties of oceanic and continental crust produces a magnetic anomaly, but this is not a direct result of the plate tectonic processes. Over marginal basins magnetic anomalies are not lined except in some rare cases. This is because the oceanic crust in the basin does not originate by sea-floor spreading at a ridge, but by diffuse intrusion throughout the basin.

1.2.6.3 Conservative margins

Transform faults are strike-slip faults with steeply dipping fault planes. They may link segments of subduction zones, but they are mostly observed at constructive plate margins where they connect oceanic ridge segments. Transform faults are the most seismically active parts of a ridge system, because here the relative motion between neighboring plates is most pronounced. Seismic studies have confirmed that the displacements on transform faults agree with the relative motion between the adjacent plates.

The trace of a transform fault may extend away from a ridge on both sides as a fracture zone. Fracture zones are among the most dramatic features of ocean-floor topography. Although only some tens of kilometers wide, a fracture zone can be thousands of kilometers long. It traces the arc of a small circle on the surface of the globe. This important characteristic allows fracture zones to be used for the deduction of relative plate motions, which cannot be obtained from the strike of a ridge or trench segment, where oblique spreading or subduction is possible (note, for example, the direction of plate convergence relative to the strike of the Aleutian island arc in Fig. 1.11).

Any displacement on the surface of a sphere is equivalent to a small rotation about a pole. The motion of one plate relative to the other takes place as a rotation about the Euler pole of relative rotation between the plates (see Section 1.2.9). This pole can be located from the orientations of fracture zones, because the strike of a transform fault is parallel to the relative motion between two adjacent plates. Thus a great circle normal to a transform fault or fracture zone must pass through the Euler pole of relative rotation between the two plates. If several great circles are drawn at different places on the fracture zone (or normal to different transform faults offsetting a ridge axis) they intersect at the Euler pole. The current model of relative plate motions NUVEL-1 was obtained by determining the Euler poles of rotation between pairs of plates using magnetic anomalies, the directions of slip on earthquake fault planes at plate boundaries, and the topography that defines the strikes of transform faults. The rates of relative motion at different places on the plate boundaries (Fig. 1.11) were computed from the rates of rotation about the appropriate Euler poles.

There may be a large change in elevation across a fracture zone; this is related to the different thermal histories of the plates it separates. As a plate cools, it becomes more dense and less buoyant, so that it gradually sinks. Consequently, the depth to the top of the oceanic lithosphere increases with age, i.e., with distance from the spreading center. Places facing each other across a transform fault are at different distances from their respective spreading centers. They have different ages and so have subsided by different amounts relative to the ridge. This may result in a noticeable elevation difference across the fracture zone.

Ultrabasic rocks are found in fracture zones and there may be local magnetic anomalies. Otherwise, the magnetic effect of a transform fault is to interrupt the oceanic magnetic lineations parallel to a ridge axis, and to offset them in the same measure as ridge segments. This results in a very complex pattern of magnetic lineations in some ocean basins (e.g., in the northeast Pacific).

A transform fault can also connect subduction zones. Suppose a consuming plate boundary consisted originally of two opposed subduction zones (Fig. 1.19a). Plate Y is consumed below plate X along the segment ab of the boundary, whereas plate X is consumed beneath plate Y along segment bc. The configuration is unstable, because a trench cannot sustain subduction in opposite directions. Consequently, a dextral transform fault develops at the point b. After some time, motion on the fault displaces the lower segment to the position b’c (Fig. 1.19b). An example of such a transform boundary is the Alpine fault in New Zealand (Fig. 1.19c). To the northeast of North Island, the Pacific plate is being subducted at the Tonga–Kermadec trench. To the southwest of South Island, the Pacific plate overrides the Tasman Sea at the anomalous Macquarie Ridge (earthquake analysis has
shown that the plate margin at this ridge is compressive; the compression may be too slow to allow a trench to develop. The Alpine fault linking the two opposed subduction zones is therefore a dextral transform fault.

1.2.7 Triple junctions

It is common, although imprecise, to refer to a plate margin by its dominant topographic feature, rather than by the nature of the margin. A ridge (R) represents a constructive margin or spreading center, a trench (T) refers to a destructive margin or subduction zone, and a transform fault (F) stands for a conservative margin. Each margin is a location where two tectonic plates adjoin. Inspection of Fig. 1.11 shows that there are several places where three plates come together, but none where four or more plates meet. The meeting points of three plate boundaries are called triple junctions. They are important in plate tectonics because the relative motions between the plates that form a triple junction are not independent. This may be appreciated by considering the plate motions in a small plane surrounding the junction.

Consider the plate velocities at an RTF junction formed by all three types of boundary (Fig. 1.20a). If the plates are rigid, their relative motions take place entirely at their margins. Let \( V_A \) denote the velocity of plate B relative to plate A, \( V_C \) the velocity of plate C relative to plate B, and \( V_A \) the velocity of plate A relative to plate C. Note that these quantities are vectors; their directions are as important as their magnitudes. They can be represented on a vector diagram by straight lines with directions parallel to and lengths proportional to the velocities. In a circuit about the triple junction an observer must return to the starting point. Thus, a vector diagram of the interplate velocities is a closed triangle (Fig. 1.20b). The velocities are related by

\[
A V_B + B V_C + C V_A = 0 \tag{1.7}
\]

This planar model is a “flat Earth” representation. As discussed in Section 1.2.9, displacements on the surface of a sphere are rotations about Euler poles of relative motion. This can be taken into account by replacing each linear velocity \( V \) in Eq. (1.7) by the rotational velocity \( \omega \) about the appropriate Euler pole.

1.2.7.1 Stability of triple junctions

The different combinations of three plate margins define ten possible types of triple junction. The combinations correspond to all three margins being of one type (RRR, TTT, FFF), two of the same type and one of the other (RTT, RRF, FFT, FFR, TTR, TTF), and all different (RTF). Different combinations of the sense of subduction at a trench increase the number of possible junctions to sixteen. Not all of these junctions are stable in time. For a junction to preserve its geometry, the orientations of the three plate boundaries must fulfill conditions which allow the relative velocities to satisfy Eq. (1.7). If they do so, the junction is stable and can maintain its shape. Otherwise, the junction is unstable and must evolve in time to a stable configuration.

The stability of a triple junction is assessed by considering how it can move along any of the plate boundaries...
Fig. 1.21 Plate margin geometry (left) and locus \( ab \) of a triple junction in velocity space (right) for (a) a trench, (b) a transform fault, and (c) a ridge (after Cox and Hart, 1986).

velocity lines form the sides of a triangle, which can never meet in a point (Fig. 1.22b). The other types of triple junction are conditionally stable, depending on the angles between the different margins. For example, in an RTF triple junction the velocity lines of the trench \( ac \) and transform fault \( bc \) must both pass through \( C \), because this plate is common to both boundaries. The junction is stable if the velocity line \( ab \) of the ridge also passes through \( C \), or if the trench and transform fault have the same trend (Fig. 1.22c). By similar reasoning, the FFT triple junction is only stable if the trench has the same trend as one of the transform faults (Fig. 1.22d).

In the present phase of plate tectonics only a few of the possible types of triple junction appear to be active. An RRR-type is formed where the Galapagos Ridge meets the East Pacific Rise at the junction of the Cocos, Nazca and Pacific plates. A TTT-type junction is formed by the Japan trench and the Bonin and Ryukyu arcs. The San Andreas fault in California terminates in an FFT-type junction at its northern end, where it joins the Mendocino Fracture Zone.

1.2.7.2 Evolution of triple junctions in the northeast Pacific

Oceanic magnetic anomalies in the northeast Pacific form a complex striped pattern. The anomalies can be identified
by interpreting their shapes. Their ages can be found by comparison with a geomagnetic polarity timescale such as that shown in Fig. 5.78, which gives the age of each numbered chron since the Late Jurassic. In the northeast Pacific the anomalies become younger toward the North American continent in the east, and toward the Aleutian trench in the north. The anomaly pattern produced at a ridge is usually symmetric (as in Fig. 1.13), but in the northeast Pacific only the western half of an anomaly pattern is observed. The plate on which the eastern half of the anomaly pattern was formed is called the Farallon plate. It and the ridge itself are largely missing and have evidently been subducted under the American plate. Only two small remnants of the Farallon plate still exist: the Juan de Fuca plate off the coast of British Columbia, and the Rivera plate at the mouth of the Gulf of California. The magnetic anomalies also indicate that another plate, the Kula plate, existed in the Late Mesozoic but has now been entirely consumed under Alaska and the Aleutian trench. The anomaly pattern shows that in the Late Cretaceous the Pacific, Kula and Farallon plates were diverging from each other and thus met at an RRR-type triple junction. This type of junction is stable and preserved its shape during subsequent evolution of the plates. It is therefore possible to reconstruct the relative motions of the Pacific, Kula and Farallon plates in the Cenozoic (Fig. 1.23a–c).

The anomaly ages are known from the magnetic timescale so the anomaly spacing allows the half-rates of spreading to be determined. In conjunction with the trends of fracture zones, the anomaly data give the rates and directions of spreading at each ridge. The anomaly pattern at the mouth of the Gulf of California covers the last 4 Ma and gives a mean half-rate of spreading of 3 cm yr\(^{-1}\) parallel to the San Andreas fault. This indicates that the Pacific plate has moved northward past the American plate at this boundary with a mean relative velocity of about 6 cm yr\(^{-1}\) during the last 4 Ma. The half-rate of spreading on the remnant of the Farallon–Pacific ridge is 5 cm yr\(^{-1}\), giving a relative velocity of 10 cm yr\(^{-1}\) between the plates. A vector diagram of relative velocities at the Farallon–Pacific–American triple junction (Fig. 1.23d) shows convergence of the Farallon plate on the American plate at a rate of 7 cm yr\(^{-1}\). Similarly, the spacing of east–west trending magnetic anomalies in the Gulf of Alaska gives the half-rate of spreading on the Kula–Pacific ridge, from which it may be inferred that the relative velocity between the plates was 7 cm yr\(^{-1}\). A vector diagram combining this value with the 6 cm yr\(^{-1}\) northward motion of the Pacific plate gives a velocity of 12 cm yr\(^{-1}\) for the Kula plate relative to the American plate.

Using these velocities the history of plate evolution in the Cenozoic can be deduced by extrapolation. The interpretation is tenuous, as it involves unverifiable assumptions. The most obvious is that the Kula–Pacific motion in the late Cretaceous (80 Ma ago) and the American–Pacific motion of the past 4 Ma have remained constant throughout the Cenozoic. With this proviso, it is evident that triple junctions formed and migrated along the American plate margin. The Kula–American–Farallon RTF junction was slightly north of the present location of San Francisco 60 Ma ago (Fig. 1.23c); it moved to a position north of Seattle 20 Ma ago (Fig. 1.23a). Around that time in the Oligocene an FFT junction formed between San Francisco and Los Angeles, while the Farallon–Pacific–American RTF junction evolved to the south. The development of these two triple junctions is due to the collision and subduction of the Farallon–Pacific ridge at the Farallon–American trench.

At the time of magnetic anomaly 13, about 34 Ma ago, a north–south striking ridge joined the Mendocino and Murray transform faults as part of the Farallon–Pacific
plate margin to the west of the American trench (Fig. 1.24a). By the time of anomaly 9, about 27 Ma ago, the ridge had collided with the trench and been partly consumed by it (Fig. 1.24b). The Farallon plate now consisted of two fragments: an FFT junction developed at point 1, formed by the San Andreas fault system, the Mendocino fault and the consuming trench to the north; and an RTF junction formed at point 2. Both junctions are stable when the trenches are parallel to the transform fault along the San Andreas system. Analysis of the velocity diagrams at each triple junction shows that point 1 migrated to the northwest and point 2 migrated to the southeast at this stage. Later, when the southern segment of the Farallon–Pacific ridge had been subducted under the American plate, the Murray transform fault changed the junction at point 2 to an FFT junction, which has subsequently also migrated to the northwest.

1.2.8 Hotspots

In 1958 S. W. Carey coined the term “hot spot” – now often reduced to “hotspot” – to refer to a long-lasting center of surface volcanism and locally high heat flow. At one time more than 120 of these thermal anomalies were proposed. Application of more stringent criteria has reduced their number to about 40 (Fig. 1.25). The hotspots may occur on the continents (e.g., Yellowstone), but are more common in the ocean basins. The oceanic hotspots are associated with depth anomalies. If the observed depth is compared with the depth predicted by cooling models of the oceanic lithosphere, the hotspots are found to lie predominantly in broad shallow regions, where the lithosphere apparently swells upward. This elevates denser mantle material, which creates a mass anomaly and disturbs the geoid; the effect is partially mitigated by reduced density of material in the hot, rising plume. The geoid surface is also displaced by subduction zones. The residual geoid obtained by removing the effects associated with cold subducting slabs shows a remarkable correlation with the distribution of hotspots (Fig. 1.25). The oceanic hotspots are found in conjunction with intraplate island chains, which provide clues to the origin of hotspots and allow them to be used for measuring geodynamic processes.

Two types of volcanic island chains are important in plate tectonics. The arcuate chains of islands associated with deep oceanic trenches at consuming plate margins are related to the process of subduction and have an arcuate shape. Nearly linear chains of volcanic islands are observed within oceanic basins far from active plate margins. These intraplate features are particularly evident on a bathymetric map of the Pacific Ocean. The Hawaiian, Marquesas, Society and Austral Islands form subparallel chains that trend approximately perpendicular to the axis of ocean-floor spreading on the East Pacific rise. The most closely studied is the Hawaiian Ridge (Fig. 1.26a). The volcanism along this chain decreases from present-day activity at the southeast, on the island of Hawaii, to long extinct seamounts and guyots towards the northwest along the Emperor Seamount chain. The history of development of the chain is typical of other linear volcanic island chains in the Pacific basin (Fig. 1.26b). It was explained in 1963 by J. T. Wilson, before the modern theory of plate tectonics was formulated.
1.2 THE DYNAMIC EARTH

A hotspot is a long-lasting magmatic center rooted in the mantle below the lithosphere. A volcanic complex is built up above the magmatic source, forming a volcanic island or, where the structure does not reach sea-level, a seamount. The motion of the plate transports the island away from the hotspot and the volcanism becomes extinct. The upwelling material at the hotspot elevates the ocean floor by up to 1500 m above the normal depth of the ocean floor, creating a depth anomaly. As they move away from the hotspot the by now extinct volcanic islands sink beneath the surface; some are truncated by erosion to sea-level and become guyots. Coral atolls may accumulate on some guyots. The volcanic chain is aligned with the motion of the plate.

Confirmation of this theory is obtained from radiometric dating of basalts samples from islands and seamounts along the Hawaiian Ridge portion of the Hawaiian–Emperor chain. The basalts increase in age with distance from the active volcano Kilauea on the island of Hawaii (Fig. 1.27). The trend shows that the average rate of motion of the Pacific plate over the Hawaiian hotspot has been about 10 cm yr\(^{-1}\) during the last 20–40 Ma. The change in trend between the Hawaiian Ridge and the Emperor Seamount chain indicates a change in direction and speed of the Pacific plate about 43 Ma ago, at which time there was a global reorganization of plate motions. The earlier rate of motion along the Emperor chain is less well determined but is estimated to be about 6 cm yr\(^{-1}\).

Radiometric dating of linear volcanic chains in the Pacific basin gives almost identical rates of motion over their respective hotspots. This suggests that the hotspots form a stationary network, at least relative to the lithosphere. The velocities of plate motions over the hotspots...
are therefore regarded as absolute velocities, in contrast to the velocities derived at plate margins, which are the relative velocities between neighboring plates. The assumption that the hotspots are indeed stationary has been contested by studies that have yielded rates of inter-hotspot motion of the order of 1.5–2 cm yr\(^{-1}\) (comparable to present spreading rates in the Atlantic). Thus, the notion of a stationary hotspot reference frame may only be valid for a limited time interval. Nevertheless, any motions between hotspots are certainly much slower than the motions of plates, so the hotspot reference frame provides a useful guide to absolute plate motions over the typical time interval (~10 Ma) in which incremental sea-floor spreading is constant.

As well as geophysical evidence there are geochemical anomalies associated with hotspot volcanism. The type of basalt extruded at a hotspot is different from the andesitic basalts formed in subduction zone magmatism. It also has a different petrology from the mid-oceanic ridge basalts (MORB) formed during sea-floor spreading and characteristic of the ocean floor. The hotspot source is assumed to be a mantle plume that reaches the surface. Mantle plumes are fundamental features of mantle dynamics, but they remain poorly understood. Although they are interpreted as long-term features it is not known for how long they persist, or how they interact with convective processes in the mantle. Their role in heat transport and mantle convection, with consequent influence on plate motions, is believed to be important but is uncertain. Their sources are controversial. Some interpretations favor a comparatively shallow origin above the 670 km discontinuity, but the prevailing opinion appears to be that the plumes originate in the D\(^{''}\) layer at the core–mantle boundary. This requires the mantle plume to penetrate the entire thickness of the mantle (see Fig. 4.38). In either case the stationary nature of the hotspot network relative to the lithosphere provides a reference frame for determining absolute plate motions, and for testing the hypothesis of true polar wander.

1.2.9 Plate motion on the surface of a sphere

One of the great mathematicians of the eighteenth century was Leonhard Euler (1707–1783) of Switzerland. He made numerous fundamental contributions to pure mathematics, including to complex numbers (see Box 2.6) and spherical trigonometry (see Box 1.4). A corollary of one of his theorems shows that the displacement of a rigid body on the surface of a sphere is equivalent to a rotation about an axis that passes through its center. This is applicable to the motion of a lithospheric plate.

Any motion restricted to the surface of a sphere takes place along a curved arc that is a segment of either a great circle (centered, like a “circle of longitude,” at the Earth’s center) or a small circle. Small circles are defined relative to a pole of rotational symmetry (such as the geographical pole, when we define “circles of latitude”). A point on the surface of the sphere can be regarded as the end-point of a radius vector from the center of the Earth to the point. Any position on the spherical surface can be speci-
The sides of a triangle on a plane surface are straight lines and the sum of its internal angles is 180° (or \( \pi \) radians). Let the angles be \( A \), \( B \) and \( C \) and the lengths of the sides opposite each of these angles be \( a \), \( b \) and \( c \), as in Fig. B1.4a. The sizes of the angles and the lengths of the sides are governed by the sine law:

\[
\frac{\sin A}{a} = \frac{\sin B}{b} = \frac{\sin C}{c}
\]

(1)

The length of any side is related to the lengths of the other two sides and to the angle they include by the cosine law, which for the side \( a \) is

\[ a^2 = b^2 + c^2 - 2bc \cos A \]

(2)

with similar expressions for the sides \( b \) and \( c \).

The sides of a triangle on a spherical surface are great circle arcs and the sum of the internal angles is greater than 180°. The angle between two great circles at their point of intersection is defined by the tangents to the great circles at that point. Let the angles of a spherical triangle be \( A \), \( B \) and \( C \), and let the lengths of the sides opposite each of these angles be \( a \), \( b \) and \( c \), respectively, as in Fig. B1.4b. The lengths of the sides may be converted to the angles they subtend at the center of the Earth. For example, the distance from pole to equator on the Earth's surface may be considered as 10,007 km or as 90 degrees of arc. Expressing the sides of the spherical triangle as angles of arc, the law of sines is

\[
\frac{\sin A}{\cos a} = \frac{\sin B}{\cos b} = \frac{\sin C}{\cos c}
\]

(3)

and the law of cosines is

\[ \cos a = \cos b \cos c + \sin b \sin c \cos A \]

(4)

1.2.9.1 Euler poles of rotation

Geophysical evidence does not in itself yield absolute plate motions. Present-day seismicity reflects relative motion between contiguous plates, oceanic magnetic anomaly patterns reveal long-term motion between neighboring plates, and paleomagnetism does not resolve displacements in longitude about a paleopole. The relative motion between plates is described by keeping one plate fixed and moving the other one relative to it; that is, we rotate it away from (or toward) the fixed plate (Fig. 1.28). The geometry of a rigid plate on the surface of a sphere is outlined by a set of bounding points, which maintain fixed positions relative to each other. Provided it remains rigid, each point of a moving plate describes an arc of a different small circle about the same Euler pole. Thus, the motion between plates is equivalent to a relative rotation about their mutual Euler rotation pole.

The traces of past and present-day plate motions are recorded in the geometries of transform faults and fracture zones, which mark, respectively, the present-day and earlier locations of conservative plate margins. A segment of a transform fault represents the local path of
relative motion between two plates. As such, it defines a small circle about the Euler pole of relative rotation between the plates. Great circles drawn normal to the strike of the small circle (transform fault) should meet at the Euler pole (Fig. 1.29a), just as at the present day circles of longitude are perpendicular to circles of latitude and converge at the geographic pole. In 1968, W. J. Morgan first used this method to locate the Euler rotation pole for the present-day plate motion between America and Africa (Fig. 1.29b). The Caribbean plate may be absorbing slow relative motion, but the absence of a well-defined seismic boundary between North and South America indicates that these plates are now moving essentially as one block. The great circles normal to transform faults in the Central Atlantic converge and intersect close to $58^\circ 110^34_1$ N $36^\circ 110^34_1$ W, which is an estimate of the Euler pole of recent motion between Africa and South America. The longitude of the Euler pole is determined more precisely than its latitude, the errors being $\pm 2^\circ$ and $\pm 5^\circ$, respectively. When additional data from earthquake first motions and spreading rates are included, an Euler pole at $62^\circ 110^34_1$ N $36^\circ 110^34_1$ W is obtained, which is within the error of the first location.

The “Bullard-type fit” of the African and South American coastlines (Section 1.2.2.2) is obtained by a rotation about a pole at $44^\circ 110^34_1$ N $31^\circ 110^34_1$ W. This pole reflects the average long-term motion between the continents. A rotation which matches a starting point with an endpoint is a finite rotation. As the difference between the present-day and age-averaged Euler poles illustrates, a finite rotation is a mathematical formality not necessarily related to the actual motion between the plates, which may consist of a number of incremental rotations about different poles.

Box 1.5: Direction cosines

It is often useful to express a direction with the aid of direction cosines. These are the cosines of the angles that the direction makes with the reference axes. Define the z-axis along the Earth’s spin axis, the x-axis along the Greenwich meridian and the y-axis normal to both of these, as in Fig. B1.5. If a line makes angles $\alpha_x$, $\alpha_y$, and $\alpha_z$ to the x-, y- and z-axes, respectively, its direction cosines with respect to these axes are

$$\begin{align*}
l &= \cos \alpha_x \\
m &= \cos \alpha_y \\
n &= \cos \alpha_z.
\end{align*}$$

Consider a position P on the Earth’s surface with latitude $\lambda$ and longitude $\phi$. A line of length $R$ from the center of the Earth to the point P has projections $R \cos \alpha_z (= R \sin \lambda)$ on the z-axis and $R \cos \alpha_x (= R \cos \lambda)$ in the equatorial plane. The latter has projections $(R \cos \alpha \cos \phi)$ and $(R \cos \alpha \sin \phi)$ on the x- and y-axes, respectively. The direction cosines of the line are thus

$$\begin{align*}
l &= \cos \lambda \cos \phi \\
m &= \cos \lambda \sin \phi \\
n &= \sin \lambda
\end{align*}$$

The angle $\Delta$ between two lines with direction cosines $(l_1, m_1, n_1)$ and $(l_2, m_2, n_2)$ is given by

$$\cos \Delta = l_1 l_2 + m_1 m_2 + n_1 n_2$$

These relationships are useful for computing great circle distances and the angular relationships between lines.

**Fig. B1.5** The definition of direction cosines.

**Fig. 1.28** Illustration that the displacement of a rigid plate on the surface of a sphere is equivalent to the rotation of the plate about an Euler pole (after Morgan, 1968)
1.2.9.2 Absolute plate motions

The axial dipole hypothesis of paleomagnetism states that the mean geomagnetic pole – averaged over several tens of thousands of years – agrees with the contemporaneous geographic pole (i.e., rotation axis). Paleomagnetic directions allow the calculation of the apparent pole position at the time of formation of rocks of a given age from the same continent. By connecting the pole positions successively in order of their age, an apparent polar wander (APW) path is derived for the continent. Viewed from the continent it appears that the pole (i.e., the rotation axis) has moved along the APW path. In fact, the path records the motion of the lithospheric plate bearing the continent, and differences between APW paths for different plates reflect motions of the plates relative to each other.

During the displacement of a plate (i.e., when it rotates about an Euler pole), the paleomagnetic pole positions obtained from rocks on the plate describe a trajectory which is the arc of a small circle about the Euler pole. The motion of the plate over an underlying hotspot leaves a trace that is also a small circle arc about the same hotspot. The paleomagnetic record gives the motion of plates relative to the rotation axis, whereas the hotspot record shows the plate motion over a fixed point in the mantle. If the mantle moves relative to the rotation axis, the network of hotspots – each believed to be anchored to the mantle – shifts along with it. This motion of the mantle deeper than the mobile lithosphere is called true polar wander (TPW). The term is rather a misnomer, because it refers to motion of the mantle relative to the rotation axis.

Paleomagnetism provides a means of detecting whether long-term true polar wander has taken place. It involves comparing paleomagnetic poles from hotspots with contemporaneous poles from the stable continental cratons. Consider first the possibility that TPW does not take place: each hotspot maintains its position relative to the rotation axis. A lava that is magnetized at an active hotspot acquires a direction appropriate to the distance from the pole. If the plate moves from north to south over the stationary hotspot, a succession of islands and seamounts (Fig. 1.31a, A–D) is formed, which, independently of their age, have the same magnetization direction. Next, suppose that true polar wander does take place: each hotspot moves with time relative to the rotation axis. For simplicity, let the hotspot migration also be from north to south (Fig. 1.31b). Seamount A is being formed at present and its magnetization direction corresponds to the present-day distance from the pole. If the plate moves from north to south over the stationary hotspot, a succession of islands and seamounts (Fig. 1.31a, A–D) is formed, which, independently of their age, have the same magnetization direction. Next, suppose that true polar wander does take place: each hotspot moves with time relative to the rotation axis. For simplicity, let the hotspot migration also be from north to south (Fig. 1.31b). Seamount A is being formed at present and its magnetization direction corresponds to the present-day distance from the pole. However, older seamounts B, C and D were formed closer to the pole and have progressively steeper inclinations the further south they are. The change in paleomagnetic direction with age of the volcanism along the hotspot trace is evidence for true polar wander.

To test such a hypothesis adequately a large number of data are needed. The amount of data from a single plate, such as Africa, can be enlarged by using data from other
The Earth as a planet

plates. For example, in reconstructing Gondwanaland, South America is rotated into a matching position with Africa by a finite rotation about an Euler pole. The same rotation applied to the APW path of South America allows data from both continents to be combined. Likewise, rotations about appropriate Euler poles make the paleomagnetic records for North America and Eurasia accessible. Averaging the pooled data for age-windows 10 Ma apart gives a reconstructed paleomagnetic APW path for Africa (Fig. 1.32a). The next step is to determine the motions of plates over the network of hotspots, assuming the hotspots have not moved relative to each other. A “hotspot” apparent polar wander path is obtained, which is the track of an axis in the hotspot reference frame presently at the north pole. The appearance of this track relative to Africa is shown in Fig. 1.32b.

We now have records of the motion of the lithosphere relative to the pole, and of the motion of the lithosphere relative to the hotspot reference frame. The records coincide for the present time, both giving pole positions at the present-day rotation axis, but they diverge with age as a result of true polar wander. A paleomagnetic pole of a given age is now moved along a great circle (i.e., rotated about an Euler pole in the equatorial plane) until it lies on the rotation axis. If the same rotation is applied to the hotspot pole of the same age, it should fall on the rotation axis also. The discrepancy is due to motion of the hotspot reference frame relative to the rotation axis. Joining locations in order of age gives a true polar wander path (Fig. 1.32c). This exercise can be carried out for only the last 200 Ma, in which plate reconstructions can be confidently made. The results show that TPW has indeed taken place but that its amplitude has remained less than 15° for the last 150 Ma.

1.2.10 Forces driving plate tectonic motions

An unresolved problem of plate tectonics is what mechanism drives plate motions. The forces acting on plates may be divided into forces that act on their bottom surfaces and forces that act on their margins. The bottom forces arise due to relative motion between the lithospheric plate and the viscous asthenosphere. In this context it is less important whether mantle flow takes place by whole-mantle convection or layered convection. For plate tectonics the important feature of mantle rheology is that viscous flow in the upper mantle is possible. The motion vectors of lithospheric plates do not reveal directly the mantle flow pattern, but some general inferences can be drawn. The flow pattern must include the mass transport involved in moving lithosphere from a ridge to a subduction zone, which has to be balanced by return flow deep in the mantle. Interactions between the plates and the viscous substratum necessarily influence the plate motions. In order to assess the importance of these effects we need to compare them to the other forces that act on plates, especially at their boundaries (Fig. 1.33).

1.2.10.1 Forces acting on lithospheric plates

Some forces acting on lithospheric plates promote motion while others resist it. Upper mantle convection could fall into either category. The flow of material beneath a plate exerts a mantle drag force \( F_{DP} \) on the base of the plate. If the convective flow is faster than plate velocities, the plates are dragged along by the flow, but if the opposite is true the mantle drag opposes the plate motion. Plate velocities are observed to be inversely related to the area of continent on the plate, which suggests that the greater lithospheric thickness results in an additional continental drag force \( F_{CD} \) on the plate. The velocity of a plate also depends on the length of its subduction zone but not on the length of its spreading ridge. This suggests that subduction forces may be more important than spreading forces. This can be evaluated by considering the forces at all three types of plate margin.

At spreading ridges, upwelling magma is associated with the constructive margin. It was long supposed that this process pushes the plates away from the ridge. It also elevates the ridges above the oceanic abyss, so that potential energy encourages gravitational sliding toward the trenches. Together, the two effects make up the ridge push force \( F_{RP} \).

At transform faults, high seismicity is evidence of interactive forces where the plates move past each other. A transform force \( F_{TR} \) can be envisioned as representing frictional resistance in the contact zone. Its magnitude may be different at a transform connecting ridge segments,
where the plates are hot, than at a transform between subduction zones, where the plates are cold.

At subduction zones, the descending slab of lithosphere is colder and denser than the surrounding mantle. This creates a positive mass anomaly – referred to as negative buoyancy – which is accentuated by intraplate phase transitions. If the descending slab remains attached to the surface plate, a slab pull force ($F_{SP}$) ensues that pulls the slab downwards into the mantle. Transferred to the entire plate it acts as a force toward the subduction zone. However, the subducting plate eventually sinks to depths where it approaches thermal equilibrium with the surrounding mantle, loses its negative buoyancy and experiences a slab resistance force ($F_{SR}$) as it tries to penetrate further into the stiffer mantle.

Plate collisions result in both driving and resistive forces. The vertical pull on the descending plate may cause the bend in the lower plate to migrate away from the subduction zone, effectively drawing the upper plate toward the trench. The force on the upper plate has also been termed “trench suction” ($F_{SU}$). The colliding plates also impede each other’s motion and give rise to a collision-resistance force ($F_{CR}$). This force consists of separate forces due to the effects of mountains or trenches in the zone of convergence.

At hotspots, the transfer of mantle material to the lithosphere may result in a hotspot force ($F_{HS}$) on the plate.

In summary, the driving forces on plates are slab pull, slab suction, ridge push and the trench pull force on the upper plate. The motion is opposed by slab resistance, collision resistance, and transform fault forces. Whether the forces between plate and mantle (mantle drag, continental drag) promote or oppose motion depends on the sense of
implies that upwelling at ridges is a passive feature, with sion of local processes in the oceanic lithosphere. This are observed, indicating that the topography is an expres-
sets beyond the ends of ridge segments. Instead, sharp o
ff exhibit discontinuities at the faults but would bulge due to buoyant upwelling, the fluid mantle could not is o
ff set by transform faults. If the ridge topography were is o
importance. Moreover, the topography of oceanic ridges verging margin, and it is considered to be of secondary
ridge push force is much smaller than the forces at a con-
gible in comparison to the other forces (Fig. 1.34). The hotspots and the resistance at transform faults are negli-
der relative magnitudes of the torques. The push exerted by these several forces. Some can be shown to be more impor-
table motions. The torque analysis shows that the strongest force driving plate motions is the pull of a descending slab on its plate; the force that pulls the upper plate toward a trench may also be considerable. The opposing force due to the collision between the plates is consistently smaller than the upper plate force. The resistance experienced by some slabs to deep mantle penetration may diminish the slab pull force. However, seismic evidence has shown that some slabs may become detached from their parent plate, and apparently sink all the way to the core–mantle boundary. The descending motion contributes to mantle circulation, and thus acts indirectly as a driving force for plate motions; it is known as slab suction. However, analysis of this force has shown that it is less important than slab pull, which emerges as the most important force driving plate motions.

1.3 SUGGESTIONS FOR FURTHER READING

Introductory level


Intermediate level


Advanced level


1.5 EXERCISES

1.5 REVIEW QUESTIONS

1. Write down Kepler’s three laws of planetary motion. Which law is a result of the conservation of momentum? Which law is a result of the conservation of energy?

2. The gravitational attraction of the Sun on an orbiting planet is equal to the centripetal acceleration of the planet. Show for a circular orbit that this leads to Kepler’s third law of motion.

3. What causes the precession of the Earth’s rotation axis? Why is it retrograde?

4. What other long-term changes of the rotation axis or the Earth’s orbit occur? What are the periods of these motions? What are their causes?

5. If a planet existed in place of the asteroid belt, what would Bode’s law predict for the radius of its orbit? What would be the period of its orbital motion around the Sun?

6. What is the nebular hypothesis for the origin of the solar system?

7. What geological evidence is there in support of continental drift? What is the essential difference between older models of continental drift and the modern theory of plate tectonics?

8. What was Pangaea? When and how did it form? When and how did it break up?

9. What is the Earth’s crust? What is the lithosphere? How are they distinguished?

10. What are the major discontinuities in the Earth’s internal structure? How are they known?

11. Distinguish between constructive, conservative and destructive plate margins.

12. Make a brief summary, using appropriate sketches, of the oceanic and continental collision zones. How does this age compare to the age of the crust?

13. What kind of plate margin is a continental collision zone? How does it differ from a subduction zone?


15. Explain how sea-floor spreading can be used to determine the age of the oceanic crust. Where are the oldest parts of the oceans? How old are they? How does this age compare to the age of the Earth?

16. What are the names of the 12 major tectonic plates and where do their plate margins lie?

17. With the aid of a globe or map, estimate roughly a representative distance across one of the major plates. What is the ratio of this distance to the thickness of the plate? Why are the tectonic units called plates?

18. What is a triple junction? Explain the role of triple junctions in plate tectonics.

19. What is a hotspot? Explain how the Hawaiian hotspot provides evidence of a change in motion of the Pacific plate.

20. How may the Euler pole of relative rotation between two plates be located?

1.5 EXERCISES

1. Measured from a position on the Earth’s surface at the equator, the angle between the direction to the Moon and a reference direction (distant star) in the plane of the Moon’s orbit is 11°57’ at 8 p.m. one evening and 14°32’ at 4 a.m. the following morning. Assuming that the Earth, Moon and reference star are in the same plane, and that the rotation axis is normal to the plane, estimate the approximate distance between the centres of the Earth and Moon.

2. The eccentricity $e$ of the Moon’s orbit is 0.0549 and the mean orbital radius $r_1 = (ab)^{1/2}$ is 384,100 km.

(a) Calculate the lengths of the principal axes $a$ and $b$ of the Moon’s orbit.

(b) How far is the center of the Earth from the center of the elliptical orbit?

(c) Calculate the distances of the Moon from the Earth at perigee and apogee.

3. If the Moon’s disk subtends a maximum angle of 0° 31’ 36.8” at the surface of the Earth, what is the Moon’s radius?

4. Bode’s Law (Eq. (1.3)) gives the orbital radius of the $n$th planet from the Sun (counting the asteroid belt) in astronomical units. It fits the observations well except for Neptune ($n = 9$) and Pluto ($n = 10$). Calculate the orbital radii of Neptune and Pluto predicted by Bode’s Law, and compare the results with the observed values (Table 1.2). Express the discrepancies as percentages of the predicted distances.

5. An ambulance passes a stationary observer at the side of the road at a speed of 60 km h$^{-1}$. Its dual tone siren emits alternating tones with frequencies of 700 and 1700 Hz. What are the dual frequencies heard by the observer (a) before and (b) after the ambulance passes? [Assume that the speed of sound, $c$, in m s$^{-1}$ at the temperature $T(°C)$ is given by $c = 331 + 0.607T$.]

6. A spacecraft landing on the Moon uses the Doppler effect on radar signals transmitted at a frequency of $5$ GHz to determine the landing speed. The pilot discovers that the precision of the radar instrument has deteriorated to ±100 Hz. Is this adequate to ensure a safe landing? [Speed of light $300,000$ km s$^{-1}$.]

7. Explain with the aid of a sketch the relationship between the length of a day and the length of a year on the planet Mercury (see Section 1.1.3.2).

8. The rotations of the planet Pluto and its moon Charon about their own axes are synchronous with the revolution of Charon about Pluto. Show with the
aid of simple sketches that Pluto and Charon always present the same face to each other.

9. The barycenter of a star and its planet – or of a planet and its moon – is the center of mass of the pair. Using the mass and radius of primary body and satellite, and the orbital radius of the satellite, as given in Tables 1.1–1.3 or below, calculate the location of the barycenter of the following pairs of bodies. In each case, does the barycenter lie inside or outside the primary body?
   (a) Sun and Earth.
   (b) Sun and Jupiter.
   (c) Earth and Moon.
   (d) Pluto (mass \(1.27 \times 10^{22}\) kg, radius 1137 km) and Charon (mass \(1.9 \times 10^{21}\) kg, radius 586 km); the radius of Charon's orbit is 19,640 km.

10. A planet with radius \(R\) has a mantle with uniform density \(\rho_m\) enclosing a core with radius \(r_c\) and uniform density \(\rho_c\). Show that the mean density of the planet \(\rho\) is given by

\[
\frac{\rho - \rho_m}{\rho_c - \rho_m} = \left(\frac{r_c}{R}\right)^3
\]

11. The radius of the Moon is 1738 km and its mean density is 3347 kg m\(^{-3}\). If the Moon has a core with radius 400 km and the uniform density of the overlying mantle is 3300 kg m\(^{-3}\), what is the density of the core?

12. Summarize the geological and geophysical evidence resulting from plate tectonic activity in the following regions: (a) Iceland, (b) the Aleutian islands, (c) Turkey, (d) the Andes, (e) the Alps?

13. Using the data in Fig 5.77, compute the approximate spreading rates in the age interval 25–45 Ma at the oceanic ridges in the S. Atlantic, S. Indian, N. Pacific and S. Pacific oceans.

14. Three ridges A, B and C meet at a triple junction. Ridge A has a strike of 329° (N31W) and a spreading rate of 7.0 cm yr\(^{-1}\); ridge B strikes at 233° (S53W) and has a spreading rate of 5.0 cm yr\(^{-1}\). Determine the strike of ridge C and its spreading rate.

15. Three sides of a triangle on the surface of a sphere measure 900 km, 1350 km, and 1450 km, respectively. What are the internal angles of the triangle? If this were a plane triangle, what would the internal angles be?

16. An aircraft leaves a city at latitude \(\lambda_1\) and longitude \(\phi_1\) and flies to a second city at latitude \(\lambda_2\) and longitude \(\phi_2\). Derive an expression for the great circle distance between the two cities.

17. Apply the above formula to compute the great circle distances between the following pairs of cities:
   (a) New York \((\lambda = 40^\circ 43'\ N, \phi_1 = 73^\circ 1'\ W)\) – Madrid \((40^\circ 25'\ N, 3^\circ 43'\ E)\);
   (b) Seattle \((\lambda = 47^\circ 21'\ N, \phi_1 = 122^\circ 12'\ W)\) – Sydney \((\lambda = 33^\circ 52'\ S, \phi_1 = 151^\circ 13'\ E)\);
   (c) Moscow \((\lambda = 55^\circ 45'\ N, \phi_1 = 37^\circ 35'\ E)\) – Paris \((\lambda = 48^\circ 52'\ N, \phi_1 = 2^\circ 20'\ E)\);
   (d) London \((\lambda = 51^\circ 30'\ N, \phi_1 = 0^\circ 10'\ W)\) – Tokyo \((\lambda = 35^\circ 42'\ N, \phi_1 = 139^\circ 46'\ E)\).

18. Calculate the heading (azimuth) of the aircraft’s flight path as it leaves the first city in each pair of cities in the previous exercise.
2.1 THE EARTH’S SIZE AND SHAPE

2.1.1 Earth’s size

The philosophers and savants in ancient civilizations could only speculate about the nature and shape of the world they lived in. The range of possible travel was limited and only simple instruments existed. Unrelated observations might have suggested that the Earth’s surface was upwardly convex. For example, the Sun’s rays continue to illuminate the sky and mountain peaks after its disk has already set, departing ships appear to sink slowly over the horizon, and the Earth’s shadow can be seen to be curved during partial eclipse of the Moon. However, early ideas about the heavens and the Earth were intimately bound up with concepts of philosophy, religion and astrology. In Greek mythology the Earth was a disk-shaped region embracing the lands of the Mediterranean and surrounded by a circular stream, Oceanus, the origin of all the rivers. In the sixth century BC the Greek philosopher Anaximander visualized the heavens as a celestial sphere that surrounded a flat Earth at its center. Pythagoras (582–507 BC) and his followers were apparently the first to speculate that the Earth was a sphere. This idea was further propounded by the influential philosopher Aristotle (384–322 BC). Although he taught the scientific principle that theory must follow fact, Aristotle is responsible for the logical device called syllogism, which can explain correct observations by apparently logical accounts that are based on false premises. His influence on scientific methodology was finally banished by the scientific revolution in the seventeenth century.

The first scientifically sound estimate of the size of the terrestrial sphere was made by Eratosthenes (275–195 BC), who was the head librarian at Alexandria, a Greek colony in Egypt during the third century BC. Eratosthenes had been told that in the city of Syene (modern Aswan) the Sun’s noon rays on midsummer day shone vertically and were able to illuminate the bottoms of wells, whereas on the same day in Alexandria shadows were cast. Using a sun-dial Eratosthenes observed that at the summer solstice the Sun’s rays made an angle of one-fiftieth of a circle (7.2°) with the vertical in Alexandria (Fig. 2.1). Eratosthenes believed that Syene and Alexandria were on the same meridian. In fact they are slightly displaced; their geographic coordinates are 24° 5’N 32° 56’E and 31° 13’N 29° 55’E, respectively. Syene is actually about half a degree north of the tropic of Cancer. Eratosthenes knew that the approximate distance from Alexandria to Syene was 5000 stadia, possibly estimated by travellers from the number of days (“10 camel days”) taken to travel between the two cities. From these observations Eratosthenes estimated that the circumference of the global sphere was 250,000 stadia. The Greek stadiam was the length (about 185 m) of the U-shaped racecourse on which footraces and other athletic events were carried out. Eratosthenes’ estimate of the Earth’s circumference is equivalent to 46,250 km, about 15% higher than the modern value of 40,030 km.

Estimates of the length of one meridian degree were made in the eighth century AD during the Tang dynasty in China, and in the ninth century AD by Arab astronomers in Mesopotamia. Little progress was made in Europe until the early seventeenth century. In 1662 the Royal Society was founded in London and in 1666 the Académie Royale des Sciences was founded in Paris. Both organizations provided support and impetus to the scientific revolution. The invention of the telescope enabled more precise geodetic surveying. In 1671 a French astronomer, Jean Picard
(1620–1682), completed an accurate survey by triangulation of the length of a degree of meridian arc. From his results the Earth’s radius was calculated to be 6372 km, remarkably close to the modern value of 6371 km.

2.1.2 Earth’s shape

In 1672 another French astronomer, Jean Richer was sent by Louis XIV to make astronomical observations on the equatorial island of Cayenne. He found that an accurate pendulum clock, which had been adjusted in Paris precisely to beat seconds, was losing about two and a half minutes per day, i.e., its period was now too long. The error was much too large to be explained by inaccuracy of the precise instrument. The observation aroused much interest and speculation, but was only explained some 15 years later by Sir Isaac Newton in terms of his laws of universal gravitation and motion.

Newton argued that the shape of the rotating Earth should be that of an *oblate ellipsoid*; compared to a sphere, it should be somewhat flattened at the poles and should bulge outward around the equator. This inference was made on logical grounds. Assume that the Earth does not rotate and that holes could be drilled to its center along the rotation axis and along an equatorial radius (Fig. 2.2). If these holes are filled with water, the hydrostatic pressure at the center of the Earth sustains equal water columns along each radius. However, the rotation of the Earth causes a centrifugal force at the equator but has no effect on the axis of rotation. At the equator the outward centrifugal force of the rotation opposes the inward gravitational attraction and pulls the water column upward. At the same time it reduces the hydrostatic pressure produced by the water column at the Earth’s center. The reduced central pressure is unable to support the height of the water column along the polar radius, which subsides. If the Earth were a hydrostatic sphere, the form of the rotating Earth should be an oblate ellipsoid of revolution. Newton assumed the Earth’s density to be constant and calculated that the flattening should be about 1:230 (roughly 0.5%). This is somewhat larger than the actual flattening of the Earth, which is about 1:298 (roughly 0.3%).

The increase in period of Richer’s pendulum could now be explained. Cayenne was close to the equator, where the larger radius placed the observer further from the center of gravitational attraction, and the increased distance from the rotational axis resulted in a stronger opposing centrifugal force. These two effects resulted in a lower value of gravity in Cayenne than in Paris, where the clock had been calibrated.

There was no direct proof of Newton’s interpretation. A corollary of his interpretation was that the degree of meridian arc should subtend a longer distance in polar regions than near the equator (Fig. 2.3). Early in the eighteenth century French geodesists extended the standard meridian from border to border of the country and found a puzzling result. In contrast to the prediction of Newton, the degree of meridian arc *decreased* northward. The French interpretation was that the Earth’s shape was a *prolate ellipsoid*, elongated at the poles and narrowed at the equator, like the shape of a rugby football. A major scientific controversy arose between the “flatteners” and the “elongators.”
To determine whether the Earth’s shape was oblate or prolate, the Académie Royale des Sciences sponsored two scientific expeditions. In 1736–1737 a team of scientists measured the length of a degree of meridian arc in Lapland, near the Arctic Circle. They found a length appreciably longer than the meridian degree measured by Picard near Paris. From 1735 to 1743 a second party of scientists measured the length of more than 3 degrees of meridian arc in Peru, near the equator. Their results showed that the equatorial degree of latitude was shorter than the meridian degree in Paris. Both parties confirmed convincingly the prediction of Newton that the Earth’s shape is that of an oblate ellipsoid.

The ellipsoidal shape of the Earth resulting from its rotation has important consequences, not only for the variation with latitude of gravity on the Earth’s surface, but also for the Earth’s rate of rotation and the orientation of its rotational axis. These are modified by torques that arise from the gravitational attractions of the Sun, Moon and planets on the ellipsoidal shape.

## 2.2 GRAVITATION

### 2.2.1 The law of universal gravitation

Sir Isaac Newton (1642–1727) was born in the same year in which Galileo died. Unlike Galileo, who relished debate, Newton was a retiring person and avoided confrontation. His modesty is apparent in a letter written in 1675 to his colleague Robert Hooke, famous for his experiments on elasticity. In this letter Newton made the famous disclaimer “if I have seen further (than you and Descartes) it is by standing upon the shoulders of Giants.” In modern terms Newton would be regarded as a theoretical physicist. He had an outstanding ability to synthesize experimental results and incorporate them into his own theories. Faced with the need for a more powerful technique of mathematical analysis than existed at the time, he invented differential and integral calculus, for which he is credited equally with Gottfried Wilhelm von Leibnitz (1646–1716) who discovered the same method independently. Newton was able to resolve many issues by formulating logical thought experiments; an example is his prediction that the shape of the Earth is an oblate ellipsoid. He was one of the most outstanding synthesizers of observations in scientific history, which is implicit in his letter to Hooke. His three-volume book *Philosophiae Naturalis Principia Mathematica*, published in 1687, ranks as the greatest of all scientific texts. The first volume of the *Principia* contains Newton’s famous *Laws of Motion*, the third volume handles the *Law of Universal Gravitation*.

The first two laws of motion are generalizations from Galileo’s results. As a corollary Newton applied his laws of motion to demonstrate that forces must be added as vectors and showed how to do this geometrically with a parallelogram. The second law of motion states that the rate of change of momentum of a mass is proportional to the force acting upon it and takes place in the direction of the force. For the case of constant mass, this law serves as the definition of force ($F$) in terms of the acceleration ($a$) given to a mass ($m$):

$$F = ma$$ \hspace{1cm} (2.1)

The unit of force in the SI system of units is the newton (N). It is defined as the force that gives a mass of one kilogram (1 kg) an acceleration of 1 m s$^{-2}$.

His celebrated observation of a falling apple may be a legend, but Newton’s genius lay in recognizing that the type of gravitational field that caused the apple to fall was the same type that served to hold the Moon in its orbit around the Earth, the planets in their orbits around the Sun, and that acted between minute particles characterized only by their masses. Newton used Kepler’s empirical third law (see Section 1.1.2 and Eq. (1.2)) to deduce that the force of attraction between a planet and the Sun varied with the “quantities of solid matter that they contain” (i.e., their masses) and with the inverse square of the distance between them. Applying this law to two particles or point masses $m$ and $M$ separated by a distance $r$ (Fig. 2.4a), we get for the gravitational attraction $F$ exerted by $M$ on $m$

$$F = -\frac{GMm}{r^2}$$ \hspace{1cm} (2.2)

In this equation $\hat{i}$ is a unit vector in the direction of increase in coordinate $r$, which is directed away from the center of reference at the mass $M$. The negative sign in the equation indicates that the force $F$ acts in the opposite direction, toward the attracting mass $M$. The constant $G$, which converts the physical law to an equation, is the constant of universal gravitation.

There was no way to determine the gravitational constant experimentally during Newton’s lifetime. The method to be followed was evident, namely to determine the force between two masses in a laboratory experiment. However, seventeenth century technology was not yet up to this task. Experimental determination of $G$ was extremely difficult, and was first achieved more than a century after the publication of *Principia* by Lord Charles Cavendish (1731–1810). From a set of painstaking measurements of the force of attraction between two spheres of lead Cavendish in 1798 determined the value of $G$ to be $6.754 \times 10^{-11}$ m$^3$ kg$^{-1}$ s$^{-2}$. A modern value (Mohr and Taylor, 2005) is $6.674 \times 10^{-11}$ m$^3$ kg$^{-1}$ s$^{-2}$. It has not yet been possible to determine $G$ more precisely, due to experimental difficulty. Although other physical constants are now known with a relative standard uncertainty of much less than $1 \times 10^{-6}$, the gravitational constant is known to only $150 \times 10^{-6}$.

### 2.2.1.1 Potential energy and work

The law of conservation of energy means that the total energy of a closed system is constant. Two forms of
The work done by the x-component of the force when it is displaced along the x-axis is $F_x dx$, and there are similar expressions for the displacements along the other axes. The change in potential energy $dE_p$ is now given by

$$dE_p = -dW = -(F_x dx + F_y dy + F_z dz)$$

(2.4)

The expression in brackets is called the scalar product of the vectors $F$ and $dr$. It is equal to $F \cdot dr \cos \theta$, where $\theta$ is the angle between the vectors.

### 2.2.2 Gravitational acceleration

In physics the field of a force is often more important than the absolute magnitude of the force. The field is defined as the force exerted on a material unit. For example, the electrical field of a charged body at a certain position is the force it exerts on a unit of electrical charge at that location. The gravitational field in the vicinity of an attracting mass is the force it exerts on a unit mass. Equation (2.1) shows that this is equivalent to the acceleration vector.

In geophysical applications we are concerned with accelerations rather than forces. By comparing Eq. (2.1) and Eq. (2.2) we get the gravitational acceleration $a_G$ of the mass $m$ due to the attraction of the mass $M$:

$$a_G = -GM/r^2$$

(2.5)

The SI unit of acceleration is the m s$^{-2}$; this unit is unpractical for use in geophysics. In the now superseded c.g.s. system the unit of acceleration was the cm s$^{-2}$, which is called a g (gal) in recognition of the contributions of Galileo. The small changes in the acceleration of gravity caused by geological structures are measured in thousandths of this unit, i.e., in milligal (mgal). Until recently, gravity anomalies due to geological structures were surveyed with field instruments accurate to about one-tenth of a milligal, which was called a gravity unit. Modern instruments are capable of measuring gravity differences to a millionth of a gal, or microgal (µgal), which is becoming the practical unit of gravity investigations.

The value of gravity at the Earth’s surface is about 9.8 m s$^{-2}$, and so the sensitivity of modern measurements of gravity is about 1 part in 10$^6$.

#### 2.2.2.1 Gravitational potential

The gravitational potential is the potential energy of a unit mass in a field of gravitational attraction. Let the potential be denoted by the symbol $U_G$. The potential energy $E_G$ of a mass $m$ in a gravitational field is thus equal to $(m U_G)$. Thus, a change in potential energy $(dE_G)$ is equal to $(m dU_G)$. Equation (2.3) becomes, using Eq. (2.1),

$$m dU_G = F dr = -ma_G dr$$

(2.6)

Rearranging this equation we get the gravitational acceleration.
2.2 GRAVITATION

\[ a_G = - \frac{dU_G}{dr^2} \]  (2.7)

In general, the acceleration is a three-dimensional vector. If we are using Cartesian coordinates \((x, y, z)\), the acceleration will have components \((a_x, a_y, a_z)\). These may be computed by calculating separately the derivatives of the potential with respect to \(x, y \) and \(z\):

\[ a_x = -\frac{\partial U_G}{\partial x}, \quad a_y = -\frac{\partial U_G}{\partial y}, \quad a_z = -\frac{\partial U_G}{\partial z} \]  (2.8)

Equating Eqs. (2.3) and (2.7) gives the gravitational potential of a point mass \(M\):

\[ \frac{dU_G}{dr} = G \frac{M}{r} \]  (2.9)

the solution of which is

\[ U_G = -G \frac{M}{r} \]  (2.10)

2.2.2.2 Acceleration and potential of a distribution of mass

Until now, we have considered only the gravitational acceleration and potential of point masses. A solid body may be considered to be composed of numerous small particles, each of which exerts a gravitational attraction at an external point \(P\) (Fig. 2.5a). To calculate the gravitational acceleration of the object at the point \(P\) we must form a vector sum of the contributions of the individual discrete particles. Each contribution has a different direction. Assuming \(m_i\) to be the mass of the particle at distance \(r_i\) from \(P\), this gives an expression like

\[ a_G = -G \frac{m_1}{r_1^2} - G \frac{m_2}{r_2^2} - G \frac{m_3}{r_3^2} - \cdots \]  (2.11)

Depending on the shape of the solid, this vector sum can be quite complicated.

An alternative solution to the problem is found by first calculating the gravitational potential, and then differentiating it as in Eq. (2.5) to get the acceleration. The expression for the potential at \(P\) is

\[ U_G = -G \frac{m_1}{r_1^2} - G \frac{m_2}{r_2^2} - G \frac{m_3}{r_3^2} - \cdots \]  (2.12)

This is a scalar sum, which is usually more simple to calculate than a vector sum.

More commonly, the object is not represented as an assemblage of discrete particles but by a continuous mass distribution. However, we can subdivide the volume into discrete elements; if the density of the matter in each volume is known, the mass of the small element can be calculated and its contribution to the potential at the external point \(P\) can be determined. By integrating over the volume of the body its gravitational potential at \(P\) can be calculated. At a point in the body with coordinates \((x, y, z)\) let the density be \(\rho(x, y, z)\) and let its distance from \(P\) be \(r(x, y, z)\) as in Fig. 2.5b. The gravitational potential of the body at \(P\) is

\[ U_G = -G \int \int \int p(x, y, z) \frac{dxdydz}{r(x, y, z)} \]  (2.13)

The integration readily gives the gravitational potential and acceleration at points inside and outside a hollow or homogeneous solid sphere. The values outside a sphere at distance \(r\) from its center are the same as if the entire mass \(E\) of the sphere were concentrated at its center (Fig. 2.4b):

\[ U_G = -GE \]  (2.14)

\[ a_G = -GE \frac{r}{r^2} \]  (2.15)

2.2.2.3 Mass and mean density of the earth

Equations (2.14) and (2.15) are valid everywhere outside a sphere, including on its surface where the distance from the center of mass is equal to the mean radius \(R\) (Fig. 2.4c). If we regard the Earth to a first approximation as a sphere with mass \(E\) and radius \(R\), we can estimate the Earth’s mass by rewriting Eq. (2.15) as a scalar equation in the form

\[ E = \frac{R^2 a_G}{G} \]  (2.16)

The gravitational acceleration at the surface of the Earth is only slightly different from mean gravity, about 9.81 m s\(^{-2}\), the Earth’s radius is 6371 km, and the gravitational constant is \(6.674 \times 10^{-11}\) m\(^3\) kg\(^{-1}\) s\(^{-2}\). The mass of the Earth is found to be \(5.974 \times 10^{24}\) kg. This large number is not so meaningful as the mean density of the Earth, which may be calculated by dividing the Earth’s mass by its volume \((\frac{4}{3} \pi R^3)\). A mean density of 5515 kg m\(^{-3}\) is obtained,
which is about double the density of crustal rocks. This indicates that the Earth's interior is not homogeneous, and implies that density must increase with depth in the Earth.

2.2.3 The equipotential surface

An equipotential surface is one on which the potential is constant. For a sphere of given mass the gravitational potential (Eq. (2.15)) varies only with the distance \( r \) from its center. A certain value of the potential, say \( U_1 \), is realized at a constant radial distance \( r_1 \). Thus, the equipotential surface on which the potential has the value \( U_1 \) is a sphere with radius \( r_1 \); a different equipotential surface \( U_2 \) is the sphere with radius \( r_2 \). The equipotential surfaces of the original spherical mass form a set of concentric spheres (Fig. 2.6a), one of which (e.g., \( U_o \)) coincides with the surface of the spherical mass. This particular equipotential surface describes the figure of the spherical mass.

By definition, no change in potential takes place (and no work is done) in moving from one point to another on an equipotential surface. The work done by a force \( F \) in a displacement \( dr \) is \( Fdr \cos \theta \) which is zero when \( \cos \theta \) is zero, that is, when the angle \( \theta \) between the displacement and the force is 90°. If no work is done in a motion along a gravitational equipotential surface, the force and acceleration of the gravitational field must act perpendicular to the surface. This normal to the equipotential surface defines the vertical, or plumb-line, direction (Fig. 2.6b). The plane tangential to the equipotential surface at a point defines the horizontal at that point.

2.3 THE EARTH’S ROTATION

2.3.1 Introduction

The rotation of the Earth is a vector, i.e., a quantity characterized by both magnitude and direction. The Earth behaves as an elastic body and deforms in response to the forces generated by its rotation, becoming slightly flattened at the poles with a compensating bulge at the equator. The gravitational attractions of the Sun, Moon and planets on the spinning, flattened Earth cause changes in its rate of rotation, in the orientation of the rotation axis, and in the shape of the Earth’s orbit around the Sun. Even without extra-terrestrial influences the Earth reacts to tiny displacements of the rotation axis from its average position by acquiring a small, unsteady wobble. These perturbations reflect a balance between gravitation and the forces that originate in the Earth’s rotational dynamics.

2.3.2 Centripetal and centrifugal acceleration

Newton’s first law of motion states that every object continues in its state of rest or of uniform motion in a straight line unless compelled to change that state by forces acting on it. The continuation of a state of motion is by virtue of the inertia of the body. A framework in which this law is valid is called an inertial system. For example, when we are travelling in a car at constant speed, we feel no disturbing forces; reference axes fixed to the moving vehicle form an inertial frame. If traffic conditions compel the driver to apply the brakes, we experience decelerating forces; if the car goes around a corner, even at constant speed, we sense sideways forces toward the outside of the corner. In these situations the moving car is being forced to change its state of uniform rectilinear motion and reference axes fixed to the car form a non-inertial system.

Motion in a circle implies that a force is active that continually changes the state of rectilinear motion. Newton recognized that the force was directed inwards, towards the center of the circle, and named it the centripetal (meaning “center-seeking”) force. He cited the example of a stone being whirled about in a sling. The inward centripetal force exerted on the stone by the sling holds it in a circular path. If the sling is released, the restraint of the centripetal force is removed and the inertia of the stone causes it to continue its motion at the point of release. No longer under the influence of the restraining force, the stone flies off in a straight line. Arguing that the curved path of a projectile near the surface of the Earth was due to the effect of gravity, which caused it constantly to fall toward the Earth, Newton postulated that, if the speed of the projectile were exactly right, it might never quite reach the Earth’s surface. If the projectile fell toward the center of the Earth at the same rate as the curved surface of the Earth fell away from it, the projectile would go into orbit around the Earth. Newton suggested that the Moon was held in
orbit around the Earth by just such a centripetal force, which originated in the gravitational attraction of the Earth. Likewise, he visualized that a centripetal force due to gravitational attraction restrained the planets in their circular orbits about the Sun.

The passenger in a car going round a corner experiences a tendency to be flung outwards. He is restrained in position by the frame of the vehicle, which supplies the necessary centripetal acceleration to enable the passenger to go round the curve in the car. The inertia of the passenger’s body causes it to continue in a straight line and pushes him outwards against the side of the vehicle. This outward force is called the centrifugal force. It arises because the car does not represent an inertial reference frame. An observer outside the car in a fixed (inertial) coordinate system would note that the car and passenger are constantly changing direction as they round the corner. The centrifugal force feels real enough to the passenger in the car, but it is called a pseudo-force, or inertial force. In contrast to the centripetal force, which arises from the gravitational attraction, the centrifugal force does not have a physical origin, but exists only because it is being observed in a non-inertial reference frame.

### 2.3.2.1 Centripetal acceleration

The mathematical form of the centripetal acceleration for circular motion with constant angular velocity \( \omega \) about a point can be derived as follows. Define orthogonal Cartesian axes \( x \) and \( y \) relative to the center of the circle as in Fig. 2.7a. The linear velocity \( \omega \) at any point where the radius vector makes an angle \( \theta = (vt) \) with the \( x \)-axis has components

\[
\begin{align*}
    v_x &= -r \sin(\omega t) = -r \omega \sin(\omega t) \\
    v_y &= r \cos(\omega t) = r \omega \cos(\omega t) 
\end{align*}
\]

The \( x \) - and \( y \) -components of the acceleration are obtained by differentiating the velocity components with respect to time. This gives

\[
\begin{align*}
    a_x &= -r \omega \cos(\omega t) = -r \omega^2 \cos(\omega t) \\
    a_y &= -r \omega \sin(\omega t) = -r \omega^2 \sin(\omega t) 
\end{align*}
\]

These are the components of the centripetal acceleration, which is directed radially inwards and has the magnitude \( \omega^2 r \) (Fig. 2.7b).

### 2.3.2.2 Centrifugal acceleration and potential

In handling the variation of gravity on the Earth’s surface we must operate in a non-inertial reference frame attached to the rotating Earth. Viewed from a fixed, external inertial frame, a stationary mass moves in a circle about the Earth’s rotation axis with the same rotational speed as the Earth.

However, within a rotating reference frame attached to the Earth, the mass is stationary. It experiences a centrifugal acceleration \( (a_c) \) that is exactly equal and opposite to the centripetal acceleration, and which can be written in the alternative forms

\[
\begin{align*}
    a_c &= \omega^2 r \\
    a_c &= \frac{v^2}{r} 
\end{align*}
\]

The centrifugal acceleration is not a centrally oriented acceleration like gravitation, but instead is defined relative to an axis of rotation. Nevertheless, potential energy is associated with the rotation and it is possible to define a centrifugal potential. Consider a point rotating with the Earth at a distance \( r \) from its center (Fig. 2.8). The angle \( \theta \) between the radius to the point and the axis of rotation is called the colatitude; it is the angular complement of the latitude \( \lambda \). The distance of the point from the rotational axis is \( x \) \((= r \sin \theta)\), and the centrifugal acceleration is \( \omega^2 x \) outwards in the direction of increasing \( x \). The centrifugal potential \( U_c \) is defined such that

\[
a_c = -\frac{\partial U}{\partial x} \hat{k} = (\omega^2 x) \hat{k}
\]

where \( \hat{k} \) is the outward unit vector. On integrating, we obtain

\[
U_c = -\frac{1}{2} \omega^2 x^2 = -\frac{1}{2} \omega^2 r^2 \cos^2 \lambda = -\frac{1}{2} \omega^2 r^2 \sin^2 \theta
\]
2.3.2.4 Verification of the inverse square law of gravitation

Let the corresponding angular rate of rotation be \( \omega \). Setting these moments equal we get

\[
\frac{r_p^3}{T_p^2} = \frac{GS}{4\pi^2} = \text{constant} \tag{2.23}
\]

2.3.2.4 Verification of the inverse square law of gravitation

Newton realized that the centripetal acceleration of the Moon in its orbit was supplied by the gravitational attraction of the Earth, and tried to use this knowledge to confirm the inverse square dependence on distance in his law of gravitation. The sidereal period \( T_L \) of the Moon about the Earth, a sidereal month, is equal to 27.3 days. Let the corresponding angular rate of rotation be \( \omega_L \). We can equate the gravitational acceleration of the Earth at the Moon with the centripetal acceleration due to \( \omega_L \):

\[
G \frac{E}{r_L^2} = \omega_L^2 r_L \tag{2.24}
\]

This equation can be rearranged as follows

\[
\left( \frac{G E}{R^2} \right)^2 = \omega_L^2 \frac{r_L^3}{R^3} \tag{2.25}
\]

Comparison with Eq. (2.15) shows that the first quantity in parentheses is the mean gravitational acceleration on the Earth’s surface, \( a_G \). Therefore, we can write

\[
a_G = G \frac{E}{R^2} = \omega_L^2 R \left( \frac{r_L}{R} \right)^3 \tag{2.26}
\]

In Newton’s time little was known about the physical dimensions of our planet. The distance of the Moon was known to be approximately 60 times the radius of the Earth (see Section 1.1.3.2) and its sidereal period was known to be 27.3 days. At first Newton used the accepted value 5500 km for the Earth’s radius. This gave a value of only 8.4 m s\(^{-2}\) for gravity, well below the known value of 9.8 m s\(^{-2}\). However, in 1671 Picard determined the Earth’s radius to be 6372 km. With this value, the inverse square character of Newton’s law of gravitation was confirmed.

2.3.3 The tides

The gravitational forces of Sun and Moon deform the Earth’s shape, causing tides in the oceans, atmosphere and solid body of the Earth. The most visible tidal effects are the displacements of the ocean surface, which is a hydrostatic equipotential surface. The Earth does not react rigidly to the tidal forces. The solid body of the Earth deforms in a like manner to the free surface, giving rise to so-called bodily Earth-tides. These can be observed with specially designed instruments, which operate on a similar principle to the long-period seismometer.

The height of the marine equilibrium tide amounts to only half a meter or so over the free ocean. In coastal areas the tidal height is significantly increased by the shallowing of the continental shelf and the confining shapes of bays and harbors. Accordingly, the height and variation of the tide at any place is influenced strongly by complex local factors. Subsequent subsections deal with the tidal deformations of the Earth’s hydrostatic figure.

2.3.3.1 Lunar tidal periodicity

The Earth and Moon are coupled together by gravitational attraction. Their common motion is like that of a pair of ballroom dancers. Each partner moves around the center of mass (or barycenter) of the pair. For the Earth–Moon pair the location of the center of mass is easily found. Let \( E \) be the mass of the Earth, and \( m \) that of the Moon; let the separation of the centers of the Earth and Moon be \( r_L \) and let the distance of their common center of mass be \( d \) from the center of the Earth. The moment of the Earth about the center of mass is \( Ed \) and the moment of the Moon is \( m(r_L - d) \). Setting these moments equal we get

\[
d = \frac{m}{E + m} r_L \tag{2.27}
\]
The mass of the Moon is 0.0123 that of the Earth and the distance between the centers is 384,400 km. These figures give $d/H_{11005} = 4600$ km, i.e., the center of revolution of the Earth–Moon pair lies within the Earth.

It follows that the paths of the Earth and the Moon around the Sun are more complicated than at first appears. The elliptical orbit is traced out by the barycenter of the pair (Fig. 2.9). The Earth and Moon follow wobbly paths, which, while always concave towards the Sun, bring each body at different times of the month alternately inside and outside the elliptical orbit.

To understand the common revolution of the Earth–Moon pair we have to exclude the rotation of the Earth about its axis. The “revolution without rotation” is illustrated in Fig. 2.10. The Earth–Moon pair revolves about S, the center of mass. Let the starting positions be as shown in Fig. 2.10a. Approximately one week later the Moon has advanced in its path by one-quarter of a revolution and the center of the Earth has moved so as to keep the center of mass fixed (Fig. 2.10b). The relationship is maintained in the following weeks (Fig. 2.10c, d) so that during one month the center of the Earth describes a circle about S. Now consider the motion of point number 2 on the left-hand side of the Earth in Fig. 2.10. If the Earth revolves as a rigid body and the rotation about its own axis is omitted, after one week point 2 will have moved to a new position but will still be the furthest point on the left. Subsequently, during one month point 2 will describe a small circle with the same radius as the circle described by the Earth’s center. Similarly points 1, 3 and 4 will also describe circles of exactly the same size. A simple illustration of this point can be made by chalking the tip of each finger on one hand with a different color, then moving your hand in a circular motion while touching a blackboard; your fingers will draw a set of identical circles.

The “revolution without rotation” causes each point in the body of the Earth to describe a circular path with identical radius. The centrifugal acceleration of this motion has therefore the same magnitude at all points in the Earth and, as can be seen by inspection of Fig. 2.10(a–d), it is directed away from the Moon parallel to the Earth–Moon line of centers. At C, the center of the Earth (Fig. 2.11a), this centrifugal acceleration exactly balances the gravitational attraction of the Moon. Its magnitude is given by

$$a_L = \frac{G m}{r_L^2}$$  \hspace{1cm} (2.28)

At B, on the side of the Earth nearest to the Moon, the gravitational acceleration of the Moon is larger than at the center of the Earth and exceeds the centrifugal acceleration $a_L$. There is a residual acceleration toward the Moon, which raises a tide on this side of the Earth. The magnitude of the tidal acceleration at B is
Gravity, the figure of the Earth and geodynamics

At points D and D’ the direction of the gravitational acceleration due to the Moon is not exactly parallel to the line of centers of the Earth–Moon pair. The residual tidal acceleration is almost along the direction toward the center of the Earth. Its effect is to lower the free surface in this direction.

The free hydrostatic surface of the Earth is an equipotential surface (Section 2.2.3), which in the absence of the Earth’s rotation and tidal effects would be a sphere. The lunar tidal accelerations perturb the equipotential surface, raising it at A and B while lowering it at D and D’, as in Fig. 2.11a. The tidal deformation of the Earth produced by the Moon thus has an almost prolate ellipsoidal shape, like a rugby football, along the Earth–Moon line of centers. The daily tides are caused by superposing the Earth’s rotation on this deformation. In the course of one day a point rotates past the points A, D, B and D’ and an observer experiences two full tidal cycles, called the semi-diurnal tides. The extreme tides are not equal at every latitude, because of the varying angle between the Earth’s rotational axis and the Moon’s orbit (Fig. 2.11b). At the equator E the semi-diurnal tides are equal; at an intermediate latitude F one tide is higher than the other; and at latitude G and higher there is only one (diurnal) tide per day. The difference in height between two successive high or low tides is called the diurnal inequality.

In the same way that the Moon deforms the Earth, so the Earth causes a tidal deformation of the Moon. In fact, the tidal relationship between any planet and one of its moons, or between the Sun and a planet or comet, can be treated analogously to the Earth–Moon pair. A tidal acceleration similar to Eq. (2.31) deforms the smaller body; its self-gravitation acts to counteract the deformation. However, if a moon or comet comes too close to the planet, the tidal forces deforming it may overwhelm the gravitational forces holding it together, so that the moon or comet is torn apart. The separation at which this occurs is called the Roche limit (Box 2.1). The material of a disintegrated moon or comet enters orbit around the planet, forming a system of concentric rings, as around the great planets (Section 1.1.3.3).

**2.3.3.2 Tidal effect of the Sun**

The Sun also has an influence on the tides. The theory of the solar tides can be followed in identical manner to the lunar tides by again applying the principle of “revolution without rotation.” The Sun’s mass is 333,000 times greater than that of the Earth, so the common center of mass is close to the center of the Sun at a radial distance of about 450 km from its center. The period of the revolution is one year. As for the lunar tide, the imbalance between gravitational acceleration of the Sun and centrifugal acceleration due to the common revolution leads to a prolate ellipsoidal tidal deformation. The solar effect is smaller than that of the Moon. Although the mass of the Sun is vastly
greater than that of the Moon, its distance from the Earth is also much greater and, because gravitational acceleration varies inversely with the square of distance, the maximum tidal effect of the Sun is only about 45% that of the Moon.

2.3.3.3 Spring and neap tides

The superposition of the lunar and solar tides causes a modulation of the tidal amplitude. The ecliptic plane is defined by the Earth’s orbit around the Sun. The Moon’s orbit around the Earth is not exactly in the ecliptic but is...
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Fig. 2.12 The orientations of the solar and lunar tidal deformations of the Earth at different lunar phases.

inclined at a very small angle of about 5° to it. For discussion of the combination of lunar and solar tides we can assume the orbits to be coplanar. The Moon and Sun each produce a prolate tidal deformation of the Earth, but the relative orientations of these ellipsoids vary during one month (Fig. 2.12). At conjunction the (new) Moon is on the same side of the Earth as the Sun, and the ellipsoidal deformations augment each other. The same is the case half a month later at opposition, when the (full) Moon is on the opposite side of the Earth from the Sun. The unusually high tides at opposition and conjunction are called spring tides. In contrast, at the times of quadrature the waxing or waning half Moon causes a prolate ellipsoidal deformation out of phase with the solar deformation. The maximum lunar tide coincides with the minimum solar tide, and the effects partially cancel each other. The unusually low tides at quadrature are called neap tides. The superposition of the lunar and solar tides causes modulation of the tidal amplitude during a month (Fig. 2.13).

2.3.3.4 Effect of the tides on gravity measurements

The tides have an effect on gravity measurements made on the Earth. The combined effects of Sun and Moon cause an acceleration at the Earth’s surface of approximately 0.3 mgal, of which about two-thirds are due to the Moon and one-third to the Sun. The sensitive modern instruments used for gravity exploration can readily detect gravity differences of 0.01 mgal. It is necessary to compen-

sate gravity measurements for the tidal effects, which vary with location, date and time of day. Fortunately, tidal theory is so well established that the gravity effect can be calculated and tabulated for any place and time before beginning a survey.

2.3.3.5 Bodily Earth-tides

A simple way to measure the height of the marine tide might be to fix a stake to the sea-bottom at a suitably sheltered location and to record continuously the measured water level (assuming that confusion introduced by wave motion can be eliminated or taken into account). The observed amplitude of the marine tide, defined by the displacement of the free water surface, is found to be about 70% of the theoretical value. The difference is explained by the elasticity of the Earth. The tidal deformation corresponds to a redistribution of mass, which modifies the gravitational potential of the Earth and augments the elevation of the free surface. This is partially counteracted by a bodily tide in the solid Earth, which deforms elastically in response to the attraction of the Sun and Moon. The free water surface is raised by the tidal attraction, but the sea-bottom in which the measuring rod is implanted is also raised. The measured tide is the difference between the marine tide and the bodily Earth-tide.

In practice, the displacement of the equipotential surface is measured with a horizontal pendulum, which reacts to the tilt of the surface. The bodily Earth-tides also affect gravity measurements and can be observed with sensitive gravimeters. The effects of the bodily Earth-tides are incorporated into the predicted tidal corrections to gravity measurements.

2.3.4 Changes in Earth’s rotation

The Earth’s rotational vector is affected by the gravitational attractions of the Sun, Moon and the planets. The rate of rotation and the orientation of the rotational axis change with time. The orbital motion around the Sun is also affected. The orbit rotates about the pole to the plane of the ecliptic and its ellipticity changes over long periods of time.

2.3.4.1 Effect of lunar tidal friction on the length of the day

If the Earth reacted perfectly elastically to the lunar tidal forces, the prolate tidal bulge would be aligned along the line of centers of the Earth–Moon pair (Fig. 2.14a). However, the motion of the seas is not instantaneous and the tidal response of the solid part of the Earth is partly anelastic. These features cause a slight delay in the time when high tide is reached, amounting to about 12 minutes. In this short interval the Earth’s rotation carries the line of the maximum tides past the line of centers by a small angle of approximately 2.9° (Fig. 2.14b). A point on the rotating Earth passes under the line of maximum
tides 12 minutes after it passes under the Moon. The small phase difference is called the tidal lag.

Because of the tidal lag the gravitational attraction of the Moon on the tidal bulges on the far side and near side of the Earth ($F_1$ and $F_2$, respectively) are not collinear (Fig. 2.14b). $F_2$ is stronger than $F_1$ so a torque is produced in the opposite sense to the Earth's rotation (Fig. 2.14c). The tidal torque acts as a brake on the Earth's rate of rotation, which is gradually slowing down.

The tidal deceleration of the Earth is manifested in a gradual increase in the length of the day. The effect is very small. Tidal theory predicts an increase in the length of the day of only 2.4 milliseconds per century. Observations of the phenomenon are based on ancient historical records of lunar and solar eclipses and on telescopically observed occultations of stars by the Moon. The current rate of rotation of the Earth can be measured with very accurate atomic clocks. Telescopic observations of the daily times of passage of stars past the local zenith are recorded with a camera controlled by an atomic clock. These observations give precise measures of the mean value and fluctuations of the length of the day.

The occurrence of a lunar or solar eclipse was a momentous event for ancient peoples, and was duly recorded in scientific and non-scientific chronicles. Untimed observations are found in non-astronomical works. They record, with variable reliability, the degree of totality and the time and place of observation. The unaided human eye is able to decide quite precisely just when an eclipse becomes total. Timed observations of both lunar and solar eclipses made by Arab astronomers around 800–1000 AD and Babylonian astronomers a thousand years earlier give two important groups of data (Fig. 2.15). By comparing the observed times of alignment
of Sun, Moon and Earth with times predicted from the theory of celestial mechanics, the differences due to change in length of the day may be computed. A straight line with slope equal to the rate of increase of the length of the day inferred from tidal theory, 2.4 ms per century, connects the Babylonian and Arab data sets. Since the medieval observations of Arab astronomers the length of day inferred from tidal theory, 2.4 ms per century, connects the Babylonian and Arab data sets. Since the data are equally compatible with a smoothly changing polynomial. The slope is that non-tidal causes have opposed the deceleration of the Earth’s rotation since about 950 AD. It would be wrong to infer that some sudden event at that epoch caused an abrupt change, because the data are equally compatible with a smoothly changing polynomial. The observations confirm the importance of tidal braking, but they also indicate that tidal friction is not the only mechanism affecting the Earth’s rotation.

The short-term fluctuations in rotation rate are due to exchanges of angular momentum with the atmosphere. On a longer timescale of decades, the changes in length of the day may be related to changes in the angular momentum of the core. The fluid in the outer core has a speed of the order of 0.1 mm s\(^{-1}\) relative to the overlying mantle. The mechanism for exchange of angular momentum between the fluid core and the rest of the Earth depends on the way the core and mantle are coupled. The coupling may be mechanical if topographic irregularities obstruct the flow of the core fluid along the core–mantle interface. The core fluid is a good electrical conductor so, if the lower mantle also has an appreciable electrical conductivity, it is possible that the core and mantle are coupled electromagnetically.

### 2.3.4.2 Increase of the Earth–Moon distance

Further consequences of lunar tidal friction can be seen by applying the law of conservation of momentum to the Earth–Moon pair. Let the Earth’s mass be \(E\), its rate of rotation be \(\omega\) and its moment of inertia about the rotation axis be \(C\); let the corresponding parameters for the Moon be \(m\), \(\Omega\), and \(C\), and let the Earth–Moon distance be \(r\). Further, let the distance of the common center of revolution be \(d\) from the center of the Earth, as given by Eq. (2.27). The angular momentum of the system is given by

\[
C \omega + EM_d^2 + mL(r - d)^2 + C_L = \text{constant} \tag{2.34}
\]

The fourth term is the angular momentum of the Moon about its own axis. Tidal deceleration due to the Earth’s attraction has slowed down the Moon’s rotation until it equals its rate of revolution about the Earth. Both \(\Omega\) and \(C_L\) are very small and the fourth term can be neglected. The second and third terms can be combined so that we get

\[
C \omega + \left(\frac{E}{E + M}\right)mL = \text{constant} \tag{2.35}
\]

The gravitational attraction of the Earth on the Moon is equal to the centripetal acceleration of the Moon about the common center of revolution, thus

\[
\frac{GM_e}{r_L^2} = \Omega_L^2 (r - d) = \Omega_L^2 r_L \left(\frac{E}{E + M}\right) \tag{2.36}
\]

from which

\[
\Omega_L r_L = \sqrt{G(E + m)}r_L \tag{2.37}
\]

Inserting this in Eq. (2.35) gives

\[
C \omega + \frac{Em}{\sqrt{(E + m)}} \sqrt{Gr_L} = \text{constant} \tag{2.38}
\]

The first term in this equation decreases, because tidal friction reduces \(\omega\). To conserve angular momentum the second term must increase. Thus, lunar tidal braking of the
Earth’s rotation causes an increase in the Earth–Moon distance, $r_L$. At present this distance is increasing at about 3.7 cm yr$^{-1}$. As a further consequence Eq. (2.37) shows that the Moon’s rate of revolution about the Earth ($\Omega_L$) and consequently also its synchronous rotation about its own axis – must decrease when $r_L$ increases. Thus, tidal friction slows down the rates of Earth rotation, lunar rotation, and lunar orbital revolution and increases the Earth–Moon distance.

Eventually a situation will evolve in which the Earth’s rotation has slowed until it is synchronous with the Moon’s own rotation and its orbital revolution about the Earth. All three rotations will then be synchronous and equivalent to about 48 present Earth days. This will happen when the Moon’s distance from Earth is about 88 times the Earth’s radius ($r_L = 88R$; it is presently equal to about $60R$). The Moon will then be stationary over the Earth, and Earth and Moon will constantly present the same face to each other. This configuration already exists between the planet Pluto and its satellite Charon.

2.3.4.3 The Chandler wobble

The Earth’s rotation gives it the shape of a spheroid, or ellipsoid of revolution. This figure is symmetric with respect to the mean axis of rotation, about which the moment of inertia is greatest; this is also called the axis of figure (see Section 2.4). However, at any moment the instantaneous rotational axis is displaced by a few meters from the axis of figure. The orientation of the total angular momentum vector remains nearly constant but the axis of figure changes location with time and appears to meander around the rotation axis (Fig. 2.16).

The theory of this motion was described by Leonhard Euler (1707–1783), a Swiss mathematician. He showed that the displaced rotational axis of a rigid spheroid would execute a circular motion about its mean position, now called the Euler nutation. Because it occurs in the absence of an external driving torque, it is also called the free nutation. It is due to differences in the way mass is distributed about the axis of rotational symmetry and an axis at right angles to it in the equatorial plane. The mass distributions are represented by the moments of inertia about these axes. If $C$ and $A$ are the moments of inertia about the rotational axis and an axis in the equatorial plane, respectively, Euler’s theory shows that the period of free nutation is $A/(C-A)$ days, or approximately 305 days.

Astronomers were unsuccessful in detecting a polar motion with this period. In 1891 an American geodesist and astronomer, S. C. Chandler, reported that the polar motion of the Earth’s axis contained two important components. An annual component with amplitude about 0.10 seconds of arc is due to the transfer of mass between atmosphere and hydrosphere accompanying the changing of the seasons. A slightly larger component with amplitude 0.15 seconds of arc has a period of 435 days. This polar motion is now called the Chandler wobble. It corresponds to the Euler nutation in an elastic Earth. The increase in period from 305 days to 435 days is a consequence of the elastic yielding of the Earth. The superposition of the annual and Chandler frequencies results in a beat effect, in which the amplitude of the
The forces are not collinear. A torque results, which acts about a line in the equatorial plane, normal to the Earth–Sun line and normal to the spin axis. The magnitude of the torque changes as the Earth orbits around the Sun. It is minimum (and zero) at the spring and autumn equinoxes and maximum at the summer and winter solstices.

The response of a rotating system to an applied torque is to acquire an additional component of angular momentum parallel to the torque. In our example this will be perpendicular to the angular momentum \( \mathbf{h} \) of the spinning Earth. The torque has a component \( \tau \) parallel to the line of equinoxes (Fig. 2.17b) and a component normal to this line in the equatorial plane. The torque \( \tau \) causes an increment \( \Delta h \) in angular momentum and shifts the angular momentum vector to a new position. If this exercise is repeated incrementally, the rotation axis moves around the surface of a cone whose axis is the pole to the ecliptic (Fig. 2.17a). The geographic pole \( P \) moves around a circle in the opposite sense from the Earth’s spin. This motion is called retrograde precession. It is not a steady motion, but pulsates in sympathy with the driving torque. A change in orientation of the rotation axis affects the location of the line of equinoxes and causes the timing of the equinoxes to change slowly. The rate of change is only 50.4 seconds of arc per year, but it has been recognized during centuries of observation. For example, the Earth’s rotation axis now points at Polaris in the constellation Ursa Minor, but in the time of the Egyptians around 3000 BC the pole star was Alpha Draconis, the brightest star in the constellation Draco. Hipparchus is credited with discovering the precession of the equinoxes in 120 BC by comparing his own observations with those of earlier astronomers.

The theory of the phenomenon is well understood. The Moon also exerts a torque on the spinning Earth and contributes to the precession of the rotation axis (and equinoxes). As in the theory of the tides, the small size of the Moon compared to the Sun is more than compensated by its nearness, so that the precessional contribution of the Moon is about double the effect of the Sun. The theory of precession shows that the period of 25,700 yr is proportional to the Earth’s dynamical ellipticity, \( H \) (see Eq. (2.45)). This ratio (equal to 1/305.457) is an important indicator of the internal distribution of mass in the Earth.

The component of the torque in the equatorial plane adds an additional motion to the axis, called nutation, because it causes the axis to nod up and down (Fig. 2.17a). The solar torque causes a semi-annual nutation, the lunar torque a semi-monthly one. In fact the motion of the axis exhibits many forced nutations, so-called because they respond to external torques. All are tiny perturbations on the precessional motion, the largest having an amplitude of only about 9 seconds of arc and a period of 18.6 yr. This nutation results from the fact that the plane of the lunar orbit is inclined at 5.145° to the plane of the ecliptic and (like the motion of artificial Earth
satellites) precess retrogradely. This causes the inclination of the lunar orbit to the equatorial plane to vary between about 18.4° and 28.6°, modulating the torque and forcing a nutation with a period of 18.6 yr.

It is important to note that the Euler nutation and Chandler wobble are polar motions about the rotation axis, but the precession and forced nutations are displacements of the rotation axis itself.

2.3.4.5 Milankovitch climatic cycles

Solar energy can be imagined as flowing equally from the Sun in all directions. At distance \( r \) it floods a sphere with surface area \( 4\pi r^2 \). The amount of solar energy falling per second on a square meter (the insolation) therefore decreases as the inverse square of the distance from the Sun. The gravitational attractions of the Moon, Sun, and the other planets – especially Jupiter – cause cyclical changes of the orientation of the rotation axis and variations in the shape and orientation of Earth’s orbit. These variations modify the insolation of the Earth and result in long-term periodic changes in Earth’s climate.

The angle between the rotational axis and the pole to the ecliptic is called the obliquity. It is the main factor determining the seasonal difference between summer and winter in each hemisphere. In the northern hemisphere, the insolation is maximum at the summer solstice (currently June 21) and minimum at the winter solstice (December 21–22). The exact dates change with the precession of the equinoxes, and also depend on the occurrence of leap years. The solstices do not coincide with extreme positions in Earth’s orbit. The Earth currently reaches aphelion, its furthest distance from the Sun, around July 4–6, shortly after the summer solstice, and passes perihelion around January 2–4. About 13,000 yr from now, as a result of precession, the summer solstice will occur when Earth is close to perihelion. In this way, precession causes long-term changes in climate with a period related to the precession.

The gravitational attraction of the other planets causes the obliquity to change cyclically with time. It is currently equal to 23° 26’ 21.4° but varies slowly between a minimum of 21° 55’ and a maximum of 24° 18’. When the obliquity increases, the seasonal differences in temperature become more pronounced, while the opposite effect ensues if obliquity decreases. Thus, the variation in obliquity causes a modulation in the seasonal contrast between summer and winter on a global scale. This effect is manifest as a cyclical change in climate with a period of about 41 kyr.

A further effect of planetary attraction is to cause the eccentricity of the Earth’s orbit, at present 0.017, to change cyclically (Fig. 2.18). At one extreme of the cycle, the orbit is almost circular, with an eccentricity of only 0.005. The closest distance from the Sun at perihelion is then 99% of the furthest distance at aphelion. At the other extreme, the orbit is more elongate, although with an eccentricity of 0.058 it is only slightly elliptical. The perihelion distance is then 89% of the aphelion distance.

These slight differences have climatic effects. When the orbit is almost circular, the difference in insolation between summer and winter is negligible. However, when the orbit is most elongate, the insolation in winter is only 78% of the summer insolation. The cyclical variation in eccentricity has a dominant period of 404 kyr and lesser periodicities of 95 kyr, 99 kyr, 124 kyr and 131 kyr that together give a roughly 100 kyr period. The eccentricity variations generate fluctuations in paleoclimatic records with periods around 100 kyr and 400 kyr.

Not only does planetary attraction cause the shape of the orbit to change, it also causes the perihelion–aphelion axis of the orbit to precess. The orbital ellipse is not truly closed, and the path of the Earth describes a rosette with a period that is also around 100 kyr (Fig. 2.18). The precession of perihelion interacts with the axial precession and modifies the observed period of the equinoxes. The 26 kyr axial precession is retrograde with a rate of 0.038 cycles/kyr; the 100 kyr orbital precession is prograde, which speeds up the effective precession rate to 0.048 cycles/kyr. This is equivalent to a retrograde precession with a period of about 21 kyr. A corresponding climatic fluctuation has been interpreted in many sedimentary deposits.

Climatic effects related to cyclical changes in the Earth’s rotational and orbital parameters were first studied between 1920 and 1938 by a Yugoslavian astronomer, Milutin Milanković (anglicized to Milankovitch). Periodicities of 21 kyr, 41 kyr, 100 kyr and 400 kyr – called the
Milankovitch climatic cycles – have been described in various sedimentary records ranging in age from Quaternary to Mesozoic. Caution must be used in interpreting the cyclicities in older records, as the characteristic Milankovitch periods are dependent on astronomical parameters that may have changed appreciably during the geological ages.

2.3.5 Coriolis and Eötvös accelerations

Every object on the Earth experiences the centrifugal acceleration due to the Earth’s rotation. Moving objects on the rotating Earth experience additional accelerations related to the velocity at which they are moving. At latitude $\lambda$ the distance $d$ of a point on the Earth’s surface from the rotational axis is equal to $R \cos \lambda$, and the rotational spin $\omega$ translates to an eastwards linear velocity $v$ equal to $\omega R \cos \lambda$. Consider an object (e.g., a vehicle or projectile) that is moving at velocity $v$ across the Earth’s surface. In general $v$ has a northward component $v_N$ and an eastward component $v_E$. Consider first the effects related to the eastward velocity, which is added to the linear velocity of the rotation. The centrifugal acceleration increases by an amount $\Delta a_c$, which can be obtained by differentiating $a_c$ in Eq. (2.19) with respect to $\omega$

$$\Delta a_c = 2\omega (R \cos \lambda) \Delta \omega = 2\omega v_E$$  \hspace{1cm} (2.39)

The extra centrifugal acceleration $\Delta a_c$ can be resolved into a vertical component and a horizontal component (Fig. 2.19a). The vertical component, equal to $2\omega v_E \cos \lambda$, acts upward, opposite to gravity. It is called the *Eötvös acceleration*. Its effect is to decrease the measured gravity by a small amount. If the moving object has a westward component of velocity the Eötvös acceleration increases the measured gravity. If gravity measurements are made on a moving platform (for example, on a research ship or in an airplane), the measured gravity must be corrected to allow for the Eötvös effect. The correction depends on the eastward velocity of the object.

The horizontal component of the extra centrifugal acceleration due to $v_N$ is equal to $2\omega v_E \sin \lambda$. In the northern hemisphere it acts to the south. If the object moves westward, the acceleration is northward. In each case it acts horizontally to the right of the direction of motion. In the southern hemisphere the sense of this acceleration is reversed; it acts to the left of the direction of motion. This acceleration is a component of the *Coriolis acceleration*, another component of which derives from the northward motion of the object.

Consider an object moving northward along a meridian of longitude (Fig. 2.19b, point 1). The linear velocity of a point on the Earth’s surface decreases poleward, because the distance from the axis of rotation ($d = R \cos \lambda$) decreases. The angular momentum of the moving object must be conserved, so the eastward velocity $v_E$ must increase. As the object moves to the north its eastward velocity is faster than the circles of latitude it crosses and its trajectory deviates to the right. If the motion is to the south (Fig. 2.19b, point 2), the inverse argument applies. The body crosses circles of latitude with faster eastward velocity than its own and, in order to maintain angular momentum, its trajectory must deviate to the west. In each case the deviation is to the right of the direction of motion. A similar argument applied to the southern hemisphere gives a Coriolis effect to the left of the direction of motion (Fig. 2.19b, points 3 and 4).

The magnitude of the Coriolis acceleration is easily evaluated quantitatively. The angular momentum $h$ of a mass $m$ at latitude $\lambda$ is equal to $m \omega R^2 \cos^2 \lambda$. Conservation of angular momentum gives

$$\frac{dh}{dt} = mR^2 \cos^2 \lambda \frac{d\omega}{dt} + ma_R (2 \cos \lambda \sin \lambda) \frac{d\lambda}{dt} = 0$$  \hspace{1cm} (2.40)

Rearranging and simplifying, we get
(R\cos\lambda)\frac{\partial \omega}{\partial t} = 2\omega \sin\lambda (\frac{\partial \lambda}{\partial t}) \tag{2.41}

The expression on the left of the equation is an acceleration, \(a_E\), equal to the rate of change of the eastward velocity. The expression in brackets on the right is the northward velocity component \(v_N\). We can write this component of the Coriolis acceleration as \(2\omega v_N \sin\lambda\). The north and east components of the Coriolis acceleration are therefore:

\[ a_N = 2\omega v_N \sin\lambda \]
\[ a_E = 2\omega v_N \sin\lambda \tag{2.42} \]

The Coriolis acceleration deflects the horizontal path of any object moving on the Earth's surface. It affects the directions of wind and ocean currents, eventually constraining them to form circulatory patterns about centers of high or low pressure, and thereby plays an important role in determining the weather.

### 2.4 THE EARTH’S FIGURE AND GRAVITY

#### 2.4.1 The figure of the Earth

The true surface of the Earth is uneven and irregular, partly land and partly water. For geophysical purposes the Earth's shape is represented by a smooth closed surface, which is called the figure of the Earth. Early concepts of the figure were governed by religion, superstition and non-scientific beliefs. The first circumnavigation of the Earth, completed in 1522 by Magellan’s crew, established that the Earth was probably round. Before the era of scientific awakening the Earth’s shape was believed to be a sphere. As confirmed by numerous photographs from spacecraft, this is in fact an excellent first approximation to Earth’s shape that is adequate for solving many problems. The original suggestion that the Earth is a spheroid flattened at the poles is credited to Newton, who used a hydrostatic argument to account for the polar flattening. The slightly flattened shape permitted an explanation of why a clock that was precise in Paris lost time near the equator (see Section 2.1).

Earth’s shape and gravity are intimately associated. The figure of the Earth is the shape of an equipotential surface of gravity, in particular the one that coincides with mean sea level. The best mathematical approximation to the figure is an oblate ellipsoid, or spheroid (Fig. 2.20). The precise determination of the dimensions of the Earth (e.g., its polar and equatorial radii) is the main objective of the science of geodesy. It requires an exact knowledge of the Earth’s gravity field, the description of which is the goal of gravimetry.

Modern analyses of the Earth’s shape are based on precise observations of the orbits of artificial Earth satellites. These data are used to define a best-fitting oblate ellipsoid, called the International Reference Ellipsoid. In 1930 geodesists and geophysicists defined an optimum reference ellipsoid based on the best available data at the time. The dimensions of this figure have been subsequently refined as more exact data have become available. In 1980 the International Association of Geodesy adopted a Geodetic Reference System (GRS80) in which the reference ellipsoid has an equatorial radius \(a\) equal to 6378.137 km and a polar radius \(c\) equal to 6356.752 km. Subsequent determinations have resulted in only minor differences in the most important geodetic parameters. Some current values are listed in Table 2.1. The radius of the equivalent sphere \(R\) is found from \(R = (a^2 c)^{1/3}\) to be 6371.000 km. Compared to the best-fitting sphere the spheroid is flattened by about 14.2 km at each pole and the equator bulges by about 7.1 km. The polar flattening \(f\) is defined as the ratio

\[ f = \frac{a-c}{a} \tag{2.43} \]

The flattening of the optimum reference ellipsoid defined in 1930 was exactly 1/297. This ellipsoid, and the variation of gravity on its surface, served as the basis of gravimetric surveying for many years, until the era of satellite geodesy and highly sensitive gravimeters showed it to be too inexact. A recent best estimate of the flattening is \(f = 3.352 \times 10^{-3}\) (i.e., \(f = 1/298.256\)).

If the Earth is assumed to be a rotating fluid in perfect hydrostatic equilibrium (as assumed by Newton’s theory), the flattening should be 1/299.5, slightly smaller than the observed value. The hydrostatic condition assumes that the Earth has no internal strength. A possible explanation for the tiny discrepancy in \(f\) is that the Earth has sufficient strength to maintain a non-hydrostatic figure, and the present figure is inherited from a time of more rapid rotation. Alternatively, the slightly more flattened form of the
Earth may be due to internal density contrasts, which could be the consequence of slow convection in the Earth’s mantle. This would take place over long time intervals and could result in a non-hydrostatic mass distribution.

The cause of the polar flattening is the deforming effect of the centrifugal acceleration. This is maximum at the equator where the gravitational acceleration is smallest. The parameter \( m \) is defined as the ratio of the equatorial centrifugal acceleration to the equatorial gravitational acceleration:

\[
m = \frac{\omega^2 a}{G\varepsilon_a^2} = \frac{\omega^2 a^3}{G\varepsilon_a^3}
\]  

The value of \( m \) based on current geodetic values (Table 2.1) is \( 3.4619 \times 10^{-3} \) (i.e., \( m = 1/288.901 \)).

As a result of the flattening, the distribution of mass within the Earth is not simply dependent on radius. The moments of inertia of the Earth about the rotation axis (C) and any axis in the equatorial plane (A) are unequal. As noted in the previous section the inequality affects the way the Earth responds to external gravitational torques and is a determining factor in perturbations of the Earth’s rotation. The principal moments of inertia define the dynamical ellipticity:

\[
H = \frac{C - \frac{1}{2}(A + B)}{C} = \frac{C - A}{C}
\]  

The dynamical ellipticity is obtained from precise observations of the orbits of artificial satellites of the Earth (see Section 2.4.5.1). The current optimum value for \( H \) is \( 3.2737875 \times 10^{-3} \) (i.e., \( H = 1/305.457 \)).

### 2.4.2 Gravitational potential of the spheroidal Earth

The ellipsoidal shape changes the gravitational potential of the Earth from that of an undeformed sphere. In 1849 J. MacCullagh developed the following formula for the gravitational potential of any body at large distance from its center of mass:

\[
U_G = -G\frac{F}{2r^3} \left( A + B + C - 3I \right) - \cdots
\]

---


<table>
<thead>
<tr>
<th>Parameter</th>
<th>Symbol</th>
<th>Value</th>
<th>Units</th>
<th>Reference</th>
</tr>
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<td><strong>Terrestrial parameters (2004)</strong></td>
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<tr>
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<td>m(^3) kg(^{-1}) s(^{-2})</td>
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<td>Geocentric gravitational constant</td>
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<tr>
<td>Earth’s polar radius: ( c = a(1 - f) )</td>
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<tr>
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2.4 THE EARTH’S FIGURE AND GRAVITY

The first term, of order $r^{-1}$, is the gravitational potential of a point mass or sphere with mass $E$ (Eqs. (2.10) and (2.14)); for the Earth it describes the potential of the undeformed globe. If the reference axes are centered on the body’s center of mass, there is no term in $r^{-2}$. The second term, of order $r^{-2}$, is due to deviations from the spherical shape. For the flattened Earth it results from the mass displacements due to the rotational deformation. The parameters $A$, $B$, and $C$ are the principal moments of inertia of the body and $I$ is the moment of inertia about the line OP joining the center of mass to the point of observation (Fig. 2.21). In order to express the potential accurately an infinite number of terms of higher order in $r$ are needed. In the case of the Earth these can be neglected, because the next term is about 1000 times smaller than the second term.

For a body with planes of symmetry, $I$ is a simple combination of the principal moments of inertia. Setting $A$ equal to $B$ for rotational symmetry, and defining the angle between OP and the rotation axis to be $\theta$, the expression for $I$ is

$$I = A\sin^2\theta + C\cos^2\theta$$

(2.47)

MacCullagh’s formula for the ellipsoidal Earth then becomes

$$U_G = -\frac{G}{r} \left( \frac{C-A}{2} \right) \left( 3\cos^2\theta - 1 \right)$$

(2.48)

The function $(3\cos^2\theta - 1)/2$ is a second-order polynomial in $\cos\theta$, written as $P_2(\cos\theta)$. It belongs to a family of functions called Legendre polynomials (Box 2.2). Using this notation MacCullagh’s formula for the gravitational potential of the oblate ellipsoid becomes

$$U_G = -\frac{G}{r} \left( \frac{C-A}{2} \right) P_2(\cos\theta)$$

(2.49)

This can be written in the alternative form

$$U_G = -\frac{G}{r} \left( 1 - \left( \frac{C-A}{ER} \right) \frac{b^2}{a^2} \right) P_2(\cos\theta)$$

(2.50)

Potential theory requires that the gravitational potential of the spheroidal Earth must satisfy an important equation, the Laplace equation (Box 2.3). The solution of this equation is the sum of an infinite number of terms of increasing order in $1/r$, each involving an appropriate Legendre polynomial:

$$U_G = -\frac{G}{r} \left( 1 - \sum_{n=2}^{\infty} \left( \frac{b}{a} \right)^n J_n P_n(\cos\theta) \right)$$

(2.51)

In this equation the coefficients $J_n$, multiplying $P_n(\cos\theta)$, determine the relative importance of the term of nth order. The values of $J_n$ are obtained from satellite geodesy: $J_2 = 1082.6 \times 10^{-6}$; $J_4 = -2.54 \times 10^{-6}$; $J_6 = -1.59 \times 10^{-6}$; higher orders are insignificant. The most important coefficient is the second order one, the dynamical form-factor $J_2$, which describes the effect of the polar flattening on the Earth’s gravitational potential. Comparison of terms in Eqs. (2.48) and (2.51) gives the result

$$J_2 = \frac{C-A}{ER^2}$$

(2.52)

The term of next higher order ($n = 3$) in Eq. (2.51) describes the deviations from the reference ellipsoid which correspond to a pear-shaped Earth (Fig. 2.22). These deviations are of the order of 7–17 m, a thousand times smaller than the deviations of the ellipsoid from a sphere, which are of the order of 7–14 km.

2.4.3 Gravity and its potential

The potential of gravity ($U_G$) is the sum of the gravitational and centrifugal potentials. It is often called the geopotential. At a point on the surface of the rotating spheroid it can be written

$$U_g = U_G - \frac{1}{2} \omega^2 r^2 \sin^2 \theta$$

(2.53)

If the free surface is an equipotential surface of gravity, then $U_g$ is everywhere constant on it. The shape of the equipotential surface is constrained to be that of the spheroid with flattening $f$. Under these conditions a simple relation is found between the constants $f$, $m$ and $J_2$:

$$J_2 = \frac{1}{3} (2f - m)$$

(2.54)

By equating Eqs. (2.52) and (2.54) and re-ordering terms slightly we obtain the following relationship

$$\frac{C-A}{ER^2} = \frac{1}{3} (2f - m)$$

(2.55)

This yields useful information about the variation of density within the Earth. The quantities $f$, $m$ and $(C-A)/C$ are each equal to approximately $1/300$. Inserting their values in the equation gives $C = 0.33ER^2$. Compare this value with the principal moments of inertia of a hollow spherical shell $(0.66ER^2)$ and a solid sphere with uniform density $(0.4ER^2)$. The concentration of mass near the center causes a reduction in the multiplying factor from 0.66 to 0.4. The value of 0.33 for the Earth implies that, in comparison with a uniform solid sphere, the density must increase towards the center of the Earth.

2.4.4 Normal gravity

The direction of gravity at a point is defined as perpendicular to the equipotential surface through the point. This defines the vertical at the point, while the plane tangential to the equipotential surface defines the horizontal (Fig. 2.20). A consequence of the spheroidal shape of the Earth is that the vertical direction is generally not radial, except on the equator and at the poles.
In the triangle depicted in Fig. B2.2 the side $u$ is related to the other two sides $r$ and $R$ and the angle $\theta$ they enclose by the cosine law. The expression for $1/u$ can then be written:

$$\frac{1}{u} = \left( \frac{R^2 + r^2 - 2rr \cos \theta}{R^2 - 2rr \cos \theta} \right)^{1/2}$$

which on expanding becomes

$$\frac{1}{u} = \frac{1}{R} \left[ 1 + \left( \frac{r}{R} \cos \theta + \left( \frac{r}{R} \right)^2 \frac{3 \cos^2 \theta - 1}{2} \right) + \left( \frac{r}{R} \right)^2 \frac{5 \cos^3 \theta - 3 \cos \theta}{2} + \cdots \right]$$

This infinitely long series of terms in $(r/R)$ is called the reciprocal distance formula. It can be written in shorthand form as

$$\frac{1}{u} = \frac{1}{R} \sum_{n=0}^{\infty} \left( \frac{r}{R} \right)^n P_n(\cos \theta)$$

The angle $\theta$ in this expression describes the angular deviation between the side $r$ and the reference side $R$. The functions $P_n(\cos \theta)$ in the sum are called the ordinary Legendre polynomials of order $n$ in $\cos \theta$. They are named after a French mathematician Adrien Marie Legendre (1752–1833). Each polynomial is a coefficient of $(r/R)^n$ in the infinite sum of terms for $(1/u)$, and so has order $n$. Writing $\cos \theta = x$, and $P_n(\cos \theta) = P_n(x)$, the first few polynomials, for $n = 0, 1, 2, 3$, respectively, are as follows

$$P_0(x) = 1 \quad P_2(x) = \frac{1}{2} (3x^2 - 1)$$
$$P_1(x) = x \quad P_3(x) = \frac{1}{2} (5x^3 - 3x)$$

By substituting $\cos \theta$ for $x$ these expressions can be converted into functions of $\cos \theta$. Legendre discovered that the polynomials satisfied the following second-order differential equation, in which $n$ is an integer and $y = P_n(x)$:

$$\frac{\partial}{\partial x} (1 - x^2) \frac{\partial y}{\partial x} + n(n + 1)y = 0$$

On a spherical Earth there is no ambiguity in how we define latitude. It is the angle at the center of the Earth between the radius and the equator, the complement to the polar angle $\theta$. This defines the geocentric latitude $\lambda'$. However, the geographic latitude in common use is not defined in this way. It is found by geodetic measurement of the angle of elevation of a fixed star above the horizon. But the horizontal plane is tangential to the ellipsoid, not to a sphere (Fig. 2.20), and the vertical direction (i.e., the local direction of gravity) intersects the equator at an angle $\lambda$ that is slightly larger than the geocentric latitude $\lambda'$ (Fig. 2.23). The difference $(\lambda - \lambda')$ is zero at the equator.

This, named in his honor, is the Legendre equation. It plays an important role in geophysical potential theory for situations expressed in spherical coordinates that have rotational symmetry about an axis. This is, for example, the case for the gravitational attraction of a spheroid, the simplified form of the Earth’s shape.

The derivation of an individual polynomial of order $n$ is rather tedious if the expanded expression for $(1/u)$ is used. A simple formula for calculating the Legendre polynomials for any order $n$ was developed by another French mathematician, Olinde Rodrigues (1794–1851). The Rodrigues formula is

$$P_n(x) = \frac{1}{2^n n!} \frac{d^n}{dx^n} (x^2 - 1)^n$$

A relative of this equation encountered in many problems of potential theory is the associated Legendre equation, which written as a function of $x$ is

$$\frac{\partial}{\partial x} (1 - x^2) \frac{\partial y}{\partial x} + \left( n(n + 1) - \frac{m^2}{1 - x^2} \right) y = 0$$

The solutions of this equation involve two integers, the order $n$ and degree $m$. As in the case of the ordinary Legendre equation the solutions are polynomials in $x$, which are called the associated Legendre polynomials and written $P_n^m(x)$. A modification of the Rodrigues formula allows easy computation of these functions from the ordinary Legendre polynomials:

$$P_n^m(x) = (1 - x^2)^{m/2} \frac{\partial}{\partial x} mP_n(x)$$

To express the associated Legendre polynomials as functions of $\theta$, i.e. as $P_n^m(\cos \theta)$, it is again only necessary to substitute $\cos \theta$ for $x$. 

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**Box 2.2: Legendre polynomials**

![Fig. B2.2 Reference triangle for derivation of Legendre polynomials.](image)
2.4 THE EARTH’S FIGURE AND GRAVITY

Box 2.3: Spherical harmonics

Many natural forces are directed towards a central point. Examples are the electrical field of a point charge, the magnetic field of a single magnetic pole, and the gravitational acceleration toward a center of mass. The French astronomer and mathematician Pierre Simon, marquis de Laplace (1749–1827) showed that, in order to fulfill this basic physical condition, the potential of the field must satisfy a second-order differential equation, the Laplace equation. This is one of the most famous and important equations in physics and geophysics, as it applies to many situations in potential theory. For the gravitational potential \( U_G \) the Laplace equation is written in Cartesian coordinates \((x, y, z)\) as

\[
\frac{\partial^2 U_G}{\partial x^2} + \frac{\partial^2 U_G}{\partial y^2} + \frac{\partial^2 U_G}{\partial z^2} = 0
\]

(1)

In spherical polar coordinates \((r, \theta, \phi)\) the Laplace equation becomes

\[
\frac{1}{r^2} \frac{\partial}{\partial r} \left( r^2 \frac{\partial U_G}{\partial r} \right) + \frac{1}{r^2 \sin \theta} \frac{\partial}{\partial \theta} \left( \sin \theta \frac{\partial U_G}{\partial \theta} \right) + \frac{1}{r^2 \sin^2 \theta} \frac{\partial^2 U_G}{\partial \phi^2} = 0
\]

(2)

The variation with azimuth \(\phi\) disappears for symmetry about the rotational axis. The general solution of the Laplace equation for rotational symmetry (e.g., for a spheroidal Earth) is

\[
U_G = \sum_{n=0}^{\infty} \left( A_n r^n + \frac{B_n}{r^{n+1}} \right) Y_n^m(\cos \theta)
\]

(3)

where \( Y_n^m(\cos \theta) \) is an ordinary Legendre polynomial of order \(n\) and the coordinate \(\theta\) is the angular deviation of the point of observation from the reference axis (see Box 2.1). In geographic coordinates \(\theta\) is the co-latitude.

If the potential field is not rotationally symmetric — as is the case, for example, for the geoid and the Earth’s magnetic field — the solution of the Laplace equation varies with azimuth \(\phi\) as well as with radius \(r\) and axial angle \(\theta\) and is given by

\[
U_G = \sum_{n=0}^{\infty} \left( A_n r^n + \frac{B_n}{r^{n+1}} \right) \sum_{m=0}^{n} (a_n^m \cos m \phi + b_n^m \sin m \phi) P_n^m(\cos \theta)
\]

(4)

where in this case \( P_n^m(\cos \theta) \) is an associated Legendre polynomial of order \(n\) and degree \(m\) as described in Box 2.2. This equation can in turn be written in modified form as

\[
U_G = \sum_{n=0}^{\infty} \left( A_n r^n + \frac{B_n}{r^{n+1}} \right) \sum_{m=0}^{n} Y_n^m(0, \phi) P_n^m(\cos \theta)
\]

(5)

The function

\[
Y_n^m(0, \phi) = (a_n^m \cos m \phi + b_n^m \sin m \phi) P_n^m(\cos \theta)
\]

(6)

is called a spherical harmonic function, because it has the same value when \(\theta\) or \(\phi\) is increased by an integral multiple of \(2\pi\). It describes the variation of the potential with the coordinates \(\theta\) and \(\phi\) on a spherical surface (i.e., for which \(r\) is a constant). Spherical harmonic functions are used, for example, for describing the variations of the gravitational and magnetic potentials, geoid height, and global heat flow with latitude and longitude on the surface of the Earth.

and poles and reaches a maximum at a latitude of \(45^\circ\), where it amounts to only 0.19% (about 12').

The International Reference Ellipsoid is the standardized reference figure of the Earth. The theoretical value of gravity on the rotating ellipsoid can be computed by differentiating the gravity potential (Eq. (2.53)). This yields the radial and transverse components of gravity, which are then combined to give the following formula for gravity normal to the ellipsoid:

\[
g_n = g_0 (1 + \beta_1 \sin^2 \lambda + \beta_2 \sin^2 2\lambda)
\]

(2.56)

Where, to second order in \(f\) and \(m\),

\[
g_n = g_0 \left(1 + f - \frac{3}{2} m + f^2 - \frac{27}{14} fm\right)
\]

\[
\beta_1 = \frac{5}{2} m - f + \frac{15}{4} m^2 - \frac{17}{14} fm
\]

\[
\beta_2 = \frac{1}{8} f^2 - \frac{5}{8} fm
\]

(2.57)
Equation (2.56) is known as the normal gravity formula. The constants in the formula, defined in 1980 for the Geodetic Reference System (GRS80) still in common use, are:

\[
g_{e} = 9.780 \, 327 \, \text{m} \, \text{s}^{-2}; \quad \beta_1 = 5.30244 \times 10^{-3}; \quad \beta_2 = -5.8 \times 10^{-6}.
\]

They allow calculation of normal gravity at any latitude with an accuracy of 0.1 mGal. Modern instruments can measure gravity differences with even greater precision, in which case a more exact formula, accurate to 0.0001 mGal, can be used. The normal gravity formula is very important in the analysis of gravity measurements on the Earth, because it gives the theoretical variation of normal gravity \((g)\) with latitude on the surface of the reference ellipsoid.

The normal gravity is expressed in terms of \(g_e\), the value of gravity on the equator. The second-order terms \(\beta_2\) and \(\beta_3\) are about 300 times smaller than the first-order terms \(\beta_1\). If we drop second-order terms and use \(\lambda = 90^\circ\), the value of normal gravity at the pole is \(g_p = g_e\), so by rearranging and retaining only first-order terms, we get

\[
g_p - g_e = \frac{5}{2} \beta_1 - f
\]

This expression is called Clairaut’s theorem. It was developed in 1743 by a French mathematician, Alexis-Claude Clairaut, who was the first to relate the variation of gravity on the rotating Earth with the flattening of the spheroid. The normal gravity formula gives \(g_p = 9.832\) 186 m s \(^{-2}\). Numerically, this gives an increase in gravity from equator to pole of approximately 5.186 \(\times 10^{-2}\) m s \(^{-2}\), or 5186 mGal.

There are two obvious reasons for the poleward increase in gravity. The distance to the center of mass of the Earth is shorter at the the poles than at the equator. This gives a stronger gravitational acceleration \((a_G)\) at the poles. The difference is

\[
\Delta a_G = \frac{G M_e}{c^2} \left( \frac{1}{R_e^2} - \frac{1}{R_p^2} \right)
\]

This gives an excess gravity of approximately 6600 mGal at the poles. The effect of the centrifugal force in diminishing gravity is largest at the equator, where it equals \((ma_G)\), and is zero at the poles. This also results in a poleward increase of gravity, amounting to about 3375 mGal. These figures indicate that gravity should increase by a total of 9975 mGal from equator to pole, instead of the observed difference of 5186 mGal. The discrepancy can be resolved by taking into account a third factor. The computation of the difference in gravitational attraction is not so simple as indicated by Eq. (2.59). The equatorial bulge places an excess of mass under the equator, increasing the equatorial gravitational attraction and thereby reducing the gravity decrease from equator to pole.

2.4.5 The geoid

The international reference ellipsoid is a close approximation to the equipotential surface of gravity, but it is really a mathematical convenience. The physical equipotential surface of gravity is called the geoid. It reflects the true
2.4 THE EARTH’S FIGURE AND GRAVITY

distribution of mass inside the Earth and differs from the theoretical ellipsoid by small amounts. Far from land the geoid agrees with the free ocean surface, excluding the temporary perturbing effects of tides and winds. Over the continents the geoid is affected by the mass of land above mean sea level (Fig. 2.24a). The mass within the ellipsoid causes a downward gravitational attraction toward the center of the Earth, but a hill or mountain whose center of gravity is outside the ellipsoid causes an upward attraction. This causes a local elevation of the geoid above the ellipsoid. The displacement between the geoid and the ellipsoid is called a geoid undulation; the elevation caused by the mass above the ellipsoid is a positive undulation.

2.4.5.1 Geoid undulations

In computing the theoretical figure of the Earth the distribution of mass beneath the ellipsoid is assumed to be homogeneous. A local excess of mass under the ellipsoid will deflect and strengthen gravity locally. The potential of the ellipsoid is achieved further from the center of the Earth. The equipotential surface is forced to warp upward while remaining normal to gravity. This gives a positive geoid undulation over a mass excess under the ellipsoid (Fig. 2.24b). Conversely, a mass deficit beneath the ellipsoid will deflect the geoid below the ellipsoid, causing a negative geoid undulation. As a result of the uneven topography and heterogeneous internal mass distribution of the Earth, the geoid is a bumpy equipotential surface.

The potential of the geoid is represented mathematically by spherical harmonic functions that involve the associated Legendre polynomials (Box 2.3). These are more complicated than the ordinary Legendre polynomials used to describe the gravitational potential of the ellipsoid (Eqs. (2.49)–(2.51)). Until now we have only considered variation of the potential with distance \( r \) and with the co-latitude angle \( \theta \). This is an oversimplification, because density variations within the Earth are not symmetrical about the rotation axis. The geoid is an equipotential surface for the real density distribution in the Earth, and so the potential of the geoid varies with longitude as well as co-latitude. These variations are taken into account by expressing the potential as a sum of spherical harmonic functions, as described in Box 2.3. This representation of the geopotential is analogous to the simpler expression for the gravitational potential of the rotationally symmetric Earth using a series of Legendre polynomials (Eq. (2.51)).

In modern analyses the coefficient of each term in the geopotential – similar to the coefficients \( J_n \) in Eq. (2.51) – can be calculated up to a high harmonic degree. The terms up to a selected degree are then used to compute a model of the geoid and the Earth’s gravity field. A combination of satellite data and surface gravity measurements was used to construct the Goddard Earth Model (GEM) 10. A global comparison between a reference ellipsoid with flattening 1/298.257 and the geoid surface computed from the GEM 10 model shows long-wavelength geoid undulations (Fig. 2.25). The largest negative undulation (–105 m) is in the Indian Ocean south of India, and the largest positive undulation (+73 m) is in the equatorial Pacific Ocean north of Australia. These large-scale features are too broad to be ascribed to shallow crustal or lithospheric mass anomalies. They are thought to be due to heterogeneities that extend deep into the lower mantle, but their origin is not yet understood.

2.4.6 Satellite geodesy

Since the early 1960s knowledge of the geoid has been dramatically enhanced by the science of satellite geodesy.
The motions of artificial satellites in Earth orbits are influenced by the Earth’s mass distribution. The most important interaction is the simple balance between the centrifugal force and the gravitational attraction of the Earth’s mass, which determines the radius of the satellite’s orbit. Analysis of the precession of the Earth’s rotation axis (Section 2.3.4.4) shows that it is determined by the dynamical ellipticity \( H \), which depends on the difference between the principal moments of inertia resulting from the rotational flattening. In principle, the gravitational attraction of an artificial satellite on the Earth’s equatorial bulge also contributes to the precession, but the effect is too tiny to be measurable. However, the inverse attraction of the equatorial bulge on the satellite causes the orbit of the satellite to precess around the rotation axis. The plane of the orbit intersects the equatorial plane in the line of nodes. Let this be represented by the line CN1 in Fig. 2.26. On the next passage of the satellite around the Earth the precession of the orbit has moved the nodal line to a new position CN2. The orbital precession in this case is retrograde; the nodal line regresses. For a satellite orbiting in the same sense as the Earth’s rotation the longitude of the nodal line shifts gradually westward; if the orbital sense is opposite to the Earth’s rotation the longitude of the nodal line shifts gradually eastward. Because of the precession of its orbit the path of a satellite eventually covers the entire Earth between the north and south circles of latitude defined by the inclination of the orbit. The profusion of high-quality satellite data is the best source for calculating the dynamical ellipticity or the related parameter \( J_2 \) in the gravity potential. Observations of satellite orbits are so precise that small perturbations of the orbit can be related to the gravitational field and to the geoid.

2.4.6.1 Satellite laser-ranging

The accurate tracking of a satellite orbit is achieved by satellite laser-ranging (SLR). The spherical surface of the target satellite is covered with numerous retro-reflectors. A retro-reflector consists of three orthogonal mirrors that form the corner of a cube; it reflects an incident beam of light back along its path. A brief pulse of laser light with a wavelength of 532 nm is sent from the tracking station on Earth to the satellite, and the two-way travel-time of the reflected pulse is measured. Knowing the speed of light, the distance of the satellite from the tracking station is obtained. The accuracy of a single range measurement is about 1 cm.

America’s Laser Geodynamics Satellite (LAGEOS 1) and France’s Starlette satellite have been tracked for many years. LAGEOS 1 flies at 5858–5958 km altitude, the inclination of its orbit is 110° (i.e., its orbital sense is opposite to the Earth’s rotation), and the nodal line of the orbit advances at 0.343° per day. Starlette flies at an altitude of 806–1108 km, its orbit is inclined at 50°, and its nodal line regresses at 3.95° per day.
The track of a satellite is perturbed by many factors, including the Earth’s gravity field, solar and lunar tidal effects, and atmospheric drag. The perturbing influences of these factors can be computed and allowed for. For the very high accuracy that has now been achieved in SLR results, variations in the coordinates of the tracking stations become detectable. The motion of the pole of rotation of the Earth can be deduced and the history of changes in position of the tracking station can be obtained. LAGEOS 1 was launched in 1976 and has been tracked by more than twenty laser-tracking stations on five tectonic plates. The relative changes in position between pairs of stations can be compared with the rates of plate tectonic motion deduced from marine geophysical data. For example, a profile from the Yaragadee tracking station in Australia and the tracking station in Hawaii crosses the converging plate boundary between the Indo-Australian and Pacific plates (Fig. 2.27). The results of four years of measurement show a decrease of the arc distance between the two stations at a rate of $63 \pm 3$ mm yr$^{-1}$. This is in good agreement with the corresponding rate of $67$ mm yr$^{-1}$ inferred from the relative rotation of the tectonic plates.

Satellite altimeters are best suited for marine surveys, where sub-meter accuracy is possible. The satellite GEOS-3 flew from 1975–1978, SEASAT was launched in 1978, and GEOSAT was launched in 1985. Specifically designed for marine geophysical studies, these satellite altimeters revealed remarkable aspects of the marine geoid. The long-wavelength geoid undulations (Fig. 2.25) have large amplitudes up to several tens of meters and are maintained by mantle-wide convection. The short-wavelength features are accentuated by removing the computed geoid elevation up to a known order and degree. The data are presented in a way that emphasizes the elevated and depressed areas of the sea surface (Fig. 2.28).

There is a strong correlation between the short-wavelength anomalies in elevation of the mean sea surface and features of the sea-floor topography. Over the ocean ridge systems and seamount chains the mean sea surface (geoid) is raised. The locations of fracture zones, in which one side is elevated relative to the other, are clearly discernible. Very dark areas mark the locations of deep ocean trenches, because the mass deficiency in a trench depresses the geoid. Seaward of the deep ocean trenches the mean sea surface is raised as a result of the upward flexure of the lithosphere before it plunges downward in a subduction zone.

### 2.4.6.2 Satellite altimetry

From satellite laser-ranging measurements the altitude of a spacecraft can be determined relative to the reference ellipsoid with a precision in the centimetre range. In satellite altimetry the tracked satellite carries a transmitter and receiver of microwave (radar) signals. A brief electromagnetic pulse is emitted from the spacecraft and reflected from the surface of the Earth. The two-way travel-time is converted using the speed of light to an estimate of the height of the satellite above the Earth’s surface. The difference between the satellite’s height above the ellipsoid and above the Earth’s surface gives the height of the topography relative to the reference ellipsoid. The precision over land areas is poorer than over the oceans, but over smooth land features like deserts and inland water bodies an accuracy of better than a meter is achievable.

### 2.4.6.3 Satellite-based global positioning systems (GPS)

Geodesy, the science of determining the three-dimensional coordinates of a position on the surface of the Earth, received an important boost with the advent of the satellite era. The first global satellite navigation system, the US Navy Navigation Satellite System known as TRANSIT consisted of six satellites in polar orbits about 1100 km above the surface of the Earth. Signals transmitted from these satellites were combined in a receiver on Earth with a signal generated at the same frequency in the receiver. Because of the motion of the satellite the frequency of its signal was modified by the Doppler effect and was thus slightly different from the receiver-generated signal, producing a beat frequency. Using the speed of light, the beat signal was
converted to the oblique distance between the satellite and receiver. By integrating the beat signal over a chosen time interval the change in range to the satellite in the interval was obtained. This was repeated several times. The orbit of the satellite was known precisely from tracking stations on the ground, and so the position of the receiver could be calculated. Originally developed to support ballistic missile submarines in the 1960s, the system was extended to civilian navigation purposes, especially for fixing the position of a ship at sea. The TRANSIT program was terminated in 1996, and succeeded by the more precise GPS program.

The Navigation Satellite Timing and Ranging Global Positioning System (NAVSTAR GPS, or, more commonly, just GPS) utilizes satellites in much higher orbits, at an altitude of around 20,200 km (i.e., a radial distance of 26,570 km), with an orbital period of half a sidereal day. The GPS system consists of 24 satellites. There are four satellites in each of six orbital planes, equally spaced at 60° intervals around the equator and inclined to the equator at about 55°. Between five and eight GPS satellites are visible at any time and location on Earth. Each satellite broadcasts its own predetermined position and reference signal every six seconds. The time difference between emission and reception on Earth gives the “pseudo-range” of the satellite, so-called because it must be corrected for errors in the clock of the receiver and for tropospheric refraction. Pseudo-range measurements to four or more satellites with known positions allows computation of the clock error and the exact position of the receiver. The precision with which a point can be located depends on the quality of the receiver and signal processing. Low-cost single civilian-quality receivers have about 100 m positioning accuracy. In scientific and military missions a roving receiver is used in conjunction with a base station (a fixed receiver), and differential signal processing improves the accuracy of location to around 1 cm.

The GPS system allows very precise determination of changes in the distance between observation points. For example, a dense network of GPS measurements was made in southeastern Italy, the Ionian Islands and western Greece in 1989 and 1993. The differences between the two measuring campaigns show that southwestern Greece moved systematically to the southwest relative to Matera in southeastern Italy at mean annual rates of 20–40 mm yr⁻¹ (Fig. 2.29).

2.4.6.4 Measurement of gravity and the geoid from orbiting satellites

The equipotential surface of gravity, the geoid (Section 2.4.5), is characterized by undulations caused by inhomogeneous distribution of mass in the Earth. Until recently, construction of a global model of the geoid was very laborious, as it required combining data from many different sources of variable precision. Surface gravity measurements made on land or at sea were augmented by data from a large number of Earth-orbiting satellites. The resulting figure showed large-scale features (Fig. 2.25), but fine details were impossible to define accurately. Satellites in comparatively low orbits, a few hundreds of
2.4 THE EARTH’S FIGURE AND GRAVITY

kilometers above the Earth’s surface, can now be used in conjunction with the GPS satellites orbiting at high altitudes (20,200 km) to measure the global gravity field and geoid with a precision that is several orders of magnitude better than was previously possible.

In 2000 the German CHAMP (Challenging Mini-satellite Payload) satellite was inserted into a nearly circular, almost polar orbit with an initial altitude of 450 km. At this altitude the thin atmosphere is still capable of exerting drag, which lowers the altitude of the satellite to about 300 km over a 5 year interval. Sensitive accelerometers on the board allow correction for non-gravitational forces, such as atmospheric drag or the pressure of solar radiation. A highly precise GPS receiver on board the CHAMP satellite, using position data from up to 12 GPS satellites simultaneously, allows retrieval of CHAMP’s position with an accuracy of a few centimeters. Whereas the orbits of earlier satellites were compiled from many comparatively short tracks measured when the satellite was in view of different ground stations, the CHAMP orbit is continuously tracked by the GPS satellites. Small perturbations of CHAMP’s orbit may be tracked and modelled. The models of the Earth’s gravity field and of the global geoid derived from CHAMP data were greatly improved in accuracy and definition over previous models.

Building on the experience gained from CHAMP, a joint American–German project, the Gravity Recovery and Climate Experiment (GRACE), was launched in 2002. The GRACE mission uses two nearly identical satellites in near-circular polar orbits (inclination 89.5° to the equator), initially about 500 km above Earth’s surface. The twin satellites each carry on-board GPS receivers, which allow precise determination of their absolute positions over the Earth at any time. The satellites travel in tandem in the same orbital plane, separated by approximately 220 km along their track. Changes in gravity along the orbit are determined by observing small differences in the separation of the two satellites. This is achieved by using a highly accurate microwave ranging system. Each satellite carries a microwave antenna transmitting in the K-band frequency range (wavelength ~1 cm) and directed accurately at the other satellite. With this ranging system the separation of the two satellites can be measured with a precision of one micrometer (1 μm).

As the satellite-pair orbits the Earth, it traverses variations in the gravity field due to the inhomogeneous mass distribution in the Earth. If there is a mass excess, the equipotential surface bulges upward, and gravity is enhanced locally. The leading satellite encounters this anomaly first and is accelerated away from the trailing satellite. Tiny changes in separation between the two satellites as they move along-track are detected by the accurate microwave ranging system. In conjunction with exact location of the satellite by the on-board GPS devices, the GRACE satellites provide fine-scale definition of the gravity field, and determination of the geoid from a single source. Moreover, the satellites measure the gravity field completely in about 30 days. Thus, comparison of data from selected surveys of a region can reveal very small, time-dependent changes in gravity resulting, for example, from transient effects such as changes in groundwater level, or the melting of glaciers, in the observed region. Other instruments on board the GRACE satellites make further observations for atmospheric and ionospheric research.

2.4.6.5 Observation of crustal deformation with satellite-borne radar

Among the many satellites in Earth orbit, some (identified by acronyms such as ERS1, ERS2, JERS, IRS, RADARSAT, Envisat, etc.) are specifically designed to detect beams of radar waves at the Earth and record the reflections from the Earth’s surface. Synthetic aperture radar (SAR) is a remote sensing technique that has made it possible to record features of the Earth’s surface in remarkable detail based on these radar reflections. In a typical SAR investigation enormous amounts of radar data are gathered and subjected to complex data-processing. This requires massive computational power, and so is usually performed on the ground after the survey has been carried out.

Radar signals, like visible light, are subject to reflection, refraction and diffraction (these phenomena are explained in Section 3.6.2 for seismic waves). Diffraction (see Fig. 3.55) bends light passing through a lens in such a way that a point source becomes diffuse. When two adjacent point sources are observed with the lens, their diffuse images overlap and if they are very close, they may not be seen as distinct points. The resolving power of an optical instrument, such as a lens, is defined by the smallest angular separation (θ) of two points that the instrument can distinguish clearly. For a given lens this angle is dependent inversely on the diameter (d) of the aperture that allows light to pass through the lens, and directly on the wavelength (λ) of the light. It is given by the approximate relationship \( θ \sim \frac{λ}{d} \). High resolution requires that closely spaced details of the object can be distinguished, i.e., the angular resolution θ should be a small number. Thus, the larger the aperture of the lens, the higher is the optical resolution.

The same principle applies to radar. Instead of being dependent on the diameter of an optical lens, the resolution of a radar system is determined by the antenna length. When mounted in a satellite, the physical dimensions of the antenna are limited to just a few meters. SAR makes use of the motion of the antenna and powerful data-processing to get around this limitation.

The radar antenna is mounted so that it directs its beam at right angles to the direction of motion of the host spacecraft. The beam “illuminates” a swathe of ground surface, each particle of which reflects a signal to the antenna. Hundreds of radar pulses are sent out per second (e.g., the European Radar Satellites (ERS) emit 1700 pulses per
second); this results in huge amounts of reflected signals. As the craft moves forward, the illuminated swathe moves across the target surface. Each particle of the target reflects hundreds of radar pulses from the time when it is first energized until it is no longer covered by the beam. During this time the craft (and real antenna) move some distance along the track. In subsequent data-processing the signals reflected from the target are combined and corrected for the changing position of the antenna in such a way that they appear to have been gathered by an antenna as long as the distance moved along the track. This distance is called the synthetic aperture of the radar. For example, a SAR investigation with an ERS1 satellite in orbit 800 km above the Earth’s surface created a synthetic aperture of about 4 km. The high resolving power achieved with this large aperture produced SAR images of ground features with a resolution of about 30 m.

An important aspect of the data reduction is the ability to reconstruct the path of each reflection precisely. This is achieved using the Doppler effect, the principle of which is described in Box 1.2. Reflections from target features ahead of the moving spacecraft have elevated frequencies; those from behind have lowered frequencies. Correcting the frequency of each signal for its Doppler shift is necessary to obtain the true geometry of the reflections.

A further development of the SAR method is Interferometric SAR (InSAR). This technique analyzes the phase of the reflected radar signal to determine small changes in topography between repeated passages of the satellite over an area. The phase of the wave is a measure of the time-delay the wave experiences in transit between transmitter and receiver. To illustrate this point, picture the shape of a waveform as an alternation of crests and troughs, which leaves the satellite at the instant its amplitude is maximum (i.e., at a crest). If the reflected signal returns to the satellite as a crest, it has the same phase as the transmitted signal. Its amplitude could be expressed by the equation \( y = A \cos(\omega t) \). This will be the case if the path of the reflected wave to and from the target is an exact number of complete wavelengths. On the other hand, if the reflection arrives back at the satellite as a trough, it is exactly out-of-phase with the original wave. This happens when the length of its path is an odd number of half-wavelengths. More generally, the path length is not an exact even or odd number of half-wavelengths, and the equation of the reflected wave must be written \( y = A \cos(\omega t + \Delta) \), where the phase difference \( \Delta \) depends on the path length. The InSAR technique developed in the 1990s is based on analysis of the phases inherent in each reflection recorded by the satellite.

If a SAR image is made of a target area during one orbit, it should be reproduced exactly on a later orbit that revisits the same location (this is not exactly possible, but paths that repeat within a few hundred meters can be corrected for geometric differences). In particular, because each point of the target is the same distance from the transmitter, the phases of the imaged signals should be identical. However, if geological events have caused surface displacements between the times of the two images there will be phase differences between the two images. These are made visible by combining the two images so that they interfere with each other.

When harmonic signals with different phases are mixed, they interfere with each other. Constructive interference occurs when the signals have the same phase; if they are superposed, the combined signal is strengthened. Destructive interference occurs when out-of-phase signals are mixed; the combined signal is weakened. The interference pattern that results from mixing the two waveforms consists of alternating zones of reinforcement and reduction of the signal, forming a sequence of so-called “interference fringes.” The use of color greatly enhances the visual impact of the interference fringes.

When this procedure is carried out with SAR images, the resulting interference pattern makes it possible to interpret ground motion of large areas in much greater detail than would be possible from ground-based observations. The method has been used to record various large-scale ground displacements related, for example, to earthquakes, tectonic faulting, and volcanism. Figure 2.30 shows an interference pattern superposed on the background topography of Mount Etna, in Sicily, following a cycle of eruptions in 1992 and 1993. Successive radar images from a common vantage point were obtained 13 months apart by the ERS1 satellite, which transmitted radar signals with wavelength 5.66 cm. In order to change the along-path distance to and from the target by a full wavelength, the ground must move by a half-wavelength perpendicular to the path, in this case by 2.83 cm. The concentric dark and light fringes around the

![Fig. 2.30 Interferometric Synthetic Aperture Radar (InSAR) pattern of interference fringes showing changes in elevation of Mount Etna, Sicily, following the 1992–1993 eruptive cycle. The four pairs of light and dark fringes correspond to subsidence of the mountaintop by about 11 cm as magma drains out of the volcano (after Massonnet, 1997).](image)
2.5 GRAVITY ANOMALIES

2.5.1 Introduction

The mean value of gravity at the surface of the Earth is approximately 9.80 m s\(^{-2}\), or 980,000 mgal. The Earth’s rotation and flattening cause gravity to increase by roughly 5300 mgal from equator to pole, which is a variation of only about 0.5%. Accordingly, measurements of gravity are of two types. The first corresponds to determination of the absolute magnitude of gravity at any place; the second consists of measuring the change in gravity from one place to another. In geophysical studies, especially in gravity prospecting, it is necessary to measure accurately the small non-tidal LOD variations are associated with changes in the angular momentum of the atmosphere due to shifts in the east–west component of the wind patterns. To conserve the total angular momentum of the Earth a change in the angular momentum of the atmosphere must be compensated by an equal and opposite change in the angular momentum of the crust and mantle. The largely seasonal transfers of angular momentum correlate well with high-frequency variations in the LOD obtained from VLBI results (Fig. 2.32).

If the effects of marine tidal braking and non-tidal transfers of atmospheric angular momentum variations are taken into account, small residual deviations in the LOD remain. These are related to the tides in the solid Earth (Section 2.3.3.5). The lunar and solar tidal forces deform the Earth elastically and change its ellipticity slightly. The readjustment of the mass distribution necessitates a corresponding change in the Earth’s rate of rotation in order to conserve angular momentum. The expected changes in LOD due to the influence of tides in the solid Earth can be computed. The discrepancies in LOD values determined from VLBI results agree well with the fluctuations predicted by the theory of the bodily Earth-tides (Fig. 2.32).

By combining VLBI observations from different stations the orientation of the Earth to the extra-galactic inertial coordinate system known for geodetic measurements is obtained. Repeated determinations yield a record of the Earth’s orientation and rotational rate with unprecedented accuracy. Motion of the rotation axis (e.g., the Chandler wobble, Section 2.3.4.3) can be described optically with a resolution of 0.5–1 m; the VLBI data have an accuracy of 3–5 cm. The period of angular rotation can be determined to better than 0.1 millisecond. This has enabled very accurate observation of irregularities in the rotational rate of the Earth, which are manifest as changes in the length of the day (LOD).

The most important, first-order changes in the LOD are due to the braking of the Earth’s rotation by the lunar and solar marine tides (Section 2.3.4.1). The most significant variations expected from changes in the angular momentum of the Earth’s orientation and rotational rate with unprecedented accuracy. Motion of the rotation axis (e.g., the Chandler wobble, Section 2.3.4.3) can be described optically with a resolution of 0.5–1 m; the VLBI data have an accuracy of 3–5 cm. The period of angular rotation can be determined to better than 0.1 millisecond. This has enabled very accurate observation of irregularities in the rotational rate of the Earth, which are manifest as changes in the length of the day (LOD).

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Fig. 2.31 Fine-scale fluctuations in the LOD observed by VLBI, and LOD variations expected from changes in the angular momentum of the atmosphere (after Carter, 1989).

Fig. 2.32 High-frequency changes in the LOD after correction for the effects due to atmospheric angular momentum (points) and the theoretical variations expected from the solid bodily Earth-tides (after Carter, 1989).
changes in gravity caused by underground structures. These require an instrumental sensitivity of the order of
0.01 mgal. It is very difficult to design an instrument to measure the absolute value of gravity that has this high
precision and that is also portable enough to be used easily in different places. Gravity surveying is usually carried
out with a portable instrument called a gravimeter, which determines the variation of gravity relative to one or more
reference locations. In national gravity surveys the relative variations determined with a gravimeter may be converted
to absolute values by calibration with absolute measurements made at selected stations.

2.5.2 Absolute measurement of gravity

The classical method of measuring gravity is with a pendulum. A simple pendulum consists of a heavy weight
suspended at the end of a thin fiber. The compound (or reversible) pendulum, first described by Henry Kater in
1818, allows more exact measurements. It consists of a stiff metal or quartz rod, about 50 cm long, to which is
attached a movable mass. Near each end of the rod is fixed a pivot, which consists of a quartz knife-edge resting
on a flat quartz plane. The period of the pendulum is measured for oscillations about one of the pivots. The
pendulum is then inverted and its period about the other pivot is determined. The position of the movable mass is
adjusted until the periods about the two pivots are equal. The distance \( L \) between the pivots is then measured ac-
curately. The period of the instrument is given by

\[
T = 2\pi \sqrt{\frac{I}{mgh}} = 2\pi \sqrt{\frac{L}{g}}
\]  

(2.60)

where \( I \) is the moment of inertia of the pendulum about a pivot, \( h \) is the distance of the center of mass from the pivot, and \( m \) is the mass of the pendulum. Knowing the length \( L \) from Kater’s method obviates knowledge of \( I \), \( m \) and \( h \).

The sensitivity of the compound pendulum is found by differentiating Eq. (2.60). This gives

\[
\frac{\Delta g}{g} = -2\Delta T
\]

(2.61)

To obtain a sensitivity of about 1 mgal it is necessary to determine the period with an accuracy of about 0.5 \( \mu \)s. This can be achieved easily today with precise atomic clocks. The compound pendulum was the main instrument for gravity prospecting in the 1930s, when timing the swings precisely was more difficult. It was necessary to time as accurately as possible a very large number of swings. As a result a single gravity measurement took about half an hour.

The performance of the instrument was handicapped by several factors. The inertial reaction of the housing to
the swinging mass of the pendulum was compensated by mounting two pendulums on the same frame and swing-
ing them in opposite phase. Air resistance was reduced by housing the pendulum assemblage in an evacuated ther-
ostatically controlled chamber. Friction in the pivot was

\[
\text{minimized by the quartz knife-edge and plane, but due to}
\]

minor unevenness the contact edge was not exactly repeat-
able if the assemblage was set up in a different location, which affected the reliability of the measurements. The
apparatus was bulky but was used until the 1950s as the

main method of making absolute gravity measurements.

2.5.2.1 Free-fall method

Modern methods of determining the acceleration of

gravity are based on observations of falling objects. For

an object that falls from a starting position \( z_0 \) with initial

velocity \( u \) the equation of motion gives the position \( z \) at
time \( t \) as

\[
z = z_0 + ut + \frac{1}{2}gt^2
\]

(2.62)

The absolute value of gravity is obtained by fitting a quad-
ratic to the record of position versus time.

An important element in modern experiments is the
accurate measurement of the change of position with a

Michelson interferometer. In this device a beam of mono-

chromatic light passes through a beam splitter, consisting

of a semi-silvered mirror, which reflects half of the light

incident upon it and transmits the other half. This divides the incident ray into two subrays, which subsequently
travel along different paths and are then recombined to
give an interference pattern. If the path lengths differ by a

full wavelength (or a number of full wavelengths) of the

monochromatic light, the interference is constructive. The
recombined light has maximum intensity, giving a bright

interference fringe. If the path lengths differ by half a

wavelength (or by an odd number of half-wavelengths) the

recombined beams interfere destructively, giving a dark

fringe. In modern experiments the monochromatic light
source is a laser beam of accurately known wavelength.

In an absolute measurement of gravity a laser beam is split
along two paths that form a Michelson interferome-

ter (Fig. 2.33). The horizontal path is of fixed length,

while the vertical path is reflected off a corner-cube retro-

reflector that is released at a known instant and falls

down. The path of free-fall is about 0.5 m long. The cube

falls in an evacuated chamber to minimize air resistance.

A photo-multiplier and counter permit the fringes within

any time interval to be recorded and counted. The intensity

of the recombined light fluctuates sinusoidally with

increasing frequency the further and faster the cube falls.

The distance between each zero crossing corresponds to

half the wavelength of the laser light, and so the distance

travelled by the falling cube in any time interval may be

obtained. The times of the zero crossings must be mea-

sured with an accuracy of 0.1 ns (10\(^{-10}\) s) to give an ac-

curacy of 1 \( \mu \)gal in the gravity measurement.

Although the apparatus is compact, it is not quite

portable enough for gravity surveying. It gives measure-
ments of the absolute value of gravity with an accuracy of
about 0.005–0.010 mgal (5–10 μgal). A disadvantage of
the free-fall method is the resistance of the residual air
molecules left in the evacuated chamber. This e
eff
fect is
reduced by placing the retro-reflector in a chamber that
falls simultaneously with the cube, so that in e
eff
ect the
cube falls in still air. Air resistance is further reduced in
the rise-and-fall method.

2.5.2.2 Rise-and-fall method

In the original version of the rise-and-fall method a glass
sphere was fired vertically upward and fell back along the
same path (Fig. 2.34). Timing devices at two different levels
registered the times of passage of the ball on the upward
and downward paths. In each timer a light beam passed
through a narrow slit. As the glass sphere passed the slit it
acted as a lens and focussed one slit on the other. A photo-
multiplier and detector registered the exact passage of the
ball past the timing level on the upward and downward
paths. The distance \( h \) between the two timing levels (around
1 m) was measured accurately by optical interferometry.

Let the time spent by the sphere above the first timing
level be \( T_1 \) and the time above the second level be \( T_2 \);
further, let the distances from the zenith level to the
timing levels be \( z_1 \) and \( z_2 \), respectively. The corresponding
times of fall are \( t_1 = T_1/2 \) and \( t_2 = T_2/2 \). Then,

\[
T_1^2 = \frac{1}{4g}\left(\frac{T_1}{2}\right)^2 \tag{2.63}
\]

with a similar expression for the second timing level.
Their separation is

\[
h = z_1 - z_2 = \frac{1}{8g}(T_1^2 - T_2^2) \tag{2.64}
\]

The following elegantly simple expression for the value of
gravity is obtained:

\[
g = \frac{8h}{(T_1^2 - T_2^2)} \tag{2.65}
\]

Although the experiment is conducted in a high
vacuum, the few remaining air molecules cause a drag
that opposes the motion. On the upward path the air drag
is downward, in the same direction as gravity; on the
downward path the air drag is upward, opposite to the
direction of gravity. This asymmetry helps to minimize
the effects of air resistance.

In a modern variation Michelson interferometry is used
as in the free-fall method. The projectile is a corner-cube

\[
laser-detector
\]

Fig. 2.34 The rise-and-fall method of measuring absolute gravity.
retro-reflector, and interference fringes are observed and counted during its upward and downward paths. Sensitivity and accuracy are comparable to those of the free-fall method.

2.5.3 Relative measurement of gravity: the gravimeter

In principle, a gravity meter, or gravimeter, is a very sensitive balance. The first gravimeters were based on the straightforward application of Hooke’s law (Section 3.2.1). A mass $m$ suspended from a spring of length $s_0$ causes it to stretch to a new length $s$. The extension, or change in length, of the spring is proportional to the restoring force of the spring and so to the value of gravity, according to:

$$F = mg = -k(s - s_0) \tag{2.66}$$

where $k$ is the elastic constant of the spring. The gravimeter is calibrated at a known location. If gravity is different at another location, the extension of the spring changes, and from this the change in gravity can be computed.

This type of gravimeter, based directly on Hooke’s law, is called a stable type. It has been replaced by more sensitive unstable or astatic types, which are constructed so that an additional force acts in the same direction as gravity and opposes the restoring force of the spring. The instrument is then in a state of unstable equilibrium. This condition is realized through the design of the spring. If the natural length $s_0$ can be made as small as possible, ideally zero, Eq. (2.66) shows that the restoring force is then proportional to the physical length of the spring instead of its extension. The zero-length spring, first introduced in the LaCoste–Romberg gravimeter, is now a common element in modern gravimeters. The spring is usually of the helical type. When a helical spring is stretched, the fiber of the spring is twisted; the total twist along the length of the fiber equals the extension of the spring as a whole. During manufacture of a zero-length spring the helical spring is given an extra twist, so that its tendency is to uncoil. An increase in gravity stretches the spring against its restoring force, and the extension is augmented by the built-in pre-tension.

The operation of a gravimeter is illustrated in Fig. 2.35. A mass is supported by a horizontal rod to which a mirror is attached. The position of the rod is observed with a light-beam reflected into a microscope. If gravity changes, the zero-length spring is extended or shortened and the position of the rod is altered, which deflects the light-beam. The null-deflection principle is utilized. An adjusting screw changes the position of the upper attachment of the spring, which alters its tension and restores the rod to its original horizontal position as detected by the light-beam and microscope. The turns of the adjusting screw are calibrated in units of the change in gravity, usually in mgal.

The gravimeter is light, robust and portable. After initially levelling the instrument, an accurate measurement of a gravity difference can be made in a few minutes. The gravimeter has a sensitivity of about 0.01 mgal (10 μgal). This high sensitivity makes it susceptible to small changes in its own properties.
2.5 GRAVITY ANOMALIES

2.5.3.1 Gravity surveying

If a gravimeter is set up at a given place and monitored for an hour or so, the repeated readings are found to vary smoothly with time. The changes amount to several hundredths of a mgal. The instrumental drift is partly due to thermally induced changes in the elastic properties of the gravimeter spring, which are minimized by housing the critical elements in an evacuated chamber. In addition, the elastic properties of the spring are not perfect, but creep slowly with time. The effect is small in modern gravimeters and can be compensated by making a drift correction. This is obtained by repeated occupation of some measurement stations at intervals during the day (Fig. 2.36). Gravity readings at other stations are adjusted by comparison with the drift curve. In order to make this correction the time of each measurement must be noted.

During the day, while measurements are being made, the gravimeter is subject to tidal attraction, including vertical displacement due to the bodily Earth-tides. The theory of the tides is known well (see Section 2.3.3) and their time-dependent effect on gravity can be computed precisely for any place on Earth at any time. Again, the tidal correction requires that the time of each measurement be known.

The goal of gravity surveying is to locate and describe subsurface structures from the gravity effects caused by their anomalous densities. Most commonly, gravimeter measurements are made at a network of stations, spaced according to the purpose of the survey. In environmental studies a detailed high-resolution investigation of the gravity expression of a small area requires small distances of a few meters between measurement stations. In regional gravity surveys, as used for the definition of hidden structures of prospective commercial interest, the distance between stations may be several kilometers. If the area surveyed is not too large, a suitable site is selected as base station (or reference site), and the gravity differences between the surveyed sites and this site are measured. In a gravity survey on a national scale, the gravity differences may be determined relative to a site where the absolute value of gravity is known.

2.5.4 Correction of gravity measurements

If the interior of the Earth were uniform, the value of gravity on the international reference ellipsoid would vary with latitude according to the normal gravity formula (Eq. (2.56)). This provides us with a reference value for gravity measurements. In practice, it is not possible to measure gravity on the ellipsoid at the place where the reference value is known. The elevation of a measurement station may be hundreds of meters above or below the ellipsoid. Moreover, the gravity station may be surrounded by mountains and valleys that perturb the measurement. For example, let P and Q represent gravity stations at different elevations in hilly terrain (Fig. 2.37a).

The theoretical value of gravity is computed at the points R on the reference ellipsoid below P and Q. Thus, we must correct the measured gravity before it can be compared with the reference value.

The hill-top adjacent to stations P and Q has a center of mass that lies higher than the measurement elevation (Fig. 2.37a). The gravimeter measures gravity in the vertical direction, along the local plumb-line. The mass of the hill-top above P attracts the gravimeter and causes an acceleration with a vertically upward component at P. The measured gravity is reduced by the presence of the hill-top; to compensate for this a terrain (or topographic) correction is calculated and added to the measured gravity. A similar effect is observed at Q, but the hill-top above Q is smaller and the corresponding terrain correction is smaller. These corrections effectively level the topography to the same elevation as the gravity station. The presence of a valley next to each measurement station also requires a terrain correction. In this case, imagine that we could fill the valley up to the level of each station with rock of the same density p as under P and Q. The downward attraction on the gravimeter would be increased, so the terrain correction for a valley must also be added to the measured gravity, just as for a hill. Removing the effects of the topography around a gravity station requires making positive terrain corrections (\(\Delta g_t\)) for both hills and valleys.

After levelling the topography there is now a fictive uniform layer of rock with density \(\rho\) between the gravity station and the reference ellipsoid (Fig. 2.37b). The gravitational acceleration of this rock-mass is included in the
measured gravity and must be removed before we can compare with the theoretical gravity. The layer is taken to be a flat disk or plate of thickness $h_P$ or $h_Q$ under each station; it is called the Bouguer plate. Its gravitational acceleration can be computed for known thickness and density $\rho$, and gives a Bouguer plate correction ($\Delta g_{BP}$) that must be subtracted from the measured gravity, if the gravity station is above sea-level. Note that, if the gravity station is below sea-level, we have to fill the space above it up to sea-level with rock of density $\rho$; this requires increasing the measured gravity correspondingly. The Bouguer plate correction ($\Delta g_{BP}$) is negative if the station is above sea-level but positive if it is below sea-level. Its size depends on the density of the local rocks, but typically amounts to about 0.1 mgal m$^{-1}$.

Finally, we must compensate the measured gravity for the elevation $h_P$ or $h_Q$ of the gravity station above the ellipsoid (Fig. 2.37c). The main part of gravity is due to gravitational attraction, which decreases proportionately to the inverse square of distance from the center of the Earth. The gravity measured at $P$ or $Q$ is smaller than it would be, if measured on the ellipsoid at $R$. A free-air correction ($\Delta g_{FA}$) for the elevation of the station must be added to the measured gravity. This correction ignores the effects of material between the measurement and reference levels, as this is taken care of in $\Delta g_{BP}$. Note that, if the gravity station were below sea-level, the gravitational part of the measured gravity would be too large by comparison with the reference ellipsoid; we would need to subtract $\Delta g_{FA}$ in this case. The free-air correction is positive if the station is above sea-level but negative if it is below sea-level (as might be the case in Death Valley or beside the Dead Sea). It amounts to about 0.3 mgal m$^{-1}$.

The free-air correction is always of opposite sense to the Bouguer plate correction. For convenience, the two are often combined in a single elevation correction, which amounts to about 0.2 mgal m$^{-1}$. This must be added for gravity stations above sea-level and subtracted if gravity is measured below sea-level. In addition, a tidal correction ($\Delta g_{T}$) must be made (Section 2.3.3), and, if gravity is measured in a moving vehicle, the Eötvös correction (Section 2.3.5) is also necessary.

After correction the measured gravity can be compared with the theoretical gravity on the ellipsoid (Fig. 2.37d). Note that the above procedure reduces the measured gravity to the surface of the ellipsoid. In principle it is equally valid to correct the theoretical gravity from the ellipsoid upward to the level where the measurement was made. This method is preferred in more advanced types of analysis of gravity anomalies where the possibility of an anomalous mass between the ellipsoid and ground surface must be taken into account.

### 2.5.4.1 Latitude correction

The theoretical gravity at a given latitude is given by the normal gravity formula (Eq. 2.56). If the measured gravity is an absolute value, the correction for latitude is made by subtracting the value predicted by this formula. Often, however, the gravity survey is made with a gravimeter, and the quantity measured, $g_m$, is the gravity difference relative to a base station. The normal reference gravity $g_n$ may then be replaced by a latitude correction, obtained by differentiating Eq. (2.56):

$$\frac{dg}{d\lambda} = g_n (\beta_1 \sin 2\lambda + \beta_2 \sin 4\lambda)$$  \hspace{1cm} (2.67)

After converting $d\lambda$ from radians to kilometers and neglecting the $\beta_1$ term, the latitude correction ($\Delta g_{lat}$) is $0.8140 \sin 2\lambda$ mgal per kilometer of north-south displacement. Because gravity decreases towards the poles, the correction for stations closer to the pole than the base station must be added to the measured gravity.
2.5 GRAVITY ANOMALIES

2.5.4.2 Terrain corrections

The terrain correction (ΔgT) for a hill adjacent to a gravity station is computed by dividing the hill into a number of vertical prisms (Fig. 2.38a). The contribution of each vertical element to the vertical acceleration at the point of observation P is calculated by assuming cylindrical symmetry about P. The height of the prism is h, its inner and outer radii are ri and ro, respectively, the angle subtended at P is φi, and the density of the hill is ρ (Fig. 2.38b). Let the sides of a small cylindrical element be dr, dz and r dφ; its mass is dm = ρ r dφ dr dz and its contribution to the upward acceleration caused by the prism at P is

\[ Δg = G \frac{d\rho}{r^2 + z^2} \cos(θ) = G \frac{d\rho r dz dφ}{r^2 + z^2} \frac{z}{\sqrt{(r^2 + z^2)^2}} \]  

(2.68)

Combining and rearranging terms and the order of integration gives the upward acceleration at P due to the cylindrical prism:

\[ Δg_T = G ρ \int_0^{φ_i} \left( \int_{r_i}^{r_o} \frac{z dz}{(r^2 + z^2)^{3/2}} \right) dr \]  

(2.69)

The integration over φ gives φi; after further integration over z we get:

\[ Δg_T = G ρ φ_i \int_{r_i}^{r_o} \left( \frac{r}{\sqrt{(r^2 + h^2)}} - 1 \right) dr \]  

(2.70)

Integration over r gives the upward acceleration produced at P by the cylinder:

\[ Δg_T = G ρ φ_i \left( \sqrt{(r_o^2 + h^2)} - \sqrt{(r_i^2 + h^2)} \right) \]  

(2.71)

The direction of ΔgT in Fig. 2.38b is upward, opposite to gravity; the corresponding terrain correction must be added to the measured gravity.

In practice, terrain corrections can be made using a terrain chart (Fig. 2.38c) on which concentric circles and radial lines divide the area around the gravity station into sectors that have radial symmetry like the cross-section of the element of a vertical cylinder in Fig. 2.38b. The inner and outer radii of each sector correspond to ri and ro, and the angle subtended by the sector is φi. The terrain correction for each sector within each zone is pre-calculated using Eq. (2.71) and tabulated. The chart is drawn on a transparent sheet that is overlaid on a topographic map at the same scale and centered on the gravity station. The mean elevation within each sector is estimated as accurately as possible, and the elevation difference (i.e., h in Eq. (2.71)) of the sector relative to the station is computed. This is multiplied by the correction factor for the sector to give its contribution to the terrain correction. Finally, the terrain correction at the gravity station is obtained by summing up the contributions of all sectors. The procedure must be repeated for each gravity station. When the terrain chart is centered on a new station, the mean topographic relief within each sector changes and must be computed anew. As a result, terrain corrections are time consuming and tedious. The most important effects come from the topography nearest to the station. However, terrain corrections are generally necessary if a topographic difference within a sector is more than about 5% of its distance from the station.

2.5.4.3 Bouguer plate correction

The Bouguer plate correction (ΔgBP) compensates for the effect of a layer of rock whose thickness corresponds to the elevation difference between the measurement and reference levels. This is modelled by a solid disk of density ρ and infinite radius centered at the gravity station P. The correction is computed by extension of the calculation for the terrain correction. An elemental cylindrical prism is defined as in Fig. 2.38b. Let the angle φ subtended by the prism increase to 2π and the inner radius decrease to zero; the first term in brackets in Eq. (2.71) reduces to h. The gravitational acceleration at the center of a solid disk of radius r is then

\[ Δg_T = 2\pi G ρ \left( h - \sqrt{(r^2 + h^2)} - r \right) \]  

(2.72)

Now let the radius r of the disk increase. The value of h gradually becomes insignificant compared to r; in the limit, when r is infinite, the second term in Eq. (2.72) tends to zero. Thus, the Bouguer plate correction (ΔgBP) is given by

\[ Δg_{BP} = 2\pi G ρ h \]  

(2.73)

Inserting numerical values gives 0.0419 × 10⁻³ ρ mgal m⁻¹ for ΔgBP, where the density ρ is in kg m⁻³ (see Section 2.5.5). The correct choice of density is very important in computing ΔgBP and ΔgT. Some methods of determining the optimum choice are described in detail below.

An additional consideration is necessary in marine gravity surveys. ΔgBP requires uniform density below the surface of the reference ellipsoid. To compute ΔgBP over an oceanic region we must in effect replace the sea-water with rock of density ρ. However, the measured gravity contains a component due to the attraction of the sea-water (density 1030 kg m⁻³) in the ocean basin. The Bouguer plate correction in marine gravity surveys is therefore made by replacing the density ρ in Eq. (2.73) by (ρ – 1030) kg m⁻³. When a shipboard gravity survey is made over a large deep lake, a similar allowance must be made for the depth of water in the lake using an assumed density of (ρ – 1000) kg m⁻³.

2.5.4.4 Free-air correction

The free-air correction (ΔgFA) has a rather colorful, but slightly misleading title, giving the impression that the measurement station is floating in air above the ellipsoid. The density of air at standard temperature and pressure is around 1.3 kg m⁻³ and a mass of air between
the observation and reference levels would cause a detectable gravity effect of about 50 µgal at an elevation of 1000 m. In fact, the free-air correction pays no attention to the density of the material between the measurement elevation and the ellipsoid. It is a straightforward correction for the decrease of gravitational acceleration with distance from the center of the Earth:

\[
\frac{\Delta g}{r^2} = \frac{\partial G_E^3}{\partial r} = -2G_E^3 \frac{\Delta r}{r^3} = -2g
\]

(2.74)

On substituting the Earth’s radius (6371 km) for \(r\) and the mean value of gravity (981,000 mgal) for \(g\), the value of \(\Delta g_{FA}\) is found to be 0.3086 mgal m\(^{-1}\).

### 2.5.4.5 Combined elevation correction

The free-air and Bouguer plate corrections are often combined into a single elevation correction, which is \((0.3086 - (0.0419p \times 10^{-3}))\) mgal m\(^{-1}\). Substituting a typical density for crustal rocks, usually taken to be 2670 kg m\(^{-3}\), gives a combined elevation correction of 0.197 mgal m\(^{-1}\). This must be added to the measured gravity if the gravity station is above the ellipsoid and subtracted if it is below.

The high sensitivity of modern gravimeters allows an achievable accuracy of 0.01–0.02 mgal in modern gravity surveys. To achieve this accuracy the corrections for the variations of gravity with latitude and elevation must be made very exactly. This requires that the precise coordinates of a gravity station must be determined by accurate geodetic surveying. The necessary precision of horizontal positioning is indicated by the latitude correction. This is maximum at 45° latitude, where, in order to achieve a survey accuracy of ±0.01 µgal, the north–south positions of gravity stations must be known to about ±10 m. The requisite precision in vertical positioning is indicated by the combined elevation correction of 0.2 mgal m\(^{-1}\). To achieve a survey accuracy of ±0.01 mgal the elevation of the gravimeter above the reference ellipsoid must be known to about ±5 cm.

The elevation of a site above the ellipsoid is often taken to be its altitude above mean sea-level. However, mean sea-level is equated with the geoid and not with the ellipsoid. Geoid undulations can amount to tens of meters (Section 2.4.5.1). They are long-wavelength features. Within a local survey the distance between geoid and ellipsoid is unlikely to vary much, and the gravity differences from the selected base station are unlikely to be strongly affected. In a national survey the discrepancies due to geoid undulations may be more serious. In the event that geoid undulations are large enough to affect a survey, the station altitudes must be corrected to true elevations above the ellipsoid.

#### 2.5.5 Density determination

The density of rocks in the vicinity of a gravity profile is important for the calculation of the Bouguer plate and terrain corrections. Density is defined as the mass per unit volume of a material. It has different units and different numerical values in the c.g.s. and SI systems. For example, the density of water is 1 g cm\(^{-3}\) in the c.g.s. system, but 1000 kg m\(^{-3}\) in the SI system. In gravity prospecting c.g.s. units are still in common use, but are slowly being replaced by SI units. The formulas given for \(\Delta g_{FA}\) and \(\Delta g_{BP}\) in Eq. (2.71) and Eq. (2.73), respectively, require that density be given in kg m\(^{-3}\).

A simple way of determining the appropriate density to use in a gravity study is to make a representative collection of rock samples with the aid of a geological map. The specific gravity of a sample may be found directly by weighing it first in air and then in water, and applying Archimedes’ principle. This gives its density \(\rho_r\) relative to that of water:

\[
\rho_r = \frac{W_a}{W_a - W_w}
\]

(2.75)

Typically, the densities found for different rock types by this method show a large amount of scatter about their means, and the ranges of values for different rock types overlap (Fig. 2.39). The densities of igneous and metamorphic rocks are generally higher than those of sedimentary rocks. This method is adequate for reconnaissance of an area. Unfortunately, it is often difficult to ensure that the surface collection of rocks is representative of the rock types in subsurface structures, so alternative methods of determining the appropriate density are usually employed. Density can be measured in vertical boreholes, drilled to explore the nature of a presumed structure. The density determined in the borehole is used to refine the interpretation of the structure.

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**Fig. 2.39** Typical mean values and ranges of density for some common rock types (data source: Dobrin, 1976).
2.5 GRAVITY ANOMALIES

2.5.5.2 Gamma–gamma logging

The density of rock formations adjacent to a borehole can be determined from an instrument in the borehole. The principle makes use of the Compton scattering of $\gamma$-rays by loosely bound electrons in the rock adjacent to a borehole. An American physicist, Arthur H. Compton, discovered in 1923 that radiation scattered by loosely bound electrons experienced an increase in wavelength. This simple observation cannot be explained at all if the radiation is treated as a wave; the scattered radiation would have the same wavelength as the incident radiation. The Compton effect is easily explained by regarding the radiation as particles or photons, i.e., particles of quantized energy, rather than as waves. The energy of a photon is inversely proportional to its wavelength. The collision of a $\gamma$-ray photon with an electron is like a collision between billiard balls; part of the photon’s energy is transferred to the electron. The scattered photon has lower energy and hence a longer wavelength than the incident photon. The Compton effect was an important verification of quantum theory.

The density logger, or gamma–gamma logger (Fig. 2.41), is a cylindrical device that contains a radioactive source of $\gamma$-rays, such as $^{137}$Cs, which emits radiation through a narrow slit. The $\gamma$-ray photons collide with the loosely bound electrons of atoms near the hole, and are scattered. A scintillation counter to detect and measure the intensity of $\gamma$-rays is located in the tool about 45–60 cm above the emitter; the radiation reaching it also passes through a slit. Emitter and detector are shielded with lead, and the tool is pressed against the wall of the borehole by a strong spring, so that the only radiation registered is that resulting from the Compton scattering in the surrounding formation. The intensity of detected radiation is determined by the density of electrons, and so by the density of rock near to the logging tool. The $\gamma$-rays penetrate only about 15 cm into the rock.

Calibrated gamma–gamma logs give the bulk density of the rock surrounding a borehole. This information is also needed for calculating porosity, which is defined as the fractional volume of the rock represented by pore spaces. Most sedimentary rocks are porous, the amount depending on the amount of compaction experienced. Igneous and metamorphic rocks generally have low porosity, unless they have been fractured. Usually the pores are filled with air, gas or a fluid, such as water or oil. If the densities of the matrix rock and pore fluid are known, the bulk density obtained from gamma–gamma logging allows the porosity of the rock to be determined.

2.5.5.3 Borehole gravimetry

Modern instrumentation allows gravity to be measured accurately in boreholes. One type of borehole gravimeter is a modification of the LaCoste–Romberg instrument, adapted for use in the narrow borehole and under conditions of elevated temperature and pressure. Alternative
The usage for down-hole density determination is based on instruments that have a comparable sensitivity of about 0.01 mgal. Their usage for down-hole density determination is based on application of the free-air and Bouguer plate corrections.

Gravity stations

Gravity, the figure of the Earth and geodynamics

Fig. 2.41 (a) The design of a gamma–gamma logging device for determining density in a borehole (after Telford et al., 1990), and (b) a schematic gamma–gamma log calibrated in terms of the rock density.

Fig. 2.42 Geometry for computation of the density of a rock layer from gravity measurements made in a vertical borehole.

Let \( g_1 \) and \( g_2 \) be the values of gravity measured in a vertical borehole at heights \( h_1 \) and \( h_2 \), respectively, above the reference ellipsoid (Fig. 2.42). The difference between \( g_1 \) and \( g_2 \) is due to the different heights and to the material between the two measurement levels in the borehole. The value \( g_2 \) will be larger than \( g_1 \) for two reasons. First, because the lower measurement level is closer to the Earth’s center, \( g_2 \) will be greater than \( g_1 \) by the amount of the combined elevation correction, namely \((0.3086 - 0.0419p \times 10^{-3}) \Delta h \) mgal, where \( \Delta h = h_1 - h_2 \). Second, at the lower level \( h_2 \) the gravimeter experiences an upward Bouguer attraction due to the material between the two measurement levels. This reduces the measured gravity at \( h_2 \) and requires a compensating increase to \( g_2 \) of amount \((0.0419p \times 10^{-3}) \Delta h \) mgal. The difference \( \Delta g \) between the corrected values of \( g_1 \) and \( g_2 \) after reduction to the level \( h_2 \) is then

\[
\Delta g = (0.3086 - 0.0419p \times 10^{-3}) \Delta h - 0.0419p \times 10^{-3} \Delta h = (0.3086 - 0.0838p \times 10^{-3}) \Delta h
\]

Rearranging this equation gives the density \( \rho \) of the material between the measurement levels in the borehole:

\[
\rho = (3.683 - 11.93 \frac{\Delta g}{\Delta h}) \times 10^3 \text{kg m}^{-3}
\]
for a distance $\Delta h = 10\, \text{m}$ between measurement levels). This is much larger than the lateral range penetrated by gamma–gamma logging. As a result, effects related to the borehole itself are unimportant.

### 2.5.5.4 Nettleton’s method for near-surface density

The near-surface density of the material under a hill can be determined by a method devised by L. Nettleton that compares the shape of a Bouguer gravity anomaly (see Section 2.5.6) with the shape of the topography along a profile. The method makes use of the combined elevation correction ($\Delta g_{\text{FA}} + \Delta g_{\text{BP}}$) and the terrain correction ($\Delta g_{\text{T}}$), which are density dependent. The terrain correction is less important than the Bouguer plate correction and can usually be neglected.

A profile of closely spaced gravity stations is measured across a small hill (Fig. 2.43). The combined elevation correction is applied to each measurement. Suppose that the true average density of the hill is 2600 $\text{kg m}^{-3}$. If the value assumed for $\rho$ is too small (say, 2400 $\text{kg m}^{-3}$), $\Delta g_{\text{BP}}$ at each station will be too small. The discrepancy is proportional to the elevation, so the Bouguer gravity anomaly is a positive image of the topography. If the value assumed for $\rho$ is too large (say, 2800 $\text{kg m}^{-3}$), the opposite situation occurs. Too much is subtracted at each point, giving a computed anomaly that is a negative image of the topography. The optimum value for the density is found when the gravity anomaly has minimum correlation with the topography.

### 2.5.6 Free-air and Bouguer gravity anomalies

Suppose that we can measure gravity on the reference ellipsoid. If the distribution of density inside the Earth is homogeneous, the measured gravity should agree with the theoretical gravity given by the normal gravity formula. The gravity corrections described in Section 2.5.4 compensate for the usual situation that the point of measurement is not on the ellipsoid. A discrepancy between the corrected, measured gravity and the theoretical gravity is called a gravity anomaly. It arises because the density of the Earth’s interior is not homogeneous as assumed. The most common types of gravity anomaly are the Bouguer anomaly and the free-air anomaly.

The Bouguer gravity anomaly ($\Delta g_B$) is defined by applying all the corrections described individually in Section 2.5.4:

$$\Delta g_B = g_m + (\Delta g_{\text{FA}} - \Delta g_{\text{BP}} + \Delta g_{\text{T}} + \Delta g_{\text{tide}}) - g_n \quad (2.78)$$

In this formula $g_m$ and $g_n$ are the measured and normal gravity values; the corrections in parentheses are the free-air correction ($\Delta g_{\text{FA}}$), Bouguer plate correction ($\Delta g_{\text{BP}}$), terrain correction ($\Delta g_{\text{T}}$) and tidal correction ($\Delta g_{\text{tide}}$).

The free-air anomaly $\Delta g_f$ is defined by applying only the free-air, terrain and tidal corrections to the measured gravity:

$$\Delta g_f = g_m + (\Delta g_{\text{FA}} + \Delta g_{\text{T}} + \Delta g_{\text{tide}}) - g_n \quad (2.79)$$

The Bouguer and free-air anomalies across the same structure can look quite different. Consider first the topographic block (representing a mountain range) shown in Fig. 2.44a. For this simple structure we neglect the terrain and tidal corrections. The difference between the Bouguer anomaly and the free-air anomaly arises from the Bouguer plate correction. In computing the Bouguer anomaly the simple elevation of the measurement station is taken into account together with the free-air correction. The measured gravity contains the attraction of the landmass above the ellipsoid, which is compensated with the Bouguer plate correction. The underground structure
does not vary laterally, so the corrected measurement agrees with the theoretical value and the Bouguer anomaly is everywhere zero across the mountain range. In computing the free-air anomaly only the free-air correction is applied; the part of the measured gravity due to the attraction of the landmass above the ellipsoid is not taken into account. Away from the mountain-block the Bouguer and free-air anomalies are both equal to zero. Over the mountain the mass of the mountain-block increases the measured gravity compared to the reference value and results in a positive free-air anomaly across the mountain range.

In fact, seismic data show that the Earth’s crust is usually much thicker than normal under a mountain range. This means that a block of less-dense crustal rock projects down into the denser mantle (Fig. 2.44b). After making the free-air and Bouguer plate corrections there remains a Bouguer anomaly due to a block that represents the “root-zone” of the mountain range. As this is less dense than the adjacent and underlying mantle it constitutes a mass deficit. The attraction on a gravimeter at stations on a profile across the mountain range will be less than in Fig. 2.44a, so the corrected measurement will be less than the reference value. A strongly negative Bouguer anomaly is observed along the profile. At some distance from the mountain-block the Bouguer and free-air anomalies are equal but they are no longer zero, because the Bouguer anomaly now contains the effect of the root-zone. Over the mountain-block the free-air anomaly has a constant positive offset from the Bouguer anomaly, as in the previous example. Note that, although the free-air anomaly is positive, it falls to a very low value over the center of the block. At this point the attraction of the mountain is partly cancelled by the missing attraction of the less-dense root-zone.

### 2.6 INTERPRETATION OF GRAVITY ANOMALIES

#### 2.6.1 Regional and residual anomalies

A gravity anomaly results from the inhomogeneous distribution of density in the Earth. Suppose that the density of rocks in a subsurface body is \( \rho \) and the density of the rocks surrounding the body is \( \rho_0 \). The difference \( \Delta \rho = \rho - \rho_0 \) is called the density contrast of the body with respect to the surrounding rocks. If the body has a higher density than the host rock, it has a positive density contrast; a body with lower density than the host rock has a negative density contrast. Over a high-density body the measured gravity is augmented; after reduction to the reference ellipsoid and subtraction of the normal gravity a positive gravity anomaly is obtained. Likewise a negative anomaly results over a region of low density. The presence of a gravity anomaly indicates a body or structure with anomalous density; the sign of the anomaly is the same as that of the density contrast and shows whether the density of the body is higher or lower than normal.

The appearance of a gravity anomaly is affected by the dimensions, density contrast and depth of the anomalous body. The horizontal extent of an anomaly is often called its apparent “wavelength.” The wavelength of an anomaly is a measure of the depth of the anomalous mass. Large, deep bodies give rise to broad (long-wavelength), low-amplitude anomalies, while small, shallow bodies cause narrow (short-wavelength), sharp anomalies.

Usually a map of Bouguer gravity anomalies contains superposed anomalies from several sources. The long-wavelength anomalies due to deep density contrasts are called regional anomalies. They are important for understanding the large-scale structure of the Earth’s crust under major geographic features, such as mountain ranges, oceanic ridges and subduction zones. Short-wavelength residual anomalies are due to shallow anomalous masses that may be of interest for commercial exploitation. Geological knowledge is essential for interpreting the residual anomalies. In eroded shield areas, like Canada or Scandinavia, anomalies with very short wavelengths may be due to near-surface mineralized bodies. In sedimentary basins, short- or intermediate-wavelength anomalies may arise from structures related to reservoirs for petroleum or natural gas.

#### 2.6.2 Separation of regional and residual anomalies

The separation of anomalies of regional and local origin is an important step in the interpretation of a gravity map. The analysis may be based on selected profiles across some structure, or it may involve the two-dimensional distribution of anomalies in a gravity map. Numerous techniques have been applied to the decomposition of a gravity anomaly into its constituent parts. They range in sophistication from simple visual inspection of the anomaly pattern to advanced mathematical analysis. A few examples of these methods are described below.
2.6.2.1 Visual analysis

The simplest way of representing the regional anomaly on a gravity profile is by visually fitting the large-scale trend with a smooth curve (Fig. 2.45). The value of the regional gravity given by this trend is subtracted point by point from the Bouguer gravity anomaly. This method allows the interpreter to fit curves that leave residual anomalies with a sign appropriate to the interpretation of the density distribution.

This approach may be adapted to the analysis of a gravity map by visually smoothing the contour lines. In Fig. 2.46a the contour lines of equal Bouguer gravity curve sharply around a local abnormality. The more gently curved contours have been continued smoothly as dotted lines. They indicate how the interpreter thinks the regional gravity field (Fig. 2.46b) would continue in the absence of the local abnormality. The values of the regional and original Bouguer gravity are interpolated from the corresponding maps at points spaced on a regular grid. The regional value is subtracted from the Bouguer anomaly at each point and the computed residuals are contoured to give a map of the local gravity anomaly (Fig. 2.46c). The experience and skill of the interpreter are important factors in the success of visual methods.

2.6.2.2 Polynomial representation

In an alternative method the regional trend is represented by a straight line or, more generally, by a smooth polynomial curve. If \( x \) denotes the horizontal position on a gravity profile, the regional gravity \( g_R \) may be written

\[
g_R = g_0 + g_1x + g_2x^2 + g_3x^3 + \ldots + g_nx^n \quad (2.80)
\]

The polynomial is fitted by the method of least squares to the observed gravity profile. This gives optimum values for the coefficients \( g_n \). The method also has drawbacks. The higher the order of the polynomial, the better it fits the observations (Fig. 2.47). The ludicrous extreme is when the order of the polynomial is one less than the number of observations; the curve then passes perfectly through all the data points, but the regional gravity anomaly has no meaning geologically. The interpreter’s judgement is important in selecting the order of the polynomial, which is usually chosen to be the lowest possible order that represents most of the regional trend. Moreover, a curve fitted by least squares must pass through the mean of the gravity values, so that the residual anomalies are divided equally between positive and negative values. Each residual anomaly is flanked by anomalies of opposite sign (Fig. 2.47), which are due to the same anomalous mass that caused the central anomaly and so have no significance of their own.

Polynomial fitting can also be applied to gravity maps. It is assumed that the regional anomaly can be represented by a smooth surface, \( \Delta g(x, y) \), which is a low order polynomial of the horizontal position coordinates \( x \) and \( y \). In the simplest case the regional anomaly is expressed
### Box 2.4: Fourier analysis

If integrated over a full cycle, the resulting value of a sine or cosine function is zero. The integrated value of the product of a sine function and a cosine function is also zero. Mathematically, this defines the sines and cosines as orthogonal functions. The squared values of sines and cosines do not integrate to zero over a full cycle; this property can be used to normalize functions that can be expressed in terms of sines and cosines. These observations may be summarized as follows for the functions $\sin(n\theta)$ and $\cos(n\theta)$

\[
\int_0^{2\pi} \sin(n\theta)\cos(n\theta) d\theta = \frac{1}{2} \left( 0 \quad \text{if} \quad m \neq n \quad \text{or} \quad \pi \quad \text{if} \quad m = n \right)
\]

These relationships form the basis of Fourier analysis.

Geophysical signals that vary periodically in time (e.g., seismic waves) or space (e.g., gravity, magnetic or thermal anomalies) can be expressed as the superposition of harmonics of a fundamental frequency or wave number. For example, consider a gravity anomaly $\Delta g(x)$ along a profile of length $L$ in the $x$-direction. The fundamental “wavelength” $\lambda$ of the anomaly is equal to $2L$, i.e., twice the profile length. A corresponding fundamental wave number is defined for the profile as $k = (2\pi/\lambda)$, so that the argument $\theta$ in the sine and cosine functions in Eqs. (1) and (2) is replaced by $(kx)$. The observed anomaly is represented by adding together components corresponding to harmonics of the fundamental wavelength:

\[
\Delta g(x) = a_1\cos(kx) + b_1\sin(kx) + a_2\cos(2kx) + b_2\sin(2kx) + a_3\cos(3kx) + b_3\sin(3kx) + \ldots
\]

\[
\Delta g(x) = \sum_{n=1}^{N} (a_n\cos(nkx) + b_n\sin(nkx))
\]

This expression for $\Delta g(x)$ is called a Fourier series. In Eq. (3) the summation is truncated after $N$ sine and cosine terms. The value of $N$ is chosen to be as large as necessary to describe the gravity anomaly adequately. The importance of any individual term of order $n$ is given by the values of the corresponding coefficients $a_n$ and $b_n$, which act as weighting functions.

The coefficients $a_n$ and $b_n$ can be calculated using the orthogonal properties of sine and cosine functions summarized in Eqs. (1) and (2). If both sides of Eq. (3) for the gravity anomaly $\Delta g(x)$ are multiplied by $\cos(nkx)$, we get

\[
\int_0^L \Delta g(x)\cos(nkx) dx = \frac{1}{2} [ a_n \int_0^L \cos(nkx) \cos(nkx) dx + b_n \int_0^L \sin(nkx) \cos(nkx) dx ]
\]

Each product on the right-hand side of this equation can be written as the sum or difference of two sines or cosines. Thus,

\[
\int_0^L \Delta g(x)\cos(nkx) dx = \frac{1}{2} \sum_{m=-\infty}^{\infty} (a_m \cos((n + m)kx) + b_m \sin((n + m)kx))
\]

Integration of $\Delta g(x) \cos(2mkx)$ over a full wavelength of $x$ causes all terms on the right side of Eq. (5) to vanish unless $n = m$, when the term $\cos((n - m)kx) = \cos(0) = 1$. This allows us to determine the coefficients $a_n$ in Eq. (2):

\[
\int_0^L \Delta g(x)\cos(nkx) dx = \frac{1}{2} a_n \int_0^L \cos(nkx) dx = \frac{\pi}{2} a_n
\]

\[
a_n = \frac{2}{L} \int_0^L \Delta g(x)\cos(nkx) dx = \frac{\pi}{2} \int_0^{2\pi} \Delta g(x)\cos(n\theta) d\theta
\]

The coefficients $b_n$ in Eq. (3) are obtained similarly by multiplying $\Delta g(x)$ by $\sin(nkx)$ and integrating over a full cycle of the signal. This gives

\[
b_n = \frac{2}{L} \int_0^L \Delta g(x)\sin(nkx) dx = \frac{\pi}{2} \int_0^{2\pi} \Delta g(x)\sin(n\theta) d\theta
\]
2.6 INTERPRETATION OF GRAVITY ANOMALIES

Box 2.5: Double Fourier series

The two-dimensional variation of a mapped gravity anomaly can be analyzed with the aid of double Fourier series. In this case the gravity anomaly is a function of both the x- and y-coordinates and can be written

\[ \Delta g(x, y) = \sum_{n=1}^{N} \sum_{m=1}^{M} (a_{nm} C_n S_m + b_{nm} C_n S_m + c_{nm} S_n C_m + d_{nm} S_n C_m) \]  

where

\[ C_n = \cos \left( 2n \pi \frac{x}{\lambda_x} \right) \quad S_n = \sin \left( 2n \pi \frac{x}{\lambda_x} \right) \]

\[ C_m = \cos \left( 2m \pi \frac{y}{\lambda_y} \right) \quad S_m = \sin \left( 2m \pi \frac{y}{\lambda_y} \right) \]  

In these expressions the fundamental wavelengths \( \lambda_x \) and \( \lambda_y \) express the extent of the anomaly in the x- and y-directions, respectively. The derivation of the coefficients \( a_{nm}, b_{nm}, c_{nm}, d_{nm} \) is similar in principle to the one-dimensional case, relying on the orthogonality of the individual sine and cosine terms, and the property that the products of two sine terms, two cosine terms or a sine term with a cosine term, can be expressed as the sum or difference of other sine or cosine functions. As might be expected, the analysis of double Fourier series is somewhat more complicated than in the one-dimensional case, but it delivers results that characterize the two-dimensional variation of the regional gravity anomaly.

as a first-order polynomial, or plane. To express changes in the gradient of gravity a higher-order polynomial is needed. For example, the regional gravity given by a second-order polynomial is written

\[ \Delta g(x, y) = \Delta g_0 + \Delta g_{1x}x + \Delta g_{1y}y 
+ \Delta g_{2x}x^2 + \Delta g_{2y}y^2 + \Delta g_{xy}xy \]  

As in the analysis of a profile, the optimum values of the coefficients \( \Delta g_0, \Delta g_{1x}, \Delta g_{1y}, \) etc., are determined by least-squares fitting. The residual anomaly is again computed point by point by subtracting the regional from the original data.

2.6.2.3 Representation by Fourier series

The gravity anomaly along a profile can be analyzed with techniques developed for investigating time series. Instead of varying with time, as the seismic signal does in a seismometer, the gravity anomaly \( \Delta g(x) \) varies with position

\( x \) along the profile. For a spatial distribution the wave number, \( k = 2\pi/\lambda \), is the counterpart of the frequency of a time series. If it can be assumed that its variation is periodic, the function \( \Delta g(x) \) can be expressed as the sum of a series of discrete harmonics. Each harmonic is a sine or cosine function whose argument is a multiple of the fundamental wave number. The expression for \( \Delta g(x) \) is called a Fourier series (Box 2.4).

The breakdown of a complex anomaly (or time series) in terms of simpler periodic variations of different wavelengths is called Fourier analysis and is a powerful method for resolving the most important components of the original signal.

The two-dimensional variation of a mapped gravity anomaly can be expressed in a similar way with the aid of double Fourier series (Box 2.5). As in the simpler one-dimensional case of a gravity anomaly on a profile, the expression of two-dimensional gravity anomalies by double Fourier series is analogous to summing weighted sinusoidal functions. These can be visualized as corrugations of the x–y plane (Fig. 2.48), with each corrugation weighted according to its importance to \( \Delta g(x, y) \).

2.6.2.4 Anomaly enhancement and filtering

The above discussion shows how a function that is periodic can be expressed as a Fourier sum of harmonics of a
In determining the roots of cubic and higher order polynomials it is sometimes necessary to use the square roots of negative numbers. A negative number can be written as the positive number multiplied by \((-1)\), so the square root of a negative number contains the imaginary unit \(i\), defined as the square root of \((-1)\). A complex number \(z\) consists of a real part \(x\) and an imaginary part \(y\), and is written

\[ z = x + iy \]  

(1)

The number \(z^* = x - iy\) is called the complex conjugate of \(z\). The product \(zz^* = x^2 + y^2\) gives the squared amplitude of the complex number.

The great Swiss mathematician Leonhard Euler showed in 1748 how the trigonometric functions, \(\cos \theta\) and \(\sin \theta\), are related to a complex exponential function:

\[ \cos \theta + i \sin \theta = e^{i\theta} \]  

(2)

where \(e\) is the base of the natural logarithms and \(i\) is the imaginary unit. This formula is fundamental to the study of functions of a complex variable, a branch of mathematics known as complex analysis that can be used to solve a wide range of practical and theoretical problems. In this expression, \(\cos \theta\) is called the real part of \(e^{i\theta}\) and \(\sin \theta\) is called the imaginary part. It follows that

\[ i = e^{i\pi/2} \]

\[ \sqrt{-1} = e^{i\pi/4} = \frac{1}{\sqrt{2}}(1 + i) \]  

(3)

Complex numbers can be depicted geometrically on an Argand diagram, (also called the complex plane) invented in 1806 by another Swiss mathematician, Jean-Robert Argand, as an aid to their visualization. In this diagram (Fig. B2.6) the real part of the number, \(x\) in our case, is plotted on the horizontal axis and the imaginary part, \(y\), on the vertical axis. If the distance of the point \(P\) at \((x, y)\) from the origin of the plot at \(O\) is \(r\), and the angle between \(OP\) and the \(x\)-axis is \(\theta\), then the real part of the complex number is \(x = r \cos \theta\) and the imaginary part is \(y = r \sin \theta\), so that, using Euler’s formula

\[ z = x + iy = r(\cos \theta + i \sin \theta) = re^{i\theta} \]  

(4)

Complex numbers are combined in the same way as real numbers. Two complex numbers, \(z_1\) and \(z_2\), combine to form a new complex number, \(z\). A special case is when the complex number is combined with its complex conjugate; this results in a real number. Some examples of possible combinations of complex numbers are as follows:

\[ z = z_1 + z_2 = (x_1 + x_2) + i(y_1 + y_2) \]

\[ z = r_1e^{i\theta_1} + r_2e^{i\theta_2} \]  

(5)

\[ z = z_1z_2 = (x_1 + iy_1)(x_2 + iy_2) \]

\[ = (x_1x_2 - y_1y_2) + i(x_1y_2 - y_1x_2) \]

\[ z = r_1r_2e^{i(\theta_1 + \theta_2)} \]  

(6)

To handle this kind of problem the spatial variation of gravity is represented by a Fourier integral, which consists of a continuous set of frequencies or wave numbers instead of a discrete set. The Fourier integral can be used to represent non-periodic functions. It uses complex numbers (Box 2.6), which are numbers that involve \(i\), the square-root of \(-1\). If the gravity anomaly is analyzed in two dimensions (instead of on a single profile), a two-dimensional integral is needed, analogous to the double Fourier series representation described in Section 2.6.2.3 and Box 2.5. The observed gravity can then be manipulated using the techniques of Fourier transforms (Box 2.7). These techniques
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Box 2.7: Fourier transforms

Functions that cannot be expressed as the Fourier series of individual terms, as in Boxes 2.4 and 2.5, may be replaced by a Fourier integral. This consists of a continuous set of frequencies or wave numbers instead of a discrete set. The Fourier integral can be used to represent non-periodic functions. The gravity anomaly $\Delta g(x)$ is now written as an integral instead of as a sum of discrete terms:

$$\Delta g(x) = \int_{-\infty}^{\infty} G(u) e^{iux} du$$

(1)

where, by using the properties of complex numbers, it can be shown that

$$G(u) = \frac{1}{2\pi} \int_{-\infty}^{\infty} \Delta g(x) e^{-iux} dx$$

(2)

The complex function $G(u)$ is the Fourier transform of the real-valued function $\Delta g(x)$. An adequate treatment of Fourier transforms is beyond the scope of this book. However, the use of this powerful mathematical technique can be illustrated without delving deeply into the theory.

A map of gravity anomalies can be represented by a function $\Delta g(x, y)$ of the Cartesian map coordinates. The Fourier transform of $\Delta g(x, y)$ is a two-dimensional complex function that involves wave numbers $k_x$ and $k_y$ defined by wavelengths of the gravity field with respect to the $x$- and $y$-axes ($k_x = 2\pi/\lambda_x, k_y = 2\pi/\lambda_y$). It is

$$G(x, y) = \int_{-\infty}^{\infty} \int_{-\infty}^{\infty} \Delta g(x, y) \cos(k_x x + k_y y)$$

$$- i\sin(k_x x + k_y y) \, dx \, dy$$

(3)

This equation assumes that the observations $\Delta g(x, y)$ can be represented by a continuous function defined over an infinite $x$-$y$ plane, whereas in fact the data are of finite extent and are known at discrete points of a measurement grid. In practice, these inconsistencies are usually not important. Efficient computer algorithms permit the rapid computation of the Fourier transform $G(x, y)$ of the gravity anomaly $\Delta g(x, y)$. The transformed signal can be readily manipulated in the Fourier domain by convolution with a filter function and the result deconvoluted back into the spatial domain. This allows the operator to examine the effects of low- and high-pass filtering on the observed anomaly, which can greatly help in its interpretation.

Box 2.7 concludes with a statement about the interpretability of gravity anomalies. It states that the interpretation of gravity anomalies is not unique; different density distributions can give the same anomaly. Inconsistencies are usually not important, but they can be problematic. The book suggests that the interpretation of gravity anomalies is not unique; different density distributions can give the same anomaly. Inconsistencies are usually not important, but they can be problematic. The book suggests that the interpretation of gravity anomalies is not unique; different density distributions can give the same anomaly.

2.6.3 Modelling gravity anomalies

After removal of regional effects the residual gravity anomaly must be interpreted in terms of an anomalous density distribution. Modern analyses are based on iterative modelling using high-speed computers. Earlier methods of interpretation utilized comparison of the observed gravity anomalies with the computed anomalies of geometric shapes. The success of this simple approach is due to the insensitivity of the shape of a gravity anomaly to minor variations in the anomalous density distribution. Some fundamental problems of interpreting gravity anomalies are traditional. Modern analyses are based on iterative modelling using high-speed computers. Earlier methods of interpretation utilized comparison of the observed gravity anomalies with the computed anomalies of geometric shapes. The success of this simple approach is due to the insensitivity of the shape of a gravity anomaly to minor variations in the anomalous density distribution. Some fundamental problems of interpreting gravity anomalies are learned from the computed effects of geometric models. In particular, it is important to realize that the interpretation of gravity anomalies is not unique; different density distributions can give the same anomaly. Inconsistencies are usually not important, but they can be problematic. The book suggests that the interpretation of gravity anomalies is not unique; different density distributions can give the same anomaly.
Diapiric structures introduce material of different density into the host rock. A low-density salt dome (ν = 2150 kg m⁻³) intruding higher-density carbonate rocks (ρ = 2500 kg m⁻³) has a density contrast Δρ = -350 kg m⁻³ and causes a negative gravity anomaly. A volcanic plug (ρ = 2800 kg m⁻³) intruding a granite body (ρ = 2600 kg m⁻³) has a density contrast Δρ = +200 kg m⁻³, which causes a positive gravity anomaly. The contour lines on a map of the anomaly are centered on the diapir, so all profiles across the center of the structure are equivalent. The anomalous body can be modelled equally by a vertical cylinder or by a sphere, which we will evaluate here because of the simplicity of the model.

Assume a sphere of radius R and density contrast Δρ with its center at depth z below the surface (Fig. 2.50). The attraction Δg of the sphere is as though the anomalous mass M of the sphere were concentrated at its center. If we measure horizontal position from a point above its center, at distance x the vertical component Δgz is given by

\[ Δg_z = Δg \sin θ = \frac{GMz}{r^2} \]  

(2.82)

where

\[ M = \frac{4}{3} \pi R^3 Δρ \quad \text{and} \quad r^2 = z^2 + x^2 \]

Fig. 2.50 Gravity anomalies for buried spheres with the same radius R and density contrast Δρ but with their centers at different depths z below the surface. The anomaly of the deeper sphere B is flatter and broader than the anomaly of the shallower sphere A.
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This equation shows how the depth to the center of the sphere affects the peak amplitude of the anomaly; the greater the depth to the center, the smaller the amplitude (Fig. 2.50). For a given depth the same peak anomaly can be produced by numerous combinations of $\Delta \rho$ and $R$; a large sphere with a low density contrast can give an identical anomaly to a small sphere with a high density contrast. The gravity data alone do not allow us to resolve this ambiguity.

The terms in the second pair of parentheses in Eq. (2.83) describe how the amplitude of the anomaly varies with distance along the profile. The anomaly is symmetrical with respect to $x$, reaches a maximum value $\Delta g_y$ over the center of the sphere ($x = 0$) and decreases to zero at great distances ($x = \infty$). Note that the larger the depth $z$, the more slowly the amplitude decreases laterally with increasing $x$. A deep source produces a smaller but broader anomaly than the same source at shallower depth. The width $w$ of the anomaly where the amplitude has one-half its maximum value is called the "half-height width." The depth $z$ to the center of the sphere is deduced from this anomaly width from the relationship $z = 0.652w$.

2.6.3.2 Horizontal line element

Many geologically interesting structures extend to great distances in one direction but have the same cross-sectional shape along the strike of the structure. If the length along strike were infinite, the two-dimensional variation of density in the area of cross-section would suffice to model the structure. However, this is not really valid as the lateral extent is never infinite. As a general rule, if the length of the structure normal to the profile is more than twenty times its width or depth, it can be treated as two-dimensional (2D). Otherwise, the end effects due to the limited lateral extent of the structure must be taken into account in computing its anomaly. An elongate body that requires end corrections is sometimes referred to as a 2.5D structure. For example, the mass distribution under elongated bodies like anticlines, synclines and faults should be modelled as 2.5D structures. Here, we will handle the simpler two-dimensional models of these structures.

Let an infinitely long linear mass distribution with mass $m$ per unit length extend horizontally along the $y$-axis at depth $z$ (Fig. 2.51). The contribution $d(\Delta g_y)$ to the vertical gravity anomaly $\Delta g_y$ at a point on the $x$-axis due to a small element of length $dy$ is

$$d(\Delta g_y) = Gm \frac{dy}{r^2} \sin \theta = Gm \frac{dy}{r^2} \frac{z}{\sqrt{r^2 + y^2}} \quad (2.85)$$

The line element extends to infinity along the positive and negative $y$-axis, so its vertical gravity anomaly is found by integration:

$$\Delta g_y = Gmz \int_0^\infty dy \frac{y}{(u^2 + y^2)^{3/2}} = Gmz \int_0^\infty dy \frac{1}{(u^2 + y^2)^{3/2}} \quad (2.86)$$

$$where \, u^2 = x^2 + z^2. \, The \, integration \, is \, simplified \, by \, changing \, variables, \, so \, that \, y = u \tan \varphi, \, then \, dy = u \sec^2 \varphi \, d\varphi \, and \, (u^2 + y^2)^{3/2} = u^3 \sec^3 \varphi. \, This \, gives \,$$

$$\Delta g_y = \frac{Gmz}{u^2} \int_{-w/2}^{w/2} \cos \varphi \, d\varphi \quad (2.87)$$

which, after evaluation of the integral, gives

$$\Delta g_y = \frac{2Gmz}{z^2 + x^2} \quad (2.88)$$

This expression can be written as the derivative of a potential function $\Psi$:

$$\Delta g_y = \frac{Gmz^2}{u^2} = -\frac{\partial \Psi}{\partial z} \quad (2.89)$$

$$\Psi = Gm \log \left( \frac{1}{r} \right) = Gm \log_e \left( \frac{1}{\sqrt{x^2 + z^2}} \right) \quad (2.90)$$

$\Psi$ is called the logarithmic potential. Equations (2.88) and (2.90) are useful results for deriving formulas for the gravity anomaly of linear structures like an anticline (or syncline) or a fault.

2.6.3.3 Horizontal cylinder: model for anticline or syncline

The gravity anomaly of an anticline can be modelled by assuming that the upward folding of strata brings rocks with higher density nearer to the surface (Fig. 2.52a), thereby causing a positive density contrast. A syncline is modelled by assuming that its core is filled with strata of lower density that cause a negative density contrast. In each case the geometric model of the structure is an infinite horizontal cylinder (Fig. 2.52b).

A horizontal cylinder may be regarded as composed of numerous line elements parallel to its axis. The cross-sectional area of an element (Fig. 2.53) gives a mass anomaly per unit length, $m = \Delta \rho r \, d\theta \, dr$. The contribution $d\Psi$ of a line element to the potential at the surface is

$$d\Psi = 2G \Delta \rho \log_e \left( \frac{1}{r} \right) \, d\rho \, d\theta \quad (2.91)$$
The anomaly of a horizontal cylinder decreases laterally less rapidly than that of a sphere, due to the long extent of the cylinder normal to the profile. The “half-height width” \( w \) of the anomaly is again dependent on the depth \( z \) to the axis of the cylinder; in this case the depth is given by \( z = 0.5w \).

### 2.6.3.4 Horizontal thin sheet

We next compute the anomaly of a thin horizontal ribbon of infinite length normal to the plane of the profile. This is done by fictively replacing the ribbon with numerous infinitely long line elements laid side by side. Let the depth of the sheet be \( z \), its thickness \( t \) and its density contrast \( \Delta \rho \) (Fig. 2.54a); the mass per unit length in the \( y \)-direction of a line element of width \( dx \) is \( \Delta \rho dx \). Substituting in Eq. (2.88) gives the gravity anomaly of the line element; the anomaly of the thin ribbon is then computed by integrating between the limits \( x_1 \) and \( x_2 \) (Fig. 2.54b)

\[
\Delta g_z = 2\pi G \int_0^t \frac{R^2 dx}{x^2 + z^2}
\]

Writing \( \tan^{-1}(x_1/z) = \phi_1 \) and \( \tan^{-1}(x_2/z) = \phi_2 \) as in Fig. 2.54b the equation becomes

\[
\Delta g_z = 2\pi G \Delta \rho \left[ \phi_2 - \phi_1 \right] \tag{2.97}
\]

i.e., the gravity anomaly of the horizontal ribbon is proportional to the angle it subtends at the point of measurement.

The anomaly of a semi-infinite horizontal sheet is a limiting case of this result. For easier reference, the origin of \( x \) is moved to the edge of the sheet, so that distances to the left are negative and those to the right are positive (Fig. 2.54c). This makes \( \phi_1 = -\tan^{-1}(x/z) \). The remote
end of the sheet is at infinity, and \( \phi_3 = \pi/2 \). The gravity anomaly is then

\[
\Delta g_z = 2G\Delta \rho t \left[ \frac{\pi}{2} + \tan^{-1}\left(\frac{x}{z}\right) \right] \tag{2.98}
\]

A further example is the infinite horizontal sheet, which extends to infinity in the positive and negative \( x \) and \( y \) directions. With \( \phi_3 = \pi/2 \) and \( \phi_1 = -\pi/2 \) the anomaly is

\[
\Delta g_z = 2\pi G\Delta \rho t \tag{2.99}
\]

which is the same as the expression for the Bouguer plate correction (Eq. (2.73)).

### 2.6.3.5 Horizontal slab: model for a vertical fault

The gravity anomaly across a vertical fault increases progressively to a maximum value over the uplifted side (Fig. 2.55a). This is interpreted as due to the upward displacement of denser material, which causes a horizontal density contrast across a vertical step of height \( h \) (Fig. 2.55b). The faulted block can be modelled as a semi-infinite horizontal slab of height \( h \) and density contrast \( \Delta \rho \) with its mid-point at depth \( z_0 \) (Fig. 2.55c).

Let the slab be divided into thin, semi-infinite horizontal sheets of thickness \( dz \) at depth \( z \). The gravity anomaly of a given sheet is given by Eq. (2.98) with \( dz \) for the thickness \( t \). The anomaly of the semi-infinite slab is found by integrating with respect to \( z \) over the thickness of the slab; the limits of integration are \( z = -h/2 \) and \( z = +h/2 \). After slightly rearranging terms this gives

\[
\Delta g_z = 2G\Delta \rho h \left[ \frac{\pi}{2} + \tan^{-1}\left(\frac{x}{z_0}\right) \right] \tag{2.100}
\]

The second expression in the brackets is the mean value of the angle \( \tan^{-1}(x/z) \) averaged over the height of the fault step. This can be replaced to a good approximation by the value at the mid-point of the step, at depth \( z_0 \). This gives

\[
\Delta g_z = 2G\Delta \rho h \left[ \frac{\pi}{2} + \tan^{-1}\left(\frac{z_0}{h}\right) \right] \tag{2.101}
\]

Comparison of this expression with Eq. (2.98) shows that the anomaly of the vertical fault (or a semi-infinite thick horizontal slab) is the same as if the anomalous slab were replaced by a thin sheet of thickness \( h \) at the mid-point of the vertical step. Equation (2.101) is called the “thin-sheet approximation.” It is accurate to about 2% provided that \( z_0 > 2h \).
2.6.3.6 Iterative modelling

The simple geometric models used to compute the gravity anomalies in the previous sections are crude representations of the real anomalous bodies. Modern computer algorithms have radically changed modelling methods by facilitating the use of an iterative procedure. A starting model with an assumed geometry and density contrast is postulated for the anomalous body. The gravity anomaly of the body is then computed and compared with the residual anomaly. The parameters of the model are changed slightly and the computation is repeated until the discrepancies between the model anomaly and the residual anomaly are smaller than a determined value. However, as in the case of the simple models, this does not give a unique solution for the density distribution.

Two- and three-dimensional iterative techniques are in widespread use. The two-dimensional (2D) method assumes that the anomalous body is infinitely long parallel to the strike of the structure, but end corrections for the possibly limited horizontal extent of the body can be made. We can imagine that the cross-sectional shape of the body is replaced by countless thin rods or line elements aligned parallel to the strike. Each rod makes a contribution to the vertical component of gravity at the origin (Fig. 2.56a). The gravity anomaly of the structure is calculated by adding up the contributions of all the line elements; mathematically, this is an integration over the end surface of the body. Although the theory is beyond the scope of this chapter, the gravity anomaly has the following simple form:

\[ \Delta g_z = 2G \Delta \rho z \, d\theta \]  

(2.102)

The angle \( \theta \) is defined to lie between the positive \( x \)-axis and the radius from the origin to a line element (Fig. 2.56a), and the integration over the end-surface has been changed to an integration around its boundary. The computer algorithm for the calculation of this integral is greatly speeded up by replacing the true cross-sectional shape of the body with an \( N \)-sided polygon (Fig. 2.56a). Apart from the assumed density contrast, the only important parameters for the computation are the \( (x, z) \) coordinates of the corners of the polygon. The origin is now moved to the next point on a profile across the structure. This move changes only the \( x \)-coordinates of the corners of the polygon. The calculations are repeated for each successive point on the profile. Finally, the calculated gravity anomaly profile across the structure is compared to the observed anomaly and the residual differences are evaluated. The coordinates of the corners of the polygon are adjusted and the calculation is reiterated until the residuals are less than a selected tolerance level.

The gravity anomaly of a three-dimensional (3D) body is modelled in a similar way. Suppose that we have a contour map of the body; the contour lines show the smooth outline of the body at different depths. We could construct a close replica of the body by replacing the material between successive contour lines with thin laminae. Each lamina has the same outline as the contour line and has a thickness equal to the contour separation. As a further approximation the smooth outline of each lamina is replaced by a multi-sided polygon (Fig. 2.56b). The gravity anomaly of the polygon at the origin is computed as in the 2D case, using the \( (x, y) \) coordinates of the corners, the thickness of the lamina and an assumed density contrast. The gravity anomaly of the 3D body at the origin is found by adding up the contributions of all the laminae. As in the simpler 2D example, the origin is now displaced to a new point and the computation is repeated. The calculated and observed anomalies are now compared and the coordinates of the corners of the laminae are adjusted accordingly; the assumed density distribution can also be adjusted. The iterative procedure is repeated until the desired match between computed and observed anomalies is obtained.
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2.6.4 Some important regional gravity anomalies

Without auxiliary information the interpretation of gravity anomalies is ambiguous, because the same anomaly can be produced by different bodies. An independent data source is needed to restrict the choices of density contrast, size, shape and depth in the many possible gravity models. The additional information may be in the form of surface geological observations, from which the continuation of structures at depth is interpreted. Seismic refraction or reflection data provide better constraints.

The combination of seismic refraction experiments with precise gravity measurements has a long, successful history in the development of models of crustal structure. Refraction seismic profiles parallel to the trend of elongate geological structures give reliable information about the vertical velocity distribution. However, refraction profiles normal to the structural trend give uncertain information about the tilt of layers or lateral velocity changes. Lateral changes of crustal structure can be interpreted from several refraction profiles more or less parallel to the structural trend or from seismic reflection data. The refraction results give layer velocities and the depths to refracting interfaces. To compute the gravity effect of a structure the velocity distribution must first be converted to a density model using a P-wave velocity–density relationship like the curve shown in Fig. 2.40. The theoretical gravity anomaly over the structure is computed using a 2D or 3D method. Comparison with the observed gravity anomaly (e.g., by calculating the residual differences point by point) indicates the plausibility of the model. It is always important to keep in mind that, because of the non-uniqueness of gravity modelling, a plausible structure is not necessarily the true structure. Despite the ambiguities, some characteristic features of gravity anomalies have been established for important regions of the Earth.

2.6.4.1 Continental and oceanic gravity anomalies

In examining the shape of the Earth we saw that the ideal reference figure is a spheroid, or ellipsoid of rotation. It is assumed that the reference Earth is in hydrostatic equilibrium. This is supported by observations of free-air and isostatic anomalies which suggest that, except in some unusual locations such as deep-sea trenches and island arcs in subduction zones, the continents and oceans are in approximate isostatic equilibrium with each other. By applying the concepts of isostasy (Section 2.7) we can understand the large-scale differences between Bouguer gravity anomalies over the continents and those over the oceans. In general, Bouguer anomalies over the continents are negative, especially over mountain ranges where the crust is unusually thick; in contrast, strongly positive Bouguer anomalies are found over oceanic regions where the crust is very thin.

The inverse relationship between Bouguer anomaly amplitude and crustal thickness can be explained with the aid of a hypothetical example (Fig. 2.57). Continental crust that has not been thickened or thinned by tectonic processes is considered to be “normal” crust. It is typically 30–35 km thick. Under location A on an undeformed continental coastal region a thickness of 34 km is assumed. The theoretical gravity used in computing a gravity anomaly is defined on the reference ellipsoid, the surface of which corresponds to mean sea-level. Thus, at coastal location A on normally thick continental crust the Bouguer anomaly is close to zero. Isostatic compensation of the mountain range gives it a root-zone that increases the crustal thickness at location B. Isostatic evidence shows that continental crustal density increases with depth from about 2700 kg m\(^{-3}\) in the upper granitic crust to about 2900 kg m\(^{-3}\) in the lower gabbroic crust. Thus, the density in the root-zone is much lower than the typical mantle density of 3300–3400 kg m\(^{-3}\) at the same depth under A. The low-density root beneath B causes a negative Bouguer anomaly, which typically reaches −150 to −200 mGal.

At oceanic location C the vertical crustal structure is very different. Two effects contribute to the Bouguer anomaly. A 5 km thick layer of sea-water (density 1030 kg m\(^{-3}\)) overlies the thin basic oceanic crust (density 2900 kg m\(^{-3}\)) which has an average thickness of only about 6 km. To compute the Bouguer anomaly the seawater must be replaced by oceanic crustal rock. The attraction of the water layer is inherent in the measured gravity so the density used in correcting for the Bouguer plate and the topography of the ocean bottom is the reduced density of the oceanic crust (i.e., 2900 – 1030 = 1870 kg m\(^{-3}\)). However, a more important effect is that the top of the mantle is at a depth of only 11 km. In a vertical section below this depth the mantle has a density of 3300–3400 kg m\(^{-3}\), much higher than the density of the continental crust at equivalent depths below coastal site.

![Fig. 2.57 Hypothetical Bouguer anomalies over continental and oceanic areas. The regional Bouguer anomaly varies roughly inversely with crustal thickness and topographic elevation (after Robinson and Çoruh, 1988).](image-url)
A. The lower 23 km of the section beneath C represents a large excess of mass. This gives rise to a strong positive Bouguer anomaly, which can amount to 300–400 mgal.

2.6.4.2 Gravity anomalies across mountain chains

The typical gravity anomaly across a mountain chain is strongly negative due to the large low-density root-zone. The Swiss Alps provide a good example of the interpretation of such a gravity anomaly with the aid of seismic refraction and reflection results. A precise gravity survey of Switzerland carried out in the 1970s yielded an accurate Bouguer gravity map (Fig. 2.58). The map contains effects specific to the Alps. Most obviously, the contour lines are parallel to the trend of the mountain range. In the south a strong positive anomaly overrides the negative anomaly. This is the northern extension of the positive anomaly of the so-called Ivrea body, which is a high-density wedge of mantle material that was forced into an uplifted position within the western Alpine crust during an earlier continental collision. In addition, the Swiss gravity map contains the effects of low-density sediments that fill the Molasse basin north of the Alps, the Po plain to the south and the major Alpine valleys.

In the late 1980s a coordinated geological and geophysical study – the European Geotraverse (EGT) – was made along and adjacent to a narrow path stretching from northern Scandinavia to northern Africa. Detailed reflection seismic profiles along its transect through the Central Swiss Alps complemented a large amount of new and extant refraction data. The seismic results gave the depths to important interfaces. Assuming a velocity–density relationship, a model of the density distribution in the lithosphere under the traverse was obtained. Making appropriate

---

**Fig. 2.58** Bouguer gravity map of Switzerland (after Klingelé and Olivier, 1980).

**Fig. 2.59** Lithosphere density model for the Central Swiss Alps along the European Geotraverse transect, compiled from seismic refraction and reflection profiles. The 2.5D gravity anomaly calculated for this lithospheric structure is compared to the observed Bouguer anomaly after removal of the effects of the high-density Ivrea body and the low-density sediments in the Molasse basin, Po plain and larger Alpine valleys (after Holliger and Kissling, 1992).
corrections for end effects due to limited extent along strike, a 2.5D gravity anomaly was calculated for this lithospheric structure (Fig. 2.59). After removal of the effects of the high-density Ivrea body and the low-density sediments, the corrected Bouguer gravity profile is reproduced well by the anomaly of the lithospheric model. Using the geometric constraints provided by the seismic data, the lithospheric gravity model favors a subduction zone, dipping gently to the south, with a high-density wedge of deformed lower crustal rock (“mélange”) in the middle crust. As already noted, the fact that a density model delivers a gravity anomaly that agrees well with observation does not establish the reality of the interpreted structure, which can only be confirmed by further seismic imaging. However, the gravity model provides an important check on the reasonableness of suggested models. A model of crustal or lithospheric structure that does not give an appropriate gravity anomaly can reasonably be excluded.

2.6.4.3 Gravity anomalies across an oceanic ridge

An oceanic ridge system is a gigantic submarine mountain range. The difference in depth between the ridge crest and adjacent ocean basin is about 3 km. The ridge system extends laterally for several hundred kilometers on either side of the axis. Gravity and seismic surveys have been carried out across several oceanic ridge systems. Some common characteristics of continuous gravity profiles across a ridge are evident on a WNW–ESE transect crossing the Mid-Atlantic Ridge at 32°N (Fig. 2.60). The free-air gravity anomalies are small, around 50 mgal or less, and correlate closely with the variations in ocean-bottom topography. This indicates that the ridge and its flanks are nearly compensated isostatically. As expected for an oceanic profile, the Bouguer anomaly is strongly positive. It is greater than 350 mgal at distances beyond 1000 km from the ridge, but decreases to less than 200 mgal over the axis of the ridge.
The depths to important refracting interfaces and the P-wave layer velocities are known from seismic refraction profiles parallel to the ridge. The seismic structure away from the ridge is layered, with P-wave velocities of 4–5 km s\(^{-1}\) for the basalts and gabbros in oceanic Layer 2 and 6.5–6.8 km s\(^{-1}\) for the meta-basalts and meta-gabbros in Layer 3; the Moho is at about 11 km depth, below which typical upper-mantle velocities of 8–8.4 km s\(^{-1}\) are found.

However, the layered structure breaks down under the ridge at distances less than 400 km from its axial zone. Unusual velocities around 7.3 km s\(^{-1}\) occurred at several places, suggesting the presence of anomalous low-density mantle material at comparatively shallow depth beneath the ridge. The seismic structure was converted to a density model using a velocity–density relationship. Assuming a 2D structure, several density models were found that closely reproduced the Bouguer anomaly. However, to satisfy the Bouguer anomaly on the ridge flanks each model requires a flat body in the upper mantle beneath the ridge; it extends down to about 30 km depth and for nearly 1000 km on each side of the axis (Fig. 2.60).

The density of the anomalous structure is only 3150 kg m\(^{-3}\) instead of the usual 3400 kg m\(^{-3}\). The model was proposed before the theory of plate tectonics was accepted. The anomalous upper mantle structure satisfies the gravity anomaly but has no relation to the known physical structure of a constructive plate margin. A broad zone of low upper-mantle seismic velocities was not found in later experiments, but narrow low-velocity zones are sometimes present close to the ridge axis.

A further combined seismic and gravity study of the Mid-Atlantic Ridge near 46°N gave a contradictory density model. Seismic refraction results yielded P-wave velocities of 4.6 km s\(^{-1}\) and 6.6 km s\(^{-1}\) for Layer 2 and Layer 3, respectively, but did not show anomalous mantle velocities beneath the ridge except under the median valley. A simpler density model was postulated to account for the gravity anomaly (Fig. 2.61). The small-scale free-air anomalies are accounted for by the variations in ridge topography seen in the bathymetric plot. The large-scale free-air gravity anomaly is reproduced well by a wedge-shaped structure that extends to 200 km depth. Its base extends to hundreds of kilometers on each side of the axis. A very small density contrast of only \(-40\) kg m\(^{-3}\) suffices to explain the broad free-air anomaly. The model is compatible with the thermal structure of an accreting plate margin. The low-density zone may be associated with hot material from the asthenosphere, which rises...
beneath the ridge, melts and accumulates in a shallow magma chamber within the oceanic crust.

### 2.6.4.4 Gravity anomalies at subduction zones

Subduction zones are found primarily at continental margins and island arcs. Elongate, narrow and intense isostatic and free-air gravity anomalies have long been associated with island arcs. The relationship of gravity to the structure of a subduction zone is illustrated by the free-air anomaly across the Chile trench at 23°S (Fig. 2.62). Seismic refraction data define the thicknesses of the oceanic and continental crust. Thermal and petrological data are integrated to give a density model for the structure of the mantle and the subducting lithosphere.

The continental crust is about 65 km thick beneath the Andes mountains, and gives large negative Bouguer anomalies. The free-air gravity anomaly over the Andes is positive, averaging about +50 mgal over the 4 km high plateau. Even stronger anomalies up to +100 mgal are seen over the east and west boundaries of the Andes. This is largely due to the edge effect of the low-density Andean crustal block (see Fig. 2.44b and Section 2.5.6).

A strong positive free-air anomaly of about +70 mgal lies between the Andes and the shore-line of the Pacific ocean. This anomaly is due to the subduction of the Nazca plate beneath South America. The descending slab is old and cool. Subduction exposes it to higher temperatures and pressures, but the slab descends faster than it can be heated up. The increase in density accompanying greater depth and pressure outweighs the decrease in density due to hotter temperatures. There is a positive density contrast between the subducting lithosphere and the surrounding mantle. Also, petrological changes accompanying the subduction result in mass excesses. Peridotite in the upper lithosphere changes phase from plagioclase-type to the higher-density garnet-type. When oceanic crust is subducted to depths of 30–80 km, basalt changes phase to eclogite, which has a higher density (3560–3580 kg m⁻³) than upper mantle rocks. These effects combine to produce the positive free-air anomaly.

The Chile trench is more than 2.5 km deeper than the ocean basin to the west. The sediments flooring the trench have low density. The mass deficiency of the water and sediments in the trench cause a strong negative free-air anomaly, which parallels the trench and has an amplitude greater than −250 mgal. A small positive anomaly of about +20 mgal is present about 100 km seaward of the trench axis. This anomaly is evident also in the mean level of the ocean surface as mapped by SEASAT (Fig. 2.28), which shows that the mean sea surface is raised in front of deep ocean trenches. This is due to upward flexure of the lithosphere before its downward plunge into the subduction zone. The flexure elevates higher-density mantle rocks and thereby causes the small positive free-air anomaly.

### 2.7 ISOSTASY

#### 2.7.1 The discovery of isostasy

Newton formulated the law of universal gravitation in 1687 and confirmed it with Kepler’s laws of planetary motion. However, in the seventeenth and eighteenth centuries the law could not be used to calculate the mass or mean density of the Earth, because the value of the gravitational constant was not yet known (it was first determined by Cavendish in 1798). Meanwhile, eighteenth century scientists attempted to estimate the mean density of the Earth
Gravity, the figure of the Earth and geodynamics

by various means. They involved comparing the attraction of the Earth with that of a suitable mountain, which could be calculated. Inconsistent results were obtained.

During the French expedition to Peru in 1737–1740, Pierre Bouguer measured gravity with a pendulum at different altitudes, applying the elevation correction term which now bears his name. If the density of crustal rocks is \( \rho \) and the mean density of the Earth is \( \rho_0 \), the ratio of the Bouguer-plate correction (see Section 2.5.4.3) for elevation \( h \) to mean gravity for a spherical Earth of radius \( R \) is

\[
\frac{\Delta g_{BP}}{g} = \frac{2\pi G\rho h}{3\pi G\rho_0 R} = \frac{2}{3} \left( \frac{\rho h}{\rho_0 R} \right)
\]

(2.103)

From the results he obtained near Quito, Bouguer estimated that the mean density of the Earth was about 4.5 times the density of crustal rocks.

The main method employed by Bouguer to determine the Earth’s mean density consisted of measuring the deflection of the plumb-line (vertical direction) by the mass of a nearby mountain (Fig. 2.63). Suppose the elevation of a known star is measured relative to the local vertical direction at points N and S on the same meridian. The elevations should be \( \alpha_N \) and \( \alpha_S \), respectively. Their sum is \( \alpha \), the angle subtended at the center of the Earth by the radii to N and S, which corresponds to the difference in latitude. If N and S lie on opposite sides of a large mountain, the plumb-line at each station is deflected by the attraction of the mountain. The measured elevations of the star are \( \beta_N \) and \( \beta_S \), respectively, and their sum is \( \beta \). The local vertical directions now intersect at the point D instead of at the center of the (assumed spherical) Earth. The difference \( \delta = \beta - \alpha \) is the sum of the deviations of the vertical direction caused by the mass of the mountain.

The horizontal attraction \( f \) of the mountain can be calculated from its shape and density, with a method that resembles the computation of the topographic correction in the reduction of gravity measurements. Dividing the mountain into vertical cylindrical elements, the horizontal attraction of each element is calculated and its component \( (G\rho h) \) towards the center of mass of the mountain is found. Summing up the effects of all the cylindrical elements in the mountain gives the horizontal attraction \( f \) towards its center of mass. Comparing \( f \) with mean gravity \( g \), we then write

\[
\tan \delta = \frac{f}{g} = \frac{G \sum h_i}{\frac{2}{3} \pi G\rho_0 R} = \frac{\sum h_i (\rho/\rho_0)}{\frac{2}{3} \pi R}
\]

(2.104)

For very small angles, \( \tan \delta \) is equal to \( \delta \), and so the deflection of the vertical is proportional to the ratio \( \rho/\rho_0 \) of the mean densities of the mountain and the Earth. Bouguer measured the deflection of the vertical caused by Mt. Chimborazo (6272 m), Ecuador’s highest mountain. His results gave a ratio \( \rho/\rho_0 \) of around 12, which is unrealistically large and quite different from the values he had obtained near Quito. The erroneous result indicated that the deflection of the vertical caused by the mountain was much too small for its estimated mass.

In 1774 Bouguer’s Chimborazo experiment was repeated in Scotland by Neville Maskelyne on behalf of the Royal Society of London. Measurements of the elevations of stars were made on the north and south flanks of Mt. Schiehallion at sites that differed in latitude by 42.9° of arc. The observed angle between the plumb-lines was 54.6°. The analysis gave a ratio \( \rho/\rho_0 \) equal to 1.79, suggesting a mean density for the Earth of 4500 kg m\(^{-3}\). This was more realistic than Bouguer’s result, which still needed explanation.

Further results accumulated in the first half of the nineteenth century. From 1806 to 1843 the English geodesist George Everest carried out triangulation surveys in India. He measured by triangulation the separation of a site at Kalianpur on the Indo-Ganges plain from a site at Kalian in the foothills of the Himalayas. The distance differed substantially from the separation of the sites
computed from the elevations of stars, as in Fig. 2.63. The discrepancy of 5.23° of arc (162 m) was attributed to
deflection of the plumb-line by the mass of the
Himalayas. This would affect the astronomic
determination but not the triangulation measurement. In 1855 I. H.
Pratt computed the minimum deflection of the plumb-
line that might be caused by the mass of the Himalayas
and found that it should be 15.89° of arc, about three
times larger than the observed deflection. Evidently the
attraction of the mountain range on the plumb-line was
not as large as it should be.

The anomalous deflections of the vertical were first
understood in the middle of the nineteenth century, when
it was realized that there are regions beneath mountains —
"root-zones" — in which rocks have a lower density than
expected. The deflection of a plumb-line is not caused
only by the horizontal attraction of the visible part of a
mountain. The deficiency of mass at depth beneath the
mountain means that the "hidden part" exerts a reduced
lateral attraction, which partly offsets the effect of the
mountain and diminishes the deflection of the vertical. In
1889 C. E. Dutton referred to the compensation of a
topographic load by a less-dense subsurface structure as
isostasy.

2.7.2 Models of isostasy

Separate explanations of the anomalous plumb-line
deflections were put forward by G. B. Airy in 1855 and
J. H. Pratt in 1859. Airy was the Astronomer Royal and
director of the Greenwich Observatory. Pratt was an
archdeacon of the Anglican church at Calcutta, India,
and a devoted scientist. Their hypotheses have in common
the compensation of the extra mass of a mountain
above sea-level by a less-dense region (or root) below
sea-level, but they differ in the way the compensation is
achieved. In the Airy model, when isostatic compensation
is complete, the mass deficiency of the root equals
the excess load on the surface. At and below a certain
compensation depth the pressure exerted by all overlying
vertical columns of crustal material is then equal. The
pressure is then hydrostatic, as if the interior acted like a
fluid. Hence, isostatic compensation is equivalent to
applying Archimedes’ principle to the uppermost layers of
the Earth.

The Pratt and Airy models achieve compensation
locally by equalization of the pressure below vertical
columns under a topographic load. The models were very
successful and became widely used by geodesists, who
developed them further. In 1909–1910, J. F. Hayford in the
United States derived a mathematical model to describe
the Pratt hypothesis. As a result, this theory of isostasy is
often called the Pratt–Hayford scheme of compensation.
Between 1924 and 1938 W. A. Heiskanen derived sets of
tables for calculating isostatic corrections based on the
Airy model. This concept of isostatic compensation has
since been referred to as the Airy–Heiskanen scheme.

Fig. 2.64 Local isostatic compensation according to (a) the
Airy–Heiskanen model and (b) the Pratt–Hayford model; (c) regional
compensation according to the elastic plate model of Vening Meinesz.

It became apparent that both models had serious deficien-
cies in situations that required compensation over a
larger region. In 1931 F. A. Vening Meinesz, a Dutch geo-
physicist, proposed a third model, in which the crust acts
as an elastic plate. As in the other models, the crust floats
buoyantly on a substratum, but its inherent rigidity
spreads topographic loads over a broader region.

2.7.2.1 The Airy–Heiskanen model

According to the Airy–Heiskanen model of isostatic
compensation (Fig. 2.64a) an upper layer of the Earth
“floats” on a denser magma-like substratum, just as ice-
bergs float in water. The upper layer is equated with
the crust and the substratum with the mantle. The height of a
mountain above sea-level is much less than the thickness of the
crust underneath it, just as the visible tip of an
iceberg is much smaller than the subsurface part. The
densities of the crust and mantle are assumed to be con-
stant; the thickness of the root-zone varies in proportion
to the elevation of the topography.

The analogy to an iceberg is not exact, because
underland at sea-level the “normal” crust is already
about 30–35 km thick; the compensating root-zone of a
mountain lies below this depth. Oceanic crust is only about 10 km thick, thinner than the “normal” crust. The mantle between the base of the oceanic crust and the normal crustal depth is sometimes called the anti-root of the ocean basin.

The Airy–Heiskanen model assumes local isostatic compensation, i.e., the root-zone of a mountain lies directly under it. Isostasy is assumed to be complete, so that hydrostatic equilibrium exists at the compensation depth, which is equivalent to the base of the deepest mountain root. The pressure at this level is due to the weight of the rock material in the overlying vertical column (of basal area one square meter) extending to the Earth’s surface. The vertical column for the mountain of height \( h \) in Fig. 2.64a contains only crustal rocks of density \( \rho_c \). The pressure at \( CC' \) due to the mountain, “normal” crust of thickness \( t \), and a root-zone of thickness \( r \), amounts to \( (h + r + t) \rho_c \). The vertical column below the “normal” crust contains a thickness \( t \) of crustal rocks and thickness \( r \) of mantle rocks; it exerts a pressure of \( (r \rho_m + t \rho_c) \). For hydrostatic equilibrium the pressures are equal. Equating, and noting that each expression contains the term \( t \rho_c \), we get

\[
r_1 = \frac{\rho_m - \rho_c}{\rho_m - \rho_c} D_1
\]

with a similar expression for the root of depth \( r_2 \) under the hill of height \( h \). The thickness \( r_0 \) of the anti-root of the oceanic crust under an ocean basin of water depth \( d \) and density \( \rho_w \) is given by

\[
r_0 = \frac{\rho_c - \rho_w}{\rho_m - \rho_c} D
\]

The Airy–Heiskanen model assumes an upper layer of constant density floating on a more dense substratum. It has root-zones of variable thickness proportional to the overlying topography. This scenario agrees broadly with seismic evidence for the thickness of the Earth’s crust (see Section 3.7). The continental crust is much thicker than the oceanic crust. Its thickness is very variable, being largest below mountain chains, although the greatest thickness is not always under the highest topography. Airy-type compensation suggests hydrostatic balance between the crust and the mantle.

2.7.2.2 The Pratt–Hayford model

The Pratt–Hayford isostatic model incorporates an outer layer of the Earth that rests on a weak magmatic substratum. Differential expansion of the material in vertical columns of the outer layer accounts for the surface topography, so that the higher the column above a common base the lower the mean density of rocks in it. The vertical columns have constant density from the surface to their base at depth \( B \) below sea-level (Fig. 2.64b). If the rock beneath a mountain of height \( h \) \((i = 1, 2, \ldots)\) has density \( \rho_r \) the pressure at \( CC' \) is \( \rho_r (h_i + D) \).

Beneath a continental region at sea-level the pressure of the rock column of density \( \rho_r \) is \( \rho_r D \). Under an ocean basin the pressure at \( CC' \) is due to water of depth \( d \) and density \( \rho_w \) on top of a rock column of thickness \( (D - d) \) and density \( \rho_c \): it is equal to \( \rho_w d + \rho_c (D - d) \). Equating these pressures, we get

\[
\rho_c = \frac{D}{h_0 + D} \rho_c
\]

for the density below a topographic elevation \( h_0 \), and

\[
\rho_0 = \frac{\rho_c D - \rho_w d}{D - d}
\]

for the density under an oceanic basin of depth \( d \). The compensation depth \( D \) is about 100 km.

The Pratt–Hayford and Airy–Heiskanen models represent local isostatic compensation, in which each column exerts an equal pressure at the compensation level. At the time these models were proposed very little was yet known about the internal structure of the Earth. This was only deciphered after the development of seismology in the late nineteenth and early twentieth century. Each model is idealized, both with regard to the density distributions and the behavior of Earth materials. For example, the upper layer is assumed to offer no resistance to shear stresses arising from vertical adjustments between adjacent columns. Yet the layer has sufficient strength to resist stresses due to horizontal differences in density. It is implausible that small topographic features require compensation at large depths; more likely, they are entirely supported by the strength of the Earth’s crust.

2.7.2.3 Vening Meinesz elastic plate model

In the 1920s F. A. Vening Meinesz made extensive gravity surveys at sea. His measurements were made in a submarine to avoid the disturbances of wave motions. He studied the relationship between topography and gravity anomalies over prominent topographic features, such as the deep sea trenches and island arcs in southeastern Asia, and concluded that isostatic compensation is often not entirely local. In 1931 he proposed a model of regional isostatic compensation which, like the Pratt–Hayford and Airy–Heiskanen models, envisages a light upper layer that floats on a denser fluid substratum. However, in the Vening Meinesz model the upper layer behaves like an elastic plate overlaying a weak fluid. The strength of the plate distributes the load of a surface feature (e.g., an island or seamount) over a horizontal distance wider than the feature (Fig. 2.64c). The topographic load bends the plate downward into the fluid substratum, which is pushed aside. The buoyancy of the displaced fluid forces it upward, giving support to the bent plate at distances well away from the central depression. The bending of the plate which accounts for the regional compensation in the Vening Meinesz model depends on the elastic properties of the lithosphere.


2.7 ISOSTASY

(a) complete

(b) overcompensation

(c) undercompensation

Fig. 2.65 Explanation of the isostatic gravity anomaly ($\Delta g$) as the difference between the Bouguer gravity anomaly ($\Delta g_B$) and the computed anomaly ($\Delta g_R$) of the root-zone estimated from the topography for (a) complete isostatic compensation, (b) isostatic overcompensation and (c) isostatic undercompensation.

2.7.3 Isostatic compensation and vertical crustal movements

In the Pratt–Hayford and Airy–Heiskanen models the lighter crust floats freely on the denser mantle. The system is in hydrostatic equilibrium, and local isostatic compensation is a simple application of Archimedes' principle. A “normal” crustal thickness for sea-level coastal regions is assumed (usually 30–35 km) and the additional depths of the root-zones below this level are exactly proportional to the elevations of the topography above sea-level. The topography is then completely compensated (Fig. 2.65a). However, isostatic compensation is often incomplete. The geodynamic imbalance leads to vertical crustal movements. Mountains are subject to erosion, which can disturb isostatic compensation. If the eroded mountains are no longer high enough to justify their deep root-zones, the topography is isostatically overcompensated (Fig. 2.65b). Buoyancy forces are created, just as when a wooden block floating in water is pressed downward by a finger: the underwater part becomes too large in proportion to the amount above the surface. If the finger pressure is removed, the block rebounds in order to restore hydrostatic equilibrium. Similarly, the buoyancy forces that result from overcompensation of mountainous topography cause vertical uplift.

The opposite scenario is also possible. When the visible topography has roots that are too small, the topography is isostatically undercompensated (Fig. 2.65c). This situation can result, for example, when tectonic forces thrust crustal blocks on top of each other. Hydrostatic equilibrium is now achieved by subsidence of the uplifted region.

The most striking and best-observed examples of vertical crustal movements due to isostatic imbalance are related to the phenomenon of glacial rebound observed in northern Canada and in Fennoscandia. During the last ice-age these regions were covered by a thick ice-cap. The weight of ice depressed the underlying crust. Subsequently melting of the ice-cap removed the extra load on the crust, and it has since been rebounding. At stations on the Fennoscandian shield, modern tide-gauge observations and precision levelling surveys made years apart allow the present uplift rates to be calculated (Fig. 2.66). The contour lines of equal uplift rate are inexact over large areas due to the incompleteness of data from inaccessible regions. Nevertheless, the general pattern of glacial rebound is clearly recognizable, with uplift rates of up to 8 mm yr$^{-1}$.

2.7.4 Isostatic gravity anomalies

The different degrees of isostatic compensation find expression in gravity anomalies. As explained in Section 2.5.6 the free-air gravity anomaly $\Delta g_f$ is small near the center of a large region that is isostatically compensated; the Bouguer anomaly $\Delta g_B$ is strongly negative. Assuming complete isostatic compensation, the size and shape of the root-zone
can be determined from the elevations of the topography. With a suitable density contrast the gravity anomaly $\Delta g_B$ of the modelled root-zone can be calculated; because the root-zone has lower density than adjacent mantle rocks $\Delta g_B$ is also negative. The isostatic gravity anomaly $\Delta g_I$ is defined as the difference between the Bouguer gravity anomaly and the computed anomaly of the root-zone, i.e.,

$$\Delta g_I = \Delta g_B - \Delta g_R \quad (2.109)$$

Examples of the isostatic gravity anomaly for the three types of isostatic compensation are shown schematically in Fig. 2.65. When isostatic compensation is complete, the topography is in hydrostatic equilibrium with its root-zone. Both $\Delta g_B$ and $\Delta g_R$ are negative but equal; consequently, the isostatic anomaly is everywhere zero ($\Delta g_I = 0$). In the case of overcompensation the eroded topography suggests a root-zone that is smaller than the real root-zone. The Bouguer anomaly is caused by the larger real root, so $\Delta g_B$ is numerically larger than $\Delta g_R$. Subtracting the smaller negative anomaly of the computed root-zone leaves a negative isostatic anomaly ($\Delta g_I < 0$). On the other hand, with undercompensation the topography suggests a root-zone that is larger than the real root-zone. The Bouguer anomaly is caused by the smaller real root, so $\Delta g_B$ is numerically smaller than $\Delta g_R$. Subtracting the larger negative anomaly of the root-zone leaves a positive isostatic anomaly ($\Delta g_I > 0$).

A national gravity survey of Switzerland carried out in the 1970s gave a high-quality map of Bouguer gravity anomalies (see Fig. 2.58). Seismic data gave representative parameters for the Central European crust and mantle: a crustal thickness of 32 km without topography, and mean densities of 2670 kg m$^{-3}$ for the topography, 2810 kg m$^{-3}$ for the crust and 3310 kg m$^{-3}$ for the mantle. Using the Airy–Heiskanen model of compensation, a map of isostatic gravity anomalies in Switzerland was derived (Fig. 2.67) after correcting the gravity map for the effects of low-density sediments in the Molasse basin north of the Alps and high-density material in the anomalous Ivrea body in the south.

The pattern of isostatic anomalies reflects the different structures beneath the Jura mountains, which do not have a prominent root-zone, and the Alps, which have a low-density root that extends to more than 55 km depth in places. The dominant ENE–WSW trend of the isostatic gravity anomaly contour lines is roughly parallel to the trends of the mountain chains. In the northwest, near the Jura mountains, positive isostatic anomalies exceed 20 mgal. In the Alps isostatic anomalies are mainly negative, reaching more than $\pm$30 mgal in the east.

A computation based on the Vening Meinesz model gave an almost identical isostatic anomaly map. The agreement of maps based on the different concepts of isostasy is somewhat surprising. It may imply that vertical crustal columns are not free to adjust relative to one another without friction as assumed in the Airy–Heiskanen model. The friction is thought to result from horizontal compressive stresses in the Alps, which are active in the on-going mountain-building process.

Comparison of the isostatic anomaly map with one of recent vertical crustal movements (Fig. 2.68) illustrates the relevance of isostatic gravity anomalies for tectonic interpretation. Precise levelling surveys have been carried out since the early 1900s along major valleys transecting and parallel to the mountainous topography of Switzerland. Relative rates of uplift or subsidence are computed from the differences between repeated surveys. The results have not been tied to absolute tide-gauge observations and so are relative to a base station at Aarburg in the canton of Aargau, in the northeast.
The rates of relative vertical movement in northeastern Switzerland are smaller than the confidence limits on the data and may not be significant, but the general tendency suggests subsidence. This region is characterized by mainly positive isostatic anomalies. The rates of vertical movement in the southern part of Switzerland exceed the noise level of the measurements and are significant. The most notable characteristic of the recent crustal motions is vertical uplift of the Alpine part of Switzerland relative to the central plateau and Jura mountains. The Alpine uplift rates are up to 1.5 mm yr\(^{-1}\), considerably smaller than the rates observed in Fennoscandia. The most rapid uplift rates are observed in the region where isostatic anomalies are negative. The constant erosion of the mountain topography relieves the crustal load and the isostatic response is uplift. However, the interpretation is complicated by the fact that compressive stresses throughout the Alpine region acting on deep-reaching faults can produce non-isostatic uplift of the surface. The separation of isostatic and non-isostatic vertical crustal movements in the Alps will require detailed and exact information about the structure of the lithosphere and asthenosphere in this region.

2.8 RHEOLOGY

2.8.1 Brittle and ductile deformation

Rheology is the science of the deformation and flow of solid materials. This definition appears at first sight to contradict itself. A solid is made up of particles that cohere to each other; it is rigid and resists a change of shape. A fluid has no rigidity; its particles can move about comparatively freely. So how can a solid flow? In fact, the way in which a solid reacts to stress depends on how large the stress is and the length of time for which it is applied. Provided the applied stress does not exceed the yield stress (or elastic limit) the short-term behavior is elastic. This means that any deformation caused by the stress is completely recoverable when the stress is removed, leaving no permanent change in shape. However, if the applied stress exceeds the yield stress, the solid may experience either brittle or ductile deformation.

Brittle deformation consists of rupture without other distortion. This is an abrupt process that causes faulting in rocks and earthquakes, accompanied by the release of elastic energy in the form of seismic waves. Brittle fracture occurs at much lower stresses than the intrinsic strength of a crystal lattice. This is attributed to the presence of cracks, which modify the local internal stress field in the crystal. Fracture occurs under either extension or shear. Extensional fracture occurs on a plane at right angles to the direction of maximum tension. Shear fracture occurs under compression on one of two complementary planes which, reflecting the influence of internal friction, are inclined at an angle of less than 45° (typically about 30°) to the maximum principal compression. Brittle deformation is the main mechanism in tectonic processes that involve the uppermost 5–10 km of the lithosphere.

Ductile deformation is a slow process in which a solid acquires strain (i.e., it changes shape) over a long period of time. A material may react differently to a stress that is applied briefly than to a stress of long duration. If it experiences a large stress for a long period of time a solid can slowly and permanently change shape. The time-dependent deformation is called plastic flow and the capacity of the solid to flow is called its ductility. The ductility of a solid above its yield stress depends on temperature and confining pressure, and materials that are brittle under ordinary conditions may be ductile at high temperature and pressure. The behavior of rocks and minerals in
The deep interior of the Earth is characterized by ductile deformation. The transition from brittle to ductile types of deformation is thought to occur differently in oceanic and continental lithosphere (Fig. 2.69). The depth of the transition depends on several parameters, including the composition of the rocks, the local geothermal gradient, initial crustal thickness and the strain rate. Consequently it is sensitive to the vertically layered structure of the lithosphere. The oceanic lithosphere has a thin crust and shows a gradual increase in strength with depth, reaching a maximum in the upper mantle. In the upper mantle the strength increases again, leading to a second brittle–ductile transition at about 100 km depth. The continental crust is much thicker than the oceanic crust and has a more complex layering. The upper crust is brittle, but the minerals of the lower crust are weakened by high temperature. As a result the lower crust becomes ductile near to the Moho at about 30–35 km depth. In the upper mantle the strength increases again, leading to a second brittle–ductile transition at about 40–50 km depth. The difference in rheological layering of continental and oceanic lithosphere is important in collisions between plates. The crustal part of the continental lithosphere may detach from the mantle. The folding, underthrusting and stacking of crustal layers produce folded mountain ranges in the suture zone and thickening of the continental crust. For example, great crustal thicknesses under the Himalayas are attributed to underthrusting of crust from the Indian plate beneath the crust of the Eurasian plate.

2.8.2 Viscous flow in liquids

Consider the case when a liquid or gas flows in thin layers parallel to a flat surface (Fig. 2.70). The laminar flow exists as long as the speed stays below a critical value, above which the flow becomes turbulent. Turbulent flow does not interest us here, because the rates of flow in solid earth materials are very slow.

Suppose that the velocity of laminar flow along the horizontal x-direction increases with vertical height z above the reference surface. The molecules of the fluid may be regarded as having two components of velocity. One component is the velocity of flow in the x-direction, but in addition there is a random component with a variable velocity whose root-mean-square value is determined by the temperature (Section 4.2.2). Because of the random component, one-sixth of the molecules in a unit volume are moving upward and one-sixth downward on average at any time. This causes a transfer of molecules between adjacent layers in the laminar flow. The number of transfers per second depends on the size of the random velocity component, i.e., on temperature. The influx of molecules from the slower-moving layer reduces the momentum of the faster-moving layer. In turn, molecules transferred downward from the faster-velocity layer increase the momentum of the slower-velocity layer. This means that the two layers do not move freely past each other. They exert a shear force – or drag – on each other and the fluid is said to be viscous.

The magnitude of the shear force $F_{xz}$ depends on how much momentum is transferred from one layer to the next. If all the molecules in a fluid have the same mass, the momentum transfer is determined by the change in the velocity $v_z$ between the layers; this depends on the vertical gradient of the flow velocity ($\frac{dv_z}{dz}$). The momentum exchange depends also on the number of molecules that cross the boundary between adjacent layers and so is proportional to the surface area, $A$. We can bring these observations together, as did Newton in the seventeenth century, and derive the following proportionality relationship for $F_{xz}$:

$$F_{xz} \propto A \frac{dv_z}{dz}$$ (2.110)

If we divide both sides by the area $A$, the left side becomes the shear stress $\sigma_{xz}$. Introducing a proportionality constant $\eta$ we get the equation

$$\sigma_{xz} = \frac{dv_z}{dz}$$ (2.111)
This equation is Newton’s law of viscous flow and \( \eta \) is the coefficient of viscosity. If \( \eta \) is constant, the fluid is called a Newtonian fluid. The value of \( \eta \) depends on the transfer rate of molecules between layers and so on temperature. Substituting the units of stress (pascal) and velocity gradient \( (\text{ms}^{-1}/\text{m} = \text{s}^{-1}) \) we find that the unit of \( \eta \) is a pascal-second (Pa s). A shear stress applied to a fluid with low viscosity causes a large velocity gradient; the fluid flows easily. This is the case in a gas (\( \eta \) in air is of the order \( 2 \times 10^{-5} \text{ Pa s} \)) or in a liquid (\( \eta \) in water is \( 1.005 \times 10^{-3} \text{ Pa s} \) at 20 °C). The same shear stress applied to a very viscous fluid (with a large value of \( \eta \)) produces only a small velocity gradient transverse to the flow direction. The layers in the laminar flow are reluctant to move past each other. The viscous fluid is “sticky” and it resists flow. For example, \( \eta \) in a viscous liquid like engine oil is around 0.1–10 Pa s, three or four orders of magnitude higher than in water.

### 2.8 RHEOLOGY

**2.8.3 Flow in solids**

The response of a solid to an applied load depends upon whether the stress exceeds the elastic limit (Section 3.2.1) and for how long it is applied. When the yield stress (elastic limit) is reached, a solid may deform continuously without further increase in stress. This is called plastic deformation. In perfectly plastic behavior the stress–strain curve has zero slope, but the stress–strain curves of plastically deformed materials usually have a small positive slope (see Fig. 3.2a). This means that the stress must be increased above the yield stress for plastic deformation to advance. This effect is called strain-hardening. When the stress is removed after a material has been strain-hardened, a permanent residual strain is left.

Consider the effects that ensue if a stress is suddenly applied to a material at time \( t_0 \), held constant until time \( t_1 \) and then abruptly removed (Fig. 2.71a). As long as the applied stress is lower than the yield stress, the solid deforms elastically. The elastic strain is acquired immediately and remains constant as long as the stress is applied. Upon removal of the stress, the object at once recovers its original shape and there is no permanent strain (Fig. 2.71b).

If a constant load greater than the yield stress is applied, the resulting strain consists of a constant elastic strain and a changing plastic strain, which increases with time. After removal of the load at \( t_1 \) the plastic deformation does not disappear but leaves a permanent strain (Fig. 2.71c). In some plastic materials the deformation increases slowly at a decreasing rate, eventually reaching a limiting value for any specific value of the stress. This is called viscoelastic deformation (Fig. 2.71d). When the load is removed at \( t_1 \), the elastic part of the deformation is at once restored, followed by a slow decrease of the residual strain. This phase is called recovery or delayed elasticity. Viscoelastic behavior is an important rheological process deep in the Earth, for example in the asthenosphere and deeper mantle.

![Fig. 2.71](image)

The analogy to the viscosity of liquids is apparent by inspection of Eq. (2.111). Putting \( \text{d}x/\text{d}t \) and changing the order of differentiation, the equation becomes

\[
\sigma_{xz} = \eta \frac{\text{d} \varepsilon}{\text{d} t} = \eta \frac{\text{d} \varepsilon}{\text{d} x} \frac{\text{d} x}{\text{d} t} = \eta \frac{\text{d} \varepsilon}{\text{d} x}
\]

(2.112)

This equation resembles the elastic equation for shear deformation, which relates stress and strain through the shear modulus (see Eq. (3.16)). However, in the case of the “viscous flow” of a solid the shear stress depends on the strain rate. The parameter \( \eta \) for a solid is called the viscosity modulus, or dynamic viscosity. It is analogous to the viscosity coefficient of a liquid but its value in a solid is many orders of magnitude larger. For example, the viscosity of the asthenosphere is estimated to be of the order of \( 10^{20}–10^{21} \) Pa s. Plastic flow in solids differs from true flow in that it only occurs when the stress exceeds the yield stress of the solid. Below this stress the solid does not flow. However, internal defects in a metal or crystal can be mobilized and reorganized by stresses well below the yield stress. As a result the solid may change shape over a long period of time.

#### 2.8.3.1 Viscoelastic model

Scientists have tried to understand the behavior of rocks under stress by devising models based on mechanical analogs. In 1890 Lord Kelvin modelled viscoelastic deformation by combining the characteristics of a perfectly elastic solid and a viscous liquid. An applied stress causes both elastic and viscous effects. If the elastic strain is \( \varepsilon \), the corresponding elastic part of the stress is \( E \varepsilon \), where \( E \) is Young’s modulus. Similarly, if the rate of change of strain with time is \( d\varepsilon/dt \), the viscous part of the applied
stress is $\eta \frac{d\epsilon}{dt}$, where $\eta$ is the viscosity modulus. The applied stress $\sigma$ is the sum of the two parts and can be written

$$\sigma = E\epsilon + \eta \frac{d\epsilon}{dt} \quad (2.113)$$

To solve this equation we first divide throughout by $E$, then define the retardation time $\tau = \eta/E$, which is a measure of how long it takes for viscous strains to exceed elastic strains. Substituting and rearranging the equation we get

$$\epsilon + \frac{d\epsilon}{dt} = \frac{\sigma}{E} \quad (2.114)$$
$$\tau + \frac{d\epsilon}{dt} = \frac{\sigma_m}{\tau} \quad (2.115)$$

where $\sigma_m = \sigma/E$. Multiplying throughout by the integrating factor $e^{\tau t}$ gives

$$\frac{d(\epsilon e^{\tau t})}{dt} = \frac{\sigma_m e^{\tau t}}{\tau} \quad (2.116)$$
$$\frac{d}{dt}(\epsilon e^{\tau t}) = \frac{\sigma_m e^{\tau t}}{\tau} \quad (2.117)$$

Integrating both sides of this equation with respect to $t$ gives

$$\epsilon e^{\tau t} = \epsilon_m e^{\tau t} + C \quad (2.118)$$

where $C$ is a constant of integration determined by the boundary conditions. Initially, the strain is zero, i.e., at $t = 0$, $\epsilon = 0$; substituting in Eq. (2.118) gives $C = \epsilon_m$. The solution for the strain at time $t$ is therefore

$$\epsilon = \epsilon_m (1 - e^{-\tau t}) \quad (2.119)$$

The strain rises exponentially to a limiting value given by $\epsilon_m = \sigma/E$. This is characteristic of viscoelastic deformation.

### 2.8.4 Creep

Many solid materials deform slowly at room temperature when subjected to small stresses well below their brittle strength for long periods of time. The slow time-dependent deformation is known as creep. This is an important mechanism in the deformation of rocks because of the great intervals of time involved in geological processes. It is hard enough to approximate the conditions of pressure and temperature in the real Earth, but the time factor is an added difficulty in investigating the phenomenon of creep in laboratory experiments.

The results of observations on rock samples loaded by a constant stress typically show three main regimes of creep (Fig. 2.72a). At first, the rock at once strains elastically. This is followed by a stage in which the strain initially increases rapidly (i.e., the strain rate is high) and then levels off. This stage is known as primary creep or delayed elastic creep. If the stress is removed within this regime, the deformation drops rapidly as the elastic strain recovers, leaving a deformation that sinks progressively to zero. Beyond the primary stage creep progresses at a slower and nearly constant rate. This is called secondary creep or steady-state creep. The rock deforms plastically, so that if the stress is removed a permanent strain is left after the elastic and delayed elastic recoveries. After the secondary stage the strain rate increases ever more rapidly in a stage called tertiary creep, which eventually leads to failure.

Fig. 2.72 Hypothetical strain-time curve for a material exhibiting creep under constant stress, and (b) model creep curve that combines elastic, viscoelastic and viscous elements after Ramsay, 1967.

The primary and secondary stages of the creep curve can be modelled by combining elastic, viscoelastic and viscous elements (Fig. 2.72b). Below the yield stress only elastic and delayed elastic (viscoelastic) deformation occur and the solid does not flow. The strain flattens off at a limiting value $\epsilon_m = \sigma/E$. The viscous component of strain rises linearly with time, corresponding to a constant strain rate. In practice the stress must exceed the yield stress $\sigma_y$ for flow to occur. In this case the viscous component of strain is proportional to the excess stress $(\sigma - \sigma_y)$ and to the time $t$. Combining terms gives the expression

$$\epsilon = \frac{\sigma}{E} + \epsilon_m (1 - e^{-\tau t}) + \frac{(\sigma - \sigma_y)}{\eta} t \quad (2.120)$$

This simple model explains the main features of experimentally observed creep curves. It is in fact very difficult to ensure that laboratory observations are representative of creep in the Earth. Conditions of pressure and temperature
in the crust and upper mantle can be achieved or approximated. The major problems arise from the differences in timescales and creep rates. Even creep experiments conducted over months or years are far shorter than the lengths of time in a geological process. The creep rates in nature (e.g., around $10^{-11}$ s$^{-1}$) are many orders of magnitude slower than the slowest strain rate used in laboratory experiments (around $10^{-8}$ s$^{-1}$). Nevertheless, the experiments have provided a better understanding of the rheology of the Earth’s interior and the physical mechanisms active in different depths.

2.8.4.1 Crystal defects

Deformation in solids does not take place homogeneously. Laboratory observations on metals and minerals have shown that crystal defects play an important role. The atoms in a metal or crystal are arranged regularly to form a lattice with a simple symmetry. In some common arrangements the atoms are located at the corners of a cube or a hexagonal prism, defining a unit cell of the crystal. The lattice is formed by stacking unit cells together. Occasionally an imperfect cell may lack an atom. The space of the missing atom is called a vacancy. Vacancies may be distributed throughout the crystal lattice, but they can also form long chains called dislocations.

There are several types of dislocation, the simplest being an edge dislocation. It is formed when an extra plane of atoms is introduced in the lattice (Fig. 2.73). The edge dislocation terminates at a plane perpendicular to it called the glide plane. It clearly makes a difference if the extra plane of atoms is above or below the glide plane, so edge dislocations have a sign. This can be represented by a T-shape, where the cross-bar of the T is parallel to the glide plane and the stalk is the extra plane of atoms. If oppositely signed edge dislocations meet, they form a complete plane of atoms and the dislocations annihilate each other. The displacement of atoms in the vicinity of a dislocation increases the local internal stress in the crystal. As a result, the application of a small external stress may be enough to mobilize the dislocation, causing it to glide through the undisturbed part of the crystal (Fig. 2.73). If the dislocation glide is not blocked by an obstacle, the dislocation migrates out of the crystal, leaving a shear deformation.

Another common type of dislocation is the screw dislocation. It also is made up of atoms that are displaced from their regular positions, in this case forming a spiral about an axis.

The deformation of a crystal lattice by dislocation glide requires a shear stress; hydrostatic pressure does not cause plastic deformation. The shear stress needed to actuate dislocations is two or three orders of magnitude less than the shear stress needed to break the bonds between layers of atoms in a crystal. Hence, the mobilization of dislocations is an important mechanism in plastic deformation at low stress. As deformation progresses it is accompanied by an increase in the dislocation density (the number of dislocations per unit area normal to their lengths). The dislocations move along a glide plane until it intersects the glide plane of another set of dislocations. When several sets of dislocations are mobilized they may interfere and block each other, so that the stress must be increased to mobilize them further. This is manifest as strain-hardening, which is a thermodynamically unstable situation. At any stage of strain-hardening, given enough time, the dislocations redistribute themselves to a configuration with lower energy, thereby reducing the strain. The time-dependent strain relaxation is called recovery.

Recovery can take place by several processes, each of which requires thermal energy. These include the annihilation of oppositely signed edge dislocations moving on parallel glide planes (Fig. 2.74a) and the climb of edge dislocations past obstacles against which they have piled up (Fig. 2.74b). Edge dislocations with the same sign may align to form walls between domains of a crystal that have low dislocation density (Fig. 2.74c), a process called polygonization. These are some of the ways in which thermal energy promotes the migration of lattice defects,
eventually driving them out of the crystal and leaving an annealed lattice.

2.8.4.2 Creep mechanisms in the Earth

Ductile flow in the Earth’s crust and mantle takes place by one of three mechanisms: low-temperature plastic flow; power-law creep; or diffusion creep. Each mechanism is a thermally activated process. This means that the strain rate depends on the temperature $T$ according to an exponential function with the form $e^{-E_a/kT}$. Here $k$ is Boltzmann’s constant, while $E_a$ is the energy needed to activate the type of flow; it is called the activation energy. At low temperatures, where $T \ll E_a/k$, the strain rate is very slow and creep is insignificant. Because of the exponential function, the strain rate increases rapidly with increasing temperature above $T = E_a/k$. The type of flow at a particular depth depends on the local temperature and its relationship to the melting temperature $T_{mp}$. Above $T_{mp}$ the interatomic bonds in the solid break down and it flows as a true liquid.

Plastic flow at low temperature takes place by the motion of dislocations on glide planes. When the dislocations encounter an internal obstacle or a crystal boundary they pile up and some rearrangement is necessary. The stress must be increased to overcome the obstacle and reactivate dislocation glide. Plastic flow can produce large strains and may be an important mechanism in the bending of the oceanic lithosphere near some subduction zones. It is likely to be most effective at depths below the brittle–ductile transition.

Power-law creep, or hot creep, also takes place by the motion of dislocations on glide planes. It occurs at higher temperatures than low-temperature plastic flow, so internal obstacles to dislocation migration are thermally activated and diffuse out of the crystal as soon as they arise. The strain rate in power-law creep is proportional to the $n$th power of the stress $\sigma$ and has the form

$$\frac{d\varepsilon}{dt} = A(\frac{\sigma}{\mu})^n e^{-E_a/kT}$$

(2.121)

where $\mu$ is the rigidity modulus and $A$ is a constant with the dimensions of strain rate; typically $n \approx 3$. This relationship means that the strain rate increases much more rapidly than the stress. From experiments on metals, power-law creep is understood to be the most important mechanism of flow at temperatures between $0.55T_{mp}$ and $0.85T_{mp}$. The temperature throughout most of the mantle probably exceeds half the melting point, so power-law creep is probably the flow mechanism that permits mantle convection. It is also likely to be the main form of deformation in the lower lithosphere, where the relationship of temperature to melting point is also suitable.

Diffusion creep consists of the thermally activated migration of crystal defects in the presence of a stress field. There are two main forms. Nabarro–Herring creep consists of diffusion of defects through the body of a grain; Coble creep takes place by migration of the defects along grain boundaries. In each case the strain rate is proportional to the stress, as in a Newtonian fluid. It is therefore possible to regard ductile deformation by diffusion creep as the slow flow of a very viscous fluid. Diffusion creep has been observed in metals at temperatures $T > 0.85T_{mp}$. In the Earth’s mantle the temperature approaches the melting point in the asthenosphere. As indicated schematically in Fig. 2.69 the transition from the rigid lithosphere to the soft, viscous underlying asthenosphere is gradational. There is no abrupt boundary, but the concept of rigid lithospheric plates moving on the soft, viscous asthenosphere serves well as a geodynamic model.

2.8.5 Rigidity of the lithosphere

Lithospheric plates are thin compared to their horizontal extents. However, they evidently react rigidly to the forces that propel them. The lithosphere does not easily buckle under horizontal stress. A simple analogy may be made to a thin sheet of paper resting on a flat pillow. If pushed on one edge, the page simply slides across the pillow without crumpling. Only if the leading edge encounters an obstacle does the page bend, buckling upward some distance in...
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front of the hindrance, while the leading edge tries to burrow under it. This is what happens when an oceanic lithospheric plate collides with another plate. A small forebulge develops on the oceanic plate and the leading edge bends downward into the mantle, forming a subduction zone.

The ability to bend is a measure of the rigidity of the plate. This is also manifest in its reaction to a local vertical load. If, in our analogy, a small weight is placed in the middle of the page, it is pressed down into the soft pillow. A large area around the weight takes part in this process, which may be compared with the Vening Meinesz type of regional isostatic compensation (Section 2.7.2.3). The weight of a seamount or chain of islands has a similar effect on the oceanic lithosphere. By studying the flexure due to local vertical loads, information is obtained about a static property of the lithosphere, namely its resistance to bending.

In our analogy the locally loaded paper sheet would not bend if it lay on a hard flat table. It is only able to flex if it rests on a soft, yielding surface. After the weight is removed, the page is restored to its original flat shape. The restoration after unloading is a measure of the properties of the pillow as well as the page. A natural example is the rebound of regions (such as the Canadian shield or Fennoscandia) that have been depressed by now-vanished ice-sheets. The analysis of the rates of glacial rebound information is obtained about elastic parameter of the mantle beneath the lithosphere. The pressure of the surface forces mantle material to flow away laterally to make way for it; when the load is removed, the return mantle flow presses the concavity back upward. The ease with which the mantle material flows is described by its dynamic viscosity.

The resistance to bending of a thin elastic plate overlying a weak fluid is expressed by an elastic parameter called the flexural rigidity and denoted $D$. For a plate of thickness $h$,

$$D = \frac{E}{12(1 - \nu^2)} h^3 \quad \text{(2.122)}$$

where $E$ is Young’s modulus and $\nu$ is Poisson’s ratio (see Sections 3.2.3 and 3.2.4 for the definition of these elastic constants). The dimensions of $E$ are N m$^{-2}$ and $\nu$ is dimensionless; hence, the dimensions of $D$ are those of a bending moment (N m). $D$ is a fundamental parameter of the elastic plate, which describes how easily it can be bent; a large value of $D$ corresponds to a stiff plate.

Here we consider two situations of particular interest for the rigidity of the oceanic lithosphere. The first is the bending of the lithosphere by a topographic feature such as an oceanic island or seamount; only the vertical load on the elastic plate is important. The second is the bending of the lithosphere at a subduction zone. In this case vertical and horizontal forces are located along the edge of the plate and the plate experiences a bending moment which deflects it downward.

2.8.5.1 Lithospheric flexure caused by oceanic islands

The theory for elastic bending of the lithosphere is derived from the bending of thin elastic plates and beams. This involves a fourth-order differential equation, whose derivation and solution are beyond the scope of this book. However, it is instructive to consider the forces involved in setting up the equation, and to examine its solution in a simplified context.

Consider the bending of a thin isotropic elastic plate of thickness $h$ carrying a surface load $L(x, y)$ and supported by a substratum of density $\rho_i$ (Fig. 2.75a). Let the deflection of the plate at a position $(x, y)$ relative to the center of the load be $w(x, y)$. Two forces act to counteract the downward force of the load. The first, arising from Archimedes’ principle, is a buoyant force equal to $(\rho_m - \rho_i)gw$, where $\rho_i$ is the density of the material that fills in the depression caused by the deflection of the plate. The second force arises from the elasticity of the beam. Elasticity theory shows that this produces a restoring force proportional to a fourth-order differential of the deflection $w$. Balancing the elastic and buoyancy forces against the deforming load leads to the equation

$$D \left[ \frac{\partial^4 w}{\partial x^4} + 2 \frac{\partial^4 w}{\partial x^2 \partial y^2} + \frac{\partial^4 w}{\partial y^4} \right] + (\rho_m - \rho_i)gw = L(x, y) \quad \text{(2.123)}$$
For a linear topographic feature, such as a mountain range, oceanic island or chain of seamounts, the bending geometry is the same in any cross-section normal to its length and the problem reduces to the two-dimensional elastic bending of a thin beam. If the load is a linear feature in the $y$-direction, the variation of $w$ with $y$ disappears and the differential equation becomes:

$$D\frac{d^4w}{dx^4} + (\rho_m - \rho_l)gw = L \tag{2.124}$$

An important example is a linear load $L$ concentrated along the $y$-axis at $x = 0$. The solution is a damped sinusoidal function:

$$w = w_0 e^{-x\alpha}(\cos \frac{x}{\Delta} + \sin \frac{x}{\Delta}) \tag{2.125}$$

where $w_0$ is the amplitude of the maximum deflection underneath the load (at $x = 0$). The parameter $\alpha$ is called the flexural parameter; it is related to the flexural rigidity $D$ of the plate by

$$\alpha^2 = \frac{4D}{(\rho_m - \rho_l)g} \tag{2.126}$$

The elasticity of the plate (or beam) distributes the load of the surface feature over a large lateral distance. The fluid beneath the load is pushed aside by the penetration of the plate. The buoyancy of the displaced fluid forces it upward, causing uplift of the surface adjacent to the central depression. Equation (2.125) shows that this effect is repeated with increasing distance from the load, the wavelength of the fluctuation is $\lambda = 2\pi\alpha$. The amplitude of the disturbance diminishes rapidly because of the exponential attenuation factor. Usually it is only necessary to consider the central depression and the first uplifted region. The wavelength $\lambda$ is equal to the distance across the central depression. Substituting in Eq. (2.126) gives $D$, which is then used with the parameters $E$ and $\nu$ in Eq. (2.122) to obtain $h$, the thickness of the elastic plate. The computed values of $h$ are greater than the thickness of the crust, i.e., the elastic plate includes part of the upper mantle. The value of $h$ is equated with the thickness of the elastic lithosphere.

The difference between the deflection caused by a two-dimensional load (i.e., a linear feature) and that due to a three-dimensional load (i.e., a feature that has limited extent in the $x$- and $y$-directions) is illustrated in Fig. 2.75b. If the length of the three-dimensional load normal to the cross-section is more than about ten times its width, the deflection is the same as for a two-dimensional load. A load with a square base (i.e., extending the same distance along both $x$- and $y$-axes) causes a central depression that is less than a quarter the effect of the linear load.

The validity of the lithospheric flexural model of isostasy can be tested by comparing the computed gravity effect of the model with the observed free-air gravity anomaly $\Delta g$. The isostatic compensation of the Great Meteor seamount in the North Atlantic provides a suitable test for a three-dimensional model (Fig. 2.76). The shape of the free-air gravity anomaly obtained from detailed marine gravity surveys was found to be fitted best by a model in which the effective flexural rigidity of the deformed plate was assumed to be $6 \times 10^{22}$ N m.

### 2.8.5.2 Lithospheric flexure at a subduction zone

The bathymetry of an oceanic plate at a subduction zone is typified by an oceanic trench, which can be many kilometers deep (Fig. 2.77a). Seaward of the trench axis the plate develops a small upward bulge (the outer rise) which can extend for 100–150 km away from the trench and reach heights of several hundred meters. The lithospheric plate bends sharply downward in the subduction zone. This bending can also be modelled with a thin elastic plate.

In the model, a horizontal force $P$ pushes a plate of thickness $h$ toward the subduction zone, the leading edge carries a vertical load $L$, and the plate is bent by a bending moment $M$ (Fig. 2.77b). The horizontal force $P$ is negligible in comparison to the effects of $M$ and $L$. The vertical deflection of the plate must satisfy Eq. (2.124), with the same parameters as in the previous example. Choosing the origin to be the point nearest the trench where the deflection is zero simplifies the form of the solution, which is...
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\[ w = Ae^{-\alpha x} \sin \frac{x}{\alpha} \]  \hspace{1cm} (2.127)

where \( A \) is a constant and \( \alpha \) is the flexural parameter as in Eq. (2.126). The value of the constant \( A \) is found from the position \( x_b \) of the forebulge, where \( dw/dx = 0 \):

\[ \frac{dw}{dx} = A \left\{ -\frac{1}{\alpha} e^{-\alpha x} \sin \frac{x}{\alpha} + \frac{1}{\alpha} e^{-\alpha x} \cos \frac{x}{\alpha} \right\} = 0 \]  \hspace{1cm} (2.128)

from which

\[ x_b = \frac{\pi}{4} \alpha \quad \text{and} \quad A = w_b \sqrt{2} e^{-\pi^2 / 4} \]  \hspace{1cm} (2.129)

It is convenient to normalize the horizontal distance and vertical displacement: writing \( x' = x/x_b \) and \( w' = w/w_b \), the generalized equation for the elastic bending at an oceanic trench is obtained:

\[ w' = \sqrt{2} \sin \left( \frac{\pi}{4} x' \right) \exp \left[ \frac{\pi}{4} (1 - x') \right] \]  \hspace{1cm} (2.130)

The theoretical deflection of oceanic lithosphere at a subduction zone obtained from the elastic bending model agrees well with the observed bathymetry on profiles across oceanic trenches (Fig. 2.78a, b). The calculated thicknesses of the elastic lithosphere are of the order 20–30 km and the flexural rigidity is around \( 10^{23} \) N m. However, at some trenches the assumption of a completely elastic upper lithosphere is evidently inappropriate. At the Tonga trench the model curve deviates from the observed bathymetry inside the trench (Fig. 2.78c). It is likely that the elastic limit is exceeded at parts of the plate where the curvature is high. These regions may yield, leading to a reduction in the effective rigidity of the plate. This effect can be taken into account by assuming that the inelastic deformation is perfectly plastic. The deflection calculated within the trench for an elastic–perfectly plastic model agrees well with the observed bathymetry.

2.8.5.3 Thickness of the lithosphere

The rheological response of a material to stress depends on the duration of the stress. The reaction to a short-lasting stress, as experienced during the passage of a seismic wave, may be quite different from the reaction of the same material to a steady load applied for a long period of time. This is evident in the different thicknesses obtained for the lithosphere in seismic experiments and in elastic plate modelling. Long-period surface waves penetrate well into the upper mantle. Long wavelengths are slowed down by the low rigidity of the asthenosphere, so the dispersion of surface waves allows estimates of the seismic thickness of the lithosphere. For oceanic lithosphere the seismic thickness increases with age of the lithosphere (i.e., with distance from the spreading center), increasing to more than 100 km at ages older than 100 Ma (Fig. 2.79). The lithospheric thicknesses obtained from elastic modelling of the bending caused by seamounts and island chains or at subduction zones also increase with distance from the ridge, but are only one-third to one-half of the corresponding seismic thickness. The discrepancy shows that only the upper part of the lithosphere is elastic. Indeed, if the entire lithosphere had the flexural rigidity found in elastic models \( (D = 10^{21} - 10^{22} \text{ N m}) \), it would bend by only small amounts under topographic loads or at subduction zones. The base of the elastic lithosphere agrees well with the modelled depths of the 300–600 °C oceanic isotherms. At greater
depths the increase in temperature results in inelastic behavior of the lower lithosphere.

The elastic thickness of the continental lithosphere is much thicker than that of the oceanic lithosphere, except in rifts, passive continental margins and young orogenic belts. Precambrian shield areas generally have a flexural thickness greater than 100 km and a high flexural rigidity of around $10^{25}$ N m. Rifts, on the other hand, have a flexural thickness less than 25 km. Both continental and oceanic lithosphere grade gradually into the asthenosphere, which has a much lower rigidity and is able to flow in a manner determined by mantle viscosity.

### 2.8.6 Mantle viscosity

As illustrated by the transition from brittle to ductile behavior (Section 2.8.1), the Earth’s rheology changes with depth. The upper part of the lithosphere behaves elastically. It has a constant and reversible response to both short-term and long-term loads. The behavior is characterized by the rigidity or shear modulus $\mu$, which relates the strain to the applied shear stress and so has the dimensions of stress (N m$^{-2}$, or Pa). The resistance of the lithosphere to flexure is described by the flexural rigidity $D$, which has the dimensions of a bending moment (N m). In a subduction zone the tight bending may locally exceed the elastic limit, causing parts of the plate to yield.

The deeper part of the lithosphere does not behave elastically. Although it has an elastic response to abrupt stress changes, it reacts to long-lasting stress by ductile flow. This kind of rheological behavior also characterizes the asthenosphere and the deeper mantle. Flow takes place with a strain rate that is proportional to the stress or a power thereof. In the simplest case, the deformation occurs by Newtonian flow governed by a viscosity coefficient $\eta$, whose dimensions (Pa s) express the time-dependent nature of the process.

Under a surface load, such as an ice-sheet, the elastic lithosphere is pushed down into the viscous mantle (Fig. 2.80a). This causes an outflow of mantle material away from the depressed region. When the ice-sheet melts, removing the load, hydrostatic equilibrium is restored and there is a return flow of the viscous mantle material (Fig. 2.80b). Thus, in modelling the time-dependent reaction of the Earth to a surface load at least two and usually three layers must be taken into account. The top layer is an elastic lithosphere up to 100 km thick; it has infinite viscosity (i.e., it does not flow) and a flexural rigidity around $5 \times 10^{24}$ N m. Beneath it lies a low-viscosity “channel” 75–250 km thick that includes the asthenosphere, with a low viscosity of typically $10^{19}$–$10^{20}$ Pa s. The deeper mantle which makes up the third layer has a higher viscosity around $10^{21}$ Pa s.

Restoration of the surface after removal of a load is accompanied by uplift, which can be expressed well by a simple exponential relaxation equation. If the initial depression of the surface is $w_0$, the deflection $w(t)$ after time $t$ is given by

$$w(t) = w_0 e^{-t/\tau} \quad (2.131)$$

Here $\tau$ is the relaxation time, which is related to the mantle viscosity by

$$\tau = \frac{4\pi}{\rho_g g \lambda \eta} \quad (2.132)$$

where $\rho_m$ is the mantle density, $g$ is gravity at the depth of the flow and $\lambda$ is the wavelength of the depression, a
viscosity models, which indicate that its viscosity must be about 100–150 km thick, whose thickness and seismic velocity are variable from one locality to another. Interpreted as the result of softening or partial melting due to temperature, the seismic low-velocity channel beneath the lithosphere is called the asthenosphere. It is not sharply bounded, yet it must be represented by a distinct layer in viscosity models, which indicate that its viscosity must be at least 25 times less than in the deeper mantle.

The larger the ice-sheet (or other type of surface load), the deeper the effects reach into the mantle. By studying the uplift following removal of different loads, information is obtained about the viscosity at different depths in the mantle.

### 2.8.6.1 Viscosity of the upper mantle

About 18,000–20,000 years ago Lake Bonneville, the predecessor of the present Great Salt Lake in Utah, USA, had a radius of about 95 km and an estimated depth of about 305 m. The water mass depressed the lithosphere, which was later uplifted by isostatic restoration after the lake drained and dried up. Observations of the present heights of ancient shorelines show that the central part of the lake has been uplifted by about 65 m. Two parameters are involved in the process: the flexural rigidity of the lithosphere, and the viscosity of the mantle beneath. The elastic response of the lithosphere is estimated from the geometry of the depression that would be produced in isostatic equilibrium. The maximum flexural rigidity that would allow a 65 m deflection under a 305 m water load is found to be about $5 \times 10^{23}$ N m.

The surface load can be modelled as a heavy vertical right cylinder with radius $r$, which pushes itself into the soft mantle. The underlying viscous material is forced aside so that the central depression is surrounded by a circular uplifted “bulge.” After removal of the load, restorative uplift takes place in the central depression, the peripheral bulge subsides and the contour of zero-uplift migrates outward. The wavelength $\lambda$ of the depression caused by a load with this geometry has been found to be about 2.6 times the diameter of the cylindrical load. In the case of Lake Bonneville $2r = 192$ km, so $\lambda$ is about 500 km. The mantle viscosity is obtained by assuming that the response time of the lithosphere was short enough to track the loading history quite closely. This implies that the viscous relaxation time $\tau$ must have been 4000 yr or less. Substitution of these values for $\lambda$ and $\tau$ in Eq. (2.132) suggests a maximum mantle viscosity $\eta$ of about $2 \times 10^{20}$ Pa s in the top 250 km of the mantle. A lower value of $\eta$ would require a thinner low-viscosity channel beneath the lithosphere.

The Fennoscandian uplift can be treated in the same way (Fig. 2.82), but the weight and lateral expanse of the load were much larger. The ice-cap is estimated to have been about 1100 m thick and to have covered most of Norway, Sweden and Finland. Although the load was therefore somewhat elongate, it is possible to model it satisfactorily by a vertical cylinder of radius $r = 550$ km centered on the northern Gulf of Bothnia (Fig. 2.83). The load was applied for about 20,000 yr before being removed 10,000 yr ago. This caused an initial depression of about 300 m, which has subsequently been relaxing (Fig. 2.82). The uplift rates allow upper-mantle viscosities to be estimated. The data are compatible with different models of mantle structure, two of which are compared in Fig. 2.83. Each model has an elastic lithosphere with a flexural rigidity of $5 \times 10^{24}$ N m underlain by a low-viscosity channel...
and the rest of the mantle. The first model has a 75 km thick channel ($\eta = 4 \times 10^{19}$ Pa s) over a viscous mantle ($\eta = 10^{21}$ Pa s). The alternative model has a 100 km thick channel (viscosity coefficient $\eta = 1.3 \times 10^{19}$ Pa s) over a rigid mantle. Both models yield uplift rates that agree quite well with the observed uplift rates. However, results from North America indicate that the lower mantle is not rigid and so the first model fits the data better.

2.8.6.2 Viscosity of the lower mantle

Geologists have developed a coherent picture of the last glacial stage, the Wisconsin, during which much of North America was covered by an ice-sheet over 3500 m thick. The ice persisted for about 20,000 yr and melted about 10,000 yr ago. It caused a surface depression of about 600 m. The subsequent history of uplift in the James Bay area near the center of the feature has been reconstructed using geological indicators and dated by the radiocarbon method (see Section 4.1.4.1). For modelling purposes the ice-sheet can be represented as a right cylindrical load with radius $r = 1650$ km (Fig. 2.84, inset). A load as large as this affects the deep mantle. The central uplift following removal of the load has been calculated for several Earth models. Each model has elastic parameters and density distribution obtained from seismic velocities, including a central dense core. The models differ from each other in the number of viscous layers in the mantle and the amount by which the density gradient departs from the adiabatic gradient (Table 2.2). The curvature of the observed uplift curve only fits models in which the viscosity of the lower mantle is around $10^{21}$ Pa s (Fig. 2.84). A highly viscous lower mantle ($\eta = 10^{23}$ Pa s, model 4) is
incompatible with the observed uplift history. The best fit is obtained with model 1 or 5. Each has an adiabatic density gradient, but model 1 has a uniform lower mantle with viscosity $10^{21}$ Pa s, and model 5 has $\eta$ increasing from $10^{21}$ Pa s below the lithosphere to $3 \times 10^{21}$ Pa s just above the core.

The viscoelastic properties of the Earth’s interior influence the Earth’s rotation, causing changes in the position of the instantaneous rotation axis. The motion of the axis is traced by repeated photo-zenith tube measurements, in which the zenith is located by photographing the stars vertically above an observatory. Photo-zenith tube measurements reveal systematic movements of the rotation axis relative to the axis of figure (Fig. 2.85). Decomposed into components along the Greenwich meridian ($X$-axis) and the 90°W meridian ($Y$-axis), the polar motion exhibits a fluctuation with cyclically varying amplitude superposed on a linear trend. The amplitude modulation has a period of approximately seven years and is due to the interference of the 12-month annual wobble and the 14-month Chandler wobble. The linear trend represents a slow drift of the pole toward northern Canada at a rate of 0.95 degrees per million years. It is due to the melting of the Fennoscandian and Laurentide ice-sheets. The subsequent uplift constitutes a redistribution of mass which causes modifications to the Earth’s moments and products of inertia, thus affecting the rotation.

The observed polar drift can be modelled with different viscoelastic Earth structures. The models assume a 120 km thick elastic lithosphere and take into account different viscosities in the layers bounded by the seismic discontinuities at 400 km and 670 km depths (Section 3.7, Table 3.4) and the core–mantle boundary. An Earth that is homogeneous below the lithosphere (without a core) is found to give imperceptible drift. Inclusion of the core, with a density jump across the core–mantle boundary and assuming the mantle viscosity to be around $1 \times 10^{21}$ Pa s, gives a drift that is perceptible but much slower than that observed. Introduction of a density change at the 670 km discontinuity increases the drift markedly; the 400 km discontinuity does not noticeably change the drift further. The optimum model has an upper-mantle viscosity of about $1 \times 10^{21}$ Pa s and a lower-mantle viscosity of about $3 \times 10^{21}$ Pa s. The model satisfies both the rate of drift and its direction (Fig. 2.85). The viscosities are comparable to values found by modelling post-glacial uplift (Table 2.2, model 5).

### Table 2.2 Parameters of Earth models used in computing uplift rates. All models are for an elastic Earth with a dense core (after Cathles, 1975)

<table>
<thead>
<tr>
<th>Model</th>
<th>Density gradient</th>
<th>Viscosity $[10^{21}$ Pa s$]$</th>
<th>Depth interval</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>adiabatic</td>
<td>1</td>
<td>entire mantle</td>
</tr>
<tr>
<td>2</td>
<td>adiabatic,</td>
<td>1</td>
<td>entire mantle</td>
</tr>
<tr>
<td></td>
<td>except 335–635 km</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>adiabatic,</td>
<td>0.1</td>
<td>0–335 km</td>
</tr>
<tr>
<td></td>
<td>except 335–635 km</td>
<td></td>
<td>335 km to core</td>
</tr>
<tr>
<td>4</td>
<td>adiabatic</td>
<td>1</td>
<td>0–985 km</td>
</tr>
<tr>
<td></td>
<td></td>
<td>100</td>
<td>985–2185 km</td>
</tr>
<tr>
<td>5</td>
<td>adiabatic</td>
<td>1</td>
<td>0–985 km</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3</td>
<td>985–2185 km</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3</td>
<td>2185 km to core</td>
</tr>
</tbody>
</table>

![Fig. 2.85 Changes in the position of the instantaneous rotation axis from 1900 to 1975 relative to axes defined in the inset (after Peltier, 1989).](image_url)

### 2.9 SUGGESTIONS FOR FURTHER READING

### Introductory level


118  Gravity, the figure of the Earth and geodynamics

**Intermediate level**


**Advanced level**


### 2.10 REVIEW QUESTIONS

1. Describe the principle of operation of a gravimeter.
2. Explain why a gravimeter only gives relative measurements of gravity.
3. What is the geoid? What is the reference ellipsoid? How and why do they differ?
4. What is a geoid anomaly? Explain how a positive (or negative) anomaly arises.
5. What is normal gravity? What does the word normal imply? Which surface is involved?
6. Gravitational acceleration is directed toward a center of mass. With the aid of a sketch that shows the directions of gravity’s components, explain why gravity is not a centrally directed acceleration.
7. Write down the general expression for the normal gravity formula. Explain which geophysical parameters determine each of the constants in the formula.
8. What is the topographic correction in the reduction of gravity data? Why is it needed?
9. What are the Bouguer plate and free-air gravity corrections?
10. What is a free-air gravity anomaly? How does it differ from a Bouguer anomaly?
11. Sketch how the Bouguer gravity anomaly might vary on a continuous profile that extends from a continental mountain range to an oceanic ridge.
12. What is the Coriolis acceleration? How does it originate? How does it affect wind patterns in the North and South hemispheres?
13. What is the Eötvös gravity correction? When is it needed? How does it originate?
14. Describe two borehole methods for determining the density of rocks around the borehole.
15. Describe and explain the Nettleton profile method to determine the optimum density for interpreting a gravity survey.
16. Explain how to calculate the position of the common center of mass of the Earth–Moon system.
17. Explain with the aid of diagrams that show the forces involved, why there are two lunar tides per day.
18. Why are the lunar tides almost equal on opposite sides of the Earth? Why are they not exactly equal?
19. What is isostasy? What is an isostatic gravity anomaly?
20. Why is the free-air gravity anomaly close to zero at the middle of a large crustal block that is in isostatic equilibrium?
21. Describe the three models of isostasy, and explain how they differ from each other.
22. What would be the geodynamic behavior of a region that is characterized by a negative isostatic gravity anomaly?

### 2.11 EXERCISES

1. Using the data in Table 1.1 calculate the gravitational acceleration on the surface of the Moon as a percentage of that on the surface of the Earth.
2. An Olympic high-jump champion jumps a record height of 2.45 m on the Earth. How high could this champion jump on the Moon?
3. (a) Calculate the escape velocity of an object on the Earth, assuming a mean gravitational acceleration of 9.81 m s$^{-1}$ and mean Earth radius of 6371 km. (b) What is the escape velocity of the same object on the Moon?
4. The equatorial radius of the Earth is 6378 km and gravity at the equator is 9.780 m s$^{-2}$. Compute the ratio $m$ of the centrifugal acceleration at the equator to the gravitational acceleration at the equator. If the ratio $m$ is written as $1/k$, what is the value of $k$?
5. Given that the length of a month is 27.32 days, the mean gravity on Earth is 9.81 m s\(^{-2}\) and the Earth's radius is 6371 km, calculate the radius of the Moon’s orbit.

6. A communications satellite is to be placed in a geostationary orbit.
   (a) What must the position and orientation of the orbit be?
   (b) What is the radius of the orbit?
   (c) If a radio signal is sent to the satellite from a transmitter at latitude 45°N, what is the shortest time taken for its reflection to reach the Earth?

7. Calculate the centrifugal acceleration due to the Earth’s rotation of an object at rest on the Earth’s surface in Paris, assuming a latitude of 48° 52’ N. Express the result as a percentage of the gravitational attraction on the object.

8. A solid angle (\(\Omega\)) is defined as the quotient of the area (\(A\)) of the part of a spherical surface subtended by the angle, divided by the square of the spherical radius (\(r\)): \(\Omega = A/r^2\) (see Box 5.4). Show with the aid of a diagram that the gravitational acceleration at any point inside a thin homogeneous spherical shell is zero.

9. Assuming that the gravitational acceleration inside a homogeneous spherical shell is zero, show that the gravitational acceleration inside a homogeneous uniform solid sphere is proportional to the distance from its center.

10. Show that the gravitational potential \(U_G\) inside a homogenous uniform solid sphere of radius \(R\) at a distance \(r\) from its center is given by
    \[
    U_G = -\frac{2\pi}{3} G\rho (3R^2 - r^2)
    \]

11. Sketch the variations of gravitational acceleration and potential inside and outside a homogeneous solid sphere of radius \(R\).

12. A thin borehole is drilled through the center of the Earth, and a ball is dropped into the borehole. Assume the Earth to be a homogenous solid sphere. Show that the ball will oscillate back and forth from one side of the Earth to the other. How long does it take to traverse the Earth and reach the other side?

13. The Roche limit is the closest distance an object can approach a planet before being torn apart by the tidal attraction of the planet. For a rigid spherical moon the Roche limit is given by Eq. (6) in Box 2.1.
   (a) Using the planetary dimensions in Table 1.1, calculate the Roche limit for the Moon with respect to the Earth. Express the answer as a multiple of the Earth’s radius.
   (b) Show that, for a planet whose mean density is less than half that of its rigid moon, the moon would collide with the planet before being torn apart by its gravity.
   (c) Given that the Sun’s mass is \(1.989 \times 10^{30}\) kg and that its radius is 695,500 km, calculate the Roche limit for the Earth with respect to the Sun.
   (d) The mean density of a comet is about 500 kg m\(^{-3}\). What is the Roche limit for comets that might collide with the Earth?
   (e) The mean density of an asteroid is about 2000 kg m\(^{-3}\). If an asteroid on collision course with the Earth has a velocity of 15 km s\(^{-1}\), how much time will elapse between the break-up of the asteroid at the Roche limit and the impact of the remnant pieces on the Earth’s surface, assuming they maintain the same velocity as the asteroid?

14. The mass \(M\) and moment of inertia \(C\) of a thick shell of uniform density \(\rho\), with internal radius \(r\) and external radius \(R\) are given by
    \[
    M = 4\pi\rho (R^3 - r^3) \quad C = \frac{8}{15}\pi\rho (R^5 - r^5)
    \]
    The Earth has an internal structure consisting of concentric spherical shells. A simple model with uniform density in each shell is given in the following figure.

<table>
<thead>
<tr>
<th>Layer</th>
<th>Radius (km)</th>
<th>Density (kg m(^{-3}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>upper mantle</td>
<td>6370</td>
<td>3300</td>
</tr>
<tr>
<td>lower mantle</td>
<td>5700</td>
<td>5000</td>
</tr>
<tr>
<td>outer core</td>
<td>3480</td>
<td>11000</td>
</tr>
<tr>
<td>inner core</td>
<td>1220</td>
<td>13000</td>
</tr>
</tbody>
</table>

   (a) Compute the mass and moment of inertia of each spherical shell.
   (b) Compute the total mass and total moment of inertia of the Earth.
   (c) If the moment of inertia can be written \(C = kMR^2\), where \(M\) is Earth’s mass and \(R\) its radius, what is the value of \(k\)?
   (d) What would the value of \(k\) be if the density were uniform throughout the Earth?

15. By differentiating the normal gravity formula given by Eq. (2.56) develop an expression for the change in gravity with latitude. Calculate the gravity change in milligals per kilometer of northward displacement at latitude 45°.

16. The following gravity measurements were made on a traverse across a rock formation. Use the combined elevation correction to compute the apparent density of the rock.
17. Show that the “half-width” \( w \) of the gravity anomaly over a sphere and the depth \( z \) to the center of the sphere are related by \( z / H \approx 0.652 w \).

18. Assume the “thin-sheet approximation” (Eq. (2.101)) for the gravity anomaly over a vertical fault of density contrast \( \Delta \rho \) and height \( h \) with midpoint at depth \( z_0 \).

(a) What is the maximum slope of the anomaly and where does it occur?

(b) Determine the relationship between the depth \( z_0 \) and the horizontal distance \( w \) between the positions where the slope of the anomaly is one-half the maximum slope.

19. Calculate the maximum gravity anomaly at ground level over a buried anticlinal structure, modelled by a horizontal cylinder with radius 1000 m and density contrast 200 kg m\(^{-3}\), when the depth of the axis is (a) 1500 m and (b) 5000 m.

20. The peak \( A \) of a mountain is 1000 meters above the level CD of the surrounding plain, as in the diagram. The density of the rocks forming the mountain is 2800 kg m\(^{-3}\), that of the surrounding crust is 3000 kg m\(^{-3}\). Assuming that the mountain and its “root” are symmetric about \( A \) and that the system is in isostatic equilibrium, calculate the depth of \( B \) below the level CD.

21. A crustal block with mean density 3000 kg m\(^{-3}\) is initially in isostatic equilibrium with the surrounding rocks whose density is 3200 kg m\(^{-3}\), as in the figure (a). After subsequent erosion the above-surface topography is as shown in (b). The distance \( L \) remains constant (i.e. there is no erosion at the highest point \( A \)) and Airy-type isostatic equilibrium is maintained. Calculate in terms of \( L \) the amount by which the height of \( A \) is changed. Explain why \( A \) moves in the sense given by your answer.

22. An idealized mountain-and-root system, as in the figure, is in isostatic equilibrium. The densities in kg m\(^{-3}\) are as shown. Express the height \( H \) of the point \( A \) above the horizontal surface RS in terms of the depth \( D \) of the root \( B \) below this surface.
3 Seismology and the internal structure of the Earth

3.1 INTRODUCTION

Seismology is a venerable science with a long history. The Chinese scientist Chang Heng is credited with the invention in 132 AD, nearly two thousand years ago, of the first functional seismoscope, a primitive but ingenious device of elegant construction and beautiful design that registered the arrival of seismic waves and enabled the observer to infer the direction they came from. The origins of earthquakes were not at all understood. For centuries these fearsome events were attributed to supernatural powers. The accompanying destruction and loss of life were often understood in superstitious terms and interpreted as punishment inflicted by the gods on a sinful society. Biblical mentions of earthquakes–e.g., in the destruction of Sodom and Gomorrah–emphasize this vengeful theme.

Although early astronomers and philosophers sought to explain earthquakes as natural phenomena unrelated to spiritual factors, the belief that earthquakes were an expression of divine anger prevailed until the advent of the Age of Reason in the eighteenth century. The path to a logical understanding of natural phenomena was laid in the seventeenth century by the systematic observations of scientists like Galileo, the discovery and statement of physical laws by Newton and the development of rational thought by contemporary philosophers.

In addition to the development of the techniques of scientific observation, an understanding of the laws of elasticity and the limited strength of materials was necessary before seismology could progress as a science. In a pioneering study, Galileo in 1638 described the response of a beam to loading, and in 1660 Hooke established the law of the spring. However, another 150 years passed before the generalized equations of elasticity were set down by Navier. During the early decades of the nineteenth century Cauchy and Poisson completed the foundations of modern elasticity theory.

Early descriptions of earthquake characteristics were necessarily restricted to observations and measurements in the “near-field” region of the earthquake, i.e. in comparatively close proximity to the place where it occurred. A conspicuous advance in the science of seismology was accomplished with the invention of a sensitive and reliable seismograph by John Milne in 1892. Although massive and primitive by comparison with modern instruments, the precision and sensitivity of this revolutionary new device permitted accurate, quantitative descriptions of earthquakes at large distances from their source, in their “far-field” region. The accumulation of reliable records of distant earthquakes (designated as “teleseismic” events) made possible the systematic study of the Earth’s seismicity and its internal structure.

The great San Francisco earthquake of 1906 was intensively studied and provided an impetus to efforts at understanding the origin of these natural phenomena, which were clarified in the same year by the elastic rebound model of H. F. Reid. Also, in 1906, R. D. Oldham proposed that the best explanation for the travel-times of teleseismic waves through the body of the Earth required a large, dense and probably fluid core; the depth to its outer boundary was calculated in 1913 by B. Gutenberg. From the analysis of the travel-times of seismic body waves from near earthquakes in Yugoslavia, A. Mohorovicic in 1909 inferred the existence of the crust–mantle boundary, and in 1936 the existence of the solid inner core was deduced by I. Lehmann. The definitions of these and other discontinuities associated with the deep internal structure of the Earth have since been greatly refined.

The needs of the world powers to detect incontrovertibly the testing of nuclear bombs by their adversaries provided considerable stimulus to the science of seismology in the 1950s and 1960s. The amount of energy released in a nuclear explosion is comparable to that of an earthquake, but the phenomena can be discriminated by analyzing the directions of first motion recorded by seismographs. The accurate location of the event required improved knowledge of seismic body-wave velocities throughout the Earth’s interior. These political necessities of the cold war led to major improvements in seismological instrumentation, and to the establishment of a new world-wide network of seismic stations with the same physical characteristics.

These developments had an important feedback to the earth sciences, because they resulted in more accurate location of earthquake epicenters and a better understanding of the Earth’s structure. The pattern of global seismicity, with its predominant concentration in narrow active zones, was an important factor in the development of the theory of plate tectonics, as it allowed the identification of plate margins and the sense of relative plate motions.

The techniques of refraction and reflection seismology, using artificial, controlled explosions as sources, were developed in the search for petroleum. Since the 1960s these
methods have been applied with notable success to the resolution of detailed crustal structure under continents and oceans. The development of powerful computer technology enabled refinements in earthquake location and in the determination of travel-times of seismic body waves. These advances led to the modern field of seismic tomography, a powerful and spectacular technique for revealing regions of the Earth’s interior that have anomalous seismic velocities. In the field of earthquake seismology, the need to protect populations and man-made structures has resulted in the investment of considerable effort in the study of earthquake prediction and the development of construction codes to reduce earthquake damage.

To appreciate how seismologists have unravelled the structure of the Earth’s interior it is necessary to understand what types of seismic waves can be generated by an earthquake or man-made source (such as a controlled explosion). The propagation of a seismic disturbance through the Earth is governed by physical properties such as density, and by the way in which the material of the Earth’s interior reacts to the disturbance. Material within the seismic source suffers permanent deformation, but outside the source the passage of a seismic disturbance takes place predominantly by elastic displacement of the medium; that is, the medium suffers no permanent deformation. Before analyzing the different kinds of seismic waves, it is important to have a good grasp of elementary elasticity theory. This requires understanding the concepts of stress and strain, and the various elastic constants that relate them.

### 3.2 ELASTICITY THEORY

#### 3.2.1 Elastic, anelastic and plastic behavior of materials

When a force is applied to a material, it deforms. This means that the particles of the material are displaced from their original positions. Provided the force does not exceed a critical value, the displacements are reversible; the particles of the material return to their original positions when the force is removed, and no permanent deformation results. This is called elastic behavior.

The laws of elastic deformation are illustrated by the following example. Consider a right cylindrical block of height \( h \) and cross-sectional area \( A \), subjected to a force \( F \) which acts to extend the block by the amount \( \Delta h \) (Fig. 3.1). Experiments show that for elastic deformation \( \Delta h \) is directly proportional to the applied force and to the unstretched dimension of the block, but is inversely proportional to the cross-section of the block. That is, \( \Delta h \propto \frac{F/h}{A} \), or

\[
\frac{F}{A} \propto \frac{\Delta h}{h} \tag{3.1}
\]

When the area \( A \) becomes infinitesimally small, the limiting value of the force per unit area \( F/A \) is called the stress, \( \sigma \). The units of stress are the same as the units of pressure. The SI unit is the pascal, equivalent to a force of 1 newton per square meter (1 Pa = 1 N m\(^{-2}\)); the c.g.s. unit is the bar, equal to 106 dyne cm\(^{-2}\).

When \( h \) is infinitesimally small, the fractional change in dimension \( \Delta l/hl \) is called the strain \( \varepsilon \), which is a dimensionless quantity. Equation (3.1) states that, for elastic behavior, the strain in a body is proportional to the stress applied to it. This linear relationship is called Hooke’s law. It forms the basis of elasticity theory.

Beyond a certain value of the stress, called the proportionality limit, Hooke’s law no longer holds (Fig. 3.2a). Although the material is still elastic (it returns to its original shape when stress is removed), the stress–strain relationship is non-linear. If the solid is deformed beyond a certain point, known as the elastic limit, it will not recover its original shape when stress is removed. In this range a small increase in applied stress causes a disproportionately large increase in strain. The deformation is said to be plastic. If the applied stress is removed in the plastic range, the strain does not return to zero; a permanent strain has been produced. Eventually the applied stress exceeds the strength of the material and failure occurs. In some rocks failure can occur abruptly within the elastic range; this is called brittle behavior.

The non-brittle, or ductile, behavior of materials under stress depends on the timescale of the deformation (Fig. 3.2b). An elastic material deforms immediately upon application of a stress and maintains a constant strain until the stress is removed, upon which the strain returns to its original state. A strain–time plot has a box-like shape. However, in some materials the strain does not reach a stable value immediately after application of a stress, but rises gradually to a stable value. This type of strain response is characteristic of anelastic materials. After removal of the stress, the time-dependent strain returns reversibly to the original level. In plastic deformation the strain keeps increasing as long as the stress is applied. When the stress is removed, the strain does not return to the original level; a permanent strain is left in the material.
3.2 ELASTICITY THEORY

Our knowledge of the structure and nature of the Earth's interior has been derived in large part from studies of seismic waves released by earthquakes. An earthquake occurs in the crust or upper mantle when the tectonic stress exceeds the local strength of the rocks and failure occurs. Away from the region of failure seismic waves spread out from an earthquake by elastic deformation of the rocks through which they travel. Their propagation depends on elastic properties that are described by the relationships between stress and strain.

3.2.2 The stress matrix

Consider a force \( F \) acting on a rectangular prism \( P \) in a reference frame defined by orthogonal Cartesian coordinate axes \( x, y, \) and \( z \) (Fig. 3.3a). The component of \( F \) which acts in the direction of the \( x \)-axis is designated \( F_x \); the force \( F \) is fully defined by its components \( F_x, F_y, \) and \( F_z \). The size of a small surface element is characterized by its area \( A \), while its orientation is described by the direction normal to the surface (Fig. 3.3b). The small surface with area normal to the \( x \)-axis is designated \( A_x \). The component of force \( F_x \) acting normal to the surface \( A_x \) produces a normal stress, denoted by \( \sigma_{xx} \). The components of force along the \( y \)- and \( z \)-axes result in shear stresses \( \sigma_{yx} \) and \( \sigma_{zx} \) (Fig. 3.3c), given by

\[
\sigma_{xx} = \lim_{\Delta x \to 0} \left( \frac{F_x}{A_x} \right) \quad \sigma_{yx} = \lim_{\Delta y \to 0} \left( \frac{F_y}{A_y} \right) \quad \sigma_{zx} = \lim_{\Delta z \to 0} \left( \frac{F_z}{A_z} \right) \tag{3.2}
\]

Similarly, the components of the force \( F \) acting on an element of surface \( A_y \) normal to the \( y \)-axis define a normal stress \( \sigma_{yy} \) and shear stresses \( \sigma_{yx} \) and \( \sigma_{zy} \), while the components of \( F \) acting on an element of surface \( A_z \) normal to the \( z \)-axis define a normal stress \( \sigma_{zz} \) and shear stresses \( \sigma_{zx} \) and \( \sigma_{yz} \). The nine stress components completely define the state of stress of a body. They are described conveniently by the stress matrix

\[
\begin{bmatrix}
\sigma_{xx} & \sigma_{xy} & \sigma_{xz} \\
\sigma_{yx} & \sigma_{yy} & \sigma_{yz} \\
\sigma_{zx} & \sigma_{zy} & \sigma_{zz}
\end{bmatrix} \tag{3.3}
\]

If the forces on a body are balanced to give no rotation, this \( 3 \times 3 \) matrix is symmetric (i.e., \( \sigma_{xy} = \sigma_{yx}, \sigma_{xz} = \sigma_{zx}, \sigma_{yz} = \sigma_{zy} \)) and contains only six independent elements.

3.2.3 The strain matrix

3.2.3.1 Longitudinal strain

The strains produced in a body can also be expressed by a \( 3 \times 3 \) matrix. Consider first the one-dimensional case shown in Fig. 3.4 of two points in a body located close together at the positions \( x \) and \( x + \Delta x \). If the point \( x \) is displaced by an infinitesimally small amount \( u \) in the
direction of the $x$-axis, the point $(x + \Delta x)$ will be displaced by $(u + \Delta u)$, where $\Delta u$ is equal to $(\partial u/\partial x)\Delta x$ to first order. The longitudinal strain or extension in the $x$-direction is the fractional change in length of an element along the $x$-axis. The original separation of the two points was $\Delta x$; one point was displaced by $u$, the other by $(u + \Delta u)$, so the new separation of the points is $(\Delta x + \Delta u)$. The component of strain parallel to the $x$-axis resulting from a small displacement parallel to the $x$-axis is denoted $\varepsilon_{xx}$, and is given by

$$\varepsilon_{xx} = \frac{(\Delta x + \partial u/\partial x\Delta x) - \Delta x}{\Delta x} = \frac{\partial u}{\partial x}$$  \hspace{1cm} (3.4)

The description of longitudinal strain can be expanded to three dimensions. If a point $(x, y, z)$ is displaced by an infinitesimal amount to $(x + u, y + v, z + w)$, two further longitudinal strains $\varepsilon_{yy}$ and $\varepsilon_{zz}$ are defined by

$$\varepsilon_{yy} = \frac{\partial v}{\partial y} \quad \text{and} \quad \varepsilon_{zz} = \frac{\partial w}{\partial z}$$  \hspace{1cm} (3.5)

In an elastic body the transverse strains $\varepsilon_{yy}$ and $\varepsilon_{zz}$ are not independent of the strain $\varepsilon_{xx}$. Consider the change of shape of the bar in Fig. 3.5. When it is stretched parallel to the $x$-axis, it becomes thinner parallel to the $y$-axis and parallel to the $z$-axis. The transverse longitudinal strains $\varepsilon_{yy}$ and $\varepsilon_{zz}$ are of opposite sign but proportional to the extension $\varepsilon_{xx}$, and can be expressed as

$$\varepsilon_{yy} = -\nu \varepsilon_{xx} \quad \text{and} \quad \varepsilon_{zz} = -\nu \varepsilon_{xx}$$  \hspace{1cm} (3.6)

The constant of proportionality $\nu$ is called Poisson's ratio. The values of the elastic constants of a material constrain $\nu$ to lie between 0 (no lateral contraction) and a maximum value of 0.5 (no volume change) for an incompressible fluid. In very hard, rigid rocks like granite $\nu$ is about 0.45, while in soft, poorly consolidated sediments it is about 0.05. In the interior of the Earth, $\nu$ commonly has a value around 0.24–0.27. A body for which the value of $\nu$ equals 0.25 is sometimes called an ideal Poisson body.

3.2.3.2 Dilatation

The dilatation is defined as the fractional change in volume of an element in the limit when its surface area decreases to zero. Consider an undeformed volume element (as in the description of longitudinal strain) which has sides $\Delta x$, $\Delta y$, and $\Delta z$ and undistorted volume $V = \Delta x \Delta y \Delta z$. As a result of the infinitesimal displacements $\Delta u$, $\Delta v$, and $\Delta w$ the edges increase to $\Delta x + \Delta u$, $\Delta y + \Delta v$, and $\Delta z + \Delta w$, respectively. The fractional change in volume is

$$\frac{\Delta V}{V} = \frac{(\Delta x + \Delta u)(\Delta y + \Delta v)(\Delta z + \Delta w) - \Delta x \Delta y \Delta z}{\Delta x \Delta y \Delta z}$$

$$\frac{\Delta V}{V} = \frac{\Delta x \Delta y \Delta z + \Delta u \Delta y \Delta z + \Delta x \Delta z \Delta v + \Delta u \Delta z \Delta v - \Delta x \Delta y \Delta z}{\Delta x \Delta y \Delta z}$$

where very small quantities like $\Delta u \Delta v$, $\Delta v \Delta w$, $\Delta u \Delta w$ and $\Delta u \Delta v \Delta w$ have been ignored. In the limit, as $\Delta x$, $\Delta y$, and $\Delta z$ all approach zero, we get the dilatation

$$\theta = \frac{\partial u}{\partial x} + \frac{\partial v}{\partial y} + \frac{\partial w}{\partial z}$$

$$\theta = \varepsilon_{xx} + \varepsilon_{yy} + \varepsilon_{zz}$$  \hspace{1cm} (3.8)

3.2.3.3 Shear strain

During deformation a body generally experiences not only longitudinal strains as described above. The shear components of stress ($\sigma_{xy}$, $\sigma_{yz}$, $\sigma_{zx}$) produce shear strains, which are manifest as changes in the angular relationships between parts of a body. This is most easily illustrated in two dimensions. Consider a rectangle ABCD with sides $\Delta x$ and $\Delta y$ and its distortion due to shear stresses acting in the $x$–$y$ plane (Fig. 3.6). As in the earlier example of longitudinal strain, the point A is displaced parallel to the $x$-axis by an amount $u$ (Fig. 3.6a). Because of the shear deformation, points between A and D experience larger $x$-displacements the further they are from A.
The point D which is at a vertical distance \( \Delta y \) above A is displaced by the amount \( \frac{\partial u}{\partial y} \Delta y \) in the direction of the \( x \)-axis. This causes a clockwise rotation of side AD through a small angle \( \phi_1 \) given by

\[
\tan \phi_1 = \frac{\partial u}{\partial y} \Delta y = \frac{\partial u}{\partial y}
\]  

(3.9)

Similarly, the point A is displaced parallel to the \( y \)-axis by an amount \( u \) (Fig. 3.6b), while the point B which is at a horizontal distance \( \Delta x \) from A is displaced by the amount \( \frac{\partial v}{\partial x} \Delta x \) in the direction of the \( y \)-axis. As a result side AB rotates counterclockwise through a small angle \( \phi_2 \) given by

\[
\tan \phi_2 = \frac{\partial v}{\partial x} \Delta x = \frac{\partial v}{\partial x}
\]  

(3.10)

Elastic deformation involves infinitesimally small displacements and distortions, and for small angles we can write \( \tan \phi_1 = \phi_1 \) and \( \tan \phi_2 = \phi_2 \). The shear strain in the \( x-y \) plane \( (\varepsilon_{xy}) \) is defined as half the total angular distortion (Fig. 3.6c):

\[
\varepsilon_{xy} = \frac{1}{2} \left( \frac{\partial u}{\partial y} + \frac{\partial u}{\partial x} \right)
\]  

(3.11)

By transposing \( x \) and \( y \), and the corresponding displacements \( u \) and \( v \), the shear component \( \varepsilon_{yx} \) is obtained:

\[
\varepsilon_{yx} = \frac{1}{2} \left( \frac{\partial u}{\partial y} + \frac{\partial v}{\partial x} \right)
\]  

(3.12)

This is identical to \( \varepsilon_{xy} \). The total angular distortion in the \( x-y \) plane is \( (\varepsilon_{xy} + \varepsilon_{yx}) = 2\varepsilon_{xy} = 2\varepsilon_{yx} \). Similarly, strain components \( \varepsilon_{yz} \) \( (= \varepsilon_{zy}) \) and \( \varepsilon_{zx} \) \( (= \varepsilon_{xz}) \) are defined for angular distortions in the \( y-z \) and \( z-x \) planes, respectively:

\[
\varepsilon_{yz} = \varepsilon_{zy} = \frac{1}{2} \left( \frac{\partial u}{\partial z} + \frac{\partial v}{\partial z} \right)
\]

\[
\varepsilon_{zx} = \varepsilon_{xz} = \frac{1}{2} \left( \frac{\partial u}{\partial x} + \frac{\partial v}{\partial x} \right)
\]
\[ e_{xx} = e_{yy} = e_{zz} = \frac{1}{2} \left( \frac{\partial u}{\partial x} + \frac{\partial w}{\partial x} \right) \quad (3.13) \]

The longitudinal and shear strains define the symmetric $3 \times 3$ strain matrix
\[ \begin{pmatrix} e_{xx} & e_{xy} & e_{xz} \\ e_{yx} & e_{yy} & e_{yz} \\ e_{zx} & e_{zy} & e_{zz} \end{pmatrix} \quad (3.14) \]

### 3.2.4 The elastic constants

According to Hooke’s law, when a body deforms elastically, there is a linear relationship between stress and strain. The ratio of stress to strain defines an elastic constant, or elastic modulus. Strain is itself a ratio of lengths and therefore dimensionless. The elastic moduli, defined for different types of deformation, are Young’s modulus, the rigidity modulus and the bulk modulus.

**Young’s modulus** is defined from the extensional deformations. Each longitudinal strain is proportional to the corresponding stress component, that is,
\[ \sigma_{xx} = E e_{xx} \quad \sigma_{yy} = E e_{yy} \quad \sigma_{zz} = E e_{zz} \quad (3.15) \]

where the constant of proportionality, $E$, is Young’s modulus.

**The rigidity modulus** (or shear modulus) is defined from the shear deformation. Like the longitudinal strains, the total shear strain in each plane is proportional to the corresponding shear stress component:
\[ \sigma_{xy} = 2 \mu e_{xy} \quad \sigma_{yz} = 2 \mu e_{yz} \quad \sigma_{zx} = 2 \mu e_{zx} \quad (3.16) \]

where the proportionality constant, $\mu$, is the rigidity modulus and the factor 2 arises as explained for Eqs. (3.11) and (3.12).

The **bulk modulus** (or incompressibility) is defined from the dilatation experienced by a body under hydrostatic pressure. Shear components of stress are zero for hydrostatic conditions ($\sigma_{yy} = \sigma_{zz} = \sigma_{xx} = 0$), and the inwards pressure (negative normal stress) is equal in all directions ($\sigma_{xx} = \sigma_{yy} = \sigma_{zz} = -p$). The bulk modulus, $K$, is the ratio of the hydrostatic pressure to the dilatation, that is,
\[ p = -K \theta \quad (3.17) \]

The inverse of the bulk modulus ($K^{-1}$) is called the **compressibility**.

#### 3.2.4.1 Bulk modulus in terms of Young’s modulus and Poisson’s ratio

Consider a rectangular volume element subjected to normal stresses $\sigma_{xx}$, $\sigma_{yy}$ and $\sigma_{zz}$ on its end surfaces. Each longitudinal strain $e_{xx}$, $e_{yy}$ and $e_{zz}$ results from the combined effects of $\sigma_{xx}$, $\sigma_{yy}$ and $\sigma_{zz}$. For example, applying Hooke’s law, the stress $\sigma_{xx}$ produces an extension equal to $\sigma_{xx}/E$ in the $x$-direction. The stress $\sigma_{yy}$ causes an extension $\sigma_{yy}/E$ in the $y$-direction, which results in an accompanying transverse strain $\nu(\sigma_{yy}/E)$ in the $x$-direction, where $\nu$ is Poisson’s ratio. Similarly, the stress component $\sigma_{zz}$ makes a contribution $\nu(\sigma_{zz}/E)$ to the total longitudinal strain $e_{xx}$ in the $x$-direction. Therefore,
\[ e_{xx} = \frac{\sigma_{xx}}{E} - \nu \frac{\sigma_{yy}}{E} - \nu \frac{\sigma_{zz}}{E} \quad (3.18) \]

Similar equations describe the total longitudinal strains $e_{yy}$ and $e_{zz}$. They can be rearranged as
\[ E(e_{xx} + e_{yy} + e_{zz}) = (1 - 2\nu)(\sigma_{xx} + \sigma_{yy} + \sigma_{zz}) \quad (3.20) \]

Consider now the effect of a constraining hydrostatic pressure, $p$, where $\sigma_{xx} = \sigma_{yy} = \sigma_{zz} = -p$. Using the definition of dilatation ($\theta$) in Eq. (3.8) we get
\[ E\theta = (1 - 2\nu)(-3p) \]

\[ E = (1 - 2\nu) \left( -\frac{3p}{\theta} \right) \quad (3.21) \]

from which, using the definition of bulk modulus ($K$) in Eq. (3.17),
\[ K = \frac{E}{3(1 - 2\nu)} \quad (3.22) \]

#### 3.2.4.2 Shear modulus in terms of Young’s modulus and Poisson’s ratio

The relationship between $\mu$ and $E$ can be appreciated by considering the shear deformation of a rectangular prism that is infinitely long in one dimension and has a square cross-section in the plane of deformation. The shear causes shortening of one diagonal and extension of the other. Let the length of the side of the square be $a$ (Fig. 3.7a) and that of its diagonal be $d_0 (= a\sqrt{2})$. The small shear through the angle $\phi$ displaces one corner by the amount ($a\tan \phi$) and stretches the diagonal to the new length $d$ (Fig. 3.7b), which is given by Pythagoras’ theorem:
\[ d^2 = a^2 + (a + a\tan \phi)^2 = a^2 + a^2 + a^2 \tan^2 \phi + 2a^2 \tan \phi = 2a^2 \left( 1 + \tan \phi + \frac{1}{2} \tan^2 \phi \right) \]
The extension of the diagonal is $d$. 

(b) When the square is deformed by shear

\[
\frac{\Delta d}{d_0} = \frac{d - d_0}{d_0} = \frac{\phi}{2}
\]

where for an infinitesimally small strain tan $\phi = \phi$, and powers of $\phi$ higher than first order are negligibly small. The extension of the diagonal is

\[
\frac{\Delta d}{d_0} = \frac{d - d_0}{d_0} = \frac{\phi}{2}
\]

This extension is related to the normal stresses $\sigma_{xx}$ and $\sigma_{yy}$ in the $x$-$y$ plane of the cross-section (Fig. 3.8a), which are in general unequal. Let $p$ represent their average value: $p = (\sigma_{xx} + \sigma_{yy})/2$. The change of shape of the square cross-section results from the differences $\Delta p$ between $p$ and $\sigma_{xx}$ and $\sigma_{yy}$, respectively (Fig. 3.8b). The outwards stress difference $\Delta p$ along the $x$-axis produces an $x$-extension equal to $\Delta p/E$, while the inwards stress difference along the $y$-axis causes contraction along the $y$-axis and a corresponding contribution to the $x$-extension equal to $\nu(\Delta p/E)$, where $\nu$ is Poisson’s ratio as before. The total $x$-extension $\Delta x/x$ is therefore given by

\[
\frac{\Delta x}{x} = \frac{\Delta p}{E}(1 + \nu)
\]

Let each edge of the square represent an arbitrary area $A$ normal to the plane of the figure. The stress differences $\Delta p$ produce forces $f = \Delta p A$ on the edges of the square, which resolve to shear forces $f/\sqrt{2}$ parallel to the sides of the inner square defined by joining the mid-points of the sides of the original square (Fig. 3.8c). Normal to the plane of the figure the surface area represented by each inner side is $A/\sqrt{2}$, and therefore the tangential (shear) stress acting on these sides simply equals $\Delta p$ (Fig. 3.8d). The inner square shears through an angle $\phi$, and so we can write

\[
\Delta p = \mu \phi
\]

One diagonal becomes stretched in the $x$-direction while the other diagonal is shortened in the $y$-direction. The extension of the diagonal of a sheared square was shown above to be $\phi/2$. Thus,

\[
\frac{\Delta p}{E}(1 + \nu) = \frac{\phi}{2} - \frac{\Delta p}{2\mu}
\]

Rearranging terms we get the relationship between $\mu$, $E$ and $\nu$.

\[
\mu = \frac{E}{2(1 + \nu)}
\]

### 3.2.4.3 The Lamé constants

The first line of Eq. (3.19) can be rewritten as

\[
E \varepsilon_{xx} = (1 + \nu)\sigma_{xx} - \nu(\sigma_{xx} + \sigma_{yy} + \sigma_{zz})
\]

and from Eq. (3.20) we have

\[
(\sigma_{xx} + \sigma_{yy} + \sigma_{zz}) = \frac{E}{(1 - 2\nu)}(\varepsilon_{xx} + \varepsilon_{yy} + \varepsilon_{zz})
\]

\[
= \frac{E}{(1 - 2\nu)}\theta
\]

where $\theta$ is the dilatation, as defined in Eq. (3.8). After substituting Eq. (3.30) in Eq. (3.29) and rearranging we get

\[
\sigma_{xx} = \frac{\nu E}{(1 + \nu)(1 - 2\nu)}\theta + \frac{E}{(1 + \nu)}\varepsilon_{xx}
\]

Writing

\[
\lambda = \frac{\nu E}{(1 + \nu)(1 - 2\nu)}
\]

and substituting from Eq. (3.28) we can write Eq. (3.31) in the simpler form

\[
\sigma_{xx} = \lambda \theta + 2\mu \varepsilon_{xx}
\]

with similar expressions for $\sigma_{yy}$ and $\sigma_{zz}$.

The constants $\lambda$ and $\mu$ are known as the Lamé constants. They are related to the elastic constants defined physically above. $\mu$ is equivalent to the rigidity modulus, while the bulk modulus $K$, Young’s modulus $E$ and Poisson’s ratio $\nu$ can each be expressed in terms of both $\lambda$ and $\mu$ (Box 3.1).

### 3.2.4.4 Anisotropy

The foregoing discussion treats the elastic parameters as constants. In fact they are dependent on pressure and temperature and so can only be considered constant for specified conditions. The variations of temperature and pressure in the Earth ensure that the elastic parameters vary with depth. Moreover, it has been assumed that the relationships between stress and strain hold equally for all directions, a property called isotropy. This condition is not fulfilled in many minerals. For example, if a mineral has uniaxial symmetry in the arrangement of the atoms in its unit cell, the physical properties of the mineral parallel and perpendicular to the axis of symmetry are different. The mineral is anisotropic. The relations between components of stress and strain in an anisotropic substance are
Fig. 3.8 (a) Unequal normal stresses \( \sigma_{xx} \) and \( \sigma_{yy} \) in the \( x-y \) plane, and their average value \( p \). (b) Stress differences \( \Delta p \) between \( p \) and \( \sigma_{xx} \) and \( \sigma_{yy} \) respectively, cause elongation parallel to \( x \) and shortening parallel to \( y \). (c) Forces \( T = \Delta p A \) along the sides of the original square give shear forces \( f/\sqrt{2} \) along the edges of the inner square, each of which has area \( A/2 \). (d) The shear stress on each side of the inner square has value \( \Delta p \) and causes extension of the diagonal of the inner square and shear deformation through an angle \( \phi \).

3.2.5 Imperfect elasticity in the Earth

A seismic wave passes through the Earth as an elastic disturbance of very short duration lasting only some seconds or minutes. Elasticity theory is used to explain seismic wave propagation. However, materials may react differently to brief, sudden stress than they do to long-lasting steady stress. The stress response of rocks and minerals in the Earth is affected by various factors, including temperature, hydrostatic confining pressure, and time. As a result, elastic, anelastic and plastic behavior occur with various degrees of importance at different depths.

Anelastic behavior in the Earth is related to the petrophysical properties of rocks and minerals. If a material is not perfectly elastic, a seismic wave passing through it loses energy to the material (e.g., as frictional heating) and the amplitude of the wave gradually diminishes. The decrease in amplitude is called *attenuation*, and it is due to anelastic *damping* of the vibration of particles of the material (see Section 3.3.2.7). For example, the passage of seismic waves through the asthenosphere is damped owing...
3.2 ELASTICITY THEORY

Box 3.1: Elastic parameters in terms of the Lamé constants

1. Bulk modulus ($K$)

The bulk modulus describes volumetric shape changes of a material under the effects of the normal stresses $\sigma_{xx}$, $\sigma_{yy}$, and $\sigma_{zz}$. Writing Hooke’s law for each normal stress gives

$$\sigma_{xx} = \lambda \theta + 2\mu e_{xx}$$  \hspace{1cm} (1a)
$$\sigma_{yy} = \lambda \theta + 2\mu e_{yy}$$  \hspace{1cm} (1b)
$$\sigma_{zz} = \lambda \theta + 2\mu e_{zz}$$  \hspace{1cm} (1c)

Adding equations (1a), (1b), and (1c) together gives

$$\sigma_{xx} + \sigma_{yy} + \sigma_{zz} = 3\lambda \theta + 2\mu (e_{xx} + e_{yy} + e_{zz})$$  \hspace{1cm} (2)

The dilatation $\theta$ is defined by Eq. (3.8) as

$$(e_{xx} + e_{yy} + e_{zz}) = 0$$  \hspace{1cm} (3)

For hydrostatic conditions we can write $\sigma_{xx} = \sigma_{yy} = \sigma_{zz} = -\rho$ and substitute in Eq. (2), which can now be rearranged in the form

$$-3\rho = 3\lambda \theta + 2\mu \theta$$  \hspace{1cm} (4)

Using the definition of the bulk modulus as $K = -\rho/\theta$, we get the result

$$K = \lambda + \frac{2\mu}{3}$$  \hspace{1cm} (5)

2. Young’s modulus ($E$)

Young’s modulus describes the longitudinal strains when a uniaxial normal stress is applied to a material. When only the longitudinal stress $\sigma_{xx}$ is applied (i.e., $\sigma_{yy} = \sigma_{zz} = 0$), Hooke’s law becomes

$$\sigma_{xx} = \lambda \theta + 2\mu e_{xx}$$  \hspace{1cm} (6a)
$$0 = \lambda \theta + 2\mu e_{yy}$$  \hspace{1cm} (6b)
$$0 = \lambda \theta + 2\mu e_{zz}$$  \hspace{1cm} (6c)

Adding equations (6a), (6b), and (6c) together gives

$$\sigma_{xx} = 3\lambda \theta + 2\mu (e_{xx} + e_{yy} + e_{zz}) = 3\lambda \theta + 2\mu \theta$$  \hspace{1cm} (7)
$$\sigma_{xx} = (3\lambda + 2\mu) \theta$$  \hspace{1cm} (8)
$$0 = \frac{\sigma_{xx}}{3\lambda + 2\mu}$$  \hspace{1cm} (9)

This expression is now substituted in Eq. (6a), which becomes

$$\sigma_{xx} = \lambda \frac{\sigma_{xx}}{3\lambda + 2\mu} + 2\mu e_{xx}$$  \hspace{1cm} (10)

Gathering and rearranging terms gives the following succession:

$$\sigma_{xx} \left(1 - \frac{\lambda}{3\lambda + 2\mu}\right) = 2\mu e_{xx}$$  \hspace{1cm} (11)
$$\sigma_{xx} \left(\frac{\lambda + \mu}{3\lambda + 2\mu}\right) = \mu e_{xx}$$  \hspace{1cm} (12)
$$\sigma_{xx} = \mu \left(\frac{3\lambda + 2\mu}{\lambda + \mu}\right) e_{xx}$$  \hspace{1cm} (13)

The definition of Young’s modulus is $E = \sigma_{xx}/e_{xx}$ and so in terms of the Lamé constants

$$E = \mu \left(\frac{3\lambda + 2\mu}{\lambda + \mu}\right)$$  \hspace{1cm} (14)

3. Poisson’s ratio ($\nu$)

Poisson’s ratio is defined as $\nu = -e_{yy}/e_{xx} = -e_{zz}/e_{xx}$. It relates the bulk modulus $K$ and Young’s modulus $E$ as developed in Eq. (3.22):

$$K = \frac{E}{3(1-2\nu)}$$  \hspace{1cm} (15)

Substituting the expressions derived above for $K$ and $E$ we get

$$\frac{3\lambda + 2\mu}{3} = \frac{1}{3(1-2\nu)^2} \left(\frac{3\lambda + 2\mu}{\lambda + \mu}\right)$$  \hspace{1cm} (16)

Rearranging terms leads to the expression for Poisson’s ratio $\nu$ in terms of the Lamé constants:

$$\frac{\lambda + \mu}{\mu} = \frac{1}{(1-2\nu)}$$  \hspace{1cm} (17)
$$1 - 2\nu = \frac{\mu}{\lambda + \mu}$$  \hspace{1cm} (18)
$$\nu = \frac{\lambda}{2(\lambda + \mu)}$$  \hspace{1cm} (19)

The values of $\lambda$ and $\mu$ are almost equal in some materials, and it is possible to assume $\lambda = \mu$, from which it follows that $\nu = 0.25$. This approximation is called Poisson’s relation; it applies to most rocks in the Earth.

A material that reacts elastically to a sudden stress may deform and flow plastically under a stress that acts over a long time interval. Plastic behavior in the asthenosphere and in the deeper mantle may allow material to

to anelastic behavior at the grain level of the minerals. This may consist of time-dependent slippage between grains; alternatively, fluid phases may be present at the grain boundaries.
flow, perhaps due to the motion of dislocations within crystal grains. The flow takes place over times on the order of hundreds of millions of years, but it provides an efficient means of transporting heat out of the deep interior.

### 3.3 SEISMIC WAVES

#### 3.3.1 Introduction

The propagation of a seismic disturbance through a heterogeneous medium is extremely complex. In order to derive equations that describe the propagation adequately, it is necessary to make simplifying assumptions. The heterogeneity of the medium is often modelled by dividing it into parallel layers, in each of which homogeneous conditions are assumed. By suitable choice of the thickness, density and elastic properties of each layer, the real conditions can be approximated. The most important assumption about the propagation of a seismic disturbance is that it travels by elastic displacements in the medium. This condition certainly does not apply close to the seismic source. In or near an earthquake focus or the shot point of a controlled explosion the medium is destroyed. Particles of the medium are displaced permanently from their neighbors; the deformation is anelastic. However, when a seismic disturbance has travelled some distance away from its source, its amplitude decreases and the medium deforms elastically to permit its passage. The particles of the medium carry out simple harmonic motions, and the seismic energy is transmitted as a complex set of wave motions.

When seismic energy is released suddenly at a point P near the surface of a homogeneous medium (Fig. 3.9), part of the energy propagates through the body of the medium as seismic body waves. The remaining part of the seismic energy spreads out over the surface as a seismic surface wave, analogous to the ripples on the surface of a pool of water into which a stone has been thrown.

#### 3.3.2 Seismic body waves

When a body wave reaches a distance \( r \) from its source in a homogeneous medium, the wavefront (defined as the surface in which all particles vibrate with the same phase) has a spherical shape, and the wave is called a spherical wave. As the distance from the source increases, the curvature of the spherical wavefront decreases. At great distances from the source the wavefront is so flat that it can be considered to be a plane and the seismic wave is called a plane wave. The direction perpendicular to the wavefront is called the seismic ray path. The description of the harmonic motion in plane waves is simpler than for spherical waves, because for plane waves we can use orthogonal Cartesian coordinates. Even for plane waves the mathematical description of the three-dimensional displacements of the medium is fairly complex. However, we can learn quite a lot about body-wave propagation from a simpler, less rigorous description.

#### 3.3.2.1 Compressional waves

Let Cartesian reference axes be defined such that the \( x \)-axis is parallel to the direction of propagation of the plane wave; the \( y \)- and \( z \)-axes then lie in the plane of the wavefront (Fig. 3.10). A generalized vibration of the medium can be reduced to components parallel to each of the reference axes. In the \( x \)-direction the particle motion is back and forward parallel to the direction of propagation. This results in the medium being alternately stretched and condensed in this direction (Fig. 3.11a). This harmonic motion produces a body wave that is transmitted as a sequence of rarefactions and condensations parallel to the \( x \)-axis.

Consider the disturbance of the medium shown in Fig. 3.11b. The area of the wavefront normal to the \( x \)-direction is \( A_x \), and the wave propagation is treated as one-dimensional. At an arbitrary position \( x \) (Fig. 3.11c), the passage of the wave produces a displacement \( u \) and a force \( F_x \) in the \( x \)-direction. At the position \( x + dx \) the displacement is \( u + du \) and the force is \( F_x + dF_x \). Here \( dx \) is the infinitesimal length of a small volume element which has mass \( \rho \, dx \, A_x \). The net force acting on this element in the \( x \)-direction is given by

\[
(F_x + dF_x) - F_x = dF_x = \frac{\partial F_x}{\partial x} dx
\]

(3.33)

The force \( F_x \) is caused by the stress element \( \sigma_{xx} \) acting on the area \( A_x \), and is equal to \( \sigma_{xx} \, A_x \). This allows us to write the one-dimensional equation of motion

\[
(\rho \, dx \, A_x) \frac{\partial^2 u}{\partial t^2} = dx \, A_x \frac{\partial \sigma_{xx}}{\partial x}
\]

(3.34)
The definitions of Young's modulus, \( E \) (Eq. (3.15)) and the normal strain \( e_{xx} \) (Eq. (3.4)) give, for a one-dimensional deformation

\[
\sigma_{xx} = E e_{xx} = \frac{E x u}{\partial x}
\]  

(3.35)

Substitution of Eq. (3.35) into Eq. (3.34) gives the one-dimensional wave equation

\[
\frac{\partial^2 u}{\partial t^2} = V^2 \frac{\partial^2 u}{\partial x^2}
\]  

(3.36)

where \( V \) is the velocity of the wave, given by

\[ V = \sqrt{\frac{E}{\rho}} \]  

(3.37)

A one-dimensional wave is rather restrictive. It represents the stretching and compressing in the \( x \)-direction as effects that are independent of what happens in the \( y \) - and \( z \)-directions. In an elastic solid the elastic strains in any direction are coupled to the strains in transverse directions by Poisson's ratio for the medium. A three-dimensional analysis is given in Appendix A that takes into account the simultaneous changes perpendicular to the direction of propagation. In this case the area \( A_x \) can no longer be considered constant. Instead of looking at the displacements in one direction only, all three axes must be taken into account. This is achieved by analyzing the changes in volume. The longitudinal (or compressional) body wave passes through a medium as a series of dilatations and compressions. The equation of the compressional wave in the \( x \)-direction is

\[
\frac{\partial^2 u}{\partial t^2} = \alpha^2 \frac{\partial^2 u}{\partial x^2}
\]  

(3.38)

where \( \alpha \) is the wave velocity and is given by

\[ \alpha = \sqrt{\frac{\lambda + 2\mu}{\rho}} = \sqrt{\frac{K + 4/3 \mu}{\rho}}. \]  

(3.39)

The longitudinal wave is the fastest of all seismic waves. When an earthquake occurs, this wave is the first to arrive at a recording station. As a result it is called the primary wave, or \( P \)-wave. Eq. 3.39 shows that \( P \)-waves can travel through solids, liquids and gases, all of which are compressible (\( K \neq 0 \)). Liquids and gases do not allow shear. Consequently, \( \mu = 0 \), and the compressional wave velocity in a liquid or gas is given by

\[ \alpha = \sqrt{\frac{K}{\rho}} \]  

(3.40)

### 3.3.2.2 Transverse waves

The vibrations along the \( y \)- and \( z \)-axes (Fig. 3.10) are parallel to the wavefront and transverse to the direction of
propagation. If we wish, we can combine the \( y \)- and \( z \)-components into a single transverse motion. It is more convenient, however, to analyze the motions in the vertical and horizontal planes separately. Here we discuss the disturbance in the vertical plane defined by the \( x \)- and \( z \)-axes; an analogous description applies to the horizontal plane.

The transverse wave motion is akin to that seen when a rope is shaken. Vertical planes move up and down and adjacent elements of the medium experience shape distortions (Fig. 3.12a), changing repeatedly from a rectangle to a parallelogram and back. Adjacent elements of the medium suffer vertical shear.

Consider the distortion of an element bounded by vertical planes separated by a small horizontal distance \( dx \) (Fig. 3.12b) at an arbitrary horizontal position \( x \). The passage of a wave in the \( x \)-direction produces a displacement \( w \) and a force \( F_x \) in the \( z \)-direction. At the position \( x + dx \) the displacement is \( w + dw \) and the force is \( F_x + dF_x \). The mass of the small volume element bounded by the vertical planes is \( \rho \ dx A_z \), where \( A_z \) is the area of the bounding plane. The net force acting on this element in the \( z \)-direction is given by

\[
(F_z + dF_z) - F_z = dF_x = \frac{\partial F_x}{\partial x} \ dx
\]

The force \( F_z \) arises from the shear stress \( \sigma_{xz} \) on the area \( A_z \), and is equal to \( \sigma_{xz} A_z \). The equation of motion of the vertically sheared element is

\[
\left( \rho dx A_z \right) \frac{\partial^2 w}{\partial t^2} = \frac{\partial}{\partial x} \left( \sigma_{xz} \frac{\partial w}{\partial x} \right)
\]

We now have to modify the Lamé expression for Hooke’s law and the definition of shear strain so that they apply to the passage of a one-dimensional shear wave in the \( x \)-direction. In this case, because the areas of the parallelograms between adjacent vertical planes are equal, there is no volume change. The dilatation \( \theta \) is zero, and Hooke’s law is as given in Eq. (3.16):

\[
\sigma_{xz} = 2\mu e_{xz}
\]

Following the definition of shear-strain components in Eq. (3.12) we have

\[
e_{xz} = \frac{1}{2} \left( \frac{\partial w}{\partial x} + \frac{\partial w}{\partial z} \right)
\]

For a one-dimensional shear wave there is no change in the distance \( dx \) between the vertical planes; \( dw \) and \( \partial w/\partial z \) are zero and \( e_{xz} \) is equal to \( (\partial w/\partial x)/2 \). On substitution into Eq. (3.43) this gives

\[
\sigma_{xz} = \mu \frac{\partial w}{\partial x}
\]

and on further substitution into Eq. (3.42) and rearrangement of terms we get

\[
\frac{\partial^2 w}{\partial t^2} = \beta^2 \frac{\partial^2 w}{\partial x^2}
\]

where \( \beta \) is the velocity of the shear wave, given by

\[
\beta = \sqrt{\frac{\mu}{\rho}}
\]

The only elastic property that determines the velocity of the shear wave is the rigidity or shear modulus, \( \mu \). In liquids and gases \( \mu \) is zero and shear waves cannot propagate. In solids, a quick comparison of Eqs. (3.39) and (3.47) gives

\[
\frac{\alpha^2}{\beta^2} = \frac{K}{\rho}
\]

By definition, the bulk modulus \( K \) is positive (if it were negative, an increase in confining pressure would cause an increase in volume), and therefore \( \alpha \) is always greater than \( \beta \). Shear waves from an earthquake travel more slowly than P-waves and are recorded at an observation station as later arrivals. Shear waves are often referred to as secondary waves or S-waves.

The general shear-wave motion within the plane of the wavefront can be resolved into two orthogonal components, one being horizontal and the other lying in the vertical plane containing the ray path (Fig. 3.10). Equation (3.46) describes a one-dimensional shear wave which travels in the \( x \)-direction, but which has particle displacements \( (w) \) in the \( z \)-direction. This wave can be considered
3.3 SEISMIC WAVES

to be polarized in the vertical plane. It is called the SV-wave. A similar equation describes the shear wave in the x-direction with particle displacements (v) in the y-direction. A shear wave that is polarized in the horizontal plane is called an SH-wave.

As for the description of longitudinal waves, this treatment of shear-wave transmission is over-simplified; a more rigorous treatment is given in Appendix A. The passage of a shear wave involves rotations of volume elements within the plane normal to the ray path, without changing their volume. For this reason, shear waves are also sometimes called rotational (or equivoluminal) waves. The rotation is a vector, \( \psi \), with x-, y-, and z-components given by

\[
\psi_x = \frac{\partial v_y}{\partial y} - \frac{\partial v_z}{\partial z}, \quad \psi_y = \frac{\partial v_x}{\partial x} - \frac{\partial v_z}{\partial z}, \quad \psi_z = \frac{\partial v_x}{\partial x} - \frac{\partial v_y}{\partial y}
\]  

(3.49)

A more appropriate equation for the shear wave in the x-direction is then

\[
\frac{\partial^2 \psi}{\partial t^2} = \beta \frac{\partial^2 \psi}{\partial x^2}
\]  

(3.50)

where \( \beta \) is again the shear-wave velocity as given by Eq. (3.47).

Until now we have chosen the direction of propagation along one of the reference axes so as to simplify the mathematics. If we remove this restriction, additional second-order differentiations with respect to the y- and z-coordinates must be introduced. The P-wave and S-wave equations become, respectively,

\[
\frac{\partial^2 u}{\partial t^2} = \alpha^2 \left( \frac{\partial^2 u}{\partial x^2} + \frac{\partial^2 u}{\partial y^2} + \frac{\partial^2 u}{\partial z^2} \right)
\]  

(3.51)

\[
\frac{\partial^2 \psi}{\partial t^2} = \beta^2 \left( \frac{\partial^2 \psi}{\partial x^2} + \frac{\partial^2 \psi}{\partial y^2} + \frac{\partial^2 \psi}{\partial z^2} \right)
\]  

(3.52)

3.3.2.3 The solution of the seismic wave equation

Two important characteristics of a wave motion are: (1) it transmits energy by means of elastic displacements of the particles of the medium, i.e., there is no net transfer of mass, and (2) the wave pattern repeats itself in both time and space. The harmonic repetition allows us to express the amplitude variation by a sine or cosine function. As the wave passes any point, the amplitude of the disturbance is repeated at regular time intervals, \( T \), the period of the wave. The number of times the amplitude is repeated per second is the frequency, \( f \), which is equal to the inverse of the period (\( f = 1/T \)). At any instant in time, the disturbance in the medium is repeated along the direction of travel at regular distances, \( \lambda \), the wavelength of the wave. During the passage of a P-wave in the x-direction, the harmonic displacement \((u)\) of a particle from its mean position can be written

\[
u = A \sin 2\pi \left( \frac{x}{\lambda} - \frac{t}{T} \right) \]  

(3.53)

where \( A \) is the amplitude.

The quantity in brackets is called the phase of the wave. Any value of the phase corresponds to a particular amplitude and direction of motion of the particles of the medium. The wave number \((k)\), angular frequency \((\omega)\) and velocity \((c)\) are defined and related by

\[
k = \frac{2\pi}{\lambda}, \quad \omega = 2\pi f = \frac{2\pi}{T}, \quad c = \frac{\omega}{k}
\]  

(3.54)

Equation (3.53) for the displacement \((u)\) can then be written

\[
u = A \sin(kx - \omega t) = A \sin(kx - ct)
\]  

(3.55)

The velocity \( c \) introduced here is called the phase velocity. It is the velocity with which a constant phase (e.g., the “peak” or “trough,” or one of the zero displacements) is transmitted. This can be seen by equating the phase to a constant and then differentiating the expression with respect to time, as follows:

\[
kx - \omega t = \text{constant}
\]

\[
k \frac{dx}{dt} - \omega = 0
\]  

(3.56)

To demonstrate that the displacement given by Eq. (3.55) is a solution of the one-dimensional wave equation (Eq. (3.38)) we must partially differentiate \( u \) in Eq. (3.55) twice with respect to time (\( t \)) and twice with respect to position (\( x \)):

\[
\frac{\partial u}{\partial t} = A k \cos(kx - \omega t) \quad \frac{\partial^2 u}{\partial t^2} = -A k^2 \sin(kx - \omega t) = -k^2 u
\]

\[
\frac{\partial u}{\partial x} = A \omega \sin(kx - \omega t) \quad \frac{\partial^2 u}{\partial x^2} = -A \omega^2 \sin(kx - \omega t) = -\omega^2 u
\]

\[
\frac{\partial^2 u}{\partial t^2} - \frac{k^2 \partial^2 u}{\partial x^2} = c^2 \frac{\partial^2 u}{\partial x^2}
\]  

(3.57)

For a P-wave travelling along the x-axis the dilatation \( \theta \) is given by an equation similar to Eq. (3.57), with substitution of the P-wave velocity \( (\alpha) \) for the velocity \( c \). Similarly, for an S-wave along the x-axis the rotation \( \psi \) is given by an equation like Eq. (3.57) with appropriate substitutions of \( \psi \) for \( u \) and the S-wave velocity \( (\beta) \) for the velocity \( c \). However, in general, the solutions of the three-dimensional compressional and shear wave equations (Eqs. (3.51) and (3.51), respectively) are considerably more complicated than those given by Eq. (3.55).

3.3.2.4 D’Alembert’s principle

Equation (3.55) describes the particle displacement during the passage of a wave that is travelling in the direction of the positive x-axis with velocity \( c \). Because the velocity enters the wave equation as \( c^2 \), the one-dimensional wave equation is also satisfied by the displacement

\[
u = B \sin(k(x + ct))
\]  

(3.58)
which corresponds to a wave travelling with velocity $c$ in the direction of the negative $x$-axis.

In fact, any function of $(x \pm ct)$ that is itself continuous and that has continuous first and second derivatives is a solution of the one-dimensional wave equation. This is known as D'Alembert's principle. It can be simply demonstrated for the function $F(x-ct) = f(\phi)$ as follows:

$$\frac{\partial F}{\partial x} = \frac{\partial F}{\partial \phi} \frac{\partial \phi}{\partial x} = \frac{\partial F}{\partial \phi}$$

$$\frac{\partial^2 F}{\partial x^2} = \frac{\partial^2 F}{\partial \phi^2} \left(\frac{\partial \phi}{\partial x}\right)^2$$

$$\frac{\partial^2 F}{\partial t^2} = -c^2 \frac{\partial^2 F}{\partial \phi^2}$$

$$\frac{\partial^2 F}{\partial x^2} - \frac{\partial^2 F}{\partial t^2} = \frac{\partial^2 F}{\partial \phi^2}$$

(3.60)

Because Eq. (3.60) is valid for positive and negative values of $c$, its general solution $F$ represents the superposition of waves travelling in opposite directions along the $x$-axis, and is given by

$$F = f(x-ct) + g(x+ct)$$

(3.61)

### 3.3.2.5 The eikonal equation

Consider a wave travelling with constant velocity $c$ along the axis $x'$ which has direction cosines $(l, m, n)$. If $x'$ is measured from the center of the coordinate axes $(x, y, z)$ we can substitute $x' = lx + my + nz$ for $x$ in Eq. (3.60). If we consider for convenience only the wave travelling in the direction of $+x'$, we get as the general solution to the wave equation

$$F = f(lx + my + nz - ct)$$

(3.62)

The wave equation is a second-order differential equation. However, the function $F$ is also a solution of a first-order differential equation. This is seen by differentiating $F$ with respect to $x, y, z,$ and $t$, respectively, which gives

$$\frac{\partial F}{\partial x} = \frac{\partial F}{\partial \phi} \frac{\partial \phi}{\partial x} = \frac{\partial F}{\partial \phi}$$

$$\frac{\partial^2 F}{\partial x^2} = \frac{\partial^2 F}{\partial \phi^2} \left(\frac{\partial \phi}{\partial x}\right)^2$$

$$\frac{\partial^2 F}{\partial t^2} = -c^2 \frac{\partial^2 F}{\partial \phi^2}$$

(3.62)

The direction cosines $(l, m, n)$ are related by $l^2 + m^2 + n^2 = 1$, and so, as can be verified by substitution, the expressions in Eq. (3.62) satisfy the equation

$$\left(\frac{\partial F}{\partial x}\right)^2 + \left(\frac{\partial F}{\partial y}\right)^2 + \left(\frac{\partial F}{\partial z}\right)^2 = \left(\frac{1}{c}\right)^2 \left(\frac{\partial F}{\partial t}\right)^2$$

(3.63)

In seismic wave theory, the progress of a wave is described by successive positions of its wavefront, defined as the surface in which all particles at a given instant in time are moving with the same phase. For a particular value of $t$ a constant phase of the wave equation solution given by Eq. (3.61) requires that

$$lx + my + nz = \text{constant}$$

(3.64)

From analytical geometry we know that Eq. (3.64) represents a family of planes perpendicular to a line with direction cosines $(l, m, n)$. We began this discussion by describing a wave moving with velocity $c$ along the direction $x'$, and now we see that this direction is normal to the plane wavefronts. This is the direction that we defined earlier as the ray path of the wave.

In a medium like the Earth the elastic properties and density -- and therefore also the velocity -- vary with position. The ray path is no longer a straight line and the wavefronts are not planar. Instead of Eq. (3.61) we write

$$F = f[S(x,y,z) - ct]$$

(3.65)

where $S(x, y, z)$ is a function of position only and $c_0$ is a constant reference velocity. Substitution of Eq. (3.65) into Eq. (3.63) gives

$$\left(\frac{\partial S}{\partial x}\right)^2 + \left(\frac{\partial S}{\partial y}\right)^2 + \left(\frac{\partial S}{\partial z}\right)^2 = \left(\frac{c_0}{\xi}\right)^2 = \xi^2$$

(3.66)

where $\xi$ is known as the refractive index of the medium. Equation (3.66) is called the eikonal equation. It establishes the equivalence of treating seismic wave propagation by describing the wavefronts or the ray paths. The surfaces $S(x, y, z) = \text{constant}$ represent the wavefronts (no longer planar). The direction cosines of the ray path (normal to the wavefront) are in this case given by

$$\lambda = \xi \frac{\partial S}{\partial x} \quad \mu = \xi \frac{\partial S}{\partial y} \quad v = \xi \frac{\partial S}{\partial z}$$

(3.67)

### 3.3.2.6 The energy in a seismic disturbance

It is important to distinguish between the velocity with which a seismic disturbance travels through a material and the speed with which the particles of the material vibrate during the passage of the wave. The vibrational speed ($v_p$) is obtained by differentiating Eq. (3.55) with respect to time, which yields

$$v_p = \frac{\partial u}{\partial t} = -\omega A \cos(kx - \omega t)$$

(3.68)

The intensity or energy density of a wave is the energy per unit volume in the wavefront and consists of kinetic and potential energy. The kinetic part is given by

$$I = \frac{1}{2} \rho v_p^2 = \frac{1}{2} \rho A^2 \omega^2 \cos^2(kx - \omega t)$$

(3.69)

The energy density averaged over a complete harmonic cycle consists of equal parts of kinetic and potential energy; it is given by...
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\[ I_{av} = \frac{1}{4} \rho v^2 A^2 \]  
(3.70)

i.e., the mean intensity of the wave is proportional to the square of its amplitude.

3.3.2.7 Attenuation of seismic waves

The further a seismic signal travels from its source the weaker it becomes. The decrease of amplitude with increasing distance from the source is referred to as attenuation. It is partly due to the geometry of propagation of seismic waves, and partly due to anelastic properties of the material through which they travel.

The most important reduction is due to geometric attenuation. Consider the seismic body waves generated by a seismic source at a point P on the surface of a uniform half-space (see Fig. 3.9). If there is no energy loss due to friction, the energy \( E_s \) in the wavefront at distance \( r \) from its source is distributed over the surface of a hemisphere with area \( 2\pi r^2 \). The intensity (or energy density, \( I_s \)) of the body waves is the energy per unit area of the wavefront, and at distance \( r \) is:

\[ I_s(r) = \frac{E_s}{2\pi r^2} \]  
(3.71)

The surface wave is constricted to spread out laterally. The disturbance affects not only the free surface but extends downwards into the medium to a depth \( d \), which we can consider to be constant for a given wave (Fig. 3.9). When the wavefront of a surface wave reaches a distance \( r \) from the source, the initial energy \( E_s \) is distributed over a circular cylindrical surface with area \( 2\pi rd \). At a distance \( r \) from its source the intensity of the surface wave is given by:

\[ I_s(r) = \frac{E_s}{2\pi rd} \]  
(3.72)

These equations show that the decrease in intensity of body waves is proportional to \( 1/r^2 \) while the decrease in surface wave intensity is proportional to \( 1/r \). As shown in Eq. (3.70), the intensity of a wave-form, or harmonic vibration, is proportional to the square of its amplitude. The corresponding amplitude attenuations of body waves and surface waves are proportional to \( 1/r \) and \( 1/\sqrt{r} \), respectively. Thus, seismic body waves are attenuated more rapidly than surface waves with increasing distance from the source. This explains why, except for the records of very deep earthquakes that do not generate strong surface waves, the surface-wave train on a seismogram is more prominent than that of the body waves.

Another reason for attenuation is the absorption of energy due to imperfect elastic properties. If the particles of a medium do not react perfectly elastically with their neighbors, part of the energy in the wave is lost (reappearing, for example, as frictional heat) instead of being transferred through the medium. This type of attenuation of the seismic wave is referred to as anelastic damping.

The damping of seismic waves is described by a parameter called the quality factor \( Q \), a concept borrowed from electric circuit theory where it describes the performance of an oscillatory circuit. It is defined as the fractional loss of energy per cycle

\[ \frac{2\pi}{Q} = -\frac{\Delta E}{E} \]  
(3.73)

In this expression \( \Delta E \) is the energy lost in one cycle and \( E \) is the total elastic energy stored in the wave. If we consider the damping of a seismic wave as a function of the distance that it travels, a cycle is represented by the wavelength \( (\lambda) \) of the wave. Equation (3.73) can be rewritten for this case as

\[ \frac{2\pi}{Q} = -\frac{1}{E} \frac{dE}{dr} \]

\[ \frac{dE}{E} = -\frac{2\pi r}{Q \lambda} \]  
(3.74)

It is conventional to measure damping by its effect on the amplitude of a seismic signal, because that is what is observed on a seismic record. We have seen that the energy in a wave is proportional to the square of its amplitude \( A \) (Eq. (3.70)). Thus we can write \( dE/E = 2dA/A \) in Eq. (3.74), and on solving we get the damped amplitude of a seismic wave at distance \( r \) from its source:

\[ A = A_0 \exp \left( -\frac{\pi r}{Q \lambda} \right) \]

\[ = A_0 \exp \left( -\frac{r}{D} \right) \]  
(3.75)

In this equation \( D \) is the distance within which the amplitude falls to \( 1/e \) (36.8%, or roughly a third) of its original value. The inverse of this distance \( (D^{-1}) \) is called the absorption coefficient. For a given wavelength, \( D \) is proportional to the \( Q \)-factor of the region through which the wave travels. A rock with a high \( Q \)-factor transmits a seismic wave with relatively little energy loss by absorption, and the distance \( D \) is large. For body waves \( D \) is generally of the order of 10,000 km and damping of the waves by absorption is not a very strong effect. It is slightly stronger for seismic surface waves, for which \( D \) is around 3000 km.

Equation (3.75) shows that the damping of a seismic wave is dependent on the \( Q \)-factor of the region of the Earth that the wave has travelled through. In general the \( Q \)-factor for P-waves is higher than the \( Q \)-factor for S-waves. This may indicate that anelastic damping is determined primarily by the shear component of strain. In solids with low rigidity, the shear strain can reach high levels and the damping is greater than in materials with high rigidity. In fluids the \( Q \)-factor is high and damping is low, because shear strains are zero and the seismic wave is purely compressional. The values of \( Q \) are quite variable in the Earth: values of around \( 10^2 \) are found for the mantle, and around \( 10^3 \) for P-waves in the liquid core. Because \( Q \) is a measure of the deviation from perfect elasticity, it is also encountered in the theory of natural oscillations of the...
Earth, and has an effect on fluctuations of the Earth’s free rotation, as in the damping of the Chandler wobble.

It follows from Eq. (3.75) that the absorption coefficient ($D^{-1}$) is inversely proportional to the wavelength $\lambda$. Thus the attenuation of a seismic wave by absorption is dependent upon the frequency of the signal. High frequencies are attenuated more rapidly than are lower frequencies. As a result, the frequency spectrum of a seismic signal changes as it travels through the ground. Although the original signal may be a sharp pulse (resulting from a shock or explosion), the preferential loss of high frequencies as it travels away from the source causes the signal to assume a smoother shape. This selective loss of high frequencies by absorption is analogous to removing high frequencies from a sound source using a filter. Because the low frequencies are not affected so markedly, they pass through the ground with less attenuation. The ground acts as a low-pass filter to seismic signals.

### 3.3.3 Seismic surface waves

A disturbance at the free surface of a medium propagates away from its source partly as seismic surface waves. Just as seismic body waves can be classified as P- or S-waves, these waves are subdivided into Rayleigh waves ($L_R$) and Love waves ($L_Q$), which are distinguished from each other by the types of particle motion in their wavefronts. In the description of body waves, the motion of particles in the wavefront was resolved into three orthogonal components – a longitudinal vibration parallel to the ray path (the P-wave motion), a transverse vibration in the vertical plane containing the ray path (the vertical shear or SV-wave) and a horizontal transverse vibration (the horizontal shear or SH-wave). These components of motion, restricted to surface layers, also determine the particle motion and character of the two types of surface waves.

#### 3.3.3.1 Rayleigh waves ($L_R$)

In 1885 Lord Rayleigh described the propagation of a surface wave along the free surface of a semi-infinite elastic half-space. The particles in the wavefront of the Rayleigh wave are polarized to vibrate in the vertical plane. The resulting particle motion can be regarded as a combination of the P- and SV-vibrations. If the direction of propagation of the Rayleigh wave is to the right of the viewer (as in Fig. 3.13), the particle motion describes a retrograde ellipse in the vertical plane with its major axis vertical and minor axis in the direction of wave propagation. If Poisson’s relation holds for a solid (i.e., Poisson’s ratio $\nu = 0.25$) the theory of Rayleigh waves gives a speed ($V_{LR}$) equal to $\sqrt{(2 - 2/\sqrt{3})} = 0.9194$ of the speed ($\beta$) of S-waves (i.e., $V_{LR} = 0.9194\beta$). This is approximately the case in the Earth.

The particle displacement is not confined entirely to the surface of the medium. Particles below the free surface are also affected by the passage of the Rayleigh wave; in a uniform half-space the amplitude of the particle displacement decreases exponentially with increasing depth. The penetration depth of the surface wave is typically taken to be the depth at which the amplitude is attenuated to (e$^{-1}$) of its value at the surface. For Rayleigh waves with wavelength $\lambda$ the characteristic penetration depth is about 0.4$\lambda$.

#### 3.3.3.2 Love waves ($L_Q$)

The boundary conditions which govern the components of stress at the free surface of a semi-infinite elastic half-space prohibit the propagation of SH-waves along the surface. However, A. E. H. Love showed in 1911 that if a horizontal layer lies between the free surface and the semi-infinite half-space (Fig. 3.14a), SH-waves within the layer that are reflected at supercritical angles (see Section 3.6) from the top and bottom of the layer can interfere constructively to give a surface wave with horizontal particle motions (Fig. 3.14b). The velocity ($\beta_L$) of S-waves in the near-surface layer must be lower than in the underlying half-space ($\beta_S$). The velocity of the Love waves ($V_{LQ}$) lies between the two extreme values: $\beta_S < V_{LQ} < \beta_S$.

Theory shows that the speed of Love waves with very short wavelengths is close to the slower velocity $\beta_S$ of the upper layer, while long wavelengths travel at a speed close to the faster velocity $\beta_L$ of the lower medium. This dependence of velocity on wavelength is termed dispersion. Love waves are always dispersive, because they can only propagate in a velocity-layered medium.

#### 3.3.3.3 The dispersion of surface waves

The dispersion of surface waves provides an important tool for determining the vertical velocity structure of the lower crust and upper mantle. Love waves are intrinsically...
dispersive even when the surface layer and underlying half-space are uniform. Rayleigh waves over a uniform half-space are non-dispersive. However, horizontal layers with different velocities are usually present or there is a vertical velocity gradient. Rayleigh waves with long wavelengths penetrate more deeply into the Earth than those with short wavelengths. The speed of Rayleigh waves is proportional to the shear-wave velocity \( V_{LR} \approx 0.92\beta \), and in the crust and uppermost mantle \( \beta \) generally increases with depth. Thus, the deeper penetrating long wavelengths travel with the phase velocity \( c \) faster than the short wavelengths (i.e., the longer wavelengths propagate as the envelope of the wave packet due to normal dispersion as the faster-moving long wavelengths pass through the packet). At large distances from the source, the long wavelengths arrive as the first part of the surface-wave record at large distances from the seismic source.

The situation in which phase velocity increases with increasing wavelength (i.e., the longer wavelengths propagate faster than the short wavelengths) is called normal dispersion. In this case, because \( \beta \) is positive, the group velocity \( U \) is slower than the phase velocity \( c \). The shape of the wave packet changes systematically as the faster moving long wavelengths pass through the packet (Fig. 3.15b). As time elapses, an initially concentrated pulse becomes progressively stretched out into a long train of waves. Consequently, over a medium in which velocity increases with depth, the long wavelengths arrive as the first part of the surface-wave record at large distances from the seismic source.

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**Fig. 3.14** In a Love wave the particle motion is horizontal and perpendicular to the direction of propagation. The amplitude of the wave decreases with depth below the free surface.

**Fig. 3.15** (a) The surface-wave energy propagates as the envelope of the wave packet with the group velocity \( U \), while the individual wavelengths travel with the phase velocity \( c \). (b) Change of shape of a wave packet due to normal dispersion as the faster-moving long wavelengths pass through the packet. At large distances from the source, the long wavelengths arrive as the first part of the surface-wave train (modified from Telford et al., 1990).

### 3.3.4 Free oscillations of the Earth

When a bell is struck with a hammer, it vibrates freely at a number of natural frequencies. The combination of natural oscillations that are excited gives each bell its particular sonority. In an analogous way, the sudden release of energy in a very large earthquake can set the entire Earth into vibration, with natural frequencies of oscillation that are determined by the elastic properties and structure of the Earth’s interior. The free oscillations involve three-dimensional deformation of the Earth’s spherical shape and can be quite complex. Before discussing the Earth’s free oscillations it is worth reviewing some concepts of vibrating systems that can be learned from the one-dimensional excitation of a vibrating string that is fixed at both ends.

Any complicated vibration of the string can be represented by the superposition of a number of simpler vibrations, called the normal modes of vibration. These arise when travelling waves reflected from the boundaries at the ends of the string interfere with each other to give a standing wave. Each normal mode corresponds to a particular sonority. In an analogous way, the sudden release of energy peak
mode of vibration has no nodes. The second normal mode (sometimes called the first overtone) has one node; its wavelength and period are half those of the fundamental mode. The third normal mode (second overtone) has three times the frequency of the first mode, and so on. Modes with one or more node are called higher-order modes.

The concepts of modes and nodes are also applicable to a vibrating sphere. The complex general vibration of a sphere can be resolved into the superposition of a number of normal modes. The nodes of zero displacement become nodal surfaces on which the amplitude of the vibration is zero. The free oscillations of the Earth can be divided into three categories. In radial oscillations the displacements are purely radial, in spheroidal oscillations they are partly radial and partly tangential, and in toroidal oscillations they are purely tangential.

3.3.4.1 Radial oscillations

The simplest kind of free oscillations are the radial oscillations, in which the shape of the Earth remains “spherical” and all particles vibrate purely radially (Fig. 3.17a). In the fundamental mode of this type of oscillation the entire Earth expands and contracts in unison with a period of about 20.5 minutes. The second normal mode (first overtone) of radial oscillations has a single internal spherical nodal surface. While the inner sphere is contracting, the part outside the nodal surface is expanding, and vice versa. The nodal surfaces of higher modes are also spheres internal to the Earth and concentric with the outer surface.

3.3.4.2 Spheroidal oscillations

A general spheroidal oscillation involves both radial and tangential displacements that can be described by spherical harmonic functions (see Box 2.3). These functions are referred to an axis through the Earth at the point of interest (e.g., an earthquake epicenter), and to a great circle which contains the axis. With respect to this reference frame they describe the latitudinal and longitudinal variations of the displacement of a surface from a sphere. They allow complete mathematical description and concise identification of each mode of oscillation with the aid of three indices. The longitudinal order \( m \) is the number of nodal lines on the sphere that are great circles, the order \( l \) is determined from the \((l - m)\) latitudinal nodal lines, and the overtone number \( n \) describes the number of internal nodal surfaces. The notation \( \ell l n \) denotes a spheroidal oscillation of order \( l \), longitudinal order \( m \), and overtone number \( n \). In practice, only oscillations with longitudinal order \( m = 0 \) (rotationally symmetric about the reference axis) are observed, and this index is usually dropped. Also, the oscillation of order \( l = 1 \) does not exist; it would have only a single equatorial nodal plane and the vibration would involve displacement of the center of gravity. The spheroidal oscillations \( \ell l 0 \) and \( \ell l 1 \) are shown in Fig. 3.17b. Spheroidal oscillations displace the Earth’s surface and alter the internal density distribution. After large earthquakes they produce records on highly sensitive gravity meters used for bodily Earth-tide observations, and also on strain gauges and tilt meters.

The radial oscillations can be regarded as a special type of spheroidal oscillation with \( l = 0 \). The fundamental radial oscillation is also the fundamental spheroidal oscillation, denoted \( \ell 0 0 \); the next higher mode is called \( 1 0 0 \) (Fig. 3.17a).

3.3.4.3 Toroidal oscillations

The third category of oscillation is characterized by displacements that are purely tangential. The spherical shape and volume of the Earth are unaffected by a toroidal oscillation, which involves only longitudinal displacements about an axis (Fig. 3.18a). The amplitude of the longitudinal displacement varies with latitude. (Note that, as for spheroidal oscillations, “latitude” and “longitude” refer to the symmetry axis and are different from their geographic definitions.) The toroidal modes have nodal planes that intersect the surface on circles of “latitude,” on which the toroidal displacement is zero. Analogously to the spheroidal oscillations, the notation \( \ell j j \) is used to describe the spatial geometry of toroidal modes. The mode \( \ell 1 j \) has zero displacement and \( \ell 0 j \) does not exist, because it describes a constant azimuthal twist of the entire Earth, which would change its angular momentum. The simplest toroidal mode is \( \ell 2 j \), in which two hemispheres oscillate in opposite senses across a single nodal plane (Fig. 3.18a). Higher toroidal modes of order \( l \) oscillate across \((l - 1)\) nodal planes perpendicular to the symmetry axis.

The amplitudes of toroidal oscillations inside the Earth change with depth. The displacements of internal
3.3 SEISMIC WAVES

spherical surfaces from their equilibrium positions are zero at internal nodal surfaces (Fig. 3.18b). The nomenclature for toroidal oscillations also represents these internal nodal surfaces. Thus, $T_1$ denotes a general toroidal mode with one internal nodal surface; the internal sphere twists in an opposite sense to the outer spherical shell; $T_2$ has two internal nodal surfaces, etc. (Fig. 3.18b).

The twisting motion in toroidal oscillations does not alter the radial density distribution in the Earth, and so they do not show in gravity meter records. They cause changes in strain and displacement parallel to the Earth’s surface and can be recorded by strain meters. Toroidal oscillations are dependent on the shear strength of the Earth’s interior. The Earth’s fluid core cannot take part in these oscillations, and they are therefore restricted to the Earth’s rigid mantle and crust.

3.3.4.4 Comparison with surface waves

The higher-order free oscillations of the Earth are related directly to the two types of surface wave.

(i) In a Rayleigh wave the particle vibration is polarized in the vertical plane and has radial and tangential components (see Fig. 3.13). The higher-order spheroidal oscillations are equivalent to the standing wave patterns that arise from the interference of trains of long-period Rayleigh waves travelling in opposite directions around the Earth.

(ii) In a Love wave the particle vibration is polarized horizontally. The toroidal oscillations may be regarded as the standing wave patterns due to the interference of oppositely travelling Love waves.

The similarity between surface waves and higher-order natural oscillations of the Earth is evident in the variations of displacement with depth. Like any vibration, a train of surface waves is made up of different modes. Theoretical analysis of surface waves shows that the amplitudes of different modes decay with depth in the Earth (Fig. 3.19) in an equivalent manner to the depth attenuation of natural oscillations (Fig. 3.18b).

The periods of the normal modes of free oscillations were calculated before they were observed. They have long periods - the period of $S_0$ is 20.5 minutes, that of $T_2$ is 43.9 minutes and that of $S_2$ is 53.9 minutes – and pendulum seismographs are not suitable for recording them. Their recognition had to await the development of long-period seismographs. The spheroidal oscillations have a radial component of displacement and can be recorded with long-period, vertical-motion seismographs. Continually recording gravimeters used for the observation of bodily Earth-tides also record the spheroidal oscillations but not the toroidal oscillations, which have no vertical component. These usually must be recorded.
with an instrument that is sensitive to horizontal displacements, such as the strain meter designed by H. Benioff in 1935. Long-period, horizontal-motion seismographs are capable of recording the toroidal oscillations induced by great earthquakes.

Strain meter records of the November 4, 1952, magnitude 8 earthquake in Kamchatka exhibited a long-period surface wave with a period of 57 minutes (Benioff, 1958). This is much longer than the known periods of travelling surface waves, and was interpreted as a free oscillation of the Earth. Several independent investigators, using bodily Earth-tide gravity meters and di

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Figure 3.20 illustrates some of the normal modes of free oscillation of the Earth set up by the huge 2004 Sumatra–Andaman Islands earthquake ($M_s = 9.0$). Vertical motions were recorded during 240 hours by the Canberra, Australia, station of the Geoscope global network of digital broadband seismic stations (see Section 3.4.3.2). The power spectrum in Fig. 3.20 is a plot of the energy associated with different frequencies of vibration. Some spheroidal and toroidal free oscillations are identified and illustrated schematically. The splitting of some normal modes is caused by the non-spherical shape and rotation of the Earth.

The study of the natural oscillations of the Earth set up by large earthquakes is an important branch of seismology, because the normal modes are strongly dependent on the Earth’s internal structure. The low-order modes are affected by the entire interior of the Earth, while the higher-order modes react primarily to movements of the upper mantle. The periods of free oscillation are determined by the radial distributions of elastic properties and densities in the Earth. Comparison of the observed periods of different modes with values computed for different models of the Earth’s velocity and density structure provides an important check on the validity of the Earth model. The oscillations are damped by the anelasticity of the Earth. The low-order mode oscillations with periods up to 40–50 minutes persist much longer than the higher-order modes with short periods of only a few minutes. By studying the decay times of different modes, a profile of the anelastic quality factor ($Q$) within the Earth can be calculated that is compatible with results obtained for body waves.

### 3.4 THE SEISMOGRAPH

#### 3.4.1 Introduction

The earliest known instrument for indicating the arrival of a seismic tremor from a distant source is reputed to have been invented by a Chinese astronomer called Chang Heng in 132 AD. The device consisted of eight inverted dragons placed at equal intervals around the rim of a vase. Under each dragon sat an open-mouthed metal toad. Each dragon held a bronze ball in its mouth. When a slight tremor shook the device, an internal mechanism opened the mouth of one dragon, releasing its bronze ball, which fell into the open mouth of the metal toad beneath, thereby marking the direction of arrival of the tremor. The principle of this instrument was used in eighteenth-century European devices that consisted of brimful bowls of water or mercury with grooved rims under which tiny collector bowls were placed to collect the overflow occasioned by a seismic tremor. These instruments gave visible evidence of a seismic event but were unable to trace a permanent record of the seismic wave itself. They are classified as seismoscopes.

The science of seismology dates from the invention of the seismograph by the English scientist John Milne in 1892. Its name derives from its ability to convert an unfelt ground vibration into a visible record. The seismograph consists of a receiver and a recorder. The ground vibration is detected and amplified by a sensor, called the seismometer or, in exploration seismology, the geophone. In modern instruments the vibration is amplified and filtered electronically. The amplified ground motion is converted to a visible record, called the seismogram.

The seismometer makes use of the principle of inertia. If a heavy mass is only loosely coupled to the ground (for example, by suspending it from a wire like a pendulum as in Fig. 3.21), the motion of the Earth caused by a seismic wave is only partly transferred to the mass. While the ground vibrates, the inertia of the heavy mass assures that it does not move as much, if at all. The seismometer amplifies and records the relative motion between the mass and the ground.

Early seismographs were undamped and reacted only to a limited band of seismic frequencies. Seismic waves
with inappropriate frequencies were barely recorded at all, but strong waves could set the instrument into resonant vibration. In 1903, the German seismologist Emil Wiechert substantially increased the accuracy of the seismograph by improving the amplification method and by damping the instrument. These early instruments relied on mechanical levers for amplification and recording signals on smoked paper. This made them both bulky and heavy, which severely restricted their application.

A major technological improvement was achieved in 1906, when Prince Boris Galitzin of Russia introduced the electromagnetic seismometer, which allowed galvanometric recording on photographic paper. This electrical method had the great advantage that the recorder could now be separated from the seismometer. The seismometer has evolved constantly, with improvements in seismometer design and recording method, culminating in modern broadband instruments with digital recording on magnetic tape, hard disk, or solid state memory device.

3.4.2 Principle of the seismometer

Seismometers are designed to react to motion of the Earth in a given direction. Mechanical instruments record the amplified displacement of the ground; electromagnetic instruments respond to the velocity of ground motion. Depending on the design, either type may respond to vertical or horizontal motion. Some modern electromagnetic instruments are constructed so as to record simultaneously three orthogonal components of motion. Most designs employ variations on the pendulum principle.

3.4.2.1 Vertical-motion seismometer

In the mechanical type of vertical-motion seismometer (Fig. 3.22a), a large mass is mounted on a horizontal bar hinged at a pivot so that it can move only in the vertical plane. A pen attached to the bar writes on a horizontal rotating drum that is fixed to the housing of the instrument. The bar is held in a horizontal position by a weak spring. This assures a loose coupling between the mass and the housing, which is connected rigidly to the ground. Vertical ground motion, as sensed during the passage of a seismic wave, is transmitted to the housing but not to the inertial mass and the pen, which remain stationary. The pen inscribes a trace of the vertical vibration of the housing on a paper fixed to the rotating drum. This trace is the vertical-motion seismogram of the seismic wave.
The electromagnetic seismometer responds to the relative motion between a magnet and a coil of wire. One of these members is fixed to the housing of the instrument and thereby to the Earth. The other is suspended by a spring and forms the inertial member. Two basic designs are possible. In the moving-magnet type, the coil is fixed to the housing and the magnet is inertial. In the moving-coil type the roles are reversed (Fig. 3.22b). A coil of wire fixed to the inertial mass is suspended between the poles of a strong magnet, which in turn is fixed to the ground by the rigid housing. Any motion of the coil within the magnetic field induces a voltage in the coil proportional to the rate of change of magnetic flux. During a seismic arrival the vibration of the ground relative to the mass is converted to an electrical voltage by induction in the coil. The voltage is amplified and transmitted through an electrical circuit to the recorder.

3.4.2.2 Horizontal-motion seismometer

The principle of the mechanical type of horizontal-motion seismometer is similar to that of the vertical-motion instrument. As before the inertial mass is mounted on a horizontal bar, but the fulcrum is now hinged almost vertically so that the mass is confined to swing sideways in a nearly horizontal plane (Fig. 3.23). The behavior of the system is similar to that of a gate when its hinges are slightly out of vertical alignment. If the hinge axis is tilted slightly forward, the stable position of the gate is where its center of mass is at its lowest point. In any displacement of the gate, the restoring gravitational forces try to return it to this stable position. Similarly, the horizontal-motion seismometer swings about its equilibrium position like a horizontal pendulum (in fact it is the housing of the instrument that moves and not the inertial mass). As in the vertical-motion seismometer, a pen or light-beam attached to the stationary inertial mass writes on a rotating drum (which in this case has a horizontal axis) and records the relative motion between the mass and the instrument housing. The trace of the ground motion detected with this instrument is the horizontal-motion seismogram of the seismic wave.

The design of an electromagnetic horizontal-motion seismometer is similar to that of the vertical-motion type, with the exception that the axis of the moving member (coil or magnet) is horizontal.

3.4.2.3 Strain seismometer

The pendulum seismometers described above are inertial devices, which depend on the resistance of a loosely coupled mass to a change in its momentum. At about the same time that he developed the inertial seismograph, Milne also conducted experiments with a primitive strain seismograph that measured the change in distance between two posts during the passage of a seismic wave. The gain of early strain seismographs was low. However, in 1935 H. Benioff invented a sensitive strain seismograph from which modern versions are descended.

The principle of the instrument is shown in Fig. 3.24. It can record only horizontal displacements. Two collinear
horizontal rods made of fused quartz so as to be insensitive to temperature change are attached to posts about 20 m apart, fixed to the ground; their near ends are separated by a small gap. The changes in separation of the two fixed posts result in changes in the gap width, which are detected with a capacitance or variable-reluctance transducer. In modern instruments the variation in gap width may be observed optically, using the interference between laser light-beams reflected from mirrors attached to the opposite sides of the gap. The strain instrument is capable of resolving strains of the order of $10^{-8}$ to $10^{-10}$.

### 3.4.3 The equation of the seismometer

Inertial seismometers for recording horizontal and vertical ground motion function on the pendulum principle. When the instrument frame is displaced from its equilibrium position relative to the inertial mass, a restoring force arises that is, to first order, proportional to the displacement. Let the vertical or horizontal displacement, dependent on the type of seismometer, be $u$ and the restoring force $-ku$, and let the corresponding displacement of the ground be $q$. The total displacement of the inertial mass $M$ is then $(u + q)$, and the equation of motion is

$$M\frac{d^2}{dt^2}(u + q) = -ku$$  \hspace{1cm} (3.77)

We now divide throughout by $M$, write $k/M = \omega_0^2$, and after rearranging the equation, we get the familiar equation of forced simple harmonic motion

$$\frac{d^2u}{dt^2} + \omega_0^2u = -\frac{d^2q}{dt^2}$$  \hspace{1cm} (3.78)

In this equation $\omega_0$ is the natural frequency (or resonant frequency) of the instrument. For a ground motion with this frequency, the seismometer would execute large uncontrolled vibrations and the seismic signal could not be recorded accurately. To get around this problem, the seismometer motion is damped by providing a velocity-dependent force that opposes the motion. A damping term enters into the equation of motion, which becomes

$$\frac{d^2u}{dt^2} + 2\lambda\omega_0\frac{du}{dt} + \omega_0^2u = -\frac{d^2q}{dt^2}$$  \hspace{1cm} (3.79)

The constant $\lambda$ in this equation is called the damping factor of the instrument. It plays an important role in determining how the seismometer responds to a seismic wave.

A seismic signal is generally composed of numerous superposed harmonic vibrations with different frequencies. We can determine how a seismometer with natural frequency $\omega_0$ responds to a signal with any frequency $\omega$ by solving Eq. (3.78) with $q = A\cos(\omega t)$ (Box 3.2). Here, $A$ is the magnified amplitude of the ground motion, equal to the true ground motion multiplied by a magnification factor that depends on the sensitivity of the instrument. Let the displacement $u$ recorded by the seismometer be

$$u = U\cos(\omega t - \Delta)$$  \hspace{1cm} (3.80)

where $U$ is the amplitude of the recorded signal and $\Delta$ is the phase difference between the record and the ground motion. As derived in Box 3.2, the phase lag $\Delta$ is given by

$$\Delta = \tan^{-1}\left(\frac{2A\omega\omega_0}{\omega_0^2 - \omega^2}\right)$$  \hspace{1cm} (3.81)

The solution of the equation of motion (Eq. (3.79)) gives the displacement $u$ on the seismic record as

$$u = \frac{A\omega^2}{(\omega_0^2 - \omega^2)^2 + 4\lambda^2\omega_0^2}\omega^2\cos(\omega t - \Delta)$$  \hspace{1cm} (3.82)

### 3.4.3.1 Effect of instrumental damping

The ground motion caused by a seismic wave contains a broad spectrum of frequencies. Equation (3.82) shows that the response of the seismometer to different signal frequencies is strongly dependent on the value of the damping factor $\lambda$ (Fig. 3.25). A completely undamped seismometer has $\lambda = 0$, and for small values of $\lambda$ the response of the seismometer is said to be underdamped. An undamped or greatly underdamped seismometer preferentially amplifies signals near the natural frequency, and therefore cannot make an accurate record of the ground motion; the undamped instrument will resonate at its natural frequency $\omega_0$. For all damping factors $\lambda < 1/\nu$ the instrument response function has a peak, indicating preferential amplification of a particular frequency.

The value $\lambda = 1$ corresponds to critical damping, so-called because it delineates two different types of seismometer response in the absence of a forcing vibration. If $\lambda < 1$, the damped, free seismometer responds to a disturbance by swinging periodically with decreasing amplitude about its rest position. If $\lambda > 1$, the disturbed seismometer behaves aperiodically, moving smoothly back to its rest position. However, if the damping is too severe ($\lambda \gg 1$), the instrument is overdamped and all frequencies in the ground motion are suppressed.

The optimum behavior of a seismometer requires that the instrument should respond to a wide range of frequencies in the ground motion, without preferential amplification or excessive suppression of frequencies.
Box 3.2: The seismometer equation

Equation (3.79) is the damped equation of motion for the signal \( u \) recorded by a seismometer with natural frequency \( \omega_0 \) and damping factor \( \lambda \), when the ground displacement is \( q \). Let the ground displacement be \( q = A \cos(\omega_0 t - \Delta) \) and the recorded seismic signal be \( u = U \cos(\omega t - \Delta) \). Substituting in Eq. (3.79) we get the following equation:

\[- \omega^2 U \cos(\omega t - \Delta) - 2 \lambda \omega \omega_0 U \sin(\omega t - \Delta) + \omega_0^2 U \cos(\omega t - \Delta) = A \omega^2 \cos(\omega t - \Delta)\]

i.e.,

\[U[(\omega_0^2 - \omega^2) \cos(\omega t - \Delta) - 2 \lambda \omega \omega_0 \sin(\omega t - \Delta)] = A \omega^2 \cos(\omega t - \Delta)\]

If we now write

\[(\omega_0^2 - \omega^2) = R \cos \phi,\quad 2 \lambda \omega \omega_0 = R \sin \phi\]

where

\[R = [(\omega_0^2 - \omega^2)^2 + 4 \lambda^2 \omega^2 \omega_0^2]^{1/2}, \quad \tan \phi = \frac{2 \lambda \omega \omega_0}{\omega_0^2 - \omega^2}\]

the equations reduce to

\[U[R \cos \phi \cos(\omega t - \Delta) - R \sin \phi \sin(\omega t - \Delta)] = A \omega^2 \cos(\omega t - \Delta)\]

\[U R \cos(\omega t - \Delta + \phi) = A \omega^2 \cos(\omega t - \Delta)\]

Complex numbers (see Box 2.6) allow a simple solution to this equation. The function \( \cos \theta \) is the real part of the complex number (i.e., \( \cos \theta = \text{Re} \{e^{i \theta}\} \)). Therefore,

\[U \text{Re} \{e^{i(\omega t - \Delta + \phi)}\} = A \omega^2 \text{Re} \{e^{i \omega t}\}\]

\[U = \frac{A \omega^2}{R} \text{Re} \{e^{i \omega t} e^{-i(\Delta + \phi)}\} = \frac{A \omega^2}{R} \text{Re} \{e^{i(\Delta - \phi)}\}\]

\[U = \frac{A \omega^2}{R} \cos(\Delta - \phi)\]

The maximum amplitude of the record, \( U \), is when \( \cos(\Delta - \phi) = 1 \). The corresponding phase lag \( \Delta \) between the recorded signal and the ground motion is

\[\Delta = \tan^{-1} \left(\frac{2 \lambda \omega \omega_0}{\omega_0^2 - \omega^2}\right)\]

and the equation for the amplitude of the seismic record is given by

\[u = \frac{A \omega^2}{[(\omega_0^2 - \omega^2)^2 + 4 \lambda^2 \omega^2 \omega_0^2]^{1/2}} \cos(\omega t - \Delta)\]

This requires that the damping factor should be close to the critical value. It is usually chosen to be in the range 70% to 100% of critical damping (i.e., \( 1/\sqrt{2} \leq \lambda < 1 \)). At critical damping the response of the seismometer to a periodic disturbing signal with frequency \( \omega \) is given by

\[u = \frac{A \omega^2}{(\omega_0^2 + \omega^2)} \cos(\omega t - \Delta)\]  

(3.83)

### 3.4.3.2 Long-period and short-period seismometers

The natural period \( (2\pi/\omega) \) of a seismometer is an important factor in determining what it actually records. Two examples of special interest correspond to instruments with very long and very short natural periods, respectively.

The **long-period seismometer** is an instrument in which the resonant frequency \( \omega_0 \) is very low. For all but the lowest frequencies we can write \( \omega_0 \ll \omega \). The phase lag \( \Delta \) between the seismometer and the ground motion becomes zero (see Box 3.2), and the amplitude of the seismometer displacement becomes equal to the amplified ground displacement \( q \):

\[u = A \cos(\omega t - \Delta) = q\]  

(3.84)

The long-period seismometer is sometimes called a **displacement meter**. It is usually designed to record seismic signals with frequencies of 0.01–0.1 Hz (i.e., periods in the range 10–100 s).
3.4 THE SEISMOGRAPH

The short-period seismometer is constructed to have a very short natural period and a correspondingly high resonant frequency $\omega_0$, which is higher than most frequencies in the seismic wave. Under these conditions we have $\omega_0 \gg \omega$, the phase difference $\Delta$ is again small and Eq. (3.83) becomes

$$u = \frac{\omega_0^2}{\omega_0^2} A \cos \omega t = -\frac{1}{\omega_0^2} A \cos \omega t$$  (3.85)

This equation shows that the displacement of the short-period seismometer is proportional to the acceleration of the ground, and the instrument is accordingly called an accelerometer. It is usually designed to respond to seismic frequencies of 1–10 Hz (periods in the range 0.1–1 s). An accelerometer is particularly suitable for recording strong motion earthquakes, when the amplitude of the ground motion would send a normal type of displacement seismometer off-scale.

3.4.3.3 Broadband seismometers

Short-period seismometers operate with periods of 0.1–1 s and long-period instruments are designed for periods greater than 10 s. The resolution of seismic signals with intermediate frequencies of 0.1–1 Hz (periods of 1–10 s) is hindered by the presence in this range of a natural form of seismic background noise. The noise derives from a nearly continuous succession of small ground movements that are referred to as microseisms. Some microseismic noise is of local origin, related to such effects as vehicular traffic, rainfall, wind action on trees, etc. However, an important source is the action of storm waves at sea, which is detectable on seismic records far inland. The drumming of rough surf on a shoreline and the interference of sea waves over deep water are thought to be the principal causes of microseismic noise. The microseismic noise has a low amplitude on a seismogram, but it may be as strong as a weak signal from a distant earthquake, which cannot be selectively amplified without also magnifying the noise. The problem is exacerbated by the limited dynamic range of short- or long-period seismometers. Short-period instruments yield records dominated by high frequencies while long-period devices smooth these out, giving a record with only a low-frequency content (Fig. 3.26a).

The range between the strongest and weakest signals that can be recorded without distortion by a given instrument is called its dynamic range. Dynamic range is measured by the power (or energy density) of a signal, and is expressed in units of decibels (dB). A decibel is defined as $10 \log_{10}$(signal power). Because power is proportional to the square of amplitude (Section 3.3.2.6), a decibel is equivalent to $20 \log_{10}$(amplitude). So, for example, a range of 20 dB in power corresponds to a factor 10 variation in acceleration, and a dynamic range of 100 dB corresponds to a $10^3$ variation in amplitude. Short- and long-period seismometers have narrow dynamic ranges because they are designed to give optimum performance in limited frequency ranges, below or above the band of ground noise. This handicap was overcome by the design of broadband seismometers that have high sensitivity over a very wide dynamic range.

The broadband seismometer has basically an inertial pendulum-type design, with enhanced capability due to a force-feedback system. This works by applying a force proportional to the displacement of the inertial mass to prevent it from moving significantly. The amount of feedback force applied is determined using an electrical transducer to convert motion of the mass into an electrical signal. The force needed to hold the mass stationary corresponds to the ground acceleration. The signal is digitized with 16-to-24 bit resolution, synchronized with accurate time signals, and recorded on magnetic tape, hard disk, or solid state memory. The
feedback electronics are critical to the success of this instrument.

Broadband design results in a seismometer with great bandwidth and linear response. It is no longer necessary to avoid recording in the 1–10 s bandwidth of ground noise avoided by short-period and long-period seismometers. The recording of an earthquake by a broadband seismometer contains more useable information than can be obtained from the short-period or long-period recordings individually or in combination (Fig. 3.26a).

Broadband seismometers can be used to register a wide range of signals (Fig. 3.26b). The dynamic range extends from ground noise up to the strong acceleration that would result from an earthquake with magnitude 9.5, and the periods that can be recorded range from high-frequency body waves to the very long period oscillations of the ground associated with bodily Earth-tides (Section 2.3.3.5). The instrument is employed world-wide in modern standardized seismic networks, replacing short-period and long-period seismometers.

3.4.4 The seismogram

A seismogram represents the conversion of the signal from a seismometer into a time record of a seismic event. The commonest method of obtaining a directly visible record, in use since the earliest days of modern seismology, uses a drum that rotates at a constant speed to provide the time axis of the record, as shown schematically in Fig. 3.22a and Fig. 3.23. In early instruments a mechanical linkage provided the coupling between sensor and record. The invention of the electromagnetic seismometer by Galitzin allowed transmission of the seismic signal to the recorder as an electrical signal. For many years, a galvanometer was used to convert the electrical signal back to a mechanical form for visual display.

In a galvanometer a small coil is suspended on a fine wire between the poles of a magnet. The current in the coil creates a magnetic field that interacts with the field of the permanent magnet and causes a deflection of the coil. The electrical circuitry of the galvanometer is designed with appropriate damping so that the deflection of the galvanometer is a faithful record of the seismic signal. The deflection can be transferred to a visible record in a variety of ways.

Mechanical and electromagnetic seismometers deliver continuous analog recordings of seismic events. These types of seismometer are now of mainly historic interest, having been largely replaced by broadband seismometers. Galvanometer-based analog recording has been superseded by digital recording.

3.4.4.1 Analog recording

In an early method of recording, a smoked paper sheet was attached to the rotating drum. A fine stylus was connected to the pendulum by a system of levers. The point of the stylus scratch a fine trace on the smoked paper. Later instruments employed a pen instead of the stylus and plain paper instead of the smoked paper. These methods make a "wiggly line" trace of the vibration.

In a further development of galvanometric recording a light-beam was reflected from a small mirror attached to the coil or its suspension to trace the record on photographic paper attached to the rotating drum. Photographic methods of recording are free of the slight friction of mechanical contact and allow inventive modifications of the form of the record. In the variable density method, the galvanometer current modulated the intensity of a light-bulb, so that the fluctuating signal showed on the photographic record as successive light and dark bands. A variable area trace was obtained by using the galvanometer current to vary the aperture of a narrow slit, through which a light-beam passed on to the photographic paper.

Every seismogram carries an accurate record of the elapsed time. This may be provided by a tuned electrical circuit whose frequency of oscillation is controlled by the natural frequency of vibration of a quartz crystal. At regular intervals the timing system delivers a short impulse to the instrument, causing a time signal to be imprinted on the seismogram. Modern usage is to employ a time signal transmitted by radio. The timing lines appear as regularly spaced blips on a wiggly line trace, or as bright lines on photographic records.

An important development after the early 1950s, especially in commercial seismology, was the replacement of photographic recording by magnetic recording. The electrical current from the seismometer was sent directly to the recording head of a tape recorder. The varying current in the recording head imprinted a corresponding magnetization on a magnetic tape. Magnetic recorders might have 24 to 50 parallel channels, each able to record from a different source.

Many years of development resulted in sophisticated methods of analog recording, but they have now been superseded universally by digital methods.

3.4.4.2 Digital recording

In digital recording, the analog signal from a seismometer is passed through an electronic device called an analog-to-digital converter, which samples the continuous input signal at discrete, closely spaced time intervals and represents it as a sequence of binary numbers. Conventional representation on the familiar decimal scale expresses a number as a sum of powers of 10 multiplied by the digits 0–9. In contrast, a number is represented on the binary scale as a sum of powers of 2 multiplied by one of only two digits, 0 or 1. For example, in decimal notation the number 153 represents $1 \times 10^2 + 5 \times 10^1 + 3 \times 10^0$. In binary notation, the same number is represented as...
3.4 THE SEISMOGRAPH

\[ 153 = 128 + 16 + 8 + 1 = 1 \times 2^7 + 0 \times 2^6 + 0 \times 2^5 + 1 \times 2^4 + 1 \times 2^3 + 0 \times 2^2 + 0 \times 2^1 + 1 \times 2^0 = 10011001. \]

Each of the digits in a binary number is called a *bit* and the combination that expresses the digitized quantity is called a *word*; the binary number 10011001 is an *eight-bit word*. A binary number is much longer and thus more cumbersome for everyday use than the decimal form, but it is suitable for computer applications. Because it involves only two digits, a binary number can be represented by a simple condition, such as whether a switch is on or off, or the presence or absence of an electrical voltage or current.

Digital recording of seismic signals was developed in the 1960s and since the early 1970s it has virtually replaced analog recording. The analog method records the signal, usually employing the galvanometer principle, on photographic film or on magnetic tape as a continuous time-varying voltage whose amplitude is proportional to a characteristic of the ground disturbance (displacement, velocity or acceleration). The dynamic range of the analog method is limited and the system has to be adapted specifically to the characteristics of the signal to be recorded. For example, an analog device for recording strong signals lacks the sensitivity needed to record weak signals, whereas an analog recorder of weak signals will be overloaded by a strong signal. The digital recording technique samples the amplified output of the seismometer at time increments of a millisecond or so, and writes the digitized voltage directly to magnetic tape or to a computer hard-disk. This avoids possible distortion of the signal that can result in mechanical or optical recording. Digital recording has greater fidelity than the analog method, its dynamic range is wide, and the data are recorded in a form suitable for numerical processing by high-speed computers.

After processing, the digital record is usually converted back to an analog form for display and interpretation. The processed digital signal is passed through a digital-to-analog converter and displayed as a wiggle trace or variable density record. The familiar continuous paper record is still a common form of displaying earthquake records. However, instead of employing galvanometers to displace the pen in response to a signal, modern devices utilize a motor-driven pen; in this case the electrical signal from the earthquake record powers a small servo-motor which controls the pen displacement on the paper record.

### 3.4.4.3 Phases on a seismogram

The seismogram of a distant earthquake contains the arrivals of numerous seismic waves that have travelled along different paths through the Earth from the source to the receiver. The appearance of the seismogram can therefore be very complicated, and its interpretation demands considerable experience. The analysis of seismic waves that have been multiply reflected and refracted in the Earth’s interior will be treated in Section 3.7. Each event that is recorded in the seismogram is referred to as a *phase*.

As described in Section 3.3.2.1, the fastest seismic waves are the longitudinal waves. The first phase on the seismogram corresponds to the arrival of a longitudinal body wave, identified as the primary wave, or *P-wave* (Fig. 3.27). The next phase is the bodily shear wave, referred to as the secondary wave or *S-wave*, which usually has a larger amplitude than the P-wave. It is followed by the large-amplitude disturbance associated with surface waves, which are sometimes designated *L-waves* because of their much longer wavelengths. Dispersion (Section 3.3.3.3) causes the wavelengths at the head of the surface-wave train to be longer than those at the tail. Conventionally, Rayleigh waves are referred to as L\(_R\) waves, while Love waves are called L\(_Q\) waves.

The arrivals recorded on any seismogram depend on the type of sensor used. For example, a vertical-component seismometer responds to P-, SV- and Rayleigh waves but does not register SH- or Love waves; a horizontal-component seismometer can register P-, SH-, Rayleigh and Love waves. The amplitudes of the different phases on a seismogram are influenced by several factors: the orientation of the instrument axis to the wave path, the epicentral distance (see Section 3.5.2), the focal mechanism and the structure traversed by the waves.

Representative seismograms for a distant earthquake are shown in Fig. 3.27. They were recorded at Harvard, Massachusetts, for an earthquake that occurred deep beneath Peru on May 24, 1991. The seismic body waves from this earthquake have travelled through deep regions of the mantle and several seismic phases are recorded.
The upper seismogram is the trace of a horizontal-component seismometer oriented almost transverse to the seismic path, so the P-wave arrival is barely discernible. The first strong signal is the S body wave (in this case, an SH-wave), closely followed by several other phases (defined in Section 3.7.2) and the Love ($L_Q$) surface-wave train. The lower seismogram, recorded by a vertical-component seismometer, shows the arrivals of P and SV body waves and the Rayleigh ($L_R$) surface-wave train. Love waves travel along the surface at close to the near-surface S-wave velocity ($V_{\text{LQ}} = \beta$); Rayleigh waves are slower, having $V_{\text{LR}} = 0.92\beta$, so they reach the recording station later than the Love waves.

### 3.5 EARTHQUAKE SEISMOLOGY

#### 3.5.1 Introduction

Most of the earthquakes which shake the Earth each year are so weak that they are only registered by sensitive seismographs, but some are strong enough to have serious, even catastrophic, consequences for mankind and the environment. About 90% of all earthquakes result from tectonic events, primarily movements on faults. The remaining 10% are related to volcanism, collapse of subterranean cavities, or man-made effects.

Our understanding of the processes that lead to earthquakes derives to a large extent from observations of seismic events on the San Andreas fault in California. The average relative motion of the plates adjacent to the San Andreas fault is about 5 cm yr$^{-1}$, with the block to the west of the fault moving northward. On the fault-plane itself, this motion is not continuous but takes place spasmodically. According to modern plate tectonic theory this extensively studied fault system is a transform fault. This is a rather special type, so it cannot be assumed that the observations related to the San Andreas fault are applicable without reservation to all other faults. However, the *elastic rebound model*, proposed by H. F. Reid after the 1906 San Francisco quake, is a useful guide to how an earthquake may occur.

The model is illustrated in Fig. 3.28 by the changes to five parallel lines, drawn normal to the trace of the fault in the unstrained state and intersecting it at the points A–E. Strain due to relative motion of the blocks adjacent to the fault accumulates over several years. Far from the trace of the fault the five lines remain straight and parallel, but close to it they are bent. When the breaking point of the crustal rocks at C is exceeded, rupture occurs and there is a violent displacement on the fault-plane. The relative displacement that has been taken place progressively between the adjacent plates during years or decades is achieved on the fault-plane in a few seconds. The strained rocks adjacent to the fault “rebound” suddenly. The accumulated strain energy is released with the seismic speed of the ruptured rocks, which is several kilometers per second. The segments BC and C′D undergo compression, while CD and BC′ experience dilatation. The points A and E do not move; the stored strain energy at these points is not released. The entire length of the fault-plane is not displaced, only the region in which the breaking point has been exceeded. The greater the length of the fault-plane that is activated, the larger is the ensuing earthquake.

The occurrence of a large earthquake is not necessarily as abrupt as described in the preceding paragraph, although it can be very sudden. In 1976 a major earthquake with magnitude 7.8 struck a heavily populated area of northern China near the city of Tangshan. Although there were known faults in the area, they had long been seismically inactive, and the large earthquake struck without warning. It completely devastated the industrial region and caused an estimated 243,000 fatalities. However, in many instances the accumulating strain is partially released locally as small earthquakes, or foreshocks. This is an indicator that strain energy is building up to the rupture level and is sometimes a premonition that a larger earthquake is imminent.

When an earthquake occurs, most of the stored energy is released in the main shock. However, for weeks or
months after a large earthquake there may be numerous lesser shocks, known as *aftershocks*, some of which can be comparable in size to the main earthquake. Structures weakened by the main event often collapse in large aftershocks, which can cause physical damage as severe as the main shock. The death toll from aftershocks is likely to be less, because people have evacuated damaged structures.

Although in fact the earthquake involves a part of the fault-plane measuring many square kilometers in area, from the point of view of an observer at a distance of hundreds or even thousands of kilometers the earthquake appears to happen at a point. This point is called the *focus* of the earthquake (Fig. 3.29). It generally occurs at a *focal depth* many kilometers below the Earth’s surface. The point on the Earth’s surface vertically above the focus is called the *epicenter* of the earthquake.

### 3.5.2 Location of the epicenter of an earthquake

The distance of a seismic station from the epicenter of an earthquake (the *epicentral distance*) may be expressed in kilometers $\Delta_{em}$ along the surface, or by the angle $\Delta = (180/\pi)(\Delta_{em}/R)$ subtended at the Earth’s center. The travel-times of P- and S-waves from an earthquake through the body of the Earth to an observer are dependent on the epicentral distance (Fig. 3.30a). The travel-time versus distance plots are not linear, because the ray paths of waves travelling to distant seismographs are curved. However, the standard seismic velocity profile of the Earth’s interior is well enough known that the travel-times for each kind of wave can be tabulated or graphed as a function of epicentral distance. In computing epicentral distance from earthquake data the total travel-time is not at first known, because an observer is rarely at the epicenter to record the exact time of occurrence $t_0$ of the earthquake. However, the difference in travel-times for P- and S-waves $(t_s - t_p)$ can be obtained directly from the seismogram; it increases with increasing epicentral distance (Fig. 3.30a).

For local earthquakes we can assume that the seismic velocities $\alpha$ and $\beta$ are fairly constant in the near-surface layers. The time when the earthquake occurred, $t_0$, can then be obtained by plotting the differences $(t_s - t_p)$ against the arrival times $t_p$ of the P-wave at different stations. The plot, called a *Wadati diagram*, is a straight line (Fig. 3.30b). If $D$ is the distance travelled by the seismic wave, the travel-times of P- and S-waves are respectively $t_p = D/\alpha$ and $t_s = D/\beta$, so

$$t_s - t_p = D\left(\frac{1}{\beta} - \frac{1}{\alpha}\right) = t_p\left(\frac{\alpha}{\beta} - 1\right)$$

(3.86)

The intercept $t_0$ of the straight line with the arrival-time axis is the time of occurrence of the earthquake and the slope of the line is $[(\alpha/\beta) - 1]$. Knowing the P-wave velocity $\alpha$, the distance to the earthquake is obtained from $D = \alpha(t_p - t_0)$.

In order to determine the location of an earthquake, epicenter travel-times of P- and S-waves to at least three seismic stations are necessary (Fig. 3.31). The data from one station give only the distance of the epicenter from that station. It could lie anywhere on a circle centered at the station. The data from an additional station define a second circle which intersects the first circle at two points,
each of which could be the true epicenter. Data from a third station remove the ambiguity: the common point of intersection of the three circles is the epicenter.

Generally the circles do not intersect at a point, but form a small spherical triangle. The optimum location of the epicenter is at the center of the triangle. If data are available from more than three seismic stations, the epicentral location is improved; the triangle is replaced by a small polygon. This situation arises in part from observational errors, and because the theoretical travel-times are imperfectly known. The interior of the Earth is neither homogeneous nor isotropic, as must be assumed. The exact location of earthquake epicenters requires detailed knowledge of the seismic velocities along the entire path, but especially under the source area and the receiving station. The main reason for the intersection triangle or polygon is, however, that the seismic rays travel to the seismograph from the focus, and not from the epicenter. The focal depth of the earthquake, $d$, which may be up to several hundred kilometers, must be taken into account. It can be estimated from simple geometry. If $\Delta_{km}$ is the epicentral distance and $D$ the distance travelled by the wave, to a first approximation $d = (D^2 - \Delta_{km}^2)^{1/2}$. Combining several values of $d$ from different recording stations gives a reasonable estimate of the focal depth.

### 3.5.3 Global seismicity

The epicenters of around 30,000 earthquakes are now reported annually by the International Seismological Center. The geographical distribution of world seismicity (see Fig. 1.10) dramatically illustrates the tectonically active regions of the Earth. The seismicity map is important evidence in support of plate tectonic theory, and delineates the presently active plate margins.

Earthquake epicenters are not uniformly distributed over the Earth’s surface, but occur predominantly along rather narrow zones of interplate seismic activity. The circum-Pacific zone, in which about 75–80% of the annual release of seismic energy takes place, forms a girdle that encompasses the mountain ranges on the west coast of the Americas and the island arcs along the east coast of Asia and Australasia. The Mediterranean-transasianic zone, responsible for about 15–20% of the annual seismic energy release, begins at the Azores triple junction in the Atlantic Ocean and extends along the Azores–Gibraltar ridge; after passing through North Africa it makes a loop through the Italian peninsula, the Alps and the Dinarides; it then runs through Turkey, Iran, the Himalayan mountain chain and the island arcs of southeast Asia, where it terminates at the circum-Pacific zone. The system of oceanic ridges and rises forms the third most active zone of seismicity, with about 3–7% of the annually released seismic energy. In addition to their seismicity, each of these zones is also characterized by active volcanism.

The remainder of the Earth is considered to be aseismic. However, no region of the Earth can be regarded as completely earthquake-free. About 1% of the global seismicity is due to intraplate earthquakes, which occur remote from the major seismic zones. These are not necessarily insignificant: some very large and damaging earthquakes (e.g. the New Madrid, Missouri, earthquakes of 1811 and 1812 in the Mississippi river valley) have been of the intraplate variety.

Earthquakes can also be classified according to their focal depths. Earthquakes, with shallow focal depths less than 70 km, occur in all the seismically active zones; only shallow earthquakes occur on the oceanic ridge systems. The largest proportion (about 85%) of the annual release of seismic energy is liberated in shallow-focus earthquakes. The remainder is set free by earthquakes with intermediate focal depths of 70–300 km (about 12%) and by earthquakes with deep focal depths greater than 300 km (about 3%). These occur only in the circum-Pacific and Mediterranean-transasianic seismic zones, and accompany the process of plate subduction.
The distributions of epicentral locations and focal depths of intermediate and deep earthquakes give important evidence for the processes at a subduction zone. When the earthquake foci along a subduction zone are projected onto a cross-section normal to the strike of the plate margin, they are seen to define a zone of seismicity about 30–40 km thick in the upper part of the 80–100 km thick subducting oceanic plate, which plunges at roughly 30–60° beneath the overriding plate (Fig. 3.32). For many years the inclined seismic zone was referred to in Western literature as a Benioff zone in recognition of the Californian scientist, Hugo Benioff. In the years following World War II Benioff carried out important pioneering studies that described the distribution of deep earthquakes on steeply dipping surfaces of seismicity. Many characteristics of the occurrence of deep earthquakes had been described in the late 1920s by a Japanese seismologist, Kiyoo Wadati. He discovered that the closer the epicenters of earthquakes lay to the Asian continent, the greater were their focal depths; the deep seismicity appeared to lie on an inclined plane. It was Benioff, however, who in 1954 proposed as an explanation of the phenomenon that the ocean floor was being “subducted” underneath the adjacent land. This was a bold proposal well in advance of the advent of plate tectonic theory. Today the zone of active seismicity is called a Wadati–Benioff zone in recognition of both discoverers.

In three dimensions the Wadati–Benioff zone constitutes an inclined slab dipping underneath the overriding plate. It marks the location and orientation of the upper surface of the subducting plate. The dip-angle of the zone varies between about 30° and 60°, becoming steeper with increasing depth, and it can extend to depths of several hundred kilometers into the Earth. The deepest reliably located focal depths extend down to about 680 km. Important changes in the crystalline structure of mantle minerals occur below this depth.

The structure of a subducting plate is not always as simple as described. A detailed study of the subducting Pacific plate revealed a double Wadati–Benioff zone under northeast Honshu, Japan (Fig. 3.33). The seismicity at depths below 100 km defines two parallel planes about 30–40 km apart. The upper plane, identified with the top of the subducting plate, is in a state of compression; the lower plane, in the middle of the slab, is in a state of extension. These stress states are the result of unbending of the subducting plate, which had previously undergone sharp bending at shallow depth below the trench axis. This information is inferred from analysis of the mechanisms by which the earthquakes occur.

3.5.4 Analysis of earthquake focal mechanisms

During an earthquake the accumulated elastic energy is released suddenly by physical displacement of the ground, as heat and as seismic waves that travel outwards from the focus. By studying the first motions recorded by seismographs at distant seismic stations, the focal mechanism of the earthquake can be inferred and the motion on the fault-plane interpreted.

Consider a vertical section perpendicular to the plane of a normal fault on which the hypocenter of an earthquake is located at the point H (Fig. 3.34). When the region above the fault moves up-slope, it produces a region of compression ahead of it and a region of dilatation (or expansion) behind it. In conjunction with the compensatory down-slope motion of the lower block, the earthquake produces two regions of compression and
two regions of dilatation surrounding the hypocenter. These are separated by the fault-plane itself, and by an auxiliary plane through the focus and normal to the fault-plane. When a seismic P-wave travelling out from a region of compression reaches an observer at C, its first effect is to push the Earth’s surface upwards; the initial effect of a P-wave that travels out from a region of dilatation to an observer at D is to tug the surface downwards. The P-wave is the earliest seismic wave to reach a seismograph at C or D and therefore the initial motion recorded by the instrument allows us to distinguish whether the first arrival was compressional or dilatational.

3.5.4.1 Single-couple and double-couple radiation patterns

The amplitudes of P-waves and S-waves vary with distance from their source because of the effects of physical damping and geometric dispersion. The amplitudes also depend geometrically on the angle at which the seismic ray leaves the source. This geometric factor can be calculated mathematically, assuming a model for the source mechanism. The simplest is to represent the source by a single pair of antiparallel motions. Analysis of the amplitude of the P-wave as a function of the angle \( \theta \) between a ray and the plane of the fault (Fig. 3.35a) gives an equation of the form

\[
A(r, t, \alpha, \theta) = A_0(r, t, \alpha) \sin^2 \theta
\]  

(3.87)

in which \( A_0(r, t, \alpha) \) describes the decrease in amplitude with distance \( r \), time \( t \), and seismic P-wave velocity \( \alpha \). A plot of the amplitude variation with \( \theta \) is called the radiation pattern of the P-wave amplitude, which for the single-couple model has a quadrupolar character (Fig. 3.35a). It consists of four lobes, two corresponding to the angular variation of amplitude where the first motion is compressional, and two where the first motion is dilatational.
dilatational. The lobes are separated by the fault-plane and the auxiliary plane.

The radiation pattern for S-waves from a single-couple source is described by an equation of the form

\[ B(r,t,\beta,0) = B_0(r,t,\beta)\sin^2\beta \]

where the amplitude \( B \) is now dependent on the S-wave velocity \( \beta \). The radiation pattern has a dipolar character consisting of two lobes in which the first motions are of opposite sense.

An alternative model of the earthquake source is to represent it by a pair of orthogonal couples (Fig. 3.35b). The double-couple source gives the same form of radiation pattern for P-waves as the single-couple source, but the radiation pattern for S-waves is quadrupolar instead of dipolar. This difference in the S-wave characteristics enables the seismologist to determine which of the two earthquake source models is applicable. S-waves arrive later than P-waves, so their first motions must be resolved from the background noise of earlier arrivals. They can be observed and are consistent with the double-couple model.

Note that the maximum P-wave amplitudes occur at 45° to the fault plane. The directions of maximum amplitude in the compressional and dilatational fields define the T-axis and P-axis, respectively. Here T and P imply "tension" and "compression," respectively, the stress conditions before faulting. Geometrically the P- and T-axes are the bisectors of the angles between the fault-plane and auxiliary plane. The orientations of these axes and of the fault-plane and auxiliary plane can be obtained even for distant earthquakes by analyzing the directions of first motions recorded in seismograms of the events. The analysis is called a fault-plane solution, or focal mechanism solution.

3.5.4.2 Fault-plane solutions

The ray path along which a P-wave travels from an earthquake to the seismogram is curved because of the variation of seismic velocity with depth. The first step in the fault-plane solution is to trace the ray back to its source. A fictitious small sphere is imagined to surround the focus (Fig. 3.36a) and the point at which the ray intersects its surface is computed with the aid of standardized tables of seismic P-wave velocity within the Earth. The azimuth and dip of the angle of departure of the ray from the earthquake focus are calculated and plotted as a point on the lower hemisphere of the small sphere. This direction is then projected onto the horizontal plane through the epicenter. The projection of the entire lower hemisphere is called a stereogram. The direction of the ray is marked with a solid point if the first motion was a push away from the focus (i.e., the station lies in the field of compression). An open point indicates that the first motion was a tug towards the focus (i.e., the station lies in the field of dilatation).

First-motion data of any event are usually available from several seismic stations that lie in different directions from the focus. The solid and open points on the stereogram usually fall in distinct fields of compression and dilatation (Fig. 3.36b). Two mutually orthogonal planes are now drawn so as to delineate these fields as well as possible. The fit is best made mathematically by a least-squares technique, but often a visual fit is obvious and sufficient. The two mutually orthogonal planes correspond to the fault-plane and the auxiliary plane, although it is not possible to decide which is the active fault-plane from the seismic data alone. The regions of the stereogram corresponding to compressional first motions are usually shaded to distinguish them from the regions of dilatational first motions (Fig. 3.36c). The P- and T-axes are the lines that bisect the angles between the fault-plane
and auxiliary plane in the fields of dilatation and compression, respectively. To attach a physical meaning to the P- and T-axes we will have to take a closer look at the mechanics of faulting.

### 3.5.4.3 Mechanics of faulting

As discussed in elasticity theory, the state of stress can be represented by the magnitudes and directions of the three principal stresses $\sigma_1 > \sigma_2 > \sigma_3$. The directions of these principal stresses are by definition parallel to the coordinate axes and are therefore positive for tensional stress. The theory of faulting of homogeneous materials has been developed by studying the failure of materials under compressional stress, which is directed inwards toward the origin of the coordinate axes. The minimum tensional stress corresponds to the maximum compressional stress, and vice versa. The reason for taking this view is that geologists are interested in the behavior of materials within the Earth, where pressure builds up with increasing depth and faulting occurs under high confining pressures.

We can combine both points of view if we consider stress to consist of a part that causes change of volume and a part that causes distortion. The first of these is called the hydrostatic stress, and is defined as the mean $(\sigma_m)$ of the three principal stresses: $\sigma_m = (\sigma_1 + \sigma_2 + \sigma_3)/3$. If we now subtract this value from each of the principal stresses we get the deviatoric stresses: $\sigma'_1 > \sigma'_2 > \sigma'_3$. Note that $\sigma'_1$ is positive; it is a tensional stress, directed outward. However, $\sigma'_3$ is negative; it is a compressional stress, directed inward.

Failure of a material occurs on the plane of maximum shear stress. For a perfectly homogeneous material this is the plane that contains the intermediate principal stress axis $\sigma_2$ (or $\sigma_3$). The fault-plane is oriented at 45° to the axes of maximum and minimum compressional stress, i.e., it bisects the angle between the $\sigma'_1$ and $\sigma'_3$ axes. In real materials inhomogeneity and the effects of internal friction result in failure on a fault-plane inclined at 20°–30° to the axis of maximum compression. Seismologists often ignore this complication when they make fault-plane analyses. The axis of maximum tensional stress $\sigma'_1$ is often equated with the T-axis, in the field of compressional first motions on the bisector of the angle between the two principal planes. The axis of maximum compressional stress $\sigma'_3$ is equated with the P-axis, in the field of dilatational first motions. In reality the direction of $\sigma'_3$ will lie between the P-axis and the fault plane.

The locations of P and T may at first seem strange, for the axes appear to lie in the wrong quadrants. However, one must keep in mind that the orientations of the principal stress axes correspond to the stress pattern before the earthquake, while the fault-plane solution shows the ground motions that occurred after the earthquake. The focal mechanism analysis makes it possible to interpret the directions of the principal axes of stress in the Earth’s crust that led to the earthquake.

![Fig. 3.37](image-url) The three main types of fault and their focal mechanisms. Left: the orientations of each fault-plane and the principal deviatoric stresses, $\sigma'_1$ and $\sigma'_3$. Right: focal mechanisms and orientations of P- and T-axes.

There are only three types of tectonic fault. These can be distinguished by the orientations of the principal axes of stress to the horizontal plane (Fig. 3.37). The focal solutions of earthquakes associated with each type of fault have characteristic geometries. When motion on the fault occurs up or down the fault plane it is called a dip-slip fault, and when the motion is horizontal, parallel to the strike of the fault, it is called a strike-slip fault.

Two classes of dip-slip fault are distinguished depending on the sense of the vertical component of motion. In a normal fault, the block on the upper side of the fault drops down an inclined plane of constant steepness relative to the underlying block (Fig. 3.37a). The corresponding fault-plane solution has regions of compression at the margins of the stereogram. The T-axis is horizontal and the P-axis is vertical. A special case of this type is the listric fault, in which the steepness of the fault surface is not constant but decreases with increasing depth.

In the second type of dip-slip fault, known as a reverse fault or thrust fault, the block on the upper side of the fault moves up the fault-plane, overriding the underlying block (Fig. 3.37b). The fault-plane solution is typified by a central compressional sector. The orientations of the axes of maximum tension and compression are the inverse of the case for a normal fault. The T-axis is now vertical and the P-axis is horizontal. When the fault-plane is inclined at a very flat angle, the upper block can be transported over large horizontal distances. This special type of overthrust fault is common in regions of continental collision, as for example in the Alpine–Himalayan mountain belts.
The simplest type of strike-slip fault is the transcurrent fault, in which the fault-plane is steep or vertical (Fig. 3.37c). To cause this type of fault the T- and P-axes must both lie in the horizontal plane. The fault-plane solution shows two compressional and two dilatational quadrants. Each side of the fault moves in opposite horizontal directions. If the opposite side of the fault to an observer is perceived to move to the left, the fault is said to be sinistral, or left-handed; if the opposite side moves to the right, the fault is dextral, or right-handed.

A variant of the strike-slip fault plays a very important role in the special context of plate tectonics. A transform fault allows horizontal motion of one plate relative to its neighbor. It joins segments of a spreading ocean ridge, or segments of a subducting plate margin. It is important in plate tectonics because it constitutes a conservative plate margin at which the tectonic plates are neither formed nor destroyed. The relative motion on the transform fault therefore reveals the direction of motion on adjacent segments of an active plate margin. The sense of motion is revealed by the pattern of compressional and dilatational sectors on the fault-plane solution.

3.5.4.4 Focal mechanisms at active plate margins

Some of the most impressive examples of focal mechanism solutions have been obtained from active plate margins. The results fully confirm the expectations of plate tectonic theory and give important evidence for the directions of plate motions. We can first ask what types of focal mechanism should be observed at each of the three types of active plate margin. In the theory of plate tectonics these are the constructive (or “accreting”), conservative (or “transform”) and destructive (or “consuming”) margins.

Oceanic spreading systems consist of both constructive and conservative margins. The seismicity at these plate margins forms narrow belts on the surface of the globe. The focal depths are predominantly shallow, generally less than 10 km below the ocean bottom. Active ridge segments are separated by transform faults (Fig. 3.38). New oceanic lithosphere is generated at the spreading oceanic ridges; the separation of the plates at the spreading center is accompanied by extension. The plates appear to be pulled apart by the plate tectonic forces. The extensional nature of the ridge tectonics is documented by fault-plane solutions indicative of normal faulting, as is seen for some selected earthquakes along the Mid-Atlantic Ridge (Fig. 3.39).
If we ignore changes in plate motion (which can, however, occur sporadically on some ridge systems), the offset of the neighboring ridge segments is a permanent feature that reflects how the plates first split apart. The orientations of the transform faults are very important, because the plates must move parallel to these faults. Thus the transform faults provide the key to determining the directions of plate motion. Where a ridge axis is not perpendicular to the transform fault the plate motion will have a component parallel to the ridge segment, which gives an oblique focal mechanism on the ridge.

A destructive (or consuming) plate margin is marked by a subduction zone, where a plate of oceanic lithosphere is destroyed by plunging under another plate of oceanic or continental lithosphere. Because this is a margin of convergence of the adjacent plates, the earthquake fault-plane solutions are typical of a compressional regime (Fig. 3.38). The regions of compressional first motion are in the center of the stereogram, indicating reverse faulting; the P-axes of maximum compressive stress are perpendicular to the strike of the subduction zone.

The type of focal mechanism observed at a subduction zone is dependent on the focal depth of the earthquake. This is because the state of stress varies within the subducting plate. At first the overriding plate is thrust at a shallow angle over the subducting plate. In the seismic zone at the contact between the two plates earthquake focal mechanisms are typical of low-angle reverse faulting. The focal mechanisms of earthquakes along the west coast of Mexico illustrate the first of these characteristics (Fig. 3.41). The selected earthquakes have large magnitudes (6.9 ≤ $M_s$ ≤ 7.8) and shallow focal depths. The strikes of the fault-planes follow the trend of the oceanic trench along the Mexican coastline. The focal mechanisms have a central sector of compressional first motions and the fault-plane is at a low angle to the northeast, typical of overthrusting tectonics. The seismicity pattern documents the subduction of the Cocos plate under the North American plate.

As a result of the interplate collision the subducting oceanic plate is bent downwards and its state of stress changes. Deeper than 60–70 km the seismicity does not arise from the contact between the converging plates. It is
caused by the stress pattern within the subducted plate itself. The focal mechanisms of some intermediate-depth earthquakes (70–300 km) show down-dip extension (i.e., the T-axes are parallel to the surface of the dipping slab) but some show down-dip compression (i.e., the P-axes are parallel to the dip of the slab). At great depths in most Wadati–Benioff zones the focal mechanisms indicate down-dip compression.

### 3.5.4.5 Focal mechanisms in continental collisional zones

When the continental portions of converging plates collide, they resist subduction into the denser mantle. The deformational forces are predominantly horizontal and lead to the formation of folded mountain ranges. The associated seismicity tends to be diffuse, spread out over a large geographic area. The focal mechanisms of earthquakes in the folded mountain belt reflect the ongoing collision. The collision of the northward-moving Indian plate with the Eurasian plate in the Late Tertiary led to the formation of the Himalayas. Focal mechanisms of earthquakes along the arcuate mountain belt show that the present style of deformation consists of two types (Fig. 3.42). Two fault-plane solutions south of the main mountain belt correspond to an extensional regime with normal faulting. To the north, in southern Tibet, the fault-plane solutions also show normal faulting on north–south oriented fault planes. Beneath the Lesser Himalaya seismic events distributed along the entire 1800 km length of the mountain chain indicate a deformational regime with some strike-slip faulting but predominantly low-angle thrust faulting. The Indian continental crust appears to be thrusting at a shallow angle under the Tibetan continental crust. This causes crustal thickening toward the north. In
the main chain of the Lesser Himalaya the minimum compressive stress is vertical. The increased vertical load due to crustal thickening causes the directions of the principal stresses to change, so that under southern Tibet the maximum compressive stress is vertical.

A different type of collisional tectonics is shown by fault-plane solutions from the Alpine mountain belt in south-central Europe. The Alps were formed during the collision between the African and European plates, starting in the Early Tertiary. The focal mechanisms of many, mostly small earthquakes show that they are predominantly associated with strike-slip faults (Fig. 3.43). The horizontal projections of the compressional axes are oriented almost perpendicular to the strike of the Alps and rotate along the arc. The fault-plane solutions for the modern seismicity indicate that the alpine fold-belt is a region of continuing interplate collision.

3.5.5 Secondary effects of earthquakes: landslides, tsunami, fires and fatalities

Before discussing methods of estimating the size of earthquakes, it is worth considering some secondary effects that can accompany large earthquakes: landslides, seismic sea waves and conflagrations. These rather exceptional effects cannot be included conveniently in the definition of earthquake intensity, because their consequences cannot be easily generalized or quantified. For example, once a major fire has been initiated, other factors not related directly to the size of the earthquake (such as aridity of foliage, combustibility of building materials, availability and efficiency of fire-fighting equipment) determine how it is extinguished.

A major hazard associated with large earthquakes in mountainous areas is the activation of landslides, which can cause destruction far from the epicenter. In 1970, an earthquake in Peru with magnitude 7.8 and shallow focal depth of 40 km caused widespread damage and a total death toll of about 66,000. High in the Cordillera Blanca mountains above the town of Yungay, about 15 km away, an enormous slide of rock and ice was released by the tremors. Geologists later speculated that a kind of “air cushion” had been trapped under the mass of rock and mud, enabling it to acquire an estimated speed of 300–400 km h⁻¹, so that it reached Yungay less than five minutes later. Over 90% of the town was buried under the mud and rock, in places to a depth of 14 m, and 20,000 lives were lost.

When a large earthquake occurs under the ocean, it can activate a seismic sea wave known as a tsunami, which in Japanese means “harbor wave.” This special type of sea wave (Box 3.3) is a long-wavelength disturbance of the ocean surface that can be triggered by large-scale collapse or uplift of part of the ocean floor, by an underwater landslide, or by submarine volcanism. The entire water column must be displaced to set off a tsunami, so only earthquakes with magnitude larger than about 7.5 are capable of doing so. The magnitude 9.0 Sumatra–Andaman Islands earthquake on December 26, 2004, had a rupture zone 1300 km in length and 150 km in width, with displacements on the fault of 20 m (Lay et al., 2005). This probably resulted in uplift of the ocean bottom by several meters, which initiated a disastrous tsunami in the Indian Ocean.

A tsunami propagates throughout an ocean basin as a wave with period $T$ of around 15–30 min. The entire water column participates in the motion. As a result, the velocity of the wave, $v$, is dependent on the water depth, $d$, and the acceleration due to gravity, $g$, and is given by:

$$v = \sqrt{gd}$$ (3.89)

Over an ocean basin with water depth greater than 4 km, the tsunami velocity is higher than 200 m s⁻¹ (720 km
3.5 EARTHQUAKE SEISMOLOGY

Box 3.3: Tsunami

When the ocean floor is abruptly lifted or dropped by a submarine earthquake or landslide, the entire water column is pushed up and down. More than 95% of the potential energy of the displaced water is gravitational and less than 5% is elastic energy resulting from compression of the ocean floor or water column. The potential energy of the vertical motion is converted to kinetic energy and propagates away from the source as a tsunami. The propagation of ocean waves is a complex non-linear problem that cannot be handled here, but approximate solutions are useful for the special case of a tsunami.

Ocean surface waves with periods shorter than 50 s are confined to the top kilometer of the ocean and cannot be excited by motions of the sea floor. The propagation of ocean waves is in general dispersive, i.e., the wave velocity depends on its wavelength and period (see section 3.3.3.3). Figure B3.3 shows the computed wave velocities for different periods in water of different depths. The wave speeds are dispersive for periods shorter than about 300 s (5 minutes). These waves have very long wavelengths, much greater than the ocean depth.

For a wave with period $T$ (angular velocity $\omega = 2\pi/T$) and wavelength $\lambda$ (wave number $k = 2\pi/\lambda$), the dispersion in an ocean of depth $d$ is governed by the relation

$$\omega^2 = gk\tanh(kd)$$  \hspace{1cm} (1)

where $g$ is the acceleration of gravity and $\tanh(x)$ is the hyperbolic tangent of $x$:

$$\tanh(x) = \frac{e^x - e^{-x}}{e^x + e^{-x}} = \begin{cases} x, & \text{if } x \text{ small} \\ 1, & \text{if } x \text{ large} \end{cases}$$  \hspace{1cm} (2)

If the wavelength $\lambda$ is very much greater than the ocean depth, $\tanh(kd)$ can be replaced in Eq. (1) by $(kd)$ and the dispersion relation becomes

$$\omega^2 = gk(kd) = k^2gd$$  \hspace{1cm} (3)

$$\omega = k\sqrt{gd}$$

Analogously to surface waves, a tsunami travels across an ocean as a packet of waves of different period. The phase velocity $c$ of a wave is the speed of an individual phase in the packet. Using Eq. (3) and Eq. (3.62), the phase velocity for a tsunami is given by

$$c = \frac{\omega}{k} = \sqrt{gd}$$  \hspace{1cm} (4)

The group velocity $U$ is the propagation speed of the envelope of a wave packet, and thus is the speed with which the energy of the tsunami travels. Using Eq. (3) and Eq. (3.82), the group velocity for a tsunami is given by

$$U = \frac{\partial U}{\partial k} = 0$$  \hspace{1cm} (5)

For water depths that are much less than the wavelength the phase and group velocities are equal and the propagation of a tsunami is non-dispersive. These relations are valid for wavelengths greater than about 11 times the water depth. Over an ocean basin that is 4 km deep the propagation velocity is 200 m s$^{-1}$ (720 km h$^{-1}$) and the wavelengths for periods of 1000–2000 s (around 15–30 min) are 200–400 km. When the tsunami approaches shore and the water depth decreases, its velocity slows. The advancing wave loses kinetic energy, part of which is lost as friction with the ocean bottom and part converted into potential energy. This is evident as in increase in wave height, from a few tens of centimeters over the open ocean to many meters close to land. The contact with land is less often as a ferocious breaking wave than as a rapid increase in sea-level – as during high tides – accompanied by fierce currents which may sweep far inland and withdraw in the same way.

h$^{-1}$, 450 m.p.h), and the wavelength (equal to the product $vT$) may measure 200 km (Box 3.3). The amplitude of a tsunami over the open ocean is comparatively small; the Sumatra tsunami measured about 80–100 cm from crest to trough in the open Indian Ocean. Despite the speed of the wave, an observer on a ship would be scarcely aware of the passage of such a low-amplitude, long-wavelength disturbance. However, on approaching shallower water the leading part of the tsunami slows down and tends to be overridden by the following water mass, so that the height of the wave increases. The wave height may be amplified by the shapes of the sea-bottom and the coastline to several meters. In 1896 a large earthquake raised the ocean-bottom off the southern shore of Japan and initiated a
tsunami that raced ashore with an estimated wave height of more than 20 m, causing 26,000 fatalities. One of the best-studied tsunami was set off by a great earthquake in the Aleutian islands in 1946. It travelled across the Pacific and several hours later reached Hilo, Hawaii, where it swept ashore and up river estuaries as a wave 7 m high. A consequence of the devastation by this tsunami around the Pacific basin was the formation of the Tsunami Warning System. When a major earthquake is detected that can produce a tsunami, a warning is issued to alert endangered regions to the imminent threat. The system works well far from the source. It may take several hours for the tsunami to arrive at a distant location, as illustrated by records of tsunami propagation for the 1960 Chilean and 1964 Alaskan earthquakes (Fig. 3.44). This allows time to warn and evacuate the population in many places far from the epicenter. However, tsunami casualties may still occur near to the epicenter of the earthquake.

The tsunami warning system is currently effective only in the Pacific Ocean. The 2004 Sumatra earthquake triggered the worst tsunami in recorded history with probably more than 250,000 fatalities in Indonesia, Thailand, Sri Lanka and other countries bordering the Indian Ocean as far away as Somalia. As a consequence of this disaster tsunami warning systems are planned for the Indian and other marine basins.

In addition to causing direct damage to man-made structures, an earthquake can disrupt subterranean supply routes (e.g., telephone, electrical and gas lines) which in turn increases the danger of explosion and fire. Aqueducts and underground water pipelines may be broken, with serious consequences for the inhibition or suppression of fires. The San Francisco earthquake of 1906 was very powerful. The initial shock caused widespread damage, including the disruption of water supply lines. But a great fire followed the earthquake, and because the water supply lines were broken by the tremor, it could not be extinguished. The greatest damage in San Francisco resulted from this conflagration.

There are two methods of describing how large an earthquake is. The intensity of the earthquake is a subjective parameter that is based on an assessment of visible effects. It therefore depends on factors other than the actual size of the earthquake. The magnitude of an earthquake is determined instrumentally and is a more objective measure of its size, but it says little directly about the seriousness of the ensuing effects. Illogically, it is usually the magnitude that is reported in news coverage of a major earthquake, whereas the intensity is a more appropriate parameter for describing the severity of its effects on mankind and the environment.

Large earthquakes produce alterations to the Earth’s natural surface features, or severe damage to man-made structures such as buildings, bridges and dams. Even small earthquakes can result in disproportionate damage to these edifices when inferior constructional methods or materials have been utilized. The intensity of an earthquake at a particular place is classified on the basis of the local character of the visible effects it produces. It depends very much on the acuity of the observer, and is in principle subjective. Yet, intensity estimates have proved to be a viable method of assessing earthquake size, including historical earthquakes.

The first attempt to grade earthquake severity was made in the late eighteenth century by Domenico Pignataro, an Italian physician, who classified more than 1000 earth-
Section 3.5 Earthquake Seismology

Table 3.1 Abridged and simplified version of the European Macroseismic Scale 1998 (European Seismological Commission, 1998) for earthquake intensity

The scale focuses especially on the effects on people and buildings. It takes into account classifications of both the vulnerability of a structure (i.e., the materials and method of construction) and the degree of damage.

<table>
<thead>
<tr>
<th>Intensity</th>
<th>Description of effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>I–IV</td>
<td>light to moderate earthquakes</td>
</tr>
<tr>
<td>I</td>
<td>Not felt.</td>
</tr>
<tr>
<td>II</td>
<td>Scarcely felt. Felt only by a few individual people at rest in houses.</td>
</tr>
<tr>
<td>III</td>
<td>Weak. Felt indoors by a few people. People at rest feel a swaying or light trembling.</td>
</tr>
<tr>
<td>IV</td>
<td>Largely observed. Felt indoors by many people; outdoors by very few. A few people are awakened. Windows, doors and dishes rattle.</td>
</tr>
<tr>
<td>V–VIII</td>
<td>moderate to severe earthquakes</td>
</tr>
<tr>
<td>V</td>
<td>Strong. Felt indoors by most, outdoors by few. Many sleeping people awake. A few are frightened. Buildings tremble throughout. Hanging objects swing considerably. Small objects are shifted. Doors and windows swing open or shut.</td>
</tr>
<tr>
<td>VI</td>
<td>Slightly damaging. Many people are frightened and run outdoors. Some objects fall. Many houses suffer slight non-structural damage like hair-line cracks and fall of small pieces of plaster.</td>
</tr>
<tr>
<td>VII</td>
<td>Damaging. Most people are frightened and run outdoors. Furniture is shifted and objects fall from shelves in large numbers. Many well built ordinary buildings suffer moderate damage: small cracks in walls, fall of plaster, parts of chimneys fall down; older buildings may show large cracks in walls and failure of fill-in walls.</td>
</tr>
<tr>
<td>VIII</td>
<td>Heavily damaging. Many people find it difficult to stand. Many houses have large cracks in walls. A few well built ordinary buildings show serious failure of walls, while weak older structures may collapse.</td>
</tr>
<tr>
<td>IX–XII</td>
<td>severe to destructive earthquakes</td>
</tr>
<tr>
<td>X</td>
<td>Very destructive. Many ordinary well built buildings collapse.</td>
</tr>
<tr>
<td>XI</td>
<td>Devastating. Most ordinary well built buildings collapse, even some with good earthquake resistant design are destroyed.</td>
</tr>
<tr>
<td>XII</td>
<td>Completely devastating. Almost all buildings are destroyed.</td>
</tr>
</tbody>
</table>

Earthquakes that devastated the southern Italian province of Calabria in the years 1783–1786. His crude analysis classified the earthquakes according to whether they were very strong, strong, moderate or slight. In the mid-nineteenth century an Irish engineer, Robert Mallet, produced a list of 6831 earthquakes and plotted their estimated locations, producing the first map of the world’s seismicity and establishing that earthquakes occurred in distinct zones. He also used a four-stage intensity scale to grade earthquake damage, and constructed the first isoseismal maps with lines that outlined areas with broadly equal grades of damage. The Rossi–Forel intensity scale, developed in the late nineteenth century by the Italian scientists M. S. de Rossi and the Swiss scientist F. Forel, incorporated ten stages describing effects of increasing damage. In 1902 an Italian seismologist, G. Mercalli, proposed a still more extensive, expanded intensity scale which reclassified earthquake severity in twelve stages. A variation, the Modified Mercalli (MM) scale, was developed in 1931 to suit building conditions in the United States, where a later modification is in common use. The Medvedev–Sponheuer–Karnik (MSK) scale, introduced in Europe in 1964, and modified in 1981, also has twelve stages and differs from the MM scale mainly in details. A new European Macroseismic Scale (EMS-98) was adopted in 1998; an abridged version is shown in Table 3.1. The new 12-stage EMS scale is based on the MSK scale but takes into account the vulnerability of buildings to earthquake damage and incorporates more rigorous evaluation of the extent of damage to structures with different building standards.

In order to evaluate the active seismicity of a region questionnaires may be distributed to the population, asking for observations that can be used to estimate the intensity experienced. The questionnaires are evaluated with the aid of an intensity scale, and the intensity recorded at the location of each observer is plotted on a map. Continuous lines are then drawn to outline places with the same intensity (Fig. 3.45), in the same way that contour lines are used on topographic maps to show elevation. Comparison of the isoseismal maps with geological maps helps explain the response of the ground to the shake of an earthquake. This is valuable information for understanding earthquake risk. The foundation on which structures are erected plays a vital role in their survival of an earthquake. For example, soft sediments can amplify the ground motion, enhancing the damage caused. This is even more serious when the sediments have a high water content, in which case liquefaction of the sediments can occur, robbing structures built on them of support and promoting their collapse.
There are numerous examples of this having occurred. In the great Alaskan earthquake in 1964 a section of Anchorage that was built on a headland underlain by a wet clay substratum collapsed and slid downslope to the sea. In 1985 a very large earthquake with magnitude 8.1 struck the Pacific coast of Mexico. About 350 km away in Mexico City, despite the large distance from the epicenter, the damage to buildings erected on the alluvium of a drained lakebed was very severe while buildings set on hard rock foundations on the surrounding hills suffered only minor damage. In the Loma Prieta earthquake of 1989 severe damage was caused to houses built on landfill in San Francisco’s Mission district, and an overhead freeway built on pillars on young alluvium north of Oakland collapsed dramatically. Both regions of destruction were more than 70 km from the epicenter in the Santa Cruz mountains. Similarly, in the San Francisco earthquake of 1906 worse damage occurred to structures built on landfill areas around the shore of the bay than to those with hard rock foundations in the hills of the San Francisco peninsula.

Intensity data play an important role in determining the historic seismicity of a region. An earthquake has dramatic consequences for a population; this was especially the case in the historic past, when real hazards were augmented by superstition. The date (and even the time) of occurrence of strong earthquakes and observations of their local effects have been recorded for centuries in church and civil documents. From such records it is sometimes possible to extract enough information for a given earthquake to estimate the intensity experienced by the observer. If the population density is high enough, it may be possible to construct an isoseismal map from which the epicenter of the tremor may be roughly located. An interesting example of this kind of analysis is the study of the New Madrid earthquakes of 1811–1812, which caused devastation in the Mississippi valley and were felt as far away as the coastlines along the Atlantic and Gulf of Mexico (Fig. 3.45). There were probably three large earthquakes, but the events occurred before the invention of the seismograph so details of what happened are dependent on the subjective reports of observers. Historical records of the era allow development of an intensity map for the settled area east of the Mississippi, but the pioneering population west of the river was at that time too sparse to leave adequate records for intensity interpretation. On the basis of the available evidence these earthquakes are estimated to have had magnitudes of 7.8–8.1.

Earthquake intensity data are valuable for the construction of seismic risk maps, which portray graphically the estimated earthquake hazard of a region or country. The preparation of a seismic risk map is a lengthy and involved task, which typically combines a study of the present seismicity of a region with analysis of its historic seismicity. A map of the maximum accelerations experienced in recent seismicity helps to identify the areas that are most likely to suffer severe damage in a large earthquake. The likelihood of an earthquake happening in a given interval of time must also be taken into account. Even in a region where large earthquakes occur, they happen at irregular intervals, and so knowledge of local and regional earthquake frequency is important in risk estimation. A seismic risk map of the United States (Fig. 3.46) shows the peak accelerations that are likely to be experienced at least once in a 50-year period.

Devastating earthquakes can occur even in countries with relatively low seismic risk. Switzerland, in the center of Europe, is not regarded as a country prone to earthquakes. The seismic activity consists mostly of small earthquakes, mainly in the Alps in the collision zone of the African and European plates. Yet, in 1356 the northern Swiss town of Basel was destroyed by a major earthquake. Seismic risk maps are useful in planning safe sites for important edifices like nuclear power plants or high dams for hydroelectric power, which supply a substantial proportion of Switzerland’s energy needs. Risk maps are also valuable to insurance companies, which must know the seismic risk of a region in order to assess the costs of earthquake insurance coverage for private and public buildings.

### 3.5.6.2 Earthquake magnitude

Magnitude is an experimentally determined measure of the size of an earthquake. In 1935 C. F. Richter attempted to grade the sizes of local earthquakes in Southern California on the basis of the amplitude of the ground vibrations they produced at a known distance from the
3.5 EARTHQUAKE SEISMOLOGY

**Fig. 3.46** Seismic risk map of the United States. The numbers on contour lines give the maximum acceleration (in percent of g) that might be expected to be exceeded with a probability of 1 in 10 during a 50-year period (after Bolt, 1988).

epicenter. The vibrations were recorded by seismographs, which were standardized to have the same response to a given stimulus. Richter’s original definition of magnitude was based on surface-wave amplitudes ($A_s$) recorded by seismographs at an epicentral distance of 100 km. Because seismographs were located at various distances from the earthquake, an extra term was added to compensate for attenuation of the signal with increasing epicentral distance. Increasingly sensitive instruments allowed the recording of signals from distant earthquakes; those from events with epicentral distances greater than 20° are known as *telesismic* signals. Originally, the magnitude was determined from the horizontal ground motion, because seismological stations were equipped mainly with horizontal-motion seismometers. However, the surface waves recorded by these instruments consist of superposed Love and Rayleigh waves, which complicates the theoretical interpretation of the records. Vertical-motion seismometers record only the Rayleigh waves (together with P- and SV-waves), and so progressively the definition of surface-wave magnitude has come to be based on the vertical component of motion. The majority of surface-wave magnitudes assigned to earthquakes worldwide are now based on vertical-motion records.

The International Association for Seismology and Physics of the Earth’s Interior (IASPEI) has adopted the following definition of the surface-wave magnitude ($M_s$) of an earthquake:

$$M_s = \log_{10}\left(\frac{A_s}{T}\right) + 1.66\log_{10}(\Delta) + 3.3 \quad (3.90)$$

where $A_s$ is the vertical component of the ground motion in micrometers ($\mu$m) determined from the maximum Rayleigh-wave amplitude, $T$ is the period of the wave (18–22 seconds), $\Delta$ is the epicentral distance in degrees ($20^\circ \leq \Delta \leq 160^\circ$), and where the earthquake has a focal depth of less than 50 km. A similar equation to Eq. (3.90) applies to broadband recordings; in this case $(A_s/T)_{max}$ corresponds to the maximum ground velocity. The surface-wave magnitudes of some important historical earthquakes are given in Table 3.2.

The depth of the source affects the nature of the seismic wave train, even when the same energy is released. An earthquake with a deep focus may generate only a small surface-wave train, while shallow earthquakes cause very strong surface waves. Equation (3.90) for $M_s$ was derived from the study of shallow earthquakes, observed at a distance greater than 20°. Therefore, corrections must be made to the computed value of $M_s$ to compensate for the effects of a focal depth greater than 50 km or epicentral distance less than 20°.

The amplitude of body waves is not sensitive to the focal depth. As a result, earthquake magnitude scales have also been developed for use with body waves. An equation, proposed by B. Gutenberg in 1945, can be used to calculate a *body-wave magnitude* ($m_b$) from the maximum amplitude ($A_p$) of the ground motion associated with P-waves having a period ($T$) of less than 3 s:

$$m_b = \log_{10}(\frac{A_p}{T}) + Q(\Delta, h) \quad (3.91)$$

Where $Q(\Delta, h)$ is an empirical correction for signal attenuation due to epicentral distance ($\Delta$) and focal depth ($h$) that is made by reading directly from a graph or table of values.

For some earthquakes both $M_s$ and $m_b$ can be calculated. Unfortunately, the different estimates of magnitude often do not agree well, except for small earthquakes. This is due to the way the ground responds to a seismic event, and to the different natures of body waves and surface waves. Body waves have a different dependence of amplitude on frequency than do surface waves. $m_b$ is estimated from a high-frequency (1 Hz) phase whereas $M_s$ is determined from
Table 3.2 Some important historical earthquakes, with their surface-wave magnitudes $M_s$, moment magnitudes $M_w$, and the numbers of fatalities

<table>
<thead>
<tr>
<th>Year</th>
<th>Epicenter</th>
<th>$M_s$</th>
<th>$M_w$</th>
<th>Fatalities</th>
</tr>
</thead>
<tbody>
<tr>
<td>1906</td>
<td>San Francisco, California</td>
<td>8.3</td>
<td>7.8</td>
<td>3,000</td>
</tr>
<tr>
<td>1908</td>
<td>Messina, Italy</td>
<td>7.2</td>
<td>—</td>
<td>70,000</td>
</tr>
<tr>
<td>1923</td>
<td>Kanto, Japan</td>
<td>8.2</td>
<td>7.9</td>
<td>143,000</td>
</tr>
<tr>
<td>1952</td>
<td>Kamchatka, Russia</td>
<td>8.2</td>
<td>9.0</td>
<td></td>
</tr>
<tr>
<td>1957</td>
<td>Andreanof Islands, Alaska</td>
<td>8.1</td>
<td>8.6</td>
<td></td>
</tr>
<tr>
<td>1960</td>
<td>Valdivia, Chile</td>
<td>8.5</td>
<td>9.5</td>
<td>5,700</td>
</tr>
<tr>
<td>1960</td>
<td>Agadir, Morocco</td>
<td>5.9</td>
<td>5.7</td>
<td>10,000</td>
</tr>
<tr>
<td>1964</td>
<td>Prince William Sound, Alaska</td>
<td>8.6</td>
<td>9.2</td>
<td>125</td>
</tr>
<tr>
<td>1970</td>
<td>Chimbote, Peru</td>
<td>7.8</td>
<td>7.9</td>
<td>66,000</td>
</tr>
<tr>
<td>1971</td>
<td>San Fernando, California</td>
<td>6.5</td>
<td>6.7</td>
<td>65</td>
</tr>
<tr>
<td>1975</td>
<td>Haicheng, China</td>
<td>7.4</td>
<td>7.0</td>
<td>300</td>
</tr>
<tr>
<td>1976</td>
<td>Tangshan, China</td>
<td>7.8</td>
<td>7.5</td>
<td>243,000</td>
</tr>
<tr>
<td>1980</td>
<td>El Asnam, Algeria</td>
<td>7.3</td>
<td>—</td>
<td>2,590</td>
</tr>
<tr>
<td>1985</td>
<td>Michoacan, Mexico</td>
<td>8.1</td>
<td>8.0</td>
<td>9,500</td>
</tr>
<tr>
<td>1989</td>
<td>Loma Prieta, California</td>
<td>7.1</td>
<td>6.9</td>
<td>62</td>
</tr>
<tr>
<td>1994</td>
<td>Northridge, California</td>
<td>6.8</td>
<td>6.7</td>
<td>60</td>
</tr>
<tr>
<td>1999</td>
<td>Izmit, Turkey</td>
<td>—</td>
<td>7.6</td>
<td>17,100</td>
</tr>
<tr>
<td>2004</td>
<td>Sumatra–Andaman Islands</td>
<td>—</td>
<td>9.0</td>
<td>250,000</td>
</tr>
</tbody>
</table>

The seismic moment can be used to define a moment magnitude ($M_w$). The definition adopted by the responsible commission of IASPEI is:

$$M_w = \frac{2}{3}(\log_{10}M_0 - 9.1) \quad (3.93)$$

In this equation $M_0$ is in N m. If c.g.s. units are used instead of SI units, $M_0$ is expressed in dyne cm and the corresponding equation is:

$$M_w = \frac{2}{3}(\log_{10}M_0 - 16.1) \quad (3.94)$$

$M_w$ is more appropriate for describing the magnitudes of very large earthquakes. It has largely replaced $M_s$ in scientific evaluation of earthquake size, although $M_w$ is often quoted in reports in the media. The moment magnitudes and surface-wave magnitudes of some historical earthquakes are listed in Table 3.2.

The magnitude scale is, in principle, open ended. Negative Richter magnitudes are possible, but the limit of sensitivity of seismographs is around −2. The maximum possible magnitude is limited by the shear strength of the crust and upper mantle, and since the beginning of instrumental recording none has been observed with a surface-wave magnitude $M_s$ as high as 9. However, this is largely due to a saturation effect resulting from the method by which surface-wave magnitudes $M_s$ are computed. Seismic moment magnitudes $M_w$ of 9 and larger have been computed for some giant earthquakes (Table 3.2). The largest in recorded history was the 1960 Chile earthquake with $M_w = 9.5$.

When an earthquake occurs on a fault, the ruptured area spreads in size from the point of initial failure, akin to opening a zip fastener. If the fault ruptures along a length $L$, and the fractured segment has a down-dip dimension $w$ (referred to as the width of the faulted zone), the area $S$ is equal to $wL$. Assuming that the aspect ratio of faults is constant (i.e., $w$ is proportional to $L$) then the ruptured area $S$ is proportional to $L^2$. This is a generalization, because different faults have different aspect ratios. Similarly, if the stress drop in an earthquake is constant, the displacement on the fault, $D$, can be assumed proportional to $L$. Together, these assumptions imply that the seismic moment $M_0$ scales as $L^3$. Assuming that $S \propto L^2$, the seismic moment scales as $S^{3/2}$. This inference is supported...
by the correlation between seismic moment and the ruptured area $S$ for many shallow earthquakes (Fig. 3.47).

The length $L$ of the rupture zone determines how long the ground motion lasts in an earthquake. This factor has a significant bearing on the extent of damage to structures during moderate to strong earthquakes. Theory gives the speed of propagation of the rupture as 75–95% of the shear wave velocity. Assuming that the rupture propagates along the fault at about $2.5–3.0 \text{ km s}^{-1}$, the duration of the ground motion can be estimated for an earthquake of a given magnitude or seismic moment (Fig. 3.48). For example, close to the epicenter of a magnitude 5 earthquake the vibration may last a few seconds, whereas in a magnitude 8 earthquake it might last about 50 s. However, there is a lot of scatter about the ideal correlation. In the great Sumatra earthquake, with magnitude 9.0, the rupture continued for some 500 s.

The seismic moment $M_0$ of an earthquake is determined in practice from the analysis of seismograms at a large number of seismic stations (currently 137) distributed worldwide and linked to form the Global Seismographic Network (GSN). The waveform of each seismogram is a product of the epicentral distance, focal depth and radiation pattern of the earthquake (Section 3.5.4.1). It is also affected by source parameters such as the area, orientation and amount of slip of the ruptured segment, factors which determine the seismic moment $M_0$. Inversion of the seismogram data leads to an understanding of the earthquake source. At large distances and for long-period components of the seismogram (e.g., $T > 40$ s) that correspond to wavelengths much longer than the dimensions of the faulted area, the source can be considered as a point source. Assuming a radiation model for the source and a velocity model for the propagation, a synthetic seismogram (see also Section 3.6.5.4) can be computed for each station and compared with the observed waveform. Iterative adjustment of the source parameters to give a best fit to the waveforms at a number of receivers defines mathematically a moment tensor. The tensor can be visualized geometrically as an ellipsoid in which the lengths and orientations of the axes correspond to the moment of the earthquake and the directions of the tensional (T) axis and compressional (P) axis at the source. The location of the system of couples modelled by the moment tensor is the optimum point source location for the earthquake and is called its centroid. The centroid moment tensor (CMT) analyses of earthquakes are made rapidly and give the source parameters of an earthquake within a few hours. The centroid location may differ from the hypocenter of the earthquake (the place where rupture first occurred). The former is based on analysis of the full seismogram waveform, the latter only on the first arrival times. So, for example, the centroid of the 2004 Sumatra–Andaman earthquake was located about 160 km west of its epicenter.

### 3.5.6.3 Relationship between magnitude and intensity

The intensity and magnitude scales for estimating the size of an earthquake are defined independently but they have some common features. Intensity is a measure of earthquake size based on the extent of local damage it causes at the location of an observer. The definition of magnitude is based on the amplitude of ground motion inferred from the signal recorded by the observer’s seismograph, and of course it is the nature of the ground motion – its amplitude,
velocity and acceleration – which produce the local damage used to classify intensity. However, in the definition of magnitude the ground-motion amplitude is corrected for epicentral distance and converted to a focal characteristic. Isoseismal maps showing the regional distribution of damage give the maximum intensity ($I_{\text{max}}$) experienced in an earthquake, which, although influenced by the geographic patterns of population and settlement, is usually near to the epicenter. A moderately strong, shallow-focus earthquake under a heavily populated area can result in higher intensities than a large deep focus earthquake under a wilderness area (compare, for example, the death tolls for the 1960 Agadir magnitude 5.7 and 1964 Alaskan magnitude 9.2 earthquakes in Table 3.2). However, for earthquakes with focal depth $h < 50$ km the dependence of $I_{\text{max}}$ on the focal depth can be taken into account, and it is possible to relate the maximum intensity to the magnitude with an empirical equation (Karnik, 1969):

$$I_{\text{max}} = 1.5M_s - 1.8\log h + 1.7 \quad (3.95)$$

This type of equation is useful for estimating quickly the probable damage that an earthquake causes. For example, it predicts that in the epicentral region of an earthquake with magnitude 5 and a shallow focal depth of 10 km, the maximum MSK intensity will be VII (moderately serious damage), whereas, if the focal depth is 100 km, a maximum intensity of only IV–V (minor damage) can be expected.

### 3.5.7 Earthquake frequency

Every year there are many small earthquakes, and only a few large ones. According to a compilation published by Gutenberg and Richter in 1954, the mean annual number of earthquakes in the years 1921–1945 with magnitudes 4–4.9 was around 6000, while there were only an average of about 100 earthquakes per year with magnitudes 6–6.9. The relationship between annual frequency ($N$) and magnitude ($M_s$) is logarithmic and is given by an equation of the form:

$$\log N = a - bM_s \quad (3.96)$$

The value of $a$ varies between about 8 and 9 from one region to another, while $b$ is approximately unity for regional and global seismicity. The mean annual numbers of earthquakes in different magnitude ranges are listed in Table 3.3; the frequency decreases with increasing magnitude (Fig. 3.49a), in accordance with Eq. (3.96). The annual number of large earthquakes with magnitude $M_s \geq 7$ in the years 1900–1989 has varied between extremes of about 10 and 40, but the long-term average is about 20 per year (Fig. 3.49b).

### 3.5.8 Energy released in an earthquake

The definition of earthquake magnitude relates it to the logarithm of the amplitude of a seismic disturbance. Noting that the energy of a wave is proportional to the square of its amplitude it should be no surprise that the magnitude is also related to the logarithm of the energy. Several equations have been proposed for this relationship. An empirical formula worked out by Gutenberg and Richter (1954), relates the energy release $E$ to the surface-wave magnitude $M_c$:
3.5 EARTHQUAKE SEISMOLOGY

\[ \log_{10} E = 4.8 + 1.5M_s \]  \hspace{1cm} (3.97)

where \( E \) is in joules. The logarithmic nature of the formula means that the energy release increases very rapidly with increasing magnitude. For example, when the magnitudes of two earthquakes differ by 1, their corresponding energies differ by a factor \( 10^1 \), which is equivalent to a factor \( 10^{1.5} \). Hence, a magnitude 7 earthquake releases about 1000 times the energy of a magnitude 5 earthquake. Another way of regarding this observation is that it takes 1000 magnitude 5 earthquakes to release the same amount of energy as a single large earthquake with magnitude 7. Multiplying the mean number of earthquakes per year by their estimated energy (using one of the energy–magnitude equations) gives an impression of the importance of very large earthquakes. Table 3.3 shows that the earthquakes with \( M_s \geq 7 \) are responsible for most of the annual seismic energy. In a year in which a very large earthquake \( (M_s \geq 8) \) occurs, most of the annual seismic energy is released in that single event.

It is rather difficult to appreciate the amount of energy released in an earthquake from the numerical magnitude alone. A few examples help illustrate the amounts of energy involved. Earthquakes with \( M_s = 1 \) are so weak that they can only be recorded instrumentally; they are referred to as microearthquakes. The energy associated with one of these events is equivalent to the kinetic energy of a medium sized automobile weighing 1.5 tons which is travelling at 130 km h\(^{-1} \) (80 m.p.h.). The energy released by explosives provides another means of comparison, although the conversion of energy into heat, light and shock waves is proportionately different in the two phenomena. One ton of the explosive trinitrotoluene (TNT) releases about \( 4.2 \times 10^6 \) joules of energy. Equation (3.97) shows that the 11 kiloton atomic bomb which destroyed Hiroshima released about the same amount of energy as an earthquake with magnitude 5.9. The energy released in a 1 megaton nuclear explosion is equivalent to an earthquake with magnitude 7.2; the 2004 Sumatra earthquake (magnitude 9.0) released an amount of energy equivalent to the detonation of 475 megaton bombs.

3.5.9 Earthquake prediction

The problem of earthquake prediction is extremely difficult and is associated with sundry other problems of a sociological nature. To predict an earthquake correctly means deciding, as far in advance as possible, exactly where and when it will occur. It is also necessary to judge how strong it will be, which means realistically that people want to know what the likely damage will be, a feature expressed in the earthquake intensity. In fact the geophysicist is almost helpless in this respect, because at best an estimate of the predicted magnitude can be made. As seen above, even if it is possible to predict accurately the magnitude, the intensity depends on many factors (e.g., local geology, construction standards, secondary effects like fires and floods) which are largely outside the influence of the seismologist who is asked to presage the seriousness of the event. The problem of prediction rapidly assumes sociological and political proportions. Even if the approximate time and place of a major earthquake can be predicted with reasonable certainty, the question then remains of what to do about the situation. Properly, the threatened area should be evacuated, but this would entail economic consequences of possibly enormous dimension.

The difficulties are illustrated by the following possible scenario. Suppose that seismologists conclude that a very large earthquake, with probable magnitude 7 or greater, will take place sometime in a known month of a given year under a specific urban center. No more precise details are possible; in particular, the time of occurrence cannot be determined more exactly. Publication of this kind of prediction would cause great consternation, even panic, which for some of the population could be as disastrous as the earthquake itself. Should the warning be withheld? If the entire area, with its millions of inhabitants, were to be evacuated, the economic dislocation would be enormous. How long should the evacuation last, when every day is economically ruinous? Clearly, earthquake prediction is only useful in a practical sense when it is accurate in both place and time. The responsible authorities must also be provided with a reasonable estimate of the expected magnitude, from which the maximum intensity \( I_{\text{max}} \) may be gauged with the aid of a relationship like that of Eq. (3.95). The problem then passes out of the domain of the scientist and into that of the politician, with the scientist retaining only a peripheral role as a consultant. But if the earthquake prediction is a failure, the consequences will rebound with certainty on the scientist.

3.5.9.1 Prediction of the location of an earthquake

It is easier to predict where a major earthquake is likely to occur than when it will occur. The global seismicity patterns demonstrate that some regions are relatively aseismic. They are not completely free of earthquakes, many of which appear to occur randomly and without warning in these regions. Some intraplate earthquakes have a history of repeated occurrence at a known location, which can sensibly be expected to be the locus of a future shock. However, because most earthquakes occur in the seismically active zones at plate margins, these are the prime areas for trying to predict serious events. Predicting the location of a future earthquake in these zones combines a knowledge of the historical seismicity pattern with the elastic rebound model of what causes an earthquake.

The seismic gap theory is based on the simple idea that global plates move under the influence of forces which affect the plates as entities. The interactions at a plate margin therefore act along the entire length of the interplate boundary. Models of plate tectonic reconstructions
Fig. 3.50 Seismic gaps along the Aleutian island arc. Shaded regions mark the areas of rupture of very large historic earthquakes (\(M_\text{w} \geq 7.4\)). Three large gaps in seismicity are potential locations of a future large earthquake (after Sykes et al., 1981).

Problematic returns do not lend themselves to convenient statistical evaluation. However, prediction by scientific methods is still uncertain, so the usefulness of alternative premonitory phenomena cannot be rejected out of hand.

Several scientific methods have been tested as possible ways of predicting the time of earthquake occurrence. They are based on detecting ground changes, or effects related to them, that accompany the progression of strain. For example, a geochemical method which has had some degree of success is the monitoring of radon. Some minerals in the Earth's crust contain discrete amounts of uranium. The gas radon is a natural product of the radioactive decay of uranium. It migrates through pores and cracks and because of its own radioactivity it is a known environmental health hazard in buildings constructed in some geographic areas. Radon gas can become trapped in the Earth's crust, and in many areas it forms a natural radioactive background. Prior to some earthquakes anomalous levels of radon have been detected. The enhanced leakage of radon from the crust may be due to porosity changes at depth in response to the accumulating strains.

The build-up of strain is manifest in horizontal and vertical displacements of the Earth's surface, depending on the type of faulting involved. These displacements can be measured geodetically by triangulation or by modern techniques of trilateration which employ laser-ranging devices to measure the time taken for a laser beam to travel to a reflecting target and back to its source. The travel-time of the beam is measured extremely accurately, and converted into the distance between emitter and reflector. A shift on the fault will change this distance. With laser techniques the constant creep of one side of a horizontal fault relative to the other can be observed accurately. In one method the source and receiver of a laser beam are placed on one side of a fault, with a reflector on the opposite side. In an alternative method a laser source and receiver are placed on each side of the fault. Pulsed signals are beamed from each unit to an orbiting reflecting satellite, the position of which is known accurately. For each ground station, differences in the elapsed time of the pulse are converted by computer into ground movement, and into differential motion on the fault.

Several methods are suited to detecting differential vertical motion. Sensitive gravimeters on opposite sides of a fault can detect vertical displacement of as little as one centimeter of one instrument relative to the other. Distension...
of the Earth can be monitored with a tiltmeter, which is an instrument designed on the principle of a water-level. It consists of a long tube about 10 m in length, connecting two water-filled containers, in which the difference in water levels is monitored electronically. A tiltmeter is capable of determining tilt changes of the order of 10–7 degrees. Tiltmeters and gravimeters installed near to active faults have shown that episodes of ground uplift and tilt can sometimes precede major earthquakes.

Geodetic and geophysical observations, such as changes in the local geomagnetic field or the electrical resistivity of the ground, are of fundamental interest. However, the most promising methods of predicting the time and size of an imminent earthquake are based on seismic observations. According to the elastic rebound model, the next earthquake on an active fault (or fault segment) will occur when the stress released in the most recent earthquake again builds up to the local breaking point of the rocks. Thus the probability at any time that an earthquake will occur on the fault depends on the magnitude of the latest earthquake, the time elapsed since it occurred and the local rate of accumulation of stress. A symptom of the stress build-up before a major earthquake can be an increase in foreshock activity, and this was evidently a key parameter in the successful prediction of a large earthquake in Liaoning province, China, in February, 1975. The frequency of minor shocks increased, at first gradually but eventually dramatically, and then there was an ominous pause in earthquake activity. Chinese seismologists interpreted this “time gap” as an indication of an impending earthquake. The population was ordered to vacate their homes several hours before the province was struck by a magnitude 7.4 earthquake that destroyed cities and communities. Because of the successful prediction of this earthquake, the epicenter of which was near to the city of Haicheng, the death toll among the 3,000,000 inhabitants of the province was very low.

A technique, no longer in favor, but which at one time looked promising for predicting the time and magnitude of an earthquake is the dilatancy hypothesis, based upon systematic variations in the ratio of the travel-times of P-waves and S-waves which originated in the focal volume of larger shocks. Russian seismologists noticed that prior to an earthquake the travel-time ratio \( t_p/t_s \) changed systematically: at first it decreased by up to 5%, then it returned to normal values just before the earthquake.

The observations have been attributed to changes in the dilatancy of the ground. Laboratory experiments have shown that, before a rock fractures under stress, it develops minute cracks which cause the rocks to dilate or swell. This dilatancy alters the P-wave velocity, which drops initially (thereby increasing \( t_p \)) as, instead of being water filled, the new volume of the dilated pores at first fills with air. Later, water seeps in under pressure, replaces the air and the \( t_p/t_s \) ratio returns to normal values. At this point an earthquake is imminent. The time for which the ratio remains low is a measure of the strain energy that is stored and therefore a guide to the magnitude of the earthquake that is eventually unleashed. Initial success in predicting small earthquakes with the dilatancy model led to a period of optimism that an ultimate solution to earthquake prediction had been found. Unfortunately, it has become apparent that the dilatancy effect is not universal, and its importance appears to be restricted only to certain kinds of earthquakes.

In summary, present seismicity patterns, in conjunction with our knowledge of where historic earthquakes have occurred, permit reasonable judgements of where future earthquakes are most likely to be located. However, despite years of effort and the investigation of various scientific methods, it is still not possible to predict reliably when an earthquake is likely to happen in an endangered area.

### 3.5.10 Earthquake control

Earthquakes constitute a serious natural environmental hazard. Despite great efforts by scientists in various countries, successful prediction is not yet generally possible. Consequently, the protection of people against seismic hazard depends currently on the identification of especially perilous areas (such as active faults), the avoidance of these as the sites of constructions, the development and enforcement of appropriate building codes, and the education and training of the population in emergency procedures to be followed during and in the aftermath of a shock. Unfortunately, many densely populated regions are subject to high seismic risk. It is impossible to prevent the cumulation of strain in a region subject to tectonic earthquakes; the efforts of the human race are not likely to have much effect on the processes of plate tectonics! However, it may be possible to influence the manner in which the strain energy is released. The catastrophic earthquakes are those in which a huge amount of strain energy that has accumulated over a long period of time is suddenly released in a single event. If the energy could be released progressively over a longer period of time in many smaller shocks, the violence and disastrous consequences of a major earthquake might be avoided. The intriguing possibility of this type of earthquake control has been investigated in special situations.

In 1962 the US Army began to dispose of liquid toxic waste from the manufacture of chemical weapons by injection into a well, more than 3 km deep, near Denver, Colorado. Although the region had been devoid of earthquake activity for the preceding 80 years, earthquakes began to occur several weeks after pumping started. Until 1965, when waste injection was halted, more than 1000 earthquakes were recorded. They were mostly very small, of the microearthquake category, but some had Richter magnitudes as high as 4.6. When pumping was halted, the seismicity ceased; when pumping was resumed, the earthquake activity started anew. It was conjectured that the liquid waste had seeped into old faults, and by acting as a kind of lubricant or by increasing the pore pressure, had repeatedly permitted slippage, with an accompanying small earthquake. This incident suggested that it might be
possible to control fault motion, either by injecting fluids to lubricate sections of the fault plane, or by pumping out fluid to lock the fault. This opened the intriguing possibility that, by making alternate use of the two processes, the slippage on a fault might be controlled so that it took place by a large number of small earthquakes rather than by a few disastrous earthquakes.

In 1969 the US Geological Survey carried out a test of this lubrication effect in the depleted Rangely oil field in western Colorado. There were many disused wells in the oil field, through which fluids were pumped in and out of the ground over a considerable area. Meanwhile the local seismicity was monitored. This controlled test agreed with the observations at the Denver site, confirming that the earthquake activity correlated with the amount of fluid injected. Moreover, it was established that the earthquake activity increased when the pore pressure exceeded a critical threshold value, and it ceased when the pressure dropped below this value as the fluid was withdrawn.

Despite the apparent success of this experiment, it was agreed that further testing was necessary to explore the validity of the method. An obvious difficulty of testing the modification of seismic activity on critical faults is that the tests must be made in remote areas so as to avoid costly damage caused by the testing. The conditions under which the method may be applicable have not been established definitively.

### 3.5.11 Monitoring nuclear explosions

Since 1963 most nuclear explosions have been conducted underground to prevent the dangerous radioactive fallout that accompanies nuclear explosions conducted under-water or in the atmosphere. The detection and monitoring of such testing activity became important tasks for seismologists. The bilateral Threshold Test Ban Treaty of 1974 between the former Soviet Union and the USA prohibited underground testing of nuclear devices with a yield greater than 150 kilotons of TNT equivalent, which corresponds roughly to an earthquake with magnitude about 6. Current efforts are underway to establish a global monitoring system to verify compliance with a future Comprehensive Test Ban Treaty. The system should detect nuclear explosions with a yield as low as one kiloton (a well-coupled kiloton explosion has a magnitude of around 4). Detection of these events at distances of several thousand kilometers, and discriminating them from the approximately 7000 earthquakes that occur annually with magnitudes of 4 or above (Table 3.3) poses a monumental challenge to seismologists.

In order to achieve the high detection capability needed to monitor underground testing, many so-called seismic arrays have been set up. An array consists of several individual seismometers, with spacing on the order of a kilometer or less, that feed their output signals in parallel into a centralized data-processing center. By filtering, delaying and summing the signals of the individual instruments, the incoherent noise is reduced and the coherent signal is increased, thus improving the signal-to-noise ratio significantly over that for a single sensor. A local seismic disturbance arrives at the array on an almost horizontal path and triggers the individual seismometers at successively different times, whereas a teleseismic arrival from a very distant source reaches all seismometers in the array at nearly the same time along a steeply inclined path. The development of seismic arrays permitted the analysis of distant weak events. The enhanced sensitivity led to several advances in seismology. Features of the deep structure of the Earth (e.g., the inner core) could be investigated, earthquake location became more accurate and the analysis of focal mechanisms received a necessary impetus. These improvements were essential because of the need to identify correctly the features that distinguish underground nuclear explosions from small earthquakes.

An earthquake is the result of sudden motion of crustal blocks on opposite sides of a fault-plane. The radiation pattern of P-wave amplitude has four lobes of alternating compression and dilatation (Fig. 3.51). The first motions at the surface of the Earth are either pushes away from the source or tugs toward it, depending on the geometry of the
3.6 SEISMIC WAVE PROPAGATION

3.6.1 Introduction

A seismic disturbance is transmitted by periodic elastic displacements of the particles of a material. The progress of the seismic wave through a medium is determined by the advancement of the wavefront. We now have to consider how the wave behaves at the boundary between two media. Historically, two separate ways of handling this problem developed independently in the seventeenth century. One method, using Huygens’ principle, describes the behavior of wavefronts; the other, using Fermat’s principle, handles the geometry of ray paths at the interface. The eikonal equation (Section 3.3.2.5) establishes that these two methods of treating seismic wave propagation are equivalent.

In the Earth’s crust the velocities of P- and S-waves are often proportional to each other. This follows from Eqs. 3.39 and 3.47, which give the body-wave velocities in terms of the Lamé constants $\lambda$ and $\mu$. For many rocks, Poisson’s relation $\lambda = \mu$ applies (see Box 3.1), and so

$$\frac{\alpha}{\beta} = \sqrt{\frac{\lambda + 2\mu}{\mu}} = \sqrt{3} \quad (3.98)$$

For brevity, the following discussion handles P-waves only, which are assumed to travel with velocities $\alpha_1$ and $\alpha_2$ in the two media. However, we can equally apply the analyses to S-waves, by substituting the appropriate shear-wave velocities $\beta_1$ and $\beta_2$ for the media.

3.6.2 Huygens’ principle

The passage of a wave through a medium and across interfaces between adjacent media was first explained by the seventeenth century Dutch mathematician and physicist, Christiaan Huygens, who formulated a principle for the propagation of light as a wave, rather than as the stream of particles visualized by his great and influential contemporary, Sir Isaac Newton. Although derived for the laws of optics, Huygens’ principle (1678) can be applied equally to any kind of wave phenomenon. The theory is based on simple geometrical constructions and permits the future position of a wavefront to be calculated if its present position is known. Huygens’ principle can be stated: “All points on a wavefront can be regarded as point sources for the production of new spherical waves; the new wavefront is the tangential surface (or envelope) of the secondary wavelets.”

This principle can be illustrated simply for a plane wavefront (Fig. 3.52), although the method also applies to curved wavefronts. Let the wavefront initially occupy the position AB and let the open circles represent individual particles of the material in the wavefront. The particles are agitated by the arrival of the wavefront and act as sources of secondary waves. If the seismic velocity of the material is $V$, the distance travelled by each wavelet after time $t$ is $Vt$ and it describes a small sphere around its source particle. If the original wavefront contained numerous closely spaced particles instead of a discrete number, the plane CD tangential to the small wavelets would represent the new position of the wavefront. It is also planar, and lies at a perpendicular distance $Vt$ from the original wavefront. In their turn the particles in the wavefront CD act as sources for new secondary wavelets, and the process is repeated. This principle can be used to derive the laws of reflection and refraction of seismic waves at an interface, and also to describe the process of diffraction by which a wave is deflected at a corner or at the edge of an object in its path.

3.6.2.1 The law of reflection using Huygens’ principle

Consider what happens to a plane P-wave travelling in a medium with seismic velocity $\alpha_1$ when it encounters the boundary to another medium in which the P-wave velocity is $\alpha_2$ (Fig. 3.53). At the boundary part of the energy of
the incident wave is transferred to the second medium, and the remainder is reflected back into the first medium. If the incident wavefront AC first makes contact with the interface at A it agitates particles of the first medium at A and simultaneously the particles of the second medium in contact with the first medium at A. The vibrations of these particles set up secondary waves that travel away from A, back into the first medium as a reflected wave with velocity $\alpha_1$ (and onward into the second medium as a refracted wave with velocity $\alpha_2$).

By the time the incident wavefront reaches the interface at B all particles of the wavefront between A and B have been agitated. Applying Huygens’ principle, the wavefront of the reflected disturbance is the tangent plane to the secondary wavelet in the first medium. In Fig. 3.53 this is represented by the tangent BD from B to the circle centered at A, the first point of contact with the boundary. In the time $t$ that elapses between the arrival of the plane wave at A and its arrival at B, the incident wavefront travels a distance $AB$ and the secondary wavelet from A travels the equal distance $AD$. The triangles $ABC$ and $ABD$ are congruent. It follows that the reflected wavefront makes the same angle with the interface as the incident wave. It is customary to describe the orientation of a plane wave by the direction of its normal. The angle between the normal to the interface and the normal to the incident wavefront is called the angle of incidence ($i$); the angle between the normal to the interface and the normal to the reflected wavefront is called the angle of reflection ($r$). This application of Huygens’ principle to plane seismic waves shows that the angle of reflection is equal to the angle of incidence ($i = r$). This is known as the law of reflection.

**3.6.2.2 The law of refraction using Huygens’ principle**

The discussion of the interaction of the incident wave with the boundary can be extended to cover the part of the disturbance that travels into the second medium (Fig. 3.54). This disturbance travels with the velocity $\alpha_2$ of the second medium. Let $t$ be the time taken for the incident wavefront in the first medium to advance from $C$ to $B$; then $BC = \alpha_1 t$. In this time all particles of the second medium between A and B have been agitated and now act as sources for new wavelets in the second medium. When the incident wave reaches B, the wavelet from A in the second medium has spread out to the point E, where $AE = \alpha_2 t$. The wavefront in the second medium is the tangent BE from B to the circle centered at A. The angle of incidence ($i$) is defined as before; the angle between the normal to the interface and the normal to the transmitted wavefront is called the angle of refraction ($r$). Comparison of the triangles ABC and ABE shows that $BC = AB \sin i$, and $AE = AB \sin r$. Consequently,

$$\frac{AB \sin i}{AB \sin r} = \frac{BC}{AE} = \frac{\alpha_1}{\alpha_2}$$

Equation (3.100) is called the law of refraction for plane seismic waves. Its equivalent in optics is often called Snell’s law, in recognition of its discoverer, the Dutch mathematician Willebrod Snellius (or Snell).

**3.6.2.3 Diffraction**

The laws of reflection and refraction derived above with the aid of Huygens’ principle apply to the behavior of plane seismic waves at plane boundaries. When a plane or spherical seismic wave encounters a pointed obstacle or discontinuous surface, it experiences diffraction. This phenomenon allows the wave to bend around the obstacle, penetrating what otherwise would be a shadow zone for the wave. It is the diffraction of sound waves, for example, that allows us to hear the voices of people who are still invisible to us around a corner, or on the other side of a high fence. Huygens’ principle also gives an explanation for diffraction, as illustrated by the following simple case.

Consider the normal incidence of a plane wave on a straight boundary that ends at a sharp corner B
3.6 SEISMIC WAVE PROPAGATION

(Fig. 3.55). The incident wave is reflected along the entire length AB, with each particle of AB acting as a secondary source according to Huygen’s principle. Beyond the edge B the incident wavefronts cannot be reflected. The plane wavefront passes by the edge B, so that the point C should lie in the shadow of AB. However, the corner B also acts as a source of secondary wavelets, part of which contribute to the reflected wavefront and part pass into the shadow zone. The intensity of the wave diffracted into the shadow zone is weaker than in the main wavefront, and it decreases progressively with increasing angle away from the direction of travel of the incident wavefront.

3.6.3 Fermat’s principle

The behavior of seismic ray paths at an interface is explained by another principle of optics that was formulated – also in the seventeenth century – by the French mathematician Pierre de Fermat. As applied to seismology, Fermat’s principle states that, of the many possible paths between two points A and B, the seismic ray follows the path that gives the shortest travel-time between the points. If \( ds \) is the element of distance along a ray path and \( c \) is the seismic velocity over this short distance, then the travel-time \( t \) between A and B is minimum. Thus,

\[
t = \int \frac{ds}{c} = \text{minimum}
\]  

(3.101)

Generally, when the velocity varies continuously with position, the determination of the ray path is intricate. In the case of a layered medium, in which the velocity is constant in each layer, Fermat’s principle provides us with an independent method for determining the laws of reflection and refraction.

3.6.3.1 The law of reflection using Fermat’s principle

Consider the reflection of a seismic ray in a medium with constant P-wave velocity \( \alpha \), at the boundary to another medium (Fig. 3.56). For convenience we take the boundary to be horizontal. Let A be a point on the incident ray at a vertical distance \( h \) from the boundary and let B be the corresponding point on the reflected ray. Let C and D be the nearest points on the boundary to A and B, respectively. Further, let \( d \) be the horizontal separation AB, and let O be the point of reflection on the interface at a horizontal distance \( x \) from C; then OD is equal to \( (d-x) \) and we can write for the travel-time \( t \) from A to B:

\[
t = \frac{AO}{c_1} + \frac{OB}{c_2} = \frac{1}{c_1} \left[ \sqrt{h^2 + x^2} + \sqrt{h^2 + (d-x)^2} \right]
\]  

(3.102)

According to Fermat’s principle the travel-time \( t \) must be a minimum. The only variable in Eq. (3.102) is \( x \). To find the condition that gives the minimum travel-time we differentiate \( t \) with respect to \( x \) and set the result equal to zero:

\[
\frac{dt}{dx} = \frac{1}{c_1} \left( \frac{x}{\sqrt{h^2 + x^2}} - \frac{(d-x)(d-x)}{\sqrt{h^2 + (d-x)^2}} \right) = 0
\]  

(3.103)

By inspection of Fig. 3.56 the relationships of these expressions to the angle of incidence \( i \) and the angle of reflection \( i' \) are evident. The first expression inside the brackets is \( \sin i \) and the second is \( \sin i' \). The condition for the minimum travel-time is again \( i' = i \); the angle of reflection equals the angle of incidence.
3.6.3.2 The law of refraction using Fermat’s principle

We can use a similar approach to determine the law of refraction. This time we study the passage of the seismic ray from a medium with velocity $v_1$ into a medium with higher velocity $v_2$ (Fig. 3.57). Let A again be a point on the incident ray at a vertical distance $h$ from a point C on the interface. The ray traverses the boundary at O, a horizontal distance $x$ from C. Let B now be a point on the ray in the second medium at a distance $h$ from D, the closest point on the interface. The distance CD is $d$, so that again OD is equal to $(d - x)$. The travel-time $t$ which we have to minimize is given by

$$t = \frac{AO}{v_1} + \frac{OB}{v_2} = \frac{\sqrt{h^2 + x^2}}{v_1} + \frac{\sqrt{h^2 + (d - x)^2}}{v_2}$$

(3.104)

Differentiating Eq. (3.104) with respect to $x$ and setting the result equal to zero gives us the condition for the minimum value of $t$:

$$\frac{x}{\sqrt{h^2 + x^2}} = \frac{(d - x)}{\sqrt{h^2 + (d - x)^2}} = 0$$

(3.105)

By reference to Fig. 3.57 we can write this expression in terms of the sines of the angles of incidence ($i$) and refraction ($r$). This application of Fermat’s principle to the seismic ray paths gives again the law of refraction that we derived by applying Huygens’ principle to the wavefronts (Eq. (3.100)). It can also be stated as

$$\frac{\sin i_1}{\alpha_1} = \frac{\sin r_1}{\alpha_2}$$

(3.106)

In this example we have assumed that $\alpha_2 > \alpha_1$. As it passes from the medium with lower velocity into the medium with higher velocity the refracted ray is bent away from the normal to the boundary, giving an angle of refraction that is greater than the angle of incidence ($r > i$). Under the opposite conditions, if $\alpha_1 < \alpha_2$, the refracted ray is bent back toward the normal and the angle of refraction is less than the angle of incidence ($r < i$).

3.6.4 Partitioning of seismic body waves at a boundary

The conditions that must be fulfilled at a boundary are that the normal and tangential components of stress, as well as the normal and tangential components of the displacements, must be continuous across the interface. If the normal (or tangential) stress were not continuous, the point of discontinuity would experience infinite acceleration. Similarly, if the normal displacements were not continuous, a gap would develop between the media or parts of both media would overlap to occupy the same space; discontinuous tangential displacements would result in relative motion between the media across the boundary. These anomalies are impossible if the boundary is a fixed surface that clearly separates the media.

As a result of the conditions of continuity, a P-wave incident on a boundary energizes the particles on each side of the boundary at the point of incidence, and sets up four waves. The energy of the incident P-wave is partitioned between P- and S-waves that are reflected from the boundary, and other P- and S-waves that are transmitted into the adjacent layer. The way in which this takes place may be understood by considering the particle motion that is induced at the interface.

The particle motion in the incident P-wave is parallel to the direction of propagation. At the interface the vibration of particles of the lower layer can be resolved into a component perpendicular to the interface and a component parallel to it in the vertical plane containing the incident P-wave. In the second layer each of these motions can in turn be resolved into a component parallel to the direction of propagation (a refracted P-wave) and a component perpendicular to it in the vertical plane (a refracted SV-wave). Because of continuity at the interface, similar vibrations are induced in the upper layer, corresponding to a reflected P-wave and a reflected SV-wave, respectively.

Let the angles between the normal to the interface and the ray paths of the P- and S-waves in medium 1 be $r_p$ and $i_p$, respectively, and the corresponding angles in medium 2 be $r_s$ and $i_s$ (Fig. 3.58). Applying Snell’s law to both the reflected and refracted P- and S-waves gives

$$\frac{\sin i_1}{\alpha_1} = \frac{\sin r_1}{\alpha_2} = \frac{\sin r_s}{\beta_2} = \frac{\sin i_s}{\beta_1}$$

(3.107)

By similar reasoning it is evident that an incident SV-wave also generates vibrations that have components normal and parallel to the interface, and will set up refracted and reflected P- and SV-waves. The situation is different for an incident SH-wave, which has no component of motion normal to the interface. In this case only refracted and reflected SH-waves are created.
3.6 SEISMIC WAVE PROPAGATION

3.6.4.1 Subcritical and supercritical reflections, and critical refraction

Let O be a seismic source near the surface of a uniformly thick horizontal layer with P-wave velocity $\alpha_1$ that lies on top of a layer with higher velocity $\alpha_2$ (Fig. 3.59). Consider what happens to seismic rays that leave O and arrive at the boundary with all possible angles of incidence. The most simple ray is that which travels vertically to meet the boundary with zero angle of incidence at the point N. This normally incident ray is partially reflected back along its track, and partially transmitted vertically into the next medium without change of direction. As the angle of incidence increases, the point of incidence moves from N towards C. The transmitted ray experiences a change of direction according to Snell’s law of refraction, and the ray reflected to the surface is termed a subcritical reflection.

The ray that is incident on the boundary at C is called the critical ray because it experiences critical refraction. It encounters the boundary with a critical angle of incidence. The corresponding refracted ray makes an angle of refraction of 90°; with the normal to the boundary. As a result, it travels parallel to the boundary in the top of the lower layer with faster velocity $\beta_2$. The sine of the angle of refraction of the critical ray is unity, and we can calculate the critical angle, $i_c$, by applying Snell’s law:

$$\sin i_c = \frac{\alpha_1}{\alpha_2}$$  \hspace{1cm} (3.108)

The critical ray is accompanied by a critical reflection. It reaches the surface at a critical distance ($x_c$) from the source at O. The reflections that arrive inside the critical distance are called subcritical reflections. At angles up to the critical angle, refracted rays pass into the lower medium, but for rays incident at angles greater than the critical angle, refraction is no longer possible. The seismic rays that are incident more obliquely than the critical angle are reflected almost completely. These reflections are termed supercritical reflections, or simply wide-angle reflections. They lose little energy to refraction, and are thus capable of travelling large distances from the source in the upper medium. Supercritical reflections are recorded with strong amplitudes on seismograms at distant stations.

3.6.5 Reflection seismology

Reflection seismology is directed primarily at finding the depths to reflecting surfaces and the seismic velocities of subsurface rock layers. The techniques of acquiring and processing reflection seismology data have been developed and refined to a very high degree of sophistication as a result of the intensive application of this method in the search for petroleum. The principle is simple. A seismic signal (e.g., an explosion) is produced at a known place at a known time, and the echoes reflected from the boundaries between rock layers with different seismic velocities and densities are recorded and analyzed. Compact, designed, robust, electromagnetic seismometers – called “geophones” in industrial usage – are spread in the region of subcritical reflection, within the critical distance from the shot-point, where no refracted arrivals are possible. Within this distance the only signals received are the wave that travels directly from the shot-point to the geophones and the waves reflected at subsurface interfaces. Surface waves are also recorded and constitute an important disturbing “noise,” because they interfere with the reflected signal. The closer the geophone array is located to the shot-point, the more nearly the paths of the reflected rays travel vertically. Reflection seismic data are most usually acquired along profiles that cross geological structures as nearly as possible normal to the strike of the structure. The travel-times recorded at the geophones along a profile are plotted as a two-dimensional cross-section of the structure. In recent years, three-dimensional
surveying, which covers the entire subsurface, has become more important.

Several field procedures are in common use. They are distinguished by different layouts of the geophones relative to the shot-point. The most routine application of reflection seismology is in continuous profiling, in which the geophones are laid out at discrete distances along a profile through the shot-point. To reduce seismic noise, each recording point is represented by a group of interconnected geophones. After each shot the geophone layout and shot-point are moved a predetermined distance along the profile, and the procedure is repeated. Broadly speaking, there are two main variations of this method, depending on whether each reflection point on the reflector is sampled only once (conventional coverage) or more than once (redundant coverage).

The most common form of conventional coverage is a split–spread method (Fig. 3.60), in which the geophones are spread symmetrically on either side of the shot-point. If the reflector is flat-lying, the point of reflection of a ray recorded at any geophone is below the midpoint of the ray path. For a shot-point at Q the rays QAP and QBR that are reflected to geophones at P and R represent extreme cases. The two-way travel-time of the ray QAP gives the depth of the reflection point A, which is plotted below the mid-point of QP. Similarly, B is plotted below the mid-point of QR. The split–spread layout around the shot-point Q gives the depths of reflection points along AB, which is half the length of the geophone spread PR. The shot-point is now moved to the point R, and the geophones between P and Q are moved to cover the segment RS. From the new shot-point R the positions of reflection points in the segment BC of the reflector are obtained. The ray RBQ from shot-point R to the geophone at Q has the same path as the ray QBR from shot-point Q to the geophone at R. By successively moving the shot-point and half of the split–spread geophone layout a continuous coverage of the subsurface reflector is obtained.

Redundant coverage is illustrated by the common-midpoint method, which is routinely employed as a means of reducing noise and enhancing the signal-to-noise ratio. Commonly 24 to 96 groups of geophones feed recorded signals into a multi-channel recorder. The principle of common-midpoint coverage is illustrated for a small number of 11 geophone groups in Fig. 3.61. When a shot is fired at A, the signals received at geophones 3–11 give subsurface coverage of the reflector between points a and e. The shot-point is now moved to B, which coincides with the position occupied by geophone 2 for the first shot, and the geophone array is moved forward correspondingly along the direction of the profile to positions B–E. From shot-point B the subsurface coverage of the reflector is between points b and f. The reflector points b to e are common to both sets of data. By repeatedly moving the shot-point and geophone array in the described manner, each reflecting point of the interface is sampled multiply. For example, in Fig. 3.61 the reflecting point d is sampled multiply by the rays Ad9, Bd8, Cd7, etc. The lengths of these ray paths are different. During subsequent data-processing the reflection travel-times are corrected for normal moveout, which is a geometrical effect related to geophone distance from the shot-point. The records are then stacked, which is a procedure for enhancing the signal-to-noise ratio.

### 3.6.5.1 Reflection at a horizontal interface

The simplest case of seismic reflection is the two-dimensional reflection at a horizontal boundary (Fig. 3.62). Let the reflecting bed be at depth d below the shot-point S. The ray that strikes the boundary at R is reflected to the surface and recorded by a geophone at the point G, so that the angles of incidence and reflection are equal. Let G be at a horizontal distance x from the shot-point. If the P-wave velocity is $V$, the first signal received at G is from the direct wave that travels directly along SG. Its travel-time is given by $t_d = x/V$. It is important to keep in mind that the direct wave is not a surface wave but a body wave that travels parallel to and just below the surface of the top layer. The travel-time $t$ of the reflected ray SRG is $t = \frac{SR + RG}{V}$. However, SR and RG are equal and therefore

$$ t = \frac{2d}{V} \sqrt{d^2 + \frac{h^2}{4}} \quad \text{(3.109)} $
symmetrical about the vertical time axis, which it intersects
horizontal boundary is a hyperbola. The vertical reflection time \( t_0 \) is the
time corresponding to the vertical intercept of the hyperbola with the travel-time axis.

Fig. 3.62 The travel-time versus distance curve for reflections from a
horizontal boundary is a hyperbola. The vertical reflection time \( t_0 \) is the
intercept of the hyperbola with the travel-time axis.

\[
t = \frac{2d}{V} \sqrt{1 + \frac{x^2}{4d^2}} = t_0 \sqrt{1 + \frac{x^2}{4d^2}} \quad (3.110)
\]

At \( x = 0 \) the travel-time corresponds to the vertical echo from the reflector; this “echo-time” is given by \( t_0 = 2d/V \). The quantity under the square root in Eq. (3.110)
determines the curvature of the \( t-x \) curve and is called the normal moveout factor. It arises because the ray reaching a geophone at a horizontal distance \( x \) from the shot-point has not travelled vertically between it and the reflector. Squaring both sides of Eq. (3.110) and rearranging terms gives

\[
t^2 = \frac{t_0^2}{4d^2} + \frac{x^2}{4d^2} = 1 \quad (3.111)
\]

This is the equation of a hyperbola (Fig. 3.62) that is symmetrical about the vertical time axis, which it intersects at \( t_0 \). For large distances from the shot-point \((x \gg 2d)\) the travel-time of the reflected ray approaches the travel-time of the direct ray and the hyperbola is asymptotic to the two lines \( t = \pm x/V \).

A principle goal of seismic reflection profiling is usually to find the vertical distance \( d \) to a reflecting interface. This can be determined from \( t_0 \), the two-way reflection travel-time recorded by a geophone at the shot-point, once the velocity \( V \) is known. One way of determining the velocity is by comparing \( t_0 \) with the travel-time \( t_x \) to a geophone at distance \( x \). In reflection seismology the geophones are laid out close to the shot-point and the assumption is made that the geophone distance is much less than the depth of the reflector \((x \ll d)\). Equation (3.110) becomes

\[
t_x = t_0 \left( 1 + \frac{x}{2d} \right)^{2} = t_0 \left( 1 + \frac{1}{2} \frac{x}{V} \right)^{2} + \ldots
\]

\[
t_x = t_0 \left( 1 + \frac{1}{2} \frac{x}{V} \right)^{2} \quad (3.112)
\]

The difference between the travel-time \( t_x \) and the shot-
point travel-time \( t_0 \) is the normal moveout, \( \Delta t_n = t_x - t_0 \). By
rearranging Eq. (3.112) we get

\[
\Delta t_n = \frac{x^2}{2V^2 t_0} \quad (3.113)
\]

The echo time \( t_0 \) and the normal moveout time \( \Delta t_n \) are
found from the reflection data. The distance \( x \) of the geophone from the shot-point is known and therefore the layer velocity \( V \) can be determined. The depth \( d \) of the reflecting horizon can then be found by using the formula for the echo time.

An alternative way of interpreting reflection arrival times becomes evident when Eq. (3.111) is rearranged in the form

\[
t^2 = t_0^2 + \frac{x^2}{V^2} \quad (3.114)
\]

A plot of \( t^2 \) against \( x^2 \) is a straight line that has slope \( 1/V^2 \). Its intercept with the \( t^2 \)-axis gives the square of the echo time, \( t_0 \), from which the depth \( d \) to the reflector can be found once the velocity \( V \) is known. The record at each geophone will contain reflections from several reflectors. For the first reflector the velocity determined by the \( t^2-x^2 \) method is the true interval velocity of the uppermost layer, \( V_1 \), which, in conjunction with \( t_{01} \), the first echo time, gives the thickness \( d_1 \) of the top layer. However, the ray reflected from the second interface has travelled through the first layer with interval velocity \( V_1 \) and the second layer with interval velocity \( V_2 \). The velocity interpreted in the \( t^2-x^2 \) method for this reflection, and for reflections from all deeper interfaces, is an average velocity. If the incident and reflected rays travel nearly vertically, the average velocity \( V_{a,n} \) for the reflection from the \( n \)th reflector is given by

\[
V_{a,n} = \frac{d_1 + d_2 + d_3 + \ldots + d_n}{t_1 + t_2 + t_3 + \ldots + t_n} = \frac{\sum_i d_i}{\sum_i t_i} \quad (3.115)
\]

where \( d_i \) is the thickness and \( t_i \) the interval travel-time for
the \( i \)th layer.

The \( t^2-x^2 \) method is a simple way of estimating layer
thicknesses and average velocities for a multi-layered Earth (Fig. 3.63). The slope of the second straight line gives \( V_{a,2} \) which is used with the appropriate echo time \( t_{02} \) to find the combined depth \( D_2 \) to the second interface,
given by \( D_2 = d_1 + d_2 = (V_{a,2})(t_{02}); d_1 \) is known, and so \( d_2 \)
can be calculated. The two-way travel-time in the second layer is \((t_{02} - t_{01})\) and thus the interval velocity \(V_2\) can be found. In this way the thicknesses and interval velocities of deeper layers can be determined successively.

In fact, of course, the rays do not travel vertically but are bent as they pass from one layer to another (Fig. 3.63). Moreover, the elastic properties of a layer are rarely homogeneous so that the seismic velocity is variable and the ray path in the layer is curved. Exploration seismologists have found it possible to compensate for these effects by replacing the average velocity with the root-mean-square velocity \(V_{\text{rms}}\) defined by

\[
V_{\text{rms}} = \sqrt{\frac{\sum V_i^2 t_i}{\sum t_i}}
\]

where \(V_i\) is the interval velocity and \(t_i\) the travel-time for the \(i\)th layer.

### 3.6.5.2 Reflection at an inclined interface

When the reflecting interface is inclined at an angle \(\theta\) to the horizontal, as in Fig. 3.64, the shortest distance \(d\) between the shot-points and the reflector is the perpendicular distance to the inclined plane. The paths of reflected rays on the down-dip side of the shot-point are longer than those on the up-dip side; this has corresponding effects on the travel-times. The rays obey the laws of reflection optics and appear to return to the surface from the point \(S'\), which is the image point of the shot-point with respect to the reflector. The travel-time \(t\) through the layer with velocity \(V\) is readily found with the aid of the image point. For example for the ray SRG, recorded by a geophone on the surface at \(G\), we get

\[
t = \frac{SR + RG - S'R + RG - S'G}{V}
\]

The image point \(S'\) is as far behind the reflector as the shot-point is in front: \(S'S = 2d\). In triangle \(S'SG\) the side \(SG\) equals the geophone distance \(x\) and the obtuse angle \(S'\) equals \((90^\circ + \theta)\). If we apply the law of cosines to the triangle \(S'SG\) we can solve for \(S'G\) and substitute the answer in Eq. (3.117). This gives the travel-time of the reflection from the inclined boundary:

\[
t = \frac{1}{V} \sqrt{(x^2 + 4x\sin\theta + 4d^2)}
\]
Equation (3.118) is the equation of a hyperbola whose axis of symmetry is vertical, parallel to the \( t \)-axis. For a flat reflector the hyperbola was symmetric about the \( t \)-axis (Fig. 3.62) and the minimum travel-time (echo time) corresponded to the vertical reflection below the shot-point \((x = 0)\). For an inclined reflector the minimum travel-time \( t_m \) is no longer the perpendicular path to the reflector, which would give the travel-time \( t_0 \) in Fig. 3.64. Although the perpendicular path is the shortest distance from shot-point to reflector, it is not the shortest path of a reflected ray between the shot-point and a geophone. The shortest travel-time is recorded by the geophone at a horizontal distance \( x_m \) on the up-dip side of the shot-point. The coordinates \((x_m, t_m)\) of the minimum point of the travel-time hyperbola are

\[
x_m = -2dsin\theta \\
t_m = \frac{2dcos\theta}{V}
\]  

(3.119)

In practice it is not known a priori whether a reflector is horizontal or inclined. If reflection records are not corrected for the effect of layer-dip, an error results in plotting the positions of dipping beds. The shot-point travel-time \( t_0 \) gives the direct distance to a reflector, but the path along which the echo has travelled is not known. Consider the geometry of the inclined boundary in Fig. 3.65. First arrival reflections recorded for shot-points P, Q, and R come from the true reflection points A, B, and C. If the computed reflector depths are plotted directly below the shot-points at \( A'\), \( B'\) and \( C'\), the dipping boundary will appear to lie at a shallower depth than its true position, and the apparent dip of the reflector will be less steep than the true dip. This leads to a distorted picture of the underground structure. For example, an anticline appears broader and less steep-sided than it is. Similarly, if the limbs of a syncline dip steeply enough, the first arrivals from the dipping limbs can conceal the true structure (Fig. 3.66).

This happens when the radius of curvature of the bottom of the syncline is less than the subsurface depth of its axis. Over the axis of the syncline, rays reflected from the dipping flanks may be the first to reach the shot-point geophone. The bottom of the syncline is seen as an upwardly convex reflection between two cusps (Fig. 3.67).
On an uncorrected reflection record, the appearance of a tight syncline resembles a diffraction.

Reflection seismic records must be corrected for non-vertical reflections. The correctional process is called migration. It is an essential part of a reflection seismic study. When the reflection events on seismic cross-sections are plotted vertically below control points on the surface (e.g., as the two-way vertical travel-time to a reflector below the shot-point), the section is said to be unmigrated. As discussed above, an unmigrated section misrepresents the depth and dip of inclined reflectors. A migrated section is one which has been corrected for non-vertical reflections. It gives a truer picture of the positions of subsurface reflectors.

The process of migration is complex, and requires prior knowledge of the seismic velocity distribution, which in an unexplored or tectonically complicated region is often inadequately known. Several techniques, mostly computer-based, can be used but to treat them adequately is beyond the scope of this text.

### 3.6.5.3 Reflection and transmission coefficients

The partitioning of energy between refractions and reflections at different angles of incidence on a boundary is rather complex. For example, an incident P-wave may be partially reflected and partially refracted, or it may be totally reflected, depending how steeply the incident ray encounters the boundary. The fraction of the incident P-wave energy that is partitioned between reflected and refracted P- and S-waves depends strongly on the angle of incidence (Fig. 3.68). In the case of oblique incidence at less than the critical angle, the amplitudes of the different waves are given by complicated functions of the wave velocities and the angles of incidence, reflection and refraction. The relative amounts of energy in the refracted and reflected P- and S-waves do not change much for angles of incidence up to about 15°. Beyond the critical angle, the refracted P-wave ceases, so that the incident energy is partly reflected as a P-wave and partly converted to refracted and reflected S-waves.

In practice, reflection seismology is carried out at comparatively small angles of incidence. At normal incidence on an interface a P-wave excites no tangential stresses or displacements, and no shear waves are induced. The partitioning of energy between the reflected and refracted P-waves then becomes much simpler. It depends on a property of each medium known as its **acoustic impedance**, Z, which is defined as the product of the density \( \rho \) of the medium and its P-wave velocity \( \alpha \); thus \( Z = \rho \alpha \). The solution of the equations for the amplitudes \( A_1 \) and \( A_2 \) of the reflected and refracted P-waves, respectively, in terms of the amplitude \( A_0 \) of the incident wave are given by:

\[
RC = \frac{A_1}{A_0} = \frac{Z_2 - Z_1}{Z_2 + Z_1} \frac{\rho_2 \alpha_2 - \rho_1 \alpha_1}{\rho_2 \alpha_2 + \rho_1 \alpha_1},
\]

\[
TC = \frac{A_2}{A_0} = \frac{2Z_1}{Z_2 + Z_1} = \frac{2\rho_1 \alpha_1}{\rho_2 \alpha_2 + \rho_1 \alpha_1} \quad (3.120)
\]

The amplitude ratios RC and TC are called the reflection coefficient and the transmission coefficient, respectively. As shown earlier (see Eq. (3.70)), the energy of a wave is proportional to the square of its amplitude. The fraction \( E_t \) of the incident energy that is reflected is given by the square of RC, the fraction \( E_t \) that is transmitted is equal to \( (1 - E_t) \).

When the incident wave is reflected at the surface of a medium with higher seismic impedance \( Z_2 > Z_1 \), the reflection coefficient RC is positive. This means that the
reflected wave is in phase with the incident wave. However, if the wave is incident on a medium with lower seismic impedance \((Z_2 < Z_1)\), the reflection coefficient will be negative. This implies that the reflected wave is \(180^\circ\) out of phase with the incident wave. The fraction of energy reflected from an interface is equal to \(RC^2\), and therefore does not depend on whether the incidence is from the medium of higher or lower seismic impedance.

### 3.6.5.4 Synthetic seismograms

The travel-time of a reflection from a deep boundary in a multi-layered Earth is determined by the thicknesses and seismic velocities of the layers the seismic ray traverses. The amplitude of the recorded reflection is determined by the transmission and reflection coefficients for the subsurface interfaces. If the densities and seismic velocities of subsurface layers are known (for example, from sonic and density logs in conveniently located boreholes), it is possible to reconstruct what the seismogram should look like. Sometimes the density variations are ignored, and reflection and transmission coefficients are calculated simply on the basis of seismic velocities. This approximation can often be very useful, for example in the exploration of the deep structure of the Earth’s crust with the seismic reflection method. The densities of deep layers are inaccessible directly, although they can be inferred from seismic velocities. A vertical model of seismic velocities may be available from a related refraction study. These data can be incorporated in a deep seismic reflection study to calculate a synthetic seismogram. The comparison of actual and synthetic seismograms is useful for correlating reflection events and for separating real reflections from noise signals such as multiples.

The principle is simple, but the construction is laborious. A vertically incident wave on the first boundary is resolved into reflected and transmitted components, with amplitudes corresponding to the seismic impedances above and below the interface. The transmitted wave is further subdivided at the next deeper interface into other reflected and transmitted components, and this is repeated at each subsequent boundary. Each wave is followed as it is reflected and refracted at subsurface interfaces until it eventually returns to the surface. The theoretical record consists of the superposition of the numerous events, and represents the total travel-time and amplitude of each event. Whether in a high-resolution reflection study of near-surface sedimentary layering for petroleum exploration or in an analysis of deep crustal structure, the construction of a synthetic seismogram is an exacting chore that requires the use of fast modern computers.

### 3.6.5.5 Seismic noise

Controlled-source seismology allows fine resolution of a layered underground through analysis of the seismic travel-times. However, the seismic record contains not only primary reflections, or signals, from subsurface interfaces but also spurious secondary events, or noise, that interfere with the desired signals. The ratio of the energy in the signal to that in the noise, called the signal-to-noise ratio, is a measure of the quality of a seismic record. The higher the signal-to-noise ratio the better the record; records that have a ratio less than unity are unlikely to be usable.

There are many ways in which seismic noise can be excited. They can be divided into incoherent (or random) noise and coherent noise. Incoherent noise is local in origin and is caused by shallow inhomogeneities such as boulders, roots or other non-uniformities that can scatter seismic waves locally. As a result of its local nature incoherent noise is different on the records from adjacent geophones unless they are very close. In reflection seismology it is reduced by arranging the geophones in groups or arrays, of typically 16 geophones per group, and combining the individual outputs to produce a single record. When \(n\) geophones form a group, this practice enhances the signal-to-noise ratio by the factor \(\sqrt{n}\).

Coherent noise is present on several adjacent traces of a seismogram. Two common forms result from surface waves and multiple reflections, respectively. A near-surface explosion excites surface waves (especially Rayleigh waves) that can have strong amplitudes. They travel more slowly than P-waves but often reach the geophones together with the train of subsurface reflections. The resultant “ground roll” can obscure the reflections, particularly if they are weak. The problem can be minimized by the geometry of the geophone layout (Fig. 3.69a). For example, if the 16 geophones in a group are laid out at equal distances to cover a complete wavelength of the Rayleigh wave, the signals of individual geophones are effectively integrated to a low value. This procedure is only partially effective. Although most of the ground roll is related to Rayleigh waves, part is thought to have more complex near-surface origins. Another method of reducing the effects of ground roll is frequency filtering. The frequency of the ground roll is often lower than that of the reflected P-waves, which allows attenuation of this type of coherent noise by including a high-pass filter in the geophone circuit or in the subsequent processing to cut out low frequencies.

Multiple reflections are a very common source of coherent noise in a layered medium. They can originate in several ways, the most serious of which is the surface multiple (Fig. 3.69b). The reflection coefficient at the free surface of the Earth is high (in principle, \(RC=-1\)), and multiple reflections can occur between the surface and a reflecting interface. The travel-time for a first-order multiple at near-vertical incidence is double that of the primary signal. A copy of the reflector is observed on the seismogram at twice the real depth (or travel-time). Higher-order multiples produce additional apparent reflectors. A further advantage of using the common-mid-point method of
3.6.6 Refraction seismology

The method of seismic refraction can be understood by applying Huygens' principle to the critical refraction at the interface between two layers. The seismic disturbance travels immediately below the interface with the higher velocity of the lower medium. It is called a head wave (or Mintrop wave, after the German seismologist who patented its use in seismic exploration in 1919). The upper and lower media are in contact at the interface and so the upper medium is forced to move in phase with the lower medium. The vibration excited at the boundary by the passage of the head wave acts as a moving source of secondary waves in the upper layer. The secondary waves interfere constructively (in the same way as a reflected wave is formed) to build plane wavefronts; the ray paths return to the surface at the critical angle within the region of supercritical reflections (Fig. 3.71). The doubly refracted waves are especially important for the information they reveal about the layered structure of the deep interior of the Earth.

3.6.6.1 Refraction at a horizontal interface

The method of refraction seismology is illustrated for the case of the flat interface between two horizontal layers in Fig. 3.71. Let the depth to the interface be \( d \) and the seismic velocities of the upper and lower layers be \( V_1 \) and \( V_2 \) respectively (\( V_1 < V_2 \)). The direct ray from the shot-point at \( S \) is recorded by a geophone \( G \) at distance \( x \) on the surface after time \( x/V_1 \). The travel-time curve for the direct ray is a straight line through the origin with slope \( m_1 = 1/V_1 \). The hyperbolic \( t-x \) curve for the reflected ray intersects the time axis at the two-way vertical reflection ("echo") time \( t_0 \). At great distances from the shot-point the reflection hyperbola is asymptotic to the straight line for the direct ray.

The doubly refracted ray travels along the path \( SC \) with the velocity \( V_1 \) of the upper layer, impinges with critical angle \( i_c \) on the interface at \( C \), passes along the segment \( CD \) with velocity \( V_2 \) of the lower layer, and returns to the surface along \( DG \) with velocity \( V_1 \). The segments \( SC \) and \( DG \) are equal, \( CD = x - 2SA \) and the travel-time for the path \( SCDG \) can be written

\[
t = \frac{2SC}{V_1} + \frac{CD}{V_2}
\]  

(3.121)

i.e.,

\[
t = \frac{2d}{V_1 \cos i_c} + \frac{(x - 2d \tan i_c)}{V_2}
\]  

(3.122)

Rearranging terms and using Snell's law, \( \sin i_c = V_1/V_2 \), we get for the travel-time of the doubly refracted ray

\[
t = \frac{x}{V_2} + \frac{2d}{V_1 \cos i_c}
\]  

(3.123)
The equation represents a straight line with slope \( m_2 = 1/V_2 \). The doubly refracted rays are only recorded at distances greater than the critical distance \( x_c \). The first arrival recorded at \( x_c \) can be regarded as both a doubly refracted ray and a reflection; the travel-time line for the head wave is tangential to the reflection hyperbola at \( x_c \).

By backward extrapolation the refraction \( t-x \) curve is found to intersect the time axis at the intercept time \( t_i \), given by

\[
(3.124)
\]

Close to the shot-point the direct ray is the first to be recorded. However, the doubly refracted ray travels part of its path at the faster velocity of the lower layer, so that it eventually overtakes the direct ray and becomes the first arrival. The straight lines for the direct and doubly refracted rays cross each other at this distance, which is accordingly called the crossover distance, \( x_{cr} \). It is computed by equating the travel-times for the direct and refracted rays:

\[
(3.125)
\]

\[
(3.126)
\]

Refraction seismology gives the velocities of subsurface layers directly from the reciprocal slopes of the straight lines corresponding to the direct and doubly refracted rays. Once these velocities have been determined it is possible to compute the depth \( d \) to the interface by using either the intercept time \( t_i \) or the crossover distance \( x_{cr} \), which can be read directly from the \( t-x \) plot.
\[ d = \frac{1}{2} \sqrt{V_1^2 - V_2^2} \quad (3.127) \]
\[ d = \frac{1}{2} \sqrt{V_2^2 - V_1^2} \quad (3.128) \]

### 3.6.6.2 Refraction at an inclined interface

In practice, the refracting interface is often not horizontal. The assumption of flat layers then leads to errors in the velocity and depth estimates. When the refractor is suspected to have a dip, the velocities of the beds and the dip of the interface can be obtained by shooting a second complementary profile in the opposite direction. Suppose a refractor dips at an angle \( \theta \) as in Fig. 3.72. Shot-points A and B are located at the ends of a geophone layout that covers AB. The ray ACDB from the shot-point A strikes the interface at the critical angle \( i_c \) at C, runs as a head wave with velocity \( V_1 \) along the dipping interface, and the ray emerging at D eventually reaches a geophone at the end of the profile at B. During reverse shooting, the ray from the shot-point at B to a geophone at A traverses the same path in the reverse direction. However, the \( t-x \) curves are different for the up-dip and down-dip shots. Let \( d_A \) and \( d_B \) be the perpendicular distances from the shot-points A and B to the interface at P and Q, respectively. For the down-dip shot at A the travel-time to distance \( x \) is given by

\[ t_u = \frac{AC + DB}{V_1} + \frac{CD}{V_2} \quad (3.129) \]

The geometry of Fig. 3.72 gives the following trigonometric relationships:

\[ AC = d_A \cos i_c \quad DB = d_B \cos i_c \]
\[ PC = d_A \tan i_c \quad DQ = d_B \tan i_c \]
\[ CD = x \cos \theta - (PC + DQ) \quad (3.130) \]

we have

\[ t_u = \frac{d_A}{V_1 \cos i_c} + \frac{x \cos \theta - (d_A + d_B) \tan i_c}{V_2} \]

\[ = \frac{x \cos \theta}{V_2} + \frac{(d_A + d_B) \tan i_c}{V_1 \cos i_c} \quad (3.131) \]

Equation (3.131) can be simplified by noting that

\[ d_B = d_A + x \sin \theta \quad \text{and} \quad \frac{1}{V_2} = \frac{\sin \theta}{V_1} \quad (3.132) \]

After substitution and gathering terms the down-dip travel-time is given by

\[ t_u = \frac{x \sin \theta \cos \theta}{V_1} + \frac{x \cos \theta \sin \theta}{V_1} + \frac{2d_A \cos i_c}{V_1} \]

\[ = \frac{x}{V_1} \sin(i_c + \theta) + t_{id} \quad (3.133) \]

where \( t_{id} \) is the intercept time for the down-dip shot:

\[ t_{id} = \frac{2d_A \cos i_c}{V_1} \quad (3.134) \]

The analysis for shooting in the up-dip direction is analogous and gives

\[ t_u = \frac{x}{V_1} \sin(i_c - \theta) + t_{iu} \quad (3.135) \]

where \( t_{iu} \) is the intercept time for the up-dip shot:

\[ t_{iu} = \frac{2d_B \cos i_c}{V_1} \quad (3.136) \]

If the upper layer is homogeneous, the segments for the direct ray will have equal slopes, the reciprocals of which give the velocity \( V_1 \) of the upper layer. The segments of the \( t-x \) curves corresponding to the doubly refracted ray are different for up-dip and down-dip shooting. The total travel-times in either direction along ACDB must be equal, but the \( t-x \) curves have different intercept times. As these are proportional to the perpendicular distances to the refractor below the shot points, the up-dip intercept time \( t_{iu} \) is larger than the down-dip intercept time \( t_{id} \). This means that the slope of the up-dip refraction in Fig. 3.72 is flatter than the down-dip slope. If we interpret the reciprocal of the slope as the velocity of the lower medium, we get two apparent velocities, \( V_d \) and \( V_u \) given by
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\[ \frac{1}{V_d} = \frac{1}{V_1} \sin(i_1 + \theta) \]
\[ \frac{1}{V_u} = \frac{1}{V_1} \sin(i_1 - \theta) \]  \hspace{1cm} (3.137)

Once the real velocity \( V_1 \) and the apparent velocities \( V_d \) and \( V_u \) have been determined from the \( t-x \) curves, the dip of the interface \( \theta \) and the critical angle \( i_c \) (and from it the true velocity \( V_2 \) of the lower layer) can be computed:

\[ \theta = \frac{1}{2} \left[ \sin^{-1} \left( \frac{V_1}{V_d} \right) - \sin^{-1} \left( \frac{V_1}{V_u} \right) \right] \]  \hspace{1cm} (3.138)
\[ i_c = \frac{1}{2} \left[ \sin^{-1} \left( \frac{V_1}{V_d} \right) + \sin^{-1} \left( \frac{V_1}{V_u} \right) \right] \]  \hspace{1cm} (3.139)

If the reciprocal apparent velocities (Eq. (3.137)) are added, a simple approximation for the true velocity of the lower layer is obtained:

\[ \frac{1}{V_2} = \frac{1}{2} \left( \frac{1}{V_d} + \frac{1}{V_u} \right) \]  \hspace{1cm} (3.140)

If the refractor dip is small, \( \cos \theta \approx 1 \) (for example, if \( \theta < 15^\circ \), \( \cos \theta > 0.96 \)) and an approximate formula for the true velocity of the second layer is

\[ \frac{1}{V_2} \approx \frac{1}{2} \left( \frac{1}{V_1} + \frac{1}{V_m} \right) \]  \hspace{1cm} (3.141)

3.6.6.3 Refraction with continuous change of velocity with depth

Imagine the Earth to have a multi-layered structure with numerous thin horizontal layers, each characterized by a constant seismic velocity, which increases progressively with increasing depth (Fig. 3.73). A seismic ray that leaves the surface with angle \( i_1 \) will be refracted at each interface until it is finally refracted critically. The ray that finally returns to the surface will have an emergence angle equal to \( i_1 \). Snell’s law applies to each successive refraction (e.g., at the top surface of the \( n \)th layer, which has velocity \( V_n \))

\[ \frac{\sin i_1}{V_1} = \frac{\sin i_2}{V_2} = \cdots = \frac{\sin i_n}{V_n} = \text{constant} = p \]  \hspace{1cm} (3.142)

The constant \( p \) is called the ray parameter. It is characteristic for a particular ray with emergence angle \( i_1 \) and velocity \( V_1 \) in the surface layer. If \( V_m \) is the velocity of the deepest layer, along whose surface the ray is eventually critically refracted (\( \sin i_m = 1 \)), then the value of \( p \) must be equal to \( 1/V_m \).

As the number of layers increases and the thickness of each layer decreases, the situation is approached in which the velocity increases constantly with increasing depth. Each ray then has a smoothly curved path. If the vertical increase in velocity is linear with depth, the curved rays are circular arcs.

In the above we have assumed that the refracting interfaces are horizontal. This type of analysis is common in seismic prospecting, where only local structures and comparatively shallow depths are evaluated. The passage of seismic body waves through a layered spherical Earth can be treated to a first approximation in the same way. We can represent the vertical (radial) velocity structure by subdividing the Earth into concentric shells, each with a faster body-wave velocity than the shell above it (Fig. 3.74). Snell’s law of refraction applies to the interface between each pair of shells. For example, at point A we can write

\[ \frac{\sin i_1}{V_1} = \frac{\sin i_2}{V_2} \]  \hspace{1cm} (3.143)

Multiplying both sides by \( r_1 \) gives

\[ \frac{r_1 \sin i_1}{V_1} = \frac{r_1 \sin i_2}{V_2} \]  \hspace{1cm} (3.144)

In triangles ACD and BCD, respectively, we have

\[ d = r_1 \sin a_1 = r_2 \sin a_2 \]  \hspace{1cm} (3.145)

Combining Eqs. (3.143), (3.144) and (3.145) gives the result

\[ \frac{r_1 \sin i_1}{V_1} = \frac{r_1 \sin i_2}{V_2} = \cdots = \frac{r_n \sin i_n}{V_n} = p \]  \hspace{1cm} (3.146)
The constant \( p \) is again called the *ray parameter*, although it has different dimensions than in Eq. (3.142) for flat horizontal layers. Here the seismic ray is a straight line within each spherical layer with constant velocity. If the velocity increases continuously with depth, the seismic ray is refracted continuously and its shape is curved concavely upward. It reaches its deepest point when \( \sin i = 1 \), at radius \( r_0 \) where the velocity is \( V_0^p \); these parameters are related by the Benndorf relationship:

\[
\frac{r_{ini}}{V} = \frac{r_0}{V_0} = p
\]  

(3.147)

Determination of the ray parameter is the key to determining the variation of seismic velocity inside the Earth. Access to the Earth’s interior is provided by analysis of the travel-times of seismic waves that have traversed the various internal regions and emerge at the surface, where they are recorded. We will see in Section 3.7.3.1 that the velocity \( V \) at the deepest point of the path. The theory applies for P- and S-waves, the general velocity \( V \) being replaced by the appropriate velocity \( \alpha \) or \( \beta \), respectively.

### 3.7 Internal Structure of the Earth

#### 3.7.1 Introduction

It is well known that the Earth has a molten core. What is now general knowledge was slow to develop. In order to explain the existence of volcanoes, some nineteenth century scientists postulated that the Earth must consist of a rigid outer crust around a molten interior. It was also known in the last century that the mean density of the Earth is about 5.5 times that of water. This is much larger than the known specific density of surface rocks, which is about 2.5–3. From this it was inferred that density increased towards the Earth’s center under the effect of gravitational pressure. The density at the Earth’s center was estimated to be comparatively high, greater than 7000 kg m\(^{-3}\) and probably in the range 10,000–12,000 kg m\(^{-3}\).

It was known that some meteorites had a rock-like composition, while others were much denser, composed largely of iron. In 1897 E. Wiechert, who subsequently became a renowned German seismologist, suggested that the interior of the Earth might consist of a dense metallic core, cloaked in a rocky outer cover. He called this cloak the “Mantel,” which later became anglicized to mantle.

The key to modern understanding of the interior of the Earth – its density, pressure and elasticity – was provided by the invention of the Milne seismograph. The progressive refinement of this instrument and its systematic employment world-wide led to the rapid development of the modern science of seismology. Important results were obtained early in the twentieth century. The Earth’s fluid core was first detected seismologically in 1906 by R. D. Oldham. He observed that, if the travel-times of P-waves observed at epicentral distances of less than 100° were extrapolated to greater distances, the expected travel-times were less than those observed. This meant that the P-waves arriving at large epicentral distances were delayed in their passage through the Earth. Oldham inferred from this the existence of a central core in which the P-wave velocity was reduced. He predicted that there would be a region of epicentral distances (a “shadow zone”) in which P-waves could not arrive. About this time it was found that P- and S-waves passed through the mantle but that no S-waves arrived beyond an epicentral distance of 105°. In 1914, B. Gutenberg verified the existence of a shadow zone for P-waves in the range of epicentral distances between 105° and 143°. Gutenberg also located the depth of the core-mantle boundary with impressive accuracy at about 2900 km. A modern estimate of the radius of the core is 3485 ± 3 km, giving a mantle 2885 km thick. Gutenberg also predicted that P-waves and S-waves would be reflected from the core-mantle boundary. These waves, known today as PcP and ScS waves, were not observed until many years later. In honor of Gutenberg the core-mantle boundary is known today as the *Gutenberg seismic discontinuity*.

While studying the P-wave arrivals from an earthquake in Croatia in 1909 Andrija Mohorovicic found only a single arrival (P\(_g\)) at distances close to the epicenter. Beyond about 200 km there were two arrivals; the P\(_g\) event was overtaken by another arrival (P\(_n\)) which had evidently travelled at higher speed. Mohorovicic identified P\(_g\) as the direct wave from the earthquake and P\(_n\) as a doubly refracted wave (equivalent to a head wave) that travelled partly in the upper mantle. Mohorovicic calculated velocities of 5.6 km s\(^{-1}\) for P\(_g\) and 7.9 km s\(^{-1}\) for P\(_n\) and estimated that the sudden velocity increase occurred at a depth of 54 km. This seismic discontinuity is now called the *Mohorovicic discontinuity*, or Moho for short. It represents the boundary between the crust and mantle.
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The crustal thickness is known to be very variable. It averages about 33 km, but measures as little as 5 km under oceans and as much as 60–80 km under some mountain ranges.

The **seismological Moho** is commonly defined as the depth at which the P-wave velocity exceeds 7.6 km s\(^{-1}\). This seismic definition is dependent on the density and elastic properties of crustal and mantle rocks, and need not correspond precisely to a change of rock type. An alternative definition of the Moho as the depth where the rock types change is called the **petrological Moho**. For most purposes the two definitions of the crust–mantle boundary are equivalent.

It is now known that the crust is not homogeneous but has a vertically layered structure. In 1925 V. Conrad separated arrivals from a Tauern (Eastern Alps) earthquake of 1923 into \(P_2\) and \(S_2\) waves in an upper crustal layer and faster \(P^*\) and \(S^*\) waves that travelled with velocities 6.29 km s\(^{-1}\) and 3.57 km s\(^{-1}\), respectively, in a deeper layer. Because the \(P^*\) and \(S^*\) velocities are significantly slower than corresponding upper mantle velocities, Conrad deduced that they were head waves from a lower crustal layer. The interface separating the continental crust into an upper crustal layer and a lower crustal layer is called the **Conrad discontinuity**. Influenced by early petrological models of crustal composition and by comparison with seismic velocities in known materials, seismologists referred to the upper and lower crustal layers as the granitic layer and the basaltic layer, respectively. This petrological separation is now known to be overly simplistic. In contrast to the Moho, which is found everywhere, the Conrad discontinuity is poorly defined or absent in some areas.

The appearance of some of these arrivals on refraction seismic records from the European continental crust is illustrated in Fig. 3.75. The vertical axis in this figure shows a reduced travel-time, obtained by dividing the shot-point to receiver distance by a representative crustal velocity (6 km s\(^{-1}\) in this case) and subtracting this from the observed travel-time. This method of displaying the data prevents the plot from becoming unwieldy at large distances. The crustal direct wave \(P_6\) is represented as a nearly horizontal arrival, \(PmP\) is a P-wave reflected from the Moho, and \(P_n\) is the upper mantle head wave along the Moho. Additional intracrustal reflections are labeled \(P2P\), \(P3P\), etc.

Similar designations are used for events in seismic sections of oceanic crust. The oceanic crustal layers consist of sedimentary Layer 1, basaltic Layer 2, and gabbroic Layer 3 (Section 3.7.5.1). The direct wave in Layer 2 is called \(P_2\); the head wave at the basalt–gabbro interface is referred to as \(P_1\), and the Moho head wave is \(P_n\).

The core shadow-zone and its interpretation in terms of a fluid core were well established in 1936 when Inge Lehmann, a Danish seismologist, reported weak P-wave arrivals within the shadow zone. She interpreted these in terms of an inner core with higher seismic velocity. However, the existence of the inner core remained controversial for many years. Improved seismometer design, digital signal treatment and the setting up of seismic arrays have provided corroborating evidence. The existence of a solid inner core is also supported by analyses of the Earth’s natural vibrations.

The gross internal structure of the Earth is modelled as a set of concentric shells, corresponding to the inner core, outer core and mantle (Fig. 3.76). An important step in understanding this layered structure has been the development of travel-time curves for seismic rays that pass through the different shells. To facilitate identification of the arrivals of these rays on seismograms, a convenient shorthand notation is used. A P- or S-wave that travels from an earthquake directly to a seismometer is labelled with the appropriate letter P or S; until the margin of the core shadow-zone, the P- and S-waves follow identical curved paths. (The curvature, as explained in Section 3.6.6.3, arises from the increase in seismic velocity with depth.) A wave that reaches the seismometer after being reflected once from the crust is labelled PP (or PS), as its path consists of two identical P- or S-segments.

The energy of an incident P- or S-wave is partitioned at an interface into reflected and refracted P- and S-waves (see Section 3.6.4). A P-wave incident on the boundary between mantle and fluid outer core is refracted towards the normal to the interface, because the P-wave velocity drops from about 13 km s\(^{-1}\) to about 8 km s\(^{-1}\) at the boundary. After a second refraction it emerges beyond...
the shadow zone and is called a PKP wave (the letter K stands for Kern, the German word for core). An S-wave incident at the same point has a lower mantle velocity of about 7 km s$^{-1}$. Part of the incident energy is converted to a P-wave in the outer core, which has a higher velocity of 8 km s$^{-1}$. The refraction is away from the normal to the interface. After a further refraction the incident S-wave reaches the surface as an SKS phase. A P-wave that travels through mantle, fluid core and inner core is labelled PKIKP. Each of these rays is refracted at an internal interface. To indicate seismic phases that are reflected at the outer core boundary the letter c is used, giving rise, for example, to PcP and ScS phases (Fig. 3.76). Reflections from the inner core are designated with the letter i, as for example in the phase PKiKP.

### 3.7.2 Refractions and reflections in the Earth’s interior

If it possesses sufficient energy, a seismic disturbance may be refracted and reflected – or converted from a P-wave to an S-wave, or vice versa – many times at the several seismic discontinuities within the Earth and at its free surface. As a result, the seismogram of a large earthquake contains numerous overlapping seismic signals and the identification of individual phases is a difficult task. Late-arriving phases that have been multiply reflected or that have travelled through several regions of the Earth’s interior are difficult to resolve from the disturbance caused by earlier arrivals. In the period 1932–1939 H. Jeffreys and K. E. Bullen analyzed a large number of good records of earthquakes registered at a world-wide, though sparse, distribution of seismic stations. In 1940 they published a set of tables giving the travel-times of P- and S-waves through the Earth. A slightly different set of tables was reported by B. Gutenberg and C. F. Richter. The good agreement of the independent analyses confirmed the reliability of the results. The Jeffreys–Bullen seismological tables were used by the international seismic community as the standard of reference for many years.

The travel-time of a seismic wave to a given epicentral distance is affected by the focal depth of the earthquake, which may be as much as several hundred kilometers. The travel-time versus distance curves of some important phases are shown in Fig. 3.77 for an earthquake occurring at the Earth’s surface. The model assumes that the Earth is spherically symmetric, with the same vertical structure underneath each place on the surface. This assumption works fairly well, although it is not quite true. Lateral variations of seismic velocity have been found at many depths within the Earth. For example, there are lateral differences in seismic velocity between oceanic and continental crust, and between oceanic and continental lithosphere. At even greater depths significant lateral departures from the spherical model have been detected. These discrepancies form the basis of the branch of seismology called seismic tomography, which we will examine later.

#### 3.7.2.1 Seismic rays in a uniformly layered Earth

It is important to understand clearly the relationship between the travel-time ($t$) versus epicentral distance ($\Delta$) curves and the paths of seismic waves in the Earth, like those shown in Fig. 3.76. Consider first an Earth that
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Fig. 3.78 Seismic wave paths and their \( t-\Delta \) curves for P-waves passing through a spherical Earth with constant velocities in the mantle, outer core and inner core, respectively. (a) Development of a shadow zone when the mantle velocity \( (V_1) \) is higher than the outer core velocity \( (V_2) \). The figure shows the paths of 18 rays that leave a surface source at angular intervals of 5°. Rays 1–12 travel directly through the mantle as P-waves and emerge at progressively greater epicentral distances. The convex upwards shape of the \( t-\Delta \) curve is here due to the curved outer surface, the layer velocity being constant. Ray 13 is partially refracted into the core, and partially reflected (not shown in the figure). Because \( V_2 < V_1 \) the refracted ray is bent towards the Earth's radius, which is normal to the refracting interface. This ray is further bent on leaving the core and reaches the Earth's surface as a PKP phase at an epicentral distance greater than 180°. Rays 14 and 15 impinge more directly on the core and are refracted less severely; their epicentral distances become successively smaller and their travel-times become shorter than that of ray 13, as indicated by branch 13–15 of the \( t-\Delta \) curve. This branch is offset in time from the extrapolation of branch 1–12 because of the lower velocity in the core. The paths of rays 16, 17 and 18 (which is a straight line through mantle and core and emerges at an epicentral distance of 180°) become progressively longer, and a second branch 16–18 develops on the \( t-\Delta \) curve. The two branches meet at a sharp point, or cusp. No P-waves reach the surface in the gap between rays 12 and 15 in this simple model. There is a shadow zone between the last P-wave that just touches the core and the PKP-wave with the smallest epicentral distance. The existence of a shadow zone for P-waves is evidence for a core with lower P-wave velocities than the mantle. S-waves in the mantle follow the same ray paths as the P-waves. However, no direct S-waves arrive in the shadow zone, which indicates that the core must be fluid.

Now suppose an inner core with a constant velocity \( V_3 \) that is higher than the velocity \( V_2 \) in the outer core (Fig. 3.78b). The paths of rays 1–15 are the same as before, through the mantle and outer core. The segments 1–12 and 13–15 of the \( t-\Delta \) curve are the same as previously. Ray 16 impinges on the inner core and is sharply refracted away from the Earth's radius; on returning to the outer core it is again refracted, back towards the radius. After further refraction at the core–mantle interface this ray emerges at the Earth's surface at a smaller epicentral distance than ray 15, within the P-wave shadow-zone, as a PKIKP event. Successive PKIKP rays are bent less strongly. The PKIKP rays map out a new branch 16–18 of the \( t-\Delta \) curve (see also Fig. 3.77).

3.7.2.2 Travel-time curves for P-, PKP- and PKIKP-waves

In general the velocities of P-waves and S-waves increase with depth. As described in Section 3.6.6.3 and illustrated in Fig. 3.73a, the ray paths are curved lines, concave towards the surface. However, the explanation of the paths of P, PKP, and PKIKP phases shown in Fig. 3.79 closely follows the preceding discussion. There is a shadow zone for direct P-waves between about 103° and 143°, and no direct S-waves are found beyond 103°. The shallowest PKP ray (A in Fig. 3.79) is deviated the furthest, emerging at an epicentral distance greater than 180°. Successively deeper PKP rays (B–E) emerge at ever-smaller epicentral distances until about 143°, after which the epicentral distance increases again to almost 170° (rays F, G). It was long believed that the boundary between inner and outer core was a transitional region (called region F in standard Earth models) with higher P-wave velocity, and that PKP rays traversing this region would again emerge at smaller epicentral distances (ray H). The first rays penetrating the inner core are sharply refracted and emerge in the P-wave shadow zone. The most strongly deviated (ray I) is observed at an epicentral distance of about 110°; deeper rays (J–P) arrive at
ever-greater distances up to 180°. There are at least two branches of the $t-\Delta$ curve for $\Delta > 143^\circ$, corresponding to the PKP and PKIKP phases, respectively (Fig. 3.77). In fact, depending how the transitional region F is modelled, the $t-\Delta$ curve near 143° can have several branches.

The edges of the shadow zone defined by P and PKP phases are not sharp. One reason is the intrusion of PKIKP phases at the 143° edge. Another is the effect of diffraction of P-waves at the 103° edge (Fig. 3.79). The bending of plane waves at an edge into the shadow of an obstruction was described in Section 3.6.2.3, and explained with the aid of Huygen's principle. The diffraction of plane waves is called *Fraunhofer diffraction*. When their source is not at infinity, waves must be handled as spherical waves. Spherical wavefronts that pass an obstacle are also diffracted. This type of behavior is called *Fresnel diffraction*, and it is also explainable with Huygen's principle as the product of interference between the primary wavefront and secondary waves generated at the obstacle. Wave energy penetrates into the shadow of the obstacle, as though the wavefront were bent around the edge. In this way very deep P-waves are diffracted around the core and into the shadow zone. The intensity of the diffracted rays falls off with increasing angular distance from the diffracting edge, in this case the core–mantle boundary. Modern instrumentation enables detection of long-period diffracted P-waves to large epicentral distances (Figs. 3.77, 3.79). The velocity structure above the core–mantle boundary, in particular in the D\textsuperscript{\textit{r}}-layer (Section 3.7.5.3), has a strong influence on the ray paths, travel-times and waveforms of the diffracted waves.

### 3.7.3 Radial variations of seismic velocities

Models of the radial variations of physical parameters inside the Earth implicitly assume spherical symmetry. They are therefore “average” models of the Earth that do not take into account lateral variations (e.g., of velocity or density) at the same depth. This is a necessary first step in approaching the true distributions as long as lateral variations are relatively small. This appears to be the case; although geophysically significant, the lateral variations in physical properties remain within a few percent of the average value at any depth.

There are two main ways to determine the distributions of body-wave velocities in a spherically symmetric Earth. They are referred to as forward and inverse modeling. Both methods have to employ the same sets of observations, which are the travel-times of different seismic phases to known epicentral distances. The forward technique starts with a known or assumed variation of seismic velocities and calculates the corresponding travel-times. The inversion method starts with the observed $t-\Delta$ curves and computes a model of the velocity distributions that could produce the curves. The inversion method is the older one, in use since the early part of the twentieth century, and forms an important branch of mathematical theory. Forward modelling is a more recent method that has been successfully employed since the advent of powerful computers.

#### 3.7.3.1 Inversion of travel-time versus distance curves

In 1907 the German geophysicist E. Wiechert, building upon an evaluation of the Benndorf problem (Eq. (3.147)) by the mathematician G. Herglotz, developed an analytical method for computing the internal distributions of seismic velocities from observations made at the Earth’s surface. The technique is called inversion of travel-times, and it is considered one of the classical methods of geophysics. The observational data consist of the $t-\Delta$ curves for important seismic phases (Fig. 3.77). The clues to deciphering the velocity distributions were the Benndorf relationship for the ray
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The Herglotz–Wiechert inversion is valid for regions of the Earth in which $p$ varies monotonically with $\Delta$. It cannot be used in the Earth's crust, because conditions are too inhomogeneous. Seismic velocity distributions in the crust are deduced empirically from long seismic refraction profiles. The Herglotz–Wiechert method cannot be used where a low-velocity zone is present, because the ray does not bottom in the zone. It works well for the Earth’s mantle, but care must be taken where a seismic discontinuity is present. In the Earth’s core the refraction of P-waves at the core–mantle boundary means that no PKP waves reach their deepest point in the outer layers of the core (Fig. 3.79). However, SKS-waves bottom in the outer core. Inversion of SKS-wave travel-times complements the inversion of PKP-wave data to give the P-wave velocity distribution in the core.

3.7.3.2 Forward modelling: polynomial parametrization

The forward modelling method starts with a presupposed dependence of seismic velocity with depth. The method assumes that the variation of velocity can be expressed by a smooth polynomial function of radial distance within limited depth ranges. This procedure is called polynomial parametrization and in constructing models of the Earth’s interior it is applied to the P-wave and S-wave velocities, the seismic attenuation, and the density.

The travel-times of P- and S-waves to any epicentral distance are calculated on the basis of the spherically symmetric, layered model. The computed travel-times are compared with the observed $t-\Delta$ curves, and the model is adjusted to account for differences. The procedure is repeated as often as necessary until an acceptable agreement between the computed and real travel-times is achieved. The method requires good travel-time data for many seismic phases and involves intensive computation.

In 1981 A. M. Dziewonski and D. L. Anderson constructed a Preliminary Reference Earth Model (acronym: PREM) in which the distributions of body-wave velocities in important layers of the Earth were represented by cubic or quadratic polynomials of normalized radial distance; in thin layers of the upper mantle linear relationships were used. A similar, revised parametrized velocity model (iasp91) was proposed by B. L. N. Kennett and E. R. Engdahl in 1991. The variations of P- and S-wave velocities with depth in the Earth according to the iasp91 model are shown in Fig. 3.81.

3.7.4 Radial variations of density, gravity and pressure

In order to determine density, gravity and pressure in the Earth’s interior several simplifying assumptions must be made, which appear to be warranted by experience. The Earth is assumed to be spherically symmetric and composed of concentric homogeneous shells or layers (e.g., inner core, outer core, mantle, etc.). Possible effects of chemical and phase changes within a shell are not taken

\[
\sin i = \frac{V}{R} \frac{dr}{d\Delta} \quad (3.148)
\]

\[
\frac{R \sin i}{V} = p = \frac{dr}{d\Delta} \quad (3.149)
\]

This means that the value of $p$ for the ray emerging at epicentral distance $\Delta$ can be obtained by calculating the slope $(dr/d\Delta)$ of the travel-time curve for that distance. This is an important step in finding the velocity $V_0$ at the deepest point of the ray, at radius $r_0$, because $V_0 = r_0/p$. However, before we can find the velocity we need to know the value of $r_0$. The continuation of the analysis – known as Herglotz–Wiechert inversion – is an intricate mathematical procedure, beyond the scope of this text, which fortunately results in a fairly simple formula:

\[
\ln \frac{R}{R_0} = \frac{1}{4} \int_0^{\Delta} \cosh^{-1} \left( \frac{p(\Delta)}{p(\Delta_1)} \right) d\Delta \quad (3.150)
\]

where $p(\Delta_1)$ is the slope of the $t-\Delta$ curve at $\Delta_1$, the epicentral distance of emergence, and $p(\Delta)$ is the slope at any intermediate epicentral distance $\Delta$. Equation (3.150) can be used to integrate numerically along the ray to give the value of $r_0$ for the ray, and $V_0 = r_0/p$.
into account. Pressure and density are assumed to increase purely hydrostatically. If the distributions of seismic body-wave velocities $\alpha$ and $\beta$ are known, an important seismic parameter $\Phi$ can be defined:

$$\Phi = \alpha^2 - \frac{4}{3} \rho \beta^2 \quad (3.151)$$

By comparing Eq. (3.151) and Eq. (3.48) we see that $\Phi$ is equal to $K/p$, where $K$ is the bulk modulus and $\rho$ the density.

### 3.7.4.1 Density inside the Earth

Consider a vertical prism between depths $z$ and $dz$ (Fig. 3.82). The hydrostatic pressure increases from $p$ at depth $z$ to $(p + dp)$ at depth $(z + dz)$ because of the extra pressure due to the extra material in the small prism of height $dz$. The pressure increase $dp$ is equal to the weight $w$ of the prism divided by the area $A$ of the base of the prism, over which the weight is distributed.

$$dp = \frac{w}{A} = \frac{(\text{volume} \times \rho)g}{A} = \frac{(Adz \times \rho)g}{A} = \rho g dz = -\rho g dr \quad (3.152)$$

From the definition of bulk modulus, $K$ (Eq. (3.17)) we can write

$$K = -V \frac{d\rho}{d\rho} = \rho \frac{dp}{d\rho} \quad (3.153)$$

Combining Eqs. (3.151), (3.152) and (3.153) gives

$$\frac{-dp}{\rho g dr} = \frac{\rho}{K} = \frac{1}{\Phi} \quad (3.154)$$

Equation (3.155) is known as the Adams–Williamson equation. It was first applied to the estimation of density in the Earth in 1923 by E. D. Williamson and L. H. Adams. It yields the density gradient at radius $r$, when the quantities on the right-hand side are known. The seismic parameter $\Phi$ is known accurately, but the density $\rho$ is unknown; it is in fact the goal of the calculation. The value of gravity $g$ used in the equation must be computed separately for radius $r$. It is due only to the mass contained within the sphere of radius $r$, because external (homogeneous) shells of the Earth do not contribute to gravitation inside them. This mass is the total mass $E$ of the Earth minus the cumulative mass of all spherical shells external to $r$.

The procedure requires that a starting value for $\rho$ be assumed at a known depth. Analyses of crustal and upper mantle structure in isostatically balanced areas give estimates of upper mantle density around 3300 kg m$^{-3}$. Using this as a starting value at an initial radius $r_1$ the density gradient in the uppermost mantle can be calculated from Eq. (3.155). Linear extrapolation of this gradient to a chosen greater depth gives the density $\rho$ at radius $r_2$; the corresponding new value of $g$ can be calculated by subtracting the mass of the shell between the two depths; together with the value of $\Phi(r_2)$ the density gradient can be computed at $r_2$. This iterative type of calculation gives the variation of density with depth (or radius). Fine steps in the extrapolations give a smooth density distribution (Fig. 3.83). There are two important boundary conditions on the computed density distribution. Integrated over the Earth’s radius it must give the correct total mass ($E = 5.974 \times 10^{24}$ kg). It must also fulfil the relationship $C = 0.3308ER^2$ between Earth’s moment of inertia ($C$), mass ($E$) and radius ($R$), as explained in Section 2.4.3.

The density changes abruptly at the major seismic discontinuities (Fig. 3.83), showing that it is affected principally by changes in composition. If it could be mea-
sured at normal sea-level pressure and temperature, the density would be found to be around 4200 kg m$^{-3}$ in the mantle, 7600 kg m$^{-3}$ in the outer core and 8000 kg m$^{-3}$ in the inner core. The smooth increase in density between the major compositional discontinuities is the result of the increases in pressure and temperature with depth.

### 3.7.4.2 Gravity and pressure inside the Earth

The radial variation of gravity can be computed from the density distribution. As stated above, the value of $g(r)$ is due only to the mass $m(r)$ contained within the sphere of radius $r$. Let the density at radius $x (\leq r)$ be $\rho(x)$; the gravity at radius $r$ is then given by

$$g(r) = - \frac{G m(r)}{r^2} = - \frac{G}{r^2} \int_0^r 4\pi x^2 \rho(x) \, dx \tag{3.156}$$

A remarkable feature of the internal gravity (Fig. 3.84) is that it maintains a value close to 10 m s$^{-2}$ throughout the mantle, rising from 9.8 m s$^{-2}$ at the surface to 10.8 m s$^{-2}$ at the core–mantle boundary. It then decreases almost linearly to zero at the Earth’s center. Hydrostatic pressure is due to the force (N) per unit area (m$^2$) exerted by overlying material. The SI unit of pressure is the pascal (1 Pa = 1 N m$^{-2}$). In practice this is a small unit. The high pressures in the Earth are commonly quoted in units of gigapascal (1 GPa = 10$^9$ Pa), or alternatively in kilobars or megabars (1 bar = 10$^5$ Pa; 1 kbar = 10$^6$ Pa; 1 Mbar = 10$^{11}$ Pa = 100 GPa).

Within the Earth the hydrostatic pressure $p(r)$ at radius $r$ is due to the weight of the overlying Earth layers between $r$ and the Earth’s surface. It can be computed by integrating Eq. (3.152) using the derived distributions of density and gravity. This gives

$$p(r) = \int_r^{\infty} \rho(r) g(r) \, dr \tag{3.157}$$

The pressure increases continuously with increasing depth in the Earth (Fig. 3.84). The rate of increase (pressure gradient) changes at the depths of the major seismic discontinuities. The pressure reaches a value close to 380 GPa (3.8 Mbar) at the center of the Earth, which is about 4 million times atmospheric pressure at sea-level.

### 3.7.5 Models of the Earth’s internal structure

Once the velocity distributions of P- and S-waves inside the Earth were known the broad internal structure of the Earth – crust, mantle, inner and outer core – could be further refined. In 1940–1942 K. E. Bullen developed a model of the internal structure consisting of seven concentric shells. The boundaries between adjacent shells were located at sharp changes in the body-wave velocities or the velocity gradients. For ease of identification the layers were labelled A–G (Table 3.4); this nomenclature has been carried over into more modern models.

In the original Model A the density distribution was not well constrained. Two different density distributions ($A$ and $A'$) that fitted the known mass and moment of inertia gave disparate central densities of 17,300 kg m$^{-3}$ and 12,300 kg m$^{-3}$, respectively. In 1950 Bullen presented Earth Model B, in which the bulk modulus ($K$) and seismic parameter ($\Phi$) were assumed to vary.
smoothly with pressure below a depth of 1000 km. The model suggested a central density around 18,000 kg m$^{-3}$.

In the 1950s the development of long-period seismographs made possible the observation of the natural oscillations of the Earth. After very large earthquakes numerous modes of free oscillation are excited with periods up to about one hour (Section 3.3.4). These were first observed unambiguously after the huge Chilean earthquake of 1960. The free oscillations form an independent constraint on Earth models. The lowest-frequency spheroidal modes involve radial displacements that take place against the restoring force of gravitation and are therefore affected by the density distribution. Starting from a spherically symmetric Earth model with known distributions of density and elastic properties, the forward problem consists of calculating how such a model will reverberate. The calculated and observed normal modes of oscillation are compared and the model is adjusted until the required fit is obtained. The inverse problem consists of computing the model of density and elastic properties by inverting the frequency spectrum of the free oscillations. The parametrized model PREM, based upon the inversion of body-wave, surface-wave and free-oscillation data, is the current standard model of the Earth’s internal structure. It predicts a central density of 13,090 kg m$^{-3}$.

### 3.7.5.1 The crust

The Earth’s crust corresponds to Bullen’s region A. The structures of the crust and upper mantle are complex and show strong lateral variations. This prohibits using the inversion of body-wave travel-times to get a vertical distribution of seismic velocities. The most reliable information on crustal seismic structure comes from seismic refraction profiles and deep crustal reflection sounding. The variation of seismic velocity with depth in the crust differs according to where the profiles are carried out. Ancient continental shield domains have different vertical velocity profiles than younger continental or oceanic domains. In view of this variability any generalized petrological model of crustal structure is bound to be an oversimplification. However, with this reservation, it is still possible to summarize some general features of crustal structure, and the corresponding petrological layering.

A generalized model of the structure of oceanic crust is shown in Fig. 3.85. Oceanic crust is only 5–10 km thick. Under a mean water depth of about 4.5 km the top part of the oceanic crust consists of a layer of sediments that increases in thickness away from the oceanic ridges. The igneous oceanic basement consists of a thin (~0.5 km) upper layer of superposed basaltic lava flows underlain by a complex of basaltic intrusions, the sheeted dike complex. Below this the oceanic crust consists of gabbroic rocks.
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The vertical structure of continental crust is more complicated than that of oceanic crust, and the structure under ancient shield areas differs from that under younger basins. It is more difficult to generalize a representative model (Fig. 3.86). The most striking difference is that the continental crust is much thicker than oceanic crust. Under stable continental areas the crust is 35–40 km thick and under young mountain ranges it is often 50–60 km thick. The continental Moho is not always a sharp boundary. In some places the transition from crust to mantle may be gradual, with a layered structure. Originally, the Conrad seismic discontinuity was believed to separate an upper crustal (granitic) layer from a lower crustal (basaltic) layer. However, the Conrad discontinuity is not found in all places and there is some doubt as to its real nature; it may represent a more complicated compositional or phase boundary. Crustal velocity studies have defined two anomalous zones that often disrupt the otherwise progressive increase of velocity with depth. A low-velocity layer within the middle crust is thought to be due to intruded granitic laccoliths; it is called the sialic low-velocity layer. Together with the crust, the lid forms the lithosphere, the rigid outer shell of the Earth that takes part in plate tectonic processes (see Section 1.2). The Moho is the transition zone of positive velocity gradient, in the depth range 100–200 km, between the crust and upper mantle. Lateral differences in structure are important down to depths of at least 400 km. The uppermost mantle between the Moho and a depth of 80–120 km is rigid, with increasing P- and S-wave velocities. This layer is sometimes called the lid of the underlying low-velocity layer. With increasing depth and pressure, the velocities decrease with increasing depth. The uppermost mantle is called the sialic low-velocity layer (LVL). Its nature cannot be evaluated from body waves, because they do not bottom discontinuity (Moho), below which the P-wave velocity exceeds 7.6 km s\(^{-1}\). The Moho depth is very variable, with a global mean value around 30–40 km. A weighted average of oceanic and continental structures equal to 24.4 km is used in model PREM. The assumption of a spherically symmetric Earth does not hold well for the crust and upper mantle. Lateral differences in structure are important down to depths of at least 400 km. The uppermost mantle between the Moho and a depth of 80–120 km is rigid, with increasing P- and S-wave velocities. This layer is sometimes called the lid of the underlying low-velocity layer. Together with the crust, the lid forms the lithosphere, the rigid outer shell of the Earth that takes part in plate tectonic processes (see Section 1.2). The lithosphere is subdivided laterally into tectonic plates that may be as large as 10,000 km across (e.g., the Pacific plate) or as small as a few thousand kilometers (e.g., the Philippines plate). The plates are very thin in comparison to their horizontal extent.

An abrupt increase of P- and S-wave velocities by 3–4% has been observed at around 220 ± 30 km depth; it is called the Lehmann discontinuity. Like the Conrad discontinuity in the crust it is not found everywhere and its true meaning is in question. Between the lid and the Lehmann discontinuity, in the depth range 100–200 km, body-wave velocity gradients are weakly negative, i.e., the velocities decrease with increasing depth. This layer is called the low-velocity layer (LVL). Its nature cannot be evaluated from body waves, because they do not bottom

3.7.5.2 The upper mantle

In his 1942 model of the Earth’s interior (see Table 3.4) Bullen made a distinction between the upper mantle (layers B and C) and the lower mantle (layer D). The upper mantle is characterized by several discontinuities of body-wave velocities and steep velocity gradients (Fig. 3.87). The top of the mantle is defined by the Mohorovicic

Fig. 3.85 Generalized petrological model and P-wave velocity–depth profile for oceanic crust.

The Moho depth is very variable, with a global mean value around 30–40 km. A weighted average of oceanic and continental structures equal to 24.4 km is used in model PREM. The assumption of a spherically symmetric Earth does not hold well for the crust and upper mantle. Lateral differences in structure are important down to depths of at least 400 km. The uppermost mantle between the Moho and a depth of 80–120 km is rigid, with increasing P- and S-wave velocities. This layer is sometimes called the lid of the underlying low-velocity layer. Together with the crust, the lid forms the lithosphere, the rigid outer shell of the Earth that takes part in plate tectonic processes (see Section 1.2). The lithosphere is subdivided laterally into tectonic plates that may be as large as 10,000 km across (e.g., the Pacific plate) or as small as a few thousand kilometers (e.g., the Philippines plate). The plates are very thin in comparison to their horizontal extent.

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in the layer and its lower boundary is not sharp. The evidence for understanding the LVL comes from the inversion of surface-wave data. Only long-period surface waves with periods longer than about 200 s can penetrate to the depths of the base of the LVL. The data from surface waves are not precise. Their depth resolution is poor and only the S-wave velocity can be determined. Thus, the top and bottom of the LVL are not sharply defined.

The LVL is usually associated with the asthenosphere, which also plays an important role in plate tectonic theory. The decreases in seismic velocities are attributed to reduced rigidity in this layer. Over geological time intervals the mantle reacts like a viscous medium, with a viscosity that depends on temperature and composition. From the point of view of plate tectonics the asthenosphere is a viscous layer that decouples the lithosphere from the deeper mantle; by allowing slow convection, it permits or promotes the relative motions of the global plates. The LVL and asthenosphere reflect changes of rheological properties of the upper mantle material. Brittle behavior in the crust and lithosphere gives way with increasing depth to ductile behavior in the low-rigidity asthenosphere. The brittle–ductile transition is gradual and depends on properties such as the rock composition, geothermal gradient, initial crustal thickness and strain rate. It probably occurs differently in oceanic and continental lithosphere, as suggested in Fig. 2.69.

The composition of the upper mantle is generally taken to be peridotitic, with olivine \([\text{Mg},\text{Fe}]_2\text{SiO}_4\] as the dominant mineral. With increasing depth the hydrostatic pressure increases and eventually causes high-pressure transformation of the silicate minerals. This is reflected in the seismic properties. Travel-time–Δ curves of body waves show a distinct change in slope at epicentral distances of about 20°. This is attributed to a discontinuity in mantle velocities at a depth of around 400 km (Fig. 3.87). The 400 km discontinuity (or 20° discontinuity) is interpreted as due to a petrological change from an olivine-type lattice to a more closely packed spinel-type lattice.

The lid, low-velocity layer and the zone down to the 400 km discontinuity together correspond to layer B in Bullen's Earth Model A. A further seismic discontinuity occurs at a depth of 650–670 km. This is a major feature of mantle structure that has been observed world-wide. In the transition zone between the 400 km and 670 km discontinuities there is a further change in structure from \(\beta\)-spinel to \(\gamma\)-spinel, but this is not accompanied by appreciable changes in physical properties. At 670 km the spinel transforms to perovskite. This transition zone corresponds to the upper part of Bullen's layer C.

### 3.7.5.3 The lower mantle

The lower mantle is now classified as the part below the important seismic discontinuity at 670 km. Its composition is rather poorly known, but it is thought to consist of oxides of iron and magnesium as well as iron–magnesium silicates with a perovskite structure. The uppermost part of the lower mantle between 670 and 770 km depth has a high positive velocity gradient and corresponds to the lower part of Bullen's layer C. Beneath it lies Bullen's layer D', which represents a great thickness of normal mantle, characterized by smooth velocity gradients and the absence of seismic discontinuities.

Just above the core–mantle boundary an anomalous layer, approximately 150–200 km thick, has been identified in which body-wave velocity gradients are very small and may even be negative. Although part of the lower mantle, it evidently serves as a boundary layer between the mantle and core. It is labelled D'' to distinguish it from the normal mantle above it. The structure and role of the D'' layer are not yet known with confidence, but it is the focus of intensive current research. Models of the internal structure of D'' have been proposed with positive velocity gradients, others with negative velocity gradients, and some with small velocity discontinuities. The latter possibility is important because it would imply some stratification within D''.

The most interesting aspect of D'' is the presence – revealed by seismic tomographic imaging (Section 3.7.6.2)
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– of velocity variations of several percent that take place over lateral distances comparable in size to the continents and oceans in Earth's crust. The term “grand structure” has been coined for these regions; the thicker parts have also been termed “crypto-continents” and the thinner parts “crypto-oceans” (see Fig. 4.38). Moreover, the seismically fast regions (in which temperatures are presumed to be cooler than normal) lie beneath present subduction zones. This suggests that cold subducted lithosphere may eventually sink to the bottom of the mantle where it is colder and more rigid than the surrounding mantle and hence has higher body-wave velocities. A large low-velocity (hot) region underlies the Pacific basin, in which many centers of volcanism and locally high heat flow (“hotspots”, see Section 1.2.8) are located. The D’ layer is suspected of being the source of the mantle plumes that cause these anomalies. Exceptionally hot material from D’ rises in thin pipe-like mantle plumes to the 670 km discontinuity, which opposes further upward motion in the same way that it resists the deeper subduction of cold lithospheric slabs. Occasionally a hot plume is able to break through the 670 km barrier, producing a surface hotspot.

If our current understanding of the D’ layer is correct, it plays an important role in geodynamic and geothermal behavior. On the one hand, D’ serves as the source of material for the mantle plumes that give rise to hotspots, which are important in plate tectonics. On the other hand, the thermal properties of D’ could influence the outward transport of heat from the Earth’s core: in turn, this could affect the intricate processes that generate the Earth’s magnetic field.

3.7.5.4 The core

Early in Earth’s history, dense metallic elements are thought to have settled towards the Earth’s center, forming the core, while lighter silicates ascended and solidified to form the mantle. Studies of the compositions of meteorites and of the behavior of metals at high pressure and temperature give a plausible picture of the composition and formation of the core. It consists mainly of iron, with perhaps up to 10% nickel. The observed pressure-density relationships suggest that some less-dense non-metallic elements (Si, S, O) may be present in the outer core. It is not known whether small amounts of the more common radioactive elements ($^{40}$K, $^{232}$Th, $^{235}$U and $^{238}$U) are present in large enough abundances to contribute to the heat supply of the core.

The core has a radius of 3480 km and consists of a solid inner core (layer G in Bullen’s Earth Model A) surrounded by a liquid outer core (Bullen layer E) that is 1220 km thick. The transitional layer (F) was modelled by Bullen as a zone in which the P-wave velocity gradient is negative (i.e., $\alpha$ decreases with increasing depth). Not all seismologists agreed with this interpretation of the t-$\Delta$ curves and so the nature of layer F remained controversial for many years. The need for a layer F has now been discarded. Improved seismographic resolution has yielded a large quantity of high-quality data for reflections from the boundaries of the outer core (PcP, ScS) and the inner core (PKiKP) which have helped clear up the nature of these boundaries. The PKiKP phase contains high frequencies; this implies that the inner core boundary is sharp, probably no more than 5 km thick. The seismic events earlier interpreted as due to a layer F are now regarded as rays that have been scattered by small-scale features at the bottom of the mantle.

The inner core transmits P-waves (PKiKP phase) but S-waves in the inner core (PKJKP phase), although in principle possible, have not yet been observed unequivocally. Body-wave travel-times do not constrain the rigidity of the inner core. However, the amplitude spectrum of the frequencies of higher modes of the Earth’s free oscillations show that the inner core is likely solid. However, it is possible that it is not completely solid. Rather, it may be a mixture of solid and liquid phases at a temperature close to the solidification temperature. An analogy can be made with the mushy, semi-frozen state that water passes through on freezing to ice.

The outer core is fluid, with a viscosity similar to that of water. It is assumed to be homogeneous and its thermal state is supposed to be adiabatic. These are the conditions to be expected in a fluid that has been well mixed, in the Earth’s case by convection and differential rotation. One theory of core dynamics holds that the iron-rich inner core is solidifying from the fluid outer core, leaving behind its lighter elements. These constitute a less-dense, therefore gravitationally buoyant, fluid, which rises through the denser overlying liquid. This compositional type of buoyancy could be an important contributor to convection in the outer core, and therefore to the dynamics of the core and generation of the Earth’s magnetic field.

The core–mantle boundary (CMB) is also called the Gutenberg discontinuity. It is characterized by very large changes in body-wave velocities and is the most sharply defined seismic discontinuity. Seismic data show that the boundary is not smooth but has a topography of hills and valleys. Anomalies in the travel-times of PKKP phases – which are reflected once internally at the CMB – have been attributed to scattering by topographic features with a relief of a few hundred meters. However, depending on conditions in the hot D’ layer of the lower mantle immediately above the CMB some topographic features may be up to 10 km high. Interference between the CMB topography and fluid motions in the outermost core may couple the core and mantle to each other dynamically.

3.7.6 Seismic tomography

A free translation of the term tomography is “representation in cross-section.” Neighboring two-dimensional...
cross-sections can be combined to give a three-dimensional model. The use of computer-aided tomography (CAT) in medical diagnosis is well known as a non-invasive method of examining internal organs for abnormal regions. X-rays or ultrasonic rays are absorbed unequally by different materials. CAT consists of studying the attenuation of x-rays or ultrasonic waves that pass through the body in distinctly controlled planar sections. Seismic tomography uses the same principles, with the difference that the travel-times of the signals, as well as their attenuation, are observed. Hence the technique may be described as the three-dimensional modelling of the velocity or attenuation distribution of seismic waves in the Earth. The technique requires powerful computational facilities and sophisticated programming.

The travel-time of a seismic wave from an earthquake focus to a seismograph is determined by the velocity distribution along its path. For an idealized, spherically symmetric Earth model the radial distributions of velocity are known. The velocities in the model are mean values, which average out lateral fluctuations. If such a velocity model were used to compute travel-times of different phases to any epicentral distance, a set of curves indistinguishable from Fig. 3.77 would result. In reality, the observed travel-times usually show small deviations from the calculated times. These discrepancies are called travel-time residuals or anomalies, and they can have several causes. An obvious cause is that the focal depth of an earthquake is not zero, as assumed in Fig. 3.77, but may be up to several hundred kilometers. The parametrized Earth model *iasp91* takes this into account and tabulates travel-times for several focal depths. Clearly, precise determination of earthquake focal parameters (epicentral location, depth and time of occurrence) are essential prerequisites for seismic tomography. An important factor is the assumption of spherical symmetry, which is not perfectly valid, and so the ellipticity of the Earth's figure must be taken into account.

A local source of travel-time residuals is the particular velocity–depth distribution under an observational network. Ideally, the local vertical velocity profile should be known, so as to allow compensation of an observed travel-time for local anomalous structure. In practice, the signals from a selected earthquake are averaged for several stations in a given area of the surface (e.g., about 3° square) to reduce local perturbations.

The lateral variations in P- and S-wave velocity at any given depth may amount to a few percent of the “average” velocity for that depth assumed in the reference model. If, at a certain depth, a seismic ray passes through a region in which the velocity is slightly faster than average, the wave will arrive slightly sooner than expected at the receiver; if the anomalous velocity is slower than average, the wave will arrive late. This permits classification of travel-time as “early” or “late” depending on whether a ray has traversed a region that is “fast” or “slow” with respect to the assumed model.

The velocity of a seismic wave is determined by elastic parameters and density, which are affected by temperature. Thus, the velocity anomalies obtained from seismic tomography on a global scale are generally interpreted in terms of abnormal temperature and rigidity. A “slow” region is commonly associated with above-average temperature and lower rigidity, while a “fast” region is attributed to lower temperature and higher rigidity.

### 3.7.6.1 Travel-time residuals and velocity anomalies

The velocity distribution in the Earth is more heterogeneous than in a standard velocity-model. The time *(t)* taken for a seismic ray to travel along the path from an earthquake to a recording station thus differs from the value *(t)* predicted by the model. The difference *(Δt = t - t)* is a travel-time anomaly. A travel-time residual *(Δtt)* for the path is obtained by expressing the anomaly as a percentage of the expected travel-time *(t)*. If a given path is subdivided into segments with different velocities, the travel time is equivalent to the sum of the travel-times through the individual segments. This is like an equation with one known value *(Δtt)* and several unknown terms (the velocity anomalies in each segment). For a given earthquake several seismic stations may record rays that traverse the region of interest. Each station may also have recorded rays that crossed the region from different earthquakes. A large set of travel-time residuals results, equivalent to a large number of equations. A mathematical procedure called matrix inversion is used to solve the set of equations and obtain the velocity anomalies. The sophisticated analysis involves intensive data processing, and is beyond the scope of this text. However, Box 3.5 illustrates for a simple pattern of travel-time residuals (Box 3.4, Fig. B3.4) how a distribution of velocity anomalies may be deduced by back projection. In this method successive adjustments are made to an initial velocity distribution to account for the observed travel-time residuals.

Tomographic imaging of the Earth's three-dimensional velocity distribution is able to resolve small differences in seismic velocities. In some studies the signals are of local origin, generated by earthquakes or explosions within or near the volume of interest. Other studies are based on teleseismic signals, which originated in earthquakes at more than 20° epicentral distance. The analysis requires precise location of the earthquake source, corrections for local effects (such as crustal structure close to each measurement station), and reconstruction of the seismic rays between source and seismometer. Seismic tomography can be based on either body waves or surface waves. The database for P-wave studies consists of hundreds of thousands of first arrival times from earthquakes that occurred in the past 30 years, which
Box 3.4: Calculation of travel-time residuals

Consider the passage of seismic rays in six directions through a square region containing four equal areas, each a square with edge 5 km, in which the P-wave velocities ($V$) are 4.9 km s$^{-1}$, 5.0 km s$^{-1}$, 5.1 km s$^{-1}$ and 5.2 km s$^{-1}$ (Fig. B3.4). Let the expected velocity ($V_0$) throughout the region be 5.0 km s$^{-1}$. The velocity difference ($\Delta V$) in each area is found by subtracting the reference value, and the velocity anomaly ($\Delta V/V_0$) is obtained by expressing the difference as a percentage of the expected value. This gives zones that are 2% and 4% fast, a zone that is 2% slow and a zone with no anomaly.

Suppose that six seismic rays traverse the square region as in Fig. B3.4. If the velocity were 5.0 km s$^{-1}$ in each area, the expected travel-time ($t_0$) would be 2.0 s for each of the horizontal and vertical rays and $2\sqrt{2} = 2.828$ s for the longer diagonal rays. However, the real velocity is different in each area. As a result, some of the observed travel-times are shorter and some are longer than the expected value. A travel-time anomaly ($\Delta t$) is computed by subtracting the observed travel-times ($t$) from the expected value (Fig. B3.4d). The travel-time residual ($\Delta t/t_0$) is obtained by expressing the anomaly as a percentage of the expected travel-time (Fig. B3.4e). Note that the residuals are not simple averages of the velocity anomalies along each path.

![Fig. B3.4 Computation of relative travel-time residuals (in percent) for a simple four-block structure (modified after Kissling, 1993).](image)

The inversion of body-wave data provides evidence for the lateral variations in velocity in the deeper interior. The lateral variations at a given depth are equivalent to the variations on the surface of a sphere at that depth, and so can be depicted with the aid of spherical harmonic functions. The contoured velocity anomalies obtained by spherical harmonic analysis for a depth of 2500 km in the lower mantle are dominated by a ring of fast P-wave velocities around the Pacific and a slow-velocity region in the center (Fig. 3.88). Slow velocities are also present at depth under Africa. The pattern is present at nearly all depths in the lower mantle (i.e., below 1000 km). This is shown clearly by a vertical cross-section along a profile around the equator (Fig. 3.89, bottom frame). The slow velocities are interpreted as the expression of warm mantle material that may be rising from the core–mantle boundary as so-called “super-plumes.” The role of these super-plumes in the circulation pattern of mantle convection is not yet understood. Deep fast-velocity (“cold”) regions under America and Indonesia and slow-velocity (“warm”) regions under the Pacific and Atlantic oceans extend from about 1000 km depth to the core–mantle boundary. The lower mantle velocity anomalies show little correlation with plate tectonic elements, which are features of the lithosphere.

Velocity anomalies in the upper mantle, on the other hand, are clearly related to plate tectonic features. The upper mantle velocity structures can be modelled by inverting teleseismic body waves, but these depths can also be probed by long-period surface waves, which are more sensitive to variations in rigidity (and thus temperature). The inversion of long-period surface-wave data is an effective technique for modelling upper mantle S-wave velocities, especially under oceanic areas, where the resolution of body-wave anomalies is poor. The pattern of S-wave velocity anomalies in the upper mantle along the equatorial cross-section (Fig. 3.89, middle frame) shows generally elevated velocities under the “cold” continents and reduced velocities under the “warm” oceanic ridge systems. In these analyses the large local variations in S-wave velocities in the heterogeneous crust constitute an impediment to greater resolution.

Important advances in understanding the geodynamical state of the Earth’s mantle have been made as a result of numerous regional seismic tomography studies, which have focussed on specific tectonic environments such as zones of tectonic plate convergence. In particular,
Box 3.5: Calculation of velocity anomalies

To illustrate how a velocity structure can be deduced from observed travel-time residuals, we take as starting point the set of travel-time residuals $\Delta t/t_0$ computed for the velocity distribution in Box 3.4. The expected travel-time along a seismic path of length $L$ is $t_0 = L/V_0$, where $V_0$ is the reference velocity. If the true velocity is $V = V_0 + \Delta V$, the observed travel-time is

$$t = t_0 + \frac{L}{V_0 + \Delta V} = t_0 \left( 1 + \frac{\Delta V}{V_0} \right)^{-1}$$

$$= t_0 \left( 1 - \frac{\Delta V}{V_0} + \frac{\Delta V}{V_0} \right)^{-1} + \cdots$$

$$= t_0 \left( 1 - \frac{\Delta V}{V_0} + \frac{\Delta V}{V_0} \right)^{-1} + \cdots$$

(1)

Next, we assume that the percentage velocity anomalies $(\Delta V/V_0)$ are the same as the percentage travel-time residuals. As shown by Eq. (2), this is not strictly true; the discrepancy is of the order of $(\Delta V/V_0)$. However, the velocity anomalies are usually very small (see Figs. 3.88–3.90), so assuming that the velocity anomaly is equal to the travel-time residual is a reasonable approximation.

Consider a horizontal ray that traverses the upper two “fast” blocks in Fig. B3.5. To account for the (early) travel-time anomaly of $+2.9\%$ let each upper block be allocated a (fast) velocity anomaly of $+2.9\%$. Similarly, let each of the bottom two blocks be allocated velocity-antimy of $-1.02\%$. This simple velocity distribution (Fig. B3.5a) satisfies the travel-time anomalies for the horizontal rays. However, in the vertical direction it gives travel-time anomalies of $+0.94\%$ (the mean of $+2.90\%$ and $-1.02\%$) for each ray (Fig. B3.5b). This does not agree with the observed anomalies for the two vertical rays ($+0.98\%$ and $+0.90\%$, respectively); one vertical anomaly is $0.04\%$ too large, the other is $0.04\%$ too small. The velocities in the blocks are adjusted accordingly by making a correction of $+0.04\%$ to the left-hand blocks and $-0.04\%$ to the right-hand blocks (Fig. B3.5c). This gives a new distribution of velocity anomalies which satisfies the horizontal and vertical rays (Fig. B3.5d).

The model now gives travel-time anomalies along each of the diagonal rays of $+0.94\%$, compared to observed anomalies of $+1.92\%$ and $-0.04\%$, respectively (Fig. B3.5e). Further corrections of $+0.98\%$ are now made to the upper right and lower left blocks, and $-0.98\%$ to the lower right and upper left blocks (Fig. B3.5f). The resulting distribution of velocity anomalies (Fig. B3.5g) satisfies all six rays through the anomalous region, and is close to the original distribution of velocity anomalies (Fig. B3.4b).

Investigations of subduction zones have provided new insights into the effects on mantle convection of the seismic discontinuities at 400 km and 670 km depth. As discussed in more detail in Section 4.2.9.3, two models of mantle convection are prevalent. They differ mainly in the roles played by these seismic discontinuities. In the first – whole mantle convection – the entire mantle participates in the convection process. In the second – layered convection – the seismic discontinuities bound a transition zone that separates a system of convection cells in the upper mantle from a system of convection cells in the lower mantle.

However, the reality is more complex, and neither model satisfies all the observational constraints. At a subduction zone, the subducting plate is colder and denser than the overriding plate and sinks into the mantle (Fig. 3.90). Evidence from regional seismic tomography shows that, at some subduction zones (e.g., the Aegean arc, Fig. 3.90a) the subducting cold slab is able to penetrate through the transition zone and to sink deeper into the lower mantle. In other cases (e.g., the Kurile arc, Fig. 3.90b) the 670 km discontinuity appears to block the downward motion of the subducting slab, which is then caused to deflect horizontally. Some slabs that penetrate the 670 km discontinuity appear to sink deep into the lower mantle. Some may in fact reach the core–mantle boundary.

Seismic tomography is a potent technique for describing conditions in the mantle. Many questions, such as the nature of the “super-plume” structures beneath the Pacific and Africa, remain unanswered. Despite convective mixing there may be more heterogeneity in mantle composition than is often supposed. Indeed, the assumption that fast and slow seismic velocities imply cold and hot temperatures, respectively, may be overly simplistic. Yet, seismic tomography is the best tool currently
available for investigating geodynamic processes in the Earth’s interior.

### 3.8 SUGGESTIONS FOR FURTHER READING

#### Introductory level


**Fig. 3.88** Map of P-wave velocity anomalies in the lower mantle (2500 km depth). The deviations are plotted as percent faster or slower than the reference velocity at this depth (after Dziewonski, 1984, 1989).

**Fig. 3.89** Seismic tomographic section through the mantle along an equatorial profile. Middle frame: S-wave anomalies in the upper mantle to a depth of 670 km. Bottom frame: P-wave anomalies in the lower mantle in depths between 670 and 2890 km (after Woodhouse and Dziewonski, 1984).
Intermediate level


Advanced level


3.9 REVIEW QUESTIONS

1. Describe the principle of the seismometer.
2. Describe the particle motions relative to the direction of propagation of the two seismic body waves and the two seismic surface waves.
3. What are the Lamé constants in seismic theory?
4. What is meant by dispersion with regard to the propagation of surface waves?
5. Sketch how the train of Rayleigh waves from a distant earthquake would appear on a seismogram.
6. What are the free oscillations of the Earth? What kinds are possible? What is meant by the normal modes? What are higher modes? How do they relate to surface waves?
7. How does the elastic rebound model explain the origin of a tectonic earthquake?
8. What is the epicenter of an earthquake? Explain how the epicenter of an earthquake can be located? What is the minimum number of seismic records needed?
9. Explain the difference between earthquake intensity and magnitude.
10. If the magnitude of an earthquake is 0.5 greater than that of another, how much greater is the amount of energy it releases?
11. How does a tsunami originate? Why are tsunami barely noticeable over the open ocean but very dangerous near shore?
12. Describe the geographical distribution of the Earth’s seismically active zones.
13. Describe with the aid of sketches the distribution of earthquakes at the three major plate boundaries:
3.10 EXERCISES

(a) spreading ridge, (b) transform fault, and (c) subduction zone.

14. Sketch the fault-plane solutions that characterize earthquakes at each type of plate boundary. Describe their significance with regard to plate tectonic motions.

15. How do the distribution and focal solutions of earthquakes at a transform fault differ from those at a strike-slip fault?

16. What is the Mohorovičić discontinuity? What is the seismic evidence for this feature? What is its average depth under continents, under oceans, and for the entire world?

17. How do seismic wave velocities change at the major discontinuities in the Earth’s internal structure? How are these discontinuities characterized?

18. What is the brittle–ductile transition in the Earth? What physical properties determine the depth and nature of this transition?

19. On a cross-section of the Earth, sketch the paths of the following seismic rays: (i) PKP, (ii) SKS, (iii) PnP, (iv) PPP.

20. What is a PKIKP wave? Describe the refraction of this wave at each discontinuity it crosses.

21. What is the critical distance in seismic refraction surveying? What is the crossover distance? What is a head wave? What are supercritical reflections?

22. A seismic survey is conducted over level ground consisting of two horizontal layers. Sketch a travel-time diagram that shows the arrivals of the direct wave, the reflected wave, and the doubly refracted wave. How can the seismic velocity of each layer be determined from the diagram?

23. What is normal move-out in seismic reflection profiling?

24. What is meant by the migration of reflection seismic records? Why is it necessary?

25. Describe the split-spread and common-mid-point methods in reflection profiling? What is achieved by each of these methods?

2. The table below gives the densities and seismic P- and S-wave velocities at various depths in the Earth.

<table>
<thead>
<tr>
<th>Depth [km]</th>
<th>Density [kg m(^{-3})]</th>
<th>P-wave [km s(^{-1})]</th>
<th>S-wave [km s(^{-1})]</th>
</tr>
</thead>
<tbody>
<tr>
<td>100</td>
<td>3380</td>
<td>7.95</td>
<td>4.45</td>
</tr>
<tr>
<td>500</td>
<td>3890</td>
<td>9.55</td>
<td>5.28</td>
</tr>
<tr>
<td>1000</td>
<td>4680</td>
<td>11.42</td>
<td>6.36</td>
</tr>
<tr>
<td>1500</td>
<td>4970</td>
<td>12.13</td>
<td>6.68</td>
</tr>
<tr>
<td>2000</td>
<td>5240</td>
<td>12.79</td>
<td>6.93</td>
</tr>
<tr>
<td>2500</td>
<td>5490</td>
<td>13.39</td>
<td>7.16</td>
</tr>
<tr>
<td>2890</td>
<td>5680</td>
<td>13.64</td>
<td>7.30</td>
</tr>
<tr>
<td>2900</td>
<td>9430</td>
<td>8.10</td>
<td>0</td>
</tr>
<tr>
<td>3500</td>
<td>10230</td>
<td>8.90</td>
<td>0</td>
</tr>
<tr>
<td>4000</td>
<td>10760</td>
<td>9.51</td>
<td>0</td>
</tr>
<tr>
<td>4500</td>
<td>11190</td>
<td>9.97</td>
<td>0</td>
</tr>
<tr>
<td>5000</td>
<td>11540</td>
<td>10.44</td>
<td>0</td>
</tr>
</tbody>
</table>

(a) From these quantities calculate the rigidity modulus, \(\mu\), bulk modulus, \(K\), and Poisson’s ratio, \(\gamma\), at each depth.

(b) Discuss in your own words the information that these curves give about the deep interior of the Earth.

3. A strong earthquake off the coast of Japan sets off a tsunami that propagates across the Pacific Ocean (average depth \(d = 5\) km).

(a) Calculate the velocity of the wave in km hr\(^{-1}\) and the corresponding wavelength, when the wave has a dominant period of 30 min.

(b) How long does the wave take to reach Hawaii, which is at an angular distance of 54° from the epicenter?

4. The dispersion relation between frequency \(\omega\) and wave number \(k\) of seismic water waves for water depth \(d\) is (Box 3.3)

\[\omega^2 = gktanh(kd)\]

(a) Modify this expression for wavelengths that are much shorter than the water depth.

(b) Determine the phase velocity of these waves.

(c) Show that the group velocity of the waves is half the phase velocity.

5. In a two-layer Earth the mantle and core are each homogeneous and the radius of the core is one-half the radius of the Earth. Derive a formula for the travel-time curve for the arrival time \(t\) of the phase PnP at epicentral distance \(\Delta\). Verify the formula for the maximum possible value of \(\Delta\) in this model.

6. Why might one expect an interface with a small critical angle to be a good reflector of seismic energy?

7. The P-wave from an earthquake arrives at a seismograph station at 10:20 a.m. and the S-wave arrives at...
10:25 a.m. Assuming that the P-wave velocity is 5 km s\(^{-1}\) and that Poisson’s ratio is 0.25, compute the time at which the earthquake occurred and its epicentral distance in degrees from the seismograph station.

8. The following table gives arrival times of P-waves (\(t_p\)) and S-waves (\(t_s\)) from a nearby earthquake:

<table>
<thead>
<tr>
<th>Recording Station</th>
<th>Time of day [hr:min]</th>
<th>(t_p) [s]</th>
<th>(t_s) [s]</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>23:36</td>
<td>54.65</td>
<td>57.90</td>
</tr>
<tr>
<td>B</td>
<td>23:36</td>
<td>57.34</td>
<td>62.15</td>
</tr>
<tr>
<td>C</td>
<td>23:37</td>
<td>00.49</td>
<td>07.55</td>
</tr>
<tr>
<td>D</td>
<td>23:37</td>
<td>01.80</td>
<td>10.00</td>
</tr>
<tr>
<td>E</td>
<td>23:37</td>
<td>01.90</td>
<td>10.10</td>
</tr>
<tr>
<td>F</td>
<td>23:37</td>
<td>02.25</td>
<td>10.70</td>
</tr>
<tr>
<td>G</td>
<td>23:37</td>
<td>03.10</td>
<td>12.00</td>
</tr>
<tr>
<td>H</td>
<td>23:37</td>
<td>03.50</td>
<td>12.80</td>
</tr>
<tr>
<td>I</td>
<td>23:37</td>
<td>06.08</td>
<td>18.30</td>
</tr>
<tr>
<td>J</td>
<td>23:37</td>
<td>07.07</td>
<td>19.79</td>
</tr>
<tr>
<td>K</td>
<td>23:37</td>
<td>08.32</td>
<td>21.40</td>
</tr>
<tr>
<td>L</td>
<td>23:37</td>
<td>11.12</td>
<td>26.40</td>
</tr>
<tr>
<td>M</td>
<td>23:37</td>
<td>11.50</td>
<td>26.20</td>
</tr>
<tr>
<td>N</td>
<td>23:37</td>
<td>17.80</td>
<td>37.70</td>
</tr>
</tbody>
</table>

(a) Plot the arrival-time differences (\(t_s - t_p\)) against the arrival times of the P-wave to produce a Wadati diagram.
(b) Determine the ratio \(\alpha/\beta\) of the seismic velocities.
(c) Determine the time of occurrence (\(t_o\)) of the earthquake.

9. A plane seismic wave, travelling vertically downwards in a rock of density 2200 kg m\(^{-3}\) with seismic velocity 2000 m s\(^{-1}\), is incident on the horizontal top surface of a rock layer of density 2400 kg m\(^{-3}\) and seismic velocity 3300 m s\(^{-1}\).
(a) What are the amplitude ratios of the transmitted and reflected waves?
(b) What fraction of the energy of the incident wave is transmitted into the lower medium?
(c) Using the listed densities calculate the reflection coefficient for each interface (except the base of the dolomite). Which interface gives the strongest reflection and which the weakest? At which interfaces does a change in phase occur? What does this mean?

10. A plane seismic wave travels vertically downwards at a velocity of 4800 m s\(^{-1}\) through a salt layer with density 2100 kg m\(^{-3}\). The wave is incident upon the top surface of a sandstone layer with density 2400 kg m\(^{-3}\) and seismic velocity 3300 m s\(^{-1}\). The phase of the reflected wave is changed by 180° and the reflected amplitude is 2% of the incident amplitude. What is the seismic velocity of the sandstone?

(a) Calculate the minimum arrival times for seismic reflections from each of the reflecting interfaces in the following section. Consider the base of the lowermost bed to be a reflector as well.
(b) What is the average velocity of the section for a reflection from the base of the dolomite?
(c) Using the listed densities calculate the reflection coefficient for each interface (except the base of the dolomite). Which interface gives the strongest

11. A reflection seismic record in an area of relatively flat dips gave the following data:

(a) Plot the \(t-x\) curves for these reflections to show the “moveout” effect.
(b) On a different graph, plot the \(t^2-x^2\) curves (i.e., squared data) for the reflections.
(c) Determine the average vertical velocity from the surface to each reflecting bed.
(d) Use these velocities to compute the depths to the reflecting beds.

12. The following table gives two-way travel times of seismic waves reflected from different reflecting interfaces in a horizontally layered medium:

<table>
<thead>
<tr>
<th>Distance shot-point to detector [m]</th>
<th>Travel-time ((t_1), 1st reflection) [s]</th>
<th>Travel time ((t_2), 2nd reflection) [s]</th>
</tr>
</thead>
<tbody>
<tr>
<td>30</td>
<td>1.000</td>
<td>1.200</td>
</tr>
<tr>
<td>90</td>
<td>1.002</td>
<td>1.201</td>
</tr>
<tr>
<td>150</td>
<td>1.003</td>
<td>1.201</td>
</tr>
<tr>
<td>210</td>
<td>1.007</td>
<td>1.202</td>
</tr>
<tr>
<td>270</td>
<td>1.011</td>
<td>1.205</td>
</tr>
<tr>
<td>330</td>
<td>1.017</td>
<td>1.207</td>
</tr>
<tr>
<td>390</td>
<td>1.023</td>
<td>1.207</td>
</tr>
</tbody>
</table>

(a) Plot the \(t^2-x^2\) against (distance)\(^2\) to show the “moveout” effect.
(b) Determine the vertical two-way travel-time (“echo-time”) and average velocity to each reflecting interface.
(c) Compute the depth of each reflector and the thickness of each layer.
(d) Compute the true velocity (interval velocity) of each layer.
(e) Verify your results by computing the total vertical travel-time for a wave reflected from the deepest interface.

14. Assume the horizontally layered structure from the previous problem.
   (a) If a seismic ray leaves the surface at an angle of 15° to the vertical, how long does it take to return to the surface after reflecting from the basement?
   (b) At what horizontal distance from the shot-point does this ray reach the surface?

15. Assume that the three horizontal homogeneous rock layers in the previous problems have densities of 1800, 2200, and 2500 kg m⁻³ respectively. The lowest layer overlies basement with velocity 5.8 km s⁻¹ and density 2700 kg m⁻³.
   (a) Compute the reflection and transmission coefficients at each interface for a plane P-wave travelling vertically downwards.
   (b) Calculate what fraction of the initial energy of the wave is transmitted into the basement.
   (c) Calculate the fraction of the initial energy carried in the reflection that returns to the surface from the basement.

16. An incident P-wave is converted into refracted and reflected P- and S-waves at an interface. Calculate all the critical angles in the following three cases, where \( \alpha \) and \( \beta \) are the P-wave and S-wave velocities, respectively:

<table>
<thead>
<tr>
<th>Layer</th>
<th>Seismic wave</th>
<th>Case (a) [km s⁻¹]</th>
<th>Case (b) [km s⁻¹]</th>
<th>Case (c) [km s⁻¹]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Above interface</td>
<td>( \alpha )</td>
<td>3.5</td>
<td>4.0</td>
<td>5.5</td>
</tr>
<tr>
<td></td>
<td>( \beta )</td>
<td>2.0</td>
<td>2.3</td>
<td>3.1</td>
</tr>
<tr>
<td>Below interface</td>
<td>( \alpha )</td>
<td>8.5</td>
<td>6.0</td>
<td>7.0</td>
</tr>
<tr>
<td></td>
<td>( \beta )</td>
<td>5.0</td>
<td>3.5</td>
<td>4.0</td>
</tr>
</tbody>
</table>

17. An incident P-wave is converted into refracted and reflected P- and S-waves at an interface that is inclined at 20° to the horizontal, as in the figure below. The respective P- and S-wave velocities are 5 km s⁻¹ and 3 km s⁻¹ above the interface and 7 km s⁻¹ and 4 km s⁻¹ below the interface. If the incident P-wave strikes the interface at an angle of 40° to the horizontal, calculate the angles to the horizontal made by the reflected and refracted P- and S-waves.

18. A seismic refraction survey gave the following data for the first arrival times at various distances from the shot-point.

<table>
<thead>
<tr>
<th>Distance [km]</th>
<th>Time [s]</th>
<th>Distance [km]</th>
<th>Time [s]</th>
</tr>
</thead>
<tbody>
<tr>
<td>3.1</td>
<td>1.912</td>
<td>13.1</td>
<td>6.678</td>
</tr>
<tr>
<td>5.0</td>
<td>3.043</td>
<td>14.8</td>
<td>7.060</td>
</tr>
<tr>
<td>6.5</td>
<td>3.948</td>
<td>16.4</td>
<td>7.442</td>
</tr>
<tr>
<td>8.0</td>
<td>4.921</td>
<td>18.0</td>
<td>7.830</td>
</tr>
<tr>
<td>9.9</td>
<td>5.908</td>
<td>19.7</td>
<td>8.212</td>
</tr>
<tr>
<td>11.5</td>
<td>6.288</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(a) Plot the travel-time curve for the first arrivals.
(b) Calculate the seismic velocities of the layers.
(c) Calculate the minimum depth to the refracting interface.
(d) Calculate the critical angle of refraction for the interface.
(e) Calculate the critical distance for the first arrival of refracted rays.
(f) Calculate the crossover distance beyond which the first arrivals correspond to head waves.

19. A seismic refraction survey is carried out over a layered crust with flat-lying interfaces. In one case the crust is homogeneous and 30 km thick with a P-wave velocity 6 km s⁻¹ and overlies mantle with P-wave velocity of 8 km s⁻¹. In the other case the crust consists of an upper layer 20 km thick with P-wave velocity 6 km s⁻¹ overlying a lower layer 10 km thick with P-wave velocity 5 km s⁻¹. The upper mantle P-wave velocity is again 8 km s⁻¹. On the same graph, plot the first arrival time curves for the two cases. What is the effect of the low-velocity layer on the estimation of depth to the top of the mantle?

20. The table below gives up-dip and down-dip travel-times of P-wave arrivals for refraction profiles over an inclined interface. The geophones are laid out in a straight line passing through the alternate shot-points A and B, which are 2700 m apart on the profile.
   (a) Plot the travel-time curves for each shot-point.
   (b) Calculate the true velocity of the upper layer.
   (c) Calculate the apparent velocities of the layer below the refraactor.
   (d) In which direction does the refracting interface dip?
   (e) What is the angle of dip of the interface?
   (f) What is the true velocity of the layer below the refraactor?
(g) What are the closest distances to the refractor below A and B?
(h) What are the vertical depths to the refractor below A and B?

<table>
<thead>
<tr>
<th>Distance from shot-point [m]</th>
<th>Travel-time [s] from A</th>
<th>Travel-time [s] from B</th>
</tr>
</thead>
<tbody>
<tr>
<td>300</td>
<td>0.139</td>
<td>0.139</td>
</tr>
<tr>
<td>600</td>
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<td>1.339</td>
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</tr>
<tr>
<td>3600</td>
<td>1.424</td>
<td>1.329</td>
</tr>
</tbody>
</table>
4.1 GEOCHRONOLOGY

4.1.1 Time

Time is both a philosophical and physical concept. Our awareness of time lies in the ability to determine which of two events occurred before the other. We are conscious of a present in which we live and which replaces continually a past of which we have a memory; we are also conscious of a future, in some aspects predictable, that will replace the present. The progress of time was visualized by Sir Isaac Newton as a river that flows involuntarily at a uniform rate. The presumption that time is an independent entity underlies all of classical physics. Although Einstein’s Theory of Relativity shows that two observers moving relative to each other will have different perceptions of time, physical phenomena are influenced by this relationship only when velocities approach the speed of light. In everyday usage and in non-relativistic science the Newtonian notion of time as an absolute quantity prevails.

The measurement of time is based on counting cycles (and portions of a cycle) of repetitive phenomena. Prehistoric man distinguished the differences between day and night, he observed the phases of the Moon and was aware of the regular repetition of the seasons of the year. From these observations the day, month and year emerged as the units of time. Only after the development of the clock could the day be subdivided into hours, minutes and seconds.

4.1.1.1 The clock

The earliest clocks were developed by the Egyptians and were later introduced to Greece and from there to Rome. About 2000 BC the Egyptians invented the water clock (or clepsydra). In its primitive form this consisted of a container from which water could escape slowly by a small hole. The progress of time could be measured by observing the change of depth of the water in the container (using graduations on its sides) or by collecting and measuring the amount of water that escaped. The Egyptians (or perhaps the Mesopotamians) are also credited with inventing the sundial. Early sundials consisted of devices – poles, upright stones, pyramids or obelisks – that cast a shadow; the passage of time was observed by the changing direction and length of the shadow. After trigonometry was developed dials could be accurately graduated and time precisely measured. Mechanical clocks were invented around 1000 AD but reliably accurate pendulum clocks first came into use in the seventeenth century. Accurate sundials, in which the shadow was cast by a fine wire, were used to check the setting and calibration of mechanical clocks until the nineteenth century.

4.1.1.2 Units of time

The day is defined by the rotation of the Earth about its axis. The day can be defined relative to the stars or to the Sun (Fig. 4.1). The time required for the Earth to rotate through 360° about its axis and to return to the same meridian relative to a fixed star defines the sidereal day. All sidereal days have the same length. Sidereal time must be used in scientific calculations that require rotational velocity relative to the absolute framework of the stars. The time required for the Earth to rotate about its axis and return to the same meridian relative to the Sun defines the solar day. While the Earth is rotating about its axis, it is also moving forward along its orbit. The orbital motion about the Sun covers 360° in about 365 days, so that in one day the Earth moves forward in its orbit by approximately 1°. To return to the solar meridian the Earth must rotate this extra degree. The solar day is therefore slightly longer than the sidereal day. Solar days are not equal in length. For example, at perihelion the Earth
Earth's age, thermal and electrical properties

is moving faster forward in its orbit than at aphelion (see Fig. 1.2). At perihelion the higher angular rate about the Sun means that the Earth has to rotate through a larger than average angle to catch up with the solar meridian. Thus, at perihelion the solar day is longer than average; at aphelion the opposite is the case. The obliquity of the ecliptic causes a further variation in the length of the solar day. Mean solar time is defined in terms of the mean length of the solar day. It is used for most practical purposes on Earth, and is the basis for definition of the hour, minute and second. One mean solar day is equal to exactly 86,400 seconds. The length of the sidereal day is approximately 86,164 seconds.

The sidereal month is defined as the time required for the Moon to circle the Earth and return to the celestial longitude of a given star. It is equal to 27.32166 (solar) days. To describe the motion of the Moon relative to the Sun, we have to take into account the Earth's motion around its orbit. The time between successive alignments of the Sun, Earth and Moon on the same meridian is the synodic month. It is equivalent to 29.530 59 days.

The sidereal year is defined as the time that elapses between successive occupations by the Earth of the same point in its orbit with respect to the stars. It is equal to 365.256 mean solar days. Two times per year, in spring and autumn, the Earth occupies positions in its orbit around the Sun where the lengths of day and night are equal at any point on the Earth. The spring occurrence is called the vernal equinox; that in the autumn is the autumnal equinox. The solar year (correctly called the tropical year) is defined as the time between successive vernal equinoxes. It equals 365.242 mean solar days, slightly less than the sidereal year. The small difference (0.014 days, about 20 minutes) is due to the precession of the equinoxes, which takes place in the retrograde sense (i.e., opposite to the revolution of the Earth about the Sun) and thereby reduces the length of the tropical year.

Unfortunately the lengths of the sidereal and tropical years are not constant but change slowly but measurably. In order to have a world standard the fundamental unit of scientific time was defined in 1956 in terms of the length of the tropical year 1900, which was set equal to 31,556,925.9747 seconds of ephemeris time. Even this definition of the second is not constant enough for the needs of modern physics. Highly stable atomic clocks have been developed that are capable of exceptional accuracy. For example, the alkali metal cesium has a sharply defined atomic spectral line whose frequency can be determined accurately by resonance with a tuned radio-frequency circuit. This provides the physical definition of the second of ephemeris time as the duration of 9,192,631,770 cycles of the cesium atomic clock.

Other units of time are used for specific purposes. Astronomers use a practical unit of time synchronized to the Earth's rotation. This gives a uniform timescale called universal time and denoted UT2; it is defined for a particular year by the Royal Observatory at Greenwich, England. The particular ways of defining the basic units of time are important for analyzing some problems in astronomy and satellite geodesy, but for most geophysical applications the minor differences between the different definitions are negligible.

4.1.1.3 The geological timescale

Whereas igneous rocks are formed in discrete short-lived eruptions of magma, sequences of sedimentary rocks take very long periods of time to form. Many sedimentary formations contain fossils, i.e., relics of creatures that lived in or near the basin in which the sediments were deposited. Evolution gives a particular identifiable character to the fossils in a given formation, so that it is possible to trace the evolution of different fossil families and correlate their beginnings and extinctions. These characteristics allow the formation of the host rock to be correlated with others that contain part or all of the same fossil assemblage, and permit the development of a scheme for dating sediments relative to other formations. Gradually a biostratigraphical timescale has been worked out, which permits the accurate determination of the relative age of a sedimentary sequence. This information is intrinsic to any geological timescale.

A geological timescale combines two different types of information. Its basic record is a chronostatigraphical scale showing the relationship between rock sequences. These are described in detail in stratigraphical sections that are thought to have continuous sedimentation and to contain complete fossil assemblages. The boundary between two sequences serves as a standard of reference and the section in which it occurs is called a boundary stratotype. When ages are associated with the reference points, the scale becomes a geological timescale. Time is divided into major units called epochs, which are subdivided into eras; these in turn are subdivided into periods containing several epochs. The lengths of time units in early timescales, before ages were determined with radioactive isotopes, were expressed as multiples of the duration of the Eocene epoch. Modern geological timescales are based on isotopic ages, which are calibrated in normal time units.

The geological timescale is constantly being revised and updated. Improved descriptions of important boundaries in global stratotype sections, more refined correlation with other chronostratigraphical units, and better calibration by more accurate isotopic dating lead to frequent revisions. Figure 4.2 shows an example of a current geological timescale.

4.1.2 Estimating the Earth's age

Early estimates of the Earth's age were dominated by religious beliefs. Some oriental philosophers in antiquity believed that the world had been in existence for millions of years. Yet, western thought on this topic was dominated
for centuries by the tenets of the Jewish and Christian faiths. Biblical estimates of the world’s age were made on the basis of genealogy by adding up the lengths of lifetimes and generations mentioned in the Old Testament. Some estimates also incorporate information from other ancient scriptures or related sources. The computed age of Creation invariably gave an age of the Earth less than 10,000 yr.

The best-known biblical estimate of the date of Creation was made by James Ussher (1581–1656), an Irish archbishop. His analysis of events recorded in the Old Testament and contemporary ancient scrolls led Ussher to proclaim that the exact time of Creation was at the beginning of night on October 22, in the year 4004 BC, which makes the age of the Earth approximately 6000 yr. Other biblical scholars inferred similar ages. This type of “creationist” age estimate is still favored by many fundamen-
talist Christians, whose faith is founded on literal interpretation of the Bible. However, biblical scholars with a more broadly based faith recognize age estimates based upon scientific methodology and measurement.

In the late nineteenth century the growth of natural philosophy (as physics was then called) fostered calculations of the Earth’s age from physical properties of the solar system. Among these were estimates based on the cooling of the Sun, the cooling of the Earth, and the slow increase in the Earth–Moon distance. Chemists tried to date the Earth by establishing the time needed for the seas to acquire their salinity, while geologists conjectured how long it would take for sediments and sedimentary rocks to accumulate.

4.1.2.1 Cooling of the Sun

Setting aside the biblical “sequence” chronicled in the book of Genesis, modern-day philosophers opine that the Earth cannot be older than the Sun. Cooling of the Sun takes place by continuous radiation of energy into space. The amount of solar energy falling on a square meter per second at the Earth’s distance from the Sun (1 AU) is called the solar constant; it equals 1360 W m$^{-2}$. The amount of energy lost by the Sun per second is obtained by multiplying this value by the surface area of a sphere whose radius is one astronomical unit. This simple calculation shows that the Sun is losing energy at the rate of $3.83 \times 10^{26}$ W. In the nineteenth century, before the dis-
covey of radioactivity and nuclear reactions, the source of this energy was not known. A German scientist, H. L. F. von Helmholtz, suggested in 1856 that it might result from the change of potential energy due to gravitational condensation of the Sun from an originally more dis-
tended body. The condensational energy $E_s$ of a mass $M_s$ of uniform density and radius $R_s$ is given by

$$E_s = \frac{3}{5} \frac{M_s^2}{R_s^2} = 2.28 \times 10^{44} \text{ J} \tag{4.1}$$

In this equation $G$ is the gravitational constant (see Section 2.2.1). The factor $3/5$ in the result arises from the assumption of a uniform density distribution inside the Sun. Dividing the condensational energy by the rate of energy loss gives an age of 19 million years for the Sun. Allowing for the increase in density towards the center...
causes an approximately threefold increase in the gravitational energy and the inferred age of the Sun.

4.1.2.2 Cooling of the Earth

In 1862 William Thomson, who later became Lord Kelvin, examined the cooling of the Sun in more detail. Unaware (although suspicious) of other sources of solar energy, he concluded that gravitational condensation could supply the Sun’s radiant energy for at least 10 million years but not longer than 500 million years (500 Ma). In the same year he investigated the cooling history of the Earth. Measurements in deep wells and mines had shown that temperature increases with depth. The rate of increase, or temperature gradient, was known to be variable but seemed to average about 1 °F for every 50 ft (0.036 °C m⁻¹). Kelvin inferred that the Earth was slowly losing heat and assumed that it did so by the process of conduction alone (Section 4.2.4). This enabled him to deduce the Earth’s age from a solution of the one-dimensional equation of heat conduction (see Eq. 4.57 and Box 4.2), which relates the temperature $T$ at time $t$ and depth $z$ to physical properties of the cooling body, such as its density ($\rho$), specific heat ($c$) and thermal conductivity ($k$).

These parameters are known for particular rock types, but Kelvin had to adopt generalized values for the entire Earth. The model assumes that the Earth initially had a uniform temperature $T_0$ throughout, and that it cooled from the outside leaving the present internal temperatures higher than the surface temperature. After a time $t$ has elapsed the temperature gradient at the surface of the Earth is given by

$$\left(\frac{dT}{dz}\right)_{z=0} = \sqrt{\frac{\rho c T_0}{\pi k \rho}}$$

(4.2)

Kelvin assumed an initial temperature of 7000°F (3871°C, 4144 K) for the hot Earth, and contemporary values for the surface temperature gradient and thermal parameters. The calculation for $t$ yielded an age of about 100 Ma for the cooling Earth.

4.1.2.3 Increase of the Earth–Moon separation

The origin of the Moon is still uncertain. George H. Darwin, son of the more famous Charles Darwin and himself a pioneer in tidal theory, speculated that it was torn from the Earth by rapid rotation. Like other classical theories of the Moon’s origin (e.g., capture of the Moon from elsewhere in the solar system, or accretion in Earth orbit) the theory is flawed. In 1898 Darwin tried to explain the Earth’s age from the effects of lunar tidal friction. The tidal bulges on the Earth interact with the Moon’s gravitation to produce a decelerating torque that slows down the Earth’s rotation and so causes an increase in the length of the day. The equal and opposite reaction is a torque exerted by the Earth on the Moon’s orbit that increases its angular momentum. As explained in Section 2.3.4.2, this is achieved by an increase in the distance between the Moon from the Earth and a decrease in the rotation rate of the Moon about the Earth, which increases the length of the month. The Earth’s rotation decelerates more rapidly than that of the Moon, so that eventually the angular velocities of the Earth and Moon will be equal. In this synchronous state the day and month will each last about 47 of our present days and the Earth–Moon distance will be about 87 Earth radii; the separation is presently 60.3 Earth radii.

Similar reasoning suggests that earlier in Earth’s history, when the Moon was much closer to the Earth, both bodies rotated faster so that an earlier synchronous state may be conjectured. The day and month would each have lasted about 5 of our present hours and the Earth–Moon distance would have been about 2.3 Earth radii. However, at this distance the Moon would be inside the Roche limit, about 3 Earth radii, at which the Earth’s gravitation would tear it apart. Thus, it is unlikely that this condition was ever realized.

Darwin calculated the time needed for the Earth and Moon to progress from an initially unstable close relationship to their present separation and rotation speeds, and concluded that a minimum of 56 Ma would be needed. This provided an independent estimate of the Earth’s age, but it is unfortunately as flawed in its underlying assumptions as other models.

4.1.2.4 Oceanic salinity

Several determinations of the age of the Earth have been made on the basis of the chemistry of sea water. The reasoning is that salt is picked up by rivers and transported into lakes and seas; evaporation removes water but the vapor is fresh, so the salt is left behind and accumulates with time. If the accumulation rate is measured, and if the initial salt concentration was zero, the age of the sea and, by inference, the Earth can be calculated by dividing the present concentration by the accumulation rate.

Different versions of this method have been investigated. The most noted is that of an Irish geologist, John Joly, in 1899. Instead of measuring salt he used the concentration of a pure element, sodium. Joly determined the total amount of sodium in the oceans and the annual amount brought in by rivers. He concluded that the probable maximum age of the Earth was 89 Ma. Later estimates included corrections for possible sodium losses and non-linear accumulation but gave similar ages less than about 100 Ma.

The principal flaw in the chemical arguments is the assumption that sodium accumulates continuously in the ocean. In fact, all elements are withdrawn from the ocean at about the same rate as they are brought in. As a result, sea water has a chemically stable composition, which is not changing significantly. The chemical methods do not
4.1 GEOCHRONOLOGY

measure the age of the ocean or the Earth, but only the average length of time that sodium resides in the seas before it is removed.

4.1.2.5 Sedimentary accumulation

Not to be outdone by the physicists and chemists, late nineteenth century geologists tried to estimate Earth’s age using stratigraphical evidence for the accumulation of sediments. The first step involved determining the thicknesses of sediment deposited during each unit of geological time. The second step was to find the corresponding sedimentation rate. When these parameters are known, the length of time represented in each unit can be calculated. The age of the Earth is the sum of these times.

The geological estimates are fraught with complications. Sediments with a silicate matrix, such as sandstones and shales, are deposited mechanically, but carbonate rocks form by precipitation from sea water. To calculate the mechanical rate of sedimentation the rate of input has to be known. This requires knowing the area of the depositional basin, the area supplying the sediments and its rate of land erosion. Rates used in early studies were largely intuitive. The calculations for carbonate rocks required knowing the rate of solution of calcium carbonate from the land surfaces, but could not correct for the variation of solubility with depth in the depositional basin. The number of unknown or crudely known parameters led to numerous divergent geological estimates for the Earth’s age, ranging from tens of millions to hundreds of millions of years.

4.1.3 Radioactivity

In 1896 a French physicist, Henri Becquerel, laid a sample of uranium ore on a wrapped, undeveloped photographic plate. After development the film showed the contour of the sample. The exposure was attributed to invisible rays emitted by the uranium sample. The phenomenon, which became known as radioactivity, provides the most reliable methods yet known of dating geological processes and calculating the age of the Earth and solar system. To appreciate radioactivity we must consider briefly the structure of the atomic nucleus.

The nucleus of an atom contains positively charged protons and electrically neutral neutrons. The electrostatic Coulomb force causes the protons to repel each other. It decreases as the inverse square of the separation (see Section 4.3.2) and so can act over distances that are large compared to the size of the nucleus. An even more powerful nuclear force holds the nucleus together. This attractive force acts between protons, between neutrons, and between proton and neutron. It is only effective at short distances, for a particle separation less than about $3 \times 10^{-15}$ m.

Suppose that the nucleus of an atom contains $Z$ protons and is surrounded by an equal number of negatively charged electrons, so that the atom is electrically neutral; $Z$ is the atomic number of the element and defines its place in the periodic table. The number $N$ of neutrons in the nucleus is its neutron number, and the total number $A$ of protons and neutrons is the mass number of the atom. Atoms of the same element with different neutron numbers are called isotopes of the element. For example, uranium contains 92 protons but may have 142, 143, or 146 neutrons. The different isotopes are distinguished by appending the mass number to the chemical symbol, giving $^{234}$U, $^{235}$U and $^{238}$U.

The Coulomb force of repulsion acts between every pair of protons in a nucleus, while the short-range nuclear force acts only on nearby protons and neutrons. To avoid flying apart due to Coulomb repulsion all nuclei with atomic number $Z$ greater than about 20 have an excess of neutrons ($N > Z$). This helps to dilute the effects of the repulsion-producing protons. However, nuclei with $Z \geq 83$ are unstable and disintegrate by radioactive decay. This means that they break up spontaneously by emitting elementary particles and other radiation.

At least 28 distinct elementary particles are known to nuclear physics. The most important correspond to the three types of radiation identified by early investigators and called α-, β-, and γ-rays. An α-ray (or α-particle) is a helium nucleus that has been stripped of its surrounding electrons; it is made up of two protons and two neutrons and so has atomic number 2 and mass number 4. A β-particle is an electron. Some reactions emit additional energy in the form of γ-rays, which have a very short wavelength and are similar in character to x-rays.

4.1.3.1 Radioactive decay

A common type of radioactive decay is when a neutron $n_0$ in the nucleus of an atom spontaneously changes to a proton $p^+$, and a β-particle (non-orbital electron). The β-particle is at once ejected from the nucleus along with another elementary particle called an antineutrino, $\nu$, which has neither mass nor charge and need not concern us further here. The reaction can be written

$$n_0 \Rightarrow p^+ + \beta^- + \nu \quad (4.3)$$

Radioactive decay is a statistical process. It is customary to call the nucleus that decays the parent and the nucleus after decay the daughter. It is not possible to say in advance which nucleus will spontaneously decay. But the probability that any one will decay per second is a constant, called the decay constant or decay rate. The number of disintegrations per second is called the activity of the nucleus. The derived unit of radioactivity, corresponding to one disintegration per second, is the becquerel (Bq).

Statistical behavior is really only applicable to large numbers, but the process of radioactive decay can be illustrated with a simple example. Suppose we start with 1000 nuclei, and the chance per second of a decay is 1 in 10, or 0.1. In the first second 10% of the parent nuclei spontaneously decay (i.e., 100 decays take place); the
number of parent nuclei is reduced to 900. The probability that any parent decays in the next second is still 1 in 10, so 90 further decays can be expected. Thus after 2 seconds the number of parent nuclei is reduced to 810, after 3 seconds to 729, and so on. The total number of parent nuclei constantly gets smaller but in principle it never reaches zero, although after a long time it approaches zero asymptotically. The decay is described by an exponential curve.

If the decay rate is equal to \( \lambda \), then in a short time-interval \( dt \) the probability that a given nucleus will decay is \( \lambda dt \); if at any time we have \( P \) parent nuclei the number that decay in the following interval \( dt \) is \( P\lambda dt \). The change \( dP \) in the number of \( P \) parent nuclei in a time interval \( dt \) due to spontaneous decays is

\[
\frac{dP}{dt} = -\lambda P
\]

which has the solution

\[
P = P_0 e^{-\lambda t}
\]  

(4.5)

Equation (4.5) describes the exponential decay of the number of parent nuclides, starting from an initial number \( P_0 \). While the number of parent nuclides diminishes, the number of daughter nuclides \( D \) increases (Fig. 4.3). \( D \) is the difference between \( P \) and \( P_0 \) and so is given by

\[
D = P_0 - P = P_0 (1 - e^{-\lambda t})
\]

(4.6)

The original amount \( P_0 \) of the parent nuclide is not known; a rock sample contains a residual amount \( P \) of the parent nuclide and an amount \( D \) of the daughter product. Eliminating the unknown \( P_0 \) from Eq. (4.5) and Eq. (4.6) gives

\[
D = P(e^{\lambda t} - 1)
\]

(4.7)

Equation (4.4) shows that the number of nuclear disintegrations per second – the activity of the nucleus – is proportional to the number of parent nuclei. The activity \( A \) at any given time is thus related to the initial activity \( A_0 \) by an equation similar to Eq. (4.5)

\[
A = \frac{dP}{dt} = -\lambda P_0 e^{-\lambda t}
\]

\[
A = A_0 e^{-\lambda t}
\]

(4.8)

The experimental description of radioactive decay by Ernest (later Lord) Rutherford and Frederick Soddy in 1902 was based on the observations of times needed for the activity of radioactive materials to decrease by steps of one-half. This time is known as the half-life of the decay. In the first half-life the number of parent nuclides decreases to a half, in the second half-life to a quarter, in the third to an eighth, etc. The number of daughter nuclides increases in like measure, so that the sum of parent and daughter nuclides is always equal to the original number \( P_0 \). Letting

\[
\frac{P}{P_0} \text{ equal } 1/2 \text{ in Eq. (4.5) we get the relationship between half-life } t_{1/2} \text{ and decay constant } \lambda:
\]

\[
t_{1/2} = \frac{\ln 2}{\lambda}
\]

(4.9)

The decay rates and half-lives are known for more than 1700 radioactive isotopes. Some are only produced in nuclear explosions and are so short-lived that they last only a fraction of a second. Other short-lived isotopes are produced by collisions between cosmic rays and atoms in the upper atmosphere and have short half-lives lasting minutes or days. A number of naturally occurring isotopes have half-lives of thousands of years (kiloyear, ka), millions of years (megayear, Ma) or billions of years (gigayear, Ga), and can be used to determine the ages of geological events.

4.1.4 Radiometric age determination

Each age-dating scheme involves precise measurement of the concentration of an isotope. This is usually very small. If the radioactive decay has advanced too far, the resolution of the method deteriorates. The best results for a given isotopic decay scheme are obtained for ages less than a few half-lives of the decay. The decay constants and half-lives of some radioactive isotopes commonly used in dating geological events are listed in Table 4.1 and illustrated in Fig. 4.4. Historical and archeological artefacts can be dated by the radioactive carbon method.

4.1.4.1 Radioactive carbon

The Earth is constantly being bombarded by cosmic radiation from outer space. Collisions of cosmic particles with
atoms of oxygen and nitrogen in the Earth's atmosphere produce high-energy neutrons. These in turn collide with nitrogen nuclei, transforming them into $^{14}\text{C}$, a radioactive isotope of carbon. $^{14}\text{C}$ decays by $\beta$-particle emission to $^{14}\text{N}$ with a half-life of 5730 yr. The production of new $^{14}\text{C}$ is balanced by the loss due to decay, so that a natural equilibrium exists. Photosynthesis in animals and plants replenishes living tissue using carbon dioxide, which contains a steady proportion of $^{14}\text{C}$. When an organism dies, the renewal stops and the residual $^{14}\text{C}$ in the organism decays radioactively.

The radioactive carbon method is a simple decay analysis based on Eq. (4.5). The remaining proportion $P$ of $^{14}\text{C}$ is measured by counting the current rate of $\beta$-particle activity, which is proportional to $P$. This is compared to the original equilibrium concentration $P_0$. The time since the onset of decay is calculated by solving Eq. (4.5) using the decay rate for $^{14}\text{C}$ ($\lambda = 1.21 \times 10^{-4} \text{ yr}^{-1}$).

The radioactive carbon method has been valuable in dating events in the Holocene epoch, which covers the last 10,000 yr of geological time, as well as events related to human prehistory. Unfortunately, human activity has disturbed the natural equilibrium of the replenishment and decay scheme. The concentration of $^{14}\text{C}$ in atmospheric carbon has changed dramatically since the start of the industrial age. Partly this is due to the combustion of fossil fuels like coal and oil as energy sources; they have long lost any $^{14}\text{C}$ and dilute its presence in atmospheric carbon. In the past half-century atmospheric testing of nuclear weapons doubled the concentration of $^{14}\text{C}$ in the atmosphere. Older materials may still be dated by the $^{14}\text{C}$ method, although natural fluctuations in $P_0$ must be taken into account. These are due to variations in intensity of the geomagnetic field, which acts as a partial shield against cosmic radiation.

4.1.4.2 The mass spectrometer

In the years before World War II, physicists invented the mass spectrometer, an instrument for measuring the mass of an ion. The instrument was further refined during the development of the atomic bomb. After the war it was adopted into the earth sciences to determine isotopic ratios and became a vital part of the process of isotopic age determination.

The mass spectrometer (Fig. 4.5) utilizes the different effects of electrical and magnetic fields on a charged particle, or ion. First, the element of interest is extracted from selected mineral fractions or from pulverized whole

Table 4.1  Decay constants and half-lives of some naturally occurring, radioactive isotopes commonly used in geochronology

<table>
<thead>
<tr>
<th>Parent isotope</th>
<th>Daughter isotope</th>
<th>Decay constant $[10^{-10} \text{ yr}^{-1}]$</th>
<th>Half-life [Ga]</th>
</tr>
</thead>
<tbody>
<tr>
<td>$^{40}\text{K}$</td>
<td>$^{40}\text{Ca}$</td>
<td>5.543</td>
<td>1.25</td>
</tr>
<tr>
<td>87% $^{40}\text{Rb}$</td>
<td>13% $^{40}\text{Ar}$</td>
<td>0.1420</td>
<td>48.8</td>
</tr>
<tr>
<td>$^{147}\text{Sm}$</td>
<td>$^{147}\text{Nd}$</td>
<td>0.0654</td>
<td>106.0</td>
</tr>
<tr>
<td>$^{232}\text{Th}$</td>
<td>$^{208}\text{Pb}$</td>
<td>0.4948</td>
<td>14.01</td>
</tr>
<tr>
<td>$^{235}\text{U}$</td>
<td>$^{207}\text{Pb}$</td>
<td>9.5485</td>
<td>0.704</td>
</tr>
<tr>
<td>$^{238}\text{U}$</td>
<td>$^{206}\text{Pb}$</td>
<td>1.5513</td>
<td>4.468</td>
</tr>
</tbody>
</table>

Fig. 4.4 Normalized radioactive decay curves of some important isotopes for dating the Earth and solar system. The arrows indicate the respective half-lives in 10$^9$ yr (data source: Dalrymple, 1991).

Fig. 4.5 Schematic design of a mass spectrometer, and hypothetical mass spectrum for analysis of strontium isotopes (after York and Farquhar, 1972).
rock. The extract is purified chemically before being introduced into the mass spectrometer, where it is ionized. For the analysis of a gas like argon, bombardment by a stream of electrons may be used. Solid elements, such as potassium, rubidium, strontium or uranium, are vaporized by depositing a sample on an electrically heated filament, or by heating with a high-energy laser beam so that the sample is vaporized directly. The latter process is known as “laser ablation.” The ions then enter an evacuated chamber and pass through an “ion gun,” where they are accelerated by an electrical field and filtered by a velocity selector. This device uses electrical and magnetic fields at right angles to the ion beam to allow only the passage of ions with a selected velocity \( v \). The ion beam is next subjected to a powerful uniform magnetic field \( B \) at right angles to its direction of motion. An ion with charge \( q \) experiences a Lorentz force (see Section 5.2.4) equal to \( (qvB) \) perpendicular to its velocity and to the magnetic field. Its trajectory is bent to form a circular arc of radius \( r \); the centrifugal force on the particle is equal to \( (mv^2/r) \). The curved path focusses the beam on a collector device that measures the intensity of the incident beam. The radius of the circular arc is given by equating the Lorentz and centrifugal forces:

\[
r = \frac{mv}{qB} \tag{4.10}
\]

The focal point of the path is determined by the mass of the ion and the strength of the magnetic field \( B \). In the case of a strontium analysis, the ion beam leaving the ion gun contains the four isotopes \(^{86}\text{Sr}, \(^{87}\text{Sr}, \(^{88}\text{Sr} \text{, and } \(^{84}\text{Sr} \text{. The beam splits along four paths, each with a different curvature. The magnetic field is adjusted so that only one isotope at a time falls on the collector. The incident current is amplified electronically and recorded. A spectrum is obtained with peaks corresponding to the incidence of individual isotopes (Fig. 4.5, inset). The intensity of each peak is proportional to the abundance of the isotope, which can be measured with a precision of about 0.1%. However, the relative peak heights give the relative abundances of the isotopes to a precision of about 0.001%.

Significant improvements in the design of mass spectrometer systems have increased their sensitivity further. Following laser ablation, the vaporized sub-micrometer-sized particles are mixed with an argon plasma. This is a gas consisting of positively charged argon ions and an equal number of unbound electrons at a temperature of 6000–10,000 K. The plasma ionizes the particles, which are extracted into a high vacuum before passing through an ion gun into the mass spectrometer. A drawback of the single-collector design described above is that it is vulnerable to fluctuations in the beam intensity. This problem has been overcome with the introduction of multiple-collector instruments, which allow simultaneous measurement of the split isotopic beams. The inductively coupled plasma mass spectrometer (ICP-MS) is a highly sensitive, versatile instrument for determining trace element abundances with a detection level in the region of parts per trillion. It can be used for analyzing both liquid and solid samples, and has many applications in fields outside the earth and environmental sciences.

An important development in the field of mass spectrometry is the ion microprobe mass analyzer. In conventional mass spectrometry the analysis of a particular element is preceded by separating it chemically from a rock sample. The study of individual minerals or the variation of isotopic composition in a grain is very difficult. The ion microprobe avoids contamination problems that can arise during the chemical separation and its high resolution permits isotopic analysis of very small volumes. Before a sample is examined in the instrument its surface is coated with gold or carbon. A narrow beam of negatively charged oxygen ions, about 3–10 \( \mu \)m wide, is focussed on a selected grain. Ions are sputtered out of the surface of the mineral grain by the impacting ion beam, accelerated by an electrical field and separated magnetically as in a conventional mass spectrometer. The ion microprobe allows description of isotopic concentrations and distributions in the surface layer of the grain and provides isotopic ratios. The instrument allows isotopic dating of individual mineral grains in a rock, and is especially well suited to the analysis of very small samples.

### 4.1.4.3 Rubidium–strontium

The use of a radioactive decay scheme as given by Eq. (4.6) assumes that the amount of daughter isotope in a sample has been created only by the decay of a parent isotope in a closed system. Usually, however, an unknown initial amount of the daughter isotope is present, so that the amount measured is the sum of the initial concentration \( D_0 \) and the fraction derived from decay of the parent \( P_0 \). The decay equation is modified to

\[
D = D_0 + P(e^{t/T} - 1) \tag{4.11}
\]

The need to know the amount of initial daughter isotope \( D_0 \) is obviated by the analytical method, which makes use of a third isotope of the daughter element to normalize the concentrations of daughter and parent isotopes. The rubidium–strontium method illustrates this technique.

Radioactive rubidium (\(^{87}\text{Rb} \)) decays by \( \beta \)-particle emission to radiogenic strontium (\(^{87}\text{Sr} \)). The non-radiogenic, stable isotope \(^{86}\text{Sr} \) has approximately the same abundance as the radiogenic product \(^{87}\text{Sr} \), and is chosen for normalization. Writing \(^{87}\text{Rb} \) for \( P \), \(^{87}\text{Sr} \) for \( D \) in Eq. (4.11) and dividing both sides by \(^{86}\text{Sr} \) gives

\[
\left( \frac{^{87}\text{Sr}}{^{86}\text{Sr}} \right) = \left( \frac{^{87}\text{Sr}}{^{86}\text{Sr}} \right)_0 + \left( \frac{^{87}\text{Rb}}{^{86}\text{Sr}} \right)(e^{t/T} - 1) \tag{4.12}
\]

In a magmatic rock, the isotopic ratio \( \left( \frac{^{87}\text{Sr}}{^{86}\text{Sr}} \right)_0 \) is uniform in all minerals precipitated from the melt because
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the isotopes are chemically identical (i.e., they have the same atomic number). However, the proportion of the different elements Rb and Sr varies from one mineral to another. Equation (4.12) can be compared with the equation of a straight line, such as

\[ y = y_0 + mx \]  

(4.13)

The ratio \( ^{87}\text{Sr} / ^{86}\text{Sr} \) is the dependent variable, \( y \), and the ratio \( ^{87}\text{Rb} / ^{86}\text{Sr} \) is the independent variable, \( x \). If we measure the isotopic ratios in several samples of the rock and plot the ratio \( ^{87}\text{Sr} / ^{86}\text{Sr} \) against the ratio \( ^{87}\text{Rb} / ^{86}\text{Sr} \), we get a straight line, called an isochron (Fig. 4.6). The intercept with the ordinate axis gives the initial ratio of the daughter isotope. The slope \( m \) of the line gives the age of the rock, using the known decay constant \( \lambda \):

\[ t = \frac{1}{\lambda} \ln(1 + m) = 7.042 \times 10^{10} \ln(1 + m) \]  

(4.14)

Because of its long half-life of 48.8 Ga (Fig. 4.4) the Rb–Sr method is well suited for dating very old events in Earth's history. It has been used to obtain the ages of meteorites and lunar samples, as well as some of the oldest rocks on Earth. For example, the slope of the Rb–Sr isochron in Fig. 4.6 yields an age of 3.62 Ga for the Early Precambrian (Archean) Uivak gneisses from eastern Labrador.

The Rb–Sr and other methods of isotopic dating can be applied to whole rock samples or to individual minerals separated from the rock. The decay equation applies only to a closed system, i.e., to rocks or minerals which have undergone no loss or addition of the parent or daughter isotope since they formed. A change is more likely in a small mineral grain than in the rock as a whole. Individual mineral isochrons may express the age of postformational metamorphism, while the whole rock isochron gives an older age. On the other hand, spurious ages may result from whole rock analyses if samples of different origin, and hence differing composition, are inadvertently used to construct an isochron.

4.1.4.4 Potassium–argon

For several reasons the potassium–argon (K–Ar) method is probably the age-dating technique most commonly used by geologists. The parent isotope, potassium, is common in rocks and minerals, while the daughter isotope, argon, is an inert gas that does not combine with other elements. The half-life of 1250 Ma (1.25 Ga) is very convenient. On the one hand, the Earth's age is equal to only a few half-lives, so radiogenic \(^{40}\text{K} \) is still present in the oldest rocks; on the other hand, enough of the daughter isotope \(^{40}\text{Ar} \) accumulates in 10^7 yr or so to give fine resolution. In the late 1950s the sensitivity of mass spectrometers was improved by constructing instruments that could be preheated at high temperature to drive off contaminating atmospheric argon. This made it possible to use the K–Ar method for dating lavas as young as a few million years.

Radioactive \(^{40}\text{K} \) constitutes only 0.01167% of the K in rocks. It decays in two different ways: (a) by \( \beta \)-particle emission to \(^{40}\text{Ca} \), with decay rate \( \lambda_{\text{Ca}} = 4.962 \times 10^{-10} \) yr\(^{-1} \) and (b) by electron capture to \(^{40}\text{Ar} \), with decay rate \( \lambda_{\text{Ar}} = 0.581 \times 10^{-10} \) yr\(^{-1} \). The combined decay constant \( \lambda = \lambda_{\text{Ca}} + \lambda_{\text{Ar}} \) is equal to \( 5.543 \times 10^{-10} \) yr\(^{-1} \). The decay schemes are, respectively:

(a) \( ^{40}\text{K} \) \( \Rightarrow ^{40}\text{Ca} + \text{e}^- \)  
(b) \( ^{40}\text{K} + \text{e}^- \Rightarrow ^{40}\text{Ar} \)  

(4.15)

Electron capture by a nucleus is more difficult and rarer than \( \beta \)-particle emission, so the decay of \(^{40}\text{K} \) to \(^{40}\text{Ca} \) is more common than the formation of \(^{40}\text{Ar} \). The ratio of electron capture to \( \beta \)-particle decay, \( \lambda_{\text{Ar}} / \lambda_{\text{Ca}} \), is called the branching ratio; it equals 0.117. Thus, only the fraction \( \lambda_{\text{Ar}} / (\lambda_{\text{Ar}} + \lambda_{\text{Ca}}) \), or 10.5%, of the initial radioactive potassium decays to argon. The initial amount of radiogenic \(^{40}\text{Ca} \) usually cannot be determined, so the decay to Ca is not used. Allowing for the branching ratio, the K–Ar decay equation is analogous to Eq. (4.7) with \(^{40}\text{Ar} \) for the accumulated amount of the daughter product \( D \) and \(^{40}\text{K} \) for the residual amount of the parent product \( P \):

\[ ^{40}\text{Ar} = \frac{\lambda_{\text{Ar}}}{\lambda_{\text{Ar}} + \lambda_{\text{Ca}}} \times ^{40}\text{K}(e^{\lambda t} - 1) \]

\[ = 0.1048 \times ^{40}\text{K}(e^{\lambda t} - 1) \]  

(4.16)

The potassium–argon method is an exception to the need to use isochrons. It is sometimes called an accumulation clock, because it is based on the amount of \(^{40}\text{Ar} \) that has accumulated. It involves separate measurements of the concentrations of the parent and daughter isotopes.
The amount of $^{40}\text{K}$ is a small but constant fraction ($0.01167\%$) of the total amount of $\text{K}$, which can be measured chemically. The $^{40}\text{Ar}$ is determined by mixing with a known amount of another isotope $^{38}\text{Ar}$ before being introduced into the mass spectrometer. The relative abundance of the two argon isotopes is measured and the concentration of $^{40}\text{Ar}$ is found using the known amount of $^{38}\text{Ar}$. By re-ordering Eq. (4.16) and substituting for the decay constant, the age of the rock is obtained from the K–Ar age equation:

$$e^{\lambda t} = 1 + 9.54 \frac{^{40}\text{Ar}}{^{40}\text{K}}$$

$$t = 1.804 \times 10^9 \ln \left( 1 + 9.54 \frac{^{40}\text{Ar}}{^{40}\text{K}} \right)$$  (4.17)

The $^{40}\text{Ar}$ in a molten rock easily escapes from the melt. It may be assumed that all of the radiogenic $^{40}\text{Ar}$ now present in a rock has formed and accumulated since the solidification of the rock. The method works well on igneous rocks that have not been heated since they formed. It cannot be used in sedimentary rocks that consist of the detritus of older rocks. Often it is unsuccessful in metamorphic rocks, which may have complicated thermal histories. A heating phase may drive out the argon, thereby re-setting the accumulation clock. This problem limits the usefulness of the K–Ar method for dating meteorites (which have a fiery entry into Earth’s atmosphere) and very old terrestrial rocks (because of their unknown thermal histories). The K–Ar method can be used for dating lunar basalts, as they have not been reheated since their formation.

4.1.4.5 Argon–argon

Some uncertainties related to post-formational heating of a rock are overcome in a modification of the K–Ar method that uses the $^{40}\text{Ar}/^{39}\text{Ar}$ isotopic ratio. The method requires conversion of the $^{39}\text{K}$ in the rock to $^{39}\text{Ar}$. This is achieved by irradiating the sample with fast neutrons in an atomic reactor.

The terms slow and fast refer to the energy of the neutron radiation. The energy of slow neutrons is comparable to their thermal energy at room temperature; they are also referred to as thermal neutrons. Slow neutrons can be captured and incorporated into a nucleus, changing its size without altering its atomic number. The capture of slow neutrons can increase the size of unstable uranium nuclei beyond a critical value and initiate fission. In contrast, fast neutrons act on a nucleus like projectiles. When a fast neutron collides with a nucleus it may eject a neutron or proton, while itself being captured. If the ejected particle is another neutron, no effective change results. But if the ejected particle is a proton (with a $\beta$-particle to conserve charge), the atomic number of the nucleus is changed. For example, bombarding $^{39}\text{K}$ nuclei in a rock sample with fast neutrons converts a fraction of them to $^{39}\text{Ar}$.

To determine this fraction a control sample of known age is irradiated at the same time. By monitoring the change in isotopic ratios in the control sample the fraction of $^{39}\text{K}$ nuclei converted to $^{39}\text{Ar}$ can be deduced. The age equation is similar to that given in Eq. (4.17) for the K–Ar method. However, $^{39}\text{Ar}$ replaces $^{40}\text{K}$ and an empirical constant $J$ replaces the constant 9.54. The value of $J$ is found from the control sample whose age is known. The $^{40}\text{Ar}/^{39}\text{Ar}$ age equation is

$$t = 1.804 \times 10^9 \ln \left( 1 + J \frac{^{40}\text{Ar}}{^{39}\text{Ar}} \right)$$  (4.18)

In the $^{40}\text{Ar}/^{39}\text{Ar}$ method the sample is heated progressively to drive out argon at successively higher temperatures. The $^{40}\text{Ar}/^{39}\text{Ar}$ isotopic ratio of the argon released at each temperature is determined in a mass spectrometer. The age computed for each increment is plotted against the percentage of Ar released. This yields an age spectrum. If the rock has not been heated since it was formed, the argon increments given out at each heating stage will yield the same age (Fig. 4.7a). An isochron can be constructed as in the Rb–Sr method by measuring the abundance of a non-radiogenic $^{36}\text{Ar}$ fraction and comparing the isotopic ratios $^{40}\text{Ar}/^{36}\text{Ar}$ and $^{39}\text{Ar}/^{36}\text{Ar}$. In an unheated sample all points fall on the same straight line (Fig. 4.7b).

If the rock has undergone post-formational heating, the argon formed since the heating is released at lower temperatures than the original argon (Fig. 4.7c). It is not certain why this is the case. The argon probably passes out of the solid rock by diffusion, which is a thermally activated process that depends on both the temperature and
the duration of heating. Unless the post-formational heating is long and thorough, only the outside parts of grains may be forced to release trapped argon, while the argon located in deeper regions of the grains is retained. In a sample that still contains original argon the results obtained at high temperatures form a plateau of consistent ages (Fig. 4.7c) from which the optimum age and its uncertainty may be calculated. On an isochron diagram reheated points deviate from the straight line defined by the high-temperature isotopic ratio (Fig. 4.7d).

The high precision of 40Ar/39Ar dating is demonstrated by the analysis of samples of melt rock from the Chicxulub Cretaceous–Tertiary impact crater in the Yucatan peninsula of Mexico. The crater is the favored candidate for the impact site of a 10 km diameter meteorite which caused global extinctions at the close of the Cretaceous period. Laser heating was used to release the argon isotopes. The age spectra of three samples show no argon loss (Fig. 4.8). The plateau ages are precisely defined and give a weighted mean age of 64.98 ± 0.05 Ma for the impact. This agrees closely with the 65.01 ± 0.08 Ma age of tektite glass found at a Cretaceous–Tertiary boundary section in Haiti, and so ties the impact age to the faunal extinctions.

The age spectrum obtained in Ar–Ar dating of the Menow meteorite is more complicated and shows the effects of ancient argon loss. Different ages are obtained during incremental heating below about 1200 °C (Fig. 4.9). The plateau ages above this temperature indicate a mean age of 4.48 ± 0.06 Ga. The shape of the age spectrum suggests that about 25% of the Ar was lost much later, about 2.5 Ga ago.

4.1.4.6 Uranium–lead: the concordia–discordia diagram

Uranium isotopes decay through a series of intermediate radioactive daughter products, but eventually they result in stable end-product isotopes of lead. Each of the decays is a multi-stage process but can be described as though it had a single decay constant. We can describe the decay of 238U to 206Pb by

\[
\frac{206\text{Pb}}{238\text{U}} = e^{\lambda_{238}t} - 1 \quad (4.19)
\]

Likewise the decay of 235U to 207Pb can be written

\[
\frac{207\text{Pb}}{235\text{U}} = e^{\lambda_{235}t} - 1 \quad (4.20)
\]

The 235U and 238U isotopes have well known decay constants: \(\lambda_{235} = 9.8485 \times 10^{-10}\) yr\(^{-1}\), \(\lambda_{238} = 1.55125 \times 10^{-10}\) yr\(^{-1}\). A graph of the 208Pb/238U ratio against the 207Pb/235U ratio is a curve, called the concordia line (Fig. 4.10). All points on this line satisfy Eqs. (4.19) and (4.20) simultaneously. The concordia line has a curved shape because the decay rates of the uranium isotopes, \(\lambda_{238}\) and \(\lambda_{235}\) respectively, are different. Calculations of the isotopic ratios given by Eqs. (4.19) and (4.20) defining the concordia line for the past 5 Ga are listed in Table 4.2.
The amounts of the daughter lead isotopes accumulate at different rates. Lead is a volatile element and is easily lost from minerals, but this does not alter the isotopic ratio of the lead that remains. Loss of lead will cause a point to deviate from the concordia line. However, because the isotopic ratio remains constant the deviant point lies on a straight line between the original age and the time of the lead-loss. This line corresponds to points that do not agree with the concordia curve; it is called the discordia line (Fig. 4.10). Different mineral grains in the same rock experience different amounts of lead loss, so the isotopic ratios in these grains give different points B, C and D on the discordia. The intersection of the concordia and discordia lines at A gives the original age of the rock, or in the case that it has lost all of its original lead, the age of this event. The intersection of the lines at E gives the age of the event that caused the lead loss.

The U–Pb method has been used to date some of the oldest rocks on Earth. The Duerrer formation in the Pilbara Supergroup in Western Australia contains Early Precambrian greenstones. U–Pb isotopic ratios for zircon grains separated from a volcanic rock in the Duerrer formation define a discordia line that intercepts the theoretical concordia curve at 3.45 ± 0.02 Ga (Fig. 4.11).
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Fig. 4.11 U–Pb concordia–discordia diagram for zircon grains from a volcanic rock in the Precambrian Duffer formation in the Pilbara Block, Western Australia (after Pidgeon, 1978).

4.1.4.7 Lead–lead isochrons

It is possible to construct isochron diagrams for the U–Pb decay systems. This is done (as in the Rb/Sr decay scheme) by expressing the radiogenic isotopes $^{206}\text{Pb}$ and $^{207}\text{Pb}$ as ratios of $^{204}\text{Pb}$, the non-radiogenic isotope of lead. The decay of $^{238}\text{U}$ to $^{206}\text{Pb}$ (or of $^{235}\text{U}$ to $^{207}\text{Pb}$) can then be described as in Eq. (4.11), where the initial value of the daughter product is unknown. This gives the decay equations

$$\frac{^{207}\text{Pb}}{^{204}\text{Pb}} \text{ vs } \frac{^{206}\text{Pb}}{^{204}\text{Pb}} = \frac{^{235}\text{U}}{^{208}\text{Pb}} \left( e^{\lambda t} - 1 \right)$$

(4.21)

$$\frac{^{206}\text{Pb}}{^{204}\text{Pb}} \text{ vs } \frac{^{207}\text{Pb}}{^{204}\text{Pb}} = \frac{^{238}\text{U}}{^{208}\text{Pb}} \left( e^{\lambda t} - 1 \right)$$

(4.22)

These equations can be combined into a single isochron equation:

$$\frac{^{207}\text{Pb}}{^{204}\text{Pb}} \text{ vs } \frac{^{206}\text{Pb}}{^{204}\text{Pb}} = \frac{^{235}\text{U}}{^{208}\text{Pb}} \left( e^{\lambda t} - 1 \right) \frac{^{238}\text{U}}{^{208}\text{Pb}} \left( e^{\lambda t} - 1 \right)$$

(4.23)

The ratio of $^{235}\text{U}$ to $^{238}\text{U}$ as measured today has been found to have a constant value 1/137.88 in lunar and terrestrial rocks and in meteorites. The decay constants are well known, so for a given age $t$ the right side in Eq. (4.23) is constant. The equation then has the form

$$y - y_0 = m (x - x_0)$$

(4.24)

This is the equation of a straight line of slope $m$ through the point $(x_0, y_0)$. The initial values of the lead isotope ratios are not known, but a plot of the isotopic ratio $^{207}\text{Pb}/^{204}\text{Pb}$ against the ratio $^{206}\text{Pb}/^{204}\text{Pb}$ is a straight line. The age of the rock cannot be found algebraically. Values for $t$ must be inserted successively on the right side of Eq. (4.23) until the observed slope is obtained.

4.1.5 Ages of the Earth and solar system

An important source of information concerning the Earth’s age is the radiometric dating of meteorites. A meteor is a piece of solid matter from space that penetrates the Earth’s atmosphere at a hypersonic speed of typically 10–20 km/s. Atmospheric friction causes it to become incandescent. Outside the Earth’s atmosphere it is known as a meteoroid; any part that survives passage through the atmosphere and reaches the Earth’s surface is called a meteorite. Most meteorites are thought to originate in the asteroid belt between the orbits of Mars and Jupiter (see Section 1.1.3.2), although tracking of entry paths shows that before colliding with Earth they have highly elliptical counterclockwise orbits about the Sun (in the same sense as the planets). Meteorites are often named after the place on Earth where they are found. They can be roughly divided into three main classes according to their composition. Iron meteorites consist of an alloy of iron and nickel; stony meteorites consist of silicate minerals; and iron–stony meteorites are a mixture of the two. The stony meteorites are further subdivided into chondrites and achondrites. Chondrites contain small spherules of high-temperature silicates, and constitute the largest fraction (more than 85%) of recovered meteorites. The achondrites range in composition from rocks made up essentially of single minerals like olivine to rocks resembling basaltic lava. Each category is further subdivided on the basis of chemical composition. All main types have been dated radiometrically, with most studies being done on the dominant chondrite fraction. There are no obvious age differences between the meteorites of the various groups. Chondrites, achondrites and iron meteorites consistently yield ages of around 4.45–4.50 Ga (Fig. 4.12).

In 1969, a large carbonaceous chondritic meteorite entered the Earth’s atmosphere and exploded, scattering its fragments over a large area around Pueblo del Allende in Mexico. The Allende meteorite contained refractory inclusions enriched in uranium relative to lead. Refractory inclusions are small aggregates of material that solidify and vaporize at high temperature, and thus can preserve information from the early history of the meteorite, and hence of the solar system. The high U/Pb ratios of the Allende inclusions gave precise $^{207}\text{Pb}/^{206}\text{Pb}$ dates averaging 4.566 ± 0.002 Ga (Allègre et al., 1995), which is the current best estimate of the age of the solar system.

A further bound on the age of the Earth is placed by the age of the Moon, which is thought to have formed during the accretion of the Earth. Six American manned missions and three Russian unmanned missions obtained samples of the Moon’s surface, which have been dated by several isotopic techniques. The ages obtained differ according to
the source areas of the rocks. The dark areas of the Moon's surface, the so-called lunar seas or mare, were formed when enormous outpourings of basaltic lava filled up low-lying areas – such as the craters formed by large meteoritic impacts – and so created flat surfaces. The lunar volcanism may have persisted until 1 Ga ago. The light areas on the Moon's surface are rough, extensively cratered highlands that reach elevations of 3000–4000 m above the maria. They represent the top part of the lunar crust and are the oldest regions of the Moon. Frequent collisions with large asteroids early in the Moon's history produced numerous craters and pulverized the lunar crust, leaving impact breccia, rock fragments and a dust layer a few meters thick, called the lunar regolith. Age dates from the highland rocks range from about 3.5–4.5 Ga, but the oldest age reported for a lunar rock is 4.51 ± 0.07 Ga, obtained by the Rb–Sr method.

In contrast to the ages of meteorites and the Moon, no rock of comparable age has been preserved on the Earth. Among the oldest terrestrial rocks are the Isua metasediments from western Greenland. They have been extensively studied and yield an age of 3.77 Ga that is consistent with different decay schemes. Even older are the Acasta gneisses in the northwestern part of the Canadian shield, which gave a U–Pb age of 3.96 Ga. Terrestrial rocks from the oldest Precambrian shield areas in Australia, South Africa, South America and Antarctica give maximum ages of 3.4–3.8 Ga.

Older ages are obtained from individual grains of zircon (ZrSiO₄), which forms as a trace mineral in granitic magmas. Zircon is a very durable mineral and can survive erosion that destroys the original rock. The zircon grains then become incorporated in sedimentary rocks. Zircon grains from sedimentary rocks in Australia have yielded U–Pb ages of 4.1–4.2 Ga; a fragment of a zircon grain, only 200 μm in diameter, has been dated by the U–Pb method at 4.404 ± 0.008 Ga, making it the oldest dated terrestrial solid. It is also possible to carry oxygen-isotope analyses on these ancient zircon grains. Although fragmentary, the results from zircon grains are important for understanding conditions on the primeval Earth. The scanty evidence suggests a possible history for the development of the hot early Earth, as follows.

The Sun and its accretionary disk, from which the solar system eventually evolved, formed 4.57 Ga ago. The accretion of the Earth and formation of its iron core lasted for about the first 100 Ma of Earth's early history. The accretion of Mars, which has only about one-ninth the mass of Earth, would have been completed in the first 30 Ma. Later in Earth's accretion, after perhaps 60 Ma, the Moon was formed as a result of the collision – called the “Giant Impact” – of a Mars-sized planetesimal with the proto-Earth, which was still much smaller than its present size. In the interval up to 4 Ga ago – called the Hadean (for hell-like) era – the early Earth was hot and covered by magma oceans; any crust would melt or be destroyed in the intense meteoritic bombardment. Water, if present, would be vaporized. After about 150 Ma the proto-Earth might have cooled sufficiently so that an early granitic crust, and possibly liquid water, could form. Only a few zircon grains that survived this period remain as possible witnesses. Repeated bombardments may have destroyed the crust and vaporized the water repeatedly in the following 400 Ma. Around 4 Ga ago the oldest surviving continental crust was formed.

This scenario is speculative and it is not unique. Some oxygen-isotope studies on zircon grains have been interpreted as indicating a cool early Earth. High δ²⁸O values (see Box 5.2) measured in these zircons imply surface temperatures low enough for liquid water; uniform conditions throughout the Archean era (4.4–2.6 Ga) were deduced. This model of early Earth suggests that meteoritic impacts may have been less intense than usually hypothesized. The model of a hot early Earth is more popularly accepted but the true history is not yet known. The events that happened in the first few hundred million years of its history are among Earth's best-kept secrets.

4.2.1 introduction

The radiant energy from the Sun, in conjunction with gravitational energy, determines almost all natural processes that occur at or above the Earth's surface. The hot incandescent Sun emits radiation in a very wide range of wavelengths. The radiation incident on the Earth is largely reflected into space, part enters the atmosphere and is reflected by the clouds or is absorbed and re-radiated into space. A very small part reaches the surface, where it is also partly reflected, especially from the water
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surfaces that cover three-quarters of the globe. Some is absorbed (e.g., by vegetation) and serves as the source of power for various natural cycles. A small fraction is used to heat up the Earth’s surface, but it only penetrates a short distance, some tens of centimeters in the case of the daily cycle and a few tens of meters for the annual changes. As a result, solar energy has negligible influence on internal terrestrial processes. Systems as diverse as the generation of the geomagnetic field and the motion of global lithospheric plates are ultimately powered by the Earth’s internal heat.

The Earth is constantly losing heat from its interior. Although diminutive compared to solar energy, the loss of internal heat is many times larger than the energy lost by other means, such as the change in Earth’s rotation and the energy released in earthquakes (Table 4.3). Tidal friction slows down the Earth’s rotation, and the change can be monitored accurately with modern technology such as very long baseline interferometry (VLBI) and the satellite-based geodetic positioning system (GPS). The associated loss of rotational energy can be computed accurately. The elastic energy released in an earthquake can be estimated reliably, and it is known that most of the energy is released in a few large shocks. However, the annual number of large earthquakes is very variable. The number with magnitude $M_r > 7$ varies between about 10 and 40 (see Fig. 3.49), giving estimates of the annual energy release from about $5 \times 10^{17}$ J to $4 \times 10^{19}$ J. The energies of tidal deceleration and earthquakes are small fractions of the geothermal flux, which is the most important form of energy originating in the body of the Earth.

The Earth’s internal heat derives from several sources (Section 4.2.5). For the past 4 Ga or so the Earth’s heat has been obtained from two main sources. One is the cooling of the Earth since its early history, when internal temperatures were much higher than they now are. The other is the heat produced by the decay of long-lived radioactive isotopes. This is the main source of the Earth’s internal heat, which, in turn, powers all geodynamic processes.

### Table 4.3 Estimates of notable contributions to the Earth’s annual energy budget

<table>
<thead>
<tr>
<th>Energy source</th>
<th>Annual energy [J]</th>
<th>Normalized [geothermal flux $= 1$]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reflection and re-radiation of solar energy</td>
<td>$5.4 \times 10^{24}$</td>
<td>$\approx 4000$</td>
</tr>
<tr>
<td>Geothermal flux from Earth’s interior</td>
<td>$1.4 \times 10^{21}$</td>
<td>1</td>
</tr>
<tr>
<td>Rotational deceleration by tidal friction</td>
<td>$\approx 10^{20}$</td>
<td>$\approx 0.1$</td>
</tr>
<tr>
<td>Elastic energy in earthquakes</td>
<td>$10^{19}$</td>
<td>$\approx 0.01$</td>
</tr>
</tbody>
</table>

portional to the temperature of the gas. If we add up the kinetic energies of all molecules in the container we obtain the amount of heat it contains. If heat is added to the container from an external source, the gas molecules speed up, their mean kinetic energy increases and the temperature of the gas rises.

The change of temperature of a gas is accompanied by changes of pressure and volume. If a solid or liquid is heated, the pressure remains constant but the volume increases. Thermal expansion of a suitable solid or liquid forms the principle of the thermometer for measuring temperature. Although Galileo reputedly invented an early and inaccurate “thermoscope,” the first accurate thermometers – and corresponding temperature scales – were developed in the early eighteenth century by Gabriel Fahrenheit (1686–1736), Ferchault de Réamur (1683–1757) and Anders Celsius (1701–1744). Their instruments utilized the thermal expansion of liquids and were calibrated at fixed points such as the melting point of ice and the boiling point of water. The Celsius scale is the most commonly used for general purposes, and it is closely related to the scientific temperature scale.

Temperature apparently has no upper limit. For example, the temperature of the surface of the Sun is less than 10,000 K but the temperature at its center is around 10,000,000 K and temperatures greater than 100,000,000 K have been achieved in physics experiments. But as heat is removed from an object it becomes more and more difficult to lower its temperature further. The limiting low temperature is often called “absolute zero” and is taken as the zero of the Kelvin temperature scale, named in honor of Lord Kelvin. Its divisions are the same as the Celsius scale and the temperature unit is called a kelvin. The scale is defined so that the triple point of water – where the solid, liquid and gaseous phases of water can coexist in equilibrium – is equal to 273.16 kelvins, written 273.16 K.

Heat was imagined by early investigators to be exchanged between bodies by the flow of a mystic fluid, called caloric. However, in the mid nineteenth century James Joule, an English brewer, demonstrated in a series of careful experiments that mechanical energy could be converted into heat. In his famous experiment, falling
Earth's age, thermal and electrical properties

weights drove a paddle wheel in a container of water, raising its temperature. The increase was tiny, less than 0.3 K, yet Joule was able to compute the amount of energy needed to raise the temperature by 1 K. His estimate of this energy – called the mechanical equivalent of heat – was within 5% of our modern value. The unit of energy is called the joule in recognition of his pioneering efforts. Originally, however, the unit of heat energy was defined as the amount needed to raise the temperature of one gram of water from 14.5 °C to 15.5 °C. This unit, the calorie (cal), is equivalent to 4.1868 J.

In physics and engineering it is often important to know the change of heat energy in a unit of time, known as the power. The unit of power is the watt, named after James Watt, the Scottish engineer who played an important role in harnessing thermal energy as a source of mechanical power. In geothermal problems we are usually concerned with the loss of heat from the Earth per unit area of its surface. This quantity is called the heat flux (or more commonly heat flow); it is the amount of heat that flows per second across a square meter of surface. The mean heat flow from the Earth is very small and is measured in units of milliwatt per square meter (mW m⁻²).

The addition of a quantity of heat ΔQ raises the temperature by an amount ΔT, which is proportional to ΔQ. The larger the mass m of the body, the smaller is the temperature change, and a given amount of heat produces different temperature changes in different materials. The amount of heat needed to raise the temperature of 1 kg of a material by 1 K is called its specific heat, denoted c_p for a process that occurs at constant pressure (and c_v when it happens at constant volume). These observations are summarized in the equation

\[ \Delta Q = c_p m \Delta T \]  

(4.25)

The added heat causes a fractional change of volume that is proportional to the temperature change but which differs from one material to another. The material property is called the volume coefficient of expansion α, and is defined by the equation

\[ \alpha = \frac{1}{V} \left( \frac{\partial V}{\partial T} \right)_p \]  

(4.26)

When thermal energy is added to a system, part is used to increase the internal energy of the system – i.e., the kinetic energy of the molecules – and part is expended as work, for example, by changing the volume. If the change in total energy ΔQ occurs at constant temperature T, we can define a new thermodynamic parameter, the entropy S, which changes by an amount ΔS equal to ΔQ/T. Thus we can write

\[ \Delta Q = T \Delta S = \Delta U + \Delta W \]  

(4.27)

where ΔU is the change of internal energy and ΔW is the work done externally. A thermodynamic process in which heat cannot enter or leave the system is said to be adiabatic.

The entropy of an adiabatic reaction remains constant: ΔS = 0. This is the case when a process occurs so rapidly that there is no time for heat transfer. An example is the passage of a seismic wave in which the compressions and rarefactions occur too rapidly for heat to be exchanged. The adiabatic temperature gradient in the Earth serves as an important reference for estimates of the actual temperature gradient and for determining how heat is transferred.

4.2.3 Temperature inside the Earth

In contrast to the radial distributions of density, seismic velocity and elastic parameters, which are known with a good measure of reliability, our knowledge of the temperature inside the Earth is still imprecise. The temperature can only be measured in the immediate vicinity of the Earth's surface, in boreholes and deep mines. As early as 1530 Georgius Agricola (the latinized name of Georg Bauer, a German physician and pioneer in mineralogy and mining) noted that conditions were warmer in deep mines. In fact, near-surface temperatures increase rapidly with depth by roughly 30 K km⁻¹. At this rate, linear extrapolation would give a temperature around 200,000 K at the center of the Earth. This is greater than the temperature of the surface of the Sun and is unrealistically high.

The conditions of high temperature and pressure in the deep interior can be inferred from experiments, and the adiabatic and melting-point temperatures can be computed with reasonable assumptions. Nevertheless, the temperature–depth profile is poorly known and conjectured temperatures have ranged widely. Limits are placed on the actual temperature by the known physical state of the Earth's interior deduced from seismology. The temperature in the solid inner core must be lower than the melting point, while the temperature of the molten outer core is above the melting point. Similarly the temperature in the solid mantle and crust are below the melting point; the asthenosphere has low rigidity because its temperature comes close to the solidus (“softening point”). The relationship of the actual temperature to the melting point determines how different parts of the Earth's interior behave rheologically (see Section 2.8).

The experimental approach to estimating the variation of temperature with depth combines knowledge obtained from seismology with laboratory results. The travel-times of seismic body waves show that changes in mineral structure (phase transitions) occur at certain depths (see Section 3.7.5). Important examples are the olivine–spinel transition at 400 km depth and the spinel–perovskite transition at 670 km depth in the upper mantle. The conditions of temperature and pressure (and hence depth) at which these phase transitions take place can be observed in laboratory experiments, so that the temperatures at the transition depths in the Earth can be determined. Similarly, the depth variation of the melting points of mantle rocks and the iron–nickel core can be inferred from laboratory observations at high pressure and temperature. Seismic
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velocities in the Earth are now so well known that deviations from normal velocities can be determined by seismic tomography (see Section 3.7.6) and interpreted in terms of temperature anomalies.

4.2.3.1 The adiabatic temperature gradient

An alternative way of estimating temperature inside the Earth is by using physical equations in which the parameters are known from other sources. In the late nineteenth century James Clark Maxwell expressed the laws of thermodynamics in four simple equations involving entropy (\(S\)), pressure (\(p\)), temperature (\(T\)) and volume (\(V\)). One of these equations is

\[
\left( \frac{\partial T}{\partial p} \right)_S = \left( \frac{\partial V}{\partial S} \right)_p
\]  

(4.28)

The left side is the adiabatic change in temperature with pressure, from which we obtain the adiabatic change in temperature with depth by substituting \(dp = \rho \partial \rho \partial T\), as in Section 3.7.4. Substituting from Eqs. (4.25) and (4.26) we get

\[
\left( \frac{\partial T}{\partial p} \right)_S = \frac{T \alpha V \Delta T}{c_p m \Delta \rho} = \frac{T \alpha}{c_p \rho}
\]  

(4.29)

from which we obtain

\[
\left( \frac{\partial T}{\partial z} \right)_{\text{adiabatic}} = \frac{T \alpha}{c_p \rho}
\]  

(4.30)

The dependence of density and gravity on depth \(z\) are known from seismic travel-times, and the profiles of \(\alpha\) and \(c_p\) can be estimated from laboratory observations (Fig. 4.13). For example, in the lower mantle at a depth of 1500 km, \(g = 9.9 \text{ m s}^{-2}\), \(c_p = 1200 \text{ J kg}^{-1} \text{ K}^{-1}\), \(\alpha = 14 \times 10^{-6} \text{ K}^{-1}\), and \(T = 2400 \text{ K}\). This gives an adiabatic temperature gradient of about 0.3 K km\(^{-1}\). In the outer core at about 3300 km depth the corresponding values are: \(g = 10.1 \text{ m s}^{-2}\), \(c_p = 700 \text{ J kg}^{-1} \text{ K}^{-1}\), \(\alpha = 14 \times 10^{-6} \text{ K}^{-1}\), and \(T = 4000 \text{ K}\) and the adiabatic temperature gradient is about 0.8 K km\(^{-1}\).

Approximate estimates of adiabatic temperatures inside the Earth can also be obtained with the aid of the Grüneisen thermodynamic parameter, \(\gamma\). This is a dimensionless parameter, defined as

\[
\gamma = \frac{\alpha K_s}{\rho c_p}
\]  

(4.31)

where \(K_s\) is the adiabatic incompressibility or bulk modulus. It is defined in Section 3.2.4 and Eq. (3.17), which, by writing \(dp\) instead of \(p\) and \(dV/V\) for the dilatation \(\theta\), becomes

\[
dp = - K_s V \theta = K_s \rho c_p \partial \rho
\]  

(4.32)

where \(\rho\) is the density. Substituting \(K_s\) \(\partial \rho \partial \rho\) in Eq. (4.29) gives

\[
\frac{dT}{dp} = T \frac{\gamma}{\rho c_p}
\]  

(4.33)

\[
T = T_0 \frac{p}{\rho c_p} \gamma
\]  

(4.34)

With this equation, and knowing the temperature \(T_0\) and density \(\rho_0\) at a given depth, the adiabatic temperature can be computed from the density profile in a region where the Grüneisen parameter \(\gamma\) is known. Fortunately, \(\gamma\) is fairly constant within large regions of the Earth’s interior (Fig. 4.13). Clearly, Eq. (4.34) cannot be applied across a boundary between these domains, where \(\gamma\) is discontinuous. If \(T_0\) and \(\rho_0\) are known at calibration points, the adiabatic temperature profile may be computed iteratively within a particular depth interval. A current estimate of the temperature profile in the Earth (Fig. 4.14) has steep gradients in the lithosphere, asthenosphere and in the D’ layer above the core–mantle boundary. It indicates a temperature near 3750 K at the core–mantle boundary and a central temperature of about 5100 K.

4.2.3.2 The melting point gradient

Another of Maxwell’s thermodynamic equations is

\[
\left( \frac{\partial S}{\partial p} \right)_T = - \left( \frac{\partial V}{\partial T} \right)_p
\]  

(4.35)
This equation can be applied to the effect of pressure on the melting point of a substance \( T_{mp} \). The heat required to melt a unit mass of the substance is its latent heat of fusion \( L \), so the change in entropy on the left-hand side of the equation is equal to \( (mL/T_{mp}) \). The volume change is the difference between that of the solid phase \( (V_s) \) and that of the liquid phase \( (V_L) \), so Eq. (4.35) can be rewritten

\[
\frac{dT_{mp}}{dp} = \frac{T_{mp}}{mL} (V_s - V_L)
\]

This is known to physicists as the Clausius–Clapeyron equation. It describes the effect of pressure on the melting point, and it is of interest to us because we can easily convert it to give the variation of melting point with depth, assuming that the pressure is hydrostatic so that \( dp = \rho g dz \) as previously. For a given mass \( m \) of the substance we can replace the volumes \( V_s \) and \( V_L \) with the corresponding densities \( \rho_s \) and \( \rho_L \) of the solid and liquid phases, respectively, so that

\[
\frac{1}{T_{mp}} \frac{dT_{mp}}{dz} = \frac{g}{L} \left( \frac{\rho_s}{\rho_L} - 1 \right)
\]

Again, to obtain the depth distribution of the melting point the variations of density and gravity with depth in the Earth are needed. At outer core pressures the densities of the solid and liquid phases of iron are about 13,000 kg m\(^{-3}\) and 11,000 kg m\(^{-3}\), respectively, and the latent heat of fusion of iron is about \( 7 \times 10^6 \) J kg\(^{-1}\), so the gradient of the melting point curve in the outer core is about 1 K km\(^{-1}\), i.e., the melting point in the core increases more steeply with depth than the adiabatic temperature. The computations of the adiabatic and melting temperature curves depend on parameters (e.g., \( L, \alpha, c_v \)) that are not known with a great degree of reliability in the Earth so the temperature profiles (Fig. 4.14) will undoubtedly change and become more secure as basic knowledge improves.

One factor that must still be evaluated is the role of phase transitions in the mantle. The D\(^\prime\) layer just above the core–mantle boundary evidently plays a crucial role in transferring heat from the core to the mantle. It constitutes a thermal boundary layer. Likewise the lithosphere forms a thermal boundary layer that conveys mantle heat to the Earth's surface. It appears unlikely that the phase transition at 400 km constitutes a thermal boundary layer but the phase transition at 670 km depth may do so. In the model used to derive the temperature profiles in Fig. 4.14 the phase transitions do not act as thermal boundary layers. Throughout most of the mantle the temperature gradient is assumed to equal the adiabatic gradient, but the mantle is bounded at top and bottom by thermal boundary layers (the lithosphere and D\(^\prime\)-layer, respectively) in which the temperature gradient greatly exceeds the adiabatic gradient.

### 4.2.4 Heat transport in the Earth

Heat can be transported by three processes: conduction, convection and radiation. Conduction and convection require the presence of a material; radiation can pass through space or a vacuum. Conduction is the most significant process of heat transport in solid materials and thus it is very important in the crust and lithosphere. However, it is an inefficient form of heat transport, and when the molecules are free to move, as in a fluid or gas, the process of convection becomes more important. Although the mantle is solid from the standpoint of the rapid passage of seismic waves, the temperature is high enough for the mantle to act as a viscous fluid over long time intervals. Consequently, convection is a more important form of heat transfer than conduction in the mantle. Convection is also the most important form of heat transport in the fluid core, where related changes in the geomagnetic field show that the turnover of core fluid is rapid in geological terms. Radiation is the least important process of heat transport in the Earth. It is only significant in the hottest regions of the core and lower mantle. The absorption of radiant energy by matter increases its temperature and thereby the temperature gradient. Hence, thermal radiation can be taken into account as a modification of the ability of the material to transfer heat by conduction.

#### 4.2.4.1 Conduction

Thermal conduction takes place by the transfer of kinetic energy between molecules or atoms. A true understanding
of the processes involved would require us to invoke quantum theory and the so-called “band theory of solids,” but a general understanding is possible without resorting to such measures. The electrons in an atom that are most loosely bound—the valence electrons—are essentially free of the ionic cores and can move through a material, so transferring kinetic energy. Hence they are called conduction electrons. Because electrons are electrically charged, the net movement of conduction electrons also causes an electrical current. Not surprisingly materials that are good electrical conductors (e.g., silver, copper) also conduct heat well. In this atomic view the conduction electrons move at very high speeds (~1000 km s⁻¹) but in random directions so that there is no net energy transfer in any particular direction. In an electrical or temperature field the conduction electrons drift systematically down the slope of the field (i.e., in the direction of the electrical field or temperature gradient). The additional drift velocity is very small (about 0.1 mm s⁻¹) but it passes kinetic energy through the material. This form of conduction is possible in liquids, gases or solids.

An additional mechanism plays an important role in conduction in solids. The atoms in a solid occupy definite positions that form a lattice with a certain symmetry. The atoms are not stationary but vibrate at a frequency that is temperature dependent. The lattice vibrational energy is quantized, forming units called phonons. An increase in temperature at one end of a solid raises the lattice vibrational frequency there. Due to the coupling between atoms the increased vibration is eventually passed through the lattice as an increase in temperature.

The relative importance of electrons and phonons in conducting heat differs from one solid to another. In metals, which contain large numbers of conduction electrons, thermal transport is due largely to the electrons; the lattice conductivity is barely measurable. In an insulator or poor conductor, such as the minerals of the crust and mantle, there are few conduction electrons and thermal conductivity is largely determined by lattice vibrations (phonons).

The transport of heat by conduction in a solid is governed by a simple equation. Consider a solid bar of length \( L \) and cross-sectional area \( A \) with its ends maintained at temperatures \( T_1 \) and \( T_2 \), respectively (\( T_2 > T_1 \)). Assuming that heat flows only along the bar (i.e., there are no side losses) the net amount of heat \( Q \) that passes in a given time from the hot end to the cold end depends directly on the temperature \( T_1 \) (and \( T_2 \)), the area of cross-section \( A \) and the temperature gradient \( (dT/dz) \), and inversely on the length of the bar \( L \). These observations can be summarized in the equation

\[
\Delta Q = kA\frac{T_2 - T_1}{L}\Delta t
\]

The constant of proportionality, \( k \), is the thermal conductivity, which is a property of the material of the bar. If the length of the bar is very small or the temperature change across it is uniform, the ratio \((T_2 - T_1)/L\) is the temperature gradient. We can modify the equation to describe the vertical flow of heat out of the Earth by substituting the vertical temperature gradient, \((dT/dz)\), which is also called the geothermal gradient. Equation (4.38) can then be rearranged as follows

\[
q_z = -\frac{1}{A} \frac{dQ}{dt} = -k \frac{dT}{dz}
\]

In this equation \( q_z \) is the heat flux, defined as the flow of heat per unit area per second. The negative sign is needed to account for the direction of the heat flow; if temperature increases in the downward direction of the \( z \)-axis, the flow of heat from high to low temperature is upward. The mean value of the undisturbed geothermal gradient near to the Earth’s surface is about 30 °C km⁻¹, with low values of around 10 °C km⁻¹ in ancient crust and high values of around 50 °C km⁻¹ in young active zones.

The change of temperature within a body is described by the heat conduction equation, which is solved in Section 4.2.6 for special situations that are of interest for the transfer of thermal energy in the Earth. Conduction is a slow, and less effective means of heat transport than convection. It is important in the rigid crust and lithosphere, where convection cannot take place. However, it cannot be neglected in the fluid core, which is metallic and therefore a good conductor. A significant part of the core’s heat is conducted out of the core along the adiabatic temperature gradient. The remainder, in excess of the conductive heat flow, is transported by convection currents.
4.2.4.2 Convection

Suppose that a small parcel of material at a certain depth in the Earth is in thermal equilibrium with its surroundings. If the parcel is displaced vertically upward without gaining or losing heat, it experiences a drop in pressure accompanied by a corresponding loss in temperature. If the new temperature of the parcel is the same as that of its surroundings at the new depth, the conditions at each depth are in adiabatic equilibrium. The variation of temperature with depth then defines the adiabatic temperature curve.

Now suppose that the real temperature increases with depth more rapidly than the adiabatic temperature gradient. The temperature loss of the upwardly displaced parcel is due to the change in pressure, which will be the same as in the previous case. But the real temperature has dropped by a larger amount, so the parcel is now hotter and therefore less dense than its surroundings. Its buoyancy causes it to continue to rise until it reaches a level where it is in equilibrium or can rise no further. Meanwhile, the volume vacated by the displaced parcel is occupied by adjacent material. Conversely, if a parcel of material is displaced downward, it experiences adiabatic increases in pressure and temperature. The temperature increase is less than required by the real temperature gradient, so the parcel remains cooler than its surroundings and sinks further. A pattern of cyclical behavior arises in which material is heated up and rises, while cooler material sinks to take its place, and is in turn heated up and rises, and so on. The process is called thermal convection and the physical transportation of material and heat is called a convection current.

The difference between the real and adiabatic temperature gradients is the superadiabatic temperature gradient, $\theta$. For thermal convection to take place in a fluid, $\theta$ must be positive. Suppose that the temperature at a certain depth exceeds the adiabatic temperature by an amount $\Delta T$. The temperature excess causes a volume $V$ of fluid to expand by an amount proportional to the volume coefficient of expansion, $\alpha$; this causes a mass deficiency of $(V \rho g \alpha \Delta T)$. Archimede’s principle applies, so the hot volume $V$ experiences a buoyancy force given by:

$$F_B = V \rho g \alpha \Delta T$$  \hspace{1cm} (4.40)

Two effects inhibit the hot volume from rising. First, some of the heat that would contribute to the buoyancy is removed by thermal conduction; the efficacy of this process is expressed by the thermal diffusivity $\kappa$ of the material, which depends on its density $\rho$, thermal conductivity $k$ and specific heat $c_p$ (see Section 4.2.6). Second, as soon as the overheated volume of fluid begins to rise, it experiences a resisting drag due to the viscosity $\eta$ of the fluid. The effects combine to produce a force, proportional to $\kappa \eta$, which opposes convection. If the volume $V$ involved in the convection has a typical dimension $D$, so that $V \sim D^3$, we can define a dimensionless number $Ra$, the Rayleigh number, which is proportional to the ratio of the buoyancy force to the diffusive–viscous force:

$$Ra = \frac{g \rho c_p \alpha \Delta T}{\kappa \eta \rho g}$$  \hspace{1cm} (4.41)

Initially, heat passes through the material by conduction, but the diffusion takes some time. If the heat flux is large enough, it cannot diffuse entirely. The temperature rises above the adiabatic and buoyancy forces develop. For convection to occur, the buoyancy forces must dominate the resisting forces. This does not happen until the Rayleigh number exceeds a critical value, which is determined additionally by the boundary conditions and the geometry of the convection. For example, the condition for the onset of convection in a thin horizontal fluid layer, heated from below and with the top and bottom surfaces free from stress, was shown by Lord Rayleigh in 1916 to depend on the value of $Ra$ given by:

$$Ra = \frac{g \alpha \theta D^4}{\kappa \eta}$$  \hspace{1cm} (4.42)

Here $D$ is the layer thickness, $\theta$ is the superadiabatic temperature gradient and $r$ (equal to $\eta \rho g$) is the kinematic viscosity. Convection begins in the flat layer if $Ra$ is greater than $27 \pi^4/4 = 658$. In cases with different boundary conditions, or for convection to occur in a spherical shell, the critical Rayleigh number is higher. However, convection generally originates if $Ra$ is of the order of $10^3$ and when $Ra$ reaches around $10^5$ heat transport is almost entirely by convection with little being transferred by diffusion.

For convection to occur, the real temperature gradient must exceed the adiabatic gradient. However, the loss of heat by convection reduces the difference between the gradients. Accordingly, the adiabatic gradient evolves as a convection fluid cools. An important effect of convection is to keep the temperature gradient close to the adiabatic gradient. This condition is realized in the Earth’s fluid core, where convection is the major mechanism of heat transport. Thermal convection is augmented by compositional convection related to the solidification of the inner core. The core fluid is made up of iron, nickel and lower-density elements, e.g., sulfur. Solidification of the inner core separates the dense iron from the lower-density elements at the inner core boundary. Being less dense than the core fluid, the residual materials experience an upward buoyancy force, resulting in a cycle of compositionally driven convection. Both thermal and compositional convection in the Earth's core each acts as a source of the energy needed to drive the geomagnetic field, with compositional convection the more important type.

Convection is the most important process of thermal transport in the fluid core, but it is also important in the mantle. The material of the Earth's mantle is rigid to the short-lived passage of seismic waves but is believed to yield slowly over long periods of time (section 2.8.6) Although the mantle viscosity is high, the time-scale of geological processes is so long that long-term flow can...
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4.2.4.3 Radiation

Atoms can exist in many distinct energy states. The most stable is the ground state, in which the energy is lowest. When an atom changes from an excited state to a lower-energy state, it is said to undergo a transition. Energy corresponding to the difference in energy between the states is emitted as an electromagnetic wave, which we call radiation. Quantum physics teaches that the radiant energy emitted consists of a discrete number of fundamental units, called quanta. The particular wavelength of the electromagnetic radiation associated with a transition is proportional to the energy difference between the two states. If several different transitions are taking place simultaneously, the body emits a spectrum of wavelengths. Radio signals, heat, light, and x-rays are examples of electromagnetic radiation that have different wavelengths. The electromagnetic wave consists of fluctuating electric and magnetic fields, which need no medium for their passage. For this reason, radiation can travel through space or a vacuum. In materials it may be scattered or absorbed, depending on its wavelength. Heat radiation corresponds to the infrared part of the electromagnetic spectrum with wavelengths just longer than those of visible light.

The radiation of a commonplace hot object depends on factors that are difficult to assess. Classical physics fails to explain adequately the absorption and emission of radiation. To provide an explanation physicists introduced the concept of a black body as a perfect absorber and emitter of radiation. At any temperature it emits a continuous spectrum of radiation; the frequency content of the spectrum does not depend on the material composition of the body but only on its temperature. An ideal black body does not exist in practice, but it can be approximated by a hollow container that has a small hole in its wall. When the container is heated, the radiation escaping through the hole – so-called cavity radiation – is effectively black-body radiation. In 1879 Josef Stefan pointed out that the loss of heat by radiation from a hot object is proportional to the fourth power of the absolute temperature. If \( R \) represents the radiant energy per second emitted per unit area of the surface of the body at temperature \( T \), then

\[
R = \sigma T^4
\]

(4.43)

where \( \sigma \), known as Stefan’s constant or the Stefan-Boltzmann constant, has the value \( 5.6704 \times 10^{-8} \text{ W m}^{-2} \text{K}^{-4} \).

In 1900, Max Planck, professor of physics at the university of Berlin, proposed that an oscillator could only have discrete amounts of energy. This was the birth of quantum theory. The energy of an oscillator of frequency \( \nu \) is equal to the product \( h \nu \), where the universal constant \( h \) (known as Planck’s constant) has the value \( 6.626 \times 10^{-34} \text{ J s} \). The application of quantum principles to black-body radiation provides a satisfactory explanation of Stefan’s law and allows Stefan’s constant to be expressed in terms of other fundamental physical constants.

Radiation is reflected and refracted in a transparent medium wherever the refractive index \( n \) changes; energy is transferred to the medium in each of these interactions. The transparency of the medium is determined by the opacity \( e \), which describes the degree of absorption of electromagnetic radiation. The opacity is wavelength dependent. In an ionic crystal the absorption of infrared radiation is large. It alters the vibrational frequency, and thereby influences the ability of the crystal lattice to transport heat by conduction. Thus the effect can be taken into account by increasing the conductivity by an extra radiative amount, \( k_r \), given by

\[
k_r = \frac{16}{3} \frac{n^2 \sigma}{e} T^3
\]

(4.44)

The \( T^4 \)-dependence in this expression suggests that radiation might be more important than lattice conductivity in the hotter regions of the Earth. In fact other arguments lead to the conclusion that this is probably not the case in the upper mantle, because the effect of increasing temperature is partly offset by an increase in the opacity, \( e \). The lower mantle is believed to have a high density of free electrons, which efficiently absorb radiation and raise the opacity. This may greatly reduce the efficacy of heat transfer by radiation in the mantle.

4.2.5 Sources of heat in the Earth

The interior of the Earth is losing heat via geothermal flux at a rate of about \( 4.4 \times 10^{13} \text{ W} \), which amounts to \( 1.4 \times 10^{21} \text{ J yr}^{-1} \) (see Table 4.3). The heat is brought to the surface in different ways. The creation of new lithosphere at oceanic ridges releases the largest fraction of the thermal energy. A similar mechanism, the spreading of the sea-floor, releases heat in the marginal basins behind island arcs. Rising plumes of magma originating deep in the mantle bring heat to the surface where they break through the oceanic or continental lithosphere at “hotspots,” characterized by intense localized volcanic
The early thermal history of the Earth is obscure and a matter of some speculation. According to the cold accretion model of the formation of the planets (see Section 1.1.4), colliding bodies in a primordial cloud of dust and gas coalesced by self-gravitation. The gravitational collapse released energy that heated up the Earth. When the temperature reached the melting point of iron, a liquid core formed, incorporating also nickel and possibly sulfur or another light element associated with iron. The differentiation of a denser core and lighter mantle from an initially homogeneous fluid must have released further gravitational energy in the form of heat. The dissipation of Earth's initial heat still has an important effect on internal temperatures.

Energy released by short-lived radioactive isotopes may have contributed to the initial heating, but the short-lived isotopes would be consumed quite quickly. The heat generated by long-lived radioactive isotopes has been an important heat source during most of Earth's history. These isotopes separated into two fractions: some, associated with heavy elements, sank into the core; some, associated with lighter elements, accumulated in the crust. The present distribution of radiogenic sources within the differentiated Earth is uneven. The highest concentrations are in the rocks and minerals of the Earth's crust, while the concentrations in mantle and core materials are low. However, continuing generation of heat by radioactivity in the deep interior, though small, may influence internal temperatures.

### 4.2.5.1 Radioactive heat production

When a radioactive isotope decays, it emits energetic particles and γ-rays. The two particles that are important in radioactive heat production are α-particles and β-particles. The α-particles are equivalent to helium nuclei and are positively charged, while, β-particles are electrons. In order to be a significant source of heat a radioactive isotope must have a half-life comparable to the age of the Earth, the energy of its decay must be fully converted to heat, and the isotope must be sufficiently abundant. The main isotopes that fulfil these conditions are 238U, 235U, 232Th and 40K.

The isotope 235U has a shorter half-life than 238U (see Table 4.1) and releases more energy in its decay. In natural uranium the proportion of 235U is 99.28%, that of 238U is about 0.71%, and the rest is 234U. The abundance of the radioactive isotope 40K in natural potassium is only 0.01167%, but potassium is a very common element and its heat production is not negligible. The amounts of heat generated per second by these elements (in μW kg⁻¹) are: natural uranium, 95.2; thorium, 25.6; and natural potassium, 0.00348 (Rybach, 1976, 1988). The heat \( Q \) produced by radioactivity in a rock that has concentrations \( C_U \), \( C_{Th} \) and \( C_K \), respectively, of these elements is

\[
Q = 95.2C_U + 25.6C_{Th} + 0.00348C_K 
\] (4.45)

Rates of radioactive heat production computed with this equation are shown for some important rock types in Table 4.4. Chondritic meteorites, made up of silicate minerals like olivine and pyroxene, are often taken as a proxy for the initial composition of the mantle; likewise, the olivine-dominated rock dunite represents the ultramafic rocks of the upper mantle. It is apparent that very little heat is produced by radioactivity in the mantle or in the basaltic rocks that dominate the oceanic crust and lower continental crust. The greatest concentration of radiogenic heat sources is in the granitic rocks in the upper continental crust. Multiplying the radioactive heat production values in the last column of Table 4.4 by the rock density gives the radiogenic heat generated in a cubic meter of the rock, \( A \). If we assume that all the heat generated in a rock layer of thickness \( D \) meters escapes vertically, the amount crossing a square meter at the surface per second (i.e., the radioactive component of the heat flow) is \( D \cdot A \). For example, a layer of granite 1 km thick contributes about 3 mW m⁻² to the continental heat flow. The figures suggest that the

### Table 4.4 Estimates of radioactive heat production in selected rock types, based on heat production rates (from Rybach, 1976, 1988) and isotopic concentrations

<table>
<thead>
<tr>
<th>Rock type</th>
<th>Concentration [p.p.m. by weight]</th>
<th>Heat production ( [10^{-11} \text{ W kg}^{-1}] )</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>( U )</td>
<td>( Th )</td>
</tr>
<tr>
<td>Granite</td>
<td>4.6</td>
<td>18</td>
</tr>
<tr>
<td>Alkali basalt</td>
<td>0.75</td>
<td>2.5</td>
</tr>
<tr>
<td>Tholeiitic basalt</td>
<td>0.11</td>
<td>0.4</td>
</tr>
<tr>
<td>Peridotite, dunite</td>
<td>0.006</td>
<td>0.02</td>
</tr>
<tr>
<td>Chondrites</td>
<td>0.015</td>
<td>0.045</td>
</tr>
<tr>
<td>Continental crust</td>
<td>1.2</td>
<td>4.5</td>
</tr>
<tr>
<td>Mantle</td>
<td>0.025</td>
<td>0.087</td>
</tr>
</tbody>
</table>
4.2 THE EARTH’S HEAT

10–20 km thick upper crust produces about one-half of the mean continental heat flow, which is 65 mW m$^{-2}$.

In fact the relative importance of radiogenic heat in the crust is variable from one region to another. A region in which the heat flow is linearly related to the heat produced by radioactivity is called a heat-flow province. Some examples of heat-flow provinces are Western Australia, the Superior Province in the Canadian Shield, and the Basin-and-Range Province in the western United States. As shown in Fig. 4.16 each province is characterized by a different linear relation between $q$ and $A$, such that

$$q = q_1 + DA$$

(4.46)

The parameters $q_1$ and $D$ typify the heat-flow province. The intercept of the straight line with the heat-flow axis, $q_1$, is called the reduced heat flow. This is the heat flow that would be observed in the province if there were no radiogenic crustal heat sources. It is due in part to the heat flowing from deeper regions of the Earth into the base of the crustal layer, and partly to cooling of the originally hotter upper crustal layer. Investigations in different heat-flow provinces show that the reduced heat flow averages about 55% of the mean measured heat flow in a province (Fig. 4.17).

The simplest interpretation of $D$ is to regard it as a characteristic thickness of crust involved in radioactive heat production. This assumes that the radiogenic heat sources are distributed uniformly in a crustal slab of constant thickness, which is an unlikely situation. A more likely model is that the radioactive heat generation decreases with depth. Assuming an exponential decrease, the heat production $A(z)$ at depth $z$ is related to the surface heat generation $A_0$ as

$$A(z) = A_0 e^{-z/D}$$

(4.47)

where $D$ is a characteristic depth (the depth at which $A(z)$ has decreased to $e^{-1}$ of its surface value). Integrating from the surface to infinite depth gives the total radioactive heat production:

$$\varphi \int_{0}^{\varphi} A(z) dz = A_0 \int_{0}^{\infty} e^{-z/D} dz = DA_0$$

(4.48)

which is the same as for the uniform distribution. The infinite lower limit to the exponential distribution is obviously unrealistic. If the radiogenic sources are distributed in a layer of finite thickness $s$, the integration becomes

$$A_0 \int_{0}^{s} e^{-z/D} dz = DA_0(1 - e^{-s/D})$$

(4.49)

If $s$ is greater than three times $D$, this expression differs from $DA_0$ by less than 5%. The value of $D$ estimated from studies of heat-flow and radioactive heat generation averages about 10 km, but varies from $\sim 4$ km to $\sim 16$ km (Fig. 4.17).

The three main sources of the Earth’s surface heat flow are (i) heat flowing into the base of the lithosphere from the deeper mantle, (ii) heat lost by cooling of the lithosphere with time, and (iii) radiogenic heat production in the crust. The contributions are unequal and different in the oceans and continents (Table 4.5). The most obvious disparity is in the relative importance of lithospheric cooling and radioactivity. The lithosphere is hot when created at oceanic ridges and cools slowly as it ages. The

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**Fig. 4.16** Dependence of surface heat flux on radioactive heat generation in two heat-flow provinces: (a) Sierra Nevada, (b) eastern United States (data source: Roy et al., 1968).

**Fig. 4.17** Mean heat flow, reduced heat flow and characteristic thickness of the layer of radioactive heat production in several heat-flow provinces (data source: Vitorello and Pollack, 1980).
loss of heat by lithospheric cooling is most pronounced in the oceanic crust, which moreover contains few radiogenic heat sources. In contrast, the older continental lithosphere has lost much of the early heat of formation, and the higher concentration of radioactive minerals increases the importance of radiogenic heat production. Regardless of its source, the passage of heat through the rigid outer layers takes place predominantly by conduction, although in special circumstances, such as the flow of magma in the crust or hydrothermal circulation near to oceanic ridges, convection also plays an important role.

### 4.2.6 The heat conduction equation

Jean Baptiste Joseph Fourier (1768–1830), a noted French mathematician and physicist, developed the theory of heat conduction in 1822. Here we consider the example of one-dimensional heat flow, which typifies many interesting problems involving the flow of heat in a single direction. The equation of heat conduction is most easily developed for this case, from which it can be readily extended to three-dimensional heat flow.

Consider the flow of heat in the negative direction of the z-axis through a small rectangular prism with sides dx, dy and dz (see Fig. 4.15b). We will assume that there are no sources of heat inside the box. Let the amount of heat entering the prism at (z + dz) be \( Q_z \). This is equal to the heat flow \( q_y \) multiplied by the area of the surface it flows across (dx dy) and by the duration of the flow (dt). The heat leaving the box at z is \( Q_z - dQ_z \), which can be written as \( -dQ_z / dz \). The increase in heat in the small box is the difference between these amounts:

\[
\frac{dQ_z}{dz} = \frac{dq_y}{dz}(dx dy)dt = k \frac{dT}{dz} dV dt \tag{4.50}
\]

where \( dV \) is the volume of the box (dx dy dz). Note that \( Q_z, q_y, T \) and \( dT \) are all understood to decrease in the direction of flow, so we have substituted for \( q_y \) from Eq. (4.39) without using the negative sign. The heat increase in the box causes its temperature to rise by an amount \( dT \), determined by the specific heat at constant pressure \( (c_p) \) and the mass of material \( (m) \) in the box. Using Eq. (4.25), we write

\[
c_p \rho dT = c_p \rho dV dT \tag{4.51}
\]

where \( \rho \) is the density of the material in the box. If we equate Eq. (4.50) and Eq. (4.51) for the amount of heat left in the box, we get the equation of heat conduction

\[
\frac{dT}{dt} = \frac{k}{\rho c_p} \frac{d^2T}{dz^2}
\]

where \( k (= k/\rho c_p) \) is called the thermal diffusivity; it has the dimensions \( m^2 s^{-1} \). The equation is written with partial differentials because the temperature \( T \) is a function of both time and position: \( T = T(z, t) \). This just means that, on the one hand, the temperature at a certain position changes with time, and, on the other hand, the temperature at any given time varies with position in the body.

The same arguments can be applied to the components of heat flow through the box in the x- and y-directions. We obtain the three-dimensional heat flow equation (also called the diffusion equation):

\[
\frac{\partial T}{\partial t} = \kappa \left( \frac{\partial^2 T}{\partial x^2} + \frac{\partial^2 T}{\partial y^2} + \frac{\partial^2 T}{\partial z^2} \right) \tag{4.53}
\]

The equation may be solved for any set of boundary conditions using the method of separation of variables (Box 4.1). Two important situations involving the flow of heat across the Earth’s surface are the heating of Earth’s surface by solar energy and the cooling of hot lithosphere. These can be handled to a first approximation as problems of one-dimensional heat conduction.

### 4.2.6.1 Penetration of external heat into the Earth

Solar energy is by far the greatest of the Earth’s energy sources (Table 4.4). In order to determine the geothermal flux from the Earth’s interior, it is important to understand the effects of solar energy that reaches the Earth’s surface. Rocks at Earth’s surface heat up during the day and cool down at night. The effect is not restricted to the immediate surface, but affects a volume of rock near the surface. Similarly, the mean surface temperature varies throughout the year with the changing seasons. The heat conduction equation allows us to estimate what depths are affected by these cyclic temperature variations.

Suppose that the surface temperature of the Earth varies cyclically with angular frequency \( \omega \), so that at time \( t \) it is equal to \( T_0 \cos \omega t \), where \( T_0 \) is the peak temperature during a cycle. The temperature at time \( t \) and depth \( z \) is obtained by solving the one-dimensional heat equation (Box 4.2) and is given by

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**Table 4.5** Approximate relative contributions (in %) of the main sources of heat flow in oceanic and continental lithosphere (from Bott, 1982)

<table>
<thead>
<tr>
<th>Heat source</th>
<th>Contribution to heat flow in:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Continents</td>
</tr>
<tr>
<td>Cooling of the lithosphere</td>
<td>20%</td>
</tr>
<tr>
<td>Heat flow from below the lithosphere</td>
<td>25%</td>
</tr>
<tr>
<td>Radiogenic heat:</td>
<td></td>
</tr>
<tr>
<td>upper crust</td>
<td>55%</td>
</tr>
<tr>
<td>rest of lithosphere</td>
<td>40%</td>
</tr>
<tr>
<td>rest of continental crust</td>
<td>15%</td>
</tr>
</tbody>
</table>

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4.2 THE EARTH’S HEAT

Box 4.1: Method of separation of variables
The method of separation of variables is encountered in the solution of several geophysical problems. It may be illustrated by the case of one-dimensional heat flow, described by Eq. (4.52). The coordinates in this equation are independent, i.e., each can take any value regardless of the other. Assume that the temperature \( T(z,t) \) can be written as the product of \( Z(z) \), a function of position only, and \( \theta(t) \), a function of time only:

\[
T(z,t) = Z(z)\theta(t)
\]  

(1)

The actual solution for \( T(z,t) \) usually does not have this final form, but once \( Z(z) \) and \( \theta(t) \) are known they can be combined to fit the boundary conditions. On substituting in Eq. (4.52) we get

\[
\frac{d^2 \theta}{dz^2} = \kappa \frac{d^2 Z}{dz^2}
\]

(2)

Note that we can use full differentials here because \( \theta \) only depends on \( t \) and \( Z \) only on \( z \). Dividing both sides by \( Z\theta \) gives

\[
\frac{1}{\theta} \frac{d\theta}{dt} = \frac{\kappa}{Z} \frac{d^2 Z}{dz^2}
\]

(3)

Note that in Eq. (3) the left side depends only on \( t \) and the right side depends only on \( z \), and that \( z \) and \( t \) are independent variables. On substituting a particular value for the time \( t \) the left side becomes a numerical constant. We have not restricted \( z \), which varies independently. Equation (3) requires that the variable right side remains equal to the numerical constant for any value of \( z \). Conversely, if we substitute a particular value for \( z \), the right side becomes a (new) numerical constant and the variable left side must equal this constant for any \( t \). The identity inherent in Eq. (3) implies that each side must be equal to the same separation constant. Let this constant be \( C \). Then,

\[
\frac{1}{\theta} \frac{d\theta}{dt} = C
\]

\[
\kappa \frac{1}{Z} \frac{d^2 Z}{dz^2} = C
\]

(4)

The value of the separation constant in a particular problem is determined by the boundary conditions of the problem.

\[
T(z,t) = T_0 e^{-z/d} \cos \left( \frac{\omega t - z}{d} \right)
\]

(4.54)

where \( d = (2\pi \omega / \omega_i)^{1/2} \) is the decay depth of the temperature. At a depth of \( 5d \) the amplitude is less than 1% of the surface value and is effectively zero. Note that \( d \) depends inversely on the frequency, so long-period fluctuations penetrate more deeply than rapid fluctuations. This is illustrated by a quick comparison of the decay depths for daily and annual temperature variations in the same ground

\[
d_{\text{annual}} = \sqrt{\frac{(2\pi/1)}{(2\pi/365)}} = \sqrt{365} = 19.1
\]

(4.55)

i.e., the annual variation penetrates about 19 times the depth of the daily variation (Fig. 4.18). Moreover, the temperature at any depth varies with the same frequency \( \omega \) as the surface variation, but it experiences a phase shift or delay, reaching its maximum value progressively later than the surface effect (Fig. 4.18).

As representative values for crustal rocks we take: density, \( \rho = 2650 \text{ kg m}^{-3} \); thermal conductivity, \( k = 2.5 \text{ W m}^{-1} \text{ K}^{-1} \); and specific heat \( c_v = 700 \text{ J kg}^{-1} \text{ K}^{-1} \). These give a thermal diffusivity, \( \kappa = 1.25 \times 10^{-6} \text{ m}^2 \text{ s}^{-1} \). The daily temperature variation (period 86,400 s) has \( \omega = 7.27 \times 10^{-3} \text{ s}^{-1} \), so its penetration depth is about 20 cm. The daily variation has negligible effect deeper than about a meter. Similarly, the annual temperature variation accompanying seasonal changes has a penetration depth of 3.8 m and is negligible deeper than about 19 m. Heat-flow measurements made within 20 m of the surface will be contaminated by the daily and annual variations of surface temperature. This is not a problem in the deep oceans, where the Sun’s rays never reach the bottom, but it must be taken into account in continental heat-flow measurements. A serious effect is the role of the ice ages, which recur on a timescale of about 100,000 yr and have a penetration depth of several kilometers. The measured temperature gradient must be corrected appropriately.

4.2.6.2 Cooling of the oceanic lithosphere
A thermodynamic problem commonly encountered in geology is the sudden heating or cooling of a body. For example, molten lava intrudes cool host rocks as a dike or sill virtually instantaneously, but the heat is conducted to the adjacent rocks over a long period of time until it slowly dissipates. An important case in the context of plate tectonics is the cooling of the oceanic lithosphere, with its base at the temperature of the hot mantle, \( T_m \), and its top in contact with cold ocean-bottom sea-water.

Assume that the fresh hot lithosphere is created at a ridge axis as a thin vertical prism with a uniform initial temperature equal to that of the hot mantle, \( T_m \). Sea-floor spreading transports the lithosphere horizontally, so, for a constant spreading rate, distance from the ridge axis is proportional to the cooling time \( t \). Except in the immediate vicinity of a spreading ridge the heat loss may be assumed to be solely in the vertical \((-z)\) direction through the surface at \( z = 0 \), which is in contact with cold ocean-bottom sea-water at a temperature of 0 °C. Although the lithosphere has finite vertical extent, the temperature in the cooling slab may be approximated as the one-dimensional cooling of a semi-infinite half-space extending to infinity in the \( z \)-direction. The error introduced by
Box 4.2: One-dimensional heat conduction

The surface-temperature variation $T_0 \cos \omega t$ can be expressed with the aid of complex numbers (Box 2.5) as the real part of $T_0 e^{i \omega t}$. Let $z$ represent the depth of a point below the surface. The heat conduction equation, Eq. (4.52), can be separated and written as two parts equal to the same constant. We want to match boundary conditions with the surface disturbance $T_0 e^{i \omega t}$, so we write the separation constant as $i \omega$. Applying the method of separation of variables (Box 4.1) we get for the two parts of the solution

$$\frac{d \theta}{d t} = i \omega \quad (1)$$
$$\frac{1}{\kappa} \frac{d^2 Z}{d z^2} = i \omega \quad (2)$$

The solution of Eq. (1) gives the time dependence of the temperature as

$$\theta = \theta_0 e^{i \omega t} \quad (3)$$

In order to find the depth dependence, we can rewrite Eq. (2) as

$$\frac{d^2 Z}{d z^2} - i \omega \kappa Z = 0 \quad (4)$$

Writing $- n^2 = i \omega / \kappa$, this is equivalent to the harmonic equation

$$\frac{d^2 Z}{d z^2} + n^2 Z = 0 \quad (5)$$

which has the solutions

$$Z = Z_0 e^{inz} \quad \text{and} \quad Z = Z_1 e^{-inz} \quad (6)$$

where

$$n = \sqrt{\frac{i \omega}{\kappa}} = \sqrt{\frac{\omega}{2 \kappa}} (1 + i) \quad (7)$$

We get two possible solutions for the depth variation:

$$Z = Z_0 e^{inz} = Z_1 e^{(i \omega / 2 \kappa)(1 + i)z} \quad \text{and}$$

$$Z = Z_1 e^{inz} = Z_0 e^{-(i \omega / 2 \kappa)(1 + i)z} \quad (8)$$

The temperature must decrease with increasing depth $z$ below the surface, so only the second solution is acceptable. Combining the solutions for $\theta$ and $Z$ we get

$$T(z, \omega t) = Z_0 e^{-(i \omega / 2 \kappa)z} e^{i \omega t} \quad (9)$$
$$T(z, t) = Z_0 \theta_0 e^{-(i \omega / 2 \kappa)z} e^{i \omega t} \quad (10)$$

The surface-temperature variation $T_0 \cos \omega t$ was expressed as the real part of $T_0 e^{i \omega t}$. Taking the real part of Eq. (10) we get for the temperature $T(z, t)$ at time $t$ and depth $z$

$$T(z, t) = Z_0 \theta_0 e^{-(i \omega / 2 \kappa)z} \cos (\omega t - \sqrt{\frac{\omega}{2 \kappa}}) \quad (11)$$

Writing $T_0 = Z_0 \theta_0$ and the equation reduces to

$$T(z, t) = T_0 e^{-z/d} \cos \left( \frac{\omega t - z}{d} \right) \quad (12)$$

The parameter $d$ is called the decay depth of the temperature. At this depth the amplitude of the temperature fluctuation is attenuated to $1/e$ of its value on the surface. Eq. (12) can also be written in the form

$$T(z, t) = T_0 e^{-z/d} \cos \phi (t - \tau) \quad (13)$$

where the phase difference, or delay time,

$$\tau = \frac{z}{\omega d} \quad (14)$$

represents the length of time by which the temperature at depth $z$ lags behind the surface temperature.

This simplification is small, and the simple model can be used to estimate the heat flow from the cooling lithosphere with acceptable accuracy.

The one-dimensional cooling of a semi-infinite half-space is described in Appendix B. The solution with the stated boundary conditions involves the error function, which is described in Box 4.3. The error function (erf) and complementary error function (erfc) of the parameter $x$ are defined as

$$\text{erf}(x) = \frac{2}{\sqrt{\pi}} \int_0^x e^{-u^2} du$$
$$\text{erfc}(x) = 1 - \text{erf}(x) \quad (4.56)$$

The shapes of these functions are shown in Fig. 4.19; their values for any particular value of $x$ are obtained from tables, just like other statistical or trigonometric functions. The complementary error function sinks asymptotically to zero, and is effectively zero for $x \geq 2$. The temperature $T$ at depth $z$ and time $t$ after the half-space starts to cool is given by

$$T = T_m \text{erf} \left( \frac{z}{2 \sqrt{\kappa t}} \right) = T_m \text{erf} (\eta) \quad \text{with} \quad \eta = \frac{z}{2 \sqrt{\kappa t}} \quad (4.57)$$

where $T_m$ is the temperature of the hot mantle, $\kappa$ is the thermal diffusivity of the half-space, and the surface $(z = 0)$ is at the temperature of the ocean floor, which is taken to be 0°C.
The heat loss from the cooling lithosphere can be computed from the temperature distribution in Eq. (4.57). The heat flow is proportional to the temperature gradient (Eq. (4.39)). The heat flowing out of a semi-infinite half-space is thus obtained by differentiating Eq. (4.57) with respect to \( z \):

\[
q_z = -\frac{kT_m}{\sqrt{\pi kt}} \frac{\partial}{\partial \eta} \left\{ T_m \text{erf}(\eta) \right\}
\]

which simplifies to

\[
q_z = -\frac{kT_m}{\sqrt{\pi kt}} \frac{\partial}{\partial \eta} \left\{ e^{-\eta^2} \right\}
\]

At the surface, \( z = 0, \eta = 0, \) and \( \exp(-\eta^2) = 1 \). The surface heat flow at time \( t \) is

\[
q_z = -\frac{kT_m}{\sqrt{\pi kt}} \frac{\partial}{\partial \eta} \left\{ e^{-\eta^2} \right\}
\]

The negative sign here indicates that the heat flows upward, in the direction of decreasing \( z \). The semi-infinite half-space is quite a good model for the cooling of oceanic lithosphere. The oceanic heat flow indeed varies with distance from an oceanic ridge as \( 1/\sqrt{t} \), where \( t \) is the corresponding age of the lithosphere. Models for the cooling of oceanic lithosphere are discussed further in Section 4.2.8.3.

### 4.2.7 Continental heat flow

The computation of heat flow at a locality requires two measurements. The thermal conductivities of a representative suite of samples of the local rocks are measured in the laboratory. The temperature gradient is measured in the field at the investigation site. At continental sites this is usually carried out in a borehole (Fig. 4.20). There are several ways of determining the temperature in the borehole. During commercial drilling the temperature of the drilling fluid can be measured as it returns to the surface. This gives a more or less continuous record, but is influenced strongly by the heat generated during the drilling. At times when drilling is interrupted, the bottom-hole temperature can be measured. Both of these methods give data of possibly commercial interest but they are too inaccurate for heat-flow determination.

In-hole measurements of temperature for heat-flow analyses are made by lowering a temperature-logging tool into the borehole and continuously logging the temperature during its descent. The circulation of drilling fluids redistributes heat in the hole, so it is necessary to allow some time after drilling has ceased for the hole to return to thermal equilibrium with the penetrated formations. The temperature of the water in the hole is taken to be the ambient temperature of the adjacent rocks, provided there are no convection currents.

The most common devices for measuring temperature are the platinum resistance thermometer and the thermistor. A thermistor is a ceramic solid-state device with an electrical resistance that is strongly dependent on temperature. Its resistance depends non-linearly on temperature, requiring accurate calibration, but the sensitivity of the device makes feasible the measurement of temperature differences of 0.001–0.01 K. The platinum resistance thermometer and the thermistor are used in two basic ways. In one method the sensor element constitutes an arm of a sensitive Wheatstone bridge, with which its resistance is measured directly. The other common method uses the thermal sensor as the resistive element in a tuned electrical circuit. The tuned frequency depends on the resistance of the sensor element, which is related in a known way to temperature.

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**Fig. 4.18** Temperature variations at various depths in a sandy soil: (a) daily fluctuations, (b) annual (seasonal) variations.

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\[
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\]

At the surface, \( z = 0, \eta = 0, \) and \( \exp(-\eta^2) = 1 \). The surface heat flow at time \( t \) is

\[
q_z = -\frac{kT_m}{\sqrt{\pi kt}} \frac{\partial}{\partial \eta} \left\{ e^{-\eta^2} \right\}
\]

The negative sign here indicates that the heat flows upward, in the direction of decreasing \( z \). The semi-infinite half-space is quite a good model for the cooling of oceanic lithosphere. The oceanic heat flow indeed varies with distance from an oceanic ridge as \( 1/\sqrt{t} \), where \( t \) is the corresponding age of the lithosphere. Models for the cooling of oceanic lithosphere are discussed further in Section 4.2.8.3.

### 4.2.7 Continental heat flow

The computation of heat flow at a locality requires two measurements. The thermal conductivities of a representative suite of samples of the local rocks are measured in the laboratory. The temperature gradient is measured in the field at the investigation site. At continental sites this is usually carried out in a borehole (Fig. 4.20). There are several ways of determining the temperature in the borehole. During commercial drilling the temperature of the drilling fluid can be measured as it returns to the surface. This gives a more or less continuous record, but is influenced strongly by the heat generated during the drilling. At times when drilling is interrupted, the bottom-hole temperature can be measured. Both of these methods give data of possibly commercial interest but they are too inaccurate for heat-flow determination.

In-hole measurements of temperature for heat-flow analyses are made by lowering a temperature-logging tool into the borehole and continuously logging the temperature during its descent. The circulation of drilling fluids redistributes heat in the hole, so it is necessary to allow some time after drilling has ceased for the hole to return to thermal equilibrium with the penetrated formations. The temperature of the water in the hole is taken to be the ambient temperature of the adjacent rocks, provided there are no convection currents.

The most common devices for measuring temperature are the platinum resistance thermometer and the thermistor. A thermistor is a ceramic solid-state device with an electrical resistance that is strongly dependent on temperature. Its resistance depends non-linearly on temperature, requiring accurate calibration, but the sensitivity of the device makes feasible the measurement of temperature differences of 0.001–0.01 K. The platinum resistance thermometer and the thermistor are used in two basic ways. In one method the sensor element constitutes an arm of a sensitive Wheatstone bridge, with which its resistance is measured directly. The other common method uses the thermal sensor as the resistive element in a tuned electrical circuit. The tuned frequency depends on the resistance of the sensor element, which is related in a known way to temperature.
Errors may be of two types: systematic and random. A systematic error results, for example, when the measuring device is wrongly calibrated (e.g., if times are measured with a clock that runs “fast” or “slow”). Random errors occur naturally when a value is measured a large number of times. The observations will be distributed randomly about their mean value. Usually there will be a few large deviations from the mean and a lot of small deviations, and there will be as many negative as positive deviations. The scatter of the results can be described by the standard deviation of the mean.

The observations will be distributed randomly about their mean value. Usually there will be a few large deviations from the mean and a lot of small deviations, and there will be as many negative as positive deviations. The scatter of the results can be described by the standard deviation of the mean, which is often called a “bell curve” because of its shape (Fig. B4.3). If the mean of the distribution of the parameter is zero, the standard deviation is described by the probability density function.

\[ f(u) = \frac{1}{\sqrt{2\pi}} e^{-u^2/2} \]

The standard normal distribution is defined so that it has a mean of zero and standard deviation \( \sigma = 1 \). When integrated from \(-\infty\) to \(+\infty\), the area under this curve is

\[ \int_{-\infty}^{\infty} f(u) du = \frac{1}{\sqrt{2\pi}} \int_{-\infty}^{\infty} e^{-u^2/2} du = 1 \]

The error function is closely related to the standard normal distribution. However, only positive values are considered, so the graph of the defining function is similar to the right half of the curve in Fig. B4.3a. It is given as

\[ f(u) = \frac{1}{\sqrt{\pi}} e^{-u^2} \]

The area under this curve from the origin at \( u = 0 \) to the value \( u = x \) (Fig. B4.3b) defines the error function \( \text{erf}(x) \):

\[ \text{erf}(x) = \frac{2}{\sqrt{\pi}} \int_{0}^{x} e^{-u^2} du \]

The complementary error function, \( \text{erfc}(x) \), is defined as

\[ \text{erfc}(x) = 1 - \text{erf}(x) = \frac{2}{\sqrt{\pi}} \int_{x}^{\infty} e^{-u^2} du \]

The value of \( \text{erf}(x) \) or \( \text{erfc}(x) \) for any particular value of \( x \) may be obtained from tables. Some useful properties of the error function and complementary error function are:

\[ \text{erf}(\infty) = \frac{2}{\sqrt{\pi}} \int_{0}^{\infty} e^{-u^2} du = 1 \]

\[ \frac{d}{dx}(\text{erf}(x)) = \frac{2}{\sqrt{\pi}} e^{-x^2} \]

Some values of \( \text{erf}(x) \) are listed in the following table.

<table>
<thead>
<tr>
<th>( x )</th>
<th>( \text{erf}(x) )</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.05</td>
<td>0.05637</td>
</tr>
<tr>
<td>0.1</td>
<td>0.11246</td>
</tr>
<tr>
<td>0.15</td>
<td>0.16800</td>
</tr>
<tr>
<td>0.2</td>
<td>0.22270</td>
</tr>
<tr>
<td>0.25</td>
<td>0.27633</td>
</tr>
<tr>
<td>0.3</td>
<td>0.32863</td>
</tr>
<tr>
<td>0.4</td>
<td>0.42839</td>
</tr>
<tr>
<td>0.5</td>
<td>0.52050</td>
</tr>
<tr>
<td>0.6</td>
<td>0.60386</td>
</tr>
<tr>
<td>0.7</td>
<td>0.67780</td>
</tr>
<tr>
<td>0.8</td>
<td>0.74210</td>
</tr>
<tr>
<td>0.9</td>
<td>0.79691</td>
</tr>
<tr>
<td>1.0</td>
<td>0.84270</td>
</tr>
<tr>
<td>1.2</td>
<td>0.91031</td>
</tr>
<tr>
<td>1.4</td>
<td>0.95229</td>
</tr>
<tr>
<td>1.5</td>
<td>0.97635</td>
</tr>
<tr>
<td>1.6</td>
<td>0.99532</td>
</tr>
</tbody>
</table>
4.2 THE EARTH’S HEAT

From the measured temperature distribution, the average temperature gradient is computed for a geological unit or a selected depth interval (Fig. 4.20). The gradient is then multiplied by the mean thermal conductivity of the rocks to obtain the interval (or formation) heat flow. Thermal conductivity can show large variations even between adjacent samples, so the harmonic mean of at least four samples is usually used. The interval heat-flow values may then be averaged to obtain the mean heat flow for the borehole (Table 4.6).

The computation of continental heat flow from borehole data requires the implementation of several corrections. An important assumption is that heat flow is only vertical. Well below the surface the isotherms (surfaces of constant temperature) are flat lying and the flow of heat (normal to the isotherms) is vertical. However, the surface of the Earth is also presumed to be locally isothermal (i.e., to have constant temperature) near to the borehole. The near-surface isotherms adapt to the topography (Fig. 4.21) so that the direction of heat flow is deflected and acquires a horizontal component, while the vertical temperature gradient is also modified. Consequently, heat flow measured in a borehole must be corrected for the effects of local topography.

The need for a topographic correction was recognized late in the nineteenth century. Further corrections must be applied for long-term effects such as the penetration of external heat related to cyclical climatic changes, for example the ice ages. Erosion, sedimentation and changes in the thermal conductivities of surface soils are other long-term effects that may require compensation.

4.2.7.1 Reconstruction of ground surface temperature changes from borehole temperature profiles

The main variation of temperature with depth in the Earth is the geothermal gradient related to the outflow of heat from the Earth’s deep interior. Over limited depth intervals the temperature profile is linear, its slope varying by region with local conditions. However, changes in temperature at Earth’s surface affect the subsurface temperature distribution close to the surface (Section 4.2.6.1). Rapid temperature changes have shallow penetration, but slow changes can modify subsurface temperatures well below the surface. For example, daily variations do not reach below about a meter, but temperature variations over a century may extend to about 200 m depth.

The temperature profiles in boreholes from eastern Canada (Fig. 4.22) show effects of surface temperature variations above depths of 180–250 m, below which the regular geothermal gradient is evident. The curved upper segment contains information about the history of temperature changes at the surface. This history can be retrieved by inversion of the borehole temperature data, which is a complex and sophisticated process. A large database of borehole temperature measurements, used for the determination of the global heat flow pattern (Section 4.2.8.2), has been analyzed to obtain the variation of mean surface temperature since 1500 (Fig. 4.23). Instrumental measurements of air temperature since 1860 agree well with the long-term surface temperatures estimated from the borehole data. The results show a consistent increase in surface temperature over the past five centuries.

4.2.7.2 Variation of continental heat flow with age

Many processes contribute to continental heat flow. Apart from the heat generated by radioactive decay, the most important sources are those related to tectonic events. During an orogenic episode various phenomena may introduce heat into the continental crust. Rocks may be deformed and metamorphosed in areas of continental collision. In extensional regions the crust may be thinned, with intrusion of magma. Uplift and erosion of elevated areas and deposition in sedimentary basins also affect the surface heat flow. After a tectonic event

Table 4.6 Computation of heat flow from temperature measurements in WSR 1, a 570 m deep borehole, and thermal conductivity measurements on cored samples (after Powell et al., 1988).

<table>
<thead>
<tr>
<th>Depth interval [m]</th>
<th>Temperature gradient [mK m⁻¹]</th>
<th>Thermal conductivity [W m⁻¹ K⁻¹]</th>
<th>Interval heat flow [mW m⁻²]</th>
</tr>
</thead>
<tbody>
<tr>
<td>45–105</td>
<td>15.0</td>
<td>3.96</td>
<td>60</td>
</tr>
<tr>
<td>105–245</td>
<td>18.0</td>
<td>3.43</td>
<td>62</td>
</tr>
<tr>
<td>245–320</td>
<td>24.8</td>
<td>2.75</td>
<td>68</td>
</tr>
<tr>
<td>320–455</td>
<td>16.0</td>
<td>4.18</td>
<td>67</td>
</tr>
<tr>
<td>455–515</td>
<td>17.2</td>
<td>4.20</td>
<td>72</td>
</tr>
<tr>
<td>515–575</td>
<td>16.5</td>
<td>3.86</td>
<td>64</td>
</tr>
<tr>
<td></td>
<td>Mean heat flow = 65 mW m⁻²</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
convective cooling takes place efficiently in circulating fluids, while some excess heat is lost by conductive cooling. Consequently, the variation of continental heat flow with time is best understood in terms of the tectonothermal age, which is the age of the last tectonic or magmatic event at a measurement site. The continental heat-flow values comprise a broad spectrum. Even when grouped in broad age categories there is a large degree of overlap (Fig. 4.24). The greatest scatter is seen in the youngest regions. The mean heat flow decreases with increasing crustal or tectonothermal age (Fig. 4.25), falling from 70–80 mW m$^{-2}$ in young provinces to a steady-state value of 40–50 mW m$^{-2}$ in Precambrian regions older than $\sim$800 Ma.

4.2.7.3 Heat transfer through porous crustal rocks

The transfer of heat through the continental crust takes place not only by conduction but also by advection. A sediment or rock is composed of mineral grains closely packed in contact with each other. The pore spaces between the grains can represent an appreciable fraction of the total volume of a rock sample. This fraction, sometimes expressed as a percentage, is the porosity $\phi$ of the rock. A highly porous rock, for example, may have a porosity of 0.3, or 30%, implying that only 70% of the rock is solid mineral. The porosity depends on how the mineral grains are arranged, how well they are cemented, and on their degree of sorting. Well sorted sediments have fairly uniform grain and pore sizes; in poorly sorted sediments, there is a range of grain sizes, so the finer grains may “block” the voids between the larger grains, reducing the porosity. Igneous and metamorphic rocks often contain cracks and fissures, which, if sufficiently numerous, may give these rocks a low porosity. The degree to which the pore spaces are connected with each other determines the permeability of the rock, which is its ability to transmit fluids such as water and petroleum.

Permeability is defined by an empirical relationship observed in 1856 by Henry Darcy, a French hydraulic engineer. He observed that the volume of fluid per second crossing a surface was proportional to the area of the surface ($A$) and the gradient of the hydraulic pressure head ($\frac{dp}{dx}$) driving the flow, assumed here to be in the $x$-direction, and inversely to the viscosity ($\mu$) of the liquid.
If the velocity of the flow is $v$, then in one second the volume crossing a surface $A$ is $(vA)$, so we have

$$ (vA) \frac{dp}{dx} = -k \frac{dA}{dx} $$

$$ v = -k \frac{dp}{dx} $$

$$ v = -K \frac{dp}{dx} \tag{4.61} $$

This equation, in which $K = k/\mu$ defines the hydraulic conductivity, is known as Darcy’s law. Its derivation and form are analogous to the law of heat conduction (Eq. (4.38), Fig. 4.15) and Ohm’s law for electrical currents (Eq. (4.71), Fig. 4.40). The negative sign in the equation indicates that the flow is in the direction of decreasing pressure. The constant $k$ is the permeability, which has dimensions $m^2$. However, the unit of permeability used in hydraulic engineering is called the darcy (equivalent to $9.87 \times 10^{-13} m^2$) or, more practically, the millidarcy (md). For example, an approximate range for the permeability of gravel is $10^5$–$10^6$ md, that of sandstone is $1$–$100$ md, and that of granite is $10^{-3}$–$10^{-5}$ md.

The ability of fluids to flow through crustal rocks enables them to transmit heat. In this case the process of heat transfer is not by convection, because the fluid motion is not driven by temperature differences but by the pressure gradient. The heat transfer “piggybacks” on the fluid motion, and the process is called advection (Section 4.2.4.2). The motion of water through the continental crust provides a source of geothermal energy, which can be tapped in several interesting ways for commercial purposes.

### 4.2.8 Oceanic heat flow

Whereas the mean altitude of the continents is only 840 m above sea-level, the mean depth of the oceans is 3880 m; the abyssal plains in large ocean basins are 5–6 km deep.
At these depths extra-terrestrial heat sources have no effect on heat-flow measurements and the flatness of the ocean bottom (except near ridge systems or seamounts) obviates the need for topographic corrections. Measuring the heat flow through the ocean bottom presents technical difficulties that were overcome with the development of the Ewing piston corer. This device, intended for taking long cores of marine sediment from the ocean floor, enables in-situ measurement of the temperature gradient. It consists of a heavily weighted, hollow sampling pipe (Fig. 4.26a), commonly about 10 m long although in special cases cores over 20 m in length have been taken (very long coring pipes tend to bend before they reach maximum penetration). A plunger inside the pipe is displaced by sediment during coring and makes a seal with the sediment surface, so that sample loss and core deformation are minimized when the core is withdrawn from the ocean floor. Thermistors are mounted on short arms a few centimeters from the body of the pipe, and the temperatures are recorded in a water-tight casement. The instrument is lowered from a surface ship until a free-dangling trigger-weight makes contact with the bottom (Fig. 4.26b). This releases the corer, which falls freely and is driven into the sediment by the one-ton lead weight. The friction accompanying this process generates heat, but the ambient temperatures in the sediments can be measured and recorded before the heat reaches the offset sensors (Fig. 4.26c). The sediment-filled corer is hauled back on board the ship, where the thermal conductivity of the sediment can be determined. The recovered core is used for paleontological, sedimentological, geochemical, magnetostratigraphic and other scientific analyses.

Special probes have been devised explicitly for in-situ measurement of heat flow. They consist of two parallel tubes about 3–10 m in length and 5 cm apart. One tube is about 5–10 cm in diameter and provides strength; the other, about 1 cm in diameter, is oil filled and contains arrays of thermistors. After penetration of the ocean-bottom sediments, as described for the Ewing corer, the equilibrium temperature gradient is measured. A known electrical current, either constant in value or pulsed, is then passed along a heating wire and the temperature response is recorded. The observations allow the thermal heat flow to be determined.
conductivity of the sediment to be found. In this way a complete determination of heat flow is obtained without having to recover the contents of the corer.

4.2.8.1 Variation of oceanic heat flow and depth with lithospheric age

The most striking feature of oceanic heat flow is the strong relationship between the heat flow and distance from the axis of an oceanic ridge. The heat flow is highest near to the ridge axis and decreases with increasing distance from it. For a uniform sea-floor spreading rate the age of the oceanic crust (and lithosphere) is proportional to the distance from the ridge axis, and so the heat flow decreases with increasing age (Fig. 4.27). The lithospheric plate accretes at the spreading center, and as the hot material is transported away from the ridge crest it gradually cools. Model calculations for the temperature in the cooling plate are discussed in the next section: they all predict that the heat flow \( q \) (caused by cooling of the plate) decreases with age \( t \) as \( \frac{1}{\sqrt{t}} \), when the age of the plate is less than about 55–70 Ma. Older lithosphere cools slightly less rapidly. Currently the decrease in heat flow with age is best explained by a global model called the Global Depth and Heat Flow model (GDH1). The model predicts the following relationships between heat flow \( q \) (mW m\(^{-2}\)) and age \( t \) (Ma):

\[
q = \begin{cases} \frac{510}{\sqrt{t}} & (t \leq 55\text{Ma}) \\ q_s [1 + 2\exp(-\kappa \pi^2 t/a^2)] & (t > 55\text{Ma}) \\ 48 + 96\exp(-0.0278t) \end{cases}
\]  

(4.62)

Here \( q_s \) is the asymptotic heat flow, to which the heat flow decreases over very old oceanic crust (\( \approx 48\text{ mW m}^{-2} \)), \( a \) is the asymptotic thickness of old oceanic lithosphere (\( \approx 95\text{ km} \)), and \( \kappa \) is its thermal diffusivity (\( \approx 0.8 \times 10^{-6} \text{ m}^2\text{s}^{-1} \)).

Close to a ridge axis the measured heat flow is unpredictable: extremely high values and very low values have been recorded. Over young lithosphere the observed heat flow is systematically less than the values predicted by
cooling models (Fig. 4.27). The divergence is related to the process of accretion of the new lithosphere. At a ridge crest magma erupts in a narrow zone through feeder dikes and/or supplies horizontal lava flows. Very hot material is brought in contact with sea water, which cools and fractures the fresh rock. The water is in turn heated rapidly and a hydrothermal circulation is set up, which transports heat out of the lithosphere by convection. The eruption of hot hydrothermal currents has been observed directly from manned submersibles in the axial zones of oceanic ridges. The expeditions witnessed strong outpourings of mineral-rich hot water (called “black smokers” and “white smokers”) in the narrow axial rift valley. The heat output of these vents is high: the power associated with a single vent has been estimated to be about 200 MW.

About 30% of the hydrothermal circulation takes place very near to the ridge axis through crust younger than 1 Ma. The rest is due to off-ridge circulation, which is possible because the fractured crust is still permeable to sea-water at large distances from the ridge axis. As it moves away from the ridge, sedimentation covers the basement with a progressively thicker layer of low permeability sediments, inhibiting the convective heat loss. The hydrothermal circulation eventually ceases, perhaps because it is sealed by the thick sediment cover, but probably also because the cracks and pore spaces in the crust become closed with increasing age. This is estimated to take place by about 55–70 Ma, because for greater ages the observed decrease in heat flow is close to that predicted by plate cooling models. The hydrothermal circulation in oceanic crust is an important part of the Earth’s heat loss. It accounts for about a third of the total oceanic heat flow, and a quarter of the global heat flow.

The free-air gravity anomaly over an oceanic ridge system is generally small and related to ocean-bottom topography (Section 2.6.4.3), which suggests that the ridge system is isostatically compensated. As hot material injected at the ridge crest cools, its volume contracts and its density increases. To maintain isostatic equilibrium a vertical column sinks into the supporting substratum as it cools. Consequently, the depth of the ocean floor (the top surface of the column) is expected to increase with age of the lithosphere. The cooling half-space model predicts an increase in depth proportional to \( \sqrt{t} \), where \( t \) is the age of the lithosphere, and this is observed up to an age of about 80 Ma (Fig. 4.28). However, the square-root relationship is not the best fit to the observations. Other cooling models fit the observations more satisfactorily, although the differences from one model to another are small. Beyond 20 Ma the data are better fitted by an exponential decay. The optimum relationships between depth \((d, m)\) and age \((t, \text{Ma})\) can be written:

\[
\begin{align*}
  d &= 2600 + 365\sqrt{t} \quad (t < 20\text{Ma}) \\
  d &= d_i + [1 - (8/\pi^2)] \exp(-k\pi^2/t/a^2)] \quad (t \geq 20\text{Ma})
\end{align*}
\]

where \( d_i \) is the mean depth of the ocean floor at ridge crests, \( d_i \) is the asymptotic subsidence of old lithosphere and the other parameters are as before.

**Fig. 4.27** Comparison of observed and predicted heat flow as a function of age of oceanic lithosphere. (a) Schematic summary for all oceans, showing the influence of hydrothermal heat flow at the ocean ridges (after Anderson and Skilbeck, 1981). Comparisons with the reference cooling models PSM (Parsons and Sclater, 1977) and GDH1 (Stein and Stein, 1992) for (b) the Pacific, (c) Atlantic and (d) Indian oceans.
Oceanic heat flow has been measured routinely in oceanographic surveys since the 1950s and in situ profiles have been made since the 1970s. In contrast to the measurement of continental heat flow it is not necessary to have an available (and usually expensive) drillhole. However, the areal coverage of the oceans by heat-flow measurements is uneven. A large area in the North Pacific Ocean is still unsurveyed, and most of the oceanic areas south of about latitude 35°S (the approximate latitude of Cape Town or Buenos Aires) are unsurveyed or only sparsely covered. The uneven data distribution is dense along the tracks of research vessels and absent or meager between them. The sites of measured continental heat flow are even more irregularly distributed. Antarctica, most of the interiors of Africa and South America, and large expanses of Asia are either devoid of heat-flow data or are represented by only a few sites.

In recent years a global data set of heat-flow values has been assembled, representing 20,201 heat-flow sites. The data set is almost equally divided between observations on land (10,337 sites) and in the oceans (9,864 sites). Histograms of the heat-flow values are spread over a wide range for each domain (Fig. 4.29). The distributions have similar characteristics, extending from very low, almost zero values to more than 200 mW m$^{-2}$. The high values on the continents are from volcanic and tectonically active regions, while the highest values in the oceans are found near to the axes of oceanic ridges. Both on the continents (Fig. 4.25) and in the oceans (Fig. 4.27), heat flow varies with crustal age. To determine global heat-flow statistics, the fraction of the Earth’s surface area having a given age is multiplied by the mean heat flow measured for that age domain. The weighted sum gives a mean heat flow of 65 mW m$^{-2}$ for the continental data set. The oceanic data must be corrected for hydrothermal circulation in young crust; the areally weighted mean heat flow is then 101 mW m$^{-2}$ for the oceanic data set. The oceans cover 60.6% and the continents 39.4% of the Earth’s surface, the latter figure including 9.1% for the continental shelves and other submerged continental crust. The weighted global mean heat flow is 87 mW m$^{-2}$. The heat-flow values in both continental and oceanic domains are found to depend on crustal age and geological characteristics. These relationships make it possible to create a map of global heat flow that allows for the uneven distribution of actual measurements (Fig. 4.30). The procedure in creating this map was as follows. First, the Earth’s surface was divided into 21 geological domains, of which 12 are in the oceans and 9 on the continents. Next, relationships between heat flow and age were used to associate a representative heat flow with each domain (Table 4.7). This made it possible to estimate heat flow for regions that have no measurement sites. Allowance was also made for the loss of heat by hydrothermal circulation near to ridge systems. The surface of the globe was then divided into a grid of 1° elements (i.e., each element measures 1° × 1°), and the mean heat flow through each element was estimated. This gave a complete data set (partly observed and partly synthesized) covering the entire globe. The gridded data were fitted by spherical harmonic functions (as in the representation of the geoid, Section 2.4.5) up to degree and
order 12. The results of the analysis were used to compute smooth contours of equal heat flow, which were then plotted as the global heat-flow map (Fig. 4.30). If the global mean value is subtracted, the Earth’s surface can be divided into regions with above-average and below-average heat flow, respectively (Fig. 4.31). The regions with above-average heat flow are notably associated with the oceanic ridge systems. About half of the Earth’s heat is lost by the
cooling of oceanic lithosphere of Cenozoic age (younger than 65 Ma).

One must keep in mind that this global model is based on a mixture of actual heat-flow measurements in regions where they are available, and estimated values in inaccessible regions. Moreover, the measured data near ocean ridges are replaced with values predicted by cooling models to compensate for the known loss of heat by hydrothermal circulation. Nevertheless, these global heat-flow maps are the best available representations of the geographical pattern and flux of the heat flowing out of the Earth’s interior. Although details may eventually need modification, the main features are not in doubt.

4.2.8.3 Models for the cooling of oceanic lithosphere

The variations of heat flow and ocean depth with time constrain the possible thermal models for cooling of the lithosphere in different ways. The predicted heat flow is contingent on the temperature gradient in a model, but the oceanic depth is defined by the vertical distribution of density, which, in turn, depends on the volume coefficient of expansion and the temperature profile in the plate. Thus, oceanic bathymetry depends on the temperature integrated over depth.

Several cooling models have been proposed, all of which satisfy the decrease in heat flow and increase in ocean depth with age. The simplest model represents the cooling lithosphere as a semi-infinite half-space (Fig. 4.32a). Initially, the temperature inside the half-space is uniform and higher than on its upper surface, which is maintained at the temperature of cold ocean-bottom seawater. As long as the lithosphere is thin – which it is near the ridge – horizontal heat conduction can be neglected. The heat flow in the uniform half-space is vertical, along the z-axis, and is equivalent to the one-dimensional flow in a thin vertical column (Fig. 4.33a). The spreading process can be envisioned as transporting the column away from the ridge axis, during which conductive cooling takes place and the temperature distribution in the column changes.

This model allows us to compute the temperature distribution in the oceanic lithosphere. We need to compute the depth z at which a given temperature T is reached after time t, when the vertical column has moved at velocity v to a distance vt from the ridge. First, the desired temperature T is expressed as a fraction of the mantle temperature $T_m$. Using Eq. (4.57) and the appropriate table, the argument $\eta_T$ is found which gives an error function equal to $(T/T_m)$. Setting the numerical value $\eta_T$ equal
predicts that the lithosphere becomes thicker with age. The thermal model is an isotherm (Fig. 4.33b). The model shear waves are attenuated, the base of the lithosphere in being defined mechanically as the depth where seismic thermal model of the lithosphere (Fig. 4.32b). Instead of a chosen fraction of the observed heat flow (Fig. 4.27) and the ocean depth from each spreading center it gives very good fits to both. For ages greater than about 70 Ma the ocean depths in particular are less than predicted by the $\sqrt{t}$ relationship (Fig. 4.28). This suggests that the source of heat from below the lithosphere may be shallower than in the half-space model at large ages. As an alternative to the half-space models the oceanic lithosphere has been modelled as a flat layer or plate of finite thickness, bounded above by cold sea-water and with a constant temperature on its lower surface. Far from a ridge axis this model brings hot mantle temperatures nearer to the surface than in the half-space model. Below the ridge axis the vertical edge of the new plate has the same high temperature of its lower surface (Fig. 4.32c), which results in heat being conducted horizontally through the plate. This is not a serious problem as long as the plate is much thinner than its horizontal extent away from a ridge. This condition is clearly met for the main lithospheric plates, which are several thousand kilometers across and only of the order of a hundred kilometers thick.

The half-space and boundary-layer models fit the observed variations of heat flow and ocean depth with age for young lithosphere. For ages greater than about 70 Ma the ocean depths in particular are less than predicted by the $\sqrt{t}$ relationship (Fig. 4.28). This suggests that the source of heat from below the lithosphere may be shallower than in the half-space model at large ages. As an alternative to the half-space models the oceanic lithosphere has been modelled as a flat layer or plate of finite thickness, bounded above by cold sea-water and with a constant temperature on its lower surface. Far from a ridge axis this model brings hot mantle temperatures nearer to the surface than in the half-space model. Below the ridge axis the vertical edge of the new plate has the same high temperature of its lower surface (Fig. 4.32c), which results in heat being conducted horizontally through the plate. This is not a serious problem as long as the plate is much thinner than its horizontal extent away from a ridge. This condition is clearly met for the main lithospheric plates, which are several thousand kilometers across and only of the order of a hundred kilometers thick.

The plate model is not intended to model the vertical mechanical structure of the plate, but only to explain in a phenomenological way the typical age dependence of both ocean depth and heat flow. The plate thickness in the model is the asymptotic thermal thickness of old oceanic lithosphere and reflects the combined effects of temperature and rheology. Its horizontal isothermal base requires additional deep heat sources that prevent the lithosphere from cooling as a half-space at great ages. The model allows simple computation of the thermal cooling history. The best known version, proposed by Parsons and Sclater (1977), assumed a plate thickness of 125 km and a basal temperature of 1350 °C. At large distances from each spreading center it gives very good fits to both the observed heat flow (Fig. 4.27) and the ocean depth (Fig. 4.28). The most recent update, the GDH1 plate model, has a thickness of 95 km and a basal temperature of 1450 °C. It fits the observations even better.

The isotherms in the cooling lithosphere have a parabolic shape with respect to the time (or horizontal distance) axis, of which only the part for $z > 0$ is of interest. The surface heat flow for this model is given by Eq. (4.60), and so is inversely proportional to $\sqrt{t}$.

$$z = \frac{z}{2\sqrt{kt}}$$

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The half-space model has some unrealistic aspects. It predicts infinitely large heat flow at the ridge axis, and the initial mantle temperature $T_m$ is approached asymptotically and is only reached at infinite depth. The distances between successive isotherms for equal increments in temperature get progressively larger. The near-surface layer in which the temperature changes are significant has been called a thermal boundary layer. Its base is defined arbitrarily as the depth at which the temperature reaches a chosen fraction of $T_m$. The layer can be regarded as a thermal model of the lithosphere (Fig. 4.32b). Instead of being defined mechanically as the depth where seismic shear waves are attenuated, the base of the lithosphere in the thermal model is an isotherm (Fig. 4.33b). The model predicts that the lithosphere becomes thicker with increasing age, as also inferred from seismic data, and the thickness is proportional to $\sqrt{t}$. As it cools and thickens, the lithosphere sinks deeper into the asthenosphere, so that the ocean depth increases away from a ridge axis. Together with Pratt-type, thermally influenced isostasy the half-space model of lithosphere cooling predicts a depth increase that is also proportional to $\sqrt{t}$ (Box 4.4).

Parker and Oldenburg (1973) proposed a modification of the boundary-layer model in which a solid lithosphere overlies a fluid asthenosphere. The base of the lithosphere is taken to be the solid–liquid phase boundary of the material. It is defined by the melting-point isotherm, and denotes a phase change. This is probably a closer representation of the real situation, although, by treating the asthenosphere as a fluid, it exaggerates the change in rheology. The temperature of the asthenosphere lies close to the solidus temperature, but its condition is only partially molten (perhaps about 5%).

The half-space and boundary-layer models fit the observed variations of heat flow and ocean depth with age for young lithosphere. For ages greater than about 70 Ma the ocean depths in particular are less than predicted by the $\sqrt{t}$ relationship (Fig. 4.28). This suggests that the source of heat from below the lithosphere may be shallower than in the half-space model at large ages. As an alternative to the half-space models the oceanic lithosphere has been modelled as a flat layer or plate of finite thickness, bounded above by cold sea-water and with a constant temperature on its lower surface. Far from a ridge axis this model brings hot mantle temperatures nearer to the surface than in the half-space model. Below the ridge axis the vertical edge of the new plate has the same high temperature of its lower surface (Fig. 4.32c), which results in heat being conducted horizontally through the plate. This is not a serious problem as long as the plate is much thinner than its horizontal extent away from a ridge. This condition is clearly met for the main lithospheric plates, which are several thousand kilometers across and only of the order of a hundred kilometers thick.

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Box 4.4: Variation of ocean depth with age

As oceanic lithosphere moves away from the ridge it cools, thickens, and becomes denser (Fig. 4.31). It sinks progressively into the underlying asthenosphere with time, so that the depth of an oceanic basin increases with age $t$ of the oceanic lithosphere. A simple model accounts for the depth change $w$ by assuming that the lithosphere and asthenosphere are in Pratt-type isostatic balance. The isostatic model is sometimes referred to as thermal isostasy.

Compare the composition of two vertical columns of unit cross-sectional area above a compensation level in the asthenosphere at depth $D$ (Fig. B4.4). The column below R on the ridge axis consists of hot asthenosphere, of assumed constant density $\rho_a$ and temperature $T_a$, and the depth $d_r$ of sea-water (density $\rho_w$) above the ridge. The column below B over the adjacent ocean basin is a section through oceanic lithosphere. The density $\rho_L$ of the lithosphere depends on its temperature $T_L$ and the lithosphere thickness $L$ increases with age $t$. The sea-water layer of depth $d_s$, above the ridge is present in both columns, as is the thickness $A$ of asthenosphere between the base of the lithosphere and the compensation depth. The isostatic balance is determined by equating the weights of $w$ km of seawater and $L$ km of lithosphere with the corresponding weight of $(w+L)$ km of asthenosphere:

$$ (w+L)\rho_w g = w\rho_w g + g \int_0^{L} \rho_L \, dz \tag{1} $$

$$ w(\rho_a - \rho_w) = \int_0^{L} (\rho_L - \rho_a) \, dz \tag{2} $$

Thermal isostasy assumes that the lithosphere changes density as it cools. The volume coefficient of expansion $\alpha$ is defined by Eq. (4.26) as

$$ \alpha = \frac{1}{V} \frac{dV}{dT} = -\frac{1}{\rho} \frac{d\rho}{dT} \tag{3} $$

where density $\rho = M/V$ and thus $dV/V = -d\rho/\rho$. Rewriting Eq. (3), we get

$$ \alpha \rho_a (T_a - T_L) = (\rho_L - \rho_a) \tag{4} $$

The expression for the density difference between the lithosphere and asthenosphere is now substituted into Eq. (2):

$$ w = \frac{\alpha \rho_a}{(\rho_a - \rho_w)} \int_0^{L} (T_a - T_L) \, dz \tag{5} $$

The temperature of the lithosphere $T_L$ is given by Eq. (4.57) with $T_a$ instead of $T_m$. Substituting in Eq. (5), we find

$$ w = \frac{\alpha \rho_a}{(\rho_a - \rho_w)} \int_0^{L} (T_a - T_L) \, dz \tag{6} $$

$$ dz = \frac{\alpha \rho_L T_a}{(\rho_a - \rho_w) 2\sqrt{\pi}t} \int_0^{\infty} \text{erfc}(\eta) \, d\eta \tag{7} $$

From Box 4.3, Eqs. (8) and (9), we have $\int_0^{\infty} \text{erfc}(\eta) \, d\eta = 1/\sqrt{\pi}$. Thus,

$$ w = \frac{\alpha \rho_L T_a}{\sqrt{\pi} (\rho_a - \rho_w)} \sqrt{\pi}t \tag{8} $$

This is the amount by which the ocean deepens away from the ridge axis. The total depth of the ocean, taking into account the depth $d_r$ at the ridge, is $d = (d_r + w)$. Optimum values for the parameters of the lithosphere in Eq. (8) are given in the Global Depth and Heat Flow model (GDH1) of Stein and Stein (1992): $\alpha = 3.1 \times 10^{-5}$ K$^{-1}$, $\rho_a = 3300$ kg m$^{-3}$, $\kappa = (k/\rho_a) = 8.04 \times 10^{-7}$ m$^2$ s$^{-1}$. Assuming a mean depth of 2600 m over the ridge axis, a temperature $T_a = 1450$ C for the asthenosphere, and $\rho_w = 1030$ kg m$^{-3}$ for the density of sea-water, the depth of the ocean over crust of age $t$ in Ma) is given by

$$ d = (d_r + w) = 2600 + 370 \sqrt{t} \tag{9} $$

This computed result is close to the depth–age relationship in Eq. (4.63) predicted by the GDH1 model for young lithosphere (age $< 20$ Ma).
4.2.8.4 Structure of oceanic lithosphere

The plate models explain observed thermal data better than the boundary-layer models. The boundary-layer model is most appropriate near to a ridge axis, and agrees better with other geophysical data, which show that the lithosphere thickens with distance from a ridge. However, the plate model is needed at great distances to explain heat flux and ocean depths over old lithosphere. To reconcile these contrasting attributes a two-layered model of the lithosphere has been proposed (Fig. 4.34). The upper layer is rigid and has a mechanically defined lower boundary, above which heat transfer is by conduction. Below this level the increasing temperature causes a change in mechanical properties. The lower lithosphere is plastic enough to permit material movement, and so behaves like a viscous solid. The base of the upper layer is an isotherm, representing the temperature at which rigidity is lost. The base of the lower lithosphere is a thermally defined boundary, and is also an isotherm. Several suggestions have been made as to how this structure may approximate the plate model for old lithosphere. They include additional heat sources such as radiogenic heating, frictional heating as a result of shear at the base of the lithosphere, and reheating of old lithosphere due to the intrusion of mantle plumes at hotspots. It has also been postulated that, at lithosphere ages greater than about 70 Ma, small-scale convection currents in the lower lithosphere may augment thermal conduction. This would bolster the transfer of heat from the convection asthenosphere into the lithosphere, effectively giving a thinner lithosphere than in the half-space models. Analysis of the dispersion of seismic surface waves indicates that there are differences in structure between continents and oceans down to about 200 km. This is compatible with the thermal model of a rigid mechanical layer underlain by a convecting thermal boundary layer extending to about 150–200 km.

4.2.8.5 Heat flow at subduction zones

The oceanic lithosphere is bent sharply downward beneath the overriding plate in a subduction zone. It extends as an inclined slab deep into the upper mantle, which it penetrates at a rate of a few centimeters per year. The old lithosphere is cold, having lost much of its original heat of formation at the ridge axis. By the time it reaches an ocean trench the isotherms in the plate are far apart and the temperature gradient is small. The separation of the isotherms is increased by the downward bending of the plate. Since the heat flow is proportional to the temperature gradient, very low heat-flow values ($\approx 35$ mW m$^{-2}$) are measured in oceanic trenches. After bending downward the plate is subducted to great depths, subjecting it to increases in pressure and temperature. Heat is conducted into the plate from the adjacent mantle. This process is so slow that the interior of the subducting slab remains colder than its environment (Fig. 4.35). A temperature of 800 °C is normally reached at about 70 km depth in the oceanic plate but in the descending slab this temperature exists to deeper than 500 km. Above this depth the coldest part of the slab has a horizontal temperature deficit of 800–1000 K.

Heat conducted from the mantle is not the only heat source that must be taken into account in modelling the thermal structure of the subducting slab. An important additional source is the frictional heating that results from shear deformation at the surfaces of the slab where it is in contact with the mantle. In the upper part of a subduction zone the shear heating melts the basaltic layer of the oceanic lithosphere and forms a layer of eclogite in the top of the slab. The high density of the eclogite causes a positive gravity anomaly (see Fig. 2.62), and adds to the forces propelling the slab downward. The phase transition in which the open structure of olivine-type minerals converts to a denser spinel-type structure normally takes place at a depth of 400 km. The phase transition depends on temperature and pressure. Laboratory experiments indicate that it takes place at lower pressure at low temperature than at high temperature. Consequently it occurs at shallower depths within the cold plate than in the adjacent mantle. As a result the transition depth is deflected upward by about 100 km. The transition is exothermic and the latent heat given out in the transition is an additional heat source that contributes to the thermal structure of the subduction
zone. The transition also results in a density increase, which adds to the forces driving the plate downward.

The deeper transition at 670 km is less well understood. High temperature apparently causes it to take place at higher pressure, and so the depth of occurrence is deflected downward inside the subducting slab. It is uncertain whether the transition is endothermic, absorbing heat from the environment, or exothermic as assumed in the model in Fig. 4.35. An endothermic phase change has the effect of reducing the density, and acts against the other downward forces on the slab.

Although other, slightly different models have been derived for the temperature distribution in the descending slab, they all have in common the downward deflection of isotherms in the cold descending slab. The heat flow can be computed for a given thermal model. When compared with the observed heat flow on a profile across the subduction zone, the models fail to explain adequately the high heat flow observed on the overriding plate (Fig. 4.35). Volcanic activity is partly responsible, fed by magmas produced by partial melting of oceanic crustal material in the descending slab and of the upper mantle in the overriding plate. Shallow melting is promoted by water from the subducting plate and generates basaltic magma; deeper melting involves less water and results in andesitic magma. When the overriding plate is continental, volcanic chains form along the continental margin parallel to the deep oceanic trench. The volcanicity is typified by the eruption of both basaltic and andesitic lavas. The lavas are more felsic than those formed when two oceanic plates collide, which may imply that they include melted material from the upper mantle of the overriding continental plate.

When two oceanic plates converge, a volcanic arc is formed on the overriding plate. Behind the arc, high heat flow on the overriding plate is related to back-arc spreading, in which new oceanic crust is generated by the intrusion of basaltic magma from partial melting in the upper mantle. This form of sea-floor spreading produces a marginal basin behind the island arc. The intrusion of magma is not confined to a single location, as at a ridge axis, but is spread diffusely in the basin. Consequently, the stripes of linearbed oceanic magnetic anomalies characteristic of sea-floor spreading at ridge systems are missing or at best weakly defined in a marginal basin.

4.2.9 Mantle convection

It has gradually become accepted that thermally driven convection takes place in the mantle and that it is probably the most important mechanism in geodynamic processes. There are several reasons for these conclusions. The evidence summarized in Section 2.8 demonstrates that the mantle has a viscoelastic rheology. The passage of seismic compressional and shear waves through the mantle attests that it reacts as a solid to abrupt stress changes. Yet, observations of post-glacial isostatic uplift and long-term movements of the rotation axis indicate that the mantle is capable of viscous flow when stressed over long time intervals. The surmised temperature distribution in the mantle implies that, although conduction is mainly responsible for heat transfer in the lithosphere, convection is the predominant process deeper in the mantle, involving mass transfer by sub-solidus creep. Applying the theory of thermal convection to the mantle and using the best available estimates of physical parameters indicates that robust convection must be taking place.

4.2.9.1 Thermal convection

The conditions for convection to occur (see Section 4.2.4.2) reflect a balance between causal forces due to thermal expansion and resistive effects due to viscosity and thermal diffusivity. When a fluid is heated, thermal expansion gives rise to an upward buoyancy force. This produces instability, which is partly counteracted by diffusion of heat into the surrounding fluid by thermal conduction. As soon as a volume of the fluid starts to rise in response to the buoyancy force its motion is resisted by viscous forces. The effects are familiar to anyone who has heated a pan of thick soup or porridge. If the pan is heated too rapidly, or the instructions to “stir constantly” are ignored, the soup may stick to the bottom of the pan and become charred. This happens because the viscosity of the fluid is initially too large to allow convection. Despite the large temperature gradient between the hot bottom of the pan and the cool surface of the liquid, conduction is unable to
Thermal diffusivity and viscosity act as stabilizing influences in a heated fluid. If heating is slow enough, the temperature gradient adjusts to transfer the heat by conduction, remaining close to the adiabatic gradient. Convection becomes possible when the real temperature gradient exceeds the adiabatic gradient; the difference \( \theta \) is called the superadiabatic gradient. The excess heat expands the fluid, causing the buoyancy force. When this becomes larger than the viscous resistance, convection ensues. The ratio of the competing forces is embodied in the Rayleigh number (Eq. (4.42)). The Rayleigh number \( (Ra) \) for convection due to the superadiabatic temperature gradient in a fluid layer of thickness \( D \) is

\[
Ra = \frac{g \alpha \theta}{\kappa \nu} D^4
\]  

(4.66)

where \( g \) is gravity and \( \alpha \) is the coefficient of thermal expansion.

The superadiabatic gradient is not the only source of power for convection. Although radioactive heat generation in mantle materials is small (see Section 4.2.5.1), it can still contribute to convection. If \( Q \) is the radiogenic heat production in a layer of thickness \( D \), we can invoke Eq. (4.39) and Eq. (4.45) and replace \( \theta \) in the above equation by \((QD/k)\), where \( k \) is the thermal conductivity. This allows us to define a second Rayleigh number \( (Ra_Q) \) for convection driven by radiogenic heat:

\[
Ra_Q = \frac{g \alpha Q}{\kappa \nu} D^5
\]  

(4.67)

Convection is initiated when the Rayleigh number exceeds a critical value, \( Ra_c \), which is dependent on the geometry of the flow and the boundary conditions on the upper and lower surfaces. In Rayleigh–Bénard convection the top and bottom of the horizontal layer are stress free; the critical Rayleigh number is \( Ra_c = 658 \). If the top and bottom of the layer are rigid boundaries at which the horizontal velocity vanishes, \( Ra_c = 1708 \).

Table 4.8 shows computed Rayleigh numbers \( Ra_T \) for convection driven by the superadiabatic temperature gradient for viscous flow in the upper, lower and whole mantle, assuming representative values from the literature for the parameters in Eq. (4.66). Reasonable estimates of the radiogenic heat produced in the mantle give even larger values for \( Ra_Q \).

4.2.9.2 Convection at high Rayleigh numbers

The computed Rayleigh numbers greatly exceed the critical values for convection throughout the entire mantle or in separate layers. Thus, each region of the sub-lithospheric mantle is capable of convection. The Rayleigh number for whole-mantle convection is so much larger than the critical value \( Ra_c \) that vigorous mantle convection must be expected. This does not imply rapid flow in normal terms. The speed of flow in the mantle is usually assumed to be of the same order as the rate of motion of tectonic plates, about 5–10 cm yr\(^{-1} \) on average. As long as the flow rate \( \nu \) is
4.2 THE EARTH’S HEAT

Table 4.8 Some physical parameters for mantle convection models (mostly from Jarvis and Peltier, 1989)

The critical Rayleigh numbers (Ra) for the onset of convection in each part of the mantle are calculated assuming a superadiabatic temperature gradient $\theta = 0.1 \text{ K km}^{-1}$ and a mean gravity $g = 10 \text{ m s}^{-2}$. Lower mantle parameters are interpolated from the upper- and whole-mantle values.

<table>
<thead>
<tr>
<th>Physical parameter</th>
<th>Units</th>
<th>Upper mantle (70–670 km)</th>
<th>Lower mantle (670–2890 km)</th>
<th>Whole mantle (70–2890 km)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Layer thickness ($H$)</td>
<td>km</td>
<td>600</td>
<td>2220</td>
<td>2820</td>
</tr>
<tr>
<td>Expansion coefficient ($\alpha$)</td>
<td>K$^{-1}$</td>
<td>$2 \times 10^{-5}$</td>
<td>$1.0 \times 10^{-5}$</td>
<td>$1.4 \times 10^{-5}$</td>
</tr>
<tr>
<td>Density ($\rho$)</td>
<td>kg m$^{-3}$</td>
<td>3700</td>
<td>5500</td>
<td>4700</td>
</tr>
<tr>
<td>Specific heat ($c_p$)</td>
<td>J kg$^{-1}$ K$^{-1}$</td>
<td>1260</td>
<td>1260</td>
<td>1260</td>
</tr>
<tr>
<td>Thermal conductivity ($k$)</td>
<td>W m$^{-1}$ K$^{-1}$</td>
<td>6.7</td>
<td>20</td>
<td>15</td>
</tr>
<tr>
<td>Thermal diffusivity ($\alpha$)</td>
<td>m$^2$ s$^{-1}$</td>
<td>$1.4 \times 10^{-6}$</td>
<td>$3 \times 10^{-6}$</td>
<td>$2.5 \times 10^{-6}$</td>
</tr>
<tr>
<td>Dynamic viscosity ($\eta$)</td>
<td>kg m$^{-1}$ s$^{-1}$</td>
<td>$1 \times 10^{11}$</td>
<td>$2.5 \times 10^{10}$</td>
<td>$2 \times 10^{11}$</td>
</tr>
<tr>
<td>Kinematic viscosity ($\nu$)</td>
<td>m$^2$ s$^{-1}$</td>
<td>$2.7 \times 10^{27}$</td>
<td>$4.5 \times 10^{17}$</td>
<td>$4.3 \times 10^{17}$</td>
</tr>
<tr>
<td>Rayleigh number ($Ra_T$)</td>
<td>—</td>
<td>7000</td>
<td>180,000</td>
<td>820,000</td>
</tr>
</tbody>
</table>

low, adjacent lamina of the fluid move past each other under the conditions for Newtonian viscosity (Section 2.8.2). At faster flow rates this condition breaks down, and the flow becomes turbulent. The conditions favoring turbulence are high momentum ($\rho v$) and large scale $D$ of the flow, whereas it is inhibited by high viscosity $\eta$. These factors are contained in the Reynolds number, $Re$, defined as

$$Re = \frac{\rho v D}{\eta} \quad (4.68)$$

Reasonable values for the mantle are $\rho = 5000 \text{ kg m}^{-3}$, $D = 2900 \text{ km} = 2.9 \times 10^6 \text{ m}$, $v = 5 \text{ cm yr}^{-1} = 1.5 \times 10^{-9} \text{ m s}^{-1}$, and $\eta = 1.5 \times 10^{21} \text{ Pa s}$. The Reynolds number is found to be $Re = 1.5 \times 10^{20}$, which is so small that turbulence is negligible. Similar results are found by considering the upper or lower mantle alone. Clearly, although mantle convection involves high Rayleigh numbers (implying vigorous convection on a geological timescale), it takes place by laminar flow.

The effect of convection is to replace conduction as the principal mechanism of heat transfer. A measure of the relative effectiveness of the two processes of heat transfer is the Nusselt number, $Nu$. This is defined as the ratio of the heat transport in the presence of convection to the heat transport without convection. In the absence of radiogenic heat sources, the heat transport with convection is determined by the Rayleigh number $Ra_T$, while the non-convective heat transport is expressed by the critical Rayleigh number $Ra_c$. The Nusselt number depends on the ratio of these two numbers and can be written

$$Nu = \beta \left(\frac{Ra_T}{Ra_c}\right)^S \quad (4.69)$$

where the coefficient $\beta$ and the exponent $S$ are functions of the aspect ratio of the convection cells. Mathematical evaluation of the problem of Rayleigh–Bénard convection with stress-free upper and lower boundaries gives $\beta = 1$ and $S = 1/3$, and, since in this case $Ra_c = 10^3$, the Nusselt number has the simpler form

$$Nu = 0.1 (Ra_T)^{1/3} \quad (4.70)$$

Using the estimated values of $Ra_T$ in Table 4.8 gives Nusselt numbers of 19 for layered convection in the upper mantle and 97 for whole-mantle convection. Hence, heat transfer by convection is dominant in the mantle.

Once convection has been initiated the boundary conditions determine the shapes of the convection cells. In Rayleigh–Bénard convection the aspect ratio of a cell – the ratio of its horizontal dimension to its vertical one – is $2^{1/2} = 1.41$; when the layer has rigid boundaries the cell aspect ratio is 1.01. Hence, the horizontal extent of a convection cell is comparable with the layer thickness. This has implications for convection in the mantle. If we assume that the scale of mantle convection is represented by the pattern of the plate boundaries (Fig. 1.11), we can estimate the horizontal dimensions of the convection cells. It is evident that they must have very different sizes. The ridge-to-trench horizontal distances across major plates are in the range 2000–10,000 km, with an average of about 5000 km. This is larger than the maximum thickness of the convecting layer, whether we assume convection to be restricted to the upper mantle or to occupy the entire 2900 km thickness of the sub-lithospheric mantle. Thus, if convection is uniform through the whole mantle, the aspect ratios of at least some cells must be much larger than unity (Table 4.9).
reason for this is the rigidity of the cold upper boundary formed by the lithosphere, which inhibits the breakup of the fluid flow into cells with smaller horizontal extents.

4.2.9.3 Models of mantle convection

The feasibility of mantle convection is accepted but there is still some doubt as to the form it takes. This is in part due to uncertainty as to the role played by the seismic discontinuities at 400 km and 670 km depth, which bound the upper-mantle transition zone. The discontinuities are not sharp, and are understood to represent mineral phase changes rather than compositional differences (as, for example the crust–mantle and core–mantle boundaries). The upper discontinuity marks the olivine–spinel phase change, the lower one represents the phase change from spinel to perovskite structure (Section 3.7.5.2), with accompanying changes in density and elastic parameters. In principle, mass can be carried by convection currents across these discontinuities. The 670 km discontinuity is close to the maximum depth of seismicity in subduction zones, and may be where the subducting plate is absorbed into the mantle.

There are two main models of mantle convection, each with an interface at the 670 km seismic discontinuity. An important change in viscosity occurs at this level. In whole-mantle convection (Fig. 4.37a) the viscosity doubles from the upper mantle to the lower mantle (see Table 4.8) and there is a net flow of material across the boundary. In this model, convection ensures that the entire mantle is well mixed mechanically, and the phase changes at 400 and 670 km have only a small effect on the temperature gradient. This model agrees with much of the available evidence.

The alternative layered convection model has distinct convecting layers in the upper and lower mantle (Fig. 4.37b). There are two ways in which this can take place. The upper and lower convection patterns in a vertical section may represent circulations in the same sense (e.g., both clockwise or both counterclockwise) or in opposite senses (e.g., one clockwise and the other counterclockwise). In each case the radial velocity is zero at 670 km depth and there is no mass transfer across the discontinuity, the material in each flow pattern spreads out along the boundary. However, the models imply different types of coupling between the layers. Opposite senses of circulation in the layers would cause little or no shear between the tangential flows at the boundary, resulting in mechanical coupling between the layers. Cold material sinking in the upper mantle would overly hot material rising in the lower mantle. However, if the layered flow patterns have the same sense of circulation (as in Fig. 4.37b), hot material rising in the upper mantle overlies hot material rising in the lower mantle, so that the flow regimes are coupled thermally. This model has a strong velocity shear across the 670 km discontinuity, which requires a large and abrupt change in viscosity at this depth; viscosity in the lower mantle would need to be at least two orders of magnitude smaller than in the upper mantle. Estimates of mantle viscosity (Section 2.8.6) indicate the opposite; viscosity is higher in the lower mantle than in the upper mantle.

A model of layered convection assumes that there is no mass transfer across the discontinuity. The upper and lower mantles are well mixed individually, but the separation of the flow patterns at the discontinuity means that they may have distinct chemical compositions. Because
there is no convective flow across it, heat can only cross the boundary by conduction. The 670 km discontinuity therefore acts as a thermal boundary, with a large temperature change of perhaps 500–1000 K across it. Thus, the temperature profile in the lower mantle, although maintained adiabatic by the convection, would be 500–1000 K higher than in whole-mantle convection. This would result in a smaller temperature change across the core–mantle boundary, a less-steep temperature gradient in the D^* layer, and so a lower heat flux from the core. The long-term rate of cooling of the Earth would thereby be reduced.

The problem of understanding mantle convection is complicated by the non-uniform structure and rheology of the mantle. As yet, there is no complete picture of how the various factors that influence convection act together. The convection pattern depends strongly on what happens physically and thermodynamically at the 670 km discontinuity. This can only be inferred indirectly. Our understanding of the discontinuity is incomplete, but it is essential to resolving the real pattern of mantle convection.

### 4.2.9.4 Mantle plumes

The viscosity in the upper mantle is inferred from post-glacial rebound studies to be around $10^{21}$ Pa s, but lower-mantle viscosity is less well known. The sub-solidus creep in the mantle implies a temperature-dependent viscosity, which allows thermal boundary layers at the top and bottom of the mantle to influence the patterns of convective flow.

The lithosphere constitutes an upper, cold boundary layer. It accretes at high temperature at spreading ridges, where upwelling magma from the mantle reaches the surface. The eruptive lavas issue from magma chambers beneath ridge crests, in which magma from the deeper mantle undergoes differentiation. As part of the plate tectonic cycle the lithosphere rapidly cools and hardens as it spreads away from the ridge. Its high viscosity (i.e., rigidity) inhibits internal convection, but at subduction zones the plate (by now old) flexes downward and carries cold material into the underlying mantle, altering its thermal balance. Seismic tomography (Section 3.7.6) has revealed broad regions of raised seismic velocity in the deep mantle below subduction zones, giving rise to the surmise that the material in the cold subducted plate eventually sinks to the bottom of the mantle. The material must eventually take part in a broad-scale return flow, completing the convective cycle, but how this takes place is not clear.

The core–mantle boundary (CMB) at 2890 km depth constitutes a lower, hot boundary layer. The D^* layer at the base of the mantle (Section 3.7.5.3) is characterized by reductions in seismic velocities between about 2740 km and the CMB. It evidently has different physical properties than the mantle above it, and appears to play an important thermodynamic role. The heat flux from the core to the mantle diminishes the rigidity of the layer, thus reducing the seismic velocities. The viscosity of the hot thin D^* layer is presumed to be much lower than that of the overlying mantle. The topography of the CMB has been explored by seismic waves reflected from the core or passing it at grazing incidence. The thickness of the D^* layer appears to be uneven, and has been interpreted by analogy to the crust. Thick segments have been designated as crypto-continents and thinner regions as crypto-oceans (Fig. 4.38).

The low-viscosity material in the D^* layer is thought to supply relatively fast-flowing narrow mantle plumes. This name is given to vertical features, thin in cross-section, that facilitate the upwelling of low-viscosity hot magma through the more viscous mantle. A new plume melts its way to the surface behind a larger head. Some plumes may not reach the surface but intrude their material into the asthenosphere or lower lithosphere. Other mature plumes may penetrate the entire mantle and reach the surface, where they are evident as places of persistent volcanism, high regional topography and local high heat flow, called hotspots. These areas of anomalous volcanism are found in the oceans and on the continents, within plates and on plate margins. The plumes that feed them are thought to remain fixed in position for long periods of
time, and so the hotspots are anchored to the mantle below the lithosphere. As a result they have important consequences for studies of plate tectonic motions.

4.3 GEOELECTRICITY

4.3.1 Introduction

Electric charge – together with mass, length and time – is a fundamental property of nature. The name electric derives from the Greek word for amber (“elektron”), the naturally occurring fossilized resin of coniferous trees that has been used since antiquity in the making of jewelry. The Greek philosopher Thales of Miletus (ca. 600 BC) is credited with first reporting the power of amber, when rubbed with a cloth, to attract light objects. The ancient sages could not understand this behavior in terms of their everyday world, and so, together with the power of magnetism possessed by natural lodestone (see section 5.1.1), electricity remained a wonderful but unknown phenomenon for more than two millennia. In 1600 AD the English physician William Gilbert summarized previous investigations and extant knowledge in the first systematic study of these phenomena.

In the following century it was established that there were two types of electric charge, now referred to as positive and negative. Objects that carried like types of charge were observed to repel each other, and those that carried opposite types were attracted to each other. In 1752 the American statesman, diplomat and scientist Benjamin Franklin performed a celebrated experiment; by flying a kite during a thunderstorm, he established that lightning is an electrical phenomenon. Having survived this risky endeavor Franklin developed the far-sighted theory that electricity consisted of an omnipresent fluid, and that the different types of charge represented surplus and scarcity of this fluid. This view strikingly resembles modern theory, in which the “fluid” consists of electrons.

The laws of electrostatic attraction and repulsion were established in 1785 as a result of careful experiments by a French scientist, Charles Augustin de Coulomb (1736–1806), who also established the laws of magnetostatic force (Section 5.1.3). Coulomb invented a sensitive torsion balance, with which he could measure the magnetostatic force (Section 5.1.3). Coulomb established that the force of attraction or repulsion between two charged spheres is proportional to the product of the individual electric charges and inversely proportional to the square of the distance between the centers of the spheres. His law can be written as the following equation:

\[ F = k \frac{Q_1 Q_2}{r^2} \]

(4.71)

where \( Q_1 \) and \( Q_2 \) are the electric charges, \( r \) is their separation and \( k \) is a constant. This inverse-square law strongly resembles the law of universal gravitation (Eq. (2.2)), formulated by Newton more than a century before Coulomb's law. However, in gravitation the force is always attractive, whereas in electricity it may be attractive or repulsive, depending on the nature of the charges. In the law of gravitation the units of mass, distance and force are already defined, so that the gravitational constant is predetermined; only its numerical value needed to be measured. In Coulomb's law, \( F \) and \( r \) are defined from mechanics (as the newton and meter, respectively), but the units of \( Q \) and \( K \) are undefined. The value of \( K \) was originally set equal to unity, thereby defining the unit of electric charge. This definition led to unfortunate complications when the magnetic effects of electric currents were analyzed. The alternative is to define independently the unit of charge, thereby fixing the meaning of the constant \( K \).

The unit of charge is the coulomb (C), defined as the amount of charge that passes a point in an electrical circuit when an electric current of one ampère (A) flows for one second (i.e., 1 C = 1 A s). In turn, the ampère is defined from the magnetic effects of a current (see Section 5.2.4). When a current flows in the same direction through two parallel long straight conductors, magnetic fields are produced around the conductors, which cause them to attract each other. If the current flows through the conductors in opposite directions they repel each other. The ampère is defined as the current that produces a force of \( 2 \times 10^{-7} \) N per meter of length between infinitely long thin conductors that are one meter apart in vacuum. Thus, the unit of charge is defined precisely, if rather indirectly. In the Système Internationale (SI) units \( K \) is written as \( (4\pi \varepsilon_0)^{-1} \), so that Coulomb's law becomes

\[ F = \frac{1}{4\pi \varepsilon_0} \frac{Q_1 Q_2}{r^2} \]

(4.72)
where the constant \( \varepsilon_0 \) is called the *permittivity constant*. It is approximately equal to \( 8.854 \times 10^{-12} \text{ C}^2 \text{ N}^{-1} \text{ m}^{-2} \).

Modern electrical theory descends from the discovery in 1897 by the English physicist Joseph J. Thomson of the electron as the basic elementary unit of electric charge. It has a negative charge of \( 1.602 \times 10^{-19} \text{ C} \). A proton in the nucleus of an atom has an equal positive charge. Normally an atom contains as many electrons as it has protons in its nucleus and is electrically neutral. If an atom or molecule loses one or more electrons, it has a net positive charge and is called a *positive ion*; similarly, a negative ion is an atom or molecule with a surplus of electrons.

In metals, some electrons are only loosely bound to the atoms. They can move with relative ease through the material, which is called an electrical conductor. Metals like copper and silver are good conductors. In other materials, called *insulators*, the electrons are tightly bound to the atoms. Glass, rubber, and dry wood are typical insulators. A perfect insulator does not allow electrons to move through it, whereas a perfect conductor offers no opposition to the passage of electrons. Real conductors offer different degrees of opposition.

A flow of charge, or *electric current*, results when the free electrons in a conductor move in a common direction. A current of one ampère corresponds to a flow of about 6,250,000,000,000,000,000 electrons per second past any point of a circuit! The direction of an electric current is defined to be the direction of flow of positive charge, which is opposite to the direction of motion of the electrons.

### 4.3.2.1 Electric field and potential

The force exerted on a unit electric charge by another charge \( Q \) is called the *electric field* of the charge \( Q \). Thus, if we let \( Q_1 = Q \) and \( Q_2 = 1 \) in Eq. (4.72), we obtain the equation for the electric field \( E \) at distance \( r \) from a charge \( Q \)

\[
E = \frac{Q}{4\pi\varepsilon_0 r^2} \quad (4.73)
\]

According to this definition \( E \) has the dimensions of newton/coulomb (N C\(^{-1}\)).

The term “field” also has another connotation, introduced by Michael Faraday (1791–1867) to refer to the geometry of the lines of force near a charge. Around a positive point charge the field lines are directed radially outward, describing the (divergent) direction along which a free positive charge would move (Fig. 4.39a); around a negative point charge they are directed radially inward (convergent) (Fig. 4.39b). The field lines of a pair of opposite point charges diverge from the positive charge, spread apart and converge on the negative charge (Fig. 4.39c); they give the appearance of drawing the opposite charges together.

The combined field of two positive point charges is characterized by field lines that leave each charge and diverge in the space between (Fig. 4.39d); the field lines appear visibly to push the like charges apart. The direction of the electric field at any point is tangential to the electric field line. The strength of the field is represented by the spatial concentration of the field lines. Close to either electrical charge the field is strong and it weakens with increasing distance from the charge. Consequently, work is required to move a charged particle from one point in the field to another. This work contributes to the potential energy of the system.

For example, at an infinite distance from a positive charge \( Q \) the repulsive force on a unit positive charge is zero, but at a distance \( r \) it is given by Eq. (4.73). The potential energy of the unit charge at \( r \) is called the *electric potential* at \( r \); we will denote it \( U \). The units of \( U \) are energy per unit charge, i.e. joules/coulomb. If we move a distance \( dr \) against the field \( E \), the potential changes by an amount \( dU \) equal to the work done against \( E \), which is \( -Edr \). i.e., \( dU = -Edr \), so that

\[
E = -\frac{dU}{dr} \quad (4.74)
\]

We can readily compute the electric potential \( U \) at \( r \) by integration:
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\[ U = - \int E \, dr = - \int_{r_0}^{r_f} \frac{Q}{4\pi \epsilon_0} \, dr \]  
(4.75)

from which

\[ U = \frac{Q}{4\pi \epsilon_0} \]  
(4.76)

The energy needed to move a unit charge from one point to another in the electric field of \( Q \) is the potential difference between the two points. The unit of potential difference is the same as that of \( U \) (i.e., joules/coulomb) and is called a volt. From Eq. (4.74) we obtain the more common alternative units of volt/meter (V m\(^{-1}\)) for the electric field \( E \).

Electric charge flows from a point with higher potential to a point with lower potential. The situation is analogous to the flow of water through a pipe from one level to a lower level. The rate of flow of water through the pipe is determined by the difference in gravitational potential between the two levels. Likewise, the electric current in a circuit depends on the potential difference in the circuit.

4.3.2.2 Ohm’s law

The German scientist Georg Simon Ohm established in 1827 that the electric current \( I \) in a conducting wire is proportional to the potential difference \( V \) across it. The linear relationship is expressed by the equation

\[ V = IR \]  
(4.77)

where \( R \) is the resistance of the conductor. The unit of resistance is the ohm (\( \Omega \)). The inverse of resistance is called the conductance of a circuit; its unit is the reciprocal ohm (\( \Omega^{-1} \)), variously also called a mho or siemens (\( S \)).

Experimental observations on different wires of the same material showed that a long wire has a larger resistance than a short wire, and a thin wire has a larger resistance than a thick wire. Formulated more precisely, for a given material the resistance is proportional to the length \( L \) and inversely proportional to the cross-sectional area \( A \) of the conductor (Fig. 4.40). These relationships are expressed in the equation

\[ R = \frac{\rho L}{A} \]  
(4.78)

The proportionality constant \( \rho \) is the resistivity of the conductor. It is a physical property of the material of the conductor, which expresses its ability to oppose a flow of charge. The inverse of \( \rho \) is called the conductivity of the material, denoted \( \sigma \). The unit of resistivity is the ohm-meter (\( \Omega \text{m} \)); the unit of conductivity is the reciprocal ohm-meter (\( \Omega^{-1} \text{m}^{-1} \)).

If we substitute Eq. (4.78) for \( R \) in Eq. (4.77) and rearrange the terms we get the following expression:

\[ \frac{V}{L} = \frac{\rho L}{A} \]  
(4.79)

4.3.2.3 Types of electrical conduction

Electric current passes through a material by one of three different modes: by electronic, dielectric, or electrolytic conduction. Electronic (or ohmic) conduction occurs in metals and crystals, dielectric conduction in insulators, and electrolytic conduction in liquids.

Electronic conduction is typical of a metal. The free electrons in a metal have a high average speed (about \( 1.6 \times 10^6 \text{ m s}^{-1} \) in copper). They collide with the atoms of the metal, which occupy fixed lattice sites, and bounce off in random directions. When an electric field is applied, the electrons acquire a common drift velocity, which is superposed on their random motions, so that they move at a much smaller speed (about \( 4 \times 10^{-5} \text{ m s}^{-1} \) in copper) in the direction of the field. The resistivity is determined by the mean free time between collisions. If the atomic arrangement causes frequent collisions, the resistivity is high, whereas a large mean free time between collisions results in low resistivity. The energy lost in the collisions appears in the form of heat.

A form of semiconduction is important in some crystals, such as the silicate minerals. The resistivity of the mineral is higher than that of a conductor but lower than that of an insulator, and it is called a semiconductor. Different types of semiconduction are possible. Silicates contain fewer conduction electrons than a metal, but the electrons are not rigidly bound to atoms as in an insulator. The energy needed to liberate additional electrons from...
their atoms is not large, and thermal excitation is enough to allow them to take part in electronic semiconduction. The liberated electron leaves a vacancy or hole in the valence level of the atomic structure, which behaves as a positive charge. Natural crystals also contain impurity atoms, which may have a different valency than that required by the lattice for charge balance. The impurity is a source of holes or excess electrons which take part in impurity semiconduction. At high temperature ions may detach from the lattice; they behave like ions in an electrolyte and give rise to electrical currents by ionic semiconduction. A potential difference across a semiconductor produces an electric current made up of opposite flows of negative electrons and positive holes. If most of the current is carried by the negative electrons, the semiconductor is called n-type; if the positive holes predominate, the semiconductor is said to be p-type.

Dielectric conduction occurs in insulators, which contain no free electrons. Normally, the electrons are distributed symmetrically about a nucleus. However, an electric field displaces the electrons in the direction opposite to that of the field, while the heavy nucleus shifts slightly in the direction of the field. The atom or ion acquires an electric polarization and acts like an electric dipole. The net effect is to change the permittivity of the material from $\varepsilon_0$ to a different value $\varepsilon$, given by

$$\varepsilon = \varepsilon_0$$

(4.81)

Here, $\varepsilon_0$ is called the relative permittivity. When it is measured in a constant electric field, the relative permittivity is called the dielectric constant, $\kappa$, of the material. It is dimensionless, and has a value commonly in the range 3–80. Examples of $\kappa$ for some natural materials are: air 1.00059; mica 3; glass 5; sandstone 5–12; granite 3–19; diorite 6; basalt 12; water 80. Dielectric effects are unimportant in constant current situations. However, in an alternating electric field the polarization changes with the frequency of the field, and thus the relative permittivity is frequency dependent. The fluctuating polarization of the electric charge contributes to the alternating current, and so modifies the effective conductivity or resistivity. In practice, this effect depends strongly on the frequency of the inducing alternating field. The higher the frequency, the greater is the effect of dielectric conduction. Some geoelectric methods utilize signals in the audio-frequency range, where dielectric conduction is insignificant, but ground-penetrating radar uses frequencies in the MHz to GHz range and depends on dielectric contrasts.

Electrolytic conduction occurs in aqueous solutions that contain free ions. The water molecule is polar (i.e., it has a permanent electric dipole moment) with a strong electric field which breaks down molecules of dissolved salts into positively and negatively charged ions. For example, in a saline solution the molecule of sodium chloride (NaCl) dissociates into separate Na$^+$ and Cl$^-$ ions. The solution is called an electrolyte. The ions in the electrolyte are mobilized by an electric field, which causes a current to flow.

Electric charge is transported by positive ions in the direction of the field and negative ions in the opposite direction. The resistivity of an electrolyte may be understood by analogy with the flow of water through a partially blocked pipe. The electric current in the electrolyte involves the physical transport of material (ions), which results in collisions with the molecules of the medium (electrolyte), causing resistance to the flow. Ionic conduction is consequently slower than electronic conduction.

### 4.3.3 Electrical properties of the Earth

In our daily lives we experience frequent reminders of the Earth’s gravity field. It is less obvious that the Earth also has an electric field. Its presence mainly becomes evident during thunderstorms, when electrical discharges take place as lightning. The Earth’s electric field acts radially inward, so that the Earth behaves like a negatively charged sphere. At its surface the vertically downward electric field amounts to about 200 $V\,m^{-1}$. The atmosphere has a net positive charge, equal and opposite to that of the Earth, and resulting from the distribution of positively and negatively ionized air molecules. The charges originate from the continual bombardment of the Earth by cosmic rays.

Cosmic rays are subatomic particles with very high energy. Primary cosmic rays reach the Earth from outer space, travelling at velocities close to that of light. They consist largely of protons (hydrogen nuclei) and $\alpha$-particles (helium nuclei), with lesser amounts of other ions. Their origin is still unknown. Some are emitted by the Sun at the time of solar flares, but these occur too infrequently to be the main source. This source lies elsewhere in our galaxy. It is thought that a large proportion of the galactic cosmic rays are accelerated to high speed by supernova explosions. The path of a cosmic ray is easily deflected by a magnetic field. Even the weak interstellar magnetic field is enough to disperse fast-moving cosmic rays, so that they reach the Earth equally from all directions. The incoming particles collide with nuclei in the upper atmosphere, producing showers of secondary cosmic rays, consisting of protons, neutrons, electrons and other elementary particles. Consequently, at any given time a fraction of the molecules of the atmosphere are electrically charged. The Earth’s electric field accelerates positive particles downward to the Earth’s surface, where they neutralize negative surface charges. This would rapidly eliminate the negative surface charge, which is maintained by thunderstorm activity.

Thunderstorms, and the causes of lightning, are not yet fully understood. A possible scenario is the following. In a storm-cloud, droplets of water vapor become electrically charged. The Earth’s downward electric field may cause polarization within a droplet, with positive charge on the bottom and negative on the top. When the drop is heavy enough to fall, negative charges from molecules pushed out of its path are attracted to the bottom while fewer positive charges gather on the top. As a result the droplet becomes negatively charged. The wind action in storm-clouds causes
the negative charge to accumulate at the base of the cloud, while a corresponding positive charge gathers in its upper extent. When the potential difference between the two charges exceeds the break-down voltage of the atmosphere, a brief but powerful electric current flows. Most lightning strokes occur within the storm-cloud. However, the negative charge on the base of the cloud repels the negative charge on the ground surface beneath it. Once again, if the potential difference between the cloud and the ground becomes large enough to overcome the break-down voltage of the air, a lightning stroke ensues. This carries negative charge to the ground. In this way, the numerous daily lightning storms that occur worldwide maintain the negative charge of the Earth.

To a first approximation the Earth may be regarded as a uniform electrical conductor. Electric charges on the surface of a conductor disperse so that the electric potential is the same at all points on the surface, i.e., it is an electrical equipotential surface. The surface potential is commonly used as the reference level for electrical potential energy and is defined to be zero. Thus a positively charged body has a positive potential difference (voltage) with respect to ground, while a negatively charged body has a negative voltage.

### 4.3.3.1 Electrical surveying

As with other physical parameters, geoelectrical properties are utilized in both applied and general geophysics. They are exploited commercially in the search for valuable orebodies, which may be located by their anomalous electrical conductivities. Deep electrical sounding provides valuable information about the internal structure of the Earth’s crust and mantle. Electrical surveys may be based on natural sources of potential and current. More commonly, they involve the detection of signals induced in subsurface conducting bodies by electric and magnetic fields generated above ground. Investigations in this category include resistivity and electromagnetic methods. These techniques have long been used in commercial geophysical surveying. In recent years they have also become important in the scientific investigation of environmental problems. The electrical differences require the measurement of potential differences in the ground between suitably implanted electrodes. The electromagnetic techniques detect subsurface conductivity anomalies remotely; they do not need contact with the ground. As well as being employed in surface surveys they are especially suited to airborne use.

The important physical properties of rocks for electrical surveying are the permittivity (for georadar) and the resistivity (or conductivity), on which several techniques are based. Anomalies arise, for example, when a good conductor (such as a mineralized dike or orebody) is present in rocks that have higher resistivities. The resistivity contrast between orebody and host rock is often large, because the resistivities of different rocks and minerals vary widely (Fig. 4.41). In metallic ores the resistivity can be very low, but igneous rocks that contain no water can have a very high resistivity. For example, in a high-grade pyrrhotite ore $\rho$ is of the order of $10^{-5} \Omega \text{m}$, while in dry marble it is around $10^5 \Omega \text{m}$. The range between these extremes spans 13 orders of magnitude. Moreover, the resistivity range of any given rock type is wide and overlaps with other rock types (Fig. 4.41).

The resistivity of rocks is strongly influenced by the presence of groundwater, which acts as an electrolyte. This is especially important in porous sediments and sedimentary rocks. The minerals that form the matrix of a rock are generally poorer conductors than groundwater, so the conductivity of a sediment increases with the amount of groundwater it contains. This depends on the fraction of the rock that consists of pore spaces (the porosity, $\phi$), and the fraction of this pore volume that is water filled (the water saturation, $S$). The conductivity of the rock is proportional to the conductivity of the groundwater, which is quite variable because it depends on the concentration and type of dissolved minerals and salts it contains. These observations are summarized in an empirical formula, called Archie’s law, for the resistivity $\rho$ of the rock

$$\rho = \frac{a}{\phi^m} S^w$$  \hspace{1cm} (4.82)

By definition $\phi$ and $S$ are fractions between 0 and 1, $\rho_w$ is the resistivity of the groundwater, and the parameters $a$, $m$ and $n$ are empirical constants that have to be determined for each case. Generally, $0.5 \leq a \leq 2.5$, $1.3 \leq m \leq 2.5$ and $n = 2$. 

![Fig. 4.41 Ranges of electrical resistivity for some common rocks, soils and ores (data source: Ward, 1990; augmented by data from Telford et al., 1990).](image-url)
4.3 GEOELECTRICITY

4.3.4 Natural potentials and currents

Electrical investigations of natural electrical properties are based on the measurement of the voltage between a pair of electrodes implanted in the ground. Natural differences in potential occur in relation to subsurface bodies that create their own electric fields. The bodies act like simple voltaic cells; their potential arises from electrochemical action. Natural currents (called telluric currents) flow in the crust and mantle of the Earth. They are induced electromagnetically by electric currents in the ionsphere (described in Section 5.4.3.2). In studying natural potentials and currents the scientist has no control over the source of the signal. This restricts the interpretation, which is mostly only qualitative. The natural methods are not as useful as controlled induction methods, such as resistivity and electromagnetic techniques, but they are inexpensive and fast.

4.3.4.1 Self-potential (spontaneous potential)

A potential that originates spontaneously in the ground is called a self-potential (or spontaneous potential). Some self-potentials are due to man-made disturbances of the environment, such as buried electrical cables, drainage pipes or waste disposal sites. They are important in the study of environmental problems. Other self-potentials are natural effects due to mechanical or electrochemical action. In every case the groundwater plays a key role by acting as an electrolyte.

Some self-potentials have a mechanical origin. When an electrolyte is forced to flow through a narrow pipe, a potential difference (voltage) may arise between the ends of the pipe. Its amplitude depends on the electrical resistivity and viscosity of the electrolyte, and on the pressure difference that causes the flow. The voltage is due to differences in the electrokinetic or streaming potential, which in turn is influenced by the interaction between the liquid and the surface of the solid (an effect called the zeta-potential). The voltage can be positive or negative and may amount to some hundreds of millivolts. This type of self-potential can be observed in conjunction with seepage of water from dams, or the flow of groundwater through different lithological units.

Most self-potentials have an electrochemical origin. For example, if the ionic concentration in an electrolyte varies with location, the ions tend to diffuse through the electrolyte so as to equalize the concentration. The diffusion is driven by an electric diffusion potential, which depends on the temperature as well as the difference in ionic concentration. When a metallic electrode is inserted in the ground, the metal reacts electrochemically with the electrolyte (i.e., groundwater), causing a contact potential. If two identical electrodes are inserted in the ground, variations in concentration of the electrolyte cause different electrochemical reactions at each electrode. A potential difference arises, called the Nernst potential. The combined diffusion and Nernst potentials are called the electrochemical self-potential. It is temperature sensitive and may be either positive or negative, amounting to at most a few tens of millivolts.

The self-potentials that originate by the above mechanisms are attracting increased attention in environmental and engineering situations. However, in the exploration for subsurface regions of mineralization they are often smaller than the potentials associated with orebodies and are classified accordingly as “background potentials.” The self-potential associated with an orebody is called its “mineralization potential.” Self-potential (SP) anomalies across orebodies are invariably negative, amounting usually to a few hundred millivolts. They are most commonly associated with sulfide ores, such as pyrite, pyrrhotite, and chalcopyrite, but also with graphite and some metallic oxides.

The origin of the mineralization type of self-potential is still obscure, despite decades of applied investigations. At one time it was thought that the effect arose from galvanic action. This occurs when dissimilar metal electrodes are placed in an electrolyte. Unequal contact potentials are formed between the metals and the electrolyte, giving rise to a potential difference between the electrodes. According to this model an orebody behaves like a simple voltaic cell, with groundwater acting as the electrolyte. It was believed that oxidation of the part of the orebody above the water table produced a potential difference between the upper and lower parts, causing a spontaneous electric polarization of the body. Oxidation involves the addition of electrons, so the top of the orebody becomes negatively charged, explaining the observed negative anomalies. Unfortunately, this simple model does not explain many of the observed features of self-potential anomalies and has proved to be untenable.

Another mechanism for self-potential depends on variations in oxidation (redox) potential with depth (Fig. 4.42). The ground above the water table is more accessible to oxygen than the submerged part, so moisture above the water table contains more oxidized ions than that below it. An electrochemical reaction takes place at the surface between the orebody and the host rock above the water table. It results in reduction of the oxidized ions in the
adjacent solution. An excess of negative ions appears above the water table. A simultaneous reaction between the submersed part of the orebody and the groundwater causes oxidation of the reduced ions present in the groundwater. This produces excess positive ions in the solution and liberates electrons at the surface of the orebody, which acts as a conductor connecting the two half-cells. Electrons flow from the deep part to the shallow part of the orebody. Outside the orebody, positive ions move from bottom to top along the electric field lines. The equipotential surfaces are normal to the field lines. The self-potential is measured where they intersect the ground surface (Fig. 4.42).

The redox model is inadequate for the same reason as the galvanic model; it fails to account for many of the observed features of self-potential anomalies. In particular, the association of self-potential models with the water table has been cast in doubt. Moreover, sulfide orebodies appear to persist for geological lengths of time, so that a mechanism involving permanent flow of charge appears unlikely. Self-potential is a feature of a stable system that is perturbed by making an electrical connection between the host rock and the sulfide conductor through the inserted electrodes and their connecting wire. The observed potential difference appears to be due to the difference in oxidation potential between the locations of the measurement electrodes, one inside and the other outside the zone of mineralization.

4.3.4.2 SP surveying

The equipment needed for an SP survey is very simple. It consists of a sensitive high-impedance digital voltmeter to measure the natural potential difference between two electrodes implanted in the ground. Simple metal stakes are inadequate as electrodes. Electrochemical reactions take place between the metal and moisture in the ground, causing the build-up of spurious charges on the electrodes, which can falsify or obscure the small natural self-potentials. To avoid or minimize this effect non-polarizable electrodes are used. Each electrode consists of a metal rod submersed in a saturated solution of its own salt; a common arrangement is a copper rod in copper sulfate solution. The combination is contained in a ceramic pot which allows the electrolyte to leak slowly through its porous walls, thereby making electrical contact with the ground.

Two field methods are in common use (Fig. 4.43). The gradient method employs a fixed separation between the electrodes, of the order of 10 m. The potential difference is measured between the electrodes, then the pair is moved forward along the survey line until the trailing electrode occupies the location previously occupied by the leading electrode. The total potential at a measurement station relative to a starting point outside the study area is found by summing the incremental potential differences. Some electrode polarization is unavoidable, even with non-polarizable electrodes. This gives rise to a small error in each measurement; these add up to a cumulative error in the total potential. The polarization effects can sometimes be reduced by interchanging the leading and trailing electrodes. In this “leapfrog” technique the leading electrode for one measurement is kept in place and becomes the trailing electrode for the next measurement; meanwhile the previous trailing electrode is moved ahead to become the leading electrode. Cumulative error is the most serious disadvantage of the fixed electrode configuration. A practical advantage of the technique is that only a short length of connecting wire must be moved along with the electrodes.

The total field method utilizes a fixed electrode at a base station outside the area of exploration and a mobile measuring electrode. With this method the total potential is measured directly at each station. The wire connecting the electrodes has to be long enough to allow good coverage of the area of interest. This necessitates a long wire that must be wound or unwound on a reel for each measurement station. However, the total field method results in smaller cumulative error than the gradient method. It allows more flexibility in placing the mobile electrode and usually gives data of better quality. Hence, the total field method is usually preferred except in difficult terrain.

The surveying procedure with each technique consists of measuring potential at discrete stations along a profile. As in gravity and magnetic surveys, the data are mapped (Fig. 4.44) and interpretations of anomalies are based on their geometry. Methods used to interpret self-potential anomalies are often qualitative or are based on simple geometric models. Visual inspection of mapped anomalies
may reveal trends related to elongation of the orebody; crowding of contour lines can indicate its orientation. Profiles plotted in known directions across the anomaly can be compared with curves generated from simple models of the source. For example, a polarized sphere may be used to model the source of approximately circular anomalies, while a horizontal line source (or polarized cylinder) may be used to model an elongate anomaly. A common and effective method is to model SP anomalies with point sources; complex anomalies are modelled with combinations of sources and sinks.

4.3.4.3 Telluric currents

Ultraviolet radiation from the Sun ionizes molecules of air in the thin upper atmosphere of the Earth. The ions accumulate in several layers, forming the ionosphere (see Section 5.4.3.2) at altitudes between about 80 km and 1500 km above the Earth’s surface. Electric currents in the ionosphere arise from systematic motions of the ions, which are affected by various factors such as the daily and monthly tides, seasonal variations in insolation and the periodic fluctuation in ionization related to the 11-yr sunspot cycle. The currents produce varying magnetic fields with the same frequencies, which are observed at the surface of the Earth and can be analyzed from long-term continuous records of the geomagnetic field. The ionospheric effects show up in the energy spectrum of the geomagnetic field as distinct peaks representing periods that range from fractions of a second (geomagnetic pulsations) to several years (Fig. 4.45). The magnetic fields induce fluctuating electric currents, called telluric currents, that flow in horizontal layers in the crust and mantle. The current pattern consists of several huge whorls, thousands of kilometers across, which remain fixed with respect to the Sun and thus move around the Earth as it rotates.

The distribution of telluric current density depends on the variation of resistivity in the horizontal conducting layers. At shallow crustal depths the lines of current flow are disturbed by subsurface structures which cause contrasts in resistivity. These could arise from geological structures or the presence of mineralized zones. Consider, for example, a buried anticline which has a highly resistive rock (such as granite) as its core and is overlain by a conducting layer of porous sedimentary rocks saturated with groundwater. The horizontal flow of telluric current across the anticline chooses the less-resistive path through the conducting sediments. The current lines bunch together over the axis of the anticline, increasing the horizontal current density (Fig. 4.46). The equipotential surfaces normal to the current lines intersect the ground surface, where potential differences can be measured with a high-impedance voltmeter.

The field equipment for measuring telluric current density is simple. The sensors are a pair of non-polarizable electrodes with a fixed separation $L$ of the order of 10–100 m. The potential difference $V$ between the electrodes is measured with a high-impedance voltmeter. The
electric field $E$ at a point mid-way between the electrodes can be assumed to be $V/L$. Using Ohm’s law (Eq. (4.80)) and assuming that the telluric current flows in conducting rock layer with resistivity $\rho_1$, the telluric current density $J$ at each measurement station along a profile is given by

$$J = \frac{V}{\rho_1 L}$$  \hspace{1cm} (4.83)$$

The direction of the telluric current is not known, so two pairs of electrodes oriented perpendicular to each other are used. One pair is aligned north–south, the other east–west. Telluric currents vary unpredictably with time, but they change only slowly within a homogeneous region. To keep track of the temporal changes an orthogonal pair of electrodes is set up at a fixed base station outside the area to be explored. Another orthogonal pair is moved across the survey area. The potential difference between equipotential surfaces can be measured where they intersect the ground surface, using a second pair of electrodes. A simple direct current can cause charges to accumulate on the potential electrodes, which results in spurious signals. A common practice is to commutate the direct current so that its direction is reversed every few seconds; alternatively a low-frequency alternating current may be used. In multi-electrode investigations the current electrode-pair and potential electrode-pair are usually interchangeable.

### 4.3.5 Resistivity surveying

The large contrast in resistivity between orebodies and their host rocks (see Fig. 4.41) is exploited in electrical resistivity prospecting, especially for minerals that occur as good conductors. Representative examples are the sulfide ores of iron, copper and nickel. Electrical resistivity surveying is also an important geophysical technique in environmental applications. For example, due to the good electrical conductivity of groundwater the resistivity of a sedimentary rock is much lower when it is waterlogged than in the dry state.

Instead of relying on natural currents, two electrodes are used to supply a controlled electrical current to the ground. As in the telluric method, the lines of current flow adapt to the subsurface resistivity pattern so that the potential difference between equipotential surfaces can be measured where they intersect the ground surface, using a second pair of electrodes. A simple direct current can cause charges to accumulate on the potential electrodes, which results in spurious signals. A common practice is to commutate the direct current so that its direction is reversed every few seconds; alternatively a low-frequency alternating current may be used. In multi-electrode investigations the current electrode-pair and potential electrode-pair are usually interchangeable.

#### 4.3.5.1 Potential of a single electrode

Consider the flow of current around an electrode that introduces a current $I$ at the surface of a uniform half-space (Fig. 4.47a). The point of contact acts as a current source, from which the current disperses outward. The electric field lines are parallel to the current flow and normal to the equipotential surfaces, which are hemispherical in shape. The current density $J$ is equal to $I$ divided by the surface area, which is $2\pi r^2$ for a hemisphere of radius $r$. The electric field $E$ at distance $r$ from the input electrode is obtained from Ohm’s law (Eq. (4.80))

$$E = \rho J = \rho_1 \frac{I}{2\pi r^2}$$  \hspace{1cm} (4.84)$$

Putting this expression in Eq. (4.74) yields the electric potential $U$ at distance $r$ from the input electrode:

$$\frac{dU}{dr} = -\rho_1 \frac{I}{2\pi r^2}$$

$$U = \rho_1 \frac{I}{2\pi r}$$  \hspace{1cm} (4.85)$$

If the ground is a uniform half-space, the electric field lines around a source electrode, which supplies current to the ground, are directed radially outward (Fig. 4.47b). Around a sink electrode, where current flows out of the ground, the field lines are directed radially inward (Fig. 4.47c). The equipotential surfaces around a source or sink electrode are hemispheres, if we regard the electrode
in isolation. The potential around a source is positive and diminishes as 1/r with increasing distance. The sign of I is negative at a sink, where the current flows out of the ground. Thus, around a sink the potential is negative and increases (becomes less negative) as 1/r with increasing distance from the sink. We can use these observations to calculate the potential difference between a second pair of electrodes at known distances from the source and sink.

### 4.3.5.2 The general four-electrode method

Consider an arrangement consisting of a pair of current electrodes and a pair of potential electrodes (Fig. 4.48). The current electrodes A and B act as source and sink, respectively. At the detection electrode C the potential due to the source A is + \( \rho I (2\pi r) \), while the potential due to the sink B is − \( \rho I (2\pi r) \). The combined potential at C is

\[
U_C = \frac{\rho I}{2\pi} \left( \frac{1}{r_{AC}} - \frac{1}{r_{CB}} \right) \tag{4.86}
\]

Similarly, the resultant potential at D is

\[
U_D = \frac{\rho I}{2\pi} \left( \frac{1}{r_{AD}} - \frac{1}{r_{DB}} \right) \tag{4.87}
\]

The potential difference measured by a voltmeter connected between C and D is

\[
V = \frac{\rho I}{2\pi} \left( \frac{1}{r_{AC}} - \frac{1}{r_{CB}} \right) - \left( \frac{1}{r_{AD}} - \frac{1}{r_{DB}} \right) \tag{4.88}
\]

All quantities in this equation can be measured at the ground surface except the resistivity, which is given by

\[
\rho = 2\pi \frac{V}{I} \left( \frac{1}{r_{AC}} - \frac{1}{r_{CB}} - \frac{1}{r_{AD}} - \frac{1}{r_{DB}} \right)^{-1} \tag{4.89}
\]

### 4.3.5.3 Special electrode configurations

The general formula for the resistivity measured by a four-electrode method is simpler for some special geometries of the current and potential electrodes. The most commonly used configurations are the Wenner, Schlumberger and double-dipole arrangements. In each configuration the four electrodes are collinear but their geometries and spacings are different.

In the *Wenner configuration* (Fig. 4.49a) the current and potential electrode pairs have a common mid-point and the distances between adjacent electrodes are equal, so that \( r_{AC} = r_{DB} = a \), and \( r_{CB} = r_{AD} = 2a \). Inserting these values in Eq. (4.89) gives

\[
\rho = 2\pi \frac{V}{I} \left( \frac{1}{a} - \frac{1}{2a} \right)^{-1} \tag{4.90}
\]

\[
\rho = 2\pi a \frac{V}{I} \tag{4.91}
\]

In the *Schlumberger configuration* (Fig. 4.49b) the current and potential pairs of electrodes often also have a common mid-point, but the distances between adjacent electrodes differ. Let the separations of the current and potential electrodes be \( L \) and \( a \), respectively. Then \( r_{AC} = r_{DB} = (L - a)/2 \) and \( r_{CB} = r_{AD} = (L + a)/2 \). Substituting in the general formula, we get

\[
\rho = 2\pi \frac{V}{I} \left( \frac{2}{L - a} - \frac{2}{L + a} - \frac{2}{L + 2a} - \frac{2}{L - 2a} \right)^{-1}
\]

\[
= \frac{\pi V}{4I} \left( \frac{L^2 - a^2}{a} \right) \tag{4.92}
\]

In this configuration the separation of the current electrodes is kept much larger than that of the potential electrodes \((L \gg a)\). Under these conditions, Eq. (4.92) simplifies to
rocks with a strong resistivity contrast and good conductivity are suited to locating steeply dipping contacts between units of contrasting resistivity within an area at a particular depth. It is best to adapt the Wenner configuration to lateral profiling. The Wenner configuration is best adapted to determining depth and resistivity for flat-lying layered rock structures, such as sedimentary beds, or the depth to the water table. The Schlumberger configuration is most commonly used for vertical electrical sounding (VES) investigations. The mid-point of the array is kept fixed while the distance between the current electrodes is progressively increased. This causes the current lines to penetrate to ever greater depths, depending on the vertical distribution of conductivity.

**4.3.5.4 Current distribution**

The current pattern in a uniform half-space extends laterally on either side of the profile line. Viewed from above, the current lines bulge outward between source and sink with a geometry similar to that shown in Fig. 4.39c. In a vertical section the current lines resemble half of a dipole geometry. In three dimensions the current can be visualized as flowing through tubes that fatten as they leave the source and narrow as they converge towards the sink. Figure 4.50 shows the flow pattern of the current in a vertical section through the “tubes” in a uniform half-space.

In order to evaluate the depth penetration of current in a uniform half-space we define orthogonal Cartesian coordinates with the x-axis parallel to the profile and the z-axis vertical (Fig. 4.51a). Let the spacing of the current electrodes be \( L \) and the resistivity of the half-space be \( \rho \). The horizontal electric field \( E_y \) at \((x, y, z)\) is

\[
E_y = -\frac{\partial U}{\partial x} = -\frac{\partial}{\partial x} \left( \frac{\rho F}{2\pi r_1} \left( \frac{1}{r_1} - \frac{1}{r_2} \right) \right)
\]

where \( r_1 = (x^2 + y^2 + z^2)^{1/2} \) and \( r_2 = ((L - x)^2 + y^2 + z^2)^{1/2} \). Differentiating and using Ohm’s law (Eq. (4.80)) gives the horizontal current density \( J_y \) at \((x, y, z)\):
d/dx in the median vertical plane is \( \zeta \) above the depth \( H \). The fraction of current between any two depths is found from the difference in the fractions above each depth calculated with Eq. (4.101).

\[
J_x = \frac{I}{2\pi} \left( \frac{x}{r_1} + \frac{L-x}{r_2} \right)
\]

If \((x,y,z)\) is on the vertical plane mid-way between the current electrodes, \(x = L/2\), \(r_1 = r_2\) and the current density is given by

\[
J_x = \frac{IL}{2\pi} \left( \frac{1}{r_1^3} + \frac{1}{r_2^3} \right)
\]

The horizontal current \(dI_x\) across an element of area \((dydz)\) in the median vertical plane is \(dI_x = J_x dydz\). The fraction of the input current \(I\) that flows across the median plane above a depth \(z\) is obtained by integration:

\[
\frac{I_x}{I} = \frac{L}{2\pi} \int_0^z \frac{dz}{\left( (L/2)^2 + y^2 + z^2 \right)^{3/2}}
\]

\[
\frac{I_x}{I} = \frac{L}{2\pi} \int_0^z \frac{dz}{\left( (L/2)^2 + z^2 \right)^{1/2}}
\]

\[
\frac{I_x}{I} = \frac{2}{\pi} \tan^{-1} \frac{z}{L}
\]

Equation (4.101) shows that \(I_x\) depends upon the current-electrode spacing \(L\) (Fig. 4.51b). Half the current crosses the plane above a depth \(z = L/2\), and almost 90% passes above the depth \(z = 3L\). The fraction of current between any two depths is found from the difference in the fractions above each depth calculated with Eq. (4.101).

4.3.5.5 Apparent resistivity

In the idealized case of a perfectly uniform conducting half-space the current flow lines resemble a dipole pattern (Fig. 4.50), and the resistivity determined with a four-electrode configuration is the true resistivity of the half-space. But in real situations the resistivity is determined by different lithologies and geological structures and so may be very inhomogeneous. This complexity is not taken into account when measuring resistivity with a four-electrode method, which assumes that the ground is uniform. The result of such a measurement is the apparent resistivity of an equivalent uniform half-space and generally does not represent the true resistivity of any part of the ground.

Consider a horizontally layered structure in which a layer of thickness \(d\) and resistivity \(\rho_1\) overlies a conducting half-space with a lower resistivity \(\rho_2\) (Fig. 4.52). If the current electrodes are close together, so that \(L \ll d\), all or most of the current flows in the more resistive upper layer, so that the measured resistivity is close to the true value of the upper layer, \(\rho_1\). With increasing separation of the current electrodes the depth reached by the current lines increases. Proportionally more current flows in the less resistive layer, so the measured resistivity decreases. Conversely, if the upper layer is a better conductor than the lower layer, the apparent resistivity increases with...
increasing electrode spacing. When the electrode separation is much larger than the thickness of the upper layer \((L \gg d)\) the measured resistivity is close to the value \(\rho_1\) of the bottom layer. Between the extreme situations the apparent resistivity determined from the measured current and voltage is not related simply to the true resistivity of either layer.

### 4.3.5.6 Vertical electrical sounding

A two-layer situation is encountered often in electrical prospecting, for example when a conducting overburden overlies a resistive basement. It is also common in environmental applications, when the conducting water table lies under drier, more resistive soil or rocks. Before the advent of portable computers, two-layer cases were interpreted with the aid of characteristic curves. These theoretical curves, calculated for a particular four-electrode array, take into account the change in depth penetration when current lines cross the boundary to a layer with different resistivity. The electrical boundary conditions require continuity of the component of current density \(J\) normal to the interface and of the component of electric field \(E\) tangential to the interface. At a boundary the current lines behave like optical or seismic rays, and are guided by similar laws of reflection and refraction. For example, if \(\theta\) is the angle between a current line and the normal to the interface, the electrical “law of refraction” is

\[
\tan \theta_1 = \frac{\rho_2}{\tan \theta_2 / \rho_1} \quad (4.102)
\]

In a set of characteristic curves the apparent resistivity \(\rho_a\) is normalized by the resistivity \(\rho_1\) of the upper layer and the electrode spacing is expressed as a multiple of the layer thickness. The shape of the curve of apparent resistivity versus electrode spacing depends on the resistivity contrast between the two layers, and a family of characteristic curves is calculated for different ratios of \(\rho_2/\rho_1\) (Fig. 4.53). The resistivity contrast is conveniently expressed by a \(k\)-factor defined as

\[
k = \frac{\rho_2 - \rho_1}{\rho_2 + \rho_1} \quad (4.103)
\]

The \(k\)-factor ranges between \(-1\) and \(+1\) as the resistivity ratio \(\rho_2/\rho_1\) varies between 0 and \(\infty\). The characteristic curves, drawn as full logarithmic plots on a transparent overlay, are compared graphically with the field data to find the best-fitting characteristic curve. The comparison yields the resistivities \(\rho_1\) and \(\rho_2\) of the upper and lower layers, respectively, and the layer thickness, \(d\).

Although characteristic curves can also be computed for the interpretation of structures with more than two layers, modern VES analyses take advantage of the flexibility offered by small computers with graphic outputs on which the apparent resistivity curves can be assessed visually. The first step in the analysis consists of classifying the shape of the vertical sounding profile.

The apparent resistivity curve for a three-layer structure generally has one of four typical shapes, determined by the vertical sequence of resistivities in the layers (Fig. 4.54). The type \(K\) curve rises to a maximum then decreases, indicating that the intermediate layer has higher resistivity than the top and bottom layers. The type \(H\) curve shows the opposite effect; it falls to a minimum then increases again due to an intermediate layer that is a better conductor than the top and bottom layers. The type \(A\) curve may show some changes in gradient but the apparent resistivity generally increases continuously with increasing electrode separation, indicating that the true resistivities increase with depth from layer to layer. The type \(Q\) curve exhibits the opposite effect; it decreases continuously along with a progressive decrease of resistivity with depth.

Once the observed resistivity profile has been identified as of \(K\), \(H\), \(A\) or \(Q\) type, the next step is equivalent to one-dimensional inversion of the field data. The technique involves iterative procedures that would be very time-consuming without a fast computer. The method assumes the equations for the theoretical response of a multi-layered ground. Each layer is characterized by its thickness and resistivity, each of which must be determined. A first estimate of these parameters is made for each layer and the predicted curve of apparent resistivity versus electrode spacing is computed. The discrepancies between the...
The observed and theoretical curves are then determined point by point. The layer parameters used in the governing equations are next adjusted, and the calculation is repeated with the corrected values, giving a new predicted curve to compare with the field data. Using modern computers the procedure can be reiterated rapidly until the discrepancies are smaller than a predetermined value.

The inversion method is equivalent to matching automatically the observed and theoretical curves. A one-dimensional analysis accommodates only the variations of resistivity and layer thickness with depth. The response of a vertically layered structure has an analytical solution, so efficient inversion algorithms can be established. In recent years, procedures have been proposed that also take into account lateral heterogeneities. The response of two- or three-dimensional structures must be approximated by a numerical solution, based on the finite-difference or finite-element techniques. The number of unknown quantities increases, as do the computational difficulties of the inversion.

4.3.5.7 Induced polarization

If commutated direct current is used in a four-electrode resistivity survey, the sequence of positive and negative flow may be interspersed with periods when the current is off. The inducing current then has a box-like appearance (Fig. 4.55a). When the current is interrupted, the voltage across the potential electrodes does not drop immediately to zero. After an initial abrupt drop to a fraction of its steady-state value it decays slowly for several seconds (Fig. 4.55b). Conversely, when the current is switched on, the potential rises suddenly at first and then gradually approaches the steady-state value. The slow decay and growth of part of the signal are due to induced polarization, which results from two similar effects related to the rock structure: membrane polarization and electrode polarization.
Membrane polarization is a feature of electrolytic conduction. It arises from differences in the ability of ions in pore fluids to migrate through a porous rock. The minerals in a rock generally have a negative surface charge and thus attract positive ions in the pore fluid. They accumulate on the grain surfaces and extend into the adjacent pores, partially blocking them. When an external voltage is applied, positive ions can pass through the “cloud” of positive charge but negative ions accumulate, unless the pore size is large enough to allow them to bypass the blockage. The effect is like a membrane, which selectively allows the passage of one type of ion. It causes temporary accumulations of negative ions, giving a polarized ionic distribution in the rock. The effect is most pronounced in rocks that contain clay minerals; firstly, because the grain and pore sizes are small, and, secondly, because clay grains are relatively strongly charged and adsorb ions on their surfaces. The ionic build-up takes a short time after the voltage is switched on; when the current is switched off, the ions drift back to their original positions.

Electrode polarization is a similar effect that occurs when ore minerals are present. The metallic grains conduct charge by electronic conduction, while electrolytic conduction takes place around them. However, the flow of electrons through the metal is much faster than the flow of ions in the electrolyte, so opposite charges accumulate on facing surfaces of a metallic grain that blocks the path of ionic flow through the pore fluid. An overvoltage builds up for some time after the external current is switched on. The size of the effect is commensurate with the metallic concentration. After the current is switched off, the accumulated ions disperse and the overvoltage decays slowly.

The two effects responsible for induced polarization are indistinguishable at measurement level. The field method for an induced polarization (IP) survey is most often based on the double-dipole array. The current electrodes form a transmitter pair, while the potential electrodes form a receiver pair. The steady-state voltage \( V_0 \) is recorded and compared with the amplitude of the decaying residual voltage \( V(t) \) at time \( t \) after the current is interrupted (Fig. 4.55c). The ratio \( V(t)/V_0 \) is expressed as a percentage, which decays during the 0.1–10s between switching the current on and off. If the decay curve is sampled at many points, its shape and the area under the curve may be obtained (Fig. 4.55d). The area under the decay curve, expressed as a fraction of the steady-state voltage, is called the chargeability \( M \), defined as

\[
M = \frac{1}{V_0} \int_{0}^{\infty} V(t) \, dt \tag{4.104}
\]

\( M \) has the dimensions of time and is expressed in seconds or milliseconds. It is the most commonly used parameter in IP studies.

The induced polarization determines the length of the potential decay time. If it is shorter than the time when the inducing current is off, successive half-cycles of the potential will not interfere. However, if a disseminated conductor is present, the decay time increases, causing overlap and distortion of the half-cycles. The higher the signal frequency the more pronounced is the effect. It increases the ratio \( V(t)/V_0 \) giving the impression of a better conductor than is really present (i.e., the apparent resistivity decreases with increasing frequency). Clearly, IP and resistivity surveys with alternating current are also influenced. The frequency dependence of the IP effect is exploited by measuring apparent resistivity at two low frequencies. Let these be \( f \) and \( F (> f) \). Commonly \( f \approx 0.05–0.5 \) Hz and \( F \approx 1–10 \) Hz. Then \( \rho_i > \rho_F \) and we can define a frequency effect as

\[
FE = \frac{\rho_i - \rho_F}{\rho_F} \tag{4.105}
\]

The ratio FE is often multiplied by 100 to express it as a percentage (PFE). If no IP effect is present the resistivity will be the same at both frequencies. The larger the value of FE or PFE, the greater is the induced polarization in the ground. At frequencies above 10 Hz, mutual inductance effects between the cables of the primary and detection circuits can produce troublesome potentials, which must be avoided by the field procedure (such as restricting \( F \) or minimized analytically).

The presence of metallic conductors is expressed by a similar parameter to FE, the metallic factor (MF). This is proportional to the difference in conductivities at the two measurement frequencies.

\[
MF = A(\sigma_i - \sigma_F) = A(\rho_F^{-1} - \rho_i^{-1})
\]

\[
= A\left(\frac{\rho_i - \rho_F}{\rho_F}\right) \tag{4.106}
\]

The constant \( A \) is equal to \( 2\pi \times 10^8 \); the units of MF are those of conductivity (i.e., \( \Omega^{-1} \text{ m}^{-1} \) or \( \text{S m}^{-1} \)).

An IP survey includes both lateral profiling and vertical sounding with the expanding spread method. Using a double-dipole array the distance between nearest electrodes of the transmitter and receiver pairs is a multiple \( na \) of the electrode spacing \( a \) in each pair. Measurements are made at several discrete positions as the receiver pair is moved incrementally away from the fixed transmitter pair. The transmitter pair is then moved by one increment along the profile and the procedure is repeated. The value of \( \rho_o \) FE or MF obtained in each measurement is plotted below the mid-point of the array at the intersection of two lines inclined at 45° (Fig. 4.56a). Information is obtained from increasingly greater depths as the transmitter–receiver array expands (i.e., as \( n \) increases). The plotted value is, however, not the real value of the parameter at the indicated depth. (Recall, for example, that a measurement of apparent resistivity represents an equivalent half-space beneath the array.) A twodimensional picture of the variation of the IP parameter beneath the profile is synthesized by contouring the results (Fig. 4.56b). The plot is called a pseudo-section; it provides a convenient (though artificial)
4.3 GEOELECTRICITY

Resistivity anomalies depend on the presence of continuous conductors, such as groundwater or massive orebodies. If mineralization is disseminated through a rock it may not cause a significant resistivity anomaly. The good response of the IP method for disseminated concentrations of conducting ore minerals led to its development in base-metal exploration, where large low-grade orebodies may be commercially important. However, the IP effect also depends on the porosity and saturation of the rock. As a result, it can also be used in the search for groundwater and in other environmental applications.

4.3.5.8 Electrical resistivity tomography

The availability of fast, inexpensive computers and the development of efficient algorithms has led to the development of electrical tomographic methods akin to the technique of electrical tomography described in Section 3.7.6. Seismic tomography based upon teleseismic arrivals from earthquakes is used to describe regions deep in the Earth’s mantle that have anomalous seismic velocities. Seismic tomography using refracted and reflected signals from controlled sources can likewise be used to describe velocity perturbations due to shallow features in the Earth’s crust. In a similar way, electrical tomography is used to describe the resistivity structure of near-surface regions to depths of several tens of meters. In seismic tomography the observed travel-times are inverted to obtain the velocity structure along the path of a seismic ray. Analogously, in resistivity tomography, an inversion procedure is applied to the electrical potentials measured between electrode pairs to obtain the resistivity structure along the current flow lines.

The methods of direct-current resistivity and induced polarization are readily adapted to tomographic analysis. Instead of deploying a single pair of current-electrodes and a single pair of potential-electrodes, an array of regularly spaced electrodes is deployed. For two-dimensional surveys a linear arrangement of electrodes is used; for three-dimensional investigations the electrodes form an areal array. Various combinations of current-electrode pairs and potential-electrode pairs are analyzed. The inversion computation is both complex and computer intensive. It yields a two- or three-dimensional vertical cross-section of the true resistivities beneath the electrode array. As in standard resistivity methods, the resolution and maximum depth of investigation depend on the separation and geometry of the electrodes.

The application of direct-current resistivity tomography to an environmental problem is illustrated by an investigation of the extent of ice and permafrost in a buried Alpine rock glacier. The Murtel rock glacier on Mt. Corvatsch in Switzerland is a creeping, permanently frozen (permafrost) body. Its vertical structure is known from a drillhole through the body down to about 50 m depth below the surface. A resistivity tomographic survey employing more than 30 electrodes in Wenner configurations gave a vertical profile of resistivity in good agreement with the drillhole results, and described the lateral extent of the subsurface structure of the permafrost body in detail (Fig. 4.57). Ice has a much higher resistivity than sand or gravel. High resistivities of about 2 MΩ m, corresponding to massive ice, are found between 5 m and 15 m depths in the rock glacier. Above and below the ice body, the resistivities are lower. The surface layer resistivity of ~10 kΩ m is two orders of magnitude less than in the ice. The region below the ice body is interpreted to consist of frozen sand containing about 30% ice. Its resistivity is an order of magnitude lower than in the ice, but the lower boundary of the ice-block is not clearly defined. Resistivities are less than 5 kΩ m in front of the rock glacier, marking the sharp transition to permafrost-free material.
velocity produces a changing magnetic field, which in turn proportional to the speed of the charge. If the electric field moving in a conductor produces a magnetic field proportional to the current. Ampère’s law shows that an electric charge (or current) which classifies light as an electromagnetic phenomenon.

Unfortunately, Maxwell died while still in the prime of his career, before his theoretical predictions were verified. The German physicist Heinrich Hertz established the existence of electromagnetic waves experimentally in 1887, eight years after Maxwell’s untimely death. Maxwell’s equations are written as:

\[ \Delta^2 B = \mu_0 \mu_r \frac{\partial B}{\partial t} + \mu_0 \mu_r \varepsilon_0 \frac{\partial^2 B}{\partial t^2} \]  \hspace{1cm} (4.107)

\[ \Delta^2 E = \mu_0 \eta \frac{\partial E}{\partial t} + \mu_0 \eta \varepsilon_0 \frac{\partial^2 E}{\partial t^2} \]  \hspace{1cm} (4.108)

where, in Cartesian coordinates,

\[ \Delta^2 = \frac{\partial^2}{\partial x^2} + \frac{\partial^2}{\partial y^2} + \frac{\partial^2}{\partial z^2} \]

\[ \Delta^2 = \frac{\partial^2}{\partial x^2} + \frac{\partial^2}{\partial y^2} + \frac{\partial^2}{\partial z^2} \]

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4.3 GEOELECTRICITY

In these equations \( \sigma \) is the electrical conductivity, \( \mu_0 \) is the magnetic permeability, which in most materials (unless they are ferromagnetic) is very close to 1, \( \mu_0 \) is the permeability constant (Section 5.2.2.1), or permeability of free space \( (\mu_0 = 4\pi \times 10^{-7} \text{ N A}^{-2}) \), \( \epsilon_r \) is the relative permittivity of the material, and \( \epsilon_0 \) is the permittivity constant, or permittivity of free space \( (\epsilon_0 = 8.854187 \times 10^{-12} \text{ C}^2 \text{ N}^{-1} \text{ m}^{-2}) \). In constant electrical fields, \( \epsilon_r \) is known as the dielectric constant, \( \kappa \). The value of \( \kappa \) is 5–20 in most rocks and minerals and 80 in water (Section 4.3.2.3). In sediments and sedimentary rocks, the water content plays an important role in determining the value of \( \kappa \). The value of \( \epsilon_r \) increases with the frequency of the electrical field.

Electromagnetic radiation encompasses a wide frequency spectrum. It extends from very high-frequency (short-wavelength) \( \gamma \)-rays and x-rays to low-frequency (long-wavelength) radio signals (Fig. 4.59). Visible light constitutes a narrow part of the spectrum. Two ranges of electromagnetic radiation are of particular importance in solid Earth geophysics: a high-frequency range in the radar part of the spectrum, and a broad range of low frequencies extending from audio frequencies to signals with periods of hours, days or years. The electromagnetic equations reduce to simpler forms for these two particular frequency ranges.

The left side of Eq. (4.108) describes the variation of the E-component of the electromagnetic wave in space. The right side describes its variation with time and so its frequency dependence. The first term is related to the familiar conduction of electricity in a conductor. Maxwell introduced the second term and called it the displacement current. It originates when charges are displaced but not separated from their atoms, causing an electric polarization; fluctuations in the polarization have the effect of an alternating displacement current. Suppose that the electric dipole emitting \( E \) and \( B \) oscillates sinusoidally with angular frequency \( \omega \). Then \( \partial \mathbf{E}/\partial t = -\omega \mathbf{E} \) and \( \partial^2 \mathbf{E}/\partial t^2 = -\omega^2 \mathbf{E} \), so the magnitude ratio \( (MR) \) of the second (displacement) term to the first (conduction) term on the right side of Eq. (4.108) is

\[
MR = \frac{\mu_0 \epsilon_0 \mathbf{E} \partial^2 \mathbf{E}/\partial t^2}{\epsilon_0 \partial \mathbf{E}/\partial t} = \frac{\epsilon_0 \omega^2 \mathbf{E}^2}{\sigma \mathbf{E}} = \frac{2\pi f\rho_{ef}}{|r|} \tag{4.109}
\]

where \( f \) is the frequency of the signal. The conductivity \( \sigma \) of rocks and soils is generally in the range \( 10^{-5} \) to \( 10^{-1} \Omega^{-1} \text{ m}^{-1} \); in orebodies \( \sigma \) may be as large as \( 10^{4} \) \( \Omega^{-1} \text{ m}^{-1} \) (see Fig. 4.41). Electromagnetic induction surveying is usually carried out at frequencies below \( 10^4 \) Hz, for which the magnitude ratio \( MR \) is much less than unity in both good and bad conductors. At these frequencies the electromagnetic signal passes through the ground in a diffusive manner, by conversion of the changing magnetic fields to electric currents and vice versa. High-frequency surveying employs radar signals with frequencies around \( 10^8 \) Hz, for which the magnitude ratio \( MR \) is very small in an orebody but can be much greater than unity in rocks and soils. Under these conditions the electromagnetic signal propagates like a wave, and so is subject to diffraction, refraction and reflection.

4.3.6.1 Electromagnetic induction

Electromagnetic (EM) surveys carried out at frequencies below 50 kHz are based on the principle of electromagnetic induction. An alternating magnetic field in a coil or cable induces electric currents in a conductor. The conductivity of rocks and soils is too poor to permit significant induction currents, but when a good conductor is present a system of eddy-currents is set up. In turn, the eddy currents produce secondary magnetic fields that are superposed on the primary field and can be measured at the ground surface (Fig. 4.60a).

Suppose that a low-frequency plane wave propagates along the vertical \( z \)-axis. The displacement current is now
negligible compared to the conduction current and Eq. (4.107) becomes

$$\frac{\partial^2 B}{\partial z^2} = \mu_0 \sigma \frac{\partial B}{\partial t}$$  \hspace{1cm} (4.110)$$

where the magnetic field has components \( B_x \) and \( B_z \). The form of this equation is reminiscent of the one-dimensional equation of diffusion or heat conduction (Eq. (4.52)), whose solution (Eq. (4.54)) describes how the temperature changes with time and position when a fluctuating temperature acts on the surface. By analogy, the solution of Eq. (4.110) for the components \( B_x \) or \( B_z \) of an alternating magnetic field with angular frequency \( \omega \) (= \( 2\pi f \)) in a conductor with conductivity \( \sigma \) is

$$B_{x,z}(z,t) = B_0 e^{-z/d} \cos(\omega t - \frac{z}{d})$$  \hspace{1cm} (4.111)$$

where \( d = \sqrt{\frac{2}{\mu_0 \sigma}} \)  \hspace{1cm} (4.112)$$

Here, \( d \) is called the skin depth. At this depth the magnetic field is attenuated to \( e^{-1} \) (\( \approx 37\% \)) of its value outside the conductor. The skin depth is dependent on the conductivity of the body and the frequency of the field. The skin depth in normal ground (\( \sigma \approx 10^{-3} \ \Omega^{-1} \ m^{-1} \)) for a low-frequency alternating magnetic field (\( f \approx 10^3 \) Hz) is about 500 m but in an orebody (\( \sigma \approx 10^2 \ \Omega^{-1} \ m^{-1} \)) it is only \( \approx 16 \) cm. The comparable figures for a high-frequency radar signal (\( f \approx 10^9 \) Hz) are 50 cm and 0.16 mm, respectively. Note that the skin depth is not the maximum depth of penetration of the magnetic field. It helps to indicate how rapidly the field is attenuated, but the magnetic field is effective at depths that are many times the skin depth. However, it decays to \( \approx 1\% \) at a depth \( z = 5d \) and to \( \approx 0.1\% \) at \( z = 7d \), effectively limiting the practical depth of exploration with the induction method.

The many field methods of EM induction have a common principle. A coil or cable is used as transmitter of the primary alternating magnetic field, while another coil serves as receiver of both the primary signal and a secondary signal from the eddy currents induced in a conductor (Fig. 4.60a). The magnetic field in the conductor experiences a phase shift (equal to \( zd/\mu_0 \), Eq. (4.111)) due to the conductivity. This results in a phase difference \( \phi \) between the secondary and primary signals in the receiver (Fig. 4.60b). The exact theory of EM induction is complicated, even in a simple situation, but we can obtain a simple qualitative appreciation by applying some concepts from electrical circuit theory. As in Section 4.2.6.1 we will use complex numbers involving \( i = \sqrt{-1} \).

Let the current systems in transmitter, receiver and conductor be represented by simple loops carrying currents \( I_t \), \( I_s \), and \( I_c \), respectively. If the currents are sinusoidal, each has the form \( I = I_0 e^{i\omega t} \), so that \( dI/dt = i\omega I_0 \). Let the resistance of the conductor be \( R \) and its self-inductance be \( L \). The voltage \( V_c \) in the conductor is composed of two parts. A resistive part due to the current \( I_c \) in the resistance \( R \) is equal to \( I_c R \). An inductive part due to the change of current is equal to \( L dI_c/dt \). The complete voltage in the conductor is then

$$V_c = I_c R + L \frac{dI_c}{dt} = I_c (R + i\omega L)$$  \hspace{1cm} (4.113)$$

The voltage \( V_c \) is induced in the conductor by the changing current \( I_c \) in the transmitter circuit. Let the mutual inductance between transmitter and conductor be \( M_{cr} \); then \( V_c = -M_{cr} dI_c/dt \). Similarly, the transmitter current induces a primary voltage \( V_p = -M_{tr} dI_t/dt \) in the receiver, in which the eddy currents in the conductor also induce a secondary voltage \( V_s = -M_{sc} dI_t/dt \). Here \( M_{tr} \) and \( M_{sc} \) are the mutual inductances between transmitter and receiver, respectively. The following relationships exist between the different voltages and currents:

$$V_p = -M_{tr} \frac{dI_t}{dt} = -i\omega M_{tr} I_t$$  \hspace{1cm} (4.114)$$

$$V_s = -M_{sc} \frac{dI_t}{dt} = -i\omega M_{sc} I_c$$  \hspace{1cm} (4.115)$$

$$V_c = -M_{cr} \frac{dI_t}{dt} = -i\omega M_{cr} I_t$$  \hspace{1cm} (4.116)$$

Combining Eq. (4.113) and Eq. (4.116) we get

$$I_t = \frac{-i\omega M_{sc} I_c}{R + i\omega L} = \frac{-i\omega M_{sc}}{(R^2 + \omega^2 L^2)} (R - i\omega L)$$  \hspace{1cm} (4.117)$$

From Eq. (4.114) and Eq. (4.115) the ratio of \( V_s \) to \( V_p \) in the receiver is

$$V_p = M_{tr} I_t = \frac{-M_{sc} M_{cr} (\omega^2 L + i\omega R)}{M_{tr} (R^2 + \omega^2 L^2)}$$  \hspace{1cm} (4.118)$$
which can be written in the form

$$\frac{V_s}{V_p} = \frac{M_s I_s}{M_p I_p} = -\frac{M_o M_t}{M_p L} \left( \frac{\beta^2 + i\beta}{1 + \beta^2} \right)$$  \hspace{1cm} (4.119)$$

where $\beta = \omega L / R$ is the response parameter of the conductor. The function in parentheses in Eq. (4.119) is a complex number, so the voltage ratio (or response of the measuring system) can be written $P + iQ$. The real part $P$ has the same phase as the primary signal and is called the in-phase component of the response. The imaginary part $Q$ is $90^\circ$ out of phase with the primary signal (i.e., if the primary signal is $\sim \cos \omega t$, the imaginary part is $\sim \sin \omega t = \cos [\omega t - \pi/2]$); it is called the quadrature component.

### 4.3.6.2 EM induction surveying

The resemblance of the basic equations of EM induction to the diffusion equation (Eq. (4.53)) classifies the method as a diffusive one. Diffusive techniques – for example, the gravity, magnetic field, geothermal and seismic surface-wave methods – respond to a volumetric average of the specific physical parameter and do not show fine detail of its distribution. Hence, EM induction yields an average value of the electrical conductivity in a particular volume, but the resolution is better than that of potential field methods.

The EM induction method is very suitable for airborne surveys. These were carried out originally with fixed wing aircraft but they now more commonly use helicopters, which can adapt better to terrain roughness while flying close to the ground. The transmitter and receiver coils are mounted (usually with their axes coaxial or parallel to the line of flight) in fixed positions in the aircraft or in a towed “bird,” using configurations similar to those of airborne magnetometer surveys (see Fig. 5.44). Alternatively, the transmitter may be in the airplane and the receiver in the bird or in another airplane. The increased separation of transmitter and receiver in this configuration gives greater depth penetration. However, in flight the bird yaws and pitches, altering the separation and parallelism of the coils, so that normally only the quadrature component is usable. The flight patterns consist of parallel profiles traversing the terrain. Lines are flown at about 100 m above ground level with fixed-wing aircraft and 30 m with helicopters.

When a potential conductivity anomaly has been located from the air, it is usual to investigate it further with a ground-based EM induction method. The transmitter may be a long cable or large horizontal loop on the ground surface, or it may be a small coil (diameter $\sim 1$ m) with its axis vertical or horizontal. The receiver is usually a similar small coil. It can be used to detect the direction, intensity or phase of the secondary signal. In its most simple application the tilt of the coil about a horizontal axis measures the dip-angle of the combined primary and secondary fields at the receiver. The method allows location, outlining and, to some extent, depth determination of a conductor. However, the dip-angle method registers only part of the available information in the secondary signal and does not describe the electrical properties of the conductor. As shown by Eq. (4.119) these properties affect the phase and relative amplitudes of the in-phase and quadrature components of the secondary signal relative to the primary. Phase-component EM measurement methods, therefore, allow more detailed interpretation.

The methods are illustrated by the horizontal loop electromagnetic method (HLEM), popularly known also by the commercial names Slingram or Ronka. The receiver and transmitter are coupled by a fixed cable about 30 to 100 m in length, and kept at a constant separation while the pair is moved along a traverse of a suspected conductor (Fig. 4.61a). The cable supplies a direct signal that exactly cancels the primary signal at the receiver, leaving only the secondary field of the conductor. This is separated into in-phase and quadrature components, which are expressed as percentages of the primary field and plotted against the position of the mid-point of the pair of coils (Fig. 4.61b). The in-phase and quadrature signals are zero far from the conductor and at the places where either the transmitter or receiver passes over the conductor. This enables the outline of a buried conductor to be charted. The signal rises to a positive peak on either side and falls to a negative peak over the middle of the conductor. The peak-to-peak responses of the in-phase and quadrature components depend on the quality of the conductor, which is expressed...
4.3.6.3 Magnetotelluric sounding

Magnetotelluric (MT) sounding is a natural-source electromagnetic method. The fluctuating electromagnetic fields that originate in the ionosphere are partly reflected at the Earth’s surface; the returning fields are again reflected off the conducting ionosphere. This happens repeatedly, so that the fields eventually have a strong vertical component and may be regarded as vertically propagating plane waves with a wide spectrum of frequencies. These fields penetrate into the ground and induce telluric electric currents (Section 4.3.4.3), which in turn generate secondary magnetic fields. The telluric currents are detected with two pairs of electrodes, usually oriented north–south and east–west. Three components of the magnetic fields are measured: the vertical component and a horizontal component parallel to each of the telluric components. The method yields conductivity information from much greater depths than artificial-source induction methods. It has been applied in the search for petroleum and deep zones of mineralization in the upper crust. Utilizing long periods in the range 10–1000s it is an important method for the investigation of the structure of the crust and upper mantle.

Consider a plane electromagnetic wave propagating in the z-direction (Fig. 4.58). Let the electric component $E_y$ be along the x-axis, so that the magnetic field $B_y$ (being normal to $E_y$) is along the y-axis. Ampère’s law, as summarized in Maxwell’s equations, relates $E_y$ to the gradient of $B_y$ in the z-direction. Because $B$ has only a y-component, Ampère’s law simplifies to

$$E_y = -\frac{1}{\mu_0 \sigma} \frac{\partial B_y}{\partial z}$$  \hspace{1cm} (4.120)

where $B_y$ has the form of Eq. (4.111). Differentiating $B_y$ by parts gives

$$E_y = -\frac{B_0}{\mu_0 \sigma d} \left[ -e^{-z/d} \cos \left( \omega t - \frac{z}{d} \right) ight]$$

$$+ e^{-z/d} \left( -1 \right) \left[ - \sin \left( \omega t - \frac{z}{d} \right) \right]$$

$$= \frac{B_0}{\mu_0 \sigma d} e^{-z/d} \cos \left( \omega t - \frac{z}{d} \right)$$

$$+ \frac{B_0}{\mu_0 \sigma d} e^{-z/d} \sin \left( \omega t - \frac{z}{d} \right)$$

$$= \frac{B_0}{\mu_0 \sigma d} e^{-z/d} \sqrt{2} \cos \left( \omega t - \frac{z}{d} + \frac{\pi}{4} \right)$$  \hspace{1cm} (4.121)

Comparison of Eq. (4.111) and Eq. (4.121) shows a phase shift of $45^\circ (\pi/4)$ between $E_y$ and $B_y$. However, the ratio of the maximum amplitudes of the two components is

$$\frac{|E_y|}{|B_y|} = \frac{\sqrt{2}}{\mu_0 \sigma d}$$  \hspace{1cm} (4.122)

If we now substitute for $d$ from Eq. (4.112) and write $\rho = 1/\sigma$, we get

$$\rho = \frac{\mu_0 |E_y|^2}{|B_y|^2}$$  \hspace{1cm} (4.123)

$$d = \frac{|E_y|}{\rho |B_y|}$$  \hspace{1cm} (4.124)

for the effective resistivity $\rho$ at depth $d$. The analysis gives similar results for an electric field along the y-axis and a corresponding magnetic field along the x-axis. In this case the ratio of the field amplitudes is $|E_y|/|B_y|$.

In addition to the horizontal magnetic fields, $B_x$ and $B_y$, the vertical component $B_z$ is also recorded for use in the interpretation of two-dimensional structures. Thus the data set from an MT site consists of two electrical components and three magnetic field components recorded continuously during a lengthy observation interval covering some hours or days. The recorded magnetic fields consist of an external part from the ionosphere and an internal part related to the induced current distribution. These components must be separated analytically. The electric and magnetic records contain numerous frequencies, some of which are simply noise and some are of geophysical interest. As a result, sophis-
4.3 GEOELECTRICITY

Fig. 4.62 Two-dimensional resistivity model of the crust and upper mantle beneath Vancouver Island and the adjacent mainland derived from magnetotelluric results (redrawn from Kurtz et al., 1986).

At high frequencies in poorly conducting media the conduction term in the electromagnetic equations is negligible compared to the displacement term. The electric field equation then becomes

$$\nabla^2 E = \mu_0 \frac{\partial^2 E}{\partial t^2}$$

with a similar equation for the magnetic field. This has the familiar form of the wave equation, which describes the propagation of an elastic disturbance (Eqs. (3.55) and (3.56)). Analogously, Eq. (4.125) describes the propagation of the electric part of an electromagnetic wave. By comparing with the seismic wave equations we see that the E and B fields in an electromagnetic wave have the same velocity $v$, where $v^2 = 1/\mu_0 \varepsilon_0$. In a vacuum the wave velocity is equal to the velocity of light $c$, given by $c^2 = 1/\mu_0 \varepsilon_0$. Using the relationship $v^2 = c^2/\kappa$, and taking into account that the dielectric constant $\kappa$ is $\sim 5-20$ in Earth materials, the velocity of an electromagnetic wave in the ground is found to be about 0.2$c$–0.6$c$.

To simplify further discussion suppose that the electromagnetic disturbance propagates along the $z$-axis (i.e., $\partial/\partial x = \partial/\partial y = 0$), so that $\nabla^2 = \partial^2/\partial z^2$. For this one-dimensional case

$$\frac{\partial^2 E}{\partial z^2} = \mu_0 \varepsilon_0 \frac{\partial^2 E}{\partial t^2} = \frac{1}{v^2} \frac{\partial^2 E}{\partial t^2}$$

If we compare this equation with Eq. (3.58) for a seismic wave, we see that the solution for a component $E_z$ of the electric field is

$$E_z = E_0 \sin 2\pi (\frac{z}{\lambda} - ft)$$

where $\lambda$ is the wavelength, $f$ the frequency and $f\lambda = v$, the velocity of the wave.
The above considerations suggest that high-frequency electromagnetic waves travel in the ground in an analogous manner to seismic waves. Instead of being determined by the elastic parameters the propagation of radar signals is dependent on the dielectric properties of the ground. A comparatively young branch of geophysical exploration has been developed to investigate underground structures with ground-penetrating radar (GPR, or georadar). GPR makes use of the familiar “echo-principle” used in reflection seismology. A very short radar pulse, lasting only several nanoseconds (i.e., $\approx 10^{-8}$ s) is emitted by a mobile antenna on the ground surface. The path of the radar signal through the ground can be traced as a ray, which experiences refractions, reflections and diffractions at boundaries where the dielectric constant changes. A second antenna, the receiver, is located close to the transmitter, as in the case of seismic reflection, so as to receive near-vertical reflections from underground discontinuities. The signal-processing techniques of reflection seismology can also be applied to the georadar signal to help minimize the effects of diffractions and other noise. Consequently, georadar provides a detailed picture of the shallow subsurface structure (Fig. 4.63). It has become an important tool in environmental studies of near-surface features, such as buried and forgotten waste deposits, fracture patterns in otherwise uniform rock bodies, or the investigation of groundwater resources.

High-frequency signals are rapidly attenuated with depth. Geometrical spreading of the signal outward from its source (spherical divergence) causes a decrease in intensity with distance. More important is absorption of the signal by ground materials, which is a function of their conductivity. Depending on the composition of the soil or rocks (e.g., the presence of clay-rich layers or groundwater), the nature of subsurface structures and the frequency of the radar signal, the effective penetration may be up to 10 m, although conditions commonly restrict it to only a few meters. However, at a radar frequency of $10^8$–$10^9$ Hz and with a velocity of $\approx 10^8$ m s$^{-1}$, the resolution is in the range 0.1–1 m. Thus, despite its limited depth penetration, the high resolution of georadar makes it a powerful tool for near-surface geophysical exploration.

4.3.7 Electrical conductivity in the Earth

The complicated structure of the crust and upper mantle results in large lateral variations in electrical conductivity. Apart from the oceans, sediments and individual anomalous conductors, the outer carapace of the Earth is generally a poor electrical conductor. The physical mechanism of conductivity in silicate rocks is by semiconduction, which can take place in three different ways (Section 4.3.2.3). Each type of semiconduction is governed by a
4.3 GEOELECTRICITY

thermally activated process, in which the conductivity $\sigma$ at temperature $T$ is given by

$$\sigma = \sigma_0 e^{-E_a/kT} \quad (4.128)$$

where $k$ is Boltzmann’s constant ($k = 1.38065 \times 10^{-23}$ J K$^{-1}$). The constant $\sigma_0$ is the hypothetical maximum value of the conductivity, reached asymptotically at very high temperatures. $E_a$ is the activation energy of the particular type of semiconduction. Its value determines the temperature range in which the thermally activated process becomes effective as a mechanism for $\sigma$. In the crust and upper mantle (i.e., in the lithosphere) impurity semiconduction is likely the main mechanism in dry rocks. Electronic semiconduction is probably dominant in the asthenosphere and deeper regions of the mantle. Ionic semiconduction is an important mechanism at high temperature, but is unlikely to be significant below about 400 km, because it is suppressed by the high pressure at greater depth.

The electrical conductivity in the Earth at great depths is inferred from four sources: deep electrical sounding, geomagnetic variations, secular variations and extrapolation from laboratory experiments. The first two methods are based on induction effects arising from changes in the external part of the geomagnetic field; these encompass a broad spectrum with peaks of energy at several periods (see Fig. 4.45). Electrical and magnetotelluric sounding use the components with periods from milliseconds to one or two days. The inversion of MT data gives a conductivity pattern that is generally concordant with seismic data and related to the broad geological structure of the crust and upper mantle.

The time spectrum of external geomagnetic field variations contains some prominent periods that are longer than a day (Fig. 4.45). The study of the longer-period geomagnetic variations provides information about conductivity in the Earth down to about 2000 km. The longer the period of the variation the deeper its penetration depth. The daily (or diurnal) variation (Section 5.4.3.3) yields conductivity information to about 900 km. Magnetic storms last several days or weeks and have a strong 48 hr component, which is used to extend conductivity information to about 1000 km. In addition to the spectrum of geomagnetic variation shown in Fig. 4.45 there is a longer-period component related to the 11-yr sunspot cycle. This results from increased solar activity and is accompanied by solar flares (see Section 5.4.7.1) and emissions of charged particles that augment the solar wind and excite ionospheric activity. Analysis of the 11-yr component allows the model of mantle conductivity to be extended to about 2000 km depth. Our knowledge of the electrical conductivity in the mantle at depths greater than 2000 km cannot be obtained from analysis of effects related to the external magnetic field.

The secular variation of the internal geomagnetic field (Section 5.4.5) originates in the upper part of the fluid outer core. It consists of fluctuations in intensity and direction with periods of the order of $10^{-4}$ yr. If the secular variation could be observed at the core–mantle boundary it would be possible to determine conductivity throughout the mantle. Unfortunately, secular variation must be observed at the Earth’s surface, after it has passed through the conducting mantle, which acts as a filter. The signal is attenuated by the skin effect, preferentially affecting the highest frequencies. Thus, observations of high-frequency changes in secular variation place an upper limit on the average conductivity of the mantle, because a greater conductivity would block them out. From time to time, abrupt changes in the rate of secular variation take place for unknown reasons. A conspicuous example of these “geomagnetic jerks” occurred in 1969–1970, when a pulse in secular variation occurred with an estimated duration of less than two years. Although not all analysts concur, the effect is widely believed to be of internal origin. Analysis of the propagation of a secular-variation pulse provides an estimate of the mean conductivity of the whole mantle, which, integrated with data from other sources, gives the conductivity in the lower mantle.

Some of the different models of mantle conductivity that have been proposed are shown in Fig. 4.64. The differences between the models reflect increases in quantity and improvements in quality of geomagnetic data as well as advances in the techniques of data-processing, especially the development of inversion methods. Although the models diverge in many respects they have some features in common. The conductivity averages about $10^{-2}$ $\Omega^{-1}$ m$^{-1}$ in the lithosphere and increases with increasing depth. Sharper rates of increase are found at depths of 400 and 670 km, where the olivine–spinel and spinel–perovskite phase changes occur, respectively (see Section 3.7.5.2). At about 700 km depth each model gives a conductivity of about 1 $\Omega^{-1}$ m$^{-1}$ which rises in the lower mantle to be about 10–200 $\Omega^{-1}$ m$^{-1}$ at the core–mantle boundary. The secular variation is not uniform over the Earth’s surface. Large areas of continental size (the Central Pacific is the best studied) are characterized by slow rates of variation. This is possibly due to the additional screening effect of features in the D$^\ast$-layer above the core–mantle boundary (Section 3.7.5.3), so-called “crypto-continents” (see Fig. 4.38) in which the conductivity may be 1000 times higher than in the overlying mantle (Stacey, 1992).

Conductivity in the outer core is estimated by extrapolation from laboratory experiments. The core has the composition of an iron alloy, with an iron content of about 83% and a concentration of the alloying elements of about 17%. The effect of pressure on the conductivity of the alloy is not large at this concentration. Measurements of resistivity at atmospheric pressure and different temperatures lead to an extrapolated resistivity of $\rho = 3.3 \times 10^{-6}$ $\Omega$ m at the temperature of the outer core, with a corresponding conductivity $\sigma = 3 \times 10^{-5}$ $\Omega^{-1}$ m$^{-1}$.
4.4 SUGGESTIONS FOR FURTHER READING

Introductory level


Intermediate level


Advanced level


4.5 REVIEW QUESTIONS

1. Define the following age-dating parameters: (a) decay constant, (b) half-life, (c) isochron.

2. The radioactive carbon method of age dating is a simple decay analysis. Explain what this statement means. Describe the principle of the method.

3. Describe the principle of a mass spectrometer. What is the Lorentz force?

4. What aspects make the $^{40}\text{K}/^{40}\text{Ar}$ method suitable for determining the ages of rocks? What advantages does $^{40}\text{Ar}/^{39}\text{Ar}$ dating have over the $^{40}\text{K}/^{40}\text{Ar}$ method?

5. What types of materials are suitable for dating with the radioactive carbon method? For what range of ages may it be applied? What are possible problems with the method?
6. Where are the oldest regions of the oceans? Where are the oldest continental regions? Compare the ages of the oldest oceanic and continental regions and account for the difference.

7. Explain the uranium–lead dating method. What is the concordia curve? What is the discordia line? Why is the U–Pb method suitable for dating very old materials, such as Precambrian rocks?

8. Why are zircons important for dating very old rocks? How do the ages of the oldest rocks on Earth compare with the ages of meteorites and the Moon?

9. What is meant by temperature? What is meant by heat?

10. What are the processes by which heat can be transferred? What is the relative importance of each process in (a) the crust, (b) the mantle, (c) the outer core, and (d) the inner core?

11. Sketch how (a) temperature and (b) the melting point (solidus) vary with depth in the Earth’s interior.

12. How is heat flow defined? How is it measured (a) on the continents and (b) in the oceans?

13. What factors determine the depth of penetration of solar energy into the earth? What precautions does this impose for measuring heat flow?

14. Why is the average oceanic heat flow higher than the average continental heat flow?

15. How does heat flow vary with distance from an oceanic ridge?

16. Which regions of the Earth have (a) the highest and (b) the lowest heat flow?

17. Discuss the statement: “The internal heat of the Earth causes the formation of mountains and the external heat of the Sun causes their destruction.”

18. Which characteristics of the ground determine its electrical resistivity?

19. Explain why geoelectrical resistivity measurements yield only an apparent resistivity.

20. What are telluric currents? How do they originate?

21. What is meant by the skin depth for the propagation of electromagnetic waves?

22. What are the in-phase and quadrature components in an electromagnetic induction survey? What causes the phase shift? Which component responds more strongly to the presence of a good conductor?

23. What is the magnetotelluric method of electromagnetic surveying? What are the merits of this method for deep Earth sounding?

24. What is ground-penetrating radar and in which part of the electromagnetic spectrum is it operative? Why can GPR signals be processed analogously to reflected and refracted seismic waves?

25. Why is GPR a powerful method for exploring shallow subsurface structure? Which properties of the ground determine the effectiveness of the method and limit its depth range?

4.6 EXERCISES

Geochronology

1. How many half-lives must elapse before the activity of a radioactive isotope decreases to 1% of its initial value? How long is this time for $^{14}$C, which has a decay rate of $1.21 \times 10^{-4}$ yr$^{-1}$?

2. Radiocarbon dating of a sample of wood from the tomb of an Egyptian pharaoh gave isotopic concentrations of 7045 p.p.m. for $^{14}$C and 144,330 p.p.m. for $^{12}$C. Assuming that the initial $^{14}$C/$^{12}$C ratio in the sample corresponded to the long-term atmospheric ratio of 1:12, determine the age of the tomb, the percentage of $^{14}$C remaining, and the original $^{14}$C concentration in the wood.

3. The decay constants of $^{235}$U and $^{238}$U are $\lambda_{235} = 9.8485 \times 10^{-10}$ yr$^{-1}$ and $\lambda_{238} = 1.55125 \times 10^{-10}$ yr$^{-1}$. Calculate the half-lives of these uranium isotopes.

4. Assuming that the isotopes $^{235}$U and $^{238}$U were created in a common event, such as a supernova, and given that their abundances are now in the ratio $^{235}$U/$^{238}$U = 1/137.88, calculate how long ago they were created.

5. The analysis of strontium and rubidium isotopes in whole rock samples from a granitic batholith gave the following concentrations in p.p.m.:

<table>
<thead>
<tr>
<th>Sample</th>
<th>$^{87}$Sr</th>
<th>$^{87}$Rb</th>
<th>$^{86}$Sr</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>2.304</td>
<td>8.831</td>
<td>2.751</td>
</tr>
<tr>
<td>B</td>
<td>0.518</td>
<td>29.046</td>
<td>0.450</td>
</tr>
<tr>
<td>C</td>
<td>1.619</td>
<td>111.03</td>
<td>1.232</td>
</tr>
<tr>
<td>D</td>
<td>1.244</td>
<td>100.60</td>
<td>0.871</td>
</tr>
</tbody>
</table>

Maximum heating temperature [°C]  $^{39}$Ar/$^{36}$Ar  $^{40}$Ar/$^{36}$Ar

| 750    | 1852     | 8855     |
| 830    | 1790     | 8439     |
| 895    | 1439     | 6867     |
| 970    | 3214     | 15380    |
| 1030   | 2708     | 12970    |

(a) Calculate the $^{87}$Rb/$^{86}$Sr and $^{87}$Sr/$^{86}$Sr isotopic ratios for these samples.
(b) Determine the age of the batholith and the initial $^{87}$Sr/$^{86}$Sr ratio.

6. Argon–argon dating of muscovite in a Late Cretaceous granite gave the following isotope ratios for the plateau stages during incremental heating:

(a) Calculate the $^{40}$Ar/$^{39}$Ar ratios for each incremental heating step.
(b) A calibration constant $J = 0.00964$ was determined for the monitor mineral. Using the $^{40}\text{Ar}^{39}\text{Ar}$ ratios from part (a) in Eq. (4.18), calculate the apparent ages at each heating step.

c) Draw an $^{40}\text{Ar}^{39}\text{Ar}$ isochron diagram by plotting each $^{40}\text{Ar}^{36}\text{Ar}$ ratio as ordinate against the corresponding $^{39}\text{Ar}^{36}\text{Ar}$ ratio as abscissa. Draw a best-fitting line – the isochron – through the data points, and determine its slope and intercept.

d) Compute the age of the muscovite from the slope of the isochron. Is the intercept on the ordinate axis significant?

7. The following isotopic ratios were measured in U–Pb age determinations on three zircon grains extracted from a granite:

<table>
<thead>
<tr>
<th>Sample</th>
<th>$^{207}\text{Pb}^{235}\text{U}$</th>
<th>$^{206}\text{Pb}^{238}\text{U}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>zircon 1</td>
<td>27.4</td>
<td>0.60</td>
</tr>
<tr>
<td>zircon 2</td>
<td>33.3</td>
<td>0.68</td>
</tr>
<tr>
<td>zircon 3</td>
<td>37.9</td>
<td>0.74</td>
</tr>
</tbody>
</table>

(a) Using the values listed in Table 4.2, plot a concordia diagram on graph paper or with a plotting routine. Enter the measurements from the above table on the graph, and draw the straight discordia line through the points.

(b) Determine the coordinates of the intersection points of the concordia and discordia lines.

(c) Using the coordinates of the upper intersection point together with Eq. (4.19) and Eq. (4.20), calculate the age of formation of the zircons.

(d) Calculate when loss of lead occurred in the zircons.

8. The following isotopic ratios were measured in a K–Ar age determination on an ignimbrite as part of a combined radiometric-paleomagnetic study of geomagnetic polarity.

<table>
<thead>
<tr>
<th>Sample</th>
<th>$^{40}\text{K}^{36}\text{Ar}$</th>
<th>$^{40}\text{Ar}^{36}\text{Ar}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>4,716,000</td>
<td>822</td>
</tr>
<tr>
<td>B</td>
<td>8,069,000</td>
<td>1200</td>
</tr>
<tr>
<td>C</td>
<td>12,970,000</td>
<td>1730</td>
</tr>
<tr>
<td>D</td>
<td>27,670,000</td>
<td>3280</td>
</tr>
</tbody>
</table>

(a) Plot the isotope ratios, draw the isochron, and compute its slope and intercept.

(b) Calculate the isochron age of the ignimbrite.

(c) Correct the observed $^{40}\text{Ar}^{36}\text{Ar}$ ratios for the initial $^{40}\text{Ar}^{36}\text{Ar}$ concentration, and compute the individual sample ages.

(d) Calculate the mean age and its standard deviation. Compare the mean age with the isochron age.

(e) With reference to the radiometric timescale in Fig. 5.74, what magnetic polarity would you expect the ignimbrite samples to have?

The Earth’s heat

9. List and compare the various factors that may influence the measured temperature gradient at a depth of 5 m in (a) a deep drillhole in oceanic sediments and (b) a continental well that encounters the groundwater table at 2 m depth.

10. A shallow circular pond 100 m in diameter freezes solid during a very cold night. The pond is in a geothermal area in which the temperature reaches 40 °C at 200 m depth. The thermal conductivity of the intervening rock is 3.75 W m$^{-1}$ K$^{-1}$ and the latent heat of fusion of ice is 334 kJ kg$^{-1}$. Neglecting other heat sources, calculate the mass of ice that melts per hour due to the geothermal gradient.

11. Assuming a constant geothermal gradient of 30 °C per kilometer, estimate what percentage of the Earth’s volume is hotter than the temperature of molten lava at atmospheric pressure. Why is the deeper interior of the Earth not entirely molten?

12. The mean global heat flow at the Earth’s surface is 82 mW m$^{-2}$. Calculate the time in years needed for the mantle and core to cool by 100 °C, with the following assumptions: (i) the Earth’s mantle and core cool as a homogeneous unit, (ii) 20% of the observed heat flow at the Earth’s surface is from the mantle, (iii) the lithospheric thickness is 100 km, (iv) thermal effects from the lithosphere itself may be ignored. Relevant properties of the mantle and core are: mean density 6000 kg m$^{-3}$, specific heat 400 J kg$^{-1}$ °C$^{-1}$.

13. A temperature gradient of 35 °C km$^{-1}$ is measured in the upper few meters of sediments covering the ocean floor. If the mean thermal conductivity of oceanic sediments is 1.7 W m$^{-1}$ °C$^{-1}$, calculate the local heat flow. How far do you think the sampling site is from the nearest active ridge?

14. What heat flow values would you expect at the locations of the oceanic magnetic anomalies with numbers C5N, C10N, C21N, C32N, M0? Interpret the ages of the anomalies from Fig. 5.78 and use the heat-flow model GDH1 (Eq. (4.62)) for the cooling of oceanic lithosphere.

15. Using the relationships in Eq. (4.63), estimate the approximate depths of the ocean at these locations? What is the thickness of the elastic lithosphere and the depth of the top of the asthenosphere at these locations (see Fig. 2.79)?

16. Assuming that the Earth initially had a uniform temperature throughout and has been cooling by con-
4.6 EXERCISES

17. The temperature in the near-surface layers of the Earth’s crust varies cyclically with daily, annual and longer periods. For a surface temperature variation given by \( T = T_0 \cos(\omega t) \), the temperature variation at depth \( z \) and time \( t \) is described by:

\[
T(z,t) = T_0 \exp\left( -\frac{z}{d} \right) \cos\left( \omega t - \frac{z}{d} \right)
\]

where \( \Phi \) is the period of the variation, \( k \) is the thermal conductivity, \( c_p \) is the specific heat, and \( p \) is the density. For surface sediments assume \( k = 2.5 \text{ W m}^{-1} \text{C}^{-1} \), \( c_p = 10^3 \text{ J kg}^{-1} \text{C}^{-1} \), and \( p = 2300 \text{ kg m}^{-3} \).

(a) Calculate the phase difference (in days) between the temperature variation at the surface and at depths of 2 m and 5 m, respectively. Perform the calculations for both the daily and annual temperature fluctuations.

(b) Assuming that the range in surface temperatures between summer and winter is 40 °C, calculate the depth at which the annual temperature range is 5 °C. How large (in weeks and days) is the phase difference between the surface temperature and the actual temperature at this depth?

18. The daily average temperature in northern Canada is +10 °C in July and −20 °C in January. Using the heat conduction equation calculate the depth of the permafrost (below which the ground is permanently frozen). Relevant physical properties of the ground are: thermal conductivity \( k = 3 \text{ W m}^{-1} \text{C}^{-1} \), specific heat \( c_p = 840 \text{ J kg}^{-1} \text{C}^{-1} \), density \( p = 2700 \text{ kg m}^{-3} \).

19. The half-spreading rate at an oceanic ridge in the middle of a symmetric ocean basin bounded by subduction zones is 44 mm yr\(^{-1}\). The ridge is 1000 km long and the distance from the ridge to each subduction zone is 2000 km. If the oceanic heat flow varies with crustal age as in Eq. (4.62), calculate how much heat is lost per year from the ocean basin.

20. At the interface between two layers with electrical resistivities \( \rho_1 \) and \( \rho_2 \), as in the figure below, the electrical boundary conditions are: (i) the component of current density \( J \), normal to the interface is continuous, and (ii) the component of electric field \( E \), tangential to the interface is continuous. A current flow-line makes angles \( \theta_1 \) and \( \theta_2 \) before and after refraction, respectively.

\[
\begin{align*}
J_1 & = J_2 \\
E_{ix} & = E_{2x}
\end{align*}
\]

Derive the electrical “law of refraction” given by Eq. (4.102):

\[
\tan \theta_1 = \frac{\rho_2}{\rho_1}
\]

21. What is the effective resistivity of a slab of thickness \( L \) composed of two half-slabs each of thickness \( L/2 \) and with resistivities \( (\rho/2) \) and \( (\rho/2) \), respectively, as in the diagram?

22. Sea-water is contaminating an aquifer that is the source of drinking water for a seaside town. The following measurements of apparent resistivity \( (\rho_a) \) were made at various electrode separations \( (a) \) with the expanding-spread Wenner method to investigate the leak.

<table>
<thead>
<tr>
<th>( a ) [m]</th>
<th>( \rho_a ) [Ω m]</th>
<th>( a ) [m]</th>
<th>( \rho_a ) [Ω m]</th>
<th>( a ) [m]</th>
<th>( \rho_a ) [Ω m]</th>
<th>( a ) [m]</th>
<th>( \rho_a ) [Ω m]</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>29.0</td>
<td>40</td>
<td>28.5</td>
<td>100</td>
<td>23.5</td>
<td>120</td>
<td>21.7</td>
</tr>
<tr>
<td>20</td>
<td>28.9</td>
<td>60</td>
<td>27.1</td>
<td>80</td>
<td>25.3</td>
<td>60</td>
<td>27.1</td>
</tr>
<tr>
<td>40</td>
<td>28.5</td>
<td>80</td>
<td>25.3</td>
<td>100</td>
<td>23.5</td>
<td>40</td>
<td>14.5</td>
</tr>
<tr>
<td>60</td>
<td>27.1</td>
<td>100</td>
<td>23.5</td>
<td>120</td>
<td>21.7</td>
<td>120</td>
<td>9.9</td>
</tr>
</tbody>
</table>
(a) Estimate the electrical resistivity of each layer.

(b) Divide the apparent resistivity at each position by the resistivity of the upper layer, then plot the normalized resistivity against electrode separation on a log–log diagram on the same scale as the model curves in Fig. 4.53.

(c) Match the measured curve with the model curves and estimate the depth to the interface.

23. In the Schlumberger resistivity method the separation of the current electrodes $L$ is much larger than the separation $a$ of the voltage electrodes. Suppose that the mid-point of the voltage pair is displaced by a distance $x$ from the mid-point of the current electrode pair. Show that, for $(L - 2x) \gg a$, the apparent resistivity is given by

$$\rho_a = \frac{\pi V}{4 T a(L^2 + 4x^2)}$$

24. In the double-dipole resistivity method it is common to keep the separation of the pairs $L$ an integer multiple $n$ of the distance $a$ between the electrodes in each pair, i.e. $L = na$.

(a) Rewrite the formula for the apparent resistivity with this assumption.

(b) If $L$ is very large compared to $a$, modify the formula to show that the apparent resistivity is proportional to $n^3$.

25. Consider a double-dipole configuration in which the electrode pairs are not collinear but are broadside to each other (i.e., normal to the line joining them). The electrode separation is $a$ and the distance between the mid-points of the pairs is $L = na$. Show that, for large values of $n$, the apparent resistivity is given in this case by

$$\rho_a = 2\pi n^3 a^2 V$$

26. Calculate the velocity of a long-wavelength electromagnetic wave in (a) basalt (dielectric constant $\kappa = 12$) and (b) water ($\kappa = 80.4$).

27. Calculate the “skin depths” of penetration in (a) granite ($\rho = 5,000 \ \Omega \ \text{m}$) and (b) a pyrrhotite ore-body ($\rho = 5 \times 10^{-5} \ \text{t} \ \text{m}$) for electromagnetic waves in surveys employing (i) electromagnetic induction ($f = 1 \ \text{kHz}$) and (ii) ground penetrating radar ($f = 100 \ \text{MHz}$). Would these methods detect the conducting bodies if they were buried under a water-saturated soil layer, 3 m thick with resistivity $100 \ \Omega \ \text{m}$?
5.1 HISTORICAL INTRODUCTION

5.1.1 The discovery of magnetism

Mankind’s interest in magnetism began as a fascination with the curious attractive properties of the mineral lodestone, a naturally occurring form of magnetite. Called loadstone in early usage, the name derives from the old English word load, meaning “way” or “course”; the loadstone was literally a stone which showed a traveller the way.

The earliest observations of magnetism were made before accurate records of discoveries were kept, so that it is impossible to be sure of historical precedents. Nevertheless, Greek philosophers wrote about lodestone around 800 BC and its properties were known to the Chinese by 300 BC. To the ancient Greeks science was equated with knowledge, and was considered an element of philosophy. As a result, the attractive forces of lodestone were ascribed to metaphysical powers. Some early animistic philosophers even believed lodestone to possess a soul. Contemporary mechanistic schools of thought were equally superstitious and gave rise to false conceptions that persisted for centuries. Foremost among these was the view that electrical and magnetic forces were related to invisible fluids. This view persisted well into the nineteenth century. The power of a magnet seemed to flow from one pole to the other along lines of induction that could be made visible by sprinkling iron filings on a paper held over the magnet. The term “flux” (synonymous with flow) is still found in “magnetic flux density,” which is regularly used as an alternative to “magnetic induction” for the fundamental magnetic field vector $B$.

One of the greatest and wealthiest of the ancient Greek city-colonies in Asia Minor was the seaport of Ephesus, at the mouth of the river Meander (modern Küçük Menderes) in the Persian province of Caria, in what is now the Turkish province of western Anatolia. In the fifth century BC the Greek state of Thessaly founded a colony on the Meander close to Ephesus called Magnesia, which after 133 BC was incorporated into the Roman empire as Magnesia ad Maeandrum. In the vicinity of Magnesia the Greeks found a ready supply of lodestone, pieces of which subsequently became known by the Latin word magneta from which the term magnetism derives.

It is not known when the directive power of the magnet – its ability to align consistently north–south – was first recognized. Early in the Han dynasty, between 300 and 200 BC, the Chinese fashioned a rudimentary compass out of lodestone. It consisted of a spoon-shaped object, whose bowl balanced and could rotate on a flat polished surface. This compass may have been used in the search for gems and in the selection of sites for houses. Before 1000 AD the Chinese had developed suspended and pivoted-needle compasses. Their directive power led to the use of compasses for navigation long before the origin of the aligning forces was understood. As late as the twelfth century, it was supposed in Europe that the alignment of the compass arose from its attempt to follow the pole star. It was later shown that the compass alignment was produced by a property of the Earth itself. Subsequently, the characteristics of terrestrial magnetism played an important role in advancing the understanding of magnetism.

5.1.2 Pioneering studies in terrestrial magnetism

In 1269 the medieval scholar Pierre Pélerin de Maricourt, who took the Latin nom-de-plume of Petrus Peregrinus, wrote the earliest known treatise of experimental physics (Epistola de Magnete). In it he described simple laws of magnetic attraction. He experimented with a spherical magnet made of lodestone, placing it on a flat slab of iron and tracing the lines of direction which it assumed. The term “flux” (synonymous with flow) is still found in “magnetic flux density,” which is regularly used as an alternative to “magnetic induction” for the fundamental magnetic field vector $B$.

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declination was the principal cause of error in contemporary map-making.

Georg Hartmann, a German cleric, discovered in 1544 that a magnetized needle assumed a non-horizontal attitude in the vertical plane. The deviation from the horizontal is now called the magnetic inclination. He reported his discovery in a letter to his superior, Duke Albrecht of Prussia, who evidently was not impressed. The letter lay unknown to the world in the royal archives until its discovery in 1831. Meanwhile, an English scientist, Robert Norman, rediscovered the inclination of the Earth's magnetic field independently in 1576.

In 1600 William Gilbert (1544–1603), an English scientist and physician to Queen Elizabeth, published De Magnete, a landmark treatise in which he summarized all that was then known about magnetism, including the results of about seventeen years of his own research. His studies extended also to the electrostatic effects seen when some materials were rubbed, for which he coined the name “electricity” from the Greek word for amber. Gilbert was the first to distinguish clearly between electrical and magnetic phenomena. His magnetic studies followed the work of Peregirus three centuries earlier. Using small magnetic needles placed on the surface of a sphere of lodestone to study its magnetic field, he recognized the poles, where the needles stood on end, and the equator, where they lay parallel to the surface. Gilbert achieved the leap of imagination that was necessary to see the analogy between the attraction of the lodestone sphere and the known magnetic properties of the Earth. He recognized that the Earth itself behaved like a large magnet. This was the first unequivocal recognition of a geophysical property, preceding the laws of gravitation in Newton’s Principia by almost a century. Although founded largely on qualitative observations, De Magnete was the most important work on magnetism until the nineteenth century.

The discovery that the declination of the geomagnetic field changed with time was made by Henry Gellibrand (1597–1637), an English mathematician and astronomer, in 1634. He noted, on the basis of just three measurements made by William Borough in 1580, Edmund Gunter in 1622 and himself in 1634, that the declination had decreased by about 7° in this time. From these few observations he deduced what is now called the secular variation of the field.

Gradually the variation of the terrestrial magnetic field over the surface of the Earth was established. In 1698–1700 Edmund Halley, the English astronomer and mathematician, carried out an important oceanographic survey with the prime purpose of studying compass variations in the Atlantic ocean. In 1702 this resulted in the publication of the first declination chart.

5.1.3 The physical origins of magnetism

By the end of the eighteenth century many characteristics of terrestrial magnetism were known. The qualitative properties of magnets (e.g., the concentration of their powers at their poles) had been established, but the accumulated observations were unable to provide a more fundamental understanding of the phenomena because they were not quantitative. A major advance was achieved by Charles Augustin de Coulomb (1736–1806), the son of a noted French family, who in 1784 invented a torsion balance that enabled him to make quantitative measurements of electrostatic and magnetic properties. In 1785 he published the results of his intensive studies. He established the inverse-square law of attraction and repulsion between small electrically charged balls. Using thin, magnetized steel needles about 24 inches (61 cm) in length, he also established that the attraction or repulsion between their poles varied as the inverse square of their separation.

Alessandro Volta (1745–1827) invented the voltaic cell with which electrical currents could be produced. The relationship between electrical currents and magnetism was detected in 1820 by Hans Christian Oersted (1777–1851), a Danish physicist. During experiments with a battery of voltaic cells he observed that a magnetic needle is deflected at right angles to a conductor carrying a current, thus establishing that an electrical current produces a magnetic force.

Oersted’s result was met with great enthusiasm and was followed at once by other notable discoveries in the same year. The law for the direction and strength of the magnetic force near a current-carrying wire was soon formulated by the French physicists Jean-Baptiste Biot (1774–1862) and Felix Savart (1791–1841). Their compatriot André Marie Ampère (1775–1836) quickly undertook a systematic set of experiments. He showed that a force existed between two parallel straight current-carrying wires, and that it was of a type different from the known electrical forces. Ampère experimented with the magnetic forces produced by current loops and proposed that internal electrical currents were responsible for the existence of magnetism in iron objects (i.e., ferromagnetism). This idea of permanent magnetism due to constantly flowing currents was audacious for its time.

At this stage, the ability of electrical currents to generate magnetic fields was known, but it fell to the English scientist Michael Faraday (1791–1867), to demonstrate in 1831 what he called “magneto-electric” induction. Faraday came from a humble background and had little mathematical training. Yet he was a gifted experimenter, and his results demonstrated that the change of magnetic flux in a coil (whether produced by introducing a magnet or by the change in current in another coil) induced an electric current in the coil. The rule that governs the direction of the induced current was formulated three years later by a Russian physicist, Heinrich Lenz (1804–1865). Unhampered by mathematical equations, Faraday made fundamental contributions to understanding magnetic processes. Instead of regarding magnetic and electrical phenomena as the effects of centers of force acting at a distance, he saw in his mind’s eye fictional lines of force.
5.2 THE PHYSICS OF MAGNETISM

traversing space. This image emphasized the role of the medium and led eventually to the concept of magnetic field, which Faraday first used in 1845.

Although much had been established by the early 1830s, it was still necessary to interpret the strengths of magnetic forces by relating magnetic units to mechanical units. This was achieved in 1832 by the German scientist and mathematician, Carl Friedrich Gauss (1777–1855), who assumed that static magnetism was carried by magnetic “charges,” analogous to the carriers of static electricity. Experiment had shown that, in contrast to electric charge, magnetic poles always occur as oppositely signed pairs, and so the basic unit of magnetic properties corresponds to the dipole. Together with Wilhelm Weber (1804–1891), Gauss developed a method of absolute determination of the intensity of the Earth’s magnetic field. They founded a geomagnetic observatory at Göttingen where the Earth’s magnetic field was observed at regular intervals. By 1837 global charts of the total intensity, inclination and declination were in existence, although the data had been measured at different times and their areal coverage was incomplete. To analyze the data-set Gauss applied the mathematical techniques of spherical harmonic analysis and the separation of variables, which he had invented. In 1839 he established that the main part of the Earth’s magnetic field was a dipole field that originated inside the Earth.

The fundamental physical laws governing magnetic effects were now firmly established. In 1872 James Clerk Maxwell (1831–1879), a Scottish physicist, derived a set of equations that quantified all known relationships between electrical and magnetic phenomena: Coulomb’s laws of force between electric charges and magnetic poles; Oersted’s and Ampère’s laws governing the magnetic effects of electric currents; Faraday’s and Lenz’s laws of electromagnetic induction and Ohm’s law relating current to electromotive force. Maxwell’s mathematical studies predicted the propagation of electric waves in space, and concluded that light is also an electromagnetic phenomenon transmitted through a medium called the luminiferous ether. The need for this light-transmitting medium was eliminated by the theory of relativity. By putting the theory of the electromagnetic field on a mathematical basis, Maxwell enabled a greater understanding of electromagnetic phenomena before the discovery of the electron.

A further notable discovery was made in 1879 by Heinrich Lorentz (1853–1928), a Dutch physicist. In experiments with vacuum tubes he observed the deflection of a beam of moving electrical charge by a magnetic field. The deflecting force acted in a direction perpendicular to the magnetic field and to the velocity of the charged particles, and was proportional to both the field and the velocity. This result now serves to define the unit of magnetic induction.

Since the time of man’s first awareness of magnetic behavior, students of terrestrial magnetism have made important contributions to the understanding of magnetism as a physical phenomenon. In turn, advances in the physics of magnetism have helped geophysicists to understand the morphology and origin of the Earth’s magnetic field, and to apply this knowledge to geological processes, such as global tectonics. The physical basis of magnetism is fundamental to the geophysical topics of geomagnetism, rock magnetism and paleomagnetism.

5.2 THE PHYSICS OF MAGNETISM

5.2.1 Introduction

Early investigators conceptualized gravitational, electrical and magnetic forces between objects as instantaneous effects that took place through direct action-at-a-distance. Faraday introduced the concept of the field of a force as a property of the space in which the force acts. The force-field plays an intermediary role in the interaction between objects. For example, an electric charge is surrounded by an electrical field that acts to produce a force on a second charge. The pattern of a field is portrayed by field lines. At any point in a field the direction of the force is tangential to the field line and the intensity of the force is proportional to the number of field lines per unit cross-sectional area.

Problems in magnetism are often more complicated for the student than those in gravitation and electrostatics. For one thing, gravitational and electrostatic fields act centrally to the source of force, which varies in each case as the inverse square of distance. Magnetic fields are not central; they vary with azimuth. Moreover, even in the simplest case (that of a magnetic dipole or a small current loop) the field strength falls off inversely as the cube of distance. To make matters more complicated, the student has to take account of two magnetic fields (denoted by B and H).

The confusion about the B-field and the H-field may be removed by recalling that all magnetic fields originate with electrical currents. This is the case even for permanent magnets, as Ampère astutely recognized in 1820. We now know that these currents are associated with the motions of electrons about atomic nuclei in the permanent magnets. The fundamental magnetic field associated with currents in any medium is B. The quantity H should be regarded as a computational parameter proportional to B in non-magnetizable materials. Inside a magnetizable material, H describes how B is modified by the magnetic polarization (or magnetization, M) of the material. The magnetic B-field is also called the magnetic induction or magnetic flux density.

Historically, the laws of magnetism were established by relating the B-field to fictitious centers of magnetic force called magnetic poles, defined by comparison with the properties of a bar magnet. Gauss showed that, in contrast to electrostatic charges, free magnetic poles cannot exist; each positive pole must be paired with a
corresponding negative pole. The most important type of magnetic field – and also the dominant component of the geomagnetic field – is that of a magnetic dipole (Fig. 5.1a). This is the field of two magnetic poles of opposite sense that are infinitesimally close to each other. The geometry of the field lines shows the paths along which a free magnetic pole would move in the vicinity of the dipole. A tiny current loop (Fig. 5.1b) and a uniformly magnetized sphere (Fig. 5.1c) also have dipole-type magnetic fields around them. Although magnetic poles do not exist physically, many problems that arise in geophysical situations can be readily solved in terms of surface distributions of poles or dipoles. So we will first examine these concepts.

5.2.2 Coulomb’s law for magnetic poles

Coulomb’s experiments in 1785 established that the force between the ends of long thin magnets was inversely proportional to the square of their separation. Gauss expanded Coulomb’s observations and attributed the forces of attraction and repulsion to fictitious magnetic charges, or poles. An inverse square law for the force $F$ between magnetic poles with strengths $p_1$ and $p_2$ at distance $r$ from each other can be formulated as

$$F(r) = K \frac{p_1 p_2}{r^2} \quad (5.1)$$

The proportionality constant $K$ was originally defined to be dimensionless and equal to unity, analogously to the law of electrostatic force. This gave the dimensions dyne 1/2 cm.

5.2.2.1 The field of a magnetic pole

The gravitational field of a given mass is defined as the force it exerts on a unit mass (Section 2.2.2). Similarly, the electric field of a given charge is the force it exerts on a unit charge. These ideas cannot be transferred directly to magnetism, because magnetic poles do not really exist. Nevertheless, many magnetic properties can be described and magnetic problems solved in terms of fictitious poles.

For example, we can define a magnetic field $B$ as the force exerted by a pole of strength $p$ on a unit pole at distance $r$. From Eq. (5.1) we get

$$B(r) = K \frac{p}{r^2} \quad (5.2)$$

Setting $K = 1$, the unit of the magnetic $B$-field has dimensions dyne 1/2 cm in c.g.s. units and is called a gauss. Geophysicists employ a smaller unit, the gamma ($\gamma$), to describe the geomagnetic field and to chart magnetic anomalies ($1 \gamma = 10^{-5}$ gauss).

Unfortunately, the c.g.s. system required units of electrical charge that had different dimensions and size in electrostatic and electromagnetic situations. By international agreement the units were harmonized and rationalized. In the modern Système Internationale (SI) units the proportionality constant $K$ is not dimensionless. It has the value $\mu_0/4\pi$, where $\mu_0$ is called the permeability constant and is equal to $4\pi \times 10^{-7}$ N A$^{-2}$ (or henry/meter, H m$^{-1}$, which is equivalent to N A$^{-2}$).

5.2.2.2 The potential of a magnetic pole

In studying gravitation we also used the concept of a field to describe the region around a mass in which its attraction could be felt by another test mass. In order to move the test mass away from the attracting mass, work had to be done against the attractive force and this was found to be equal to the gain of potential energy of the test mass. When the test mass was a unit mass, the attractive force was called the gravitational field and the gain in potential energy was called the change in potential. We calculated the gravitational potential at distance $r$ from an attracting point mass by computing the work that would have to be expended against the field to move the unit mass from $r$ to infinity.

We can define the magnetic potential $W$ at a distance $r$ from a pole of strength $p$ in exactly the same way. The magnetic field of the pole is given by Eq. (5.2). Using the value $\mu_0/4\pi$ for $K$ and expressing the pole strength $p$ in SI units, the magnetic potential at $r$ is given by

$$W = - \int_1^r B \, dr = \frac{\mu_0 p}{4\pi r} \quad (5.3)$$
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5.2.3 The magnetic dipole

In Fig. 5.1 the line joining the positive and negative poles (or the normal to the plane of the loop, or the direction of magnetization of the sphere) defines an axis, about which the field has rotational symmetry. Let two equal and opposite poles, +p and –p, be located a distance d apart (Fig. 5.2). The potential W at a distance r from the mid-point of the pair of poles, in a direction that makes an angle θ to the axis, is the sum of the potentials of the positive and negative poles. At the point (r, θ) the distances from the respective poles are \( r_+ \) and \( r_- \) and we get for the magnetic potential of the pair

\[
W = \frac{\mu_0 p}{4\pi} \left( \frac{1}{r_+} + \frac{1}{r_-} \right) \quad (5.4)
\]

\[
W = \frac{\mu_0 p}{4\pi} \left( \frac{r_- - r_+}{r_+ r_-} \right) \quad (5.5)
\]

The pair of opposite poles is considered to form a dipole when their separation becomes infinitesimally small compared to the distance to the point of observation (i.e., \( d \ll r \)). In this case, we get the approximate relations

\[
r_+ = r - \frac{d}{2} \cos \theta
\]

\[
r_- = r + \frac{d}{2} \cos \theta \quad (5.6)
\]

When \( d \ll r \), we can write \( \theta = \theta' + \text{terms of order } (d/r)^2 \) and higher can be neglected. This leads to the further simplifications

\[
r_- - r_+ = \frac{d}{4} (\cos \theta' + \cos \theta) = d \cos \theta
\]

\[
r_+ r_- = r^2 - \frac{d^2}{4} \cos^2 \theta = r^2 \quad (5.7)
\]

\[
\tau = m \times B
\]

5.2.4 The magnetic field of an electrical current

The equation used to define the magnetic \( B \)-field was formulated by Lorentz in 1879. Let \( q \) be an electrical charge that moves with velocity \( v \) through a magnetic field \( B \) (Fig. 5.4a). The charged particle experiences a deflecting force \( F \) given by Lorentz’s law, which in SI units is:

\[
F = q(v \times B) \quad (5.10)
\]
The SI unit of the magnetic $B$-field defined by this equation is called a tesla; it has the dimensions $N\ A^{-1}\ m^{-1}$.

Imagine the moving charge to be confined to move along a conductor of length $dL$ and cross-section $A$ (Fig. 5.4b). Let the number of charges per unit volume be $N$. The number inside the element $dL$ is then $NA_d$. Each charge experiences a deflecting force given by Eq. (5.10). Thus the total force transferred to the element $dL$ is

$$dF = I(dL \times B)$$

The electrical current $I$ along the conductor is the total charge that crosses $A$ per second, and is given by $I = NA_v$. From Eq. (5.11) we get the law of Biot and Savart for the force experienced by the element $dL$ of a conductor carrying a current $I$ in a magnetic field $B$:

$$dF = I(dL \times B)$$

The orienting effect of an electrical current on magnetic compass needles, reported by Oersted and Ampère in 1820, is illustrated in Fig. 5.5. The magnetic field lines around an infinitely long straight wire carry an electrical current form concentric circles in a plane normal to the wire.

The Biot–Savart law can be applied to determine the torque exerted on a small rectangular loop PQRS in a magnetic field (Fig. 5.6a). Let the lengths of the sides of the loop be $a$ and $b$, respectively, and define the $x$-axis parallel to the sides of length $a$. The area of the loop can be expressed as a vector with magnitude $A = ab$, and direction $\mathbf{n}$ normal to the plane of the loop. Suppose that a current $I$ flows in the loop and that a magnetic field $\mathbf{B}$ acts normal to the $x$-axis, making an angle $\theta$ with the
normal to the plane of the loop. Applying Eq. (5.12), a force \( F_x \) equal to \((ibB \cos \theta)\) acts on the side PQ in the direction of \(+x\); its effect is cancelled by an equal and opposite force \( F_x \) acting on side RS in the direction of \(-x\). Forces equal to \((iaB)\) act in opposite directions on the sides QR and SP (Fig. 5.6b). The perpendicular distance between their lines of action is \(b \sin \theta\), so the torque \( \tau \) experienced by the current loop is

\[
\tau = (iaB)b \sin \theta = (IA)B \sin \theta
\]

(5.14)

The quantity \( m = IA \) is a vector with direction parallel to the normal to the plane of the current loop. This expression is valid for an arbitrary small loop of area \( A \), regardless of its shape. By comparing Eqs. (5.14) and (5.9) for the torque on a dipole, it is evident that \( m \) corresponds to the magnetic moment of the current loop. At distances greater than the dimensions of the loop, the magnetic field is that of a dipole at the center of the loop (Fig. 5.1b). The definition of \( m \) in terms of a current-carrying loop shows that magnetic moment has the units of current times area (A m²).

5.2.5 Magnetization and the magnetic field inside a material

A true picture of magnetic behavior requires a quantum-mechanical analysis. Fortunately, a working understanding of the magnetic behavior of materials can be acquired without getting involved in the quantum-mechanical details. The simplified concept of atomic structure introduced by Ernest Rutherford in 1911 gives a readily understandable model for the magnetic behavior of materials. The motion of an electron around an atomic nucleus is treated like the orbital motion of a planet about the Sun. The orbiting charge forms an electrical current with which an orbital magnetic moment is associated. A planet also rotates about its axis; likewise each electron can be visualized as having a spin motion about an axis. The spinning electrical current produces a spin magnetic moment. Each magnetic moment is directly related to the corresponding angular momentum. In quantum theory each type of angular momentum of an electron is quantized. Thus the spin and orbital magnetic moments are restricted to having discrete values. The spin magnetic moment is usually more important than the orbital moment in the rock-forming minerals (see Section 5.2.6).

A simplified picture of the magnetic moments inside a material is shown in Fig. 5.7. The magnetic moment \( m \) of each atom is associated with a current loop as illustrated in Fig. 5.1b and described in the previous section. The net magnetic moment of a volume \( V \) of the material depends on the degree of alignment of the individual atomic magnetic moments. It is the vector sum of all the atomic magnetic moments in the material. The magnetic moment per unit volume of the material is called its magnetization, denoted \( M \):
acts in the opposite direction to the magnetization $\mathbf{M}$. Outside the magnet the $\mathbf{B}$-field and $\mathbf{H}$-field are parallel; the $\mathbf{H}$-field is discontinuous at the ends of the magnet.

The same situation can be portrayed in terms of current loops. The physical source of every $\mathbf{B}$-field is an electrical current, even in a permanent magnet (Fig. 5.8b). Atomic current loops give a continuous $\mathbf{B}$-field that emerges from the magnet at one end, re-enters at the other end and is closed inside the magnet. The aligned magnetic moments of the elementary current loops cancel out inside the body of the magnet, but the currents in the loops adjacent to the sides of the magnet combine to form a surface “current” that maintains the magnetization $\mathbf{M}$.

In a vacuum there is no magnetization ($\mathbf{M} = 0$); the vectors $\mathbf{B}$ and $\mathbf{H}$ are parallel and proportional ($\mathbf{B} = \mu_0 \mathbf{H}$). Inside a magnetizable material the magnetic $\mathbf{B}$-field has two sources. One is the external system of real currents that produce the magnetizing field $\mathbf{H}$; the other is the set of internal atomic currents that cause the atomic magnetic moments whose net alignment is expressed as the magnetization $\mathbf{M}$. In a general, anisotropic magnetic material $\mathbf{B}$, $\mathbf{M}$ and $\mathbf{H}$ are not parallel. However, many magnetic materials are not strongly anisotropic and the elementary atomic magnetic moments align in a statistical fashion with the magnetizing field. In this case $\mathbf{M}$ and $\mathbf{H}$ are parallel and proportional to each other

$$\mathbf{M} = k \mathbf{H} \quad (5.17)$$

The proportionality factor $k$ is a physical property of the material, called the magnetic susceptibility. It is a measure of the ease with which the material can be magnetized. Because $\mathbf{M}$ and $\mathbf{H}$ have the same units (A m$^{-1}$), $k$ is a dimensionless quantity. The susceptibility of most materials is temperature dependent, and in some materials (ferromagnets and ferrites) $k$ depends on $\mathbf{H}$ in a complicated fashion. In general, Eq. (5.16) can be rewritten

$$\mathbf{B} = \mu_0 (\mathbf{H} + \mathbf{M}) = \mu_0 \mathbf{H} (1 + k)$$

$$\mathbf{B} = \mu_0 \mathbf{H} \quad (5.18)$$

The quantity $\mu = (1 + k)$ is called the magnetic permeability of the material. The term “permeability” recalls the early nineteenth century association of magnetic powers with an invisible fluid. For example, the permeability of a material expresses the ability of the material to allow a fluid to pass through it. Likewise, the magnetic permeability is a measure of the ability of a material to convey a magnetic flux. Ferromagnetic metals have high permeabilities; in contrast, minerals and rocks have low susceptibilities and permeabilities $\mu \approx 1$.

5.2.6 The magnetic properties of materials

The magnetic behavior of a solid depends on the magnetic moments of the atoms or ions it contains. As discussed above, atomic and ionic magnetic moments are proportional to the quantized angular momenta associated with the orbital motion of electrons about the nucleus and with the spins of the electrons about their own axes of rotation. In quantum theory the exclusion principle of Wolfgang Pauli states that no two electrons in a given system can have the same set of quantum numbers. When applied to an atom or ion, Pauli’s principle stipulates that each possible electron orbit can be occupied by up to two electrons with opposite spins. The orbits are arranged in shells around the nucleus. The magnetic moments of paired opposite spins cancel each other out. Consequently, the net angular momentum
and the net magnetic moment of a filled shell must be zero. The net magnetic moment of an atom or ion arises from incompletely filled shells that contain unpaired spins. The atoms or ions in a solid are not randomly distributed but occupy fixed positions in a regular lattice, which reflects the symmetry of the crystalline structure and which controls interactions between the ions. Hence, the different types of magnetic behavior observed in solids depend not only on the presence of ions with unpaired spins, but also on the lattice symmetry and cell size.

Three main classes of magnetic behavior can be distinguished on the basis of magnetic susceptibility: \textit{diamagnetism}, \textit{paramagnetism} and \textit{ferromagnetism}. In diamagnetic materials the susceptibility is low and negative, i.e., a magnetization develops in the opposite direction to the applied field. Paramagnetic materials have low, positive susceptibilities. Ferromagnetic materials can be subdivided into three categories. True ferromagnetism is a cooperative phenomenon observed in metals like iron, nickel and cobalt, in which the lattice geometry and spacing allows the exchange of electrons between neighboring atoms. This gives rise to a \textit{molecular field} by means of which the magnetic moments of adjacent atoms reinforce their mutual alignment parallel to a common direction. Ferromagnetic behavior is characterized by high positive susceptibilities and strong magnetic properties. The crystal structures of certain minerals permit an indirect cooperative interaction between atomic magnetic moments. This \textit{indirect exchange} confers magnetic properties that are similar to ferromagnetism. The mineral may display \textit{antiferromagnetism} or \textit{ferrimagnetism}. The small group of ferrimagnetic minerals is geophysically important, especially in connection with the analysis of the Earth’s paleomagnetic field.

\subsection{5.2.6.1 Diamagnetism}

All magnetic materials show a diamagnetic reaction in a magnetic field. The diamagnetism is often masked by stronger paramagnetic or ferromagnetic properties. It is characteristically observable in materials in which all electron spins are paired.

The Lorentz law (Eq. (5.10)) shows that a change in the $B$-field alters the force experienced by an orbiting electron. The plane of the electron orbit is compelled to precess around the field direction; the phenomenon is called \textit{Larmor precession}. It represents an additional component of rotation and angular momentum. The sense of the rotation is opposite to that of the orbital rotation about the nucleus. Hence, the magnetic moment associated with the Larmor precession opposes the applied field. As a result a weak magnetization proportional to the field strength is induced in the opposite direction to the field. The magnetization vanishes when the applied magnetic field is removed. Diamagnetic susceptibility is reversible, weak and negative (Fig. 5.9a); it is independent of temperature. Many important rock-forming minerals belong to this class, amongst them quartz and calcite. They have susceptibilities around $-10^{-6}$ in SI units.

\subsection{5.2.6.2 Paramagnetism}

Paramagnetism is a statistical phenomenon. When one or more electron spins is unpaired, the net magnetic moment of an atom or ion is no longer zero. The resultant magnetic moment can align with a magnetic field. The alignment is opposed by thermal energy which favors chaotic orientations of the spin magnetic moments. The magnetic energy is small compared to the thermal energy, and in the absence of a magnetic field the magnetic moments are oriented randomly. When a magnetic field is applied, the chaotic alignment of magnetic moments is biased towards the field direction. A magnetization is induced proportional to the strength of the applied field and parallel to its direction. The susceptibility is reversible, small and positive (Fig. 5.9a). An important paramagnetic characteristic is that the susceptibility $k$ varies inversely with temperature (Fig. 5.9b) as given by the \textit{Curie law}

$$k = \frac{C}{T} \quad (5.19)$$

where the constant $C$ is characteristic of the material. Thus, a plot of $1/k$ against temperature is a straight line (Fig. 5.9c). In solids and liquids mutual interactions...
between ions may be quite strong and paramagnetic behavior is only displayed when the thermal energy exceeds a threshold value. The temperature above which a solid is paramagnetic is called the paramagnetic Curie temperature or Weiss constant, denoted by $\Theta$; it is close to zero degree kelvin in paramagnetic solids. At temperatures $T > \Theta$ the paramagnetic susceptibility $k$ is given by the Curie–Weiss law

$$k = \frac{C}{T - \Theta} \quad (5.20)$$

For a solid the plot of $1/k$ against $(T - \Theta)$ is a straight line (Fig. 5.9c). Many clay minerals and other rock-forming minerals (e.g., chlorite, amphibole, pyroxene, olivine) are paramagnetic at room temperature, with susceptibilities commonly around $10^{-5} - 10^{-4}$ in SI units.

### 5.2.6.3 Ferromagnetism

In paramagnetic and diamagnetic materials the interactions between individual atomic magnetic moments are small and often negligible. However, in some metals (e.g., iron, nickel, cobalt) the atoms occupy lattice positions that are close enough to allow the exchange of electrons between neighboring atoms. The exchange is a quantum-mechanical effect that involves a large amount of energy, called the exchange energy of the metal. The exchange interaction produces a very strong molecular field within the metal, which aligns the atomic magnetic moments (Fig. 5.10a) exactly parallel and produces a spontaneous magnetization ($M_s$). The magnetic moments react in unison to a magnetic field, giving rise to a class of strong magnetic behavior known as ferromagnetism.

A rock sample may contain thousands of tiny ferromagnetic mineral grains. The magnetization loop of a rock sample shows the effects of magnetic hysteresis (Fig. 5.11). In strong fields the magnetization reaches a saturation value (equal to $M_s$), at which the individual magnetic moments are aligned with the applied field. If the magnetizing field is reduced to zero, a ferromagnetic material retains part of the induced magnetization. The residual magnetization is called the remanence, or isothermal remanent magnetization (IRM); if the sample is magnetized to saturation, the remanence is a saturation IRM ($M_{rs}$). For a given ferromagnetic mineral, the ratio $M_{rs}/M_s$ depends on grain size. If a magnetic field is applied in the opposite direction to the IRM, it remagnetizes part of the material in the antiparallel direction. For a particular value $H_c$ of the reverse field (called the coercive force) the induced reverse magnetization exactly cancels the original remanence and the net magnetization is zero. If the reverse field is removed at this stage, the residual remanence is smaller than the original IRM. By repeating the process in ever stronger reverse fields a back-field $H_{cr}$ (called the coercivity of remanence) is found which gives a reverse remanence that exactly cancels the IRM, so that the residual remanence is zero. The ratio $H_{cr}/H_c$ also depends on grain size. Rock-forming magnetic minerals often have natural remanences with very high coercive properties.
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When a ferromagnetic material is heated, its spontaneous magnetization disappears at the ferromagnetic Curie temperature \((T_c)\). At temperatures higher than the paramagnetic Curie temperature \((\theta)\) the susceptibility \(k\) becomes the paramagnetic susceptibility, so that \(1/k\) is proportional to \((T - \theta)\) as given by the Curie-Weiss law (Eq. (5.20)). The paramagnetic Curie temperature for a ferromagnetic solid is several degrees higher than the ferromagnetic Curie temperature, \(T_c\). The gradual transition from ferromagnetic to paramagnetic behavior is explained by persistence of the molecular field due to short-range magnetic order above \(T_c\).

5.2.6.4 Antiferromagnetism

In oxide crystals the oxygen ions usually keep the metal ions far apart, so that direct exchange of electrons between the metal ions is not possible. However, in certain minerals, interaction between magnetic spins becomes possible by the exchange of electrons from one metal ion to another through the electron “cloud” of the oxygen ion. This indirect exchange (or superexchange) process results in antiparallel directions of adjacent atomic magnetic moments (Fig. 5.10b), giving two sublattices with equal and opposite intrinsic magnetic moments. As a result, the susceptibility of an antiferromagnetic crystal is weak and positive, and remanent magnetization is not possible. The antiferromagnetic alignment breaks down at the Néel temperature, \(T_N\), above which paramagnetic behavior is shown. The Néel temperature \(T_N\) of many antiferromagnetic substances is lower than room temperature, at which they are paramagnetic. A common example of an antiferromagnetic mineral is ilmenite \((\text{FeTiO}_3)\), which has a Néel temperature of 50 K.

5.2.6.5 Parasitic ferromagnetism

When an antiferromagnetic crystal contains defects, vacancies or impurities, some of the antiparallel spins are unpaired. A weak “defect moment” can result due to these lattice imperfections. Also if the spins are not exactly antiparallel but are inclined at a small angle, they do not cancel out completely and again a ferromagnetic type of magnetization can result (Fig. 5.10c). Materials that exhibit this form of parasitic ferromagnetism have the typical characteristics of a true ferromagnetic metal, including hysteresis, a spontaneous magnetization and a Curie temperature. An important geological example is the common iron mineral hematite \((\alpha-\text{Fe}_2\text{O}_3)\), in which both the spin-canted and defect moments contribute to the ferromagnetic properties. Hematite has a variable, weak spontaneous magnetization of about 2000 A m\(^{-1}\), very high coercivity and a Curie temperature around 675°C. The variable magnetic properties are due to variation in the relative importances of the defect and spin-canted moments.

5.2.6.6 Ferrimagnetism

The metallic ions in an antiferromagnet occupy the voids between the oxygen ions. In certain crystal structures, of which the most important geological example is the spinel structure, the sites of the metal ions differ from each other in the coordination of the surrounding oxygen ions. Tetrahedral sites have four oxygen ions as nearest neighbors and octahedral sites have six. The tetrahedral and octahedral sites form two sublattices. In a normal spinel the tetrahedral sites are occupied by divalent ions and the octahedral sites by \(\text{Fe}^{3+}\) ions. The most common iron oxide minerals have an inverse spinel structure. Each sublattice has an equal number of \(\text{Fe}^{3+}\) ions. The same number of divalent ions \((\text{e.g. Fe}^{2+})\) occupy other octahedral sites, while the corresponding number of tetrahedral sites is empty.

When the indirect exchange process involves antiparallel and unequal magnetizations of the sublattices (Fig. 5.10d), resulting in a net spontaneous magnetization, the phenomenon is called ferrimagnetism. Ferrimagnetic materials (called ferrites) exhibit magnetic hysteresis and retain a remanent magnetization when they are removed from a magnetizing field. Above a given temperature – sometimes called the ferrimagnetic Néel temperature but more commonly the Curie temperature – the long-range molecular order breaks down and the mineral behaves paramagnetically. The most important ferrimagnetic mineral is magnetite \((\text{Fe}_3\text{O}_4)\), but maghemite, pyrrhotite and goethite are also significant contributors to the magnetic properties of rocks.

5.2.7 Magnetic anisotropy

Anisotropy is the directional dependency of a property. The magnetism of metals and crystals is determined by the strengths of the magnetic moments associated with atoms or ions, and the distances between neighbors. Here the symmetry of the lattice plays an important role, and so the magnetic properties of most ferromagnetic materials depend on direction. Magnetic anisotropy is an important factor in the dependence on grain size of the magnetic behavior of rocks and minerals. There are three important types: magnetocrystalline, magnetostatic and magnetostrictive anisotropies.

5.2.7.1 Magnetocrystalline anisotropy

The direction of the spontaneous magnetization \((M_s)\) in a ferromagnetic metal is not arbitrary. The molecular field that produces \(M_s\) originates in the direct exchange of electron spins between neighboring atoms in a metal. The symmetry of the lattice structure of the metal affects the exchange process and gives rise to a magnetocrystalline anisotropy energy, which has a minimum value when \(M_s\) is parallel to a favored direction referred to as the easy axis (or easy direction) of magnetization. The simplest form of
magnetic anisotropy is uniaxial anisotropy, when a metal has only a single easy axis. For example, cobalt has a hexagonal structure and the easy direction is parallel to the c-axis at room temperature. Iron and nickel have cubic unit cells; at room temperature, the easy axes in iron are the edges of the cube, but the easy axes in nickel are the body diagonal directions.

Magnetocrystalline anisotropy is also exhibited by ferrites, including the geologically important ferrimagnetic minerals. The exchange process in a ferrite is indirect, but energetically preferred easy axes of magnetization arise that reflect the symmetry of the crystal structure. This gives rise to different forms of the anisotropy in hematite and magnetite.

Hematite has a rhombohedral or hexagonal structure and a uniaxial anisotropy with regard to the c-axis of symmetry. Oxygen ions form a close-packed hexagonal lattice in which two-thirds of the octahedral interstices are occupied by ferric (Fe$^{3+}$) ions. When the spontaneous magnetization makes an angle $\phi$ with the c-axis of the crystal, the uniaxial anisotropy energy density can be written to first order

$$E_a = K_u \sin^2 \phi$$

(5.21)

$K_u$ is called the uniaxial magnetocrystalline anisotropy constant. Its value in hematite is around $-10^3$ J m$^{-3}$ at room temperature. The negative value of $K_u$ in Eq. (5.21) means that $E_a$ decreases as the angle $\phi$ increases, and is minimum when $\sin^2 \phi$ is maximum, i.e., when $\phi = 90^\circ$. As a result, the spontaneous magnetization lies in the basal plane of the hematite crystal at room temperature.

Because of its inverse spinel structure, magnetite has cubic anisotropy. Let the direction of the spontaneous magnetization be given by direction cosines $\alpha_1$, $\alpha_2$, and $\alpha_3$ relative to the edges of the cubic unit cell (Box 1.5). The magnetocrystalline anisotropy energy density is then given by

$$E_a = K_1(\alpha_1^2\alpha_2^2 + \alpha_1^2\alpha_3^2 + \alpha_2^2\alpha_3^2) + K_2\alpha_1^2\alpha_2^2\alpha_3^2$$

(5.22)

The anisotropy constants $K_1$ and $K_2$ of magnetite are equal to $-1.36 \times 10^4$ J m$^{-3}$ and $-0.44 \times 10^4$ J m$^{-3}$, respectively, at room temperature. Because these constants are negative, the anisotropy energy density $E_a$ is minimum when the spontaneous magnetization is along a [111] body diagonal, which is the magnetocrystalline easy axis of magnetization at room temperature.

$5.2.7.2$ Magnetostatic (shape) anisotropy

In strongly magnetic materials the shape of the magnetized object causes a magnetostatic anisotropy. In rocks this effect is associated with the shapes of the individual grains of ferrimagnetic mineral in the rock, and to a lesser extent with the shape of the rock sample. The anisotropy is magnetostatic in origin and can be conveniently explained with the aid of the concept of magnetic poles.

The spontaneous magnetization of a uniformly magnetized material can be pictured as giving rise to a distribution of poles on the free end surfaces (Fig. 5.12). As noted above, a property of the magnetic B-field is that its demagnetizing field is that its field lines form closed loops, whereas the H-field begins and ends on boundary surfaces, at which it is discontinuous. The field lines of the magnetic field H outside a magnet are parallel to the B-field and are directed from the surface distribution of N-poles to the distribution of S-poles. In the absence of an externally applied field the H-field inside the magnet is also directed from the distribution of N-poles on one end to the S-poles on the other end. It forms a demagnetizing field $H_d$ that opposes the magnetization. The strength of the demagnetizing field varies directly with the surface density of the magnetic pole distribution on the end surfaces of the magnet, and inversely with the distance between these surfaces. $H_d$ thus depends on the shape of the magnet and the intensity of magnetization; it can be written

$$H_d = -NM$$

(5.23)

$N$ is called the demagnetizing factor. It is a dimensionless constant determined by the shape of the magnetic grain. It can be computed for a geometrical shape, such as a triaxial ellipsoid. The demagnetizing factors $N_1$, $N_2$, and $N_3$ parallel to the symmetry axes of an ellipsoid satisfy the relationship

$$N_1 + N_2 + N_3 = 1$$

(5.24)

The magnetostatic energy of the interaction of the grain magnetization with the demagnetizing field is called the
demagnetizing energy \( (E_d) \). For a grain with uniform magnetization \( M \) in a direction with demagnetizing factor \( N \),

\[
E_d = \frac{\mu_0}{2} N M^2
\]  

(5.25)

Consider the shape anisotropy of a small grain shaped like a prolate ellipsoid. When the spontaneous magnetization \( M_s \) is along the long axis of the ellipsoid (Fig. 5.12a), the opposing pole distributions are further away from each other and their surface density is lower than when \( M_s \) is parallel to the short axis (Fig. 5.12b). The demagnetizing field and energy are smallest when \( M_s \) is parallel to the long axis, which is the energetically favored direction of magnetization. The demagnetizing energy is larger in any other direction, giving a shape anisotropy. If \( N_1 \) is the demagnetizing factor for the long axis and \( N_2 \) that for the short axis \((N_1 < N_2)\), the difference in energy between the two directions of magnetization defines a magnetostatic anisotropy energy density given by

\[
E_a = \frac{\mu_0}{2}(N_2 - N_1) M^2
\]  

(5.26)

which is minimum when \( M_s \) is parallel to the longest dimension of the grain.

Shape-dependent magnetic anisotropy is important in minerals that have a high spontaneous magnetization. The more elongate the grain is, the higher the shape anisotropy will be. It is the predominant form of anisotropy in very fine grains of magnetite (and maghemite) if the longest axis exceeds the shortest axis by only about 20%.

### 5.2.7.3 Magnetostrictive anisotropy

The process of magnetizing some materials causes them to change shape. Within the crystal lattice the interaction energy between atomic magnetic moments depends on their separations (called the bond length) and on their orientations, i.e., on the direction of magnetization. If an applied field changes the orientations of the atomic magnetic moments so that the interaction energy is increased, the bond lengths adjust to reduce the total energy. This produces strains which result in a shape change of the ferromagnetic specimen. This phenomenon is called magnetostriction. A material which elongates in the direction of magnetization exhibits positive magnetostriction, while a material that shortens parallel to the magnetization shows negative magnetostriction. The maximum difference in magnetoelastic strain, which occurs between the demagnetized state and that of saturation magnetization, is called the saturation magnetostriction, denoted \( \lambda_s \).

The inverse effect is also possible. For example, if pressure is applied to one face of a cubic crystal, it will shorten elastically along the direction of the applied stress and will expand in directions perpendicular to it. These strains alter the separations of atomic magnetic moments, thereby perturbing the effects that give rise to magnetocrystalline anisotropy. Thus the application of stress to a magnetic material can change its magnetization; the effect is called piezomagnetism. On a submicroscopic scale the stress field that surrounds a vacancy, defect or dislocation in the crystal structure can locally affect the orientations of ionic magnetic spins.

Magnetostriction is a further source of anisotropy in magnetic minerals. The magnetostrictive (or magnetoelastic) anisotropy energy density \( E_a \) depends on the amount and direction of the stress \( \sigma \). If the saturation magnetization makes an angle \( \theta \) to the stress, \( E_a \) is given for a uniaxial magnetic mineral by

\[
E_a = \frac{3}{2} \lambda_s \sigma \cos^2 \theta
\]  

(5.27)

This is the simplest expression for magnetostrictive energy. It assumes that the magnetostriction is isotropic, i.e., that it has the same value in all directions. This condition is fulfilled if the magnetocrystalline axes of the ferromagnetic minerals in a rock are randomly distributed. The magnetoelastic energy of a cubic mineral is more complicated. Instead of a single magnetostriction constant \( \lambda_s \), separate constants \( \lambda_{100} \) and \( \lambda_{111} \) are required for the saturation magnetostriction along the [100] and [111] directions, respectively, corresponding to the edge and body diagonal directions of the cubic unit cell.

In magnetite the magnetoelastic energy is more than an order of magnitude less than the magnetocrystalline energy at room temperature. Consequently, magnetostriction plays only a secondary role in determining the direction of magnetization of magnetite grains. However, in minerals that have high magnetostriction (e.g., titanomagnetites (see Section 5.3.2.1) with a compositional factor \( x = 0.65 \)) the magnetoelastic energy may be significant in determining easy directions of magnetization, and the magnetization may be sensitive to modification by deformation.

### 5.3 ROCK MAGNETISM

#### 5.3.1 The magnetic properties of rocks

A rock may be regarded as a heterogeneous assemblage of minerals. The matrix minerals are mainly silicates or carbonates, which are diamagnetic in character. Interspersed in this matrix is a lesser quantity of secondary minerals (such as the clay minerals) that have paramagnetic properties. The bulk of the constituent minerals in a rock contribute to the magnetic susceptibility but are incapable of any contribution to the remanent magnetic properties, which are due to a dilute dispersion of ferrimagnetic minerals (e.g., commonly less than 0.01% in a limestone). The variable concentrations of ferrimagnetic and matrix minerals result in a wide range of susceptibilities in rocks (Fig. 5.13).

The weak and variable concentration of ferrimagnetic minerals plays a key role in determining the magnetic properties of the rock that are significant geologically and geophysically. The most important factors influencing
5.3.2 The ternary oxide system of magnetic minerals

The most important magnetic minerals are iron–titanium oxides, which are naturally occurring ferrites. The mineral structure consists of a close-packed lattice of oxygen ions, in which some of the interstitial spaces are occupied by regular arrays of ferrous (Fe\(^{2+}\)) and ferric (Fe\(^{3+}\)) iron ions and titanium (Ti\(^{4+}\)) ions. The relative proportions of these three ions determine the ferrimagnetic properties of the mineral. The composition of an iron–titanium oxide mineral can be illustrated graphically on the ternary oxide diagram (Fig. 5.14), the corners of which represent the end members of the solid-solution series are magnetite (Fe\(_3\)O\(_4\)), which is a typical strongly magnetic ferrite, and ulvöspinel (Fe\(_2\)TiO\(_4\)), which is antiferromagnetic at very low temperature but is paramagnetic at room temperature. The compositional parameter \(x\) expresses the relative proportion of titanium in the unit cell. The end members of the solid-solution series are magnetite (Fe\(_3\)O\(_4\)), which is a typical strongly magnetic ferrite, and ulvöspinel (Fe\(_2\)TiO\(_4\)), which is antiferromagnetic at very low temperature but is paramagnetic at room temperature.

An alternative form of the general formula is \(x\)Fe\(_2\)TiO\(_4\)(1 – \(x\))FeO\(_2\). Written in this way, it is apparent that the compositional parameter \(x\) describes the molecular fraction of ulvöspinel. As the amount of titanium \((x)\) increases, the cell size increases and the Curie temperature \(\theta\) and spontaneous magnetization \(M_s\) of the titanomagnetite decrease (Fig. 5.15).

Magnetite is one of the most important ferrimagnetic minerals. It has a strong spontaneous magnetization \((M_s = 4.8 \times 10^5 \text{ A m}^{-1})\) and a Curie temperature of 578 °C. Because of the high value of \(M_s\), magnetite grains can have a strong shape anisotropy. The magnetic susceptibility is the strongest of any naturally occurring mineral.

Fig. 5.14 Ternary compositional diagram of the iron–titanium oxide solid solution magnetic minerals (after McElhinny, 1973).

Fig. 5.13 (a) Median values and ranges of the magnetic susceptibility of some common rock types, and (b) the susceptibilities of some important minerals.
5.3 ROCK MAGNETISM

Fig. 5.15 Variations of Curie temperature and unit-cell size with composition in titanomagnetite (after Nagata, 1961).

(k = 1–10SI). For many sedimentary and igneous rocks the magnetic susceptibility is proportional to the magnetite content.

Maghemite (γ-Fe₂O₃) can be produced by low-temperature oxidation of magnetite. It is a strongly magnetic mineral (Mᵣ = 4.5 × 10⁵ A m⁻¹). Experiments on maghemite doped with small amounts of foreign ions indicate that it has a Curie temperature of 675 °C. However, it is metastable and reverts to hematite (α-Fe₂O₃) when heated above 300–350 °C. The low-temperature oxidation of titanomagnetite leads to a “titanomaghemite” solid-solution series.

Titanomagnetite is responsible for the magnetic properties of oceanic basalts. The basaltic layer of the oceanic crust is the main origin of the marine magnetic anomalies that are of vital importance to modern plate tectonic theory. The magnetic properties of the 0.5 km thick basaltic layer are due to the presence of very fine grained titanomagnetite (or titanomaghemite, depending on the degree of ocean-floor weathering). The molecular fraction (x) of Fe₂TiO₄ in titanomagnetite in oceanic basalts is commonly around 0.6.

5.3.2.2 The titanohematite series

The minerals of the titanohematite solid-solution series are also variously referred to as “hemoilmenite,” “hematite-ilmenite” or “ilmenohematite.” They have the general formula Feₓ₂₋ₓTi₁₋ₓO₄. The unit cell has rhombohedral symmetry. Ionic substitution is the same as for titanomagnetite, and the compositional parameter x has the same implications for the titanium content of the unit cell. The end members of the solid-solution series are hematite (Fe₂O₃) and ilmenite (FeTiO₃). The chemical formula can be written in the alternative form xFeTiO₃·(1–x)Fe₂O₃, where x represents the molecular fraction of ilmenite. As in the case of titanomagnetite, the cell size increases and the Curie point decreases as the titanium content increases. The Curie point of hematite is 675 °C, while ilmenite is antiferromagnetic at low temperature and paramagnetic at room temperature. For titanium contents 0.5 < x < 0.95 titanohematite is ferrimagnetic and for x < 0.5 it exhibits parasitic ferromagnetism.

The end member hematite (α-Fe₂O₃) is an extremely important magnetic mineral. Its magnetic properties arise from parasitic ferromagnetism due to the spin-canted magnetic moment and the possible defect moment of its otherwise antiferromagnetic lattice. Hematite has a weak spontaneous magnetization (M₀ = 2.2 × 10³ A m⁻¹) and a strong magnetocrystalline anisotropy (Kₐ = 10³ J m⁻³). Hematite is paleomagnetically important because of its common occurrence and its high magnetic and chemical stability. It often occurs as a secondary mineral, formed by oxidation of a precursor mineral, such as magnetite, or by precipitation from fluids passing through a rock.

5.3.3 Other ferrimagnetic minerals

Although the iron–titanium oxides are the dominant magnetic minerals, rocks frequently contain other minerals with ferromagnetic properties. Although pyrite (FeS₂) is a very common sulfide mineral, especially in sedimentary rocks, it is paramagnetic and therefore cannot carry a remanent magnetization. As a result it does not contribute directly to the paleomagnetic properties of rocks, but it may act as a source for the formation of goethite or secondary magnetite.

Pyrrhotite is a common sulfide mineral which can form authigenically or during diagenesis in sediments, and which can be ferrimagnetic in certain compositional ranges. It is non-stoichiometric (i.e., the numbers of anions and cations in the unit cell are unequal) and has the formula Fe₁₋ₓS. The parameter x refers to the proportion of vacancies among the cation lattice sites and is limited to the range 0 < x < 0.14. Pyrrhotite has a pseudo-hexagonal crystal structure and would be antiferromagnetic but for the presence of the cation vacancies. The Néel temperature at which the fundamental antiferromagnetism disappears is around 320 °C. Pyrrhotite with the formula Fe₀.95S is ferrimagnetic with a Curie temperature close to the Néel temperature and a strong spontaneous magnetization of about 10⁵ Am⁻¹ at room temperature. The magnetocrystalline anisotropy restricts the easy axis of magnetization to the hexagonal basal plane at room temperature.

The iron oxyhydroxide goethite (FeOOH) is another common authigenic mineral in sediments. Like hematite, goethite is antiferromagnetic, but has a weak parasitic ferromagnetism. It has a very high coercivity (with maximum values in excess of 5 T) and a low Curie point around 100 °C or less. It is thermally unstable relative to hematite under most natural conditions, and decomposes on heating above about 350 °C. It is a common (and paleomagnetically undesirable) secondary mineral in limestones and other sedimentary rocks.
5.3.4 Identification of ferrimagnetic minerals

It is often difficult to identify the ferrimagnetic minerals in a rock, because their concentration is so low, especially in sedimentary rocks. If the rock is coarse grained, ferrimagnetic minerals may be identified optically among the opaque grains by studying polished sections in reflected light. However, in many rocks that are paleomagnetically important (e.g., basaltic lava, pelagic limestone) optical examination may be unable to resolve the very fine grain size of the ferrimagnetic mineral. The ferrimagnetic mineral fraction may then be identified by its properties of Curie temperature and coercivity.

The Curie temperature is measured using a form of balance in which a strong magnetic field gradient exerts a force on the sample that is proportional to its magnetization. The field (usually 0.4–1 T) is strong enough to saturate the magnetization of many minerals. The sample is heated and the change of magnetic force (i.e., sample magnetization) is observed with increasing temperature. When the Curie point is reached, the ferromagnetic behavior disappears; at higher temperatures the sample is paramagnetic. The Curie point is diagnostic of many minerals. For example, an extract of magnetic minerals from a pelagic limestone (Fig. 5.16) shows the presence of goethite (e.g., in hematite the magnetocrystalline anisotropy constant of these minerals.

5.3.5 Grain size dependence of ferrimagnetic properties

The ferromagnetic properties of metals and ferrites vary sensitively with grain size. Consider an assemblage of uniformly magnetized grains of a ferrimagnetic mineral characterized by a spontaneous magnetization $M_0$ and uniform grain volume $v$. Let the spontaneous magnetization be oriented parallel to an easy direction of magnetization (crystalline or shape determined), defined by the anisotropy energy $K_u$ per unit volume. The energy that keeps the magnetization parallel to the easy direction is equal to $vK_u$.

Thermal energy, proportional to the temperature, has the effect of disturbing this alignment. At temperature $T$ the thermal energy of a grain is equal to $kT$, where $k$ is Boltzmann’s constant ($k = 1.381 \times 10^{-23}$ J K$^{-1}$). At any instant in time there is a chance that thermal energy will deflect the magnetic moment of a grain away from its easy direction. Progressively, the net magnetization of the material (the sum of all the magnetic moments of the numerous magnetic grains) will be randomized by the thermal energy, and the magnetization will be observed to decay. If the initial magnetization of the assemblage is $M_{0r}$, after time $t$ it will decrease exponentially to $M(t)$, according to

$$M(t) = M_{0r}e^{-t/\tau}$$

(5.28)

In this equation $\tau$ is known as the relaxation time of the grain (Box 5.1). If the relaxation time is long, the exponential decrease in Eq. (5.28) is slow and the magnetization is stable. The parameter $\tau$ depends on properties of the grain and is given by the equation

$$\tau = \frac{1}{\nu_0}\exp\left(\frac{vK_u}{kT}\right)$$

(5.29)

The constant $\nu_0$ is related to the lattice vibrational frequency and has a very large value ($\approx 10^8–10^{10}$ s$^{-1}$). The value of $K_u$ depends on whether the easy direction of the magnetic mineral is determined by the magnetocrystalline anisotropy or the magnetostatic (shape) anisotropy. For example, in hematite the magnetocrystalline anisotropy prevails because the spontaneous magnetization is very weak, and $K_u$ is equal to the magnetocrystalline...
Box 5.1: Magnetic relaxation

Relaxation behavior is characterized by the exponential return of a physical property with time from a state of elevated energy to a state of lower energy. The most familiar example is radioactive decay (Section 4.1.3.1), but the magnetizations of natural materials also exhibit relaxation behavior, as explained in Section 5.3.5.

In the absence of an external field, the easy direction of magnetization of a single domain magnetic grain with volume \( v \) and anisotropy energy \( K_u \) per unit volume is determined by the anisotropy energy \( vK_u \) (Section 5.2.7). The probability that the grain’s thermal energy \( kT \) can overcome this energy barrier and allow the grain magnetization to change direction is determined by the Maxwell-Boltzmann distribution of energies, and is proportional to \( \exp(-vK_u/kT) \). The probability per unit time, \( \lambda \), of a magnetic moment changing to a different easy axis is given by the Arrhenius equation for a thermally activated process

\[
\lambda = C \exp \left( -\frac{vK_u}{kT} \right)
\]  

(1)

The parameter \( C \) is called the frequency factor; here it is the lattice vibration frequency, \( \nu_o \). At a particular temperature, all parameters on the right side of Eq. (1) are constant and so the probability \( \lambda \) per unit time of a magnetic moment changing to a different easy axis is constant.

Now suppose an assemblage of identical non-interacting single domain particles, of which \( N_1 \) are magnetized initially in one direction (state 1) and \( N_2 \) in the opposite direction (state 2). The net magnetization is proportional to \((N_1 - N_2)\). Assuming a constant probability \( \lambda \) per unit time, the number of particles \( dN_1 \) that change from state 1 to state 2 is proportional to the time interval \( dt \) and to the number of grains \( N_1 \) in state 1. It is given by

\[
dN_1 = -\lambda N_1 dt
\]  

(2)

where the negative sign indicates a reduction in \( N_1 \). Similarly, the number changing in the opposite sense from state 2 to state 1 is

\[
dN_2 = -\lambda N_2 dt
\]  

(3)

The net change in magnetization is therefore

\[
dM = \mu dN_1 - \mu dN_2 = \mu d(N_1 - N_2)
\]  

(4)

\[
dM = -\lambda \mu (N_1 - N_2) dt = -\lambda M dt
\]  

(5)

The solution of this differential equation is

\[
M = M_o \exp(-\lambda t) = M_o \exp \left( -\frac{t}{\tau} \right)
\]  

(6)

where \( \tau \), the relaxation time, is the inverse of \( \lambda \) in Eq. (1):

\[
\tau = \frac{1}{\lambda} = \left( \frac{vK_u}{kT} \right)
\]  

(7)

Equations (6) and (7) are very important in paleomagnetism. The strong anisotropy of fine grained ferromagnetic minerals can result in very long relaxation times, and consequently the magnetizations of fine grained rocks can be extremely stable over geological lengths of time.

5.3.5.1 Superparamagnetism

In a ferromagnetic material the strong molecular fields keep the atomic spin magnetic moments uniformly aligned with each other, and the grain anisotropy requires this spontaneous magnetization to lie parallel to an “easy” direction. If the temperature is too high, thermal energy \((kT)\) may exceed the anisotropy energy \((vK_u)\) but still be too small to break up the spontaneous magnetization. The thermal energy causes the entire magnetic moment of the grain to fluctuate coherently in a manner similar to paramagnetism (the theory of which applies to individual atomic magnetic moments). The grain magnetization has no stable direction, and the behavior is said to be superparamagnetic. It is important to note that superparamagnetic grains themselves are immobile; only their uniform magnetization fluctuates relative to the grain. Whether the ferrimagnetic grain exists in a stable or superparamagnetic state depends on the grain size, the grain shape (if the origin of \( K_u \) is magnetostatic) and the temperature. If the grain volume \( v \) is very small, unstable magnetic behavior due to superparamagnetism becomes likely. Magnetite and hematite grains finer than about
5.3.5.2 Single domain particles

When the anisotropic magnetic energy ($\nu K_u$) of a grain is greater than the thermal energy ($kT$), the spontaneous magnetization direction favors one of the easy directions. The entire grain is uniformly magnetized as a single domain. This situation occurs in very fine grains of ferromagnetic minerals.

In magnetite $K_u$ is the magnetostatic energy related to the particle shape. The theoretical range of single domain sizes in magnetite is narrow, from about 0.03 to 0.1 µm in equant grains and up to about 1 µm in elongate grains (Fig. 5.17). In hematite $K_u$ is the large magnetocrystalline anisotropy energy, and the range of single domain sizes is larger, from about 0.03 to 15 µm.

The magnetization of a single domain particle is very stable, because to change it requires rotating the entire uniform spontaneous magnetization of the grain against the grain anisotropy, which requires a very strong magnetic field. The magnetic field required to reverse the direction of magnetization of a single domain grain is called its coercivity $B_c$ and is given by:

$$B_c = \frac{2K_u}{M_s} \quad (5.30)$$

The maximum coercivity of single domain magnetite is around 0.3 T for needle-shaped elongate grains. The magnetocrystalline anisotropy of hematite gives it higher maximum coercivities, in excess of 0.5 T. However, the magnetic properties of hematite are very variable and its maximum coercivity commonly exceeds 2 T. Because of their stable remanent magnetizations, single domain particles play a very important role in paleomagnetism.

5.3.5.3 Multidomain particles

Single domain behavior is restricted to a limited range of grain sizes. When a grain is large enough, the magnetic energy associated with its magnetization becomes too large for the magnetization to remain uniform. This is because the demagnetizing field of a uniformly magnetized grain (Fig. 5.18a) interacts with the spontaneous magnetization and generates a magnetostatic (or self-demagnetizing) energy. To reduce this energy, the magnetization subdivides into smaller, uniformly magnetized units, called Weiss domains after P. Weiss, who theoretically predicted domain structure in 1907. In the simplest case the magnetization divides into two, oppositely magnetized domains (Fig. 5.18b). The net magnetization is reduced to zero, and the magnetostatic energy is reduced by about a half. Further subdivision (Fig. 5.18c) reduces the magnetostatic energy correspondingly. In a grain with $n$ domains of alternately opposed spontaneous magnetizations the magnetostatic energy is reduced by a factor of
5.3 ROCK MAGNETISM

The domains are separated from one another by thin regions, about 0.1 \( \mu \text{m} \) thick, that are usually much thinner than the domains they divide. These regions are called Bloch domain walls in recognition of F. Bloch, who in 1932 proposed a theory for the structure of the domain wall on an atomic scale. Within the domain wall the magnetization undergoes small progressive changes in direction from each atom to its neighbor. The crystalline magnetic anisotropy of the material attempts to keep the atomic magnetic spins parallel to favored crystalline directions, while the exchange energy resists any change of direction from parallel alignment by the molecular field. The energy of these competing effects is expressed as a domain wall energy associated with each unit area of the wall.

The magnetization of a multidomain grain can be changed by moving the position of a domain wall, which causes some domains to increase in size and others to decrease. A large multidomain grain may contain many easily movable domain walls. Consequently, it is much easier to change the magnetization of a multidomain grain than of a single domain grain. As a result multidomain grains are less stable carriers of remanent magnetization than single domain grains.

The transition between single domain and multidomain behavior occurs when the reduction of magnetostatic energy is balanced by the energy associated with the domain wall that has been added. If magnetite grains are elongate, they can persist as single domain grains up to about 1 \( \mu \text{m} \) (Fig. 5.17). Magnetite grains larger than a few micrometers in diameter are probably multidomain. In equidimensional grains of magnetite the transition should occur in grains of about 0.05–0.1 \( \mu \text{m} \) diameter. However, at this point the grain is not physically large enough to contain a wall, which has a thickness of about 0.1 \( \mu \text{m} \). Grains in the intermediate range of sizes, and those large enough to contain only a few walls, are said to carry a pseudo-single domain magnetization. True multidomain behavior in magnetite is observed when the grain size exceeds 15–20 \( \mu \text{m} \).

In a pseudo-single domain grain that is large enough to contain only two domains, the domain wall separating them is not able to move freely. Its freedom of movement is restricted by interactions with the grain surface. A small grain in this size range has more stable magnetic properties than a multi domain particle but is not as stable as a true single domain grain. Magnetite grains between about 0.1 m and several micrometers in diameter have pseudo-single domain properties.

5.3.6 Remanent magnetizations in rocks

The small concentration of ferrimagnetic minerals in a rock gives it the properties of magnetic hysteresis. Most important of these is the ability to acquire a remanent magnetization (or remanence). The untreated remanence of a rock is called its natural remanent magnetization (NRM). It may be made up of several components acquired in different ways and at different times. The geologically important types of remanence are acquired at known times in the rock’s history, such as at the time of its formation or subsequent alteration. The remanence of a rock can be very stable against change; the high coercivity (especially of the fine grains) of the ferrimagnetic mineral assures preservation of the magnetic signal during long geological epochs.

A remanence acquired at or close to the time of formation of the rock is called a primary magnetization; a remanence acquired at a later time is called a secondary magnetization. Examples of primary remanence are thermoremanent magnetization, which an igneous rock acquires during cooling, and the remanent magnetizations acquired by a sediment during or soon after deposition. Secondary remanences may be caused by chemical change of the rock during diagenesis or weathering, or by sampling and laboratory procedures.

5.3.6.1 Thermoremanent magnetization

The most important type of remanent magnetization in igneous (and high-grade metamorphic) rocks is thermoremanent magnetization (TRM). Igneous rocks solidify at temperatures well above 1000 °C. At this temperature the grains are solid and fixed in a rigid matrix. The grains of a ferrimagnetic mineral are well above their Curie temperature, which in magnetite is 578 °C and in hematite is 675 °C. There is no molecular field and the individual atomic magnetic moments are free to fluctuate chaotically; the magnetization is paramagnetic (Fig. 5.19).

As the rock cools, the temperature eventually passes below the Curie temperature of the ferrimagnetic grains and a spontaneous magnetization appears. In single domain grains the relaxation time of the grain magnetization is governed by Eq. (5.29), which can be modified by writing the anisotropy energy density \( K_u \) in terms of the spontaneous magnetization \( M_s \) and coercivity \( B_c \) of the grain. This gives the relaxation time as follows

\[
\tau = \frac{1}{v_0} \exp \left( \frac{v M_s B_c}{2kT} \right)
\]

(5.31)

At high temperature the thermal energy \( (kT) \) is larger than the magnetic energy \( (vM_s B_c/2) \) and the magnetization is unstable. Although the individual atomic magnetic moments are forced by the molecular field to act as coherent units, the grain magnetizations are superparamagnetic. As the rock cools further, the spontaneous magnetization and the magnetic anisotropy energy \( K_u \) of the grain increase. Eventually the temperature passes below a value at which the thermal energy, whose effect is to randomize the grain magnetic moments, is no longer greater than the magnetic anisotropy energy. The spontaneous magnetization then becomes “blocked” along an easy direction of magnetization of the grain. In the absence of an external magnetic field the grain...
magnetic moments will be randomly oriented (assuming the easy axes to be randomly distributed). If the grain cools below its blocking temperature in a magnetic field, the grain magnetic moment is “blocked” along the easy axis that is closest to the direction of the field at that time (Fig. 5.19). The alignment of grain magnetic moments with the field is neither perfect nor complete; it represents a statistical preference. This means that in an assemblage of grains more of the grains have their magnetic moments aligned close to the field direction than any other direction. The degree of alignment depends on the strength of the field.

It is important to note that the mineral grains themselves are immobile throughout the process of acquisition of TRM. Only the internal magnetizations of the grains can change direction and eventually become blocked. The blocking temperatures of TRM are dependent on the grain size, grain shape, spontaneous magnetization and magnetic anisotropy of the ferrimagnetic mineral. If the rock contains a wide range of grain sizes and perhaps more than a single magnetic mineral, there may be a broad spectrum of blocking temperatures. Maximum blocking temperatures may range as high as the Curie point, and the spectrum can extend to below ambient temperature. The pressure of overlying sediment during deep burial results in compaction, which can produce further directional errors. The DRM is finally fixed in sedimentary rocks during diagenesis.

A modified form of post-depositional remanence (pDRM) is important in fine grained sediments. A water-logged slurry forms at the sediment–water interface. Fine grained magnetic minerals in the water-filled pore spaces in the sediment are subject to mechanical forces caused by the water flow. These forces can cause the magnetic particles to rotate, leading to a declination error. The particle then rolls into a stable attitude, causing an inclination error. The pressure of overlying sediment during deep burial results in compaction, which can produce further directional errors. The DRM is finally fixed in sedimentary rocks during diagenesis.

TRM is a very stable magnetization which can exist unchanged for long intervals of geological time. The ability of TRM to record accurately the field direction is demonstrated by the results from a lava that erupted on Mt. Etna at a time for which a record of the magnetic field direction is available from observatory data. The directions of the TRM in the lava samples are the same as the direction of the ambient field (Fig. 5.20).

5.3.6.2 Sedimentary remanent magnetizations

The acquisition of depositional remanent magnetization (DRM) during deposition of a sediment takes place at constant temperature. Magnetic and mechanical forces compete to produce a physical alignment of detrital ferrimagnetic particles. During settling through still water these particles are oriented by the ambient magnetic field in the same way that it orients a compass needle. The particles become aligned statistically with the Earth’s magnetic field (Fig. 5.21). The action of mechanical forces may at times spoil this alignment. Water currents cause hydromechanical forces that disturb the alignment during settling, giving rise to a declination error. On contact with the bottom of the sedimentary basin, the mechanical force of gravity rolls the particle into a stable attitude, causing an inclination error. The pressure of overlying sediment during deep burial results in compaction, which can produce further directional errors. The DRM is finally fixed in sedimentary rocks during diagenesis.
the sediment are partially in suspension and may be reoriented by the magnetic field, if they are free enough to move (Fig. 5.22a). The energy for this motion of the particles is obtained from the Brownian motion of the water molecules, which continuously and randomly collide with the particles in the pore spaces. Large particles are probably in contact with the surrounding grains and are unaffected by collisions with the water molecules. Very fine particles that are virtually floating in the pore spaces may acquire enough freedom from the Brownian agitation to align statistically with the Earth's magnetic field. Laboratory experiments have established that the direction of pDRM is an accurate record of the depositional field, without inclination error (Fig. 5.22b). The pDRM is acquired later than the actual time of sedimentation, and is fixed in the sediment during compaction and de-watering at a depth of ~10 cm. This may represent a lock-in time delay of 100 yr in lacustrine sediments or 10,000 yr in pelagic marine sediments, where it is no more important geologically than the errors involved in locating paleontological stage boundaries. The pDRM process is particularly effective in fine grained sediments containing strongly magnetic magnetite grains. For example, pDRM is the most important mechanism of primary magnetization in pelagic limestones. Compaction may cause a flattening of the inclination under some conditions.

Bioturbation may mix the sediment, typically to a depth of about 10 cm in pelagic sediments. This affects the positions of stratigraphic marker levels. First occurrence datum levels of fossils are carried deeper, and last occurrences are carried higher than the true stratigraphic levels. Agitation of bioturbated sediment by the burrowing organisms assists the Brownian motion of the magnetic particles. Under these conditions the pDRM is acquired at the base of the bioturbated zone.

5.3.6.3 Chemical remanent magnetization

Chemical remanent magnetization (CRM) is usually a secondary form of remanence in a rock. It occurs when the magnetic minerals in a rock suffer chemical alteration or when new minerals form authigenically. An example is the precipitation of hematite from a goethite precursor or from iron-saturated fluids that pass through the rock. The magnetic minerals may also experience diagenetic modification or oxidation by weathering, which usually happens on the grain surface and along cracks (Fig. 5.23). The growth of a new mineral (or the alteration of an existing one) involves changes in grain volume $v$, spontaneous
magnetization $M_s$ and coercivity $B_c$. The chemical change affects the relaxation time of the grain magnetization, according to Eq. (5.31). The grains eventually grow through a critical volume, at which the grain magnetization becomes blocked. The new CRM is acquired in the direction of the ambient field during the chemical change, and so it is usually a secondary remanence. It has stable magnetic properties similar to TRM. A common example is the formation of hematite during diagenesis or weathering. Hematite that originates in this way typically carries a secondary remanent magnetization.

5.3.6.4 Isothermal remanent magnetization

Isothermal remanent magnetization (IRM) is induced in a rock sample by placing it in a magnetic field at constant temperature. For example, rock samples are exposed to the magnetic fields of the sampling equipment and to other magnetic fields during transport to the laboratory. A common technique of rock magnetic analysis consists of deliberately inducing IRM via a known magnetic field produced in a large coil or between the poles of an electromagnet. The magnetic moments within each grain are partially aligned by the applied field. The degree of alignment depends on the field strength and on the resistance of the magnetic mineral to being magnetized, loosely referred to as its coercivity. After removing the sample from the applied field an IRM remains in the rock (see Fig. 5.11. If the rock sample is placed in progressively stronger fields the IRM increases to a maximum value called the saturation IRM, which is determined by the type and concentration of the magnetic mineral. The shape of the progressive acquisition curve and the field needed to reach saturation IRM depend on the coercivities of the magnetic minerals in the rock (Fig. 5.24).

The maximum coercivities of the most common ferromagnetic minerals in rocks are fairly well known (Table 5.1). These minerals also have distinctive maximum blocking temperatures (Section 5.3.6.1). The combination of these properties provides a method of identification of the predominant magnetic minerals in rocks. Starting with the strongest field available, IRM is imparted in successively smaller fields, chosen to remagnetize different coercivity fractions, along two or three orthogonal directions. The compound IRM is then subjected to progressive thermal demagnetization. The demagnetization characteristics of

<table>
<thead>
<tr>
<th>Ferromagnetic mineral</th>
<th>Maximum coercivity $T$</th>
<th>Maximum blocking temperature $C$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Magnetite</td>
<td>0.3</td>
<td>575</td>
</tr>
<tr>
<td>Maghemite</td>
<td>0.3</td>
<td>≈350</td>
</tr>
<tr>
<td>Titanomagnetite ($Fe_{3-x}Ti_xO_4$):</td>
<td>$x = 0.3$</td>
<td>0.2</td>
</tr>
<tr>
<td></td>
<td>$x = 0.6$</td>
<td>0.1</td>
</tr>
<tr>
<td>Pyrrhotite</td>
<td>0.5–1</td>
<td>325</td>
</tr>
<tr>
<td>Hematite</td>
<td>1.5–5</td>
<td>675</td>
</tr>
<tr>
<td>Goethite</td>
<td>&gt; 5</td>
<td>80–120</td>
</tr>
</tbody>
</table>

Table 5.1 Maximum coercivities and blocking temperatures for some common ferromagnetic minerals

Fig. 5.23 Acquisition of chemical remanent magnetization (CRM) accompanies the diagenetic modification or oxidation by weathering of magnetic minerals; this often happens on the grain surface and along cracks.

Fig. 5.24 Examples of the identification of magnetic minerals by acquisition and subsequent thermal demagnetization of IRM. Hematite is present in both (a) and (b), because saturation IRM requires fields $> 1$ T and thermal demagnetization of the hard fraction persists to $T = 675 \, ^\circ C$. In (a) the soft fraction that demagnetizes at $T = 575 \, ^\circ C$ is magnetite, while in (b) no magnetite is indicated but pyrrhotite is present in all three fractions, shown by thermal unblocking at $T = 300–330 \, ^\circ C$ (after Lowrie, 1990).
5.3 ROCK MAGNETISM

5.3.6.5 Other remanent magnetizations

If a rock containing magnetic minerals with unstable magnetic moments experiences a magnetic field, there is a finite probability that some magnetic moments opposite to the field may switch direction to be parallel to the field. As time goes on, the number of magnetic moments in the direction of the field increases and the magnetization grows logarithmically with time (Fig. 5.25). The time-dependent remanence acquired in this way is called a viscous remanent magnetization (VRM). The VRM also decreases logarithmically when the field is removed or changed and is often identifiable during laboratory analysis as an unstable time-dependent change in remanence. The direction of a VRM is often parallel to the present-day field direction, which can be useful in its identification. However, it is always a secondary remanence and it can mask the presence of geologically interesting stable components. Techniques of progressive demagnetization have been designed to remove VRM and IRM, effectively “cleaning” the remanence of a rock of undesirable components.

An important form of remanent magnetization can be produced in a rock sample by placing it in a coil that carries an alternating magnetic field, whose amplitude is then slowly reduced to zero. In the absence of another field, this procedure randomizes the orientations of grain magnetic moments with coercivities less than the peak field. If, however, the rock sample is exposed to a small constant magnetic field while the amplitude of the alternating magnetic field is decreasing to zero, the magnetic moments are not randomized. Their distribution is biased with a statistical preference for the direction of the constant field. This produces an anhysteretic remanent magnetization (ARM) in the sample. The intensity of ARM increases with the amplitude of the alternating field, and also with the strength of the constant bias field. ARM may be produced deliberately, as described, and it is commonly observed as a spurious effect during progressive alternating field demagnetization of rock samples when the shielding from external fields is imperfect (Section 5.6.3.2).

Stress and the associated strains of magnetostriction combine to produce magnetoelastic energy (see Section 5.2.7.3). This acts as a source of magnetic anisotropy that can modify the easy directions of magnetization in some magnetic minerals. As a result, stress caused by tectonic deformation or by defects in the crystal structure of constituent minerals may influence the direction of magnetization in a rock. The effect of high hydrostatic pressure, such as might be encountered at depth in the Earth’s crust, is to deflect the magnetizations of individual grains away from existing easy axes, which leads to a net demagnetization of the rock. On the other hand, the effect of non-hydrostatic stress in the presence of a magnetic field is to produce a piezoremanent magnetization (PRM) that is added to, or may replace, any pre-existing remanent magnetization. The modification of rock magnetization by deformational stress, whether elastic or plastic, can overprint the original magnetization in rocks that have been deeply buried, or subjected to high stress and temperature during deformation.

A related type of magnetization has been observed in terrestrial and lunar rocks that have been shocked by meteoritic impact. The event causes very high local stress of short duration, and gives rise to a shock remanent magnetization (SRM) by a similar mechanism to PRM. The collision of a meteorite with another planetary body imposes a characteristic shock texture on the impacted rocks, so that SRM and PRM are not exactly equivalent. Moreover, much of the energy of the collision is released as heat, which can modify and reduce the SRM over a much longer time than the brief duration of the impact. However, SRM may have played a role in the origin of anomalous crustal magnetizations on the Moon, Mars and other extraterrestrial bodies that have suffered extensive meteoritic bombardment.

5.3.7 Environmental magnetism

Since the mid-1980s rock magnetic properties have found important new applications in environmental research, where they can serve as tracers for pollution or as indicators of past climates. Heavy metals from industrial processes enter the environment, contaminating both the ground surface and the water in lakes, rivers and the groundwater. Magnetic minerals such as magnetite and hematite accompany more toxic elements in the emissions. Magnetic susceptibility surveys can provide a rapid method of determining the geographic dispersal and relative concentration of the heavy metal pollution. Motor-vehicle exhaust emissions also pollute the environment by loading it with heavy metals as well as sub-micrometer sized particles of soot, a fraction of which consists of nanoparticle sized magnetic minerals. Rock magnetic parameters and investigative techniques provide ways of...
characterizing these particles and of describing their regional distribution and concentration.

The magnetic grain fraction in a sediment or soil usually consists of hematite, magnetite, maghemite or an iron sulfide. The magnetic mineral composition may be modified by the climatic conditions at or after deposition. These affect magnetic properties of the sediment that depend on grain size and mineral composition, such as magnetic susceptibility, coercivity and the various remanent magnetizations. These magnetic parameters can act as proxies for paleoclimatic change. Numerous studies have been devoted to understanding the processes involved and their application. The following example illustrates the potential of magnetic methods for analyzing a paleoclimatic problem.

Great thicknesses of loess sediments up to 100 m in thickness occur in Central China. The loess are very fine grained (10–50 μm size) wind-blown sediments that are deposited in cold, dry conditions. Alternating with the beds of loess are layers of paleosol, the name given to fossil soils. These form by conversion of the loess to soils (a process called pedogenesis), which takes place during interglacial periods of warmer, humid conditions. In turn, later loess deposits bury the soils. The alternation of loess and paleosol is therefore a record of past climatic variations. The magnetic susceptibility correlates with the lithology of the loess–paleosol sequences. In the loess beds the susceptibility measures around 25 SI units, but in the paleosol layers the values exceed 200 SI units (Fig. 5.26). Rock magnetic analysis showed that the dominant magnetic mineral in both lithologies is magnetite. The higher susceptibility of the paleosols is thus due to higher concentrations of magnetite. The susceptibility variation correlates very well with the oxygen isotope stratigraphy measured in deep-sea sediments at site 677 of the Ocean Drilling Program (ODP). The oxygen isotope ratio (see Box 5.2) is a record of climatic variation, with the most negative values corresponding to warmer temperature, and it has been dated by comparison to magnetostratigraphy in marine sediments (Section 5.7.2). The correlation in Fig. 5.26 provides a way of dating the loess–paleosol sequence and shows that the paleosols were formed under warmer conditions than the loess. Possibly the change to a warmer, humid climate encouraged the in situ formation of a new phase of magnetite in the paleosols, with a consequent increase in susceptibility.

The new phase of magnetite in the paleosols originates by a chemical process. However, magnetite can be produced by the action of bacteria in sediments deposited in lakes and in the sea. This is referred to as a biogenic process and the bacteria that can produce magnetite are called magnetotactic bacteria. The magnetite forms as tiny crystallites called magnetosomes, less than 0.1 μm in size and thus of single domain type, which occur as chains of particles enclosed in a membrane. The most common magnetosome mineral is magnetite, but greigite, an iron sulfide with structure similar to magnetite, has also been found. The chain-like assembly imparts a dipole magnetic moment to the bacteria. This is evidently an evolutionary feature, which helps the bacteria to survive. If the sediment is disturbed, the magnetic moment of the bacteria enables them to move along the Earth’s magnetic field lines. The dipping field lines guide them back down into the sediment where they find nutrition needed for their survival. It was long believed that magnetite in deep-sea sediments was mainly of detrital origin, washed in from the continents and ocean ridges, but it is now clear that a large portion must be biomagnetic.

Magnetite of biogenic origin has been identified in many unusual settings. For example, submicroscopic magnetite occurs in the brains of dolphins and birds; whether it plays a role in their guidance during migration is unclear. Magnetite of nanometer size has been located in the human brain, where it may be related to neurological disorders such as epilepsy, Alzheimer’s disease and Parkinson’s disease. It also occurs in other human organs. The magnetic properties
5.4 GEOMAGNETISM

5.4.1 Introduction

The magnetic field of the Earth is a vector, that is, it has both magnitude and direction. The magnitude, or intensity $F$, of the field is measured in the same units as other B-fields, namely in tesla (see Eq. (5.10)). However, a tesla is an extremely strong magnetic field, such as one would observe between the poles of a powerful electromagnet. The Earth's magnetic field is much weaker; its maximum intensity is reached near to the magnetic poles, where it amounts to about $6 \times 10^{-5}$ T. Modern instruments for measuring magnetic fields (called magnetometers) have a sensitivity of about $10^{-9}$ T; this unit is called a nanotesla (nT) and has been adopted in geophysics as the practical unit for expressing the intensity of geomagnetic field intensity. There is a practical reason for adopting this unit. Most geomagnetic surveys carried out until the 1970s used the now abandoned c.g.s. system of units, in which the $B$-field was measured in gauss, equivalent to $10^{-4}$ T. The practical unit of geophysical exploration was then $10^{-5}$ gauss, called a gamma ($\gamma$). Thus, the former unit ($\gamma$) is conveniently equal to $10^{-9}$ T, which is the new unit (nT).

The magnetic vector can be expressed as Cartesian components parallel to any three orthogonal axes. The geomagnetic elements are taken to be components parallel to the geographic north and east directions and the vertically downward direction (Fig. 5.27). Alternatively, the

Box 5.2: Oxygen isotope ratio

The element oxygen has two important stable isotopes. These are a “light” isotope $^{16}\text{O}$, with 8 protons and 8 neutrons in its nucleus, and a “heavy” isotope $^{18}\text{O}$ with 8 protons and 10 neutrons. Mass spectrometers are capable of determining accurately the masses of these isotopes, and so the mass ratio $^{18}\text{O}/^{16}\text{O}$ can be measured in quite small samples. In order to describe the deviation of the $^{18}\text{O}/^{16}\text{O}$ ratio in a given sample from a standard value, a useful parameter, $\delta^{18}\text{O}$, is defined as follows:

$$\delta^{18}\text{O} = 1000 \times \left( \frac{^{18}\text{O}^{16}\text{O}}{^{16}\text{O}^{18}\text{O}} \right)_{\text{standard}} - 1$$

(1)

The standard value is taken to be the $^{18}\text{O}/^{16}\text{O}$ ratio in the modern oceans at depths in the 200–500 m range, or alternatively the ratio in a fossil belemnite known as PDB. The factor 1000 causes $\delta^{18}\text{O}$ values to be expressed in parts per thousand. The global climate at present is relatively warmer than in the past, so $^{18}\text{O}/^{16}\text{O}$ ratios measured in ancient samples tend to be smaller than the standard value and give negative $\delta^{18}\text{O}$ values.

The $^{18}\text{O}/^{16}\text{O}$ ratio in water is dependent on temperature, as evident in the plot of $\delta^{18}\text{O}$ measured in the annual precipitation at a site against the mean temperature at the site (Fig. B5.2). Two factors determine this correlation: cool air holds less moisture than warm air, and the “heavy” isotope $^{18}\text{O}$ condenses more easily than the “light” isotope $^{16}\text{O}$. From an air mass that moves polewards, precipitation in warmer regions is relatively rich in $^{18}\text{O}$ whereas in colder regions it is enriched in $^{16}\text{O}$. During global warm periods, polar ice sheets melt, adding fresh water enriched in $^{16}\text{O}$ to the oceans. As a result, low (i.e., large negative) $\delta^{18}\text{O}$ values in the oceans indicate intervals of global warming. In contrast, during glacial intervals water is transferred from the oceans to the ice sheets primarily as $^{16}\text{O}$, so the remaining ocean water is enriched in $^{18}\text{O}$, which increases the $\delta^{18}\text{O}$ value. Thus, high (i.e., small negative or even positive) $\delta^{18}\text{O}$ values in oceanic water and sediments indicate cool intervals. Conversely, the same conditions result in the opposite relationships between global temperature and $\delta^{18}\text{O}$ values measured in polar ice.

Paleoclimatology uses the $\delta^{18}\text{O}$ parameter as a guide to past climates, as illustrated by the use of the record in sediment cores from the Ocean Drilling Project to interpret the paleoclimate that affected Chinese loess–paleosol profiles (Fig. 5.26). Samples can be used from a variety of sources: polar ice cores, ocean sediments, and fossil shells. In the last case, biological and chemical processes disturb the simple relationship with temperature, but this can be taken into account and corrected.

of the magnetite, its source and the mechanism of its formation in human tissue are exciting fields of modern research.
geomagnetic elements can be expressed in spherical polar coordinates. The magnitude of the magnetic vector is given by the field strength $F$; its direction is specified by two angles. The declination $D$ is the angle between the magnetic meridian and the geographic meridian; the inclination $I$ is the angle at which the magnetic vector dips below the horizontal (Fig. 5.27). The Cartesian ($X, Y, Z$) and spherical polar ($r, \theta, \phi$) sets of geomagnetic elements are related to each other as follows:

$$X = F \cos I \cos D; \quad Y = F \sin I \cos D; \quad Z = F \sin I$$

$$F^2 = X^2 + Y^2 + Z^2$$

$$D = \arctan \left( \frac{Y}{X} \right); \quad I = \arctan \left( \frac{Z}{\sqrt{X^2 + Y^2}} \right) \quad (5.32)$$

### 5.4.3 The magnetic field of external origin

The magnetic field $B$ and its potential $W$ at any point can be expressed in terms of the spherical polar coordinates ($r, \theta, \phi$) of the point of observation. Gauss expressed the potential of the geomagnetic field as an infinite series of terms involving these coordinates. Essentially, his method divides the field into separate components that decrease at different rates with increasing distance from the center of the Earth. The detailed analysis is complicated and beyond the scope of this text. The magnitude of the potential is given by

$$W = R \sum_{n=1}^{\infty} \left( A_n + \frac{B_n}{r^{n+1}} \right) \sum_{l=0}^{\infty} Y_l^m(\theta, \phi) \quad (5.33)$$

where $R$ is the Earth’s radius.

This is a rather formidable expression, but fortunately the most useful terms are quite simple. The summation signs indicate that the total potential is made up of an infinite number of terms with different values of $n$ and $l$. We will only pay attention here to the few terms for which $n = 1$. The expression in parentheses describes the variation of the potential with distance $r$. For each value of $n$ there will be different dependences (e.g., on $r, r^2, r^{-1}, r^{-2}$, $r^{-3}$, etc.). The function $Y_n^m(\theta, \phi)$ describes the variation of the potential when $r$ is constant, i.e., on the surface of a sphere. It is called a spherical harmonic function, because it has the same value when $\theta$ or $\phi$ is increased by an integral multiple of $2\pi$ (Box 2.3). For observations made on the spherical surface of the Earth, the constants $A_n$ describe parts of the potential that arise from magnetic field sources outside the Earth, which are called the geomagnetic field of external origin. The constants $B_n$ describe contributions to the magnetic potential from sources inside the Earth. This part is called the geomagnetic field of internal origin.

The potential itself is not measured directly. The geomagnetic elements $X, \ Y$ and $Z$ (Fig. 5.27) are recorded at magnetic observatories. Ideally these should be distributed uniformly over the Earth’s surface but in fact they are predominantly in the northern hemisphere. The geomagnetic field components are directional derivatives of the magnetic potential and depend on the same coefficients $A_n$ and $B_n$. Observations of magnetic field elements at a large number of measurement stations with a world-wide distribution allows the relative importance of $A_n$ and $B_n$ to be assessed. From the sparse data-set available in 1838 Gauss was able to show that the coefficients $A_n$ are very much smaller than $B_n$. He concluded that the field of external origin was insignificant and that the field of internal origin was predominantly that of a dipole.

5.4.4 The magnetic field of external origin

The magnetic field of the Earth in space has been measured from satellites and spacecraft. The external field has a quite complicated appearance (Fig. 5.28). It is strongly affected by the solar wind, a stream of electrically charged particles (consisting mainly of electrons, protons and helium nuclei) that is constantly emitted by the Sun. The solar wind is a plasma. This is the physical term for an ionized gas of low particle density made up of nearly equal concentrations of oppositely charged ions. At the distance of the Earth from the Sun (1 AU) the density of the solar wind is about 7 ions per cm$^3$, and it produces a magnetic field of about 6 nT. The solar wind interacts with the magnetic field of the Earth to form a region called the magnetosphere. At distances greater than a few Earth radii the interaction greatly alters the magnetic field from that of a simple dipole.

The velocity of the solar wind relative to the Earth is about 450 km s$^{-1}$. At a great distance (about 15 Earth radii) from the Earth, on the day side, the supersonic solar wind...
5.4 GEOMAGNETISM

Earth's surface and extends to an altitude of about 3000 km (Fig. 5.29); the outer belt occupies a doughnut-shaped region at distances between about 3 and 4 Earth radii (20,000–30,000 km) from the center of the Earth.

5.4.3.1 The Van Allen radiation belts

Charged particles that penetrate the magnetopause are trapped by the geomagnetic field lines and form the Van Allen radiation belts. These constitute two doughnut-shaped regions coaxial with the geomagnetic axis (Fig. 5.29). The inner belt contains mainly protons, the outer belt energetic electrons. Within each belt the charged particles move in helical fashion around the geomagnetic field lines (Fig. 5.30). The pitch of the spiraling motion gets smaller as the particle comes ever closer to the Earth and the field intensity increases; eventually it reaches zero and reverses sense. This compels the particles to shuttle rapidly from one polar region to the other along the field lines. The inner Van Allen belt starts about 1000 km above the Earth's surface and extends to an altitude of about...

5.4.3.2 The ionosphere

The effects described above illustrate how the Earth's magnetic field acts as a shield against much of the extra-terrestrial radiation. The atmosphere acts as a protective blanket against the remainder. Most of the very short-wavelength fraction of the solar radiation that penetrates the atmosphere does not reach the Earth's surface. Energetic γ- and x-rays and ultraviolet radiation cause ionization of the molecules of nitrogen and oxygen in the thin upper atmosphere at altitudes between about 50 km and 1500 km, forming an ionized region called the ionosphere. It is formed of five layers, labelled the D, E, F₁, F₂ and G layers from the base to the top. Each layer can reflect radio waves. The thicknesses and ionizations of the layers change during the course of a day; all but one or two layers on the night side of the Earth disappear while they thicken and strengthen on the day side (Fig. 5.31). A radio transmitter on the day side can bounce signals off the ionosphere that then travel around the world by multiple reflections between the ground surface and the ionosphere. Consequently, radio reception of distant stations from far across the globe is best during the local night hours. The D layer is closest to the Earth at an altitude of about 80–100 km. It was first discovered in 1902, before the nature of the ionosphere was known, because of its ability to reflect long-wavelength radio waves, and is named the Kennelly–Heaviside layer in honor of its discoverers. The E layer is used by short-wave amateur radio enthusiasts. The F layers are the most intensely ionized.

5.4.3.3 Diurnal variation and magnetic storms

The ionized molecules in the ionosphere release swarms of electrons that form powerful, horizontal, ring-like electrical currents. These act as sources of external magnetic fields that are detected at the surface of the Earth. The ionization is most intense on the day side of the Earth, where extra layers develop. The Sun also causes atmospheric tides in the ionosphere, partly due to gravitational attraction but mainly because the side facing the Sun is heated up during the day. The motions of the charged particles through the Earth’s magnetic field produce an electrical field, according to Lorentz's law (Eq. (5.10)), which drives electrical currents in the ionosphere. In particular, the horizontal component of particle velocity interacts with the vertical component of the geomagnetic field to produce horizontal electrical current loops in the ionosphere. These currents cause a magnetic field at the Earth's surface. As the Earth rotates beneath the ionosphere the observed intensity of the geomagnetic field fluctuates with a range of amplitude of about 10–30 nT at the Earth's surface and a period of one day (Fig. 5.32a). This...
time-dependent change of geomagnetic field intensity is called the diurnal (or daily) variation. The magnitude of the diurnal variation depends on the latitude at which it is observed. Because it greatly exceeds the accuracy with which magnetic fields are measured during surveys, the diurnal variation must be compensated by correcting field measurements accordingly. The intensity of the diurnal variation depends on the latitude at which it is observed. Because it greatly exceeds the accuracy with which magnetic fields are measured during surveys, the diurnal variation must be compensated by correcting field measurements accordingly.
of the effect depends on the degree of ionization of the ionosphere and is therefore determined by the state of solar activity. As described in Section 5.4.7.1, the solar activity is not constant. On days when the activity of the Sun is especially low, the diurnal variation is said to be of solar quiet (Ss) type. On normal days, or when the solar activity is high, the Ss variation is overlaid by a solar disturbance (Sp) variation. Solar activity changes periodically with the 11-yr cycle of sunspots and solar flares. The enhanced emission of radiation associated with these solar phenomena increases the ionospheric currents. These give rise to rapidly varying, anomalously strong magnetic fields (called magnetic storms) with amplitudes of up to 1000 nT at the Earth’s surface (Fig. 5.32b). The ionospheric disturbance also disrupts short-wave to long-wave radio transmissions. Magnetic surveying must be suspended temporarily while a magnetic storm is in progress, which can last for hours or days, depending on the duration of the solar activity.

5.4.4 The magnetic field of internal origin

To understand how geophysicists describe the geomagnetic field of internal origin mathematically we return to Eq. (5.33). First, we follow Gauss and omit the coefficients $A_n$ of the external field. The spherical harmonic functions $Y_{nm}(\theta, \phi)$ that describe the variation of potential on a spherical surface are then written in expanded form (Box 2.3). The potential $W$ of the field of internal origin becomes

$$W = R \sum_{n=1}^{\infty} \sum_{m=-n}^{n} \left( \frac{1}{R} \right)^{n+1} (g_{nm} \cos \theta + h_{nm} \sin \theta) P_n^m(\cos \theta)$$

(5.34)

Here, $R$ is the Earth’s radius, as before, and $P_n^m(\cos \theta)$ are called Schmidt polynomials, which are related to the associated Legendre polynomials (Box 2.2).

Equation (5.34) is a multipole expression of the geomagnetic potential. It relates the potential of the measured field to the potentials of particular combinations of magnetic poles (Box 5.3). The constants $g_{nm}$ and $h_{nm}$ in the geomagnetic potential are called the Gauss (or Gauss–Schmidt) coefficients of order $n$ and degree $m$. Inspection of Eq. (5.34) shows that they have the same dimensions (nT) as the $B$-field. Their values are computed from analysis of measurements of the geomagnetic field.

Data from several sources are integrated in modern analyses of the Earth’s magnetic field. Until the dawn of the satellite era, continuous records at magnetic observatories were the principal sources of geomagnetic data. Average values of the geomagnetic elements were determined, from which optimum values for the Gauss coefficients could be derived. Currently, about 200 permanent observatories make continuous measurements of the field. However, satellites orbiting the Earth in low near-polar orbits now provide most of the high-quality data used to model the field. The Polar Orbiting Geophysical Observatory (POGO), launched in 1965, was the first to deliver field measurements, but the greatest advance came with the Magnetic Field Satellite (MAGSAT), which delivered high-quality data during a six-month mission in 1979–1980. In 1979, a Danish satellite, ØRSTED, was placed in an elliptical, low-polar orbit, with altitude between 650 km and 865 km. It carried a vector magnetometer and a total field magnetometer, each with a sensitivity of 0.5 nT; the mission was dedicated to a survey of the geomagnetic field. The German satellite CHAMP (see also Section 2.4.6.4), was launched in 2000 in a lower, near-circular orbit on a planned five-year mission. In addition to measuring the gravity field, this satellite was also equipped to make scalar and vector measurements of the magnetic field. The low orbit and improved instrumentation were designed to provide magnetic measurements with high resolution.

In principle, an infinite number of Gauss coefficients would be needed to define the field completely. The coefficients of order and degree 8 and higher are very small and the calculation of Gauss coefficients must usually be truncated. A global model of the field is provided by the International Geomagnetic Reference Field (IGRF), which is based on coefficients up to $n = 10$, although analyses of higher order have been made. It is updated at regular intervals. The IGRF also gives the rate of change of each of the Gauss coefficients (its secular variation), which permits correction of the current values between update years.

The Gauss coefficients get smaller with increasing order $n$; this decrease provides a way of estimating the origin of the internal field. The analysis involves a technique called power spectral analysis. The distance across a feature of the magnetic field (for example, a region where the field is stronger than average) is called the wavelength of the feature. As in the case of gravity anomalies, deep-seated magnetic sources produce broad (long-wavelength) magnetic anomalies, while shallow sources result in narrow (short-wavelength) anomalies. Spectral analysis consists of calculating the power (alternatively called the energy density) associated with each “frequency” in the signal. This is obtained by computing the sum of the squares of all coefficients with the same order. In the case of the geomagnetic field, the spectral analysis is based on the values of the Gauss coefficients. The spatial frequency of any part of the observed field is contained in the order $n$ of the coefficients. Low-order terms (those with small values of $n$) correspond to long-wavelength features, high-order terms are related to short-wavelength features.

Measurements of the geomagnetic field from the MAGSAT Earth-orbiting satellite at a mean altitude of 420 km above the Earth’s surface have been analyzed to yield Gauss coefficients to order $n = 66$, special techniques being invoked for orders $n > 29$. A plot of the energy density associated with each order $n$ of the geomagnetic field shows three distinct segments (Fig. 5.33). The high-frequency terms of order $n > 40$ are uncertain and the terms with $n > 50$ are in the “noise level” of the analysis.
Each term in Eq. (5.34) is the potential of the magnetic field due to a particular combination of magnetic poles. For example, Eq. (5.34) contains three terms for which \( n = 1 \). Each corresponds to a dipole field. One dipole is parallel to the axis of rotation of the Earth and the other two are at right angles to each other in the equatorial plane. The five terms with \( n = 2 \) describe a geometrically more complex field, known as a quadrupole field. Just as a dipole field results when two single poles are brought infinitesimally close to each other (see Section 5.2.3), a quadrupole field results when two opposing coaxial dipoles are brought infinitesimally close end-to-end. As its name implies, the quadrupole field derives from four magnetic poles of which two are “north” poles and two are “south” poles. The terms with \( n = 3 \) describe an octupole field which is characterized by eight (\( 2^3 \)) poles. The terms with \( n = N \) describe a field that arises from a configuration of \( 2^N \) poles.

The configurations of axial dipole, quadrupole and octupole fields relative to a reference sphere are illustrated in Fig. B5.3.1 for the case where the order \( m \) of the Gauss coefficients is zero. If \( m = 0 \) in Eq. (5.34), the potential does not vary around a circle of “latitude” defined by a chosen combination of \( \theta \) and \( r \). This kind of field is said to have zonal symmetry. Any cross-section that contains the axis of symmetry is representative for the symmetry. Fig. B5.3.1 shows axial cross-sections of the simplest field line geometries corresponding to (a) dipole, (b) quadrupole and (c) octupole fields; each field is rotationally symmetrical about the axis of the configuration. The corresponding zonal spherical harmonics are illustrated symbolically by shading the alternate zones in which magnetic field lines leave or return to the surface of a sphere.

The dipole field is horizontal at the equator. In the southern hemisphere the field lines leave the reference sphere; in the northern hemisphere they return to it. In the northern hemisphere the field of an axial quadrupole is horizontal at latitude 35.3°N. North of this latitude the field lines of the quadrupole leave the reference sphere. An equivalent circle of latitude is located in the southern hemisphere at 35.3°S; south of this latitude the quadrupole field lines also leave the Earth. The field lines re-enter the Earth in the band of latitudes between those where the field is horizontal. The symmetry of the quadrupole field is described by these three zones around the axis. The axial octupole field exhibits zonal symmetry with four zones; two zones in which the field leaves the Earth alternate with two zones in which it re-enters it.

Terms in the potential expansion for which the degree \( m \) of the Gauss coefficients is equal to the order \( n \) (e.g., \( g_1^1, h_1^1, g_2^0, h_2^0 \)) are called sectorial harmonics. Their symmetry relative to the Earth’s axis is characterized by an even number of sectors around the equator in which the field lines alternately leave and re-enter the Earth (Fig. B5.3.2).

The potential terms for which \( m \neq n \) (e.g., \( g_1^0, h_0^0, g_0^1, h_0^1 \)) are known as tesseral harmonics. Their pattern of symmetry is defined by the intersections of circles of latitude and longitude, which outline alternating domains in which the field lines leave and re-enter the Earth, respectively.

Multipole representation allows the very complex geometry of the total field to be broken down into contributions from a number of fields with simple geometries.

By superposing many terms corresponding to these simple geometries a field of great complexity can be generated. It must be kept in mind that this is not a physical expression of the magnetic field, because magnetic poles do not exist. It is a convenient technique for describing the field mathematically.
and cannot be attributed any geophysical importance. The terms \(15 < n \leq 40\) are due to short-wavelength magnetic anomalies associated with the magnetization of the Earth's crust. The terms \(n \leq 14\) dominate the Earth's magnetic field and are due to much deeper sources in the fluid core.

5.4.4.1 The dipole field

The most important part of the Earth's magnetic field at the surface of the Earth is the dipole field, given by the Gauss coefficients for which \(n = 1\). If we write only the first term of Eq. (5.34) we get the potential

\[
W = \frac{R^3 \mu_0 g_1 \cos \theta}{r^3}
\]

(5.35)

Note that the spatial variation of this potential depends on \((\cos \theta r^2)\) in the same way as the potential of a dipole, which was found (Eq. (5.8)) to be

\[
W = \frac{\mu_0 m r \cos \theta}{4\pi r^2}
\]

(5.36)

Comparison of the coefficients of \((\cos \theta r^2)\) in Eq. (5.35) and Eq. (5.36) gives the dipole moment of the Earth’s axial dipole in terms of the first Gauss coefficient:

\[
m = \frac{4\pi}{\mu_0} R^3 g_1
\]

(5.37)

The term \(g_1\) is the strongest component of the field. It describes a magnetic dipole at the center of the Earth and aligned with the Earth's rotation axis. This is called the geocentric axial magnetic dipole.

The magnetic field \(\mathbf{B}\) of a dipole is symmetrical about the axis of the dipole. At any point at distance \(r\) from the center of a dipole with moment \(m\) on a radius that makes an angle \(\theta\) to the dipole axis the field of the dipole has a radial component \(B_r\) and a tangential component \(B_\theta\) which can be obtained by differentiating the potential with respect to \(r\) and \(\theta\), respectively:

\[
B_r = -\frac{\partial W}{\partial r} = \frac{\mu_0}{4\pi} \frac{2m \cos \theta}{r^3}
\]

(5.38)

\[
B_\theta = \frac{1}{r} \frac{\partial W}{\partial \theta} = \frac{\mu_0}{4\pi} m \sin \theta
\]

(5.39)

Note that \(B_r\) vanishes at the equator (\(\theta = 90^\circ\)) and the field is horizontal; comparing Eqs. (5.37) and (5.39) we get that the horizontal equatorial field \(B_\theta\) is equal to \(g_1\). At a point \((r, \theta)\) on the surface of a uniformly magnetized sphere the magnetic field line is inclined to the surface at an angle \(I\), which is given by

\[
\tan I = \frac{B_r}{B_\theta} = 2\cot \theta = 2\tan \lambda
\]

(5.40)

The angle \(I\) is called the inclination of the field, and \(\theta\) is the angular distance (or polar angle) of the point of observation from the magnetic axis. The polar angle is the complement of the magnetic latitude, \(\lambda\), \((\lambda = 90^\circ - \theta)\). Equation (5.40) has an important application in paleomagnetism, as will be seen later.

The terms \(g_1^1\) and \(h_1^1\) are the next strongest in the potential expansion. They describe contributions to the potential from additional dipoles with their axes in the equatorial plane. The total dipole moment of the Earth is then obtained from the vector sum of all three components:

\[
m = \frac{4\pi}{\mu_0} R^3 \left( g_1^1 + g_1^2 + h_1^1 \right)
\]

(5.41)

The analysis of the geomagnetic field for the year 2005 gave the following values for the dipole coefficients: \(g_1^1 = -29,556.8\) nT; \(g_1^2 = -1671.8\) nT; \(h_1^1 = 5080.0\) nT. The strength of the Earth's dipole magnetic moment obtained by inserting these values in Eq. (5.41) is \(m = 7.7674 \times 10^{22}\) A m\(^2\). Note that the sign of \(g_1^1\) is negative. This means that the axial dipole points opposite to the direction of rotation. Taken together, the three dipole components describe a geocentric dipole inclined at about 11.2° to the Earth's rotation axis. This tilted geocentric dipole accounts for more than 90% of the geomagnetic field at the Earth's surface. Its axis cuts the surface at the north and south geomagnetic poles. For epoch 2005 the respective poles were located at 79.7°N, 71.8°W (i.e., 288.2°E) and 79.7°S, 108.2°E. The geomagnetic poles are antipodal (i.e., exactly opposite) to each other.

The magnetic poles of the Earth are defined as the locations where the inclination of the magnetic field is ±90° (i.e., where the field is vertically upward or downward). An isoclinal map (showing constant inclination values) for the year 1980 shows that the location of the north magnetic pole was at 77.3°N, 258.2°E while the south magnetic pole was at 65.6°S, 139.4°E (Fig. 5.34a). These poles are not exactly opposite one another. The discrepancy between the magnetic poles and the geomagnetic poles arises because the terrestrial magnetic field is somewhat more complex than that of a perfect dipole. The intensity of the geomagnetic field is generally stronger in high latitudes than near the equator (Fig. 5.34b). The intensity is
especially low over the South Atlantic, where it is about 20 μT weaker than expected. The cause of this “South Atlantic magnetic anomaly” is not understood. Because the geomagnetic field shields the Earth from cosmic radiation and charged particles of the solar wind, this protection is less effective over the South Atlantic. Orbiting satellites note increased impacts of extra-terrestrial particles over this region. The feature poses hazards for astronauts in low-orbiting spacecraft and also for pilots and passengers in high-altitude aircraft that pass through the region. The enhanced particle flux can interfere with their on-board computers, communications and guidance systems.

5.4.4.2 The non-dipole field

The part of the field of internal origin (about 5% of the total field), obtained by subtracting the field of the inclined geocentric dipole from the total field, is collectively called the non-dipole field. A map of the non-dipole field consists of a system of irregularly sized, long-wavelength magnetic anomalies (Fig. 5.35). To describe this field requires all the terms in the potential expansion of order \( n \geq 2 \) in Eq. (5.34).

The distribution of positive and negative non-dipole field anomalies suggests an alternative representation of the non-dipole field to the multipole portrayal. The positive and negative anomalies have been modelled by inward or outward oriented radial dipoles in the core at about one-quarter the Earth’s radius. Each dipole is presumed to be caused by a toroidal current loop parallel to the surface of the core. A single centered axial dipole and eight auxiliary radial dipoles are adequate to represent the field observed at the Earth’s surface. The secular variation at a site (Section 5.4.5.2) is explained by this model as the passage of one of the auxiliary dipoles under the site. This model may be a little closer to physical reality but it is more unwieldy to handle. The multipole representation is the most convenient way of modelling the geomagnetic potential for mathematical analysis.

5.4.5 Secular variation

At any particular place on the Earth the geomagnetic field is not constant in time. When the Gauss coefficients of the internal field are compared from one epoch to another, slow but significant changes in their values are observed. The slow changes of the field only become appreciable
over decades or centuries of observation and so they are called secular variations (from the Latin word saeculum for a long age). They are manifest as variations of both the dipole and non-dipole components of the field.

5.4.5.1 Secular variation of the dipole field

The dipole field exhibits secular variations of intensity and direction. Calculations of the Gauss coefficients for different historical epochs show a near-linear decay of the strength of the dipole moment at a rate of about 3.2% per century between about 1550 AD and 1900 AD. At the start of the twentieth century the decay became even faster and has averaged about 5.8% per century during the last 80 yr (Fig. 5.36a). If it continues at the same almost linear rate, the field intensity would reach zero in about another 2000 yr. The cause of the quite rapid decay in intensity is not known; it may simply be part of a longer term fluctuation. However, another possibility is that the dipole moment may be decreasing preparatory to the next reversal of geomagnetic field polarity.

The position of the dipole axis also shows secular variation. The changes can be traced by plotting the colatitude (the angle between the dipole axis and the rotation axis) and longitude of the geomagnetic pole as a function of time. Data are only sufficiently abundant for spherical harmonic analysis since the early nineteenth century. Less reliable data, enlarged by archeomagnetic results (see Section 5.6.2.1), allow estimates of the secular variation of the dipole axis since the middle of the sixteenth century. The earlier data suggest that in the sixteenth century the dipole axis was tilted at only about 3° to the rotation axis; a gradual increase in tilt took place between the sixteenth and nineteenth centuries. During the last 200 yr the dipole axis has maintained an almost constant tilt of about 11–12° to the rotation axis (Fig. 5.36b).

For the past 400 yr the longitude of the geomagnetic pole has drifted steadily westward (Fig. 5.36c). Before the nineteenth century the pole moved westward at about 0.14 yr⁻¹; this corresponds to a pseudo-period of 2600 yr for a complete circle about the geographic pole. However, since the early nineteenth century the westward motion of the pole has been slower, at an average rate of 0.044 yr⁻¹, which corresponds to a pseudo-period of 8200 yr.
5.4.5.2 Secular variation of the non-dipole field

Comparison of maps of the non-dipole field for different epochs (Fig. 5.35) show two types of secular variation. Some anomalies (e.g., over Mongolia, the South Atlantic and North America) appear to be stationary but they change in intensity. Other anomalies (e.g., over Africa) slowly change position with time. The secular variation of the non-dipole field therefore consists of a standing part and a drifting part.

Although some foci may have a north–south component of motion, the most striking feature of the secular variation of the recent non-dipole field is a slow westward drift. This is superposed on the westward drift of the dipole, but can be separated readily by spherical harmonic analysis. The rate of drift of the non-dipole field can be estimated from longitudinal changes in selected features plotted for different epochs. The mean rate of westward drift of the non-dipole field in the first half of the last century has been estimated to be 0.18°/yr, corresponding to a period of about 2000 yr. However, some foci drift at up to about 0.7°/yr, much faster than the average rate. Results from several geomagnetic observatories show that the rate of drift is dependent on latitude (Fig. 5.37).

Westward drift is an important factor in theories of the non-dipole field. It is considered to be a manifestation of rotation of the outer layers of the core relative to the lower mantle. Theoretical models of the geomagnetic field (discussed in the next section) presume conservation of angular momentum of the fluid core. To maintain the angular momentum of a particle of fluid that moves radially inwards (decreasing the distance from the rotational axis) its angular rate of rotation must speed up. This results in a layered structure for the radial profile of angular rate of rotation relative to the mantle (Fig. 5.38). The outer layers of the core probably rotate more slowly than the solid mantle, imparting a westward drift to features of the magnetic field rooted in the fluid motion.

5.4.6 Origin of the internal field

Analysis of the Gauss coefficients and the wavelengths of features of the non-dipole field indicate that the main field is produced in the fluid outer core of the Earth. The composition of the fluid core has been estimated from seismic and geochemical data. The major constituent is liquid iron, with smaller amounts of other less dense elements. Geochemical analyses of iron meteorites suggest that the core composition may have a few percent of nickel, while shock-wave experiments require 6–10% of non-metallic light elements such as silica, sulfur or oxygen. The solid inner core is inferred from seismic and shock-wave data to consist of almost pure iron.

For the generation of the magnetic field the important parameters of the core are its temperature, viscosity and electrical conductivity. Temperature is known very poorly inside the Earth, but probably exceeds 3000 °C in...
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The electrical conductivity of iron at 20°C is $10^7 \, \Omega^{-1} \text{m}^{-1}$ and decreases with increasing temperature. At the high temperatures and pressures in the core, the electrical conductivity is estimated to be around $(3-5) \times 10^5 \, \Omega^{-1} \text{m}^{-1}$, which corresponds to a good electrical conductor. For comparison, the conductivity of carbon (used for many electrical contacts) is $3 \times 10^6 \, \Omega^{-1} \text{m}^{-1}$ at 20°C.

5.4.6.1 Magnetostatic and electromagnetic models

As observed by Gilbert in 1600, the dipole field of the Earth resembles that of a uniformly magnetized sphere. However, permanent magnetization is an inadequate explanation for the geomagnetic dipole. The mean magnetization of the Earth would need to be many times greater than that of the most strongly magnetic crustal rocks. The Curie point of the most important minerals at atmospheric pressure is less than 700°C, which is reached at depths of about 25 km so that only a thin outer crustal shell could be permanently magnetized. The necessary magnetization of this shell is even greater than values observed in crustal rocks. Moreover, a magnetostatic origin cannot account for the temporal changes observed in the internal field, such as its secular variation.

The main magnetic field of the Earth is thought to be produced by electrical currents in the conductive core. Although the core is a good conductor, an electrical current system in the core continually loses energy through ohmic dissipation. The lost electrical energy is converted to heat and contributes to the thermal balance of the core. The equations of electromagnetism applied to the core show that an electrical current in the core would decay to zero in around 10,000–20,000 yr unless it is sustained. Paleomagnetic evidence in the form of coherently magnetized rocks supports the existence of a geomagnetic field since Precambrian time, i.e. for about 3000 Myr. This implies that it must be continuously maintained or regenerated. The driving action for the main field is called the dynamo process, by analogy to the production of electrical power in a conductor that rotates in a magnetic field.

5.4.6.2 The geomagnetic dynamo

When a charged particle moves through a magnetic field, it experiences a deflecting electrical field (called the Lorentz field) proportional to the magnetic flux density $B$ and the particle velocity $v$, and acting in the direction normal to both $B$ and $v$. The Lorentz field acts as an additional source of electrical current in the core. Its strength is dependent on the velocity of motion of the conducting fluid relative to the magnetic field lines. When this term is included in the Maxwell electromagnetic equations, a magnetohydrodynamic equation relating the magnetic field $B$ to the fluid flow $v$ and conductivity $\sigma$ in the core is obtained. It is written

$$\frac{\partial B}{\partial t} = \frac{1}{\mu_0 \sigma} \Delta B + \Delta \times (v \times B)$$

(5.42)

This vector equation, although complicated, has immediate consequences. The left side gives the rate of change of magnetic flux in the core; it is determined by two terms on the right side. The first is inversely dependent on the electrical conductivity, and determines the decay of the field in the absence of a driving potential; the better the conductor, the smaller is this diffusion term. The second, dynamo term, depends on the Lorentz electrical field, which is determined by the velocity field of the fluid motions in the core. The conductivity of the outer core $(3-5) \times 10^5 \, \Omega^{-1} \text{m}^{-1}$ is high and for a fluid velocity of about 1 mm s$^{-1}$ the dynamo term greatly exceeds the diffusion term. Under these conditions, the lines of magnetic flux in the core are dragged along by the fluid flow. This concept is called the frozen-flux theorem, and it is fundamental to dynamo theory. The diffusion term is only zero if the electrical conductivity is infinite. There is probably some diffusion of the field through the fluid, because it is not a perfect conductor. However, the frozen-flux theorem appears to approximate well the conditions in the fluid outer core.

The derivation of a solution of the dynamo theory is difficult. In addition to Maxwell’s equations with the addition of a term for the Lorentz field, the Navier–Stokes equation for the fluid flow, Poisson’s equation for the gravitational potential and the generalized equation of heat transfer must be simultaneously satisfied. The fluid flow consists of a radial component and a rotational component. The energy for the radial flow comes from two sources. The slow cooling of the Earth produces a temperature gradient in the core, which results in thermally driven convection in the iron-rich fluid of the outer core. This is augmented by latent heat released at the boundary between the inner and outer cores as the inner core solidifies. The solidification of the pure iron inner core depletes the fluid of the outer core of its heaviest component. The remaining lighter elements rise through the liquid outer core, causing a buoyancy-driven convection.

The rotational component of the fluid flow is the result of a radial velocity gradient in the liquid core, with inner layers rotating faster than outer layers (Fig. 5.38). The relative rotation of the conducting fluid drags magnetic field lines around the rotational axis to form ring-like, toroidal configurations. The toroidal field lines are parallel to the flow and therefore to the surface of the core. This means that the toroidal fields are confined to the core and cannot be measured; their strengths and configurations must be estimated from models. Their interactions with the upwelling and descending branches of convective currents create electrical current systems that produce poloidal magnetic fields. These, in their turn, escape from the core and can be measured at the surface of the Earth. The fluid motions are subject to the effects of Coriolis forces, which prove to be strong enough to dominate the resultant flow patterns.
5.4.6.3 Computer simulation of the geodynamo

Important advances in understanding the geomagnetic dynamo (or geodynamo) have been made by simulating the related core processes with supercomputers. In 1995, G. A. Glatzmaier and P. H. Roberts presented a numerical model for the generation of a magnetic field assuming a hot convective fluid outer core surrounding a solid inner core, with rotation rate akin to that of the Earth. The heat flow, electrical conductivity and other material properties were made as similar as possible to those of the Earth’s core. The simulated dynamo had dominantly dipole character and intensity similar to the Earth’s, and it exhibited a westward drift of the non-dipole field comparable to that measured at the Earth’s surface. It also reversed polarity spontaneously, with long periods of constant polarity between short polarity transitions, as is the case for the history of geomagnetic reversals in the last 160 Myr (Section 5.7). During a polarity reversal, the field intensity decreased by an order of magnitude, as is also observed in paleomagnetic studies (see Fig. 5.71).

The simulations showed that the ability to reverse polarity was increased when the heat flow across the core–mantle boundary was non-uniform, as in the Earth, showing that thermal conditions in the lower mantle influence the formation of the magnetic field in the fluid core. The solid inner core evidently plays an important role in the reversal process. Magnetic fields in the outer core can change quickly, accompanying convection, and may act to initiate reversals. However, magnetic fields in the solid inner core change more slowly by diffusion and thus the inner core may act to stabilize the field against reversals. A reversal occurs occasionally, on a longer time scale than the core processes.

A prediction of this model is that the magnetic field couples the inner core to the eastward flowing fluid above it, so that the inner core rotates slightly faster than the mantle and crust (Fig. 5.38). There is seismic evidence that this indeed occurs. The solid inner core is seismically anisotropic, with cylindrical symmetry about an axis close to the rotation axis. The axis of symmetry rotates at a rate about 1° faster than that of the mantle and crust.

5.4.7 Magnetic fields of the Sun, Moon and planets

Our knowledge of the magnetic fields of the Sun and planets derives from two types of observation. Indirect observations utilize spectroscopic effects. All atoms emit energy related to the orbital and spin motions. The energy is quantized so that the atom possesses a characteristic spectrum of energy levels. The lowest of these is called the ground state. When an atom happens to have been excited to a higher energy level, it is unstable and eventually returns to the ground state. In the process the energy corresponding to the difference between the elevated and ground states is emitted as light. For example atomic hydrogen gas in the Sun and galaxies emits radiation at 1420 MHz, with corresponding wavelength of 21 cm. This frequency is in the microwave range and can be detected by radio telescopes. If the hydrogen gas is in motion the frequency is shifted by the Doppler effect, from which the velocity of the motion can be deduced.

In the presence of a magnetic field a spectral line can split into several lines. This “hyperfine splitting” is called the Zeeman effect. When a hydrogen atom is in the ground state its hyperfine structure has only one line. However, if the atom is in a magnetic field, it acquires additional energy from the interactions between the atom and the magnetic field and hence it can exist in several energy states. As a result the spectral lines of energy emitted by the atom are split into closely spaced lines, representing transitions between the different energy states. The energy differences between these states are dependent on the strength of the magnetic field, which can be estimated from the observations.

Direct observations of extra-terrestrial magnetic fields have been carried out since the 1960s by space probes. Magnetometers mounted in these spacecraft have recorded directly the intensity of the interplanetary magnetic field as well as the magnetic fields around several planets. The manned Apollo missions to the Moon resulted in a large amount of data obtained from the orbiting spacecraft. The materials collected on the Moon’s surface and brought back to Earth by the astronauts have provided valuable information about lunar magnetic properties.

5.4.7.1 Magnetic field of the Sun

The Sun has nearly 99.9% of the mass of the solar system. About 99% of this mass is concentrated in a massive central core that reaches out to about 80% of the Sun’s radius. The remainder of the Sun consists of an outer conducting shell, which contains only 1% of the Sun’s mass but has a thickness equivalent to about 20% of the solar radius. Thermonuclear conversion of hydrogen into helium in the dense core produces temperatures of the order of 15,000,000 K. The visible solar disk is called the photosphere. Its diameter is about 2,240,000 km (about 175 times the diameter of the Earth) and its surface temperature is about 6000 K. The lower solar atmosphere is called the chromosphere; the outer atmosphere is called the corona. The chromosphere includes spike-like emissions of hydrogen gas called solar prominences that sometimes can reach far out into the corona.

The heat from the core is radiated out to the outer conducting shell, where it sets up convection currents. Some of these convection currents are small scale (about 1000 km across) and last only a few minutes; others are larger scale (30,000 km across) and persist for about a (terrestrial) day. The convection is affected by the Sun’s rotation and is turbulent.

The rotation of the Sun has been estimated spectroscopically by measuring the Doppler shift of spectral
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Lines. Independent estimates come from observing the motion of solar features like sunspots, etc. The rotational axis is tilted slightly at 7° to the pole to the ecliptic. Near the Sun's equator the rotational period is about 25 Earth days; the polar regions rotate more slowly with a period of 35–40 days. The Sun's core may rotate more rapidly than the outer regions. Turbulent convection and velocity shear in the outer conducting shell are conducive to a dynamo origin for the Sun's magnetic field. The surface field is dipolar in higher latitudes but has a more complicated pattern in equatorial latitudes.

Temperatures in the outer solar atmosphere (corona) are very high, around 1,000,000 K. The constituent particles achieve velocities in excess of the escape velocity of the Sun's gravitational field. The supersonic stream of mainly protons (H⁺ ions), electrons and α-particles (He²⁺ ions) escaping from the Sun forms the solar wind. The flow of electric charge produces an interplanetary magnetic field (IMF) of varying intensity. At the distance of the Earth from the Sun the IMF measures about 6 nT.

Sunspots have been observed since the invention of the telescope (ca. 1610 AD). A sunspot is a dark fleck on the surface of the Sun, measuring roughly 100 to 100,000 km in diameter. It has a lower temperature than the surrounding photosphere and represents a strong disturbance extending far into the Sun's interior. Sunspots last for several days or weeks and move with the Sun's rotation, providing a means of estimating the rotational speed. The frequency of sunspots changes cyclically with a period of 11 years.

Intense magnetic fields are associated with the sunspots. These often occur in unequally sized pairs of opposite polarity. The predominating magnetic polarity of sunspots changes from one period of maximum sunspot activity to the next, implying that the period is in fact 22 years. The magnetic field of each sunspot is toroidal. It can be imagined to resemble a vortex, or tornado, in which the magnetic field lines leave the solar surface in one sunspot and return to it in the other member of the pair. The polarity of the Sun's dipole field also reverses with the change of polarity of the sunspots, indicating that the features are related.

Associated with the sunspots and their strong magnetic fields are emissions of hydrogen gas, called solar flares. The charged particles ejected in the flares contribute to the solar wind, which consequently transfers the sunspot cyclicity to fluctuations in the Earth's magnetic field of external origin and to ancillary terrestrial phenomena such as magnetic storms, brilliant auroras and interference with radio transmissions.

5.4.7.2 Lunar magnetism

Classical hypotheses for the origin of the Moon – rotational fission, capture or binary accretion – have been superseded by the Giant Impact hypothesis. According to this model, the Earth experienced a catastrophic collision with a Mars-sized protoplanet early in its development. Some debris from the collision remained in orbit around the Earth, where it re-accumulated as the Moon about 4.5 Ga ago. This origin accounts for why the Earth and Moon have the same oxygen isotope composition. At the time of the Giant Impact the Earth's mantle and core had already differentiated and dense iron had settled into the core. This may also have taken place in the impactor so the Moon formed from the iron-depleted rocky mantles of the impacting bodies. The Moon has only a small core, whereas if it had originated like the other planets its core would be proportionately larger.

The early Moon was probably covered by an ocean of molten magma at least a hundred kilometers thick, the relics of which are now the major constituents of the lunar highlands. After cooling and solidification, the lunar surface was bombarded by planetesimals and meteoroids until about 4 Ga ago. The huge craters left by the impacts were later filled by molten basalt to form the lunar maria. Rock samples recovered from the maria in the manned Apollo missions have been dated radiometrically at 3.1–3.9 Ga. The volcanism may have continued after this time but probably ceased about 2.5 Ga ago. Since then the lunar surface has been pulverized by the constant bombardment by meteorites, micrometeorites and elementary particles of the solar wind and cosmic radiation. Except for the lunar highlands the surface is now covered by a layer of shocked debris several meters thick called the lunar regolith.

Our knowledge of lunar magnetism derives from magnetic field measurements made from orbiting spacecraft, magnetometers set up on the Moon and rock samples collected from the lunar surface and brought back to Earth. Measurements of the lunar surface field from orbiting spacecraft have been made by on-board magnetometers and by the electron reflection method, the principle of which is basically the same as used to explain the origin of the Van Allen belts in the Earth's magnetosphere (see Section 5.4.3.1). Electrons rain abundantly upon the lunar surface, in part from the solar wind and in part from the charges trapped in the Earth's magnetotail, which is crossed by the monthly orbit of the Moon about the Earth. In the absence of a lunar magnetic field the electrons would be absorbed by the lunar surface. As in the Earth's magnetosphere, an electron incident on the Moon is forced by the Lorentz force to spiral about a magnetic field line (see Fig. 5.30). As the electron approaches the lunar surface the magnetic field strength increases and the pitch of the helix decreases to zero. At the point of closest approach (also called the mirroring point) the electron motion is completely rotational about the field line. The electron path then spirals with increasing pitch back along the field line, away from the Moon. The effectiveness of the electron reflection is directional; for a given electron velocity there is a critical angle of incidence below which reflection ceases. By counting the number of electrons with a known energy that are reflected past a
satellite in known directions, the surface field $B_s$ can be estimated from the field $B_\theta$ measured at the altitude of the satellite.

The lunar magnetic field was surveyed extensively in 1998 by the Lunar Prospector orbiting spacecraft. The measurements show that the Moon currently has no detectable dipole moment. It has only a very weak non-dipolar magnetic field due to local magnetization of crustal rocks. Estimates of surface anomalies suggest that the largest are of the order of 100 nT. From records of meteoritic impacts and moonquakes made by seismometers left on the Moon for several years it has been inferred that, if the Moon has a core (necessary for a lunar dynamo), it must be smaller that 400–500 km in radius, representing less than 2% of the Moon’s volume and 1–3% of its mass. In contrast, the Earth’s core occupies about 16% of the planet’s volume and has about 33% of its mass. Lunar Prospector detected magnetic field changes due to electrical currents induced in the Moon as it traversed the stretched out tail of Earth’s magnetosphere. They suggest an even smaller, iron-rich metallic core, with radius 340 ± 90 km (Hood et al., 1999).

Although the Moon now has no global dipole magnetic field, samples of lunar rocks recovered in the manned Apollo missions possessed quite strong natural remanent magnetizations. Rock magnetic studies on Apollo samples with radiometric ages of ~3.6–3.9 Ga suggest that they were magnetized in fields of the order of 10–100 $\mu$T (0.1–1 gauss), much stronger than present fields on the Moon. As yet, the ancient lunar magnetic field is not well understood. The interior of the Moon may have acquired a primordial remanent magnetization in the external field of the Sun or Earth. In turn this may have provided the fields for acquisition of the observed crustal magnetizations. The small lunar metallic core is currently solid, and the core heat flux is too low to have powered an internal dynamo at the time of formation of the Apollo samples. However, models suggest that a thin layer of the Moon’s mantle adjacent to the core may have first blanketed the core, then later provided enough radioactive heating to assist dynamo activity that persisted for a limited period during a “magnetic era” about 0.5–1.0 Ga after the Moon was formed (Stegman et al., 2003).

### 5.4.7.3 Extra-terrestrial magnetic exploration

Jupiter and Saturn, like the Earth, have magnetic fields that are strong enough to trap charged particles. The motions of these charged particles generate electromagnetic radiation which is detectable as radio waves far from the planet. The magnetic field of Jupiter was first detected in this way. Moreover, the radio emissions are modulated by the rotation of the planet. Analysis of the periodicity of their modulated radio emissions provides the best estimates of the rotational rates of Jupiter and Saturn.

Most data concerning the magnetic fields of the planets (Table 5.2) have been obtained with flux-gate magnetometers (see Section 5.5.2.1) installed in passing or orbiting spacecraft. When the spacecraft traverses the magnetosphere of a planet, the magnetometer registers the passage through the bow shock and magnetopause (Fig. 5.39). A bow shock results from the supersonic collision of the solar wind with the atmosphere of a planet, just as it does for the Earth (see Fig. 5.28). Counters of energetic particles register a sudden increase in frequency and the magnetometer shows a change in magnetic field strength during passage through the bow shock into the magnetosheath. When the spacecraft leaves the magnetosheath and crosses the magnetopause it enters the region that is shielded from the solar wind by the magnetic field of the planet. The magnetopause is where the kinetic energy of the plasma is equal to the potential energy of the planetary magnetic field. The existence of a bow shock may be regarded as evidence for a planetary atmosphere, while the magnetopause is evidence that the planet has a magnetic field.

### 5.4.7.4 The magnetic fields of the planets

**Mercury** was visited by the Mariner 10 spacecraft, which made three passes of the planet in 1974 and 1975. The
on-board magnetometer detected a bow shock and a magnetopause (Fig. 5.39), which imply that the planet has a magnetic field. On the first encounter a magnetic field of about 100 nT was measured at an altitude of 700 km (radial distance of 3100 km), which was the point of closest approach. In estimating the magnetic field of the planet compensation must be made for the magnetic field of the solar wind. Different models have given estimates of the strength of the dipole moment in the range \((2–7) \times 10^{-11}\) of the Earth's magnetic moment \((m_E = 7.7674 \times 10^{22} \text{ Am}^2\) in 2005), which would give a surface equatorial field of about 100–400 nT. The source of the magnetic field is uncertain. It is possible, but unlikely, that Mercury has a global magnetic moment due to crustal remanent magnetization; it would be difficult to explain the uniformity and duration of the external field needed to produce this. Mercury is believed to have a large iron core with a radius of about 1800 km, proportionately larger than Earth's. Probably part of this core is molten, so it is possible that Mercury has a small active internal dynamo. This would however be unlike Earth's. It might be of thermolectric origin, or it could be due to dynamo processes in a thin shell-like liquid outer core surrounding a solid inner core.

**Venus** was investigated by several American and Russian spacecraft in the 1960s. The instruments on Mariner 5 in 1967 clearly detected a bow shock from the collision of the solar wind with the planetary atmosphere. Magnetometer data from later spacecraft found no evidence for a planetary magnetic field. If a magnetopause exists, it must lie very close to the planet and may wrap around it. Data from the Pioneer Venus orbiter in 1978 set an upper limit of \(10^{-5} m_E\) for a planetary dipole moment, which would give a surface equatorial field of less than 1 nT. The absence of a detectable magnetic field was not expected. As shown by Eq. (5.37) the dipole magnetic moment is proportional to the equatorial field times the cube of the planetary radius. The strength of a dynamo field is expected to be proportional to the core radius and to the rotation rate. These considerations give a scaling law whereby the dipole magnetic moment of a planet is proportional to its rotation rate and the fourth power of the core radius. The rotation of the planet is very slow compared to that of the Earth; one sidereal day on Venus lasts 243 Earth days, but the size of the planet is close to that of the Earth. It was therefore expected that Venus might have an internal dynamo with a dipole moment about 0.2% that of the Earth and an equatorial surface field of about 86 nT. It seems likely that the slow rotation does not provide enough energy for an active dynamo.

**Mars** was expected to have a magnetic moment between that of Earth and Mercury, because of its size and rotation rate. Mariner 4 in 1965 was the first American spacecraft carrying magnetometers to visit Mars; it detected a bow shock but no conclusive evidence for a magnetopause. In September 1997, the Mars Global Surveyor spacecraft entered orbit around the planet. Since 1999 it has mapped the Martian magnetic field from an almost circular orbit about 400 km above the planet's surface. The survey measurements have nearly uniform global coverage and show that there is no significant global magnetic field at present. This does not exclude the possible existence of a dynamo-generated global field in the planet's distant past. The measured magnetic field consists of large regional magnetic anomalies, which are attributed to remanent magnetization of the Martian crust. At the survey altitude the crustal magnetic anomalies have amplitudes up to 220 nT, an order of magnitude larger than the crustal anomalies of Earth at that altitude, which attests to the presence of strongly magnetic minerals in the Martian crust. The crustal magnetic field shows some features with circular geometry, some of which may be related to impact processes. However, the most striking anomalies are prominent
east–west trending linear features over a region in the planet’s southern hemisphere. These sub-parallel lines are attributed to alternating bands of strongly magnetized crust. It is tempting to interpret their origin in the same way as the generation of oceanic magnetic lines on Earth by a plate tectonic process in the presence of a reversing field. However, as yet the origin of the magnetic anomalies on Mars is not understood.

Jupiter has been known since 1955 to possess a strong magnetic field, because of the polarized radio emissions associated with it. The spacecraft Pioneer 10 and 11 in 1973–4 and Voyager 1 and 2 in 1979 established that the planet has a bow shock and magnetopause. From 1995 to 2003 the spacecraft Galileo made extensive surveys of Jupiter’s magnetosphere. The huge magnetosphere encounters the solar wind about 5,000,000 km “upwind” from the planet; its magnetotail may extend all the way to Saturn. Two reasons account for the great size of the magnetosphere compared to that of Earth. First, the solar wind pressure on the jovian atmosphere is weaker due to the greater distance from the Sun; secondly, Jupiter’s magnetic field is much stronger than that of the Earth. The dipole moment is almost 20,000$M_E$, which gives a powerful equatorial magnetic field of more than 400,000 nT at Jupiter’s surface. The quadrupole and octupole parts of the non-dipole magnetic field have been found to be proportionately much larger relative to the dipole field than on Earth. The dipole axis is tilted at 9.7° to the rotation axis, and is also displaced from it by 10% of Jupiter’s equatorial radius. The magnetic field of Jupiter results from an active dynamo in the metallic hydrogen core of the planet. The core is probably very large, with a radius up to 75% of the planet’s radius. This would explain the high harmonic content of the magnetic field near the planet.

Saturn was reached by Pioneer 11 in 1979 and the Voyager 1 and 2 spacecraft in 1980 and 1981, respectively. The on-board magnetometers detected a bow shock and a magnetopause. In 2004 the Cassini–Huygens spacecraft entered into orbit around Saturn. In 2005 the Huygens lander descended to the surface of Saturn’s largest moon Titan while Cassini continued to orbit and measure the parent planet’s properties. Saturn’s dipole magnetic moment is smaller than expected, but is estimated to be around 500$M_E$. This gives an equatorial field of 55,000 nT, almost double that of Earth. The magnetic field has a purer dipole character (i.e., the non-dipole components are weaker) than the fields of Jupiter or Earth. The simplest explanation for this is that the field is generated by an active dynamo in a conducting core that is smaller relative to the size of the planet. The axis of the dipole magnetic field lies only about 1° away from the rotation axis, in contrast to 11.4° on Earth and 9.7° on Jupiter.

Uranus is unusual in that its spin axis has an obliquity of 97.9°. This means that the rotation axis lies very close to the ecliptic plane, and the orbital planes of its satellites are almost orthogonal to the ecliptic plane. Uranus was visited by Voyager 2 in January 1986. The spacecraft encountered a bow shock and magnetopause and subsequently entered the magnetosphere of the planet, which extends for 18 planetary radii (460,000 km) towards the Sun. Uranus has a dipole moment about 50 times stronger than Earth’s, giving a surface field of 24,000 nT, comparable to that on Earth. Intriguingly, the axis of the dipole field has a large tilt of about 60° to the rotation axis; there is no explanation for this tilt. Another oddity is that the magnetic field is not centered on the center of the planet, but is displaced by 30% of the planet’s radius along the tilted rotation axis. The quadrupole component of the field is relatively large compared to the dipole. It is therefore supposed that the magnetic field of Uranus is generated at shallow depths within the planet.

Neptune, visited by Voyager 2 in 1989, has a magnetic field with similar characteristics to that of Uranus. It is tilted at 49° to the rotation axis and is offset from the planetary center by 55% of Neptune’s radius. As in the case of Uranus, the magnetic field has a large quadrupole term compared to the dipole and thus probably originates in the outer layers of the planet, rather than in the deep interior.

It is not yet known whether Pluto has a magnetic field. The planet is probably too small to have a magnetic field sustained by dynamo action.

There is reasonable confidence that Mercury, Earth, Jupiter, Saturn, and Uranus have active planetary dynamos today. The magnetic data for Mars and Neptune are inconclusive. All available data indicate that Venus and the Moon do not have active dynamos now, but possibly each might have had one earlier in its history.

5.5 MAGNETIC SURVEYING

5.5.1 The magnetization of the earth’s crust

The high-order terms in the energy density spectrum of the geomagnetic field (Fig. 5.33) are related to the magnetization of crustal rocks. Magnetic investigations can therefore yield important data about geological structures. By analogy with gravity anomalies we define a magnetic anomaly as the difference between the measured (and suitably corrected) magnetic field of the Earth and that which would be expected from the International Geomagnetic Reference Field (Section 5.4.4). The magnetic anomaly results from the contrast in magnetization when rocks with different magnetic properties are adjacent to each other, as, for example, when a strongly magnetic basaltic dike intrudes a less magnetic host rock. The stray magnetic fields surrounding the dike disturb the geomagnetic field locally and can be measured with sensitive instruments called magnetometers.

As discussed in Section 5.3.1, each grain of mineral in a rock can be classified as having diamagnetic, paramagnetic or ferromagnetic properties. When the rock is in a magnetic field, the alignment of magnetic moments by the field produces an induced magnetization ($M_I$)
5.5 MAGNETIC SURVEYING

![Diagram of remanent, induced, and total magnetizations](image)

Fig. 5.40 The remanent ($M_r$), induced ($M_i$), and total ($M_t$) magnetizations in a rock. (a) For an arbitrary case $M_i$ lies between $M_r$ and $M_t$, (b) for a very large Königsberger ratio ($Q_n > 1$) $M_i$ is close to $M_t$, and (c) for a very small Königsberger ratio ($Q_n \ll 1$) $M_i$ is almost the same as $M_r$.

Proportional to the field, the proportionality constant being the magnetic susceptibility, which can have a wide range of values in rocks (see Fig. 5.13). The geomagnetic field is able to produce a correspondingly wide range of induced magnetizations in ordinary crustal rocks. The direction of the induced magnetization is parallel to the Earth’s magnetic field in the rock.

Each rock usually contains a tiny quantity of ferromagnetic minerals. As we have seen, these grains can become magnetized permanently during the formation of the rock or by a later mechanism. The remanent magnetization ($M_r$) of the rock is not related to the present-day geomagnetic field, but is related to the Earth’s magnetic field in the geological past. Its direction is usually different from that of the present-day field. As a result the directions of $M_r$ and $M_i$ are generally not parallel. The direction of $M_i$ is the same as that of the present field but the direction of $M_r$ is often not known unless it can be measured in rock samples.

The total magnetization of a rock is the sum of the remanent and induced magnetizations. As these have different directions they must be combined as vectors (Fig. 5.40a). The direction of the resultant magnetization of the rock is not parallel to the geomagnetic field. If the intensities of $M_r$ and $M_i$ are similar, it is difficult to interpret the total magnetization. Fortunately, in many important situations $M_r$ and $M_i$ are sufficiently different to permit some simplifying assumptions. The relative importance of the remanent and induced parts of the magnetization is expressed in the Königsberger ratio ($Q_n$), defined as the ratio of the intensity of the remanent magnetization to that of the induced magnetization (i.e., $Q_n = M_r / M_i$).

Two situations are of particular interest. The first is when $Q_n$ is very large (i.e., $Q_n > 1$). In this case (Fig. 5.40b), the total magnetization is dominated by the remanent component and its direction is essentially parallel to $M_r$. Oceanic basalts, formed by extrusion and rapid underwater cooling at oceanic ridges, are an example of rocks with high $Q_n$ ratios. Due to the rapid quenching of the molten lava, titanomagnetite grains form with skeletal structures and very fine grain sizes. The oceanic basalts carry a strong thermoremanent magnetization and often have $Q_n$ values of 100 or greater. This facilitates the interpretation of oceanic magnetic anomalies, because in many cases the induced component can be neglected and the crustal magnetization can be interpreted as if it were entirely remanent.

The other important situation is when $Q_n$ is very small (i.e., $Q_n \ll 1$). This requires the remanent magnetization to be negligible in comparison to the induced magnetization. For example, coarse grained magnetite grains carry multidomain magnetizations (Section 5.3.5.3). The domain walls are easily moved around by a magnetic field. The susceptibility is high and the Earth’s magnetic field can induce a strong magnetization. Any remanent magnetization is usually weak, because it has been subdivided into antiparallel domains. These two factors yield a low value for $Q_n$.

Magnetic investigations of continental crustal rocks for commercial exploitation (e.g., in ancient shield areas) can often be interpreted as cases with $Q_n \ll 1$. The magnetization can then be assumed to be entirely induced (Fig. 5.40c) and oriented parallel to the direction of the present-day geomagnetic field at the measurement site, which is usually known. This makes it easier to design a model to interpret the feature responsible for the magnetic anomaly.

5.5.2 Magnetometers

The instrument used to measure magnetic fields is called a magnetometer. Until the 1940s magnetometers were mechanical instruments that balanced the torque of the magnetic field on a finely balanced compass needle against a restoring force provided by gravity or by the torsion in a suspension fiber. The balance types were cumbersome, delicate and slow to operate. For optimum sensitivity they were designed to measure changes in a selected component of the magnetic field, most commonly the vertical field. This type of magnetometer has now been superseded by more sensitive, robust electronic instruments. The most important of these are the flux-gate, precession and optically pumped magnetometers.
Some special nickel–iron alloys have very high magnetic susceptibility and very low remanent magnetization. Common examples are Permalloy (78.5% Ni, 21.5% Fe) and Mumetal (77% Ni, 16% Fe, 5% Cu, 2% Cr). The preparation of these alloys involves annealing at very high temperature (1100–1200 °C) to remove lattice defects around which internal stress could produce magnetostrictive energy. After this treatment the coercivity of the alloy is very low (i.e., its magnetization can be changed by a very weak field) and its susceptibility is so high that the Earth’s field can induce a magnetization in it that is a considerable proportion of the saturation value.

The sensor of a flux-gate magnetometer consists of two parallel strips of the special alloy (Fig. 5.41a). They are wound in opposite directions with primary energizing coils. When a current flows in the primary coils, the parallel strips become magnetized in opposite directions. A secondary coil wound about the primary pair detects the change in magnetic flux in the cores (Fig. 5.41b), which is zero as soon as the cores saturate. While the primary current is rising or falling, the magnetic flux in each strip changes and a voltage is induced in the secondary coil. If there is no external magnetic field, the signals due to the changing flux are equal and opposite and no output signal is recorded. When the axis of the sensor is aligned with the Earth’s magnetic field, the latter is added to the primary field in one strip and subtracted from it in the other. The phases of the magnetic flux in the alloy strips are now different; one saturates before the other. The flux changes in the two alloy strips are no longer equal and opposite. An output voltage is produced in the secondary coil that is proportional to the strength of the component of the Earth’s magnetic field along the axis of the sensor.

The flux-gate magnetometer is a vector magnetometer, because it measures the strength of the magnetic field in a particular direction, namely along the axis of the sensor. This requires that the sensor be accurately oriented along the direction of the field component to be measured. For total field measurements three sensors are employed. These are fixed at right angles to each other and connected with a feedback system which rotates the entire unit so that two of the sensors detect zero field. The magnetic field to be measured is then aligned with the axis of the third sensor.

The flux-gate magnetometer does not yield absolute field values. The output is a voltage, which must be calibrated in terms of magnetic field. However, the instrument provides a continuous record of field strength. Its sensitivity of about 1 nT makes it capable of measuring most magnetic anomalies of geophysical interest. It is robust and adaptable to being mounted in an airplane, or towed behind it. The instrument was developed during World War II as a submarine detector. After the war it was used extensively in airborne magnetic surveying.

5.5.2.2 The proton-precession magnetometer

Since World War II sensitive magnetometers have been designed around quantum-mechanical properties. The proton-precession magnetometer depends on the fact that the nucleus of the hydrogen atom, a proton, has a magnetic moment proportional to the angular momentum of its spin. Because the angular momentum is quantized, the proton magnetic moment can only have specified values, which are multiples of a fundamental unit called the nuclear magneton. The situation is analogous to the quantization of magnetic moment associated with electron spin, for which the fundamental unit is the Bohr magneton. The ratio of the magnetic moment to the spin angular momentum is called the gyromagnetic ratio ($\gamma_p$) of the proton. It is an accurately known fundamental constant with the value $\gamma_p = 2.675 13 \times 10^8 s^{-1}T^{-1}$.

The proton-precession magnetometer is simple and robust in design. The sensor of the instrument consists of a flask containing a proton-rich liquid, such as water. Around the flask are wound a magnetizing solenoid and a detector coil (Fig. 5.42); some designs use the same solenoid alternately for magnetizing and detection. When the current in the magnetizing solenoid is switched on, it creates a magnetic field of the order of 10 mT, which is about 200 times stronger than the Earth’s field. The
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The intensity of the Earth's magnetic field is in the range 30,000–60,000 nT. The corresponding precessional frequency is approximately 1250–2500 Hz, which is in the audio-frequency range. Accurate measurement of the signal frequency gives an instrumental sensitivity of about 1 nT, but requires a few seconds of observation. Although it gives an absolute value of the field, the proton-precession magnetometer does not give a continuous record. Its portability and simplicity give it advantages for field use.

The flux-gate and proton-precession magnetometers are widely used in magnetic surveying. The two instruments have comparable sensitivities of 0.1–1 nT. In contrast to the flux-gate instrument, which measures the component of the field along its axis, the proton-precession magnetometer cannot measure field components; it is a total-field magnetometer. The total field \( \mathbf{B} \), is the vector sum of the Earth’s magnetic field \( \mathbf{B}_\text{e} \) and the stray magnetic field \( \Delta \mathbf{B} \), of, say, an orebody. Generally, \( \Delta \mathbf{B} \approx \mathbf{B}_\text{e} \), so that the direction of the total field does not deviate far from the Earth’s field. In some applications it is often adequate to regard the measured total field anomaly as the projection of \( \Delta \mathbf{B} \) along the Earth’s field direction.

5.5.2.3 The absorption-cell magnetometer

The absorption-cell magnetometer is also referred to as the alkali-vapor or optically pumped magnetometer. The principle of its operation is based on the quantum-mechanical model of the atom. According to their quantum numbers the electrons of an atom occupy concentric shells about the nucleus with different energy levels. The lowest energy level of an electron is its ground state. The magnetic moment associated with the spin of an electron can be either parallel or antiparallel to an external magnetic field. The energy of the electron is different in each case. This results in the ground state splitting into two sublevels with slightly different energies. The energy difference is proportional to the strength of the magnetic field. The splitting of energy levels in the presence of a magnetic field is called the Zeeman effect.

Absorption-cell magnetometers utilize the Zeeman effect in vapors of alkali elements such as rubidium or cesium, which have only a single valence electron in the outermost energy shell. Consider the schematic representation of an alkali-vapor magnetometer in Fig. 5.43. A polarized light-beam is passed through an absorption cell containing rubidium or cesium vapor and falls on a photoelectric cell, which measures the intensity of the light-beam. In the presence of a magnetic field the ground state of the rubidium or cesium is split into two sublevels, \( G_1 \) and \( G_2 \). If the exact amount of energy is added to the vapor, the electrons may be raised from their ground state to a higher-energy level, \( H \). Suppose that we irradiate the cell with light from which we have filtered out the spectral line corresponding to the energy needed for the transition \( G_1 \rightarrow H \). The energy for the transition \( G_1 \rightarrow H \) has not been removed, so the electrons in ground state \( G_1 \) will receive energy that excites them to level \( H \), whereas those in ground state \( G_2 \) will remain in this state. The energy for these transitions comes from the incident light-beam, which is absorbed in the cell. In due
course, the excited electrons will fall back to one of the more stable ground states. If an electron in excited state H falls back to sublevel G₁ it will be re-excited into level H; but if it falls back to sublevel G₂ it will remain there. In time, this process – called “optical pumping” – will empty sublevel G₁ and fill level G₂. At this stage no more energy can be absorbed from the polarized light-beam and the absorption cell becomes transparent. If we now supply electromagnetic energy to the system in the form of a radio-frequency signal with just the right amount of energy to permit transitions between the populated G₂ and unpopulated G₁ ground sublevels, the balance will be disturbed. The optical pumping will start up again and will continue until the electrons have been expelled from the G₁ level. During this time energy is absorbed from the light-beam and it ceases to be transparent.

In the rubidium-vapor and cesium-vapor magnetometers a polarized light-beam is shone at approximately 45° to the magnetic field direction. In the presence of the Earth’s magnetic field the electrons precess about the field direction at the Larmor precessional frequency. At one part of the precessional cycle an electron spin is almost parallel to the field direction, and one half-cycle later it is nearly antiparallel. The varying absorption causes a fluctuation of intensity of the light-beam at the Larmor frequency. This is detected by the photocell and converted to an alternating current. By means of a feedback circuit the signal is supplied to a coil around the container of rubidium gas and a radio-frequency resonant circuit is created. The ambient geomagnetic field $B_t$ that causes the splitting of the ground state is proportional to the Larmor frequency, and is given by

$$ B_t = \frac{2\pi f}{\gamma_e} \quad (5.44) $$

Here, $\gamma_e$ is the gyromagnetic ratio of the electron, which is known with an accuracy of about 1 part in $10^7$. It is about 1800 times larger than $\gamma_p$, the gyromagnetic ratio of the proton. The precessional frequency is correspondingly higher and easier to measure precisely. The sensitivity of an optically pumped magnetometer is very high, about 0.01 nT, which is an order of magnitude more sensitive than the flux-gate or proton-precession magnetometer.

5.5.3 Magnetic surveying

The purpose of magnetic surveying is to identify and describe regions of the Earth’s crust that have unusual (anomalous) magnetizations. In the realm of applied geophysics the anomalous magnetizations might be associated with local mineralization that is potentially of commercial interest, or they could be due to subsurface structures that have a bearing on the location of oil deposits. In global geophysics, magnetic surveying over oceanic ridges provided vital clues that led to the theory of plate tectonics and revealed the polarity history of the Earth’s magnetic field since the Early Jurassic.

Magnetic surveying consists of (1) measuring the terrestrial magnetic field at predetermined points, (2) correcting the measurements for known changes, and (3) comparing the resultant value of the field with the expected value at each measurement station. The expected value of the field at any place is taken to be that of the International Geomagnetic Reference Field (IGRF), described in Section 5.4.4. The difference between the observed and expected values is a magnetic anomaly.

5.5.3.1 Measurement methods

The surveying of magnetic anomalies can be carried out on land, at sea and in the air. In a simple land survey an operator might use a portable magnetometer to measure the field at the surface of the Earth at selected points that
form a grid over a suspected geological structure. This method is slow but it yields a detailed pattern of the magnetic field anomaly over the structure, because the measurements are made close to the source of the anomaly. In practice, the surveying of magnetic anomalies is most efficiently carried out from an aircraft. The magnetometer must be removed as far as possible from the magnetic environment of the aircraft. This may be achieved by mounting the instrument on a fixed boom, A, several meters long (Fig. 5.44a). Alternatively, the device may be towed behind the aircraft in an aerodynamic housing, B, at the end of a cable 30–150 m long. The “bird” containing the magnetometer then flies behind and below the aircraft. The flight environment is comparatively stable. Airborne magnetometers generally have higher sensitivity ($\approx 0.01$ nT) than those used in ground-based surveying (sensitivity $\approx 1$ nT). This compensates for the loss in resolution due to the increased distance between the magnetometer and the source of the anomaly. Airborne magnetic surveying is an economical way to reconnoitre a large territory in a short time. It has become a routine part of the initial phases of the geophysical exploration of an uncharted territory.

The magnetic field over the oceans may also be surveyed from the air. However, most of the marine magnetic record has been obtained by shipborne surveying. In the marine application a proton-precession magnetometer mounted in a waterproof “fish” is towed behind the ship at the end of a long cable (Fig. 5.44b). Considering that most research vessels consist of several hundred to several thousand tons of steel, the ship causes a large magnetic disturbance. For example, a research ship of about 1000 tons deadweight causes an anomaly of about 10 nT at a distance of 150 m. To minimize the disturbance of the ship the tow-cable must be about 100–300 m in length. At this distance the “fish” in fact “swims” well below the water surface. Its depth is dependent on the length of the tow-cable and the speed of the ship. At a typical survey speed of 10 km h$^{-1}$ its operational depth is about 10–20 m.

5.5.3.2 Magnetic gradiometers

The magnetic gradiometer consists of a pair of alkali-vapor magnetometers maintained at a fixed distance from each other. In ground-based surveying the instruments are mounted at opposite ends of a rigid vertical bar. In airborne usage two magnetometers are flown at a vertical spacing of about 30 m (Fig. 5.44c). The difference in outputs of the two instruments is recorded. If no anomalous body is present, both magnetometers register the Earth’s field equally strongly and the difference in output signals is zero. If a magnetic contrast is present in the subsurface rocks, the magnetometer closest to the structure will detect a stronger signal than the more remote instrument, and there will be a difference in the combined output signals. The gradiometer emphasizes anomalies from local shallow sources at the expense of large-scale regional variation due to deep-seated sources. Moreover, because the gradiometer registers the difference in signals from the individual magnetometers, there is no need to compensate the measurements for diurnal variation, which affects each individual magnetometer equally. Proton-precession magnetometers are most commonly used in ground-based magnetic gradiometers, while optically pumped magnetometers are favored in airborne gradiometers.

5.5.3.3 The survey pattern

In a systematic regional airborne (or marine) magnetic survey the measurements are usually made according to a predetermined pattern. In surveys made with fixed-wing aircraft the survey is usually flown at a constant flight elevation above sea-level (Fig. 5.45a). This is the procedure favored for regional or national surveys, or for the investigation of areas with dramatic topographic relief. The survey focusses on the depth to the magnetic basement, which often underlies less magnetic sedimentary surface
5.5.4 Reduction of magnetic field measurements

In comparison to the reduction of gravity data, magnetic survey data require very few corrections. One effect that must be compensated is the variation in intensity of the geomagnetic field at the Earth’s surface during the course of a day. As explained in more detail in Section 5.4.3.3 this diurnal variation is due to the part of the Earth’s magnetic field that originates in the ionosphere. At any point on the Earth’s surface the external field varies during the day as the Earth rotates beneath different parts of the ionosphere. The effect is much greater than the precision with which the field can be measured. The diurnal variation may be corrected by installing a constantly recording magnetometer at a fixed base station within the survey area. Alternatively, the records from a geomagnetic observatory may be used, provided it is not too far from the survey area. The time is noted at which each field measurement is made during the actual survey and the appropriate correction is made from the control record.

The variations of magnetic field with altitude, latitude and longitude are dominated by the vertical and horizontal variations of the dipole field. The total intensity \( B_t \) of the field is obtained by computing the resultant of the radial component \( B_r \) (Eq. (5.38)) and the tangential component \( B_t \) (Eq. (5.39)):

\[
B_t = \sqrt{B_r^2 + B_t^2} = \frac{\mu_B m}{4\pi} \frac{\sqrt{1 + 3\cos^2\theta}}{r^3}
\]  
(5.45)

The altitude correction is given by the vertical gradient of the magnetic field, obtained by differentiating the intensity \( B_t \) with respect to radius, \( r \). This gives

\[
\frac{\partial B_t}{\partial r} = -3 \frac{\mu_B m}{4\pi} \frac{\sqrt{1 + 3\cos^2\theta}}{r^4} = -3 \frac{\partial B_t}{r^4}
\]  
(5.46)

The vertical gradient of the field is found by substituting \( r = R = 6371 \) km and an appropriate value for \( B_t \). It clearly depends on the latitude of the measurement site. At the magnetic equator (\( R = 30,000 \) nT) the altitude correction is about 0.015 nT m\(^{-1}\); near the magnetic poles (\( R = 60,000 \) nT) it is about 0.030 nT m\(^{-1}\). The correction is so small that it is often ignored.

In regional studies the corrections for latitude and longitude are inherent in the reference field that is subtracted. In a survey of a small area, the latitude correction is given by the north–south horizontal gradient of the magnetic field, obtained by differentiating \( B_t \) with respect to polar angle, \( \theta \). This gives for the northward increase in \( B_t \) (i.e., with increasing latitude)

\[
-\frac{1}{r^2} \frac{\partial B_t}{\partial \theta} = \frac{\mu_B m}{4\pi} \frac{1}{r^2} \frac{\partial}{\partial \theta} \sqrt{1 + 3\cos^2\theta} = \frac{3B_t \sin \theta \cos \theta}{r(1 + 3\cos^2\theta)}
\]  
(5.47)

The latitude correction is zero at the magnetic pole (\( \theta = 0^\circ \)) and magnetic equator (\( \theta = 90^\circ \)) and reaches a maximum value of about 5 nT per kilometer (0.005 nT m\(^{-1}\)) at intermediate latitudes. It is insignificant in small-scale surveys.
5.5 MAGNETIC SURVEYING

In some land-based surveys of highly magnetic terrains (e.g., over lava flows or mineralized intrusions), the disturbing effect of the magnetized topography may be serious enough to require additional topographic corrections.

5.5.5 Magnetic anomalies

The gravity anomaly of a body is caused by the density contrast \( \Delta \rho \) between the body and its surroundings. The shape of the anomaly is determined by the shape of the body and its depth of burial. Similarly, a magnetic anomaly originates in the magnetization contrast \( \Delta M \) between rocks with different magnetic properties. However, the shape of the anomaly depends not only on the shape and depth of the source object but also on its orientation to the profile and to the inducing magnetic field, which itself varies in intensity and direction with geographical location. In oceanic magnetic surveying the magnetization contrast results from differences in the remanent magnetizations of crustal rocks, for which the Königsberger ratio is much greater than unity (i.e., \( Q_n \gg 1 \)). Commercial geophysical prospecting is carried out largely in continental crustal rocks, for which the Königsberger ratio is much less than unity (i.e., \( Q_n \ll 1 \)) and the magnetization may be assumed to be induced by the present geomagnetic field. The magnetization contrast is then due to susceptibility contrast in the crustal rocks. If \( k \) represents the susceptibility of an orobody, \( k_0 \) the susceptibility of the host rocks and \( F \) the strength of the inducing magnetic field, Eq. (5.17) allows us to write the magnetization contrast as

\[
\Delta M = (k - k_0)F
\]

Some insight into the physical processes that give rise to a magnetic anomaly can be obtained from the case of a vertically sided body that is magnetized by a vertical magnetic field. This is a simplified situation because in practice both the body and the field will be inclined, probably at different angles. However, it allows us to make a few observations that are generally applicable. Two scenarios are of particular interest. The first is when the body has a large vertical extent, such that its bottom surface is at a great depth; the other is when the body has a limited vertical extent. In both cases the vertical field magnetizes the body parallel to its vertical sides, but the resulting anomalies have different shapes. To understand the anomaly shapes we will use the concept of magnetic pole distributions.

5.5.5.1 Magnetic anomaly of a surface distribution of magnetic poles

Although magnetic poles are a fictive concept (see section 5.2.2.1), they provide a simple and convenient way to understand the origin of magnetic field anomalies. If a slice is made through a uniformly magnetized object, simple logic tells us that there will be as many south poles per unit of area on one side of the slice as north poles on the opposite side; these will cancel each other and the net sum of poles per unit area of the surface of the slice is zero. This is no longer the case if the magnetization changes across the interface. On each unit area of the surface there will be more poles of the stronger magnetization than poles of the weaker one. A quantitative derivation shows that the resultant number of poles per unit area \( \sigma \) (called the surface density of poles) is proportional to the magnetization contrast \( \Delta M \).

The concept of the solid angle subtended by a surface element (Box 5.4) provides a qualitative understanding of the magnetic anomaly of a surface distribution of magnetic poles. Consider the distribution of poles on the upper surface with area \( A \) of a vertical prism with magnetization \( M \) induced by a vertical field \( B_z \), as illustrated in Fig. 5.46a. At the surface of the Earth, distant \( r \) from the distribution of poles, the strength of their anomalous magnetic field is proportional to the total number of poles on the surface, which is the product of \( A \) and the surface density \( \sigma \) of poles. Equation (5.2) shows that the intensity of the field of a pole decreases as the inverse square of distance \( r \). If the direction of \( r \) makes an angle \( \theta \) with the vertical magnetization \( M \), the vertical component of the anomalous field at \( P \) is found by multiplying by \( \cos \theta \). The vertical magnetic anomaly \( \Delta B_z \) of the surface distribution of poles is

\[
\Delta B_z = \frac{(\sigma A) \cos \theta}{r^2} \alpha(\Delta M) \Omega \quad (5.49)
\]

A more rigorous derivation leads to essentially the same result. At any point on a measurement profile, the magnetic anomaly \( \Delta B_z \) of a distribution of poles is proportional to the solid angle \( \Omega \) subtended by the distribution at the point. The solid angle changes progressively along a profile (Fig. 5.46b). At the extreme left and right ends, the radius from the observation point is very oblique to the surface distribution of poles and the subtended angles \( \Omega_1 \) and \( \Omega_2 \) are very small; the anomaly distant from the body is nearly zero. Over the center of the distribution, the subtended angle reaches its largest value \( \Omega_0 \) and the anomaly reaches a maximum. The anomaly falls smoothly on each side of its crest corresponding to the values of the subtended angles \( \Omega_1 \) and \( \Omega_2 \) at the intermediate positions. A measurement profile across an equal distribution of “north” poles would be exactly inverted. The north poles create a field of repulsion that acts everywhere to oppose the Earth’s magnetic field, so the combined field is less than it would be if the “north” poles were not there. The magnetic anomaly over “north” poles is negative.

5.5.5.2 Magnetic anomaly of a vertical dike

We can now apply these ideas to the magnetic anomaly of a vertical dike. In this and all following examples we will assume a two-dimensional situation, where the horizontal length of the dike (imagined to be into the page) is infinite. This avoids possible complications related to “end effects.” Let us first assume that the dike extends to
Box 5.4: Solid angles

A solid angle is defined by the ratio between the area of an element of the surface of a sphere and the radius of the sphere. Let the area of a surface element be $A$ and the radius of the sphere be $r$, as in Fig. B5.4. The solid angle $\Omega$ subtended by the area $A$ at the center of the sphere is defined as

$$\Omega = \frac{A}{r^2}$$  \hspace{1cm} (1)

The angle subtended by any surface can be determined by projecting the surface onto a sphere. The shape of the area is immaterial. If an element of area $A$ is inclined at angle $\alpha$ to the radius $r$ through a point on the surface, its projection normal to the radius (i.e., onto a sphere passing through the point) is $A \cos \alpha$, and the solid angle it subtends at the center of the sphere is given by

$$\Omega = \frac{A \cos \alpha}{r^2}$$  \hspace{1cm} (2)

A solid angle is measured in units of steradians, which are analogous to radians in planar geometry. The minimum value of a solid angle is zero, when the surface element is infinitesimally small. The maximum value of a solid angle is when the surface completely surrounds the center of the sphere. The surface area of a sphere of radius $r$ is $A = 4\pi r^2$ and the solid angle at its center has the maximum possible value, which is $4\pi$.

![Fig. B5.4 Definition of the solid angle $\Omega$ subtended by an area $A$ on the surface of a sphere with radius $r$.](image-url)

---

very great depths (Fig. 5.47a), so that we can ignore the small effects associated with its remote lower end. The vertical sides of the dike are parallel to the magnetization and no magnetic poles are distributed on these faces. However, the horizontal top face is normal to the magnetization and a distribution of magnetic poles can be imagined on this surface. The direction of magnetization is parallel to the field, so the pole distribution will consist of “south” poles. The magnetized dike behaves like a magnetic monopole. At any point above the dike we measure both the inducing field and the anomalous “stray field” of the dike, which is directed toward its top. The anomalous field has a component parallel to the Earth’s field and so the total magnetic field will be everywhere stronger than if the dike were not present. The magnetic anomaly is everywhere positive, increasing from zero far from the dike to a maximum value directly over it (Fig. 5.46b).

If the vertical extent of the dike is finite, the distribution of north poles on the bottom of the dike may be close enough to the ground surface to produce a measurable stray field. The upper distribution of south poles causes a positive magnetic anomaly, as in the previous example. The lower distribution of north poles causes a negative anomaly (Fig. 5.47b). The north poles are further from the magnetometer than the south poles, so their negative anomaly over the dike is weaker. However, farther along
However, as in other potential field methods, the inverse problem of determining these factors from the measured anomaly is not unique.

The profile the deeper distribution of poles subtends a larger angle than the upper one does. As a result, the strength of the weaker negative anomaly does not fall off as rapidly along the profile as the positive anomaly does. Beyond a certain lateral distance from the dike (to the left of L and to the right of R in Fig. 5.47b) the negative anomaly of the lower pole distribution is stronger than the positive anomaly of the upper one. This causes the magnetic anomaly to have negative side lobes, which asymptotically approach zero with increasing distance from the dike.

The magnetized dike in this example resembles a bar magnet and can be modelled crudely by a dipole. Far from the dike, along a lateral profile, the dipole field lines have a component opposed to the inducing field, which results in the weak negative side lobes of the anomaly. Closer to the dike, the dipole field has a component that reinforces the inducing field, causing a positive central anomaly.

5.5.5.3 Magnetic anomaly of an inclined magnetization

When an infinitely long dike is magnetized obliquely rather than vertically, its anomaly can be modelled either by an inclined dipole or by pole distributions (Fig. 5.48). The magnetization has both horizontal and vertical components, which produce magnetic pole distributions on the vertical sides of the dike as well as on its top and bottom. The symmetry of the anomaly is changed so that the negative lobe of the anomaly is enhanced on the side towards which the horizontal component of magnetization points; the other negative lobe decreases and may disappear.

The shape of a magnetic anomaly also depends on the angle at which the measurement profile crosses the dike, and on the strike and dip of the dike. The geometry, magnetization and orientation of a body may be taken into account in forward-modelling of an anomaly. However, as in other potential field methods, the inverse problem of determining these factors from the measured anomaly is not unique.
The asymmetry (or skewness) of a magnetic anomaly can be compensated by the method of reduction to the pole. This consists of recalculating the observed anomaly for the case that the magnetization is vertical. The method involves sophisticated data-processing beyond the scope of this text. The observed anomaly map is first converted to a matrix of values at the intersections of a rectangular grid overlaying the map. The Fourier transform of the matrix is then computed and convolved with a filter function to correct for the orientations of the body and its magnetization. The reduction to the pole removes the asymmetry of an anomaly (Fig. 5.49) and allows a better location of the margins of the disturbing body. Among other applications, the procedure has proved to be important for detailed interpretation of the oceanic crustal magnetizing responsible for lineated oceanic magnetic anomalies.

5.5.5.4 Magnetic anomalies of simple geometric bodies

The computation of magnetic anomalies is generally more complicated than the computation of gravity anomalies. In practice, iterative numerical procedures are used. However, the Poisson relation (Box 5.5) enables the computation of magnetic anomalies for bodies for which the gravity anomaly is known. This is most easily illustrated for vertically magnetized bodies, such as the following examples.

(1) Sphere. The gravity anomaly $\Delta g_z$ over a sphere of radius $R$ with density contrast $\Delta \rho$ and center at depth $z$ (representing a diapir or intrusion) is given by Eq. (2.83), repeated here:

$$\Delta g_z = \frac{4}{3} \pi G \Delta \rho R^3 \left( \frac{z}{z^2 + x^2} \right)^{3/2}$$

Assuming the same dimensions and a magnetization contrast $\Delta M_z$, the potential of the magnetic anomaly over a vertically magnetized sphere according to the Poisson relation is

$$W = \frac{\mu_0}{4\pi} \frac{\Delta M_z}{G \Delta \rho} \Delta g_z = \frac{1}{2} \frac{\mu_0 R^3 \Delta M_z}{G} \frac{z}{(z^2 + x^2)^{3/2}}$$  \hspace{1cm} (5.50)

By differentiating with respect to $x$ or $z$ we get the horizontal or vertical field anomaly, respectively. The vertical field magnetic anomaly $\Delta B_z$ of the sphere is

$$\Delta B_z = - \frac{\partial W}{\partial z} = - \frac{1}{2} \frac{\mu_0 R^2 \Delta M_z}{G} \frac{z}{(z^2 + x^2)^{5/2}}$$

$$= - \frac{1}{2} \frac{\mu_0 R^2 \Delta M_z}{G} \frac{(z^2 + x^2)^{3/2} - z(3/2)(z^2 + x^2)^{1/2}}{(z^2 + x^2)^{3/2}}$$  \hspace{1cm} (5.51)

$$\Delta B_z = \frac{1}{2} \frac{\mu_0 R^2 \Delta M_z}{G} \frac{(z^2 - x^2)}{(z^2 + x^2)^{3/2}}$$  \hspace{1cm} (5.52)

(2) Horizontal cylinder. The gravity anomaly $\Delta g_z$ over a cylinder of radius $R$ with horizontal axis centered at depth $z$ and with density contrast $\Delta \rho$ (representing an anticline or syncline) is given by Eq. (2.93). If the structure is vertically magnetized with magnetization contrast $\Delta M_z$, Poisson’s relation gives for the magnetic potential

$$W = \frac{\mu_0}{4\pi} \frac{\Delta M_z}{G \Delta \rho} \Delta g_z = \frac{1}{2} \frac{\mu_0 R^2 \Delta M_z}{G} \left( \frac{z}{z^2 + x^2} \right)$$  \hspace{1cm} (5.53)

The vertical magnetic field anomaly $\Delta B_z$ over the horizontal cylinder is

$$\Delta B_z = \frac{1}{2} \frac{\mu_0 R^2 \Delta M_z}{G} \left( \frac{z}{x^2} - \frac{x}{z^2} \right)$$  \hspace{1cm} (5.54)

(3) Horizontal crustal block. The gravity anomaly for a thin horizontal sheet of thickness $t$ at depth $d$ between horizontal positions $x_1$ and $x_2$ (Fig. 2.54 b), extending to infinity normal to the plane of observation, is given by Eq. (2.96). Let the width of the block be $2m$, and let the horizontal position be measured from the midpoint of the block, so that $x_1 = x - m$ and $x_2 = x + m$. Applying Poisson’s relation, we get the magnetic potential for a semi-infinite horizontal thin sheet of vertically magnetized dipoles, of thickness $t$ at depth $z$

$$W = \frac{\mu_0}{4\pi} \frac{\Delta M_z}{G \Delta \rho} \Delta g_z = \frac{\mu_0 \Delta M_z}{2\pi}$$

$$\left[ \tan^{-1} \left( \frac{x + m}{z} \right) - \tan^{-1} \left( \frac{x - m}{z} \right) \right]$$  \hspace{1cm} (5.55)
5.5 MAGNETIC SURVEYING

Box 5.5: Poisson’s relation

Poisson (1781–1840) observed a relationship between
the gravitational and magnetic potentials of a body, which allows
a simple method of computing magnetic field anomalies if
the gravity anomaly of the body is known.

Consider an arbitrary volume \( V \) with homogeneous
density and vertical magnetization (Fig. B5.5). If the
density of the body is \( \Delta \rho \) a small element with volume
d\( V \) has mass \( (\Delta \rho dV) \). The gravitational potential \( U \) at
a point on the surface at a distance \( r \) from the element is

\[
U = -\frac{G \Delta \rho dV}{r}
\]

The vertical gravity anomaly \( \Delta g_z \) of the volume
element is found by differentiating \( U \) with respect to \( z \)

\[
\Delta g_z = \frac{\partial}{\partial z} \left( -\frac{G \Delta \rho dV}{r} \right) = \frac{G \Delta \rho dV \frac{\partial}{\partial z} \left( \frac{1}{r} \right)}{r}
\]

If the body is vertically magnetized with uniform
magnetization \( \Delta M_z \), the magnetic moment of the
volume element is \( (\Delta M_z \, dV) \). The magnetic moment
is directed downward as in Fig. B5.5. The radius vector
from the element to a point on the surface makes an
angle \( (\pi - \theta) \) with the orientation of the magnetization.
The magnetic potential \( W \) at the point \( (r, \theta) \) is

\[
W = \frac{\mu_0 \Delta M_z \cos(\pi - \theta)}{4\pi} \frac{1}{r}
\]

Comparing Eq. (2) and Eq. (5) and eliminating the
volume \( \Delta V \) we get Poisson’s relation:

\[
W = \frac{\mu_0}{4\pi} \frac{\Delta M_z \, dV \frac{\partial}{\partial z} \left( \frac{1}{r} \right)}{r}
\]

This derivation for a small element is also valid for an
extended body as long as the density and magnetization
are both uniform.

Assume that the horizontal crustal block is made
up of layers of thickness \( t = dz \). If the top of the
block is at depth \( z_1 \) and its base at depth \( z_2 \), the magnetic
potential of the block is found by integrating
Eq. (5.55) between \( z_0 \) and \( z_1 \),

\[
W = \frac{\mu_0 \Delta M_z}{2\pi} \left[ \tan^{-1} \left( \frac{x + m}{z} \right) \right. \\
- \tan^{-1} \left( \frac{x - m}{z} \right) \left. \right] dz
\]

Differentiating with respect to \( z \) gives the vertical
magnetic field anomaly over the block:

\[
\Delta B_z = -\frac{\partial W}{\partial z} = -\frac{\mu_0 \Delta M_z}{2\pi} \left[ \tan^{-1} \left( \frac{x + m}{z} \right) \right. \\
- \tan^{-1} \left( \frac{x - m}{z} \right) \left. \right] dz
\]
three-dimensional bodies on solid angles subtended by surfaces of the body at the point of observation, as illustrated in Fig. 5.46. As was the case for gravity surveys, magnetic profiles normal to the strike of elongate bodies may be regarded as two-dimensional, as long as the third dimension of the body is large enough for variations normal to the profile to be negligible.

5.5.5.5 Effect of block width on anomaly shape

The effect of the width of a crustal block on anomaly shape is illustrated by use of Eq. (5.59) to model the vertical field magnetic anomaly of a vertically magnetized block with its top at depth 2.5 km and base at depth 3 km. The block is effectively a thin magnetized layer, similar to the source of oceanic magnetic anomalies. Three cases are considered here: a narrow block of width \( w = (2m) = 5 \text{ km} \) for which \( m/z_1 = 1 \), a block of width 10 km \( (m/z_1 = 2) \), and a wide block of width 40 km \( (m/z_1 = 8) \).

The narrowest block gives a sharp, positive central anomaly with negative side lobes (Fig. 5.50b), as explained in section 5.5.5.2. As the block widens with respect to its depth, the top of the central anomaly flattens (Fig. 5.50c), its amplitude over the middle of the block decreases, and the negative side lobes grow. When the block is much wider than the depth to its top (Fig. 5.50d), a dip develops over the center of the block. The positive anomalies are steep sided and are maximum just within the edges of the block, while the null values occur close to the edges of the block. The negative side anomalies are almost as large as the positive anomalies.

The pronounced central dip in the anomaly is due to the limited vertical thickness of the layer. If the layer is very wide relative to its thickness, the central anomaly may diminish almost to zero. This is because the angle \( (\alpha_1 - \alpha_2) \) subtended by the magnetized base of the layer is almost (but not quite) as large as the angle \( (\alpha_1 - \alpha_2) \) subtended by the top of the layer. For a very large width-to-thickness ratio, the central anomaly is zero, the edge anomalies separate and become equivalent to separate anomalies over the edges of the block.

Examination of Fig. 5.50a shows that the subtended angles \( (\alpha_1 - \alpha_2) \) and \( (\alpha_3 - \alpha_4) \), and thus the anomaly shape, depend also on the height of the measurement profile above the surface of the block. A low-altitude profile over the block will show a large central dip, while a high-altitude profile over the same block will show a smaller dip or none at all.

In contrast to the example of a thin layer described above, if the crustal block is very thick, extending to great depth, the angle \( (\alpha_1 - \alpha_2) \) is zero and the effects of the magnetization discontinuity (or pole distribution) on its base are absent. The shape of the anomaly is then determined by \( (\alpha_1 - \alpha_2) \) and is flat topped over a wide block.

5.5.6 Oceanic magnetic anomalies

In the late 1950s marine geophysicists conducting magnetic surveys of the Pacific ocean basin off the west coast of North America discovered that large areas of oceanic crust are characterized by long stripes of alternating positive and negative magnetic anomalies. The striped pattern is best known from studies carried out across oceanic ridge systems (see Fig. 1.13). The striped anomalies are hundreds of kilometers in length parallel to the ridge axis, 10–50 km in width, and their amplitudes amount to several hundreds of nanotesla. On magnetic profiles perpendicular to a ridge axis the anomaly pattern is found to exhibit a remarkable symmetry about the axis of the ridge. The origin of the symmetric lineated anomaly pattern cannot be explained by conventional methods of interpretation based on susceptibility contrast.

Seismic studies indicate a layered structure for the oceanic crust. The floor of the ocean lies at water depths
of 2–5 km, and is underlain by a layer of sediment of variable thickness, called seismic Layer 1. Under the sediments lie a complex of basaltic extrusions and shallow intrusions, about 0.5 km thick, forming seismic Layer 2A, under which are found the deeper layers of the oceanic crust consisting of a complex of sheeted dikes (Layer 2B) and gabbro (Layer 3). The magnetic properties of these rocks were first obtained by studying samples dredged from exposed crests and ridges of submarine topography. The rocks of Layers 2B and 3 are much less magnetic than those of Layer 2A. Samples of pillow basalt dredged near to oceanic ridges have been found to have moderate susceptibilities for igneous rocks, but their remanent magnetizations are intense. Their Königsberger ratios are commonly in the range 5–50 and frequently exceed 100. Recognition of these properties provided the key to understanding the origin of the lineated magnetic anomalies. In 1963 the English geophysicists F. J. Vine and D. H. Matthews proposed that the remanent magnetizations (and not the susceptibility contrast) of oceanic basaltic Layer 2 were responsible for the striking lineated anomaly pattern. This hypothesis soon became integrated into a working model for understanding the mechanism of sea-floor spreading (see Section 1.2.5 and Fig. 1.14).

The oceanic crust formed at a spreading ridge acquires a thermoremanent magnetization (TRM) in the geomagnetic field. The basalts in Layer 2A are sufficiently strongly magnetized to account for most of the anomaly measured at the ocean surface. For a lengthy period of time (measuring several tens of thousands to millions of years) the polarity of the field remains constant; crust formed during this time carries the same polarity as the field. After a polarity reversal, freshly formed basalts acquire a TRM parallel to the new field direction, i.e., opposite to the previous TRM. Adjacent oceanic crustal blocks of different widths, determined by the variable time between reversals, carry antiparallel remanent magnetizations.

The oceanic crust is magnetized in long blocks parallel to the spreading axis, so the anomaly calculated for a profile perpendicular to the axis is two dimensional, as in the previous examples. Consider the case where the anomalies on a profile have been reduced to the pole, so that their magnetizations can be taken to be vertical. We can apply the concept of magnetic pole distributions to each block individually to determine the shape of its magnetic anomaly (Fig. 5.51a). If the blocks are contiguous, as is the case when they form by a continuous process such as sea-floor spreading, their individual anomalies will overlap (Fig. 5.51b). The spreading process is symmetric with respect to the ridge axis, so a mirror image of the sequence of polarized blocks is formed on the other side of the axis (Fig. 5.51c). If the two sets of crustal blocks are brought together at the spreading axis, a magnetic anomaly sequence ensues that exhibits a symmetric pattern with respect to the ridge axis (Fig. 5.51d).

This description of the origin of oceanic magnetic anomalies is over-simplified, because the crustal magnetization is more complicated than assumed in the block model. For example, the direction of the remanent magnetization, acquired at the time of formation of the ocean crust, is generally not the same as the direction of the magnetization induced by the present-day field. However, the induced magnetization has uniformly the same direction in the magnetized layer, which thus behaves like a uniformly magnetized thin horizontal sheet and does not contribute to the magnetic anomaly. Moreover, oceanic rocks have high Königsberger ratios, and so the induced magnetization component is usually negligible in comparison to the remanent magnetization. An exception is when a magnetic
survey is made close to the magnetized basalt layer, in which case a topographic correction may be needed.

Unless the strike of a ridge is north–south, the magnetization inclination must be taken into account. Skewness is corrected by reducing the magnetic anomaly profile to the pole (Section 5.5.5.3). A possible complication may arise if the oceanic magnetic anomalies have two sources. The strongest anomaly source is doubtless basaltic Layer 2B, but, at least in some cases, an appreciable part of the anomaly may arise in the deeper gabbroic Layer 3. The two contributions are slightly out of phase spatially, because of the curved depth profiles of cooling isotherms in the oceanic crust. This causes a magnetized block in the deeper gabbroic layer to lie slightly further from the ridge than the corresponding block with the same polarity in the basaltic layer above it. The net effect is an asymmetry of inclined magnetization directions on opposite sides of a ridge, so that the magnetic anomalies over blocks of the same age have different skewnesses.

5.6 PALEOMAGNETISM

5.6.1 Introduction

A mountain walker using a compass to find his way in the Swiss Alps above the high mountain valley of the Engadine would notice that in certain regions (for example, south of the Septimer Pass) the compass-needle shows very large deviations from the north direction. The deflection is due to the local presence of strongly magnetized serpentinites and ultramafic rocks. Early compasses were more primitive than modern versions, but the falsification of a compass direction near strongly magnetic outcrops was known by at least the early nineteenth century. In 1797 Alexander von Humboldt proposed that the rocks in these unusual outcrops had been magnetized by lightning strikes. The first systematic observations of rock magnetic properties are usually attributed to A. Delesse (1849) and M. Melloni (1853), who concluded that volcanic rocks acquired a remanent magnetization during cooling. After a more extensive series of studies in 1894 and 1895 of the origin of magnetism in lavas, G. Folgerhaiter reached the same conclusion and suggested that the direction of remanent magnetization was that of the geomagnetic field during cooling. By 1899 he had extended his work to the record of the secular variation of inclination in ancient potteries. Folgerhaiter noted that some rocks have a remanent magnetization opposite to the direction of the present-day field. Reversals of polarity of the geomagnetic field were established decisively early in the twentieth century.

In 1922 Alfred Wegener proposed his concept of continental drift, based on years of study of paleoclimatic indicators such as the geographic distribution of coal deposits. At the time, there was no way of explaining the mechanism by which the continents drifted. Only motions of the crust were considered, and the idea of rigid continents ploughing through rigid oceanic crust was unacceptable to geophysicists. There was as yet no way to reconstruct the positions of the continents in earlier eras or to trace their relative motions. Subsequently, paleomagnetism was to make important contributions to understanding continental drift by providing the means to trace past continental motions quantitatively.

A major impetus to these studies was the invention of a very sensitive astatic magnetometer. The apparatus consists of two identical small magnets mounted horizontally at opposite ends of a short rigid vertical bar so that the magnets are oriented exactly antiparallel to each other. The assembly is suspended on an elastic fiber. In this configuration the Earth’s magnetic field has equal and opposite effects on each magnet. If a magnetized rock is brought close to one magnet, the magnetic field of the rock produces a stronger twisting effect on the closer magnet than on the distant one and the assembly rotates to a new position of equilibrium. The rotation is detected by a light beam reflected off a small mirror mounted on the rigid bar. The device was introduced in 1952 by P. M. S. Blackett to test a theory that related the geomagnetic field to the Earth’s rotation. The experiment did not support the postulated effect. However, the astatic magnetometer became the basic tool of paleomagnetism and fostered its development as a scientific discipline. Hitherto it had only been possible to measure magnetizations of strongly magnetic rocks. The astatic magnetometer enabled the accurate measurement of weak remanent magnetizations in rocks that previously had been unmeasurable.

In the 1950s, several small research groups were engaged in determining and interpreting the directions of magnetization of rocks of different ages in Europe, Africa, North and South America and Australia. In 1956 S. K. Runcorn put forward the first clear geophysical evidence in support of continental drift. Runcorn compared the directions of magnetization of Peronian and Triassic rocks from Great Britain and North America. He found that the paleomagnetic results from the different continents could be brought into harmony for the time before 200 Ma ago by closing the Atlantic ocean. The evaluation of the scientific data was statistical and at first was regarded as controversial. However, Mesozoic paleomagnetic data were soon obtained from the southern hemisphere that also argued strongly in favor of the continental drift hypothesis. In 1957 E. Irving showed that paleomagnetic data conformed better with geological reconstructions of earlier positions of the continents than with their present-day distribution. Subsequently, numerous studies have documented the importance of paleomagnetism as a chronicle of past motions of global plates and as a record of the polarity history of the Earth’s magnetic field.

5.6.2 The time-averaged geomagnetic field

A fundamental assumption of paleomagnetism is that the time-averaged geomagnetic field corresponds to that of
5.6 PALEOMAGNETISM

an axial geocentric dipole. The data in support of this important hypothesis come partly from studies of secular variation and partly from paleomagnetic observations in young rocks and sediments.

The dipole and non-dipole parts of the historic geomagnetic field are known to change slowly with time. Spherical harmonic analysis of the geomagnetic field (see Section 5.4.5.1) shows that the axis of the geocentric inclined dipole has drifted slowly westward at about 0.044–0.14° yr⁻¹ in the last 400 yr (see Fig. 5.36). This would correspond to a complete circuit about the rotation axis in 2500–8000 yr. The rates of westward drift of the historic non-dipole field are around 0.22–0.66° yr⁻¹ (see Fig. 5.37), giving a periodicity of 550–1650 yr. However, it is important to keep in mind that the historic records of the secular variation of the geomagnetic field cover only a fragment of a complete circuit, which is not enough to confirm cyclicity or to estimate a period. The record of earlier magnetic field intensity and direction must be inferred from archæomagnetism.

5.6.2.1 Archeomagnetic records of secular variation

Paleomagnetism is the study of the geomagnetic field recorded in rock magnetizations; archæomagnetism is the study of the geomagnetic field recorded in dateable historic artefacts. The age of an archæological relict, such as a pot or vase, can often be determined with reliable precision. The pot, and the oven in which it was fired, may carry a thermoremanent magnetization (TRM) acquired during cooling. The direction of the TRM can be measured easily and, if the attitude of the pot during firing is known or can be assumed, the inclination of the ancient magnetic field in which the artefact was made can be deduced. The same considerations apply to lava flows that can be dated from historic records.

The secular record of paleoinclination during the past 2000 yr is available for two regions in which many archæomagnetic studies have been carried out: southeastern Europe and southwestern Japan. These regions are 110° apart in longitude but lie in similar latitude ranges, 35–40°N. Smoothed curves through the observations show pseudo-cyclical changes with several maxima and minima (Fig. 5.52). The shapes of the curves are not distinctive, so correlation of individual extreme values is risky; however, comparison of the four numbered maxima and minima in the last 1400 yr suggests that the extreme values appear to occur about 400 yr earlier in Japan than in Europe. The equivalent period for a full circuit of the globe is 1300 yr. The pseudo-cyclicity is interpreted as the effect of westward drift of foci of the non-dipole field past the sampling site. More detailed analysis of the archæomagnetic data shows that the drift rates vary with the latitude of the observation site (see Fig. 5.37). The mean drift rate is 0.38° longitude per year, which is faster than the rate deduced from recent secular variation.

A subtle magnetic technique devised by Thellier in 1937 permits determination of the intensity of the magnetic field in which an object acquires a TRM. If it is assumed that the field was a dipole field, the strength of the dipole magnetic moment can be inferred. When applied to rocks or ancient artifacts, this type of analysis is called a paleointensity determination. The variation in strength of the geomagnetic field during the past 7000 yr is shown by the results of 3188 paleointensity measurements on dated archeological and geological samples (Fig. 5.53). Older paleointensity data exist, but their number is too small to be included without distorting the record. The number of artifacts decreases as one goes back in time, so to obtain significant mean values the data are grouped in 1000 yr intervals before 2000 BC and in 500 yr intervals after 2000 BC. This introduces a problem, because these time intervals are too short for the geomagnetic field to average to a dipole field. The archæological relicts records the total field at the time of cooling, which contains a substantial non-dipole component. However, by combining the 3188 paleointensity data with 13,080 inclination and 16,085 declination data from lake sediments, a global field model has been calculated with Schmidt coefficients (Eq. (5.34)) up to order and degree 10. The coefficients with \( n = 1 \) from this model give the dipole magnetic moment (Eq. (5.41)). The analysis yields a smooth continuous record of the variation of the strength of the geomagnetic dipole during the past 7000 yr (Fig. 5.53). This record is displaced to slightly lower values than the direct measurements of paleointensity represented by the boxes, because it is free of non-dipole components. The dipole moment has fluctuated in the past 7000 years, but the record is too short to establish whether the changes are cyclical.
Westward drift of the dipole field implies systematic changes in the equatorial components of the dipole. If the changes are approximately cyclical, the mean long-term strength of the equatorial dipole measured at any given site would average to zero within a few multiples of $10^4$ yr. Similarly, if the secular variation of the non-dipole field can be assumed to be roughly periodic its mean value should average to zero within a few multiples of $10^3$ yr. According to these arguments, the only long-term component of the geomagnetic field that persists and is not averaged to zero within a few tens of thousands of years is the axial dipole component. The long-term equivalence of the Earth’s magnetic field with that of a dipole located at the center of the Earth and oriented along the rotation axis is a fundamental tenet of paleomagnetism; it is called the axial geocentric dipole hypothesis.

### 5.6.2.2 The axial geocentric dipole hypothesis

The evidence in support of the axial geocentric dipole hypothesis comes from paleomagnetic studies in modern deep-sea sediments and young igneous and sedimentary rocks. Pelagic sediments are deposited extremely slowly in the deep ocean basins. Sedimentation rates of 1–10 m Ma$^{-1}$ are common. The sediments acquire a post-depositional remanent magnetization (pDRM), which is an accurate record of the depositional field direction. The slow deposition of deep-sea sediments ensures thorough averaging of the magnetic field recorded. For example, at pelagic sedimentation rates a typical one-inch thick sample of deep-sea sediment averages paleomagnetic directions acquired during 2500–25,000 yr of deposition. The test of the axial geocentric dipole hypothesis in modern deep-sea sediments consists of comparing the inclination observed in sediment samples with the inclination expected for the latitude of the site where the sediment was sampled. The relationship between field inclination $I$, magnetic colatitude $\theta$ and latitude $\lambda$ was developed in Eq. (5.40) and is shown in Fig. 5.54a. The mean inclinations of remanent magnetization were measured in 52 deep-sea sediment cores of Plio-Pleistocene age taken from sites at different latitudes in the northern and southern hemispheres. The observed inclinations agree well with the values predicted by the theoretical curve for the axial geocentric dipole hypothesis (Fig. 5.54b).

Assuming the direction of the magnetic field recorded at a given site to be that of a dipole field, it is possible to calculate where the geomagnetic pole would need to be in order to produce the observed declination and inclination. This location is called the virtual geomagnetic pole (VGP) position. It is useful in computing where the pole lay in ancient times, the so-called paleomagnetic pole. The difference between a VGP and a paleomagnetic pole is illustrated by the following example for a recently extruded lava. Each sample from the lava formed in a short interval of time, and the field direction it records will be that of the total geomagnetic field at the site, combining axial dipole, non-dipole and non-axial dipole components. The VGP will therefore not coincide with the rotation axis. If data are collected from several flows of different ages, each will carry a slightly different record of the field. The computed VGP position will be different from flow to flow, and so the distribution of VGP will be scattered. If samples are measured from a large number of recent lava flows covering a long enough period of time
to average the non-dipole and non-axial dipole parts to zero, the mean direction of the collection will correspond to the field of an axial geocentric dipole. The pole position calculated from the mean direction of the collection of flows will agree with the rotation axis. This pole, representing an averaged value of the field, is called a paleomagnetic pole. The VGP represents a spot estimate of the field, including non-axial dipole components; the paleomagnetic pole represents an averaged field, corresponding to the axial geocentric dipole.

The paleomagnetic pole positions determined in studies of Plio-Pleistocene to Recent volcanic and sedimentary rocks covering the past 5 Ma lend further support to the axial geocentric dipole hypothesis. The distribution of paleomagnetic poles is clustered around the geographic pole and not around the present-day geomagnetic pole (Fig. 5.55). Statistical analysis shows that the mean of the paleomagnetic poles does not differ significantly from the geographic pole.

5.6.3 Methods of paleomagnetism

The requirement that the mean paleomagnetic pole position derived for a collection of rocks should represent the axial geocentric dipole is taken into account in the methodology of paleomagnetic analysis. This begins with the sampling of a rock formation on a hierarchical scheme designed to eliminate or minimize non-systematic
errors and to average out the effects of secular variation of the paleomagnetic field. At each hierarchical level, averaging and statistical analysis are carried out on the remanent magnetization vectors. Ideally, a paleomagnetic collection should contain a large number of samples per site. In practice, about 6–10 samples are enough to define the mean direction for a site; the mean values of typically 10–20 sites from the same formation are averaged to get a mean paleomagnetic direction for a formation or region.

A further assumption of paleomagnetism is that the natural remanent magnetization (NRM) of a rock was acquired at the time of formation of the rock (or at a known time in its history), and has since remained unaltered. In fact, the NRM is usually made up of several components acquired at different times, including during the procedures of sampling and preparation. Laboratory techniques must be applied that eliminate the undesirable components and isolate the primary magnetization. This process is loosely called “magnetic cleaning.”

The presentation of paleomagnetic directions measured in rock samples is made with the help of stereographic projection. This is a way of plotting three-dimensional directions by projecting them onto a plane. These plots have already been encountered in the analysis of first-motion studies of earthquakes (see Section 3.5.4.2). A direction is identified by the point where it intersects a unit sphere centered at the observation site. This converts a set of directions to a set of points on the surface of a sphere. The intersection point is then projected onto the horizontal plane to give a stereographic plot. This can be done in different ways. The Lambert equal-area projection is usually preferred in paleomagnetism as it avoids visually distorting the dispersion of directions. In geology, all directions are plotted on a stereogram as projections on the lower hemisphere. In paleomagnetic stereograms directions with positive (downward) inclinations are plotted as lower hemisphere projections; directions with negative (upward) inclinations are plotted with a different symbol as upper hemisphere projections.

5.6.3.1 Measurement of remanent magnetization

Measurements of the natural remanent magnetization of rocks with an astatic magnetometer were laborious and time consuming and the instrument has now fallen into disuse. In modern paleomagnetic laboratories more efficient spinner magnetometers and cryogenic magnetometers are in common use.

**Spinner magnetometers** originally consisted of a large sensor coil containing many turns of wire in which an alternating signal was induced by rotating the sample at high frequency (around 100 Hz) within the coil. Rapid rotation was needed because the voltage induced was proportional to the rate of change of flux in the coil. After phase-lock detection and electronic amplification of the signal, the calibrated output yielded two components of remanence in the plane normal to the rotational axis. The instrument was susceptible to electrostatic build-up but was capable of measuring magnetizations of around 10–3 A m\(^{-1}\) in 10–15 minutes.

The flux-gate spinner magnetometer is a subsequent refinement in which the sensor coil is replaced with flux-gate sensors. These detect directly the external magnetic fields of the sample. The signal strength is not dependent on rotational speed which could be reduced to about 5–10 Hz. The rotation of the sample gives a sinusoidal output. A large number of cycles can be averaged to reduce noise. The output is commonly digitized and stored in memory in a small on-line computer. The components of magnetization in the plane normal to the rotational axis are then determined by Fourier analysis. A computer-controlled flux-gate spinner magnetometer is capable of measuring a rock magnetization around 5 × 10\(^{-5}\) A m\(^{-1}\) in standard samples under optimum conditions. The complete measurement of a sample takes only a few minutes.

The cryogenic magnetometer is the most sensitive and rapid instrument in current use. Its sensor consists of a coil immersed in liquid helium. At this temperature (4 K) the coil is superconducting. A small change of magnetic field induces a comparatively large current, which because of the superconducting condition is persistent until the sample is removed. In line with the coil is a Josephson junction, which is a quantum-mechanical device consisting of a very thin element that allows the passage of current in distinct units proportional to a quantum of magnetic flux. By counting the number of flux jumps electronically, the external magnetic field of the rock specimen can be inferred, and from this its magnetization computed. Most cryogenic magnetometers contain orthogonal sets of coils and can measure two or three axes of magnetization simultaneously within a few seconds. The sensitivity of the instrument corresponds to a rock magnetization of 5 × 10\(^{-6}\) A m\(^{-1}\) in standard samples.

5.6.3.2 Stepwise progressive demagnetization

The natural remanent magnetization (NRM) of a rock may contain several components, some related to the geological history of the rock and others to the sampling and handling procedures. It is necessary to “magnetically clean” the natural magnetization so that the structure of the NRM can be analyzed and stable components isolated. This is done in a stepwise procedure, in which progressively more and more of the original magnetization is removed. There are two main methods of doing this.

The first method is progressive alternating field (AF) demagnetization. An alternating magnetic field can be produced in a coil by passing an alternating current through it. The field fluctuates between equal and opposite peak values. When a rock sample is placed in the alternating magnetic field, the grain magnetic moments with coercivities less than the peak value of the field are remagnetized in a new direction; the field cannot affect a magnetization component with coercivity higher than the peak field. The
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intensity of the alternating field is reduced slowly and uniformly to zero. This randomizes the part of the rock magnetization that has coercivities less than the peak value of the alternating magnetic field. The AF demagnetizing coil must be surrounded by magnetic shields or special additional coils to cancel out the Earth’s magnetic field; otherwise an anhysteretic remanent magnetization (ARM) is induced along the direction of this field.

The part of the remanence that remains after a demagnetization treatment has been “magnetically cleaned.” The direction and intensity of the remanent magnetization are affected. The demagnetization procedure is repeated using successively higher values of the peak alternating field, remeasuring the remaining magnetization after each step, until the magnetization is reduced to zero. Suppose that the NRM of a sample consists of two components AB and BC with different directions and different coercivity spectra (Fig. 5.56a, b). In the early stages of progressive demagnetization (steps 1–3) the “soft” component BC is first reduced to zero. The vector measured after each step in the progressive demagnetization is the sum of the “hard” component, which has not yet been affected by the field used, and the residual part of the soft component.

If the direction of the soft component is within 90° of the hard component the intensity decreases during this demagnetization interval; otherwise it may increase (Fig. 5.56c). The direction of the resultant vector changes continually in steps 1–3 (Fig. 5.56d). After removal of the soft component in step 3, higher alternating fields (steps 4–7) progressively reduce the hard component AB. During this stage the intensity decreases constantly but the direction remains consistent with little scatter, defining the “stable end-point” direction of the magnetization.

The effectiveness of the AF demagnetization method is limited by the strongest peak field that can be produced in the demagnetizing coil. This is commonly 0.1 T, although some equipment can reach around 0.3 T. These peak fields are well below the maximum coercivity of pyrrhotite and far below the coercivities of hematite or goethite. Thus AF demagnetization is not effective in demagnetizing components carried by these minerals. The method is most commonly used for rocks that contain magnetite as the main magnetic mineral.

An alternative method of “magnetic cleaning” is progressive thermal demagnetization. When a rock sample is heated to a given temperature $T$, magnetic components that have lower blocking temperatures than $T$ are thermally randomized. If the sample is now cooled in field-free space, this part of the NRM remains demagnetized. In stepwise thermal demagnetization the heating and cooling cycle is repeated with progressively higher maximum temperatures. The progressive destruction of the magnetization reveals the components present in the NRM as in Fig. 5.56. This method is often more effective than AF demagnetization, because it is only necessary to heat a sample above the highest Curie temperature of its constituent minerals to destroy all of the NRM. However, if the rock contains thermally unstable magnetic minerals, irreversible changes may complicate the thermal demagnetization method.

5.6.3.3 Analysis of magnetization components

The stability of a remanent magnetization during stepwise demagnetization can be demonstrated by plotting the remaining intensity after each step against the corresponding temperature or AF field, as in Fig. 5.56c; the directional stability can be controlled simultaneously by plotting the direction after each step on a stereogram (Fig. 5.56d). However, analysis of magnetization with an intensity plot and stereogram is outmoded. More sophisticated methods treat the magnetization as a vector and analyze the stability of its individual components.

The most powerful method of analysis of the structure and stability of a remanent magnetization involves constructing a vector diagram. The method was introduced by J. D. A. Zijderveld, a Dutch paleomagnetist, in the early 1960s. The magnetization at each stage of demagnetization is resolved into north (N), east (E) and vertical (V) components. Plots are then made of the north component against the east component, and of a horizontal component (north or east) against the vertical component. This is equivalent to projecting the vector onto the
Components of NRM that have distinct spectra of coercivities or blocking temperatures show as linear segments on a vector demagnetization diagram. The example in Fig. 5.57 shows three distinct linear segments on the horizontal and vertical projections. The slopes of the straight lines represent NRM components with different directions. A component removed below 150 °C is directed downward to the north, and so has normal polarity. The component is probably a soft overprint (perhaps a VRM, see Section 5.3.6.5) acquired in the present day or a recent field. A vector removed between 150 °C and 300 °C may represent a more ancient overprint; it is directed in a southerly, upward direction and thus has a reversed polarity. If a stable vector is left after demagnetization of less stable fractions, it is indicated by a straight line to the origin of each half of the vector diagram. This is the case for the component removed from 300 °C to 580 °C. It is interpreted as a stable primary component acquired when the field had reversed polarity.

If more than one magnetization component is present, it is possible that the spectra of coercivity or blocking temperature of the components may overlap partially. During demagnetization of the overlapping components the vector diagram exhibits a curved trajectory. If the spectra overlap completely, no straight segment can be determined. In this case the sample does not have a single stable magnetization component.

5.6.3.4 Statistical analysis of paleomagnetic directions

For the purposes of statistical analysis each paleomagnetic direction in a collection of samples is considered to have equal value and may be regarded as a unit vector. Each vector has unit length but a different direction. The end points of the vectors lie on the surface of a unit sphere and form a distribution of points. The statistical methods for evaluating paleomagnetic directions (or the distribution of points on a sphere) were developed in 1953 by Sir Ronald Fisher. He found that the best estimate of the mean direction of a population of \( N \) unit vectors is their vector mean, \( R \). To illustrate this point, consider five paleomagnetic directions, each represented by a unit vector (Fig. 5.58a). Usually the unit vectors are not parallel and when added vectorially their resultant has length \( R \approx 5 \) (Fig. 5.58b); its direction is the best estimate of the mean of the five paleomagnetic directions.

Fisher proposed that the probability density \( P(\theta, \kappa) \) of the angle \( \theta \) between an individual sample direction and the mean direction of the distribution is:

\[
P(\theta, \kappa) = \frac{k}{4\pi \sin h k} \exp(\kappa \cos \theta) \quad (5.60)
\]

The parameter \( \kappa \) is called the “precision parameter” or “concentration parameter.” It describes the dispersion of the directions, and is akin to the inverse of the variance of the distribution. Strictly speaking, \( \kappa \) is a property of an infinitely large population of directions. However, in paleomagnetic investigations only a small number of directions are usually sampled; it is assumed that they are representative for an infinite population. The parameters that are computed are approximate estimates of the true parameters of the population. Fisher showed that the best estimate \( (k) \) of the precision parameter \( \kappa \); (valid for \( k > 3 \)) is given by

\[
k = \frac{N - 1}{N - R} \quad (5.61)
\]

where \( R \) is the vector sum of the \( N \) unit vectors, computed as in Fig. 5.58b. When \( k \) (or \( \kappa \)) is zero, the directions
are uniformly or randomly distributed. A badly scattered set of directions has a small value of \( k \); large values of \( k \) apply to tightly grouped directions (Fig. 5.58c).

As in other statistical situations, the scatter is a property of the distribution of directions. It is described by the angular standard deviation, which is proportional to \( 1/\sqrt{k} \). However, it is usually more important to describe how well the mean direction is defined. Keep in mind that we do not know the true mean direction; we have only made an estimate of it, based on the available data. The true mean may differ by several degrees from our estimate. However, if we know that with 95% certainty the true mean lies within, say, \( 7^\circ \) of our estimate, we can draw a cone with semi-angle of \( 7^\circ \) about the estimated mean direction. The cone is said to define the confidence limits of the mean at the 95% probability level. The size of the confidence limit depends on the number of directions \( N \) in the distribution and their dispersion parameter \( k \). The semi-angle of the cone of confidence is denoted \( \alpha_{95} \) and is given approximately by

\[
\alpha_{95} = \frac{140}{\sqrt{Nk}}
\]

We could select any level of confidence to describe how well the mean is defined. However, two levels are common in statistics: the 95% (significant) and the 99% (highly significant) levels. In paleomagnetism the level of 95% confidence is used. This means that there is a 95% probability that the true mean of the distribution lies within this cone about the estimated mean direction.

5.6.3.5 Field tests of magnetization stability

If possible, paleomagnetic sampling includes a field test that can establish the stability of the magnetization stability over geological time. This was especially important in the early days of paleomagnetism when the laboratory techniques of “magnetic cleaning” were not yet available. Pioneering researchers devised some ingenious tests of paleomagnetic stability based on field observations. The fold test and reversals test still serve as the best ways to demonstrate the stability of a remanent magnetization through the aeons of geological time and to verify the timing of its acquisition.

The fold test is perhaps the most important paleomagnetic field test. It is applied to samples taken from beds that were originally horizontal and have been tilted by later tectonic effects. If the paleomagnetic direction in the rock is stable, it will experience the same rigid-body rotation as the tilted strata; its direction will vary around the fold (Fig. 5.59a, layer A). This is called a pre-folding magnetization. On the other hand, if the magnetization was acquired by the rock after it was folded, it will have a uniform direction at all points of the fold (Fig. 5.59a, layer B).
post-folding magnetization. A third situation is common, in which the magnetization is acquired during the tectonic event; in this case the direction of magnetization changes around the fold but by a smaller amount than the folding. This is called a synfolding magnetization.

In practice, the fold test consists of comparing the directions before applying tilt corrections with the directions after unfolding the tilted beds. If samples with a stable magnetization are taken from all parts of the fold, their uncorrected directions should be smeared out. After correction for bedding tilt, the dispersion of directions should be reduced, and the directions should group around a common direction, which is the pre-folding direction of the formation (Fig. 5.59b, layer A). This is called a positive fold test. If the magnetization is unstable or is due to post-folding remagnetization, the tilt corrections will increase the scatter of the distribution of directions (Fig. 5.59b, layer B); this is a negative fold test.

An application of the fold test is shown in Fig. 5.60 for 12 sites of the Scaglia Rossa limestone from southern Umbria (Italy). The sites were collected on different limbs of long anticlinal structures. Mean directions were computed for about 10–12 “magnetically cleaned” samples at each site. The uncorrected directions are quite scattered, with a confidence cone ($\alpha_{95}$) equal to 11°. After correcting each site for the local tilt of the bedding the data are much better grouped, and the confidence cone is reduced to 6°. The concentration parameter increases significantly, from 14.6 to 46.3, indicating that the magnetization was acquired before the folding.

The conglomerate test is a field test of stability that is rather seldom used. Suppose that we are investigating a limestone formation and that we discover a conglomerate containing cobbles of the limestone (Fig. 5.59a, layer C). Assuming that the cobbles have been randomly re-oriented by the processes of erosion, transport and re-deposition, their paleomagnetic directions, if stable, should be randomly distributed. If a systematic direction is found, the magnetization of the limestone may have a large secondary component.

The baked contact test is important in igneous rocks. During intrusion of a dike or sill the adjacent layers of the host rock are baked by contact with the hot lava and acquire a TRM when they cool. In general the magnetic minerals in the lava will differ in composition and grain size from those in the host rock. If samples taken from the lava and contact zone of the host rock have the same magnetization direction (Fig. 5.61b), the lava carries a stable paleomagnetic vector. If they are different (Fig. 5.61b), one of the magnetizations is unstable; alternatively, either
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5.6.4 Paleomagnetism and tectonics

Paleomagnetism has made important contributions in documenting local and regional tectonic motions as well as the motions of lithospheric plates. The reason for the failure of the reversals test in Fig. 5.62c was ascribed to local tectonic disturbances within a region. To make use of paleomagnetic data on a larger scale the observed directions must be compared to suitable reference directions. A reference direction can be computed, if it is known where the paleomagnetic pole was in the geological past. The history of paleomagnetic pole positions can be established on a continental scale.

Pole movements from central and southern Europe document the effects of large-scale tectonics
Data of Late Paleozoic to Late Cretaceous age are represented in this analysis. During this long time interval large amounts of motion of the European and African plates have taken place. As a result the inclinations measured at the indicated sites show large variations. However, the reference declinations for sites in Europe do not vary much during this time; this is also true for the African reference declinations. There is a large difference between the north–northeast pointing European declinations and the northwest directed African declinations (Fig. 5.63). The paleomagnetic declinations observed at sites in central and southern Europe show distinct affinities. North and west of a crude line through the Alpine chain the paleomagnetic declinations agree with those expected for the European continent. The declinations observed south of the Alps, on the Italian peninsula and Sicily are oriented toward the northwest, in agreement with directions expected for the African continent. The pattern of paleomagnetic data supports a tectonic interpretation of the Italian peninsula and adjacent regions of the Adriatic as a northern promontory of the African plate.

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5.6.4.1 Location of the virtual geomagnetic pole

Paleomagnetic poles are computed as the average of virtual geomagnetic pole (VGP) positions calculated for a number of samples at a site. The VGP position is where the pole of a geocentric magnetic dipole would need to be in order to give the observed declination \( D \) and inclination \( I \) of the remanent magnetization measured in the sample. The method of computation of the VGP position is illustrated in Fig. 5.64. First, from the inclination of the magnetization (i.e., the paleofield) we can calculate how far away the VGP was at the time the rock magnetization was acquired. The angular distance to the pole \( p \), assuming a dipole magnetic field, is obtained by using the relationship between inclination and polar angle (Fig. 5.54a).

The value of \( p \) determines the radius of a small circle centered on the paleomagnetic sampling site at latitude \( \lambda_s \) and longitude \( \phi_s \). The circle is the locus of all possible VGP positions that could give the observed inclination \( I \) at the site. We next have to decide which point on the small circle is the VGP position. The declination of the remanent magnetization is the angle between geographic north and the horizontal direction to the ancient magnetic pole. In this case the declination defines a meridian (or great circle) which passes through the sampling site and makes an angle \( D \) with the north–south meridian (Fig. 5.64). The place where this great circle intersects the small circle with radius \( p \) is the location of the virtual geomagnetic pole. Its latitude \( \lambda_p \) and longitude \( \phi_p \) can be computed exactly from trigonometric formulas (Box 5.6):

\[
\sin \lambda_p = \sin \lambda_s \cos p + \cos \lambda_s \sin p \cos D
\]

\[
\cos \phi_p = \frac{\sin p \sin D}{\sin \lambda_p}
\]
5.6.4.2 Apparent polar wander paths

The observation that paleomagnetic poles obtained from rocks of Pleistocene and Pliocene age are closely grouped about the geographic pole (see Fig. 5.55) is in agreement with the axial geocentric dipole hypothesis. However, when paleomagnetic pole positions are calculated for old rocks from the same continent, they group far away from the geographic pole. This is illustrated by the positions of paleomagnetic poles from the stable European craton. Pliocene and Pleistocene poles group close to the geographic pole but Permian poles are located about 45° away (Fig. 5.65). If the axial dipole hypothesis is valid for rocks of all ages, the pole distributions imply that the geographic pole for Europe in the Permian period (about 250–290 Ma ago) lay far from its present position. An alternative interpretation is that the geographic pole has not changed, but the European continent has moved relative to the pole. This suggests that the position about which the Permian poles now cluster was on the rotation axis in the Permian period. The European continent has subsequently moved to its present-day position with regard to the rotation axis.
Paleomagnetic data allow us to resolve the ambiguity. If paleomagnetic pole positions are computed for rocks of different ages from the same continent, they plot systematically along an irregular, curved path. It appears as though the paleomagnetic pole has moved slowly along this path towards the present rotation axis. The apparent motion of the paleomagnetic pole is called apparent polar wander (APW) and the path is called an apparent polar wander path. The paleomagnetic data from a particular continent define a unique APW path for that continent, and each continent has a different APW path. Thus, we have a European APW path, African APW path, North American APW path, and so on.

A schematic plot of the European and North American APW paths since the Late Paleozoic shows clearly distinct curves (Fig. 5.66). Each APW path lies on the opposite side of the geographic pole from the continent to which it belongs. Keeping the axial geocentric dipole hypothesis in mind, it is obviously impossible that the paleomagnetic pole (i.e., the Earth’s rotation axis) could have moved simultaneously along two different APW paths. The two APW paths evidently represent the separate motions of the European and North American continents relative to the rotation axis. They constitute paleomagnetic evidence for “continental drift.”

5.6.4.3 Paleogeographic reconstructions using APW paths

A more detailed plot of the two APW paths for the time before the Late Jurassic (Fig. 5.67a) shows strong similarities in their shapes, particularly for the time from the Upper Carboniferous to the Upper Triassic. It is possible to overlay these two segments of the APW paths by moving Europe (including Russia west of the Ural mountains) and North America into different positions relative to each other (Fig. 5.67b). For the time represented by the overlap of the APW paths the two continents formed part of a larger “supercontinent”, called Euramerica. When the adjacent part of Asia east of the Urals is included, the continents in the northern hemisphere form an earlier landmass called Laurasia. The present separation of the APW paths (Figs. 5.66, 5.67a) is interpreted as evidence for relative plate tectonic motion between Europe and North America that has taken place since the end of the interval for which the APW paths overlap well, i.e., since the Early Jurassic.

In order to bring the two APW paths into coincidence we have to move Europe relative to North America (or vice versa) so as to close the present gap between the continents. As shown in Section 1.2.9, the relative motion of plates on the surface of the spherical Earth is equivalent to a relative rotation about an Euler pole of rotation. The computer-generated “Bullard” fit of the 500 fathom contour lines on opposites sides of the North Atlantic ocean (see Section 1.2.2.2) can be obtained by displacing Europe toward North America by a clockwise rotation through 38° about the Euler pole located at 88.5°N 27.7°E, which by coincidence is very close to the present-day geographical pole (Fig. 5.67b). The APW path of a continent is constrained to move with the continent. If the European APW path is also rotated by 38° clockwise about the same Euler pole, the observed overlap of the Upper Carboniferous to Upper Triassic sections of the European and North American APW paths is obtained. Later segments of the paths diverge, indicating relative...
motion between the continents. The Late Jurassic pole position corresponds to the Earth’s rotation axis at that time. Circles of paleolatitude about the North American paleomagnetic pole emphasize the paleogeographic reconstruction of the relative positions of Europe and America in the Late Jurassic.

The interpretation of APW paths is not always as clear-cut as in this example. Consider the situation of the paleomagnetic pole $P$ for continental plate $C$, which is rotated through an angle $\Omega$ about the Euler pole of rotation, $E$. First, let the plate lie between the paleomagnetic pole and the rotation pole (Fig. 5.68a). The rotation of the plate from $C$ to $C'$ causes the paleomagnetic pole $P$ to move to $P'$. The arc $PP'$ of the polar motion is longer than the arc $CC'$ of the true plate motion. Next, consider what happens when the paleomagnetic pole lies between the plate and the Euler pole (Fig. 5.68b). In this case the plate moves through a large distance but the paleomagnetic pole moves only a small distance. In the extreme case where the paleomagnetic and Euler poles coincide, the plate rotation does not move the paleomagnetic pole at all. Under these special conditions plate motion leaves no trace in the APW path of the plate.

Clearly, the interpretation of an APW path as a record of plate motion relative to the geographic axis must be made with caution. The rate of motion of the pole along an APW path cannot be simply equated with the rate of motion of the parent continent or global plate. It follows that similarity of APW paths does not imply a unique solution for former relative plate positions. However, if two continents once belonged to the same plate for some length of time, they should have acquired the same APW path for this time. Matching the present APW paths of the continents for the time they were on the same plate should give a unique reconstruction of the earlier positions of the continents relative to each other. To avoid
Ambiguities, additional independent evidence (such as palaeoclimatic data, or computer matching of coastlines) must be utilized in conjunction with palaeomagnetic data for making such reconstructions.

5.6.4.4 Palaeomagnetism and continental drift

The nineteenth century geologist Eduard Suess deduced the existence of a great Late Palaeozoic continent, which he called Gondwanaland (Section 1.2.1). It was composed of Africa, Antarctica, Arabia, Australia, India and South America. In 1912, Wegener went a step further by postulating that all the present continents lay close together during the Late Palaeozoic, forming a single great continent that he called Pangaea. Wegener’s concept was based on palaeoclimatic evidence, the matching of Carboniferous coal belts and of regions of Palaeozoic glaciation in the different continents. Subsequently, additional geological evidence for the existence of Gondwanaland and Pangaea during the Late Palaeozoic and Early Mesozoic accumulated from the fields of sedimentology, palaeontology and tectonics. The earlier great continents were presumed to have dispersed to their present-day location by the process of continental drift. Unfortunately, Wegener was unable to offer a satisfactory driving mechanism for continental drift, and some of his ideas were found to be extreme. Scepticism among geophysicists and geologists brought Wegener’s theories into disrepute.

Interest in continental drift was re-awakened in the 1950s by the development of palaeomagnetism. Soon thereafter, one of the most convincing palaeomagnetic evidence for continental drift was obtained from the “southern continents.” Researchers found that Mesozoic palaeomagnetic pole positions of the same age from these continents were very dissimilar. In landmark contributions E. Irving showed that the palaeomagnetic poles of the southern continents were incompatible with the present-day arrangement of these continents, but that they agreed much better when the continents were rearranged to conform with a Gondwanaland reconstruction. Numerous palaeomagnetic investigations have subsequently provided a rich database that can be used to test the validity of reconstructions of former great continents at different times in their history.

The reconstructions are generally not made on palaeomagnetic evidence alone. Usually, a model is proposed, based on geometrical or geological grounds. The congruity of palaeomagnetic pole positions from the separate continents is then evaluated in their reconstructed positions. The model is adjusted iteratively until a configuration of the continents is obtained that gives minimum dispersion of the palaeomagnetic poles.

The evaluation of palaeomagnetic data from the Gondwanic continents, North America and Europe lends convincing support to the reconstructions and thereby to the continental drift hypothesis. From the Carboniferous to the Triassic, contemporary palaeomagnetic poles from the individual continents do not agree when the continents are in their present positions, but are more consistent when the great continent is reconstructed. In fact, the palaeomagnetic data are of high enough quality to suggest refinements to the purely geometric reconstructions. The Pangaea model in Fig. 5.69a corresponds closely to computer-assisted matches of the continental coastlines (see Section 1.2.2.2); it is referred to as Pangaea A1. It places the east coast of North America adjacent to the coast of northwest Africa. This configuration is supported well by Late Triassic and Early Jurassic palaeomagnetic poles. It is the generally accepted model of Pangaea immediately prior to its breakup in the Early Jurassic. However, older palaeomagnetic data of Permian and Carboniferous age (around 280 Ma) are less compatible with the Pangaea A1 model. Results of Late Permian to Middle Triassic age agree much better with a configuration referred to as Pangaea A2 (Fig. 5.69b), first proposed by R. Van der Voo and R. French in 1974. In this model, North America is much closer to South America and its eastern coast is opposite to western Africa. The transition from from the Late Permian Pangaea A2 to the Late Triassic Pangaea A1 configuration requires a large dextral shear between the continents in this time interval.

Pangaea may have had yet another configuration earlier in its history. In 1977 E. Irving showed that results of Carboniferous and Early Permian age agree better for a Pangaea configuration in which the east coast of North America is adjacent to the west coast of South America (Fig. 5.69c). This model, Pangaea B, is possible because...
Paleomagnetic longitudes are much more poorly constrained than paleolatitudes. The change from Early Permian Pangaea B to Late Permian Pangaea A2 requires a huge dextral megashear between Laurasia and Gondwana.

The models Pangaea A1, A2 and B are each consistent with paleomagnetic data for the different times of the reconstructions, which span about 100 Ma. None of the models accounts for the apparent polar wander paths of the individual continents over the whole interval of time from Early Permian to Late Triassic or Early Jurassic. Instead, the differences between the models imply that Pangaea was not a static great continent for this entire time interval, but that internal motions took place between the constituent continents.

Paleomagnetic data can be used to reconstruct the relative positions of continents during any time interval with enough good paleomagnetic data. APW paths (e.g., Fig. 5.66) can be determined fairly precisely by averaging the best available pole positions in 20–40 Ma time windows. Optimum fitting of APW paths of different continents allows reconstructions to be made for the time represented by the matching segments (Fig. 5.67). When this procedure is applied to paleomagnetic data covering the last 375 Ma, a picture of continental drift since the Middle Devonian is obtained (Section 1.2.2.3). According to this scenario, the supercontinents Laurasia and Gondwana, which were still separated by the Hercynian Ocean (Fig. 5.70) in the Devonian, collided in the Carboniferous to form Pangaea. The paleomagnetic reconstruction of continental drift for older epochs becomes tenuous because reliable paleomagnetic data become scarcer. The derivation of durable reconstructions for the Early Paleozoic and Precambrian will be a long and painstaking process.

The positions of continents since the breakup of Pangaea can also be obtained from analysis of APW paths. However, more precise reconstructions can be made by using a different form of paleomagnetic data, namely the record of geomagnetic polarity. Sea-floor spreading has imprinted this record in the oceanic crust, creating lineated magnetic anomalies. Matching coeval anomalies allows us to trace the motions of the lithospheric plates since the Middle Jurassic and describe the drift of the continents which they transport.

5.7 GEOMAGNETIC POLARITY

5.7.1 Introduction

The earliest demonstration that the geomagnetic field has changed polarity in the past was made by the French scientists P. David and B. Brunhes. In 1904–6 they described the magnetic properties of young lava flows in the Massif Central region of France. They found that clays baked by the lava flows had the same direction of remanent magnetization as the lavas. Moreover, when the magnetization direction in the lava was opposite to that of the present-day field, the same was the case in the baked clay. The opposite polarities were interpreted as evidence that the geomagnetic field can reverse its polarity.

A Japanese scientist, M. Matuyama, was the first to associate the polarity of remanent magnetization in lavas with their age, determined stratigraphically. In 1929 he reported finding young Quaternary lavas with magnetization directions close to the present-day field direction, whereas the directions in older Quaternary and Pleistocene lavas were clustered about an antipodal direction. He also found that one of three samples of Miocene basalt was magnetized oppositely to the other two. Matuyama’s interpretation was that geomagnetic polarity had changed several times during Late Tertiary time.

The idea that geomagnetic polarity could change was controversial, and for many years sceptics sought alternative interpretations. Scientists realized that the observed reversed polarities might have a mineralogical explanation. Indeed, some ferromagnetic minerals, because of their
composition and structure, can acquire a thermoremanent magnetization exactly opposite to the field direction. This mechanism is called self-reversal of magnetization. It has been described in lavas in which the ferromagnetic minerals are particular forms of titanohematite. Fortunately, it is a rather rare phenomenon. Most records of polarity reversals have been found to be a feature of the geomagnetic field.

To envisage what a reversal of geomagnetic polarity means, imagine that the geomagnetic dipole inverts its direction. At present the axial geocentric dipole points from the northern hemisphere towards the southern hemisphere; a polarity reversal would orient the dipole in the opposite direction. At each point on the surface the magnetic inclination $I$ changes sign and the declination $D$ changes by $180^\circ$; for example, a normal direction $\{I = 40^\circ, D = 30^\circ\}$ might change to a reverse direction $\{I = -40^\circ, D = 210^\circ\}$. A polarity reversal is a global event, experienced simultaneously all over the Earth. Thus, geomagnetic reversals provide a convenient means of stratigraphic correlation and dating.

5.7.1.1 Geomagnetic polarity transitions

The change of polarity from one sense to the opposite one is called a polarity transition. Paleomagnetic records of polarity transitions have been observed in radiometrically dated lava sequences and in deep-sea sediments with known deposition rates. These records indicate that the duration of a polarity transition is about 3.5–5 ka. This is much shorter than the length of the interval of constant polarity before or after the transition, which may last for hundreds of thousands or millions of years.

It is not yet known for sure how the geomagnetic field behaves during a polarity transition. The dipole field is dominant before and after a transition, but it is not certain that this is the case during the transition. Detailed analyses of field behavior during a polarity transition usually show a notable decrease in field intensity (Fig. 5.71); this is observed in volcanic and sedimentary records of reversals. Possibly the dipole component disappears, granting more importance to higher-order quadrupole or octupole field configurations. There seems to be stronger evidence that, even though its intensity decreases, the transitional field is still dominantly that of a dipole. If this can be assumed, the position of the virtual geomagnetic pole (VGP) of the transitional dipole field can be calculated. During a polarity transition the VGP position changes progressively. It appears to move systematically relative to the Earth’s rotation axis, defining a path from one polar region to the opposite one. The transitional paths of many reversals appear to define two longitudinal belts, one over the Americas and an antipodal belt over Southeast Asia. However, many other transitions do not pass over these two belts. It has not been established conclusively that a path over the Americas or Southeast Asia is a preferred feature of polarity transitions.

5.7.1.2 Geomagnetic polarity intervals

Long intervals of constant normal or reversed polarity, originally called polarity epochs, are referred to as polarity epochs (Fig. 5.72); they last typically from 50 ka to 5 Ma. The polarity chron is interrupted at irregular intervals by shorter polarity subchrons (originally called events) lasting for 20–50 ka. At times the polarity record
5.7 GEOMAGNETIC POLARITY

shows large departures of the magnetic pole from normal or reversed polarity, but the polarity does not change completely; the pole wanders into equatorial latitudes but returns to its initial location on the rotation axis. The departure is short lived, lasting less than 10 ka, and the phenomenon is called a magnetic excursion. The irregular pattern of polarity intervals in any sequence provides a kind of geological fingerprint which can be used under favorable circumstances to date and correlate some types of sedimentary rocks. This procedure is called magnetic polarity stratigraphy, or magnetostratigraphy.

5.7.2 Magnetostratigraphy in lavas and sediments

In the 1950s the methods of dating rocks took a giant step forward with the development of improved techniques for dating rocks radiometrically. The potassium–argon method (Section 4.1.4.4) was applied to the determination of accurate ages for Pliocene and Pleistocene lava samples from flows that were also sampled for paleomagnetic purposes. The polarity of the thermoremanent magnetization of the lava was found to correlate with its age. There were distinct intervals of time in which the field polarity was the same as at present, and these were separated by intervals of exactly opposite polarity. At first the data were sparse, and in the earliest interpretations it was thought that the field changed polarity quite regularly, roughly once every million years (Fig. 5.73). Gradually, however, a more complex history evolved. Long epochs of a given polarity were found to contain much shorter events of opposite polarity. The polarity epochs were named after important investigators of paleomagnetic polarity (Brunhes, Matuyama) and geomagnetism (Gauss, Gilbert) while the polarity events were named after the geographical location where they were first discovered (Jaramillo creek in New Mexico, Olduvai gorge in Africa, etc.). Countless studies of magnetic polarity in radiometrically dated lavas have established the history of geomagnetic polarity in the last 5 Ma (Fig. 5.74). If the polarity record in a rock sequence can be identified, its age can be determined by comparison with the dated sequence. For this reason a dated polarity sequence is called a geomagnetic polarity timescale.

There is a practical limit to the application of this technique. As can be seen by quick inspection, some of the polarity events last less than 50 ka. If a reasonable precision of 1–2% is assumed for potassium–argon dating, the error in determining the age of a lava sample that is about 5 Ma old amounts to 50–100 ka. This is longer than the duration of many short events. The dating error makes it impossible to associate the lava sample unambiguously with the correct polarity event. Extension of the magnetic polarity timescale beyond 5 Ma requires other methods.

In the middle 1960s the polarity record from lavas was augmented by a large amount of high-quality data acquired from young deep-sea sediments. The deep ocean basins provide a tranquil depositional environment, where sediments are deposited at rather uniform rates. Marine geologists routinely take cores of sediment with a special coring device (see Fig. 4.26) for sedimentological, geophysical and paleontological studies. The magnetostratigraphy
of a core is studied by measuring the direction of magnetization in small oriented samples from different depths in the core. Although deep-sea cores have vertical axes, they are not oriented in azimuth, so the declinations can only be determined relative to an arbitrary reference value. Polarity determinations are often based only on the inclination records. The boundaries between normal and reverse magnetozones are interpolated at the depths where the inclination is zero (Fig. 5.75). In equatorial cores, where inclinations are nearly zero, the relative changes in declination often give a good polarity record. The polarity records of numerous cores correlate well with the sequence found in contemporaneous young lavas. The sediment magnetic polarity records are independent of lithology; the same reversal occurs at different depths from core to core because of different sedimentation rates (Fig. 5.76). By correlating reversals with the radiometrically dated lava record, the sediment ages at the reversal depths are obtained. From the depths and ages it is a simple matter to calculate the incremental sedimentation rates in the core. In addition to providing sedimentation rates, magnetic polarity stratigraphy also yields the absolute ages of the first and last appearances of key fossils, and so gives absolute dates for paleontological fossil zones. A recent innovation is the use of astrochronology, based on the identification of Milankovich cycles (Section 2.3.4.5), to provide refined dating of sediments and their polarity record.

The magnetostratigraphic data from deep-sea sediment cores eliminated the lingering doubt that reversals observed in lavas may be due to a self-reversal mechanism. Lavas and sediments acquire their magnetizations by quite different mechanisms; the thermoremanent magnetization of a lava is acquired rapidly during cooling from high temperature, whereas the depositional or post-depositional remanent magnetization in a sediment is acquired slowly at constant ambient temperature. Although a self-reversal mechanism might be invoked to cast doubt on polarity changes in lavas, the argument is invalid for the remanence of deep-sea sediments. The common polarity sequence in lavas and sediments can only be explained as a record of the alternations of polarity of the Earth's magnetic field. Moreover, the same pattern of reversals is found regardless of geographical location, emphasizing that reversals are a global phenomenon.

5.7.3 Marine magnetic anomalies and geomagnetic polarity history

The striped magnetic anomaly patterns formed at oceanic ridges contribute to the compelling geophysical evidence in favor of the theory of global plate tectonics (see Section 1.2.5). Marine magnetic surveys and independent investigations of the rock magnetic properties of marine rocks and sediments have identified the source of the magnetic anomalies to be the basaltic Layer 2A of the oceanic crust.

Seismic evidence indicates that the oceanic crust has a vertically stratified structure (see Fig. 3.85). The uppermost part, seismic Layer 1, consists of a layer of slowly accumulating marine sediments; the thickness of the sediments increases progressively away from the ridge crest. The sediments are so weakly magnetic that they are essentially transparent to the Earth's magnetic field. The seismic Layer 2A consists of a 500 m thick layer of oceanic basalts that are extruded as submarine lava flows or intruded as dikes. These basalts are strongly magnetic and are chiefly responsible for the strong magnetic anomalies observed at the ocean surface. The metamorphosed basalts of the underlying Layer 2B are too weakly magnetic to have much signature. The rocks of the deeper gabbroic Layer 3 may be sufficiently magnetic to add to the skewness of the magnetic anomalies.

5.7.3.1 Marine magnetic anomalies

The origin of marine magnetic anomalies was explained by the Vine–Matthews–Morley hypothesis in 1963 (Section 1.2.5.1). Oceanic basalts were found to have strong and stable remanent magnetizations. Their Königsberger ratios are much larger than unity, so the remanent magnetizations are more important than the magnetization induced by the present-day geomagnetic field. The conventional method of interpreting surveyed magnetic anomalies assumed that the anomaly was due to the susceptibility contrast between adjacent crustal blocks. According to the Vine–Matthews–Morley hypothesis the oceanic magnetic anomalies arise from the contrast in remanent magnetizations between adjacent, oppositely magnetized crustal blocks. The remanent magnetization is acquired thermally by the basalts in oceanic crustal Layer 2A.

![Variations in magnetic inclination and inferred polarity with depth in two deep-sea sediment cores](image-url)
The main magnetic mineral in oceanic basalts is titanomagnetite (Section 5.3.2.1). A basaltic lava is initially at a temperature well above 1000 °C. Its titanomagnetite grains frequently have skeletal structures, indicating that they cooled and solidified so rapidly that there was not enough time for the formation of normal crystals. Eventually the temperature of the lava sinks below the Curie point of the titanomagnetite (around 200–300 °C) and the lava acquires a thermoremanent magnetization (TRM) in the direction of the Earth’s magnetic field at that time. Basalts formed contemporaneously along an active spreading ridge acquire the same polarity of magnetization. Long thin strips of similarly magnetized crust form on opposite sides of the spreading center. These elongated “crustal blocks” may be hundreds of kilometers in length parallel to the ridge axis and several tens of kilometers wide normal to the ridge, while Layer 2A – the strongly magnetic upper part – is only 0.5 km thick.

Sea-floor spreading persists for millions of years at an oceanic ridge. During this time the magnetic field changes polarity many times. The alternating field polarity leaves some blocks of oceanic crust normally magnetized while their neighbors are reversely magnetized. When the total intensity of the field is measured from a survey ship or aircraft, an alternating sequence of positive and negative anomalies is observed (see Fig. 1.13), which can be interpreted in terms of the crustal magnetization. The anomalies can be correlated almost linearly between parallel profiles across a ridge system; consequently, the stripe-like anomalies are often referred to as magnetic lineations.

5.7.3.2 Uniformity of sea-floor spreading

Each anomaly in a set of magnetic lineations derives from a crustal block (or stripe) that formed at a ridge and was subsequently transported away from the spreading center. A magnetized crustal block forms during a period of sea-floor spreading when the geomagnetic polarity was constantly normal or reversed, and therefore represents a polarity chron or subchron. The width of a particular block depends on the duration of the chron and the spreading rate at the ocean ridge.

The spreading rate can be determined easily close to a ridge (see Section 1.2.5.2). The edges of the magnetized crustal stripes correspond to the occurrences of polarity reversals, which can be correlated directly with the radiometrically dated sequence for the last 3–4 Ma determined in lavas on the continents or islands. A plot of the distance of a given polarity reversal from the spreading axis against the age of the reversal is nearly linear near the ridge; the slope of a best-fitting straight line gives the average half-rate of spreading at the ridge (see Fig. 1.15). This is half the full rate of plate separation, assuming that sea-floor spreading has been symmetric on each side of the ridge, which is often the case.

Accumulated evidence from marine magnetic profiles allows us to assess the constancy of sea-floor spreading at different ridge systems. It is thought to have been uniform for the longest time in the South Atlantic. A plot of the distance to a given anomaly in the South Atlantic against the distance to the same anomaly in the Indian, North Pacific and South Pacific oceans contains several long linear segments, representing constant rates of sea-floor spreading in both oceans defining the line (Fig. 5.77). A
change in gradient indicates a change in spreading rate in one ocean relative to the other. The plot does not exclude a synchronous worldwide change of spreading rate in all oceans, but this would be a rather unlikely occurrence. Clearly the rate of sea-floor spreading changes from time to time, but for long intervals it is a remarkably constant process.

5.7.3.3 The marine record of geomagnetic polarity history

Investigations of magnetic anomalies in all major oceanic areas have given a clear, consistent record of the history of geomagnetic polarity during the past 155–160 Ma. It consists of two sequences of polarity reversals represented by magnetic lineations and a long interval of constant normal polarity (Fig. 5.78). The sequences of chrons derived from magnetic lineations have been confirmed by magnetostratigraphic research.

The most prominent positive magnetic anomalies are numbered in turn from the youngest (anomaly 1 at an active ridge axis) to the oldest (anomalies 33–34 in the Late Cretaceous). The associated polarity chrons are identified by the same number and a letter to indicate the polarity. The polarity chrons in the latest sequence fall largely in the Cenozoic (see Fig. 4.2) and are identified by a leading letter C, which may be taken to stand for either “chron” or “Cenozoic.” The current Brunhes normal polarity interval corresponds to chron C1N; the reversed interval older than C1N is labelled chron C1R.

A phase of alternating polarity giving lineated magnetic anomalies precedes the Cretaceous Quiet Interval. It began late in the Middle Jurassic and continued until the middle of the Early Cretaceous. These Late Mesozoic oceanic anomalies are referred to as the M-sequence. To distinguish them from the later sequence the numbered chrons identified by a leading letter M. The youngest anomalies M0–M8 are numbered sequentially regardless of the magnetization polarity; older than M9 only reverse...
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Polarity chronos are numbered. The oldest securely identified chron in the sequence is M29N. Some older anomalies with low amplitudes have been interpreted as polarity chronos, but it has not yet been established that they represent polarity reversals rather than geomagnetic intensity fluctuations.

The oldest regions of the modern oceans correspond to oceanic crust formed approximately 180 Ma ago, when Pangaea broke up and the current episode of sea-floor spreading was initiated. The marine magnetic anomalies over these areas have subdued amplitudes and they do not form lineations. Either no reversals happened during the period of initial spreading or the oceanic crust has not been able to retain the record. The character of the geomagnetic field during this part of the Early Jurassic has not yet been definitively established.

5.7.4 Geomagnetic polarity timescales

The interpretation of marine magnetic anomalies provides the most continuous and reliable record of geomagnetic polarity since the Middle Jurassic. The length of the record greatly exceeds the length of the securely established, radiometrically dated magnetic polarity timescale, which covers only about the past 5 Ma. Thus, it is not possible to date most of the marine magnetic anomalies by direct correlation with a radiometrically dated polarity sequence. Knowledge of the spreading rate at a ridge system provides an alternative way of determining the age of the oceanic crust. Assuming that the rate of sea-floor spreading at the spreading center is constant, the age of a given anomaly can be computed by dividing its distance from the spreading center by the spreading rate. However, this is an unsatisfactory method because the extrapolation is many times longer than the baseline. A further method of dating the polarity record is by establishing the same polarity sequence in sedimentary rocks that are dated paleontologically. This has been achieved in several investigations of the magnetic polarity stratigraphy in pelagic carbonate rocks. Using known absolute ages of major stage boundaries as tie-levels, the ages of magnetic polarity chronos that are too old to be dated directly are calculated by interpolation.

5.7.4.1 Magnetostratigraphic calibration of polarity sequences

The correlation of the radiometrically dated polarity sequence in continental lavas (Fig. 5.74 left) with the magnetic anomaly record near to a spreading ridge was the original method used to date young marine magnetic anomalies. Subsequently, the polarity record was refined by detailed analysis of marine magnetic anomaly sequences. The polarity reversals were located in sediments where they could be dated precisely by counting Milankovitch cycles in the sediments. The combination of improved marine magnetic record and astrochronological dating gave a more reliable polarity timescale for 0–5 Ma (Fig. 5.74 right).

The independent confirmation and dating of older marine magnetic anomalies required combined magnetostratigraphical and paleontological studies in suitable rock formations. The dating of anomalies 29–34, which are the oldest found in the Cenozoic to Late Cretaceous sequence, illustrates this method. Along magnetic profiles in the Indian, North Pacific and South Atlantic oceans, the shapes of anomalies 29–34 are very different because of the different directions of the survey profiles and the orientations of the spreading axes. However, interpretation of the anomalies gives nearly identical crustal magnetization patterns (Fig. 5.79).

The Scaglia Rossa pelagic limestone in the Umbrian Apennines of Central Italy was deposited almost continuously from the Late Cretaceous to the Eocene. Rock magnetic analysis showed that the limestone contained an easily defined stable component of characteristic remanent magnetization. Samples were taken at approximately 0.5–1 m stratigraphic intervals in a long section through the limestone exposed in the Bottaccione gorge near Gubbio. The declinations and inclinations of the stable magnetization, after simple tectonic corrections, were used to calculate the latitude of the virtual geomagnetic pole (VGP) during deposition of the limestone. In times of normal polarity the VGP latitude is near 90°N, during reversed polarity it is near 90°S. The fluctuations of VGP latitude clearly define magnetozones of normal and reversed polarity (Fig. 5.80). The Gubbio polarity record in a 200 m thick section of pelagic limestone correlates almost perfectly with the oceanic polarity sequence derived from anomalies 29–34, measured in
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Fig. 5.80 Magnetoostratigraphy and biostratigraphy in the Bottaccione section at Gubbio, Italy (after Lowrie and Alvarez, 1977).

different oceans on profiles that are hundreds of kilometers long. The limestone magnetostratigraphy confirms independently this part of the oceanic magnetic polarity record.

Paleontological studies in the Gubbio section gave the locations of important planktonic foraminiferal zones and enabled the positions of the major stage boundaries to be located relative to the polarity sequence. The absolute ages of some important stage boundaries are known from independent radiometric and stratigraphic work. This enabled calculation of the absolute ages of the polarity reversals in the Gubbio section and, by correlation, the ages of the corresponding parts of the ocean floor. For example, the Santonian–Campanian boundary (with an age of about 83 Ma) lies close to the old edge of the reversed polarity chron C33R; the geologically important Cretaceous–Tertiary boundary (age 65 Ma) falls within reversed polarity chron C29R.

In this way the geomagnetic polarity sequence in the Late Cretaceous and Paleogene has been confirmed in magnetostratigraphic sections, and the locations of many major and minor stage boundaries have been correlated to the polarity sequence paleontologically (Fig. 5.81). Reliable absolute ages are available for some of the stage boundaries, which can then be used as calibration levels. The ages of magnetic reversals between the dated tie-points are computed by interpolation, giving a numerical geomagnetic polarity timescale (see Fig. 5.78). In the same way, overlapping magnetostratigraphic sections of Early Cretaceous and Late Jurassic age have permitted independent confirmation and dating of the M-sequence polarity record.

5.7.4.2 Reconstruction of plate tectonic motions

Once the ages of magnetic anomalies are known, a map showing the positions of dated anomalies is equivalent to a chronological map of the ocean basins (Fig. 5.82). The different rates of sea-floor spreading are evident from the separations of the isochrons. The oldest domains of the oceans (about 180 Ma) are found in the Atlantic ocean adjacent to the coastlines of North America and Africa, and in the North Pacific ocean. They are very much younger than the oldest rocks from the continents, which are up to 3.6 Ga old. The oceanic crust has been entirely produced since the onset of sea-floor spreading and the anomaly patterns reflect plate motions.

The past motions of the major plates can be obtained in some detail from the anomaly ages given by a geomagnetic polarity timescale. The relative positions of the continents at any time since the late Mid-Jurassic can be reconstructed by matching coeval marine magnetic anomalies formed at the same spreading center. The procedure is similar to the reconstruction of supercontinents.
by matching coastlines, 500 fathom depth contours or apparent polar wander paths.

The method is illustrated by the sea-floor spreading between North America and Africa in the Central Atlantic (Fig. 5.83). Anomaly 33 forms a long stripe on each side of the Mid-Atlantic Ridge and subparallel to it. The anomaly is due to the magnetization contrast between chron C33N and the reverse polarity chron C33R which marks the end of the Cretaceous Quiet Interval. Its age is about 81 Ma. The anomaly on the west side of the ridge was formed at the same time as the anomaly on the east side, when the newly formed crust was magnetized at the ridge axis. If the African and North American plates are moved closer together until the east and west anomalies overlap, or until they match along their lengths with minimum misfit, the continents will be brought into the same positions relative to each other that they occupied around 81 Ma ago. By repeating this process of matching dated anomalies it is possible to reconstruct the successive relative positions of the African and North American plates as they separated.

Marine magnetic anomalies in the North Atlantic can be used likewise to describe the relative plate motions between Europe and North America. A picture evolves of the separate histories of separation of Africa and Europe from North America. The differences between the European–North American and African–North American
plate motions permit the history of relative motion between Africa and Europe to be inferred. In the Late Cretaceous and Early Tertiary Africa moved eastwards in a giant shear motion relative to Europe, but since the Middle Tertiary the motion of Africa has been one of convergence and collision with the European plate. This is compatible with the formation of the Alpine fold belt and the present-day seismicity in the Alpine region.

5.7.5 Frequency of polarity reversals

Visual examination of the magnetic polarity record in Fig. 5.78 shows that the rate of polarity reversals has been quite variable in the last 160 Ma. A simple way to portray the reversal rate is to count the numbers of reversals in successive bins of equal duration (e.g., 4 Myr) and compute the average number per Myr. The plot of reversal rate against age (Fig. 5.84) is uneven, but has some distinct features.

The reversal frequency during the M-sequence averaged about 3–4 reversals/Myr. From about 130 Ma ago the reversal rate slowed down until the beginning of the Cretaceous Normal Polarity Superchron (CNPS) at about 121 Ma ago. There were no reversals for the following 38 Myr. After the end of the CNPS about 83 Ma ago the reversal rate was at first very low, but gradually speeded up. It reached a peak of about 5 reversals/Myr in the Late Miocene, about 10 Ma ago, and has since decreased slightly to about 4 reversals/Myr. The current interval of normal polarity (known as the Brunhes or chron C1N) has lasted 0.78 Myr, which is much longer than the mean length of polarity chron in the Late Tertiary.

In 1968 A. Cox theorized that reversals are random events, triggered by unknown mechanisms that affect the fluid motions in the Earth’s liquid outer core. With a random reversal process there would be no continuity between successive reversals; as soon as a reversal was completed, the next one would be as likely to occur immediately as at any time later. Because there would be no waiting period, this type of mechanism would generate a large number of very short polarity chron and a small number of long chron. The frequency distribution of polarity chron durations should decrease exponentially with increasing chron duration. In fact this model does not fit the observed lengths of polarity intervals precisely. The polarity sequence contains comparatively few short chron, a lot of medium duration chron and few long chron. This may imply that the process that causes reversals is not completely random. A distinct length of time may elapse after a reversal for the fluid motions to recover sufficiently to allow the next reversal to happen. However, the mechanism that causes a reversal is inadequately understood.

5.7.6 Early Mesozoic and Paleozoic reversal history

Because of the availability of the excellent marine magnetic anomaly record, it has been possible to construct a well dated history of geomagnetic polarity in the last 155–160 Ma (see Fig. 5.78). The determination of a geomagnetic polarity timescale for eras older than the Middle Jurassic is more complicated, because no comparable oceanic record exists. Paleomagnetic results show numerous reversals during the Triassic, but the Permian and Late Carboniferous were dominated by reversed polarity. A long interval of constant reversed polarity – the Kiaman interval – is a distinctive feature of the Paleozoic polarity record. Earlier in the Paleozoic reversals were common. Although magnetostratigraphic investigations of many formations are in progress, no unique record of the polarity succession is available for the Early Mesozoic or Paleozoic.

At present it is only possible to analyze Paleozoic polarity history in terms of the bias toward normal or reverse polarity. The polarity bias broadly defines superchrons (Fig. 5.85). When as many reversed as normal magnetizations are found it is assumed that the field polarity has been reversing; the interval is called a mixed polarity superchron. The Late Cretaceous and Cenozoic and the Triassic illustrate mixed polarity superchrons. Sometimes, for unknown reasons, the polarity was constant for long intervals. This was the case during the Cretaceous Quiet Interval, which in terms of polarity bias is called the Cretaceous Normal Polarity Superchron. The Kiaman reversed polarity interval is the same as the Permo-Carboniferous Reversed Polarity Superchron. The derivation of a more detailed history of geomagnetic polarity for Early Mesozoic and Paleozoic time is a massive task for paleomagnetists and biostratigraphers. Because of the absence of a marine magnetic record each polarity sequence will have to be confirmed by repetition in several magnetostratigraphic
sections before it can be accepted as globally significant and representative of dipole field behavior.

5.8 SUGGESTIONS FOR FURTHER READING

Introductory level


Intermediate level


Advanced level


5.9 REVIEW QUESTIONS

1. What is the evidence that the geomagnetic field originates inside the Earth?
2. Why is the geomagnetic field at the Earth’s surface mainly a dipole field?
3. What is the non-dipole geomagnetic field? How large is it compared to the dipole field at the Earth’s surface? Is its importance relative to the dipole field greater or less at the core–mantle boundary?
4. What is the magnetosphere? How does it originate?
5. What are the Van Allen belts? How are they formed?
6. Why do electrically charged particles from the solar wind have curved paths in the Earth’s magnetic field?
7. The geomagnetic field has a large anomaly over the South Atlantic, where its intensity is weakened by about 20%. What effects could this anomaly have on extra-terrestrial radiation that impinges on the Earth? What might be the consequences for (a) Earth-orbiting satellites, (b) astronauts, (c) passengers and crew in high-flying aircraft.
8. What is the ionosphere? What effect does it have on the magnetic field measured at the Earth’s surface?
9. What is a magnetic storm? What is its cause and what are its effects?
10. Describe the principle of operation of (a) the fluxgate magnetometer, and (b) the proton-precession magnetometer.


5.10 EXERCISES

1. Assuming that the geomagnetic field corresponds to a geocentric axial dipole, calculate the latitude of a site where the field inclination is 45°.

2. With the same assumption, calculate the inclination of the geocentric axial dipole field at latitude 45°N.

3. The magnetic moment of the Earth’s geocentric dipole is \(7.7674 \times 10^{22}\) A m². Assuming an axial dipole, calculate the total geomagnetic field intensity as a function of latitude. What is the latitudinal variation of the total field in nT km⁻¹ at 30°N?

4. Compute the inclination and declination of the magnetic field that would be observed in Boulder, Colorado (40°N, 105°W) if the Earth’s field corresponded to a perfect geocentric dipole whose axis penetrates the Earth’s surface at 80°N, 72°W.

5. Show that, for a small displacement along a meridian of magnetic longitude at magnetic latitude 45°N, the change of inclination is exactly 4/5 the change in latitude.

6. The north magnetic pole is at 77°N 102°W, the south magnetic pole is at 66°S 139°E.
   (a) Why are the poles not antipodal (exactly opposite)?
   (b) What is the closest distance between the center of the Earth and the magnetic axis joining the poles?

7. Measurements of the magnetic field elements at a geomagnetic observatory gave the following results:
   - N-component 27,000 nT;
   - E-component 40,000 nT;
   - V-component −1800 nT.
   (a) Is the observatory in the northern or southern hemisphere?
   (b) What is the total field intensity at the site?
   (c) What are the local values of inclination and declination of the field?

8. The IGRF for 2005 gives the values of the Gauss coefficients for the dipole and quadrupole components of the geomagnetic field shown in the following table.

<table>
<thead>
<tr>
<th>Gauss coefficients [nT]</th>
</tr>
</thead>
<tbody>
<tr>
<td>(g_1^0)</td>
</tr>
<tr>
<td>(g_2^0)</td>
</tr>
<tr>
<td>(g_1^1)</td>
</tr>
<tr>
<td>(g_2^1)</td>
</tr>
<tr>
<td>(h_1^1)</td>
</tr>
<tr>
<td>(h_2^1)</td>
</tr>
<tr>
<td>(g_2^2)</td>
</tr>
<tr>
<td>(h_2^2)</td>
</tr>
</tbody>
</table>

(a) Calculate the intensity of the dipole component of the field at the Earth’s surface.

(b) Calculate the corresponding intensity of the quadrupole component and express it as a percentage of the dipole field intensity.

9. The values of the Gauss coefficients in the previous exercise are given for the Earth’s surface. Recalculate the intensities of the dipole and quadrupole field components at the core–mantle boundary (Earth’s radius 6371 km, core radius 3485 km).

10. What are the values of \(n\) and \(m\) in the designation \(Y_n^m\) for the spherical harmonic functions illustrated in...
5.10 EXERCISES

Fig. B5.3.2 in Box 5.3? Sketch how these patterns would appear on the opposite side of the reference sphere to the one you are looking at?

11. In an aeromagnetic survey at a flight altitude of 2000 m above sea-level, the maximum total field anomaly over an orebody was 30 nT. In a repeated measurement at 2500 m altitude the maximum amplitude of the anomaly was 20 nT. Calculate the depth of the orebody below sea-level assuming (i) a monopole source and (ii) a dipole source for the anomaly.

12. The vertical field magnetic anomaly \( \Delta B_z \) over a vertically magnetized anticline (represented by a horizontal cylinder) is given by Eq. (5.54). Draw a sketch of the anomaly on a profile normal to the structure. Observe the horizontal positions where the anomaly is zero.
   (a) Calculate the horizontal positions where the anomaly has extreme values.
   (b) Calculate the peak-to-peak values of the anomaly.

13. Assume that the core of an anticline is made of basalt, the host formation is limestone, and the rocks are vertically magnetized with susceptibilities given by the median values in Fig. 5.13.
   (a) Compute the induced magnetization contrast when the vertical magnetic field intensity is 40,000 nT.
   (b) If the anticline is modelled by a horizontal cylinder whose radius is one fifth the depth of its axis, calculate the maximum amplitude of the vertical field anomaly over the structure.

14. Assuming the gravity anomaly over an anticline as given by Eq. (2.94), apply the Poisson relationship (Box 5.5) to obtain the horizontal field magnetic anomaly \( \Delta B_x \) of a vertically magnetized anticline with radius \( R \) and magnetization contrast \( \Delta M_z \).

15. The north \((N)\), east \((E)\) and vertical \((V)\) components of a magnetization can be calculated from its intensity \((M)\), declination \((D)\) and inclination \((I)\) using the following relationships:
   \[ N = M \cos D \cos I \]
   \[ E = M \cos D \sin I \]
   \[ V = M \sin I \]

   In the progressive thermal demagnetization of a sample of Cretaceous limestone for a paleomagnetic study the following remanent magnetizations were measured at different temperatures \(T\):
   (a) Calculate the north \((N)\), east \((E)\) and vertical \((V)\) components of the magnetizations at each demagnetization stage.
   (b) Plot the \(N\)-components against the \(E\)-components, fit a straight line, and determine the optimum declination for the stable magnetization direction.

   (c) Plot the \(V\)-components against either the \(N\)- or \(E\)-components, fit a straight line, and compute the optimum inclination for the stable magnetization direction.

   (d) The straight lines do not pass through the origin of the plot. What does this imply?

16. In the same paleomagnetic study, the following stable directions of remanent magnetization, corrected for the dip of bedding in the limestone formation, were measured in five samples from the same site:

<table>
<thead>
<tr>
<th>Sample</th>
<th>(D[\circ])</th>
<th>(I[\circ])</th>
</tr>
</thead>
<tbody>
<tr>
<td>SR-04</td>
<td>329.7</td>
<td>40.6</td>
</tr>
<tr>
<td>SR-05</td>
<td>336.6</td>
<td>24.7</td>
</tr>
<tr>
<td>SR-07</td>
<td>326.2</td>
<td>46.0</td>
</tr>
<tr>
<td>SR-10</td>
<td>321.1</td>
<td>40.9</td>
</tr>
<tr>
<td>SR-12</td>
<td>322.7</td>
<td>44.9</td>
</tr>
</tbody>
</table>

   (a) Calculate the direction cosines \((\lambda_N, \lambda_E, \lambda_V)\) of each direction, using the relationships
   \[ \lambda_N = \cos I \cos D \]
   \[ \lambda_E = \cos I \sin D \]
   \[ \lambda_V = \sin I \]

   (b) Add up the values for each direction cosine. Let the sums be \(X, Y\) and \(Z\), where
   \[ X = \sum \lambda_N \]
   \[ Y = \sum \lambda_E \]
   \[ Z = \sum \lambda_V \]

   Calculate the vector sum of the directions, \(R\), and the declination \(D_m\) and inclination \(I_m\) of the mean direction using the relationships
   \[ R = \sqrt{X^2 + Y^2 + Z^2} \quad \tan D_m = Y/X \quad \sin I_m = Z/R \]

   (c) Using the computed value of \(R\), calculate the precision parameter \(k\) of the data and the 95% confidence error \((a_{95})\) for the mean direction.

17. The samples analyzed in the previous exercise were gathered at a site in Italy with latitude 43.4°N, longitude 12.6°E. Calculate the latitude and longitude of the paleomagnetic pole position for the Italian limestone.
Assume that the location of the Late Cretaceous paleomagnetic pole for the European plate is at 72°N, 154°E and the corresponding pole for the African plate is at 67°N, 245°E.

(a) Calculate the expected ‘European’ and ‘African’ directions at the Italian site in the previous exercise.

(b) Compare the expected directions with the observed directions and explain how the paleomagnetic results from the Italian limestone should be interpreted.
Appendix A  The three-dimensional wave equations

Derivation of general equations of motion

The development of the equations for P-waves and S-waves in three dimensions begins like the one-dimensional cases. In this case, however, the displacement of a particle during the passage of a seismic wave is a vector \( \mathbf{U} \) with components \( u, v, \) and \( w \) in the \( x, y, \) and \( z \)-directions, respectively.

Consider the forces acting on the faces of the small box shown in Fig. A1. The faces normal to the \( x, y, \) and \( z \)-directions have areas \( A_x, A_y, \) and \( A_z, \) respectively. Let the force on the left face of area \( A_x, \) acting in the \( x \)-direction, be \( F_{xx} \) and that on the right face be \( (D F_{xx})_x \). If these forces are equal (\( D F_{xx} \)) they will cause no motion in the \( x \)-direction, if they are unequal, motion will result. If the box is small enough for the change in force to take place uniformly, we can write

\[
\Delta F_x = \frac{\partial (F_{xx})}{\partial x} \Delta x = \frac{\partial (\sigma_{xx} A_x)}{\partial x} \Delta x = \frac{\partial \sigma_{xx}}{\partial x} (\Delta y \Delta z) \Delta x \quad (A1)
\]

The forces acting on the faces normal to the \( z \)-direction also have \( x \)-components. Treating them in the same way, the force on the base of the box of area \( A_x \), acting in the \( x \)-direction is \( (F_{xx})_x \) and that on the top face is \( (F_{xx})_x + \Delta F_x \). The difference is

\[
\Delta (F_{xx}) = \frac{\partial (F_{xx})}{\partial z} \Delta z = \frac{\partial (\sigma_{xx} A_z)}{\partial z} \Delta z = \frac{\partial \sigma_{xx}}{\partial z} (\Delta x \Delta y) \Delta z \quad (A2)
\]

Analogously (not shown in the figure), the resulting force in the \( x \)-direction from the stresses on the faces normal to the \( y \)-direction is given by

\[
\Delta (F_{xx}) = \frac{\partial (F_{xx})}{\partial y} \Delta y = \frac{\partial (\sigma_{xx} A_y)}{\partial y} \Delta y = \frac{\partial \sigma_{xx}}{\partial y} (\Delta z \Delta x) \Delta y \quad (A3)
\]

Combining these results, we get the net force \( \Delta F_x \) acting on the box in the \( x \)-direction:

\[
\Delta F_x = \left( \frac{\partial \sigma_{xx}}{\partial x} + \frac{\partial \sigma_{yx}}{\partial y} + \frac{\partial \sigma_{zx}}{\partial z} \right) \Delta V \quad (A4)
\]

where \( \Delta V = \Delta x \Delta y \Delta z \) is the volume of the elementary box. If the density of the medium is \( \rho \) and the \( x \)-component of the displacement is \( u \), we can apply Newton’s law of motion to obtain the equation of motion

\[
\frac{\partial \sigma_{xx}}{\partial x} = \lambda \frac{\partial u}{\partial x} + 2 \mu \frac{\partial^2 u}{\partial x^2}
\]

From the definitions of the Lamé constants (Section 3.2.4.3) and the relationships between stress and strain (Eq. (3.32)), we have

\[
\sigma_{xx} = \lambda \theta + 2 \mu \varepsilon_{xx} = \lambda \left( \frac{\partial u}{\partial x} + \frac{\partial v}{\partial y} + \frac{\partial w}{\partial z} \right) + 2 \mu \frac{\partial u}{\partial x}
\]

\[
\sigma_{yx} = 2 \mu \varepsilon_{yx} = \mu \left( \frac{\partial v}{\partial x} + \frac{\partial u}{\partial y} \right)
\]

\[
\sigma_{zx} = 2 \mu \varepsilon_{zx} = \mu \left( \frac{\partial w}{\partial x} + \frac{\partial u}{\partial z} \right) \quad (A6)
\]

Differentiating each of these equations in turn, as in the equation of motion, gives
After substitution and grouping terms, the equation of motion for the $x$-displacement $u$ becomes
\[\lambda \frac{\partial^2 u}{\partial x^2} + 2\mu \frac{\partial^2 u}{\partial y \partial x} + \mu \left(\frac{\partial^2 u}{\partial y^2} + \frac{\partial^2 u}{\partial z^2}\right) = \rho \frac{\partial^2 u}{\partial t^2}\]  
(A7)

\[\lambda \frac{\partial^2 \theta}{\partial x^2} + 2\mu \frac{\partial \theta}{\partial y \partial x} + \mu \left(\frac{\partial^2 \theta}{\partial y^2} + \frac{\partial^2 \theta}{\partial z^2}\right)\]

\[= \rho \frac{\partial^2 \theta}{\partial t^2}\]  
(A8)

\[\lambda \frac{\partial^2 \phi}{\partial x^2} + \mu \frac{\partial^2 \phi}{\partial y^2} + \mu \frac{\partial^2 \phi}{\partial z^2} = \rho \frac{\partial^2 \phi}{\partial t^2}\]  
(A9)

\[\lambda \frac{\partial^2 \theta}{\partial x^2} + \mu \frac{\partial \theta}{\partial y \partial x} + \mu \left(\frac{\partial^2 \theta}{\partial y^2} + \frac{\partial^2 \theta}{\partial z^2}\right) + \mu \frac{\partial^2 \theta}{\partial z^2} = \rho \frac{\partial^2 \theta}{\partial t^2}\]  
(A10)

It is convenient to introduce the mathematical operator $\Delta^2 = \frac{\partial^2}{\partial x^2} + \frac{\partial^2}{\partial y^2} + \frac{\partial^2}{\partial z^2}$. This allows a more concise presentation of Eq. (A10) for the $x$-displacement, which becomes
\[(\lambda + \mu) \Delta^2 u = \rho \frac{\partial^2 u}{\partial t^2}\]  
(A11)

Similar analysis of the $y$- and $z$-displacements, $v$ and $w$ respectively, gives the additional equations of motion:
\[(\lambda + \mu) \Delta^2 v = \rho \frac{\partial^2 v}{\partial t^2}\]  
(A12)

\[(\lambda + \mu) \Delta^2 w = \rho \frac{\partial^2 w}{\partial t^2}\]  
(A13)

Equations (A11)–(A13) can now be resolved into two wave motions representing longitudinal (P-) waves and transverse (S-) waves.

**Equation of longitudinal waves**

Differentiating each of Eqs. (A11), (A12) and (A13) in turn by $x$, $y$, and $z$, respectively, gives
\[(\lambda + \mu) \frac{\partial^2 \theta}{\partial x^2} + \mu \Delta^2 \left(\frac{\partial u}{\partial x}\right) = \rho \frac{\partial^2 \theta}{\partial t^2}\]  
(A14)

\[\lambda \frac{\partial^2 \phi}{\partial y^2} + \mu \Delta^2 \left(\frac{\partial v}{\partial y}\right) = \rho \frac{\partial^2 \phi}{\partial t^2}\]  
(A15)

\[\lambda \frac{\partial^2 \phi}{\partial z^2} + \mu \Delta^2 \left(\frac{\partial w}{\partial z}\right) = \rho \frac{\partial^2 \phi}{\partial t^2}\]  
(A16)

Adding these equations together, and using the definitions of $\Delta^2$ given above and $\theta = \partial u/\partial x + \partial v/\partial y + \partial w/\partial z$ we obtain
\[(\lambda + \mu) \left(\frac{\partial^2 \theta}{\partial x^2} + \frac{\partial^2 \phi}{\partial y^2} + \frac{\partial^2 \phi}{\partial z^2}\right) + \mu \Delta^2 \left(\frac{\partial u}{\partial x} + \frac{\partial v}{\partial y} + \frac{\partial w}{\partial z}\right) = \rho \frac{\partial^2 \theta}{\partial t^2}\]  
(A17)

\[(\lambda + \mu) \Delta^2 \theta + \mu \Delta^2 \phi = \rho \frac{\partial^2 \theta}{\partial t^2}\]  
(A18)

Comparison with the one-dimensional compressional wave equation (Eq. (3.41)) shows that Eq. (A18) describes the three-dimensional propagation of a volumetric change (dilatation) $\theta$ with velocity $\alpha$ given by Eq. (A19).

This wave travels as a succession of dilatations and compressions parallel to the direction of propagation and is referred to as a *longitudinal wave*. Because it is the first to arrive at a seismometer it is more commonly called the *primary wave* or *P-wave*.

**Equation of transverse waves**

By differentiating Eq. (A11) with respect to $y$ we get
\[(\lambda + \mu) \frac{\partial^2 \theta}{\partial y^2} = \rho \frac{\partial^2 \theta}{\partial t^2}\]  
(A20)

Likewise, differentiating Eq. (A12) with respect to $x$ gives
\[(\lambda + \mu) \frac{\partial^2 \phi}{\partial x^2} = \rho \frac{\partial^2 \phi}{\partial t^2}\]  
(A21)

Subtracting Eq. (A20) from Eq. (A21) we get
\[\mu \Delta^2 \left(\frac{\partial v}{\partial x} - \frac{\partial u}{\partial y}\right) = \rho \frac{\partial^2)}{\partial t^2}\]  
(A22)

By combining Eq. (A12) with Eq. (A13), and Eq. (A13) with Eq. (A11), in the same way, we obtain the further wave equations
\[\mu \Delta^2 \left(\frac{\partial w}{\partial z} - \frac{\partial u}{\partial y}\right) = \rho \frac{\partial^2}{\partial t^2}\]  
(A23)

\[\mu \Delta^2 \left(\frac{\partial v}{\partial y} - \frac{\partial w}{\partial z}\right) = \rho \frac{\partial^2}{\partial t^2}\]  
(A24)
Define

\[ \Psi_x = \left( \frac{\partial w}{\partial y} - \frac{\partial v}{\partial z} \right) \quad \Psi_y = \left( \frac{\partial u}{\partial z} - \frac{\partial w}{\partial x} \right) \quad \Psi_z = \left( \frac{\partial v}{\partial x} - \frac{\partial u}{\partial y} \right) \]  

(A25)

The quantities \( \partial v/\partial z \), etc., are equivalent to small rotational distortions that combine to give shear strains (see Section 3.2.3.3). Thus \( \partial v/\partial z - \partial w/\partial y \) is a net rotation in the \( y-z \) plane, i.e., normal to the \( x \)-direction. The quantities defined in Eq. (A25) are the components of a vector \( \Psi = (\Psi_x, \Psi_y, \Psi_z) \). Each component of \( M \) describes a net rotation in the plane normal to the specific coordinate axis.

Equations. (A22)–(A24) show that the components of \( \Psi \) satisfy the equation

\[ \mu \Delta^2 \Psi = \rho \frac{\partial^2 \Psi}{\partial t^2} \]  

(A26)

which can be written as

\[ \beta^2 \Delta^2 \Psi = \frac{\partial^2 \Psi}{\partial t^2} \]  

(A27)

where

\[ \beta^2 = \frac{\mu}{\rho} \]  

(A28)

Equation (A27) has the form of the wave equation. It describes the propagation of a rotational or shear disturbance \( \Psi \) with velocity \( \beta \) given by Eq. (A28). Because the displacements in this wave are in the plane normal to the direction of propagation it is referred to as a transverse wave. Its velocity \( \beta \) is slower than the P-wave velocity \( a \), so it arrives at a detector later than the P-wave from the same event and is called the secondary wave, or \( S \)-wave. As the disturbance consists of shear motions of particles in the wavefront, it is also called a shear wave.
Appendix B  Cooling of a semi-infinite half-space

A semi-infinite half-space extending to infinity in the z-direction is initially at a uniform temperature $T_0$. Let the temperature at time $t$ and depth $z$ inside the cooling half-space be $T(z,t)$. The temperature of the upper surface at $z = 0$ is changed suddenly and maintained at 0 °C. The boundary conditions are therefore $T(0,t) = 0$ and $T(z,0) = T_0$. The temperature in the cooling half-space must satisfy the heat conduction equation, which can be written (Box 4.1)

$$\frac{1}{r^2} \frac{d}{dr} \left( r^2 \frac{dT}{dr} \right) = \frac{1}{\rho c_p} \frac{d}{dt} T \tag{B1}$$

Separating the variables and setting the separation constant equal to $-n^2$, we get

$$\frac{1}{r^2} \frac{d}{dr} \left( r^2 \frac{dT}{dr} \right) = -n^2 \tag{B2}$$

The particular solutions of these equations are

$$T = \sum_n \frac{e^{-n^2 t}}{n} B_n \sin nz \tag{B3}$$

The boundary condition on the upper surface at $z = 0$ is $T(0,t) = 0$, which requires $A_n = 0$. The general solution can then be written

$$T(z,t) = \sum_n \frac{e^{-n^2 t}}{n} B_n \sin nz \tag{B4}$$

If the temperature distribution is continuous, the summation can be replaced by an integral and the discrete values $B_n$ by a continuous function $B(n)$

$$T(z,t) = \int_0^\infty B(n) \sin nz \, dn \tag{B5}$$

If at $t=0$ the cooling half-space has an initial temperature distribution $T(z)$,

$$T(z,0) = T(z) = \int_0^\infty B(n) \sin nz \, dn \tag{B6}$$

By using the orthogonality properties of sine and cosine functions (see Box 2.3), we get

$$B(n) = \frac{2}{\pi} \int_0^\infty T(z) \sin nz \, dz = \frac{2}{\pi} \int_0^\infty T(\xi) \sin(n\xi) \, d\xi \tag{B7}$$

where the integration variable is changed to avoid confusion. Substituting in Eq. (B5) we get

$$T(z,t) = \frac{2}{\pi} \int_0^\infty T(\xi) \left[ e^{-nu^2 \sin(nz) \sin(n\xi)} \, dn \right] \tag{B8}$$

Invoking the trigonometric relationship $2\sin(nz)\sin(n\xi) = \cos(n(\xi-z)) - \cos(n(\xi+z))$ changes the integral in Eq. (B8) to

$$T(z,t) = \frac{1}{\pi} \int_0^\infty T(\xi) \left[ e^{-nu^2 \cos(n(\xi-z))} - e^{-nu^2 \cos(n(\xi+z))} \right] \, dn \tag{B9}$$

The integration can be simplified by writing $\kappa t - \alpha$, $(\xi-z) = u$, and $(\xi+z) = v$:

$$T(z,t) = \frac{1}{\pi} \int_0^\infty T(\xi) \left[ e^{-nu^2 \cos(mu)} \, dn \right] \tag{B10}$$

Before proceeding with the solution of Eq. (B10) it is first necessary to evaluate the integral

$$Y = \int_0^\infty e^{-nu^2 \cos(mu)} \, dn \tag{B11}$$

Note that

$$\frac{\partial Y}{\partial u} = \int_0^\infty (-nu e^{-nu^2}) \sin(mu) \, dn \tag{B12}$$

Integrating Eq. (B12) by parts gives

$$\frac{\partial Y}{\partial u} = \frac{\sin(mu)}{2u} \left[ \frac{e^{-nu^2} \sin(mu)}{2u} - \frac{e^{-nu^2} \cos(mu)}{2u} \right]_0^\infty - \frac{\sin(mu)}{2u} \frac{e^{-nu^2} \cos(mu)}{2u} = -\frac{nu}{2u} Y \tag{B13}$$

$$\frac{1}{\rho c_p} \frac{\partial Y}{\partial u} = \frac{1}{\rho c_p} \frac{\partial}{\partial u} \left[ \int_0^\infty e^{-nu^2 \sin(mu)} \, dn \right] = -\frac{\partial}{\partial u} \left[ \frac{n^2}{4u} \right] Y \tag{B14}$$

$$\ln Y = -\frac{n^2}{4u} + c = -\frac{n^2}{4u} + \ln(Y_0) \tag{B15}$$

$$Y = Y_0 e^{nu^2/4u} \tag{B16}$$
COOLING OF A SEMI-INFINITE HALF-SPACE

The constant $Y_0$ is the value of the integral $Y$ defined in Eq. (B11) for $u = 0$. This integration (evaluated below) has the value

$$Y_0 = \int_0^\infty e^{-u^2} du = \frac{1}{\sqrt{\pi}}$$  \hspace{1cm} (B17)

Thus $Y = \int_0^\infty e^{-u^2} \cos(\eta u) du = \frac{1}{\sqrt{\pi}} e^{-\frac{\eta^2}{4\alpha}}$  \hspace{1cm} (B18)

Inserting this solution in Eq. (B10) gives

$$T(z,t) = \frac{1}{\pi} \int_0^\infty T(\xi) \frac{\xi}{\sqrt{\pi \kappa t}} \left[ e^{-\xi^2/4\alpha} - e^{-\xi^2/4\alpha} \right] d\xi$$

$$= \frac{T_0}{\sqrt{\pi \kappa t}} \left[ \int_0^\infty e^{-(z-\xi)^2/4\alpha} d\xi - \int_0^\infty e^{-(z+\xi)^2/4\alpha} d\xi \right]$$  \hspace{1cm} (B19)

If the cooling body has initially a uniform constant temperature $T_0$, then $T(z,t) = T_0$ and the integration in Eq. (B19) can be written

$$T(z,t) = \frac{T_0}{\sqrt{\pi \kappa t}} \left[ \int_0^\infty e^{-(z-\xi)^2/4\alpha} d\xi - \int_0^\infty e^{-(z+\xi)^2/4\alpha} d\xi \right]$$  \hspace{1cm} (B20)

In the first integration we can write $w = (\xi - z)/2\sqrt{\kappa t}$; then $dw = 1/\sqrt{\kappa t} d\xi$ and the upper and lower limits of the integration change to $\infty$ and $-z/2\sqrt{\kappa t}$, respectively. Similarly, in the second integration we can write $w = (\xi + z)/2\sqrt{\kappa t}$, with corresponding changes to the integration limits, which become $\infty$ and $z/2\sqrt{\kappa t}$, respectively. Equation (B20) can be rewritten as

$$T(z,t) = \frac{T_0}{\sqrt{\pi}} \left[ \int_{-z/2\sqrt{\kappa t}}^{\infty} e^{-w^2} dw - \int_{z/2\sqrt{\kappa t}}^{\infty} e^{-w^2} dw \right]$$  \hspace{1cm} (B21)

$$T(z,t) = \frac{T_0}{\sqrt{\pi}} \left[ \int_{-\infty}^{-z/2\sqrt{\kappa t}} e^{-w^2} dw + \int_{z/2\sqrt{\kappa t}}^{\infty} e^{-w^2} dw \right] = \frac{2T_0}{\sqrt{\pi}} \int_{z/2\sqrt{\kappa t}}^{\infty} e^{-w^2} dw$$  \hspace{1cm} (B22)

$$T(z,t) = T_0 \left[ \sqrt{\frac{\pi}{\kappa t}} \int_0^{\infty} e^{-w^2} dw \right]$$  \hspace{1cm} (B23)

The expression in brackets is the error function (Box 4.3), defined as

$$\text{erf}(\eta) = \frac{2}{\sqrt{\pi}} \int_0^\eta e^{-\xi^2} d\xi$$  \hspace{1cm} (B24)

The solution for the temperature distribution in the cooling plate is therefore

$$T(z,t) = T_0 \text{erf}\left( \sqrt{\frac{z}{2\kappa t}} \right)$$  \hspace{1cm} (B25)

Evaluation of the integral $Y_0$ in Eq. (B17)

$$Y_0 = \int_0^\infty e^{-ax^2} dx = \int_0^\infty e^{-ay^2} dy$$  \hspace{1cm} (B26)

$$(Y_0)^2 = \int_0^\infty e^{-ax^2} (\int_0^\infty e^{-ay^2} dy) = \int_0^\infty e^{-a(x^2+y^2)} dy dx$$  \hspace{1cm} (B27)

This is an integration over an area enclosed by the positive $x$- and $y$-axes ($0 \leq x \leq \infty; 0 \leq y \leq \infty$). Change to polar coordinates $x = r \cos\varphi$ and $y = r \sin\varphi$ and write the element of area as $(dx dy) = (r dr d\varphi)$. The limits of the integration change to $0 \leq r \leq \infty; 0 \leq \varphi \leq (\pi/2)$:

$$(Y_0)^2 = \int_0^\infty \int_0^\infty e^{-ar^2} r dr d\varphi = \int_0^{\pi/2} \left[ \int_0^\infty e^{-ar^2} r dr \right] d\varphi$$  \hspace{1cm} (B28)

$$(Y_0)^2 = \int_0^\infty \int_0^\infty e^{-ar^2} r dr d\varphi = \int_0^{\pi/2} \left[ \int_0^\infty e^{-ar^2} r dr \right] d\varphi$$  \hspace{1cm} (B29)

The value of the integral in Eq. (B17) is thus

$$Y_0 = \frac{1}{2} \sqrt{\frac{\pi}{2\alpha}}$$  \hspace{1cm} (B30)
Bibliography


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