THE PHALABORWA (PALABORA) DEPOSIT AND ITS POTENTIAL CONNECTION TO IRON-OXIDE COPPER-GOLD DEPOSITS OF OLYMPIC DAM TYPE

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Abstract - Phalaborwa is the second largest copper mine in the world and the largest in Africa. The orebody is hosted by the Lookekop pipe within the Phalaborwa Complex, and is also mined for magnetite, apatite, vermiculite with a large by-products including gold, silver, phosphate, rare earth elements and uranium. The Phalaborwa Complex intruded Archaean basement at the edge of the Kaapvaal Craton in early Proterozoic times (2060±1 Ma) and consists of concentrically zoned, multiple intrusions which decrease in age from the margin to the core. The outer parts are predominantly clinopyroxenites, which have been variably metasomatized. Younger pegmatoidal pyroxenites intruded at three centres, including Lookekop, where fischerite and a banded carbonatite were also emplaced, followed by a transgressive carbonatite that intruded as the last magmatic phase along fracture and shear zones. Economic copper mineralisation is hosted predominantly within the transgressive carbonatite as disseminated carbonatite and veinlets of chalcopyrite, with lesser bornite and cubanite. Magnetite is a primary igneous phase in all rocks and is paragenetically earlier than the copper sulphides. The quantity and quality of magnetite is zoned and its distribution is antithetic to that of copper. Ore fluids are high temperature, highly saline, CO2-rich, magmatic-water dominated brines. The Complex and the mineralisation are interpreted to be products of the interaction of multiple pyroxenite to carbonatite magmas and their volatiles, which were ultimately derived from decompression melting of metasomatic mantle during extension at a transition from thick Archaean to thinner post-Archaean lithosphere. The orebody at Lookekop has many features including its age, giant size, pipe-like form, low ore grade, minor and major element associations and ore-fluid properties that are consistent with it being a proximal endmember of the widely recognised iron-oxide copper-gold deposit group. As such it helps explain characteristics such as the pipe-like brecciation as well as the common siteding of these deposits at craton edges or other lithospheric boundaries.

Introduction

The iron-oxide copper-gold deposit group is now recognised worldwide, with world-class or giant Proterozoic examples discovered in the Stuart Shelf and Con栾urry districts of central Australia and the Carajas Region of northeastern Brazil, and other Proterozoic to Phanerozoic examples in North and South America (e.g. Hauck, 1990; Hitzman et al., 1992). Although there is some controversy regarding their genesis, most authors favour an association between the clearly epigenetic, commonly breccia-hosted mineralisation and igneous intrusive activity, commonly anorogenic, which is widespread in mineralised regions. However, nowhere are the orebodies sited in major intrusive rock-bodies considered to be both coeval and co-genetic with the mineralisation. Attention is drawn here to the Phalaborwa (Palabora) deposit as a possible end-member of this group.

The Phalaborwa Complex, located in the Mpmulanga province in northeast Transvaal Province of the Republic of South Africa (Figure 1) at latitude 24° 00’S and longitude 31° 07’E, represents the oldest known carbonatite complex of the southern African continent and is host to one of the world’s major copper deposits. Copper is currently mined by open cut, and it is planned to continue mining to an approximate depth of 775m before going underground in 2002. By this time, an estimated 3.4Mt of copper metal will have been extracted, along with millions of tons of titaniferous magnetite and significant amounts of nickel, uranium, thorium, zirconium, baddeleyite, gold, silver, rare-earth elements and platinum-group metals, as well as sulphuric acid, sulphate-monohydrate and hexahydrate. In addition, it is estimated that the Complex contains 298Mt of merchant grade phosphate (in the form of apatite) per 100m of depth; it is also the world’s largest resource of high-grade vermiculite. Thus, it is enriched in many elements (e.g. Fe, Cu, Au, REE, P, F, U, Th) that typify the iron-oxide copper-gold deposit group.

The hosting Phalaborwa Complex is mainly composed of ultramafic rocks, largely dunite and pyroxenite, with a core
Figure 1. Geological setting of the Phalaborwa orebody. A) Generalised map of southern Africa showing position of Phalaborwa in relation to the edge of the Kaapvaal Craton (simplified from Reeve et al., 1990). B) Simplified maps of the Phalaborwa Complex with an enlargement of the Loolokop pipe (modified after Hanekom et al., 1965 and Palabora Mining Company, 1976).
of carbonatite. Minor rock types include glimmerite, foskorkite, syenite, and fenite (Verwoerd, 1993). Importantly, it is a dense, magnetic body, producing a coincident gravity and magnetic anomaly of regional magnitude, such as that which led to the exploration drilling for the Olympic Dam deposit (e.g. Reeve et al., 1990).

Regional Geology

The Palaeoproterozoic Phalaborwa Complex is the result of multi-stage (Table 1), ultramafic to peralkaline magmatic and metasomatic activity around three centres within Archaean granites, gneisses, quartzites, granulites, amphibolites and talc- and serpentine-schists, at the eastern margin of the Kaapvaal Craton. This igneous complex is a large elongate, irregularly shaped body (Figure 1), comprising clinopyroxenites and syenites and is described in detail by the L Alexandra Mining Company Ltd Mine Geological and Mineralogical Staff (L Alexandra Mining Company; 1976), Verwoerd (1986) and Eriksson (1989). The earliest recognised intrusion is a vertically extensive, pipe-like body of pyroxenite (diopside-phlogopite-apatite rock), which was accompanied by localised felsicisation of the Archaean country rocks to produce feldspathic pyroxenites. The pyroxenites now make up about 70 percent of the main Phalaborwa Complex and, as a result of flow differentiation, vary in composition from massive pyroxenite to glimmerite. Mineralogical relationships and textures indicate that clinopyroxene, phlogopite and apatite crystallised simultaneously. Local crystal settling resulted in monomineralic assemblages of each of these minerals (Eriksson, 1989). Subsequent intrusion of syenitic plugs and dykes at the periphery of the pyroxenite was associated with intense brecciation of the pyroxenite. This dynamic intrusive phase was followed by more passive metasomatic and formation of ultramafic pegmatoids at three igneous centres, north Phalaborwa, south Phalaborwa and at Lookekop. A magnepite-olivine-apatite rock, known as foskorkite, and banded carbonatite are also interpreted to have been emplaced at this time at the Lookekop pipe. Field evidence suggests that the foskorkite is extremely heterogeneous and may, at least in part, be the result of metasomatic pre-existing rocks during carbonatite emplacement. The two intrusive cycles culminated in the emplacement of the economically most important, transgressive carbonatite into the core of the Lookekop pipe along fractures and shear zones. These multiple surges of igneous activity have resulted in complex cross-cutting and gradational (probably in part metasomatic) contacts between the lithologies (Fourie and De Jager, 1986; Eriksson, 1989).

Swarms of NE-striking dolerite dykes cut all rocks within the Phalaborwa Complex and its vicinity. The dykes are interpreted to be late-Palaeoproterozoic in age and do not appear to have had any thermal effect. As described by Eriksson (1989), an early phase of dolerite magmatism, prior to intrusion of the Complex, is evident as xenoliths in the main pyroxenite, as well as truncated dolerite dykes.

**Table 1** Simplified geological history of the Phalaborwa Complex (data from L Alexandra Mining Company, 1976 and Fourie and De Jager, 1986). Note that magmatism related to cycle 1 was more extensive and voluminous than cycle 2 which was focussed along a network of fractures and shear zones formed during the first cycle.

<table>
<thead>
<tr>
<th>Cycle</th>
<th>Stage</th>
<th>Event</th>
<th>Assoc. Mineralisation</th>
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<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>Intrusion of apatite-poor pyroxenite along same conduit as dunite followed by glimmerite</td>
<td>Apatite</td>
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<td></td>
<td>2</td>
<td>Three intrusion centres including emplacement of two pipes of apatite-rich pyroxenite in south of Complex and at Lookekop, plus ring-like intrusion to the north. Local pegmatoids.</td>
<td>Main magnetite mineralisation; Cu-sulphides, mainly bornite; Apatite</td>
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<td></td>
<td>3</td>
<td>Syenite plugs controlled by fractures formed during earlier magmatism</td>
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<td></td>
<td>4</td>
<td>Crystallisation of foskorkite and banded carbonatite as last magmatic phase within central conduit at Lookekop.</td>
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<tr>
<td>2</td>
<td>1</td>
<td>Apatite-poor, phlogopite-pyroxene pegmatoid dykes intruded in north of complex</td>
<td>Apatite</td>
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<tr>
<td></td>
<td>2</td>
<td>Intrusion of apatite-phlogopite-pyroxene dykes throughout complex</td>
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<td></td>
<td>3</td>
<td>Intrusion of syenite dykes</td>
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<td>4</td>
<td>Emplacement of transgressive carbonatite at intersection of shear zones and stockwork of transgressive carbonatite veinlets along preferred trends in older rocks</td>
<td>Uranothorianite; Minor magnetite; Major Cu-mineralisation including chalcopyrite with bornite &amp; cubanite</td>
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<td></td>
<td></td>
<td>Infiltration of late fluids along fracture network (most intense in core of intrusion)</td>
<td>Valéryite; Vermiculite after phlogopite</td>
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The Phalaborwa Complex covers an area of some 16 km$^2$ and gravity data indicate that it is a pipe-like body, plunging 76-80° to the east, to a depth of at least 3km (Hanekom et al., 1965). Isotopic dating, using Rb-Sr in phlogopite, indicates an age of 212±19 Ma for the emplacement of the Phalaborwa Complex (Eriksson, 1989). However, an isochron age of 2047±11 Ma (Eriksson, 1984) from uranothorianahites and baddeleyites in foskorite and carbonatite was considered a better estimate (Palabora Mining Company, 1976; Eriksson, 1989; Verwoerd, 1993). Conventional U-Pb analyses of the baddeleyites using thermal ionisation mass spectrometry (TIMS) indicates an age of 2060±15 Ma (Heaman and Le Cheminant, 1993), which has been subsequently supported by U-Pb analyses using SHRIMP (Wingate, 1997). Palaeomagnetic data of several syenite pipes peripheral to the main Complex reveal ages ranging from 2047 Ma to 1950 Ma (Meier and Klemd, 1997).

**Local Geological Setting**

The Loolekop pipe, within the larger Phalaborwa Complex (Figure 1), hosts the majority of the economic iron-oxide copper-gold mineralisation, and is an elliptical (1.4 km x 0.8 km), vertical pipe, elongated in an E-W direction. The Loolekop pipe comprises a core of carbonatite surrounded by foskorite (a metasomatic halo?) within pegmatoidal phlogopite-apatite-bearing pyroxenite. Drilling by Palabora Mining Company indicates that the carbonatite reaches a depth of at least 1500 m (Eriksson, 1989).

The foskorite is a coarse-grained, basic/ultrabasic rock composed of variably serpentinised olivine, magnetite (up to 50 wt%), apatite (up to 25 wt%), carbonate (≈6 wt%) and phlogopite, which decreases inwards towards the contact with the banded carbonatite (Palabora Mining Company, 1976; Eriksson, 1989). Minerals form roughly vertical bands sub-parallel to the overall shape of the body, and modal variations vary considerably from 100% magnetite to 100% olivine. Locally, fluor-silicates of magnesium, including chlorandrite and clinohumite, occur. There are carbonate patches, or lenses, within the foskorite, and their size and abundance increase towards the centre of the pipe. The contacts of foskorite with the host pyroxenite and inner banded carbonatite vary from sharp to gradational and are concentrically interbanded, possibly indicating a passive emplacement (Eriksson, 1989) or metasomatic replacement.

The inner zone of carbonatite comprises two phases:

i) an outer, elliptical, concentrically zoned, medium-to coarse-grained carbonatite comprising various proportions of magnetite-bornite-carbonate (Mg calcite-dolomite) ± olivine, which is only weakly serpentinised. Olivine is fractured and partially replaced by phlogopite, monazite or clinohumite. There are traces of chondrodite and biotite. Lombard et al. (1964) recognised two generations of banded carbonatite distinguishable only in the orientation of the banding.

ii) an inner, younger, transgressive, dyke-like, carbonate (Mg calcite ± dolomite) – phlogopite – apatite – magnetite-bearing carbonatite that is located centrally in NW-(290) and NE-(070) striking fracture sets, as well as veins in banded carbonatite, foskorite and massive mica-feldspar pyroxenite. Two curvilinear, vertically extensive, mica-rich carbonatite intrusions are also a part of the eastern portion of banded carbonatite (Figure 1). Grain-size varies from coarse-grained in the central parts of the main body to very fine-grained and sugary in narrow veins (Palabora Mining Company, 1976; Eriksson, 1989). Banding is not evident within the transgressive carbonatite unlike the foskorite and the banded carbonatite, which both display crude vertical, elliptical mineralogical banding sub-parallel to the lithological contacts, due to the alignment of magnetite/phlogopite/apatite-rich zones.

Northeast-striking, steeply dipping, anastomosing dolerite dykes cut the complex and are interpreted to be Palaeoproterozoic in age (~1880±25 Ma; Palabora Mining Company, 1976). They are interpreted to be genetically unrelated to the carbonatite (Verwoerd, 1993).

**Nature and Distribution of Mineralisation**

Both the foskorite and carbonatite host economic copper in chalcopyrite, bornite, cubanite, and chalcolite, phosphorus in apatite, and iron-oxide as magnetite mineralisation. Other sulphides that have been reported from rocks in the Loolekop pipe include pyrrhotite, pentlandite, millerite, bravoite, linnaeite, violarite, covellite, tetrahedrite, sphalerite, galena, pyrite, marcasite and molybdenite. In addition to copper and iron-oxide, minor gold, silver, platinum and palladium are produced (Palabora Mining Company, 1976; Eriksson, 1989; Verwoerd, 1993).

**Copper**

**Distribution:** The transgressive carbonatite is richest in copper with a consistent grade of about 1 wt% copper, compared to an average grade of 0.5 wt% over the whole orebody (Leroy, 1992). Chalcopyrite is the major sulphide in transgressive carbonatite as disseminated grains as well as massive blebs along fractures. Locally, there are exsolution lamellae of bornite and cubanite and pyrrhotite is present in places. Like the transgressive carbonatite itself, lenses of copper mineralisation are strongly controlled by the NW- and NE-striking fracture sets. Mineralisation zones comprise sulphide veinlets, up to 1 cm wide, which are discontinuous along strike and down dip. These zones can be up to 10 m apart (Eriksson, 1989), and are commonly associated with thin coatings of late-stage vallerite which significantly reduce the recovery of Cu from the ore (Verwoerd, 1986; Eriksson, 1989).

The banded carbonatite contains less sulphides than the transgressive carbonatite, but copper recovery is good due to the low vallerite content (Verwoerd, 1986). Bornite is the dominant sulphide phase and forms disseminated grains, droplet-like inclusions within olivine, magnetite and calcite.
(Van Rensburg, 1965), as well as massive patches and lenses which are, in places, parallel to primary mineralogical bands defined by magnetite,apatite and silicates (Eriksson, 1989).

The foskorite contains sulphides, mainly bornite and minor late-stage chalcopyrite, as randomly disseminated grains that locally appear to replace earlier minerals such as interstitial calcite (Verwoerd, 1986; Eriksson, 1989). Locally, minute elongate sulphide inclusions are aligned parallel to elongation axis of apatite. Such crystallographic control of bornite-chalcopyrite intergrowths suggests exsolution in apatite within a copper- and sulphur-rich environment (Aldous, 1980). Bornite, chalcopyrite and pentlandite also occur as small inclusions, as well as in fluid inclusions in pyroxenes, and are interpreted to be products of early sulphide-rich melts (cf Aldous, 1986). In turn, olivine is also enclosed within chalcopyrite and bornite (Eriksson, 1989).

**Paragenesis:** Three generations of copper mineralisation are recognised at Loolekop (Verwoerd, 1986): i) early stage of orthonmagmatic disseminated grains of sulphides in foskorite and banded carbonatite, associated with early sulphide-bearing magmas (Aldous, 1965); ii) main stage of postmagmatic sulphide veinslets associated with the emplacement of the transgressive carbonatite; and, iii) late stage films of vallerite on mineral cleavage planes, grain boundaries, fracture surfaces and shear planes of broad, cross-cutting shear zones.

**Iron Oxide**

**Distribution:** Economically, magnetite is the most important co-product of the copper mining at the Loolekop pipe. The average magnetite content of the ore is in the order of 27 weight percent. It is invariably Ti-bearing, but, overall, contents of Cr₂O₃ and TiO₂ are low. However, the distribution of magnetite is zoned, both in quality and quantity. Essentially, the TiO₂ content in the magnetites decreases inwards from up to 4 wt% TiO₂ in the magnetite in foskorite, to <1 wt% TiO₂ in magnetite in the transgressive carbonatite. However, additional isolated ilmenite grains also occur in the carbonatite phases (Eriksson, 1989). The magnesium contents of magnetite also vary from up to 1.5wt% MgO in magnetite in foskorite to about 0.3wt% magnetite in carbonatite (Eriksson, 1989). Similarly, the abundance of magnetite decreases from up to 50wt% magnetite in foskorite to 15-30wt% in carbonatite (Eriksson, 1989). However, the grades of magnetite are very erratic within each rock type and can vary from <1 to >90% of rock over as little as a 1m interval. Magnetite forms disoriented clots and lenses with silicate phases in the transgressive carbonatite, and also as disseminated idiomorphic crystals (mm- to cm-scale) that locally contain exsolution lamellae of ilmenite. There is virtually no magnetite in the surrounding pyroxenite (Eriksson, 1989).

**Paragenesis:** Two generations of magnetite can be distinguished in foskorite. First, there is primary magnetite which forms anhedral to euhedral grains (<1mm to 1cm), with the development of ilmenite-rich rims on some of the magnetite grains (Eriksson, 1989); in places, ulvospinel and ilmenite also formed as exsolution products in magnetite.

In the banded carbonatite, primary igneous magnetite forms discrete idiomorphic grains, which are aligned parallel to the outer contact of the carbonatite, giving it the banded appearance. A later generation of magnetite, locally replacing baddeleyite, forms rims on sulphide minerals and also along cleavage planes in philogopite (Eriksson, 1989).

**Phosphate and REE**

Apatite is the only phosphate ore mineral at Phalaborwa and occurs as a primary igneous phase in all rock types. However, at Loolekop, the highest phosphate grade is found in foskorite where apatite can constitute up to 56% of the rock as disseminated grains or in vertical bands with sharp centimetre-scale variations in grade (Eriksson, 1989). There is an overall positive correlation between phosphate and philogopite distribution, and although magnetite is generally enriched in apatite-rich zones, there is no consistent relationship between apatite and magnetite distribution (Fouque and De Jager, 1986). Apatite also controls the total rare earth element (REE) concentration which has a resource in excess of >1.6Mt. Concentrates are unusually rich in europium, and to a lesser extent, cerium, neodymium and samarium. Apatite within the carbonatite contains the highest total REE, in particular LREE, concentrations (up to 1% REE oxides), which progressively decrease to pyroxene and foskorite. In contrast, the fluorine contents increase and fluorapatite predominates in the foskorite with only minor hydroxyapatite and rare chlorapatite (Verwoerd, 1986). Phosphate mineralisation extends to at least 1000m in depth but grades are very erratic. There is little to no apparent structural control, but the ore distribution reflects the overall pipe-like structure.

**Uranium**

Uranium oxide is recovered from uranothorite, a minor accessory phase in the transgressive carbonatite, and from thorulite, a mixture of hydrocarbon, uraninite and sulphides, which is present along shears and fractures in the centre of the transgressive carbonatite. At about 30ppm UO₂, the grades at Loolekop are the lowest of any uranium orebody in the world. However, tonnages are high and up to 258t of uranium oxide is produced annually (Verwoerd, 1986).

**Zirconium**

Baddeleyite is the main zirconium mineral at Loolekop where it is an accessory phase in all rock types. It is most abundant in the foskorite but is also recovered from the carbonatites (Verwoerd, 1986; Eriksson, 1989). Baddeleyite in the carbonatites has a higher uranium content compared to that in the foskorite (Eriksson, 1984), but baddeleyite in the foskorite is particularly enriched in Fe and is locally replaced by magnetite (Eriksson, 1989). Total annual production of baddeleyite is typically in the order of 15,000 to 20,000t (Verwoerd, 1986).

**Gold, Silver and Platinum Group Metals**

Over the past 20 years, the second-biggest profit at Loolekop next to copper has been from the combined output of gold, silver and platinum group elements (PGE), which
are by-products of the copper mining. These precious metals are present in the anode sludge from electrolytic refining of copper ore, and grades vary with copper content. The typical annual production of precious metals is in the order of 20t, containing Ag, Au and PGE at a ratio of approximately 50:2:1 respectively (Verwoerd, 1986).

Others

The Phalaborwa Complex supports the world’s second largest vermiculite mine. Approximately 180,000t of vermiculite concentrate are produced per annum from the mining of two pegmatoidal pyroxenite bodies, which contain an average grade of 22% vermiculite. The two pyroxenites, in the north and south of the Complex (Figure 1b), were emplaced late during the first magmatic cycle (Table 1), and vermiculite is interpreted to be the alteration product of phlogopite due to circulation of ground water. (Verwoerd, 1986). Other by-products of the copper mining at Phalaborwa include:

i) nickel, which is associated with sulphide minerals, including pentlandite, cubanite, millerite, bravoite, violarite, siegenite and talnakhite (Verwoerd, 1986);

ii) lead, selenium, tellurium, arsenic, antimony and bismuth, which are present in minute quantities together with precious metals in the electrolytic refinery sludge (Verwoerd, 1986); and iii) sulphuric acid.

Formation of Iron Oxide

There are two generations of magnetite recognised in the Loolekop orebody. As described above, both generations are intimately intergrown and form bands with primary olivine, apatite and carbonate in the foskerite-banded carbonatite and in the younger transgressive carbonatite. Moore (1984) proposed that this banding was the result of a diffusion-controlled mechanism around crystal nuclei in a non-turbulent magma. However, Eriksson et al. (1985) suggested in-situ accumulation of crystals, maybe undercooled and accompanied by gravity slumping and sheath folds was the cause. Either way, magnetite is interpreted to be the result of orthomagmatic differentiation processes in each of the magmatic cycles (Table 1). The chemical variations observed in the magnetite generations (e.g. titanium and magnesium concentrations) can be explained in terms of changes in the depth of emplacement, bulk chemistry and/or oxygen fugacity of the respective magma (Eriksson, 1989).

Relationship of Iron Oxide to Copper Mineralisation

Iron-oxides (magnetite) and copper-sulphides are concentrated in the last phases of each of the two magmatic cycles, i.e. the foskerite-banded carbonatite and in the transgressive carbonatite (Table 1). In both instances, the copper sulphides are paragenetically later than the magnetite (Eriksson, 1989) and the first cycle is characterised by relatively higher concentrations of magnetite than copper, whereas cycle two is more enriched in copper. Hence, there is an overall antithetic relationship between the distributions of magnetite and copper mineralisation (Leroy, 1992).

Although magnetite mineralisation is generally accepted to be orthomagmatic, the paragenetically later copper-sulphides have been interpreted to be either introduced by hydrothermal solutions (Park and MacDiarmuid, 1970), or the result of magmatic segregation (Aldous, 1986). However, the presence of chalcopyrite in primary fluid inclusions in the neighbouring pyroxenite (Aldous, 1986) indicates that copper was an intrinsic early phase in the magmatic cycle and is strong support for a magmatic origin of the copper mineralisation whereby magnetite and copper-sulphides where segregated into different phases in a parental magma. Redistribution of some of the sulphides during late magmatic fracturing, brecciation and recrystallisation (Eriksson, 1989) may also have taken place. In addition, the formation of vallervite along fractures and shear zones is interpreted to be the result of infiltration of late oxidising fluids after the cessation of magmatism.

P-T Conditions during Emplacement

The depth of emplacement of the Phalaborwa Complex is estimated to be at a maximum of about 12km (Buton, 1976). Eriksson (1982) estimated pressures in the order of 450MPa, consistent with the lack of extensive metasomatism around the main Complex. Geothermometry using non-sulphide assemblages yield a wide spread of temperatures between 150-800°C, which has been interpreted by Eriksson (1989) to reflect re-equilibration at lower temperatures. Geothermometrical studies based mainly on sulphide and phlogopite compositions suggest temperatures in excess of approximately 1000°C, consistent with what can be expected for the high-potassic, ultrabasic, and carbonatitic liquids at Phalaborwa (cf Eriksson, 1989).

Genesis of Ore System

Both the magnetite and copper mineralisations are products of multiple pulses of oxidised, halogen-rich pyroxenitic and carbonatitic magmas which differentiated due to liquid immiscibility early from a residual melt that was left after a syenitic fraction separated from a mantle-derived parent alkaline peridotite magma (Fourire and De Jager, 1986; Mutschler et al., 1994). It has been proposed that the parent alkaline magma was the product of decompression melting of metasomatised mantle (e.g. Mutschler et al., 1994) resulting from extension at boundaries where there is a transition from thick Archaean to thinner post-Archaean mantle lithosphere (Groes and Viehreicher, in press). Such processes would produce alkaline magmatism enriched in incompatible and compatible elements (e.g. Meen 1987; Harmer 1999). Further crustal level fractionation could account for the formation of several magmas including a generally undifferentiated silicate, carbonate and sulphide-bearing fractions which interacted with each other and their derived fluids to produce the mineralisation at Phalaborwa. The depleted initial 143Nd/144Nd ratio values of the
carbonatites support this interpretation and characterise the source to the carbonatites as enriched in light ion lithophile elements (LILE) relative to bulk earth (Eriksson, 1989). Processes such as flow differentiation, in-situ accumulation and gravitational settling are also evident by textures in the main pyroxenite intrusion and in the magnetite ore. Complex cross-cutting relationships within and between the pyroxenite and carbonatites, as well as the spatial variation in magnesium and titanium contents of the magnetite ore support multiple pulses of magma. However, the initial $^{87}Sr/^{86}Sr$ ratios of the carbonatites differ significantly from those of the pyroxenites, suggesting a complex magmatic history rather than differentiation alone to form the carbonatites (Eriksson, 1989). Intense texture-destructive fenitisation and other metasomatic effects indicate that the magmas were volatile rich. Mineralisation styles include orthomagmatic sulphides, late-magmatic-pneumatolytic sulfides in veinlets and fracture fills. Ore fluids were high temperature (~1000°C), highly saline, CO$_2$-rich and magmatic water-dominated chloride brines. Retrograde meteoric water-dominated fluids infiltrated fracture networks after magmatism, and resulted in the formation of vallereite as well as potentially causing some redistribution of sulphides and gold (e.g. Mutschler et al., 1994).

Connection to Iron-Oxide Copper-Gold Deposits

The group of structurally controlled, epigenetic, iron-oxide copper-gold deposits typically includes deposits which contain >100Mt of ore at 0.8-1.6% Cu and 0.2-0.8 g/t Au. The type example is the Olympic Dam deposit which formed at about 1590Ma in the Stuart Shelf area of South Australia (e.g. Campbell et al., 1998; Reeve et al., 1990). Other examples of this type include Ernest Henry, Australia (e.g. Pollard et al., 1998) and Carajas, Brazil (e.g. Huhn and Nascimento, 1998). Although deposits of this class have different characteristics in detail, the larger deposits have many similarities (Table 2; e.g. Hitzman et al., 1992; Hauck, 1990; Williams, 1998; Huhn and Nascimento, 1998). Some features such as the distribution of metals, alteration mineralogy, as well as $O_2$, of the ore-fluid can be zoned laterally and/or vertically (Hitzman et al., 1992; Hauck, 1990; Groves and Vielbrecher, in press).

A number of such features also characterise the oreybody at Loolekop in the Phalaborwa Complex (Table 2; Groves and Vielbrecher, in press), including:

i) the age of mineralisation (2060Ma), classified as part of a more widespread Palaeoproterozoic magmatic event in Africa (2.5-1.6Ga; Verwoerd, 1993);

ii) the tectonic setting at the edge of a stable Craton in an anorogenic setting associated with syenitic magmatism;

iii) the dimensions of the orebody (about 1km$^2$), its low ore grade and high tonnage (approximately 850Mt at 0.5% Cu; Leroy, 1992);

iv) the overall pipe-like shape and iron oxide-copper zonation, with higher copper grades in the centre of the deposit and magnetite in distal parts;

v) multiple phases of mineralisation, with copper sulphides paragenetically later than magnetite, and

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<thead>
<tr>
<th>Feature</th>
<th>Iron-Oxide Cu-Au Deposits</th>
<th>Loolekop Pipe, Phalaborwa</th>
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<tr>
<td>Age Range</td>
<td>Mostly 1900-1400Ma</td>
<td>ca 2000Ma</td>
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<tr>
<td>Tectonic Setting</td>
<td>Adjacent to craton margins and/or lithospheric boundaries</td>
<td>Close to eastern margin of Archaean Kaapvaal Craton</td>
</tr>
<tr>
<td>Mineralisation Style</td>
<td>Commonly pipe-like to ring-like, near vertical pipes of brecciated country rock</td>
<td>Near-vertical pipe-like carbonatite body with concentric zoning (ring-like form)</td>
</tr>
<tr>
<td>Dimensions of Breccias</td>
<td>1-20km$^2$</td>
<td>~16km$^2$ for whole complex</td>
</tr>
<tr>
<td></td>
<td></td>
<td>~1km$^2$ for orebody – Loolekop pipe</td>
</tr>
<tr>
<td>Host Rocks</td>
<td>Archaean gneisses and granitoid, greenstones, metamorphosed Proterozoic rocks, anorogenic granitoids</td>
<td>Pyroxenite-carbonatite complex intruding Archaean granitoids, gneisses, granulites, amphibolites and talc-serpentine schists</td>
</tr>
<tr>
<td>Alteration</td>
<td>Intense, texture-destructive alkali and iron metasomatism, haematite and/or magnetite, also iron silicates</td>
<td>Intense alkali metasomatism in country rocks spatially associated with carbonatite pipes. Abundant magnetite</td>
</tr>
<tr>
<td>Opaque Phases Major</td>
<td>Magnetite and/or haematite</td>
<td>Magnetite</td>
</tr>
<tr>
<td>Minor</td>
<td>Chalcopyrite ± bornite ± chalcocite ± pyrrhotite ± pyrite</td>
<td>Chalcopyrite - bornite ± cubanite ± chalcocite ± pyrrhotite</td>
</tr>
<tr>
<td>Element Associations</td>
<td>Cu-Au-REE ± Ag, As, Ba, Co, F, Fe, Mo, Nb, Ni, P, Th, U</td>
<td>Cu-Fe-P-REE-F-U-Th-Au-Ni-PGE±Mo</td>
</tr>
<tr>
<td>Cu/Au Ratio</td>
<td>~18,000/1 to 27,000/1</td>
<td>~170,000/1 to 200,000/1</td>
</tr>
</tbody>
</table>
carbonate is early and is, in places, replaced by bornite;
vi) the style of mineralisation, characterised by a
dominance of iron-oxide in the form of magnetite,
together with relatively low-sulphur copper minerals,
including chalcocite and bornite, and a lack of
quartz;
vii) the mineralisation changes in style from disseminated
to fracture-controlled with time;
viii) the characteristic mineral and element associations
(Table 2), in particular the high, but erratic, P and F
concentrations, and anomalous (also erratic) U, Th
and REE, especially LREE, concentrations, also
significant concentrations of Au, Ag, Pt and Pd
(typical production per year of about 15t, of which
approximately 641 kg is gold: Verwoerd 1986), as well
as a lack of base-metals, (i.e. high Cu/Cu+Zn+Pb
ratio);
ix) the nature of the ore fluids as hot, saline, oxidising,
and CO₂-and magmatic-water bearing brines.

However, there are two major differences between
Phalaborwa and the recognised group of iron-oxide copper-
gold deposits. Firstly, the mineralisation at Phalaborwa is
intimately associated with, and hosted in, an igneous
intrusion, whereas none of the typical iron-oxide copper-
gold deposits are sited in, or adjacent to, a recognisable
intrusion large enough to be the source of the fluids.
However, available fluid inclusion, isotope, mineral-
and other thermodynamic data support the
involvement of a hot, deep, CO₂-bearing, oxidising, acidic,
aqueous ore-fluid, at least for the early stages of
mineralisation. The exact nature of the fluid is controversial,
although all the characteristics of the fluid and the style of
mineralisation (i.e. pipe-like shape, repetitive nature of ore
paragenesis, distinctive association of both compatible and
non-compatible elements) are consistent with a magmatic
origin of mineralisation, in particular an alkaline source
(Meyer, 1988; Hauck, 1990; Campbell et al., 1998; Groves
and Vielreicher, in press). As such, it is proposed that
Phalaborwa represents a proximal endmember of this
deposit class (Groves and Vielreicher, in press).

The second major difference is that the Cu/Au ratio at
Loolekop is about 200,000/1, an order of magnitude greater
than at deposits included in the iron-oxide copper-gold
group. This is discussed by Groves and Vielreicher (in press)
who concluded that the low gold concentrations at Loolekop
are a consequence of the deep crustal level of formation
and the proximity to the causal intrusion.

Concluding Statement and
Implication for Exploration

The interpretation that the orebody at Loolekop within the
Phalaborwa Igneous Complex belongs to the iron-oxide
copper-gold deposit group is consistent with most of its
characteristics including age, size and shape, grade, as well
as mineral and element associations. This style of deposit
can be explained by decompression melting of
metasomatised mantle at a lithospheric boundary to produce
volatile rich alkaline melts, enriched in compatible and
incompatible elements. The volatiles generated from such
melts can be deficient in sulphur, enriched in Cu and Au,
amongst other elements and capable of causing intense
metasomatism and pipe-like phreatic brecciation, the most
common features of the iron-oxide copper-gold deposit
group. Such a tectonic setting is also compatible with that
of the majority of Proterozoic iron-oxide copper-gold
deposits. The alkaline association also fits well with the
available constraints on the genesis of this deposit type,
and can explain the coincident gravity and magnetic
anomalies at Olympic Dam, which first led Western Mining
Corporation to explore the area. The interpretation that
Phalaborwa represents a proximal example of the iron-oxide
copper-gold deposit group implies that there could be more
conventional, distal Olympic Dam type deposits elsewhere
around the margin of the Kaapvaal Craton. Vargenho may
indeed be another example of this deposit type, hence
supporting the prospectivity for such deposits at Craton
margins.

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