THE MINERALISATION AND GEOLOGY OF THE PATAWARTA DIAPIR
NORTHERN FLINDERS RANGES, SOUTH AUSTRALIA.

By
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National Grid Reference SH 549-6638 (1:100,000)
View of Patawarta Diapir looking north east towards Patawarta Gap
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The Patawarta Diapir is located in the northern Flinders Ranges, approximately 17 km north-east of Blinman. It has intruded into the late Proterozoic Wilpena Group sediments of the Adelaide Geosyncline. The elements occurring within the diapir consist of disrupted blocks of a variety of rock types, most of which can be correlated with rocks of Willouran age. The blocks are enclosed by a matrix of carbonate breccias.

Widespread Cu mineralisation is present in the diapir, mainly as epigenetic veins. Anomalous levels of Zn, Pb, As, Co and Ni are also found in rock chips and stream sediment samples.

Fluid inclusion data for quartz suggest a temperature of formation of the veins of approximately 150°C. Freezing point determinations indicate greater than 20 wt % NaCl equivalent on average.

Sulphur isotope studies suggest that sulphur in the mineralisation was essentially derived from evaporites, but a small contribution may have come from a mafic igneous source.

Origin of the mineralising brines is postulated to be through compaction of Callanna Bed sequences and leaching of base metals from Adelaidean (and possibly basement) rocks. Reduced sulphur and base metals are interpreted to have travelled together in the same brine and precipitation occurred in open fractures in the diapir.
1. INTRODUCTION

1.1 Location and Physiography

The Patawarta Diapir lies in rugged country on Moolooloo Station, 17 km north-east of Blinman, in the Northern Flinders Ranges. It is located at latitude 30° 56' 05'', longitude 138° 40' 00'' on the Copley 1:250,000 field sheet. Moolooloo Station is 15 km north of Blinman and the diapir is located 12 km north east of the homestead. The study area lies in a class B environmental zone (Figure 1). Access to the diapir is only possible using four-wheel-drive vehicles.

The diapir is well vegetated, supporting stunted growths of erenophillia and salt bush with sparse larger eucalypts in the carbonate lithologies, while native pines are abundant in the siltstone and shale lithologies. Surface water is available only in good seasons and only one water bore occurs in the area (Pendulum Well). The diapir shows a complex drainage pattern which accentuates the general, moderate exposure of the lithologies.

1.2 Aims

The aims of this project were to map and describe the geology of the Patawarta Diapir and to investigate the mineralisation of the area, using mineralogical and chemical data, fluid inclusion measurements and sulphur isotope studies. This information is used to discuss probable geologic processes which led to the formation of the mineralised veins in the diapir.

1.3 Previous Investigations

To date, most reports on the Patawarta Diapir have concentrated on the economic prospects of the region. The first report from the
FIGURE 1 Locality Map
area was undertaken by Austin (1863), who recorded Cu ore with a high Bi content from the Mount Rugged workings. The Inspector of Mines (1901) reported Cu ores with Au contents up to 53 g/tonne from the Mosley's workings (in Brown, 1908).

Dickinson (1944) described the Lady Lehmann workings and based on the Cu assay results concluded that the vein was uneconomic. Admin Exploration drilled 13 percussion holes along the line of the Lady Lehmann workings (Shackleton, 1968).

Read (1971) carried out a stream sediment survey for base metals on behalf of North Flinders Mines. He recorded no unusual anomalies except for Cu which was associated with known Cu occurrences.

Golding (1983) reinvestigated the major mineral occurrences of the diapir for E.Z. Exploration and currently E.Z. are still undertaking work in the area.
2. GEOLOGICAL SETTING

2.1 General

The term 'diapir' is an Anglo-derivation of the Greek word 'diapirein' which means "to pierce". O'Brien (1968) has defined diapirism as "a process in which earth materials from deeper levels have pierced, or appear to have pierced shallower materials". The term is applied to physical emplacement of mainly sedimentary material at low temperatures.

The Patawarta Diapir was introduced along the Narina Fault Zone (Coates 1973) where the fault trends west-north-west, producing an elongate structure. Callen (1970) regarded the Narina Fault as a vertical strike slip fault and suggested that the Patawarta Diapir intruded and filled spaces caused by opening up along the western end of the fault where it splits into a "horsetail zone consisting of several short curved faults".

The diapir is elongated in a west-north-west, east-south-east orientation parallel to the strike of the host Adelaidean sediments which dip generally north-eastwards.

2.2 Host Rock Lithology

The Patawarta Diapir has been intruded into host sedimentary rocks comprising shales, siltstones, limestones and quartzites of the late Proterozoic Wilpena Group (Figure 2). The oldest rocks intruded by the diapir are the red and purple shales of the Bunyeroo Formation. These are overlain by the green-grey calcareous shales and grey sandy limestones of the Wonoka Formation. The Wonoka Formation is overlain by the Pound Quartzite, a white cross bedded and ripple marked feldspathic quartzite which forms a prominent ridge running in a northwest direction.
The ABC Range Quartzite, a white ripple marked sandy quartzite, together with the purple flaggy siltstones of the Brachina Formation, both stratigraphically underlie the Bunyeroo Formation, all of which have been folded around so that they lie in contact with the northwest portion of the diapir.

The regional dip of the host rocks, produced by the Delamerian Orogeny is approximately 25° to the northeast.

2.3 Geology of the Diapir

The elements occurring within the core of the diapir consist of disrupted blocks of a great variety of rock types, including calcareous siltstones and shales, limestones, sandstones and igneous rocks, enclosed by a matrix of brecciated sediment (Figure 2). These blocks or "rafts" show great variation in size with the largest having a length of over 2 km and many are overturned. Detailed thin section descriptions of the diapiric lithologies are given in Appendix 1.

2.3.1. Sedimentary Lithologies

The major sedimentary lithologies recognised in the diapir are as follows:

(i) Heavy mineral laminated sandstone and quartzite.

This unit is characterised by cross bedding and ripple marks. The sandstones are fine to coarse grained and contain quartz, feldspar, coarse detrital mica and abundant hematite. Bedding ranges from 1 cm thick up to 1 m thick. The coarsest beds are best interpreted as the result of storm activity and were laid down very quickly as indicated by flat laminations and minor graded bedding in the poorly sorted sands. Finer sand beds show small scale cross bedding, ripple marks and load casts. Cross bedding indicates that several of these rafts are overturned. Halite casts, ranging from 2 to 10 mm, are present in thinner beds of siltstone occurring with the sandstones.
(ii) Calcareous and dolomitic siltstone and shale.

This is the most common lithology in the diapir and comprises grey-green calcareous siltstone and shale which are dolomitic in part. These sediments are characterised by abundant halite casts showing hopper face morphology and cubic form. Gypsum pseudomorphs are also recognised but are not nearly as common as halite casts. Other sedimentary features are heavy mineral laminations, minor slump folds, casts of mud cracks and load casts. Disseminated fine grained hematite is very common and its distribution and variation in grain size suggest that it is of detrital, sedimentary origin.

(iii) Interbedded sandstone, siltstone and sandstone.

Micaeous and feldspathic, fine to coarse grained sandstones are interbedded on a centimetre scale with calcareous siltstone and shale. Halite casts and heavy mineral laminations are common.

(iv) Black limestone.

Thinly bedded black limestones occur in association with calcareous siltstone and shale. This unit is intensely folded in part and may indicate deposition in an environment with episodic tectonic instability since the surrounding siltstones are undeformed.

(v) Dolomite.

Dolomite rafts occurring in the diapir include both bedded and massive types and are generally weakly brecciated. Many small dolomite rafts are incorporated in the carbonate breccia and are too small to be mapped. They contain halite casts and gypsum pseudomorphs (Plate 2).
FIGURE 3  INTERPRETIVE CROSS SECTION THROUGH PATAWARTA DIAPIR
(vi) Rim Dolomite.

An unusual formation in the study area is the Rim Dolomite so named by Read (1971). This lithology outcrops along the southwest rim of the diapir and is a prominently outcropping carbonate rock. It contains numerous subrounded to subangular clasts of diapiric detritus and has a sedimentary bedded aspect. Red and white chert fragments are common and have been attributed to the breaking up of silica bands. (Read, 1971). Read has equated this dolomite with dolomites occurring on the margins of the Blinman Diapir and which Coates (1964) attributed to "metamorphic differentiation of dolomitic material from carbonate breccia" with the bedding being interpreted as a flow structure developed during emplacement of the diapir.

This study suggests that the Rim Dolomite forms part of the Bunyeroo Formation. It cannot be a raft in the diapir since the red shales of the Bunyeroo Formation are conformable with the unit and no carbonate breccia is observed between the shales and the dolomite. The proposed mode of formation of the Rim Dolomite is shown in Figure 4. At the time of deposition of the dolomite bed, which is interpreted to be a local facies of the Bunyeroo Formation, intrusion of the diapir commenced and as a sedimentary high, shed off diapiric rock fragments which were later incorporated in the dolomite. The clasts can be clearly recognised as belonging to lithologies contained within the diapir. The amount of rounding of the clasts reflects their degree of reworking before incorporation in the dolomite. Several minor phases of diapiric uplift may have occurred during deposition of the dolomite as indicated by irregular beds where clasts are abundant alternating with beds where clasts are rare. (Plate 1).

Diapirism is then inferred to have ceased until deposition of the Wonoka Formation. At this time uplift of the diapir recommenced and diapiric detritus was incorporated in the grey limestones of the Wonoka Formation along the northwest rim of the diapir (See Figure 2). This later movement caused an upturning and faulting of the Rim Dolomite and the red shales of the Bunyeroo Formation.
FIGURE 4  INFERRED STAGES OF FORMATION OF RIM DOLOMITE

- Diapir
- Rim Dolomite
- Bunyeroo Formation
- Diapiric Clasts
- Wonoka Formation
- Pound Quartzite

(not to scale)
Subsequent overprinting by the Delamerian Orogeny produced the regional dip of 25° to the north-east in the host rocks and the formation of an apparent syncline about the axis of upturning adjacent to the diapir (Figure 2).

In order to produce the almost vertical dip in the Rim Dolomite seen at the present day, the unit must have been overturned by about 25°, during the second major phase of diapiric movement. The Delamerian Orogeny would have had the effect of tilting the dolomite bed and adjacent shales to the dip seen today. Since intrusion of the diapir would produce steepening of the dolomite to vertical with only minor overturning, (N. Lemmon, pers. comm.), an element of thrusting during diapirism is inferred to produce the overturning required for the model. (Figure 4).

All sedimentary lithologies contained within the diapir show features of a shallow water to subaerial depositional environment. Most of these units can be correlated with units of the Lower Callanna Beds of the Willouran Series occurring elsewhere, in the Flinders Ranges, in stratigraphic sequence (Table 1).

Rowlands et al. (1980) have interpreted similar Callanna Bed sequences in the Willouran Ranges as forming in sabkha and playa environments.

2.3.2. Igneous Rocks.

In addition to the sedimentary rocks, mafic intrusive and extrusive rocks have been recognised.

The extrusive rocks are represented by fine grained dark green-grey amygdaloidal and non-amygdaloidal altered basalts. Two amygdaloidal basalts were noted and had vesicles predominantly filled with calcite although some amygdales included hematite and siderite. Thin sections show the basalts to consist of laths of plagioclase in a matrix of calcite, epidote, albite, chlorite, actinolite and abundant iron oxide. Coates (1964) noted similar rocks in the Blinman Diapir and suggested that these rocks were
not emplaced after diapiric intrusion but were fragments of volcanic rock equivalent to the Wooltana Volcanics of Willouran age. Whole rock analyses confirm the similarity of the extrusive rocks at Patawarta and those of Blinman and Wooltana (Table 2).

The intrusive rocks are represented by fine to medium grained dark green dolerites, almost identical in appearance to the non-amygdaloidal basalts. The dolerites are also altered with original laths of labradorite being decalcified and producing albite. Epidote and calcite are evident and Spry (1952), in a study of igneous rocks in the Worumba region, attributes this to the process of saussuritisation. Chlorite and minor amphibole (fibrous actinolite and riebeckite) are minor assemblages.

In two localities dolerites have caused local contact metamorphism of the surrounding shale and siltstone to hornfels but dolerites enclosed by carbonate breccia show a general absence of contact metamorphic effects. This may imply that the majority of dolerites in the diapir were dyke feeders for the basalts of Willouran age and now form diapiric xenoclasts. One locality in the south-west portion of the diapir suggests an intrusive relationship into Willouran sediments which were themselves subsequently disrupted. Whole rock analyses show the similarity of the intrusive and extrusive rocks (Table 2).

A contact metamorphic zone of chloritic marble would be expected if a dolerite was intruded post-diapirically into carbonate breccia. No evidence of this type of contact metamorphism has been noted in the Patawarta Diapir although reports of post-diapiric dolerites have been reported from other diapirs, (Preiss, 1980) implying a post Willouran phase of igneous activity in the Adelaide Geosyncline. On the other hand, none of the Patawarta dolerites show evidence of brecciation or dolerite fragments in the surrounding carbonate breccia, as would be expected if they were of Willouran age and subsequently intruded with the diapir.

It should be noted that the dolerites only occur within the diapiric core complex and not in the surrounding host rocks which may imply that the dolerites do not post date diapirism.
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<th>TABLE 2. GEOCHEMICAL ANALYSES OF BASALTS &amp; DOLERITES</th>
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<td>II Dolerite, Patawarta Diapir</td>
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<td>III Basalt, Wooltana: Mawson (1926)</td>
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<td>IV Basalt, Blinman Diapir; Coates (1964)</td>
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<td>V Dolerite, Blinman Diapir; Coates (1964)</td>
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| TOTAL | 100.25 | 99.88 | 99.68 | 99.79 | 100.43 |
2.3.3. **Diapiric Breccia.**

The diapiric breccia is distinguished by its high carbonate content which consists essentially of dolomite with subordinate calcite. Approximately 60% of the breccia is made up of a wide variety of rock types varying in size from very small clasts (1 cm) up to large blocks of several metres. The rock fragments, comprising partly recrystallised siltstone, shale, sandstone, dolomite and quartzite, are cemented by a carbonate matrix of dolomite, calcite and minor quartz. Accessory constituents are specular hematite, talc, rutile, pyrite, limonite, microcline and siderite.

The reason for the high carbonate content in the breccia matrix is uncertain as it seems unlikely that the fractured and crushed calcareous beds from low down in the Adelaideon System would be a significant source for all the cementing carbonate.
(a) Rim Dolomite, south west portion of diapir. Note bedded aspect

(b) Rim Dolomite, showing subrounded to subangular clasts of diapiric detritus, defining bedding

(c) Purple brecciated siltstone containing portions of brecciated basalt. Some original bedding can still be recognised
(a) Basalt block sitting on top of a sedimentary breccia containing basalt clasts.

(b) Typical carbonate breccia.

(c) Halite casts and gypsum pseudomorphs in a dolomite raft.
(a) Cross bedding in hematite-bearing siltstone. Much of the bedding is marked by concentrations of hematite (Transmitted light) (30x)

(b) Carbonate breccia showing interlocking dolomite grain and limonite staining (dark area) (Transmitted light) (40x)

(c) Altered amygdaloidal basalt

(d) Contact rock against a dolerite plug. The rock has recrystallised and the former sedimentary bedding is now shown only by concentrations of recrystallised hematite.
3. **MINERALISATION**

3.1 **General**

The widespread Cu mineralisation in the Patawarta Diapir appears to be patchy and is irregularly exposed (Figure 2). The mineralisation is of epigenetic type, characterised by narrow cross cutting veins of calcite, siderite and rarely hematite, containing pyrite, chalcopyrite and malachite. The veins occur in calcareous shales and siltstones which are often weakly brecciated. Cu is principally present as malachite and disseminated chalcopyrite in the veins. Cu is the major anomalous metal found in the diapir but anomalous levels of Zn, Pb, As, Co and Ni in various parts of the diapir are indicated by stream sediment sampling carried out by E.Z. Exploration.

In a few instances, minor sulphides occur with malachite in a stratiform fashion but it is doubtful that this was of syngenetetic origin. Binks (1968) suggested that this was due to permeation of mineralising solutions along bedding planes with subsequent selective replacement of certain laminae.

The coarsely crystalline calcite and siderite veins show an orientation of approximately north-south with a variation of 45° either side.

Vein-type mineral occurrences are also present in the Rim Dolomite in the form of coarsely crystalline barite veins and thin 1 - 2 cm calcite veins containing minor sulphides and malachite.

3.2 **Description of Major Mine Workings**

There are 5 main groups of workings in the Patawarta Diapir plus various isolated pits, all of which show anomalous Cu values (Figure 2). These workings have been previously described by Read (1971) and Golding (1983).
3.2.1 Lady Lehmann Workings

In addition to Read (1971) and Golding (1983), this group of workings has also been described by Dickinson (1944) and Shackleton (1968). The lode comprises a massive calcite vein up to 1.5m wide, bearing grains of disseminated chalcopyrite and pyrite. The vein occurs at the contact between a raft of calcareous and dolomitic siltstone and diapiric breccia (Plate 4). Workings along the northern boundary of the siltstone raft would seem to indicate that the lode continues for about 700m but Dickinson reported that the lode was not continuous. Dickinson concluded that "the lode contained 1.7% Cu over an average width of 3' 6" with a tonnage yield of 100 - 150 tonnes per vertical foot."

Adit F shows excellent exposure of the calcite vein and the host rocks (Figure 5). Along this adit the calcite lode strikes 40° with almost vertical dip and is not conformable as the host siltstones strike 80°. The hanging wall siltstones are well defined and have a sharp contact with the lode. The vein gangue is made up of a mixture of both coarse grained and finer grained fibrous calcite with minor quartz and very occasional anhedral grains of albite.

The footwall rock is a diapiric carbonate breccia containing rock fragments of several types. Midway down the adit the breccia contains large clasts of pyritic black shales in which both fine grained and coarse grained pyrite are present. Pyrite is disseminated through both the breccias and the siltstones.

The calcite vein does not extend more than 1.5m from the hanging wall and shows variable thickness along the adit. Where the calcite pinches out the breccias are in direct contact with the siltstones. Slickensides pitching south-west at 50° occur in the hanging wall and Dickinson suggested that the vein thickness was controlled by "the varying degree of opening movements which took place along the hanging wall fissure". Golding suggested the form of the lode indicated "that hydrothermal solutions dammed up against the siltstones after moving through the footwall breccia."

FIGURE 5 LADY LEHMANN MINE

ADIT F

LEGEND
- Mineralized calcite lode
- Mineralized calcite lode as fragments in breccia
- Carbonate breccia usually with disseminated mineralization
- Calcareous siltstone
- Pyritic black shales

SCALE
0 1 2 3 4 5 6 7 8 9
metres

This wall has very clean, planar contact with siltstone and lode
LODE IS 2m THICK

Fragments of mineralized calcite lode in breccia

ADIT ENTRANCE

D.J. HALL 1984

(Adapted from W.G. Shackleton (1968))
The first 10m of the lode in the adit indicate that it was brecciated subsequent to emplacement as there is no continuity to the vein in this locality, although fragments of mineralised calcite lode occur in the footwall breccia.

Malachite staining is particularly predominant in the hanging wall siltstones and extends across the lode into the footwall breccia. The sulphide minerals have penetrated beyond the limits of the calcite vein into the breccia and although their concentration diminishes rapidly with increasing distance from the hanging wall, assay results indicate that the disseminated mineralisation in the breccia is comparable to that of the calcite lode in places.

Thin sections show the hanging wall siltstones to consist predominantly of quartz, orthoclase, sericite and calcite with minor quantities of hematite and chlorite and muscovite (Appendix 1). Alteration of the siltstones was apparently due to the highly saline brines which formed the lode and caused alteration of orthoclase to sericite (Appendix 1).

Other workings extend to the east along the northern edge of the siltstone raft and show similar mineralisation although the calcite lode is nowhere as well exposed as in adit F.

3.2.2. Lady Lennon Workings

These workings are sited on ferruginous calcite and siderite veins carrying minor amounts of disseminated chalcopyrite and pyrite, occurring in fractures in a calcareous siltstone raft over a distance of 600m. Malachite is the predominant Cu mineralisation and has diffused into the host siltstones. The main working consists of a small open cut with two short adits at different levels on the side of the hill. Thin veins of siderite, containing disseminated sulphides and malachite cut across the host siltstones. Gold has been reported from this locality and one sample of vein material yielded an average gold value of 28 ppm (Golding 1983).

Other workings in the group occur to the east of the main working and are sited on massive calcite veins up to 1m across. In all of these, malachite is contained along joint planes and bedding plane fractures in the adjacent siltstones.
3.2.3 Mosleys Workings

The Mosleys group of workings consists of a number of narrow costeans and pits in host rocks of grey-green calcareous siltstones and shales. The costeans follow Cu rich veins with specular hematite, ferruginous calcite and siderite varying up to a half metre in width, which cut through the host rocks in a general north-south orientation. Fractures in the siltstones are stained by malachite. These workings are characterised by the abundance of specular hematite. Brown (1908) reported up to 53 ppm Au from vein material taken from the 10 m deep main shaft but more recent assays have shown much lower gold values in the order of 5 ppm.

3.2.4 Home Rule Flat Workings

The Home Rule Flat Workings occupy an area with very little outcrop but, as with the other workings, the pits are sited on coarse grained calcite veins containing malachite and minor sulphides. The host rocks are calcareous shales and siltstones of a lithology similar to those at the Lady Lehmann and Lady Lennon workings. Gold has been reported from these diggings and in one polished section (831-126) one coarse grain of electrum (argentiferous gold) has been found (Plate 6).

3.2.5 Mount Rugged Workings

The two pits assumed to be the Mount Rugged workings are sited in the contact zone of a small dolerite plug. Hematite, pyrite, chalcopyrite and malachite are contained in a small calcite vein in contact metamorphosed breccia. Another pit in the vicinity is located in manganiferous rocks which, as well as showing anomalous Cu values, yielded very high Ni values (1.8%).

3.3 Mineralisation in the Igneous Rocks

The igneous rocks of the Patawarta Diapir do not show prominent Cu mineralisation, unlike those of other diapirs in the Flinders Ranges (eg. Blinman Diapir).
Only one dolerite occurrence, near the Mount Rugged Mine, showed characteristic malachite staining. Since this dolerite is sited very close to a sulphide bearing calcite vein the copper is likely to be derived from this vein rather than the dolerite. Analysis of four dolerite samples show extremely low values of Cu, Pb and Zn. (<1 ppm). Ni contents in the dolerites range up to 150 ppm which are not unexpected for a mafic rock (Rose et al., 1979).

Basalts show only slightly higher values of Cu (up to 5 ppm) with Pb and Zn still being very low (<1 ppm). Malachite staining was noted in only two non amygdaloidal basalts in the study area. Ni contents range up to 50 ppm in amygdaloidal basalts and up to 70 ppm in nonamygdaloidal basalts.

3.4 Barite Veins

Three barite veins are present in the Rim Dolomite along the south-western margin of the diapir. These range in thickness from 0.5 up to 2 m in width. These veins have infilled small faults, presumably generated during formation of the diapir. The veins consist of coarsely crystalline barite with minor quartz and opaques and may have formed from the precipitation of barium sulphate from barium-saturated brine. No barite veins were noted within the core of the diapir suggesting that the formation of the barite veins and the sulphide bearing calcite veins were not related and presumably occurred at different times.

3.5 Mineral Descriptions and Textural Relationships

Examination of polished thin sections of the vein material shows the major sulphides are pyrite and chalcopyrite with grains ranging up to 5 mm in diameter. These are unevenly distributed throughout the gangue material and generally comprise less than 5% of the vein. Chalcopyrite typically comprises 60% of the total sulphides present. Small anhedral to subhedral grains of sphalerite are found in association with the major sulphides and most of the sphalerite is contained as inclusions within pyrite and chalcopyrite. This explains the zinc anomalies indicated through stream sediment sampling, by E2. Exploration. Other inclusions found within the major sulphides are calcite and very
occasional euhedral grains of marcasite. Chalcopyrite contains pyrite grains of varying sizes (Plate 5) and in some cases chalcopyrite appears to partially replace pyrite.

Secondary alteration effects are very noticeable in all samples. Subsequent to the emplacement of the sulphide bearing veins, weathering has resulted in the oxidation and leaching of the sulphide minerals and redistribution of much of the copper has occurred. Many of the sulphide grains have been replaced by limonite which is common in all samples. Limonite, when it has replaced the sulphides, shows a botryoidal texture which implies a rhythmic replacement of the sulphides (Plate 6). Electron microprobe studies of limonite show up to 1% Cu dispersed through it and so much of the copper present may have been deposited from oxidising solutions of supergene origin.

Covellite and chalcocite, the products of supergene enrichment of copper ore, occur in association with chalcopyrite and to a minor extent with pyrite (Plate 5). Covellite coatings occur around the chalcopyrite grains and are more widespread than chalcocite. Both of these secondary sulphides occur as a compact mass of uniform fine polygonal grains. In one case covellite occurs in a cleavage crack in a larger sphalerite grain. Much of the copper is now present as malachite with very minor azurite; these oxidation products are to be expected in such a calcareous environment.

No arsenic minerals (eg. arsenopyrite) were noted in any of the thin sections and consequently the observed As anomaly in the area cannot be explained.

3.6 Mineral Chemistry of Selected Sulphides

Using the electron microprobe, samples of pyrite and chalcopyrite have been analysed for Co, Ni, Ti and As. Results are shown in Appendix 3. For pyrite, Co values are quite high, ranging from just over 1% down to 0.9%. Ni and Ti values are considerably lower, averaging about 30 ppm. As values range up to 200 ppm. This value may contribute, in part, to the observed As anomaly but does not account for As values of up to 3000 ppm found in selected rock samples (Golding, 1983).
Results for chalcopyrite are approximately the same as above, for Ti and As. Ni levels fall below the detection limit of the electron microprobe and Co levels average about 200 ppm, both considerably lower than for pyrite.

For both sulphides Co/Ni is greater than 1 which is indicative of a hydrothermal source for pyrite and chalcopyrite (Mookherjee & Philip, 1979).
(a) Adit F, Lady Lehmann Workings
    showing contact of the calcite vein
    with calcareous siltstones and breccia

(b) Calcite vein, Adit F, containing sulphides,
    malachite and limonite staining

(c) Lady Lennon Workings
    A typical outcrop of disseminated
    mineralisation. Host rocks are
    calcareous shales
(a) Photomicrograph. Reflected light (20x)
Chalcopyrite enclosing inclusion of pyrite.
Rim of chalcopyrite shows alteration to covellite
and chalcocite (top right hand corner).

(b) Photomicrograph. (Reflected Light) (20x)
Chalcopyrite showing alteration to covellite (light blue)
and chalcocite (dull brown).
Calcite gangue (dark brown)
(a) Photomicrograph
Limonite replacement of sulphide showing botryoidal texture
(Transmitted light), (20x)

[b] Photomicrograph of grain of Electrum (gold) enclosed in gangue groundmass
(Reflected light), (25x)
4. **FLUID INCLUSION STUDY**

4.1 **Introduction**

The use of fluid inclusions to determine the formation temperature of minerals has long been acknowledged and provides one of the most reliable geothermometers available. Basically, a primary inclusion is formed when an imperfection in the crystal face develops during growth and traps a portion of the ore forming fluid, as the crystal grows around this flaw. On cooling the liquid contracts, leaving a near vacuum vapour phase in the inclusion. Heating the inclusion until phase homogeneity is accomplished provides a minimum temperature of formation. A pressure correction is then added to the homogenisation temperature to obtain the trapping temperature. Fluid inclusions may also be used to obtain information on the composition of ore forming fluids. A commonly used technique is to determine the freezing point of the fluid in the inclusions and from this to calculate the salinity of the fluid.

All fluid inclusions examined were found in quartz, as the calcite was not translucent enough to permit suitable observation of the inclusions. Samples were taken from the Lady Lehmann and Home Rule Flat workings. Inclusions varied between 10 and 20 um in diameter and generally were irregular in shape. Only inclusions which appeared to be primary and are assumed to form at the same time as the crystal host were investigated.

All inclusions examined were $H_2O$ - brine dominated with relatively low vapour/liquid ratios. The inclusions can be separated into 2 different types (i) Type I which contain $H_2O$ vapour and liquid $H_2O$ and at least one daughter mineral and (ii) Type II which contain $H_2O$ vapour and liquid $H_2O$. Type I inclusions comprise about 80% of the inclusions examined. The most common daughter mineral is cubic shaped, isotropic, shows low birefringence and occupies between 10 and 20% of the inclusion. This is thought to be halite, although sylvite is another alternative. Other minor daughter minerals show high birefringence and relief and a rhombohedral form.
4.2 Freezing Measurements

Roedder (1962) defined the freezing temperature (final melting temperature) "as that temperature at which the last ice crystal melts in the inclusion, under reversible equilibrium conditions" (i.e. increasing temperature). For H₂O-brine systems the freezing temperature is depressed below the freezing temperature for pure H₂O (-0.015°C) and determination of this temperature provides an estimate of the salinity and composition of the fluid within the inclusion. Depending on the temperature recorded, different systems are used to derive the salinity. The H₂O-NaCl system (Figure 6) is generally used if the freezing temperature is between -0.015°C and -21.1°C (the H₂O-NaCl eutectic) and salinity is then expressed in terms of weight % NaCl equivalent. For temperatures less than -21.1°C the CaCl₂-H₂O system (Figure 7) may be used and the salinities expressed in terms of weight % CaCl₂ equivalent.

The final melting temperatures from which the salinities were obtained are shown graphically in Figure 8 and listed in detail in Appendix 4.

The Type I inclusions would seem to fall into two groups of final melting temperatures. The lower temperature group range between -32.4°C and -22.6°C, which correspond to salinities of 22 to 26 wt % CaCl₂ equivalent. The other group have temperatures ranging from -15.4°C to 2.4°C. Since both groups are from the same inclusion type and sample, it is possible that the higher temperatures are due to melting of a NaCl hydrate phase (e.g. NaCl.2H₂O).

Using the H₂O-NaCl system the higher temperature group correspond to salinities of 24 to 25 wt % NaCl equivalent. However, Sourirajan and Kennedy (1962) state that if daughter minerals are present, then the salinity should be greater than 30 wt % NaCl equivalent.
FIGURE 6  The NaCl - H₂O System
(after Roedder, 1962)

FIGURE 7  The CaCl₂ - H₂O System
(after Ypma, 1979)
FIGURE 6

FIGURE 7
Type II inclusions cover a range of values similar to Type I inclusions although they cannot be split into 2 groups (Figure 8). One inclusion shows a positive melting temperature of +1.8°C. This may indicate that the melting phase was NaCl hydrate, existing metastably above its melting point (+0.1°C). This corresponds to a salinity of approximately 26 wt % NaCl equivalent.

The first melting temperatures of Type I and II inclusions show a range of -45.2°C to -19.2°C, and with the exception of three inclusions, all lie below the H₂O-NaCl eutectic (-21.1°C). This implies the presence of CaCl₂ and/or MgCl₂ (Ypma, 1979). However, electron microprobe studies of some of the inclusions indicate that Na, Cl and K are present but Mg is not, using the inclusion analysis method outlined by Bone et. al. 1984. This suggests that the fluid compositions could be accommodated in the CaCl₂ + NaCl + H₂O system (Figure 9).

4.3 Temperature of Homogenisation

For this study, the homogenisation runs were duplicated on each inclusion. About 10% of the inclusions failed to regain their bubbles immediately on cooling. Because of the general small size and optically poor quality of the inclusions, the temperatures recorded have an uncertainty of ± 2.0°C, due to visual limitations of the observer.

The homogenisation data are reproduced in Figure 10 in histogram form. Temperatures ranged from 142°C to 176°C with the average temperature being approximately 155°C. On heating of the inclusions, over 80% of daughter minerals remain undissolved, well past the homogenisation temperature. This indicates a low thermal coefficient of stability which is indicative of halite. The pressure correction is considered to be minimal (5°C) as the veins are thought to have formed in open fractures and hydrostatic pressure would dominate (Jouett, 1975).
Figure 8: Final Melt Temperatures

- Type I Inclusion
- Type II Inclusion

Figure 10: Homogenisation Temperatures

- Type I Inclusion
- Type II Inclusion
FIGURE 9  CaCl$_2$-NaCl-H$_2$O System

(after Ypma, 1979)
(a) Schematic diagram of inclusions examined

(b) Both Type I and Type II inclusions are evident

(c) Typical inclusions contained within quartz samples

(d) Type I inclusion

(e) Regularly shaped Type II inclusion with unusually large daughter mineral and vapour bubble.
TYPE I INCLUSION

(a) 15 μm

V$_{H_2O}$

L$_{H_2O}$

daughter mineral

TYPE II INCLUSION

(b) 10 μm

V$_{H_2O}$

L$_{H_2O}$

(c) 50 μm

V$_{H_3O}$

L$_{H_3O}$

daughter mineral

(d) 10 μm

V$_{H_3O}$

L$_{H_3O}$

(e) daughter minerals
5. SULPHUR ISOTOPE STUDIES

5.1 Introduction

Sulphur in hydrothermal ore deposits has originated ultimately from (i) an igneous source or (ii) a sea water source. Sulphur from an igneous source includes sulphur carried in magmatic fluids and sulphur obtained by leaching of sulphur-bearing minerals in igneous rocks. Sulphur in sea water occurs as aqueous sulphates and may be incorporated into sulphide deposits through various paths. Near the earth's surface in-situ reduction of sulphate to $\text{H}_2\text{S}$ by sulphur reducing bacteria may generate sedimentary sulphides. These may be leached by hydrothermal fluids or they may be replaced by other sulphide minerals when metal bearing brines are encountered. Reduction of aqueous sulphates to aqueous sulphides can take place non-bacterially at elevated temperatures. Therefore sulphate-bearing sea water, sulphate-rich connate or meteoric water bearing sulphate dissolved from evaporites can evolve into sulphide-bearing hydrothermal fluids. Fractionation of sulphur isotopes may occur in a variety of situations in the history of an ore fluid and so the observed $\delta^{34}\text{S}$ values of hydrothermal minerals may reflect a complex geochemical history of the sulphur in hydrothermal fluids. Sulphides analysed in the Patawarta Diapir were chalcopyrite and pyrite from the Lady Lehmann and Lady Lennon workings. The results are shown below in Table 3.
TABLE 3  RESULTS OF SULPHUR ISOTOPE MEASUREMENTS

\[ \delta^{34}S \text{ per mil} \]

<table>
<thead>
<tr>
<th></th>
<th>-10</th>
<th>0</th>
<th>10</th>
<th>20</th>
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<tbody>
<tr>
<td>Pyrite</td>
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<tr>
<td>Chalcopyrite</td>
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</tr>
<tr>
<td>Mafic Igneous Rocks</td>
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5.2 Geothermometry

Sulphur isotope geothermometry is based on the equilibrium sulphur isotope fractionation between co-existing sulphur-bearing compounds, as the fractionation factor is inversely proportional to the temperature under equilibrium conditions. The sulphur isotope fractionations among sulphur species are given in Figure 13 (Ohmoto & Rye, 1979). The greater the separation of the curves for any two minerals the better the pair will be as an isotopic thermometer.

For application in geothermometry, one requires co-existing sulphides formed at the same time and in isotopic equilibrium and this is often difficult to achieve. For example, relationships between co-existing sulphide minerals in which later minerals were formed by replacement of earlier minerals do not always indicate
(b) $\delta^{34}S_{\text{ww}}$ (the sulphur isotopic composition of the fluid), which is controlled by the source of sulphur.

(c) the proportions of oxidised and reduced (i.e. $H_2S$ or $HS^-$) sulphur species in solution.

The origin of sulphur can be discussed only on the basis of the interpreted isotopic composition of the total sulphur in solution ($\delta^{34}S_{\text{SS}}$). Pyrite, for example, which along with chalcopyrite is one of the major sulphides found in the Patawarta Diapir, can obtain values of $\delta^{34}S$ (per mil) similar to the value of $\delta^{34}S_{\text{SS}}$ if:

(i) the sulphur in the fluids is dominated by one species eg. $H_2S$.

(ii) the relative isotopic enrichment factor between pyrite and the aqueous species is small.

Fluid inclusion studies show that homogenisation temperatures for the veins range from 142°C to 176°C (see figure 10) and for the purposes of this study a temperature of formation of 150°C will be assumed. The hydrothermal solutions are assumed to be slightly acidic or neutral (Anderson, 1973). Figure 11, (Crerar and Barnes, 1976), which shows the predominant regions of $H_2S$ and $HS^-$ and the temperature dependent curve for neutral pH, indicates a neutral pH for $H_2S$ of 5.9 for a temperature of 150°C.

The Fe-S-O stability fields and the sulphur species stability fields are plotted in Figure 12 for 150°C and for $\Sigma S = 0.001$ moles/kg $H_2O$. It can be seen that the pyrite stability field lies mainly in the $H_2S$ dominated sulphur species field for pH values less than 5.9. Thus condition (i) above is fulfilled.

Figure 13 shows that the isotopic enrichment factor between pyrite and $H_2S$ at 150°C is approximately +2 per mil, thus fulfilling condition (ii) above. (for chalcopyrite the corresponding value is -1 per mil).

Therefore the $\delta^{34}S$ values for the sulphide minerals can be assumed to approximate the $\delta^{34}S_{\text{SS}}$ values and so the results can now be interpreted on this basis.
FIGURE 11  Temperature dependent curve for neutral pH & predominance regions for HS\(^-\) & H\(_2\)S
(after Czeran and Barnes (1976))

FIGURE 13  Equilibrium Isotopic Fractionation Factors Among Sulfur Compounds Relative To H\(_2\)S
(after Rye and Ohmoto (1979))
FIGURE 12  Stability fields of Fe-S-O Minerals & sulphur species
(IS=0.001 moles/kg H₂O) (7°C = 450°)

FeS₂  FeS  Fe₂O₃  Fe₃O₄

Sulphur species boundary
The results fall into 2 groups and may indicate a dual source of sulphur in the deposits. The smaller group, consisting of only two pyrite samples, has slightly positive values which fall within the range for mafic igneous rocks. These values may reflect sulphur derived largely from igneous rocks, i.e. sulphur released from melts or sulphur leached from sulphides in igneous rocks.

At least the greater contribution of sulphur comes from the second group which have values ranging from 8.5 to 14.4 per mil. This may indicate a source from local country rocks either from sedimentary rocks or evaporites.

Figure 14 (Claypool et. al. 1980) summarizes the relationship between age and sulphur isotopic composition of sea water sulphates. A comparison of $^{34}$S of evaporites of Precambrian age shows them to be slightly higher that the $^{34}$S results for this study. This could be due to a sulphur component from a mafic igneous sulphur source which would lower the $^{34}$S values slightly.
FIGURE 14: Sulphur Isotope Age Curve For Sulphate

(after Claypool et al.(1980))
Figure 15 was constructed in an attempt to define the approximate physical and chemical conditions (T, fO₂, pH, δ³⁴S₈ S) of the hydrothermal fluids that precipitated mineralisation in the Patawarta Diapir as determined from a study of mineral assemblages and fluid inclusion data. Equilibrium constants for reactions used in the calculation of the figure are listed in Appendix 7. The figure has been constructed for a temperature of 150°C and ΣS = 0.001 moles/kg H₂O. This temperature is based on fluid inclusion measurements of homogenisation temperature and sulphur isotope fractionation between pyrite and chalcopyrite. (see sections 4.3 and 5.3) The ΣS concentration is considered reasonable for these small deposits. Any increase in ΣS will increase the area of pyrite stability field by increasing the fO₂ and pH range for pyrite. Pyrite and chalcopyrite have been precipitated throughout formation of the veins and so fO₂ and pH conditions are assumed to have been restricted to the common area shared by the pyrite and chalcopyrite stability fields. Bornite has not been observed in the mineral assemblages and so the chemical environment is assumed to have always been within the area in which chalcopyrite is the stable Cu-Fe-S species.

The diagram has been contoured for δ³⁴S₂py values by assuming a sulphur isotopic composition of the fluid of δ³⁴S₈ S = +10 per mil. The basis for this assumption is that the mean value of δ³⁴S₂py values (excluding the two lowest values) is approximately 12 per mil. Since the fractionation between pyrite and H₂S at 150°C is approximately +2 per mil (Ohmoto & Rye, 1979),
a contour of $\delta^{34}S_{H_2S} = +10$ per mil is equivalent to a contour of $\delta^{34}S_{P_Y} = +12$ per mil. Under the $fO_2 - pH$ conditions that existed during formation of the Patawarta mineralisation $\delta^{34}S_{H_2S}$ approximates $\delta^{34}S_{\Sigma S}$, since $H_2S$ would have been the dominant sulphur species.

It should be noted that $\delta^{34}S_i$ (species i) contour values have an uncertainty of $\pm 10\%$ due to uncertainties in $\Delta i$ values, but probably fall within $\pm 0.5 \log fO_2$ units for a given $pH$ (Ohmoto, 1972).

Approximate $pH$ constraints on the mineralising fluid chemistry can be derived from silicate alteration assemblages in the wall rocks of the veins. $K$-feldspar in the siltstones shows alteration to sericite and using the sericite boundary reaction,

$$3KAlSi_3O_8 + 2H^+ = KAl_2AlSi_3O_{10}(OH)_2 + 2K^+ + 6SiO_2$$

(orthoclase) (sericite)

an upper $pH$ limit can be given, assuming a $K^+$ concentration. Taking $ak^+ = 0.01$ gives a $pH$ of 7.1 which seems a realistic upper limit for hydrothermal brines. For comparative purposes $pH$ is also derived for an assumed $ak^+$ of 0.1.

Constraints on the $fO_2$ of mineralisation can be determined by considering the concentration of Fe in sphalerite. The diagram shows the pyrite field contoured for mole percent FeS in sphalerite. Electron microprobe analyses show that the FeS content of sphalerite in the veins varies between 0.2 and 0.6 mole percent (Appendix 6), indicating $log fO_2$ conditions of between -48.7 and -49.7.

A lower $pH$ constraint cannot be accurately determined but is unlikely to be less than 4.4 as experimental studies indicate that chloride brines co-existing with muscovite bearing assemblages are weakly acidic (Hemley et al, 1967) and neutral $pH$ at 150$^\circ$C is about 5.9.
FIGURE 15  Constraints on mineralisation ($T^°C \approx 150^°$)

Fe-S-0 mineral boundaries at
$\delta S = 0.001$ mole/kg H$_2$O

Bornite/Chalcopyrite boundary

Cu solubilities (ppm) calculated for $aCl^- = 3.0$
(fluid inclusion study & Helgeson, 1969)

mole% FeS in sphalerite

orthoclase/sericite boundaries for a given $aK^+$

$\delta^{34}S_{py}$ contours for $\delta^{34}S_{is}$ approximately 10%

proposed pH - log($IO_2$) conditions for mineralisation

FeS$_2$  FeS  Fe$_3$O$_4$  Fe$_2$O$_3$
7. GENESIS OF THE MINERALISED VEINS

In considering the genesis of the vein deposits in the Patawarta Diapir, four aspects have to be considered:

(i) a source for the constituents of the mineralisation must exist, i.e. magmatic or sedimentary rocks.

(ii) the dissolution of the mineralising constituents in the brine where the brine may be magmatic water, connate water, etc.

(iii) migration of the metal-bearing fluid.

(iv) selective precipitation of the mineralising constituents in favourable environments.

The source of Cu deposited in the diapir does not seem to pose any problems. Almost all diapirs in the Flinders Ranges show anomalous concentrations of Cu. Many other small deposits occur in Precambrian and Cambrian rocks throughout the Flinders Ranges and Mount Lofty Ranges and large Cu deposits occur at Burra and Kanmantoo. Uneconomic amounts of Cu are known to occur in some Adelaidean System sediments (e.g. Bunyeroo Formation) over distances of hundreds of kilometres and small Cu-bearing veins occur locally near many of the major faults. Therefore the whole region is clearly a Cu anomalous province and it is highly likely that the basement rocks, which underlie the Proterozoic sediments in the Adelaide Geosyncline, are enriched in Cu.

Fluid inclusions in minerals in the veins contain greater than 20 wt. % dissolved solids and are rich in Ca, Na, K and Cl ions. In a study of Mississippi Valley type brines which have similar constituents to those described above, Davidson (1968),
suggested that the brines were comparable with "connate waters which have derived the contents of salts by the diagenesis and leaching of evaporitic deposits". This agrees with the interpretation of the sulphur isotope results which indicate a predominant sulphur source from evaporites (section 5.3). In the Adelaide Geosyncline, evaporites are common in the Callanna Beds which represent the earliest Adelaidean sediments. This lithology is interpreted as forming in a sabkha and playa environment (section 2.3.1).

Bush (1970), has discussed the chemical composition of brines from a sabkha environment and comparison of these brines with the composition of fluid inclusions in the veins of the diapir show them to be quite similar. The only major difference is the Mg concentration which falls below the detection limits of the electron microprobe for the diapiric inclusions. The Mg contents of the sabkha brine are considerably higher. Therefore the origin of the vein forming solutions in the Patawarta Diapir could be through the compaction of Callanna Bed sediments. Compaction of sediments and escape of interstitial water start at the time of deposition and continue for millions of years where the initial driving force is lithostatic pressure (White, 1968). Fine grained sediments act as semipermeable membranes allowing selective escape of water and concentration dissolved components in the remaining pore fluids (White, 1968). The brine is later squeezed out of the pores and its reaction with carbonate rocks exchanges Na and Mg for Ca (White, 1968).

It has long been recognised that saline Na-Ca-Cl brines are capable of leaching base metals from the rocks and minerals through which they pass and transport them as chloride complexes over long distances. However, this ability is greatly reduced if sulphur species are present in the brines in any appreciable concentrations.

Over the years, many different mechanisms have been proposed to account for the transportation and precipitation of base metals and sulphur in hydrothermal deposits. These mechanisms can be grouped into three alternative models.
The first, known as a mixing model suggests that reduced sulphur is transported in one solution, base metals are transported in another solution and precipitation takes place where the two solutions mix. Variations on this model suggest that base metals are transported to the site of deposition where they encounter a source of reduced sulphur, such as diagenetic iron sulphides, which cause precipitation.

The second, a sulphate reduction model, suggests that base metals were transported with sulphate in solution and precipitation takes place due to a reaction which causes sulphate to be reduced to sulphide, for example, through the oxidation of organic matter (Barton, 1967).

The third, a reduced sulphur model involves base metals and reduced sulphur (H₂S or HS⁻) being transported together in the same solution at low concentrations. The cause of precipitation for this model could be (i) change of pH from acid to neutral (Helgeson, 1970) (ii) cooling or (iii) dilution (Anderson, 1973).

The reduced sulphur model would appear to provide a possible explanation for the Cu mineralisation in the diapir. Cu solubilities have been plotted for a total sulphur content of 0.001 moles/kg H₂O (Figure 16). The 0.001 ppm₅₀₀ contours intersect the log fO₂ - pH field indicated for mineralisation. Although this value for Cu solubility is low it should be remembered that the Cu content in the veins is quite low and is disseminated. Therefore, 0.01 ppm Cu in solution may provide sufficient Cu to generate a deposit of this size.

Dunham (1970) pointed out that one of the conditions necessary for the formation of a high concentration of sulphide is "long continuous flow along restricted paths which may be controlled by stratigraphy or structure". He noted that "if the paths are unrestricted, the mineralisation is likely to be dissipated" such may have been the case for the mineralisation in the Patawarta Diapir. White (1968) also discussed the behaviour of high chloride brines which will dissolve heavy metals and stated that "channels of flow eventually converge upwards towards any through going structures such as faults and fractures".
Many authors have noted that diapiric breccia zones occur in the cores of anticlines and domes which were zones of low pressure during folding. It has been suggested that these become zones of low hydrostatic pressure and this may have played a major role in directing the upward flow of deep formation brines through the rocks. Also, heat from possible post-Willouran igneous activity would have contributed to the increased circulation of deep formation brines through the breccias. Cu would be deposited where the brines encountered an increase in pH or decrease in temperature or underwent dilution.
CONCLUSIONS

1. Primary mineralisation consists of very disseminated chalcopyrite, pyrite, calcite with minor quantities of sphalerite, marcasite and quartz. Secondary alteration products are malachite, limonite, chalcocite and covellite.

2. Fluid inclusion studies indicate:
   (i) Uncorrected temperatures of mineralisation ranging from 142°C to 176°C with an average of 155°C.
   (ii) Salinities greater than 20 wt % CaCl₂ + NaCl equivalent.

3. Sulphur isotope studies indicate:
   (i) Some pyrite-chalcopyrite pairs approached sulphur isotopic equilibrium.
   (ii) A dual source for sulphur is probable with the major component derived from evaporites and a minor component derived from a mafic igneous source.

4. A homogeneous sulphur source $^{34}S_{\Sigma S} \approx +10$ per mil, together with $pH$ and $fO_2$ were stable during deposition of sulphides.

5. (i) The mineralising brines formed through compaction of evaporitic sequences of the Callanna Beds of Willouran age.
   (ii) The highly saline brines leached base metals from the basement complex and surrounding country rocks and transportation occurred through the zone of weakness caused by diapiric intrusion.
   (iii) Precipitation occurred in open fractures in blocks in the diapir due to change in $pH$, change in temperature or dilution.
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SHADE, J.W. (1974): Hydrolysis reactions in the SiO\textsubscript{2} - excess portion of the system K\textsubscript{2}O - Al\textsubscript{2}O\textsubscript{3} - SiO\textsubscript{2} - H\textsubscript{2}O in chloride fluids at magmatic conditions. Econ. Geol. v. 69, pp. 218-228.


APPENDIX 1

SELECTED ROCK DESCRIPTIONS

Sample 831-150  Diapiric carbonate breccia

Hand Specimen: brown-pale yellow weathered feldspathic
  dolomitic breccia

Micro Description:  
  Dolomite  60%  
  Quartz   10%  
  Feldspar 15%  
  Calcite  5%  
  Talc     3%  
  Hematite 5%  
  Limonite 2%

The rock is an obvious breccia with fragments of rock up to 2 cm
  across, making up to 60% of the sample with the carbonate matrix
  enclosing the fragments. Feldspar includes both microcline and
  plagioclase, occurring as anhedral grains. Quartz occurs as anhedral
  interlocking grains. Limonite is associated with dolomite in fractures and has stained the rock.

Sample 831-146  Rim Dolomite

Hand Specimen: buff coloured massive rock containing numerous clasts

Micro Description:  
  Dolomite (Matrix)  95%  
  Quartz           1%  
  Ferruginous material 4%
The bulk of the rock is made up of Callanna Bed clasts. The matrix dolomite forms a granular interlocked mosaic with variations in grain size in different parts of the slide. Grains vary up to 0.05 mm in size. Quartz is present as subhedral grains up to 0.3 mm in size.

Sample 831-55 **Diapiric Dolomite**

Hand Specimen: light brown fine to medium grained carbonate with minor quartz

**Micro Description:**

Dolomite 90%
Quartz 8%
Limonite 2%

The rock is composed of interlocking medium grained carbonate and quartz, clouded by limonite staining. Grains of dolomite are anhedral to polygonal interlocking crystals up to 0.3 mm in size. Quartz occurs on scattered anhedral grains. One grain of subhedral pyrite was noted.

Sample 831-01 **Altered Dolerite**

Hand Specimen: dark green fine grained altered rock

**Micro Description:**

Epidote 20%
Labradorite 10%
Albite 35%
Chlorite 5%
Calcite 15%
Pyroxene 10%
Actinolite 5%
Riebeckite 1%
Opales 4%

The rock is a medium grained saussuratised dolerite with euhedral laths of labradorite set in fine grained alteration minerals. Pyroxene occurs as euhedral relic fragments with a pink colouration suggesting titaniferous augite. Chlorite occurs as fine grained fibrous clusters. Fine grained actinolite appears to be replacing pyroxene. Epidote shows yellow pleochroism and occurs as clusters replacing pyroxene. 3 grains of riebeckite were noted.
Sample 831-75 Siliceous Mg-Fe rich rock

Hand Specimen: brown siliceous diapiric rock

Micro Description: Quartz 90%
Limonite 10%

A medium to fine grained rock with anhedral interlocking quartz. Limonite occurs around boundaries of quartz. Minute carbonate blebs within the quartz grains indicate a former abundance of carbonate which may have been leached.

Sample 831-43 Green-brown calcareous siltstone

Hand Specimen: Fine grained green-brown calcareous siltstone

Micro Description: Quartz 30%
Feldspar 40%
Calcite 15%
Opaques 15%

The rock consists of silt-sized quartz and feldspar grains with medium grained calcite grains. The quartz and feldspar grains are subangular to subrounded and appear to be detrital. Feldspar is predominately orthoclase with minor plagioclase. Opaques occur in bands as anhedral detrital grains.

Sample 831-44 Altered Basalt

Hand Specimen: Brecciated dark green medium grained basalt

Micro Description: Plagioclase 10%
Calcite 10%
Albite 30%
Chlorite 5%
Actinolite 20%
Hematite 15%
Epidote 10%
Laths of plagioclase (labradorite) show alteration to albite which is associated with epidote. Epidote occurs as clusters probably replacing pyroxene (?). Chlorite occurs as a fine fibrous form, occasionally intergrown with epidote. Hematite is disseminated and fine grained.

Sample 831-27 Heavy mineral laminated sandstone

Hand Specimen: Heavy mineral laminated sandstone, thinly bedded.

Micro Description: Quartz 80%
Feldspar 10%
Hematite 5%
Carbonate 1%
Mica 5%

Quartz occurs as poorly sorted fine grains containing overgrowths. Feldspar is thought to be detrital microcline. Hematite occurs as detrital grains and occurs in bands.

Sample 831-150 Hanging Wall Siltstone, Adit F, Lady Lehmann Workings

Hand Specimen: Green, brown siltstone, thinly bedded, with malachite staining

Micro Description: Quartz 10%
Feldspar 35%
Sericite 20%
Limonite 2%
Pyrite 3%
Malachite 10%
Chlorite 3%
Muscovite 2%
The sample consists of fine - medium grained detrital quartz and feldspar grains cemented by a fine grained phyllosilicate matrix. Quartz and feldspar grains are angular to subrounded. Many orthoclase grains show alteration to sericite, which comprises most of the phyllosilicate matrix. Trace muscovite and chlorite grains are randomly distributed and appear to be detrital. Pyrite grains are disseminated. Malachite forms fine acicular crystals and infills interstices between limonite and quartz and feldspar fragments. Limonite appears to have replaced some of the pyrite grains with a botryoidal form.

Sample 831-143 Vein material, Adit F, Lady Lehmann Workings

Hand Specimen: Massive calcite with disseminated sulphides

Micro Description: Calcite 75%
Quartz 10%
Feldspar 5%
Sulphides 10%

The vein gangue is made up of both coarse and fine grained fibrous calcite. Subhedral quartz grains occur in clusters and are medium grained. Occasional albite grains occur in association with quartz. The major sulphides are pyrite and chalcopyrite with grains ranging up to 5 mm in diameter. Pyrite shows an octahedral form.

Chalcopyrite typically comprises 60% of the total sulphides present. Small grains of sphalerite and marcasite are found in association with the major sulphides. Chalcopyrite contains inclusions of pyrite which is not a replacement feature and inclusions of calcite occur in both chalcopyrite and pyrite. Covellite and chalcocite occur as a compact mass of uniform polygonal grains occurring around the rims of the primary sulphides.
Sample 831-145  As above

Hand Specimen: As above

Micro Description:

This rock is very similar to 831-143, except that secondary alteration effects are far more noticeable. Limonite has replaced most of the sulphide grains and shows a botryoidal texture. Malachite forms fine crystals along the boundaries of the altered sulphides.

Sample 831-92  Siderite vein, Lady Lennon Workings

Hand Specimen: Light brown coarse siderite vein with abundant malachite staining and minor sulphides.

Micro Description:

<table>
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<tr>
<th>Mineral</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Siderite</td>
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<td>Calcite</td>
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<tr>
<td>Quartz</td>
<td>10%</td>
</tr>
<tr>
<td>Sulphides</td>
<td>5%</td>
</tr>
<tr>
<td>Malachite</td>
<td>10%</td>
</tr>
<tr>
<td>Limonite</td>
<td>5%</td>
</tr>
</tbody>
</table>

Coarse grained siderite containing minor quartz and calcite forms the gangue. Medium anhedral grains of chalcopyrite appear disseminated throughout the gangue. Minor anhedral disseminated grains of pyrite occur and pyrite is also present as inclusions within chalcopyrite. Limonite has replaced most of the sulphides and malachite occurs around the rims of the limonite as fine crystals.
Sample 831-85  Calcite-hematite vein, Mosleys Workings

Hand Specimen: Calcite material containing abundant hematite and blebs of malachite.

Micro Description:  
Calcite  60%  
Hematite  20%  
Malachite  10%  
Sulphides  5%  
Quartz  5%

Coarse grained calcite and disseminated hematite from the matrix in which very minor pyrite and chalcopyrite are contained. Limonite replacement is evident and malachite is abundant through the slide.

Sample 831-124  Calcite vein, Home Rule Flat Workings

Hand Specimen: Ferruginous coarsely crystalline calcite with malachite staining.

Micro Description:  
Calcite  70%  
Malachite  10%  
Sulphides  1%  
Quartz  2%  
Feldspar  5%  
Limonite  12%

Sample consists essentially of large calcite crystals up to several mm in size intergrown with plagioclase and very minor quartz. Semi opaque limonite is present throughout the slide and has replaced sulphides. Much of the calcite exhibits a colloform texture. Malachite is disseminated throughout the slide. One anhedral grain of electrum is surrounded by calcite.
## APPENDIX 2

### SELECTED ROCK ANALYSIS

<table>
<thead>
<tr>
<th>Sample</th>
<th>Description</th>
</tr>
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<tbody>
<tr>
<td>831-01</td>
<td>Dolerite</td>
</tr>
<tr>
<td>831-44</td>
<td>Basalt (non amygdaloidal)</td>
</tr>
<tr>
<td>831-19</td>
<td>Heavy mineral laminated sandstone</td>
</tr>
<tr>
<td>831-35</td>
<td>Black shale</td>
</tr>
<tr>
<td>831-139</td>
<td>Black limestone</td>
</tr>
<tr>
<td>831-62</td>
<td>Green calcareous siltstone</td>
</tr>
<tr>
<td>831-76</td>
<td>Siliceous Mg-rich rock, near Mt. Rugged Workings</td>
</tr>
<tr>
<td>831-39</td>
<td>Calcite vein, Mount Rugged</td>
</tr>
<tr>
<td>831-59</td>
<td>Siderite vein, Mosleys</td>
</tr>
<tr>
<td>831-83</td>
<td>Diapiric dolomite with malachite Staining</td>
</tr>
<tr>
<td>831-133</td>
<td>Pyritic black shale, Adit. F.</td>
</tr>
<tr>
<td>831-131</td>
<td>Footwall breccia, Adit. F.</td>
</tr>
<tr>
<td>831-61</td>
<td>Typical carbonate breccia</td>
</tr>
<tr>
<td>831-132</td>
<td>Hanging wall siltstone with malachite staining, Adit. F.</td>
</tr>
<tr>
<td>831-125</td>
<td>Calcite vein, Adit. F.</td>
</tr>
<tr>
<td>831-143</td>
<td>Vein material, Adit F.</td>
</tr>
<tr>
<td>831-92</td>
<td>Siderite vein with malachite staining, Lady Lehmann Workings</td>
</tr>
<tr>
<td>831-85</td>
<td>Calcite vein with Lemalite and malachite, Mosleys Workings</td>
</tr>
<tr>
<td>831-124</td>
<td>Calcite vein with malachite staining, Home Rule, Flat.</td>
</tr>
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</table>
APPENDIX 2 (continued)

Table A - Major Elements

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<tr>
<th>Sample</th>
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<th>19</th>
<th>35</th>
<th>139</th>
<th>62</th>
<th>76</th>
<th>39</th>
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<td>SiO₂</td>
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<td>50.25</td>
<td>64.60</td>
<td>57.31</td>
<td>27.52</td>
<td>57.62</td>
<td>76.92</td>
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<td>13.44</td>
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<tr>
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<td>4.54</td>
<td>5.39</td>
<td>12.06</td>
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<tr>
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<td>0.16</td>
<td>0.09</td>
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<td>0.30</td>
<td>0.08</td>
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<td>99.84</td>
<td>99.31</td>
<td>98.47</td>
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## APPENDIX 2 (continued)

### Table A - Major Elements

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## APPENDIX 2 (continued)

Table B - Trace Elements

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<td>61</td>
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<tr>
<td>132</td>
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<tr>
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### APPENDIX 3

**Trace Elements in Sulphides (ppm)**

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<th>Ti</th>
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### Fluid Inclusion Data Obtained From Patawarta Veins

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<th>Homogenisation Temperature °C</th>
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II

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<th>Homogenisation Temperature °C</th>
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## APPENDIX 5

### RESULTS OF THE SULPHUR ISOTOPE STUDY

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<th>SAMPLE</th>
<th>$^{34}\text{S}$ Chalcopyrite ($^{0/00}$)</th>
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<tr>
<td>LADY LEHMANN MINE:</td>
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<tr>
<td>831-136*</td>
<td>+10.4</td>
<td>+12.6</td>
</tr>
<tr>
<td>831-144*</td>
<td>+ 8.5</td>
<td>+11.4</td>
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* - coexisting sulphide pairs
# APPENDIX 6

## Electron Microprobe Analysis Of Concentration of Fe In Sphalerite

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<td>S</td>
<td>33.10</td>
<td>0.501</td>
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<tr>
<td><strong>Total</strong></td>
<td>100.21</td>
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2.

<table>
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3.

<table>
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4.

<table>
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<tr>
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<td><strong>Total</strong></td>
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APPENDIX 7

Equations and data used in construction of Figure 15

\[
\begin{align*}
\text{H}_2\text{S} &= \text{H}^+ + \text{HS}^- \\
\text{HS}^- &= \text{H}^+ + \text{S}^{2-} \\
2\text{H}^+ + \text{SO}_4^{2-} &= \text{H}_2\text{S} + 2\text{O}_2 \\
\text{HSO}_4^- &= \text{H}^+ + \text{SO}_4^{2-} \\
\text{H}_2\text{S} + 1/2 \text{O}_2 &= \text{H}_2\text{O} + 1/2 \text{S}_2 \\
\text{FeS}_2 &= \text{FeS} + 1/2 \text{S}_2 \\
3\text{FeS}_2 + 2\text{O}_2 &= \text{Fe}_3\text{O}_4 + 3\text{S}_2 \\
2\text{FeS}_2 + 3/2 \text{O}_2 &= \text{Fe}_2\text{O}_3 + 2\text{S}_2 \\
3\text{FeS} + 2\text{O}_2 &= \text{Fe}_3\text{O}_4 + 3/2 \text{S}_2 \\
\text{Cu}_5\text{FeS}_4 + 4\text{FeS}_2 &= 5\text{CuFeS}_2 + \text{S}_2 \\
\text{CuFeS}_2 + \text{H}^+ + 1/4 \text{O}_2 &= \text{FeS}_2 + \text{Cu}^+ + 1/2 \text{H}_2\text{O} \\
\text{CuCl} &= \text{Cu}^+ + \text{Cl}^- \\
3\text{KAlSi}_3\text{O}_8 + 2\text{H}^+ &= \text{KAl}_2\text{AlSi}_3\text{O}_{10}(\text{OH})_2 + 2\text{K}^+ + 6\text{SiO}_2
\end{align*}
\]

\[
\begin{array}{ccc}
\text{log K values} & \text{at 150°C} & \text{Reference} \\
\hline
-6.66 & 1 \\
-11.07 & 1 \\
78.10 & 2 \\
-3.79 & 2 \\
19.68 & 2 \\
-9.98 & 3 \\
40.30 & 4 \\
34.56 & 4 \\
70.66 & 4 \\
13.29 & 3 \\
6.90 & 5 \\
2.48 & 6 \\
10.2 & 7 \\
\end{array}
\]

Reference

1. Naumov et. al. (1971)
3. Schneeberg (1973)
4. Haas & Robie (1973)
6. Crerar & Barnes (1976)
7. Shade (1974)