Studies of VHMS-related alteration: geochemical and mineralogical vectors to ore

P439
Studies of VHMS-related alteration: geochemical and mineralogical vectors to ore

EXECUTIVE SUMMARY

AMIRA/ARC project P439
Volume 1, Final Report
May 1998
Centre for Ore Deposit Research,
School of Earth Sciences,
University of Tasmania,
GPO Box 252-79,
Hobart, Tasmania,
Australia 7001

http://www.geol.utas.edu.au/codes

another Pongratz Production 1998
Contents

Introduction .................................................................................................................. 1

Summary of geochemical and mineralogical vectors to ore, based on the seven case studies in P439
— Ross Large, Bruce Gemmell and Walter Herrmann ................................................ 5

Alteration halo model for the Rosebery VHMS deposit, western Tasmania
— Ross Large, Rod Allen, Mike Blake and Walter Herrmann ...................................... 13

Alteration model for the Hellyer VHMS deposit, western Tasmania
— J. Bruce Gemmell and Russell Fulton ..................................................................... 17

Mt Julia-Henty gold mine: Summary of alteration study — Tim Callaghan ..................... 21

Alteration halo model for the Zone 96 volcanogenic gold deposit, Henty gold mine, western Tasmania — Jason Beckton ................................................................. 27

Alteration zonation and geochemical dispersion at the Western Tharsis deposit, Mt Lyell, Tasmania: a summary — David Huston and Julianne Kamprad .......................................... 29

Alteration halo of the Thalanga VHMS deposit, north Queensland — Holger Paulick ......... 33

Alteration halo model for the Highway-Reward sub-seafloor replacement deposit, Mount Windsor Subprovince, Queensland — Mark G. Doyle .................................................... 39

Alteration case study of the Gossan Hill VHMS deposit — Robina Sharpe and Bruce Gemmell ...... 43

Carbonate alteration at the Rosebery mine: The relationships between alteration texture, paragenesis, chemistry of carbonate minerals, and distance to ore — Rodney L. Allen, Michael Blake and Ross R. Large ................................................................. 47

A molar element ratio analysis of lithogeochemical samples from the footwall andesite, Hellyer VHMS district, Tasmania, Australia — Clifford R. Stanley and J. Bruce Gemmell ........................................ 53

Chlorite alteration associated with syn-volcanic granites and Cu-Au mineralisation:
A pilot study along the Jukes Road — Bill Wyman ...................................................... 57

cont.

CODES: AMIRA/ARC Project P439 — Studies of VHMS-related alteration: geochemical and mineralogical vectors to ore. May 1998
Use of immobile elements and chemostratigraphy to determine precursor volcanics
— Walter Herrmann .................................................................................................................. 61

Development and use of the alteration box plot — Ross Large ............................................. 63

Application of PIMA and FTIR spectrometry to VHMS alteration studies — Walter Herrmann,
Michael Blake, Mark Doyle, David Huston and Julianne Kamprad ........................................... 65

P439 geochemical whole-rock and mineral database — Michael D. Blake .................................. 69

Discrimination of diagenetic, hydrothermal and metamorphic alteration
— Rodney L. Allen, Cathryn C. Gifkins, Ross Large and Walter Herrmann ............................. 73

Geochemical modelling of low temperature (5° to 150°C) seawater–andesite interaction:
Implications for regional alteration assemblages in VHMS districts
— Stephen B. Bodon and David R. Cooke ................................................................................... 79

Application of ironstone geochemistry to exploration for VHMS deposits in the
Mt Windsor Volcanic Belt — Garry Davidson ........................................................................... 81

Volcanic facies and alteration — J McPhie .................................................................................. 85

Alteration in different glassy volcanics with emphasis on early diagenetic alteration;
a case study from the Mount Black Volcanics — Cathryn C Gifkins ........................................... 91

Influence of volcanic facies on hydrothermal and diagenetic alteration: Evidence from the
Highway–Reward deposit, Mount Windsor Subprovince, Queensland — Mark G. Doyle ............... 95

Bibliography .............................................................................................................................. 101
Introduction

Project objectives P439

The five project objectives of P439 as outlined in the original research proposal are:

• To characterise the mineralogy and geochemistry for the various styles of hydrothermal alteration throughout the Mount Read Volcanics (MRV) and the Mount Windsor Volcanics (MWV). This will be based on mapping supported by whole-rock and trace element geochemistry, mineral chemistry, REE and stable isotope geochemistry.

• To determine the relationship between geochemical alteration patterns and sub-volcanic intrusions that are coeval with VHMS formation.

• To undertake case studies of alteration halos related to specific VHMS deposits with particular emphasis on hangingwall alteration, and the relationship between alteration patterns and volcanic facies.

• To develop a set of vectors towards ore based on the regional studies and ore deposit specific studies that can be applied in the exploration for VHMS deposits in submarine volcanic sequences throughout Australia. The vector matrix will include whole-rock, trace element, mineral chemistry, REE, isotope and volcanic facies factors.

• To produce a book on “Alteration mineralogy, geochemistry and textures in volcanics related to VHMS deposits” as a follow-up to the successful publication by CODES of “Volcanic Textures”.

At the first progress meeting (April 1995), and in subsequent meetings, a number of sponsors emphasised the importance of including studies on mineral chemistry in both the regional and deposit case studies. In response to this request the work on mineral chemistry was expanded in the latter part of the project at the expense of some of the planned isotopic and REE research.

Research framework

This project involved a multidisciplinary approach utilising studies in volcanic facies analysis, volcanic petrology and geochemistry, with alteration and mineralisation to develop models for the composition, style and extent of alteration throughout submarine volcanic environment hosting VHMS deposits. The venn diagram below outlines the relationships between the different modules of the project.
The research was undertaken at the regional scale and the deposit scale in order to develop criteria useful for both regional exploration and mine scale exploration. The regional studies concentrated in the Mt Read Volcanics with lesser work in the Mt Windsor Volcanics. The emphasis was (a) to determine the relationship between volcanic facies and alteration, and (b) to develop criteria to distinguish amongst diagenetic alteration, metamorphic-related alteration and hydrothermal alteration (VHMS related). The deposit case studies included research at Mt Lyell, Hellyer, Rosebery and Henty in the Mt Read Volcanics; Thalanga and Highway-Reward in the Mt Windsor Volcanics and Gossan Hill in the Murchison Volcanics (WA). The emphasis in the deposit studies was (a) to map out alteration assemblages and zonation, (b) relate alteration mineralogy and geochemistry, (c) investigate trace element halos, and (d) to study variations in mineral chemistry and their relationship to alteration zonation. Because considerable research has been published about VHMS footwall alteration, then the focus of the P439 deposit studies was on hangingwall alteration. Our ultimate objective was to combine the regional and deposit scale studies to characterise hydrothermal alteration and develop vectors to ore.

Achievements of project

This project has made a major advance in our understanding and interpretation of alteration in submarine volcanics related to VHMS hydrothermal systems. The principle achievements listed in the order that they appear in this final report are given below.

1. Detailed halo alteration models have been developed for the following seven VHMS deposits; Hellyer, Rosebery, Henty, Western Tharsis, Thalanga, Highway-Reward and Gossan Hill.

Each model includes a set of criteria and vectors that are useful for regional and mine scale exploration. This is the first time that a set of VHMS alteration models has been presented which include volcanic facies, chemostratigraphy, lithogeochemistry, alteration zonation, halo geochemistry, mineral chemistry and vectors to ore.

2. The deposit case study reports (Volume 2) include a series of Alteration Data Sheets which combine geological, geochemical, textural and mineralogical information, including photos on the one page, to allow comparison of the main characteristics of alteration. This format was developed during the project and emphasises the advantages of a multidisciplinary approach in the study of alteration.

3. An in-depth evaluation of the application of PIMA to alteration studies around VHMS deposits has been completed with emphasis on the spectral characteristics of muscovite and their relationship to VHMS deposits (Western Tharsis, Rosebery and Highway-Reward). This work combines PIMA spectral data, with micro-probe mineral chemistry, whole-rock chemistry and thin section petrography. No study of this type has been previously completed for VHMS deposits.

4. A set of criteria have been developed to distinguish diagenetic alteration, metamorphic alteration and hydrothermal (VHMS-related) alteration in submarine volcanics. The criteria combine studies on volcanic facies, volcanic textures, petrology and lithogeochemistry.

5. An alteration box plot has been developed and tested, which enables comparison of alteration mineralogy with geochemistry to provide a classification of alteration facies related to VHMS systems.

6. A Pearce element ratio analysis of altered and unaltered rock compositions within the andesitic footwall to the Hellyer VHMS deposit was revealed several new concepts regarding the hydrothermal alteration. In addition, new innovative exploration vectors have been developed.
7. Preliminary thermodynamic modelling of low temperature seawater-volcanic rock interaction has been completed, and demonstrates the value of this approach in determining the temperature and fluid chemical controls on diagenetic and hydrothermal alteration.

8. A data base of whole rock and trace element geochemistry for samples from the Mt Read Volcanics, Mt Windsor Volcanics and the seven ore deposit case studies has been compiled. The data base includes 1730 rock analyses (this project), 2845 MRV rock analyses from other sources (Tas Uni, MRT and Pasminco), and 1020 microprobe mineral analyses from the deposit studies. The data base is provided to all sponsors on a compact disc located in the back envelope of this report (Volume 1).

9. A study of ironstone geochemistry, including REE and isotopes has shown the application of this approach to exploration in the Mt Windsor Volcanics.

10. Controls on both diagenetic alteration and hydrothermal alteration by volcanic facies and volcanic textures has been evaluated and shown to be critically important, especially at the diagenetic stage. This work includes regional studies especially the Mt Black Volcanics, and focussed research at Rosebery and Highway-Reward.

Acknowledgements

This project has been supported by the following companies and organisations:

Aberfoyle Resources Limited
Copper Mines of Tasmania
Denehurst Limited
Mineral Resources Tasmania
Normandy Exploration
Pasminco Exploration
Queensland Metals Corporation Ltd
RGC Exploration
Rio Tinto Exploration

On behalf of the research team I would like to thank all the company representatives, collaborators and AMIRA for their support and scientific interaction throughout this three year program. In particular we are indebted to those companies that provided access to their deposits for the case studies: RGC, Pasminco, Murchisen Zinc, Copper Mines of Tasmania and Aberfoyle.

This project would not have been possible without the very significant contributions by a number of dedicated and hardworking postgraduate students: Catherine Gifkins, Robina Sharpe, Mark Doyle, Holger Paulick, Bill Wyman and Russell Fulton. Thanks also for the tireless work of Mike Blake in organising the alteration data sheets, and to Nilar Hlaing for printing and collating the papers. As usual June Pongratz has done a great job in coordinating and producing this final report.

We hope that the final outcome of this project is the discovery of a new VHMS deposit in the near future.

Ross R. Large
Director
Centre for Ore Deposit Research.
Summary of geochemical and mineralogical vectors to ore, based on
the seven case studies in P439

Ross Large, Bruce Gemmell and Walter Herrmann
Centre for Ore Deposit Research

Although there are many features of similarity in the
alteration halos surrounding the seven volcanic
hosted deposits involved in this study, there are also
significant differences that need to be taken into
account in the development of VHMS exploration
programs.

The most useful general vectors to ore are listed
below. Not all vectors are useful for each deposit and
a subset is commonly required.

Whole-rock vectors
• Ishihawa alteration index

\[
AI = \frac{100(K_2O + MgO)}{(K_2O + MgO + Na_2O + CaO)}
\]

• Chlorite/carbonate/pyrite index

\[
CCPI = \frac{100(MgO + FeO)}{(MgO + FeO + Na_2O + K_2O)}
\]

• Thallium and Antimony.
• S/Na_2O ratio.
• Ba/Sr ratio and/or Rb/Sr.

Mineral chemistry vectors
• Mn and/or Fe content of carbonate measured by
  microprobe or whole-rock calculation (calibrated
  by microprobe).
• (Mg + Fe) content of muscovite (phengicity)
  measured by ALOH I using PIMA or FTIR.
• Be content of muscovite (measured by microprobe).
• Mg/Mg+Fe molar ratio of chlorite (magnesium
  number) measured by microprobe. Does not
  appear to correlate with any PIMA spectral
  parameters.

The reader is referred to the individual ore deposit
summaries that follow in selecting the appropriate
basket of vectors for each style of VHMS.

Ishihawa Alteration Index: All deposits show a
zone of feldspar destruction accompanied by sodium
depletion and increased AI values. A diagrammatic
comparison of the shape and extent of the AI halos
is shown in Figure 1. These halos typically show a
zonation of sericite-quartz \(\rightarrow\) chlorite + sericite \(\rightarrow\)
chlorite + sericite approaching the ore.

Chlorite-Carbonate-Pyrite Index (CCPI) Chlorite
and pyrite typically increase approaching the ore
zone, leading to an increase in the CCPI. The most
common pattern is increasing AI (due to sericite
alteration) followed by increasing CCPI (due to
chlorite + pyrite alteration) as the ore zone is
approached. Chemical trends on the AI-CCPI box
plot are discussed in detail by Herrmann and Large
(this volume).

Carbonate Alteration: The various patterns of
carbonate alteration in the deposits studied are shown
in Figure 2. The carbonate generally occurs in dissemi-
|inations or spots from 2 to 20 wt%, although massive
carbonate alteration zones occur at the ore position
in Rosebery, Hercules and Thalanga. At Henty
carbonate occurs with silica in the gold-rich MQ zone
and lenses of massive carbonate are present toward
the stratigraphic top of the mineralised unit. Ore-
related carbonates are typically Fe, Mg and Mn
varieties while later overprinting metamorphic
carbonates are typically of calcitic composition. At
Rosebery the Mn content of alteration carbonate
Figure 1. Comparative sketches showing the extent of alteration halos of sericite±chlorite±quartz surrounding VHMS deposits investigated in this project.
increases toward ore (ankerite → ktnahorite → ferroan rhodochrosite → rhodochrosite), at Western Tharsis the Fe-content of carbonate increases to ore (ankerite & siderite), at Hellyer the alteration dolomite becomes more Mn-Fe-rich toward ore in the hangingwall, and at Gossan Hill the cre zone carbonates are manganese, however there is no overall vector to ore.

**Footwall Alteration:** All deposits studied exhibit intense footwall alteration trending toward high Al (>50) and high CCPI (>80) as the ore is approached. In most cases the mineral zonation is sericite dominant → chlorite dominant → quartz dominant. Hellyer is the only deposit in the group with a well defined and zoned alteration pipe. The other deposits have more diffuse, stratabound footwall alteration zones that are thickest and most intense below the centre of the orebody and thin to the margins (e.g., Thalanga and Rosebery).

**Hangingwall Alteration:** The stratiform, sheet-like deposits at Rosebery and Thalanga show no obvious visual hangingwall alteration in drill core. However lithochemical studies at Rosebery (Large, et al., this volume) have shown that a weak chemical halo depicted by Ba/Sr and Mn content of carbonate extends up to 100 m into the hangingwall. No similar chemical halo in the HW could be defined at Thalanga (Paulick, this volume).

By contrast the Cu-Au deposits at Western Tharsis and Highway-Reward (Fig. 1) exhibit extensive zones of hangingwall alteration (up to 200 m thick) of similar mineralogy, but generally less intensity, than the footwall alteration. At Hellyer alteration is focussed into a hangingwall plume (Fig. 1) (core to rim: fuchsite → chlorite-carbonate → albite-quartz), while at Gossan Hill the copper-magnetite orebody is encased by strong quartz-chlorite ± carbonate alteration and the upper zinc lens has a weak zone of hangingwall alteration associated with the Au marker.

**Albite Alteration:** Plagioclase (including albite) is commonly replaced by sericite, carbonate or quartz in the footwall zones to VHMS deposits, leading to complete sodium depletion. However this study has shown that hydrothermal albite alteration may develop in the hangingwall to massive sulfides. Hellyer, Henty and Thalanga all contain albite-quartz alteration zones in the hangingwall. This style of alteration can form a halo several hundred meters, both vertically and laterally away from the mineralisation and can develop in coherent volcanics (Hellyer and Thalanga) or volcaniclastics and sediments (Henty). Sodium, for the formation of albite, comes from the breakdown of plagioclase feldspar and/or from the hydrothermal fluids responsible for alteration (Na likely from destruction of feldspar and Na-pletion in the footwall). Rarely paragonite (Na-rich muscovite) is developed in hangingwall alteration zones (e.g. Rosebery and distal areas at Hellyer). Sodium addition in the albite ± quartz zones leads to a decrease in Al and CCPI values, even though the rocks may be intensely altered.

**Thallium and Antimony Halos:** Our research has shown that both thallium and antimony are particularly useful vectors for the stratiform zinc-rich styles of VHMS such as Rosebery, Hellyer and Thalanga. They are less useful for copper-rich and cross cutting systems such as Western Tharsis and Highway Reward.

A comparison of all Tl and Sb data for six of the deposits is shown in Figure 3 and an outline of the best developed halos (Rosebery, Hellyer and Thalanga) is shown in Figure 4. Rosebery shows the best developed Tl and Sb halo, with values ranging up to 100 ppm Tl and Sb proximal to ore and 1-10 ppm within the extensive halo zone. Hellyer also shows a well developed Sb and Tl halo within the HSV host horizon and extending into the hangingwall basalt which can be traced for up to 350 m west of the deposit.

At Thalanga Tl shows anomalous values (>1 ppm) up to 50 m into the HW and FW and along the favourable horizon, and a lower Sb/Tl ratio than the other deposits. The copper-rich orebodies at Gossan Hill, Highway Reward and Western Tharsis do not show significant halos of either Tl or Sb, compared with the zinc rich ores. This probably relates to the higher temperature of emplacement of copper mineralisation and the near complete dispersion of Tl and Sb.
a ROSEBERY

- Mn-rich carbonate
  (kutnahorite, rhochochrosite)
- Mn content increases towards ore

b HELLYER

- Ferroan dolomite & ankerite
- Mn content increases towards ore in hangingwall

c THALANGA

- Carbonate-territe
  - chlorite lenses

- Calcite overprinting dolomite
- no chemical trend to ore

d WESTERN THARSIS

- Ankerite & siderite
- no trends to ore
- most Mn-rich carbonate
  in zinc ore & veins

- Ankerite & siderite
- Fe content of carbonate increases to ore

Figure 2. Comparative sketches showing extent of carbonate alteration at Rosebery, Hellyer, Thalanga, Western Tharsis and Gossan Hill.
Muscovite Chemistry: A combination of microprobe and PIMA studies have enabled a detailed analysis of muscovite chemistry in five of the deposits.

Rosebery
- proximal to ore muscovite is phengitic with 0.5 to 1.1 Fe + Mg ions in octahedral site AIOH λ >2210 nm.
- Ba content increases (0.03 to 0.1 ions) proximal to ore.
- Hangingwall zone of anomalous sodic muscovite with Na/Na+K = 0.1 to 0.4, Al(OH) λ <2200 nm.

Hellyer
- Ba content of muscovite from the hangingwall basalt increases towards the deposit.
- Low Cr content (up to 0.3%) in white micas from intensely fuchsite altered hangingwall basalts.
- Fe + Mg content of hangingwall muscovite has limited range above deposit, but erratic values away from deposit.
- PIMA analyses indicate that muscovite in the footwall alteration pipe is enriched in Fe and Mg compared to muscovite outside the pipe.

Western Tharsis
- Phengite content of muscovite in the alteration zone decreases systematically towards ore.
- Ore zone is characterised by muscovite with AIOH λ = 2195 nm, <0.2 Fe + Mg ions in octahedral site.
- Outside ore zone AIOH λ = 2200 to 2210 nm.

Highway/Reward
- Same trends as shown by Western Tharsis.
- Smooth trend of decreasing AIOH λ of muscovite from a background of 2202 to 2222 nm down to 2195 to 2204 nm in the sericite-quartz-pyrite alteration zone adjacent to ore.

Thalanga
- No trend in muscovite composition was observed relative to the ore zone.

Chlorite Chemistry: Limited microprobe studies of chlorite have been undertaken on selected deposits. The PIMA has not contributed to a better understanding of chlorite chemistry (see Herrmann et al., 1998, vol. 2).

Rosebery
- Ore zone chlorite has Mg number from 50 to 75, the same as the hangingwall volcanic sandstone unit.
- Footwall chlorite alteration is dominantly Fe-rich, while ore zone and hangingwall chlorite is Mg-rich.
Hellyer
- chlorites in hangingwall alteration plume have Mg\# from 40–80, while chlorites from outside plume have Mg\# over limited range of 63–70.
- chlorites in hangingwall have lowest Mg\# closest to orebody.
- some hangingwall chlorites are enriched in Cr (up to 1.9%), higher than white micas (e.g., “fuchsite”).
- highest Cr values in chlorites within hangingwall alteration plume.

Western Tharsis
- Possible trend of decreasing Mg number towards ore (i.e., chlorite becomes more iron-rich).
- Chlorite in hangingwall alteration is more Mg-rich than chlorite in footwall alteration zone. This is the same pattern as Rosebery.

Thalanga
- Chlorite in footwall alteration zone exhibits a systematic increase in Mg number towards ore from values of 40 to 50 in the least altered rhyolite to 85 to 95 in the altered rhyolite close to ore.
- Biotite shows the same change in Mg number as chlorite.

Henty and Gossan Hill: Both the Henty and Gossan Hill deposits do not follow the alteration patterns displayed by either the Zn-Pb or Cu-Au styles studied. Henty displays more intense silica and carbonate alteration with an unusual zone of hangingwall albisation. Gossan Hill copper ore is encased within an intense silica-chlorite alteration zone of regional extension with strong alteration in both the footwall and hangingwall positions.

Conclusions

The extent and nature of the hydrothermal alteration halos surrounding VHMS deposits depends on such factors as the local structure and fluid pathways, composition of the host volcanics, attitude of the ore (i.e., stratiform or cross cutting) and the temperature and pH of the hydrothermal fluids.

The high temperature, sub seafloor Cu-Au replacement systems at Western Tharsis and Highway-Reward have extensive alteration zones through both footwall and hangingwall lithotypes. Both systems are dominated by quartz-sericite-pyrite alteration halos. Within these zones of about 300 to 400 m across, the muscovite chemistry changes systematically toward ore; becoming less phengitic (i.e., decreasing Fe + Mg ions in octahedral site). At Western Tharsis (and possibly also Highway-Reward?) the quartz-sericite-pyrite zone is surrounded by an outer halo of carbonate alteration which extends from 200 to 500 m from ore. Carbonate minerals in the halo increase in iron content (ankerite to siderite) approaching the ore. An unusual zone of propylitic alteration and extreme K₂O depletion occurs within 70 m of the ore at Western Tharsis, and is interpreted by Huston and Kamprod (this volume) as a magmatic-hydrothermal high sulfidation alteration zone.

The lower temperature and more stratiform Zn-Pb-rich VHMS deposits at Rosebery and Thalanga exhibit more restricted alteration halos that are generally aligned parallel to stratigraphy. In these systems the most intense and extensive development of alteration occurs in the footwall volcanics and along the base of the ore position away from the deposits. Carbonate alteration occurs proximal to ore and may be strongly manganiferous (e.g., Rosebery and Hercules).

In contrast to Rosebery and Thalanga, the mound-style massive sulfide at Hellyer has a restricted zoned footwall alteration pipe, which cross cuts stratigraphy. The pipe has lateral dimensions similar to the overlying deposit and extends downwards for several hundreds of metres.

Hanging wall alteration in the Zn-Pb ore systems is variable:
- at Hellyer an irregular alteration plume of fuchsite-carbonate alteration extends up to 200 m above the deposit. This alteration is surrounded by zones of chlorite-carbonate and albite-quartz. This hangingwall plume can extend for up to 250 m laterally from the deposit.
- at Rosebery, hangingwall alteration is very weak but can be detected by the Ba/Sr ratio and Mn content of carbonate up to 100 m above the deposit.
- at Thalanga no systematic hangingwall alteration could be defined.
Figure 4. Sketch of the extent of thallium halos at Rosebery, Hellyer and Thalanga.
Alteration halos of the volatile elements Tl and Sb are extremely variable across the seven deposits studied: Rosebery have extensive and highly anomalous Tl halos, while Thalanga has a less well developed smaller halo. The copper-rich orebodies at Highway Reward and Gossan Hill show no obvious Tl or Sb halos. A very weak Tl halo (0.5 to 1 ppm) occurs remote from Western Tharsis in the outer sericite-carbonate alteration zone.

Acknowledgements

Thanks to Robina Sharpe, Holger Paulick, David Huston, Mark Doyle, Tim Callaghan, Jason Beckton and Russell Fulton for providing the information which contributed to this Summary.

References


Alteration halo model for the Rosebery VHMS deposit, western Tasmania

Ross Large, Rod Allen, Mike Blake and Walter Herrmann

Centre for Ore Deposit Research

Summary

A detailed study of alteration mineralogy, mineral chemistry and lithogeochemistry in the host rocks surrounding the A-B and K lenses at the north-end of the Rosebery mine has revealed a series of overlapping alteration halos with varying mineralogy and geochemistry.

The study involved logging and sampling (255 samples) from nine drill holes spaced at varying distances from the A-B and K lenses. The stratiform Zn-Pb-Cu ore lenses, have a sheet-like attitude and are hosted by medium grained feldspar-phryic pumice and/or crystal-rich rhyolitic to dacitic volcanioclastics, overlying a thick homogenous sequence of rhyolitic pumiceous breccia mass flows.

Chemostratigraphic studies using immobile element geochemistry have revealed that the immediate host unit and overlying quartz-feldspar-biotite porphyry have a distinct Ti/Zr ratio (12–14) which is consistently different from the ratio in the footwall pumiceous mass flows (7 to 9), and the hangingwall volcanogenic sandstones (10 to 30). This result emphasises the value of chemostratigraphic studies, in parallel with conventional drill core logging, to aid identification of favourable units for mineralisation.

The major alteration minerals at Rosebery are arranged in a complex series of zones passing away from the deposit; quartz-sericite zone, Mn-carbonate zone, chlorite zone and outer sericite zone (Fig. 1). The outermost visible sericitic alteration extends about 60 to 100 m into the footwall, 10 to 20 m into the hangingwall and over 500 m along the upper contact of the rhyolitic pumiceous mass flows (Ti/Zr = 7 to 9).

A series of lithogeochemical halos with associated vectors have been developed which are critical in exploration either at the mine scale or in regional exploration (Fig. 2):

- Proximal vectors — Zn, Pb, MnO, Ba are d18O carbonate; may predict ore lenses at 200 to 500 m along strike and 3 to 50 m across strike.

- Medial vectors — S/Na2O, Ba/Sr, K2O, Na2O depletion, Ishikawa Al, CCPI and Mn content of carbonate; may predict presence of ore at 500 to 1000 m along strike, 10 to 80 m into hangingwall and 60 to 120 m into footwall.

- Distal vectors — Ti and Sb may predict presence of ore at greater than 1000 m along strike, 200–300 m into hangingwall and 60–100 m into the footwall.

Detailed studies of alteration mineral chemistry at Rosebery have revealed some important chemical relationships that may assist with exploration. The Mn content of alteration carbonate increases toward ore both along strike and across strike. Close to ore, alteration carbonates contain >20 mole % MnCO3 (kutnahorite, rhodochrosite, siderite and ankerite) while at distances of 40–60 m across strike the mole % MnCO3 in carbonate drops to below 10%. At greater than 80 m the carbonates are Mn poor calcites, and commonly located in syn-metamorphic structures.

Muscovite chemistry changes with stratigraphy and alteration assemblages and may be related to the mineralising event, although this has not been convincingly demonstrated. Proximal muscovite contains minor Ba substituting for K in the structure, and are phengetic with 0.5 to 1.0 (Fe + Mg) cations.
substituting for octahedral Al. However, except for their Ba content, these phengitic muscovites are similar to those found in non-mineralised areas of the MRV (Herrmann et al., this vol.). Sodic muscovite with up to 0.35 Na/(Na + K) and low phengite content (<0.5 Fe + Mg cations) occur in a zone of volcanic sandstones and black slates overlying the ore deposit. This may be a regional feature, although similar sodic muscovites have not been identified elsewhere on the MRV regional traverses.

Chlorites in the ore lenses, associated Mn-carbonate zone and the hangingwall volcanic sandstone unit are Mg-rich with a Mg number of 50 to 70. This compares with chlorite in the footwall and above the Black Slate which are generally more Fe-rich with Mg number from 20 to 50. More work is required on chlorites at Rosebery to test their relationship to mineralisation.

A study of carbon and oxygen isotopes in carbonates has found no extensive isotope halo at Rosebery, but demonstrates that C/O isotopes can be used to assist in the discrimination of carbonates associated with Cambrian VHMS mineralisation from carbonates related to metamorphism or granite emplacement during the Devonian.

In conclusion this study has identified a series of geochemical vectors and mineralogical/ isotopic criteria that are of considerable importance in the exploration for sheet-style VHMS deposits of the Rosebery type. Listed in approximate order from
proximal to distal the key vectors are:
Pb, Zn, MnO, Mg number of chlorite, δ18O_carb, Ba,
mole % MnCO3 in carbonate, CCPI, Ishikawa Al,
Na/(Na + K) muscovite [PIMA AlOH I], K2O, Na2O
depletion, Sb and Tl.

Bibliography

Allen R.L. and Large R.R., 1996, Rosebery Alteration
Study: AMIRA P439 Report 3, October 1996,
p143–152.
Large, R.R., 1997, The Hercules-Mt Read traverse:
Relationships between volcanic mineralogy,
alteration and geochemistry: AMIRA P439,

on the Rosebery lithoclochemical halo study:
Allen, R.L., 1997, Rosebery alteration study and
regional alteration studies in the Mount Read
Volcanics. The record of diagenetic alteration
in the strongly deformed, felsic volcaniclastic
succession enclosing the Rosebery and
Hercules massive sulphide deposits: AMIRA
Large, R.R., Allen, R.L. and Blake, M., 1997, Carbonate
and muscovite mineral chemistry, Rosebery
VHMS deposit, Tasmania: AMIRA P439,
Large, R.R., Allen, R.L., Blake, M. and Herrmann, W.,
1998, Alteration halo model for the Rosebery
VHMS deposit, western Tasmania. CODES
AMIRA/ARC Project P439, Final report.

---

Figure 3. Diagrammatic representation of the extent of the three major geochemical halos at Rosebery (based on the B and K lenses).
Alteration model for the Hellyer VHMS deposit, western Tasmania

J. Bruce Gemmell and Russell Fulton
Centre for Ore Deposit Research

The Hellyer massive sulfide deposit is a major (17 million tonnes), high grade (13.0% Zn, 6.8% Pb, 0.3% Cu, 160 g/t Ag, 2.3 g/t Au), mound-style polymetallic VHMS within the Mt Read Volcanics of Western Tasmania. Hellyer is hosted by the Que-Hellyer Volcanics, a 1 km thick suite of Late Middle Cambrian mafic to felsic lavas and volcanoclastics. Footwall to the mineralisation is andesitic lavas and polymictic debris flow sediments. The ore horizon is marked by polymictic debris flow sediments that contain clasts of massive sulphide and barite. The hangingwall consists of basalt flows, pillow lavas, breccias and interpillow sediments (Hellyer Basalt). Overlying the basalt is black shale (Que River Shale) and felsic volcanoclastics and sediments (Fig. 1).

Halo model

Alteration in district wide footwall geology is confined to patchy silicification, albisation, chloritisation, and minor sericitisation, epidotisation and hemitisation of the footwall lithologies east of Hellyer, with only patchy sericitisation and chloritisation observed to the west. Underlying Hellyer is a zoned alteration zone and stringer vein system. Mineralogical zoning exists within a footwall alteration pipe, with a central siliceous core giving way to zones of chlorite, chlorite-carbonate, sericite, and finally sericite-quartz (stringer envelope zone) on the margins. Pervasive alteration started with development of sericite-quartz that was subsequently overprinted by sericite, then chlorite and finally quartz as hydrothermal activity became focused towards the center of the alteration zone (Gemmell & Large, 1992).

The major hangingwall alteration assemblages are characterised by the dominant alteration mineral and the associated alteration assemblages. These are (i) silica-ablite, silica, silica-ablite-chlorite, (ii) fuchsite, fuchsite-carbonate, (iii) carbonate, carbonate-silica-carbonate-sericite, (iv) chlorite chlorite-sericite, chlorite-fuchsite and (v) sericite, sericite-fuchsite. In general, there is a zonation (fuchsite → carbonate-chlorite → silica-ablite) from the middle of the hangingwall alteration zone to the outsides, although there is some asymmetry. In cross section the hangingwall alteration is best developed directly above the middle and northern portions of the deposit and laterally away from the deposit to the south. Fuchsite-dominated alteration occupies the central portion of the hangingwall alteration. Chlorite and chlorite-carbonate alteration envelopes the fuchsite zone with carbonate-rich zones near to the ore deposit and chlorite-rich zones extend above and to the sides of the carbonate. The outermost alteration zone is characterised by silica-ablite alteration, which is best developed to the south and extends laterally away from the deposit for up to 250 m. To the east the silica-albite zone is best developed near the orebody.

The primary alteration phase to affect the Hangingwall Volcanoclastic Sequence (HVS) is sericite and pyrite.

Vectors to ore

A lithogeochemical halo model, which includes vectors to ore for the district-wide footwall lithologies, the alteration pipe, the Hangingwall Volcanoclastic Sequence (HVS), the Hellyer Basalt and Que River Shale (Fig. 2).
Important exploration vectors are:

**District-wide Footwall:**
- decrease in Na₂O, Cu, Zn, and As away from alteration pipe
- increase in whole rock δ¹⁸O values from 6-8 (within pipe) to 10–11‰ (within 1 km) to 12–14‰ at a distance of 2–3 km away

**Footwall Alteration Pipe:**
- distinct mineralogical zonation (from core to rim: quartz→chlorite (± carbonate)→sericite→quartz-sericite)
- decrease in Al, Ba, S, Ca, Fe, Cu, Pb and Zn from interior to edge of pipe
- increase in Na₂O, Ni, Cr, Sr, Al and δ¹⁸O in stringer veins from interior to edge of pipe

**Hangingwall Volcaniclastic Sequence (HVS)**
- decrease in Al, K₂O, Ti and Sb away from deposit (distance of 2 km)
- increase in P₂O₅, CaO and MnO away from deposit (distance of 2 km)

**Hellyer Basalt**
- basalt overlying deposit (Core Lava) has Ti/Zr = 55, basalt above and away from deposit (to a distance of 3 km) have Ti/Zr between 20 and 40
- alteration zonation (outwards from deposit: fuchsite→carbonate-chlorite→silica→ablite)
- Ba in muscovite decreases away from the deposit
- Mn in carbonate decreases away from the deposit
- decrease in Sb away from deposit (to a distance of 3 km)
- increase in whole rock δ¹⁸O values from 9–12 (within hangingwall alteration) to 10–14‰ in unaltered basalts away from the deposit

**Que River Shale (QRS)**
- no visible alteration
- increase in SiO₂, Ba and pyrite in QRS overlying hangingwall alteration
- decrease in FeO, MgO and CaO in QRS overlying hangingwall alteration zone of increased Pb, Zn, Fe, Ni, V, As and Sb lies approx. 50m vertically into QRS (source and significance of this metal enrichment is unknown)

**Mineral chemistry**

**Chlorite**
Urabe et al. (1983) determined that the Fe/(Fe+Mg) ratio of chlorite is a useful parameter for quantifying hydrothermal alteration associated with volcanic-hosted massive sulfide deposits. The Fe/(Fe+Mg) ratio in chlorites usually increases with distance from the core of the alteration pipe. However, the opposite relationship (decrease in chlorite Fe/(Fe+Mg) with distance from the alteration core) has also been documented. Jack (1989) determined that chlorites in the footwall alteration zone are Mg-rich, (Fe/(Fe+Mg) = 0.14–0.28), while chlorites in the footwall andesites outside the stringer zone were Fe-rich, (Fe/(Fe+Mg) = 0.44–0.46).

The majority of hangingwall chlorites have similar chemistries and are classified as ferro-clinochroes. The chlorites outside the hangingwall alteration plume are remarkably similar with Mg/Mg + Fe values ranging from ≈63 to 70. Chlorites from within the fuchsite-carbonate plume have a greater range of Mg/Mg + Fe values, from ≈42 to 81. The low Mg/Mg + Fe chlorites form a distinctive group and are from the most intensely fuchsite-carbonate altered core sample immediately above the ore body. These low Mg/Mg + Fe chlorites are magnesio-chamosites, previously identified as Fe-chlorites. A small group of chlorites have lower Al and higher Mg/Mg + Fe and also contain significant amounts of Cr, up to 1.91 wt % Cr₂O₃. The chromium content of chlorites is significant and is higher than the chromium contents of white micas. There is no relationship between chromium content and distance from the deposit, with high Cr-bearing chlorites occurring both above and distal to the deposit, although the highest Cr values are from chlorites within the hangingwall alteration plume.

**White micas**
Microprobe analyses have indicated that there is not as much white mica in the hangingwall basalt as indicated from core logging. Low K in whole rock geochemical analyses and absence of AlO₃ peak in PIMA spectra for many samples confirms this suggestion. Although it is therefore difficult to make meaningful comparisons of white mica chemistry down holes and between holes, the following...
observations can be made: (i) no observable trends in chemistry relating to distance from ore deposit, (ii) relatively low Cr content in white micas from rocks which visually appear to be strongly fuchsite altered. Maximum Cr$_2$O$_3$ content analysed was 0.28%, (iii) majority of white micas analysed were phengitic (Si:Al of 3:1 with Fe, Mg substitution for Al). No relationship between amount of phengite and distance from ore deposit, (iv) up to 1.20% BaO in white micas and Ba content of micas decreases with distance away from the orebody, (v) Fe + Mg content vs Al in the octahedral site in white mica has a limited range near the orebody but erratic values way from the deposit and (vi) white micas in MAC19 indicate a mixing trend with paragonite.

**Carbonates**

The carbonates analysed in the Hellyer hangingwall alteration are calcites, ferroan dolomites and ankerites. The dolomites and ankerites containing the greatest amounts of Mg and Fe are found further within the zone of intense fuchsite-carbonate and carry up to ≈6 wt % MnCO$_3$. Most of the carbonates outside the hangingwall alteration plume are calcite. In the hangingwall the composition of hydrothermal carbonates shows a systematic change of increasing Mn at roughly similar Fe+Mg (although Mg > Fe) as the orebody is approached.

Figure 1. Schematic cross section of the Hellyer deposit showing the geologic setting of the deposit. Abbreviations: SEZ-stringer envelope zone (quartz-sericite-pyrite alteration), Stringer zone - intense footwall alteration, from core to rim; quartz→chlorite→sericite. From Gemmell and Large (1993).
Bibliography


---

Figure 2. Schematic model of the lithogeochemical halo model and vectors to ore for the footwall alteration pipe, footwall lithologies within the district, Hangingwall Volcaniclastic Sequence (HVS), Hellyer Basalt and Que River Shale. Data from Gemmell (1990), Gemmell and Large (1992; 1993), Jack (1989), Sinclair (1994), Green and Taheri (1992) and this study.
Mt Julia-Henty gold mine: Summary of alteration study

Tim Callaghan
Goldfields Tasmania Ltd.

Halo model

Alteration can be divided into three zones: footwall alteration, the main mineralised zone (A-Zone) and hangingwall alteration.

The footwall consists of strongly foliated, mylonitic sericite-carbonate ± pyrite altered rhyolitic lavas, intrusives and volcaniclastics with rare fuchsite-sericite altered mafic dykes.

A-Zone alteration is composed of well zoned alteration facies with both gradational and sharp boundaries. The outer zone consists of sericite-pyrite-silica-chlorite (MZ), to an intermediate zone of sericite-silica (MZ to an intensely altered silica-carbonate zone (MQ)). There are lenses of massive carbonate contained within A-Zone, particularly towards the stratigraphic top of the zone.

Hangingwall alteration consists of intense albite, albite-carbonate, albite-chlorite and albite-chlorite-carbonate alteration. The carbonates are the same as those found within A-Zone and probably predate the mineralising event.

Vectors to ore

Distal halos and vectors
- Albite-silica, albite-silica-chlorite, albite-silica-chlorite-carbonate alteration of hangingwall (lithology dependant).
- Sericite-pyrite-carbonate alteration of footwall and A-Zone margins.
- Immobile element lithogeochemistry to identify favourable stratigraphy.
- Bedded carbonates as stratigraphic markers?
- Na₂O depletion and K₂O enrichment of the footwall.
- Na₂O enrichment and K₂O depletion of the hangingwall.

Proximal vectors
- Alteration zonation from sericite-chlorite-pyrite to sericite-silica to silica. Carbonate is ubiquitous but bedded carbonates generally occur near the stratigraphic top.
- Depletion of Al₂O₃, Y, Nb, Sc and V.
- Addition of Th (Mt Julia only).
- A-Zone metal accumulation halos.
- Broad distal As halo.
- Localised distal Zn halo
- Positive Bi-Cu-Pb correlation with Au with prominent halos surrounding local high Au zones.
- Strong Au-Ag correlation.

Bibliography


Figure 1. Mt Julia section 53350N
Figure 3. Schematic section of Henty post mineralisation but pre-deformation, demonstrating whole rock and immobile element vectors

- Lynchford member
- Mt Julia Rhyolite
- Mt Julia Member
- Silica-sericite-pyrite -chlorite
- Silica-sericite
- Silica-carbonate
- Albite-silica
- Massive sulphides
- Bedded carbonates
- Albitised carbonates

Decreasing Al₂O₃, SiO₂, Y, K₂O, V
Increasing Th, SiO₂

SiO₂, Na₂O + MgO enrichment

K₂O enrichment, Na₂O depletion
Alteration halo model for the Zone 96 volcanogenic gold deposit, Henty gold mine, western Tasmania

Jason Beckton
Goldfields Exploration, Kalgoorlie

Alteration mineralogy and zonation

The stratigraphy of the alteration system has been described in detail in the previous Amira report. In summary the system is characterised by asymmetry.

The stratigraphic footwall is moderately altered with feldspar destruction incomplete and volcaniclastic textures still recognisable. The footwall alteration is labelled MA (moderately altered) and is highly foliated. It is in contact with the Henty fault to the West and the MQ/MZ position to the East.

The main ore position comprises silica-sulphide-gold (MQ) enveloped by silica-sericite (MV) and sericite-silica-sulphide (MZ) alteration. MV alteration occurs predominately on the stratigraphic hangingwall side of the MQ and occurs along strike. In thin section the matrix of the MV resembles the MQ, however it is dominated by a sericite overprint.

MZ alteration texturally in many places closely resembles epiclastic shale seen in the stratigraphic hangingwall. It is volumetrically the most significant alteration assemblage.

Carbonate lenses occur predominantly stratigraphically above the MQ but also below. These units are interpreted from meso and micro scale to have overprinted epiclastic shales. This alteration is probably predating the main silica-sulphide and later sericite forming events. Along strike away from the main alteration zone, chlorite-calcite is the dominant assemblage of this alteration unit.

The hangingwall of the deposit has in many places been Na enriched. Albite-sericite alteration (AS) occurs in the stratigraphic hangingwall and appears to have replaced a coherent lithology such a rhyolite sill. This alteration lithology is thickest (up to 20m) directly above the thickest zones of the MQ alteration.

Halo geochemistry

A series of cross sections, Level Plans and Long Projections through the middle of the orebody illustrate Au, Ag, As, Cu, Pb, Zn, Bi and total Fe zonations.
Significant negative correlation can be seen between Au and Zn. Maximum Zn concentrations occur in the hanging and footwall positions as opposed to the main MQ/Au zone which occurs within the centre of the alteration zone.

**Lithogeochemical halo model and vectors to ore**

Henty may be loosely associated with the Cu-Au VHMS classification (e.g. Mt Lyell). Geochemical exploration for this deposit type must factor in the negative correlation of Zn. Cu, Fe and Pb have a less distinct negative correlation and are probably less useful as halo elements.

Tracing and extrapolation of structural features may be possible with As.

**Henty system genesis**

High grade gold is associated with the dominant cleavage formed during this deformation however on a deposit scale gold is largely stratabound within the MQ alteration. From structural relations noted underground (Beckton 1997), the MQ is interpreted to have formed prior to deformation.

Lithogeochemical Halo features noted above are largely thought to be the result of pre-deformation alteration.

**Bibliography**


Alteration zonation and geochemical dispersion at the Western Tharsis deposit, Mt Lyell, Tasmania: a summary

by David Huston and Julianne Kamprad
Australian Geological Survey Organisation

Summary

The Western Tharsis deposit occurs within a sequence of dominantly felsic volcaniclastic rocks with lesser clastic and coherent intermediate volcanic rocks in the Central Volcanic Complex. This sequence is overlain by a thin lens of Tyndall quartz-phyrpic rhyolitic volcanics, which, in turn is overlain by the Owen Conglomerate.

Lithogeochemical, mineral chemistry and PIMA analyses define vectors to ore that can be used in deposit and prospect scale exploration for Western Tharsis-type mineral deposits. As the studies were limited to within 500 m of the ore position, no regional or district-scale vectors were established.

Between 500 to 200 m from ore, rocks surrounding the Western Tharsis deposit are characterised by albicite rocks that have been moderately to strongly quartz-chlorite-sericite or chlorite-sericite altered (Fig. 1). This zone is characterised by abundant carbonate minerals that vary from ankeritic composition distal from ore to sideritic compositions more proximal to ore. Increasing proximity to ore is also characterised by a decrease in Mg/(Mg+Fe) ratios of chlorite. This carbonate-rich zone is characterised by positive C-Zn-Tl-Mn-Ca anomalousism (Fig. 2).

Within 200 m of ore, host rocks to the deposit have been strongly quartz-sericite altered, with the Fe-bearing carbonate minerals supplanted by pyrite. Apatite is an accessory mineral in this and more proximal alteration zones at Western Tharsis. This pyritic zone is characterised by low level Cu, As, Bi, Mo, Ni, S and Se anomaly and by depletion of the Zn-Tl-C-Mn-Ca assemblage. Positive Ba anomaly extends somewhat further than 200 m from the ore position. Although chlorite is not present in this zone, decreasing phenigite content of sericite is a good indication of approach to ore. This is also reflected in PIMA spectra, where λ_{MOH} decreases towards ore.

Within 70 m of ore, the orebodies are flanked by zones of extreme depletion in K and Cs. This anomaly is probably related to the development of acid-sulfate alteration assemblages dominated by pyrophyllite, but also including topaz, zonyte, fluorite and woodhouseite. These acid-sulphate assemblages are strongly associated with a zone of bornite-dominant mineralisation that overprints the dominant pyrite-chalcopyrite ore assemblage at Western Tharsis.

The alteration assemblage of pyrophyllite-topaz-zonyte-woodhouseite, which is associated with bornite-dominant mineralisation, is characteristic of acid-sulphate epithermal systems elsewhere in the world. Moreover, the close relationship of this alteration assemblage to the more prevalent quartz-sericite and quartz-chlorite-sericite alteration assemblages associated with disseminated pyrite-chalcopyrite mineralisation suggests that these two types of mineralisation are two stages of one mineralising event, not two separate events as accepted presently. Geological relationships and radiogenic isotope data are consistent with a Delamarian (~460 Ma) timing for Mt Lyell Cu-Au mineral deposits; this mineralisation may be related to the geologic event that formed the Haulage Unconformity. These observations potentially indicate a new exploration model for mineralisation in the Mt Read Volcanic belt and in surrounding Ordovician rocks.
Figure 1. Phyllosilicate alteration assemblages, section 8850N, Western Tharsis deposit.
Figure 2. Fe-S-O-C alteration assemblages, section 8850N, Western Tharsis deposit.
Bibliography


Alteration halo of the Thalanga VHMS deposit, north Queensland

Holger Paulick
Centre for Ore Deposit Research

The Thalanga volcanic-hosted massive sulphide deposit is located close to the western end of the E-W striking, Cambro-Ordovician Mount Windsor Subprovince in north Queensland approximately 180 km inland from Townsville. It has been deformed during regional metamorphism under upper greenschist facies conditions (biotite grade). The initial geological reserve was ~ 6 Mt of Zn-Pb-Cu-Ag-Au mineralisation in sheet-style, semi-connected ore lenses (Gregory et. al., 1990).

The Thalanga mineralisation is hosted within a felsic succession of rhyolite and dacite which were emplaced in a below wave-base, sub-aqueous environment (Paulick, 1997). The deposit is located at the stratigraphic contact of regionally extensive rhyolite in the footwall (Mount Windsor Formation), and a dacite dominated volcano-sedimentary succession in the hangingwall (Trooper Creek Formation). This stratigraphic position is referred to as the Favourable Horizon.

The aims of this study were to unravel the volcanic facies architecture of the host rock sequence, characterise the textural, mineralogical, and geochemical effects of hydrothermal alteration and to define geochemical proximity indicators to Thalanga-style massive-sulphide mineralisation. The results of this research can be summarised as follows.

Volcanic facies architecture and chemostratigraphy

The footwall rhyolite and hangingwall dacite can be divided into several groups based on textural observations and immobile element ratios. In the footwall, four different types of rhyolite occur which can be distinguished by consistent differences in phenocryst content. Rhyolite with abundant, coarse quartz and feldspar phenocrysts (type 4, also referred to as 'Quartz-Eye Porphyry') can be divided further into three sub-groups based on Ti/Zr and Ti/Th ratios, whereas rhyolite type 1, 2, and 3 have very similar immobile element ratios. In the hangingwall, consistent variation in feldspar phenocryst abundance and size range can be used to define 3 types of dacite. Dacite type 1 and 2 have lower immobile element ratios than dacite type 3. To the west, dacite type 2 and type 3 are the principal lithofacies in the hangingwall whereas dacite type 1 is dominant in east Thalanga.

Alteration

The principal alteration types in the footwall are: mottled chlorite-sericite ± pyrite alteration, intense quartz(-sericite)-pyrite and chlorite-pyrite alteration, late quartz ± sphalerite ± K-feldspar alteration and carbonate-calc-silicate (epidote-zoisite-tremolite) alteration. Massive to semi-massive assemblages of carbonate-chlorite-tremolite ('CCT') commonly occur within or close to the Favourable Horizon in west Thalanga. Texturally destructive quartz(-sercite)-pyrite and chlorite-pyrite alteration occur in distinctive zones within the footwall and in a stratabound zone (upto 50 m wide) parallel to the Favourable Horizon (Fig. 1).

Alteration of the hangingwall dacite is rarely texturally destructive and common alteration minerals are disseminated chlorite (-biotite) and epidote (± actinolite). A prominent patchy to vein controlled 'red rock' alteration type occurs frequently.
Fig. 1: Geology and alteration zonation in section 31380 E (east Thalanga) through the Thalanga sequence. Diamond drill holes shown have been examined for this study. The geometry of the ore bodies is based on mapping of underground exposure and logging of underground production drilling by Thalanga mine staff.
and is inferred to be caused by disseminated, sub-
microscopic hematite microcrysts. Geochemical data
of dacite indicate that a large proportion of the
hangingwall dacite experienced Na-metasomatism
(albite alteration, Paulick, 1997).

The geochemical changes associated with different
footwall alteration types were investigated by mass
change calculation using a single precursor model
(MacLean, 1990). A model for the Thalanga hydro-
thermal system shows the spatial relationship of
footwall alteration types and their principal
geochemical characteristics (Fig. 2).

In summary it can be concluded that:
• the elements Mg, S, Ba, Cu, Pb, Zn, As, Bi, Mo, and
Rb were added to the footwall rhyolite during
hydrothermal alteration,
• Na was lost, and
• the elements Si, Ca, and Sr appear to have been
redistributed within the footwall between zones
of different alteration types.

Halo geochemistry

Whole rock geochemistry
In the footwall rhyolite, several elements and indices
were found to show consistent trends with regard to
proximity to ore and may be useful vectors to
Thalanga-style mineralisation on a district or mine
scale (Fig. 3).

Geochemical proximity indicators to the Thalanga
mineralisation on a district scale are:
• elevated sulphur which is directly correlated with
the pyrite content of the rocks,
• sodium depletion,
• increase in alteration index and carbonate-chlorite-
pyrite index,
• elevated or depleted Ba depending on alteration
mineralogy,
• elevated Pb and Zn,
• elevated Mo, Bi and As (not always for mottled
chlorite-sericite± pyrite alteration), and
• increase in Rb/Sr ratio.

Geochemical proximity indicators to the Thalanga
mineralisation on a mine scale are:
• strongly elevated As, Bi, and Mo,
• elevated Tl, and
• locally strong Ba enrichment.

Mineral chemistry
The composition of chlorite and biotite changes
systematically with proximity to ore in samples from
a thick zone of mottled chlorite-sericite± pyrite
alteration below the mineralisation in west Thalanga
(section 20080E). A gradual increase from X_Mg-ratios
of 0.4 - 0.5 in least altered rhyolite to 0.85 - 0.95 just
below the mineralisation can be observed (Fig. 4).
Fluorine content of biotite is high close to the
mineralisation (1.7 F per formula unit [pfu]; Fig. 3).

Bibliography

Gregory, P. W., Hartley, J.S. & Wills, K.J.A., 1990,
Thalanga zinc-lead-copper-silver deposit, in
Hughes, F. E., ed., Geology of the mineral
deposits of Australia and Papua-New Guinea:
Melbourne, AIMM, p. 1527-1537.

MacLean, W. H., 1990, Mass change calculations in
altered rock series: Mineralium Deposita, v. 25,
p. 44-49.

architecture and alteration styles of felsic and
mafic volcanic host sequences to massive
sulphide deposits - Thalanga, northern
Queensland and Teutonic Bore, Western
Australia. CODES: AMIRA/ARC Project P439,

Paulick, H., 1997. Volcanic facies analysis, alteration
and geochemistry of the host rock sequence to
VHMS-style mineralisation at Thalanga (north
Queensland). CODES: AMIRA/ARC Project
P439, Report 4: 185-222.

Paulick, H., 1998. Alteration halo of the Thalanga
VHMS deposit, north Queensland. CODES
AMIRA/ARC P439, Final report.
Fig. 2: Model for the Thalanga hydrothermal system outlining the general spatial relationships between alteration types and associated geochemical mass changes.

Fig. 3: Summary of geochemical proximity indicators to the Thalanga mineralisation (pfu: per formula unit).
Fig. 4: Relationship between magnesium and iron content of (a) chlorite and (b) biotite in altered and least altered Thalanga footwall rhyolite (data: microprobe analyses of chlorite and biotite in least altered rhyolite from the railway cutting and variably altered rhyolite from DDH TH247, west Thalanga).
Alteration halo model for the Highway–Reward sub-seafloor replacement deposit, Mount Windsor Subprovince, Queensland

Mark G. Doyle  
Centre for Ore Deposit Research

Location
35 km south of Charters Towers in north Queensland

Resource
Highway
Primary — 1.2 m.t. @ 5.5% Cu, 6.5 g/t Ag, 1.2 g/t Au  
Oxide — 0.07 m.t. @ 6.04 g/t Au

Reward
Primary — 0.2 m.t. @ 3.5 Cu, 13 g/t Ag, 1 g/t Au  
Supergene — 0.3 m.t. @ 11.6% Cu, 21 g/t Ag, 1.8 g/t Au  
Oxide — 0.1 m.t. @ 33 g/t Ag, 6.49 g/t Au

Volcanic belt
Mt Windsor Subprovince (Highway Member, Trooper Creek Formation)

Age of volcanism
Cambro-Ordovician

Facies associations
Hosted by a rhyolitic to dacitic intrusion-dominated sequence, that includes sedimentary facies, turbiditic sandstone and pumiceous and crystal-rich sandstone-brecia. Contact relationships and phenocryst mineralogy, size and percentages indicate the presence of more than thirteen distinct porphyritic units in a volume of 1x1x0.5 km (Doyle, 1995, 1997a,b). The peperitic upper margins to most porphyritic units demonstrate their intrusion into wet unconsolidated sediment. Syn-sedimentary sills, cryptodomes and a partly extrusive cryptdome have been recognised (Fig. 1). These represent a proximal facies association from intrabasinal, non-explosive, syn-sedimentary intrusion-dominated magmatism. The Highway and Reward pyrite-chalcopyrite pipes occur within, but close to, the steep margins of intrusions in the host succession (Fig. 1A; Doyle, 1997a,b).

Ore composition
Dominantly pyrite Cu-Au type

Mineralisation types
The mineralisation can be divided into five principal types on the basis of mineralogy, textures and the relationships to the host rocks. These are: (1) discordant pyrite-chalcopyrite pipes; (2) veins, disseminations, discordant bands and strata-bound lenses of pyrite-sphalerite-galena-barite; (3) footwall quartz-pyrite stringer veins; (4) hanging wall quartz-pyrite stringer veins; and (5) gossanous breccia.

Alteration
The Highway and Reward orebodies occur within a well-developed discordant alteration envelope. The envelop extends from at least 150 m below the orebodies to over 60 m above the Highway pipe (Doyle, 1997a,b; Doyle 1998b; Fig. 1B). Broadly the alteration envelope has a mineralogical zoning which is defined by assemblages of sericite, chlorite-sericite, chlorite-sericite ± quartz, chloritohyanidrite, quartz-sericite-sericite, feldspar-sericite-chlorite-quartz and hematite ± chlorite-sericite-quartz-feldspar. A quartz-sericite ± pyrite zone is centred beneath the pyrite pipes and on some sections extends into the hanging wall succession. Small domains of intense chlorite ± anhydrite alteration occur within the footwall quartz-sericite ± pyrite zone and along some margins of the pipes. Quartz-sericite-sericite alteration gives way laterally and vertically to domains of sericite-chlorite ± quartz and chlorite-sericite alteration. Beyond the hydrothermal alteration halo, rocks of rhyolitic to dacitic composition contain various assemblages of
Figure 1. Simplified geological cross section showing the distribution of (A) lithofacies and (B) alteration at 10150N. The position of the massive sulfide bodies is also shown. Dacite D1, rhyodacites RD1-RD3 and rhyolites R2-R4 and R6 are exposed on section 10150mN (after Doyle, 1997b).
feldspar (albite, K-feldspar), calcite, sericite, chlorite, quartz and hematite (Fig. 1B).

Halo geochemistry

The Highway-Reward alteration envelope corresponds to a zone of anomalous Cu and Zn (Fig. 2). Depletion in Na2O and CaO to values < 1%, extends to over 100m below the pyrite pipes and 40 m above the pyrite pipe (Doyle, 1997c). Quartz-sericite-pyrite-altered samples are characterised by elevated K2O (1-6 wt%), coupled with depletion in MgO (0.2-6 wt%). The Ishikawa Alteration Index (AI >65), Chlorite-carbonate-pyrite index (CCPI=35-99) and box plots are an effective means of distinguishing the different alteration types. Ratios of Rb/Sr are anomalously high (> 1) in the alteration halo. S/Na,O ratios typically increase from values of <2 in background samples, to over 95 for intensely altered samples in the footwall and hanging wall to mineralisation. Also generally useful are ratios of Sr/Ba. The Sr/Ba ratio decreases systematically with increasing AI. Al(OH) wavelength values for muscovite in the Highway-Reward alteration envelope are anomalously low (2202 to 2222) in comparison with feldspar-sericite-chlorite and hematite-rich samples (2195 to 2212) (Doyle, 1998b; Herrmann et al, 1998).

Genetic considerations

The available evidence suggests that the pyrite-chalcopyrite pipes and strata-bound lenses formed together and are syn-genetic, sub-seafloor replacements of the host sediment, syn-volcanic intrusions, partly extrusive cryptodomes and volcaniclastic units (Doyle, 1995; Doyle, 1997a,b). Most of the massive sulfide ores at Highway-Reward formed by sub-seafloor replacement of rhyolite to dacite and volcaniclastic units because: (1) the mineralisation is hosted by intrusions or mass-flow emplaced units; (2) discordant and strata-bound ores contain relic patches of coherent facies or precursor volcanic particles; (3) peperite and massive sulfides are not mixed, implying sulfide deposition postdated emplacement of the enclosing succession; (4) pyrite pipes are discordant to bedding; (5) there are replacement fronts passing from strata-bound sphalerite-rich ores into discordant pyrite-pipes; (6) zones of strong quartz-sericite-alteration and pyrite veining extend into the hanging wall without any abrupt break in intensity (Doyle, 1997b). The distance below the seafloor at which infiltration and replacement took place is difficult to interpret, but was probably at least 60 m.

The location, distribution, form and shape of massive sulfide mineralisation and alteration are closely related to inferred initial patterns of permeability in the host rocks. The lavas and intrusions focussed hydrothermal fluids along the discrete mineralising pathways. Others formed an impermeable barrier promoting sub-seafloor ponding of hydrothermal fluids and replacement of the glassy and fractured margins of syn-sedimentary intrusions and lavas (Doyle, 1997a,b).

Conclusions

Highway-Reward provides a rare example of a pipe-style, sub-seafloor replacement VHMS deposit hosted by a syn-sedimentary intrusion-dominated volcanic centre. The geochemical halo vectors identified at Highway-Reward will have application to exploration for Cu-Au rich deposits elsewhere in the Mount Windsor Subprovince and in comparable volcanic successions.

Bibliography


Geological exploration criteria

- calco-alkaline felsic-intermediate volcanic succession
- hosted in the proximal facies association of non-explosive submarine (turbid) volcanic centre dominated by nph-sedimentary intrusions or lavas
- fluids focussed along fractured glassy margins of coalescing intrusions/lavas
- ponding of fluids beneath barrier (e.g., crystalline lava)
- structural control on magnetism and fluid flow
- syn-genetic sub-seafloor replacement of intrusions, lavas and volcaniclastic units
- stacked system
- halo of Pb-Zn-Ba mineralisation surrounds Cu-Au pipes
- potential for Pb-Zn lenses distal to pipes
- strong sericite-quartz-pyrite alteration in footwall and hanging wall
- alteration halo extends 10's m into hanging wall
- late structures focussed along alteration system

Geochemical exploration criteria

- footwall and hanging wall alteration halo
  - depletion Na_2O, CuO, MgO ± MnO
  - enrichment Si, B, Na_2O, F, Sr/Sm ± FeO, K_2O, MnO
  - low Sr/Ba (<0.1)
  - high fashkaw alteration index (>50)
  - REE generally higher (4-8) in HW alteration than FW (1-4.5), Values anomalous relative to background (<0.9)
  - AOH wavelength: 2195-2212 (generally <2200)
  - anomalous Cu/Pb, Zn, Au, Ag, As and Mo
  - 7.8% values in sulfides (pyrite,chalcopyrite; sphalerite) 5.0-8.7
  - Zn/Pb, Zn/Ba ratio in ore around 18.073 (±0.028)

Geophysical exploration criteria

- ground EM, IP and down hole EM will define targets
- ground EM response strong for Cu-rich massive sulfides
- pyrite pipes give good gravity response
- pyrite in FW and HW alteration zone may give significant IP response
- gravity and magnetics useful in mapping anodesites

Ore fluid conditions

- pyrite-chalcopyrite pipes
  - high temperature (>300 °C)
  - low pH (2 to 4.5)
  - high 1O_2
  - alteration halite-carbonate mineralisation
  - low temperature (<200 °C)
  - slightly more alkaline
  - oxidised

Data from Large (1991), Dean and Carr (1992), Huston (1992), Doyle (1997; 1998) and the present study

Figure 2. Exploration model: Highway-Reward type Cu-Au-rich pipes and Pb-Zn lenses.
Alteration case study of the Gossan Hill VHMS deposit

Robina Sharpe and Bruce Gemmell
Centre for Ore Deposit Studies

Gossan Hill is an Archean (2.8-3.0 Ga) Cu-Zn massive sulphide (VHMS) deposit located in the Yilgarn Block of Western Australia. The deposit is situated in the Murchison Province granite-greenstone terrane, 500 km NNE of Perth. Mineralisation at Gossan Hill and the neighboring Scuddles deposit occur at the same stratigraphic interval within the Golden Grove domain on the northeast flank of the Warrieldar Fold Belt. The Golden Grove Domain is a coherent succession of felsic to intermediate volcaniastics, lavas and breccias, as well as minor chemical "exhalite" horizons. Footwall to the mineralisation are felsic to intermediate volcaniastics while the hangingwall is dominated by coherent and brecciated dacite and rhyolite lavas. Distinguishing features which make Gossan Hill unique for an Australian VHMS deposit include; i.e. a host package of volcaniastics (100–400 m thick), ii. magnetite alteration and massive magnetite associated with Curich zones, iii. Au-rich massive sphalerite and iv. stratigraphic separation of zinc and copper-rich zones (ie stacked system). The inferred underground resource at Gossan Hill is 7.7 m.t. at 3.7% Cu and 2.4 m.t. at 12% Zn. In addition, gold and copper supergene zones overly the zinc and copperorebodies respectively.

Halo model

Hangingwall alteration
Alteration mineralogy: carbonate, sericite, quartz
Depletion strong, ubiquitous
Na₂O,
Depletion <200m from GGF contact
Sr, Ba, (Rb), Cs, SiO₂
Enrichment <100 m from GGF contact
FeO, MgO, MnO, As, Sb, CaO, K₂O, Ag

Regional alteration
Alteration mineralogy: quartz, chlorite
Depletion strong, ubiquitous
Na₂O K₂O, CaO
Enrichment strong, ubiquitous
SiO₂, FeO, ±MgO
Enrichment weak anomalismMnO

Local siliceous alteration
Alteration mineralogy: quartz, ± chlorite,
± carbonate, ± sphalerite, ± pyrite
Enrichment sporadic distribution
Cd, As, Pb, Fe, Zn, Ag, Bi, Sb, Sn
Enrichment strong to intenseSiO₂, (± FeO, MgO, CaO, MnO)
Local chlorite alteration
Alteration mineralogy: chlorite, ± carbonate,
± pyrite, ± magnetite ± chloritoid,
± apatite, ± andalusite, ± pyrrhotite,
± talc, ± rutile, ± ilmenite

Enrichment: weak anomaly TiO₂, Al₂O₃, ±Zn,
± Pb, Mo, ± Bi, ± Sb

Enrichment: pervasive strong to intense MnO,
± CaO, Cu

Enrichment: pervasive intense FeO, MgO

Vectors to ore

Alteration of the host and footwall rocks at Gossan Hill is intense and has resulted in intense geochemical modification of the tuffaceous volcanioclastic succession. Geochemistry therefore, reflects the proportional variation of a simple alteration mineralogy. An early regional siliceous alteration resulted in extensive metasomatism and effectively sealed the host and footwall rocks to later hydrothermal and metamorphic modification. As a result of impermeable wall rocks, hydrothermal alteration associated with mineralisation at Gossan Hill is constrained to the ore-depositional environment. This local alteration is characterised by:

- increasing SiO₂, FeO and MgO associated with intense chlorite alteration (proximal vector: metres);
- depletion in Na₂O and enrichment in K₂O and CaO in hangingwall volcanics (<200 metres);

Geological vectors to mineralisation include:

- M1 Marker hydrothermal chert-lithic horizons;
- discontinuity of the Golden Grove Formation due to discordant synvolcanic intrusions which reflect hydrothermal feeder sites;
- thickness variations in the Golden Grove Formation across synvolcanic intrusions, with mineralisation correlated to thickening of the volcanioclastic strata;

- magnetic anomalies.

Figure 1. Schematic model of the lithogeochemical halo model and vectors to ore within the Gossan Hill mineralising system.
Figure 2: Level plan 10150 R.L. (200 metres below surface) of the central and northern area of mineralisation at Gossan Hill showing the distribution of alteration zones. Outline of the 250 level development through the central parts of mineralisation is indicated.
Mineral chemistry

• chlorite: generally Fe-rich with footwall chlorite having a higher Mg/(Mg+Fe) ratio (0.47) than chlorites higher in the stratigraphy associated with Zn-rich mineralisation (Mg/Mg+Fe=0.28).

• carbonate: ankerite to siderite throughout alteration and mineralisation, with carbonate in Zn-rich sulphide mineralisation tending to have higher MrCO$_3$ contents (up to 16.5 wt.%).

Bibliography


Carbonate alteration at the Rosebery mine: The relationships between alteration texture, paragenesis, chemistry of carbonate minerals, and distance to ore

Rodney L. Allen, Michael Blake and Ross R. Large
Centre for Ore Deposit Research

Carbonate alteration is an integral part of the hydrothermal alteration system at the Rosebery massive sulphide deposit. A major focus of the AMIRA alteration study at Rosebery has been to systematically document and understand this carbonate alteration. Data was generated from: (1) the graphic geological logging of textural features, distribution and overprinting relationships of carbonate alteration in nine drill cores from the northern A-B and K lens area (Fig. 1), (2) detailed petrographic study of 69 samples from a proximal drill hole through K lens (120R) and a distal drill hole 500 m north of K and A-B lenses (109R), and (3) microprobe analysis of carbonate mineral chemistry at 240 points in these samples, representing the various textural types and paragenetic stages of carbonates alteration.

This study defines 12 main carbonate textural types at Rosebery (Table 1), and 20 paragenetic generations of alteration out of which nine generations have a major carbonate component. The carbonates show a considerable range in chemical and mineralogical composition. They can be classified as calcite, a spectrum of compositions between rhodochrosite [MnCO₃] and iron-rich rhodochrosite [MnFe(CO₃)₂], locally extending to manganoo siderite [(Fe> Mn)CO₃], and a spectrum of compositions between ankerite [Ca₂FeMg(CO₃)₄] and kuznachovite [CaMn(CO₃)₂] (Table 1, and Large et al., volume 2).

Table 2 summarises all the major variations and trends in the data set. This table lists representative samples and representative analyses of the various carbonate textural types and parageneses. This data

Figure 1: Long section of Rosebery Mine showing drill holes sampled in this investigation in relation to the ore lenses (modified from Graves et al., 1997).
is arranged according to proximal or distal position relative to ore (120R and 109R data respectively) and according to stratigraphic level.

The nodular or spotty carbonate alteration types are a distinctive feature of the Rosebery deposit. These spotty carbonates (1–5 in Table 1), and the massive and platey carbonate types, are the earliest generations of carbonates. They formed soon after deposition of the host pumice breccia and vitric sandstone-siltstone facies and are spatially and temporally associated with massive sulphide mineralisation. They occur only in the spotty carbonate and sericite-carbonate alteration zones (Fig. 2), within 20 m of the stratigraphic level of an ore lens and within 100 m lateral distance from the edge of an ore lens. Consequently, they are an excellent guide to ore. Subsequent, syn- and post-cleavage generations of carbonates do not show a preferred spatial association with the massive sulphide lenses.

In the spotty carbonate alteration zone, the earliest generation of carbonate spheroids are manganiferous but are very variable in composition, even within individual spheroids. Subsequent generations of spheroids and early veins have less variation in composition. This indicates that fluctuations in fluid chemistry and/or environmental conditions during early concretionary carbonate growth, resulted in great chemical and mineralogical complexity right down to the grain and hand specimen scales. This initial complexity is interpreted to have been partially homogenised by subsequent generations of carbonate growth.

Within 10–50 m of massive sulphide mineralisation, and especially in the footwall, where Mn-Fe anomalies were set up in early generations of alteration, subsequent overprinting generations perpetuate the anomalous Mn-rich character, but as noted above the compositions tend to be more homogenised. Thus although the early generations of carbonate alteration are most diagnostic of proximity to ore, sampling of later generations of carbonates in the same rocks would generally return a similar chemical signature, i.e. recognition of the textural type and paragenetic generation of carbonate is desirable, but does not preclude the use of carbonate mineral chemistry as an exploration guide.

Where early alteration was not anomalous in Mn and Fe, subsequent generations of carbonate alteration were mainly calcitic. Important exceptions are the late generations of veins (paragenesis 17, 19 and 20), which include Fe ± Mn-rich compositions even in areas with no evidence of previous FeMn-bearing alteration. This emphasises that the presence of Mn-Fe carbonates does not alone indicate close proximity to massive sulphide. The presence of early, pre-cleavage generations of MnFe-bearing carbonates, and Mn>Fe composition, are additional criteria that characterise carbonates proximal to ore at Rosebery.

The late generations of carbonate veins appear to show a large scale zonation from both Mn>Fe and Fe>Mn compositions within 100 m of ore, to only Fe>Mn compositions distal to ore. The data set in this study is not large enough and the sampling was not extended sufficiently distal from ore to be certain of this large scale zonation, and whether the MnFe contents of these late paragenetic alterations reflect partial redistribution of the early formed MnFe anomaly associated with massive sulphide mineralisation, or whether they include the influx of new MnFe unrelated to the earlier mineralisation. This should be resolved to determine whether the Mn/Fe ratio of carbonates associated with isolated distal veins can be used as a regional exploration guide, and for mine scale exploration more than 100 m from known ore.

Early carbonate alteration in the hangingwall more than 10–20 m stratigraphically above the level of massive sulphide mineralisation comprises calcitic impregnations with very low Mn and Fe contents. Apart from along the Mount Black Fault, the only FeMn-bearing carbonates recorded in the hangingwall are a late- to post-cleavage generation (paragenesis 19) of weakly manganiferous ankerite veins with less than 4 mole% MnCO₃ and Fe>Mn. These veins have distinctive bleached sericite>carbonate alteration haloes, and their wide distribution suggests they are not related to the Rosebery deposit.
Fig 2. Rosebery cross section 1700 mN alteration distribution
<table>
<thead>
<tr>
<th>Texture</th>
<th>Variations</th>
<th>Internal structure</th>
<th>Composition</th>
<th>Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>1-6. Nodular or spotty</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1. Large nodules</td>
<td>(1 cm - 2 m diameter nodules)</td>
<td>- Dispersed</td>
<td>- Massive granular, or coarse concentric layering</td>
<td>Hercules 4-level road</td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Integrown</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2. Spheroidal</td>
<td>(0.2 mm - 1 cm diameter spheroids of anhedral grains)</td>
<td>- Dispersed, distinct spheroids - Close-packed, intergrown</td>
<td>- Distinct fine concentric layering - Faint concentric layering - No visible concentric layering</td>
<td>Rosebery 120R 1361m, Rosebery 120R 1357m, Rosebery 120R 1364B</td>
</tr>
<tr>
<td>3. Spheroidal-rhombic</td>
<td>(0.5 mm - 1 cm diameter spheroids composed entirely of, or with rim of radiating rhombs)</td>
<td>as above - No concentric layering - coarse concentric layering</td>
<td>not analysed</td>
<td>Hercules South Dunnes shaft</td>
</tr>
<tr>
<td>4. Rhombic</td>
<td>(0.2 mm - 1 cm rhombs)</td>
<td>- Dispersed</td>
<td>- Concentric layering</td>
<td>not analysed</td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Random intergrowth</td>
<td>- No concentric layering</td>
<td></td>
</tr>
<tr>
<td>5. Lozenge</td>
<td>(0.2 mm - 1 cm lozenge-shaped grains)</td>
<td>- Dispersed</td>
<td>- Concentric layering</td>
<td>(Mn&gt;Fe)CO₃</td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Close-packed, intergrown</td>
<td>- No concentric layering</td>
<td></td>
</tr>
<tr>
<td>6. Feldspar pseudomorph</td>
<td>(replacement of 0.5-4 mm feldspar phenocrysts)</td>
<td>- Porphyritic distribution</td>
<td>Patchy-irregular to massive Ca(Mn&gt;Fe=Mg)(CO₃)₂(Ca&gt;Mn)CO₃</td>
<td>Rosebery 120R 1399m, Rosebery 120R 1147m</td>
</tr>
<tr>
<td>7. Platey (rare)</td>
<td>(5 mm - 3 cm tabular laths or plates ?after anhydrite, gypsum)</td>
<td>- Dispersed</td>
<td>Massive</td>
<td>not analysed</td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Interlocking network</td>
<td>- No distinct spherical texture or layering</td>
<td></td>
</tr>
<tr>
<td>8. Blebbby</td>
<td>(&lt; 10 cm irregular patches)</td>
<td>- Dispersed</td>
<td>Massive</td>
<td>(Mn&gt;Fe&gt;Mg)CO₃</td>
</tr>
<tr>
<td></td>
<td></td>
<td>- Interconnected</td>
<td>- No distinct spherical texture or layering</td>
<td>Ca(Mn&gt;Fe=Mg)(CO₃)₂(Fe&gt;Mn)CO₃</td>
</tr>
<tr>
<td>9. Massive</td>
<td>(irregular compact granular masses 5 cm - 2 m)</td>
<td>- Anhedral grains - Close-packed rhombs - Close-packed spheroids with carbonate-filled interstices</td>
<td>- Concentric layered grains - No layering in grains</td>
<td>Ca(Mn&gt;Mg&gt;Fe)(CO₃)₂</td>
</tr>
<tr>
<td>10. Impregnation</td>
<td>(filling or replacement of matrix within non-carbonate rock)</td>
<td>- Irregular patches - Pervasive</td>
<td>Anhedral, non-layered grains</td>
<td>CaCO₃ (Ca&gt;Mn)CO₃</td>
</tr>
<tr>
<td>11. Veins</td>
<td>(carbonate ± quartz veins and their alteration haloes)</td>
<td>- early pre-S₂ cleavage - pre- to syn-S₂ cleavage - syn-S₂ - post-S₂</td>
<td>- subspherical - massive to banded - massive to banded - massive</td>
<td>Ca(Mn&gt;Fe=Mg)(CO₃)₂, As above CaCO₃ (Mn,Fe)CO₃, CaCO₃</td>
</tr>
<tr>
<td>12. Limestone</td>
<td>(fine grained, compact, calcitic carbonate)</td>
<td>- Layer, bed - Clasts in mass flow beds</td>
<td>Massive to foliated - Fossiliferous (trilobites)</td>
<td>CaCO₃</td>
</tr>
<tr>
<td>Stratigraphic position</td>
<td>Texture</td>
<td>Paragenesis</td>
<td>Sample numbers</td>
<td>MnCO₃ mole%</td>
</tr>
<tr>
<td>------------------------</td>
<td>----------------</td>
<td>-------------</td>
<td>----------------</td>
<td>------------</td>
</tr>
<tr>
<td>PROXIMAL STRATIGRAPHIC TRANSECT INCLUDING ORE LENS (120R)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hangingwall Veins</td>
<td>syn-cleav (16)</td>
<td>1083.A3.2</td>
<td>3.3</td>
<td>0.9</td>
</tr>
<tr>
<td></td>
<td>1094.A4.1</td>
<td>1.9</td>
<td>0.4</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>1100.A1.2</td>
<td>1.7</td>
<td>0.4</td>
<td>0</td>
</tr>
<tr>
<td>Matrix impregnation</td>
<td>pre- to syn-cleavage (10-16)</td>
<td>1083.A2.1</td>
<td>2.2</td>
<td>0.7</td>
</tr>
<tr>
<td></td>
<td>1100.A4.2</td>
<td>1.3</td>
<td>0.4</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>1169.A3.4</td>
<td>2.1</td>
<td>1.6</td>
<td>0.4</td>
</tr>
<tr>
<td>Alteration within feldspar phenocryst</td>
<td>pre- to syn-cleavage (7-16)</td>
<td>1147.A1.1</td>
<td>3.0</td>
<td>1.3</td>
</tr>
<tr>
<td></td>
<td>1169.A1.2</td>
<td>2.4</td>
<td>1.6</td>
<td>0</td>
</tr>
<tr>
<td>Black shale Veins</td>
<td>syn-cleav (16)</td>
<td>1184.A2.1</td>
<td>1.3</td>
<td>1.8</td>
</tr>
<tr>
<td></td>
<td>1229.A1.2</td>
<td>0.7</td>
<td>1.4</td>
<td>1.4</td>
</tr>
<tr>
<td></td>
<td>1256.A1.1</td>
<td>0.9</td>
<td>0.4</td>
<td>0.5</td>
</tr>
<tr>
<td>TSV Veins</td>
<td>syn-cleav (16)</td>
<td>1237.R4.1</td>
<td>0.5</td>
<td>0.5</td>
</tr>
<tr>
<td></td>
<td>1241.A1.2</td>
<td>0.7</td>
<td>1.7</td>
<td>0.8</td>
</tr>
<tr>
<td></td>
<td>1265.A2.1</td>
<td>0.7</td>
<td>0.6</td>
<td>0.3</td>
</tr>
<tr>
<td>Matrix impregnation</td>
<td>pre- to syn-cleavage (10)</td>
<td>1237.A2.2</td>
<td>0.5</td>
<td>0</td>
</tr>
<tr>
<td></td>
<td>1265.A1.3</td>
<td>0.5</td>
<td>0.4</td>
<td>0</td>
</tr>
<tr>
<td>Alteration within feldspar phenocryst</td>
<td>pre- to syn-cleavage (7-16)</td>
<td>1241.A3.1</td>
<td>0.5</td>
<td>0.8</td>
</tr>
<tr>
<td>Hangingwall sill: top</td>
<td>Alteration within feldspar phenocryst</td>
<td>1291.A2.1</td>
<td>0.9</td>
<td>0.4</td>
</tr>
<tr>
<td></td>
<td>syn-cleav (16)</td>
<td>1315.A3.1</td>
<td>2.3</td>
<td>1.0</td>
</tr>
<tr>
<td>Hangingwall sill: base</td>
<td>Alteration within feldspar phenocryst</td>
<td>1353.A2.1</td>
<td>8.1</td>
<td>1.6</td>
</tr>
<tr>
<td></td>
<td>syn-cleav (10-15)</td>
<td>1353.A3.1</td>
<td>7.8</td>
<td>1.5</td>
</tr>
<tr>
<td>Carbonate alteration at top of ore zone</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Late veins in spotty carbonate</td>
<td>pre- to syn-cleavage (7-16)</td>
<td>1357.A1.1</td>
<td>34.0</td>
<td>7.7</td>
</tr>
<tr>
<td></td>
<td>1356.A1.3</td>
<td>55.8</td>
<td>27.2</td>
<td>13.6</td>
</tr>
<tr>
<td></td>
<td>1356.A1.6</td>
<td>36.6</td>
<td>10.4</td>
<td>8.3</td>
</tr>
<tr>
<td></td>
<td>1356.A6.1</td>
<td>41.3</td>
<td>3.6</td>
<td>3.8</td>
</tr>
<tr>
<td></td>
<td>1356.B5.2</td>
<td>36.8</td>
<td>9.7</td>
<td>7.8</td>
</tr>
<tr>
<td>Early sub-spheroidal veins in spotty carbonate</td>
<td>early pre-cleavage (6)</td>
<td>1357.A3.1</td>
<td>33.7</td>
<td>7.7</td>
</tr>
<tr>
<td></td>
<td>1361.A1.1</td>
<td>39.0</td>
<td>5.7</td>
<td>5.5</td>
</tr>
<tr>
<td></td>
<td>1364.B7.1</td>
<td>41.8</td>
<td>2.8</td>
<td>4.4</td>
</tr>
<tr>
<td></td>
<td>1365.A2.1</td>
<td>44.8</td>
<td>4.0</td>
<td>4.7</td>
</tr>
<tr>
<td>Massive</td>
<td>pre-cleavage (6)</td>
<td>1365.B2.1</td>
<td>27.4</td>
<td>3.3</td>
</tr>
<tr>
<td>Spotty-lozenge</td>
<td>pre-cleavage (6)</td>
<td>1364.A6.1</td>
<td>91.2</td>
<td>5.1</td>
</tr>
<tr>
<td>Spotty-spheroidal, faint concentric layering</td>
<td>pre-cleavage (2-6)</td>
<td>1357.A2.1</td>
<td>34.2</td>
<td>7.0</td>
</tr>
<tr>
<td></td>
<td>1364.A1.1</td>
<td>90.6</td>
<td>7.0</td>
<td>2.7</td>
</tr>
<tr>
<td></td>
<td>1364.B8.1</td>
<td>89.9</td>
<td>1.4</td>
<td>1.6</td>
</tr>
<tr>
<td>Spotty-spheroidal, strong fine concentric layering</td>
<td>pre-cleavage (2)</td>
<td>1361.A2.2</td>
<td>87.1</td>
<td>7.4</td>
</tr>
<tr>
<td></td>
<td>1361.A2.1</td>
<td>7.7</td>
<td>11.6</td>
<td>25.9</td>
</tr>
<tr>
<td></td>
<td>1361.A2.4</td>
<td>16.3</td>
<td>9.0</td>
<td>22.3</td>
</tr>
<tr>
<td>Ore</td>
<td>Blebs in remobilized ore</td>
<td>pre- to syn-cleavage (6-7)</td>
<td>1366.A3.1</td>
<td>71.2</td>
</tr>
<tr>
<td></td>
<td>1366.A3.2</td>
<td>54.6</td>
<td>30.9</td>
<td>13.2</td>
</tr>
<tr>
<td>Footwall</td>
<td>Veins, and halo</td>
<td>post-cleav (20)</td>
<td>1436.A3.1</td>
<td>70.5</td>
</tr>
<tr>
<td>------------</td>
<td>----------------</td>
<td>----------------</td>
<td>-----------</td>
<td>------</td>
</tr>
<tr>
<td>Blebs (replacing feldspar crystals)</td>
<td>pre- to syn-cleav (6-10)</td>
<td>1399.A2.1</td>
<td>28.3</td>
<td>10.2</td>
</tr>
<tr>
<td>Alteration within feldspar phenocryst</td>
<td>pre- to syn-cleav (7-16)</td>
<td>1421.A1.2</td>
<td>3.9</td>
<td>30.1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1421.A5.2</td>
<td>3.0</td>
<td>20.5</td>
</tr>
<tr>
<td></td>
<td></td>
<td>1440.A3.1</td>
<td>2.8</td>
<td>1.7</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>DISTAL TRANSECT 500 m FROM ORE (109R)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hangingwall</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Black shale</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>TSV</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td>Footwall</td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
<tr>
<td></td>
</tr>
</tbody>
</table>

**Bibliography**


A molar element ratio analysis of lithogeochemical samples from the footwall andesite, Hellyer VHMS district, Tasmania, Australia

Clifford R. Stanley and J. Bruce Gemmell

MDRU, The University of British Columbia, and Centre for Ore Deposit Research

A Pearce element ratio (PER) analysis of the Hellyer footwall alteration system was undertaken to evaluate the application of this approach to understanding the alteration process and developing vectors to ore. Because of the superior resolution offered by the PER approach, several new concepts regarding hydrothermal alteration in the footwall andesites at Hellyer have been revealed. These include:

- Zr, Nb and Y are more conserved (more incompatible and immobile) than TiO₂;
- primary compositional variations within the andesite can be attributed to fractionation of plagioclase and clinopyroxene in an approximately 8/7 molar ratio, which corresponds to an 33/20 volume ratio — this ratio is crudely consistent with the phenocryst abundances observed within the andesite;
- late calcite veins do not appear to be significant contributors to the compositional variability within the andesites;
- the composition of fractionating clinopyroxene and plagioclase are estimated to be $X_{CI} = 5/9$ and $X_{AN} = 1/6$ — this clinopyroxene composition is Fe-rich relative to the clinopyroxene phenocrysts observed in the Hellyer andesites — this plagioclase composition is relatively albic, and thus is not considered to be primary, but is likely a product of spilitic Na-Ca exchange between seawater and devitrifying andesite;
- metasomatism associated with muscovite alteration involved Na, Ca and Sr loss, K, OH, Rb and Ba addition, and did not involve significant Al or Si metasomatism — the alteration assemblage associated with muscovite alteration is muscovite plus quartz;
- pyrite (and trace chalcopyrite, galena and sphalerite) precipitation probably filled pore produced by this muscovite alteration, producing the classic QSP (quartz-sericite-pyrite) alteration assemblage observed in many VHMS camps — this reaction involved the addition of S, Fe, Cu, Pb and Zn;
- chlorite alteration of muscovite involved the loss of K, Rb and Ba, and the addition of Fe, Mg and OH — direct precipitation of chlorite from solution involved the addition of Fe, Mg, Si, Al and OH — both of these reactions involve significant volume increases, and so were restricted to proximal locations about the Jack fault, where tectonically formed open space was available to facilitate reaction progress;
- re-sericitization of chlorite occurred in the core of the alteration zone, and involved a significant volume loss coupled with the addition of K and loss of Fe, Mg and OH — it also appears to be associated with Si addition (in contrast to simple muscovite alteration, above), accommodated via quartz precipitation within the pores produced — it also is associated with S, Cu, Pb and Zn addition, as sulphides also precipitated in the new pores;
- muscovite alteration of feldspar and chlorite alteration of muscovite represent simple metasomatic reactions which buffered the hydrothermal fluid via exchange of elements between fluid and rock — these reactions were likely driven by chemical/mineralogical dis-equilibrium between the host rocks and hydrothermal fluid; and
- direct chlorite and pyrite precipitation reactions represent non-metasomatic reactions, and these were likely triggered by changes in the physico-
chemical state of the hydrothermal fluid — as such, these reactions were driven by physico-
chemical disequilibrium between the hydro-
thermal fluid and its environment.

In addition, five exploration parameters have been identified that could be used to vector toward
the most intense regions of hydrothermal alteration. These have different advantages and
disadvantages, and include:

- $(2\text{Ca}+\text{Na})/\text{Al}$, which describes the degree of
hydrolysis, cannot discriminate between muscovite
and chlorite alteration ($> 1 = \text{fresh}; 0 = \text{muscovite}
or chlorite altered) — it also suffers from a non-
unique representation of the background compositional variation due to plagioclase and clinopyroxene fractionation;

- $(2\text{Ca}+\text{Na}+\text{K})/\text{Al}$, which also describes the degree
of hydrolysis (Fig. 1), can discriminate between muscovite and chlorite alteration ($> 1 = \text{fresh}; 1/3 = \text{muscovite altered}; 0 = \text{chlorite altered}) — it also suffers from a non-unique representation of the plagioclase and clinopyroxene fractionation;

- $(2\text{Ca}+3\text{Na}+\text{K})/\text{Si}$, which also describes the degree of hydrolysis (Fig. 2), is also affected by silificationation — however, this parameter has the advantage of producing a unique fresh rock signature ($1 = \text{fresh}; 1/3 = \text{muscovite altered}; 0 = \text{chlorite altered};$ silification decreases these values);

- $(2\text{Ca}+\text{Na}+\text{K})/(\text{Al}+2\text{Fe}+2\text{Mg}-\text{S})$, which also describes the degree of hydrolysis, is also affected by Fe and Mg metasomatism — however, this metric also has the advantage of producing a unique fresh rock signature ($1 = \text{fresh}; 1/3 = \text{muscovite altered}; 0 = \text{chlorite altered};$ Fe and Mg metasomatic additions decrease these values); and

- K/Al, which describes the degree of muscovite alteration, cannot discriminate between fresh and chloritized andesite ($0 = \text{fresh or chloritized}; 1/3 = \text{muscovite altered}) — it also has a unique fresh rock signature.

Although each of these five exploration parameters responds to hydrothermal alteration effects within the visible alteration zone, three $[(2\text{Ca}+\text{Na})/\text{Al}$,
$(2\text{Ca}+\text{Na}+\text{K})/\text{Al}$ and $(2\text{Ca}+3\text{Na}+\text{K})/\text{Si}$] identify anomalies outside the alteration zone just below the contact between the footwall andesite and hangingwall basalt. As such, they do define any cryptic alteration halo that may be indicative of proximal accumulations of VHMS mineralization. These parameters are also more quantitative
measures of hydrothermal alteration than previously available. Furthermore, used in concert,
they allow recognition and discrimination of different styles of hydrothermal alteration, and
thus may provide substantial benefit where complex alteration zoning exists.

Bibliography


Figure 1 - Helleyer andesite footwall samples are plotted in bubbleplot form on an east-west cross-section at 10,750 m N. Bubble sizes are proportional to \(1-(2\text{Ca}+\text{Na}+\text{K})/\text{Al}\), one of the metrics identified as a possible exploration parameter (the ratio is subtracted from unity to ensure that large bubble sizes correspond to samples that are strongly altered). This metric distinguishes between muscovite and chlorite alteration, giving a quantitative measure of hydrolysis. Anomalous samples occur within the recognized alteration zone, and outside the alteration zone near the contact between the footwall andesite and hangingwall basalt.

Figure 2 - Helleyer andesite footwall samples are plotted in bubbleplot form on an east-west cross-section at 10,750 m N. Bubble sizes are proportional to \(1-(2\text{Ca}+3\text{Na}+\text{K})/\text{Si}\), one of the metrics identified as a possible exploration parameter (the ratio is subtracted from unity to ensure that large bubble sizes correspond to samples that are strongly altered). This metric distinguishes muscovite from chlorite alteration, and also responds to silicification. Anomalous samples occur exclusively within the recognized alteration zone.
Chlorite alteration associated with syn-volcanic granites and Cu-Au mineralisation: A pilot study along the Jukes Road

Bill Wyman
Centre for Ore Deposit Research

Hydrothermal alteration mineral assemblages containing chlorite form some of the most easily identified and mapped alteration mineral assemblages surrounding the granite related Cu-Au mineralisation at the Jukes Prospect. The Jukes Road forms an excellent cross-section through several of these chlorite bearing alteration mineral assemblages and it therefore offers an excellent opportunity to examine potential variations in chlorite mineral chemistry as a function of distance. Chlorite analyses were conducted using the Electron Microprobe at the University of Tasmania Central Science Laboratory (CSL). Five different habits of chlorite were probed as part of this study. The five habits are: (1) chlorite replacing biotite (7?) phenocrysts (some could be amphiboles). (2) chlorite in chlorite-magnetite ± sulfide veins, (3) chlorite replacing feldspar phenocrysts, (4) interstitial (intergranular) chlorite between feldspar and quartz grains, and (5) chlorite in the matrices of mineralised hydrothermal breccias.

Microprobe data was examined in relation to distance from mineralisation and also in relation to each of the five habits of chlorite. Chlorite alteration typically occurs with weak to moderate sericite alteration and occasionally with weak to intense K-feldspar alteration. Chlorite alteration is almost always accompanied by magnetite in amounts directly proportional to the degree of chlorite alteration. This study examined the mineral chemistry of chlorite within the various habits. A comparison is made between chlorites within a single relatively homogeneous feldspar-phryic rhyolite lava and chlorites within a cross-cutting quartz-feldspar ± biotite porphyry.

Figure 1 is a plot of the Mg number (Mg# = Mg\(_{\text{total}}\) / (Fe\(_{\text{total}}\) + Fe\(_{\text{total}}\) + Mg\(_{\text{total}}\)) vs distance for the five habits of chlorite in both the feldspar-phryic rhyolite and the quartz-feldspar ± biotite porphyry. Data from the chlorite microprobe analysis indicate that for both the feldspar-phryic rhyolite lava and the quartz-feldspar ± biotite porphyry:

1. There is a large variability in Mg number (Mg# = Mg\(_{\text{total}}\) / (Fe\(_{\text{total}}\) + Fe\(_{\text{total}}\) + Mg\(_{\text{total}}\)) within individual samples for a given chlorite type.
2. The range of Mg#'s from chlorite replacing feldspar phenocrysts, interstitial chlorite and chlorite in veins is the same and overlaps the range of Mg#'s from chlorite in the cores of the hydrothermal breccias.
3. There is a weakly defined decrease in the Mg# within the cores of the mineralised breccias that probably reflects an increase in iron associated with an increase in mineralisation.
4. Chlorites from within the quartz-feldspar ± biotite porphyry dykes overlap the same ranges in Mg# values as those within the feldspar-phryic rhyolite lava.
5. The Mg# of whole rocks corresponding to the probed samples, show the same range of Mg# values outside the mineralised zone. Near the boundaries of, and within the mineralised zone the Mg# of the host rocks is less than the Mg# of chlorite indicating increased iron. This supports the conclusion that magnesium in the mineralised zone is contained within chlorite and not within other mineral phases.
6. As a result of the overlap in the ranges of chlorite values within a given chlorite habit and the overlap in the ranges between various chlorite types, when
Figure 1: Plot of the Mg number ($\text{Mg\#} = \frac{\text{Mg}_{\text{total}}}{\text{Fe}_{\text{total}} + \text{Mg}_{\text{total}}}$) vs distance for the five habits of chlorite in both the feldspar-phryic rhyolite and the quartz-feldspar +/- biotite porphyry.

Figure 2: Plot of the Mg# of the various chlorite habits vs. the Mg# of the whole rock sample. Chlorite in veins and breccias were not plotted as they were excluded from the whole rock analysis.
plotted against distance, chlorite composition is not a useful vector toward mineralisation.

Based on the Al⁴⁺ content an attempt to calculate chlorite formation temperatures was made for each main type of chlorite. Table 1 summarises the calculated chlorite formation temperatures based on the Al⁴⁺ geothermometer. The determined range of average chlorite temperature values is relatively narrow (328°-362°C), and the overall range is only slightly broader (312°-386°C). Several distinct separations in the temperature data can be observed. First, the average chlorite temperature data within the quartz-feldspar ± biotite porphyry dykes (356°-358°C) is almost identical regardless of chlorite habit. Second, the average chlorite temperatures of the chlorite veins, interstitial chlorite and the breccia matrix chlorite within the rhyolite (360°-362°C) are very close to the temperatures shown in the quartz-feldspar ± biotite porphyry dykes. Third, the chlorite temperatures in the chlorite that replaced both biotite and feldspar phenocrysts within the feldspar-phyric rhyolite are almost identical (328° and 331°C) and are significantly lower than the chlorite temperatures within the quartz-feldspar ± biotite porphyry dykes. The 320°-360°C range would be consistent with temperatures expected from the later lower green-schist facies regional metamorphic event. However, equilibration of calculated chlorite temperatures and compositions would also be expected. Figure 2 is a plot of the Mg# of the various chlorite habits vs. the Mg# of the whole rock sample. Chlorite in veins and breccias was not plotted as that chlorite was excluded from the whole rock analysis. Since two clear populations of temperature exist and the data shows scatter on a plot of chlorite Mg# vs whole rock Mg#, the calculated temperatures are interpreted to represent relict hydrothermal chlorite formation temperatures.

Bibliography


Table 1. Summary of calculated chlorite formation temperatures based on the Al⁴⁺ geothermometer.

<table>
<thead>
<tr>
<th>Chlorite type</th>
<th>Feldspar-phyric rhyolite</th>
<th>Quartz-feldspar ± biotite porphyry dykes</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Biotite Phenocrysts</td>
<td>Feldspar Phenocrysts</td>
</tr>
<tr>
<td>Number of analyses</td>
<td>4</td>
<td>2</td>
</tr>
<tr>
<td>Average temp. °C</td>
<td>328</td>
<td>331</td>
</tr>
</tbody>
</table>
Use of immobile elements and chemostratigraphy to determine precursor volcanics

Walter Herrmann
Centre for Ore Deposit Studies

Summary

A number of elements, including the high field strength elements Ti, Zr, Nb, Y, Th; the heavy rare earths Lu & Yb and to a lesser degree the light rare earths; Al, Hf, Ta; and in some cases Sc, V and Cr, tend to be conserved in alteration zones associated with VHMS deposits. Although nett mass gains or losses of mobile components associated with hydrothermal alteration can result in changes of concentration of immobile elements, the inter element ratios of immobile elements will be invariable.

Immobility of element pairs should be tested, rather than assumed, by plotting data from a single precursor unit (which had primary compositional uniformity) on X-Y scatterplots. Immobile element pairs should define highly correlated linear trends which regress to the origin of the plot. (e.g. Fig. 1)

In practice, Ti and Zr have been found to be the most reliable immobile elements in VHMS systems. They can be inexpensively and accurately analysed by XRF on pressed powder pellets and they exist at easily detectable levels in most volcanics.

Some immobile elements are also incompatible* (e.g. Y and Zr), and can be interpreted to infer regional magmatic affinities, melt sources and tectonic settings.

Pairs of compatible-incompatible immobile elements (e.g. Ti and Zr) are most useful in prospect scale exploration, to identify unrecognisably altered rocks, map individual emplacement units through alteration zones, subdivide sequences by “chemostratigraphy” and infer petrogenetic relationships. Individual units from different magmatic sources or fractionated magmas may have distinctive immobile element ratios which enable their discrimination even in intensely altered or weathered zones. This is particularly so for coherent volcanic units of uniform primary composition, but is also applicable to some massive volcaniclastic units (e.g. rhyolitic pumice breccias).

Recognition of immobile elements is a prerequisite for calculating estimates of mobile component mass changes in altered rocks, relative to their least altered precursors, using the Isocon, Reconstituted Composition or Pearce Element Ratio methods.

Bibliography

Herrmann, W., 1998. Use of immobile elements and chemostratigraphy to determine precursor volcanics. CODES AMIRA/ARC P439, Final report.

* Incompatible elements are partitioned into the melt fraction during crystallisation of magmas; compatible elements are readily incorporated into early crystal phases.
Figure 1. Zr-TiO<sub>2</sub> scatterplot of 220 Thalanga volcanic samples showing separation of lithostratigraphic units and strongly correlated linear array of 134 samples of footwall rhyolite.
Development and use of the alteration box plot

Ross Large
Centre for Ore Deposit Research

The alteration box plot was introduced during the early phases of this research project (Large et al., 1996) to characterise alteration style and intensity related to VHMS systems (Fig. 1). This plot combines the power of two alteration indices; the Ishihawa alteration index (AI)* and the chlorite/carbonate/pyrite index CCPI**.

The Ishihawa alteration index measures the intensity of plagioclase and glass destruction during hydrothermal events, and the CCPI measures the importance of chlorite, pyrite and Fe–Mg–Mb carbonate alteration relative to sericite and K-feldspar alteration.

Combining the two indices in the ore plot enables a comparison of whole rock chemistry with alteration mineralogy, and leads to a classification of alteration trends and types related to VHMS ore systems (see Large et al., 1998; and Allen et al., 1998). Least altered volcanics plot in a box towards the central left of the diagram, hydrothermally altered volcanics plot toward the upper right triangle and diagenetic alteration plots in the lower left hand triangle.

Both mineral chemistry and whole-rock chemistry can be plotted on the diagram allowing a detailed analysis of their relationships. By analysing a suite of samples from a given prospect it is possible to evaluate the type of alteration (i.e. unaltered, diagenetically altered or hydrothermally altered) and to interpret their position relative to (a) the margin or (b) the centre of the hydrothermal system or (c) the ore horizon (Fig. 2).

It is recommended that the box plot be adopted as a basic lithogeochemical tool in the exploration for VHMS deposits.

Bibliography


---

* AI = 100(MgO + K₂O)/(MgO + K₂O + CaO + Na₂O)

** CCPI = 100(MgO + FeO)/(MgO + FeO + Na₂O + K₂O)
Figure 1. Alteration box plot showing the various alteration trends at the Thalanga VHMS deposit.

Figure 2. Typical alteration fields superimposed on the box plot for Thalanga. A different set of fields is required for each ore system studied (e.g. Rosebery, Hellyer, etc.).
Application of PIMA and FTIR spectrometry to VHMS alteration studies

Walter Herrmann, Michael Blake, Mark Doyle, David Huston and Julianne Kamprad
Centre for Ore Deposit Research and ASSO, Canberra

Summary

PIMA is a portable spectrometer that measures reflectance in the short wavelength infrared range. The PIMA's field portability and versatility in being able to rapidly and inexpensively spectrally analyse dry rock, core, drill cutting and soil samples, discriminate otherwise obscure clays, fine grained micas etc. and detect subtle compositional changes in some minerals (such as muscovite) opens up apparently wonderful opportunities to map and quantify hydrothermal alteration systems. Mapping the zonation of phyllosilicates, other silicates and carbonate phases, which are common products of alteration, is rarely easy, even by the practised eye, but could reveal important exploration vectors.

This study was designed to investigate the potential applications for PIMA spectrometry in interpreting hydrothermal alteration zones associated with Australian VHMS deposits.

Since most of these are deformed and metamorphosed to at least greenschist facies, the focus has been primarily on muscovite and chlorite; the early experience having shown that carbonates in alteration assemblages are not easily analysed by PIMA.

The study is based on a large number of PIMA spectra for samples from the alteration systems around the Rosebery, Western Tharsis and Highway-Reward deposits which represent contrasting styles of sulphide mineralisation hosted by felsic volcanics, in addition to spectra for background least altered rocks of the Mt Read Volcanics.

Rosebery and MRV regional traverses

Part of this study investigated the relationships between PIMA spectral characteristics, wholerock geochemistry and muscovite and chlorite compositions, in several hundred samples from regional traverses in the Mt Read Volcanics and drill holes around the northern end of the Rosebery massive sulphide deposit.

The results confirm that the wavelength of the PIMA spectral absorption feature attributed to Al-OH bonds, is systematically related to the composition of muscovite. Sodic muscovites (with partial substitution of Na for K or muscovite-paragonite mixtures?) have wavelengths below 2200 nm and phengitic muscovites (with >0.5 Fe+Mg cations substituting for octahedral Al) generally have wavelengths between 2210 and 2220 nm.

However, the composition of muscovites is not linked to alteration intensity. Phengitic muscovite from the Rosebery footwall alteration zone has a similar range of composition, and Al-OH wavelength, to muscovite in essentially unaltered rocks from elsewhere in the Mt Read Volcanics. Muscovite composition appears related to bulk rock composition in that the least phengitic, most sodic muscovites (with lowest wavelengths) exist in intermediate-mafic rocks with >10% chlorite and Ti/Zr 10–30. The hangingwall volcanioclastic sandstone unit at Rosebery contains muscovites of this type, in which the sodium content appears to be highest in proximity to sulphide lenses. Although there is probably no direct relationship to the VHMS forming hydrothermal system, this feature may have applications at the mine scale, as an exploration vector to ore lenses and to identify the favourable horizon.
requires more detailed PIMA observations on additional drill cores to establish its value as a vector.

Chlorites in the samples analysed, have considerable compositional variations that are not systematically related to the Rosebery footwall alteration zone, or alteration intensity elsewhere. In any case, PIMA spectral characteristics show no systematic relationship to chlorite compositions in these Mt Read Volcanics and Rosebery samples.

Attempts at using PIMA spectral characteristics to estimate amounts, or relative amounts, of muscovite and chlorite in Mt Read Volcanics have met with very limited success. The results are probably inferior to ordinary megascopic visual estimates. The method may produce better results in sets of samples that have compositional and spectral uniformity.

**Western Tharsis**

In contrast to Rosebery, muscovite compositions at Western Tharsis show a clear and symmetrical compositional relationship to the ore lens and alteration system in both the stratigraphic footwall and hangingwall. The composition grades from moderately phengitic (~0.5 Fe+Mg) in the outer quartz+chlorite+sericite±carbonate zones to non phengitic (<0.1 Fe+Mg) in the proximal pyrophyllite±quartz±sericite zone.

This compositional change is very closely correlated to the AIOH absorption wavelength and constitutes a fine-scale exploration vector which can be rapidly and effectively determined by PIMA analyses.

PIMA was instrumental in the recognition of several minor minerals — of major genetic importance — in the Western Tharsis alteration assemblage; they include pyrophyllite, topaz and zunyite.

Chlorites existing in the outer parts of the Western Tharsis alteration system show a weak trend of iron enrichment inwards towards the ore lens; (Mg numbers decreasing from ~45 to ~30). This compositional change is not measurable by PIMA analysis and the restriction of chlorite to the outer alteration zones limits its usefulness as an exploration vector.

**Highway–Reward**

PIMA AIOH feature wavelengths grade from intermediate to high (2200–2222 nm) in the outer zones of sericite+chlorite+feldspar alteration to significantly lower wavelengths (2195–2204 nm) in the proximal quartz+sameite+pyrite zones associated with pipe-like pyrite bodies. Although as yet unsupported by microprobe analyses, there is a strong inference that white micas in the proximal zones are non phengitic, and possibly contain some Na substitution for K. The similarity to the pattern at Western Tharsis suggests a link between Cu–Au systems and low phengite contents of muscovites in proximal alteration zones. The muscovite composition is presumably determined by physicochemical factors, perhaps related to the source of mineralising fluids.

**Comparison of FTIR and PIMA spectrometry**

FTIR and PIMA both provide qualitative information on white mica chemistry relating to the relative phengite (Fe+Mg) and paragonite (Na) composition. FTIR shows potential for measuring the Mg number variations in chlorite composition, in samples where the chlorite absorbance feature is not overshadowed by muscovite. In contrast, PIMA appears to be oblivious to chlorite chemistry, at least in mixed samples. Neither instrument has been proved to give information on carbonate chemistry in mixed assemblages, although FTIR has potential for investigation of other carbonate absorbance peaks in the low wavenumber region.

FTIR and PIMA use fundamentally different detection methods in that FTIR provides a measurement of sample transmission, whilst PIMA measures reflectivity. Because of this each may display different subtleties in particular wavelength regions, which may be complimentary.

**Bibliography**

Figure 1. Phengite and sodium content of muscovite compared to PIMA AIOH wavelength for fresh and altered volcanic from Rosebery, Hercules and MRV regional traverse samples.


Figure 2. Cross section 8850N, Western Tharsis, showing the relationship between PIMA AIOH feature wavelength and the ore lens.
P439 geochemical whole-rock and mineral database

Michael D. Blake
Centre for Ore Deposit Research

The analytical data collected during the duration of P439 has been collated and is provided on a PC format CD-ROM with this concluding report. Data is provided as Excel '95 files, each workbook containing spreadsheets with geochemistry, probe data and isotope data where available. Studies of alteration systems have been placed in a hierarchical file structure (visually represented in Fig1.), divided into two main categories - Mt Read Volcanics and Mt Windsor studies, with the Gossan Hill study independent.

The Mt Read Geochemical Database (MRChem) is a separate, relational database modelled on the Mineral Resources Tasmania (MRT) 'Rockchem' database. It contains data collected from the Tasmanian case studies, regional traverses, geochemistry donated by Pasminco, and data collated from University of Tasmania research Theses. Data provided by ISM at the beginning of the project is not included, but may be purchased from MRT with 25000 sheet GIS packages. MRChem is an ACCESS database, which has been broken down and included as CSV files for easy importation into any relational database or GIS package. The structure of MRChem is documented in Table 1., and is included on the CD as structure.doc. Definitions of fields contained in each data table are documented on the CD in the file definitions.doc.

Analytical suite

Standard major element analyses (SiO₂, TiO₂, Al₂O₃, Fe₂O₃, MnO, MgO, CaO, Na₂O, K₂O, P₂O₅, LOI, S), and trace elements Ba, Ce, Cr, Cu, La, Nb, Nd, Ni, Pb, Rb, Sc, Sr, V, Y, Zn, Zr were obtained by XRF for all samples at the University of Tasmania Geology department facility under the direction of Mr Phil Robinson.

Trace element analyses for Ag, As, Bi, Mo, Cd, Sb, Cs, Th and U were measured by ICPMS and total C by Leco at ANALAI&S facility in Perth.

Major element analyses are reported as wt% oxides with total Iron as Fe₂O₃. Trace elements are reported in ppm.

Previous work

Preliminary work was done by Large, Stolz and Duhig (1996), using the database to investigate alteration trends and to test the behaviour of alteration indices. Further work was done by Large and Blake (1997), using GIS data contouring methods to identify regional scale deposit characteristics and geochemical vectors to ore (Fig. 1).

Bibliography

Blake, M.D., 1997. P439 geochemical whole-rock and mineral databases. CODES AMIRA/ARC (439), Final report
Date, C., 1983. Database: A Primer, Addison Wesley, U.S.A.
P439, Report 2, 197-209
Large, R.R., and Blake, M.D., 1997. Surface lithochemical responses of known VHMS deposits based on the MRV geochemical database, AMIRA project P439, Report 4, 93-95

Figure 1. An example of the use of the geochemical data base from P499: Rb/Sr ratio plot of surface samples in the central MRV belt (from Large and Blake, 1997).
Discrimination of diagenetic, hydrothermal and metamorphic alteration

Rodney L. Allen, Cathryn C. Gifkins, Ross Large and Walter Herrmann
Centre for Ore Deposit Research

A requirement for the effective use of alteration studies in mineral exploration for hydrothermal ores is that the alteration associated with the mineralisation can be distinguished from diagenetic, metamorphic and tectonic alterations. In this contribution we outline methods and criteria for discriminating between these alteration types, and some examples.

Discrimination between diagenetic, hydrothermal and metamorphic alteration requires:
(1) A basic knowledge of the primary rock type, and the general textural evolution of volcanic rocks.
(2) Visible overprinting relationships at hand specimen and microscopic scale between alteration assemblages, secondary rock fabrics (such as compaction foliation and tectonic cleavage), and primary rock textures.
(3) Knowledge of the regional distribution and zonation of the alteration assemblages. Are they local or regional, and what is the zonation pattern?
(4) Knowledge of the temperature, pressure, and chemical environment under which key diagenetic, hydrothermal and metamorphic minerals are stable and unstable.
(5) Knowledge of the mineral transformations that occur during diagenesis, metamorphism and deformation, that have been documented in geologically young, well preserved regions.

Mineralogy and texture

Hydrothermal alteration is local in distribution, and commonly involves high water to rock ratios that facilitate conversion of volcanic glass and quartzofeldspathic primary mineralogy into clays, carbonate, sericite, chlorite and quartz. VHMS style alteration and mineralisation is synchronous with diagenetic alteration and pre-dates regional tectonic fabrics.

Diagenetic alteration is related to the regional geothermal gradient in the depositional basin. Temperatures range from near 0°C at the seafloor to 250°C at 2–10 km depth, the precise depths of particular isotherms depending on the geothermal gradient. The resulting diagenetic alteration pattern is generally a sequence of flat-lying zones (layers), each characterised by a particular mineral assemblage that formed within a particular temperature range. Diagenetic alteration zones are regional in distribution, and commonly involve lower water to rock ratios that facilitate conversion of volcanic glass and feldspars into smectite, zeolites, K-feldspar and albite.

As a general rule, alteration assemblages that are regional in distribution and pre-date or are synchronous with a stylolitic S1 compaction foliation can be defined as diagenetic. Alterations that show similar timing relationships but which are only local in distribution are related to discrete local hydrothermal alteration systems, or the cooling of particular volcanic emplacement units.

Alteration assemblages that are strongly foliated by regional cleavage and also overprint S1 compaction foliation are regarded as broadly syntectonic. These include regional weak sericitation associated with cleavage development, and local strong sericite...
Figure 1: Model for the evolution of, and relation between, hydrothermal alteration, diagenetic alteration, metamorphism and deformation at the Rosebery massive sulphide deposit. Post-volcanic, late to post-deformation, Devonian granite-related alteration is omitted.
4. After deformation, metamorphism, uplift and erosion

3. During deposition of hangingwall succession
and quartz alteration related to local high fluid flow zones such as faults and shears. Alteration assemblages unaffected or only very weakly affected by regional cleavage are post-main deformation in timing. Metamorphic alteration is regional in extent, post dates S1 compaction foliation, and is synchronous with or post dates regional cleavage. The boundary between diagenesis and low grade metamorphism is indistinct, but is commonly taken as the boundary between an albite-zeolite high grade diagenetic facies and a prehnite-pumpellyte or equivalent low grade metamorphic facies. The presence of zeolite in diagenetic mineral assemblages and the absence of zeolite from metamorphic assemblages is commonly the most distinct difference.

Combining data on overprinting relationships, the spatial distribution of alteration types, and mineral transformations in the Rosebery-Hercules area, allows construction of an evolutionary model for the development of diagenetic, hydrothermal and metamorphic alteration in this area (Fig. 1). The diagenetic alteration system was established during the earliest stages of hydrothermal alteration and initially preceded ore formation. The hydrothermal system then intensified, expanded and overprinted the uppermost diagenetic alteration zones. As hangingwall strata accumulated, regional isotherms and diagenetic alteration zones prograded up through the hangingwall, and a weak plume of alteration from the declining hydrothermal system formed in the hangingwall above the ore deposits. During metamorphism and deformation the diagenetic and hydrothermal mineral assemblages were recrystallised and variably replaced by metamorphic assemblages and only local relics of the earlier alteration minerals remain.

**Geochemistry**

Fresh volcanics in modern oceanic and continental margin arcs, and the least altered volcanics in ancient sequences which host massive sulphide deposits, have a fairly limited range of Na$_2$O contents (2–5%) and alteration indices* (AI = −15–65). Diagenetically and hydrothermally altered volcanics have greater compositional ranges from 0–8% Na$_2$O and 0–100 Al. This is largely attributable to albitisation in advanced diagenesis (resulting in high Na$_2$O and low Al) and the feldspar destructive sericite-chlorite-carbonate hydrothermal alteration in proximal zones associated with volcanic hosted massive sulphide deposits (resulting in low Na$_2$O and high Al).

The Na$_2$O and Al wholerock geochemical parameters permit a crude classification into least altered, diagenetically altered and hydrothermally altered volcanics. However, they do not adequately identify some early, less intense stages of diagenesis, or discriminate between sericite, chlorite and chlorite + carbonate dominated assemblages of hydrothermal alteration. A newly developed chlorite-carbonate-pyrite index (CCPI)**, used in conjunction with the AI index as a "boxplot", overcomes these deficiencies and enables simple graphic recognition of the likely major alteration phases and alteration styles.

Studies of a wide range of regional diagenetically altered and deposit associated hydrothermally altered volcanic rock samples, have shown that the AI-CCPI boxplot identifies ten alteration trends (Fig. 2). Six of these trends, lying near or above the sericite-calcite tie line on the boxplot, are related to assemblages formed in hydrothermal alteration systems. The remaining four, lying below the sericite-calcite tie line on the boxplot, are associated with texturally non destructive diagenetic alteration. Transitional zones (eg: the boundary between marginal hydrothermal and regional diagenetic) or complex overprinting (eg: diagenesis and subsequent metamorphism) should be distinguishable if geochemical data for a sufficient range of samples is available.

A combination of textural and mineralogical observations, and wholerock lithogeochemical data plotted in an AI-CCPI boxplot, enables simple discrimination of the various diagenetic and hydrothermal alteration styles in any suite of altered volcanics.

---

* AI = 100(MgO+K$_2$O)/(MgO+K$_2$O+CaO+Na$_2$O)
** CCPI = 100(MgO+FeO)/(MgO+FeO+Na$_2$O+K$_2$O)
Figure 2. Alteration trends on the alteration box plot.

**Hydrothermal trends:**
1. sericite alteration at margins of system;  
2. sericite-chlorite ± pyrite;  
3. chlorite ± sericite ± pyrite;  
4. chlorite-carbonate;  
5. Fe/Mn carbonate-sericite;  

**Diagenetic trends:**
7. albite ± chlorite;  
8. albite-calcite ± epidote;  
9. K-feldspar ± albite;  
10. paragonitic sericite ± albite.
Bibliography


Geochemical modelling of low temperature (5°C to 150°C) seawater–andesite interaction: Implications for regional alteration assemblages in VHMS districts

Stephen B. Bodon and David R. Cooke
Centre for Ore Deposit Research

Diagenetic alteration of felsic volcanioclastics hosting the Hercules and Rosebery deposits comprises early smectite-topal alteration followed by various poorly preserved zeolite stages, that are replaced by K-feldspar alteration assemblages and later albite alteration assemblages. The conditions under which K-metasomatism is progressively replaced by Na-metasomatism is not well understood, but is particularly important for gaining insights into the diagenetic processes that were occurring during and shortly after the accumulation of the volcanics. For these reasons, numerical modelling of seawater–andesite interaction was undertaken from 5°C to 150°C and rock–fluid ratios up to 7 g/l. The simulations assumed open system conditions, such that precipitated alteration minerals were not included in the subsequent iterations. This approach reflects natural systems more accurately and therefore, yields more realistic results.

Our numerical simulations predict that diagenetic alteration occurs under oxidised conditions, within the hematite stability field (Bodon & Cooke, 1998). Results indicate that secondary K-feldspar and quartz precipitate in conjunction with minor Na-rich saponite, dolomite, clinohlore, calcite, hematite, and trace muscovite (sericite) and watherite during the early stages of seawater–andesite interaction from 5°C to 75°C (Fig. 1). K-feldspar precipitation causes a progressive decrease in ΣK+ and increase in the Na+/K+ ratio in the evolving diagenetic pore water (modified seawater), coincident with decreasing pH. Low temperature K-metasomatism is predicted to result in enrichment of K and Mg, and depletion of Na, Ca and Si in the andesite.

From 75°C to limit of our heating simulations (150°C), Na-metasomatism occurs. Precipitates included albite–quartz and minor hematite, clinochlore, calcite and trace barite (Fig. 1). Albite precipitation causes a progressive decrease in ΣNa+ in the pore water coincident with decreasing pH and Na+/K+ ratios. Enrichment of Na and Mg, and depletion of Ca, Si and K are predicted to occur in the altered andesite. There is a narrow window at 75°C where K-feldspar and albite coexist in equilibrium. No zeolite minerals were precipitated during the calculations, possibly due to the low rock–water ratios used, and the concentration of ΣHCO₃⁻ in the fluid. Increasing concentrations of Cu, Pb and Zn throughout the reaction progress indicates that modified seawater has the capacity to leach metals from the andesite (Fig. 2).

In general, the predicted alteration mineral paragenesis (Fig. 1) coincides with observed diagenetic assemblages and paragenetic sequences in felsic volcanioclastics hosting the Hercules and Rosebery massive sulphide deposits. Overall, the whole-rock geochemistry predictions are close to actual variations measured in the felsic volcanics. Differences in the mineral assemblages are attributed to the assumed reaction pathway (i.e. temperature and water–rock ratio) used in the modelling, as well as the whole-rock geochemistry of the rock modelled (i.e. andesite vs rhyolite). Further simulations of various scenarios involving various rock types and reaction pathways are required to fully assess the spectrum of possible diagenetic alteration assemblages and their paragenetic relationships, as well as the associated chemical evolution of seawater and rock during diagenesis.

The ability of seawater to leach metals from the reacting rock, suggests that Zn–Cu–(Pb)-bearing fluids can be produced by diagenetic alteration of andesites. Regional diagenetic alteration of the volcanic pile most likely contributes to the overall
metal budget of mineralising solutions involved in the formation of massive sulphide deposits such as Hercules and Rosebery. In natural systems, metal concentrations and metal ratios in the mineralising solutions will depend on the initial metal contents of the host lithologies (i.e. the source rocks), and the ease by which each metal can be stripped (i.e. the relative reactivity of pyroxenes versus feldspars, etc.).

Bibliography


Figure 1. Summary of the calculated mineral paragenesis and mole% of minerals precipitated during seawater–andesite interaction as a function of the rock–water ratio and temperature (after Boden & Cooke, 1998). Minerals in italicised text constitute <1 mole%. “Rock (grams)” corresponds to grams of andesite added to 1 litre of seawater, i.e. the rock–water ratio (units are grams/litre). Reaction progress is from left to right.

Figure 2. Evolution of the calculated concentrations of total Cu²⁺, Pb⁺⁺ and Zn⁺⁺ components in seawater during seawater–andesite interaction as a function of reaction progress (after Boden & Cooke, 1998). “Rock (grams)” corresponds to grams of andesite added to 1 litre of seawater, i.e. the rock–water ratio (units are grams/litre). Reaction progress is from left to right.
Application of ironstone geochemistry to exploration for VHMS deposits in the Mt Windsor Volcanic Belt

Garry Davidson
Centre for Ore Deposit Research

This study evaluated the exploration vector potential of some iron silica deposits in the Cambro-Ordovician Mount Windsor Volcanic Belt, Queensland, complementing a similar study in the Highway-Reward area (Doyle 1997). "Ironstones", jaspilites, ferruginous cherts, barite lenses, and other styles of chemical sedimentation are generally used in exploration as stratigraphic markers for hydrothermal activity, however, their geochemistry suggests they can be far more useful than this.

Near-ore and "barren" ironstones were geochemically and texturally compared. The near-ore samples came from 3 ironstone lenses near Thalanga, very close to recently discovered base metal mineralisation, and from the Magpie area SE of Charters Towers. Barren ironstones were obtained from the Britannia and Sunrise Spur areas at the Trooper Creek Fm-Rollsion Range Fm contact. The approach was to build on previous reconnaissance work by Nathan Duhig and Joe Stolz (Duhig et al. 1992), by analysing members of these two ironstone groups in detail to determine their internal chemical variability. Numbers of samples varied from 3 to 19/ lens, depending on the size of the body. Sample spacing was 5 to 20 m across the strike of each stratiform ironstone; 20 m was determined to be the optimum spacing.

In all ironstones examined, regardless of scale, the dominant textural characteristic is early massive to inhomogeneous iron-silica deposition, overprinted by several generations of veining, in most cases becoming more silicic and metal-rich with time. However, the later generations of veining do not extend across the entire early ironstone, but tend to be more focussed around one or more smaller areas, creating more confined geochemical anomalies within the ironstone. The vein networks forming the ironstones are commonly fine and difficult to observe on uncut surfaces. This has given rise to the premise that ironstones are geochemically massive because at first sight they are texturally massive, but this is not the case. Crucially, apparently barren ironstones may contain a smaller "near-ore" signature zone, requiring very careful screening of ironstone geochemistry.

The ironstones can be genetically divided into three stages, which are not always present:

- **Stage 1** involved low temperature to microbial and inorganic precipitation of iron oxyhydroxides, and colloidal silica directly over zones of diffuse upflow.

- **Stage 2** involved selective overprinting of Stage 1 deposits by fine chert/quartz goethite networks. Geochemically, these zones were more silica-rich than Stage 1 products, also containing higher base metals, As, Mo, Ba, Se, Cd and Bi. Hydrothermal REE patterns were similar to Stage 1, with LREE enrichment and -Eu anomalies. Oxygen isotope measurements of quartz indicate that the temperature of deposition was between 85 and 190°C.

- **Stage 3** mainly characterised near-ore ironstones, and involved the formation of localised zones of barite-quartz veins, disseminated barite concentrations at % levels, elevated base-metals, and strong +Eu anomalies. The temperature of vein quartz deposition of this stage in the Thalanga and Magpie areas was between 127 and 277°C, on the basis of oxygen isotopes and an assumed broad range of potential initial water values. This unusual tendency of systems to "shrink" as they evolved to higher temperature may have reflected local heating cycles that were quickly exhausted, or
possibly was a symptom of the choking of general crustal permeability by mineral precipitation and its refocussing into one or more major channelways which formed massive sulfide deposits.

The following geochemical associations are correlated with increasing temperature in ironstones and are useful in separating near-ore from barren ironstones. Broadly this follows an evolution from Fe-Si, to Fe-Si-Ba, to Fe-Si-Ba---Eu-base metals, to Si-Ba-Eu-base metals: Fe:

**Stage 1:** \(\text{Fe}_2\text{O}_3(T) / (\text{Fe}_2\text{O}_3(T) + \text{SiO}_2) = 0.15 - 0.35\), U >0.2 ppm, \(\text{P}_2\text{O}_5 < 0.03\) wt. %, Cu + Pb + Zn <20 ppm (in weathered samples), Cu/Cu+Pb+Zn = 0.2 - 0.5, Zn/Cu+Pb+Zn = 0.2 - 0.45, Ba may be up to 800 ppm but is mainly <250 ppm, Ba/Cu+Pb+Zn variable from <1 to >50 (because the stage includes Ba-rich but base-metal poor ironstones towards higher temperatures), As < 10 ppm, Sb <4 - 6 ppm, Mo < 5 ppm, +Eu anomalies, strong -Ce anomalies.

**Stage 2:** \(\text{Fe}_2\text{O}_3(T) / (\text{Fe}_2\text{O}_3(T) + \text{SiO}_2) = 0.05 - 0.15\) but locally massive hematite around veins or coalesced veins, U <0.2 ppm, \(\text{P}_2\text{O}_5 > 0.03\) wt. %, Cu + Pb + Zn =20 - 200 ppm (in weathered samples), Cu/Cu+Pb+Zn = 0.2 - 0.5, Zn/Cu+Pb+Zn = 0.2 - 0.45, Ba may be up to ~2000 ppm, Ba/Cu+Pb+Zn >15, -Eu anomalies, -Ce anomalies, As =20 - 80 ppm, Sb up to 10 ppm, sporadic Mo up to 20 ppm; \(\text{La}^\text{iv}/\text{Sm}^\text{v} \text{iv}\) is a highly variable 2.95 - 18.31; \(\Sigma\text{REE}\) of 20 - 60, with some samples up to 150 ppm; \(\text{Na}_2\text{O}/\text{K}_2\text{O} <1\), \(\delta^{18}\text{O}\text{quartz} >14\%\).

**Stage 3:** \(\text{Fe}_2\text{O}_3(T) / (\text{Fe}_2\text{O}_3(T) + \text{SiO}_2) = <0.20\), mainly <10; Se >1 ppm, Cu + Pb + Zn >100 ppm, Cu/Cu+Pb+Zn = 0.0 - 0.9, Zn/Cu+Pb+Zn = 0.0 - 0.8. Ba >500 ppm, Ba/Cu+Pb+Zn <15 with the exception of barite lenses, MnO> As <30 ppm, \(\text{Na}_2\text{O}/\text{K}_2\text{O} <0.35 - 8.7, \delta^{18}\text{Osilicate} <14\%\), \(\text{La}^\text{iv}/\text{Sm}^\text{v} \text{iv}\) 2 - 7; \(\Sigma\text{REE} <20\) ppm although exceptions occur (see Duhig et al. 1992), +Eu anomalies, \(\text{P}_2\text{O}_5 >0.03\) wt. %, U>1 ppm, Mo <5 ppm.

In summarising this data, the key parameters for the recognition of populations of near-ore ironstones in the MWVB are (1) +Eu; (2) Ba >500 ppm; (3) Cu + Pb + Zn >100 ppm; (4) Se >1 ppm; (5) Ba/Cu + Pb + Zn <15; (6) \(\text{Fe}_2\text{O}_3(T) / (\text{Fe}_2\text{O}_3(T) + \text{SiO}_2) <0.20\); (7) a trend of increasing Mn in a given population. In analysing ironstones, care should be taken to use an ICP technique with very high resolution, because some elements of interest occur at the 0.01 - 0.1 ppm level.

**Bibliography**


### Table 1: A summary of the key geochemical features that have potential use as exploration vectors.

<table>
<thead>
<tr>
<th>Vector</th>
<th>Stage 1 ironstones</th>
<th>Stage 2 ironstones</th>
<th>Stage 3 ironstones</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mineralogy</td>
<td>Hematite-chert-quartz</td>
<td>Vein zones of hematite-goethite and quartz, with minor barite</td>
<td>Vein zones of quartz:hematite-barite-sulfides</td>
</tr>
</tbody>
</table>

**Key features**

<table>
<thead>
<tr>
<th>Element</th>
<th>Stage 1</th>
<th>Stage 2</th>
<th>Stage 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe oxide/Fe+Si oxides</td>
<td>0.15–0.35</td>
<td>0.05–0.15</td>
<td>&lt;0.20, but increasing toward mineralisation</td>
</tr>
<tr>
<td>Eu</td>
<td>negative (Eu/Eu&lt;sup&gt;+&lt;/sup&gt;*&lt;1)</td>
<td>negative (Eu/Eu&lt;sup&gt;+&lt;/sup&gt;*&lt;1)</td>
<td>positive (Eu/Eu&lt;sup&gt;+&lt;/sup&gt;*&gt;1)</td>
</tr>
<tr>
<td>Cu+Pb+Zn</td>
<td>&lt;20 ppm</td>
<td>20–200 ppm</td>
<td>&gt;200 ppm</td>
</tr>
<tr>
<td>Cu/Cu+Pb+Zn</td>
<td>0.2–0.5</td>
<td>0.2–0.5</td>
<td>0–0.9</td>
</tr>
<tr>
<td>Ba/Cu+Pb+Zn</td>
<td>&lt;1 to &gt;50</td>
<td>&gt;15</td>
<td>&lt;15</td>
</tr>
<tr>
<td>Se</td>
<td>0.001–0.054 ppm</td>
<td>0.001–0.054 ppm</td>
<td>&gt;1 ppm</td>
</tr>
<tr>
<td>As</td>
<td>&lt;10 ppm</td>
<td>20–80 ppm</td>
<td>&lt;30 ppm</td>
</tr>
<tr>
<td>ΣREE</td>
<td>20–60 ppm</td>
<td>20–60 ppm</td>
<td>&lt;20 ppm</td>
</tr>
<tr>
<td>U</td>
<td>&gt;0.2 ppm</td>
<td>&lt;0.2 ppm</td>
<td></td>
</tr>
<tr>
<td>P</td>
<td>&lt;0.03 wt. %</td>
<td>&gt;0.03 wt. %</td>
<td>0.03 wt. %</td>
</tr>
<tr>
<td>δ&lt;sup&gt;18&lt;/sup&gt;O-quartz vsn</td>
<td>&gt;14%</td>
<td>&gt;14%</td>
<td>&lt;14%</td>
</tr>
<tr>
<td>MnO</td>
<td>&lt;0.02 wt. %</td>
<td>0.02–0.05 wt. %</td>
<td>&gt;0.05 wt. %</td>
</tr>
<tr>
<td>Mo</td>
<td>&lt;5 ppm</td>
<td>&gt;5 ppm</td>
<td>&gt;5 ppm</td>
</tr>
</tbody>
</table>
Volcanic facies and alteration

J McPhie
Centre for Ore Deposit Research

Studies in this module were designed to provide a geological framework for geochemical and mineralogical research on alteration related to VHMS mineralisation, and to establish the textural, mineralogical and compositional changes that affect glassy volcanic rocks emplaced in submarine environments. The research focussed on the Cambrian Mount Read Volcanics (MRV) in western Tasmania and the Cambro-Ordovician Mount Windsor Volcanics (MWV) in Queensland. Both host successions comprise complex assemblages of texturally and compositionally diverse volcanic facies that have in addition, been variably deformed, altered and metamorphosed.

Volcanic facies mapping, facies analysis, textural studies and lithogeochemical sampling have been undertaken along four regional traverses and at selected ore deposits (Rosebery, Hercules, Jukes, Heny) in the MRV, and mainly on ore deposit scale (Thalaanga, Highway-Reward) in the MWV. Regional traverses in the MRV greatly refined knowledge of the assemblage of volcanic facies in the major lithostratigraphic units.

The influence of volcanic facies on alteration textures and alteration mineralogy

Sites of ore deposition in VHMS host successions such as the MRV and MWV are highly variable. Footwall facies include thick lava and autoclastic breccia successions (e.g. Heliyer, Thalaanga), syn-volcanic intrusions (e.g. Highway, Reward) and volcaniclastic mass-flow units (e.g. Hercules). Ore bodies may have formed in shallow subseafloor settings where they are replacements of mass-flow units or syn-volcanic intrusions, or else formed at the seafloor. In both settings, syn-volcanic faults could have controlled or influenced the sites of ore deposition.

Factors that appear to strongly control textural and compositional responses of volcanic facies to alteration include the presence of volcanic glass, the porosity and permeability, the composition and external conditions such as depth, pressure, temperature and water/rock ratio. Two very influential factors, the abundance of glass and the porosity and permeability, are closely interrelated.

1. The proportion and distribution of glassy versus crystalline domains
   The glassy domains in submarine lavas and intrusions undergo longer and more complex textural evolution, and exhibit greater compositional changes than do crystalline domains in the same facies.

2. The porosity and permeability of the volcanic facies
   Porosity and permeability vary enormously among different volcanic facies types, spatially within some volcanic facies, and also temporally, from the time of emplacement through compaction and diageneric alteration. Coherent volcanic facies are dominated by fracture-controlled (quench fractures, perlitic fractures) porosity and permeability. In clastic volcanic facies, the interparticle and intraparticle pores control porosity and permeability, so grain type (pumice or scoria versus dense), grain size and sorting are all important.
Distinguishing diagenetic and hydrothermal alteration in VHMS successions

Distinguishing textures and mineralogies specifically related to hydrothermal alteration is critical for successful exploration for VHMS mineralisation. Alteration assemblages involving sericite, carbonate, secondary feldspar and quartz are common in the host facies to massive sulfides in the MRV but are also ubiquitous elsewhere. Useful field criteria include:

- Distribution: Hydrothermal alteration is spatially restricted (metres to tens of metres) and more focussed, whereas the assemblages regarded as regional diagenetic and/or regional metamorphic in origin are widespread (hundreds of metres to kilometres).
- Textural preservation: Diagenetic alteration commonly enhances primary textures, especially in cases involving fracture-controlled alteration. In hydrothermally altered facies, apparent clastic textures are especially common.
- Foliations: In some volcanic facies, early compaction produced a stylolitic, bedding parallel foliation (S1) that affected the earliest diagenetic alteration assemblages.
- Style: Two-phase or polyphase, patchy or banded alteration styles are commonly, but not exclusively, diagenetic.

Significance of volcanic facies studies in VHMS alteration research

An understanding of volcanic facies architecture is critical for the correct interpretation of the stratigraphic, structural and temporal relationships of VHMS deposits, and hence also for the correct interpretation of alteration and mineralisation processes.

Glassy submarine volcanic facies commonly undergo hydration, compaction and devitrification during diagenesis. These processes profoundly alter the porosity, permeability, texture, mineralogy and composition, and may precede or accompany VHMS-related hydrothermal alteration.

The volcanic facies architecture can influence VHMS mineralisation by controlling the hydrology of hydrothermal systems. Submarine volcanic successions are characterised by contrasting porosities of juxtaposed volcanic facies, syn-volcanic faults and syn-volcanic sills bordered by devitrified sedimentary facies, all of which have important effects of fluid pathways and hence on favourable sites for mineralisation.

Bibliography


McPhie J, Dole M, Allen R 1993 Volcanic textures. A guide to the interpretation of textures in volcanic rocks. Centre for Ore Deposit and
Exploration Studies, University of Tasmania, Hobart, 198 p.


---

**Figure 1** Distribution of the principal lithostratigraphic formations and major massive sulfide deposits in the Cambrian Mount Read Volcanics of western Tasmania. Modified from Corbett (1992).
Figure 2 Distribution of the principal formations and major massive sulfide deposits in the Cambro-Ordovician Mount Windsor Volcanics of northern Queensland. Modified from Berry et al. (1992).
Figure 3 Schematic facies architecture of submarine volcanic sequences, such as the Mount Read Volcanics. Typically there are considerable regional variations in relative proportions of lavas, sills and volcanioclastic facies, and in volcanic versus non-volcanic facies. Modified from McPhie and Allen (1992).
Figure 4 Abundance and distribution of glassy, partly glassy and crystalline domains within the principal volcanic facies types found in submarine volcanic successions. Outlines of volcanic facies types correspond to those shown in Figure 3.

Figure 5 Distribution of fracture-controlled versus matrix-controlled porosity in the principal volcanic facies types found in submarine volcanic successions. Outlines of volcanic facies types correspond to those shown in Figure 3. The coherent parts of lavas and intrusions have fracture-controlled porosity whereas the porosity of volcaniclastic facies is matrix-controlled.
Alteration in different glassy volcanics with emphasis on early diagentic alteration; a case study from the Mount Black Volcanics

Cathryn C Gifkins
Centre for Ore Deposit Research

The Mount Black Volcanics were originally a package of dominantly glassy coherent and volcaniclastic materials. The facies distribution and contact relationships are quite complex reflecting the large volume of laterally discontinuous high relief lava domes and the high proportion of lava-like intrusive bodies (Fig. 1). An integrated approach of facies analysis, petrography and geochemistry in the Mount Black Volcanics has resulted in the recognition of a variety of alteration processes that have been active in different glassy volcanic facies. The distribution of the alteration phases generally reflects the complex and discontinuous nature of an originally glassy volcanic pile producing a complex pattern of overlapping alteration styles and intensities.

Regional diagentic alteration

Sericite, feldspar-quartz, sericite-chlorite, chlorite-epidote, chlorite-magnetite and carbonate alteration styles are related to early diagentic alteration processes. Different diagentic alteration styles preserve, enhance or destroy primary volcanic and post-depositional textures. Silicification although widespread and related to diagentic reactions also forms as distinctive hydrothermal alteration haloes around some of the more significant fault systems. A more pervasive style of sericite-quartz alteration is probably largely the result of the regional Greenschist facies metamorphism, while intense sericite, chlorite and carbonate alteration in discrete areas is the product of localised hydrothermal activity.

Early sericite alteration forms thin films that coat originally glassy surfaces. This is identical to the first stage of diagentic alteration of felsic pumice and glass shards in both marine and subaerial environments where thin films of clay and opal develop on all glass surfaces (Iijima, 1974). Sericite alteration preserves the pumice textures and also defines and enhances prelitic fractures in originally glassy lavas and sills.

The next diagentic alteration phase is represented by feldspar-quartz alteration. At least two stages of feldspar alteration, K-feldspar pre-dating Na-feldspar (albite), have been recognised. Feldspar-quartz alteration replaces uncompactd shards and pumice fragments, crystalline flowbands and spherulites preserving these primary volcanic and devitrification textures. Feldspar alteration is either an early diagentic phase or it replaces an early diagentic mineral phase, like zeolite.

Chlorite-rich alteration assemblages are more texturally destructive. In pumice-rich deposits only the porphyritic nature of pumice in the chlorite-rich fiamme is preserved. In coherent rocks chlorite-rich alteration assemblages pervasively replace the fine-grained groundmass.

Controls on the distribution of diagentic alteration styles

The complex distribution and intensity of the alteration styles in part can be related to the texture and geochemical signature of the volcanics. With weak alteration it appears that the composition of the glass is probably the most important factor in determining the type of alteration, however permeability and competency contrast also influence the
Figure 1: Schematic facies architecture of the submarine Mount Black Volcanic succession. The Mount Black Volcanics are a several kilometre thick package of feldspar-phryic massive, flow-banded and flow-breciated lavas and sills of generally rhyolitic to dacitic composition with minor andesite. Variable proportions of lithic and/or pumice-rich volcaniclastic breccias, shard-rich sandstones and siltstones, and crystal-rich sandstones are interbedded with, or intruded by the more coherent units. The volcanic architecture depicts the relationships between the main lithofacies and variations in the internal texture of many of these units.
alteration. With increasing intensity of alteration the permeability and competency contrasts have less control on the alteration.

As most of the diagenetic alteration phases are relatively weak, the alteration is strongly controlled by fluid pathways. In coherent facies fractures produced by quenching, flow and hydration and the distribution of more crystalline versus glassy textures have controlled the distribution of the diagenetic alteration assemblages. Although the original porosity of the coherent lavas and sills was low, the highly unstable nature of glass and the intensely perlitic fractured margins of the coherent units became permeable zones that have undergone subsequent diagenetic alteration. The result is that coherent units, particularly at the margins, are not necessarily less intensely altered than the originally more porous volcaniclastic units. Alteration in clastic facies is controlled by the distribution of porous matrix to less porous clasts. Early phases of alteration in clastic facies, although commonly patchy, are more pervasive and less focussed than similar alteration in the coherent facies.

**Distinguishing between diagenesis and hydrothermal alteration**

Diagenetic alteration phases are interpreted to be those that are regional in distribution and pre-date or are synchronous with the stylolitic S1 compaction foliation. The diagenetic alteration generally involves only weak to moderate intensities of alteration and subtle deviations in the chemical signature. Mass enrichment's and depletions during diagenetic alteration are an order of magnitude smaller than changes related to medial facies hydrothermal alteration. Diagenetic alteration also appears to be more complicated with enrichment's and depletions in SiO₂, Na₂O, CaO, K₂O, Fe₂O₃, MgO, MnO, Rb, Sr and Ba occurring over a small scale in association with different alteration facies. The complex and polyphase nature of the diagenetic alteration results in samples on the Al versus CCPI box plot migrating back and forth through the least altered fields (Fig. 2).

In contrast, alteration styles that are only local in distribution are related to discrete hydrothermal cells. Hydrothermal alteration appears to be more pervasive, intense and texturally destructive, obscuring both groundmass textures and primary plagioclase crystals. The enrichments and depletions associated with hydrothermal alteration are consistent over a wider area than the mass changes associated with diagenetic alteration.

**Bibliography**


Giffkins, C., 1998. Alteration in different glassy volcanics with emphasis on early diagenetic alteration; a case study from the Mount Black Volcanics. CODES: AMIRA/ARC Project P439, Final report.


Figure 2: Alteration box plots: Alteration Index (Ishikawa et al., 1976) versus Chlorite/carbonate/pyrite Index (Large, 1996) for the Mount Black and Sterling Valley Volcanics. Closed symbols represent samples that are least or only weakly altered, while open symbols represent noticeably altered samples. The degree of alteration has been determined using petrography and hand specimen analysis. The majority of samples plot in or near the least altered boxes. However, altered samples that plot in the dolomite-chlorite-sericite triangle are interpreted as hydrothermally altered samples. a) Represents least altered to strongly altered rocks of suite 1 (rhyolites). The least altered field is drawn in red. b) Depicts the variation in alteration indices for the dacites (suite 2 and 3). The least altered field is drawn in purple. c) Depicts the andesitic and basaltic rocks of suites 4 and 5. Least altered field is drawn in black.
Influence of volcanic facies on hydrothermal and diagenetic alteration: Evidence from the Highway–Reward deposit, Mount Windsor Subprovince, Queensland

Mark G. Doyle
Centre for Ore Deposit Research

Interpreting timing relationships of alteration in volcanic rocks

Volcanic facies undergo mineralogical and textural changes resulting from hydration, high temperature devitrification, diagenetic alteration and compaction, hydrothermal alteration and metamorphism. Each represents an alteration stage although the time between each stage may be very short or even overlap and several stages may be unimportant or not affect all parts of a volcanic deposit/rock.

Determination of the relative roles, timing and significance of each alteration stage in modifying a volcanic rock or facies is dependant on: (1) establishing the regional extent of mineral assemblages; (2) interpreting overprinting relationships of alteration minerals to primary volcanic textures, volcanic facies and intrusive units; (3) relation of alteration minerals and assemblages to tectonic, diagenetic and stylolitic foliations; (4) overprinting relationship between different alteration mineral assemblages; (5) consideration of mineral stability fields; and (6) knowledge of alteration assemblages in comparable modern volcanic successions (e.g., Green Tuff Belt, Japan).

Timing relationships suggest that 5 principal alteration styles are identifiable in the Highway-Reward area: (a) regional diagenetic alteration of formerly glassy volcanic facies; (b) “barren” alteration associated with local, “low temperature” hydrothermal systems; (c) syn-volcanic hydrothermal alteration concurrent with mineralisation; (d) regional metamorphic assemblages; and (e) quartz-topaz-pyrophyllite alteration. The quartz-topaz-pyrophyllite alteration is interpreted as epigenetic and of a syn- to post-D4 (Siluro-Devonian) age. A detailed summary of trace and major element variations of the principal alteration styles is presented by Doyle 1998b (this volume).

Alteration of coherent lavas and intrusions

Lavas and intrusions at Highway-Reward comprised glassy margins which pass inward to cores characterised by zones of glass, glass with coalescing spherulites and/or crystalline domains characterised by spherulitic and micropoikilitic textures. The distribution of hydrothermal and diagenetic alteration assemblages is strongly influenced by the pre-existing texture and zonation. The textural progression is briefly discussed below and summarised in Figures 1 to 3.

Glassy margins and domains

The glassy domains originally comprised coherent volcanic glass (Fig. 1.1A), in situ quench fractured glass (Fig. 1.1B) and variably matrix-rich autoclastic (hyaloclastite, autobreccia) breccia facies (Fig. 1.1C). After emplacement, perlitic cracks sometimes developed in response to hydration of the glass. Circulating fluids moved out from fractures (perlitic, quench) and the matrix progressively altering the glass in several stages. The initial alteration was either pervasive (Fig. 1.3.2) or ceased before completely affecting the whole rock, leaving domains of glass that were altered during a second alteration step or stage (Fig. 1.4.1).

In diagenetically-altered samples, perlite kernels and clasts in autoclastic breccia have often completely altered to feldspar (albite) or chlorite during initial
Figure 1: Stages in the textural evolution of the glassy margins of lenses and syn-sedimentary infusions. 1. Prior to the onset of alteration the glassy margins comprised: (A) dense glass; (B) quenched fractured glass; and (C) autostatic breccia. 2. After emplacement and in the cooling history of the glass, the cracks can be activated and (if present) will control deformation of the glass. The cracks can be: (a) in the matrix or peripheral area, (b) in the bulk glass and (c) in the breccia matrix. 3. Subsequent alteration and recrystallisation in cases where the fracture pattern was not controlled by the matrix or autostatic breccia. 4. Subsequent alteration and recrystallisation in cases where the fracture pattern was controlled by the matrix or autostatic breccia or overprinted alteration domains of similar mineralogy (4).
diagenetic alteration. Later sericite, chlorite or quartz alteration often commenced along fractures and moved out from these into the albite- or chlorite-altered domains. In some samples, feldspar alteration has been partially replaced by quartzfeldspar. Subsequent generations of sericite and/or quartz alteration overprinted earlier phyllosilicate alteration and were also fracture controlled (cf. Allen, 1988; McPhie et al., 1993).

Within the Highways-Reward hydrothermal alteration envelope, hyaloclastite clasts often pervasively sericite-chlorite-altered and feldspar phenocrysts are variably sericite, chlorite and/or carbonate-altered. Subsequent quartz-sericite alteration commenced along fractures and the matrix of autoclastic breccia (Fig. 1.4.2). In other cases, perlitic fractures and the matrix between clasts have altered to sericite, whereas perlite kernels have altered to fine quartzo-feldspathic mosaics. More advanced alteration rarely extends far from the fractures or matrix, possibly due to the reduction in porosity accompanying initial alteration stages (cf. McPhie et al., 1993).

**Mixed glassy and devitrified domains**

Prior to alteration, mixed glassy and devitrified domains comprised spherulites, lithophysae scattered in glass (Fig. 2.1A), coalescing spherulites enclosing cusparate patches of glass (Fig. 2.2B), and flow bands of glass alternating with devitrified bands (Fig. 2.1C). Hydrothermal and diagenetic alteration of the glassy domains progressed along a similar path to the glassy margins. The glassy bands now comprises sericite, chlorite, feldspar, carbonate or micropoikilitic quartz. In contrast, crystalline and devitrified domains are silicified or have recrystallised to quartzo-feldspathic mosaics (Fig. 2.4). In autoclastic breccia, clasts are often flow banded and show a similar textural progression.

**Crystalline domains**

Large parts of some lavas and intrusions have a groundmass comprising spherulites, micropoikilitic texture or granophytic texture. In many samples, hydrothermal alteration and metamorphism have recrystallised original fibrous devitrification textures to pale quartzo-feldspathic mosaics with interstitial sericite, hematite, pyrite (Fig. 3.3-3.5). The result in

hand specimen is a granular texture. In these zones, overprinting hydrothermal alteration generates mottled and patchy domains of different mineral assemblages.

**Controls on the localisation of hydrothermal alteration**

The shapes, dimensions and distribution of hydrothermal circulation (and therefore alteration and mineralisation) are closely related to the initial patterns of permeability and compositional contrasts in the host succession. Fluid conduits may include syn-volcanic faults, porus and permeable volcaniclastic facies and the glassy margins of lavas or syn-sedimentary intrusions (Doyle, 1997a,b).

By acting as a relatively impermeable barrier (cf. Einsele et al., 1980; McPhie, 1993), indurated sediment at the margins of intrusions and devitrified facies of lavas can cause fluids to be focussed along faults, local autoclastic breccia intervals, or within the fractured glassy margins of lavas and/or intrusions (cf. Large, 1992). Under these circumstances, well focussed fluid flow gives rise to lens- or pipe-shaped massive sulphide deposits and well developed, zoned alteration pipes (e.g. Highway-Reward; Doyle, 1997a,b).

Other massive sulphide deposits are hosted by sequences dominated by originally highly porous, permeable, water saturated and glassy volcaniclastic units. Ascending hydrothermal permeate through the substrate to produce widespread strata-bound alteration zones and lens- or sheet-style massive sulphide mineralisation (cf. Large, 1992). Sulphide deposition often commences below the seafloor by replacement and infilling of pore space within the volcaniclastic units (e.g. Rosebery, Hercules (Allen, 1994), Liontown (Miller, 1996; Doyle unpub. data).

In VHMS host successions comprising mixtures of relatively poorly porous rocks (e.g. lavas and shallow intrusions) and incompetent, very porous deposits (e.g. pumiceous units), a variety of mineralisation types and alteration styles may develop in close proximity.
Figure 2. Stages in the textural evolution of the mixed glassy and devitrified domains in lavas and syn-sedimentary intrusions. 1 — Prior to the onset of alteration the mixed glassy and spherulitic zones comprised: (1A) glass with scattered spherulites; (1B) coalescing spherulites with cuspatte areas of glass; and/or (1C) devitrified bands alternating with glassy bands. 2 — After emplacement perlitic cracks sometimes developed in glassy bands. 3 — Initial hydrothermal or diagenetic alteration of the glassy domains was similar to alteration in the glassy margins. Alteration was fracture controlled or pervasive and tended towards sericite- or chlorite-rich assemblages. 4 — Subsequent alteration of the glassy domains was also mainly localised along fractures or overprinted alteration domains of similar mineralogy. In contrast, single spherulites and bands or nodules of devitrification structures were recrystallised, silicified or replaced by feldspar during alteration.
Figure 3. Stages in the alteration of the crystalline and devitrified domains in lavas and syn-sedimentary intrusions. 1—These domains have a groundmass of closely packed spherulites, microlites or micropoikilitic quartz. Joints are sometimes present. 3—Diagenetic and hydrothermal alteration often commence along joints or hydraulic fractures. 4—Subsequent alteration attacked phenocrysts (e.g. feldspar) and generated patchy and mottled textures. 5—Recrystallisation and silicification of spherulites during hydrothermal alteration and metamorphism destroys fibrous devitrification structures generating a quartzo-feldspathic mosaic.
Bibliography


Bibliography of all P439 contributions


Doyle, M.W., 1995. Preliminary investigation of alteration at the Highway and Reward
deposits, Mt Windsor Volcanics, Queensland. CODES: AMIRA/ARC Project P439, Report 1: 149-155.


Herrmann, W., 1998. Use of immobile elements and chemostratigraphy to determine precursor volcanics. CODES AMIRA/ARC P439, Final report.


Large, R., Allen, R., & Blake, M., 1997. Carbonate and muscovite mineral chemistry, Rosebery VHMS deposit. CODES: AMIRA/ARC Project


geochemical characteristics of alteration from the Hall Rivulet Canal-Mt Read-Red Hills-Anthony Dam traverse, Mt Read Volcanic Belt. CODES: AMIRA/ARC Project P439, Report 3: 379-424.


