In situ multiple sulfur isotope analysis by SIMS of pyrite, chalcopyrite, pyrrhotite, and pentlandite to refine magmatic ore genetic models

La Flamme, Crystal\textsuperscript{a}, Martin, Laure\textsuperscript{b}, Jeon, Heejin\textsuperscript{b}, Reddy, Steven, M.\textsuperscript{c}, Selvaraja, Vikraman\textsuperscript{a}, Caruso, Stefano\textsuperscript{a}, Bui, Thi Hao\textsuperscript{d}, Roberts, Malcolm P.\textsuperscript{b}, Voute, Francois\textsuperscript{a}, Hagemann, Steffen\textsuperscript{a}, Wacey, David\textsuperscript{b}, Littman, Sten\textsuperscript{e}, Wing, Boswell\textsuperscript{d}, Fiorentini, Marco\textsuperscript{a}, Kilburn, Matthew R.\textsuperscript{b}

\textsuperscript{a}Centre for Exploration Targeting, ARC Research Council Centre of Excellence for Core to Crust Fluid Systems (CCFS), University of Western Australia, Australia
\textsuperscript{b}Centre for Microscopy, Characterisation, and Analysis, ARC Centre of Excellence for Core to Crust Fluid Systems (CCFS), University of Western Australia, Australia
\textsuperscript{c}The Institute for Geoscience Research, Department of Applied Geology, Curtin University of Technology, Australia
\textsuperscript{d}Department of Earth and Planetary Sciences and GEOTOP, McGill University, Canada
\textsuperscript{e}Max-Planck-Institut für Marine Mikrobiologie, Bremen, Germany

*corresponding author: crystal.laflamme@uwa.edu.au

ABSTRACT

With growing interest in the application of in situ multiple sulfur isotope analysis to a variety of mineral systems, we report here the development of a suite of sulfur isotope standards for distribution relevant to magmatic, magmatic-hydrothermal, and hydrothermal ore systems. These materials include Sierra pyrite (FeS\textsubscript{2}), Nifty-b chalcopyrite (CuFeS\textsubscript{2}), Alexo pyrrhotite (Fe\textsubscript{1-x}S), and VMSO pentlandite ((Fe,Ni)\textsubscript{x}S\textsubscript{8}) that have been chemically characterized by electron microprobe analysis, isotopically characterized for $\delta^{33}$S, $\delta^{34}$S, and $\delta^{36}$S by fluorination gas-source mass spectrometry, and tested for homogeneity at the micro-scale by secondary ion mass spectrometry. Beam-sample interaction as a function of crystallographic orientation is determined to have no effect on $\delta^{34}$S and $\Delta^{33}$S isotopic measurements of pentlandite. These new findings provided the basis for a case study on the genesis of the Long-Victor nickel-sulfide deposit located in the world class Kambalda nickel camp in the southern Kalgoorlie Terrane of Western Australia. Results demonstrate that precise multiple sulfur isotope analyses from magmatic pentlandite, pyrrhotite and chalcopyrite can better constrain genetic models related to ore-forming processes. Data indicate that pentlandite, pyrrhotite and chalcopyrite are in isotopic equilibrium and display similar $\Delta^{33}$S values +0.2‰. This isotopic equilibrium unequivocally fingerprints the isotopic signature of the
magmatic assemblage. The three sulfide phases show slightly variable δ³⁴S values
(δ³⁴S_chalcopyrite = 2.9 ± 0.3‰, δ³⁴S_pentlandite = 3.1 ± 0.2‰, and δ³⁴S_pyrrhotite = 3.9 ± 0.5‰), which
are indicative of natural fractionation. Careful in situ multiple sulfur isotope analysis of
multiple sulfide phases is able to capture the subtle isotopic variability of the magmatic
sulfide assemblage, which may help resolve the nature of the ore-forming process. Hence,
this SIMS-based approach discriminates the magmatic sulfur isotope signature from that
recorded in metamorphic- and alteration-related sulfides, which is not resolved during bulk
rock fluorination analysis. The results indicate that, unlike the giant dunite-hosted komatiite
systems that thermo-mechanically assimilated volcanogenic massive sulfides proximal to
vents and display negative Δ³³S values, the Kambalda ores formed in relatively distal
environments assimilating abyssal sulfidic shales.

HIGHLIGHTS

• Characterisation of four sulfide standards for multiple sulfur isotope analysis: pyrite, chalcopyrite, pyrrhotite, and pentlandite for distribution
• Analysis of orientation effect in pentlandite
• Natural sulfur isotope fractionation between pentlandite and pyrrhotite
• Case study multiple sulfur isotope analysis of three sulfide phases within world-class Long-Victor komatiite-hosted nickel-sulfide deposit

KEYWORDS

Multiple sulfur isotopes, SIMS, in situ, sulfide minerals, ore genesis

1. INTRODUCTION

Sulfur is a trace element in silicate melts, typically concentrated below 0.2 wt.%. However, it
is an essential element in a wide range of environments including the lithosphere, biosphere,
hydrosphere, and atmosphere. In recent years, our understanding of the sulfur cycle and its
role in the evolution of these terrestrial reservoirs has been revolutionised by the study of the
sulfur isotope composition of pyrite, the most common sulfide mineral (Farquhar et al., 2000;
Kump, 2012 and references therein; Strauss, 1997; Thomassot et al., 2015). We have gained a
fundamental understanding into the development of early Earth’s processes, in particular
those linked to the emergence of life and the development of an oxygenated atmosphere
(Farquhar et al., 2000), by the discovery of mass independent fractionation (MIF) of sulfur
isotopes.

Sulfur resides in the Earth’s mantle, crust and hydrosphere but is locally concentrated in
mineralised systems typically associated with ore deposits, where it acts as the primary
complexing ligand in the formation of sulfide minerals. Mantle- and crustally-derived
magmas have brought large quantities of economic metals from the Earth’s interior to the
near surface, and hydrothermal fluids have remobilised and re-precipitated these metals
within the crust as different sulfides. The sulfur itself may be sourced from a variety of
compositional reservoirs, each with distinct isotopic compositions. Mixing and interactions
with the mantle, crustal magmas, hydrothermal fluids, country rocks, or meteoric waters
imparts specific isotopic signatures, resulting in minerals with a range of isotopic
compositions. As such, intra-grain and inter-grain chemical and isotopic variations in sulfur-
rich mineralised systems record the interaction of these different reservoirs and offer unique
insights into the complex fluid-rock interactions within mineral systems (McCuaig et al.
2010). For example, in magmatic ore deposits, sulfur isotope data have fingerprinted the
source of the sulfur linked to ore genesis (Bekker et al., 2009; Chang et al., 2008; Fiorentini
et al., 2012a; Hiebert et al., 2013; Lesher and Groves, 1986; Penniston-Dorland et al., 2008;
Sharman et al., 2013) and constrained the geodynamic framework where these deposits
formed (e.g., Chen et al., 2015; Fiorentini et al., 2012b; Giacometti et al., 2014). Similarly,
sulfur isotope studies have proven to be vital in characterising magmatic-hydrothermal (Helt
et al., 2014; Xue et al., 2013) and hydrothermal systems (e.g., Jamieson et al., 2013; Leach et
al., 2005; Sharman et al., 2015). Constraining the sulfur isotopic signature in magmatic-
hydrothermal mineral systems is useful in delineating the source of sulfur, and is an
important parameter to understand how, when and where sulfur saturation occurs (e.g., Evans
et al., 2014). In addition, such data provides a better understanding of the geodynamic
environment in which the mineralisation process occurs which impacts on the targeting
rationale applied during exploration (e.g., Fiorentini et al., 2012a). Consequently, ore
deposits are a perfect laboratory for understanding the source and mobility of sulfur in a wide
variety of settings.

Mineral systems and ore deposits have characteristically complex microscale intra-granular
and inter-granular textures due to variations in their chemistry during formation and
subsequent re-equilibration during cooling (e.g., pentlandite exsolution in pyrrhotite; Durazzo
and Taylor, 1982). In situ sulfur isotope analysis at the microscale has the potential to
revolutionise our understanding of ore forming processes. The development of in situ
analytical techniques using laser ablation-(multi-collection)-inductively coupled plasma mass
spectrometry (LA-MC-ICPMS; see Bühn et al., 2012; Craddock et al., 2008) and large
group secondary ion mass spectrometry (SIMS; see Farquhar et al., 2013; Ireland et al.,
2014; Ushikubo et al., 2014; Whitehouse, 2013) now allows high-precision isotopic analysis
of multiple sulfur isotopes with spot sizes nearing ten microns. The ultra-high sensitivity of
SIMS in particular affords the ability to measure the least common stable isotopes of sulfur,
\(^{33}\text{S} (0.75\%)\) and \(^{36}\text{S} (0.02\%)\), together with the more abundant \(^{32}\text{S} (95.02\%)\) and \(^{34}\text{S} (4.21\%)\)
isotopes, simultaneously from the same volume of material. This gives the potential to
identify the anomalous sulfur isotopic signatures indicative of mass independent fractionation
(MIF; \(\Delta^{33}\text{S}\) and \(\Delta^{36}\text{S}\)) together with \(\delta^{34}\text{S}\).

Instrumental mass fractionation in SIMS is intrinsically linked to the composition and
crystallographic orientation of the material being analysed and the specific conditions under
which the analysis is performed. Therefore, accurate isotopic measurements require careful
standardization against a suitable matrix-matched reference material (Eiler et al., 1997; Stern,
Although a number of reference materials have been developed for acquiring in situ 2-
sulfur isotopes (e.g., Kozdon et al., 2010), the majority of published in situ 3- or 4-
sulfur isotope analyses have been acquired from the most common sulfide mineral, pyrite. As such, 
many SIMS laboratories worldwide have developed “in-house” pyrite (and to a lesser degree 
other) standards for multiple sulfur isotope analysis (Balmat, Isua 248474, Ruttan; see 
Whitehouse, 2013; Ushikubo et al., 2014; Hauri et al., 2016). With growing interest in the 
application of in situ sulfur isotope analysis to a wide variety range of mineral systems, there 
is an increased need for reference material for a variety of common sulfide minerals, which 
presently remain scarce. In addition to composition, sulfide crystallography has previously 
been demonstrated to potentially create an orientation effect-induced instrumental mass bias 
on sulfur isotope analysis for certain sulfides including galena (PbS) and sphalerite 
((Zn,Fe)S) but not others including pyrite (FeS2), pyrrhotite (Fe1-xS), and chalcopyrite 
(CuFeS2) (Kozdon et al., 2010; Kita et al., 2011). Of these sulfides, most have a diamond-
cubic crystal structure, with the exception of galena which has a cubic hexoctahedral 
structure. Here, we build on this observation to demonstrate that orientation effect does not 
induce instrumental mass fractionation on δ34S and Δ33S in another cubic hexoctahedral 
sulfide - pentlandite ((Fe,Ni)9S8).

Deviations from mass dependent fractionation are typically (but not uniquely) triggered by 
the presence of an oxygen-poor atmosphere that existed before the Great Oxidation Event at 
ca. 2.4 Ga in which ultraviolet radiation was the driver for mass independent photochemical 
separation of sulfur isotopes (e.g., Farquhar et al., 2000; Farquhar and Wing, 2003). For this 
reason, the measure of 33S has become essential to evaluate the full suite of isotopic features 
of Archean rocks (e.g., Bühn et al., 2012; Farquhar et al., 2013), and those from younger 
terranes that might be sourcing Archean rocks (e.g., Cabral et al., 2013; Selvaraja et al., in 
submission). The least abundant stable isotope of sulfur, 36S, behaves similarly to 33S, and has 
also become increasingly important for fingerprinting Archean source rocks and 
understanding early Earth processes. The Δ36S/Δ33S ratio can elucidate between mass 
dependent processes and contributions from Archean MIF sources when magnitudes of mass 
dependent fractionation deviations are small (Farquhar et al., 2007; Johnston, 2011).

We present multiple sulfur isotope data from pentlandite, pyrrhotite, and chalcopyrite from 
an Archean komatiitic massive nickel-sulfide deposit. To do so, we have developed four 
reference materials (pyrite, chalcopyrite, pyrrhotite, and pentlandite) that we have fully 
chemically and isotopically characterized for multiple sulfur isotope analysis. We present 
multiple sulfur isotope results from multiple sulfides within the deposit to constrain the 
source of the sulfur that triggered sulfide saturation and investigate the geodynamic setting in 
which this magmatic system was emplaced.

2. METHODS

To determine the chemical and isotope composition of potential sulfide reference materials 
we combined three analytical techniques. Firstly, we performed wavelength dispersive 
spectrometry (WDS) by electron probe micro-analysis (EPMA) on a multitude of grains (or 
grain fragments) of each candidate reference material to ensure that the reference material is
chemically homogeneous across a number of grains (or grain fragments). Chemical compositions (spot analyses) were linked to chemical WDS maps and scanning electron microscope backscatter electron (SEM-BSE) images to highlight potential chemical zonation, mineralogical inclusions, and fractures. Details of SEM-BSE and EPMA-WDS are presented in the Supplementary Material.

Secondly, at least five grains (or grain fragments) of each candidate reference material were analysed for in situ multiple sulfur isotopes by SIMS to determine whether an adequate level of reproducibility exists at the intra- and inter-grain scale (isotopic homogeneity). Multiple materials were tested prior to selecting the material that displayed adequate isotopic homogeneity (reproducibility on $^{34}\text{S}/^{32}\text{S}$ better than 0.5‰). For instance, five materials from different environments were tested to find a suitable chalcopyrite standard (see section 5.1 for discussion on selecting sulfide reference material). To further determine on what scale sufficient homogeneity existed once the reference material was selected, more than 40 grains for each material were analysed. If the reference material was deemed suitable, multiple grains or grain fragments (in the case of material derived from large crystals) were analysed by bulk multiple sulfur analyses. Bulk analyses were completed by fluorination gas-source mass spectrometry. In each case, sulfur was extracted from 5 to 11 separate aliquots (of extracted sulfur from separate grains or grain fragments) and analysed for its multiple sulfur isotopic composition. We further evaluated the homogeneity of the SIMS material by completing an MSWD test on the data after Wing and Farquhar, (2015). The respective analytical techniques are described in detail below.

For the case study, we investigated the chemical composition and multiple sulfur isotope signature of three phases (pentlandite, pyrrhotite, chalcopyrite). A micro-X-ray Fluorescence (XRF) elemental map was acquired to visually investigate the textural relationship between the three sulfide phases and choose samples for in situ and bulk isotope analyses (details regarding analytical set up are presented in the Supplementary Materials). EPMA analyses on pyrrhotite and pentlandite were completed to ensure a similar chemical composition to the reference materials (see Appendix A). In situ isotope signatures of pentlandite, pyrrhotite and chalcopyrite were compared to ten bulk rock fluorination values that incorporated all the co-existing sulfide phases (see Appendix B and C). We also used the case study material to assess the crystallographic orientation effect in pentlandite. To do so, Electron Backscatter Diffraction Analysis (EBSD) was completed on this sample and pentlandite crystal orientation was compared with multiple sulfur isotope results.

2.1 Secondary Ion Mass Spectrometry

2.1.1 Analytical set up and conditions

In situ sulfur isotopic ratios were measured using a CAMECA IMS1280 large-geometry ion microprobe at the CMCA, UWA. Sample mounts were made by coring 3 mm diameter pucks from rock fragments using a drill press fitted with diamond drill bits, then mounted and cast in the central portion of a 25 mm in diameter epoxy mount. Standard blocks (cast separately to be reused) were made by mounting 1-2 grain fragments of pyrite chalcopyrite and 1 mm diameter pucks of rock fragments host to pyrrhotite and pentlandite. Reference materials
were cast in epoxy ~8 mm from the edge of a mount. Sample mounts and standard blocks were trimmed to a thickness of 5 mm using a precision saw, coated with 30 nm of gold, and mounted together (after being appropriately trimmed) in the sample holder (further details provided in the Supplementary Materials). Care was taken to set the surfaces of the standard and sample blocks at the same level in the sample holder. If necessary, a small amount of carbon paint was applied to provide good conductivity between both pieces. To ensure that results are not a product of analytical artefacts due to X-Y-Z positioning, a reference material is also mounted and cast with the sample block to compare with values generated from the standard block. Additional details regarding sample preparation for SIMS analysis are presented in the Supplementary Materials.

The ion microprobe was operated in multicollection mode using a Cs\(^+\) primary beam with an intensity of \(\sim 1\)–4 nA in Gaussian mode that interacted with the sample at 20 keV. In some instances, depending on the nature of the sample (i.e., size of sulfides, surrounding material, whether sulfides are mounted as individual grains or incorporated in rock chips) a normal incidence electron flood gun was used for charge compensation. Following a 30 s pre-sputter, secondary sulfur ions from the target sample were extracted at -10 kV and admitted to the mass spectrometer with a field magnification of 133\(\times\), with automated centering of the secondary beam in the field/aperture (both x and y; aperture size 4000 μm) and entrance slit (x direction only; slit width 60 μm or 90 μm in some cases). The NMR magnetic field controller locked the axial mass at the beginning of each session, and the mass spectrometer operated at a mass resolution (M/ΔM) of about 2500 (exit slit width of 500 μm on the multicollector). Under these conditions, the hydrite interference \(^{32}\)S\(^{1}\)H on the \(^{33}\)S peak was avoided by offsetting the \(^{33}\)S peak centre to the low mass side.

For triple sulfur isotope measurements (\(^{32}\)S, \(^{33}\)S and \(^{34}\)S), a 15 μm raster was applied, and the sulfur isotopes were simultaneously detected by three Faraday Cups using amplifiers with 10\(^{10}\) Ω (L’2), 10\(^{11}\) Ω (L1), and 10\(^{11}\) Ω (FC2 or H1) resistors. Data were collected over 123 s of acquisition time in 20 integration cycles. Count rates on \(^{32}\)S varied from 1×10\(^9\) to 3.5×10\(^9\) cps on pyrite, depending on the intensity of the primary beam but is typically 2.2–2.4×10\(^9\) cps with a 2.5 nA primary beam.

Quadruple sulfur isotope (\(^{32}\)S, \(^{33}\)S, \(^{34}\)S, and \(^{36}\)S) analysis has different analytical protocols to include the measurement of \(^{36}\)S using a low-noise ion counting electron multiplier (EM) in the H2 position of the multicollector axis. A higher primary beam current (3–4 nA), larger raster (30 μm for pre-sputter and 20 μm for analysis) and longer acquisition time (279 s in 45 integration cycles) were necessary to achieve adequate repeatability on \(^{36}\)S. The count rates on \(^{36}\)S of typically 2–4×10\(^5\) cps for pyrite cause a significant gain drift for the EM. Hence, the EM high voltage was optimised using a Pulse Height Amplitude (PHA) distribution curve at the start of each session, and the gain drift was measured three times during each analysis (at the beginning, middle and end) and corrected by the CAMECA CIPS software (see Schuhmacher et al., 2004). In the case of unknown material, measurements were interspersed with matrix-matched reference material to calibrate isotope ratios and monitor internal sample repeatability.
2.1.2 *Data processing and error propagation*

In situ sulfur isotope measurements by SIMS are corrected in two steps. Firstly, the pyrite standard (Sierra; see section 3.1) is analysed once every five to eight analyses, regardless of whether the pyrite is being used as the primary standard for that particular session, allowing for the assessment of the stability of the instrument during the given analytical session, and correct for instrumental drift. Secondly, sample isotopic ratios are corrected for instrumental mass fractionation using the correction factor \( \alpha \), determined by normalising the mean of all measurements on the matrix matched reference material, \( R_{\text{std}} \), to the isotopic ratio of the reference material \( R_{\text{RM}} \) as obtained by independent bulk methods (e.g. fluorination gas-source mass spectrometry):

\[
\alpha = \frac{R_{\text{std}}}{R_{\text{RM}}}
\]

The propagated uncertainty for the \( \delta^{34}\text{S} \) value of each sample spot takes into account the internal error on the raw isotopic ratios, the uncertainty on the drift correction where necessary and the uncertainty on the standard measurement, calculated as the standard deviation on the mean isotopic ratios measured in the standards.

Identifying MIF has become increasingly important in assessing a number of geological environments because it reveals fundamental information pertaining to the age and source of sulfide mineralization (e.g., Johnston, 2011). MIF is presented as \( \Delta^{33}\text{S} \) and \( \Delta^{36}\text{S} \) to quantify the deviation from the mass dependent fractionation slope; however, these values can be small in magnitude – often much less than 1.0‰ (see Figure 1). Therefore, it is important to systematically quantify uncertainty on these values to ascertain whether a MIF signature does indeed exist.

The mass independent relationship is denoted by the \( \Delta^{33}\text{S} \) and \( \Delta^{36}\text{S} \) notation to represent the deviation between the isotopic ratios measured and those predicted according to mass dependent fractionation, and is defined as:

\[
\Delta^{x}\text{S}_i = \delta^{x}\text{S}_i - 1000 \times \left[ \left( \frac{\delta^{34}\text{S}_{\text{V-CDT}}}{1000} + 1 \right)^\lambda - 1 \right]
\]

where \( x \) is either 33 or 36 and \( \lambda \) is the slope of the mass dependent fractionation line (0.515 for \( \delta^{33}\text{S} \) and 1.91 for \( \delta^{36}\text{S} \); Hulston and Thode, 1965; Ono et al., 2006a). The \( \lambda \) values approximate the relationships for high temperature equilibrium isotopic fractionations (Farquhar and Wing, 2003).
Figure 1: Calculated magnitudes of mass independent fractionation as deviations from the mass dependent fractionation line (MDF) presented in $\delta^{33}S$ vs. $\delta^{34}S$ space. The small deviations from the MDF line represent mass independent fractionation. This highlights the importance of: 1) precise and accurate multiple sulfur isotope measurements, and 2) quantification of uncertainty on $\Delta^{33}S$. Although not expressed in this figure, quantifying $\Delta^{36}S$ is equally important.

The uncertainty on $\Delta^{33}S$ and $\Delta^{36}S$ is calculated by propagating the uncertainties on $\delta^{33}S$, $\delta^{34}S$, and $\delta^{36}S$; however, paired sulfur isotope ratios covary in $\delta^{33}S$–$\delta^{34}S$ and $\delta^{36}S$–$\delta^{34}S$ space. To account for this relationship, we determined the covariance on the reference material of the analytical session and propagated to the measurements of $\delta^{33}S$ and $\delta^{34}S$ ($\sigma_{\delta^{34}S}^{\delta^{33}S}$) and $\delta^{36}S$ and $\delta^{34}S$ ($\sigma_{\delta^{34}S}^{\delta^{36}S}$). The uncertainty on $\Delta^{3}S$ is calculated here in a similar manner to Farquhar et al. (2013) but also accounts for this covariance in the formulation of the uncertainties on $\Delta^{33}S$ and $\Delta^{36}S$ as follows:

$$
\sigma_{\Delta^{3}S} = \sqrt{\sigma_{\delta^{3}S}^{2} + \sigma_{\delta^{3}S}^{2} \times \left[ \lambda \times \left( 1 + \frac{\delta^{34}S}{1000} \right)^{-(1-\lambda)} \times \sigma_{\delta^{3}S}^{2} \right]^2 + 2 \times \sigma_{\delta^{3}S}^{2} \times \sigma_{\delta^{3}S}^{2} \times \left[ \lambda \times \left( 1 + \frac{\delta^{34}S}{1000} \right)^{-(1-\lambda)} \right]}
$$

where variables are as in equation 2.

2.2 Fluorination gas-source mass spectrometry

Samples for fluorination coupled with gas-source mass spectrometry were analysed at the Stable Isotope Laboratory of the Department of Earth and Planetary Sciences at McGill University, Montreal, Canada. Sulfide-bearing samples were microdrilled using a 1 mm
diameter hand drill. Each sample was microdrilled multiple times. Sulfur was chemically
extracted from the 15–30 mg powders to form silver sulfide by chromium reduction as
described by Canfield et al. (1986). Silver sulfide was fluorinated at 225°C in a Ni bomb
under F₂ atmosphere for nine hours to produce SF₆. The samples were purified cryogenically
and by gas chromatography and introduced by SF₆ line into a Thermo Electron MAT 253
mass spectrometer fitted with a dual inlet to measure ³²SF₅⁺, ³³SF₅⁺, ³⁴SF₅⁺, and ³⁶SF₅⁺. Sulfur
isotopic ratios are expressed on the V-CDT scale, on which the δ³⁴S, Δ³³S, and Δ³⁶S values of
the Ag₂S reference material, IAEA-S-1, are taken to be -0.3‰, 0.094‰, and -0.7‰,
respectively (Wing and Farquhar, 2015). The precision and accuracy of the bulk fluorination
system is evaluated by repeat analyses that return uncertainty (2SD) on δ³⁴S, Δ³³S, and Δ³⁶S
values as better than 0.15‰, 0.02‰ and 0.4‰, respectively. The Δ³³S and Δ³⁶S values are
calculated in the same manner as described in equation 2.

2.3 Electron Backscatter Diffraction Analysis (EBSD)
EBSD analysis of pentlandite to compare crystallographic orientation with multiple sulfur
isotope analysis was undertaken on a Tescan MIRA3 field emission SEM at the Microscopy
and Microanalysis Facility, John de Laeter Centre, Curtin University, Australia. Pentlandite
from massive sulfide ore was analysed within four pucks that were prepared into a mount as
described in the Supplementary Materials. Following standard petrographic diamond
polishing, the mount was polished chemically and mechanically to 60 nm using colloidal
silica in pH10 NaOH. Prior to analysis the sample was coated with a 5 nm carbon film.
EBSD data were acquired from a sample titled at 70° using an Oxford Instruments Aztec 3.0
system operating at 12 kV, a beam intensity of 16 nA and fixed working distance of 20 mm.
Pentlandite, pyrrhotite and magnetite orientation data were collected from each sample.
Crystallographic data required to create the theoretical match units, by which empirically
collected patterns are compared, were derived from Rajamani and Prewitt (1975) for
pentlandite, Wechsler et al. (1984) for magnetite and Alsen (1925) for pyrrhotite. EBSD data
were collected using the “mapping” mode of the Aztec EBSD software with a step size of 10
µm, a minimum of 8 bands and a Hough resolution of 60. In all cases, solutions gave mean
angular deviation (MAD) values of <1° and calculation intergrain orientations are
reproducible.

Post-processing of EBSD data was undertaken using Oxford Instruments Channel 5.12
software. Noise reduction protocols applied to the data were the Channel “wildspike”
correction and a 5 nearest neighbour zero solution algorithm. EBSD data are shown as
standard phase and inverse pole figure maps in the Supplementary Materials.

3. MATERIALS
Sulfides are inherently heterogeneous both chemically and mineralogically. Therefore,
identifying natural specimens that may be used as sulfur isotope reference material is a
difficult task whereby geological environment, pressure-temperature conditions, chemical
composition, and grain size need to be considered. In the case of selecting reference material
for pyrite and chalcopyrite, small pieces (~2 cm³) of much larger crystals from hydrothermal
deposits demonstrate a more reasonable degree of chemical and isotopic homogeneity over
small individual crystals that make up massive to sulfide-rich layers. For each of these
sulfides, although the total volume amount of reference material characterized is small (in
order to ascertain isotopic homogeneity), advancements in SIMS and LA-ICPMS capabilities
and procedures (i.e., new sample holder geometry in SIMS; Peres et al. 2012; and large
volume cells in LA-ICPMS) allow for the repeat use of the same reference material mounted
in standard blocks.

In the case of pyrrhotite and pentlandite, these two sulfide phases are commonly exsolved
from a monosulfide solid solution and typically have intra-grain associations in magmatic
environments (Durazzo and Taylor, 1982; Kelly and Vaughn, 1983). Furthermore, the
dehomogenizations of pyrrhotite (Fe_{0.1-0.5}S; x = 0.0–0.2) and pentlandite (Fe, Ni)_{0.8}S_{0.2}
are variable, reflective of conditions under which they formed. Pyrrhotite and pentlandite do not
commonly form large hydrothermally-derived crystals, and so characterized reference
material is derived from a small piece of a sulfide-rich and massive sulfide horizon,
respectively. Images of the reference materials are presented in the Supplementary Materials.
The effects of intra-grain and inter-grain textures on isotopic and chemical homogeneity of
pyrrhotite and pentlandite reference materials are assessed in sections 4.3 and 4.4. Additional
discussion on selecting reference materials is presented in section 5.1.

3.1 Pyrite – Sierra
Sierra is a 2 cm³ cube from a large 9 kg cube of pyrite sourced from a mine in the Cretaceous
stratigraphy of the Sonora region of Mexico. Although the exact provenance is unknown, the
area experienced widespread igneous activity during the late Cretaceous, resulting in the
formation of widespread porphyry copper mineralisation across the region (Barra and
Valencia, 2014). Sierra is mounted as 0.5 mm wide fragments of pyrite from the 2 cm³ cube
of Sierra for SIMS analysis.

3.2 Chalcopyrite – Nifty-b
The Nifty-b is a 2 cm³ piece of a larger 8 cm³ grain of chalcopyrite from the Nifty copper
deposit in the Proterozoic Paterson Orogen of Western Australia. Mineralization is hosted by
the Nifty stratigraphic member of the Yeneena Supergroup and occurs as hydrothermal
chalcopyrite-quartz-dolomite replacement of low grade shale (Anderson et al., 2001). Nifty-b
is mounted as 0.5 mm wide fragments of chalcopyrite pyrite from the 2 cm³ piece of Nifty-b
for SIMS analysis.

3.3 Pyrrhotite – Alexo
The Alexo monoclinic pyrrhotite forms subhedral 0.1–2 mm grains within a matrix of
peridotite. It is from the nickel-sulfide Alexo deposit within the ca. 2.7 Ga Abitibi granite-
greenstone belt of the Superior Craton, Canada. The sample comprises magmatic pyrrhotite
grains composed of disseminated to net-textured sulfides at the contact between olivine
cumulate komatiite rocks of the Munro Group and footwall andesitic rocks of the Hunter
Mine Group (Naldrett, 1966). The local metamorphic grade at Alexo is prehnite-pumpellyite-
facies. Based on stratigraphic associations, the timing of the nickel-sulfide mineralization at
Alexo is assumed to have occurred at ca. 2.7 Ga (Fyon and Green, 1991), which is coeval with other world-class komatiite-hosted nickel-sulfide systems worldwide (Barnes et al., 2013; Fiorentini et al., 2011; 2010). The Alexo sample was first isotopically defined by Bekker et al. (2009). A 2 cm x 3 cm x 0.5 cm pyrrhotite-rich layer of peridotite was trimmed from a larger hand sample. Alexo is mounted as 1 mm diameter pucks from this sample for SIMS analysis.

3.4 Pentlandite – VMSO

The VMSO pentlandite is from an amphibolite-facies massive sulfide lens from the Victor South shoot in the komatiite-hosted nickel-sulfide Long-Victor deposit of the Kambalda camp. The Kambalda camp is situated in the ca. 2.7 Ga Kalgoorlie Terrane in the Yilgarn Craton of Western Australia (Barnes et al., 2013 and references therein). At Kambalda, nickel-sulfide deposits are associated with thick channelized komatiite flow units in stratigraphic contact with a thick pile of pillowed and massive tholeiitic basalts (Lunnon Basalt Formation), locally overlain by sulfidic metasedimentary rocks (e.g., Lesher and Groves, 1986). Subsequent folding has rotated the komatiitic channels from 30° to a vertical dip to the east and 10° plunge to the south as well as resulting in the localised remobilisation of some of the original sulfides into new structurally controlled positions (Stone et al., 2005).

The VMSO sample was first isotopically defined by Bekker et al. (2009). It comprises 0.1–0.5 mm magmatic pentlandite grains interlayered at the cm-scale with pyrrhotite and pyrite from the basal sulfide layer of the Silver Lake Member of the Kambalda Formation. A 1 cm wide pentlandite rich layer trimmed from the sample. From this layer, VMSO is mounted as 1 mm diameter pucks from the host massive sulfide lens for SIMS analysis.

4. RESULTS

The following section presents chemical and isotopic results pertaining to standard development work for four new sulfide reference materials: pyrite (Sierra), chalcopyrite (Nifty-b), pyrrhotite (Alexo), and pentlandite (VMSO). A summary of mineral chemical composition collected by WDS for each reference material is presented in Table 1. The main sulfide forming elements and common trace elements are presented in the Supplementary Materials. Figure 2 shows select elemental maps (by WDS) of reference material.

**Table 1**: Summary of the chemical composition determined by wavelength dispersive spectrometry. Uncertainty is two standard deviations (2SD).

<table>
<thead>
<tr>
<th>Standard</th>
<th>Mineral</th>
<th>Formula</th>
<th>Chemical composition (wt.%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>FeS₂</td>
<td>Fe (2SD)</td>
</tr>
<tr>
<td>Sierra</td>
<td>Pyrite</td>
<td>FeS₂</td>
<td>53.3 (0.5)</td>
</tr>
<tr>
<td>Nifty-b</td>
<td>Chalcopyrite</td>
<td>CuFeS₂</td>
<td>34.5 (1.4)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>--------</td>
<td>--------</td>
<td>--------</td>
<td>--------</td>
</tr>
<tr>
<td>Alexo</td>
<td>Pyrrhotite</td>
<td>Fe$_{0.90}$S</td>
<td>60.5 (1.2)</td>
</tr>
<tr>
<td>VMSO</td>
<td>Pentlandite</td>
<td>Fe$<em>{4.1}$Ni$</em>{4.8}$Co$<em>{0.1}$S$</em>{8}$</td>
<td>29.4 (0.8)</td>
</tr>
</tbody>
</table>

**Figure 2:** Compositional maps of: A) Sierra pyrite (Fe wt.%), B) Nifty-b chalcopyrite (Cu wt.%), C) Alexo pyrrhotite (Fe wt.%), D) VMSO pentlandite (Ni wt.%). Wavelength dispersive spectrometry maps are collected by EPMA.

Table 2 shows the accumulated SIMS isotopic data to define analytical repeatability of the four reference materials. Uncertainty reported in Table 2 and the text is twice the standard deviation of the mean. A compilation of in situ sulfur isotopic measurements for reference materials and unknown samples is presented in Supplementary Material.

**Table 2:** In situ SIMS sample repeatability for four presented reference materials and their corresponding uncertainty (reported as twice the standard deviation of the mean). “$n$” is the number of measurements.
Table 3 and text presents the weighted mean bulk sulfur isotope compositions ($\delta^{33}S_{\text{V-CDT}}$, $\delta^{34}S_{\text{V-CDT}}$, $\delta^{36}S_{\text{V-CDT}}$) measured by fluorination gas-source mass spectrometry (from multiple aliquots of each reference material) and associated uncertainty, defined as two standard deviations of the mean. All sulfur isotope results presented in the text are reported on the V-CDT scale. Figure 3 demonstrates $\delta^{34}S$ isotope homogeneity of the reference materials. To demonstrate that the reference material was homogeneous within the analytical uncertainty of the fluorination gas-source mass spectrometry, an MSWD test was completed as detailed in Wing and Farquhar (2015). Briefly, a population was considered homogeneous if all values within a given sample population were consistent with a single mean value and returned within a 95% confidence interval. Because the analytical error is well defined, MSWD is used to estimate an independent “goodness of fit” to demonstrate separately that the reference material is homogeneous and consistent within a single mean value. Bulk fluorination data for four new reference materials are presented in the Supplementary Material.

Table 3: Weighted mean bulk sulfur isotope compositions ($\delta^{33}S_{\text{V-CDT}}$, $\delta^{34}S_{\text{V-CDT}}$, $\delta^{36}S_{\text{V-CDT}}$, $\Delta^{33}S_{\text{V-CDT}}$, $\Delta^{36}S_{\text{V-CDT}}$) as measured by fluorination gas-source mass spectrometry. 2SD is two standard deviations of the mean. SE is the standard error associated with the mean, and MSWD is an independent goodness-of-fit check after Wing and Farquhar (2015). “n” is the number of independent extractions and measurements of separate aliquots.

<table>
<thead>
<tr>
<th>Ref.</th>
<th>n</th>
<th>$\delta^{33}S$ (2SD)</th>
<th>1SE</th>
<th>MSWD</th>
<th>$\delta^{34}S$ (2SD)</th>
<th>1SE</th>
<th>MSWD</th>
<th>$\delta^{36}S$ (2SD)</th>
<th>1SE</th>
<th>MSWD</th>
<th>$\Delta^{33}S$ (2SD)</th>
<th>1SE</th>
<th>MSWD</th>
<th>$\Delta^{36}S$ (2SD)</th>
<th>1SE</th>
<th>MSWD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sierra (py)</td>
<td>12</td>
<td>1.09</td>
<td>0.15</td>
<td>0.02</td>
<td>0.99</td>
<td>2.17</td>
<td>0.28</td>
<td>0.04</td>
<td>0.89</td>
<td>3.96</td>
<td>0.60</td>
<td>0.09</td>
<td>1.00</td>
<td>-0.02</td>
<td>(0.01)</td>
<td>-0.18</td>
</tr>
<tr>
<td>Nifty-b (ccp)</td>
<td>5</td>
<td>-1.78</td>
<td>0.21</td>
<td>0.03</td>
<td>2.08</td>
<td>-3.58</td>
<td>0.44</td>
<td>0.07</td>
<td>2.22</td>
<td>-7.15</td>
<td>0.63</td>
<td>0.13</td>
<td>1.11</td>
<td>-0.06</td>
<td>(0.03)</td>
<td>-0.36</td>
</tr>
<tr>
<td>Alexa (po)</td>
<td>9</td>
<td>1.73</td>
<td>0.20</td>
<td>0.03</td>
<td>1.87</td>
<td>5.23</td>
<td>0.40</td>
<td>0.05</td>
<td>1.77</td>
<td>10.98</td>
<td>0.59</td>
<td>0.10</td>
<td>0.95</td>
<td>-0.96</td>
<td>(0.04)</td>
<td>1.02</td>
</tr>
<tr>
<td>VMSO (pn)</td>
<td>11</td>
<td>1.66</td>
<td>0.24</td>
<td>0.02</td>
<td>2.67</td>
<td>3.22</td>
<td>0.51</td>
<td>0.05</td>
<td>2.89</td>
<td>6.37</td>
<td>0.83</td>
<td>0.09</td>
<td>1.92</td>
<td>0.00</td>
<td>(0.02)</td>
<td>0.24</td>
</tr>
</tbody>
</table>
Figure 3: Probability density plots for $\delta^{34}$S for normalized reference material to demonstrate the low degree of uncertainty, normal distribution and lack of outliers.

4.1 Pyrite – Sierra

The chemical composition of Sierra, determined by EPMA ($n=78$ from five fragments), reveals that it is composed of $46.7 \pm 0.5$ wt.% Fe and $53.3 \pm 0.4$ wt.% S, with trace amount of Co (~400 ppm). WDS mapping further demonstrates that Sierra grain fragments are chemically homogeneous (Figure 2a) and free of inclusions. Twelve different fragments of the 2 cm$^3$ piece known as Sierra were analysed by fluorination gas-source mass spectrometry and yielded weighted averaged values of: $\delta^{33}$S = 1.09 $\pm$ 0.15‰, $\delta^{34}$S = 2.17 $\pm$ 0.28‰, and $\delta^{36}$S = 3.96 $\pm$ 0.60‰. Values for $\Delta^{33}$S and $\Delta^{36}$S are calculated as -0.02 $\pm$ 0.01‰ and 0.18 $\pm$ 0.15‰, respectively, indicating that Sierra does not contain a MIF signature. Sierra has been measured as a primary reference for triple sulfur isotopes 1417 times amongst which 861 times included quadruple sulfur isotopes over 59 analytical sessions (see Figure 3). Reproducibility of SIMS analyses on Sierra was $\delta^{33}$S = 0.15‰, $\delta^{34}$S = 0.24‰, and $\delta^{36}$S = 1.04‰. Reproducibility on $\Delta^{33}$S = 0.08‰ and $\Delta^{36}$S = 0.77‰. Different fragments of the much larger pyrite, known as ‘Sonora’, have been described by Farquhar et al. (2013), Evans et al. (2014) and Wacey et al. (2011). The Sonora fragments have slightly lighter values of $\delta^{33}$S (0.83‰), $\delta^{34}$S (1.61‰), and $\delta^{36}$S (3.25‰) than those we determine here for Sierra.

4.2 Chalcopyrite – Nifty-b
WDS analyses \((n = 352)\) reveal that Nifty-b chalcopyrite is composed of 34.5 ± 1.4 wt.% Cu, 29.8 ± 1.2 wt.% Fe, and 34.7 ± 0.5 wt.% S, with trace amounts of Zn (~300 ppm). WDS mapping demonstrates that Nifty-b is homogeneous and inclusion-free but contains fractures that can be avoided during isotope measurements (Figure 2b). Five different grain fragments of Nifty-b were analysed by fluorination gas-source mass spectrometry and yielded a weighted average of: \(\delta^{33}S = -1.78 \pm 0.21\‰\), \(\delta^{34}S = -3.58 \pm 0.44\‰\), and \(\delta^{36}S = -7.15 \pm 0.63\‰\). Values for \(\Delta^{33}S\) and \(\Delta^{34}S\) are calculated as 0.06 ± 0.03‰ and 0.36 ± 0.50‰, respectively (the large error on \(\Delta^{36}S\) prevents observation of MIF in this isotopic system). Nifty-b has a small but consistent MIF signature. Nifty-b has been measured as a primary chalcopyrite reference for triple sulfur isotopes 149 times, and for quadruple sulfur isotopes 87 times in 15 analytical sessions over the course of three years. Average measurement repeatability of SIMS analyses on Nifty-b was \(\delta^{33}S = 0.14\‰\), \(\delta^{34}S = 0.23\‰\), and \(\delta^{36}S = 0.63\‰\). Reproducibility on \(\Delta^{33}S\) = 0.08‰ and \(\Delta^{36}S\) = 0.48‰.

4.3 Pyrrhotite – Alexo

Alexo pyrrhotite has very fine exsolution lamellae, <10 µm in width, which are heterogeneously dispersed. WDS analyses of Alexo \((n = 61)\) reveal that it is composed of 60.5 ± 1.2 wt.% Fe and 38.6 ± 0.4 wt.% S. Exsolution lamellae are composed of cobalt-bearing pentlandite with a composition of 32.3 ± 1.7 wt.% Fe, 32.3 ± 1.5 wt.% Ni, 1.2 ± 0.1 wt.% Co, and 33.1 ± 0.6 wt.% S \((n = 23)\). These pentlandite lamellae are very fine and WDS mapping demonstrates that they are unevenly distributed throughout the grain (see Figure 2c), making them difficult to observe during SIMS analysis. Alexo pyrrhotite contains inclusions of magnetite (50 µm) and conjugate fracture sets; both features can be readily avoided during SIMS analysis.

Nine different grains of Alexo were analysed by fluorination gas-source mass spectrometry and yielded a weighted average of: \(\delta^{33}S = 1.73 \pm 0.20\‰\), \(\delta^{34}S = 5.23 \pm 0.40\‰\), and \(\delta^{36}S = 10.98 \pm 0.59\‰\). Uncertainty is reported as the standard error of the mean. Values for \(\Delta^{33}S\) and \(\Delta^{34}S\) are calculated as -0.96 ± 0.04‰ and 1.02 ± 0.27‰, respectively, indicating that Alexo has a MIF signature. The values represent the bulk composition of Alexo, incorporating both pyrrhotite and the pentlandite exsolution lamellae. Alexo has been measured as a primary pyrrhotite reference for triple sulfur isotopes 340 times, and for quadruple sulfur isotopes 199 times in 27 analytical sessions (see Figure 5). Reproducibility of SIMS analyses on Alexo was \(\delta^{33}S = 0.17\‰\), \(\delta^{34}S = 0.30\‰\), and \(\delta^{36}S = 0.74\‰\). Reproducibility on \(\Delta^{33}S\) = 0.11‰ and \(\Delta^{36}S\) = 0.52‰.

4.3.1 The effect of pentlandite exsolution in Alexo pyrrhotite

Unmixing of the solid solution pyrrhotite-pentlandite (Fe,Ni)\(_{1-x}\)S at temperatures below ~600 ºC via exsolution of pentlandite is a common and ubiquitous petrochemical feature of magmatic nickel-sulfide ore deposits (e.g., Kelly and Vaughn, 1983; Naldrett et al., 1967). To unravel and decipher the sulfur isotopic record hosted by pyrrhotite grains from magmatic deposits, it is important to understand the effect of potential isotope fractionation between the two phases of the Alexo pyrrhotite reference material (also derived from a magmatic ore deposit). The sulfur isotopic composition of Alexo, determined by fluorination gas-source
mass spectrometry, itself incorporates a small amount (1–5%) of pentlandite exsolution. This investigation is crucial because: 1) lamellae may be fast pathways of isotopic exchange in an open system, 2) instrumental mass fractionation may be affected by the incorporation of more than one phase (see Hervig et al., 2002), and 3) natural isotopic fractionation of pentlandite-pyrrhotite may be reflected in the bulk analysis and not represented in the SIMS analysis (when avoiding exsolution lamellae).

Because lamellae are so fine and difficult to observe even with adequate BSE imaging, it is impossible to solely analyse exsolution lamellae (Figure 4). Therefore, to evaluate the effect on sulfur isotope composition of the presence of exsolution lamellae within Alexo pyrrhotite on sulfur isotope composition, 60 in situ analyses were carried out in two groupings: 1) 30 analyses of areas with an abundance of exsolution lamellae, and 2) 30 analyses of areas free from exsolution lamellae. Alexo grains were reimaged after SIMS work to identify which analyses incorporated some component of exsolution lamellae (Figure 9). Group one yields average δ³³S, δ³⁴S, δ³⁶S values equal to 1.75 ± 0.20‰, 5.27 ± 0.43‰, and 11.12 ± 0.98‰, respectively. Group two yields average δ³³S, δ³⁴S, δ³⁶S values equal to 1.70 ± 0.22‰, 5.18 ± 0.44‰, and 10.83 ± 0.98‰, respectively. Analyses for the entire run were normalised to the bulk value of Alexo, a value that incorporates some component of pentlandite exsolution, but that returns a low MSWD (1.8) across nine separate grain extractions and analyses. These results demonstrate that exsolution of pentlandite within Alexo pyrrhotite are minimal enough to be masked by the measurement repeatability and so do not impact sulfur isotope measurements. Therefore, Alexo is an adequate reference material for the determination of sulfur isotope composition of pyrrhotite.
4.4 Pentlandite – VMSO

WDS analyses demonstrate that VMSO pentlandite is composed of 29.4 ± 0.8 wt.% Fe, 35.8 ± 0.9 wt.% Ni, 0.3 ± 0.1 wt.% Co, and 32.9 ± 0.4 wt.% S (n = 45), with trace amounts of Sb (~150 ppm), Se (~600 ppm) and Cu (~500 ppm). VMSO pentlandite contains small (~100 µm) inclusions of magnetite and is intergrown with pyrrhotite and pyrite. WDS mapping demonstrates that the pentlandite portion of VMSO is homogeneous but that fractures and sulfide intergrowths should be avoided (Figure 4d). Analysis of the intergrowths reveal compositions of 59.5 ± 0.6 wt.% Fe and 39.3 ± 0.1 wt.% S, for pyrrhotite (n = 13) and 45.6 ± 0.2 wt.% Fe, 0.9 ± 0.0 wt.% Co, and 53.5 ± 0.2 wt.% S, for pyrite (n = 11). The pyrrhotite and pyrite components and oxide inclusions are readily identified in BSE imaging and can be easily avoided during SIMS analysis.

Eleven different grains of the pentlandite portion of VMSO were analysed by fluorination gas-source mass spectrometry and yielded a weighted average of: δ33S = 1.66 ± 0.24‰, δ34S = 3.22 ± 0.51‰, and δ36S = 6.37 ± 0.83‰. Uncertainty is reported as the standard error of the mean. Values for Δ33S and Δ36S are calculated as 0.00 ± 0.02‰ and 0.24 ± 0.35‰, respectively, indicating that VMSO does not have a MIF signature. The elevated MSWD on sulfur isotope measurements of VMSO is discussed in section 5.1.3. VMSO has been measured as a primary pentlandite reference for triple sulfur isotopes 255 times, and for quadruple sulfur isotopes 98 times in 22 analytical sessions (see Figure 5). Reproducibility of SIMS analysis on VMSO was δ33S = 0.21‰, δ34S = 0.33‰, and δ36S = 0.90‰. Reproducibility on Δ33S = 0.12‰ and Δ36S = 0.72‰.

4.4.1 Investigation into the homogeneity of VMSO pentlandite

Bulk measurements of VMSO pentlandite yield larger uncertainties and greater MSWD values (δ33S = 1.66 ± 0.23‰ (MSWD 2.67), δ34S = 3.22 ± 0.51‰ (MSWD 2.89), and δ36S = 6.37 ± 0.83‰ (MSWD 1.92)) than other sulfide reference material, indicating that it is a more isotopically heterogeneous material. Pentlandite does not exist naturally as large single crystals; therefore, this reference material has a certain level of isotopic heterogeneity attributed to micro-scale intergrowths of pentlandite and pyrrhotite within the reference material, a feature typical of sulfide ore in magmatic deposits. Although these two sulfide phases are cogenetic, slightly different isotopic compositions between the two phases are expected due to natural equilibrium isotopic fractionation. The <nm-scale intergrowths are large enough to be easily avoidable during SIMS analysis; however, the incorporation of a small component of pyrrhotite during extraction of (relatively) large volumes of sulfur for bulk analysis is unavoidable (see pyrrhotite-pentlandite intergrowth in Figure 2d).

To evaluate the sulfur isotope composition of the pyrrhotite portion of VMSO, seven in situ SIMS analyses of the pyrrhotite intergrowths were normalized using Alexo pyrrhotite. Sulfur isotope analysis of pyrrhotite intergrowths yielded values of δ33S = 2.07 ± 0.26‰, δ34S =
3.86 ± 0.18‰, and δ^{36}S = 8.09 ± 0.26‰. These values are slightly elevated to the determined isotopic composition of VMSO pentlandite but within the envelope of uncertainty. Therefore, the uncertainty on the bulk values mask the incorporation of a component of pyrrhotite during the extraction process. We determine VMSO reference material to be adequate for in situ sulfur isotope analysis of pentlandite but to be used with the expectation of propagating poorer analytical precision.

5. DISCUSSION

Mineral systems are natural laboratories that provide invaluable information on the flux of metals and fluids that are cycled among the different reservoirs of our planet. Most of this information is stored as chemical and isotopic signatures, which are preserved in the magmatic and hydrothermal sulfide phases that comprise the mineralization hosted in different mineral systems. However, the interpretation of the information stored in sulfides requires care, as the mineroc-chemical features of sulfides are prone to be reset over a wide range of physical conditions. The resulting complexity recorded in the chemical and isotopic nature of any given sulfide assemblage may be cryptic to bulk rock analysis, but can be resolved with an in situ analytical approach, which allows the investigator to carefully select specific grains or crystal fractions and avoid others.

5.1 Selecting and validating sulfide reference material

In the case of selecting reference material for pyrite and chalcopyrite, small pieces (~2 cm³) of much larger crystals from low grade hydrothermal deposits yield a much higher degree of chemical and isotopic homogeneity over small individual crystals that make up sulfide-rich layers. Here, we demonstrate this fact by comparing the Sierra standard (a fragment of a large hydrothermal crystal) to other available pyrite sulfur isotope standards that occur as small individual grains within a stratigraphic horizon.

Grains of Isua 248474 pyrite (kindly supplied by Martin Whitehouse) from the Isua greenstone belt, SW Greenland (Whitehouse, 2013) were used as secondary standards to assess the adequacy of Sierra as a reference material. Isua 248474 has reported bulk sulfur isotope values of δ^{33}S = 4.33 ± 0.24‰ (1σ), δ^{34}S = 1.99 ± 0.18‰ (1σ) (Baublys et al., 2004). A total of 216 SIMS analyses (in 19 analytical sessions) have been completed on Isua 248474 pyrite as an unknown using Sierra as the primary reference material and yield an average of δ^{33}S = 4.45 ± 0.66‰, δ^{34}S = 2.60 ± 0.86‰, and δ^{36}S = 2.74 ± 1.60‰ and a calculated MIF signature: Δ^{33}S = 3.11 ± 0.33‰ and Δ^{36}S = -2.38 ± 0.81‰. Variability within the Isua 248474 pyrite standard has been observed by Whitehouse (2013) who reports some dispersion to lower values for δ^{34}S but yields an average δ^{34}S that is higher but within error of the accepted bulk measurement of Baublys et al. (2004). Here, our Isua 248474 measurements normalised to Sierra have a high uncertainty, consistent with the isotopic variability observed by Whitehouse (2013).

A grain of UW Balmat pyrite (kindly supplied by John Craven) from the Adirondack Mountains, New York was also assessed. UW Balmat pyrite was originally described by Crowe et al. (1990) who reported δ^{34}S equal to 14.63 ± 0.38‰ (2SD). Further to that,
Graham and Valley (1992) describe the UW Balmat pyrite as only being isotopically homogeneous at the ~1.0‰ level. Pyrite from a slightly different stratigraphic section of the deposit (UWPy-1) does not show any MIF anomaly (Williford et al., 2011). Two analytical sessions for triple isotopes give 31 analyses of UW Balmat pyrite with an average δ34S of 16.23 ± 0.18‰ (2SD). Our values for δ34S equal 8.30 ± 0.12‰ and yield a Δ33S of -0.03‰, confirming the lack of a MIF signature. We attribute the variability in both the Isua 248474 and UW Balmat pyrite standards to being separate grains distributed between laboratories.

In the development of a chalcopyrite standard, four separate materials were tested for isotopic homogeneity. Two materials were fragments of large hydrothermal chalcopyrite crystals which yielded a much higher degree of δ34S reproducibility than the two other materials derived from massive sulfide horizons in magmatic systems. The pyrite and chalcopyrite results demonstrate that small fragments of large hydrothermal sulfide crystals are more isotopically homogeneous and more suitable standards than individual sulfide grains in sulfide-rich layers.

In the case of pyrrhotite and pentlandite, these minerals do not commonly occur as large hydrothermal crystals and so the most homogeneous reference material were selected from massive sulfide layers from magmatic ore deposits. Preliminary investigation into how chemical heterogeneity (i.e., micro-exsolution and intergrowths between the two phases) affects isotopic homogeneity in sections 4.3 and 4.4, demonstrate that Alexo and VMSO reference materials are suitable for the determination of sulfur isotope composition of pyrrhotite and pentlandite and ideal for the investigation of magmatic deposits (in which unknown sulfides will have similar textures). However, for comparison, further effort to develop a hydrothermal pyrrhotite standard is warranted.

We have presented chemical and isotopic compositions for a suite of chemically and isotopically homogeneous sulfide reference materials that are representative of the sulfide mineralogy of magmatic and hydrothermal mineral systems. Figure 5 presents plots of: 1) δ33S_{V,CDT} vs. δ34S_{V,CDT}, and 2) Δ33S vs. δ34S_{V,CDT} for all reference material analysed in this study. Two of the four sulfide standards (pyrrhotite and chalcopyrite) contain a MIF component, a feature that is recommended to confidently assess the accuracy of Δ33S and Δ36S measurements (Whitehouse, 2013). SIMS measurements on the sulfur isotope dataset demonstrate that the reference materials are adequately isotopically homogeneous and return repeatability on the order of uncertainty on the bulk measurements. Furthermore, bulk measurements deem the reference material homogeneous by the MSWD test as detailed in Wing and Farquhar (2015).
Figure 5: Plots of: (a) $\delta^{33}\text{S}_{\text{CDT}}$ vs. $\delta^{34}\text{S}_{\text{CDT}}$ and (b) $\Delta^{33}\text{S}_{\text{CDT}}$ vs. $\delta^{34}\text{S}_{\text{CDT}}$ for all sulfide reference material (Sierra pyrite, Nifty-b chalcopyrite, Alexo pyrrhotite, and VMSO pentlandite) measured by SIMS in this study. Error crosses are the overall uncertainty (at the 2SD level) based on propagating the individual analysis repeatability with the reproducibility for the relevant analytical session. The Sierra pyrite is used as the primary reference for measuring previously characterised pyrite reference material Isua 248474 and UW Balmat pyrite (see Whitehouse, 2013; Graham and Valley, 1992).

5.2 In situ multiple sulfur isotope analysis of a nickel-sulfide deposit, Kambalda

The in situ SIMS multiple sulfur isotope data from this study are used to constrain the nature of the ore-forming process in the Moran shoot, within the Kambalda camp of Western Australia. The results are compared with existing data from the Victor South shoot along the Victor channel, where the VMSO reference material was collected and analysed, in order to reflect on the sulfur isotope architecture of the various channels exposed in the Long-Victor deposit. The integrated dataset also permits discussion of the large-scale setting of the Kambalda camp in relation to other world-class nickel-sulfide deposits in the Kalgoorlie
Terrane of the Yilgarn Craton, which hosts the largest resources and reserves of nickel-sulfide mineralisation associated with komatitites (Barnes and Fiorentini, 2012). We utilise the case study to validate the presented reference material, assess the natural fractionation factor of $\delta^{34}$S between pyrrhotite and pentlandite, and assess whether crystallographic orientation of pentlandite induces biases on $\delta^{34}$S and $\Delta^{33}$S measurements.

5.2.1 Moran shoot, Long-Victor deposit, Kambalda camp

The case study focussed on the identification of the multiple sulfur isotope signature of the major magmatic sulfide phases (pyrrhotite, pentlandite and chalcopyrite) hosted in the massive sulfide layer at the base of the Moran shoot at Kambalda, Western Australia. The spatial and genetic relationship between the Victor South shoot, where VMSO reference material was collected, and the Moran shoot, where the sample for the case study was collected, is discussed in Barnes et al. (2013) and presented in Figure 6. Briefly, the Moran shoot is hosted in the same channelized structure that hosts the larger Long shoot. This channel runs roughly parallel to the Victor channel in which the Victor South shoot is hosted. The Moran shoot attains 500 m along strike but is open to the south, where the boundaries of the mineralised body are not defined.

Figure 6: Location of the Victor-Long nickel-sulfide deposit of the Kambalda camp, Western Australia after Bekker et al. (2009) and references therein. The Victor-Long deposit is host to the Victor South and Moran shoots, discussed in this study.
The ore body largely comprises massive and matrix sulfides dominantly composed of pentlandite (38%) and pyrrhotite (58%), with minor (4%) chalcopyrite (Figure 7). Pyrrhotite and pentlandite are intergrown and are in equilibrium. Chalcopyrite forms small (<40 µm) interstitial grains in equilibrium with the other sulfides at pentlandite-pyrrhotite boundaries. Multiple sulfur isotope data were acquired from four 5 mm in diameter pucks of ore and values for δ\(^{34}\)S, Δ\(^{33}\)S, Δ\(^{36}\)S are presented in Table 4. WDS data were also acquired from the pentlandite and pyrrhotite to ensure similar chemical composition to the reference material.

Figure 7: A) Hand sample of the Moran Shoot massive sulfide horizon of the Long-Victor deposit, Kambalda camp. Numbers demarcate localities drilled for bulk analysis. B) Micro-XRF map of slab of nickel-sulfide ore from the Moran shoot, Long-Victor deposit, Kambalda camp. Massive sulfide is composed dominantly of pentlandite (light blue) and pyrrhotite (dark blue) with minor chalcopyrite (green). The proportion of sulfides is calculated from the Micro-XRF map to be 38% pentlandite (pn), 58% pyrrhotite (po), and 4% chalcopyrite ( ccp). C) EDS phase map showing distribution of NiS (pentlandite; yellow), FeS (pyrrhotite; red), and FeO (iron oxide; green). D) EDS Cu map showing chalcopyrite in green. E) Reflected light photomicrograph of pentlandite (yellow)-pyrrhotite (grey) association with small interstitial grains of chalcopyrite (orange). Area delineated by black box in C. Ion probe pits are small black boxes and are 15 µm in length.

Table 4: Multiple sulfur isotope results (δ\(^{34}\)S\(_{\text{V-CDT}}\), Δ\(^{33}\)S\(_{\text{V-CDT}}\), Δ\(^{36}\)S\(_{\text{V-CDT}}\)) acquired by SIMS using reference material presented in this manuscript to correct for instrumental mass fractionation. “n” is the number of measurements.

<table>
<thead>
<tr>
<th>Chemical composition</th>
<th>Sulfur isotope composition (%)</th>
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<tbody>
<tr>
<td></td>
<td>δ(^{34})S</td>
</tr>
<tr>
<td>pentlandite</td>
<td>Fe(<em>{4.1})Ni(</em>{4.8})Co(_{0.1})S(_8)</td>
</tr>
<tr>
<td>pyrrhotite</td>
<td>Fe(_{0.90})S</td>
</tr>
<tr>
<td>chalcopyrite</td>
<td>CuFeS(_2)</td>
</tr>
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</table>

The in situ SIMS data presented in Table 4 compare well with 10 aliquots of bulk sulfide material from the same ore shoot, which returned δ\(^{34}\)S\(_{\text{V-CDT}}\), Δ\(^{33}\)S\(_{\text{V-CDT}}\), Δ\(^{36}\)S\(_{\text{V-CDT}}\) equal to
3.10 ± 0.70‰ (2SD), 0.24 ± 0.04‰ (2SD), and -0.21 ± 0.10‰ (2SD). The large uncertainty on δ34S in the bulk data demonstrates that the yielded sulfur isotope results represent an average isotopic composition of the three sulfide phases (as depicted by the SIMS δ34S results).

The case study is aimed at emphasizing the significance of a multi-sulfide phase in situ analytical approach in the determination of the multiple sulfur isotope signature of Archean komatiite-hosted nickel-sulfide deposits. Results indicate that the magmatic sulfide assemblage of the massive sulfide horizon hosted in the Moran shoot of the Long-Victor deposit in the Kambalda region of Western Australia comprises pentlandite, pyrrhotite and chalcopyrite. These phases are in isotopic equilibrium and display similar values for Δ33S = ~+0.2‰ and Δ34S = ~−0.02‰. However, the three sulfide phases show variable δ34S values (δ34S_{pentlandite} = ~−2.9‰, δ34S_{chalcopyrite} = ~3.1‰, and δ34S_{pyrrhotite} = ~3.9‰), which are indicative of natural isotope fractionation. Natural δ34S fractionation among the sulfides that make up a multi-sulfide phase ore assemblage cannot be easily resolved through bulk rock analysis (as presented in Figure 8). This in turn may lead to misinterpretations of subtle sulfur isotope variations. In situ analysis also has the potential to highlight the presence of secondary sulfides. Therefore, the in situ approach provides the user the ability to resolve ambiguities that may arise through whole-rock isotope analysis.

![Figure 8: Probability density plots (bin size 10) of δ34S and Δ33S pyrrhotite, pentlandite, and chalcopyrite results for the Moran shoot of the Long-Victor deposit, Kambalda. A representative subset of pentlandite results are chosen. All three sulfide phases yield the same Δ33S values (+0.2‰) demonstrating they are likely cogenetic and from the same source. The variable δ34S are a result of natural isotopic fractionation between the three sulfide phases (of which pentlandite and pyrrhotite compose 96% of the ore body). Variability in δ34S chalcopyrite is a product of few analyses (13) of small interstitial grains. Whole rock bulk fluorination values (from 10 aliquots) depicted by dashed line and uncertainty (2SD) by grey box.](image)

5.2.2 Source of sulfur to trigger sulfide saturation
The multiple sulfur isotope data collected from the Moran shoot is assessed in conjunction with the multiple sulfur isotope signature of VMSO, which was collected from the Victor South shoot. Bekker et al. (2009) originally measured the multiple sulfur isotope signature of the ore sample from which the VMSO pentlandite standard was made. The bulk rock fluorination data indicated that the massive sulfide in the Victor South shoot did not display any evidence of MIF. This result was not discussed in detail in Bekker et al. (2009), but it clearly portrayed Kambalda as being radically different from the giant komatiite-hosted deposits in the northern part of the Kalgoorlie Terrane, which display a very pronounced negative MIF signature (up to -0.7‰). The bulk rock isotope signature of the VMSO ore sample is reflected in the lack of MIF recorded in the VMSO pentlandite standard presented in this study.

In the Long-Victor deposit, the relationship between the Victor South and Moran shoots is described in Barnes et al. (2013). For the sake of the argument here, the most important message to be taken away from the study of Barnes et al. (2013) is that different shoots, which originated from the emplacement of komatiite magma along spatially close but distinct volcanic channels, may have resulted from the equilibration of variable proportions of sulfide and silicate liquids. Evidence of this variable proportion, which is expressed as R Factor (Campbell and Naldrett, 1979), is not only recorded in the highly variable metal tenor of the ore shoots and associated variable degrees of chalcophile element enrichment and depletion in the host komatiite channelized units, but it is also preserved in the sulfur isotopic architecture of the deposit, as described in Lescher and Burnham (2001) and Fiorentini et al. (2012a). This should serve as a word of caution for any exploration targeting tool based on sulfur isotope analysis to track crustal assimilation as a proxy for magma fertility.

Evidence of a positive MIF anomaly (Δ^{33}S = 0.2‰) recorded in three sulfide phases of the Moran shoot suggests thermo-mechanical assimilation of sulfidic shales (Δ^{33}S = up to 0.8‰; Bekker et al., 2009) during komatiite emplacement. The magnitude of MIF as expressed by Δ^{33}S is small and so to relate the sulfur isotope signature to photochemical reactions related to an oxygen-poor atmosphere (rather than mass dependent processes as described by Johnston, 2011), we evaluate the Δ^{36}S/Δ^{33}S relationship. The Δ^{36}S/Δ^{33}S ratio ranges from −0.04 to −0.88, extending the known Archean ratio beyond −1 to −2 of Johnston (2011). The ratio is significantly different to the predicted range of mass dependent fractionation processes (Δ^{36}S/Δ^{33}S = −9 to −7; Ono et al., 2006b; Johnston et al., 2007).

The lack of any MIF anomaly in the Victor South shoot suggests higher R factor values for the channel, which was effectively flushed with higher volumes of mantle-derived silicate magma carrying no MIF anomaly, thus diluting any evidence of crustal MIF. However, it is crucial to state that in magmatic systems, near-zero δ^{34}S or Δ^{33}S values permit a mantle source (δ^{34}S = −1.28 ± 0.33‰; Δ^{33}S = 0.00‰; Labidi et al., 2013) but do not prove it if the full complexity of homogenization processes is considered. However, non-zero δ^{34}S or Δ^{33}S values prove a non-mantle source and can be further confirmed by the Δ^{36}S/Δ^{33}S ratio (Farquhar et al., 2007; Johnston, 2011). The data from two different shoots at Kambalda highlight the variation that characterizes ore-forming processes in magmatic systems. This
variation is translated downstream in the complexity of the interpretation of chemical and isotopic data that are used as proxies to target mineralisation in exploration (Fiorentini et al., 2012a).

In addition to the deposit-scale observations discussed above, the in situ multiple sulfur isotope data presented in this study also provide key insights into the geodynamic setting where the Kambalda komatiites were emplaced. Unlike the giant dunite-hosted komatiite systems that thermo-mechanically assimilated volcanogenic massive sulfides proximal to vents in the Agnew-Wiluna greenstone belt (cf. Fiorentini et al., 2012b), the integrated results from both the Moran (case study) and Victor South (VMSO sample) shoots indicate that Kambalda ores assimilated abyssal sulfidic shales. This inference is based on the very different MIF signatures that are observed: the dunite-hosted systems in the northern Kalgoorlie Terrane (Fiorentini et al 2012b) and elsewhere globally (e.g., Heggie et al., 2012a, b) commonly display negative $\Delta^{33}S$ values (up to $-1.0\%$), whereas Kambalda-style channelized systems either display no MIF or positive (up to $+1.1\%$) signatures. These variable inter-shoot isotopic results at Kambalda are reflected in the variable platinum group element signature of the various channelized systems (Barnes et al., 2013). At the broader scale, these data support the hypothesis that dunite-hosted systems are generally characterized by lower R factors, where evidence of crustal contamination from proximal volcanic massive sulfides is preserved, in contrast to Kambalda-type deposits where evidence of crustal contamination by sulfidic shales is partially to completely erased due to extreme magma flux along channelized environments.

5.2.3 In situ data to identify natural fractionation between pentlandite-pyrrhotite
The complex sulfide mineralogy hosted in any mineralized sample may document various stages of the ore forming process. However, even when sulfide phases are cogenetic and in equilibrium, natural isotopic fractionation between sulfide pairs may occur due to crystal structure, temperature of formation, and chemical composition (e.g., O’Neil, 1986). For instance, the speciation of the sulfur source and the mass of a metal speciation can affect the preference for $^{34}S$ incorporation in one sulfide over another during precipitation (Bachinski, 1969; Seal, 2006 and references therein). This means that the isotopic signature of a single sulfide phase hosted in a sulfide layer may not be representative of the true isotopic composition of that very horizon.

Although, fractionation of sulfur isotopes between cogenetic sulfide minerals has been investigated by Bachinski (1969), Li and Lui (2006), Ohmoto and Rye (1979), Seal (2006), among others, the natural isotope fractionation of sulfur isotopes between pentlandite and pyrrhotite has not yet been quantified; however it is an important element in the study of magmatic ore deposits. In the case study focussed on the Moran shoot of the Long-Victor deposit, if we interpret pyrrhotite and pentlandite to be in equilibrium we conclude that pyrrhotite has a natural affinity for $^{34}S$ over pentlandite. Our results that demonstrate enrichment in $\delta^{34}S$ in pyrrhotite over chalcopyrite are in agreement with Li and Lui (2006). This same trend is also described for VMSO pentlandite above (also derived from a
magmatic ore deposit) and has been observed in other magmatic ore deposits (LaFlamme, unpublished data).

Multiple sulfur isotope analysis of massive sulfide from the Moran shoot at Long-Victor deposit yields $\delta^{34}\text{S}$ pentlandite = 2.91 ± 0.35‰ ($n=73$) and $\delta^{34}\text{S}$ pyrrhotite = 3.90 ± 0.20‰ ($n=19$). These two phases make up >96% of sulfides in this ore body and are within error of the bulk fluorination results ($\delta^{34}\text{S} = 3.10 ± 0.70‰; n=10$). Multiple sulfur isotope analysis of the massive sulfide horizon at the Victor South shoot (from which VMSO reference material is derived) yields $\delta^{34}\text{S}$ pentlandite (measured as bulk fluorination value and incorporates a minor component of unavoidable pyrrhotite) = 3.22 ± 0.51‰ ($n=11$) and $\delta^{34}\text{S}$ pyrrhotite (normalized to Alexo pyrrhotite) = 3.86 ± 0.18‰ ($n=7$). This is a circular problem because it is impossible to get a true value for pentlandite unaccompanied by pyrrhotite for VMSO, and the values for in situ analyses from the Moran shoot rely on a correction for instrumental mass fractionation normalised to the VMSO reference material. Here though, we assume the cogenetic sulfides from both shoots are in natural equilibrium, and so we can broadly estimate $\delta^{34}\text{S}_{\text{pyrrhotite-pentlandite}}$ to be 0.7–1.0 ± 0.5‰. The natural equilibrium isotopic fractionation between pyrrhotite and chalcopyrite determined at the Moran shoot to be $\delta^{34}\text{S} = \sim 0.2‰$, is in agreement with that defined by Li and Lui (2006). Therefore, in ore bodies with multiple sulfide phases in equilibrium, $\delta^{34}\text{S}$ values are a reflection of the relative proportions of those phases. Therefore, care should be applied when small deviations in isotopic signatures derived from bulk analyses of massive sulfide bodies are used to interpret variations in the source of sulfur, volatiles and metals in ore bodies (e.g., Bekker et al., 2009; Fiorentini et al., 2012b; Li et al., 2009). In fact, these variations may solely be reflecting proportional differences of sulfide phases, or may homogenise different phases of mineralisation with different isotopic signatures. In turn this can lead to a better understanding of small spatial and temporal changes in multiple sulfur isotopes related to the magmatic and/or hydrothermal environment.

5.2.4 Test for potential crystallographic orientation effects in pentlandite

The integration of EBSD with high-spatial resolution isotopic analyses allows the potential role of crystal orientation on isotopic data quality to be assessed (Taylor et al., 2012). The potential effects of grain orientation on sulfur isotope measurements in the Moran Shoot pentlandite have been assessed by comparison of EBSD-derived crystal orientation information with sulfur isotope data (as $\delta^{34}\text{S}$ and $\Delta^{33}\text{S}$) from known ion probe analysis sites. Results demonstrate that no orientation effect on $\delta^{34}\text{S}$ and $\Delta^{33}\text{S}$ exists for pentlandite (Figure 7). Inverse pole figures (of which one sample is shown in Figure 9) relate count rates to grain orientation in which analyses are plotted based on the angular relationship between the mount x-direction and the main crystal axes. Comparison of grain orientation data from ~50 pentlandite grains and 73 multiple sulfur isotopes analyses demonstrates a consistent value of $\delta^{34}\text{S}$ and $\Delta^{33}\text{S}$, within the envelope of single analysis uncertainty irrespective of grain orientation.

Certain sulfide minerals (galena and sphalerite) have been shown to exhibit a crystal orientation effect resulting in instrumental mass bias towards $^{34}\text{S}$, while others (pyrite,
pyrrhotite and chalcopyrite) do not (Kozdon et al., 2010; Kita et al., 2011). Here, our data indicate that there is no resolvable crystallographic orientation control on either $\delta^{34}\text{S}$ or $\Delta^{33}\text{S}$ due to the interaction of the beam with the pentlandite grain. This is a completely new outcome as the orientation effect on beam-pentlandite interaction was not previously known.

**Figure 9:** Example of orientation effect in pentlandite analysed by electron backscatter diffraction (EBSD).

Pentlandite from massive sulfide lens mounted as pucks (1 puck shown here) drilled from a massive sulfide lens at the Moran shoot, Kambalda. A: EDS phase map showing distribution of NiS (pentlandite; yellow), FeS (pyrrhotite; red), and FeO (iron oxide; green). B: Inverse pole figure map, plotted with respect to the user-defined x direction. Black lines show grain boundaries with >10° misorientation. C: Inverse pole figure (lower hemisphere equal area projection) for the sample x direction showing grain orientations indicated by colours of pentlandite grains shown in B ($n=17195$ data points). D: $\delta^{34}\text{S}$ and $\Delta^{33}\text{S}$ from pentlandite with colour matched to crystallographic orientation shown in B and C.

**6. CONCLUSIONS**

Instrumental mass fractionation is dependent on the chemical matrix of the material being analysed (the so-called matrix effect). For different sulfide minerals, the matrix effect is likely due to a combination of factors including varying sulfur contents, varying cation compositions and different crystallographic systems, giving rise to different ionisation potentials and different sputtering rates (e.g., Kozdon et al., 2010). The matrix effect between conjugate pyrite-sulfide pairs is a complex problem (at least partly) related to the presence of multiple and subtle instrument conditions and analytical protocols. Hence, SIMS inter-
mineral fractionation factors (even within a single laboratory) to calculate sulfur isotope
ratios should be used with caution and only as a last resort. Multiple sulfur isotope studies
should also take caution against propagating $\delta^{33}\text{S}$ and $\delta^{36}\text{S}$ from $\delta^{34}\text{S}$ following the mass
dependent fractionation relationship for reference materials in which these values have not
been quantified. It has been shown that $\Delta^{33}\text{S}$ and $\Delta^{36}\text{S}$ anomalies can occur in a range of
environments even outside of Archean terranes (i.e., at Proterozoic craton margins, Selvaraja et al., 2016; this study (Proterozoic Nifty-b chalcopyrite reference material); LaFlamme,
unpublished data, at modern hydrothermal vents; Ono et al., 2006b; Farquhar et al., 2007; and
in other environments, Cabral et al., 2013). Therefore, for the reasons above, unknown sulfur
isotope analyses should be normalised to chemically and isotopically characterised reference
material, four sulfides of which are presented here.

At the Long-Victor nickel-sulfide deposit in the world class Kambalda nickel camp in the
southern Kalgoorlie Terrane of Western Australia, a case study demonstrates how precise
multiple sulfur isotope analyses from magmatic pentlandite, pyrrhotite and chalcopyrite can
better constrain genetic models related to ore-forming processes. Results indicate that
pentlandite, pyrrhotite and chalcopyrite are in isotopic equilibrium and yield the same value
for $\Delta^{33}\text{S}$ (+0.2‰). The three magmatic phases show variable $\delta^{34}\text{S}$ values ($\delta^{34}\text{S}_{\text{chalcopyrite}} =
2.9‰, \delta^{34}\text{S}_{\text{pentlandite}} = 3.1‰, and $\delta^{34}\text{S}_{\text{pyrrhotite}} = 3.9‰$), which are indicative of natural
fractionation. The results indicate that unlike the giant dunite-hosted komatiite systems that
thermo-mechanically assimilated volcanogenic massive sulfides proximal to vents and
display negative $\Delta^{33}\text{S}$ values, the Kambalda ores formed in relatively distal environments
assimilating abyssal sulfidic shales. Careful in situ analysis is able to image the subtle
isotopic variability of the magmatic sulfide assemblage, which may help resolve the nature of
the ore-forming process. This approach may help to discriminate the magmatic sulfur isotope
signature from that recorded in metamorphic- and alteration-related sulfides, which may not
be resolved during bulk rock fluorination analysis.

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http://www.ccfs.mq.edu.au.

8. LIST OF FIGURES
Figure 1: Calculated magnitudes of mass independent fractionation as deviations from the
mass dependent fractionation line (MDF) presented in $\delta^{33}\text{S}$ vs. $\delta^{34}\text{S}$ space. The small
deviations from the MDF line may represent mass independent fractionation. This highlights
the importance of: 1) precise and accurate multiple sulfur isotope measurements, and 2)
quantification of uncertainty on $\Delta^{33}\text{S}$. Although not expressed in this figure, quantifying $\Delta^{36}\text{S}$
is equally important.

Figure 2: Demonstration of compositional homogeneity of: A) Sierra pyrite (Fe wt.%), B)
Nifty-b chalcopyrite (Cu wt.%), C) Alexo pyrrhotite (Fe wt.%), D) VMSO pentlandite (Ni
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Figure 3: Probability density plots for $\delta^{34}\text{S}$ for normalized reference material to demonstrate
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Figure 4: Example of compositionally homogeneous area of Alexo pyrrhotite compared to
areas of significant pentlandite exsolution. Raster spots show examples of assessment of two
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Figure 5: Plots of: (a) $\delta^{33}\text{S}_{\text{V-CDT}}$ vs. $\delta^{34}\text{S}_{\text{V-CDT}}$ and (b) $\Delta^{33}\text{S}$ vs. $\delta^{34}\text{S}_{\text{V-CDT}}$ for all sulfide
reference material (Sierra pyrite, Nifty-b chalcopyrite, Alexo pyrrhotite, and VMSO
pentlandite) measured by SIMS in this study. Error crosses are the overall uncertainty (at the
2SD level) based on propagating the individual analysis repeatability with the reproducibility
for the relevant analytical session. The Sierra pyrite is used as the primary reference for
measuring previously characterised pyrite reference material Isua 248474 and Balmat pyrite
(see Whitehouse, 2013; Graham and Valley, 1992).

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Figure 9 Example of orientation effect in pentlandite analysed by electron backscatter
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