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Citation for final published version:

Martin, Andrew J., McDonald, Iain, Jamieson, John W., Jenkin, Gawen R. T., McFall, Katie A., Piercey, Glenn, MacLeod, Christopher J. and Layne, Graham D. 2021. Mineral-scale variation in the trace metal and sulfur isotope composition of pyrite: implications for metal and sulfur sources in mafic VMS deposits. *Mineralium Deposita* 57 (6) , pp. 911-933. 10.1007/s00126-021-01080-1 file

Publishers page: <http://dx.doi.org/10.1007/s00126-021-01080-1>

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1 **Mineral-scale variation in the trace metal and sulfur isotope composition of pyrite:**
2 **Implications for metal and sulfur sources in mafic VMS deposits**

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10 **Abstract**

11 The link between metal enrichment and the addition of a magmatic volatile phase in volcanogenic massive
12 sulfide deposits and actively forming seafloor massive sulfide deposits remains poorly characterized. This
13 is especially true when considering how metal, sulfur and fluid flux change with time. In this study we
14 combine *in situ* sulfur isotope ($\delta^{34}\text{S}$; n=31) measurements with trace metal chemistry of pyrite (n=143) from
15 the Mala VMS deposit, Troodos, Cyprus. The aim of our study is to assess the links between volatile influx
16 and metal enrichment and establish how, or indeed if, this is preserved at the scale of individual mineral
17 grains. We classify pyrite based on texture into colloform, granular, disseminated and massive varieties.
18 The trace metal content of different pyrite textures is highly variable and relates to fluid temperature and
19 secondary reworking that are influenced by the location of the sample within the mound. The sulfur isotope
20 composition of pyrite at Mala ranges from -17.1 to 7.5‰ (n=31), with a range of -10.9 to 2.5‰ within a
21 single pyrite crystal. This variation is attributed to changes in the relative proportion of sulfur sourced from
22 i) SO_2 disproportionation, ii) thermochemical sulfate reduction, iii) the leaching of igneous sulfur/sulfide
23 and, iv) bacterial sulfate reduction. Our data shows that there is no correlation between $\delta^{34}\text{S}$ values and the

24 concentration of volatile elements (Te, Se) and Au in pyrite at Mala indicating that remobilization of trace
25 metals occurred within the mound.

26 Introduction

27 The source of metals in volcanogenic massive sulfide (VMS) and actively forming seafloor massive sulfide
28 (SMS) deposits remains poorly constrained and actively debated (Sillitoe et al. 1996; Yang and Scott 1996;
29 Jowitt et al. 2012; Keith et al. 2018; Martin et al. 2020; Patten et al. 2020). Principally, two metal sources
30 can contribute to the metal budget of VMS deposits: leaching of metals from underlying host rocks during
31 hydrothermal alteration (Richardson et al. 1987; Jowitt et al. 2012; Banerjee et al. 2000; Patten et al. 2017);
32 and the addition of a magmatic volatile phase to the overlying hydrothermal system (Yang and Scott 1996,
33 2002; de Ronde et al. 2011; Martin et al. 2020, 2021; Patten et al. 2020). The relative contribution of metals
34 from these two sources and how they change with system maturity (i.e. time) and the subsequent
35 preservation of trace metal signatures to distinguish these sources in sulfide minerals in VMS deposits
36 remains enigmatic. Here, we apply *in situ* geochemical and sulfur isotope measurements on pyrite from the
37 92 million year old Mala VMS deposit of the Troodos ophiolite (Cyprus) to assess co-variations in trace
38 metal content and sulfur isotope ratios ($\delta^{34}\text{S}$) in an ancient mafic VMS deposit. These data will be used to
39 investigate the variability in metal and sulfur sources during the growth of individual pyrite crystals and to
40 understand if the influx of certain volatile metals can be linked to variations in sulfur isotopic composition.

41 To investigate the source(s) of metals in VMS deposits, previous studies utilized either trace metal
42 enrichment profiles (Halbach et al. 1998; Butler and Nesbitt 1999; Maslennikov et al. 2009; Wohlgemuth-
43 Ueberwasser et al. 2015; Keith et al. 2016a; Grant et al. 2018; Wang et al. 2018), the ratio of Se to S
44 (Yamamoto 1976; Huston et al. 1995; Hannington et al. 1999; Layton-Matthews et al. 2008, 2013; Martin
45 et al. 2019), or the sulfur isotopic composition of sulfide minerals (Herzig et al. 1998; Gemmell et al. 2004;
46 Huston et al. 2011; Yeats et al. 2014; Brueckner et al. 2015; Lode et al. 2017; Zeng et al. 2017). However,
47 very few studies utilize a combined approach (e.g., Rouxel et al. 2004; Sharman et al. 2015; Meng et al.

48 2020; Martin et al. 2021) that takes into consideration any systematic relationship between trace element
49 enrichment profiles and the sulfur isotopic composition of sulfide minerals; this is especially true when
50 considering the complex nature of different pyrite textures, generations (i.e. overgrowths) and variation that
51 occurs at the scale of individual mineral grains.

52 In volatile-rich, subduction influenced VMS deposits, the systematic enrichment of certain metals such as
53 Pb, As, Sb, Bi, Hg and Te (Wohlgemuth-Ueberwasser et al. 2015), Se, Cu and Te (Keith et al. 2016a), Bi
54 and Te (Mathieu 2019) or Se, Cu, Te and Au (Martin et al. 2021) have been proposed to indicate the addition
55 of a metal-rich magmatic volatile phase to the hydrothermal system from degassing of shallow magma
56 reservoirs (e.g., Huston et al. 2011). Additionally, elevated Se/S (expressed as $\text{Se/S} \cdot 10^6$) that are >500 in
57 pyrite are also interpreted as representing an increased magmatic volatile influx in VMS deposits
58 (Yamamoto 1976; Huston et al. 1995; Layton-Matthews et al. 2008, 2013). The sulfur isotopic composition
59 of sulfide minerals provides further evidence of a magmatic volatile contribution to VMS deposits with the
60 addition of degassing magmatic SO_2 that undergoes disproportionation upon mixing with hydrothermal
61 fluid, and results in a characteristic sulfur isotope composition defined by $\delta^{34}\text{S}$ values that are less than the
62 magmatic mean ($<0\%$, MORB; Herzig et al. 1998; Kusakabe et al. 2000; de Ronde et al. 2005). These low
63 $\delta^{34}\text{S}$ values are generally limited to felsic or subduction influenced environments (e.g., Herzig et al. 1998;
64 Huston et al. 2011), where magmas are more volatile-rich relative to mafic hosted mid-ocean ridge (MOR)
65 environments (Wallace 2005). However, this signature was also recorded in an ancient mafic VMS deposit
66 (Martin et al. 2021). This sulfur isotope composition contrasts signatures typical of MOR environments,
67 where $\delta^{34}\text{S}$ values in sulfide minerals are positive, indicating a combination of sulfur from thermochemical
68 sulfate reduction (TSR) of seawater at high temperatures ($>160^\circ\text{C}$) in the presence of iron-bearing minerals
69 and leaching of sulfur and primary igneous sulfide minerals from igneous host rocks (Ono et al. 2007).

70 Previous studies utilising laser ablation inductively coupled plasma mass spectrometry (LA-ICP-MS) of
71 sulfide minerals from Troodos VMS deposits indicate that deposits associated with the Solea graben, one
72 of three fossil spreading axes in the Troodos ophiolite are enriched in Se, Cu and Au (Keith et al. 2016a;

73 Fig. 1). The enrichment of these elements is attributed to the contribution of a magmatic volatile phase to
74 the hydrothermal systems of the Solea graben (Martin et al. 2019, 2020).

75 Variation in the sulfur isotope composition of pyrite has largely been constrained in Troodos VMS deposits
76 using whole-rock analytical methods of sulfide minerals with the median composition for all Troodos VMS
77 sulfide minerals of $4.6 \pm 2.7\text{‰}$ (1σ , $n=220$) indicating that sulfur is primarily sourced from both the leaching
78 of igneous sulfur/sulfide and TSR of seawater (Hannington et al. 1998; Parvaz 2014; Keith et al. 2016a;
79 Pedersen et al. 2017; Martin et al. 2020). However, significant variation is noted in the sulfur isotope
80 composition of pyrite across Troodos VMS deposits with the $\delta^{34}\text{S}$ values of pyrite ranging from -7.6‰ to
81 13.2‰ (Martin et al. 2020, 2021). Samples of disseminated pyrite that occur within the volcanic
82 stratigraphy exhibit significantly lower $\delta^{34}\text{S}$ values than those typical for Troodos VMS deposits; pyrite is
83 depleted in ^{34}S , with $\delta^{34}\text{S}$ values ranging from -22.2‰ to -6.9‰ . These values indicate that bacterial sulfate
84 reduction (BSR) occurred at low temperatures ($<120^\circ\text{C}$) in the shallow subsurface, away from areas of high
85 temperature fluid discharge (Pedersen et al. 2017).

86 Sulfide mineral chemistry and the sulfur isotopic composition of sulfide minerals analysed using bulk
87 analytical techniques leads to the homogenization of any variation in the chemical and isotopic composition
88 of the sample. Thus, there is a need to apply *in situ* analytical techniques to better understand the spatial
89 zonation in isotopic and chemical composition within individual mineral grains. Utilizing a combined
90 approach of *in situ* LA-ICP-MS analysis to measure trace metal concentrations and Secondary Ion Mass
91 Spectrometry (SIMS) to measure sulfur isotope ratios, this study aims to establish if links exist between
92 trace metal enrichment and variations in the source of sulfur during VMS deposit formation. We focus on
93 the Mala VMS deposit located in the Troodos ophiolite of Cyprus, as Mala represents an immature
94 magmatic volatile-influenced deposit where links between magmatic volatile influx, sulfur isotope ratios
95 and trace metal enrichment signatures in pyrite have not been extensively modified during fluid
96 overprinting associated with the maturation of the VMS mound (Martin et al. 2021). We present *in situ*
97 sulfur isotope data and trace element geochemistry of different pyrite textures that highlight variable trace

98 element enrichment profiles that can be related to mound-scale fluid flow. We then combine trace metal
99 geochemistry and sulfur isotope ratios to test if any systematic relationship exists between magmatic
100 volatile influx and the enrichment of Te, Se and Au in pyrite at Mala.

101 Geological Setting

102 The 92 Ma Troodos ophiolite in Cyprus comprises a complete, un-deformed oceanic pseudo-stratigraphy
103 (Gass 1968; Mukasa and Ludden 1987). Mantle peridotites are surrounded radially by gabbro and
104 plagiogranite of the plutonic sequence, a sheeted dyke complex (SDC), and the basal group (BG), which
105 consists of a transitional horizon between the SDC and overlying upper and lower pillow lavas (UPL-LPL;
106 Fig. 1) (Gass 1968). It has long been recognized that the Troodos ophiolite formed in a sediment-free
107 subduction influenced environment due to the occurrence of boninite (Pearce and Robinson 2010; Woelki
108 et al. 2018), the trace element and isotopic composition of volcanic glass (Rautenschlein et al. 1985),
109 elevated H₂O contents of parent melts (>2 wt.% H₂O; Muenow et al. 1990), and an enrichment of magmatic
110 volatile derived elements (e.g., Te, Se) in some VMS deposits (Keith et al. 2016b; Martin et al. 2019, 2021).

111 Observations from actively forming intermediate to slow seafloor spreading centers are akin to processes
112 and structures preserved in the Troodos ophiolite (Varga and Moores 1985). On the northern flank of the
113 ophiolite, three structural grabens are delineated by inversely dipping sheeted dykes along a series of
114 ~north-south normal faults (Varga and Moores 1985). The grabens from east to west are; Larnaca, Mitsero
115 and Solea (Fig. 1). The grabens are widely accepted as representing fossil axial spreading ridges (Varga
116 and Moores 1985). Associated with these graben bounding faults are VMS deposits that formed within or
117 at the contact between the BG, LPL and UPL units at the periphery of the ophiolite that once represented
118 the Cretaceous seafloor (Hannington et al. 1998; Adamides 2010; Fig. 1).

119 Hydrothermal alteration of the Troodos crust is well-characterized and the leaching of metals from the SDC
120 during spilitisation and epidosite formation is interpreted as a possible source of metal and sulfur in
121 overlying VMS deposits (Richardson et al. 1987; Jowitt et al. 2012; Patten et al. 2017). More recently, a

122 pulsed magmatic volatile component has been suggested as an additional source of some metals (Cu, Au,
123 Te and Se) in Troodos VMS deposits, especially in magmatic volatile-influenced VMS deposits such as
124 Mala and deposits associated with the Solea graben in particular (Keith et al. 2016a; Martin et al. 2020; Fox
125 et al. 2020).

126 The Mala VMS deposit

127 The Mala VMS deposit is located in the southwest Troodos ophiolite, northwest of the town of Pano Pangia
128 within the Pafos Forest region (Fig. 1). Mala is located deep in the lava stratigraphy at the BG-LPL
129 transition. Previous studies have identified an enrichment in Te, Se, Cu and Au in pyrite at Mala relative to
130 other Troodos VMS deposits (Martin et al. 2021). Mala has been historically mined for Cu and Zn (0.8 Mt
131 total; Brazilian Metals Group, 2013) leading to the exposure of a massive sulfide mound at the northern
132 end of the open pit (Fig. 2).

133 The exposed pyrite mound measures approximately 12 x 8 m (width x height) and contains pyrite and
134 gypsum (Fig. 2A). Gypsum occurs as three distinct morphologies: massive-bedded, breccia infill, and mesh
135 textured varieties (Fig. 2B-G). Within the gypsum are euhedral pyrite crystals which vary in size from <1
136 mm up to 2 cm in width (Fig. 2D-G). In the VMS mound pyrite forms as five distinct textures: massive
137 euhedral grains (Fig. 3A), dendritic growths (Fig. 3B and C), colloform bands (Fig. 3D), granular pyrite
138 (Fig 3E) and as disseminated grains within gypsum and the surrounding wall rock (Fig. 3F). A previous
139 study by Martin et al. (2021) demonstrated that the pyrite-gypsum relationships preserved at Mala are
140 analogous to those observed in actively forming SMS deposits and reflect primary hydrothermal processes
141 and are not the result of supergene weathering during uplift and exposure.

142 Methods

143 Mineral Chemistry

144 Laser ablation ICP-MS was used to determine *in situ* trace element concentrations in pyrite. Spot analyses
145 (n=143) were performed on 10 representative polished blocks. Analyte masses used were ⁵⁷Fe, ⁶⁵Cu, ⁵⁹Co,

146 ⁶⁶Zn, ⁷⁵As, ⁷⁷Se, ¹⁰⁹Ag, ¹¹¹Cd, ¹²¹Sb, ¹²⁵Te, ¹⁸⁵Re, ¹⁸⁹Os, ¹⁹³Ir, ¹⁹⁵Pt, ¹⁹⁷Au ²⁰⁶Pb and ²⁰⁹Bi (Table S1, ESM).
147 Data were collected using a New Wave Research UP213 laser coupled to a Thermo iCAP RQ-ICP-MS at
148 Cardiff University, UK. Samples were analysed in time-resolved analysis mode with a nominal spot
149 diameter of 55 µm at a frequency of 10 Hz. Each analysis lasted 40 s and a gas blank was measured for 20
150 s prior to each analysis. Data correction and the subtraction of gas blanks was performed using Thermo
151 Qtegra software. External calibration was performed on a series of synthetic NiFeS standards (see Prichard
152 et al. 2013; Smith et al. 2016) and the reproducibility of analyses was monitored through the repeat analysis
153 of UQAC FeS-1. The repeat analysis of UQAC FeS-1 yielded a relative standard deviation (RSD) of <10%
154 for Fe, Co, Ni, As, Ru, Rh, Pd, Ag, Re, Os, Ir, Pt, Au and Bi, <15% for Cu, Se, Sb and Te and <20% for
155 Zn and Pb. The RSD for Cd was 24% (Table S2, ESM). Detection limits for all elements analysed are
156 available in ESM Table S2. Sulfur-33 was used as an internal standard for all analyses and a stoichiometric
157 concentration of 53.5 wt.% sulfur was used that is within error of the measured average sulfur concentration
158 for pyrite from Troodos VMS deposits (Martin et al. 2019).

159

160 Sulfur Isotope analysis

161 Secondary ion mass spectrometry microanalysis was used to determine the sulfur isotope composition
162 ($\delta^{34}\text{S}$) of pyrite. Analysis was undertaken on the same samples analysed via LA-ICP-MS. Epoxy mounted
163 polished blocks (n=6) were first coated with 300Å of Au and were analysed using a Cameca IMS 4f SIMS
164 at the MAF-IIC Microanalysis Facility at Memorial University of Newfoundland following the procedures
165 detailed in Brueckner et al. (2015) and Lode et al. (2017). Systematic analysis of points across pyrite grains
166 was performed proximal to LA-ICP-MS ablation pits to determine variation and correlation between $\delta^{34}\text{S}$
167 values and trace element concentrations. The sample was bombarded with a primary ion beam of 350-750
168 pA of Cs^+ , accelerated through a potential of 10 keV and focused into a 5-15 µm diameter spot. The duration
169 of each analyses was 16.3 minutes including 2 minutes of pre-sputtering. To discriminate between $^{33}\text{SH}^-$
170 (and $^{32}\text{SH}_2^-$) from $^{34}\text{S}^-$ the instrument was operated with a medium contrast aperture (150 µm), with entrance

171 and exit slits paired to give flat topped peaks at a mass resolving power of 2975 (10% peak height definition;
172 Brueckner et al. 2015). Negatively charged sputtered secondary ions were accelerated into the mass
173 spectrometer using a potential of +4.5 keV. To exclude sulfur contamination from the sample surface the
174 spot was pre-sputtered for 120 s with a 15 μm square rastered beam prior to analysis. Reproducibility based
175 on the repeat analysis of standard reference material pyrite-UL9 ($\delta^{34}\text{S}= 16.3\%$) and KH87 ($\delta^{34}\text{S}= 0.4\%$),
176 is typically better than $\pm 0.4\%$ (1σ) (Table S3, ESM). All analyses are reported in standard notation ($\%$)
177 relative to Vienna-Canyon Diablo Troilite (V-CDT).

178

179 Results

180 Sample characterization

181 Samples in this study were collected from the exposed Mala VMS mound (Fig. 2A and Fig. S1, ESM). We
182 subdivide pyrite samples based on texture into massive, dendritic, colloform, granular, and disseminated
183 varieties (Fig. 3). Massive pyrite occurs in discrete meter scale pods consisting of coarse grained (mm to
184 cm) aggregates of euhedral grains (Fig. 2C and 3A and B). Multiple pyrite generations occur as overgrowths
185 within massive pyrite samples and can be delineated by inclusion-rich zones at the grain margin (Fig. 3A).
186 Within massive pyrite, dendritic horizons were common but were not analysed in this study (Fig. 3B and
187 C). Colloform pyrite is rare, occurring spatially associated with granular horizons (Fig. 3D). Granular
188 massive pyrite contains anhedral aggregates of highly resorbed and often spherical pyrite in a pyrite matrix
189 (Fig. 3E). Disseminated pyrite grains are commonly euhedral and vary in size from <1 mm to >2 cm, and
190 occur within gypsum horizons as well as within the volcanic rocks that surround the massive sulfide mound
191 (Fig. 2D and 3F).

192 Trace element geochemistry

193 To assess trace metal enrichment profiles, results are grouped based on pyrite texture, and, for disseminated
194 grains, the location of the analytical point in relation to the core or rim of the individual mineral grain that

195 was analyzed. The division between core and rim is arbitrary and depends on the number of analyses
196 performed within an individual pyrite grain, which in turn reflects the size of the pyrite grain (Table S1,
197 ESM).

198 Colloform pyrite (n=15) contains the highest median concentrations of Au, Pb and Re at 0.13, 9.8 and 0.17
199 ppm, respectively. All Ni and Cd analyses are below the detection limit, as are the majority of Cu, Zn and
200 Co (Table 1; Fig. 4A). The concentration of Se is relatively homogenous ranging from 190 to 302 ppm with
201 a median concentration of 245 ppm (n=15) (Fig. 4B, Table 1). The total measured trace metal content for
202 colloform pyrite is 371 ppm (Fig. 4D). A strong positive correlation is noted between Au and Ag ($R^2=0.95$),
203 and a moderate positive correlation between Pb and Ag ($R^2=0.72$), Te and Au ($R^2=0.56$) and Te and Bi (R^2
204 $=0.52$).

205 Granular pyrite (n=10) is depleted in most trace metals relative to other pyrite textures considered here. All
206 analyses for Ni, Ag, Zn, Cd, Re and Pb are below detection limit as are the majority of analyses for Au, Bi,
207 Cu and Sb (Table 1). Cobalt and Te are notably enriched relative to colloform pyrite with median
208 concentrations of 17.9 and 5.9 ppm, respectively (n=10; Fig. 4A; Table 1). As observed in colloform pyrite,
209 the concentration of Se is relatively homogenous, ranging from 5 to 31 ppm, with a median concentration
210 of 6.4 ppm (n=10) (Fig. 4B, Table 1). With the exception of a moderate correlation between As and Se ($R^2=$
211 0.63) no notable correlations exist. Granular pyrite contains the lowest measured total trace metal content
212 at 198 ppm (Fig. 4D).

213 Trace metal enrichment profiles in massive pyrite (n=40) are highly variable between samples. For
214 example, sample MAL 17 is enriched in Cu with a median concentration of 0.04 wt.% with a range of
215 between 0.01 to 0.11 wt.% (n=10) but is relatively depleted in Se with a median concentration of 61 ppm
216 (n=10) (Fig. 4A, Table 1). Across all massive samples, Se exhibits the most variation, ranging from 30 to
217 1253 ppm with a median concentration of 178 ppm (n=40) (Fig. 4B). With the exception of one point, all
218 analyses for Ni and Cd were below the detection limit (Table 1). Massive pyrite contains the second highest

219 total measured metal content at 566 ppm (Fig. 4D). A weak correlation is noted between Te and Se ($R^2 =$
220 0.38; Fig. 4B).

221 Grains of disseminated pyrite within gypsum (n=78) contain the highest concentration of Se, Te, As and Bi
222 relative to all other pyrite textures (Fig. 4A-C, Table 1). Trace metal concentrations are highly variable
223 within and between individual samples relative to other pyrite textures analysed in this study. For example,
224 Se concentrations range from 18 to 3261 ppm with a median concentration of 645 ppm (n=78). Similar
225 variability is also noted for Co, As and Te (Table 1; Fig. 4B-C). Sample MAL 05 is notably enriched in Se
226 and Te relative to all other samples with median concentrations of 1668 and 16.4 ppm, respectively (n=14).
227 Disseminated grains have the highest total measured metal content at 981 ppm (Fig. 4D); largely consisting
228 of Se and Cu. Disseminated pyrite grains are sub-divided further based on the location of the analytical
229 point relative to the margin of the pyrite grain to assess metal enrichment trends across individual grains
230 (Fig. 5). With the exception of Co and Cu, that are enriched at the margin of grains (n=43), pyrite cores
231 exhibit a minor enrichment in As, Se, Te and Au relative the grain margins (n=35; Fig. 5).

232 Sulfur isotopes

233 Secondary ion mass spectrometry analysis was performed on representative samples of colloform (n=6),
234 granular (n=2), massive (n=9) and disseminated (n=14) textured pyrite (Fig. 6, Table S3, ESM). The median
235 $\delta^{34}\text{S}$ value for all samples is -1.2‰ (n=31) with a range of 24.6‰. Granular pyrite exhibits the lightest
236 median sulfur isotopic composition with two analyses yielding $\delta^{34}\text{S}$ values of -10.5‰ and -4.7‰. Colloform
237 pyrite consistently exhibits $\delta^{34}\text{S}$ values <0‰ ranging from -3.8 to -0.1‰ with a median of -2.2‰ (n=6)
238 (Fig. 6). The sulfur isotopic composition of massive pyrite is highly variable, ranging from -17.1‰ to 7.5‰
239 with a median composition of -1.8‰ (n=9).

240 To assess intra-grain variation in sulfur isotopic composition of disseminated pyrite, transects of analytical
241 points were analysed across different pyrite grains (Fig. 7). The same grains were then etched with NaClO
242 for 90 s to reveal any chemical zonation. A significant amount of variability is recorded in a single pyrite

243 grain, for example MAL 05 where $\delta^{34}\text{S}$ values range from 2.5‰ to -10.9‰ (sub-grain SG1) with a median
244 of 2.0‰ (n=4, Fig. 7A). There is no systematic relationship between $\delta^{34}\text{S}$ values and the location of the
245 analysis with respect to the core or rim of the grain, for example, low values do not only occur at the margin
246 of the grain. This is best illustrated in sample MAL 05, an aggregate of three pyrite grains disseminated
247 within gypsum where opposing edges of the pyrite sub-grain SG1 have $\delta^{34}\text{S}$ values of -10.9‰ and 2.5‰,
248 respectively (Figure 7A). Individual pyrite grains that are located within the same region of the VMS mound
249 (Fig. 2 and Fig. S1, ESM) also exhibit notable variation, for example two grains from sample MAL 11 (Fig
250 7B and C) that have different median $\delta^{34}\text{S}$ values of -0.9‰ and 0.9‰, respectively (n=7). Again, there is
251 no systematic pattern of enrichment from the grain core to rim within these grains.

252 Discussion

253 Mound-scale trace element systematics

254 The trace metal content and ratio of metals in pyrite can be used as a proxy for the past physical and
255 chemical conditions of hydrothermal fluids (e.g., temperature, pH, $f\text{O}_2$ and $f\text{S}_2$) (Butler and Nesbitt 1999;
256 Hannington et al. 2005; Maslennikov et al. 2009; Genna and Gaboury 2015; Wohlgemuth-Ueberwasser et
257 al. 2015; Keith et al. 2016b; Monecke et al. 2016; Grant et al. 2018; Wang et al. 2018). Thus, variations in
258 trace metal enrichment profiles in pyrite in VMS deposits reflect changes in the physicochemistry of fluids
259 and the flux of metals and fluid entering the VMS deposit during growth of the VMS mound. At the mound
260 scale (~10s of m) steep temperature gradients occur due to variations in the relative amounts of seawater
261 ingress and hydrothermal fluid influx with depth within the mound. Increasing temperatures within the
262 mound lead to the dissolution, recrystallization and remobilization of metals associated with lower
263 temperature sulfide minerals (e.g., Zn, Au, Ag, As, Pb, Sb) from the mound interior to the cooler margin of
264 the mound during zone refining (Eldridge et al. 1983; Hannington et al. 1986; Petersen et al. 2000; Galley
265 et al. 2007). This process is preserved in active SMS deposits as systematic variations in trace metal
266 enrichment profiles between different pyrite textures, such as colloform pyrite that forms near the seawater

267 interface that is enriched in elements that are associated with low temperature sulfide minerals or those
268 sourced from seawater (e.g., Mo, Tl, Pb; Grant et al. 2018). We note the same systematic variation in the
269 distribution of trace metals between different pyrite textures at the Mala VMS deposit.

270 Colloform pyrite occurs in active SMS deposits towards the margin of the sulfide mound or during the early
271 stage of chimney growth in environments where seawater ingress is high and fluid temperatures are lower
272 producing non-equilibrium conditions (Maslennikov et al. 2009, 2017; Melekestseva et al. 2014;
273 Wohlgemuth-Ueberwasser et al. 2015; Keith et al. 2016a,b; Grant et al. 2018). Colloform pyrite from Mala
274 is enriched in Au and Pb relative to all other pyrite and contains the lowest median concentration of Co, Cu
275 and Te, with all Ni analyses below detection limit, indicating it formed at lower fluid temperatures. At these
276 low concentrations, both Au and Pb are likely incorporated into the pyrite structure as a lattice bound
277 substitution, whilst at higher concentrations inclusions may occur (Huston et al. 1995; Cook et al. 2009).
278 This trace metal enrichment profile is similar to those reported in colloform pyrite from black smoker
279 chimneys of Urals VMS deposits that are enriched in Au, Ag, Pb and elements derived from seawater such
280 as Mo and Tl that were not analysed in this study (Maslennikov et al. 2009) or colloform pyrite from
281 PACMANUS that is enriched in Au and As but depleted in Te and Se relative to massive pyrite
282 (Wohlgemuth-Ueberwasser et al. 2015). A depletion in Co and Te, which are enriched in high-temperature
283 pyrite varieties (e.g., massive pyrite) in active SMS deposits, further supports a low temperature origin for
284 colloform pyrite at Mala (Wohlgemuth-Ueberwasser et al. 2015; Monecke et al. 2016; Grant et al. 2018).
285 The trace element enrichment profile in colloform pyrite at Mala is comparable to colloform pyrite from
286 the actively forming TAG SMS deposit located at the Mid-Atlantic Ridge (Grant et al. 2018) and from
287 colloform pyrite at the Skouriotissa VMS deposit (Keith et al. 2016a). Primarily, this trace metal enrichment
288 profile reflects the location of colloform pyrite at shallow depths within the VMS mound in close proximity
289 to the seawater interface, and its formation under lower fluid temperatures generated by increased amounts
290 of seawater mixing relative to massive pyrite at deeper stratigraphic levels in the VMS mound (Monecke
291 et al. 2016).

292 Mala colloform pyrite is strongly enriched in Se (Fig. 4) relative to concentrations reported for colloform
293 pyrite at the Skouriotissa VMS deposit (Troodos) that average 12 ± 10 ppm (1σ , $n=52$; Keith et al. 2016a),
294 black smoker chimneys from Urals VMS deposits that average 9 ± 5 ppm (1σ , $n=8$; Maslennikov et al. 2009),
295 concentrations of <4.5 ppm at the Semenov vent field (Melekestseva et al. 2014), and the TAG and
296 PACMUS (Roman Ruins) deposits where colloform pyrite did not contain any detectable Se (Wohlgemuth-
297 Ueberwasser et al. 2015; Grant et al. 2018). This concentration is also notably higher than that typically
298 recorded in colloform pyrite in other Troodos VMS deposits where the median concentration is 49 ppm
299 (Martin et al. 2019, 2020). Several studies indicate that Se is preferentially enriched in pyrite that formed
300 in high-temperature regions of the VMS mound or the interior conduit of black smoker chimneys, indicating
301 an affinity for high temperature fluids (Butler and Nesbitt, 1999; Maslennikov et al. 2009; Revan et al.
302 2014; Grant et al. 2018). However, at Mala Se is also enriched in pyrite that is interpreted to have formed
303 at lower temperatures, and we suggest that the enrichment of Se in colloform pyrite is a consequence of an
304 increased magmatic volatile influx.

305 Granular textured massive pyrite occurs towards the margin of the sulfide mound and forms through the
306 collapse and physical reworking of sulfide fragments during the growth of the VMS deposit (Humphris et
307 al. 1995; Petersen et al. 2000). Granular pyrite contains the lowest total measured trace metal content but
308 is relatively enriched in Te compared to other pyrite varieties (Table 1; Fig. 4A). We suggest that metal
309 enrichment profiles in granular pyrite reflect the remobilization and subsequent loss of some metals during
310 zone refining and oxidization of primary massive pyrite at low temperatures ($<100^\circ\text{C}$) at or near the seafloor
311 towards the periphery of the VMS mound (Herzig et al. 1991; Hannington et al. 1998; Fallon et al. 2017;
312 Murton et al. 2019). Assuming that massive pyrite samples represent the primary trace metal content of
313 granular samples prior to reworking, then all elements have been depleted during reworking with the
314 exception of Te. The enrichment of Te, which is readily mobilized under oxidizing and low temperature
315 fluid conditions (e.g., McPhail 1995), possibly indicates that the granular sample analysed was initially
316 enriched in Te prior to reworking.

317 Massive pyrite at Mala occurs as discrete pods throughout the sulfide mound. The trace metal composition
318 is highly variable depending on the individual sample analysed (Table S1 and Fig. S1, ESM). Fluid flow
319 regimes evolve with time in response to permeability fluctuations related to the collapse and occlusion of
320 permeability pathways, leading to the disruption of high-temperature fluid flow pathways within the VMS
321 mound (e.g., Tivey et al. 1995; You and Bickle 1998; Tivey 2007). These local-scale fluctuations (cm to
322 m's) in fluid flow led to changes in temperature and the remobilization of metals stable at low temperatures
323 from areas proximal to zones of high temperature fluid flow (>350°C) within the mound, leading to highly
324 heterogeneous trace metal enrichment profiles between different massive pyrite samples. For example,
325 samples located proximal to high temperature (>350°C) fluid pathways (e.g., MAL 14) are relatively
326 enriched in Se, Co and Te compared with samples that have experienced zone refining during fluid
327 overprinting by later cooler fluids (<350°C) (e.g., MAL 17). Hence, such large variations in trace metal
328 concentrations between massive pyrite samples reflects the dynamic nature of fluid flow within the VMS
329 mound. Alternatively, pyrite that contains low total trace metal concentrations may have been affected by
330 a higher degree of zone refining leading to the localised leaching and remobilization of most trace metals,
331 even those associated with high temperature assemblages as suggested for the Kokkinopezula VMS deposit
332 in Troodos (Hannington et al. 1998). However, high degrees of zone refining are considered to occur in
333 mature VMS deposits, whereas the Mala deposit is suggested to represent an immature system based on the
334 elevated concentration of magmatic volatile elements and low sulfur isotope values (Martin et al. 2021)
335 and, therefore, it is unlikely that enough time has elapsed to reach the high degree of zone refining.

336 Disseminated pyrite occurs as euhedral grains within gypsum (Fig. 2D) and is strongly enriched in Se
337 compared to all other pyrite textures (Fig. 5). This enrichment in Se is most likely a function of the
338 preferential partitioning of Se into disseminated pyrite grains relative to the surrounding gypsum/anhydrite
339 matrix (Yamamoto 1976; Huston et al. 1995). Alternatively, the enrichment of Se could indicate that
340 disseminated pyrite formed under lower fluid temperatures, assuming that Se is transported as H₂Se in the
341 hydrothermal fluid, as thermodynamic models for Se in pyrite predict that it is preferentially enriched at

342 lower temperatures (150°C) and more oxidizing conditions ($\Sigma\text{SO}_4 > \Sigma\text{H}_2\text{S}$; Huston et al. 1995). However,
343 observations from the active TAG deposit indicate that anhydrite-rich zones, that are analogous to gypsum-
344 rich zones at Mala where disseminated pyrite occurs, typically form at temperatures $>350^\circ\text{C}$ (Edmond et
345 al. 1995; Grant et al. 2018; Petersen et al. 2000). Furthermore, pyrite grains appear euhedral with no
346 hematite or magnetite that would indicate more oxidized fluids. Thus, there is no evidence supporting the
347 formation of disseminated pyrite from more oxidized or low-temperature ($\sim 150^\circ\text{C}$) fluids at Mala that are
348 favorable for the incorporation of Se in pyrite (Yamamoto 1976; Huston et al. 1995).

349 As observed in massive pyrite samples, notable variation exists between individual disseminated samples
350 (Fig. 8). We use the Te and Se content of pyrite as proxies for magmatic volatile influx as these elements
351 are enriched in VMS deposits in volatile-rich subduction influenced environments (Keith et al. 2016b) and
352 Co and As as indicators of high and low temperature fluids, respectively (Grant et al. 2018) (Fig. 8).
353 Samples MAL 18 and MAL 05 are located in the upper mound region, whilst MAL 04 and MAL 11 are
354 located in the lower mound (Fig. 2A). Based on studies of the active TAG mound, spatial trends in the trace
355 metal content of pyrite exists with, for example, Ni, Co and Se enrichment in high-temperature stockwork
356 zones and As, Pb, Zn and Ag enrichment in lower temperature massive pyrite (Grant et al. 2018). Sample
357 MAL 05 contains notably higher Se and Te concentrations relative to all other grains, and is depleted in Co
358 and enriched in As, indicating that it formed at low temperatures. In contrast, sample MAL 18, which also
359 formed in the upper mound area, is relatively depleted in Se, Te and As but is strongly enriched in Co,
360 indicating that it formed from high temperature fluids ($>350^\circ\text{C}$; Metz and Trefry 2000; Keith et al. 2016a;
361 Grant et al. 2018) (Fig. 8). Some samples (e.g., MAL 04) from the lower mound are depleted in As, Co, Te
362 and Se relative to other samples, possibly providing evidence for high degrees of zone refining (Hannington
363 et al. 1998). The trace metal enrichment profile in samples from the lower mound are highly variable with
364 some analyses strongly enriched in Te, Se, Co and As, indicating a change in fluid temperature and/or
365 magmatic volatile influx as the pyrite grain grew. The sporadic variation in trace metal concentrations
366 within individual disseminated pyrite grains indicates the influence of dynamic fluid flow regimes within

367 the VMS mound that leads to strong and transient physicochemical gradients localised over cm to m scales.
368 Hence, our data indicate that the well-established zone-refining model that is modelled on the active TAG
369 deposit (Humphris et al. 1995; Grant et al. 2018) is more complex when considering the distribution of
370 trace metals, especially when accounting for the distribution of metals over individual mineral grains. This
371 notwithstanding, there are obvious similarities between actively forming SMS deposits and the zonation of
372 metals in black smoker chimneys with the ancient Mala VMS deposit, suggesting that processes that control
373 the mound scale distribution and cycling of trace metals are broadly analogous between the two and did not
374 change since the Cretaceous.

375 Sulfur isotope systematics

376 Sulfur isotope analysis of sulfide minerals, such as pyrite, in VMS deposits can be used to discriminate
377 contributions from isotopically-distinct reservoirs of sulfur in hydrothermal fluids, such as sulfur leached
378 from oceanic crust, seawater sulfate, and, SO₂ degassing from shallow magma chambers (Lüders et al.
379 2001; Hannington et al. 2005; Ono et al. 2007; de Ronde et al. 2011; Keith et al. 2016a; Martin et al. 2020;
380 LaFlamme et al. 2021). The two primary sources of sulfur in Troodos VMS deposits are the bulk Troodos
381 crust that has a sulfur isotope composition of 1 to 0‰ (Alt 1994), Cretaceous seawater sulfate at $\delta^{34}\text{S} = 18$
382 to 19‰ (Kampschulte and Strauss 2004), and at the Mala VMS deposit, the disproportionation of SO₂
383 (Martin et al. 2021). The mixing of sulfur sourced from TSR of seawater and the leaching of primary
384 igneous sulfur/sulfide from host rocks (Woodruff and Shanks 1988; Shanks 2001) lead to an average $\delta^{34}\text{S}$
385 value in pyrite of $4.6 \pm 2.8\%$ (1σ , $n=160$), indicating that, on average, ~34% of sulfur is sourced from TSR
386 during VMS deposit formation in Troodos (Martin et al. 2020). The Skouriotissa and Sha, and most notably
387 the Mala VMS deposits, contain pyrite with a sulfur isotopic composition that is less than <0‰. Previously,
388 the low $\delta^{34}\text{S}$ values in pyrite have been attributed to the disproportion of SO₂ during the degassing of
389 volatiles from a shallow magma chamber (Keith et al. 2016a; Martin et al. 2020). At Mala,
390 disproportionation generates sulfate with a sulfur isotopic composition of 14.5‰ in gypsum ($n=26$), and
391 pyrite that is depleted in ³⁴S on average by -4.3‰ ($n=28$) (Martin et al. 2021). Bacterial sulfate reduction

392 of seawater at low temperatures (<120°C) can also form negative $\delta^{34}\text{S}$ values in pyrite as low as -22.2‰ in
393 altered oceanic crust in Troodos, such as those found in pyrite veins in the pillow lavas, indicating that BSR
394 occurred distally to high-temperature VMS deposits in the shallow lava stratigraphy (Alt 1994; Pedersen et
395 al. 2017).

396 The *in situ* analysis of different pyrite textures indicates that the source of sulfur in the Mala VMS deposit
397 was not constant and evolved with time, leading to variations in $\delta^{34}\text{S}$ values across individual pyrite grains.
398 Variation in $\delta^{34}\text{S}$ values between pyrite textures relates to mound-scale fluid flow that is directly influenced
399 by the location of specific pyrite samples within the mound that in turn influences pyrite texture (e.g.,
400 colloform vs. massive samples). Similar variations are reported across black smoker chimneys where
401 intense gradients in temperature and fluid mixing occur within the chimney wall (Haymon 1983; Crowe
402 and Valley 1992). For example, $\delta^{34}\text{S}$ varies from 2.1 to 6.4‰ in pyrite from the interior to exterior chimney
403 wall, respectively, in a black smoker chimney from 1-2°S on the EPR (Meng et al. 2020). Given the large
404 range in $\delta^{34}\text{S}$ values of pyrite at Mala, from -17.1‰ to 7.5‰, we suggest four processes that account for
405 the observed variation in the sulfur isotope composition of pyrite: i) TSR of seawater (18 to 19‰), ii) the
406 leaching of igneous sulfide (0 to 1‰), iii) the disproportionation of degassing SO_2 (0 to 1‰) and, iv) BSR.
407 Firstly, considering variation in sulfur isotope composition with pyrite texture, previous studies note a
408 systematic relationship between stratigraphic depth and $\delta^{34}\text{S}$ values in pyrite from the Skouriotissa VMS
409 deposit (Keith et al. 2016a), where enrichment of ^{34}S in shallow euhedral and colloform textured pyrite
410 leads to $\delta^{34}\text{S}$ values of 4.7‰ and 4.9‰, respectively (n=4, Keith et al. 2016a). This variation is attributed
411 to the increased proportion of sulfur that is incorporated from TSR of Cretaceous seawater in regions of the
412 VMS mound near the seawater interface. In contrast, deep, stockwork pyrite has a $\delta^{34}\text{S}$ value of -1.4‰
413 indicating lesser amounts of TSR and leaching of igneous sulfur/sulfide and an increased amount of sulfur
414 sourced from the disproportionation of degassing SO_2 , producing a sulfur isotope composition in pyrite that
415 is less than the Troodos magmatic mean (0-1‰; Alt 1994) (Keith et al. 2016a). Data presented in this study
416 for the Mala VMS deposit has a range of 24.6‰ (n=31), notably more variable than Skouriotissa with a

417 range of 7.8‰ (n=29; Keith et al. 2016a; Martin et al. 2020), with no systematic variation between $\delta^{34}\text{S}$
418 values and pyrite texture at Mala (Fig. 6 and 9). Instead, 18 of the 31 analyses across disseminated, granular,
419 colloform and massive textured pyrite contain $\delta^{34}\text{S}$ values of less than the Troodos magmatic mean of $\sim 0\%$
420 (Fig. 9; Alt 1994). This supports previous interpretations that have been proposed for the Mala VMS deposit
421 that indicates sulfur was sourced from a mixture of SO_2 disproportionation, TSR and leaching of igneous
422 sulfur/sulfide. However, the low $\delta^{34}\text{S}$ values measured in massive pyrite (down to -17.1%), which are much
423 lower than any values previously reported, could be produced either via the low temperature
424 disproportionation of SO_2 or during BSR of Cretaceous seawater. Based on the expected fractionation
425 factors between SO_2 and pyrite (after Sakai 1968), SO_2 disproportionation would have had to occur at a
426 temperature of $\sim 120^\circ\text{C}$ to account for the measured depletion in ^{34}S to -17.1% , assuming a starting SO_2
427 composition of 0% . Alternatively, at low temperatures ($<120^\circ\text{C}$) BSR of seawater sulfate can generate
428 large depletions in ^{34}S up to -38.9% in pyrite (Habicht and Canfield 1997; Wortmann et al. 2001; Farquhar
429 et al. 2003; Rouxel et al. 2008; Alt and Shanks 2011; Nozaki et al. 2020).

430 In active seafloor hydrothermal systems evidence for both the low temperature disproportionation of SO_2
431 and BSR exist. Negative $\delta^{34}\text{S}$ values attributed to the disproportionation of SO_2 down to -17.5% are found in
432 active seafloor hydrothermal systems that experience an increased influx of magmatic volatiles, such as
433 Conical seamount, Papa New Guinea (Gemmell et al. 2004), however typical median values are less
434 extreme, for example the median $\delta^{34}\text{S}$ values at Conical seamount are -1.2% (n=66) (Petersen et al. 2002;
435 Gemmell et al. 2004) at Hine Hina, Valu Fa Ridge, Lau Basin, -4.5% (n=12) (Herzig et al. 1998), at
436 Brothers volcano, Kermadec arc, -2.9% (n=87) (Kermadec arc; de Ronde et al. 2005, 2011) and -5.6%
437 (n=13) at SuSu Knolls, Manus back-arc basin (Kim et al. 2004), indicating that sulfur was sourced from a
438 combination of SO_2 disproportionation, TSR and leaching of igneous sulfur/sulfide.

439 Evidence of BSR has also been suggested as an important processes in the Agrokippia B VMS deposit
440 (Troodos) where multiple sulfur isotopes ($\Delta^{33}\text{S}$) indicate re-working of sulfide minerals in the upper part of
441 the VMS mound by microbes was an important but localised process (Pedersen et al. 2017). In lower

442 temperature regions of the Troodos stratigraphy, similar $\delta^{34}\text{S}$ values of -17.2‰ and as low as -22.2‰ (Alt
443 1994; Pedersen et al. 2017) have been reported for vein pyrite from the upper lava stratigraphy where the
444 occurrence of analcime, natrolite and phillipsite indicate the presence of low temperature fluids (<100°C;
445 Gass and Smewing 1973) where BSR could occur.

446 The negative $\delta^{34}\text{S}$ signature, down to -17.1‰, with a median composition of -13.7‰ (n=3) for the massive
447 sulfide sample at Mala, is somewhat enigmatic as both disproportionation of SO_2 and BSR require low fluid
448 temperatures <120°C to explain the observed $\delta^{34}\text{S}$ signature. Fractionation factors between SO_2 and pyrite
449 (Sakai, 1968), using an initial value of 0‰, would have had to occur at a temperature of 187°C to explain
450 the median sulfur isotopic composition in pyrite of -13.7‰; assuming that SO_2 disproportionation was the
451 only source of sulfur (i.e. a closed system), which as shown in previous studies is not the case (Ono et al.
452 2007).

453 Combining trace element geochemistry and sulfur isotope data, we suggest that the $\delta^{34}\text{S}$ signature is best
454 explained by BSR of seawater. Disproportionation of SO_2 that indicates magmatic volatile influx is, at the
455 mound scale, commonly associated with increased concentrations of Se, Te and Au in the Mala VMS
456 deposit (Martin et al. 2021); however this is not the case in massive pyrite sample MAL 17, that is depleted
457 in Te, Se and Au compared to all other massive pyrite samples. Furthermore, MAL 17 has a median Co
458 concentration of 8.4 ppm (n=10), compared to 38.7 ppm for all massive samples (n=40), further supporting
459 its formation from relatively low temperature fluids (<350°C) (Keith et al. 2016a,b; Monecke et al. 2016;
460 Grant et al. 2018). The relative depletion in Te, Se and Au is not consistent with its formation during
461 magmatic degassing, indicating it most likely formed during the BSR of seawater. This is further supported
462 by the location of the sample on the western margin of the VMS mound (Fig. 2A), and assuming that the
463 mound morphology preserved today is representative of how the Mala VMS deposit formed at the
464 Cretaceous seafloor, the sample is located near the seawater interface where seawater ingress is high and
465 the fluid temperature is expected to be low, possibly <120°C (cf. Pedersen et al. 2017).

466 The sulfur isotopic composition of colloform textured pyrite at Mala has a median $\delta^{34}\text{S}$ value of -2.2‰
467 (n=6; Fig. 4). This is significantly lower than the median $\delta^{34}\text{S}$ value recorded in colloform pyrite at the
468 Skouriotissa VMS deposit of 4.9‰ (n=2; Keith et al. 2016a), the Ice VMS deposit (Yukon, Canada) at
469 5.0‰ (n=8; McDonald et al. 2018), and the Ezuri VMS deposit (Hokuroko, Japan) at 4.5‰ (n=16; Barrie
470 et al. 2009). The positive values found in colloform pyrite in most VMS deposits demonstrate that the
471 primary source of sulfur is the leaching of igneous basement and TSR of seawater (Ohmoto 1996; Barrie
472 et al. 2009; Keith et al. 2016a; McDonald et al. 2018). By contrast, $\delta^{34}\text{S}$ values in colloform pyrite at Mala
473 are less than the Troodos magmatic mean (0-1‰; Alt 1994) indicating that sulfur was not *only* sourced via
474 TSR of Cretaceous seawater and the leaching of igneous lithologies, instead indicating that colloform pyrite
475 formed during the disproportionation of SO_2 . Using the fractionation between SO_2 and pyrite (Sakai 1968),
476 the expected fractionation at 350°C would be 8.6‰. In order to explain the observed fractionation between
477 the median $\delta^{34}\text{S}$ value in colloform pyrite (2.2‰) and primary igneous sulfur/sulfide (0‰) the fluid
478 temperature would have had to be >880°C. As previously established for bulk rock samples, this
479 temperature is unrealistically high for VMS deposit formation, hence we suggest that the disproportion of
480 SO_2 occurred at lower fluid temperatures where fractionation between SO_2 and pyrite is greater and that the
481 higher than anticipated $\delta^{34}\text{S}$ values in colloform pyrite reflects the addition of a ^{34}S enriched source from
482 both TSR and the leaching of igneous sulfur/sulfide providing an additional source of sulfur in colloform
483 pyrite.

484 Disseminated pyrite grains exhibit the largest variation in $\delta^{34}\text{S}$ values with a median of 0.7‰ and a range
485 of 16.8‰ (n=14). To assess if the pyrite analysed are singular grains or aggregates of sub-grains, etching
486 was used to reveal any internal variation (Fig. 7). Sample MAL 05 (Fig. 7A) exhibits internal variation
487 appearing as three different colored sections, indicating it is formed from three sub-grains. In contrast, the
488 etched surface of MAL 11A and B does not show any notable variation indicating that they are a single
489 continuous grain (Fig. 7B and C).

490 The sulfur isotopic composition of individual pyrite grains is highly variable across disseminated pyrite
491 grains with $\delta^{34}\text{S}$ values ranging from -10.9‰ to 2.5‰ (within a single sub-grain), suggesting that the source
492 of sulfur changed as the pyrite crystal grew. However, there is no systematic variation in sulfur isotope
493 composition between the core and rim of the grain (Fig. 7A), and the two analyses located at the margin of
494 the grain yield $\delta^{34}\text{S}$ values of 2.5‰ and -10.9‰, respectively. Instead, $\delta^{34}\text{S}$ values appear to gradually
495 decrease across the pyrite grain indicating a change in the source of sulfur as the grain grew (Fig. 7A). In
496 other pyrite grains a more systematic pattern is discernable where the rim of the grain is enriched in ^{34}S
497 relative to the core (Fig. 7B and C). This variation could be related to either pulsed fluid flow events
498 (Butterfield et al. 1994, 2011; Jamieson et al. 2013) that generate fluctuations in fluid flux, temperature,
499 and the chemical and physical (e.g., phase separation) composition of hydrothermal fluids (Lalou et al.
500 1990, 1998; Butterfield et al. 1994) or local scale variation in mound-scale fluid flow in response to the
501 collapse and occlusion of permeability pathways during growth of the VMS mound (Klienrock and
502 Humphris 1996; Humphris et al. 1995; Hannington et al. 1998; Petersen et al. 2000).

503 Pulsed fluid flow events are linked to magmatic intrusions at depth within the oceanic crust that provide a
504 renewed source of heat that increases the temperature of venting hydrothermal fluid, volatile influx (e.g.,
505 H_2S and SO_2), and the metal content of fluid vented at the seafloor (Von Damm 1990; Von Damm et al.
506 1995; Butterfield et al. 1997; Andersen et al. 2017). Several studies document temporal variations in total
507 sulfur concentrations and major anions (e.g., Cl^-) in hydrothermal vent fluids sampled from active vent sites
508 demonstrating that the flux of sulfur in VMS deposits varies temporally (Butterfield and Massoth, 1994;
509 Von Damm et al. 1995; Butterfield et al. 1997, 2011; Von Damm 1995). It does however remain enigmatic
510 exactly what these variations mean, for example variations in fluid H_2S concentration could also be
511 produced during the precipitation or dissolution of sulfide minerals below the seafloor (Reed and Palandri
512 2006) or during the segregation of H_2S into a vapor phase, thus linking these variations to magmatic
513 intrusion events is challenging (Von Damm et al. 1995; Butterfield et al. 1997). The dating of SMS deposits
514 ($^{210}\text{Pb}/\text{Pb}$, $^{230}\text{Th}/^{234}\text{U}$ and ^{14}C) also provides evidence of aperiodic fluid flow with apparent pulses of activity
515 recorded at the active TAG mound every 2,000 to 6,000 years (Lalou et al. 1993; You and Bickle 1998).

516 In Troodos VMS deposits there are no hydrothermal fluids left to sample nor are the deposits young enough
517 to be dated using radiometric isotope techniques commonly applied to active SMS deposits, however
518 variation in rare earth element profiles and Sr isotopes ($^{87}\text{Sr}/^{86}\text{Sr}$) in epidote from the sheeted dykes complex
519 and plagiogranites, that formed during hydrothermal alteration of the oceanic crust indicate the episodic
520 release of volatiles into the Troodos hydrothermal systems (Fox et al. 2020). If this was the case then
521 evidence of pulsed magmatic influx may have been recorded on the scale of individual pyrite mineral grains
522 as variations in across grain sulfur isotope composition (Fig. 7). Following a volcanic event or the liberation
523 of magmatic volatiles from the roof of an underlying magma chamber during dyking events, as
524 hypothesized for Troodos (Gillis and Roberts 1999; Fox et al. 2020), the disproportionation of SO_2 occurs
525 and progressively decreases with time from the event, reducing the flux of SO_2 and subsequent
526 disproportionation in the overlying hydrothermal system, and the primary source of sulfur changes from
527 one strongly influenced by disproportionation of SO_2 to one dominated by TSR and leaching of igneous
528 sulfur/sulfide. This would produce a pronounced shift in the sulfur isotopic composition across individual
529 pyrite grains from negative $\delta^{34}\text{S}$ values indicating SO_2 disproportionation to values $>0\text{‰}$ that indicate
530 increasing amounts of sulfur sourced from TSR and the leaching of igneous sulfur/sulfide (e.g., Fig. 7C).
531 A similar model has been proposed by earlier studies to explain the sporadic preservation of $\delta^{34}\text{S}$ values
532 down to -5.5‰ in the Sha VMS deposit (Martin et al. 2020). The zonation and aperiodic nature of these
533 fluctuations in sulfur isotope composition across each mineral grain may be linked to the pulsed nature of
534 fluid flow during VMS deposit formation, however, further high-resolution mineral-scale data including
535 mapping of individual mineral grains is required to ascertain the true significance and validity of mineral-
536 scale sulfur isotope data when applied to system scale pulsed ore-forming processes.

537 [A comparison between bulk and *in situ* sulfur isotope data](#)

538 The sulfur isotope composition of pyrite from the Mala VMS deposit has previously been assessed using
539 bulk analytical methods on powdered wholerock samples or pure mineral concentrates (Fig. 9). Using bulk
540 analytical methods, the $\delta^{34}\text{S}$ values in pyrite range from -7.6‰ to 0.1‰ with a median value of -4.3‰

541 (n=28; Martin et al. 2021) (Fig. 9). Using *in situ* sulfur isotope data collected on the same samples, the
542 sulfur isotope composition of pyrite varies from -17.1‰ to 7.5‰ with a median value of -1.0‰ (n=31);
543 variation larger than for the same samples using bulk analytical techniques (Fig. 9). When comparing
544 between whole rock and *in situ* datasets, on average the difference between the same sample analysed by
545 SIMS and wholerock methods was 4.9‰ (n=6 samples), with some samples, such as MAL 17, exhibiting
546 a difference of 7.3‰ between methods. This difference between analytical scales indicates that a significant
547 amount of variability, and therefore changes in sulfur source and by implication metal source are being
548 overlooked when using bulk analytical methods compared to *in situ* mineral-scale data. This trend is
549 comparable to those recognized at Axial Seamount where the bulk analysis of pyrite yield a range of 1.1 to
550 3.9‰ (Hannington and Scott 1988) whilst *in situ* analysis exhibits more variability with a range of 1.2 to
551 6.9‰ (Crowe and Valley 1992) or at 1-2° N on the East Pacific Rise (EPR) where bulk pyrite analyses have
552 a range of 3.0 to 5.8‰ (Zeng et al. 2017) versus *in situ* analysis of pyrite that ranges from 1.8 to 7.5‰
553 (Meng et al. 2019).

554 To illustrate the disparity between analytical scales we highlight two examples of analytical transects across
555 pyrite grains (n=23 spots, Fig. 10A). Across a coarse-grained (~1 cm) pyrite grain with three sub-grains
556 that occurs within gypsum $\delta^{34}\text{S}$ values exhibit the largest intra-sample range from -10.9‰ to 4.9‰ with a
557 median of 2.0‰ (n=7) (Fig. 10A). Bulk analysis of the same sample yielded $\delta^{34}\text{S}$ values of -0.5‰ and -
558 1.2‰ (Martin et al. 2021). The application of *in situ* analytical methods indicates a variable source of sulfur
559 with values of -10.9‰ indicating dominantly SO_2 disproportionation and values of 4.5‰ signifying a larger
560 contribution of sulfur sourced via TSR of seawater and the leaching of igneous sulfur/sulfide (Fig. 9). A
561 similar difference between analytical methods is evident in all other samples we analysed (Table S3, ESM);
562 demonstrating that data obtained at different analytical scales (i.e. mineral vs. bulk samples) can change the
563 resulting interpretation that is reached.

564 Can magmatic volatile influx be traced at the mineral-scale?

565 Previous studies have identified an enrichment of Te, Se, Au, Bi, As, Sb, Pb and Hg in SMS deposits that
566 form in subduction influenced environments relative to MOR hosted hydrothermal systems, indicating a
567 link between metal enrichment and magmatic volatile degassing (Hannington et al. 1999; de Ronde et al.
568 2011; Berkenbosch et al. 2012; Layton-Matthews et al. 2013; Wohlgemuth-Ueberwasser et al. 2015; Fuchs
569 et al. 2019; Martin et al. 2019; Mathieu 2019; Patten et al. 2020). For example, at the Brothers NW Caldera
570 site (Kermadec arc), a positive correlation exists between decreasing $\delta^{34}\text{S}$ values (to -4.6‰ in sphalerite)
571 and Au concentration, indicating a link between magmatic volatile influx and Au enrichment (de Ronde et
572 al. 2011). If certain metals were sourced from a magmatic volatile phase in the Mala VMS deposit, then
573 they would be expected to show increased concentrations at low $\delta^{34}\text{S}$ values that indicate an increased
574 magmatic volatile influx associated with SO_2 disproportionation (e.g., Herzig et al. 1998). As previously
575 established, sulfur isotopic ratios exhibit notable variation across individual pyrite grains indicating a
576 variation in sulfur source that we ultimately relate to a variable magmatic volatile influx. If magmatic
577 volatile influx occurred as aperiodic releases from the plagiogranites in Troodos into the overlying
578 hydrothermal systems, a coupled relationship between sulfur isotopes and the enrichment of trace metals
579 that are associated with increased levels of volatile influx (e.g., Te and Se) should exist in pyrite (Fox et al.
580 2020). However, our data exhibits limited correlation between Te, Se, Au, As and Co and $\delta^{34}\text{S}$ values
581 indicating that the primary magmatic signature/zonation in trace metals, if initially present, was rapidly
582 modified within the VMS mound.

583 Selenium has been widely applied as an indicator of magmatic volatile influx with Se/S ratios in pyrite used
584 as evidence of an increased magmatic volatile influx in VMS deposits (Yamamoto 1976; Huston et al. 1995;
585 Hannington et al. 1999; Layton-Matthews et al. 2008, 2013; Martin et al. 2019). The contribution of Se
586 from a magmatic volatile phase at Mala is largely supported at the deposit scale by negative $\delta^{34}\text{S}$ values in
587 pyrite and a moderate positive correlation between Te and Se ($R^2=0.67$); a trend that is not observed in any

588 other Troodos VMS deposit (Martin et al. 2021). However, mineral grain scale observations show that a
589 more complex relationship exists (Fig. 10 and 11).

590 If the Se content of pyrite is a reliable proxy for magmatic volatile influx at Mala, then a negative correlation
591 is expected between Se content and $\delta^{34}\text{S}$ values in pyrite, with high Se contents occurring in the samples
592 with low $\delta^{34}\text{S}$ values indicating a coupled relationship (Yamamoto 1976). The Se content of sample MAL
593 05, a disseminated pyrite grain in gypsum (Fig. 10A) ranges from 839 to 1727 ppm with the highest Se
594 concentration corresponding to a $\delta^{34}\text{S}$ value of 2.5‰. The two lowest Se concentrations of 839 and 1273
595 ppm correspond to the highest $\delta^{34}\text{S}$ values measured in MAL 05 of 4.9 and 5.9‰, respectively (Fig. 10A).
596 However, the two lowest $\delta^{34}\text{S}$ values of -2.0 and -10.9‰ do not correspond to the highest Se content. Thus,
597 Se content does not accurately track magmatic volatile influx, if this were the case then a strong correlation
598 between $\delta^{34}\text{S}$ values and Se concentration would occur, however this is not observed (Fig. 11B).

599 Renewed volatile influx is driven by magmatic intrusions at depth below the seafloor that lead to an increase
600 in fluid temperatures preceding the intrusive event, for example at 9° 46.5' N on the East Pacific Rise where
601 vent fluid temperature varied from 403°C directly after the intrusive event to 322°C three years later (Von
602 Damm et al. 1995). High Co concentrations in pyrite have been suggested to indicate elevated fluid
603 temperatures (>350°C), therefore Co should be enriched at low $\delta^{34}\text{S}$ values that precede the intrusive event
604 (Butterfield et al. 1997; Keith et al. 2016a,b; Grant et al. 2018). Arsenic is expected to exhibit the inverse
605 trend as it is concentrated in low temperature pyrite (<350°C) and is expected to be enriched at high $\delta^{34}\text{S}$
606 values representing periods of low magmatic volatile influx and correspondingly cooler fluids (Monecke et
607 al. 2016; Grant et al. 2018). Thus, systematic shifts in Te and Se, that indicate renewed volatile influx, and
608 As and Co that signify changes in fluid temperature should correlate with changes in $\delta^{34}\text{S}$ values in pyrite
609 if they are reliable tracers of magmatic volatile influx at the mineral-scale; we assess this relationship in
610 disseminated pyrite grains at Mala.

611 In sample MAL 05 Co concentration varies from 12 to 23 ppm (Fig. 10A), exhibiting only a minor
612 correlation ($R^2=0.3$) with $\delta^{34}\text{S}$ values (Fig. 10A). Arsenic concentrations are more variable than Co ranging

613 from 5 to 145 ppm with the highest concentration corresponding to a $\delta^{34}\text{S}$ value of -2.0‰, but with no
614 notable correlation between As and $\delta^{34}\text{S}$ values ($R^2 = 0.02$). A similar trend is evident for Te data with an
615 R^2 value of 0.1 (Fig.10 and 11).

616 In other disseminated grains (Fig. 10B; MAL 11A) the highest Se and Co content correspond to the lowest
617 $\delta^{34}\text{S}$ value of 0.2‰, whilst Te exhibits the inverse trend (Fig. 10B). A similar trend is also discernable in
618 MAL 11B where the sulfur isotope composition of pyrite exhibits a saw tooth pattern ranging from -2.5‰
619 to 0.4‰ (Fig. 10C). Selenium and Te concentrations exhibit a moderate correlation with $\delta^{34}\text{S}$ values with
620 an R^2 value of 0.6 and 0.3, respectively. Thus, no clear relationship exists between trace metals and sulfur
621 isotope composition across disseminated pyrite grains at Mala.

622 For all other samples considered in this study, our data indicate that there is no notable correlation between
623 Te, Se and Co with $\delta^{34}\text{S}$ values in individual pyrite grains or within the different textures analysed (Fig.
624 11). The highest measured Se concentration of 3261 ppm corresponds to a $\delta^{34}\text{S}$ value of 0‰, whilst lower
625 values of -2.5‰ within the same mineral grain, which suggest an increased magmatic volatile influx
626 correspond to a Se concentration of 325 ppm (Fig. 11A). We exclude massive pyrite sample MAL 17 from
627 our interpretation that is influenced by BSR of seawater leading to a low $\delta^{34}\text{S}$ value in pyrite and notably
628 lower concentrations of Se, Te and Co relative to massive and disseminated samples (Fig. 11).

629 Our investigation indicates that variations in Te, Se, Co and As concentrations at the mineral-scale cannot
630 be linked to variations in magmatic volatile influx at the Mala VMS deposit (Fig. 12). The lack of
631 correlation between $\delta^{34}\text{S}$ values and Te, Se, Co and As in the Mala VMS deposit probably reflects the
632 localized remobilization of trace elements during zone refining related to fluctuations in mound scale fluid
633 flow related to the collapse and occlusion of fluid pathways during the growth of the mound that leads to
634 localised zone refining of trace metals in pyrite (Fig. 12). Any initial zonation of trace metals in pyrite
635 related to the influx of magmatic volatiles is overprinted by later fluid flow that led to the mobilization of
636 trace metals but does not alter the original sulfur isotope composition of <0‰ related to magmatic volatile
637 influx and SO_2 disproportionation, hence, this explains the lack of correlation between $\delta^{34}\text{S}$ values and Te,

638 Se, Co and As at the mineral-scale at Mala (Fig. 12). To investigate this process, further high-resolution *in*
639 *situ* isotopic and trace element geochemistry and information on the diffusion rates of trace metals and
640 sulfur in pyrite (e.g., Cherniak 2010) is needed to understand chemical zonation at the scale of individual
641 mineral grains.

642 Summary and conclusion

643 The distribution of trace metals between different pyrite textures in the Mala VMS deposit appears broadly
644 analogous to actively forming mafic SMS deposits. Colloform pyrite that formed near the seawater interface
645 is enriched in low temperature elements such as As and Au relative to massive samples that contain euhedral
646 pyrite that are relatively enriched in high temperature metals Se, Co and Te. Granular samples that formed
647 during the reworking of massive pyrite are depleted in all metals (except Te) due to the mobilization and
648 leaching of metals during low temperature fluid flow at the margin of the VMS mound. There is no
649 systematic variation in trace metal enrichment profiles with depth in the VMS mound, instead variation
650 between individual samples can be attributed to dynamic fluid flow patterns within the VMS mound and
651 the localised (cm to m scale) zone refining of metals around individual fluid conduits that vary spatially
652 and temporally during mound growth (Fig. 12). All pyrite textures appear to be enriched in Se relative to
653 both Troodos VMS deposits and mafic hosted VMS deposits more widely indicating a Se-rich magmatic
654 volatile dominated source.

655 The large range in the $\delta^{34}\text{S}$ composition of pyrite relative to previous studies and mafic VMS deposits more
656 widely can be explained by varying combinations of four main processes: i) TSR, ii) leaching of
657 sulfur/sulfide from host rocks, iii) SO_2 disproportionation and iv) BSR. Thermochemical sulfate reduction
658 of seawater and the leaching of igneous sulfur/sulfide generates $\delta^{34}\text{S}$ values in pyrite that are higher than
659 the Troodos magmatic mean of 0-1‰, whilst low values that are less than this are produced during the
660 addition of sulfur from the disproportionation of SO_2 during the degassing of shallow magma chambers.

661 When combined with trace element geochemistry, we suggest low $\delta^{34}\text{S}$ values (to -17.1‰) formed during
662 the BSR of seawater in the upper VMS mound at temperatures of <120°C.

663 Mineral-scale sulfur isotope data exhibits notably more variation than the same samples analysed via bulk
664 rock analytical methods. We observe greater amounts of variation over a single pyrite grain than recorded
665 across all bulk rock analyses with $\delta^{34}\text{S}$ values ranging from -10.9 to 2.5‰. Analytical transects across
666 individual mineral grains indicate that the source of sulfur in the Mala VMS deposit was variable and
667 alternated between disproportionation of SO_2 , TSR and the leaching of igneous sulfur/sulfide as the pyrite
668 grain grew. Such variation is not evident when using bulk analytical techniques for sulfur isotopes and
669 warrants further detailed investigation.

670 Linking variations in trace metal enrichment profiles and sulfur isotopes is key in understanding the role of
671 magmatic volatiles as a potential metal source in mafic VMS deposits. However, findings of this study
672 indicate no simple correlation between magmatic volatile elements such as Te and Se and $\delta^{34}\text{S}$ values. We
673 attribute this decoupled relationship to local scale fluid flow within the mound that leads to the
674 remobilization of trace metals during zone refining. Pyrite retains $\delta^{34}\text{S}$ values that indicate
675 disproportionation (<0‰) but can be depleted in magmatic volatile metals.

676 [Acknowledgements](#)

677 The authors acknowledge the support of the Geological Survey Department of Cyprus, especially Costas
678 Costantinou and Andreas Zissimos. We thank Michael Green and Ifigenia Gavriel for discussion and
679 assistance in the field. This research was partly funded by the NERC SoS consortium grant
680 NE/M010848/1 “TeaSe: tellurium and selenium cycling and supply” awarded to Cardiff University and by
681 the Canadian Research Chair program awarded to John W. Jamieson. We thank Stefanie Brueckner and
682 associate editor Mostafa Fayek for their constructive reviews and editor-in-chief Georges Beaudoin for the
683 editorial handling of this manuscript.

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986 Figure Captions

987 Figure 1: Simplified geological map of the Troodos ophiolite, Cyprus. Graben axes are indicated by the
988 dashed lines. Mala is located in the SW of the ophiolite at the LPL/BG transitional horizon (after Martin et
989 al. 2018).

990 Figure 2: Field observations from the Mala VMS deposit. A) The exposed portion of the Mala mound. The
991 VMS mound contains pyrite and crudely layered gypsum capped by a veneer of leached volcanic rocks.
992 The margin of the VMS mound is denoted by the white dashed line. B) Coarse-grained euhedral pyrite with
993 infilling gypsum (white). C) Massive pyrite. D) Massive gypsum horizon located in the upper mound region
994 containing coarse (1-2 cm) euhedral pyrite grains. E) Gypsum containing finely disseminated euhedral
995 pyrite. F) Euhedral pyrite in a gypsum matrix with minor Fe oxide staining. G) Aggregates of euhedral
996 pyrite in gypsum.

997 Figure 3: Photomicrographs in reflected light of common pyrite textures at Mala. A) Massive pyrite. Note
998 porous inclusion trails delineating crystal growth zones. B) The relationship between massive and dendritic
999 pyrite textures. C) Dendritic pyrite. D) Colloform pyrite in a matrix of porous pyrite. E) Granular pyrite
1000 consisting of anhedral rounded pyrite in a matrix of Fe-oxides and pyrite. F) Finely disseminated euhedral
1001 to subhedral pyrite in surrounding altered volcanic rocks.

1002 Figure 4: Pyrite chemistry analysed via LA-ICP-MS. Pyrite analyses are divided based on grain morphology
1003 into disseminated (n=78), massive (n=40), granular (n=10) and colloform (n=15). Analyses that are below
1004 detection limit are excluded (Table S1, ESM). A) Te vs. Co, B) Te vs. Se, C) Co vs. Se. D) Total measured

1005 trace metal by pyrite texture (Co, Ni, Cu, Zn, As, Se, Ag, Cd, Sb, Te, Re, Au, Pb and Bi). Data in ESM,
1006 Table S1.

1007 Figure 5: Pyrite chemistry analysed via LA-ICP-MS from the core (n=35) and rim (n=43) of individual
1008 disseminated pyrite grains. The rim of pyrite grains are enriched in Co and Cu and relative to the margin.
1009 Data in ESM, Table S1.

1010 Figure 6: Summary of sulfur isotope analyses ($\delta^{34}\text{S}$) analysed by SIMS (n=31) and classified based on pyrite
1011 texture. Massive pyrite exhibits the largest range in its $\delta^{34}\text{S}$ composition whilst colloform pyrite clusters
1012 between 0 and -5‰. Data and standard information in ESM, Table S3.

1013 Figure 7: SIMS spot analyses across disseminated pyrite grains. Lower images are the same grain etched
1014 with NaOCl to reveal any internal zonations. A) Sample MAL 05 (in the lower image “SG” refers to
1015 different sub-grains), B) sample MAL 11-A, C) MAL 11-B.

1016 Figure 8: LA-ICP-MS analyses for individual mineral grains. A) Se vs. As, B) Se vs. Te, C) Se vs. Co. Data
1017 in Table S1 and sample description in Figure S1, ESM.

1018 Figure 9: A comparison between bulk $\delta^{34}\text{S}$ analyses of Mala samples (Martin et al. 2021) with the same
1019 sub-set of samples analysed by SIMS in this study. Samples analysed by SIMS exhibit notably more
1020 variation with extremely low (-17.1‰) and high (7.5‰) values compared to the samples analysed using
1021 bulk methods. Cretaceous seawater 18 to 19‰ (SW) (Kampschulte and Strauss 2004) and Troodos
1022 magmatic mean 0 to 1‰ (TO magmatic mean; Alt 1994).

1023 Figure 10: Coupled trace metal and $\delta^{34}\text{S}$ analyses across individual disseminated pyrite grains. A) Sample
1024 MAL 05. B) Sample MAL 11 A. C) Sample MAL 11 B. P1 on X axis refers to the specific analytical point
1025 in the image above. Selenium, As, Co and Te concentrations are shown by red lines with corresponding
1026 concentration in ppm on the secondary axis (right). Note varying Y axis between graphs.

1027 Figure 11: Se, Te and Co vs. $\delta^{34}\text{S}$ for all samples. Twinned points in disseminated pyrite (as in Fig. 10) are
1028 represented by multiple analytical points. In samples where points were not twinned an average value is
1029 reported and denoted by * next to each sample in the legend. A) Se vs. $\delta^{34}\text{S}$, B) Te vs. $\delta^{34}\text{S}$, C) Co vs. $\delta^{34}\text{S}$.
1030 Volatile trend in plot B shows expected trend between Te, Se and Co and $\delta^{34}\text{S}$. The grey box represents the
1031 primary magmatic mean for Troodos magmatic rocks (Alt 1994). Solid line represents the average bulk
1032 $\delta^{34}\text{S}$ for all Troodos VMS (4.6‰; Martin et al. 2020) and the dashed line is the bulk median $\delta^{34}\text{S}$ for Mala
1033 (-4.3‰; Martin et al. 2021).

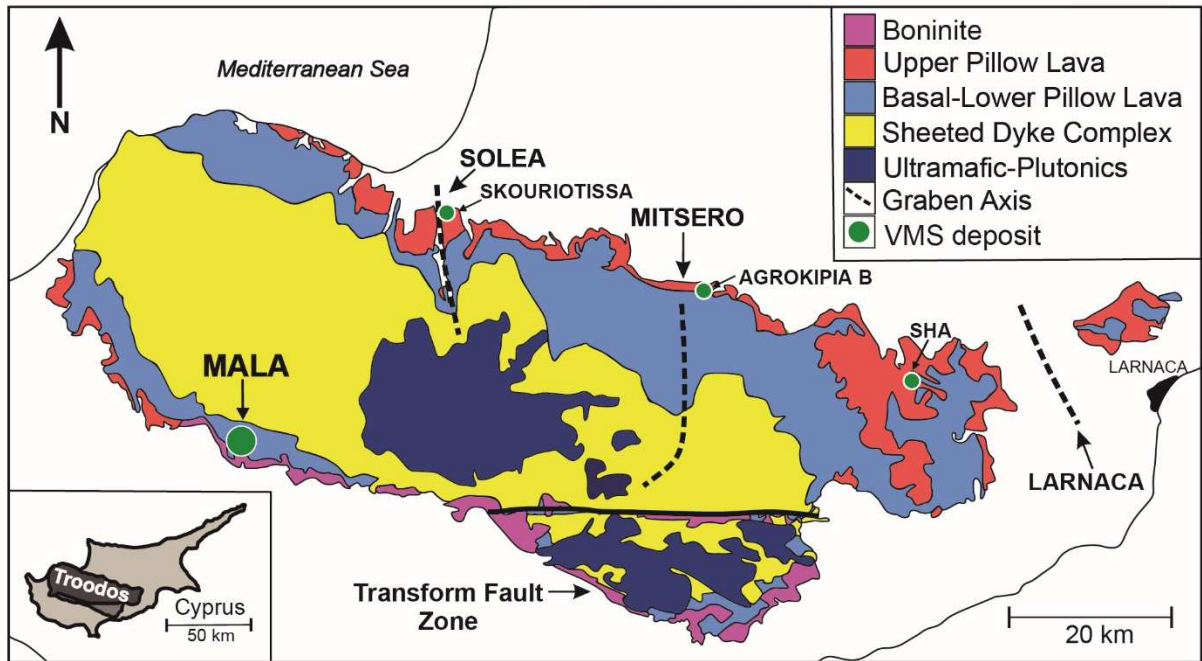
1034 Figure 12: A) Summary schematic for mound scale metal enrichment processes at the Mala VMS deposit.
1035 Granular pyrite forms from the reworking and collapse of the outer mound and chimney material, metals
1036 are remobilized during low temperature fluid flow. Disseminated pyrite forms in gypsum veins and volcanic
1037 rocks (Fig. 2). B) Colloform and dendritic textured pyrite (Fig. 3 C and D) form at the margin of the sulfide
1038 mound where seawater ingress is high creating disequilibrium textures and fluid temperatures are lower
1039 (<350°C) leading to an enrichment in Ag, Au and Pb. C) With increasing time, permeability pathways
1040 within the mound change in response to the collapse and reworking of anhydrite, creating a prominent
1041 brecciated texture leading to localised zone refining on the cm/m scale in response to new high temperature
1042 fluid pathways. D) Pyrite grains undergo zone refining leading to the decoupling of $\delta^{34}\text{S}$ and magmatic
1043 volatile elements in pyrite.

1044 Table Captions

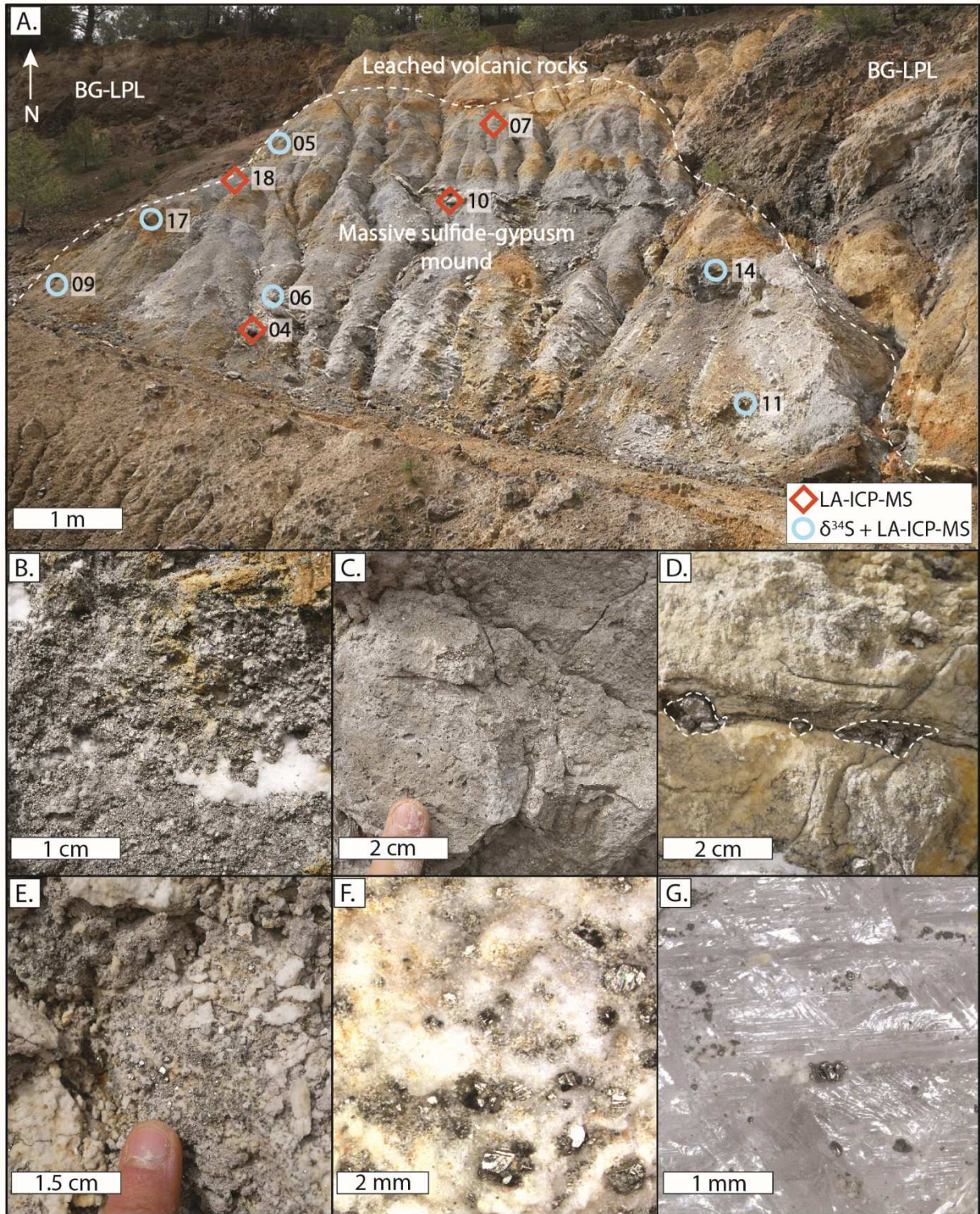
1045 Table 1: Summary of pyrite geochemistry classified by texture: colloform (n=15), granular (n=10),
1046 massive (n=40) and disseminated (n=78). Points that were below detection limit (Table S1 and S2, ESM)
1047 are excluded (n > DL = number of analyses above detection limit).

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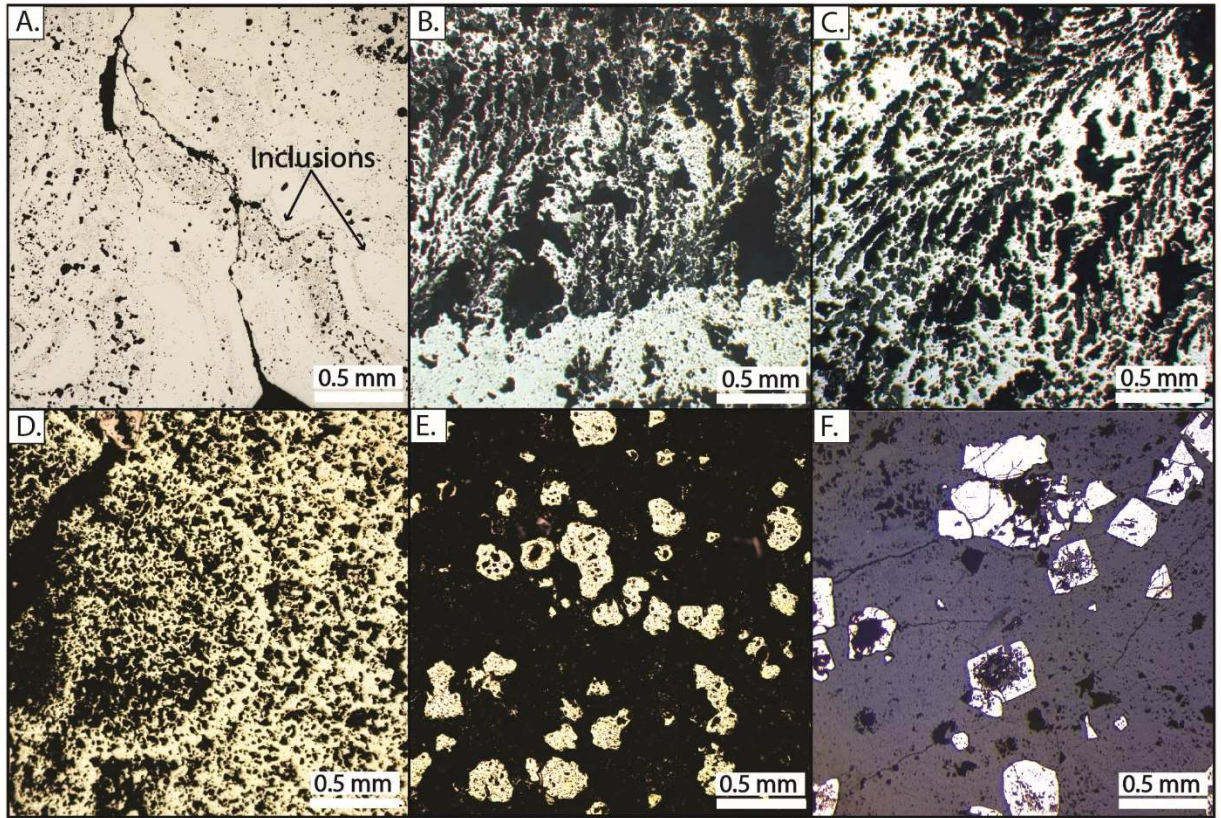
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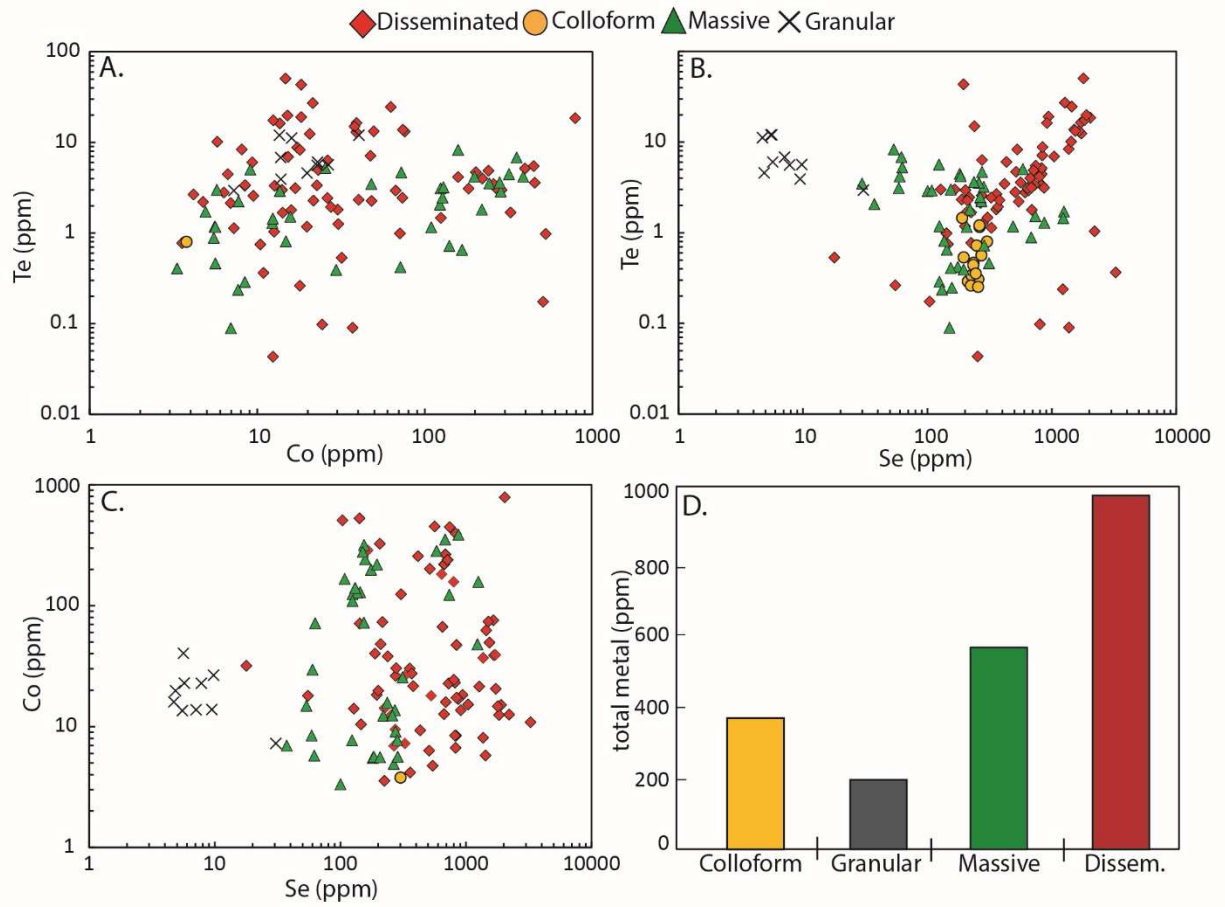
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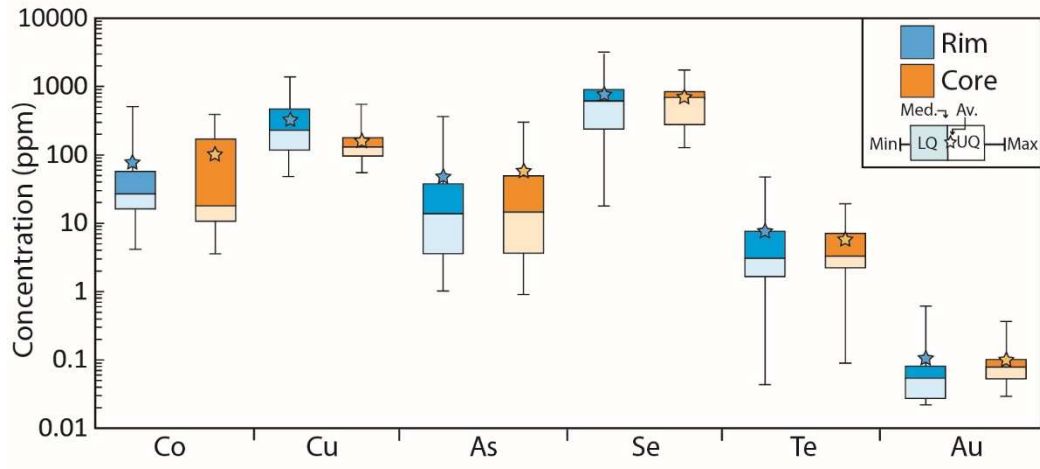
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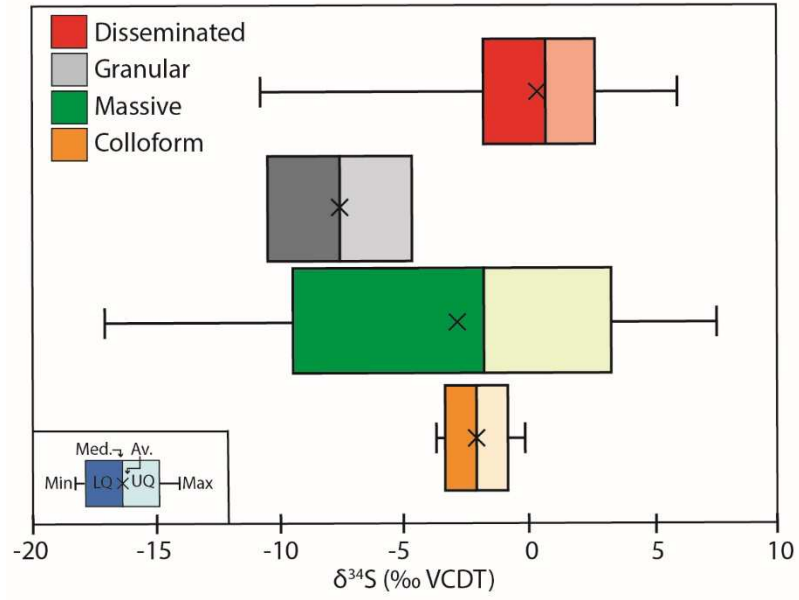
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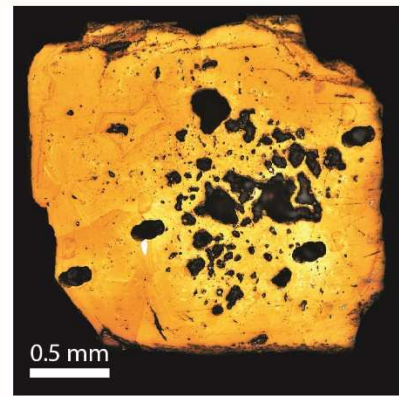
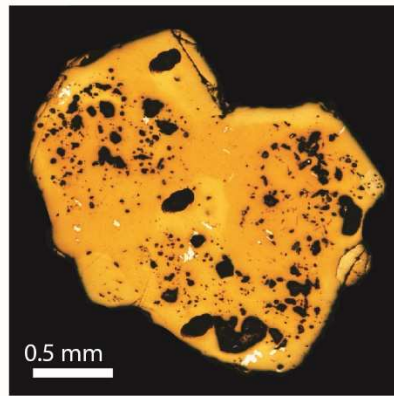
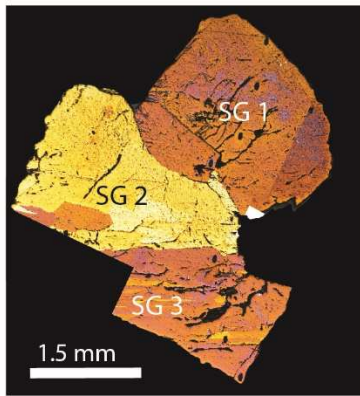
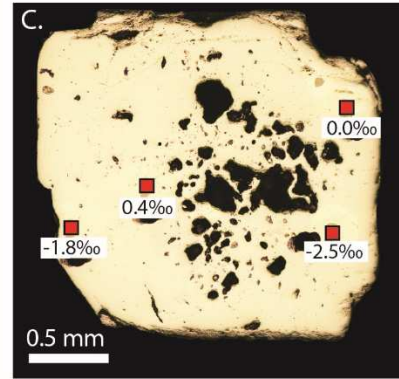
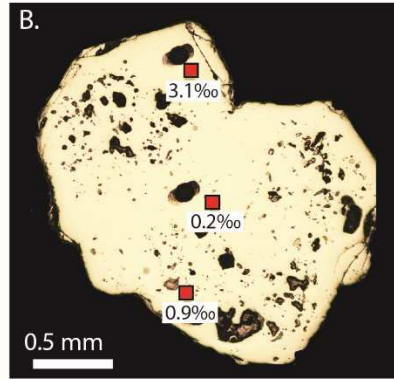
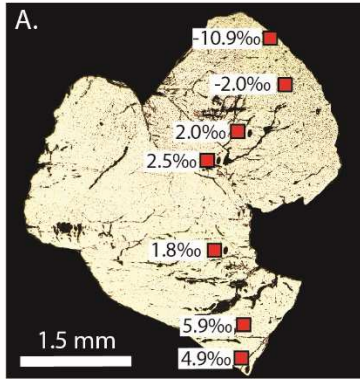
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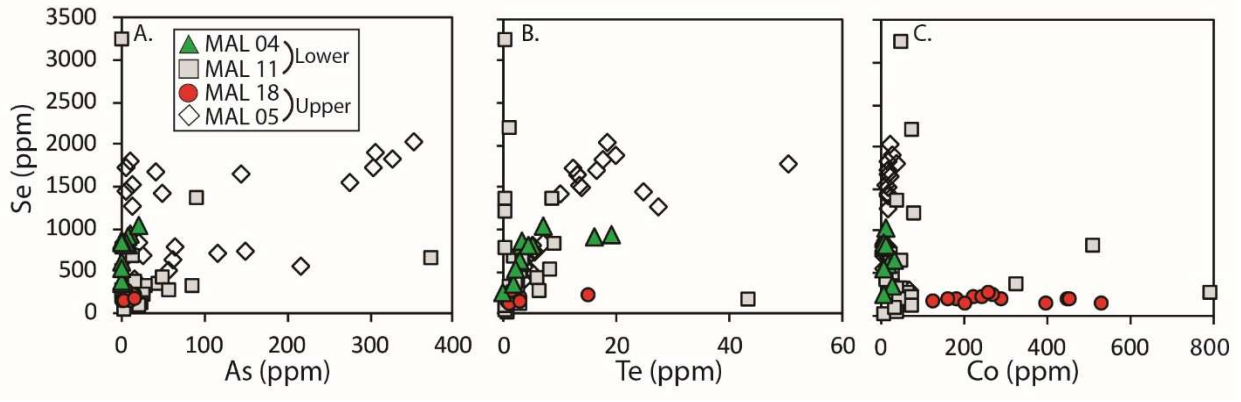
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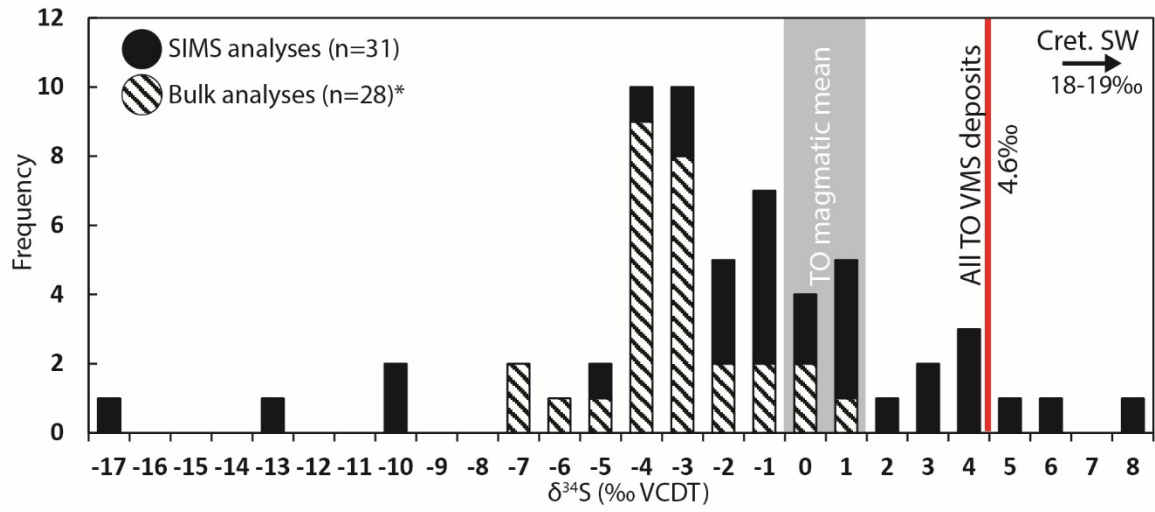
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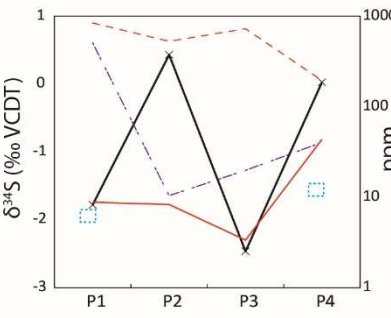
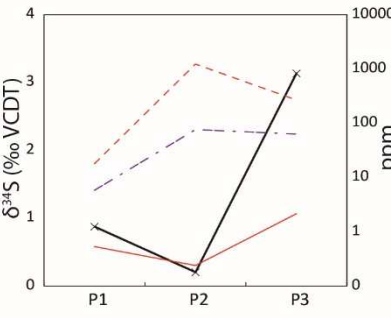
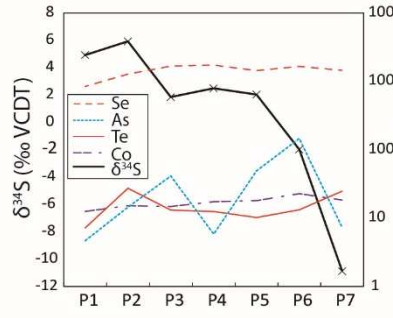
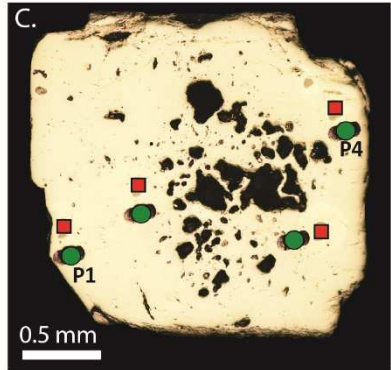
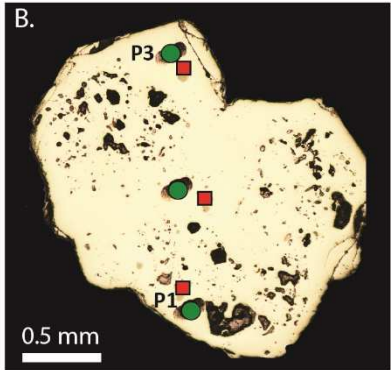
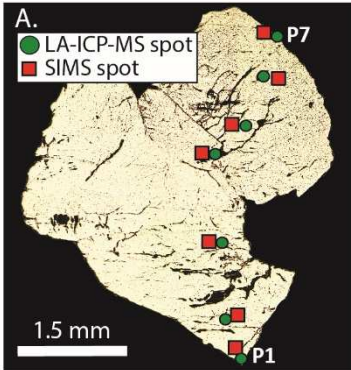
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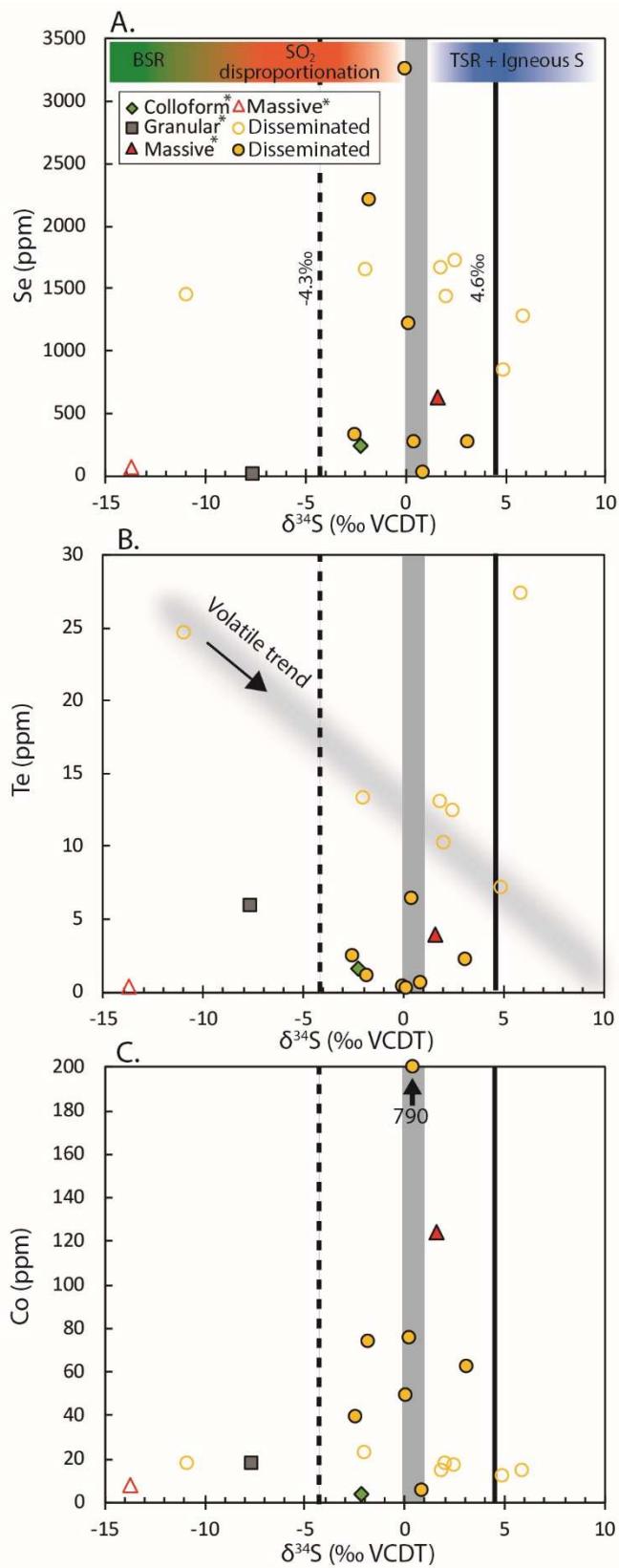
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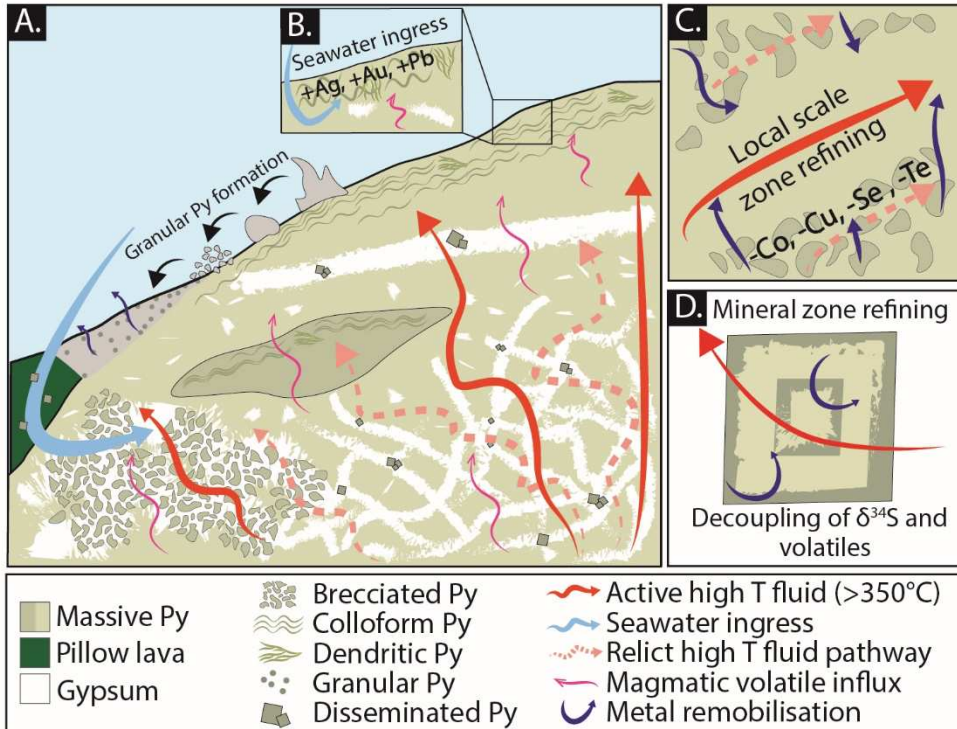


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		Colloform				Granular				Massive				Disseminated			
		Median	Min	Max	n > DL	Median	Min	Max	n > DL	Median	Min	Max	n > DL	Median	Min	Max	n > DL
Co	ppm			3.78	1	17.9	7.25	40.5	10	38.8	3.33	386.5	38	23.1	3.56	789.6	75
Ni	wt. %				0				0			0.01	1	0.01	0.01	0.07	12
Cu	wt. %	0.01	0.01	0.13	4		0.01	0.02	2	0.02	0.01	0.11	13	0.02	0.01	0.15	31
Zn	ppm		6.73	9.01	2				0	11.2	9.44	38.8	5	14.04	10.0	17.7	8
As	ppm	4.18	1.55	24.0	15	2.84	1.37	4.72	7	3.14	1.18	94.7	26	13.6	0.90	373.6	68
Se	ppm	244.6	195.9	301.7	15	6.40	4.72	30.6	10	178.3	29.8	1252.7	40	644.9	17.9	3260.7	78
Ag	ppm	0.24	0.15	0.49	15				0	0.42	0.16	0.89	11	0.24	0.19	0.33	7
Cd	ppm				0				0			0.26	1			0.67	1
Sb	ppm	0.47	0.25	1.46	15			0.35	1	0.97	0.16	5.76	14	0.25	0.22	1.14	3
Te	ppm	1.51	0.86	3.13	15	5.86	2.94	12.0	10	1.76	0.09	8.30	40	3.14	0.04	50.6	78
Re	ppm	0.17	0.11	0.25	15				0	0.15	0.03	0.83	17	0.05	0.01	1.80	35
Au	ppm	0.13	0.08	0.25	15		0.03	0.03	2	0.06	0.02	0.24	24	0.06	0.01	0.60	24
Pb	ppm	9.83	6.94	15.1	15				0	7.01	3.25	46.5	13	5.27	0.49	25.6	8
Bi	ppm	0.05	0.03	0.12	15			0.03	1	0.08	0.03	0.26	22	0.12	0.03	0.47	19

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