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5 **Seafloor massive sulfides from mid-ocean ridges: Exploring the causes of their**
6 **geochemical variability with multivariate analysis**

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17 **Highlights:**

- 18 • The geochemical variability of massive sulfides from mid-ocean ridges is explored with
19 multivariate statistical analysis.
- 20 • The observed variability is mostly related to the temperature of deposition, the ridge
21 spreading rate and zone refining.

22 • The nature of the substrate (mafic vs. ultramafic rocks) seems to play a secondary role in
23 controlling metal associations.

24

25 **Abstract**

26 The neovolcanic zones of mid-ocean ridges are host to seawater-derived hydrothermal systems
27 forming seafloor massive sulfide (SMS) deposits. These deposits have high concentrations of
28 base metals and potentially economic enrichment of a wide range of trace elements. The factors
29 controlling this enrichment are currently poorly understood. We have investigated the main
30 factors controlling SMS compositional variability through robust principal component analysis
31 and robust factor analysis of published and newly obtained bulk geochemical data for samples
32 collected from SMS deposits worldwide. We found that a large part of the observed variability is
33 produced by a combination of three independent factors, which are interpreted to reflect (in order
34 of importance): (1) the temperature of deposition, (2) the ridge spreading rate, and (3) zone
35 refining. The first and the third factors are mostly related to processes operating near the
36 seafloor, such as conductive cooling, mixing of the hydrothermal fluids with seawater and metal
37 remobilization, and determine the relative proportions of the main minerals and, thus, of Cu and
38 Zn (Co, Se, Sb, Pb). The ridge spreading rate influences the structure of the oceanic lithosphere,
39 which exerts a major control on the length and depth of the hydrothermal convection cell and on
40 the rock-to-water ratios in the reaction zone, which in turn control the behavior of the precious
41 metals Au and Ag and elements including Ni (Mo, Se). Despite the obvious role of substrate
42 rocks as metal sources, their composition (specifically mafic vs. ultramafic) does not emerge as a
43 statistically significant independent factor.

44

45 **Keywords:** seafloor massive sulfides, mid-ocean ridges, geochemistry, multivariate analysis

46 **1 Introduction**

47 Seafloor massive sulfides (SMS) are stratiform or stratabound accumulations of base
48 metal sulfides that formed on or near the seafloor by precipitation from dominantly seawater-
49 derived hydrothermal fluids (Hannington, 2014). Since their discovery in 1978 at the East Pacific
50 Rise (Francheteau et al., 1979), SMS deposits have been recognized along many neovolcanic
51 plate boundaries, including mid-ocean ridges, submarine arc volcanoes and back-arc spreading
52 centers. In all these environments, magma intrusions sustain hydrothermal convective cells, in
53 which percolating seawater leaches metals from the substrate rocks (Sleep, 1983; Tivey, 2007).
54 Upon ascent, metals precipitate primarily as sulfides below and on the seafloor, as the hot (up to
55 ~400 °C) and acidic (pH at 25 °C as low as ~3) hydrothermal fluid quenches in the presence of
56 cold (~2 °C) and alkaline (pH at 25 °C ~ 8) seawater. Considering the total length of neovolcanic
57 plate boundaries, SMS deposits could represent a significant source of metals, estimated at
58 $\sim 3 \times 10^7$ tonnes of Cu + Zn (Hannington et al., 2011). Moreover, some deposits have remarkable
59 grades of gold and silver (up to several tens of ppm; Petersen and Hein, 2013). Unraveling the
60 factors that affect the geochemistry of SMS is important to the understanding of their genesis and
61 for developing effective guidelines for the exploration and economic evaluation of both present-
62 day seafloor deposits and their ancient on-land analogues.

63 Observations on SMS deposits show that the geochemistry of the massive sulfides is
64 primarily related to the geodynamic setting in which they are formed. On mid-ocean ridges,
65 where the hydrothermal fluids dominantly leach mafic or ultramafic substrates, SMS typically
66 have high concentrations of Cu + Zn (\gg Pb) (Hannington et al., 2005). The leaching of elements
67 from substrate rocks is influenced by the structure of the oceanic lithosphere and by the nature of
68 the hydrothermal convection, which are essentially related to the spreading rate (Bougault et al.,

69 1993; Bach and Humphris, 1999; Coumou et al., 2009). On *slow-spreading* ridges, for instance,
70 the magma supply is low and part of the extension is accommodated by deep-rooted detachment
71 faults. Therefore, the lithosphere is not layered and is mainly composed of gabbros and
72 ultramafic rocks (MacLeod et al., 2009; Lowell, 2010). In this setting, low heat fluxes and deep-
73 rooted faults favor long and deep fluid pathways, which enhance fluid-rock interaction at
74 relatively high rock-to-water (r/w) ratios (Bach and Humphris, 1999). By contrast, on *fast-*
75 *spreading* ridges, the magma supply is high, the lithosphere is typically layered and the upper
76 basaltic section is highly permeable (Coumou et al., 2008). The resulting higher heat flux,
77 together with episodic diking events, favor the development of shallower and ephemeral
78 hydrothermal convection cells (Wilcock and Delaney, 1996). Therefore, fast-spreading ridges are
79 typified by less evolved seawater-dominated fluids (i.e., lower r/w ratios). Irrespective of
80 spreading rate, SMS deposits show significant geochemical variability along the same ridge and
81 even at the deposit or hand-specimen scale (Petersen and Hein, 2013), suggesting a complex
82 interplay of regional and local controls. In fact, some specific geochemical features (e.g., Au
83 enrichment and Au/Ag and Co/Ni ratios) have been variably related to the nature of the substrate
84 (e.g., mafic vs. ultramafic rocks), phase separation processes in the fluid, the presence of a
85 magmatic influx of volatiles and metals, the morphology of vent structures (tubular vs. beehive
86 chimneys generated by focused or diffuse fluid flow, respectively), the ridge spreading rate, or a
87 combination of these factors (Marques et al., 2006; Fouquet et al., 2010; German et al., 2016;
88 Melekestseva et al., 2017; Knight et al., 2018). However, the relative role of these factors in
89 controlling the geochemical variability of SMS deposits is still poorly defined. Part of this
90 uncertainty is due the fact that the nature of the substrate is inferred from two-dimensional
91 seafloor observations and different lithologies may occur in the areas surrounding the deposits.

92 Also, individual deposits may show a mixture of apparent geochemical signatures (e.g., ‘mafic’
93 and ‘ultramafic’; Marques et al., 2006; Webber et al., 2015; Melekestseva et al., 2017).
94 Moreover, experimental works on rock/seawater interactions under conditions typical of
95 subseafloor reaction zones so far mainly explored the behavior of only a few base metals (Fe,
96 Mn, Zn, Cu, Ni) in basalt-seawater systems at r/w ratios ranging from ~0.001 to 2 (Seyfried and
97 Bischoff, 1977, 1981; Seyfried and Mottl, 1982; Seyfried and Janecky, 1985; Seewald and
98 Seyfried, 1990). Experiments on peridotite-seawater systems investigated even fewer base metals
99 (Fe, Mn, Ni, Zn) and only at $r/w < 0.1$ and $T \leq 300$ °C (Seyfried and Dibble, 1980; Janecky and
100 Seyfried, 1986). Forward thermodynamic modeling of rock-seawater reactions and hydrothermal
101 fluid cooling has been used to investigate the behavior of a larger number of elements in seafloor
102 hydrothermal systems (Janecky and Seyfried, 1984; McCollom and Shock, 1998; Wetzel and
103 Shock, 2000; Palandri and Reed, 2004; Klein et al., 2013; Melekestseva et al., 2014, 2017; Fuchs
104 et al., 2019). However, the resulting models are specific to particular rock compositions reacting
105 under fixed conditions or to particular fluid compositions and are not generally applicable to all
106 hydrothermal systems on mid-ocean ridges.

107 In this work, we investigate the first-order controls on the compositions of SMS deposits
108 on mid-ocean ridges by means of multivariate statistical analysis of published and novel bulk
109 chemical analyses of SMS samples from seafloor hydrothermal fields worldwide. Attempts to
110 apply multivariate statistical analysis to SMS at various scales were previously made by
111 Hannington et al. (1991a) and Fouquet et al. (2010). Hannington et al. (1991a) applied factor
112 analysis to mafic-hosted deposits worldwide and found important geochemical associations for
113 Sb-Pb-Au-Ag and Mo-Co-Se-Cu-Fe, reflecting metal associations of low- and high-temperature
114 mineral assemblages, respectively. Fouquet et al. (2010) used principal component analysis to

115 identify site-specific geochemical groups (i.e., Cu-rich high-temperature, Zn-rich intermediate-
116 temperature, and oxide/sulfate, respectively) in ultramafic-hosted deposits at the Mid-Atlantic
117 Ridge. Compared with these previous works, the dataset considered here is more comprehensive
118 as it is built on four times the sample number analyzed by Hannington et al. (1991a), covers
119 more ridges and includes data from both mafic- and ultramafic-hosted SMS deposits. Moreover,
120 in this work we use robust principal component analysis (rPCA) and robust factor analysis (rFA)
121 to limit the influence of noise from individual anomalous samples. This is particularly useful,
122 because analyzed SMS materials mostly consist of dredged samples and are certainly affected by
123 sampling bias (Fuchs et al., 2019). Therefore, our analysis provides a better assessment of the
124 first-order factors that control the geochemical variability of ridge-hosted SMS at a global scale.
125 By comparing the results with existing experimental data and new thermodynamic models, we
126 show that temperature of deposition, seafloor spreading rate and r/w ratio, and zone refining are
127 the major factors controlling metal associations in SMS deposits and that, contrary to common
128 belief, the nature of the substrate may be of secondary importance.

129 **2 Materials and Methods**

130 Based on an initial database by Hannington et al. (2004), we have compiled published
131 and unpublished bulk chemical analyses of sulfides samples collected from SMS deposits located
132 on mid-ocean ridges. New analyses of samples from the Irinovskoe, Krasnov and Peterburgskoe
133 SMS fields at the Mid-Atlantic Ridge (Table 1) were acquired at the South Urals Federal
134 Research Center, Urals Branch of Russian Academy of Sciences, Institute of Mineralogy (Miass,
135 Russia) using atomic absorption (AA) analysis in air-acetylene flame on a Perkin Elmer 3110
136 spectrometer (Co, Ni, Cu, Zn, Pb, Au, Ag) and inductively coupled plasma-mass spectrometry
137 (ICP MS) on an Agilent 7700x mass spectrometer (Sb, Se, Mo). For AA analyses of Co, Ni, Cu,

138 Zn and Pb, 0.5–1.0 g of sample material was dissolved in a 15-ml mixture of HF, HCl, and
139 HNO₃, heated first up to 100–150°C and then to 250°C, and concentrated to the formation of wet
140 salts. This procedure was conducted three times, then the samples were dried, cooled, mixed with
141 3 ml concentrated HNO₃, and filtered. For AA analyses of Au and Ag, 1 g of sample material
142 was diluted in a mixture of 30 ml HCl and 10 ml HNO₃, concentrated on the oven to the
143 formation of wet salts, filtered using HCl solution and cooled. For ICP-MS analyses, the sample
144 material was initially digested in Teflon autoclaves using a mixture of HF, HCl and HNO₃ in a
145 SpeedWave microwave digestion system (Berghof, Germany) using a two-stage heating
146 procedure to a temperature of up to 180°C for 40 minutes. After digestion, the fluorine
147 complexes were decomposed by double evaporation of the dry residual with concentrated HNO₃
148 at 110°C in glassy carbon crucibles. The precipitates were further dissolved in hot 0.5 N HNO₃
149 and subsequently reduced to a 100-ml aliquot. All pure acids used for digestion were purified in
150 a BSB-939-IR apparatus (Berghof, Germany). The water for dilution was deionized in a Milli-
151 Q[®] Integral Water Purification System by Millipore, US. Quality control of the analytical
152 procedure was performed by analysis of state reference materials (GSO) of water metal solutions
153 (7256-96 Zn, 7252-96 Pb, 7265-96 Ni, 7268-96 Co, 7255-96 Cu, and 8402-2002 Ag) and
154 flotation concentrate of Au-bearing ore CZK-3 (2739-83 Au) for AA analysis. Indium was used
155 as an internal standard calibrated against the USGS BCR-2 rock reference material for ICP MS
156 analyses.

157 For the statistical analysis, we only considered reportedly fresh SMS samples
158 representing fragments of hydrothermal chimneys and mounds. We excluded samples which
159 contained significant proportions of weathering products or silicate gangue (e.g., Al₂O₃ and MgO
160 contents > 2 wt.%). The presence of silica phases, which do not typically contain significant

161 trace elements, was not considered critical. Since the chemical analyses were heterogeneous in
162 terms of analyzed elements and their detection limits, we only selected those records that
163 provided data for a sufficiently large number of elements (i.e., Au, Ag, Co, Mo, Ni, Sb, Se, Cu,
164 Zn, Pb). Barite-rich (Ba > 1 wt.%) samples were excluded to avoid possible bias caused by
165 elements (Pb, Co, Ni, Sb) incorporated in or adsorbed on barite (cf. Melekesteva et al., 2014;
166 Safina et al., 2016). Analyses reported only as averages of several samples were discarded, since
167 these data were not accompanied by adequate mineralogical and textural descriptions of
168 individual samples. These choices allowed us to maintain a balance between the number of
169 elements (N = 10), the number of records (N = 426) and the number of represented hydrothermal
170 sites or clusters (N = 26; Table 2). For instance, due to the incompleteness of many published
171 analyses, adding a single element such as As would have decreased the number of represented
172 sites to 25 (the Beebe hydrothermal field would be completely excluded). The SMS deposits
173 were grouped on the basis of the ridge spreading rate [fast, intermediate, and (ultra)slow] and
174 host-rock composition (mafic or ultramafic). The complete database (Table S1) includes samples
175 from 3 sites on fast, 8 sites on intermediate, 12 sites on slow, and 3 sites on ultraslow spreading
176 ridges (Table 2). The distinction between mafic-hosted and ultramafic-hosted deposits was not
177 always straightforward, because both types of rocks generally occur in the areas surrounding and
178 probably beneath nominally ultramafic-hosted deposits. In this work, deposits classified as
179 ultramafic-hosted ($N_{\text{ultramafic}} = 6$ out of a total of 26) are those sited on substrates that, based on
180 reported geological evidence, are likely to contain abundant ultramafic rocks (Table 2). The

181 resulting mafic-hosted vs. ultramafic-hosted classification is the same as that used in the recent
182 review by Fuchs et al. (2019).

183 The statistical relationships between elements were explored by using both non-
184 parametric and parametric techniques for dimension reduction, i.e., rPCA and rFA, respectively.
185 The former technique has the advantage of not being influenced by the user's decisions and is,
186 therefore, more appropriate at an exploratory stage. The latter technique is dependent on the
187 number of chosen factors, but, unlike rPCA, it does not force the factors to explain all the
188 variability (Reimann et al., 2008). Therefore, rFA may be more effective in defining statistical
189 factors that are representative of common geochemical processes (Reimann et al., 2002).

190 Accordingly, we used rPCA to guide rFA, in the sense that the number of principal components
191 that explained most of the variability in rPCA were used to determine the proper number of
192 factors for rFA (Reimann et al., 2008). The statistical analyses were performed with the R
193 software using the function "pcaCoDa" in the "robCompositions" library (Templ et al., 2011)
194 and the R script (principal factor analysis with a varimax rotation) by Filzmoser et al. (2009a).
195 Robust methods were preferred to "classical" methods, because they are less sensitive to outliers
196 (Filzmoser et al., 2009b; Filzmoser and Hron, 2011), which are common in geochemical data.
197 Isometric logratio and centered logratio transformations were applied to the data used for rPCA
198 and rFA, respectively. These transformations have the advantage of opening the data, but require
199 that no zeros are present in the data matrix. Therefore, we represented concentrations below the
200 detection limit by multiplying the detection limit by 0.65. This choice is statistically appropriate

201 since only 5.3% of the data are below detection limit (Palarea-Albaladejo and Martín-Fernández,
202 2015; Martín-Fernández et al., 2003).

203 To obtain independent constraints for the interpretation of the statistical data, we
204 simulated the basalt–seawater and peridotite–seawater reactions in a model subseafloor
205 hydrothermal reaction zone. Thermodynamic calculations were performed by Gibbs energy
206 minimization using the Selektor program (Karpov et al., 1997; Chudnenko, 2010). Rock–
207 seawater reactions were modeled at 400°C and 30 MPa assuming various r/w ratios. Complete
208 details on the modeling procedure and the choice of thermodynamic data are reported in
209 Melekestseva et al. (2017). The compositions of basaltic glass and peridotite were taken from
210 Lehnert et al. (2000) and Fouquet et al. (2010), respectively. Molybdenum content in basalt (0.31
211 ppm) and seawater (1.04×10^{-7} mol/kg H₂O) were taken from Fouquet et al. (2010) and Steele et
212 al. (2010), respectively. Seawater and reactant rock compositions are reported in Table S2.

213 **3 Results**

214 3.1 Element distribution

215 The distribution of elements in mid-ocean ridge SMS is reported in Table 2 and
216 illustrated by box-and-whiskers plots in Fig. 1. Median element concentrations include
217 ~3 wt.% Cu, ~1 wt.% Zn, ~0.02 wt.% Pb, ~270 ppb Au, ~110 ppm Co, ~60 ppm Se, ~50
218 ppm Mo, ~20 ppm Ag, ~10 ppm Ni, and ~5 ppm Sb.

219 3.2 Robust PCA and FA

220 The rPCA was used to determine the number of factors suitable to describe the
221 compositional variability of the SMS deposits. The results of the rPCA are illustrated in

222 the scree plot in Fig. 2 (see Fig. S1 for complete score plot). The line connecting the
223 eigenvalues shows a progressive decrease in slope from PC1 to PC4 and only minor
224 changes beyond PC3. Moreover, the first three components together explain most of the
225 variability (~75%). For these reasons, three factors were chosen in the subsequent
226 factorial analysis. As shown in Fig. 3, the most important variables in the first factor (F1)
227 are Pb and Sb (plus minor contributions by Zn and Ag) with positive loadings and Cu and
228 Se (Co) with negative loadings. The second factor (F2) is mainly related to Au and Ag
229 (negative loadings) and more weakly affected by Mo, Se and Ni (positive loadings). The
230 third factor (F3) is mostly influenced by Zn (Sb, Ag) (positive loadings) and Co (Mo)
231 (negative loadings). The score plots in Fig. 4 show the distribution of the samples in the
232 space defined by the factors. Figure S2 shows the same results for individual sites. The
233 samples are scattered through the plots, which indicates that rFA was effective in
234 detecting the directions of maximum variability. Most remarkably, the F1 vs. F2 plot
235 (Fig. 4A) shows that the factor F2 [Se-Mo-Ni vs. Au-Ag] discriminates well between
236 SMS samples from slow-spreading ridges (Mid-Atlantic Ridge, Mid-Cayman Rise,
237 Central Indian Ridge) and those from intermediate/fast-spreading ridges (Southern
238 Explorer Ridge, Juan de Fuca Ridge and Galapagos Rift). The great majority (87%) of
239 the samples from intermediate/fast-spreading ridges show positive F2 loadings and only
240 7% of them show F2 loadings <-0.1 . Most of these low-F2 samples are from the
241 sedimented, intermediate-spreading Gorda Ridge (Escanaba Trough) and Juan de Fuca
242 Ridge (Middle Valley) (Fig. S2A). The great majority (84%) of the samples from slow-
243 spreading ridges show negative F2 values and only 4% of them show F2 values $>+0.1$.
244 No particular cluster is apparent in the F1 vs. F3 plot (Fig. 4B), although all but one

245 samples from the Escanaba Trough and all samples from the mafic-hosted
246 Petersburgkoe field of the slow-spreading Mid Atlantic Ridge show more or less
247 negative F3 scores (Fig. 4B and Fig. S2B). The F2 vs. F3 plot (Fig. S3) does not show
248 any additional remarkable features and only confirms the good discrimination between
249 SMS from intermediate/fast- and slow-spreading ridges. Concerning samples from slow-
250 spreading ridges (low F2), the ultramafic-hosted SMS have, on average, lower values of
251 F1 relative to the mafic-hosted SMS (Fig. 4A). Nonetheless, the ranges of factors F1, F2
252 and F3 covered by the two groups of samples are similar and no clear distinction between
253 mafic- and ultramafic-hosted deposits is apparent in either plots (Fig. 4A,B). Since the
254 ultramafic-hosted deposits are undersampled in our database ($N_{\text{ultramafic}} = 64$; $N_{\text{mafic}} =$
255 362), it may be argued that these results are biased due to the disproportionate weight of
256 the mafic-hosted deposits. To test this possibility, we have performed rFA on a reduced
257 data set in which the mafic records were randomly selected to match the number of the
258 ultramafic records (Fig. S4). The resulting statistical factors F1 and F2 are very similar in
259 terms of both element associations and relative factor weights to those obtained using the
260 complete database (cf. Fig. 4). The discrimination between fast/intermediate-spreading
261 and slow-spreading ridges is somewhat less effective, probably due to the smaller number
262 of input data, but there is still substantial overlap between ultramafic-hosted and mafic-
263 hosted deposits from slow-spreading ridges. This confirms the robustness of our rFA
264 analysis.

265 3.3 Thermodynamic modeling

266 The complete results of thermodynamic modelling of basalt/seawater and
267 peridotite/seawater reactions at 400°C, 30 MPa, and log(r/w) ratios ranging between -5

268 and 0 are shown in Fig. S5 and Fig. S6. The most relevant results are summarized in Fig.
269 5. The first consequence of basalt/seawater and peridotite/seawater reactions is the
270 development of alteration mineral assemblages, the nature of which in turn affects the
271 composition of the hydrothermal fluid (Fig. S5 and S6).

272 In the basalt/seawater system, at very low $\log(r/w)$ (-5 to -3.1) the secondary
273 mineral assemblage includes chrysotile, Mg chlorite, anhydrite, hematite, and titaniferous
274 oxides, plus brucite at $\log(r/w) < -3.6$ (Fig. 5A). Within this $\log(r/w)$ range, the seawater-
275 buffered system is characterized by a high redox potential ($E_{h400^\circ\text{C}} \sim 0.2$ V). The $\text{pH}_{400^\circ\text{C}}$
276 values decrease from ~ 7.7 at $\log(r/w) -5$ to ~ 6.3 at $\log(r/w) -3.1$. With increasing
277 $\log(r/w)$, the formation of abundant secondary silicates (talc, quartz, amesite, actinolite,
278 Mg-Mn chlorite, epidote, albite) and chromite produces a jump in fluid $\text{pH}_{400^\circ\text{C}}$ to ~ 6.7
279 and then a further progressive increase to ~ 7.4 at $\log(r/w) 0$. At the same time, $E_{h400^\circ\text{C}}$
280 drops to -0.4 V at $\log(r/w) -3$ and then further decreases to below -0.6 V (Fig. 5A). At
281 $\log(r/w) > -1.8$ hematite is replaced by magnetite. At $\log(r/w) > -1.3$, when $E_{h400^\circ\text{C}}$ goes
282 below ~ -0.6 V ($\log f_{\text{O}_2} \sim -25$), several sulfides are also formed (Fig. 5A). The computed
283 alteration assemblages are consistent with natural and calculated assemblages in altered
284 oceanic basalts (Alt et al., 1986; McCollom and Shock, 1998; Wetzel and Shock, 2000).
285 Figure 5A shows the calculated concentrations of several elements in the fluid as a
286 function of the r/w ratio. The Fe and, in part, the Si contents are closely related to the Eh
287 and pH values. The chalcophile elements accumulate in the fluid until their respective
288 sulfides are stabilized in the altered rock, and then their concentrations in the fluid
289 decrease. A subsequent minor increase of Co at $\log(r/w) > -0.3$ is related to the
290 disappearance of Co-bearing pyrite. The change in the Ni curve at $\log(r/w) -2.6$ coincides

291 with a change in the Ni contents of amesite, whereas that at $\log(r/w) -1.7$ is related to the
292 disappearance of Ni-bearing amesite. Formation of Ni-bearing sulfides at $\log(r/w) -1.2$
293 produces a slight decrease in Ni concentrations to $\log(r/w) 0$. Gold and Ag progressively
294 accumulate in the fluid with increasing r/w ratios.

295 In the peridotite/seawater system, the secondary mineral assemblage is dominated
296 by chrysotile at all r/w ratios (Fig. S6). At $\log(r/w) < -4$ the seawater-buffered system is
297 characterized by relatively high $E_{h400^\circ C}$ (~ 0.1 V) and $pH_{400^\circ C}$ (~ 7.8). With increasing
298 $\log(r/w)$, the stabilization of titaniferous oxides is accompanied by a slight decrease in pH
299 to 6.8 at $\log(r/w) -2.6$. With further increasing $\log(r/w)$, the pH variations are
300 complicated due to the formation and disappearance of various secondary silicates
301 (chlorite, phlogopite–biotite s.s., talc, actinolite) and show an overall increase, reaching a
302 value of ~ 7.8 at $\log(r/w) 0$. The $E_{h400^\circ C}$ values show a nearly flat profile to $\log(r/w) -3.3$,
303 then a marked drop to -0.5 V and a further decrease to below -0.8 V at $\log(r/w) 0$ (Fig.
304 5B). Hematite is part of the alteration assemblage at $\log(r/w) < -2.5$ and is replaced by
305 magnetite at higher $\log(r/w)$. At $\log(r/w) > -2$, when $E_{h400^\circ C}$ goes below -0.6 V ($\log fO_2 \sim$
306 -25), several sulfides are also formed (Fig. 5B). The computed alteration assemblages are
307 fairly consistent with natural and calculated assemblages in altered oceanic peridotites
308 (Wetzel and Shock, 2000; Palandri and Reed, 2004; Klein and Bach, 2009; Klein et al.,
309 2013). Element contents in the fluid follow broadly similar trends vs $\log(r/w)$ as in the
310 basalt/seawater system (Fig. 5). Fe and Si contents are closely related to variations of Eh,
311 pH and silicate mineralogy, Au and Ag progressively accumulate in the fluid, and the
312 concentrations of the chalcophile elements increase until their respective sulfides are
313 formed. The stabilization of (Co, Ni)-bearing sulfides occurs at somewhat lower $\log(r/w)$

314 than in the basalt/seawater system. The $Eh_{400^\circ C}$ and fO_2 profiles are not strongly
315 dissimilar in the two systems, except at $\log(r/w)$ ratios >-0.5 , where the
316 peridotite/seawater system becomes more strongly reduced.

317 **4 Discussion**

318 4.1 The effect of temperature

319 Factor F1, which explains 49% of the variability, is dominated by the anti-
320 correlated groups of elements Cu-Se-(Co) and Pb-Sb-(Zn-Ag) (Fig. 3 and 4). We
321 interpret this anti-correlation to reflect variations in the formation temperature of the
322 mineral assemblages analyzed. Because chalcopyrite solubility in vent fluids drops below
323 350–375 °C, higher temperature fluids precipitate chalcopyrite-rich (i.e., Cu-rich)
324 assemblages, while lower temperature fluids precipitate Pb-Sb-bearing assemblages with
325 variable proportions of galena and sulfosalts (Janecky and Seyfried, 1984; Seyfried and
326 Ding, 1995). The strong correlation between Se and Cu reflects the ability of Se to
327 replace S in high-temperature chalcopyrite (Huston et al., 1995). Cobalt, which as a
328 divalent cation can substitute for Fe^{2+} and Zn^{2+} , can also be structurally hosted in
329 chalcopyrite, as well as in high-temperature (> 300 °C) Fe-sulfides (Vaughan and Rosso,
330 2006; Cook et al., 2009; Grant et al., 2018). Galena and sulfosalts are often present only
331 as (sub-)microscopic inclusions in low-temperature assemblages (e.g., Wohlgemuth-
332 Ueberwasser et al., 2015; Grant et al., 2018) and may thus not have been reported in
333 sample descriptions summarized in Table S1. Nonetheless, Grant et al. (2018) showed
334 that significant concentrations of Pb, Ag and Sb in pyrite from relatively cold and distal
335 parts of the TAG sulfide mound were in fact related to inclusions of galena and sulfosalts.

336 The small positive loading of Zn in factor F1 is consistent with the tendency of sphalerite
337 to precipitate at intermediate to low temperatures ($T < \sim 300$ °C; Janecky and Seyfried,
338 1984). The association of Sb with Pb and Zn may in part reflect substitution of Sb for Pb
339 in galena (Sharp and Buseck, 2003) or for Zn in sphalerite (Maslennikov et al., 2017).

340 The depositional temperature is mostly controlled by conductive cooling of the
341 hydrothermal fluids or mixing with cold seawater and can vary in time and space at the
342 scales of individual hydrothermal fields (e.g., among different vents or as an effect of
343 waning hydrothermal activity) and even samples (cf. the zonal structure of sulfide
344 chimneys, showing chalcopyrite-rich inner portions formed at higher temperatures and
345 pyrite-sphalerite-rich external portions formed at lower temperatures) (Hannington,
346 2014). This explains why samples from the same SMS deposit may plot at opposite ends
347 with respect to factor F1 (Fig. S2). The tendency of samples from ultramafic-hosted SMS
348 to show lower F1 loadings than mafic-hosted SMS from the same ridges may reflect the
349 more diffuse high-temperature discharge and, thus, the widespread enrichment in high-
350 temperature elements at the surface of these deposits (Fouquet et al., 2010).

351 4.2 The influence of ridge spreading rate

352 Factor F2, which explains 17% of the variability, provides a discrimination
353 between Au-Ag-rich SMS from slow-spreading ridges and Au-Ag-poor, but relatively
354 Ni-Mo-Se-rich, SMS from intermediate/fast-spreading ridges (Fig. 4A). This is in line
355 with the observation by Knight et al. (2018) that the Au content in seafloor SMS is
356 negatively correlated with the ridge spreading rate. Knight et al. (2018) suggested several
357 possible explanations for this geochemical distinction, including the intensity and

358 duration of fluid-rock interaction, variations in the Au content in the source rocks and
359 differences in fluid chemistry. Here we discuss the possible causes in the light of existing
360 geochemical data and of our statistical analysis and geochemical modeling.

361 The composition of the substrate has been invoked as an important factor in
362 controlling Au enrichment. However, the absolute average Au concentration in ultramafic
363 rocks (0.49 ppb) is not much higher than that in mid-ocean ridge basalts (MORBs) (0.34
364 ppb) and is unlikely to determine a significant difference in the amount of Au dissolved
365 in the hydrothermal fluids (Fouquet et al., 2010; Melekestseva et al., 2017; Fuchs et al.,
366 2019). Thus, even assuming a metal contribution from ultramafic rocks for all Au-rich
367 SMS from slow-spreading ridges, the Au content in the substrate alone cannot explain
368 their distinct geochemistry.

369 In our database, the samples with the lowest F2 scores are those from the
370 ultramafic-hosted Irinovskoe, Kairei, Logatchev 1 and Rainbow sites and from the mafic-
371 hosted Beebe and TAG sites. The fluids venting at Kairei, Logatchev 1 and Rainbow are
372 relatively reduced (Kairei: $\log fO_2, 350^\circ C \sim -31$ and $H_2 \sim 5$ mM; Logatchev 1: $\log fO_2, 350^\circ C$
373 ~ -32.3 and $H_2 \sim 12$ mM; Rainbow: $\log fO_2, 350^\circ C \sim -33.6$ and $H_2 \sim 16$ mM; Charlou et al.,
374 2002; Kumagai et al., 2008; Seyfried et al., 2011; Kawasumi and Chiba, 2017) and
375 similar redox states are expected for fluids from Beebe as they have similar H_2
376 concentrations ($H_2 \sim 20$ mM; McDermott et al., 2018). Higher fO_2 are reported for TAG
377 vent fluids ($\log fO_2, 350^\circ C \sim -29$; $H_2 \sim 0.2$ mM; Charlou et al., 1996; Kawasumi and Chiba,
378 2017), which have a redox state within the range of vent fluids from other mafic-hosted
379 sites on the East Pacific Rise ($\log fO_2, 350^\circ C \sim -28$ to -29 , $H_2 \sim 0.1$ mM, at $13^\circ N$; $\log fO_2,$
380 $350^\circ C \sim -29$ to -31 , $H_2 \sim 0.4$ mM, at $21^\circ N$; Tivey, 1995; Pester et al., 2011; Kawasumi

381 and Chiba, 2017), Juan de Fuca ($\log f_{\text{O}_2, 350^\circ\text{C}} \sim -29$ to -30 , $\text{H}_2 \sim 0.4$ mM, at Main
382 Endeavour Field; Seewald et al., 2003; Kawasumi and Chiba, 2017) and Central Indian
383 Ridge ($\log f_{\text{O}_2, 350^\circ\text{C}} \sim -28$ to -29 , $\text{H}_2 \sim 0.2$ mM, at Edmond Hydrothermal Field; Gallant
384 and Von Damm, 2006; Kumagai et al., 2008 Kawasumi and Chiba, 2017). Detailed data
385 on fluid chemistry are not available for Irinovskoe. A plot of Au vs. H_2 (Fig. 6) shows
386 that the median Au concentration is generally higher in SMS from slow-spreading ridges
387 and that the highest values are observed at sites where the vent fluids are the most
388 reduced, suggesting a genetic link between Au enrichment and fluid redox state. Note
389 that the number of SMS samples with reported Au concentrations that could be plotted in
390 Figure 6 for the highly variable TAG is much larger than that included in the filtered
391 database used for the multivariate rPCA and rFA (cf. Table 2 and S1), which justifies its
392 relatively low median Au (Fig. 6) in spite of its relatively low F2 scores (Fig. 4). Fluids
393 venting at sites with the highest Au and H_2 contents range from vapor-dominated (e.g.,
394 Beebe; McDermott et al., 2018) to seawater- (e.g., Logatchev; Charlou et al., 2002) or
395 brine-dominated (e.g., Kairei and Rainbow; Charlou et al., 2002; Gallant and Von Damm,
396 2006). The variable chlorinities of fluids vented at sites characterized by similar Au
397 endowment suggest that phase separation is not the primary factor in determining the
398 overall Au enrichment.

399 The apparent correlation between Au enrichment and redox state must be
400 considered cautiously. In fact, present-day vent fluid compositions may not be fully
401 representative of a specific site, as vent fluid composition can considerably vary in time
402 and space even within the same hydrothermal field (Lowell et al., 1995). This can
403 partially explain why sulfide samples from TAG, for which the most comprehensive

404 sample set is available, also show the largest compositional variability (Fig. 6). More
405 generally, relations based on absolute element concentrations are unavoidably affected by
406 sampling bias. More robust information can be obtained by considering the geochemical
407 relations of Au with other elements and particularly with those having large positive or
408 negative loadings in factor F2 (i.e., Ni, Mo, Se, and Ag).

409 In the oceanic lithosphere, Ni, Mo, Se, Au, and Ag are typically stored in sulfides
410 (Patten et al., 2016; Holwell et al., 2017). Therefore, the presence of these elements in
411 seafloor hydrothermal fluids is strictly related to alteration and re-precipitation of sulfides
412 in the substrate rocks. A high temperature in the reaction zone may enhance Au (and H₂)
413 concentrations in the fluid (McDermott et al., 2018), but it would also increase Ni
414 solubility (Liu et al., 2012). Therefore, reaction temperature cannot explain factor F2, in
415 which Au and Ni are anti-correlated (Figs. 3 and 4). Fuchs et al. (2019) showed that, in
416 ultramafic-dominated systems, strongly reduced hydrothermal fluids such as those
417 produced by peridotite serpentinization promote the deposition of Au at relatively high
418 temperatures. This may support a link between fluid redox state and Au endowment (cf.
419 Fig. 6). However, the similar F2 scores shown by mafic- and ultramafic-hosted SMS
420 from slow-spreading ridges (Fig. 4A) suggest that Au enrichment and redox conditions
421 may be largely independent of seafloor substrate composition. Thermodynamic modeling
422 of basalt–seawater and peridotite–seawater reactions shows that, at a given temperature,
423 the concentrations of metals in the fluid are chiefly controlled by r/w ratios and sulfide
424 solubility (Fig. 5). At low r/w ratios, relatively oxidized seawater-dominated fluids favor
425 sulfide dissolution (Palandri and Reed, 2004; Liu et al., 2012; Holwell et al., 2017;
426 Melekestseva et al., 2017). The enrichment in Ni relative to Au and Ag in SMS deposits

427 from intermediate/fast-spreading ridges may thus be an effect of the enhanced dissolution
428 of Ni-rich magmatic sulfides. Higher r/w ratios, which should be more typical of slow-
429 spreading ridges, would instead stabilize sulfides that may segregate Ni while the
430 concentrations of dissolved Au and Ag progressively increase (Fig. 5; see also
431 Melekestseva et al., 2017). High r/w ratios can potentially also be achieved in reaction
432 zones beneath sedimented ridges, where the low-porosity sediment cover may limit
433 seawater fluxes through the oceanic crust. This may explain why most of the samples
434 from the intermediate-spreading, but sedimented, Gorda and Juan de Fuca ridges have
435 low F2 scores similar to samples from slow-spreading ridges (Fig. 4A). Several other
436 elements, which, based on thermodynamic modeling, should also be sensitive to r/w
437 ratios (Fig. 5), do not appear to be controlled by factor F2 (Pb, Sb, Cu, Zn, Co) or have
438 small F2 loadings (< 0.5) which are opposite in sign to expectations (Se, Mo). This
439 apparent inconsistency can be explained by the fact that most of these elements are more
440 strongly controlled by other factors, such as depositional temperature (Pb, Sb, Cu, Se,
441 Co) and zone refining processes (Zn, Co; see below). Therefore, the influence of r/w ratio
442 on these elements may not be apparent in factor F2. As for Mo and Se, their model
443 concentrations at high r/w ratios (Fig. 5) are probably overestimated, because the model
444 does not allow for their incorporation in alteration silicates and sulfides, which may
445 contain significant traces of these elements (Wedepohl, 1969). Expecting an enrichment
446 in these elements at high r/w ratios is therefore unwarranted. Our thermodynamic models
447 do not consider possible kinetic effects, which may strongly influence fluid chemistry in
448 ultramafic-hosted systems at high temperature (Allen and Seyfried, 2003). However, they

449 suggest a possible mechanism for (Au, Ag)–Ni decoupling, which may be effective at
450 least under near-equilibrium conditions in both mafic and ultramafic substrates.

451 It is noteworthy that in both basalt–seawater and peridotite–seawater systems, fO_2
452 is predicted to decrease at high r/w ratios as a consequence of alteration of ferrous Fe-
453 bearing silicates (Fig. 5). Thus, the association between high dissolved H_2 , negative F2
454 loading and high Au grade in many hydrothermal sites on slow-spreading ridges (Fig. 4A
455 and 6) may at least in part be a consequence of reactions between seawater and substrate
456 rocks at relatively high r/w ratios rather than of reactions of seawater with specific
457 lithologies. The dispersion of F2 values for both slow- and intermediate/fast-spreading
458 ridges (Fig. 4A) reveals intra-ridge or even within-site geochemical variations, which can
459 be ascribed to minor changes in space and time of hydrothermal circuits and r/w ratios. A
460 similar interpretation was proposed by Bach and Humphris (1999) to explain variations in
461 Sr and O isotope compositions of seafloor hydrothermal fluids.

462 Based on the results of thermodynamic modeling at 400 °C, the r/w ratios that
463 would be the most significant in determining the observed compositional variability
464 range from ~0.002 to 1, i.e., from $\log(r/w) -2.7$ to $\log(r/w) 0$ (Fig. 5). Within this range, a
465 steady increase of dissolved Au and Ag is accompanied by constant or even decreasing
466 Ni concentrations due to its incorporation into secondary silicates and sulfides. This
467 range is compatible with the r/w ratios <1 to $\ll 1$ estimated for seafloor hydrothermal
468 systems from experiments on hydrothermal alteration of seafloor rocks (Seyfried et al.,
469 1988). Note that our r/w ratios refer to conditions in the reaction zone and thus are not
470 directly comparable with the ‘integrated’ ratios calculated from geochemical data on vent

471 fluids (B and Sr isotopes, Li/Cl ratios) and from geophysical data, which suggest a typical
472 range of ~0.3–2 (Mottl, 2003; Seyfried and Shanks, 2004; Barker et al., 2010).

473 4.3 The effect of zone refining

474 Factor F3 explains 9% of the variability and discriminates Zn-rich, Co-Mo-poor
475 assemblages from those that are poor in Zn but relatively enriched in Co and Mo (Fig. 3
476 and 4B). We interpret this anti-correlation to be related to zone refining processes.
477 During zone refining, mixing of hydrothermal fluid with seawater within the sulfide
478 mound leads to dissolution of earlier Zn-sulfides and precipitation of pyrite, anhydrite
479 and chalcopyrite, producing moderate temperature fluids enriched in Zn (and Au, Ag, Sb,
480 Pb, and Cd). These fluids may vent from white smokers, in which abundant sphalerite is
481 re-precipitated (Edmond et al., 1995; Tivey et al., 1995; Hannington, 2014). Consistently,
482 samples of white smoker chimneys from TAG show high F3 scores (Fig. S2). Cobalt and,
483 especially, Mo are mainly stored in Fe sulfides, and pyrite in particular (up to several
484 thousand or several hundred ppm, respectively; Maslennikov et al., 2009; Keith et al.,
485 2016). Chalcopyrite can also host significant Co (up to several thousand ppm;
486 Maslennikov et al., 2017). Therefore, Co and Mo are likely to remain relatively immobile
487 during zone refining and, consistently, they show negative F3 scores. On the contrary, Ag
488 and Sb, which have small positive loadings in F3, can have high concentrations in
489 sphalerite (hundreds or tens of ppm respectively; Wang et al., 2018) and may be
490 significantly remobilized and, eventually, re-concentrated in newly formed sphalerite
491 near the mound surface. In fact, sphalerite from TAG white smokers have high
492 concentrations of both Ag and Sb (up to >1000 ppm and several hundred ppm,

493 respectively; Tivey et al., 1995). Thus, F3 is likely to represent the effect of metal
494 remobilization on the geochemistry of the hydrothermal fluid.

495 The relatively Zn-rich compositions of some samples from the Main Field on the
496 Juan de Fuca Ridge, which fall in the upper left quadrant in Fig. 4B at high F1 and F3
497 loadings, may have a different origin. They have been ascribed to precipitation from
498 relatively high-temperature but (Cu, Co)-depleted fluids, characterized by a pH higher
499 than typical black smokers due to anomalously high concentrations of ammonia, possibly
500 reflecting hidden organic-rich sedimentary rocks at depth (Tivey et al., 1999). These
501 fluids precipitate Cu-Fe sulfides and then abundant Zn sulfides on conductive cooling
502 within diffusely venting structures. This particular scenario may have produced
503 geochemical associations in the final SMS deposit that are partly similar to those
504 determined by zone refining at other sites.

505 4.4 The role of phase separation, magmatic input and substrate composition

506 The results of our statistical analysis can be explained by factors such as
507 temperature of deposition, ridge spreading rate and varying r/w ratios in the reaction
508 zone, and metal remobilization. If our interpretation is correct, then processes such as
509 phase separation and the input of magmatic fluids, which have sometimes been invoked
510 to explain the compositions of seafloor vent fluids and hydrothermal precipitates
511 (Douville et al., 2002; Melekestseva et al., 2014, 2017; Tivey, 2007), do not play a major

512 role in determining the overall geochemical variability of SMS on mid-ocean ridges.

513 However, this does not mean that these processes do not occur.

514 Phase separation is believed to be commonplace in seafloor hydrothermal systems
515 (Coumou et al., 2009). On mid-ocean ridges, it may give rise to vent fluids with different
516 chlorinities and absolute metal contents (Charlou et al., 2002; Hannington et al., 2005;
517 Fouquet et al., 2010; Seyfried et al., 2011). However, experimental studies show that
518 element fractionation due to phase separation strongly depends on sulfur and chloride
519 concentrations (Nagaseki and Hayashi, 2008; Pokrovski et al., 2005, 2008) and will be
520 predictably small at the concentrations typical of seafloor vent fluids (H_2S : 0.5–13 mM,
521 Cl^- : 420–950 mM; Fouquet et al., 2010; Kawasumi and Chiba, 2017). Thus, although in
522 seafloor hydrothermal systems phase separation may certainly influence the absolute
523 concentrations of metals in the fluids, their fractionation between vapor and liquid phases
524 may not be such as to determine significant changes in the final metal-to-metal ratios,
525 which mainly remain controlled by other higher-order factors.

526 A magmatic fluid input has been specifically invoked to explain the unusual
527 characteristics of some SMS deposits at mid-ocean ridges, such as an anomalous
528 abundance of barite or of silica and Au–Ag (Melekestseva et al., 2014, 2017). Our results
529 are not inconsistent with this explanation. First, barite-rich samples were intentionally
530 discarded in the present work. Second, our statistical analysis is sensitive to overall

531 statistical trends, but does not discriminate outliers. Therefore, contributions from
532 magmatic fluids that were significant only on a local scale could not be resolved.

533 One of the most intriguing and unexpected results of this work is that the
534 composition of the substrate does not emerge as a primary factor in determining metal
535 associations in SMS at mid-ocean ridges. We do not deny a possible role of the substrate
536 in producing specific compositional features, but we suggest that, at a global scale, other
537 factors may explain a large proportion of the observed geochemical variability. The fact
538 that ultramafic substrates are restricted to (ultra)slow-spreading ridges may have
539 obscured the role of spreading rate and r/w ratios in many previous evaluations. This may
540 have led to potentially circular arguments, in which the occurrence of particular
541 geochemical features, believed to be ultramafic- (or mafic-) related, in a mafic- (or
542 ultramafic-) hosted deposit were ascribed to the involvement of concealed ultramafic (or
543 mafic) rocks at depth (Marques et al., 2006; Wang et al., 2014; Webber et al., 2015).
544 Some of these features might actually be more strongly related to the r/w ratios and,
545 hence, to the ridge spreading rate.

546 For instance, it has been argued that the Co/Ni ratio of SMS can be used to
547 discriminate between mafic-dominated and ultramafic-dominated hydrothermal systems
548 (Marques et al., 2006; Zaccarini and Garuti, 2008; Melekestseva et al., 2013). This
549 hypothesis is consistent with the fact that the Co/Ni ratio is significantly higher in
550 MORBs than in ultramafic rocks (0.35 vs. 0.05, calculated from data compiled in
551 Fouquet et al., 2010). Figure 7 shows the relationships between Ni and Co in mafic- and
552 ultramafic-hosted SMS deposits from slow- and intermediate/fast-spreading ridges. The
553 various groups overlap significantly. Thus, although the nature of the substrate (mafic on

554 intermediate/fast-spreading ridges and mafic and ultramafic on slow-spreading ridges) is
555 likely to exert some control on the distribution of Ni and Co in SMS deposits, the Co/Ni
556 ratio overall appears to have little discriminatory power. This suggests that other factors,
557 namely, depositional temperature, r/w ratios in the reaction zone, and zone refining,
558 which may influence to various extent the concentrations of Co and Ni (i.e., F1, F2 and
559 F3 in Fig. 4), may mask the effects of substrate composition when explored through a
560 simple bivariate plot. However, the Co/Ni ratio may be more sensitive in the case of
561 subseafloor stockwork and replacive mineralization, where the local contribution of Ni
562 from ultramafic host-rocks may be significant (cf. Marques et al., 2007). This type of
563 mineralization is not the focus of this work and was in fact excluded from our analysis.

564 Selenium has also been suggested as a potential ultramafic marker based on its
565 higher average content in ultramafic-hosted SMS (Melekestseva et al., 2017), but the
566 ranges of Se concentrations are similar for the mafic- and the ultramafic-hosted deposits
567 (Fuchs et al., 2019). Based on our statistical analysis, Se is strongly related to factor F1,
568 i.e., to the temperature of deposition (Fig. 4). Similar to what suggested by Fouquet et al.
569 (2010) for Cu, the higher average content of Se in sampled ultramafic-hosted SMS may
570 reflect the more diffuse high-temperature discharge and, thus, the widespread enrichment
571 in high-temperature elements at the surface of these deposits.

572 Other features that seem to be typical of ultramafic-hosted SMS cannot be easily
573 explained by factors F1, F2 and F3 and, therefore, have good potential as ultramafic
574 markers. For instance, the median Au/Ag ratio in ultramafic-hosted SMS is five times
575 higher than that in mafic-hosted SMS [$(Au/Ag)_{ultramafic} = 0.072$; $(Au/Ag)_{mafic} = 0.015$], in
576 analogy with Au/Ag in ultramafic rocks vs. MORBs [$(Au/Ag)_{ultramafics} \sim 0.20$;

577 (Au/Ag)_{MORB} ~ 0.020; Anderson, 1989]. The higher Au/Ag in ultramafic-hosted SMS
578 may also reflect the lower pH of fluids produced during peridotite alteration (Fig. 5; see
579 also Allen and Seyfried, 2003), which enhances Ag solubility in the cooling fluid (Fuchs
580 et al., 2019). Another potential marker is Sn, which is systematically enriched in
581 ultramafic-hosted SMS (Fouquet et al., 2010; Evrard et al., 2015; Melekestseva et al.,
582 2017). Because of the scarcity of data for Sn, this element could not be included in our
583 statistical analysis and its significance remains uncertain.

584 Another possibility that would be worth considering is that the fluids that formed
585 *all* SMS deposits on (ultra)slow-spreading ridges interacted with both mafic and
586 ultramafic rocks at depth. This scenario would explain the geochemical similarity
587 between several mafic-hosted and ultramafic-hosted SMS deposits. However, the relative
588 importance of ultramafic rocks vs. r/w ratios in determining the observed geochemical
589 associations would remain difficult to resolve, as these factors are likely to be highly
590 correlated with each other and with the ridge spreading rate.

591 **5 Conclusions**

592 The geochemical associations observed in SMS deposits on mid-ocean ridges can be
593 explained by a combination of the following independent factors, given in order of importance:
594 (1) temperature of deposition, (2) ridge spreading rate and (3) zone refining. The first and the
595 third factors are mostly related to processes that operate near the seafloor, such as conductive
596 cooling and/or mixing with seawater and metal remobilization, and determine the relative
597 proportions of the main minerals and, thus, of Cu and Zn (Co, Se, Sb, Pb). Thus, they are mostly
598 controlled by final depositional conditions and evolution of mound and vent structures rather

599 than by the original geochemistry of the hydrothermal fluid. The ridge spreading rate directly
600 influences the structure of the oceanic lithosphere, which in turn exerts a major control on the
601 length and depth of the hydrothermal circuit and on the r/w ratios in the reaction zone, and thus
602 on the behavior of precious metals and Ni (Mo, Se). Despite the obvious role played by substrate
603 rocks in releasing elements to hydrothermal fluids, their nature (specifically mafic vs. ultramafic)
604 does not clearly emerge as a statistically significant independent factor. Therefore, using simple
605 parameters such as metal grades or bivariate metal ratios to discriminate between mafic and
606 ultramafic hydrothermal systems may lead to erroneous evaluations. In any case, the relative
607 importance of highly correlated factors such as r/w and ultramafic/mafic volume ratios in the
608 reaction zone may be difficult to resolve using simple compositional data. The composition of
609 the substrate, however, may become relevant in subseafloor mineralization, where sulfides
610 precipitate by reaction of ascending hydrothermal fluids with substrate host-rocks. On a global
611 scale, widely recognized processes such as phase separation and magmatic fluid input do not
612 appear to play a major role in determining the overall geochemical diversity of SMS deposits on
613 mid-ocean ridges. These processes may nevertheless have a significant influence on a small
614 time-space scale, and may rather be reflected by peculiar mineral assemblages (e.g., barite-rich
615 or silica and Au-rich), which are only found at specific sites.

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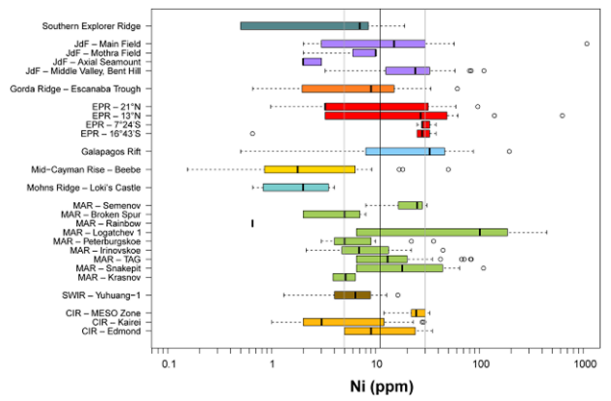
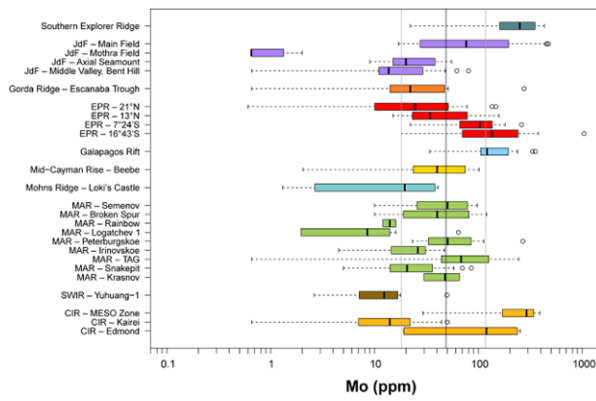
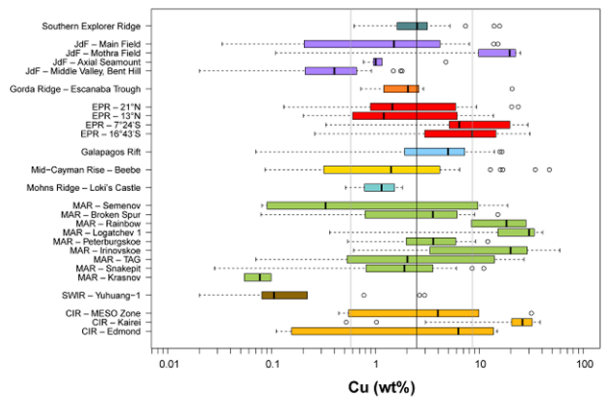
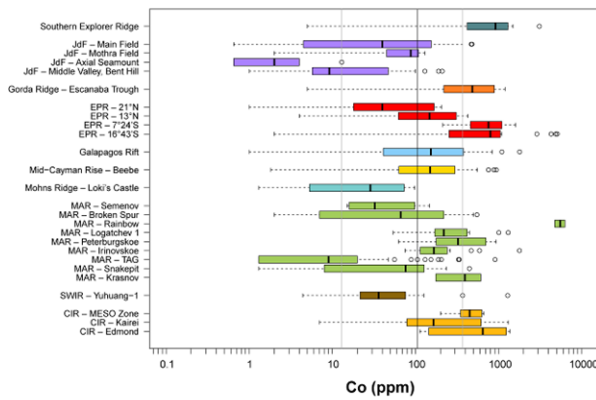
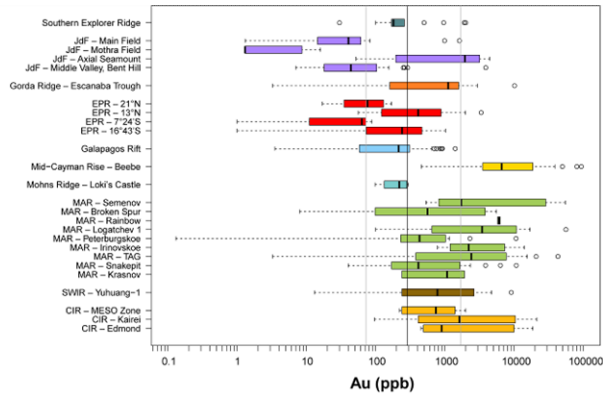
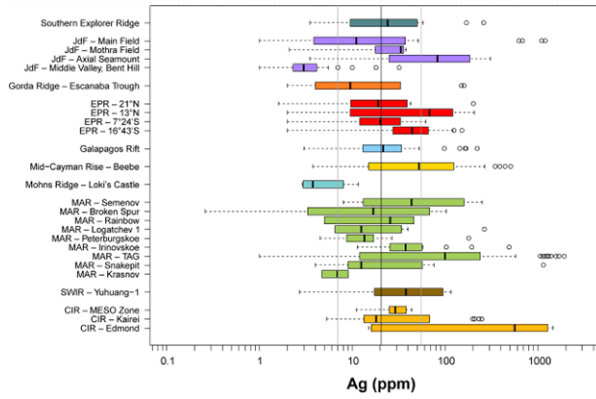
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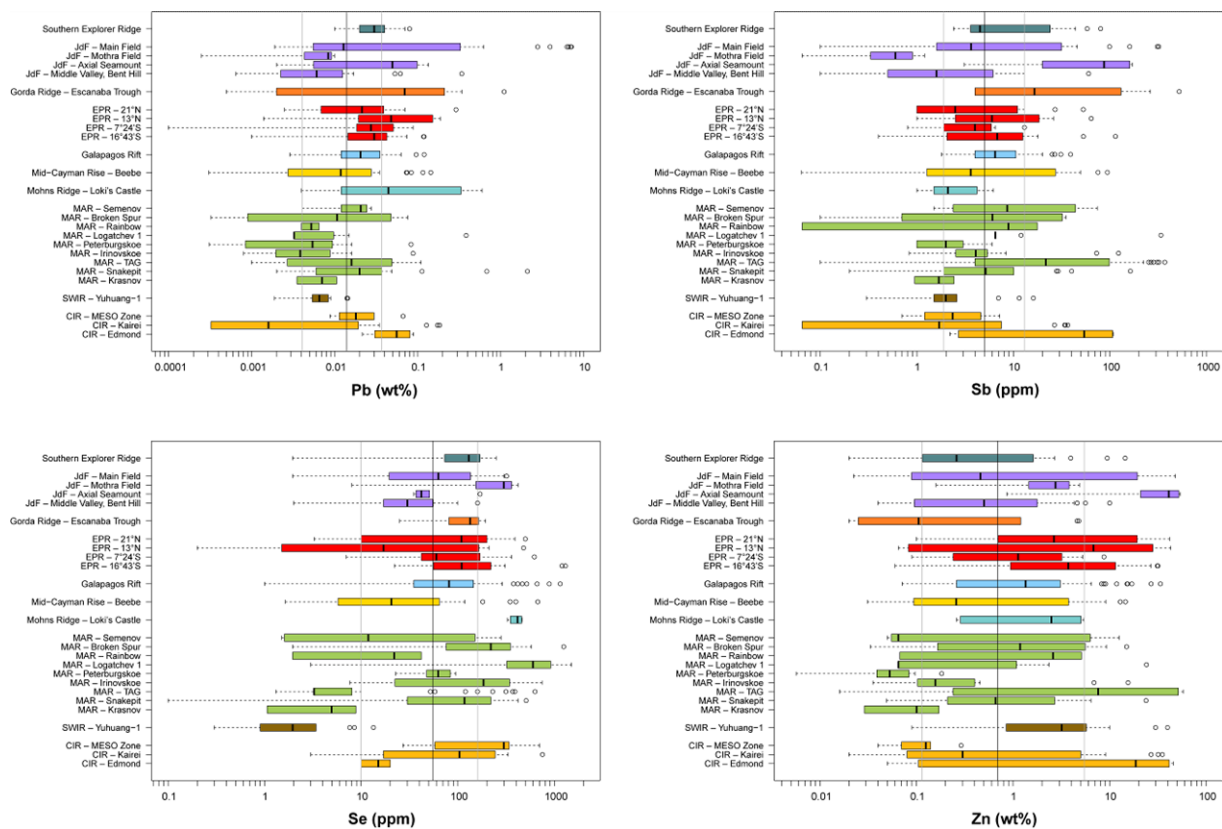
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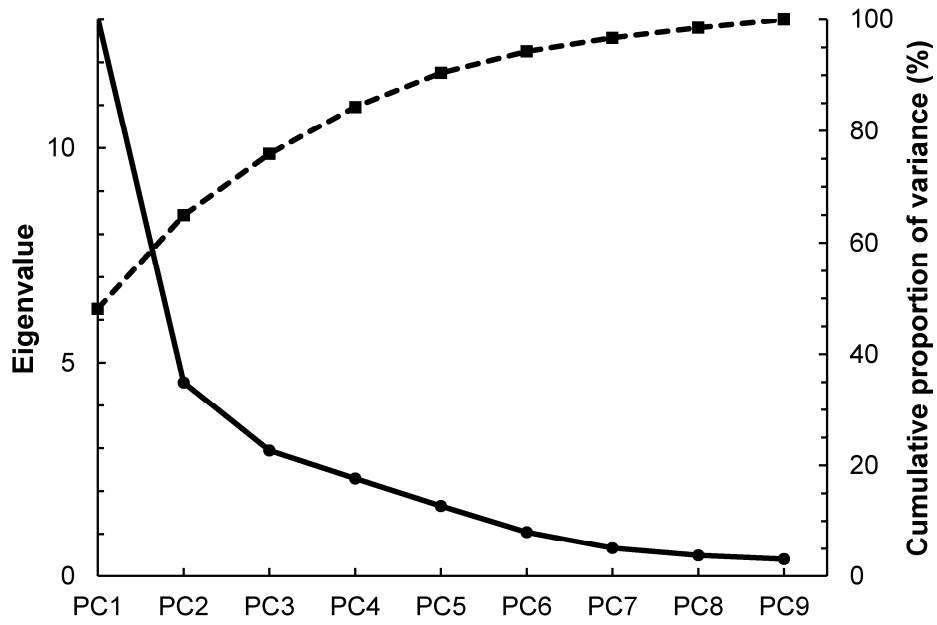


1004

1005 **Figure 1.** Box-and-whiskers plots indicating the distribution of elements in the various sites. The
 1006 boxes comprise the data between the first and third quartile (Q1 and Q3). The thick bar inside the
 1007 boxes is the median, while the whiskers are calculated as $Q1 - 1.5 \times (Q3 - Q1)$ and $Q3 + 1.5 \times (Q3 -$
 1008 $Q1)$. The circles represent the outliers. The three vertical lines (from left to right) represent the
 1009 Q1, median and Q3 of the global distribution of the specific element (see Table 2 for the values).

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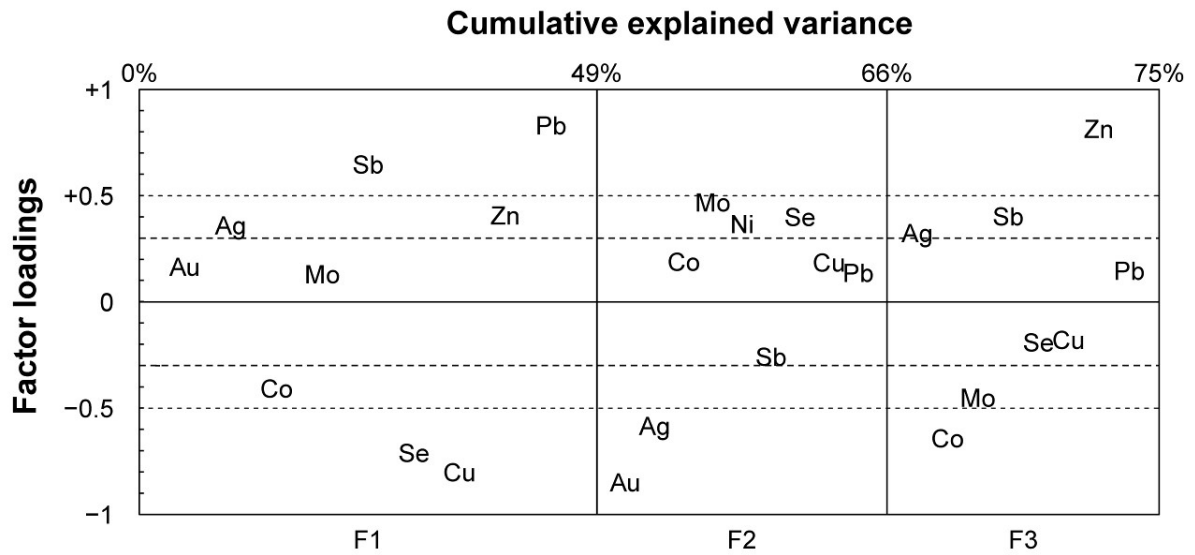
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1013 **Figure 2.** Screeplot resulting from the robust PCA. The first three components describe the
1014 greatest proportion of variance.

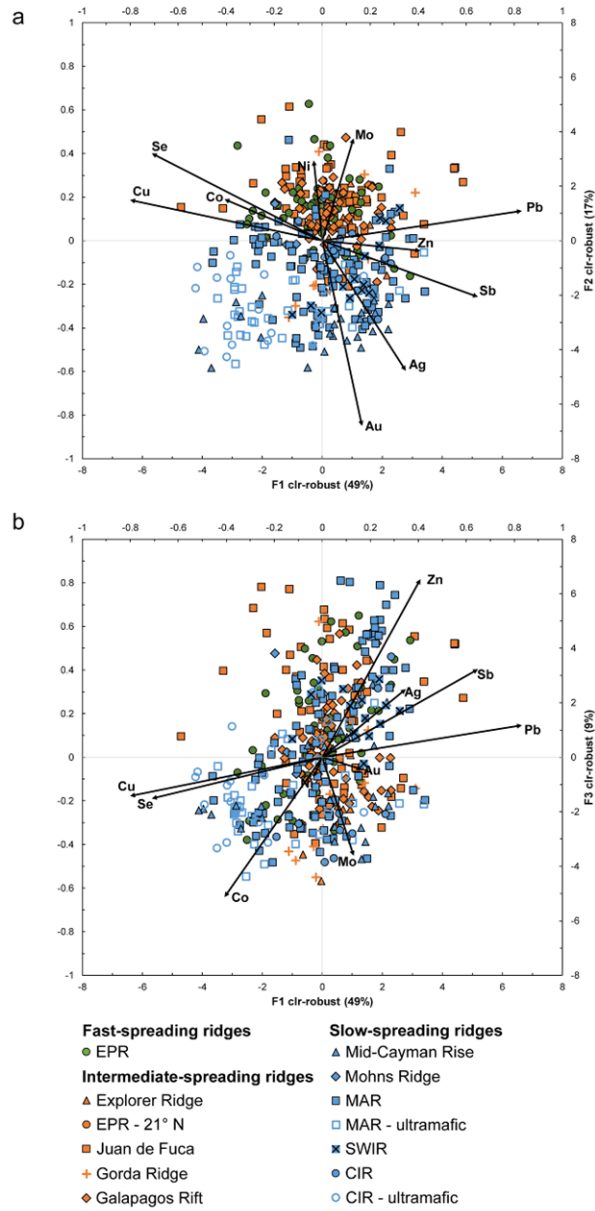
Robust FA (clr-transformed)



1015

1016 **Figure 3.** Factor loading plots for the clr-transformed MS data. The variables (elements) that
 1017 most significantly affect the factors have loadings > 0.5 (thin dashed lines). Variables with
 1018 loadings < 0.3 (thick dashed lines) are almost irrelevant and those with loadings < 0.1 are not
 1019 reported in the plot.

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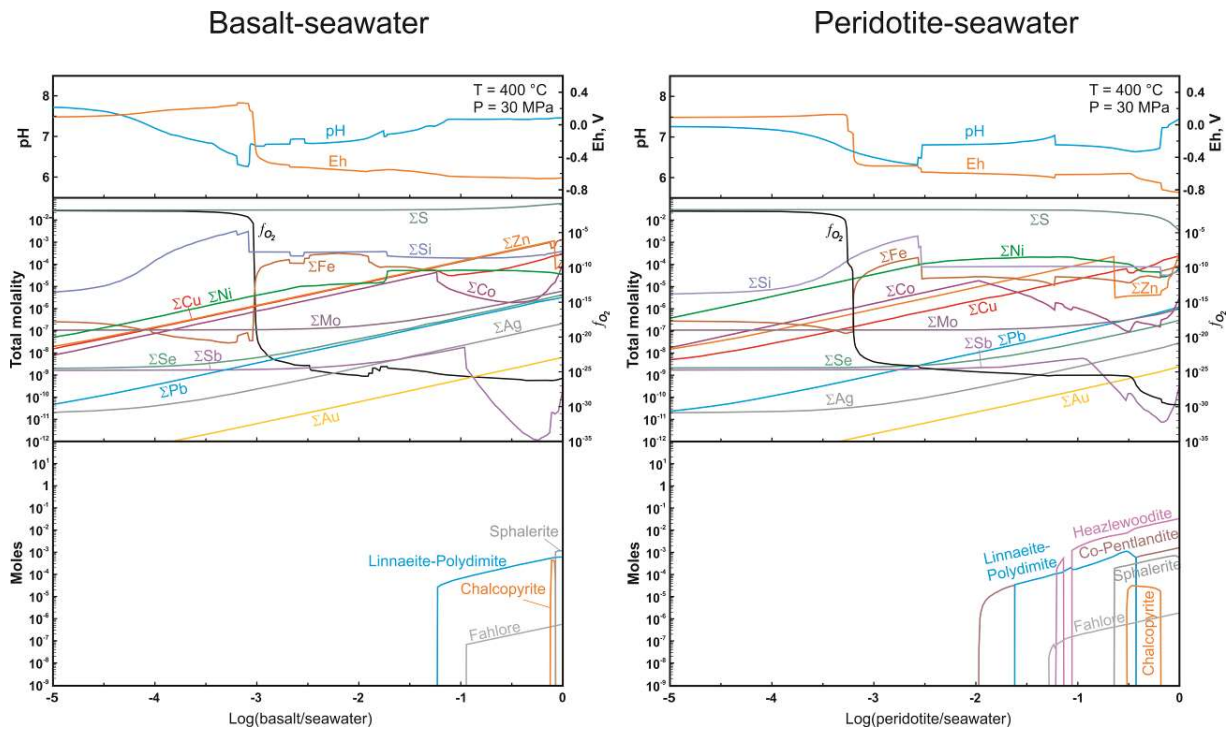
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1022 **Figure 4.** Biplots for the robust FA of clr-transformed MS data: a) F1 vs. F2; b) F1 vs. F3.

1023 Lower horizontal and right axes refer to the coordinates of the data points (scores), while upper

1024 horizontal and left axes refer to the coordinates of the variables (loadings). Data for individual
1025 sites are illustrated in Fig. S2.

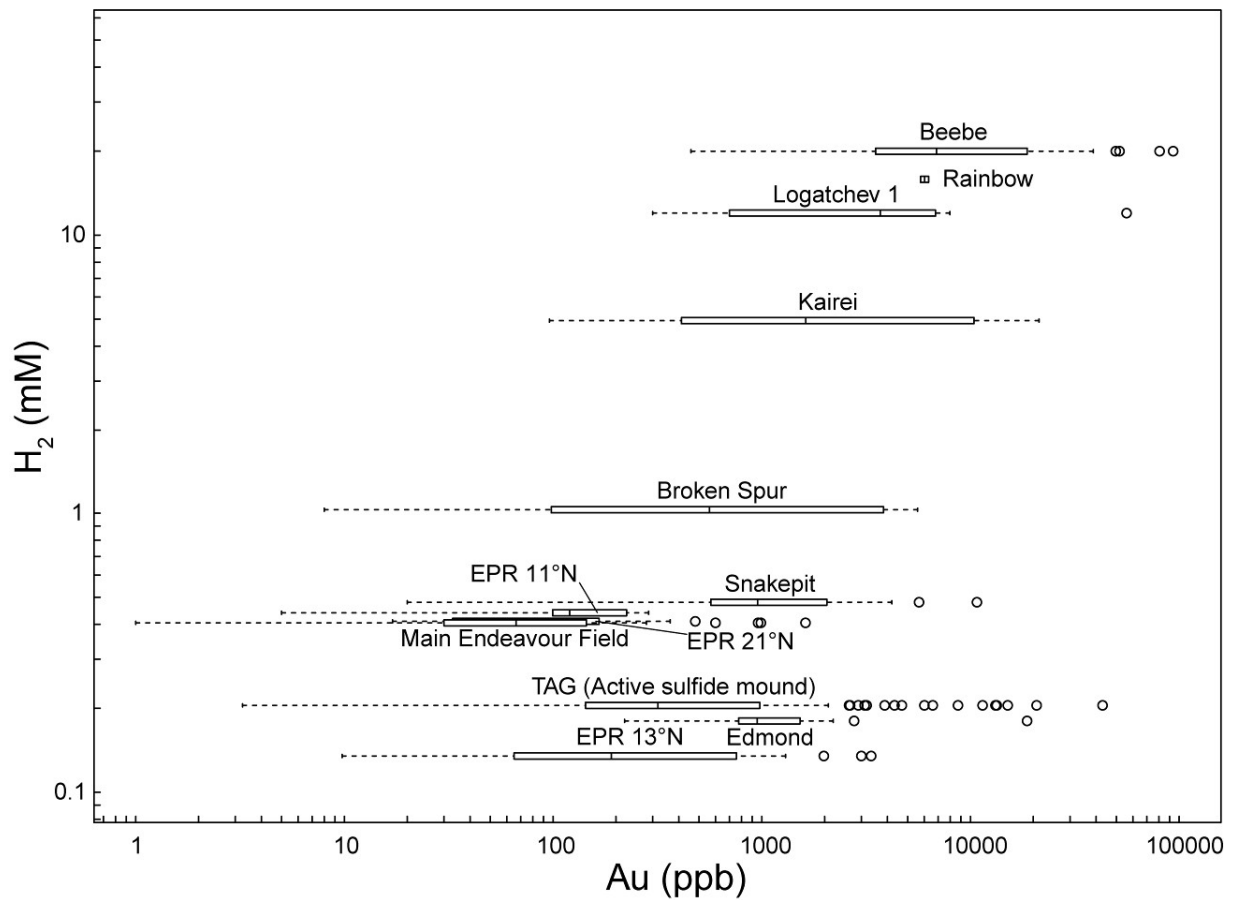
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1027

1028 **Figure 5.** Physicochemical model of the interaction of seawater with basalt (left) and peridotite
 1029 (right) at various r/w ratios (see Table SM2 for the compositions of the reactants). The
 1030 geochemical characteristics (pH, Eh, concentration of selected dissolved elements and fO_2) of the
 1031 hydrothermal fluid are reported in the upper and middle boxes. Only secondary sulfides
 1032 developed in the matrix of altered rocks are shown here (bottom boxes), since they greatly affect
 1033 the concentration of dissolved metals (see Fig. S5 and Fig. S6 for complete mineralogical data).
 1034 Note that the slight increase of Mo, and Se in solution at high r/w ratios is probably an artifact,
 1035 since the model does not allow for their isomorphic incorporation in alteration minerals.

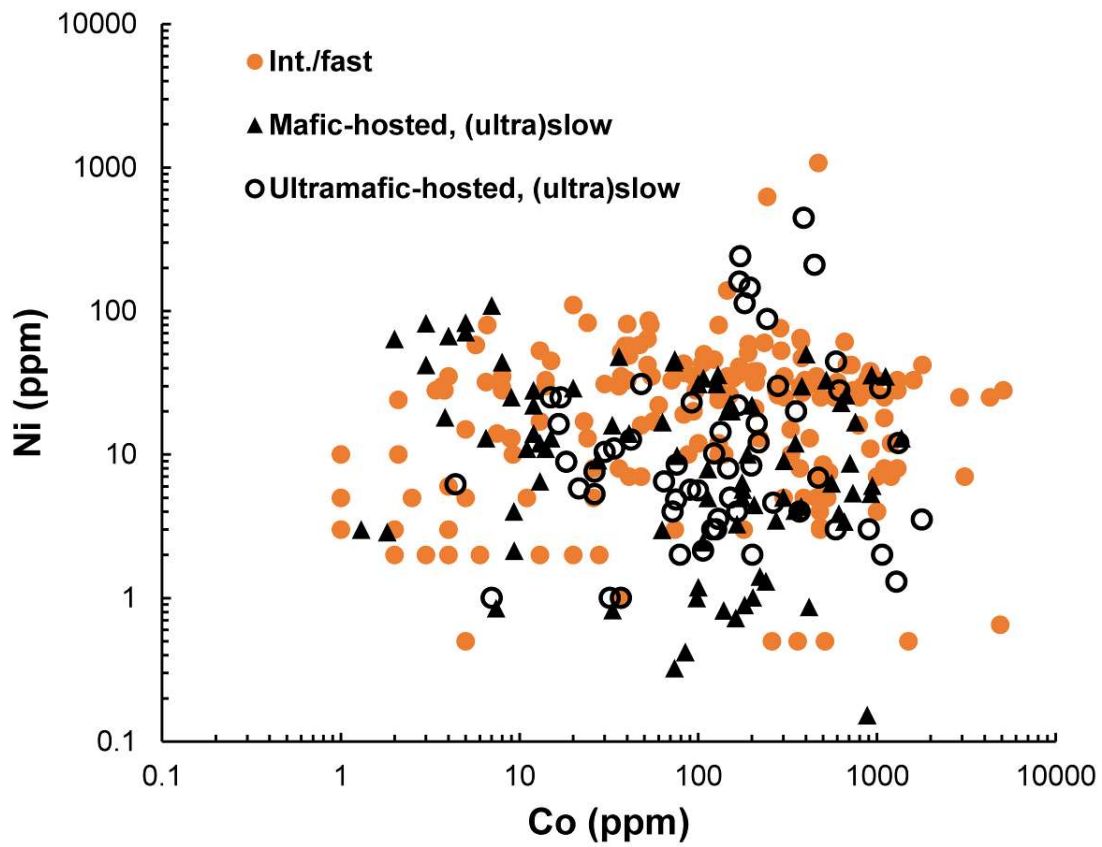
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1037

1038 **Figure 6.** Distribution of Au concentration in SMS vs. H_2 in vent fluids. The meaning of the
 1039 boxes and whiskers is the same as in Fig. 1. Data for vent fluids after Campbell et al. (1988),
 1040 Charlou et al. (1996, 2002), Kumagai et al. (2008), Fouquet et al. (2010), Gallant and Von
 1041 Damm (2006), Jean Baptiste et al. (1991), McDermott et al. (2018), Pester et al. (2011), Seewald
 1042 et al. (2003), Seyfried et al. (2011), and Tivey (1995). Data for SMS are compiled in Table S1.

1043



1044

1045 **Figure 7.** Cobalt vs. Ni concentrations in SMS from intermediate/fast-spreading ridges and slow-
 1046 spreading ridges, distinguished between MORB- and ultramafic-hosted. Low-detects are not
 1047 reported.

1048