1	Microbial strong organic ligand production is tightly coupled to iron								
2	hydrothermal plumes								
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34 Abstract. Hydrothermal vents have emerged as an-important sources of iron to seawater, yet only a subset 35 of this iron is soluble and persists long enough to impact the deep ocean iron inventory. The longevity and 36 solubility of iron in seawater is in part governed by strong organic ligands, like siderophores, that are 37 produced by microorganisms and are a part of the ocean's dissolved organic iron-binding ligand pool. These 38 ligands have been hypothesized to aid in the persistence of dissolved iron in hydrothermal environments. To 39 explore this hypothesis, we measured iron and iron-binding ligands including siderophores from 11 40 geochemically distinct sites along a 1,700 km section of the Mid-Atlantic Ridge. Siderophores were found 41 in hydrothermal plumes at all sites, with proximity to the vent playing an important role in dictating 42 siderophore type and diversity. The notable presence of amphiphilic siderophores may point to microbial 43 utilization of siderophores to access particulate hydrothermal iron, and the exchange of dissolved and 44 particulate iron. The tight coupling between strong ligands and dissolved iron within neutrally buoyant 45 plumes across distinct hydrothermal environments, and the presence of dissolved siderophores with 46 siderophore-producing microbial genera, suggests that biological production of ligands exerts a key 47 controlinfluence iron chemistry on in hydrothermal dissolved iron concentrationssystems.

### 48 **1. Introduction**

49 Over the last few decades, observations and modelling efforts have increased our understanding about the 50 critical role organic organic ligands play in the cycling, transport, and utilization of trace metals in deep water 51 (Tagliabue et al., 2017; Buck et al., 2018; Bundy et al., 2018; Moore et al., 2021; Hawkes et al., 2013b; Kleint 52 et al., 2016). Iron (Fe) binding organic ligands in deep-seawater have a wide range of sources, which are only 53 just beginning to be understood. Recent observations suggest that microbial production of siderophores, 54 humic-like substances and exopolysaccharides are some of the major contributors of marine organic ligands 55 (Hassler et al., 2017), and links microbial activity production and alteration of ligands to influences Fe cycling 56 in environments ranging from hydrothermal plumes (Cowen and Bruland, 1985; Cowen et al., 1990) to the 57 open ocean (Lauderdale et al., 2020; Whitby et al., 2024, 2020; Misumi et al., 2013). Strong Fe-binding 58 organic ligands (defined as  $L_1$  ligands) are a heterogeneous mixture of microbially produced compounds that are operationally classified based on their binding strength with Fe (defined as log  $K_{Fe',FeL}^{cond} > 12$ ). They are 59 60 thermodynamically favored to complex and stabilize external sources of Fe to prevent its scavenging and 61 removal\_(Fishwick et al., 2014; Aguilar-Islas et al., 2010). As an example, in high Fe estuarine systems, only 62 the dissolved Fe (dFe) bound to the strongest Fe binding ligands is protected from scavenging and remains 63 in solution (Bundy et al., 2015; Buck et al., 2007).

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Siderophores are the strongest known Fe-binding organic ligands. They are produced by bacteria and fungi
 to facilitate Fe uptake and solubilize otherwise inaccessible phases in the marine environment (Butler, 2005;

67 Manck et al., 2022). They have primarily been considered an important microbial strategy for Fe acquisition

68 in the low <u>dissolved</u> Fe (dFe<u>) < 0.5 nM</u>) surface ocean (Vraspir and Butler, 2009; Butler, 2005). However,

siderophore uptake and biosynthesis genes were observed in >70% of Fe-related bacterial transcripts in a
 hydrothermal environment in Guaymas Basin (Li et al., 2014), have been identified in oxygen-deficient zones

71 (Moore et al., 2021), and are a common Fe acquisition strategy within terrestrial and pathogenic ecosystems

- 72 (Sandy and Butler, 2009), all of which are environments where Fe concentrations are orders of magnitude
- 73 higher than surface seawater.
- 74

75 Previous studies have both looked atexamined unknown strong-Fe-binding ligands besides siderophores-in 76 hydrothermal plumes and throughout the deep ocean (Buck et al., 2018; Kleint et al., 2016; Hawkes et al., 77 2013b; Sander and Koschinsky, 2011), as well as siderophores observed below the euphotic zone (Bundy et 78 al., 2018; Park et al., 2023a; Moore et al., 2021; Boiteau et al., 2019). However, no previous studies have 79 ever directly measured siderophores in hydrothermal systems due to the high sample volume requirements, 80 difficulty in obtaining deep ocean trace metal samples, and the time-intensive nature of the analyses. A 81 'stabilizing agent' (i.e. ligands) has been proposed for the long-range transport of hydrothermal dFe into the 82 ocean interior. The role of strong Fe-binding ligands in hydrothermal dFe transport represents an important 83 knowledge gap in how hydrothermal vents may impact the ocean dFe inventory (Resing et al., 2015) and 84 how siderophores may influence Fe transformations in hydrothermal systems. Here, for the first time, we 85 identified siderophores and siderophore-producing microbes in 11 geochemically distinct hydrothermal environments along the slow-spreading (20-50 mm/yr) Mid-Atlantic Ridge (MAR). Four black smokers 86 87 (high temperature, high Fe), four off-axis sites, one diffuse vent (low temperature, low Fe), one alkaline vent 88 (pH 9-11, very low Fe), and one non-vent fracture zone were investigated using both competitive ligand 89 exchange-adsorptive cathodic stripping voltammetry and state-of-the-art liquid chromatography coupled to 90 electrospray ionization mass spectroscopy (Boiteau et al., 2016) in a targeted approach to-search for known 91 siderophores identify and possible discrete-compounds components of present in the  $L_1$  ligand pool in 92 hydrothermal environmentss and to search for known siderophores. Microbial community analysis was also 93 compared at three sites to understand whether microbial ligand production impacts Fe transformation the 94 supply of in hydrothermal definition of the ocean systems. Overall, our results show microbially-produced 95 siderophores were present in all sites, and that strong  $L_1$  ligands were tightly coupled to hydrothermal dFe 96 concentrations in the neutrally-buoyant plume sampless in this system. The presence of organic ligands 97 produced by bacteria in hydrothermal systems suggest that they play an important role in deep ocean Fe 98 cycling.

# 99 2. Results and Discussion

### 100 **2.1 The role of iron-binding ligands in hydrothermal plumes**

Strong organie Fe-binding ligands (defined here as L<sub>1</sub> ligands) have been previously been found to be
important in neutrally-buoyant hydrothermal plumes across a variety of systems (Tagliabue et al., 2017;
Resing et al., 2015; Buck et al., 2018; Hawkes et al., 2013b; Wang et al., 2022; Bennett et al., 2008). But
However, the relationship between organic ligands and dFe have never been investigated together

systematically across a wide variety of vents in the same study. In this work, the average binding strength

- and concentration of organic Fe-binding ligands were quantified in 11 vent systems that spanned a wide range
- 107 in dFe concentrations (0.41-90 nM) and underlying vent geology. Over 99% of dFe in the neutrally buoyant
- 108 plume sampless were complexed by L1 ligands and the ligands were almost always completely saturated with
- 109 dFe, meaning Fe-free 'excess' L<sub>1</sub> ligands capable of binding additional Fe were present in low concentrations
- 110 (< 1 nM; Fig. S1). As a result, dFe concentrations were tightly coupled to  $L_1$  ligands in a nearly 1:1 ratio
- (Fig. 1d), similar to previous studies in other neutrally buoyant plumes (Fig. 1e) (Lough et al., 2022; Buck
- 112 et al., 2018, 2015).
- 113 The strong coupling between dFe and ligands was only observed at sites where  $L_1$  ligands were detected. 114 Some samplesing locations, such as in the that were closer to the buoyant plume and vent source-or closer to the vent orifice, contained high concentrations of weaker ligands (log  $K_{Fe',FeL}^{cond}$  < 12, **Table S2<u>-S3</u>**) with whose 115 116 concentrations had no correlation to-with dFe. This is consistent with these environments likely being 117 dominated by complex Fe phases, which could include various inorganic forms (e.g. nanopyrite, Fe-118 oxyhydroxide) of Feas well as mixed organic phases of Fe as hydrothermal fluids initially mix with 119 oxygenated seawater. High concentrations of weaker ligands have also been observed in samples near the 120 vent orifice in previous studies (Hawkes et al., 2013). In this study, we are not able to discern the exact 121 chemical composition of the ligands we detect via voltammetric methods, and thus the weaker and some 122 portion of the stronger ligands we observe likely represent a mix of different inorganic and organic ligands. 123 Similar to what was described in Hawkes et al. (2013), the ligands we measure could represent multiple 124 layers of coordination bonds, forming complex Fe phases, similar to the "onion" concept t(Mackey 125 and Zirino, 1994). For example, colloidal Fe phases are common in hydrothermal vents and can form 126 aggregates that bind Fe, but not in traditional organic coordination bonds (Fitzsimmons et al., 2017; 127 Honeyman and Santschi, 1989). There are also likely processes occurring near the vent source in such a 128 complex environment that cause some Fe phases to be in various stages of disequilibria that we also measure 129 as ligands via our voltammetric methods.
- 130
- The sources of Moreover, weaker Fe-binding ligands with a(-log  $K_{Fe',FeL}^{cond} < 12$ ) that have been observed in 131 132 other hydrothermal systems <del>globally, and yet their role in Fe cycling in these environments</del> is not well 133 understood, and their impact on Fe cycling over the lifetime of neutrally-buoyant plume is unclear. Recent 134 studies have shown microbes may use siderophores or siderophore-like (strong binding ligands) ligands to 135 access the ironFe associated with weaker-type ligands — such as humic substances and thiols — and to 136 enhance the eveling and bioavailability of Fe in aquatic systems (Kuhn et al., 2014; Muller, 2018). However, 137 to date, only athe few studies that have explored ligands concentrations and binding strengths within 138 hydrothermal systems (Buck et al., 2015, 2018; Kleint et al., 2016; Hawkes et al., 2013c; Sander and 139 Koschinsky, 2011) and they have mixed results hypotheses as to the role and sources of weaker-type ligands 140 within plumes. Additional studies are needed to investigate the sources and mechanisms of weaker-type 141 ligands in hydrothermal plumes and understand their impact on the Fe cycle in hydrothermal systems.

143 144 In the neutrally buoyant plume samples, Overall, Ourour voltammetry results indicate that stronger  $L_1$  ligands were present and were cap-correlated with the dFe concentrations in neutrally buoyant plumes(Fig. 1) and 145 146 weaker ligands were no longer dominant. In other systems with a high dFe and ligand endmember such as 147 estuaries, a decrease in weaker ligands along with dFe concentrations has also been observed (Buck et al., 2007; Bundy et al., 2014). This has been interpreted as a scavenging of weaker Fe-ligand complexes, while 148 149 the dFe that remains in solution is that which is bound to stronger ligands (Bundy et al., 2014). A similar 150 control on dFe concentrations by  $L_1$  ligands has also been previously observed in estuaries (Buck et al., 2007) 151 and aerosol solubility experiments (Fishwick et al., 2014). There are a few possible explanations for the correlation of dFe and L<sub>1</sub> ligands in the neutrally-buoyant plume. One possible explanation is that both the 152 153 dFe and  $L_1$  ligands originate from the vent fluids themselves, yielding a tightly coupled hydrothermal 154 endmember. However, the concentration of  $L_1$  ligands did not correlate with excess mantle Helium-3 (<sup>3</sup>He<sub>xs</sub>, 155 Fig S2, Table S2-S3) (Lough et al., 2022), a nearly conservative tracer of the mixing of hydrothermal fluids 156 with seawater (Buck et al., 2018)., and Moreover, our samples closer to the vent source were dominated by 157 weaker organic ligands showing no correlation to dFe. This suggests the  $L_1$  ligands were not directly sourced 158 from the vent fluids along with dFe. Biological sources represent aAnother likely explanation is the source for 159 the coupling of  $L_1$  ligands and dFe, if the ligands observed in the neutrally-buoyant plume are either-from 160 bacteria that produced them in surrounding deep ocean seawater that was then entrained, local production 161 from vent-biota and/or microbial mats, diffusion from microbial production in sediments, or in-situ 162 production by bacteria within the neutrally buoyant plume (Mellett et al., submitted)(Dick et al., 2013; Li et 163 al., 2014; Sheik et al., 2015; Mellett et al., submittedn.d.). Previous work has shown the genetic potential for 164 plume associated microbial populations to produce siderophores (Li et al., 2014)

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### 166 **2.2 The presence of siderophores in hydrothermal systems**

167 Siderophores were measured in a subset of the samples to further explore the source of the  $L_1$  ligands coupled 168 to dFe in the neutrally-buoyant plume. Marine organic ligand composition changes with environmental 169 gradients (Boiteau et al., 2016; Gledhill and Buck, 2012), making the structure and functional groups of 170 siderophores identified in hydrothermal samples of particular interest. Somewhat surprisingly, siderophores 171 were found in all samples and we observed a large diversity of siderophores with high confidence using mass-172 to-charge ratio (m/z), MS/MS spectra, and specific chromatographic characteristics (Fig. 2a). On-axis 173 spreading centers contained the highest dFe concentrations (> 20 nM) and wider variety of siderophores than 174 samples from fracture zones, diffuse, and off-axis sites (dFe  $\leq 1$  nM). The greatest number of distinct 175 siderophores were identified at Lucky Strike, Broken Spur, Rainbow, and TAG (Fig. 2). On average, 13 176 compounds were identified with high confidence per on-axis spreading center sample, compared with 5 per 177 diffuse/fracture zone sample, and 2.5 per off-axis sample (Fig. 2b, Fig. S4). Mixed-type siderophores —

containing different moieties that bind to Fe(III) — were common at all sites. Hydroxamates were identified
 at and around spreading centers, yet none of these were detected with high confidence in samples from

180 diffuse/fracture zones (Fig. S4). Summed siderophore abundance in neutrally-buoyant plumes above

181 spreading centers was similarly more than twice that of samples from fracture zones or off-axis (Fig. 2c).

182 Thus, vent type and proximity played a role in the diversity and abundance of siderophore types observed,

- 183 likely related to the diversity of the microbial community and/or unique Fe acquisition strategies across sites.
- 184

185 Siderophores are <u>putatively operationally</u> part of the <u>operational</u>  $L_1$  ligand pool based on their binding 186 strength (Gledhill and Buck, 2012), and patterns in their distributions reflected were similar to those of the 187 strong ligands. The peak areas of each putative siderophore we identified were used as a proxy for 188 concentrations (section 3.3), and these concentrations significantly correlated with dFe, as observed with dFe 189 and  $L_1$  ligands (Fig. 2b). Siderophores were present in concentrations similar to the surface ocean (Boiteau 190 et al., 2016; Moore et al., 2021; Park et al., 2022; Bundy et al., 2018), and comprised-were equivalent to 191 <u>concentrations representing</u> 0.01-0.4% of the total  $L_1$  ligands (**Table 1**). This is likely a substantial 192 underestimate of siderophore contributions to the  $L_1$  ligand pool due to analytical constraints in identifying 193 unknown siderophores. Recent work on siderophore biosynthesis pathways and advances in genome mining 194 suggest that known siderophores represent a small fraction of what is expected to be produced in nature 195 (Hider and Kong, 2010; Reitz et al., 2022), and our analyses in this study were limited to only known 196 siderophores. We also restricted our reporting to compounds identified with very high confidence (Fig 2a, 197 S3). In addition, most siderophores are not commercially available to use as standards, and individual 198 siderophores have different ionization or extraction efficiencies. We restricted our reporting to compounds 199 only identified with very high confidence (Fig 2a, S3). The extraction efficiency for the solid phase extraction 200 technique is approximately 5-10% for bulk Fe-binding organics (Bundy et al., 2018) and 40% for a 201 siderophore standard (Waska et al., 2015). Employing both corrections yields siderophore contributions to 202 the total  $L_1$  pool of 0.1-4% and 0.025-1%, respectively. We are inevitably missing many naturally occurring 203 unknown compounds, and thus we consider this a lower bound. Regardless of the small percentage 204 contribution to total  $L_1$  ligands, it is evident that microbially produced siderophores were ubiquitous across 205 all vent sites and had similar distributional patterns as  $L_1$  ligands. There are also likely other compounds such as some strong binding humics that are also contributing to the L<sub>1</sub> ligand pool (Laglera and van den Berg, 206 207 2009). Future work with much larger water volumes will be able to reduce uncertainty and identify a greater 208 number of compounds. Still, tThe identification of siderophores here — and their relationship with dFe — 209 provides compelling evidence that microbial production of ligands is responsible for at least some portion of 210 the tight coupling between  $L_1$  and dFe in hydrothermal systems along the MAR.

211

212 The presence and diversity of siderophores identified in this system was surprising given the relatively high

213 Fe concentrations of hydrothermal environments, but some interesting compelling patterns were observed.

214 For example, previous work has shown that low Fe surface waters have higher concentrations of amphiphilic

215 siderophores compared to high Fe coastal waters (Boiteau et al., 2016), and amphiphilic siderophores are less 216 common in terrestrial environments (Hider and Kong, 2010). Amphiphilic siderophores have long 217 hydrocarbon tails that can be embedded into the lipid bilayer of the bacterial cell membrane providing a mechanism to shuttle Fe into the cell and prevent diffusive loss (Martinez et al., 2003). Amphiphilic 218 219 siderophores comprised 57% of the siderophores in our samples (Fig. S5), supporting the ubiquity of 220 amphiphilic siderophores in marine environments (Butler and Theisen, 2010). Amphiphilic siderophores 221 were found in concentrations between 0.3-4.7 pM, with the highest concentrations found at Rainbow (Fig. 222 **2d**, **Table S65**). These concentrations were similar to those observed in the upper ocean (Boiteau et al., 2016; 223 Bundy et al., 2018; Boiteau et al., 2019). Marine bacteria produce suites of amphiphilic siderophores as a 224 way to adapt to the change in hydrophilicity in the surrounding environment (Sandy and Butler, 2009; 225 Homann et al., 2009). Amphiphilic siderophores in plumes could be a way for bacteria to access Fe as they 226 are physically transported and cope with strong chemical gradients, similar to the production of multiple 227 siderophores in terrestrial and pathogenetic systems as a means to access inorganic particulate Fe for cellular 228 uptake and storage (Hider and Kong, 2010).

229

### 230 **2.3 Microbial sources of siderophores in hydrothermal plumes**

231 The high diversity of siderophores across a huge range of hydrothermal vent systems revealed several 232 surprising aspects of Fe cycling. The biosynthesis of a siderophore is energy-intensive and is regulated by Fe 233 concentration in the surrounding environment (Rizzi et al., 2019). Siderophore presence suggests that bacteria 234 are producing these compounds despite the overall higher Fe concentrations in the deep ocean and within 235 hydrothermal plumes. Consistent with siderophore utilization in terrestrial ecosystems (Hider and Kong, 236 2010; Sandy and Butler, 2009), one hypothesis is that siderophore production is beneficial to bacteria in the 237 plumes for transforming Fe from otherwise inaccessible forms, such as particulate nanopyrites or Fe 238 oxyhydroxides that are present close to the vent source. To explore the potential for microbial production of 239 siderophores, we examined microbial community composition around Rainbow (St. 11, 17) and Lucky Strike 240 (St. 7; Table 1, Table S1) using 16S rRNA gene-based amplicon sequencing to detect bacteria with the 241 metabolic potential to synthesize siderophores (Fig. 3, S11), where the presence of taxa encoding siderophore 242 biosynthetic gene clusters indicates whether the microbial community is genetically capable of producing the 243 compounds we observed. Bacterial genera containing known siderophore-producers were found at all three 244 MAR sites examined, and putative siderophore-producers represented 3-20% of the relative abundance of 245 the community (Fig. 3). Putative siderophore-producers were more abundant in the 3  $\mu$ m (particle-attached) 246 size fraction than in the 0.2  $\mu$ m (free-living) fraction, suggesting siderophore production is more common in 247 particle-associated bacteria in hydrothermal environments.

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We found microbial genera in our samples that can produce a subset of the siderophores identified here, including ferrioxamines, vibrioferrin, and acinetoferrin (Butler, 2005; Vraspir and Butler, 2009; Moore et al., 251 2021; Bundy et al., 2018; Boiteau et al., 2016). Genera with the genetic potential to produce ferrioxamines 252 were present at all three sites, while those known to produce vibrioferrin were present at Lucky Strike and 253 Rainbow, and those producing acinetoferrin were also present at Rainbow (Table S1, S76). Mycobactins 254 were detected with high confidence in every sample of this study, and genes encoding mycobactin have been 255 detected in a cultured organism from a hydrothermal system (Gu et al., 2019), but no mycobactin producers 256 were identified in this study. We detected woodybactin D with high confidence in 5 out of 11 sites. Although 257 these biosynthetic genes were not identified in any of the genera observed, woodybactin D is a carboxylate 258 siderophore isolated from Shewanella (Carmichael et al., 2019), and groups of deep-sea Shewanella (Kato and Nogi, 2001) were found in the dataset (Fig. S11). The biosynthesis genes for many of the siderophores 259 260 identified are unknown. Thus, finding genera capable of producing only a subset of the siderophores 261 characterized is not surprising. The observation that a significant portion of the *in-situ* microbial community 262 is capable of synthesizing siderophores (Fig 3) suggests that siderophore production is more widespread in 263 the deep ocean than previously believed.

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# 265 2.4 The impact of strong ligands and siderophores on dissolved iron in neutrally-buoyant plumes

266 Evidence that siderophores are ubiquitous in the marine environment — including higher Fe environments 267 — has been increasing (Park et al., 2023b). The high dFe associated with hydrothermal plumes may still not 268 be high enough to suppress siderophore production due to the elevated Fe requirements of heterotrophic 269 bacteria (Tortell et al., 1996). It is also likely that in hydrothermal plumes not all of the Fe is bio-accessible 270 in hydrothermal plumes. Soil microbes secrete siderophores to solubilize particulate Fe (Crowley et al., 1991) 271 and similar processes could be occurring in hydrothermal plumes, where Fe mineral phases associated with 272 organic compounds are common (Hoffman et al., 2020; Toner et al., 2009; Fitzsimmons et al., 2017; Hoffman 273 et al., 2018; German and Seyfried, 2014; Holden et al., 2012). Although our measurements suggest that dFe 274 in the neutrally-buoyant plume is likely dominated by organic complexation, the  $L_1$  measurements alone 275 cannot distinguish between purely organic phases or a mixture of inorganic and organic ligands in complex 276 aggregations or small colloids, as discussed above (section 2.1). Given the evidence from particulate Fe 277 studies in neutrally-buoyant plumes (Hoffman et al., 2020; Toner et al., 2009; Yücel et al., 2011; Fitzsimmons 278 et al., 2014, 2017; Hoffman et al., 2018)(Hoffman et al. 2020), it is highly likely that some portion of what 279 is detected in the  $L_1$  pool is a mixture of organic and inorganic Fe in small colloids which are operationally 280 in the dFe pool (Fitzsimmons et al., 2017). It is also telling that most siderophore-producing genera were 281 found to be particle-associated (Fig. 3), providing additional evidence that siderophores might be produced 282 to solubilize particulate Fe or access other colloidal phases. Further work that assesses why bacteria are 283 producing siderophores in neutrally buoyant plumes will be important for understanding microbial 284 metabolism in these systems, and the impact of siderophore production on Fe dispersal.

285

Organic Fe-binding ligands have been implicated in playing a critical role in the preservation and transport of hydrothermal dFe into the ocean interior (Hoffman et al., 2018; Resing et al., 2015; Fitzsimmons et al.,

- 288 2017; Toner et al., 2009; Bennett et al., 2011, 2008; Buck et al., 2018; Sander and Koschinsky, 2011). In this
- work, L<sub>1</sub> ligands were tightly coupled to dFe in neutrally buoyant plumes along the MAR and the presence
- 290 of siderophores in these samples provided evidence for the first time, that at least some of these ligands are
- 291 microbially produced. How these complexes may facilitate the exchange of Fe between dissolved and
- 292 particulate phases (Fitzsimmons et al., 2017), and whether siderophores are present across additional
- 293 hydrothermal vent systems will aid in constraining the biogeochemical importance of microbial feedbacks in
- impacting the hydrothermal dFe supply to the deep ocean.
- 295

### 296 **3. Appendix: Materials and Methods**

### 297 **3.1 Sampling and cruise transect**

Samples were collected as part of the 2017-2018 U.K. GEOTRACES GA13 section cruise along the Mid-298 299 Atlantic Ridge. Water samples from 11 venting and near venting locations were collected using a Seabird 911 conductivity, temperature, and depth (CTD) titanium rosette using conducting Kevlar wire with an 300 301 oxidation-reduction potential (ORP) sensor to detect plumes. Teflon coated OTE (Ocean Test Equipment) 302 bottles were pressurized to approximately 7 psi with 0.2 µm filtered air using an oil free compressor. A 303 Sartobran 300 (Sartorius) filter capsule  $(0.2 \,\mu\text{m})$  was used to collect filtered seawater samples into clean 250 304 mL LDPE sample bottles. Bottles and caps were rinsed 3 times with the filtered sample before being filled. 305 Samples were stored frozen at -20°C for Fe-organic ligand characterization by voltammetry and mass 306 spectrometry.

# 307 3.2 Fe-binding ligand concentration and binding strengths Competitive Ligand Exchange-Adsorptive 308 Cathodic Stripping Voltammetry

Fe-binding ligand concentrations and binding strengths (defined as conditional binding constants,  $\log K_{Fe',FeL}^{cond}$ 309 310 > 12) were determined by competitive ligand exchange-adsorptive cathodic stripping voltammetry (CLE-311 ACSV) with a BASi controlled growth mercury electrode (CGME) with an Ag/AgCl<sup>-</sup> reference electrode 312 and platinum auxiliary electrode (Bioanalytical Systems Incorporated). Using previously established 313 methods (Buck et al., 2015, 2018; Bundy et al., 2018; Abualhaija and van den Berg, 2014; Hawkes et al., 314 2013c), 40 frozen filtrate (<0.2 µm) samples with dFe concentrations between 0.41-11.67 nM (Table S1-315 **S2**) were thawed in a 4°C fridge prior to analysis. A 15-point titration curve was analyzed for each sample. 316 Briefly, within each titration, every point sequentially received 10 mL of sample, 7.5 mM of borate-317 ammonium buffer, 10 µM salicylaldoxime (SA) added ligand, and a dFe addition. Samples were then 318 equilibrated overnight before being measured on the BASi. - Data was collected using the Epsilon Eclipse 319 Electrochemical Analyzer (v.213) with a deposition time of 120 seconds and analyzed using ElectroChemical 320 Data Software (v2001-2014) and ProMCC (v2008-2018) to determine peak areas and Fe-binding ligand 321 parameters, respectively. All results were confirmed to fall within the analytical window of the method by 322 comparing the side reaction coefficient of the added ligand  $\alpha_{SA}$  to the side reaction coefficient of the natural

323 ligands detected ( $\alpha_L$ ). If the  $\alpha_L$  was within an order of magnitude of  $\alpha_{SA}$  then the results were deemed to fall 324 within the analytical window.

### 325 3.3 Reverse Titration-CLE-ACSV

Reverse titration-CLE-ACSV (RT-CLE-ACSV) (Hawkes et al., 2013a) was completed on 10 samples from 326 327 Broken Spur, and TAG hydrothermal vent fields with dFe concentrations between 19.01-90.25 nM (Table 328  $S_{31-S_2}$ ). Briefly, a 10-point titration curve was analyzed for each sample with each titration point consisting 329 of 10 mL of sample buffered with 7.5 mM boric acid and the competitive ligand 1-nitroso-2-napthol (NN) 330 additions. All samples were analyzed on a BASi Controlled Growth Mercury Electrode (CGME) with the 331 *Epsilon Eclipse Electrochemical Analyzer* (v.213) and deposition time of 120 seconds. For each sample, 332 competitive ligand NN additions were 0.5, 1, 2, 3, 4, 6, 9, 15, 20, and 40 µM. Samples were equilibrated 333 overnight and purged with  $N_2$  (99.99%) for 5 minutes before analysis. At the end of each titration, three Fe 334 additions (3-15 nM) were added to the final titration point to get the total concentration of Fe in equilibrium 335 with ligands. Data was analyzed using *ElectroChemical Data Software* (v2001-2014) to acquire peak areas 336 and a package in R using the model parameters of  $\beta_{\text{FeNN3}} = 5.12 \text{ x } 10^{16}$ ,  $\chi_{\text{min}} = 0.8$ ,  $\chi_{\text{max}} = 0.9$ , and c1high = 0.8337 0.75 to determine the Fe-binding ligand parameters (Hawkes et al., 2013a). These parameters were chosen 338 based on the recommendations for undersaturated samples and titrations curves where  $ip_{max}$  was not reached 339 (Hawkes et al., 2013a). All other parameters within the model we kept at the default values.

# 340 **3.4 Siderophore quantification and characterization**

341 In addition to measuring Fe-binding ligands by voltammetry, we also identified and quantified siderophores. 342 Between 0.65-1.5 L of 0.2 µm filtered seawater pooled from ligand samples at each site (described above) 343 was pumped slowly (15-20 mL min<sup>-1</sup>) onto a polystyrene-divinylbenzene (Bond Elut ENV) solid phase 344 extraction (SPE) column (Bundy et al., 2018; Boiteau et al., 2016). SPE columns were rinsed with MilliQ 345 and stored at -20°C until analysis. For the analytical measurements, samples were thawed in the dark, eluted 346 in 12 mL of distilled methanol, and dried down to between 0.2-0.5 mL of sample eluent (Table S1). Aliquots 347 were analyzed by reverse-phase liquid chromatography (LC) on a trace metal clean bio-inert LC (Thermo 348 Dionex 3000 NCS). The LC was interfaced with an electrospray ionization-mass spectrometer (ESI-MS; 349 Thermo Q-Exactive HF) to identify and quantify the compounds based on accurate mass (MS<sup>1</sup>) and the 350 fragmentation (MS<sup>2</sup>) data (Bundy et al., 2018; Boiteau et al., 2016). MSconvert (Proteowizard) was used to 351 convert MS data to an open source mzxML format, and two stages of data processing were conducted using 352 modified versions of previously reported R scripts (Bundy et al., 2018; Boiteau et al., 2016). In the first stage, mzxML files were read into R using new package "RaMS" (Kumler and Ingalls, 2022), and extracted ion 353 354 chromatograms (EICs) were generated for each targeted m/z of interest from an in-house database of siderophores. The m/z targets were the ionized apo, <sup>54</sup>Fe-bound, and <sup>56</sup>Fe-bound version of each siderophore, 355 356 with a tolerance of 7.5 ppm. Putative siderophore candidates were filtered through a series of hard thresholds, 357 such that  $MS^1$  spectra were quality controlled to contain a minimum of 25 datapoints and the maximum

intensity of each EIC was greater than 1e4 counts. Spectra meeting these criteria and containing either  ${}^{54}$ Febound and  ${}^{56}$ Fe-bound m/z peaks within 30 seconds of each other or an apo peak were displayed for the user

to further inspect peak quality and make the final decision of whether to move on to stage two of processing

361 with a given siderophore candidate.

362

363 Stage two of processing extracted MS<sup>2</sup> spectra of the apo and Fe-bound forms of candidate siderophores to compare with the predicted MS<sup>2</sup> generated by in silico fragmenter MetFrag (Ruttkies et al., 2016). The in 364 365 silico fragmenter feature was run with a tolerance of 10 ppm on "[M+H]+" and "[M+Na]+" modes. A 366 confidence level of 1-4, from highest to lowest confidence, was then assigned to putative siderophores based on the following criteria: (1) peaks were present in  $MS^1$  and  $MS^2$  spectra, and at least one of the three most-367 368 intense  $MS^2$  fragments matched *in silico* fragmentation, (2) peaks were present in  $MS^1$  and  $MS^2$  spectra, and 369 smaller-intensity fragments matched *in silico* fragmentation, (3) peaks were present in  $MS^1$  and  $MS^2$  spectra, 370 but little to no fragments matched *in silico* fragmentation, and (4) nicely shaped peaks were identified in  $MS^{1}$ 371 spectra but no MS<sup>2</sup> spectra was collected (outlined in Table S45; example spectra in Fig. S6-S9). The 372 confidence levels were modelled after reporting standards for metabolite identification (Sumner et al., 2007). 373 MetFrag pulls chemical structures from publicly-available databases like PubChem or COCONUT (Sorokina 374 et al., 2021), which contain most, but not all variations of siderophores. As such, Fe-bound candidates were 375 usually run against the apo form available in the database, and for siderophores with similar structures but 376 variations in fatty chain length or double bond placement, sometimes only one parent structure was available. 377

378 A 5-point standard curve with known concentrations of siderophore ferrioxamine E was used for 379 quantification of putative siderophores, with a limit of detection of 0.257 nM in the eluent (Fig. S10), or 380 0.07-0.21 pM in the sample depending on sample-to-eluent volume ratio at each site (**Table S1**). MS<sup>1</sup> peaks 381 were integrated for all putatively identified siderophores and peak areas were converted to concentration 382 using the standard curve and the concentration factor of sample volume to eluent volume (Fig. S10). 383 Commercial standards are not available for most siderophores, and different compounds have distinct 384 ionization efficiencies in ESI-MS. Thus, the siderophore concentrations reported here are estimates of 385 siderophore concentrations in these environments based on ferrioxamine E, chosen for its commercial 386 availability and use in prior studies (e.g., (Boiteau et al., 2016)). Additionally, 1 mM of cyanocobalamin was 387 added as an internal standard to each sample aliquot to address any changes in sensitivity during LC-ESI-MS 388 runs. All putative siderophores that were identified with peak areas less than the detection limit were 389 discarded, and all remaining putative compounds with at least confidence levels 1 and 2 at one site were 390 included in the manuscript and are referred to as siderophores throughout. Siderophore identifications remain 391 putative due to inherent uncertainty with assignments by mass, but the confidence levels were designed such 392 that high confidence candidates contain siderophore-like moieties in their fragments. Limited sample 393 volumes prevented analysis via LC-ICP-MS like previous studies, which, in addition to greater availability 394 of commercial standards and more analytical comparisons between ferrioxamine E with other siderophore

types, would allow definitive characterization in future studies. Confidence level 3 and 4 putative siderophores are only included in the Supplementary Information (**Table S56**). In a final step of quality control, EICs for <sup>13</sup>C isotopologues of candidates were inspected to verify matching peak structure.

### 398 **3.5 Microbial community analysis**

399 Microbial community composition was assessed in neutrally buoyant plumes and near venting sites at three 400 sites: Lucky Strike (Station 7; 1670 m), 10 km S of Rainbow (Station 17; 2000 m), and 200 km E of Rainbow 401 (Station 11; 600 m, 1600 m and 2250 m). A range of 1-2 L of seawater were filtered by pressure filtration 402 through sequential 25 mm membrane filters housed in polypropylene filter holders (Whatman SwinLok, GE 403 Healthcare, Pittsburgh, Pennsylvania) using a peristaltic pump and silicone tubing. Samples first passed through a 3 µm pore-size polyester membrane filter (Sterlitech, Auburn, Washington) then onto a 0.2 µm 404 405 pore-size polyethersulfone membrane filter (Supor-200, Pall Corporation, Port Washington, New York). 406 Pump tubing was acid washed with 10% hydrochloric acid and flushed with ultrapure water between each 407 sample. The filters were flash frozen in liquid nitrogen in 2 mL gasketed bead beating tubes (Fisher Scientific) 408 at sea.

409

410 Nucleic acids (DNA) were extracted as described previously (Santoro et al., 2010), with slight modifications. 411 Briefly, cells on the filters were lysed directly in the bead beating tubes with sucrose-ethylene diamine 412 tetraacetic acid (EDTA) lysis buffer (0.75 M sucrose, 20 mM EDTA, 400 mM NaCl, 50 mM Tris) and 1% 413 sodium dodecyl sulfate. Tubes were then agitated in a bead beating machine (Biospec Products) for 1 min, 414 and subsequently heated for 2 min. at 99°C in a heat block. Proteinase K (New England Biolabs) was added 415 to a final concentration of 0.5 mg/mL. Filters were incubated at 55°C for approximately 4 h and the resulting 416 lysates were purified with the DNeasy kit (Qiagen) using a slightly modified protocol (Santoro et al., 2010). 417 The purified nucleic acids were eluted in 200  $\mu$ L of DNase, RNase-free water, and quantified using a 418 fluorometer (Qubit and Quanti-T HS reagent, Invitrogen Molecular Probes).

419

420 The 16S rRNA gene was amplified in all samples using V4 primers (Apprill et al., 2015; Parada et al., 2016) 421 (515F-Y and 806RB) following a previously established protocol (Stephens et al., 2020). Amplicons were 422 sequenced using a paired-end 250bp run on an Illumina MiSeq 500 and demultiplexed by the UC Davis 423 Genome Center. The resulting 16S rRNA amplicon sequences were filtered and trimmed using the DADA2 424 pipeline in R(Callahan et al., 2016). Taxonomic assignments were made with version 138.1 of the SILVA 425 SSU database (Quast et al., 2013) (silva nr99 v138.1 wSpecies train set.fa.gz 426 doi:10.5281/zenodo.4587955; accessed March 2022). Chloroplast and mitochondrial sequences were filtered 427 out of the dataset using the 'phyloseq' R package (v 1.38.0), after which samples had read depths ranging 428 from 9375 - 65486 reads (average  $28425 \pm 20014$  reads) and represented 1010 unique amplicon sequence 429 variants (ASVs). Read counts were transformed from absolute to relative abundance and taxa were 430 aggregated to the Family level. The ten most abundant families present in each sample were visualized using431 the 'ggplot2' package (v. 3.3.5).

432

433 In order to assess the potential of the observed prokaryotic taxa to produce siderophores, we downloaded all 434 siderophore biosynthetic gene clusters (BGCs) in the antismash secondary metabolite database (n = 7909) 435 and used text-string matching to compare genera containing these BGCs to the genera found in our 16S rRNA 436 gene dataset(Blin et al., 2021). We cross-referenced the nomenclature of antismash-predicted siderophores 437 with that of the siderophores identified by LC-ESI-MS in this study, accounting for minor differences in 438 naming convention between the two databases, to determine if microbial community members present at 439 each site were predicted to make any of the siderophores that were measured at that site. Station 38 and 440 Station 12 were the closest sites with siderophore measurements for comparison against the taxonomic 441 samples taken at 200 km E of Rainbow and 10 km S of Rainbow, respectively. Samples for microbial 442 taxonomy and siderophore identity were taken from the same location at Lucky Strike and thus directly 443 compared.

444

## 445 Data Availability

The CSV data reported in this study has been deposited at Zenodo under the DOI: http://doi.org/10.5281/zenodo.7325154. The LC-ES-MS data has been deposited on Massive under the DOI: http://doi.org/doi.10.25345/C5V97ZW7N. Microbial 16S rRNA data have been deposited on GenBank under the accession number BioProject #PRJNA865382. All data is freely available on each of these data repositories.

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- def of Oceanography.
- 470
- 471 Author Contributions: Manuscript preparation, sample/data processing, CSV analysis, and interpretation
- 472 LC-ESI-MS data analysis and interpretation (C.L.H. and P.J.M.), microbial analysis and interpretation
- 473 (J.B.A. and A.E.S.), dissolved iron and derived excess <sup>3</sup>He<sub>xs</sub> measurements, sample collection (A.J.M. L. and
- 474 M.C.L.), microbial data collection and ligand data interpretation (T.M. and K.N.B.), and project design and
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- 476 in editing and revision of the manuscript.
- 477
- 478 **Competing Interest Statement:** The authors declare no competing interests.
- 479

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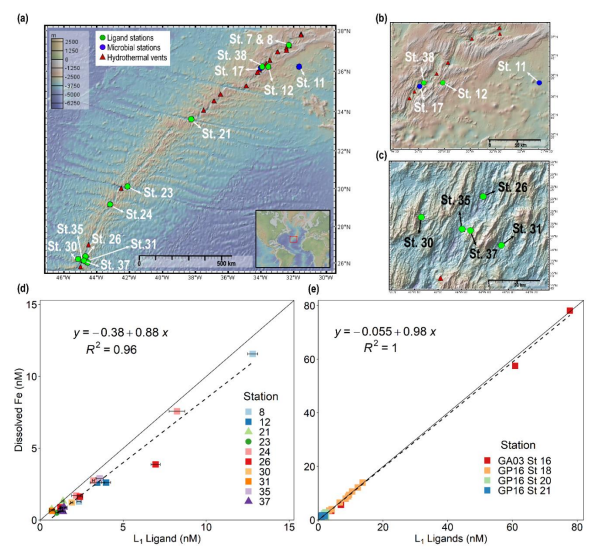
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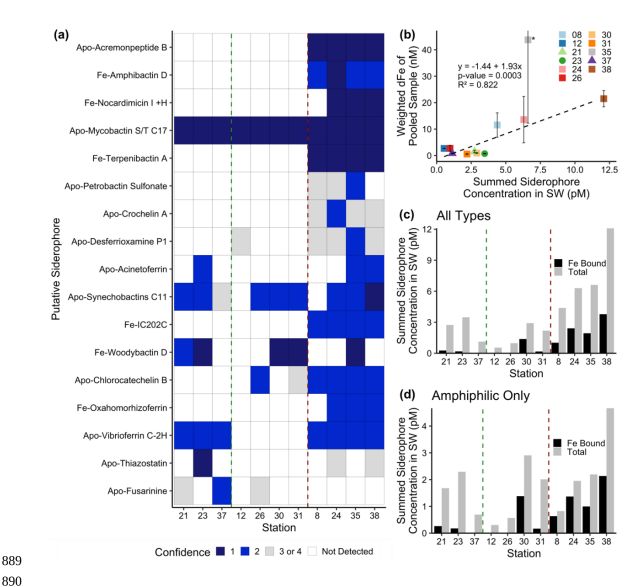


875 Figure 1. Dissolved iron is strongly correlated with L<sub>1</sub> iron-binding ligands in diverse hydrothermal 876 systems. (a) Station map showing the 11 sites investigated along the MAR. Known hydrothermal vents are 877 marked as red triangles(Beaulieu and Szafrański, 2020). Two expanded inset maps for (b) Rainbow and (c) 878 TAG hydrothermal vent fields. For additional information about vent site characteristics refer to Table 1. (d) 879 dFe versus  $L_1$  iron-binding ligands at each vent site in this study showing a ~1:1 correlation (m= 0.88,  $R^2$  = 880 0.96) with dFe in neutrally-buoyant plumes along the MAR. (e) dFe versus L<sub>1</sub> ligands from previous studies 881 over the ridge axis and ~80 km from ridge axis in the Southern East Pacific Rise hydrothermal plume(Buck 882 et al., 2018), and over TAG hydrothermal vent field(Buck et al., 2015). The solid black lines in (d) and (e) 883 are the 1:1 ratio line between dFe and ligand concentrations, and dashed lines show the linear regression for 884 the corresponding data. Square symbols refer to spreading centers, triangles refer to fracture zones, and

- circles refer to alkaline vents. Error bars represent the 95% confidence interval of the data fit as calculated
- by ProMCC(Omanović et al., 2015). The map was created using GeoMapApp version 3.6.14.

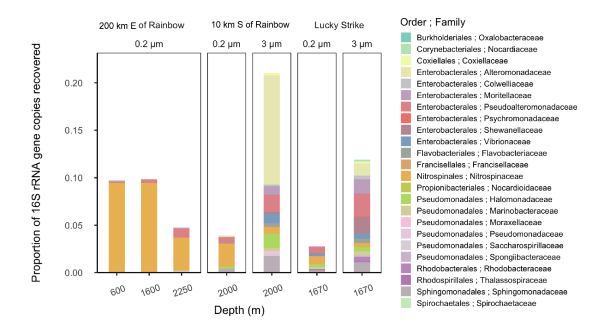
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891 Figure 2. Siderophore presence in hydrothermal plumes along the MAR. (a) Heat map of confidence 892 levels 1-2 (blue gradient, 1 = highest confidence). Gray boxes indicate a detection with lower confidence (see 893 Methods), and white boxes indicate no detection at those sites. The y-axis is ordered from top to bottom in 894 terms of descending mass of the apo (without Fe) form of the siderophore. (b) Model II ordinary least squares 895 regression on dFe versus summed siderophore concentrations (of detections in Fig. 2b), calculated from peak 896 areas, at each site. Since the siderophore analysis was performed on pooled samples, the dFe values in the 897 regression are weighted values based on measured dFe and volume of each constituent of the pooled sample. 898 The vertical error bars represent the standard deviation of dFe of the constituents. TAG (St. 35) — denoted 899 by the asterisk — was not included in the regression due to its large range of dFe values and outlier behavior. 900 (c-d) Fe bound versus total summed concentration of (c) all types of siderophores and (d) amphiphilic 901 siderophores at each station. The vertical green lines separate fracture/diffuse sites from off-axis sites and 902 vertical red lines separate off-axis from on-axis sites as defined in Table 1. Symbols follow Fig. 1.





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906 Figure 3. Relative abundance of putative siderophore-producing taxa. Bar height indicates the proportion 907 of 16S rRNA genes recovered in each sample, separated by depth from water surface, filter size fraction, and 908 site location. Colors correspond to taxonomy. Genera found in MAR vent microbial communities with 909 members in the antismash database predicted to produce siderophores are depicted at the family level. 910

Vent Names	Abbr.	Station	Geology	Host rock	Vent type	Spreading rate (mm/yr)	Summed putative siderophore concentration (pM)	Summed Siderophore concentration/ L <sub>1</sub> ligand (%)*
Lucky Strike	LS	7/8	Spreading	gabbro	Black smoker	20.2	4.38	0.034-0.19
			Center					
33 km E of Rainbow	CER	12	Spreading	-	-	-	0.537	0.013-0.017
			Center					
Rainbow	R	38	Spreading	ultramafic	Black smoker	20.6	12.1	n.a.
			Center					
Hayes Fracture	HFZ	21	Fracture	peridotites/gabbro	-	21.2	2.74	0.20-0.39
Zone			Zone					
Lost City	LC	23	Fracture	ultramafic/gabbro	Alkaline	22.6	3.47	0.27-0.35
			Zone					
Broken Spur	BS	24	Spreading	gabbro	Black	22.9	6.30	0.07-0.29
			Center		smoker/diffuse			
29 km N of TAG	CNT	26	Spreading	-	-	-	0.968	0.014-0.079
			Center					
30 km W of TAG	CWT	30	Spreading	-	-	-	2.91	0.15
			Center					
30 km E of TAG	CET	31	Spreading	-	-	-	2.19	0.31
			Center					

# 911 Table 1. Characteristics of sample locations along the Mid Atlantic Ridge.

Trans-Atlantic	TAG	35	Spreading	gabbro	Black smoker	23.6	6.61	0.18
Geotraverse			Center					
Low Temp Slope	LTS	37	-	-	Diffuse fluids	-	1.13	0.079-0.087

Spreading rates along the Mid-Atlantic Ridge were gathered from the Interridge Database v3.4. Host rock groups were determined from previously discussed classifications(Bazylev, 1997; Kelley and Shank, 2010). Off-axis sites -33 km E of Rainbow, 29 km N of TAG, 30 km E of TAG, and 30 km W of TAG– were far-field locations of their respective vent field. Low Temp Slope was a diffuse-dominated site that was sampled for the first time as a part of this study. Summed putative siderophore concentrations and the percent of L<sub>1</sub> ligand are reported for compounds detected with at least confidence level 1 and 2 at one site. These values do not take into account typical extraction efficiencies of ENV columns for Fe-binding organics. Average L<sub>1</sub> ligand and siderophore concentrations can be viewed in **Table S3** and concentrations for individual siderophores can be observed in **Table S5**.

\*The siderophore sample at each site was pooled from ligand samples, so the percentage of siderophores in the  $L_1$  pool is presented as a range based on the range of  $L_1$  concentrations at each site.

n.a.= unable to be determined

- = unknown

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