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1 Variability in iron (II) oxidation kinetics across diverse hydrothermal sites on
2 the northern Mid Atlantic Ridge

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17 Mid Atlantic Ridge.

18 **Abstract**

19 One of the recently recognized main sources of iron to the deep ocean inventory is the
20 hydrothermal activity associated with mid-ocean ridges. Little is known about the oxidation

21 kinetics of iron(II) within these environments, especially the dependence on physico-
22 chemical parameters such as temperature (T), pH, particle size-fractionation and the effect of
23 organic matter.

24 Following sample collection during the GA13 section cruise, the iron(II) oxidation at six
25 hydrothermal vent sites (Menez Gwen, Lucky Strike, Rainbow, Lost City, Broken Spur and
26 TAG) along the Mid-Atlantic Ridge were investigated, revealing high variability. The Fe(II)
27 oxidation rate constant analysis from multiple stations at two sites (Rainbow and TAG),
28 revealed that factors, other than T and pH, controlled the oxidation process. Experiments on
29 the effect of particle size-fractionation and organic matter at different pH showed that the
30 presence of organic ligands and colloidal size particles delayed the oxidation process, while
31 not affecting the overall pH dependency.

32 Extending our analysis to the broader relationship between the Fe(II) oxidation rate constants
33 across a range of temperatures (between 2 and 25 °C) and pH (between 7 and 8) on a set of
34 selected hydrothermal samples allowed us to derive a multiparametric equation to model the
35 iron(II) oxidation rate constants in the ocean. This equation covers a larger range of
36 temperatures than previous published equations, improving its applicability for global
37 biogeochemical models.

38 **1 Introduction**

39 It is now well-established that phytoplankton growth is limited by the concentration of
40 dissolved iron (Fe) in up to 30% of the world oceans (Moore et al., 2013). Iron in the ocean,
41 can be found as both ferric (Fe(III)) and ferrous (Fe(II)) iron, where 99% of the dissolved
42 Fe(III) is complexed by organic ligands (Rue and Bruland, 1995). In oxic waters, Fe(II) tends
43 to oxidize to the most thermodynamically stable form, Fe(III), in a period of minutes to hours

44 (Santana-Casiano et al., 2005). Oxidation of Fe causes the solubility to decrease (Millero et
45 al., 1995), therefore, reducing its bioavailability (Shaked et al., 2005; Morel et al., 2008).

46 Recently, in the context of the international GEOTRACES program (www.geotraces.org),
47 several studies showed that Fe supplied by hydrothermal activity from mid-ocean ridges can
48 have a substantial impact on Fe distributions at basin scales (Tagliabue et al., 2010; Conway
49 and John, 2014; Resing et al., 2015; Fitzsimmons et al., 2017; Tagliabue et al., 2017).

50 Hydrothermal vent supply of dFe is required to reproduce deep ocean observations
51 (Tagliabue et al., 2010) could represent up to 11-22 % of the dissolved iron (dFe, filtered by
52 0.2 μm) present in the deep ocean globally (Bennett et al., 2008), accounting for a similar dFe
53 flux as continental riverine runoff (Elderfield and Schultz, 1996). Therefore,
54 hydrothermalism acts as an important Fe source in the deep ocean which may affect the
55 global biological carbon pump (Tagliabue et al., 2010; Resing et al., 2015).

56 The concentration of dissolved Fe(II) (dFe(II)) in the open ocean is very low, between 0.02
57 and 2 nmol L^{-1} , due to its fast oxidation under oxic conditions following the Fenton reaction
58 mechanism (González-Davila et al., 2005; Santana-Casiano et al., 2005). However, specific
59 oceanic regions present dFe(II) concentrations which are orders of magnitude higher. In
60 marine sediment interstitial waters, concentrations can reach around 300 $\mu\text{mol L}^{-1}$ (de Baar
61 and De Jong, 2001), upwelling systems can increase Fe(II) concentrations up to 50 nmol L^{-1}
62 (Hong and Kester, 1986), while hydrothermal fluids can contain as much as 25 mmol L^{-1} of
63 dFe(II) (German and Von Damm, 2004). Water emanating from hydrothermal systems
64 contains high concentrations of Fe(II) amongst other elements (Mottl et al., 1979;
65 Mandernack and Tebo, 1993). Studies conducted at hydrothermal vent sites in different ocean
66 basins have revealed different Fe(II) oxidation rate constants, with Fe(II) half-life times
67 varying from 0.5 to 220 min, which has important implications for the dispersion of produced

68 Fe-oxide phases (Massoth et al., 1998; Field and Sherrell, 2000; Statham et al., 2005; Wang
69 et al., 2012; Santana-González et al., 2017; Rouxel et al., 2018). Recently, Gartman and
70 Findlay (2020) compiled all the available data for Fe(II) half-life from four hydrothermal
71 systems. One site was the TAG vent site in the Atlantic Ocean, while the others were located
72 in the Indian and Pacific Oceans, with Fe(II) half-life estimated using large Fe(II) spikes (500
73 nmol L⁻¹) that may affect the role played by different reactive oxygen species (González-
74 Dávila et al., 2006).

75 Temperature (T), pH, and oxygen concentration (O₂) are the primary variables known to
76 directly affect the Fe(II) oxidation (Stumm and Lee, 1961; Millero et al., 1987; Millero and
77 Sotolongo, 1989; Santana-Casiano et al., 2005). Salinity also affects the Fe(II) oxidation, as
78 reported in previous laboratory experiments (Millero et al., 1987; Santana-Casiano et al.,
79 2005). However, other factors such as changes in the inorganic complexation of Fe(II) due to
80 the specific ionic composition of the vent fluid, or organic complexation can influence these
81 oxidation rate constants (Rose and Waite, 2003; Statham et al., 2005). Therefore, the
82 oxidation rate constants must be termed “apparent” due to the effect of organic complexation
83 (Voelker and Sulzberger, 1996), as the nature and concentrations of organic ligands vary
84 between samples. Moreover, pyrite nanoparticles formed in hydrothermal vent sites, have
85 been estimated to oxidize more slowly than dissolved Fe(II) (Yücel et al., 2011), showing
86 that particle composition also influences the oxidation rate constants. The first empirical
87 equation describing the Fe(II) oxidation rate constants in seawater was reported by Millero et
88 al. (1987), and later updated by Santana-Casiano et al. (2005) for nanomolar concentrations.
89 This equation has been included in coupled physical-biogeochemical models in the ocean
90 (OGCBMs) to explore the impact of future climate change on the oxidation of Fe(II)
91 (Tagliabue and Völker, 2011). However, this equation was established for water temperatures

92 ranging from 10 to 25 °C, which leads to discrepancies for the Fe(II) oxidation rate constants
93 for cold, deep ocean water masses.

94 The aims of this study were to measure the oxidation kinetic constants (k') of Fe(II) from
95 different hydrothermal vents and to investigate whether the variations of pH, T and S can
96 explain the observed variability, or if other factors such as the presence of particles, colloids
97 or dissolved organic matter (DOM) are significant contributors. This updated version of the
98 equation incorporates a wider range of temperatures, which lowers the error associated with
99 the theoretical Fe(II) oxidation rate constants. Thus, it becomes applicable to the calculation
100 of theoretical Fe(II) oxidation for the deep waters in which hydrothermal systems are
101 typically found.

102 **2 Methods**

103 *2.1 Sampling sites and characteristics*

104 Seawater samples were collected during the GA13 section cruise (JC156) along the Mid
105 Atlantic Ridge (MAR) onboard the *RRS James Cook*. This cruise departed Southampton, UK
106 on December 20th, 2017 and arrived into Guadeloupe, France on February 1st, 2018.

107 Alongside the large scale ocean section survey, sampling focused on six hydrothermal vent
108 sites along the MAR: Menez Gwen (37° 50'N 31° 31'W; Charlou et al., 2000), Lucky Strike
109 (37° 17'N 32° 16'W; Charlou et al., 2000), Rainbow (36° 14'N 33° 54'W; Douville et al.,
110 2002), Lost City (30° 7'N 42° 7'W; Kelley et al., 2001), Broken Spur (29 °N; James et al.,
111 1995), and TAG (26°N; Gamo et al., 1996) (Fig. 1). These six hydrothermal vent sites
112 represent a wide range of possible hydrothermal conditions due to different underlying
113 geological conditions. Lost City and Menez Gwen are the shallowest sites, with the vent site
114 depths located at 700-720 m and 840-850 m, respectively. The Lucky Strike hydrothermal

115 field is situated between 1560 and 1730 m depth. The Rainbow vent site is found at 2270-
116 2280 m, within the average range (2000-3000 m) of oceanic ridge depth (Searle, 2013), while
117 Broken Spur and TAG are deeper hydrothermal systems located at 3100-3300 m and 3670-
118 3700 m, respectively. Chemically, Lost City is an alkaline hydrothermal vent, with pH
119 ranging from 9.0 to 9.8. The vent fluid is at a lower temperature of 40-75 °C, compared to
120 185-365 °C at the other vent sites and has extremely low concentrations of most trace
121 elements (e.g. Fe below the detection limit compared to 2-24000 $\mu\text{mol L}^{-1}$ at the other vent
122 sites) (Ludwig et al., 2006; Fouquet et al., 2013). All other hydrothermal vent sites are acidic
123 with pH lower than 4, and act as important sources of trace metals (Charlou et al., 2000;
124 Douville et al., 2002; Andreani et al., 2014). Geologically, Menez Gwen and Lucky Strike are
125 similar, their basement rocks are mafic, enriched-Mid-Ocean Ridge Basalt (E-MORB,
126 enriched with light rare earth elements, LREE), and present sulfide mounds. Rainbow also
127 presents sulfide mounds, but its basement rock is ultramafic. Further south, the alkaline Lost
128 City is also situated above an ultramafic base with carbonate chimneys. Finally, Broken Spur
129 and TAG have sulfide mounds and basement rocks formed by normal-MORB (N-MORB,
130 LREE depletion) (references within Fouquet et al., 2010).

131 Seawater samples were collected according to the GEOTRACES guidelines
132 (<http://www.geotraces.org/images/Cookbook.pdf>). Briefly, a titanium rosette fitted with 24 x
133 10 L trace metal-clean Teflon-coated OTE (Ocean Test Equipment) bottles, a CTD profiler
134 (Sea-bird Scientific), light scattering sensor (PMEL MAPR), and a redox potential sensor
135 (PMEL MAPR), were deployed on a conducting Kevlar wire to collect samples from the
136 water column. Sampling depths were actively chosen from the continuous T, salinity (S),
137 light scattering sensor ([https://data-](https://data-search.nerc.ac.uk:443/geonetwork/srv/api/records/8ba648f2759d57fbb2246671f8065939)
138 [search.nerc.ac.uk:443/geonetwork/srv/api/records/8ba648f2759d57fbb2246671f8065939](https://data-search.nerc.ac.uk:443/geonetwork/srv/api/records/8ba648f2759d57fbb2246671f8065939))
139 and redox potential readings.

140 Upon recovery, the OTE bottles were transported into an ISO class 6 clean air van. Unfiltered
141 samples (UF) for Fe(II) oxidation kinetics were immediately transferred into thoroughly
142 rinsed acid-cleaned 250 mL or 1 L low-density polyethylene bottles (LDPE; Nalgene)
143 following the GEOTRACES Cookbook (<http://www.geotraces.org/images/Cookbook.pdf>)
144 and frozen at -20 °C. Samples remained frozen in a -20 °C freezer for one year before their
145 analysis. pH samples were collected in 125 mL borosilicate bottles and kept in the dark until
146 the time of analysis.

147 **2.2 pH determination**

148 pH was measured onboard and during laboratory-based kinetic experiments in the free scale
149 using a Tris buffer solution (Millero, 1986). Onboard pH samples were kept in the dark,
150 inserted into a 25 °C water bath, and measured at 25 °C, within less than 1 hour after
151 collection. Laboratory-based kinetic experiments used a Titrino 719 (Metrohm™) which
152 automatically added 0.01M hydrochloric acid (HCl, PANREAC Hiperpur-plus) in order to
153 keep the pH constant. The pH at *in situ* conditions were computed considering the total
154 alkalinity for an average profile of the area (Key et al., 2015; Olsen et al., 2016). The average
155 profile was obtained from plotting all the available alkalinity data from the studied region
156 (20-30 °N and 40-50 °W). The associated error of the estimate was $\pm 6 \mu\text{mol kg}^{-1}$. Gray et al.
157 (2011) showed that computed pH values are relatively insensitive to errors in alkalinity, and
158 that pH measurements can be used to compute the magnitude and variability of inorganic
159 carbon variables. A fluctuation of $400 \mu\text{mol kg}^{-1}$ in total alkalinity values (e.g. from 2325 to
160 $2725 \mu\text{mol kg}^{-1}$) will change the pH in +0.005, which lies within the error of determination
161 for potentiometric pH. The set of constants from Lueker et al. (2000) were used.

162 All the pH data in the text and in the figures, including kinetics and profile data are expressed
163 in the free scale.

164 2.3 Fe(II) measurements

165 In order to measure the Fe(II) oxidation rate constants, continuous Fe(II) concentrations were
166 determined using the FeLume System (Waterville Analytical), with a data point acquisition
167 every 1 s. This direct Flow Injection Analysis system uses luminol (5-amino-2,3-dihydro-1,4-
168 phthalazinedione, Sigma-Aldrich) as the reagent (King et al., 1995) which reacts with Fe(II).
169 The sample was continuously aerated with pure air (Carbueros Metalicos S.A.). A 0.8 nmol L⁻¹
170 addition of Fe(II) (ammonium iron(II) sulfate hexahydrate, Sigma-Aldrich) was introduced
171 into each sample and the Fe(II) concentration was continuously monitored until the signal
172 had reduced by 70 %. This method produces apparent rate constants (k_{app}) because the
173 oxidation rate constant is measured on a Fe(II) spike rather than on the ambient Fe(II).
174 Although, the oxidation rate constant is not affected by the initial Fe(II) in solution (Millero
175 et al., 1987; Roy et al., 2008; Roy and Wells, 2011; Santana-González et al., 2018), the
176 equilibrium with the organic ligands and particle surfaces could be affected. Moreover, due to
177 the effects of organic complexation and that the concentrations of the organic ligands in each
178 sample are not constant, the apparent oxidation rate constants are consequently not true
179 constants.

180 Some previous oxidation kinetic studies have found a second-order dependency of the Fe(II)
181 concentration against time (e.g. Statham et al., 2005). However, in oxygen saturated samples
182 a pseudo-first order rate dependency was obtained, where the slope ($\ln([\text{Fe(II)}])$ against time)
183 represents the pseudo-first order rate constant k' (Eq. 1). The linearity remained longer than
184 the half-life time and was thus considered acceptable in all the analyses ($R^2 = 0.983 \pm 0.017$,
185 n samples = 298). The observed linearity confirms that the equilibrium between the added
186 Fe(II) with organic ligands and particles is achieved before 30 s (our first Fe(II)
187 determination time limit). The k' error associated with these high R^2 values was 2% for *in*

188 *situ* condition experiments, and 1% for experiments carried out at 25 °C. The half-life ($t_{1/2}$) of
189 Fe(II) in seawater can be calculated from k' (Eq. 1 and 2).

$$190 \quad d[Fe(II)]/dt = -k_{app}[O_2][Fe(II)] \quad (1)$$

191 In oxygen saturated concentrations, the corresponding pseudo-first order rate constant $k' =$
192 $k_{app}[O_2]$.

$$193 \quad t_{1/2} = \ln(2)/k' \quad (2)$$

194 The minimum energy required to start the oxidation reaction (activation energy, E_a , J mol⁻¹)
195 and the corresponding preexponential or frequency factor (A) were computed from the
196 calculated k' using the empirical Arrhenius equation, where R is the universal gas constant
197 (8.314 J mol⁻¹ K⁻¹):

$$198 \quad \log(k') = \log(A) - \frac{E_a}{R \cdot \ln(10) \cdot T} \quad (3)$$

199 Iron (II) blanks were obtained from a 2 h aerated, magnetically stirred sample which was kept
200 in the dark to achieve the oxidation of Fe(II) (Santana-Casiano et al., 2005). Iron (II) in these
201 blanks always below the detection limit. Calibration curves (0 to 1 nmol L⁻¹) were produced
202 daily and confirmed linearity in the Fe(II) signal. The detection limit, defined as three times
203 the standard deviation of the blank after being aerated for 10 minutes, was lower than 50
204 pmol L⁻¹. Reproducibility was checked by running the same sample three times throughout
205 the day after an addition of 1 nmol L⁻¹ of Fe(II) with an agreement in concentrations that was
206 better than ± 0.1 nmol L⁻¹. Accuracy could not be determined since reference materials
207 currently do not exist. Results are systematically given as $t_{1/2}$ (in min), followed by the
208 measured k' (in min⁻¹).

209 **2.4 Experimental setup**

210 Using natural seawater samples, kinetic experiments were carried out under different
211 conditions, with pH and T set to *in situ* conditions (data shown in the supplementary Table
212 S1), and with pH and T normalized to constant values, in unfiltered, filtered and filtered and
213 UV-irradiated samples. These later experiments required a high sample volume, limiting the
214 number of experiments.

215 The steps required for the Fe(II) oxidation kinetics determination are as follows. The day
216 before the experiments, three to four frozen samples were taken out of the freezer and placed
217 in a fridge set to 4 °C. The next morning, a temperature-controlled water bath was set to the
218 coldest *in situ* potential temperature (supplementary Table S1). While the water bath was
219 cooling down, each sample was subsampled into three 60 mL acid cleaned LDPE bottles and
220 placed back into the fridge. One of the subsamples was inserted into the temperature-
221 controlled water bath, which was set to the original *in situ* potential temperature of the
222 sample. The sample was left in the water bath for acclimatization during 15 min and then
223 transferred into the reaction vessel. In the reaction vessel, the sample was aerated prior to and
224 during the study. After 10 minutes of aeration, the Fe(II) addition was made and the Fe(II)
225 concentration and time were continuously monitored. After finishing with the coldest
226 subsample, the temperature of the water bath was increased to the following coldest
227 temperature, and the next sample was left to acclimate.

228 ***2.4.1 Quantification of the temperature and pH effect on the iron (II) oxidation rate*** 229 ***constant***

230 In order to improve the temperature range of the existing Fe(II) oxidation kinetic equation, 1
231 L samples of deep ocean water were collected during the JC156 cruise, at stations 13 (2339
232 m), 16 (2051 m), 24 (2829 m), 30 (2700 m), 31 (2796 m), and 35 (1999, 3350 and 3601 m).
233 The samples remained frozen at -20 °C during one year until analysis. Two days prior to the

234 analysis, samples were thawed in a 4 °C fridge. This allowed a slow melting of the samples in
235 the dark, thus limiting any photoproduction of Fe(II) and also removal of any superoxide and
236 hydroxyl radicals in the sample (Voelker and Sedlak, 1995).

237 The samples were then transferred into 60 mL LDPE bottles (one for each condition).
238 Temperature effect experiments were performed at constant pH, and those for pH effect at
239 constant temperature. The temperature effect experiments were carried out at *in situ*
240 temperature, 10, 20 and 25 °C at a fixed pH of 8. The pH effect experiments were carried out
241 at *in situ* pH, and pH 7, 7.5 and 8 at a fixed temperature of 25 °C.

242 ***2.4.2 Oxidation rate constants through the water column at the six hydrothermal vent sites***

243 To assess the expected Fe(II) oxidation rate constants at *in situ* conditions, the experiments
244 were carried out at *in situ* potential temperature and pH ($T_{is}pH_{is}$). The temperature was then
245 increased to 25 °C ($T_{25}pH_{is}$) to obtain the temperature effect on the oxidation rate constants.
246 A third set of experiments were performed at 25 °C and pH 8 ($T_{25}pH_8$). The T and pH
247 normalization of all the samples allowed for the interpretation of the effect of other variables,
248 such as organic matter and particles, in the oxidation rate constants. This allows to investigate
249 where seawater composition is influencing the oxidation rate, a process that cannot be
250 isolated when comparing two samples with different temperatures and/or pH.

251 ***2.4.3 Experimental setup to assess the effect of colloidal and organic matter***

252 The unused unfiltered (UF) sample volumes remaining from the quantification of the
253 temperature and pH effect experiments were used to assess the impact of particles, colloids
254 and organic matter. Unfiltered samples underwent sequential filtration. Three aliquots were
255 filtered through 0.2 µm (F0.2; Whatman® Anodisc), and one of them was then filtered
256 through 0.02 µm (F0.02; Whatman® Anotop®), while the last aliquot was filtered through

257 0.02 μm followed by UV-irradiation (F0.02+UV). All samples were left in the dark for 6
258 days to limit the effect of any photochemically generated reactive oxygen species (ROS)
259 produced during the UV-irradiation (Yuan and Shiller, 2001; Roy et al., 2008). All samples
260 were analyzed following the same procedure as for the assessment of the pH effect (see
261 section 2.4.2 above). In order to assess whether the generated ROS species could affect the
262 Fe(II) oxidation rate constants determined after 0.02 μm filtration and UV-irradiation, the
263 effect of the amount of hydrogen peroxide (H_2O_2) generated during this process was
264 considered (Yuan and Shiller, 2001; O'Sullivan et al., 2005). Other short-lived ROS species
265 such as hydroxyl radicals were not accounted for, since their nanosecond half-life times do
266 not allow for their determination using our analytical method.

267 **3 Results**

268 *3.1 Temperature and pH effects on the oxidation rate constants*

269 Experiments for the temperature effect study were carried out at a pH of 8. When plotting log
270 k' vs. $1/T$ (K^{-1}) (Fig. 2, Eq. 3), results showed an average slope ($-\Delta E_a/R/\ln(10)$) of $-4992 \pm$
271 232 . Notably, the data obtained from samples collected west of TAG at 2700 m depth
272 (Station 30) showed a slope of -3732 ± 216 , which is more than three standard deviations
273 away from the average value. This lower slope corresponds to a much lower activation
274 energy for this sample, with a value of $71 \pm 4 \text{ KJ mol}^{-1}$. This effect was probably caused by
275 the interaction of organic compounds or particles with the Fe(II) species which affected the
276 limiting Fe(II) oxidation step, and therefore a result of a different oxidation reaction
277 mechanism. If this sample is removed, the average slope increases to -5434 ± 183 , and the
278 activation energy of all the samples considered was $104 \pm 3 \text{ KJ mol}^{-1}$. These values are within
279 the experimental error of previous determinations from Gulf Stream water (-5362 ± 162 , and
280 an activation energy of $103 \pm 3 \text{ KJ mol}^{-1}$; Santana-Casiano et al., 2005).

281 When experiments were carried out at a constant T of 25 °C and at 5 different pH values in
282 the range from 7 to 8, $\log k'$ and pH were linearly related, with slopes of 1.00 ± 0.06 (Fig. 3)
283 and with no data points outside of 3 standard deviations. As with the temperature relationship,
284 the sample from the station West of TAG at 2700 m presented a slope (1.19 ± 0.08) out of the
285 1:1 dependence. A second sample, (TAG at 1999 m depth, over 1600 m above the vent site)
286 showed a lower slope (0.81 ± 0.12).

287 The unique use of lower temperature samples in this study, allowed a better estimation of
288 Fe(II) oxidation kinetic rate constants at temperatures lower than 10 °C, increasing the range
289 of applicability of the Fe(II) oxidation kinetic equation of Santana-Casiano et al. (2005).

290 Using the equation and dataset from Santana-Casiano et al. (2005), which had a wider range
291 in pH and salinity, together with the additional pH data and a wider range of temperature
292 from the present study, a revised equation was obtained. The pH and salinity dependence did
293 not change, while the T relationship was modified (Eq. 4). Thus, an updated equation for the
294 theoretical Fe(II) oxidation appropriate for deep waters is presented:

$$\begin{aligned} 295 \log k'(s^{-1}) = & 35.627 - 6.7109 * pH + 0.5342 * pH^2 - 5434.02/T - 0.04406 * S^{1/2} - \\ 296 & 0.002847 * S \end{aligned} \quad (4)$$

297 The T factor has been modified from $-5362.6/T$ to $-5434.02/T$ obtained from the average
298 slope of Figure 2. To assess the performance of the updated Fe(II) oxidation kinetic equation
299 (Eq. 4), it was compared to three existing equations. The chosen equations were the first
300 derived equation (Millero et al., 1987), the equation by Santana-Casiano et al. (2005), and a
301 recent equation using Labrador Seawater (Santana-González et al., 2019). The four equations
302 were compared to the measured oxidation rate constants at *in situ* pH and T conditions. The
303 residual sum of squares (RSS) and root mean square error (RMSE) between predicted and
304 observed values were calculated. Smallest values of RSS and RMSE indicate the best

305 equation fit to observations. Additionally, the difference in logarithms of the likelihood
 306 ($\Delta\log(L)$), based on a Gaussian error distribution with a constant variance of log-transformed
 307 data, was used to compare the equation performance (Armstrong et al., 2001 and references
 308 therein). The $\log(L)$ was calculated as in Moriceau et al. (2009), where a difference in $\log(L)$
 309 of at least two points is considered as a criterion of a better fit to the observations. The three
 310 most recent equations present better RSS, RMSE and $\log(L)$, with the new equation presented
 311 here providing consistently better statistical results (Table 1).

	Millero et al. (1987)	Santana-Casiano et al. (2005)	Santana- González et al. (2019)	González- Santana et al. (Eq. 4 of this manuscript)
RSS	0.019	0.003	0.006	0.002
RMSE	0.019	0.008	0.011	0.007
$\log(L)$	24.2	33.6	32.5	36.7

312 Table 1. Statistical comparison of the available Fe(II) oxidation kinetic equations with the
 313 measured oxidation rate constants for deep ocean waters without including the hydrothermal
 314 neutrally buoyant plumes. Low residual sum of squares (RSS) and root mean square error
 315 (RMSE) indicate the best equation fit to observations. Greater than two points of difference
 316 in the difference in logarithms of the likelihood ($\Delta\log(L)$) presents a better fit.

317 **3.2 Iron(II) oxidation rate constants throughout hydrothermal plumes along the MAR**

318 Figures 4 and 5 present the half-life times of Fe(II) samples collected above each
 319 hydrothermal site with Figure 5 summarizing the results using boxplots. The measured $t_{1/2}$

320 values presented a wider range than those determined using the physico-chemical parameters
321 in the improved theoretical equation. The Lost City vent site, was the only station where all
322 the measured $t_{1/2}$ at $T_{is}pH_{is}$ conditions presented measured oxidation rate constants higher
323 (shorter $t_{1/2}$) than the theoretically calculated rate constants. At $T_{25}pH_8$ conditions, all the
324 measured oxidation rate constants were higher than the theoretical ones for all stations except
325 at TAG.

326 The theoretical $t_{1/2}$ values presented in Figure 5a, show the expected high variability in the
327 Fe(II) half-life times associated with T, pH and S. These discrepancies were further
328 highlighted in the measured samples, where only Menez Gwen and Lucky Strike theoretical
329 values are within the measured results (a paired t-test did not show a statistical difference
330 between the theoretical and the measured $t_{1/2}$, with $P > 0.35$ for both).

331 At Menez Gwen (Fig. 4a), measured *in situ* $t_{1/2}$ varied between 8.3 and 32.0 min (k' between
332 0.084 and 0.022 min^{-1}), with short measured $t_{1/2}$ observed in the samples collected at 743,
333 821, and 827 m. When the temperature was increased and fixed to 25 °C, while the pH was
334 kept at *in situ* conditions (Supplementary Table S1), a significant T effect was observed in k' ,
335 with the measured $t_{1/2}$ decreasing to less than 4 min. When the pH was fixed to a constant
336 value of 8 along the water column, the measured $t_{1/2}$ was slightly affected indicating a small
337 effect, as the change in pH was always lower than 0.22. Overall, at T_{25} , samples presented
338 measured $t_{1/2}$ in the same order of magnitude as the theoretical $T_{25}pH_8$. Samples collected
339 within the lower part of the plume, at less than 30 m above the seafloor (hereafter *asf*)
340 presented higher oxidation rate constants than theoretical ones ($t_{1/2}$ decreased).

341 Lucky Strike (Fig. 5a) presented lower oxidation rate constants than Menez Gwen related to
342 lower temperatures in deeper waters (from about 8.5 °C to 4.5 °C). From 1600 m to 1680 m,
343 measured $t_{1/2}$ averaged 46.5 ± 2.5 min (k' of 0.015 ± 0.001 min^{-1} , $n = 4$) (Fig. 4b). This $t_{1/2}$

344 average is higher than the theoretical value of 39.7 min, while at 1703 m, the Fe(II) half-life
345 decreased to 26.3 min (k' to 0.026 min^{-1}). Overall, at T_{25} , samples presented measured $t_{1/2}$ in
346 the same order of magnitude as the theoretical $T_{25} \text{ pH}_8$. *In situ* pH averaged 7.89, where the
347 increase in acidity produced a lower oxidation rate constant than the measured $t_{1/2}$ at $T_{25} \text{ pH}_8$.

348 At Rainbow, station 16, the $t_{1/2}$ measured at *in situ* conditions presented a wide range of
349 variation, from 17.3 to 65.3 min (k' varied from 0.040 min^{-1} to 0.011 min^{-1}). The theoretically
350 expected values were within that range, with an average value of $39 \pm 3 \text{ min}$ ($n=8$, Fig. 4c).
351 The shortest $t_{1/2}$ were observed between 1986 m and 2108 m (175 to 295 m *asf*) averaging
352 $22.2 \pm 3.6 \text{ min}$ ($k' = 0.032 \pm 0.006 \text{ min}^{-1}$) while the shorter $t_{1/2}$ were found below the plume
353 and above 300 m *asf* (Fig. 4c). The light scattering data shows that the Rainbow plume
354 reached heights over 400 m *asf* (although samples were only collected within the first 330 m).
355 Temperature was an important factor controlling the oxidation rate constant. At T_{25}
356 conditions $t_{1/2}$ were similar to the theoretical value. When the pH was fixed to 8, small
357 changes were observed, and most of the values were close to the theoretical value except for
358 those around 1986 m and 2108 m which had shorter $t_{1/2}$.

359 Lost City (Fig. 4d), the only alkaline hydrothermal vent, was drastically different from the
360 other hydrothermal sites (Fig. 4d), with temperatures around $10 \text{ }^\circ\text{C}$ and pH_{is} of 8.07
361 (Supplementary Table S1). The theoretical $t_{1/2}$ associated with these temperatures and pH
362 ranged between 16 and 20 min. However, most of the measured $t_{1/2}$ were shorter than 3 min,
363 reaching 0.5 min (k' of 1.299 min^{-1}) at 725 m with the longest $t_{1/2}$ of 7.8 min (k' of 0.088 min^{-1})
364 at 750 m. At 750 m depth with $T_{25} \text{ pH}_{\text{is}}$ conditions, the $t_{1/2}$ decreased to 1.9 min (k' of 0.357
365 min^{-1}) and further decreased to 1.5 min (k' of 0.468 min^{-1}) at $T_{25} \text{ pH}_8$ conditions. All the
366 measured $t_{1/2}$ were significantly shorter than the theoretical $t_{1/2}$ of 3.2 min (k' of 0.218 min^{-1})
367 at normalized conditions.

368 Broken Spur presented a wide range in measured $t_{1/2}$, with values under *in situ* conditions
369 ranging from 1.0 to 65.1 min (k' ranging from 0.714 min^{-1} at 2929 m to 0.011 min^{-1} at 2858
370 m; Fig. 4e). Two different behaviors can be observed. The two shallowest samples (2820 and
371 2830 m) were collected in local light scattering minima, with $t_{1/2}$ averaging $63.2 \pm 2.7 \text{ min}$ (k'
372 of $0.011 \pm 0.001 \text{ min}^{-1}$) and values slightly higher than the theoretical ones. Deeper than 2830
373 m, the samples were collected in local light scattering maxima, where half-life times were
374 shorter with an average $t_{1/2}$ of $1.9 \pm 1.2 \text{ min}$ (averaged k' of $0.464 \pm 0.220 \text{ min}^{-1}$).
375 Experiments at $T_{25}\text{pH}_8$ also showed that samples below 2830 m presented shorter $t_{1/2}$ (high k'
376 values) than what was theoretically expected.

377 Half-life times measured at *in situ* conditions from the TAG vent site samples presented the
378 widest range in $t_{1/2}$ (Figs. 4 and 5a), varying from 35.1 to 131.2 min (k' between 0.019 min^{-1}
379 and 0.005 min^{-1}). These samples were characterized by little variability of T_{is} ($2.67 \pm 0.01 \text{ }^\circ\text{C}$)
380 and pH_{is} (8.17 ± 0.01), producing a stable theoretical $t_{1/2}$ of $43 \pm 2 \text{ min}$. Furthermore, the
381 measured $t_{1/2}$ were the longest obtained in these experiments. When the experiments were
382 repeated at both fixed T_{25} and pH_8 the measured values encompassed the theoretical value of
383 3.3 min, with values slightly higher at around 3325 m (4.3-4.4 min). Therefore, even when
384 temperature and pH affected the observed oxidation rate constants, the high range of values at
385 *in situ* conditions could not be explained by only these two physico-chemical variables.

386 ***3.3 Investigating the spatial variability of Fe(II) oxidation rate constants around Rainbow*** 387 ***and TAG sites***

388 To investigate the spatial variability in $t_{1/2}$ for Fe(II), samples were collected from four
389 stations around the Rainbow site at 0.3° (30 km) S (St 15), E (St 12) and W (St 13) and 1°
390 (120 km) E (St 10) (Fig. 1) and from five stations around TAG at 0.3° N (St 26), S (St 27), W
391 (St 30), and E (St 31) and 1° W (St 29) (Fig. 1). At both vent sites, turbidity and light

392 scattering anomalies matched and described the shape of the plume (gray shading in Fig. 4).
393 The Rainbow plume anomaly was observed between 1930-1995 m, 2060-2090 m, and most
394 strongly between 2100 and 2210 m. At the TAG site, the signal was three times lower than
395 that observed at Rainbow, indicating the presence of a lower amount of particles, which could
396 also be due to a displacement of the sampling with respect to the vent site. The largest
397 anomalies were measured between 3160-3390 m, with several minor anomalies above (e.g.
398 2980-3030 m) and below (e.g. 3430-3520 m and 3600-3625 m; Fig. 4).

399 Turbidity and light scattering were useful for tracking the plume displacement even at 30 km
400 from the vent site. The Rainbow plume was observed to the south and west (Stn. 13 and 15)
401 at depths between 2000 and 2040 m, and 2270 and 2300 m. For TAG, only stations 27 (south
402 of TAG, but inside the MAR valley) and 31 (east) showed deeper and thinner turbidity
403 anomalies between 3400 and 3690 m and a smaller signal between 2920 and 3060 m.

404 For the Rainbow field, samples collected at the vent site in the 1980-2150 m range had
405 shorter $t_{1/2}$ at *in situ* conditions (high k' values) than the theoretical $t_{1/2}$ (Fig. 6a). All the other
406 samples (except for Stn 10 at 2430 m) had higher $t_{1/2}$ values than those calculated
407 theoretically, and greater $t_{1/2}$ than observed at Rainbow at depths shallower than 1980 m,
408 indicating very low oxidation rate constants. This allowed Fe(II) to stay in solution three
409 times longer than under inorganic seawater conditions. However, when T and pH were fixed
410 (Fig. 6b), the oxidation rate constants between stations were much closer to the theoretical
411 value, indicating the T and pH control over the Fe(II) oxidation rate constant. Nevertheless,
412 plume samples at the 2100-2210 m range and around 2400 m indicated $t_{1/2}$ values that were
413 still at least double the theoretical value.

414 At the TAG hydrothermal field, stations situated towards the N and W of TAG presented
415 shorter $t_{1/2}$ (high k' values) than those at the vent site and towards the S and E (Fig. 7a).

416 Station 30, situated 30 km W of TAG, exhibited both patterns, with long $t_{1/2}$ (low k' values) at
417 depths shallower than 2600 m and short $t_{1/2}$ (high k' values) at depths below 2700 m. When
418 all the TAG hydrothermal field station samples were normalized (i.e. T_{25pH8} ; Fig. 7b), most
419 of the variability was removed, and $t_{1/2}$ averaged 3 ± 1.5 min. The same pattern as for *in situ*
420 conditions was maintained, with stations situated towards the N and W having shorter $t_{1/2}$
421 than the theoretical values. These three stations were characterized by their lower turbidity in
422 the deep waters.

423 ***3.4 The effect of particles, colloids, and organic matter on oxidation rate constants***

424 Experiments with samples from Rainbow (Stn 16, 2051 m), West of Rainbow (Stn 13, 2339
425 m), Broken Spur (Stn 24, 2829 m) and TAG (Stn 35, 3350 m) were performed to improve our
426 understanding of the impact of particulate and organic matter on the oxidation kinetics of
427 hydrothermal vent fluids. The Rainbow sample at 2051 m, located outside the plume, did not
428 show any difference in the $\log(k')$ due to the presence of particulates, with values at each pH
429 unit varying by ± 0.03 and an average slope of $\log(k')$ against pH of 0.96 ± 0.07 for UF, F0.2
430 and F0.02 treatments (Fig. 8). For the other three samples (Figs. 8b-d), oxidation rate
431 constants for UF and F0.2 did not present significant differences, with values that were
432 reproducible within ± 0.02 for the studied pH range. However, F0.02 samples had higher
433 oxidation rate constants, while also maintaining the pH dependence (0.95 ± 0.07). Therefore,
434 the presence of colloidal size particles delayed the oxidation process while not affecting the
435 pH dependence, as shown in Fig. 8, where the intercept changes but the slope is unchanged.
436 All of the UV irradiated samples in Figs. 8a-d indicate that the absence of dissolved organic
437 matter increased the oxidation rate constants. For TAG (3350 m) and Rainbow (2051 m), the
438 k' values were so high that it was impossible to determine them at pH = 8 under our
439 experimental conditions.

440 4 Discussion

441 4.1. *Derived oxidation rate constants as a function of pH and temperature*

442 In the 1990s, Fe(II) oxidation kinetics variability was reported at the TAG hydrothermal field
443 (2-3 min) (Rudnicki and Elderfield, 1993) and near the Juan de Fuca Ridge hydrothermal
444 systems (32 h) (Chin et al., 1994; Massoth et al., 1994) using the Millero et al. (1987)
445 equation. Nevertheless, the variability between those two sites was difficult to explain and
446 reflected fluctuations caused by differences in O₂, T, and pH among the different sites.

447 When combining the six hydrothermal vent sites on the MAR (Fig. 9), the relationship
448 between the measured oxidation rate constants under *in situ* conditions against the
449 theoretically derived oxidation rate constants as a function of pH and temperature (in aerated
450 water) can be obtained. Overall, equation 4 can explain 76 out of 104 data points. When
451 color-coding the data points by stations, we observed that all the outliers corresponded to
452 samples that were collected inside hydrothermal plumes (defined by turbidity, light scatter,
453 and redox potential). Consequently, the results obtained from the theoretical equation (Eq. 4)
454 and figure 9, agree well with previous results that O₂, S, T, and pH can be used to
455 characterize the Fe(II) oxidation rate constant in open ocean waters but not from within
456 hydrothermal plumes.

457 Our study clearly indicates (Figs. 8 and 9) that the presence of both colloidal particles and
458 dissolved organic matter increases the half-life of Fe(II) (decreased k'). This agrees with
459 previous work showing a decrease in the Fe(II) oxidation rate constants in seawater in the
460 presence of some natural organic compounds (Santana-González et al., 2019). However,
461 organic matter can have a positive or negative effect on the oxidation rate constants of Fe(II)
462 depending on the type of organic matter and its degree of remineralization (Rose and Waite,

2003). The variability in the effect of total organic carbon on Fe(II) oxidation proves that total organic carbon cannot be used as a variable in an equation describing k' (Santana-Casiano et al., 2000; Santana-González et al., 2018; Santana-González et al., 2019). In this sense, a more detailed characterization of organic matter would be necessary, especially in a hydrothermal environment.

4.2. Influence of organic species on Fe(II) oxidation rate constants

The Fe(II) oxidation rate constant (Eq. 5) can be expressed as a function of each Fe(II) species (Fe^{2+} , FeOH^+ , Fe(OH)_2 , FeHCO_3^+ , $\text{Fe(CO}_3\text{)}$, $\text{Fe(CO}_3\text{)}_2^{2-}$, $\text{Fe(CO}_3\text{)OH}^-$, FeCl^+ , FeSO_4 , FeH_3SiO_4 and any organically complexed Fe species). Species such as $\text{Fe(CO}_3\text{)}_2^{2-}$ and Fe(OH)_2 could become important at pH higher than 8 (Ussher et al., 2004; González-Davila et al., 2005; Santana-Casiano et al., 2005) in solution. Considering the side reaction coefficient for each inorganic (α_i) and organic (α_j) Fe(II) species in seawater and the associated second-order oxidation rate constant for the kinetically reactive Fe(II) species (k_i and k_j) (Eq. 6) (Millero and Sotolongo, 1989; King, 1998; Santana-Casiano et al., 2005), the apparent oxidation rate constant can be expressed as:

$$k_{\text{app}} = k_{\text{Fe}^{2+}}\alpha_{\text{Fe}^{2+}} + k_{\text{FeOH}^+}\alpha_{\text{FeOH}^+} + k_{\text{Fe(OH)}_2}\alpha_{\text{Fe(OH)}_2} + k_{\text{FeHCO}_3^+}\alpha_{\text{FeHCO}_3^+} + k_{\text{Fe(CO}_3\text{)}}\alpha_{\text{Fe(CO}_3\text{)}} + k_{\text{Fe(CO}_3\text{)}_2^{2-}}\alpha_{\text{Fe(CO}_3\text{)}_2^{2-}} + k_{\text{Fe(CO}_3\text{)OH}^-}\alpha_{\text{Fe(CO}_3\text{)OH}^-} + k_{\text{FeCl}^+}\alpha_{\text{FeCl}^+} + k_{\text{FeSO}_4}\alpha_{\text{FeSO}_4} + k_{\text{FeH}_3\text{SiO}_4}\alpha_{\text{FeH}_3\text{SiO}_4} + \sum_j k_{\text{FeL}_j}\alpha_{\text{FeL}_j} \quad (5)$$

$$k_{\text{app}} = \sum_i \alpha_i k_i + \sum_j \alpha_j k_j \quad (6)$$

$$\alpha_{\text{Fe}^{2+}} = 1 / \left(1 + \sum K_{L_i} [L_i] + \sum K_{L_j} [L_j] \right) \quad (7)$$

Where $\alpha_{\text{Fe}^{2+}}$ is the fraction of free Fe(II) including inorganic and organic speciation, $[L_i]$ is the concentration of each inorganic ligand with a conditional stability constant K_i and $[L_j]$ is

485 the concentration of each organic ligand with a conditional stability constant K_{Lj} . The
486 inorganic site reaction coefficient at pH 8 and 25 °C was previously determined as 0.38
487 (Santana-Casiano et al., 2006). For any inorganic or organic FeL complexes:

$$488 \quad \alpha_{FeL} = K_{FeL}[L]\alpha_{Fe^{2+}} \quad (8)$$

489 According to our results, the presence of DOM in solution generated Fe(II)-organic
490 complexes that were overall less reactive to oxidation and therefore, when removed by UV
491 irradiation the resulting oxidation rate constants were higher (Fig. 8).

492 The amount of dissolved organic carbon (DOC) in the samples ranged between 50 and 79
493 $\mu\text{mol kg}^{-1}$ (data not shown) and according to O'Sullivan et al. (2005) the maximum amount
494 of H_2O_2 produced during irradiation could be 125 to 147 nmol kg^{-1} , respectively. Following
495 González-Davila et al. (2005) and assuming the extreme case of no H_2O_2 decay after 6 days,
496 the UV generated H_2O_2 could theoretically increase the Fe(II) oxidation rate constants in
497 oxygen saturated conditions. The theoretical $t_{1/2}$ associated with the H_2O_2 oxidation was
498 between 48 % and 53 % of the $t_{1/2}$ measured for 0.02 μm filtered samples. Nevertheless, the
499 measured $t_{1/2}$ for UV-irradiated samples (Supplementary Table S1 and Fig. 8) were shorter
500 (higher k'), indicating that any H_2O_2 remaining in solution after 6 days was not the main
501 factor that increased the oxidation rate constants and that the presence of dissolved organic
502 matter was stabilizing Fe(II) in solution. Furthermore, laboratory experiments have shown
503 that the $t_{1/2}$ of H_2O_2 in oceanic waters is shorter than 5.5 days (Yuan and Shiller, 2001),
504 therefore the impact of H_2O_2 in the experiments would be further reduced.

505 The role of organically complexed Fe(II) could be calculated from the following
506 assumptions. Firstly, the Fe(II)-L complex was assumed to not oxidize in the pH range of 7.5
507 to 8. Secondly, the observed reduction in Fe(II) oxidation rate constants (Fig. 8 and

508 Supplementary Table S1) at 25 °C between the 0.02 µm filtered and the filtered and UV
509 irradiated samples was due to organic complexation. The computed role of organic
510 complexation produced a $78 \pm 6\%$ decrease in the concentration of reactive inorganic Fe(II)
511 species. The effect of organic Fe(II) species was highest at Broken Spur and TAG sites ($81 \pm$
512 5%) and lowest at Rainbow ($74.1 \pm 4\%$). Roy et al. (2008) found a value that was slightly
513 lower for the western subarctic Pacific water, with a $61 \pm 8\%$ decrease. As both the organic
514 ligand concentrations and the conditional constants are unknown, only the product $K_{Lj} \cdot [L_j]$
515 (Eq. 7) can be inferred. Knowing the inorganic distribution coefficient of Fe(II), which is pH
516 dependent but can be calculated from Santana-Casiano et al., (2005), and considering a
517 unique class of ligand (L), the product $K_{Lj} \cdot [L_j]$ should be 9 ± 2 (higher values at Broken Spur
518 and TAG, while lower values at Rainbow) while $1 + \sum K_{L_i} [L_i]$ in Eq. 7 is 2.6 ($2.6 = 1/0.38$,
519 Santana-Casiano et al., (2006)).

520 If the concentration of Fe(II) organic ligands in seawater is on the same order as Fe(III)
521 ligands (1 to 10 nmol L⁻¹), as assumed by Roy et al. (2008), the corresponding conditional
522 constant should range between 10^9 to 10^{10} , (respectively for 10 and 1 nmol L⁻¹ of ligands).
523 The strength of these Fe(II) complexing ligands compared to those for Fe(III) is in the low
524 affinity range (Gledhill and Buck, 2012; Bundy et al., 2014). However, the formation of these
525 complexes would decrease the Fe(II) oxidation rate constant through both the effects on the
526 side reaction coefficients (Eqs. 7-8) and because the complexes can be non-reactive to
527 oxidation or may oxidize slower than other inorganic species. Moreover, it should be
528 considered that organic matter might even increase the Fe(II) $t_{1/2}$ due to the reduction of
529 organically complexed Fe(III), which would increase the Fe(II) in solution, and therefore the
530 $t_{1/2}$ (Santana-Casiano et al., 2010).

531 The UV-irradiation treatment (Fig. 8) removes the organic matter effect on the Fe-
532 complexation, which for the studied samples resulted in an increase in the oxidation rate
533 constants. Consequently, in these analyzed natural samples, organic matter ultimately
534 lowered the oxidation rate constants (black data points below the straight line in Fig. 9).

535 The DOM is prone to natural and thermal degradation in hydrothermal systems (Hawkes et
536 al., 2016). Accordingly, the variety of organic-complexing ligands and their effect on
537 oxidation could change with plume transport and distance to the vent site as observed in
538 figures 6 and 7. In the proximities to the vent sites, samples presented greater than theoretical
539 oxidation rate constants (above the straight line in Fig. 9). This could be due to the presence
540 of organic matter with ligands complexing Fe that oxidizes faster than the inorganic Fe(II)
541 species. At these locations, the organic ligand effect would dominate over processes that
542 lower oxidation rate constants such as nanoparticle concentrations. Similar experiments
543 considering organic matter degradation just above the vent site should be carried out to
544 confirm this hypothesis. Nevertheless, the complexity of the organic matter pool and the
545 extent and nature of Fe-DOM interactions make the resulting effect on the oxidation rate
546 constants difficult to predict.

547 ***4.3. Effects of physical speciation***

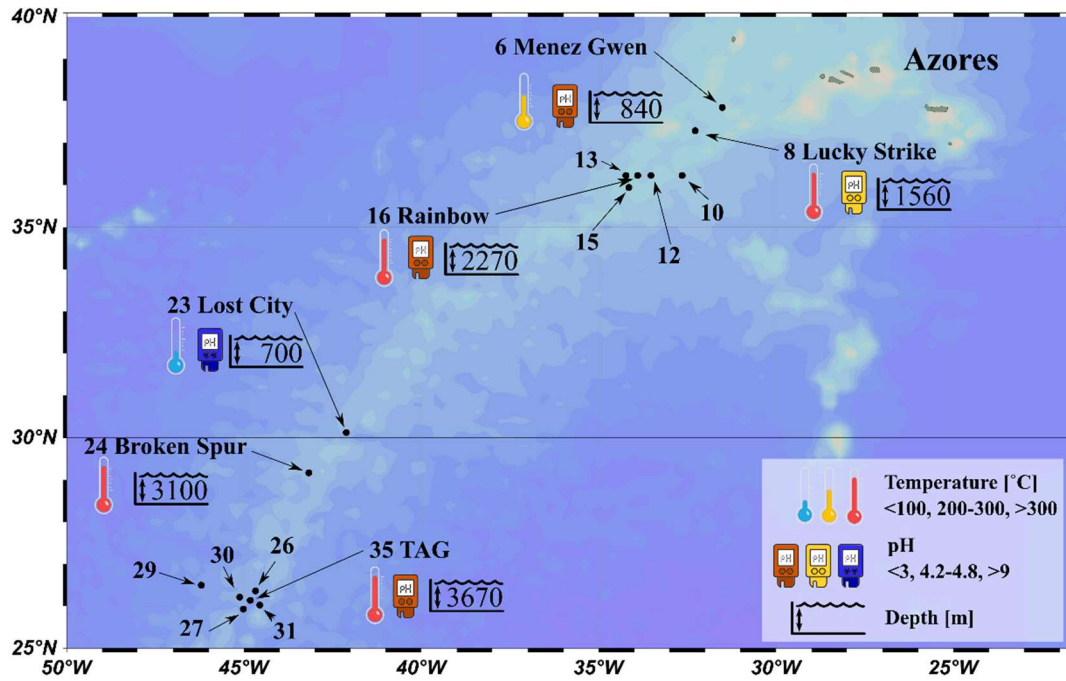
548 Lough et al. (2019) have shown that there is an Fe exchange between the different particle
549 size fractions within the hydrothermal plume. This exchange could affect Fe(II) oxidation as
550 found in this study for particles smaller than 0.2 μm and colloidal sized particles (0.2-0.02
551 μm) at different locations in TAG and in Rainbow (Fig. 8). The results shown in Figure 8
552 seem to indicate that the size-fractionation can significantly affect Fe(II) oxidation and thus,
553 is a factor that should be considered in future studies. In this sense, previous work has shown
554 that particulate Fe(II), such as pyrite nanoparticles are more resistant to oxidation, with Fe(II)

555 half-life between 4 and 48 months at 2 °C (Yücel et al., 2011). The presence of colloidal
556 particles also enhances the decay of ROS allowing for longer $t_{1/2}$ (Yuan and Shiller, 2001).
557 Therefore, nanoparticles could account for a substantial fraction of the decrease in the
558 oxidation rate constants (Fig. 8). Another consequence of filtration is the reduction in larger
559 size organic matter, which could also affect the stabilization of Fe(II) (Buck et al., 2015).

560 The variability observed in the oxidation rate constants inside the plume for the six studied
561 hydrothermal vents indicates that changes in the composition of both the DOM and colloidal
562 particles, both between sites and at different distances from the vent, can strongly affect the
563 oxidation rate constants, thus, their role requires further study.

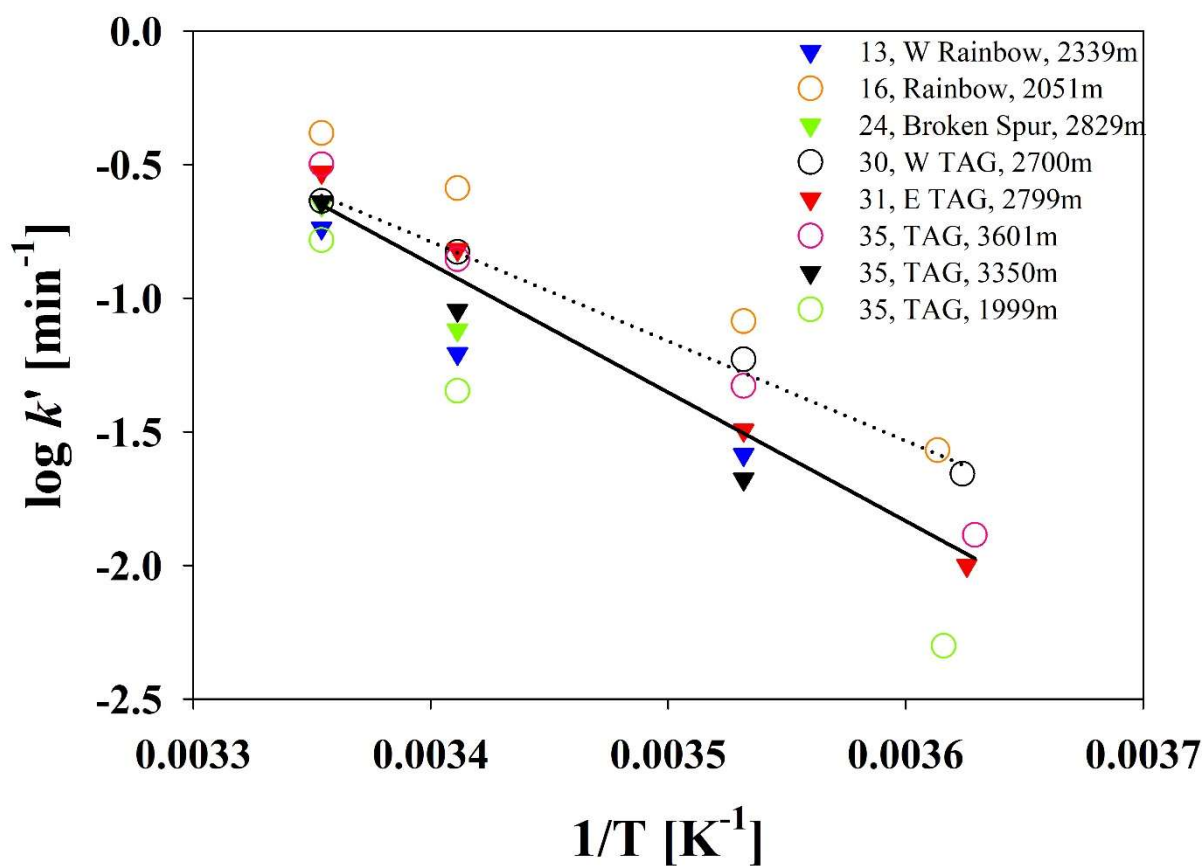
564 **5 Conclusions**

565 This study highlights the complexity of Fe(II) chemistry in the ocean, with a focus on
566 hydrothermal vent settings. Fe(II) oxidation rate constants, and therefore persistence of Fe(II)
567 in the environment, is shown to not only depend on the physico-chemical conditions (T, S,
568 pH, O₂ content) but also the physical state (soluble, colloidal and particulate) of the inorganic
569 and organic species that are present. Our results indicate that organic ligands can stabilize
570 Fe(II) and that nanoparticles present in hydrothermal plumes could increase the resistance to
571 oxidation. Both processes contribute to the observed variability in the oxidation rate constants
572 in the proximities of vents. We were able to revise the theoretical equation for the estimation
573 of k' , and to further increase the existing temperature range to include seawater temperatures
574 as low as 2 °C. Future studies on iron oxidation kinetics in the marine environment should be
575 accompanied by a characterization of the organic matter present in the medium, a size-
576 fractionation of the samples, and a detailed description of the conventional physico-chemical
577 factors (pH, T, and S).



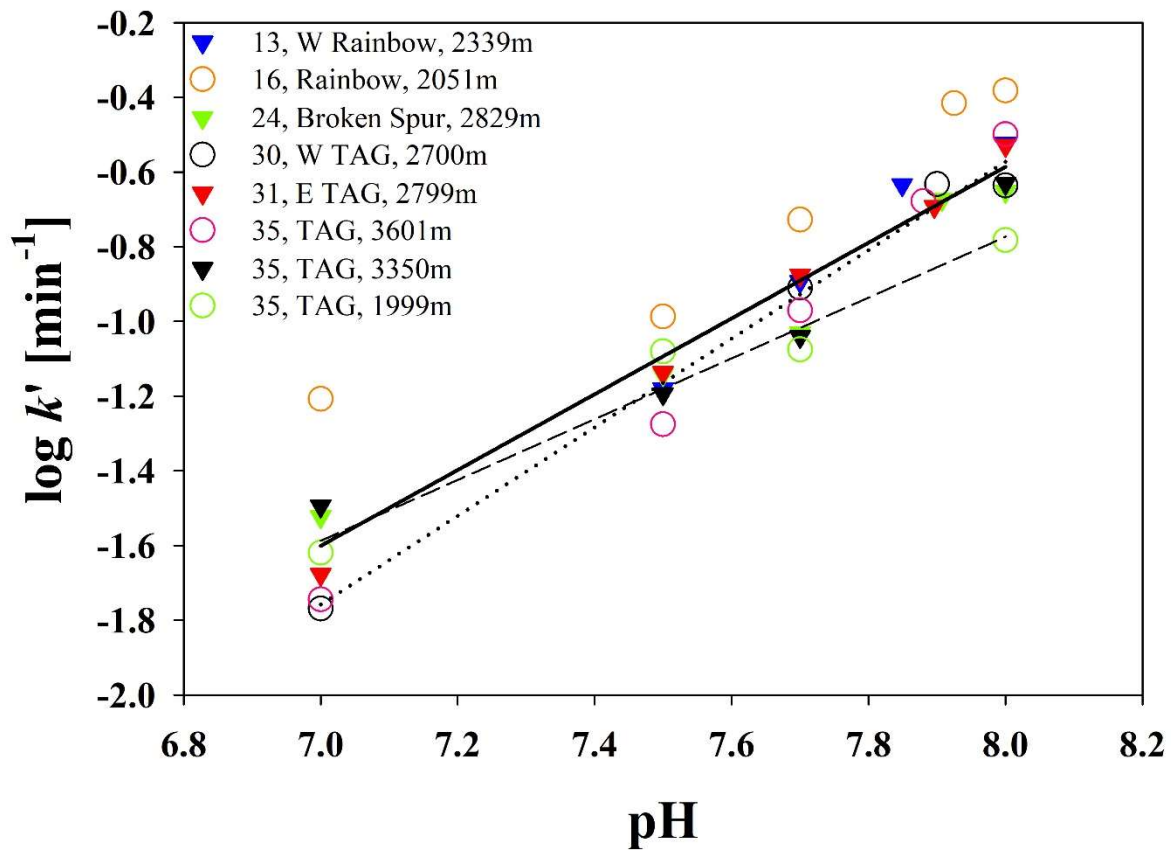
579

580 Figure 1. Map of the investigated hydrothermal vent sites along the MAR, showing the
 581 station numbers and hydrothermal vent site names for the sampling sites. Symbols represent
 582 endmember characteristics. The pH of the Broken Spur endmember was not available.



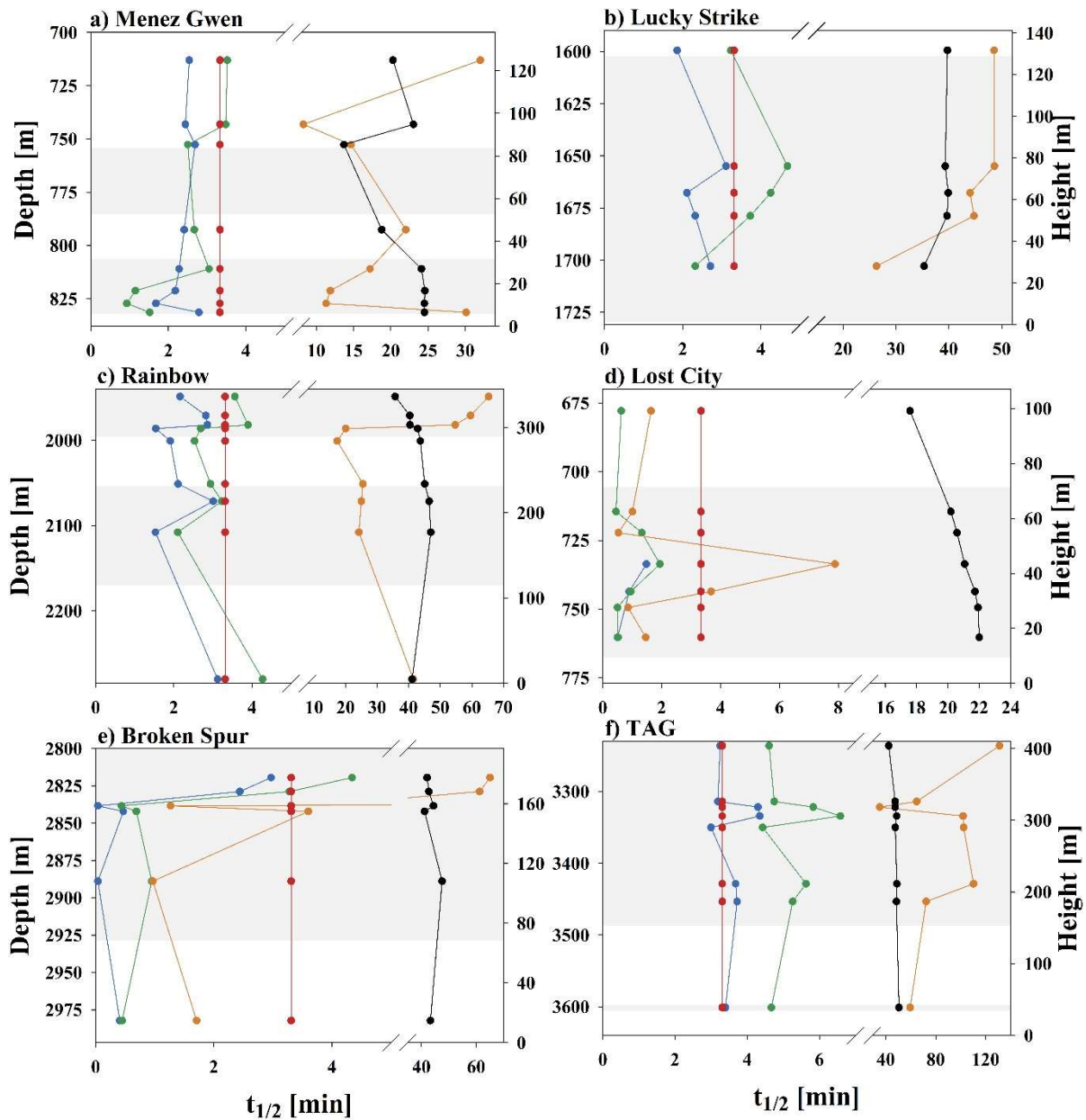
583

584 Figure 2. Arrhenius relationship between the oxidation rate constant ($\log k'$ (min^{-1})) and
 585 temperature ($1/T$ (K^{-1})) in the range 2 to 25 °C for eight selected samples. Note that the
 586 sample from station 30 (W TAG, 2700m, open squares, dotted line) shows a different
 587 behavior than samples from other stations.



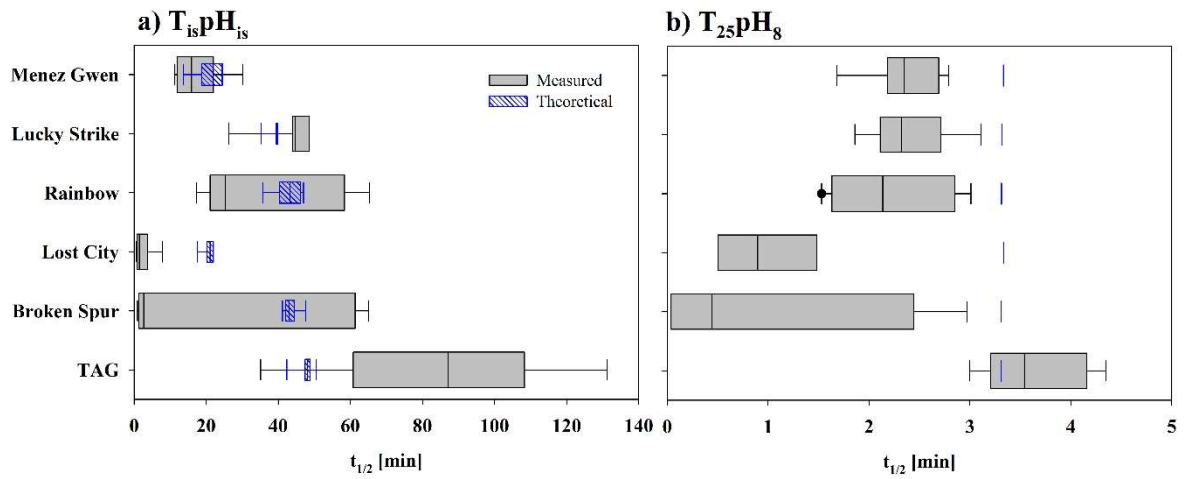
588

589 Figure 3. pH effect (from 7.0 to 8.0) on the Fe(II) oxidation rate constants ($\log k'$ (min^{-1})) for
 590 eight selected samples collected along the MAR. Note that the samples from 2700 m depth to
 591 the west of TAG (Stn 30, open squares) and from 1999 m depth at the TAG site (Stn 35, open
 592 diamonds) have different slopes than samples from other stations (data from these two
 593 stations are plotted with dotted lines).



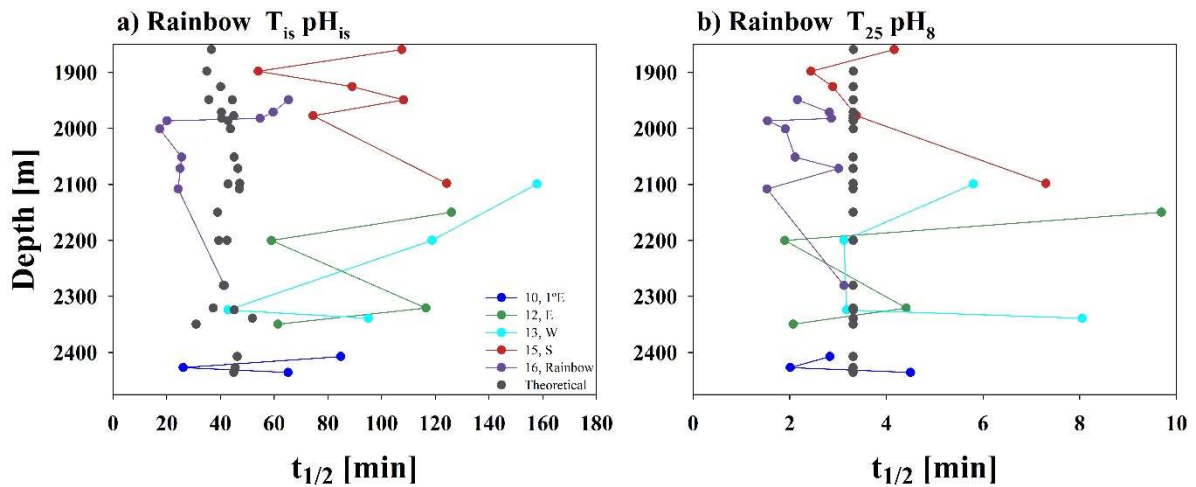
594

595 Figure 4. The measured Fe(II) half-life ($t_{1/2}$, min) at $T_{is}pH_{is}$ (orange), $T_{25}pH_{is}$ (green), and
 596 $T_{25}pH_8$ (blue), and the theoretical (Theo) values for $T_{is}pH_{is}$ (black), and $T_{25}pH_8$ (red), at the
 597 six vents. The shaded area shows the anomaly depths determined using light scattering data .
 598 Both, the depth (left y-axis) and the height above the seafloor (right y-axis) of the sampling
 599 are indicated.



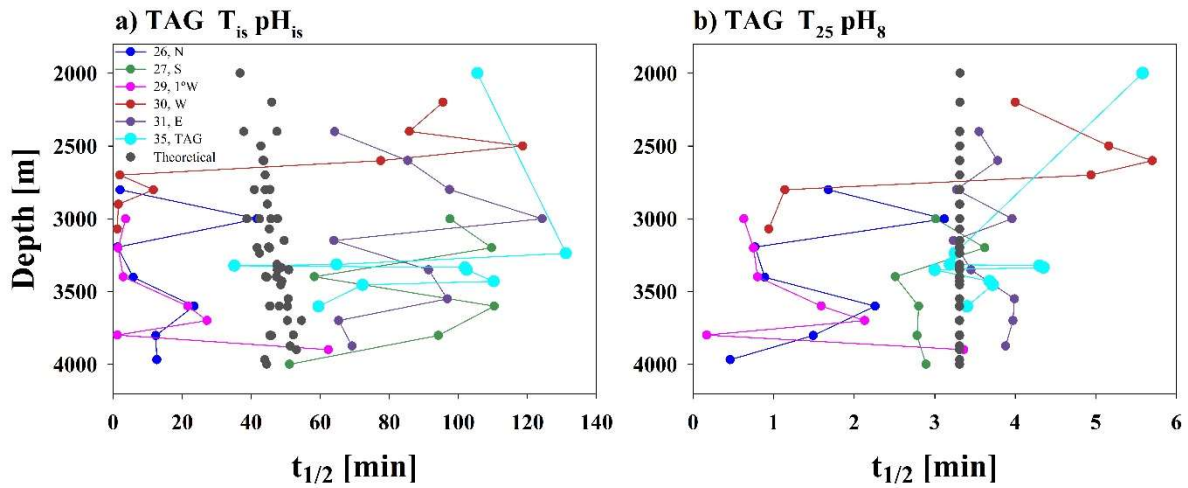
600

601 Figure 5. Boxplots of the Fe(II) half-life ($t_{1/2}$, min) within the six hydrothermal plumes. a)
 602 The measured $t_{1/2}$ at *in situ* conditions (grey) and the theoretical $t_{1/2}$ (blue; obtained using Eq.
 603 4). b) The measured $t_{1/2}$ at constant temperature (25 °C) and pH 8 (grey) and the
 604 corresponding theoretical $t_{1/2}$, the slight variances are caused by S changes (blue).



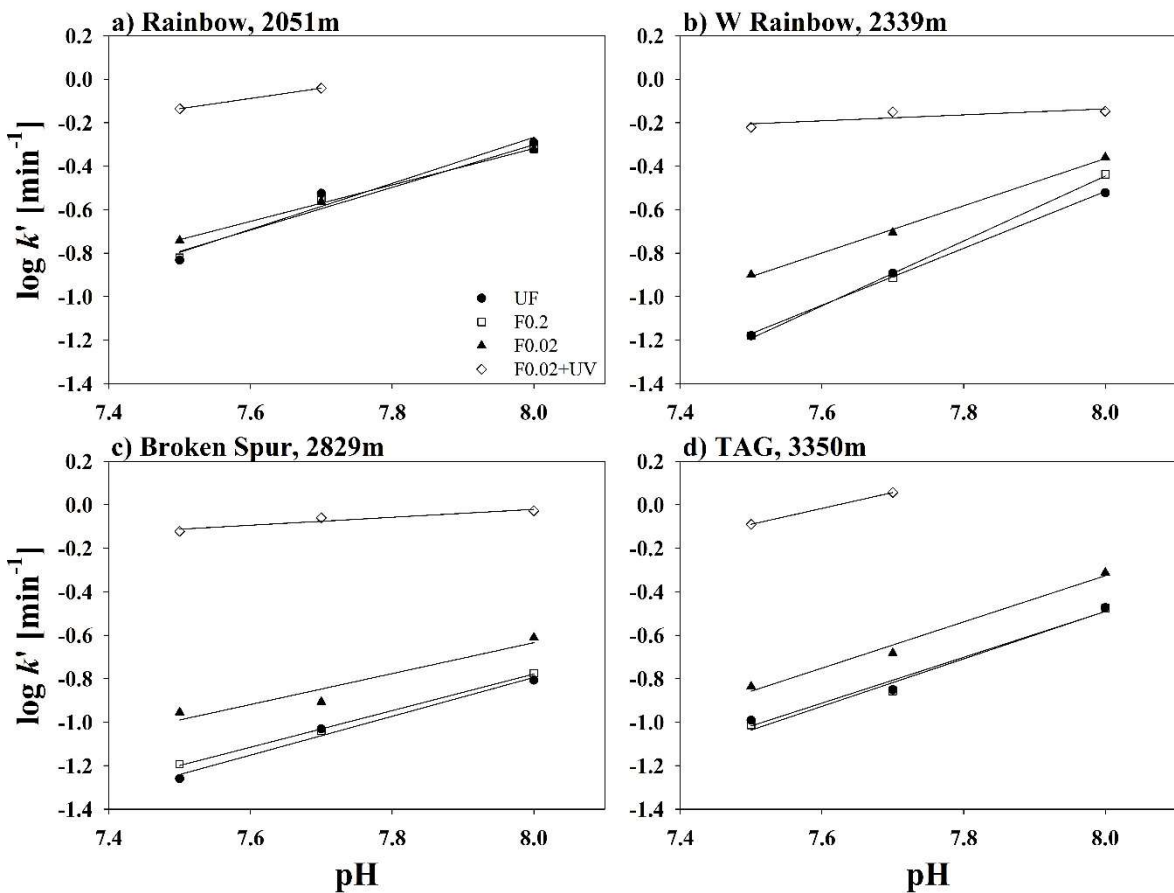
605

606 Figure 6. The Fe(II) half-life ($t_{1/2}$, min) at the Rainbow field a) under *in situ* conditions (see
 607 also Supplementary Table S1) and b) normalized to constant T of 25 °C and a pH of 8.



608

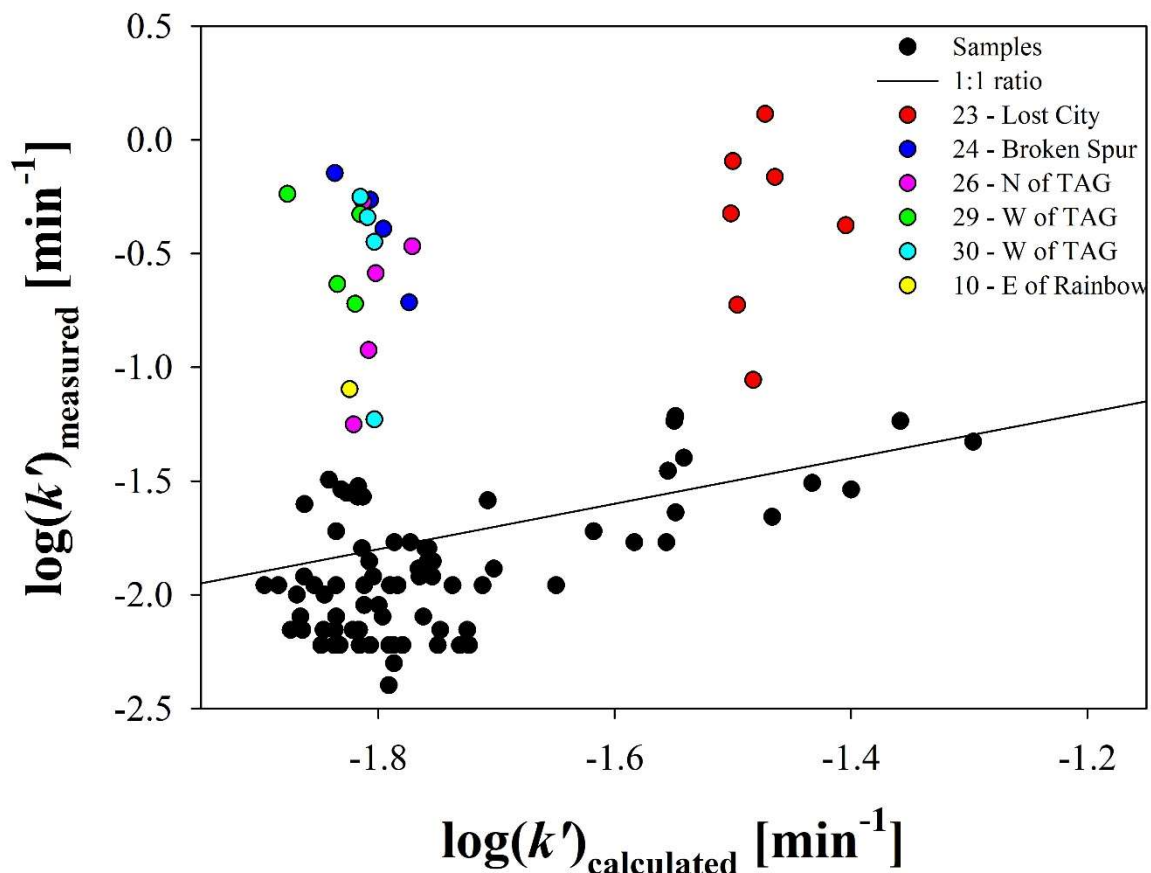
609 Figure 7. The Fe(II) half-life ($t_{1/2}$, min) at the TAG field a) under *in situ* conditions (see also
 610 Supplementary Table S1) and b) normalized to constant T of 25 °C and a pH of 8.



611

612 Figure 8. Oxidation rate constants at different pH (7.5, 7.7 and 8), for selected unfiltered (UF,
 613 filled circles), filtered through 0.2 μm (F0.2, open squares), filtered through 0.02 μm (F0.02,

614 filled triangles) and filtered through 0.02 μm followed by UV irradiation (F0.02+UV, open
 615 diamonds) samples at a) Rainbow (Stn 16, 2051 m), b) West of Rainbow (Stn 13, 2339 m), c)
 616 Broken Spur (Stn 24, 2829 m) and d) TAG (Stn 35, 3350 m).



618 Figure 9. Fe(II) oxidation rate constants ($\log(k')$) determined under *in situ* conditions against
 619 the anticipated expected theoretical value. All data points are shown in black. The black line
 620 represents a 1:1 ratio between both oxidation rate constants. Color coded data points
 621 correspond to samples with measured oxidation rate constants greater than 20% of the
 622 theoretically derived rate constants.

623 **Author contributions**

624 DGS wrote the manuscript with significant contributions from all coauthors. DGS was
625 invited to the research cruise by MCL and AT, where DGS, MCL, and LA collected the
626 samples. DGS analyzed the samples with support from MSC and MGD. Figures for the
627 results were designed by HP and GS. Fieldwork logistics and costs were overseen by MCL
628 and AT, while analysis costs were handled by MSC, MGD, GS, and HP.

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637 **Competing interests**

638 The authors declare no competing interests.

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