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1 Nickel isotopes and rare earth elements systematics in marine
2 hydrogenetic and hydrothermal ferromanganese deposits

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5 Bleuenn Gueguen^{1,2*}, Olivier Rouxel³, and Yves Fouquet³

6

7 ¹CNRS, Univ Brest, UMS 3113, F-29280 Plouzané, France

8

9 ²CNRS, Univ Brest, UMR 6538 Laboratoire Géosciences Océan, F-29280 Plouzané, France

10

11 ³IFREMER, Centre de Brest, Unité Géosciences Marines, F-29280 Plouzané, France

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**Corresponding author:*

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bleuenn.gueguen@univ-brest.fr (B. Gueguen)

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38 **Abstract**

39 Attention is now being given to Ni isotope systematics in hydrogenetic marine
40 ferromanganese (Fe-Mn) crusts as paleoceanographic proxies. Previous work focused on
41 identifying both mineralogy (post-depositional) and source effects (Gall et al., 2013; Gueguen
42 et al., 2016), in particular regarding hydrothermal inputs in the oceans and the response of Ni
43 isotope biogeochemical cycling through time. The most important sink for Ni in the oceans is
44 the Fe-Mn oxides sink, but estimation of its Ni isotope composition is only based on
45 hydrogenetic Fe-Mn crusts. In this study, we investigated a range of Fe-Mn deposits including
46 Fe-Mn deposits variably affected by hydrothermal inputs, including hydrothermal deposits
47 from the Lau back-arc basin (South West Pacific) and Lo’ihi seamount (Hawaii),
48 hydrogenetic crust and nodules from the Bauer Basin (Pacific Ocean). Nickel isotope ratios
49 were measured by multi-collector inductively coupled plasma mass spectrometer (MC-ICP-
50 MS) using a double-spike (^{61}Ni and ^{62}Ni) correction method. The combination of Ni isotopes
51 and rare earth element (REE) geochemistry show that Ni isotope fractionation in Fe-Mn
52 deposits is essentially controlled by formation processes of the deposits (such as the rate of
53 formation, the initial Mn-phase and sorption processes) which are also related to the
54 depositional environment. Consistent with previous studies, pure hydrogenetic crusts are
55 characterized by isotopically heavy Ni isotope signatures ($\delta^{60/58}\text{Ni}$ values range from ‰ 0.9
56 and 2.5 ‰) and well-developed positive Ce anomalies. In contrast, mixed hydrothermal-
57 hydrogenetic crust and nodules from the Bauer Basin (East Pacific) display negative Ce
58 anomaly and lighter $\delta^{60/58}\text{Ni}$ values (0.3 ‰ to 0.4 ‰), which are interpreted as the result of
59 far-field hydrothermal inputs of Fe-Mn precipitates from the East Pacific Rise.
60 Nickel in hydrothermal deposits from the Lau Basin (0.5 and 1.1 ‰) and Lo’ihi seamount (-
61 0.8 to -1.5 ‰) is isotopically lighter than in hydrogenetic Fe-Mn crusts. Light $\delta^{60/58}\text{Ni}$ values
62 in Lo’ihi deposits is due to the removal of Ni during Ni adsorption from seawater and from

63 the hydrothermal fluid (between 0-1.4 ‰) on Fe-oxides followed by isotope fractionation
64 between the fluid and the mineral phase. Results suggest that Ni isotopes in hydrothermal Fe-
65 rich deposits are strongly fractionated relative to the seawater/fluid source due to partial
66 removal of Ni on Fe-phases. Hydrothermal Mn-oxides deposits from the Lau Basin acquired
67 their Ni isotope signature through Ni adsorption and continuous exchange of Ni with
68 seawater. We propose that the systematic difference in Ni isotope signatures between
69 hydrogeneous and hydrothermal Fe-Mn deposits is related to the mechanisms of Ni uptake
70 into oxide minerals (e.g., birnessite vs. todorokite; Fe-oxides vs. Mn-oxides) which depend on
71 the rate of formation and the source of Mn and Fe to marine ferromanganese deposits (i.e.,
72 depositional environment) rather than Ni sources.

73 **1. Introduction**

74 Understanding the biogeochemical cycling of trace metals in the marine environment
75 is important because they are involved in key biological reactions such that variations of
76 metal distribution in time and space in the oceans could have significant impact on the
77 biological pump (Price and Morel, 1991; Saito et al., 2003; Morel et al., 2014). Recent
78 advances on hydrothermal research and metal distribution in deep water have highlighted the
79 importance of hydrothermal venting on, for example, the iron marine geochemical budget
80 (Toner et al., 2009; Tagliabue et al., 2010; Saito et al., 2013; Conway and John, 2014;
81 Fitzsimmons et al., 2014; German and Seyfried, 2014; Tagliabue et al., 2014; Horner et al.,
82 2015; Rouxel et al., 2016). Other transition metals like zinc, nickel, cadmium and copper may
83 also be impacted by deep hydrothermal venting to various extents (e.g., John et al., 2008; Gall
84 et al., 2013; Hannington, 2013; Little et al., 2014). Growing interest is now given to nickel
85 (Ni) isotope systematics in Fe-Mn crusts as a potential tracer of the evolution of the
86 biogeochemistry of trace metals in oceans through time (Gall et al., 2013; Cameron and
87 Vance, 2014; Gueguen et al., 2016).

88 Marine ferromanganese (Fe-Mn) oxide deposits are ubiquitous in the oceans, and these
89 chemically precipitated sediments represent an important sedimentary archive of trace metal
90 seawater composition. Ferromanganese oxide deposits show a range of metal enrichment (Ni,
91 Cu, Co) based on their geological settings, formation rates and sources of dissolved Fe, and
92 Mn. They are commonly classified using a ternary diagram composed of iron (Fe), manganese
93 (Mn) and (nickel (Ni) + copper (Cu) + cobalt (Co)) concentrations as the three variables in the
94 diagram (Bonatti et al., 1972). This representation allows the discrimination between the
95 hydrothermal, hydrogenetic and diagenetic fields (Bonatti et al., 1972) indicating different
96 geological settings, i.e., hydrothermal environment, strictly precipitated from seawater, and
97 dominated by diagenetic processes. Hydrogenetic ferromanganese crusts are precipitated on

98 volcanic seamounts at depths between 400 and 4000 m at very slow rates of 1-6 mm/Ma
99 (Hein et al., 2003; Hein and Koschinsky, 2014). They are generally considered to be good
100 proxies of deep seawater metal isotope composition, providing that fractionation processes
101 during metal uptake are sufficiently well constrained. Accordingly, Fe-Mn crusts have
102 attracted considerable interest for investigating seawater composition over a time-scale of
103 several million years (O'Nions et al., 1998; Frank et al., 1999; Reynolds et al., 1999; Frank,
104 2002; Frank et al., 2002; van de Flierdt et al., 2004; Klemm et al., 2005; Klemm et al., 2007;
105 Nielsen et al., 2011).

106 Initial Ni isotope composition of hydrogenetic Fe-Mn crusts reported by Gall et al.
107 (2013) and Gueguen et al. (2016) ranges from 0.9 to 2.5 ‰ for a period of more than 60 Ma,
108 suggesting that deep seawater Ni isotope composition could have changed in the past. Both
109 studies investigated the Ni isotope composition of Fe-Mn crusts from the Pacific Ocean by
110 sub-sampling Fe-Mn crusts layers through stratigraphic depth in order to evaluate the
111 variability of Ni isotopes in seawater through time. Gall et al. (2013) suggested that Ni
112 isotopes in seawater are sensitive to variations of Ni sources such that Ni isotopes in Fe-Mn
113 crusts formed close to continental margins are influenced by continental weathering, and that
114 hydrothermal inputs could also affect the Ni isotope composition of oceanic basins. It is now
115 well established that dissolved Ni in seawater is isotopically heavier ($\sim 1.44 \pm 0.15$ ‰;
116 Cameron and Vance, 2014; Wang et al., 2019; Archer et al., 2020) compared to the average
117 $\delta^{60/58}\text{Ni}$ crustal value (BSE, Bulk Silicate Earth) estimated to about 0.12 ± 0.01 ‰ (Klaver et
118 al., 2020). Nickel in the oceans is mostly coming from the riverine input and has an average
119 $\delta^{60/58}\text{Ni}$ value of 0.8 ‰ (Cameron and Vance, 2014), which is heavier than the average crustal
120 value of 0.12 ± 0.01 ‰ (Klaver et al., 2020). The affinity of Ni towards Mn oxides has been
121 well recognized (e.g.; Koschinsky and Halbach, 1995; Koschinsky and Hein, 2003; Peacock
122 and Sherman, 2007a; Peacock and Sherman, 2007b) and provide the main sink of Ni in

123 seawater. However, an estimate of the global Ni isotope composition of Fe-Mn deposits as a
124 proxy of the Mn sink yield $\delta^{60/58}\text{Ni}$ values in the range of oceanic values and even slightly
125 heavier (Gall et al., 2013; Gueguen et al., 2016) providing no leverage to affect Ni isotope
126 composition of seawater. As recently discussed by Archer et al. (2020) and (Little et al.,
127 2020), the isotopic and elemental budget of Ni in the oceans is imbalanced (Cameron and
128 Vance, 2014; Vance et al., 2016; Ciscato et al., 2018). If the ocean is at steady-state, three
129 possibilities can explain this imbalance, (1) there is a missing isotopically heavy Ni source,
130 (2) a light Ni sink is required, or (3) Fe-Mn crusts do not provide an accurate estimate of the
131 marine Mn sink for Ni isotopes. A possible sink for light Ni isotopes in the ocean could be
132 sediments rich in organic matter such as sediments deposited in continental margins and
133 anoxic basins (Takano et al., 2017; Ciscato et al., 2018; Archer et al., 2020). However, the
134 extent of Ni isotope fractionation is rather small (between 0 to 0.3 ‰), and the size of such
135 reservoir is not sufficient to exert a significant control on oceanic Ni isotope budget. Based on
136 results from Ni isotopes in metalliferous sediments from MANOP site in the East Pacific,
137 Little et al. (2020) estimated that an isotopic Ni benthic flux extremely heavy of ~3 ‰ would
138 balance the marine Ni budget leaving a residual light Ni reservoir in sedimentary authigenic
139 phases.

140 Although some significant efforts have been undertaken for measuring Ni isotope
141 composition of major oceanic Ni sources (Cameron and Vance, 2014; Vance et al., 2016;
142 Ciscato et al., 2018; Little et al., 2020) and mechanisms of isotope fractionation processes
143 (Wasylenki et al., 2015; Gueguen et al., 2018; Sorensen et al., 2020), the Ni isotope
144 composition of the oceanic Ni sink associated with Mn oxides needs to be better understood,
145 and it is yet unclear whether secular Ni isotope variations in marine Fe-Mn deposits may
146 result from source effects (e.g., circulation of water masses, local inputs (diagenetic vs.
147 hydrothermal sources)), fractionation processes during Ni incorporation in the deposits, or a

148 combination of both. More particularly, the effect of hydrothermal inputs on the Ni sink in Fe-
149 Mn deposits has to be better understood.

150 The purpose of this work is to investigate in a range of mixed hydrothermal-
151 hydrogenetic Fe-Mn deposits the Ni isotope composition of the Mn sink by evaluating, (1) the
152 processes controlling the Ni isotope composition in marine Fe-Mn deposits (e.g., depositional
153 environment, formation rates, mineralogy, source effects), and (2) the Ni isotope signature of
154 hydrothermal Fe-Mn deposits as a proxy of the impact of hydrothermal inputs on the Ni
155 biogeochemical cycling in the marine environment. To help our understanding of Ni isotopes
156 in Fe-Mn deposits, we will combine Ni isotopes and rare earth elements geochemistry. Rare
157 earth elements provide another source of information regarding metal sources and deposition
158 rates of seafloor Fe-Mn deposits. They behave coherently in natural environments with the
159 exception of Ce (and Eu), and relative REEs abundances are not fractionated from their
160 source during scavenging and precipitation processes. The contrasting behavior of Ce
161 compared to other REEs is due to the oxidation of soluble Ce(III) to insoluble Ce(IV) at the
162 oxide surface after its sorption to the mineral surface implying that the oxidized Ce(IV) no
163 longer exchanges with seawater (e.g.; Bau et al., 2014). Accordingly, the slow growth rates of
164 Fe-Mn crusts allows the accumulation of more Ce in comparison to other REEs (Bau, 1999;
165 Bau and Koschinsky, 2009; Bau et al., 2014), and they display a positive Ce anomaly.
166 Because the rate of precipitation of hydrothermal Fe-Mn oxide deposits is rapid,
167 hydrothermally precipitated REEs preserve the negative Ce anomaly characteristic of
168 seawater (De Baar et al., 1985; Elderfield et al., 1988; Kuhn et al., 1998; Mills et al., 2001).
169 Other studies also suggest that Ce oxidation could occur through similar pathways as
170 microbial Mn oxidation in the marine environment (Moffett, 1990; Moffett, 1994). Previous
171 studies have shown a trend between the magnitude of the Ce anomaly and the distance of Fe-
172 Mn deposits from hydrothermal sources, with samples located further away from the

173 hydrothermal source and precipitated more slowly displaying higher positive Ce anomalies
174 than samples formed close to the vents (Kuhn et al., 1998; Mills et al., 2001; Bau et al., 2014).
175 Excess of Ce could ultimately be used as a proxy for the deposition rate and seawater
176 exposure age of hydrothermal Fe-Mn deposits (Bau et al., 2014). Yttrium can also be used for
177 distinguishing between hydrothermal, hydrogenetic and diagenetic Fe-Mn deposits. Y_N/Ho_N
178 ratios (normalized to PAAS) < 1 are the result of preferential scavenging of Ho compared to
179 Y on Fe-Mn oxyhydroxide surfaces which produces a negative Y anomaly in hydrogenetic
180 Fe-Mn deposits (Bau et al., 2014). Finally, the Sm_N/Yb_N ratio shows the slope of the REE
181 pattern. Unless there is a change in the REE sources in seawater, the slope of the REE pattern
182 should not change. Therefore, one can use this ratio to identify a change in the REE sources in
183 seawater.

184 Our sample set forms a continuum from actively forming Fe-Mn deposits formed
185 through diffuse venting (Lau Basin, Lo'ihi seamount), and which have seen the influence
186 from near-field hydrothermal inputs (deposits near the hydrothermal site); hydrogenetic Fe-
187 Mn deposits formed on ridge flank and having recognized hydrothermal contribution (Bauer
188 Basin), that we also define as far-field hydrothermal contribution (deposits located hundreds
189 of kilometers from the vent source); and pure hydrogenetic Fe-Mn deposits from different
190 oceanic regions.

191

192 **2. Geological setting and sample description**

193

194 *2.1. Bauer Basin Fe-Mn deposits, Southeastern Pacific*

195

196 One hydrogenetic Fe-Mn crust sample and two diagenetic nodules from the Bauer
197 Basin (BB) collected during the YALOC-73 cruise on the R/V Yaquina (Oregon State

198 University) at latitude 13°40'S and longitude 102°8'W (Lyle et al., 1977; Elderfield and
199 Greaves, 1981) were selected (supplementary table S1). Bauer Basin is located in the South
200 Pacific Ocean between the East Pacific Rise and Galapagos Rise (Heath and Dymond, 1977)
201 and is therefore surrounded by active oceanic ridges. The two nodules were affected by
202 diagenetic processes (Elderfield and Greaves, 1981) and the Fe-Mn crust was formed by
203 hydrogenetic processes (Lyle et al., 1977). Previous studies have demonstrated that Fe-Mn
204 oxide deposits including Fe-Mn crusts from the BB are prone to significant influence from
205 hydrothermal inputs exported by seawater circulation through the ridge crests (Dymond and
206 Veeh, 1975; Heath and Dymond, 1977; Lyle et al., 1977; Elderfield and Greaves, 1981). High
207 metal/aluminium ratios found in sediments from the Bauer Basin and the negative Ce
208 anomaly observed in Bauer Basin nodules is interpreted as being related to the transport of
209 hydrothermal particles to the Bauer Basin (e.g., Fe-oxyhydroxides phases for rare earth
210 elements) from hydrothermal sites located on the East Pacific Rise (EPR).

211

212 *2.2. Hydrothermal Fe-Mn oxide deposits from the Lau Basin, Southwestern Pacific*

213

214 Fe-Mn oxide deposits samples were collected in the southern back-arc Lau Basin (LB)
215 during the NAUTILAU cruise on the R/V Nadir (Ifremer-Genavir) in April-May 1989 using
216 the submersible Nautile (Figure 1 and supplementary table S1). The Valu Fa Ridge in the Lau
217 Basin is an active back-arc spreading center where diffuse hydrothermal venting allows the
218 deposition of Fe- and Mn-rich deposits (Fouquet et al., 1991). Fe-Mn-rich deposits were
219 observed in both inactive and active hydrothermal vent fields including the Hine Hina site on
220 the Southern Valu Fa Ridge, Vai Lili on the Central Valu Fa Ridge, and White Church on the
221 Northern Valu Fa segment (Fouquet et al., 1993). Two samples were selected for our study:
222 (1) NL-20-05, a 5 cm thick manganese crust collected at 1650 m water depth at the Vai Lili

223 site (Central Valu Fa Ridge), (2) NL-10-09, collected at 1908 m water depth at the White
224 Church site (Northern Valu Fa Ridge) and they consist of pieces of small inactive chimneys
225 composed of porous Mn-oxides (Fouquet et al., 1993). Hydrothermal fluids in this area have
226 higher metal concentrations and are notably enriched in Zn, As, Pb, Cd, Mn compared to mid-
227 ocean ridge vent fluids (e.g.; Fouquet et al., 1993; Mottl et al., 2011).

228

229 *2.3. Fe- and Mn-rich microbial mats at Lo'ihi seamount, Northern Pacific*

230

231 Hydrothermal Fe oxide deposits variably enriched in Mn were collected at both the
232 summit and the base of Lo'ihi seamount during three oceanographic cruises: FeMO 2006 on
233 R/V Melville (Scripps Institution of Oceanography) in October-November 2006, FeMO 2008
234 on R/V Thompson (University of Washington) in September-October 2008 and FeMO 2009
235 on the R/V Kilo Moana (University of Hawaii) in October 2009 using ROV Jason II (Woods
236 Hole Oceanographic Institution) (Figure 1). Details of sample collection can be found in
237 Glazer and Rouxel (2009), Edwards et al. (2011), (Rouxel et al., 2018).

238 Continuous diffuse venting of low temperature (0-60°C) hydrothermal fluids enriched
239 in Fe and depleted in H₂S in comparison to mid-ocean ridge hydrothermal sources (Sedwick
240 et al., 1992; Wheat et al., 2000) allows thriving of abundant Fe-oxidizing bacteria that formed
241 massive biogenic Fe-rich mats observed at both the summit and base of Lo'ihi seamount
242 (Emerson and Moyer, 2002; Glazer and Rouxel, 2009; Edwards et al., 2011; Toner et al.,
243 2012; Rouxel et al., 2018). In particular, the ultra-diffuse hydrothermal venting at Ula Nui site
244 at the base of Lo'ihi seamount (~5000 m depth) produce extensive Fe-rich deposits (referred
245 as FeMO Deep mats) which are overlaid by a thin Fe-Mn crust (Edwards et al., 2011). Lo'ihi
246 Fe-Mn oxide deposits are considered to be actively forming and represent an ideal example of
247 low-temperature hydrothermal deposits.

248

249 *2.5. Atlantic and Pacific hydrogenetic Fe-Mn crusts*

250

251 Bulk hydrogenetic Fe-Mn crusts from the Pacific Ocean were collected on Apuupuu
252 seamount ~50 km south of Hawaii during FeMO 2009 cruise on the R/V Kilo Moana in
253 October 2009 (supplementary table S1). These crusts were collected by ROV Jason II, and
254 two of these Fe-Mn crusts were already investigated in Gueguen et al. (2016). Eight samples
255 were selected for our study between 2000 and 3000 m depth. Fe-Mn crusts from the Atlantic
256 Ocean were collected on the Ascension Fracture Zone (FZ), Vema FZ, Gloria FZ and the
257 Azores Triple Junction (TJ) by dredging during two CNEXO cruises on the R/V Jean Charcot
258 (cruises GIBRACO CH30 in August-September 1972 and BIOVEMA CH78 in October
259 1977) (Table 1 and Figure 1). Some subsamples of those crusts were prepared at the
260 University of Cambridge (H. Elderfield and M. Greaves, pers. comm.).

261 Continuous discrete layers for each crust were not specifically sampled, but instead,
262 we targeted Fe-Mn crusts from different oceanic regions of the globe. For each Fe-Mn crust
263 sample we selected a subsample that was generally collected in depth of the crust, and does
264 not correspond to surface scrapings. Instead, they correspond to relatively large sections, i.e.
265 several millimeters to centimeters, of the crust. Accordingly, we refer our samples as “bulk
266 crusts.” They probably span a longer period of time than samples collected from discrete
267 layers, and thus, each sample represents a range of ages, rather than a single point in time. No
268 chronological information is expected to be obtained from this sample set. The subset of
269 samples from Apuupuu seamount were previously dated and show a maximum age of 17.0
270 Ma with an average growth rate of 1.6 mm/Ma (Gueguen et al., 2016).

271

272 **3. Methods**

273 *3.1. Mineralogy*

274

275 The mineralogy of Fe-Mn deposits was determined by X-ray diffraction (XRD)
276 Brucker AXS D8 Advance and Brucker AXS D2 Phaser at Ifremer (Brest, France) on bulk
277 powder for hydrogenetic Fe-Mn crusts, Bauer Basin Fe-Mn deposits and Lau Basin Fe-Mn
278 deposits. Dried powdered samples were deposited on a XRD sample holder and flattened with
279 a glass slide. Samples were analyzed using Cu-K α radiation over 2 θ ranging from 2° to 70° at
280 40 kV and 30 mA. The mineralogy was assessed using Eva search-match software.

281

282 *3.2. Major and trace elements composition*

283

284 Major, trace and rare earth element (REE) concentrations were measured by ICP-MS-
285 Quadrupole (X-Series 2, Thermofisher Scientific) at the PSO (Pôle de Spectrométrie Océan),
286 with the exception of Lo'ihī samples which were measured by ActLabs (Activation Labs,
287 Canada) (Edwards et al., 2011). Hydrogenetic Fe-Mn crusts and Lau Basin samples were
288 dissolved in HNO₃-HCl acid mixture and nodules and Lo'ihī Fe-rich deposits in an HNO₃-
289 HCl-HF acid mixture. After evaporation residues were dissolved in 6M HCl as stock
290 solutions. A fraction of the solution was diluted for ICP-MS analysis, and geological
291 reference materials (GRMs) of Mn-nodules Nod-A-1 and Nod-P-1 were systematically run
292 during each analytical session altogether with the samples. Concentrations were calibrated
293 using a multi-elemental solution. Typical error obtained for all elements is generally better
294 than 5% based on replicate measurements of geological reference materials. Results are
295 presented in Table 1.

296 REE patterns were normalized to Post-Archean Australian Shale, the Ce anomaly is
297 defined as $Ce/Ce^* = Ce_N/(Pr_N^2/Nd_N)$ (Lawrence et al., 2006) and the Y anomaly corresponds

298 to the ratio Y_N/H_{ON} (Bau et al., 2014), where subscript N stands for normalized values to
299 PAAS (Post-Archean Australian Shale; Taylor and McLennan, 1995).

300

301 *3.3. Ni isotope measurements*

302

303 Nickel isotopes were measured by MC-ICP-MS (Neptune, Thermofisher Scientific) at
304 the PSO (Pôle de Spectrométrie Océan) at Ifremer (Brest, France) (Table 1). Detailed
305 descriptions of the analytical methods can be found in (Gueguen et al., 2013), though a brief
306 description of methodology is provided below. After a two-stage purification method through
307 ion-exchange chromatography columns using AG1-X8 (for separating Fe, Zn and most of Co
308 and Cu) and Ni-spec Eichrom resins (for complete removal of remaining element matrix),
309 $^{60}\text{Ni}/^{58}\text{Ni}$, $^{61}\text{Ni}/^{58}\text{Ni}$, and $^{62}\text{Ni}/^{58}\text{Ni}$ ratios were measured by MC-ICP-MS in medium or high-
310 mass resolution mode with an ApexQ (ESI, USA) desolvation introduction system.

311 Instrumental mass discrimination was corrected using a double-spike method. A mix
312 of ^{61}Ni and ^{62}Ni isotopes was added to each sample prior to second step of chemical
313 purification, data reduction is based on calculations from (Siebert et al., 2001). Because Ni is
314 not retained on the first column step (columns are filled with an anionic AG1-X8 resin), we
315 did not consider necessary to spike the samples prior to the first separation step. Potential Ni
316 isotope fractionation during columns procedure have already been evaluated in a previous
317 study (Gueguen et al., 2013), and spiking samples only prior to the second separation does not
318 introduce any bias in the Ni isotope composition of the samples and standards. Spiked
319 solution of the international isotopic standard of nickel NIST SRM 986 was routinely
320 analyzed during each analytical session in a similar manner as a sample-standard-bracketing
321 method, which allows monitoring of the reproducibility of each analytical session.
322 Experimental and analytical methods are described in (Gueguen et al., 2013).

323 $^{60}\text{Ni}/^{58}\text{Ni}$ ratios of samples are reported as a deviation from NIST SRM 986 standard isotope
324 composition following equation (1) in per mil:

325

$$326 \quad \delta^{60/58}\text{Ni} = (\text{R}_{\text{spl}}/\text{R}_{\text{NIST986}} - 1) \times 1000 \quad (1)$$

327

328 Where R_{spl} is the $^{60}\text{Ni}/^{58}\text{Ni}$ ratio of the sample and $\text{R}_{\text{NIST986}}$ the $^{60}\text{Ni}/^{58}\text{Ni}$ ratio of the NIST
329 SRM 986 isotopic standard. Typical measurement precision achieved on Ni isotope ratios of
330 samples is better than $\pm 0.04 \text{‰}$ (calculated with the replicate measurements of NIST SRM
331 986) and external precision (two-standard deviation) for delta measurements calculated from
332 replicate measurements of USGS (United States Geological Survey) Mn-nodules
333 geostandards Nod-A-1 and Nod-P-1 is better than $\sim \pm 0.05 \text{‰}$. Replicates include analyses of
334 repeated digestion of geostandards subsamples, repeated processing of subsamples through
335 columns for Ni separation from the matrix, and repeated runs of samples on the mass
336 spectrometer.

337

338 **4. Results**

339 *4.1. Mineralogy*

340 Results of the mineralogy are shown in the supplementary data of Table S1. The
341 mineralogy of hydrogenetic Fe-Mn crusts is difficult to obtain because they are composed of
342 poorly crystallized phases. Our results indicate that hydrogenetic Fe-Mn crusts are composed
343 of a 10 Å Mn-phase, which could be todorokite, and then birnessite. Accessory phases such as
344 quartz, calcite and fluorapatite were encountered. The samples do not present traces of
345 phosphatization (only accessory phases like calcite and fluorapatite are present).

346 Ferromanganese deposits from the Bauer Basin and the Lau Basin are also composed
347 of todorokite (or a 10 Å Mn-phase) and then birnessite. However, these two deposits show

348 that mineral phases (based on the shape of the peak) were more crystallized than hydrogenetic
349 Fe-Mn crusts from this study. Sample NL-10-09 from the Lau basin is mainly constituted by
350 birnessite and then todorokite.

351 The mineralogy of Lo'ihi samples was published in previous studies (Edwards et al.,
352 2011; Toner et al., 2012), and indicate the presence of ferrihydrite and triclinic birnessite.

353

354 *4.2. Bulk hydrogenetic ferromanganese crusts from the Atlantic and Pacific oceans*

355

356 Fe/Mn ratios are generally higher for Atlantic samples, i.e., Ascension FZ, VEMA FZ
357 and Gloria FZ, with values ranging from 1.27 to 1.87 and in the range of the average Fe/Mn
358 ratio for Atlantic crusts (i.e., 1.44; Hein et al., 2013) compared to Pacific crusts (Apuupuu
359 seamount) with Fe/Mn ratios ranging from 0.65 to 1.17 consistent with average Fe/Mn ratio
360 of the Pacific Prime crust zone (i.e., 0.74; Hein et al., 2013). Fe/Mn ratios in the Azores TJ
361 crusts are an exception because they have Fe/Mn ratios as low as 0.39. We observed that the
362 lowest Fe/Mn ratios are found for crusts recovered at shallower depths (i.e., Apuupuu and
363 Azores TJ Fe-Mn crusts). There is a significant compositional variability in the sample set.
364 Notably, the concentration of Ni is up to two times higher in Azores TJ crusts (up to 5,495
365 $\mu\text{g/g}$) than in other Fe-Mn crusts from the Atlantic (average of 2581 ppm; Hein et al., 2013).
366 Co is also four times more enriched in Azores TJ crusts (up to 12,549 $\mu\text{g/g}$) compared to the
367 average of 3,608 $\mu\text{g/g}$ in Atlantic crusts (Hein et al., 2013). Azores TJ crusts exhibit the
368 highest Ni/Mn (0.023 and 0.022) and Co/Mn ratios (0.047 and 0.054) compared to other Fe-
369 Mn crusts (i.e., between 0.009 and 0.017, and between 0.016 and 0.039 respectively). On the
370 other hand, Azores TJ samples have the lowest Zn/Mn ratios (~ 0.024) compared to other Fe-
371 Mn crusts (Zn/Mn between 0.0025 and 0.0044 but with most values > 0.0030) (Figure 3).
372 They also have the lowest Cu/Mn ratios (< 0.0030), while Fe-Mn crusts from Apuupuu,

373 Ascension FZ, VEMA FZ and Gloria FZ have values between 0.0047 and 0.0075 (with the
374 exception of two samples with Cu/Mn ratios of 0.0012 and 0.0028).

375 Rare earth elements patterns normalized to PAAS display a positive Ce anomaly and a
376 negative Y anomaly typical of hydrogenetic deposits (e.g., Bau et al., 1996) (Figure 2).

377 Results show that REE are depleted in Apuupuu and Azores TJ samples compared to other
378 Fe-Mn crusts from the study, and that the Y anomaly is less pronounced. All Fe-Mn crusts
379 samples plot in the hydrogenetic field in the Ce/Ce* vs. Y_N/Ho_N diagram (Figure 2).

380 $\delta^{60/58}\text{Ni}$ values for the hydrogenetic Fe-Mn crusts range from 0.8 ‰ to 1.8 ‰ with
381 most values clustering between 1.3 and 1.6 ‰ (Table 1 and Figure 5B). Azores TJ Fe-Mn
382 crusts show distinct elemental composition (e.g., higher Co/Mn and Ni/Mn and lower Fe/Mn)
383 compared to other Atlantic Fe-Mn crusts but have similar $\delta^{60/58}\text{Ni}$ values (Figure 4). On the
384 other hand, variable $\delta^{60/58}\text{Ni}$ values among Gloria FZ, VEMA FZ and Ascension FZ Fe-Mn
385 crusts are not associated with significant variations in elemental concentrations (Figures 3 and
386 4). Nickel isotope compositions of hydrogenetic Fe-Mn deposits reported here overlap with
387 the range of $\delta^{60/58}\text{Ni}$ values reported in the literature. Gueguen et al. (2016) reported an
388 average $\delta^{60/58}\text{Ni}$ value of 1.79 ± 0.21 ‰ for subsamples collected along stratigraphic profiles
389 acquired in two North Pacific Fe-Mn crusts (Hawaii archipelago) similar to the average value
390 of 1.73 ± 0.21 ‰ obtained for two South Pacific Fe-Mn crusts (Tahiti Archipelago). Gall et al.
391 (2013) reported a range of $\delta^{60/58}\text{Ni}$ values between 0.9 ‰ and 2.5 ‰ for Fe-Mn crusts
392 collected in various oceanic regions, yielding a best estimate for Fe-Mn crusts of 1.6 ‰
393 (Figure 5). We could not confirm the initial observations made by Gall et al. (2013) that Ni
394 isotope compositions in Fe-Mn crusts vary in relation to their distance to continental shelf
395 (Figure 5). In contrast, Ni isotope composition tends to correlate with water depth and
396 shallowest crusts exhibit the lowest Fe/Mn ratios and the heaviest $\delta^{60/58}\text{Ni}$ value (Figure 5).

397

398 *4.3. Ferromanganese oxide deposits from the Bauer Basin, Southwest Pacific*

399

400 Fe/Mn ratios in BB nodules (0.20 and 0.44) are lower than in BB Fe-Mn crust (0.98),
401 the latter being in the range of hydrogenetic Fe-Mn crusts from Apuupuu and the Atlantic.
402 Nickel concentration in BB Fe-Mn crust of 5,886 $\mu\text{g/g}$ is about twice the average of Apuupuu
403 Fe-Mn crusts average value of 2,262 $\mu\text{g/g}$ but it is closer to the average of the North Pacific
404 Prime Zone of 4,216 $\mu\text{g/g}$ (Hein et al., 2013). Nickel contents in BB Mn-nodules are 11,850
405 $\mu\text{g/g}$ and 10,054 $\mu\text{g/g}$ which are close to the average Ni concentration of 13,002 $\mu\text{g/g}$ for
406 nodules from the Clarion Clipperton Fault Zone (CCFZ) in the Central Pacific (Hein et al.,
407 2013). BB nodules and crust have higher Ni/Mn ratios (between 0.03 and 0.05), Zn/Mn ratios
408 (between 0.0046 and 0.0060) and Cu/Mn ratios (0.015 to 0.020) relative to Fe-Mn crusts
409 (Ni/Mn<0.02, Zn/Mn <0.0044 and Cu/Mn<0.0075). In contrast, Co/Mn ratios for the three
410 BB samples are lower (between 0.002 and 0.005) compared to hydrogenetic Fe-Mn crusts
411 (>0.0016). Rare earth elements patterns of BB Fe-Mn deposits are enriched in REE relative to
412 PAAS (Figure 2), they show a negative Ce anomaly and the Y anomaly is not observed, as
413 reported in previous studies (Elderfield and Greaves, 1981; Elderfield et al., 1981b; Elderfield
414 et al., 1981a). In the Ce/Ce* vs. Y_N/Ho_N diagram (Figure 2), BB nodules fall in the diagenetic
415 field and the BB crust in the hydrothermal field.

416 $\delta^{60/58}\text{Ni}$ values of the BB Fe-Mn crust of 0.33 ‰, and of 0.42 ‰ and 0.43 ‰ for BB
417 nodules stand in marked contrast compared to heavier $\delta^{60/58}\text{Ni}$ values (averaging ~1.6 ‰)
418 reported for hydrogenetic crusts and nodules of this study and previously reported values by
419 Gall et al. (2013) and Gueguen et al. (2016) (Table 1).

420

421 *4.4. Hydrothermal Fe-Mn oxide deposits from the Lau Basin, Southeast Pacific*

422

423 Fe-Mn oxide deposits from the Lau Basin are essentially composed of birnessite Mn-
424 oxides phases, which is reflected in the very low Fe/Mn ratios < 0.002 . Ni/Mn, Co/Mn,
425 Zn/Mn and Cu/Mn ratios (< 0.0003 ; < 0.00009 ; < 0.0009 ; and < 0.0012 , respectively) are also
426 extremely low compared to hydrogenetic Fe-Mn crusts and Bauer Basin deposits. Rare earth
427 elements patterns normalized to PAAS for both samples exhibit a negative Ce anomaly, a
428 positive Eu anomaly and a positive Y anomaly. The negative Ce anomaly and positive Eu
429 anomaly are more pronounced for NL-20-05 than for NL-10-09, but the positive Y anomalies
430 have the same amplitude. The overall REE patterns indicate that Lau Basin hydrothermal
431 deposits are depleted in REE relative to hydrogenetic Fe-Mn crusts of this study by 2 to 3
432 orders of magnitude but instead have the same level of enrichment as hydrothermal Fe-Mn
433 deposits from Lo'ihi (Figure 2). LB Fe-Mn deposits fall in the hydrothermal field in the
434 Ce/Ce* vs. Y_N/H_{ON} diagram. These samples display $\delta^{60/58}\text{Ni}$ values of 0.51 and 1.11 ‰ for
435 NL-10-09 and NL-20-05 respectively (Table 1). These values are lighter than the average
436 range of value (1.3 - 1.6 ‰) reported for hydrogenetic Fe-Mn crusts from this study and from
437 the literature (Gall et al., 2013; Gueguen et al., 2016).

438

439 *4.5. Hydrothermal Fe-Mn oxide deposits from Lo'ihi seamount, North Pacific*

440

441 Mineralogy of the deposits is essentially composed of ferrihydrite and triclinic
442 birnessite (Edwards et al., 2011; Toner et al., 2012), and samples are characterized by high
443 Fe/Mn ratios (between 2.84 and 16.68) and low metal/Mn ratios (i.e., Co/Mn $\sim < 0.003$;
444 Cu/Mn $\sim < 0.003$; Zn/Mn $\sim < 0.003$; and Ni/Mn $\sim < 0.008$) compared to other Fe-Mn deposits
445 described in this study (Figure 3). Geochemical composition of Lo'ihi Fe-Mn oxides is
446 reported in (Table 1). REE patterns normalized to PAAS for Lo'ihi deposits show that the
447 REE content is depleted relative to hydrogenetic Fe-Mn crusts and show flat patterns with a

448 small positive Eu anomaly (Figure 2). The Ce/Ce* vs. Y_N/H_{ON} diagram shows that Lo'ih
449 samples are in the hydrothermal field (Figure 2).

450 $\delta^{60/58}\text{Ni}$ values range from -1.49 to -0.80 ‰: with the exception of one sample (J2-244-
451 SS5), heaviest $\delta^{60/58}\text{Ni}$ values are associated with highest Ni/Mn and Fe/Mn ratios (Figure 4).
452 The absence of correlation between $\delta^{60/58}\text{Ni}$ values and Al/Fe ratios rules out the influence of
453 detrital particles like volcanic debris in Fe-rich mats on Ni isotopes.

454

455 **5. Discussion**

456

457 *5.1. Mechanisms of Ni isotope fractionation and formation of marine ferromanganese* 458 *deposits*

459 Consistent with previous studies (Gall et al., 2013; Gueguen et al., 2016),
460 hydrogenetic crusts are characterized by isotopically heavy, yet variable, Ni isotope signatures
461 ($\delta^{60/58}\text{Ni}$ values range from 0.9 ‰ and 2.5 ‰). Our results suggest that Ni isotope
462 composition of Fe-Mn crusts, may be fractionated towards both lighter to heavier values
463 relative to modern deep seawater (average ~1.4 ‰). Hydrogenetic Fe-Mn crusts are formed
464 by slow precipitation of Fe-Mn colloids from seawater, with the volcanic substrates of
465 seamounts allowing the precipitation of Fe-Mn oxyhydroxides (Koschinsky and Halbach,
466 1995; Koschinsky and Hein, 2003; Koschinsky et al., 2003). The process of Mn oxidation at
467 the base of the Oxygen Minimum Zone (OMZ) provides an efficient scavenging process of
468 seawater dissolved trace metals, such as Co, Ni, Zn, Tl and Cd, by Mn-oxide colloids, which
469 contributes to precipitation of Mn-rich layers of Fe-Mn crusts (e.g.; Mizell et al., 2020).
470 Several mechanisms may, therefore, contribute to the variability of Ni isotope composition in
471 Fe-Mn crusts, and we explore them below.

472

473 *5.1.1. The role of Ni speciation in seawater and in the water column*

474

475 The water depth and local oceanographic setting are potentially important parameters
476 affecting the chemical composition of Fe-Mn crusts. Results from Fe-Mn crusts from the Line
477 Islands (Aplin and Cronan, 1985), the Hawaii chain (De Carlo et al., 1987) and the Central
478 Pacific (Halbach et al., 1984) indicate that crusts formed in shallow waters can have four
479 times lower Fe/Mn ratios and up to ten times higher metal concentrations (Ni, Co, V, Pb, Mo,
480 Cd) than deeper crusts. In deeper waters (> 3000 m depth), Fe-Mn crusts are generally
481 characterized by higher Fe/Mn ratios due to lower inputs of Mn from the OMZ in addition to
482 an increased Fe flux through the dissolution of carbonate testas from microorganisms below
483 the CCD (Halbach and Puteanus, 1984). Other factors such as, increase of the pH favoring
484 precipitation of Fe, increase of the detrital particles in the water column (Mizell et al., 2020),
485 and input of Fe from deep hydrothermal sources (Horner et al., 2015), may explain increasing
486 Fe/Mn ratios in crusts with depth. By contrast, in shallow waters, e.g., below the OMZ (i.e.,
487 located between 500 m and 1500 m depth), the proportion of Mn-rich colloids in the water
488 column is presumably high (e.g.; Klinkhammer and Bender, 1980). The lowest Fe/Mn ratios
489 in our sample set correspond to shallowest (< 2500 m water depth) hydrogenetic Fe-Mn crusts
490 associated with the heaviest $\delta^{60/58}\text{Ni}$ values (i.e., > 1.5 ‰) (Apuupuu and Azores TJ, Figures 4
491 and 5). Azores TJ samples have the highest Co/Mn ratios (~0.5), while Apuupuu crusts have
492 ratios in the range of other crust of the study (0.2-0.3). Hydrogenetic Fe-Mn crusts from
493 Gloria, Ascension and VEMA areas collected at water depth > 2500 m have the lightest
494 $\delta^{60/58}\text{Ni}$ values (between 1.5 ‰ and 0.8 ‰) and the highest Fe/Mn ratios. More than 90% of
495 the Ni in Fe-Mn nodules and crusts is in the Mn-phase (Koschinsky and Halbach, 1995;
496 Mohwinkel et al., 2014; Wegorzewski et al., 2020), and this phase association does not
497 change with water depth. As a consequence, a change in the Fe/Mn ratio in the crusts should

498 not impact the Ni isotope composition of the crusts. In addition, the variability of Ni isotope
499 composition in deep water (i.e., < 1500 m water depth) is very limited < 0.2 ‰, while surface
500 waters (< 200 m water depth) deviate by 0.4 to 0.5 ‰ relative to deep water (Cameron and
501 Vance, 2014; Takano et al., 2017; Wang et al., 2019; Archer et al., 2020). Therefore, it is
502 unlikely that variations in Fe-Mn crusts are explained by the variability of Ni isotopes in
503 oceanic water masses.

504 An alternative explanation, but not mutually exclusive, is to consider isotopic
505 fractionation between inorganic and organic ligands. Experimental and theoretical studies on
506 Ni isotopes demonstrate fractionations up to 2.5 ‰ for $\delta^{60/58}\text{Ni}$ values during exchange
507 reactions between inorganic and organic ligands, with a range of fractionation of ~ 1.7 ‰
508 between inorganic species and a range of 2.5 ‰ between organic species (Fujii et al., 2011).
509 Accordingly, if one of these fractionated species is preferentially adsorbed onto the
510 oxyhydroxide surface it would impart a net isotope fractionation between the mineral and the
511 solution. In addition, hydrated Ni^{2+} ions are isotopically heavier than Ni-chloro complexes in
512 the order of $\text{Ni}^{2+} > \text{NiCl}^+ > \text{NiCl}_2$ (Fujii et al., 2011). Therefore, one could speculate that
513 aqueous Ni^{2+} , which is presumably preferentially adsorbed onto Fe-Mn crusts (Koschinsky
514 and Halbach, 1995), would be enriched in heavy Ni isotopes relative to other inorganic
515 species and organic species, thus consistent with heavier Ni isotopes in hydrogenetic Fe-Mn
516 crusts relative to Ni isotopes in seawater.

517

518 *5.1.2. Isotopic fractionation during Ni sorption processes*

519

520 It has been well documented that Ni is bound to Mn-oxyhydroxides and is easily
521 adsorbed onto these phases (Hem, 1978; Koschinsky and Halbach, 1995; Koschinsky and
522 Hein, 2003; Peacock and Sherman, 2007a; Peacock and Sherman, 2007b), and that Ni

523 adsorbed on birnessite is structurally incorporated within the mineral structure with time
524 (Peacock, 2009). Recent results for Ni sorption on birnessite indicate large Ni isotope
525 fractionation up to ~ -3 ‰ on the solid phase because structural vacancies in birnessite are
526 preferentially occupied by light Ni isotopes (Sorensen et al., 2020). The composition of
527 hydrogenetic Fe-Mn crusts enriched in heavier Ni isotopes compared to seawater is therefore,
528 at odds with experimental data. However, Gueguen et al. (2016) showed that light $\delta^{60/58}\text{Ni}$
529 values up to 0.25 ‰ in hydrogenetic Fe-Mn crusts from the South Pacific were the result of
530 post-depositional and alteration processes at the bottom of the crust in contact with the
531 substrate. They proposed that circulating fluids altered the substrate at the bottom of the
532 hydrogenetic crusts which result in dissolution and reprecipitation of Mn-oxyhydroxide
533 phases. This was accompanied by Ni isotope fractionation during sorption to Mn-
534 oxyhydroxide phases producing anomalously light $\delta^{60/58}\text{Ni}$ values. Consistent with these
535 results, light $\delta^{60/58}\text{Ni}$ values (-0.8 ‰ to -0.2 ‰) found in metalliferous sediments are
536 interpreted as the result of diagenetic processes affecting the Ni cycling with Mn under oxic
537 conditions (Little et al., 2020). Therefore, significant isotope fractionation during Ni sorption
538 and the importance of alteration and deposition processes highlight the importance of
539 mineralogy and formation processes for Ni isotope fractionation in marine Fe-Mn deposits.

540 Despite the poor crystallinity of our hydrogenetic Fe-Mn crusts samples, we could
541 identify two predominant Mn- phases (with the exception of Lo'ihi deposits mainly
542 constituted by Fe- phases) : a 10 Å Mn-phase (maybe todorokite) and then birnessite. By
543 contrast, LB deposits are predominantly composed of birnessite and to a lesser extent
544 todorokite and they have lighter $\delta^{60/58}\text{Ni}$ values (0.5 and 1.1 ‰) than hydrogenetic Fe-Mn
545 crusts. Identification of the mineral phases is easier for these samples due to their greater
546 crystallinity in comparison to hydrogenetic Fe-Mn crusts. Little et al. (2020) hypothesized
547 that light $\delta^{60/58}\text{Ni}$ values from -0.8 ‰ to -0.2 ‰ in metalliferous sediments from two MANOP

548 (Manganese Nodule Project) sites are related to post-depositional processes involving the
549 dissolution and reprecipitation of Mn-oxides during diagenesis (Little et al., 2020). The
550 authors propose that the diagenetic cycling of Ni associated with Mn could be a key process
551 for explaining light Ni $\delta^{60/58}\text{Ni}$ values in marine metalliferous deposits. For example, they
552 suggest that the transformation of birnessite to todorokite under oxic conditions could favor
553 the preferential incorporation of Ni in todorokite. Therefore, if we compare these results with
554 results from our study, a mineralogical change in the Mn-phase in deeper hydrogenetic Fe-Mn
555 crusts could possibly explain the lighter Ni isotope values in hydrogenetic Fe-Mn crusts
556 (Gloria, Ascension and VEMA) compared to Azores TJ and Apuupuu Fe-Mn crusts; although
557 we agree that more in-depth mineralogical analyses are necessary to confirm this assumption.
558 On the other hand, this interpretation is not consistent with our results for the Lau Basin,
559 because the lighter $\delta^{60/58}\text{Ni}$ value (0.5 ‰) observed in sample NL-10-09 has a predominance
560 of birnessite, while the sample NL-20-05 with a $\delta^{60/58}\text{Ni}$ value of 1.1 ‰ is predominantly
561 composed of todorokite. This implies that the mineralogy of Lau Basin is probably not the
562 cause for the difference in $\delta^{60/58}\text{Ni}$ values between the two samples.

563

564 *5.1.3. Are hydrothermal sources of Ni recorded in Fe-Mn deposits ?*

565

566 Before the discovery of active hydrothermal sources, low Al/(Al+Fe+Mn) ratios found
567 in deep-sea sediments from the East Pacific were interpreted as being linked to inputs from
568 the volcanic activity of the East Pacific Rise (Boström et al., 1969). This suggests that Fe-Mn
569 precipitates can travel over thousands of kilometers away from the vents. Recent evidence
570 from direct measurements of seawater during the GEOTRACES transects, shows that Fe and
571 Mn emitted at the EPR are transported several hundreds of kilometers away from the vents
572 (e.g.; Lam et al., 2018; Lee et al., 2018). The transport of Ni is not as well documented as for

573 Fe and Mn. However, results of the Ni concentration in seawater above a hydrothermal source
574 measured along the GEOTRACES transect GP16 in the South East Pacific, shows that the
575 background Ni concentration in seawater is not perturbed (Lee et al., 2018).

576 Nickel concentrations in hydrothermal fluids are generally close to seawater due to
577 negligible mobilization of Ni during alteration of the oceanic crust (Von Damm, 1995).
578 Differences are, nevertheless, observed depending on geological settings, i.e., Ni
579 concentrations of 2-3 $\mu\text{mol/L}$ were measured in Rainbow vent fluids on the mid-Atlantic
580 oceanic ridge (Douville et al., 2002), in comparison to the range of values for dissolved Ni in
581 deep seawater of 6-10 nmol/L and surface seawater of 2-5 nmol/L (Cameron and Vance,
582 2014; Takano et al., 2017; Wang et al., 2019; Archer et al., 2020). Because ultramafic-hosted
583 hydrothermal vents are not the predominant type of hydrothermal systems, the global Ni
584 hydrothermal flux to the oceans is presumably low, implying that high-temperature
585 hydrothermal fluids are probably not a major contributor to the Ni oceanic budget. However,
586 significant Ni enrichment has been observed in low-temperature hydrothermal venting
587 (Sedwick et al., 1992; Wheat et al., 2002; Wheat et al., 2003), which could possibly produce a
588 significant flux of Ni to the oceans.

589 Marine Fe-Mn deposits represent one possible way to investigate the effect of direct
590 contribution of hydrothermal sources on the Ni oceanic budget and Ni isotopes in seawater
591 through time. For example, several studies demonstrated that the geochemistry of Fe-Mn
592 crusts could be affected by hydrothermal inputs from distal (far-field) hydrothermal sources
593 (Elderfield and Greaves, 1981; van de Flierdt et al., 2004; Chu et al., 2006; Gall et al., 2013).
594 More specifically for Ni, Gall et al. (2013) attributed the range of Ni isotope values in Fe-Mn
595 crusts from 0.9 to 2.5 ‰ to variable Ni sources in seawater, in particular hydrothermal
596 sources, where lighter $\delta^{60/58}\text{Ni}$ values in a Central Pacific Fe-Mn crust were tentatively
597 ascribed to local hydrothermal inputs in the water column (Gall et al., 2013). Little et al.

598 (2020) considered that Ni scavenging onto hydrothermal Fe particles followed by their
599 deposition in the sediment, is one of the possible explanations for light Ni isotopes (up to ~ -
600 0.8 ‰) observed in metalliferous sediments from the MANOP site M close to the East Pacific
601 Rise. Therefore, hydrothermal inputs are a possible factor for explaining the variability of Ni
602 isotopes in Fe-Mn deposits, and thus a key process for estimating the Ni isotope composition
603 of the oxic Mn sink. In the following section, we intend to address this hypothesis by
604 investigating hydrothermal and mixed hydrothermal-hydrogenetic deposits using the
605 combination of rare earth element systematics and Ni isotopes.

606

607 *5.2. A classification scheme using REE and Ni isotopes*

608

609 The combination of REE geochemistry and Ni isotopes could give us insights for
610 deciphering Ni sources and formation processes in a continuum of well-characterized
611 hydrogenetic Fe-Mn crusts, hydrothermal deposits (Lo’ihi and Lau basin), and hydrogenetic
612 Fe-Mn deposits with a contribution from far-field hydrothermal inputs (i.e., Bauer basin). We
613 use three indicators in our study: Ce anomaly, Y anomaly and Sm_N/Yb_N ratio. A positive Ce
614 anomaly is an indicator of excess of accumulation of Ce in hydrogenetic Fe-Mn crusts due to
615 oxidation of Ce(III) to Ce(IV) at the surface of Fe-Mn crusts (Bau et al., 2014). By contrast,
616 hydrogenetic Fe-Mn crusts are characterized by a negative Y anomaly (Y_N/Ho_N ratio < 1), and
617 indicates preferential scavenging of Ho rather than Y on its surface. Finally, the Sm_N/Yb_N
618 ratio is an indicator of the variation of the slope in the REE pattern, and this would potentially
619 indicate changes in the REE source to the deposits. Our samples are classified according to
620 Figure 6: the hydrogenetic field (i.e., Fe-Mn crusts) is defined for Fe-Mn deposits with
621 $\delta^{60/58}Ni$ values > 0 ‰, $Ce/Ce^* > 1$, $Y_N/Ho_N < 1$ and $Sm_N/Yb_N < 0.6$; the hydrothermal field
622 (i.e., Lo’ihi deposits) is defined by $\delta^{60/58}Ni$ values < 0 ‰, $Ce/Ce^* < 1$, $Y_N/Ho_N > 1$ and

623 $\text{Sm}_N/\text{Yb}_N > 0.6$; and the mixed hydrothermal-hydrogenetic field (i.e., LB deposits and BB
624 deposits) corresponds to $\delta^{60/58}\text{Ni}$ values $> 0\text{‰}$, $\text{Ce}/\text{Ce}^* < 1$, $\text{Y}_N/\text{Ho}_N > 1$ and $\text{Sm}_N/\text{Yb}_N > 0.6$.
625 The threshold value of 0.6 for the Sm_N/Yb_N ratio is defined based on the distribution of BB
626 deposits (Figure 6). Based on these observations, we discuss below the impact of a possible
627 influence of hydrothermal inputs in our selection of Fe-Mn deposits.

628

629 *5.2.1. Ferromanganese deposits formed near seafloor hydrothermal sources:*
630 *examples from the Lo'ihī seamount (North Pacific) and the Lau Basin (Western Pacific)*

631

632 Rare earth elements, Co/Mn ratios and Fe/Mn ratios of Lo'ihī and LB deposits indicate
633 a hydrothermal origin of these deposits, i.e., they are influenced by near-field hydrothermal
634 inputs. They represent two end-members, Fe-rich and Mn-rich respectively. REE-depleted
635 patterns of Lo'ihī and LB deposits show a positive Eu anomaly which is more pronounced in
636 the LB deposits. While the negative Ce anomaly and HREE enrichment in LB deposits are
637 typical of a seawater source for REE, the positive Eu anomaly is a robust signature of REE
638 derived from high temperature hydrothermal fluids (German et al., 1990). The Ce/Ce^* vs.
639 Y_N/Ho_N diagram does not discriminate between LB deposits and Lo'ihī deposits. LB deposits
640 display different $\delta^{60/58}\text{Ni}$ values compared to Lo'ihī deposits, i.e., positive $\delta^{60/58}\text{Ni}$ values of
641 0.5 ‰ and 1.1 ‰ and negative $\delta^{60/58}\text{Ni}$ values from -1.5 ‰ to -0.7 ‰, respectively. Thus,
642 although both Lo'ihī and LB deposits are affected by hydrothermal inputs, the fractionation of
643 Ni isotopes is clearly different between the two deposits, suggesting that the effects induced
644 by near-field hydrothermal inputs are different according to the type of deposits.

645 The combination of REE and Ni isotopes data presented in Figure 6 show that Lo'ihī
646 deposits systematically fall in the hydrothermal field, with negative $\delta^{60/58}\text{Ni}$ values, negative
647 Ce anomalies, positive Y anomalies and Sm_N/Yb_N ratios > 0.6 . In contrast, as indicated by

648 Sm_N/Yb_N ratios < 0.6 and $\delta^{60/58}\text{Ni}$ values > 0 (Figure 6), LB deposits have probably a
649 contribution from both a hydrothermal source and a hydrogenetic source. Three possible
650 causes can explain the difference in Ni isotope composition between Lo’ihi and LB deposits,
651 (1) growth rates, (2) $\delta^{60/58}\text{Ni}$ value of the source fluid, and (3) mineralogy and processes of Ni
652 removal from the fluid and/or from seawater. Growth rates could be different due to the
653 geological context of the hydrothermal source (e.g., back-arc basin, slow versus fast spreading
654 ridges) or composition of the hydrothermal fluids. But because both deposits formed close to
655 hydrothermal venting, growth rates are likely of the same order. Additional measurements of
656 the growth rates of LB and Lo’ihi deposits would confirm this statement, but they are likely
657 different and faster in comparison to hydrogenetic crusts, which does not favor hypothesis 1.

658 Nickel concentrations up to 80 nmol/L have been measured in Lo’ihi hydrothermal
659 fluids (Sedwick et al., 1992), which is about 10 times the average seawater concentration
660 (Sclater et al., 1976; Bruland, 1980; Cameron and Vance, 2014; Takano et al., 2017; Wang et
661 al., 2019). However, the Ni isotope composition of Lo’ihi hydrothermal fluids has not been
662 published yet. Our best assumption is to consider that Ni isotopes in the fluid have either the
663 composition of the basalt (i.e., ~BSE, 0.1 ‰), or the composition of seawater (i.e., ~1.4 ‰).
664 LB deposits formed by low temperature diffuse hydrothermal venting through the volcanic
665 substrate, and REE-depleted but HREE-enriched patterns and the negative Ce anomalies
666 suggest that they formed by mixing between seawater and hydrothermal fluids. Therefore, we
667 speculate that the Ni isotope composition of the source fluids in both Lo’ihi and LB deposits
668 ranging between volcanic rock and seawater values (0 to 1.4 ‰), does not explain the most
669 negative $\delta^{60/58}\text{Ni}$ values from -1.5 ‰ to -0.7 ‰ at Lo’ihi. Hence, we do not favor hypothesis 2
670 implying variable Ni isotope composition of the sources.

671 Lo’ihi deposits are dominated by Fe-phases (i.e., Fe/Mn ratios range from 2.84 to
672 16.68), while LB deposits are dominated by Mn-phases (i.e., Fe/Mn ratios < 0.002). Thus, it is

673 possible that the mineralogy and processes involved in Ni removal in the deposits, influenced
674 the fractionation of Ni isotopes (hypothesis 3). Nickel isotope fractionation during sorption on
675 Fe- and Mn-oxide has been determined through experimental studies and it was shown that
676 the Fe-Mn mineral phase is systematically enriched in light Ni isotopes (Wasylenki et al.,
677 2015; Gueguen et al., 2018; Sorensen et al., 2020). The Ni isotope fractionation is much
678 larger on Mn-oxides than Fe-oxides (i.e., in the order of -2 ‰ to -3 ‰ for Mn-oxides, and -0.8
679 ‰ to -0.3 ‰ for Fe-oxides). Ni/Mn ratios are very low in LB deposits for two reasons. First,
680 LB deposits are composed of pure Mn-oxides (Fe is almost a trace element and Fe/Mn <
681 0.002). Second, Ni concentration is low (~10-12 nmol/L) in seawater and probably in the
682 hydrothermal fluid. This observation, combined with the fast rate of precipitation (e.g., low
683 Co/Mn ratios), did not allow scavenging of large amount of Ni in LB (in comparison with
684 hydrogenetic Fe-Mn crusts for example) as shown by relatively low Ni concentrations
685 (between ~50 $\mu\text{g}\cdot\text{g}^{-1}$ and ~100 $\mu\text{g}\cdot\text{g}^{-1}$) and Ni/Mn ratios <0.0003. Lighter $\delta^{60/58}\text{Ni}$ values (0.5
686 ‰ and 1.1 ‰) in LB deposits compared to seawater values (~1.4 ‰) may reflect reservoir
687 and/or mineralogical effects through scavenging of seawater trace metals in hydrothermal
688 Mn-phases (i.e., birnessite and or todorokite; Fouquet et al., 1993). One can assume that after
689 formation of LB deposits by precipitation of metals from seawater and hydrothermal fluids,
690 subsequent aging of the deposits and exposure at the seafloor allows further scavenging of
691 seawater-derived elements including Ni on Mn-oxides phases. Such mechanism have been
692 suggested for Cd isotopes sorption to birnessite (Wasylenki et al., 2014), with slow exchange
693 of Cd between seawater and birnessite.

694 By contrast, isotopically light Ni (from -1.5 ‰ to -0.8 ‰) in Lo'ihi deposits is due to
695 the removal of Ni during Ni adsorption from seawater and from the hydrothermal fluid,
696 followed by isotope fractionation between the fluid and mixed Mn/Fe-oxides based on
697 experimental results of Ni sorption to Mn-oxides (Sorensen et al., 2020) and Fe-oxides

698 (Wasylenki et al., 2015; Gueguen et al., 2018). Our results can be compared to metalliferous
699 deposits from the MANOP site M (East Pacific), which show light $\delta^{60/58}\text{Ni}$ values up to -0.8
700 ‰ and a similar range of Fe/Mn ratios from 3.8 to 44.6 (most values are between 3.6 and 6.3)
701 (Little et al., 2020) than Lo'ihi deposits (between 2.8 and 16.7). The results of Little et al.
702 (2020) were interpreted as possibly being the result of the deposition of hydrothermal Fe
703 oxyhydroxides associated with sorption of isotopically light Ni onto the hydrothermal Fe
704 oxyhydroxides.

705

706 *5.2.2. Fe-Mn deposits affected by far-field hydrothermal activity: example from the* 707 *Bauer Basin (Eastern Pacific)*

708

709 It has been proposed that oceanic circulation in the Bauer Basin allows export of
710 hydrothermal precipitates from the East Pacific Rise (EPR) (Heath and Dymond, 1977; Lyle
711 et al., 1977; Ravizza and McMurtry, 1993). Elderfield and Greaves (1981) suggested that
712 diagenetic processes in the sediment allowed the incorporation of hydrothermal Fe-oxides and
713 REE with negative Ce anomalies ($\text{Ce}/\text{Ce}^* < 0.5$) in the BB nodules, while the BB Fe-Mn
714 crust has a hydrogenetic origin (Lyle et al., 1977). This is confirmed by the Ce/Ce^* vs.
715 $\text{Y}_\text{N}/\text{Ho}_\text{N}$ diagram (Figure 2) showing that the BB crust falls in the hydrothermal field, and the
716 BB nodules in the diagenetic field. In addition to their negative Ce anomaly, REE patterns of
717 BB samples (crust and nodules) are on average depleted compared to other hydrogenetic
718 deposits (Table 1). Another important feature of the BB crust and nodules is their unusually
719 low Co/Mn ratios ($\text{Co}/\text{Mn} < 0.005$). Therefore, REE patterns and low Co/Mn ratios in BB
720 deposits could indicate hydrothermal input of Fe-Mn from the EPR in the deposits, either
721 during diagenetic reactions for the nodules, or by settling of hydrothermal plume fallouts
722 during formation of Fe-Mn crusts. The latter process has been previously suggested for Fe-

723 Mn crusts from the Central Indian Ridge (Kuhn et al., 1998). Finally, the contribution from
724 direct hydrothermal fluids in BB Fe-Mn oxide deposits can be precluded because the average
725 Ni/Mn ratios in hydrothermal fluids from the global ocean (Von Damm, 1995; Douville et al.,
726 2002) are much lower than Ni/Mn ratios in the BB Fe-Mn deposits. In addition, samples also
727 do not have a positive Eu anomaly indicative of REE precipitation from high temperature
728 hydrothermal fluids (the Bauer Basin is located ~1000 km to the East of the EPR).

729 $\delta^{60/58}\text{Ni}$ values in BB crust and nodules, i.e., 0.3 ‰ and 0.4 ‰ respectively, are
730 isotopically lighter than the average range for hydrogenetic Fe-Mn crusts of 1.3-1.6 ‰
731 (Figure 5B) and the average seawater of 1.4 ‰. The light $\delta^{60/58}\text{Ni}$ values in BB nodules could
732 possibly be explained by diagenetic remobilization of Ni within the sediment, as suggested by
733 Little et al. (2020) for Mn-nodules and metalliferous sediments from the Eastern Pacific and
734 in agreement with Elderfield and Greaves (1981) for rare earth elements in BB nodules.
735 However, it is unlikely that the same process applies to the BB Fe-Mn crust. Accordingly, in
736 agreement with the REE geochemistry, we favor the implication of far-field inputs from
737 hydrothermal sources through dispersed hydrothermal plume fallouts, such as Fe-oxides
738 precipitates which sorbed isotopically light Ni, on the formation of BB Fe-Mn crust and
739 possibly Mn-nodules. Little et al. (2020) also proposed that light $\delta^{60/58}\text{Ni}$ values up to -0.8 ‰
740 in metalliferous sediments deposited close to the East Pacific Rise could be due to scavenging
741 of light Ni isotopes on hydrothermal Fe-oxides, which were then incorporated in the
742 sediments. We suggest that a similar process could have occurred for BB Fe-Mn crust,
743 whereby scavenging of isotopically light Ni onto either hydrothermal Mn- or Fe-oxides were
744 then deposited in Fe-Mn crusts.

745

746 *5.3. Nickel isotopes in marine Fe-Mn deposits and implication for the marine Ni isotope*
747 *budget*

748

749 Investigation of rare earth elements and Ni isotopes in a continuum of marine Fe-Mn
750 deposits from typically hydrothermal, to various mixed hydrothermal and hydrogenetic and to
751 strictly hydrogenetic shows that Ni isotopes in Fe-Mn deposits are variable. These results
752 confirm that Fe-Mn crusts do not provide an accurate record of the global burial flux of Ni
753 and its isotopes via sorption to Fe-Mn oxides, implying that the Ni isotope composition of the
754 global sink of Ni Mn-oxides is not necessarily best represented by hydrogenetic Fe-Mn crusts.
755 Seafloor hydrothermal vents represent the main Mn source in seawater (Elderfield and
756 Schultz, 1996), producing widespread Mn enrichment, both in the water column and seafloor
757 sediments, up to 1000's km away from the spreading axis (e.g., Klinkhammer and Hudson,
758 1986; Resing et al., 2015). In particular, the relationship between dissolved Mn and ^3He in the
759 plume off South EPR indicates that a large fraction of hydrothermal Mn is precipitated within
760 the vicinity of ridge axis, while a minor but still significant fraction behaves conservatively
761 and disperse away in the ocean basin. We propose that the Mn sink associated with the early
762 (and rapid) precipitation of Mn oxide in the water column is characterized by isotopically
763 light Ni relative to seawater, while late-stage Mn sink, occurring over longer time scale and
764 eventually recorded in Fe-Mn crusts, yield Ni isotope composition similar to, or even heavier
765 than seawater. The results of this study also show that formation processes such as
766 precipitation and adsorption processes are the main controls of Ni isotope fractionation and
767 that these processes are dependent on conditions prevailing in the depositional environment.
768 Alternatively, heavier Ni isotopes in Fe-Mn crusts compared to seawater could be explained
769 by the selective incorporation of isotopically heavy Ni species in seawater likely produced by
770 organic complexation (see section 5.1.1.). The Mn sink is the major sink for Ni in the ocean,
771 and we have shown that the Ni isotope composition of the Mn sink is complex. Therefore, a
772 rigorous determination of Ni isotopes in fluxes associated with the Mn sink is required to

773 balance the oceanic Ni budget, in particular in the authigenic fraction of deep-sea pelagic
774 sediments.

775

776 **6. Summary**

777

778 In this study, we combined REE and Ni isotope geochemistry in a continuum of
779 samples from pure hydrothermal, hydrothermally-affected by far-field inputs to pure
780 hydrogenetic Fe-Mn deposits from different geological settings and geographical locations.
781 Results show that hydrogenetic Fe-Mn crusts in both the Atlantic and the Pacific oceans have
782 heavy Ni isotope compositions from 0.8 to 1.8 ‰ (with most values comprised between 1.3
783 and 1.6 ‰), similar to or slightly heavier than seawater value. Rare earth element
784 geochemistry (enriched REE contents, positive Ce anomaly, negative Y anomaly, $Sm_N/Yb_N >$
785 0.6) and heavy Ni isotope compositions are the result of Ni removal from seawater through
786 adsorption reactions and structural incorporation of Ni into Mn-oxide minerals.

787 Hydrothermal deposits from Lo'ihi and Lau Basin have REE-depleted patterns with
788 positive Y anomaly, negative Ce anomaly, and $Sm_N/Yb_N < 0.6$. However, Ni isotope values
789 are different between the two deposits, Lo'ihi are characterized by $\delta^{60/58}Ni$ values between -
790 1.5 ‰ and -0.7 ‰ while LB deposits have heavier $\delta^{60/58}Ni$ values from 0.5 ‰ and 1.1 ‰.
791 Extremely light $\delta^{60/58}Ni$ values in Lo'ihi are most likely due to the predominance of Fe-oxides
792 which adsorbed isotopically light Ni from seawater or the hydrothermal fluid. By contrast,
793 $\delta^{60/58}Ni$ values in LB deposits are interpreted as the result of continuous exchange of Ni with
794 seawater during aging of the deposits.

795 The Bauer Basin crust and nodules are intermediate because these deposits are
796 characterized by REE-enriched patterns with negative Ce anomaly and flat to negative Y
797 anomaly and isotopically light Ni compared to seawater, i.e., 0.3 ‰ and 0.4 ‰. We suggest

798 that the Ni isotope composition of BB deposits is the result of incorporation of isotopically
799 light Ni from hydrothermal Fe-oxides coming from the East Pacific Rise.

800 We showed that Ni isotopes in the major sink for Ni in the oceans, Fe-Mn oxide
801 phases, are variable and that a proper estimate of each sink flux for each type of deposits
802 would help explaining the imbalance in the Ni oceanic budget. The combination of Ni
803 isotopes with REE geochemistry shows that mineralogy and formation processes is a major
804 control for Ni isotope variability in Fe- and Mn-rich deposits instead of Ni sources.

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814 **Figure captions :**

815

816 **Figure 1:** Photographs of the seafloor showing outcrops of (A) Lo'ihi Fe-rich deposits, (B)
817 Apuupuu Fe-Mn crusts (sample J2-480-R11) (C) and Lau Basin Mn-rich deposits (photograph
818 is reproduced from Fouquet et al., 1993), the area showed is ~10 meters large. A section of a
819 bulk Fe-Mn crust sample collected on Apuupuu seamount is shown in (D). Sample locations
820 are shown on the world map (E). Bulk hydrogenetic Pacific and Atlantic Fe-Mn crusts are
821 indicated with red circles, hydrothermal Fe-Mn deposits from Lo'ihi and Lau Basin are
822 displayed in grey, Bauer Basin Fe-Mn deposits are shown in green.

823

824 **Figure 2 :** (A) Rare earth elements patterns normalized to PAAS (Post-Archean Australian
825 Shale; Taylor and McLennan, 1995) and (B) Ce/Ce^* vs. Y_N/Ho_N , in hydrogenetic Fe-Mn
826 crusts (red), Bauer Basin Fe-Mn deposits (green), and Lo'ihi (grey) and Lau Basin
827 hydrothermal deposits (pink).

828

829 **Figure 3 :** Plot of (A) Co/Mn, (B) Cu/Mn and (C) Zn/Mn versus Ni/Mn ratios in bulk
830 hydrogenetic Fe-Mn crusts (red) and Bauer Basin Fe-Mn deposits (green), and hydrothermal
831 Fe-Mn deposits from Lo'ihi (grey) and Lau Basin (pink).

832

833 **Figure 4:** Nickel isotope composition (‰) versus (A) Ni/Mn and (B) Fe/Mn ratios in
834 hydrogenetic Fe-Mn crusts (red), Bauer Basin Fe-Mn deposits (green), Lo'ihi hydrothermal
835 deposits (grey) and Lau Basin hydrothermal deposits (pink).

836

837 **Figure 5 :** (A) Nickel isotope composition versus depth of deposition of hydrogenetic Fe-Mn
838 crusts, and (B) distribution of Ni isotope values of hydrogenetic Fe-Mn crusts. The values

839 represented include data from Gall et al. (2013), Gueguen et al. (2016) and data from this
840 study.

841

842 **Figure 6 :** Nickel isotope composition versus (A) Ce/Ce*, (B) Y_N/Ho_N and (C) Sm_N/Yb_N in
843 hydrogenetic Fe-Mn crusts (red), Bauer Basin Fe-Mn crust (green), Lo’ihi hydrothermal
844 deposits (grey) and Lau Basin hydrothermal deposits (pink). The Ce anomaly is defined as
845 $Ce/Ce^* = Ce_N / (Pr_N^2 / Nd_N)$ (Lawrence et al., 2006). This figure allows distinguishing between
846 three groups of samples: (1) hydrogenetic Fe-Mn crusts with positive Ce anomalies and
847 $\delta^{60/58}Ni$ values falling in the range of seawater, (2) hydrothermal deposits from Lo’ihi with no
848 Ce anomalies and light $\delta^{60/58}Ni$ values, and (3) Bauer Basin and Lau Basin Fe-Mn deposits
849 displaying negative Ce anomalies and $\delta^{60/58}Ni$ values that are lighter than seawater values.

850

851 **Table captions:**

852

853 **Table 1:** Nickel isotope composition (‰), selected elemental ratios (μg/μg), and Ce/Ce* (Ce
854 anomaly) and Y_N/Ho_N (Y anomaly) of Fe-Mn deposits.

855

856 **Supplementary material:**

857

858 **Table S1:** Sampling information and mineralogy (X-Ray diffraction) of bulk hydrogenetic
859 Fe-Mn crusts and nodules, and Lau Basin Fe-Mn deposits.

860

861 **Table S2:** Geochemical composition (μg/g) of Fe-Mn deposits.

862

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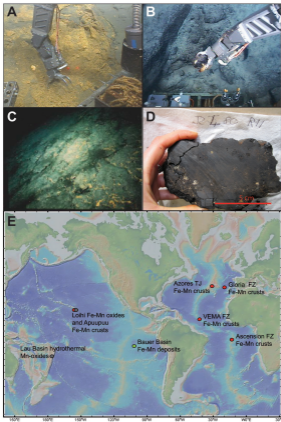


Figure 1

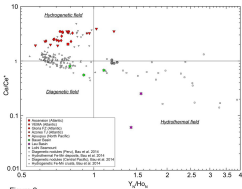
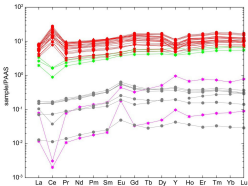


Figure 2

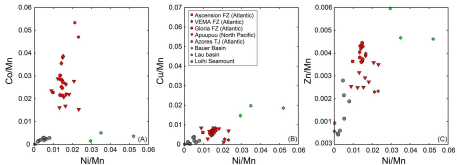


Figure 3

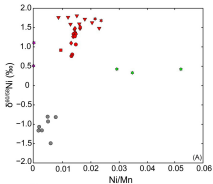
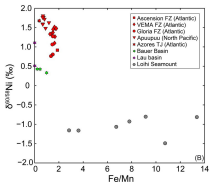


Figure 4



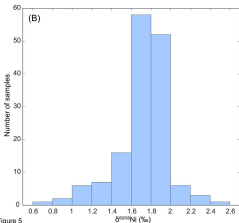
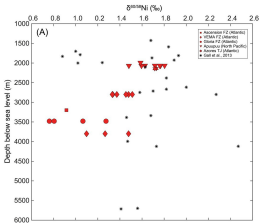


Figure 5

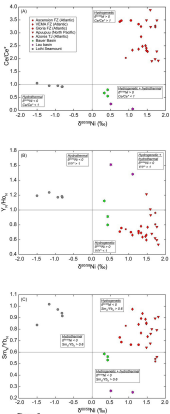


Figure 6

Table 1: Nickel isotope composition (‰), selected elemental ratios (μg/μg) and calculated Ce anomaly (Ce/Ce*) and Y anomaly (Y_N/Ho_N) of Fe-Mn deposits.

Sample Name	Location	δ ^{60/58} Ni	2se [†]	Ni/Mn	Fe/Mn	Cu/Mn	Zn/Mn	Co/Mn	Ce/Ce* ^{††}	Y _N /Ho _N
Bulk Fe-Mn crusts										
CAP BREST 03-03	Ascension FZ	1.40	0.03	0.010	1.87	0.0061	0.0036	0.023	2.03	0.78
DR01-003 4-B-5	VEMA FZ	1.48	0.02	0.017	1.70	0.0080	0.0039	0.026	2.32	0.64
DR-01-005 16-A-12	VEMA FZ	1.37	0.02	0.015	1.51	0.0061	0.0036	0.038	2.48	0.69
DR01-005 16-A-6	VEMA FZ	1.51	0.04	0.017	1.63	0.0071	0.0040	0.022	2.28	0.81
DR01-005 16-B-11	VEMA FZ	1.45	0.02	0.015	1.52	0.0051	0.0036	0.028	2.51	0.69
DR01-005 16-B-12	VEMA FZ	1.33	0.02	0.014	1.38	0.0047	0.0036	0.036	2.86	0.74
DR01-005 6-C-13	VEMA FZ	1.33	0.02	0.015	1.27	0.0052	0.0037	0.039	3.24	0.82
DR03-003 17-C-4	VEMA FZ	1.27	0.02	0.014	1.67	0.0048	0.0042	0.025	2.37	0.73
DR03-025 15#5	VEMA FZ	1.10	0.02	0.013	1.64	0.0055	0.0040	0.022	2.33	0.72
DR03-025 15-D-10	VEMA FZ	1.48	0.02	0.015	1.78	0.0075	0.0044	0.025	2.41	0.70
DR12-24 32-A-6	Gloria FZ	1.27	0.02	0.015	1.50	0.0068	0.0044	0.021	3.32	0.75
DR12-24 32-B-2	Gloria FZ	0.76	0.02	0.014	1.35	0.0056	0.0038	0.030	3.43	0.74
DR12-24 32-B-3	Gloria FZ	0.80	0.02	0.014	1.49	0.0067	0.0043	0.028	3.46	0.79
DR24-09 34-A-2	Azores TJ	1.73	0.02	0.023	0.39	0.0023	0.0023	0.047	3.28	0.79
DR24-09 34-C-2	Azores TJ	1.73	0.02	0.022	0.40	0.0030	0.0024	0.054	3.22	0.93
J2-480-R11	Apuupuu	1.48	0.03	0.023	0.90	0.0069	0.0035	0.015	3.49	1.21
J2-480-R11b	Apuupuu	1.76	0.03	0.012	0.73	0.0028	0.0025	0.029	3.48	0.77
J2-480-R13A	Apuupuu	1.72	0.03	0.018	0.81	0.0080	0.0027	0.022	1.99	0.66
J2-480-R13A	Apuupuu	1.80	0.03	0.015	0.69	0.0085	0.0025	0.021	1.93	0.57
J2-480-R13Ab	Apuupuu	1.76	0.03	0.009	0.67	0.0084	0.0025	0.024	2.28	0.63
J2-480-R13B	Apuupuu	1.63	0.04	0.021	1.18	0.0012	0.0036	0.028	1.91	0.71
J2-480-R14	Apuupuu	1.60	0.03	0.016	0.65	0.0066	0.0029	0.017	3.23	0.96
J2-480	Apuupuu	1.59	0.03	0.013	0.67	0.0067	0.0031	0.016	3.88	1.02
Average Apuupuu Fe-Mn crusts		1.67	-	0.016	0.79	0.0061	0.0029	0.021	2.77	0.82
Bauer Basin Fe-Mn deposits										
Y73-3-22D crust	Bauer Basin	0.33	0.02	0.035	0.98	0.020	0.0047	0.005	0.67	1.10
Y73-2-22D Mn NOD-1	Bauer Basin	0.42	0.03	0.052	0.44	0.019	0.0046	0.004	0.80	0.80
Y73-3-22D MnNOD-4	Bauer Basin	0.43	0.02	0.030	0.20	0.015	0.0060	0.002	0.55	0.90
Average Atlantic Ocean^{†††}		-	-	0.018	1.44	0.0059	0.0042	0.025	-	-
Average CCZ nodules^{††††}		-	-	0.046	0.22	0.0377	0.0048	0.007	-	-
Average North Pacific Prime Zone^{††††}		-	-	0.018	0.74	0.0043	0.0029	0.029	-	-
Lau basin hydrothermal deposits										
NL-10-09	Lau Basin	0.51	0.02	0.0001	0.0019	0.0001	0.0006	0.00002	0.25	1.56
NL-20-05	Lau Basin	1.11	0.02	0.0003	0.0013	0.0012	0.0009	0.00009	0.06	1.42
Loihi hydrothermal deposits										
J2-244-SS3	Ula Nui	nd	nd	0.005	16.68	0.0035	0.0028	0.003	0.99	1.13
J2-244-SS5	Ula Nui	-1.49	0.03	0.006	10.75	0.0011	nd	0.002	1.05	1.19
J2-309-SS2A	Ula Nui 2	-0.80	0.02	0.005	9.15	0.0039	0.0022	0.002	0.91	1.19
J2-309-SS2B	Ula Nui 2	-0.81	0.02	0.008	13.37	0.0019	0.0019	0.003	0.91	1.21
J2-309-SS2C	Ula Nui 2	-0.93	0.02	0.005	7.81	0.0019	0.0011	0.003	0.95	1.19
J2-313-SS1-top	Ula Nui 2	-1.16	0.03	0.002	3.61	0.0013	0.0005	0.001	0.95	1.25
J2-313-SS2	Ula Nui 2	-1.07	0.02	0.002	6.73	0.0005	0.0004	0.002	nd	nd
J2-477-SS black-green	Ula Nui	-1.16	0.03	0.003	2.84	0.0003	0.0006	0.002	nd	nd
Bulk manganese nodules USGS										
NOD-A-1	Atlantic	1.03	0.02	0.026	0.57	0.0063	0.0035	0.019	nd	nd
NOD-A-1 ^{†††}	Atlantic	1.06	0.03	0.027	0.58	0.0063	0.0037	0.019	nd	nd
NOD-P-1	Pacific	0.36	0.03	0.037	0.19	0.0415	0.0055	0.008	nd	nd
NOD-P-1 ^{†††}	Pacific	0.37	0.03	0.038	0.19	0.0407	0.0055	0.008	nd	nd

[†]2se corresponds to a two-standard error of the mean calculated on the 50 measurement cycles through MC-ICP-MS and corrected using double-spike.

^{††}Ce/Ce* = Ce_N/((Pr_N²/Nd_N), where subscript N stands for normalized values to PAAS.

^{†††}Digestion duplicate.

^{††††}Data from Hein et al. (2013).

nd: data not determined.