Persistence of deeply sourced iron in the Pacific Ocean

Tristan J. Horner^{a,1}, Helen M. Williams^{b,c,1}, James R. Hein^d, Makoto A. Saito^a, Kevin W. Burton^{b,c}, Alex N. Halliday^c, and Sune G. Nielsen^a

December 2, 2014

^aWoods Hole Oceanographic Institution, MA 02543, USA; ^bDurham University, Durham DH1 3LE, UK; ^cUniversity of Oxford,
 Oxford OX1 3AN, UK; ^dUnited States Geological Survey, Santa Cruz, CA 95060, USA

⁶ ¹To whom correspondence should be addressed. E-mail: Tristan.Horner@whoi.edu; h.m.williams2@durham.ac.uk

7 Classification PHYSICAL SCIENCES: Earth, Atmospheric, and Planetary Sciences

8 Keywords marine chemistry; micronutrient cycling; iron biogeochemistry; isotopic fingerprinting; ferromanganese oxides

9 Author contributions H.M.W., J.R.H., K.W.B., A.N.H., and S.G.N. designed research; H.M.W. and M.A.S. performed

¹⁰ research; T.J.H. and S.G.N. analyzed data; and T.J.H. wrote the paper.

¹¹ The authors declare no conflict of interest.

2

3

Abstract

Biological carbon fixation is limited by the supply of Fe in vast regions of the global ocean. Dissolved 13 Fe in seawater is primarily sourced from continental mineral dust, submarine hydrothermalism, and 14 sediment dissolution along continental margins. However, the relative contributions of these three 15 sources to the Fe budget of the open ocean remains contentious. By exploiting the Fe stable isotopic 16 fingerprints of these sources, it is possible to trace distinct Fe pools through marine environments, 17 and through time using sedimentary records. We present a reconstruction of deep-sea Fe-isotopic 18 compositions from a Pacific Fe-Mn crust spanning the past 76 Myr. We find that there have been 19 large and systematic changes in the Fe-isotopic composition of seawater over the Cenozoic that 20 reflect the influence of several, distinct Fe sources to the central Pacific Ocean. Given that deeply 21 sourced Fe from hydrothermalism and marginal sediment dissolution exhibit the largest Fe-isotopic 22 variations in modern oceanic settings, the record requires that these deep Fe sources have exerted 23 a major control over the total Fe inventory of the Pacific for the past 76 Myr. The persistence of 24 deeply sourced Fe in the Pacific Ocean illustrates that multiple sources contribute to the total Fe 25 budget of the ocean and highlights the importance of oceanic circulation in determining if deeply 26 sourced Fe is ever ventilated at the surface. 27

Significance

The vertical supply of dissolved Fe (iron) is insufficient compared to the physiological needs of marine phytoplankton in vast swathes of the open ocean. However, the relative importance of the main sources of 'new' Fe to the ocean – continental mineral dust, hydrothermal exhalations, and sediment dissolution – and their temporal evolution are poorly constrained. By analyzing the isotopic composition of Fe in marine sediments, we find that much of the dissolved Fe in the central Pacific Ocean originated from hydrothermal and sedimentary sources thousands of meters below the sea surface. As such, these data underscore the vital role of the oceans' physical mixing in determining if any deeply sourced Fe ever reaches the Fe-starved surface-dwelling biota.

28

29 Introduction

 Γ RON (Fe) is the most abundant transition metal in marine phytoplankton, reflecting its importance for 30 \mathbf{I} a range of biochemical processes such as photosynthesis and nitrogen fixation.¹ The high cellular 31 requirements for Fe, coupled with its low solubility and concentrations in seawater, render Fe a limit-32 ing nutrient in vast regions of the global ocean.² In turn, this makes the availability of dissolved Fe a 33 potential controlling factor for changes in atmospheric pCO_2 and thereby major oscillations in Earth's 34 climate. Global biogeochemical models show that more regions of the surface ocean are dominated by 35 circulation-driven dissolved Fe fluxes from below than by surface aerosol fluxes (e.g^{3,4}). This upward 36 flux of dissolved Fe is itself primarily sourced from three main pathways: dissolution of mineral dust 37 (e.g.⁵), submarine hydrothermalism (e.g.^{6–8}), and from sediment dissolution along continental margins 38 (e.g.^{9,10}), with the main removal mechanism being scavenging onto sinking particles (e.g.¹¹). However, 39 the significance of deeply-derived Fe sources – submarine sediment dissolution and hydrothermalism 40 – compared with surface Fe sources (dust dissolution), remains controversial (e.g^{12,13}). Given the key 41 role of Fe in supporting oceanic primary production, quantifying the relative importance of the various 42 Fe sources – both in the modern ocean and in the geological record – is critical to understanding how 43 micronutrient cycles are related to Earth's climatic state. 44

One promising way to trace Fe sources in the modern ocean is with measurements of stable Fe-isotopic compositions, where $\delta^{56/54}$ Fe = $({}^{56/54}$ Fe_{sample}/ ${}^{56/54}$ Fe_{IRMM-14} - 1) × 1,000. Recent studies showed that the Fe-isotopic composition of seawater is primarily controlled by the relative input of isotopically distinct Fe sources, ${}^{14-16}$ and that these source signatures can be transported and retained over thousands of kilometers within the ocean interior. 14 The large range in Fe-isotopic compositions observed between different Fe sources ($\geq 4 \%$) and in seawater (> 2 $\%^{14-18}$) should therefore also be reflected in sedimentary archives that faithfully capture the Fe-isotopic composition of seawater.

Here, we report a record of $\delta^{56/54}$ Fe from CD29-2, a mineralogically-uniform¹⁹ Fe-Mn (ferromanganese) 52 crust collected from the flank of the Karin Ridge at $16^{\circ}42.4'$ N, $168^{\circ}14.2'$ W in the central Pacific (r^{20} 53 Fig. 1). The present water depth of CD29-2 is pprox 2,000 m, though the depth at the time when Fe-Mn 54 crust formation commenced was likely $\sim 1,000$ m (owing to thermal subsidence; see SI, Supporting In-55 formation). Hydrogenetic Fe-Mn crusts are irregularly layered sedimentary deposits that form through 56 chemical precipitation of Fe- and Mn-oxides from ambient seawater, forming a mineral termed feroxy-57 hyte.²¹ Their persistence on rocky substrates away from sediment sources that might bury the crust²¹ 58 allows other metals to adsorb and become incorporated into Fe-Mn crusts via lattice replacement or 59 co-precipitation with Fe- or Mn-oxides.²² Detailed elemental stratigraphy showed that CD29-2 is hy-60 drogenetic – rather than hydrothermal or diagenetic – in origin.¹⁹ This designation means that the Fe 61 and other metals contained within CD29-2 were sourced from ambient seawater at the time of deposi-62 tion, rather than diagenetic remobilization of sedimentary metals, or through accretion of hydrothermal 63 vent-derived Fe- and Mn-oxides. 64

The extremely slow growth rate of most hydrogenetic deposits $(1 - 10 \text{ mm Myr}^{-1};^{23})$ renders Fe-Mn crusts as ideal recorders of long term changes in seawater trace element-isotopic chemistry. CD29-2 has an average growth rate of $\approx 1.4 \text{ mm Myr}^{-1};^{23}$ with each discrete sample for $\delta^{56/54}$ Fe (between 0.2 - 0.5 mm) integrating between 140 - 350 kyr of Earth history. Since the residence time of dissolved Fe in the deep ocean ($\approx 270 \text{ years}^{24}$) is less than the mixing time of the oceans ($\approx 1,000 \text{ years}^{25}$), our record provides a 'local' history of the central Pacific, rather than of global seawater $\delta^{56/54}$ Fe. Postdepositional processes such as diffusional re-equilibration with seawater²⁶ or precipitation of calcium fluorapatite in Fe-Mn crust pore spaces²⁷ have not affected the Fe-isotopic record in CD29-2 (see SI). Therefore, the **Figure 1: Map of sample locations.** Sample CD29-2 was recovered from the flanks of the Karin Ridge at $\sim 2,000$ m depth. CD29-2 is a semi-continuous hydrogenetic depositional record of many trace element isotopic compositions – including Fe – spanning the past ≈ 76 Ma (Fig. 3). The locations of other samples referred to throughout the text and in Fig. 4 are also shown: 28DSR9, a hydrogenetic Fe-Mn crust with a detailed Fe-isotopic stratigraphy for the past ≈ 10 Ma;²⁸ and DSDP (Deep Sea Drilling Project) Site 576²⁹ and LL44-GPC3,³⁰ two continuous Cenozoic records of aeolian deposition. Map drafted in Ocean Data View.³¹

Figure 2: Calculation of the Fe-isotopic offset between modern Fe-Mn crust surfaces and ambient seawater. (*A*) Fe-isotopic topology between Fe-Mn crusts (diamonds³³) and the three nearest seawater stations (squares¹⁶). The thickness of the connecting line denotes the proximity ranking, with the thickest line linking each crust with its nearest corresponding seawater station, and so on. (*Fe-Mn crust 1966.069 was excluded from the offset calculation as there are no proximal seawater data, whereas seawater data from station TAG were excluded owing to their significant hydrothermal influence.) Values of $\Delta^{56/54}$ Fe_{FeMn-SW} were calculated by comparing the Fe-isotopic composition of the surface scraping of N. Atlantic Fe-Mn crusts with the Fe-isotopic composition of the corresponding density surface (σ_{θ}) at each seawater station, for each of the three picks. The uncertainty on each pick corresponds to the propagated 2 SD measurement uncertainty on Fe-Mn crust and seawater $\delta^{56/54}$ Fe. Calculated $\Delta^{56/54}$ Fe_{FeMn-SW} compared against: (*B*) distance to nearest seawater station, (*C*) depth to corresponding density surface in the water column, and (*D*) ambient dissolved [Fe]. No relationship between $\Delta^{56/54}$ Fe_{FeMn-SW} and distance, depth, or [Fe] is evident from the data. The depths of the nine Fe-Mn crusts used in the offset calculation are shown in panel (*C*) as diamonds. The mean, unweighted fractionation factor is calculated as $\Delta^{56/54}$ Fe_{FeMn-SW} through time, it is possible to reconstruct the Fe-isotopic history of seawater from Fe-Mn crusts.

Fe-isotopic range of CD29-2 ($\delta^{56/54}$ Fe = -1.12 to +1.54 ‰, with mean and median values of -0.02

and -0.04 ‰, respectively), must reflect primary depositional signatures inherited from Fe dissolved in

⁷⁶ Estimating the Fe-isotopic fractionation factor, its driving mechanism, and variability through time

A robust reconstruction of the Fe-isotopic history of seawater from Fe-Mn crusts requires that the frac-77 tionation factor between Fe-Mn crusts and seawater, $\Delta^{56/54}$ Fe_{FeMn-SW}, is accurately known, is unaffected 78 by ambient environmental conditions, and has remained relatively constant through time. Stable iso-79 topic offsets between Fe-Mn crusts and seawater are common for many elements, and likely result from 80 differences in the relative binding strength between chemical species dissolved in seawater and incor-81 porated in Fe-Mn crusts (e.g.³²). We calculated the fractionation factor, defined as $\Delta^{56/54}$ Fe_{FeMn-SW} = 82 $\delta^{56/54}$ Fe_{FeMn} – $\delta^{56/54}$ Fe_{SW}, by comparing $\delta^{56/54}$ Fe of the surface scrapings of nine N. Atlantic Fe-Mn 83 crusts³³ with nearby seawater $\delta^{56/54}$ Fe measurements from the recent US GEOTRACES North Atlantic 84 GA03 Zonal Transect.¹⁶ Each crust was compared with linearly interpolated seawater $\delta^{56/54}$ Fe at the 85 corresponding density surface for the three nearest seawater profiles (topology shown in Fig. 2a). The 86 uncertainty on each estimate of $\Delta^{56/54}$ Fe_{FeMn-SW} refers to the propagated 2 SD external uncertainty as 87 reported in the respective original publications. The mean, unweighted fractionation factor was calcu-88 lated as $\Delta^{56/54}$ Fe_{FeMn-SW} = $-0.77 \pm 0.06 \%$ (2 SE, n = 27), and shows no obvious dependence on crust-89 seawater distance, sample depth, or ambient dissolved [Fe] (Figs. 2b, c, and d, respectively). We chose 90 to report the uncertainty about the mean value as two standard errors owing to the remarkable coher-91 ence and unidirectional nature of calculated Fe-Mn crust-seawater offsets, as well as the large number 92 of independent estimates of $\Delta^{56/54}$ Fe_{FeMn-SW} (Table 1). 93

⁹⁴ The comparison of modern Fe-Mn crust growth surfaces and nearby ambient seawater indicates that Fe

⁷⁵ seawater.

binding of Fe in seawater than in Fe-Mn crusts at equilibrium (e.g.³⁴). Given the importance of sidero-96 phore-like strong Fe-binding ligands in stabilizing dissolved Fe in seawater,^{35,36} it is extremely likely 97 that organic ligands play an important – if not dominant – role in setting $\Delta^{56/54}$ Fe_{FeMn-SW}. Several 98 studies have documented that isotopically heavy Fe will preferentially associate with organic ligands 99 during equilibration between aqueous Fe(III) and Fe–ligand complexes.^{37–39} The binding strength of 100 the Fe–ligand complex can modulate the magnitude of Fe-isotopic fractionation, with stronger ligands 101 – and thus stronger bonding environments – favoring larger equilibrium Fe-isotopic fractionation fac-102 tors. (Analogous behavior has also been identified for Cu,⁴⁰ which likely explains both the direction and 103 magnitude of Cu-isotopic fractionation between Fe-Mn crusts and seawater.⁴¹) The calculated value of 104 $\Delta^{56/54}$ Fe_{FeMn-SW} of -0.77 ± 0.06 ‰ is essentially identical to the empirically-determined $\Delta^{56/54}$ Fe_{Fe(III)-Fe(sid)} 105 between inorganic dissolved Fe(III) and Fe–siderophore complexes of -0.60 ± 0.15 ‰.³⁷ The remarkable 106 agreement between Fe-isotopic fractionation factors determined by experiments³⁷ and those observed 107 between naturally occurring Fe-Mn crusts and seawater (Fig. 2) suggests that organic ligands may exert 108

a dominant control on $\Delta^{56/54}$ Fe_{FeMn-SW}.

Interpretation of Fe-Mn crust-derived records of seawater $\delta^{56/54}$ Fe rely on $\Delta^{56/54}$ Fe_{FeMn-SW} having been 110 constant through time. If ligands are indeed exerting a significant influence on Fe-isotopic fractiona-111 tion in seawater, it is important to understand how evolutionary changes in the dominant Fe-binding 112 ligands may have also affected $\Delta^{56/54}$ Fe_{FeMn-SW}. To address this issue, we examined the evolutionary 113 history of a component from each of two common siderophore biosynthetic pathways, as siderophores 114 are thought to contribute to the oceanic Fe ligand inventory:⁴² enterobactin synthase subunit F (EntF) 115 and desferrioxamine E biosynthesis protein DesA. Whilst these are unlikely to be the only ligands in 116 seawater, these ligands – and in particular, DFO (desferrioxamine) – are good analogues to other marine 117 Fe ligands for several reasons: (1) The DFO class of ligands has been shown to exist in seawater; 43 (2) 118 DFO possesses similar conditional Fe binding constants to natural marine Fe ligands;⁴⁴ and (3) The Fe-119 isotopic fractionation factor between dissolved Fe(III) and Fe–DFO complexes of $\Delta^{56/54}$ Fe_{Fe(III)-Fe(DFO)} of 120 -0.60 ± 0.15 ‰³⁷ is identical to $\Delta^{56/54}$ Fe_{FeMn-SW}, within uncertainty. Analysis of sequence alignments of 121 the genes encoding these proteins in extant microbes demonstrates that siderophore biosynthesis genes 122 diverged from a common ancestor well before the 76 Myr timespan of interest in this study (see SI). 123 Given this finding, we contend that Fe-binding ligands have been present in seawater over the past 124 76 Myr, and likely far longer. Since the mineralogy of CD29-2 is invariant over this time period,¹⁹ it fol-125 lows that the differences in binding strength – and therefore the equilibrium $\Delta^{56/54}$ Fe_{FeMn-SW} – between 126 ligand-stabilized Fe in seawater and Fe bound in CD29-2 has also remained constant for at least 76 Myr. 127 Together with the observations that there are no resolvable Fe-isotopic effects related to Fe transport 128 distance, water depth, or ambient [Fe] on $\Delta^{56/54}$ Fe_{FeMn-SW} in the modern ocean (Fig. 2), the use of a 129 temporally-constant $\Delta^{56/54}$ Fe_{FeMn-SW} of -0.77 ± 0.06 ‰ for the past 76 Myr is justified by all available 130

¹³¹ oceanographic, experimental, and genomic data.

¹³² Controls on the Fe-isotopic composition of seawater

It is worthwhile to briefly review what is currently known about the Fe-isotopic systematics of the major Fe sources to the modern ocean, as this information is used as the interpretive framework for understanding the seawater record contained within CD29-2. The Fe-isotopic composition of seawater is thought to be primarily controlled by the relative input of local, isotopically distinct Fe sources (Fig. 3a), modulated by secondary modification processes (Fig. 3a), and mixing by oceanic circulation.^{14–16} The persistence of primary Fe-isotopic signatures along distinct water masses spanning thousands of kilometers suggests that the oceans' internal cycling of Fe through biological uptake and exchange with sinking Figure 3: The Fe-isotopic history of central Pacific seawater over the past 76 Myr. (A) Fe-isotopic compositions of the major oceanic Fe fluxes. The bold lines represent the end-member compositions of each flux: continental crust, hydrothermal fluids, and non-reductive sediment dissolution. (A second end-member for reductive sediment dissolution ~ -3 ‰ is not shown; ref.⁹) Numerous processes have been shown to modify end-member Fe-isotopic compositions; the dashed lines illustrate the observed range of Fe-isotopic compositions for each flux resulting from secondary modification processes (see text for references and discussion). (B) An Fe-isotopic history of central Pacific seawater recovered from CD29-2 spanning the past 76 Ma. The solid line links the measurements in relative chronological order; the break in the solid line between 37 and 42 Ma signifies a probable hiatus in the crust growth.²³ The surface measurement (≈ 0 Ma) is from Levasseur *et al.*³³ The gray and black error bars represent the analytical and propagated analytical and calculated uncertainties in $\Delta^{56/54}$ Fe_{FeMn-SW}, respectively. The light-colored shading indicates the boundaries between relevant geological Epochs; colors as per.⁴⁵ Within each Epoch, measurements of $\delta^{56/54}$ Fe have been binned, with the darker shading corresponding to one standard deviation either side of the mean value for that epoch. The shaded region labeled 'Modern' corresponds to the mean and standard deviation of the surfaces of globally distributed Fe-Mn crusts,³³ but has been expanded to cover the Quaternary for the sake of clarity.³³ (Pliocene and Micocene averages also include Fe-Mn crust data from Chu et al.;²⁸ see Fig. 4.) The greyed-out scale to the right of the figure shows the measured Fe-isotopic ratios for CD29-2 that have not been corrected for $\Delta^{56/54}$ Fe_{FeMn-SW} \approx -0.77 ‰.

particles exert only minimal influences on dissolved $\delta^{56/54}$ Fe.^{14,16} As such, the Fe-isotopic composition 140

of a water mass appears to be primarily governed by the Fe-isotopic composition of the dominant Fe 141

source to that water mass, in addition to any source Fe-isotopic modification processes at the time of 142

Fe addition. Iron-isotopic measurements can therefore be used to help elucidate the ultimate sources of 143 Fe to the ocean, and in particular the deep open ocean, where the dominant sources of Fe are still hotly

144

debated (e.g.^{7,46-48}). 145

The end-member Fe-isotopic composition of the three major Fe sources to the open ocean – mineral 146 aerosol or 'dust', seafloor sediment dissolution, and hydrothermalism – are summarized in Fig. 3a (bold 147 lines). The major surface Fe source, dust, is characterized by $\delta^{56/54}$ Fe $\approx +0.1 \pm 0.2 \%$,⁴⁹ identical to the 148 average Fe-isotopic composition of crustal rocks ($\delta^{56/54}$ Fe $\approx +0.1 \pm 0.1 \%^{50}$). Deeply sourced Fe from 149 dissolution of shelf sediments and hydrothermalism have distinct and variable Fe-isotopic compositions. 150 Reductive dissolution of marginal sediments delivers isotopically light Fe to seawater, with $\delta^{56/54}$ Fe \sim 151 -3.0 ‰,⁹ whereas non-reductive dissolution transfers Fe with a continental crust-like composition of 152 $\delta^{56/54}$ Fe $\approx +0.2 \pm 0.2 \%$.¹⁰ End-member hydrothermal fluid $\delta^{56/54}$ Fe has been measured $\approx -0.2 \%$.^{51,52} 153 with a small but significant fraction of this Fe escaping precipitation and becoming stabilized in seawater 154 as Fe(III).^{7,47} 155

For each of these three major oceanic Fe fluxes, secondary modification processes have been shown 156 to affect the Fe-isotopic composition of ligand-stabilized Fe in seawater (dashed lines in Fig. 3a). The 157 Fe-isotopic composition of dust-derived Fe in seawater appears to be isotopically heavier than crustal 158 rocks by $\approx +0.6$ ‰ at $\delta^{56/54}$ Fe $\approx +0.7 \pm 0.1$ ‰(Fig. 3a;¹⁶). In the absence of ligands, total digests and 159 leaching experiments on aerosol particulates have shown that Fe leached from dust possesses $\delta^{56/54}$ Fe \approx 160 $+0.1 \pm 0.2$ ‰.⁴⁹ The +0.6 ‰ offset between Fe bound in dust particles and dust-derived dissolved Fe 161 in seawater is thought to result from the equilibrium isotopic partitioning of isotopically heavy Fe into 162 strongly bound ligand-stabilized dissolved Fe(III) during dust dissolution.¹⁶ This interpretation is con-163 sistent with experimental studies of Fe-isotopic fractionation during mineral dissolution⁵³ and during 164 dissolved Fe(III)-ligand Fe-isotopic partitioning experiments.^{38,39} Since Fe-binding organic ligands have 165 an ancient biological origin that pre-dates the base of CD29-2 (see SI), it is likely that the Fe-isotopic off-166 set between dust particles and ligand-stabilized dust-derived Fe in seawater ($\Delta^{56/54}$ Fe_{dust part.-dust diss.} \approx 167 $-0.6 \pm 0.2 \ \text{\%}^{16,49,50}$) has remained constant over the course of our 76 Myr record. It is worth noting that 168 $\Delta^{56/54}$ Fe_{dust part.-dust diss.} and $\Delta^{56/54}$ Fe_{FeMn-SW} are identical, within uncertainty, lending further support 160 to the notion that a common ligand-mediated mechanism controls both Fe-isotopic offsets. 170

For deep Fe sources – hydrothermalism and sediment dissolution – Fe-isotopic modification processes 171 have also been identified, though the mechanisms involved are different than for mineral aeorsol Fe-172 isotopic modification. The most important modification processes identified in deep settings is the pre-173 cipitation of dissolved Fe in either oxide or sulfide forms, depending on local seawater conditions. These 174 two Fe-precipitating pathways impart large and distinct Fe-isotopic fractionations of opposite signs, as 175 Fe-oxides generally favor precipitation of isotopically heavy Fe (i.e. $\Delta^{56/54}$ Fe_{oxide-dissolved} > 0⁵¹, and 176 Fe-sulfides exclusively favor incorporation of isotopically light Fe (i.e. $\Delta^{56/54}$ Fe_{sulfide-dissolved} < 0.⁵⁴⁻⁵⁶). 177 Residual, dissolved Fe will thus become isotopically heavier as a result of Fe-sulfide precipitation, and 178 isotopically lighter as a result of Fe-oxide precipitation. Field studies have shown that Fe-sulfide precip-179 itation in continental margin sediments^{10,55} and at hydrothermal vent sites^{52,57} can drive the Fe-isotopic 180 composition of residual Fe, and thereby deep water Fe fluxes, toward heavier $\delta^{56/54}$ Fe by over $+2 \$ ⁵⁸ 181 compared to the end-member compositions (Fig. 3a). Furthermore, formation of isotopically heavy Fe-182 oxide precipitates around hydrothermal vents has also been shown to drive the delivery of isotopically 183 light Fe to the deep ocean (e.g. 16,51). 184

Given the large range of Fe-isotopic variability between different Fe sources (Fig. 3a) and observed in the modern ocean,^{14–17} we should naturally expect that changes in the dominant sources of Fe to the ocean with time will be accompanied by large shifts in the Fe-isotopic composition of seawater. Shifts in seawater $\delta^{56/54}$ Fe with time will thus depend on the relative input of different Fe sources to the ocean *and* the extent of their modification prior to stabilization in seawater (Fig. 3).

¹⁹⁰ An Fe-isotopic history of central Pacific seawater

Examination of our record of $\delta^{56/54}$ Fe reveals large changes in the Fe-isotopic composition of central Pacific seawater over the past 76 Myr (Fig. 3b). Though much of the record lies outside of the field defined by source 'end-member' $\delta^{56/54}$ Fe (Fig. 3a), when the aforementioned source modification processes are taken into account, even the most extreme $\delta^{56/54}$ Fe values in the Oligocene fall within the Fe-isotopic range defined by modern Fe fluxes (Fig. 3). Overall, the Fe-isotopic record of seawater reveals significant temporal variability, which suggests that the dominant Fe sources to the ocean have also varied over time, and that multiple Fe sources contribute to the total Fe budget of the central Pacific Ocean.

The large intra-epoch variation seen in past seawater $\delta^{56/54}$ Fe necessitates that the dominant Fe sources to the Pacific have changed through time (Fig. 3). Assuming a fixed dust value of $\delta^{56/54}$ Fe $\approx +0.7 \pm$ 0.1 % throughout the past 76 Myr, it is clear that more than 75 % of the record is outside of the field defined by mineral aerosol (Fig. 4). Isotopic mixing considerations³² demand that Fe-isotopic values observed outside of this narrow range must originate from mixing with other Fe sources with different Fe-isotopic compositions (Fig. 3b).

Deep sources have been documented to possess $\delta^{56/54}$ Fe that is highly variable and distinct from dust 204 (Fig. 3a). Thus, the large range of Fe-isotopic compositions observed over the last 76 Myr require the ad-205 dition of a quantitatively significant deeply sourced Fe pool to the central Pacific Ocean, such as sediment 206 dissolution or hydrothermalism (Fig. 4). Applying a constant dust-derived $\delta^{56/54}$ Fe $\approx +0.68 \pm 0.07 \%$,¹⁶ 207 we note 12 distinct events where seawater Fe-isotopic compositions cross through the dust value (Fig. 4). 208 To change seawater $\delta^{56/54}$ Fe in the past from < +0.61 to > 0.75 ‰ (and vice versa) requires the input of 209 an isotopically-distinct deep Fe source term to the central Pacific, as addition of more dust will simply 210 drive the record towards +0.7 %. That these 'events' are not restricted to any particular epoch (though 211 notably absent from the Pliocene onward), suggests deeply sourced Fe has been a significant and persis-212 tent component of the total Fe inventory of the Pacific Ocean throughout the past 76 Myr (Fig. 4). 213

Figure 4: Persistence of deeply sourced Fe in the Pacific Ocean. (*A*) Latitude of CD29-2 based on the migration of nearby DSDP locations.⁶⁰ (*B*) The Fe-isotopic history of central Pacific seawater recovered from Fe-Mn crust CD29-2 (squares). The high-resolution Fe-isotopic stratigraphy from 28DSR9²⁸ spanning the past ~ 10 Ma is also shown, renormalized to IRMM-14 and corrected for $\Delta^{56/54}$ Fe_{FeMn-SW} = $-0.77 \pm 0.06 \%$ (triangles; 28DSR9 location shown in Fig. 1). Vertical arrows indicate the 12 instances where the Fe-isotopic composition of seawater transits the Fe-isotopic composition of dust (horizontal bar; ref.¹⁶) (*C*) Reconstructed Pacific Plate seafloor generation rate.⁶¹ Both the spreading rate (mm yr⁻¹) and change in relative seafloor area (dA/dt, in km² 100 yr⁻¹) are shown, as the total amount of new crust generated depends on total ridge length *and* spreading rate. (*D*) Records of eolian deposition in the N. Pacific ocean from DSDP 576²⁹ and LL44-GPC3;³⁰ note the logarithmic scale. The interrelationships between these three records is discussed in detail in the SI.

²¹⁴ Understanding sustained changes in the Fe-isotopic composition of central Pacific seawater

The large intra-epoch variation in Fe-isotopic compositions recorded by CD29-2 necessitates a persistent 215 influence of deep Fe sources to the total Fe inventory of Pacific seawater over the past 76 Myr. It is 216 further possible to interpret some of the sustained excursions in $\delta^{56/54}$ Fe (i.e. the inter-epoch variability) 217 by understanding the location history CD29-2 and how this relates to probable changes in the supply rate 218 of the major Fe sources to the ocean (Fig. 4). A detailed paleogeography of CD29-2 is discussed in Klemm 219 *et al.*⁵⁹ Briefly, CD29-2 was situated at \approx 6 °S at the time of its formation (76 Ma), crossed the equator 220 around the K–Pg boundary (\approx 66 Ma), and has gradually progressed to its present location at \approx 16 °N 221 (Fig. 4a). Thermal subsidence of the underlying oceanic lithosphere has likely increased the water depth 222 from \sim 1,000 to \sim 2,000 m over the past 76 Myr, with the most rapid changes in depth occurring soon 223 after CD29-2 began precipitating (see SI). With these considerations in mind, we discuss below the three 224 most prominent features of the long term record: the excursion to extremely heavy $\delta^{56/54}$ Fe during the 225 Oligocene, the absence of large intra-epoch shifts after the Pliocene, and a possible shift to isotopically 226 light values during the Upper Cretaceous (Fig. 4). The Oligocene data are discussed in detail in the SI, 227 but are briefly summarized here. 228

The Oligocene data are best explained by a large and persistent increase in the hydrothermal contribu-229 tions to the total Fe budget of water masses bathing CD29-2 during this epoch (Fig. 4). The extremely 230 heavy $\delta^{56/54}$ Fe of up to +2.3 % necessitates that there were significant Fe-isotopic source modification 231 processes that were able to deliver isotopically heavy Fe to the ocean without 'choking off' the Fe supply. 232 Modification of hydrothermally-sourced Fe by precipitation of isotopically light Fe-sulfides seems an ob-233 vious candidate for such a process (e.g., 52,57 SI). Hydrothermal vents can exude fluids with μ M to mM Fe 234 concentrations,⁶² and the precipitation of Fe-sulfides from hydrothermal fluids – even at high tempera-235 tures – can result in significant Fe-isotopic modification of Fe fluxes (e.g., 52 SI). Moreover, recent studies 236 have documented distal transport of hydrothermally-sourced Fe thousands of kilometers across the Pa-237 cific that furthermore resemble the distributions of hydrothermally-derived helium anomalies (e.g^{63,64}). 238 Assuming that the modern correspondence between seafloor generation rate and hydrothermal fluid 239 fluxes [REF?] was also valid in the past, it is tempting to speculate that this shift to heavy Fe-isotopic 240 compositions in the Oligocene was driven by the approximate doubling of the rate of seafloor genera-241 tion in the Pacific basin during this epoch (;⁶¹ Fig. 4c). Conversely, CD29-2 was likely situated ≈ 11 °N 242 during the Oligocene (Fig. 4a), which is now bathed by a distal 'jet' of hydrothermally-influenced deep 243 waters between 1,500–3,000 m (evidenced by mantle-derived helium anomalies;⁶⁵). Assuming that the 244 vent systems at 9–10 °N along the EPR (East Pacific Rise) remained active during the Oligocene, and that 245 the predominantly east-to-west geostrophic flow at these depths also persisted at this time,⁶⁶ it is con-246 ceivable that CD29-2 simply moved through a plume of hydrothermally influenced deep waters. Since 247 there are no other elemental indications of hydrothermal influence on CD29-2 at this – or any other – 248 time¹⁹ during its pprox 76 Myr growth history, the Fe-isotopic systematics of CD29-2 require an unprece-240

dented degree of Fe-isotopic source modification to waters bathing CD29-2 during this epoch. However, testing whether or not this isotopically heavy reflects a basin wide increase in hydrothermally-derivd Fe during the Oligocene, or simply a local phenomenon will require a greater spatial resolution of Fe-Mn crust-derived records of seawater $\delta^{56/54}$ Fe. Nonetheless, the remarkable degree of seawater Fe-isotopic variation across the Oligocene is encouraging, as it permits the testing and tracing of Fe-isotopic provinciality – and therefore Fe sources – in the ocean interior over geological time.

An important feature of the record is that the intra-epoch shifts in $\delta^{56/54}$ Fe to values above and below 256 the dust end-member appear to cease around the start of the Pliocene (arrows in Fig. 4). The large 257 Fe-isotopic shifts seen throughout the rest of the record must be related to non dust-derived deep Fe 258 sources such as hydrothermalism and marginal sediment dissolution. However, at some point during 259 the past 10–20 Myr, the average Fe-isotopic composition of central Pacific seawater recorded by CD29-260 2 and 28DSR9 became largely invariant at $\delta^{56/54}$ Fe $\approx +0.4 \pm 0.1$ ‰ (± 2 SD). This switch to relative 261 Fe-isotopic homogeneity over the past 10–20 Myr is consistent with a reduced importance of deeply 262 sourced Fe and is coincident with the sharp increase in eolian dust deposition in the central and north 263 Pacific (Fig. $4d_{\ell}^{29,30}$). Since dust-derived Fe is thought to possess $\delta^{56/54}$ Fe $\approx +0.7 \ \%,^{16}$ the Fe-isotopic 264 chemistry of CD29-2 is consistent with the interpretation that dust has provided a significant portion 265 of the central Pacific Fe inventory from pprox 10–20 Ma to the present day. However, the small difference 266 between seawater ($\approx +0.4$ %; inferred from Fe-Mn crusts) and the dust end-member ($\approx +0.7$ %) of 267 \approx 0.3 ‰ is indicative of an influence from a secondary, isotopically light Fe source such as reductive 268 sediment dissolution⁶⁷ or Fe-oxide-influenced hydrothermalism.⁵¹ 269

The sustained shift to light Fe-isotopic ratios in the Upper Cretaceous are consistent with an increased 270 influence of continental margin-sourced Fe, or hydrothermalism modified by Fe-oxide precipitation to 271 waters bathing CD29-2. Reductive and non-reductive dissolution of sediments along continental mar-272 gins contribute Fe to the ocean with light end-member Fe-isotopic compositions of $\delta^{56/54}$ Fe ~ -3 ‰ and 273 $\approx +0.2$ ‰, respectively (;^{9,10} Fig. 3a). Analogous to Fe-sulfide modification processes described above, 274 hydrothermal modification by Fe-oxide precipitation could also facilitate the release of isotopically light 275 Fe to seawater.^{16,51} Though it is currently not possible to distinguish between these two deep sources 276 using $\delta^{56/54}$ Fe, both of these probable sources possess substantially different Fe-isotopic fingerprints 277 compared with surface dust deposition, thus ruling out a major atmospheric Fe contribution to seawater 278 bathing CD29-2 during the Upper Cretaceous (Fig. 3). During the Upper Cretaceous, CD29-2 was likely 279 at water depths $\approx 1,000$ m (SI), and situated south of the equator at $\sim 6^{\circ}$ S (;⁵⁹ Fig. 4a). This is some-280 what above than most EPR-derived helium anomalies,⁶⁵ such that a shallower, marginal sedimentary 281 Fe source is more likely. In the modern ocean, significant quantities of reduced,⁶⁸ bioavailable,⁶⁹ and 282 isotopically light Fe^{9,67} are released under low oxygen conditions associated with the highly produc-283 tive western continental margins.⁷⁰ As such, the ≈ -0.3 ‰ shift to values ≈ 0 ‰ in Upper Cretaceous 284 seawater $\delta^{56/54}$ Fe recorded by CD29-2 are best explained by a greater importance of shelf sediment disso-285 lution to the Fe budget of waters bathing CD29-2, further illustrating the power of $\delta^{56/54}$ Fe to distinguish 286 between surface and deep Fe sources through geological time. 287

288 Conclusions and outlook

The Fe-isotopic data for CD29-2 illustrate a dynamic Fe cycle in the central Pacific Ocean over the past 76 Myr. Isotopic mixing considerations demand a persistent and significant influence from deeply sourced Fe to the waters bathing CD29-2 in the central Pacific over the past 76 Myr. Deeply sourced Fe

²⁹² may have even contributed the majority of the Fe during certain epochs, such as during the Oligocene,

²⁹³ underscoring the importance of the oceans' circulation in controlling the spatial extent of deeply sourced

²⁹⁴ Fe and its contribution to basin-scale Fe budgets. However, it is clear that more records of Fe-isotopic

²⁹⁵ compositions from other Fe-Mn crusts are required to test the provinciality of oceanic Fe sources in the

²⁹⁶ past; the long-term record from CD29-2 is merely the first step towards this goal.

Reconstructions of past oceanic Fe sources can reveal much about the oceans Fe cycle in the past, but it 297 is clear that there is still much to learn. For example, what is the maximum lateral extent that dissolved 298 $\delta^{56/54}$ Fe signatures can persist across the ocean? Are there locations in the modern open ocean where dis-299 solved $\delta^{56/54}$ Fe exceeds +2 ‰? Did the deeply sourced Fe of the Oligocene ever ventilate at the surface? 300 Is deeply sourced Fe an important contributor to the total Fe inventory in other ocean basins? Are the 301 changes in the supply ratio of different Fe sources to the ocean responding to major climatic changes, or 302 driving them? All of these questions can be tackled with a greater spatial coverage of dissolved $\delta^{56/54}$ Fe 303 in the modern ocean and by performing further paleoceanographic studies of past seawater $\delta^{56/54}$ Fe 304 in other ocean basins. Coupling these currently scant Fe-isotopic observations to models of global Fe 305 biogeochemistry will help to iron-out these issues, and will refine our understanding of the role that 306 different Fe sources play in modulating global climate. 307

308 Materials and Methods

The samples of CD29-2 analyzed in this study were previously collected for TI- and Os-isotopic investigations, 309 with discrete samples taken via microdrilling. The age model for the crust was determined by matching the 310 Re decay-corrected Os-isotopic ratios for each discrete sample with the known osmium isotopic evolution of 311 seawater. Sample aliquots were then purified for Fe-isotopic analysis using anion-exchange column chemistry 312 and converted to nitrate form before mass spectrometric analysis. Iron isotopic analyses were carried out on a Nu 313 Instruments Nu Plasma HR multiple-collector inductively coupled plasma mass spectrometer at the University 314 of Oxford. Corrections for instrumental mass bias and isobaric overlap of 54 Cr on 54 Fe were performed by 315 standard-sample bracketing and monitoring ⁵³Cr, respectively. Mass dependence, reproducibility, and accuracy 316 were evaluated by analysis of various internal and external reference standards and found to be in excellent 317 agreement with previously published values (where available). Further description of methods and samples is 318 available in SI Materials and Methods. 319

ACKNOWLEDGMENTS. We thank D.N. Schmidt, T. van de Flierdt, T.M. Conway, W.B. Homoky, P.E. Lerner, J.D. Owens, S. Severmann, and – in particular – J.N. Fitzsimmons for some lively discussions; K.E. Egan and F. Poitrasson for constructive comments on an earlier version of this manuscript; and the editor and several anonymous reviewers for helping us to craft a substantially improved paper. T.J.H. is supported by the Postdoctoral Scholar Program at the Woods Hole Oceanographic Institution, with funding provided by the Doherty Foundation. H.M.W. is funded by the Natural Environment Research Council (Advanced Fellowship NE/F014295/1) and the European Research Council (Starting Grant 306655 'HabitablePlanet').

327 Supporting Information

328 SI Materials and Methods

329 Samples and sampling

Ferromanganese crust F7-86-HW CD29-2, referred to throughout the text as 'CD29-2', was recovered 330 via dredging along the Karin Ridge, a submarine volcanic feature east of Johnston Island in the central 331 Pacific.²⁰ Sample CD29-2 was recovered from the ridge flank at $16^{\circ}42.4'$ N; $168^{\circ}14.2'$ W, $\approx 200 - 600$ m 332 from the ridge summit. The age of the base of the crust is $\approx 75.5 \text{ Ma}_{2}^{23}$ based on an extrapolation of 333 the average growth rate between the PETM (≈ 55.5 Ma) and the Paleocene–Late Cretaceous boundary 334 (66 Ma;⁴⁵). Alkalic basalts from the ridge itself exhibit 40 Ar/ 39 Ar ages between 81.4 – 85.6 Ma,⁷¹ which 335 are currently situated at 1,800 m depth. The flat top of the Karin Ridge is strong evidence that it was 336 previously exposed above sea level,⁷² implying that substantial thermal subsidence has occurred to the 337 ridge over the past 81.4 - 85.6 Myr, bathing CD29-2 in progressively deeper water masses. Assuming 338 that this 1,800 m of subsidence occurred in proportion to $t^{\frac{1}{2}}$ (e.g. ref.⁷³) over \approx 83.5 Myr, CD29-2 likely 339 formed in a water depth between 750 - 1, 150 m, and has subsided ~ 16 m Myr $^{-1}$ to its current depth 340 between 1970 – 2390 m. These water depth constraints, though necessarily rough, suggest that CD29-2 341 has been submerged below $\approx 1,000$ m for its entire growth history of ≈ 76 Myr.²³ 342

The samples analyzed in this study were previously collected for Tl- and Os-isotopic investigations,^{23,74} 343 where samples were drilled from slabs of CD29-2 at a spacing of 0.2, 0.5, or 1 mm via microdrilling. 344 The age of the crust was determined by the osmium isotopic stratigraphy, where Re decay-corrected 345 Os-isotopic ratios measured on discrete samples through the entire crust⁷⁴ were matched to the known 346 Os-isotopic evolution of Cenozoic seawater (see⁷⁴ and references therein). Because portions of the Ceno-347 zoic Os-isotopic seawater curve are largely invariant, the age model for CD29-2 was constructed by 348 identifying specific events characterized by large Os-isotopic excursions (such as the K–Pg and Eocene– 349 Oligocene boundaries) and interpolating crust growth rates in between these fixed points. Therefore, 350 absolute ages may be somewhat uncertain (up to several million years uncertainty) when furthest away 351 from these fix points. However, the relative chronology of the crust is very robust as samples were 352 drilled in sequence. The use of Os-isotopic stratigraphy to date Fe-Mn crusts is discussed in Klemm et 353 al.,⁷⁴ and the revised age model used for CD29-2 is discussed in Nielsen *et al.*²³ 354

355 Analytical procedures

Powder samples of CD29-2 were leached in 6 M HCl, centrifuged, and transferred from 2 mL poly-356 propylene centrifuge tubes into acid-cleaned Savillex vials and were evaporated to dryness and oxidized 357 several times with hydrogen peroxide and concentrated nitric acid (Analar grade and Teflon-distilled, 358 respectively) after which they were converted to chloride form via several reflux cycles with 6 M Teflon-359 distilled HCl. The samples were then purified for Fe using standard anion-exchange column chemistry,⁷⁵ 360 and converted to nitrate form using hydrogen peroxide and concentrated nitric acid. The combined 361 blank for sample dissolution and column chromatography was < 5 ng Fe; this is negligible relative to 362 the amounts of Fe extracted from the samples (typically over 300 μ g total Fe). 363

³⁶⁴ Iron isotopic analyses were carried out on the Nu Instruments Nu Plasma HR multiple-collector induc ³⁶⁵ tively coupled plasma mass spectrometer at the University of Oxford following established procedures.⁷⁶

- ³⁶⁶ Samples were analyzed in 0.01 M HNO₃ and mass bias was corrected by sample–standard bracketing.
- ³⁶⁷ Measurements included collection of ion currents on m/z: 57, 56, 54, and 53 (⁵³Cr), to allow for correction
- ³⁶⁸ of any interference of ⁵⁴Cr on ⁵⁴Fe, although no difference between Cr-corrected and uncorrected ratios
- ³⁶⁹ were observed; this is due to the purification of the samples prior to analysis and effective separation of
- 370 Cr from Fe.

In order to monitor Fe-isotopic data quality and, in particular, the absence of artifacts relating to (i) in-371 complete Fe separation during column processing or (ii) sample matrix, several tests were performed. 372 Firstly, aliquots of pure IRMM-14 Fe standard (by definition, $\delta^{56/54}$ Fe of IRMM-14 $\equiv 0$ ‰) were passed 373 through columns. Column-processed IRMM-14 yielded $\delta^{56/54}$ Fe = 0.00 ± 0.06 ‰ (2 SD, n = 3) with a 374 recovery of 100 ± 2 %, demonstrating that no Fe-isotopic fractionation took place during column pro-375 cessing and that Fe yields were quantitative, respectively. Secondly, sample matrix effects were eval-376 uated by performing IRMM-doped sample matrix tests. All Fe-free sample matrix fractions were re-377 tained during column chromatography, and those corresponding to CD29-2 33.2 mm and 34.2 mm (Ta-378 ble 2) were doped with IRMM-14 to match the original Fe : matrix ratio of the corresponding sample. 379 These IRMM-14-doped samples of CD29-2 were then re-processed through column chromatography 380 and analyzed for their Fe-isotopic compositions, yielding $\delta^{56/54}$ Fe = +0.01 ± 0.07 ‰ (2 SD, n = 3) and 381 $-0.03 \pm 0.09 \$ % (2 SD, n = 2) for CD29-2 33.2 mm and 34.2 mm, respectively. 382

Reproducibility and accuracy were also evaluated by analysis of several in-house and international Fe-

³⁸⁴ isotopic reference standards:

- An in-house FeCl powder, obtained from the ETH Zürich (ETH Fe salt standard); $\delta^{56/54}$ Fe = $-0.73 \pm 0.10 \%$ (2 SD, n = 48), in agreement with published values from Mikutta *et al.*⁷⁷ of $\delta^{56/54}$ Fe = $-0.72 \pm 0.13 \%$.
- USGS Pacific Fe-Mn nodule sample (Nod-P-1); $\delta^{56/54}$ Fe = $-0.39 \pm 0.07 \%$ (2 SD, n = 7), in agreement with published values from Dideriksen *et al*.:⁷⁸ $\delta^{56/54}$ Fe = $-0.42 \pm 0.09 \%$.
- USGS Icelandic Basalt sample (BIR-1); $\delta^{56/54}$ Fe = +0.04 ± 0.07 ‰ (2 SD, n = 4), in excellent agreement with published values from Weyer *et al.*:⁷⁹ $\delta^{56/54}$ Fe = +0.05 ± 0.04 ‰.

392 SI Discussion

393 Estimating Fe diffusivity in Fe-Mn crusts

Diffusional resetting or 'smoothing' of primary signals can affect certain trace metal isotopic records recovered from Fe-Mn crusts (e.g.^{26,80}). To assess if this was an issue for records of Fe-isotopic compositions recorded by CD29-2, we estimated the diffusivity of Fe in Fe-Mn crusts by assuming that the rate of chemical exchange between fluid (seawater) and solid (Fe-Mn crusts) is governed by the same processes for Fe as for U. From this, we applied the general Fe-Mn crust diffusivity equation from Henderson and Burton:²⁶

$$D_{\rm eff.}^{\rm Fe} = \frac{D_{\rm eff.}^{\rm U} K_{\rm c}^{\rm U}}{K_{\rm c}^{\rm Fe}} \tag{1}$$

Figure 5: The ancient biological origin of siderophore production. Percent identity of BLASTp results from (*A*) enterobactin synthase subunit F (EntF) and (*B*) desferrioxamine E biosynthesis protein DesA. All sequences have been ordered by e-value (low to high), showing percent identities for sequences with e-values <1E-5 and <2E-30 for EntF and DesA, respectively (mean DesA sequence length is 419 amino acids).

where $D_{\text{eff.}}^{\text{E}}$ is the effective diffusivity of element E in Fe-Mn crusts (cm² yr⁻¹) and K_{c}^{E} is the concentration 400 ratio of E in Fe-Mn crusts relative to its porewaters (assumed to be seawater,²⁶). Assuming a range of 401 [Fe] for deep Pacific seawater between 0.4 - 0.6 nM⁸¹ and [Fe] of CD29-2 crusts between 10 - 20 wt. %,¹⁹ 402 the Fe concentration ratio between CD29-2 : seawater is between $3 - 9 \times 10^9$. As such, we calculate 403 $D_{\text{eff.}}^{\text{Fe}}$ between 4×10^{-13} and 1×10^{-12} cm² yr⁻¹. Such a slow rate of Fe diffusion – at least half the rate of Th diffusivity in Fe-Mn crusts²⁶ – suggests that our Fe-isotopic dataset is a primary record and 404 405 has not been smoothed or reset by diffusion. Diagenetic precipitation of calcium fluorapatite (calcium 406 fluorophosphate) in the lower half of CD29-2 (e.g.^{19,27}) does not appear to affect $\delta^{56/54}$ Fe. This is likely 407 because secondary calcium fluorapatite contains little, if any, Fe compared to the average 15 wt. % Fe of 408 CD29-2, and the predominantly pore-filling nature of calcium fluorapatite precipitation that primarily 409 serves to reduce Fe-Mn crust porosity. 410

⁴¹¹ The age and origin of strong Fe-binding ligands in the ocean

While neither the composition of the strong ligands in ocean seawater nor microbial siderophore biosynthetic pathways specific to abundant marine microbes are known, the global prevalence of seawater Fe ligands and the similarity of their similar conditional binding strengths to those of siderophores strongly implies a contribution of siderophores to the oceanic Fe ligand inventory.⁴⁴ Moreover, field studies have identified ferrioxamine-type ligands in seawater enrichment studies,⁴³ demonstrating the presence of siderophore biosynthetic capacity in natural assemblages and bolstering the case for their contribution to the oceanic Fe ligand reservoir.

Two types of biosynthetic pathways for siderophores are currently known, non-ribosomal peptide syn-419 thesis (NRPS) and NRPS-independent siderophore synthesis (NIS).⁸² To examine if siderophore synthe-420 sis is older than the \approx 76 Myr time period of interest in this study, we examined the diversity of a gene 421 across numerous microbial genomes from each of these two siderophore biosynthetic pathways, focus-422 ing on the specific siderophores desferrioxamine E biosynthesis protein DesA and enterobactin synthase 423 subunit F (EntF). EntF is a component of a three protein NRPS complex,^{83,84} and DesA was recently im-424 plicated in NIS siderophore synthesis.⁸² Microbial genomes were analyzed using the Standard Protein 425 Basic Local Alignment Search Tool (BLASTp;⁸⁵), which is used to identify seed amino acid sequences (the 426 query) in other organisms by searching within a specified database. For each query, regions of similarity 427 (identity) and their alignment with other amino acid sequences are identified. Each potential match is 428 scored and assigned an e-value (expected value), which decreases exponentially towards 0 the higher 429 the statistical confidence that the match is not simply chance. Using BLASTp and the non-redundant 430 sequence database with seed sequences from siderophore-producing model bacteria (E. coli and Strepto-431 *myces sp.*), sequence search and alignments showed each gene to correspond to a broad range of sequence 432 identities across diverse range of microbes, extending to as low as ≈ 30 % identity while maintaining 433 low e-values consistent with a shared ancestral origin (Fig. 5; Tables 3 and 4). 434

Amino acid sequence alignments and identities have been used to attempt to calculate evolutionary distance and divergence times of major taxa throughout Earth history, using a calibration against the

vertebrate fossil record.^{86,87} Using a conservative approach here, a comparison of the range of sequence 437 identities from the two siderophore biosynthetic gene identities with the previous approaches to con-438 strain the age of divergence major groups of organisms using amino acid distance measurements, implies 439 that the last common ancestor for both genes is extremely old. For example, the evolutionary distances 440 that corresponds to a 91 % identity is estimated to have a last common ancestor (divergence) of 100 Ma, 441 whilst a 60 % identity corresponds to more than 1,000 Ma.⁸⁸ Both of these estimates are higher in iden-442 tity than the homologous sequences examined here, hence implying these biosynthetic genes diverged 443 from a last common ancestor far longer than the \approx 76 Myr timescale of this study (Tables 3 and 4). This 444 observation remains true even if with the caveat that the calculated ages for bacterial sequences may 445 be biased several-fold faster due to a distinct synonymous substitution rate in bacteria relative to the 446 calibrated vertebrate fossil record.⁸⁹ 447

As expected, sequences with higher percentage identities are found within the same species (e.g. E. coli 448 for EntF) and hence most likely share siderophore biosynthetic function. Yet at lower identities it can 449 be more challenging to constrain protein function due to limited experimental evidence and potential 450 recruitment from related enzyme pathways. In the case of the enterobactin EntF gene, Salmonella sp. 451 are known to also produce enterobactin using this pathway,⁹⁰ providing support for this siderophore 452 biosynthetic function. EntF of Salmonella sp. has a ≈ 80 % sequence identity, implying an age of over 453 300 Ma for a divergence from a shared genetic ancestor using the approach described above (Table 3). 454 Other sequences within the lower range of sequence identity are annotated as EntF and may have sim-455 ilar siderophore biosynthesis function. For DesA, annotations of homologous sequences correspond to 456 DesA or the enzyme L-2,4-diaminobutyrate aminotransferase. Recent biochemical and genomic charac-457 terization of this enzyme have demonstrated it to have the lysine carboxylase activity required as a key 458 step in NIS siderophore biosynthesis and to be broadly disseminated in bacteria,⁸² consistent with the 459 search results in Table 4. 460

⁴⁶¹ Together, these calculated evolutionary distances for two gene representatives of two siderophore biosyn-

thetic pathways show them both to be much older than the ≈ 76 Myr time span focused on in this study. Overall, these observations lend support to the notion that the siderophore biosynthetic pathways that may contribute seawater Fe ligand production are similarly old and that the resulting ligand-mediated Fe-isotopic offsets in seawater – $\Delta^{56/54}$ Fe_{dust part.-dust diss.} and $\Delta^{56/54}$ Fe_{FeMn-SW}) – have also been stable for ≈ 76 My, if not far longer.

⁴⁶⁷ Deeply sourced Fe persists regardless of how the dust value is assigned

The interpretation that deeply sourced Fe has been important in the Pacific Ocean for the past 76 Myr is 468 robust regardless of how the dust-derived Fe-isotopic end-member is assigned. If dust-derived $\delta^{56/54}$ Fe 469 was not $+0.7 \pm 0.1$ ‰ but possessed a different value, there is no unique Fe-isotopic composition of 470 dust-derived Fe that can account for the entire record without requiring mixing with another deep Fe 471 source that has a distinct Fe-isotopic composition; such a scenario is only possible if $\Delta^{56/54}$ Fe_{FeMn-SW} also 472 changed. However, temporally-constant seawater $\delta^{56/54}$ Fe with temporally variable $\Delta^{56/54}$ Fe_{FeMn-SW} is 473 essentially precluded by the following considerations. Firstly, for seawater $\delta^{56/54}$ Fe to have remained 474 constant, $\delta^{56/54}$ Fe heavier than $\approx +0.5$ ‰ recorded by CD29-2 would necessitate periodic reversals 475 in the sense of Fe-isotopic fractionation between Fe-Mn crusts and seawater. Given the mineralogical 476 homogeneity of CD29-2,¹⁹ we consider such a scenario highly implausible. Secondly, there is no evidence 477 to suggest that $\Delta^{56/54}$ Fe_{FeMn-SW} is variable in the modern ocean, despite the large range of depths, [Fe], 478 and $\delta^{56/54}$ Fe encountered in the offset calibration ($\sim 200-4,000$ m, $\approx 0.4-1$ nM, $\approx +0.2$ to +0.7 ‰, 479

Figure 6: Pb-isotopic array for CD29-2. Plot of 207 Pb/ 206 Pb vs. 208 Pb/ 206 Pb for CD29-2 from Christensen *et al.*⁹¹ using the age model of Nielsen *et al.*²³ Given that the Pb-isotopic data were not obtained on the same slab of CD29-2 as the Os-isotopic stratigraphy, the absolute ages for the Pb-isotopic data have considerable uncertainty. To account for this uncertainty, the Pb-isotopic data have not been re-plotted against age, but instead have been binned into their respective geological Epochs to illustrate broad temporal patterns. Pb-isotopic uncertainties are shown at the 2 SD level and are discussed in Christensen *et al.*⁹¹

respectively; Fig. 2; Table 1). Thirdly, current evidence suggests that Fe-isotopic offsets between Feoxides and ligand-stabilized dissolved Fe(III), and by analogy $\Delta^{56/54}$ Fe_{FeMn-SW}, is primarily governed by ligands,⁵³ which have an ancient biological origin and are thus likely present throughout the 76 Myr record sampled by CD29-2 (SI Discussion). Whilst it is not possible to definitively demonstrate that $\Delta^{56/54}$ Fe_{FeMn-SW} did not change in the past, all available evidence supports the use of a temporallyconstant $\Delta^{56/54}$ Fe_{FeMn-SW} of -0.77 ± 0.06 ‰ for the past 76 Myr.

486 What drove the excursion to extremely heavy Fe-isotopic compositions during the Oligocene?

The processes that may have affected the Fe-isotopic composition of central Pacific seawater over the 487 past 76 Myr can be broadly subdivided into two categories based on their overall mechanism of influ-488 encing seawater Fe-isotopic chemistry: indirect and direct influences. Indirect influences on seawater 489 Fe-isotopic chemistry are those processes that affect the cycling of Fe within the ocean interior, but do 490 not necessarily alter the marine Fe inventory. Direct influences are those processes that alter the oceanic 491 inventory of Fe and/or the Fe-isotopic modification of a source as it becomes stabilized in seawater. 492 There is considerable evidence to suggest that direct factors exert the primary control on modern oceanic 493 $\delta^{56/54}$ Fe, such as the large range of Fe-isotopic compositions measured between the principal marine Fe 494 sources (Fig. 3a) and the numerous documented Fe-isotopic source modification processes that can sup-495 ply isotopically fractionated Fe to the oceans (e.g.;^{10,52,55,57} Fig. 3a). We briefly outline how indirect 496 influences are unlikely to be a major control on seawater $\delta^{56/54}$ Fe, before discussing each of the major 497 features of the Fe-isotopic record in the text below. 498

The most important of the indirect influences on central Pacific seawater Fe-isotopic compositions are 499 likely to be changes in deep water circulation patterns over the past 76 Myr. Though changes in the depth 500 of carbonate compensation⁹² and changes in atmospheric pCO_2^{93} have occurred during the record, it is 501 not clear how these processes would affect seawater $\delta^{56/54}$ Fe. However, changes in deep water circula-502 tion would result in CD29-2 being bathed in different water masses throughout its growth, which would 503 cause $\delta^{56/54}$ Fe to vary so long as different water masses possessed distinct Fe-isotopic compositions. 504 Since little is known about Pacific deep water mass geometry throughout the Cenozoic, it is only possi-505 ble to infer changes in ambient water masses through comparison with other geochemical proxy records 506 recovered from CD29-2. The high-resolution Pb-isotopic stratigraphy of Christensen *et al.*⁹¹ would ap-507 pear to support this inference, as the Pb-isotopic data for CD29-2 indicate numerous changes in central 508 Pacific Pb sources over the Cenozoic (Fig. 6). However, the three main Pb-isotopic arrays⁹¹ do not cor-509 respond to major features of the Fe-isotopic record (Fig. 3b), and it is unclear if the Pb- and Fe-isotopic 510 records from CD29-2 are directly comparable given the differing geochemical behaviors of these two 511 elements in seawater (e.g.²⁵). Ultimately a water mass structure control on central Pacific $\delta^{56/54}$ Fe still 512 implies that distinct Fe sources (i.e. direct factors) exert the dominant control over seawater $\delta^{56/54}$ Fe, but 513 that indirect factors determine the Fe-isotopic depth structure of the water column. With this in mind, 514 we interpret the major shifts in the seawater Fe-isotopic record from CD29-2 in terms of changes in the 515 relative balance and/or modifications to the dominant Fe fluxes to the ocean through time. 516

The Fe-isotopic record indicates that central Pacific seawater was as heavy as $\delta^{56/54}$ Fe $\approx +2.3$ ‰ dur-517 ing the Oligocene, with a sustained shift to values $\approx +0.8$ ‰ heavier than the mean value of the other 518 epochs (Fig. 4). The Oligocene is notable for numerous environmental changes, including: a drop in 519 global sea-level and the emplacement of major ice sheets on Antarctica, a global increase in chemical 520 weathering rates, a deepening of the carbonate compensation depth, a drop in atmospheric pCO_2 , a sig-521 nificant increase in seafloor generation rate in the Pacific basin, and the opening of the Antarctic seaways 522 (e.g.^{61,91-94}). We outline below why an increased rate of seafloor generation and associated hydrothermal 523 fluid fluxes in the Oligocene is likely responsible for the observed shift in seawater $\delta^{56/54}$ Fe, although 524 we concede that it is difficult to entirely rule out other possible explanations for the Oligocene data. 525

In the modern ocean, isotopically-heavy Fe sources share an important characteristic: modification by 526 Fe-sulfide precipitation. A shift to heavier Fe-isotopic compositions is thus inconsistent with an in-527 creased Fe flux from reductive sediment dissolution ($\sim -3 \%$;⁹ Fig. 3a) or increased source modification 528 by Fe-oxide precipitation, as both of these factors would drive marine Fe-isotopic compositions toward 529 lighter values. The sustained shift to isotopically heavy $\delta^{56/54}$ Fe in the Oligocene must relate to in-530 creased source modification by Fe-sulfide precipitation (Fig. 4), as this is the only plausible modification 531 process that can generate heavy Fe-isotopic compositions. Fe-sulfide precipitation can occur in conti-532 nental margin^{10,55} and hydrothermal settings,⁵⁷ and has been documented to drive residual, dissolved 533 Fe towards heavy $\delta^{56/54}$ Fe in field settings. In sedimentary systems with oxic bottom waters, Fe-sulfide 534 precipitation (usually as pyrite⁵⁶) occurs well below the sediment–water interface in sulfidic sedimen-535 tary horizons.^{10,55} These sulfidic redox horizons are frequently characterized by low ambient [Fe], as 536 abundant free sulfide will favor the precipitation of Fe-sulfide minerals, thus 'choking off' the benthic 537 Fe supply. Overall, these considerations suggest that the Fe-sulfide-influenced Fe flux from continental 538 margins is unlikely to have a significant influence on open ocean $\delta^{56/54}$ Fe. 539

⁵⁴⁰ Precipitation of Fe-sulfide also occurs in deep-sea hydrothermal systems, which account for large fluxes ⁵⁴¹ of Fe to the deep ocean^{6,8,63,95,96} that can be stabilized by organic compounds.⁴⁸ End-member hydrother-⁵⁴² mal fluids appear to possess $\frac{5^{56}/5^4}{5^4}$ Fe ≈ -0.2 % (Fig. $32^{51}/5^2$) although ligand stabilized Fe that on

mal fluids appear to possess $\delta^{56/54}$ Fe $\approx -0.2 \%$ (Fig. 3a;^{51,52}), although ligand-stabilized Fe that es-542 capes the local hydrothermal source may be significantly modified toward heavier or lighter $\delta^{56/54}$ Fe, 543 depending on the proportion of Fe-oxide versus Fe-sulfide precipitation ($\Delta^{56/54}$ Fe_{oxide-dissolved} > 0; 544 $\Delta^{56/54}$ Fe_{sulfide-dissolved} < 0). The partitioning of Fe between oxides and sulfides depends primarily on 545 the Fe : H₂S ratio of the vent fluids and the kinetics of Fe(II) oxidation in ambient seawater.⁵¹ In the 546 high (> 20_{i}^{51}) Fe : H₂S ultramafic-hosted Rainbow hydrothermal system in the north Atlantic, Fe precip-547 itation is near-quantitative and occurs almost exclusively as Fe-oxides that exhibit Fe-isotopic composi-548 tions similar to end-member hydrothermal fluid.⁵¹ In other basalt-hosted Atlantic vent systems, where 549 Fe : H₂S is ~ 1 ,⁹⁷ significant Fe-isotopic fractionation has been observed, resulting in a net transfer of 550 isotopically heavy dissolved Fe to seawater.⁵⁷ The hydrothermal systems of the EPR (East Pacific Rise) 551 exhibit even lower Fe : H_2S ratios ≤ 0.1 ,⁹⁸ such that dissolved Fe escaping to seawater is expected to 552 exhibit extremely heavy Fe-isotopic compositions. Unpublished Fe-isotopic profiles of seawater from 553 the southeast Pacific Ocean appear to confirm this phenomenon,⁹⁹ though open ocean seawater values 554 in excess of $\delta^{56/54}$ Fe $\sim +1$ ‰ remain to be observed. 555

Indirect evidence for an important role for hydrothermal Fe-sulfide precipitation can be obtained via examination of the Fe-rich hydrothermal precipitates at active vent sites (i.e. the reaction products of the Fe source modification processes). Fe-isotopic analyses of pyrite-rich hydrothermal chimneys from the EPR exhibit $\delta^{56/54}$ Fe between ≈ -1.3 and -0.5 ‰, thus requiring the accompanying 350 °C hydrothermal fluids to possess $\delta^{56/54}$ Fe between +1.0 and +1.5 ‰ (at equilibrium;⁵²). Even heavier Fe-isotopic compositions are theoretically possible if Fe-isotopic fractionation follows a Rayleigh distillation inside

the hydrothermal stockwork. For example, in a hydrothermal system with ≈ 97 % precipitation of an 562 initial hydrothermal fluid characterized by $\delta^{56/54}$ Fe = -0.2 %,⁵¹ the residual dissolved $\delta^{56/54}$ Fe should 563 be in excess of +2 % assuming all precipitation as Fe-sulfides when $\Delta^{56/54}$ Fe_{sulfide-dissolved} = -0.6 $\%^{57}$. 564 At 99 % precipitation, residual $\delta^{56/54}$ Fe would reach $\approx +3$ %, suggesting that the cumulative Fe-isotopic 565 composition of precipitates formed from that system would exhibit $\delta^{56/54}$ Fe ≈ -0.2 ‰. Indeed, pyrite-566 rich hydrothermal precipitates at certain EPR vent sites have been shown to exhibit $\delta^{56/54}$ Fe ≈ -0.2 ‰ in 567 field settings.⁵² The Fe-isotopic values observed in the Oligocene strongly indicate that deeply sourced 568 hydrothermal Fe dominated the total Fe inventory of the central Pacific, and that Fe-sulfide precipitation 569 was the major source modification process at that time. 570

The Oligocene shift to heavy seawater $\delta^{56/54}$ Fe is coincident with a time of increased seafloor generation 571 rate (assumed to be a proxy for hydrothermal activity;⁶¹ Fig. 4c) in the Pacific basin and extremely low 572 rates of eolian deposition (Fig. 4d). The confluence of these two factors likely explains the sustained shift 573 to heavy values during this epoch. During the Oligocene, CD29-2 was situated at an average latitude 574 of 11 °N, which would place it at the periphery of the maximum modern primordial $\delta({}^{3}\text{He})$ anomaly 575 in the central Pacific at $\sim 2,000 \text{ m}.^{65}$ (Primordial $\delta(^{3}\text{He})$ anomalies are sourced through the degassing 576 of mantle-derived ³He at hydrothermal vents.¹⁰⁰) The extremely low dust fluxes of the Oligocene ($r^{29,30}$ 577 Fig. 4d) would have provided little Fe to buffer against this predominantly hydrothermally-sourced Fe 578 anomaly. Though hydrothermal fluxes are also relatively high in the modern Pacific ocean compared to 579 the rest of the Cenozoic,⁶¹ the $\delta^{56/54}$ Fe of seawater ambient to CD29-2 has been lighter than either the 580 Oligocene values or dust value since the Pliocene (Fig. 3b). The reduced influence of deeply sourced Fe 581 on modern central Pacific $\delta^{56/54}$ Fe may reflect the significant increase in eolian-sourced Fe since the mid-582 Pliocene, the transit of CD29-2 to more northerly latitudes away from the primordial δ ⁽³He) anomaly, or 583 a change in the chemistry of EPR vent fluids since the Oligocene. 584

Deeply-derived hydrothermal Fe is increasingly recognized as an important component of the total Fe inventory of the modern deep Pacific,^{6,63,64} as well as other ocean basins (e.g.^{8,101–103}). Our Fe-isotopic record from CD29-2 suggests that this has also been true in the past, and that deeply sourced hydrothermal Fe was likely the major Fe source to water masses bathing CD29-2 during the Oligocene. More broadly, these data underscore the fact that multiple Fe sources contribute to the deep oceans' Fe inventory, and illustrate that the relative importance of these sources – and their Fe-isotopic chemistry – have also varied significantly through time. 592 Data Tables

Cruise	Sample	Distance (km)	Depth to σ_{θ} (m)	Interp. [Fe] (nM)	Interp. $\delta^{56/54}$ Fe SW	± 2 SD	$\Delta^{56/54}Fe_{FeMn-SW}$	$\pm 2 \text{ SD}$
ALV539	D2-1A	657	2423	0.61	0.53	0.05	-0.77	0.06
		681	2249	0.66	0.59	0.07	-0.83	0.07
		702	2686	0.67	0.51	0.06	-0.75	0.07
SO-154	52 CD-1	1552	863	0.53	0.52	0.07	-0.71	0.08
		1685	897	0.68	0.24	0.05	-0.43	0.07
		1782	727	0.46	0.54	0.07	-0.73	0.08
Arcl TR079	D14	1484	2074	0.63	09.0	0.08	-0.78	0.10
		1671	2294	0.61	0.56	0.06	-0.74	0.0
		1674	1900	0.66	0.59	0.06	-0.77	0.09
Gosnold 65-75	2383.00	1144	376	0.64	0.21	0.05	-0.78	0.10
		1171	243	0.39	0.55	0.09	-1.12	0.13
		1190	190	0.69	0.30	0.06	-0.87	0.11
	BM1963.897	1189	646	0.74	0.27	0.04	-0.52	0.09
		1224	393	0.43	0.51	0.09	-0.75	0.12
		1226	006	0.66	0.19	0.06	-0.44	0.10
Hudson St. 54	BM1969.05	635	1891	0.71	0.36	0.05	-0.84	0.11
		675	1761	0.68	0.44	0.05	-0.92	0.11
		707	1643	0.81	0.18	0.05	-0.67	0.11
Discovery 144	D10979	1541	4000	0.46	0.74	0.11	-1.02	0.11
		1603	3500	0.47	0.70	0.07	-0.98	0.08
		1684	3500	0.59	0.61	0.06	-0.89	0.08
VEMA CH 78	DR01-001a	1490	3217	0.48	0.71	0.07	-0.86	0.09
		1524	3771	0.50	0.60	0.06	-0.75	0.08
		1733	3317	0.49	0.74	0.08	-0.90	0.10
Atlantis II 96	D10-7	438	2488	0.47	0.70	0.06	-0.75	0.08
		677	2460	0.97	0.39	0.04	-0.44	0.06
		895	1984	0.54	0.67	0.06	-0.72	0.07
					Mean valu	ie ±2 SE	-0.77	0.06

Table 1: Fe-Mn crust-seawater data pairs used to calculate $\Delta^{56/54}$ Fe_{FeMn-SW}. Fe-Mn crust data from Levasseur *et al.*,³³ seawater Fe-isotopic data from Conway & John.¹⁶ Linear interpolation was used to estimated [Fe] and $\delta^{56/54}$ Fe for seawater samples; uncertainties are as reported in the original publications.

Table 2: Fe-isotopic record from central Pacific Fe-Mn crust CD29-2. See text in Methods for age model description. ^{*a*}Depth from top in mm ^{*b*}Fractional depth in the crust ^{*c*}Model age from²³ ^{*d*}Surface value from³³

DfT ^a	$\delta^{56/54}$ Fe	$\pm 2\text{SD}$	f^b	Age ^c (Ma)	DfT ^a	$\delta^{56/54}$ Fe	$\pm 2\text{SD}$	f^b	Age ^c (Ma)
0.5	-0.41^{d}	0.09	0.0047	0.27	43.2	+0.01	0.08	0.4075	42.26
16.5	-0.25	0.09	0.1557	8.85	57.0	+0.71	0.06	0.5377	47.44
20.5	-0.07	0.04	0.1934	11.00	60.0	+0.27	0.10	0.5660	48.56
28.5	-0.25	0.09	0.2689	22.06	65.0	+0.11	0.10	0.6132	50.44
29.0	+0.39	0.08	0.2736	22.75	67.0	+0.40	0.02	0.6321	51.19
30.0	+1.21	0.03	0.2830	24.13	69.0	-0.02	0.02	0.6509	51.94
30.5	+0.28	0.13	0.2877	24.82	69.5	-0.05	0.03	0.6557	52.13
31.5	+0.08	0.10	0.2972	26.20	70.0	+0.15	0.04	0.6604	52.31
32.5	+1.54	0.08	0.3066	27.59	70.5	+0.06	0.05	0.6651	52.50
33.2	+0.79	0.07	0.3132	28.55	71.0	+0.13	0.04	0.6698	52.69
34.2	+1.31	0.09	0.3226	29.94	71.5	-0.38	0.06	0.6745	52.88
35.2	+1.32	0.02	0.3321	31.32	73.0	-0.90	0.06	0.6887	53.44
35.4	+0.62	0.06	0.3340	31.59	73.5	-0.13	0.02	0.6934	53.63
35.6	-0.69	0.05	0.3358	31.87	74.0	-1.12	0.10	0.6981	53.81
37.0	+0.06	0.05	0.3491	33.81	75.0	-0.17	0.08	0.7075	54.19
37.2	-0.01	0.10	0.3509	34.08	80.0	-0.03	0.07	0.7547	56.70
37.8	-0.57	0.10	0.3566	34.65	83.0	-0.06	0.05	0.7830	59.10
40.0	-0.11	0.01	0.3774	35.75	86.5	+0.02	0.11	0.8160	61.90
40.4	+0.21	0.10	0.3811	35.95	90.5	+0.35	0.07	0.8538	65.10
40.6	+0.03	0.10	0.3830	36.05	91.0	-0.25	0.09	0.8585	65.50
40.8	-0.04	0.19	0.3849	36.15	91.5	-0.83	0.12	0.8632	65.90
41.0	-0.11	0.02	0.3868	36.25	93.5	-0.75	0.14	0.8821	67.50
41.2	-0.07	0.02	0.3887	36.35	95.5	-0.69	0.06	0.9009	69.10
41.4	-0.02	0.05	0.3906	36.45	96.0	-0.28	0.06	0.9057	69.50
41.6	-0.67	0.08	0.3925	36.55	97.0	-0.78	0.12	0.9151	70.30
41.8	-0.08	0.05	0.3943	36.65	100.0	-0.60	0.10	0.9434	72.69
42.0	-0.47	0.10	0.3962	36.75	102.0	-0.43	0.02	0.9623	74.29
42.4	-0.03	0.03	0.4000	36.95	102.5	+0.17	0.05	0.9670	74.69
42.6	-0.03	0.09	0.4019	42.04	104.0	-0.30	0.09	0.9811	75.89

nologous	se being	ngs.
ny hon	of the	ne belo
). Mai	many	enzym
e (NR	with	h this
tabas	30 %,	o whic
nce da	′as ≈	roup t
sequer	as low	resis g
dant	ntities	e syntl
-redun	id ider	peptid
t non	ino ac	somal
agains	ith am	n-ribo
nit F	nces w	ger no
e subu	sequer	the lar
nthase	ng to	ted to
ctin sy	extendi	ig rela
erobac	ases, é	rs beir
of ent	datab	d othe
itput (luence	sis, an
Tp or	ent seq	synthe
BLAS	e curre	tin bio
from	ing th	erobac
esults	n exist	to ent
arch r	withi	elated
ted se	stected	oeing r
Selec	are dé	ed as b
able 3:	roteins	nnotati
H	ā	ð

E-value	% identity	Taxon	BLASTp Annotation	Taxon Group
0	100.00	Escherichia coli; Escherichia coli K-12	enterobactin synthase multienzyme complex component	enterobacteria
0	98.53	Escherichia coli; Escherichia coli ATCC 8739	enterobactin synthase subunit F	enterobacteria
0	81.98	Shigella flexneri 1235-66	enterobactin synthase component F	enterobacteria
0	81.67	Citrobacter rodentium; Citrobacter rodentium ICC168	enterobactin synthetase component F	enterobacteria
0	79.21	Salmonella enterica; Oranienburg str. 0250	enterobactin synthase subunit F	enterobacteria
0	78.13	Klebsiella pneumoniae;Klebsiella pneumoniae UHKPC06	enterobactin synthetase component F	enterobacteria
0	78.13	Hafnia alvei BIDMC 31	enterobactin synthase component F	enterobacteria
0	76.89	Kosakonia radicincitans; Enterobacter radicincitans DSM 16656	enterobactin synthase subunit F	enterobacteria
0	58.37	gamma proteobacterium WG36	enterobactin synthase subunit F	g-proteobacteria
0	58.31	Pseudomonas sp. 313	enterobactin synthase subunit F	g-proteobacteria
0	49.57	Methylocystis rosea	hypothetical protein	a-proteobacteria
0	44.26	Marinomonas sp. D104	chromophore Iyase	g-proteobacteria
0	42.74	Marinomonas mediterranea MMB-1	amino acid adenylation protein	g-proteobacteria
0	40.39	Vibrio campbellii	peptide synthetase	g-proteobacteria
0	37.75	Stenotrophomonas maltophilia 5BA-I-2	enterobactin synthase	g-proteobacteria
0	36.96	Aeromonas veronii; Aeromonas veronii AER39	enterobactin synthase	g-proteobacteria
2.E-173	34.75	Bacillus amyloliquefaciens;Bacillus amyloliquefaciens DSM 7	siderophore 2	firmicutes
8.E-139	33.83	Actinokineospora sp. EG49	Siderophore biosynthesis non-ribosomal peptide synthetase module	high GC Gram+
2.E-127	32.41	Methylobacter marinus	hypothetical protein	g-proteobacteria
6.E-164	31.13	Cyanothece sp. PCC 7424	amino acid adenylation protein	cyanobacteria
1.E-88	29.18	Rhodococcus opacus;Rhodococcus opacus M213	non-ribosomal peptide synthetase	high GC Gram+

R). Many	ugh most	
oase (NF	%, althoi	
ice datal	as ≈ 30	
t sequen	as low a	
edundan	dentities	
t non-re	no acid i	
A agains	with ami	
ein Des	duences v	
esis prot	ng to sec	
biosynthe	extendi	
nine E ł	atabases	
ferrioxa	quence d	
t of des	rrent sec	oteins.
p outpu	g the cu	ietical pr
BLAST	n existin	r hypoth
lts from	ed withi	oxylase o
ch resul	re detect	to carbo
ted sear	oteins aı	e related
4: Selec	ogous pr	itions are
Table	homol	annota

BLASIP
desferrio
pyridoxa
pyridoxa
sideroph
pyridoxa
4-diamin
pyridoxal
pyridoxal
hypothetic
pyridoxal-
tyrosine de
hypothetica
Desferrioxa
hypothetica
hypothetica
cytochrome
4-diaminob
Diaminobu
L-2-diamin

593 **References**

- [1] Twining BS, Baines SB (2013) The Trace Metal Composition of Marine Phytoplankton. Ann. Rev.
 Mar. Sci. 5:191–215.
- [2] Boyd PW, et al. (2007) Mesoscale iron enrichment experiments 1993-2005: Synthesis and future directions. *Science* 315:612–617.
- ⁵⁹⁸ [3] Aumont O, Maier-Reimer E, Blain S, Monfray P (2003) An ecosystem model of the global ocean ⁵⁹⁹ including Fe, Si, P colimitations. *Global Biogeochem. Cycles* 17.
- [4] Misumi K, et al. (2014) The iron budget in ocean surface waters in the 20th and 21st centuries:
 projections by the Community Earth System Model version 1. *Biogeosci.* 11:33–55.
- [5] Jickells TD, et al. (2005) Global iron connections between desert dust, ocean biogeochemistry, and
 climate. *Science* 308:67.
- [6] Tagliabue A, et al. (2010) Hydrothermal contribution to the oceanic dissolved iron inventory. *Nat. Geosci.* 3:252–256.
- [7] Sander SG, Koschinsky A (2011) Metal flux from hydrothermal vents increased by organic complexation. *Nat. Geosci.* 4:145–150.
- [8] Saito MA, et al. (2013) Slow-spreading submarine ridges in the South Atlantic as a significant oceanic iron source. *Nat. Geosci.* 6:775–779.
- [9] Severmann S, McManus J, Berelson WM, Hammond DE (2010) The continental shelf benthic iron flux and its isotope composition. *Geochim. Cosmochim. Acta* 74:3984–4004.
- [10] Homoky WB, John SG, Conway TM, Mills RA (2013) Distinct iron isotopic signatures and supply
 from marine sediment dissolution. *Nat. Comm.* 4.
- [11] Wu J, Boyle EA, Sunda W, Wen LS (2001) Soluble and colloidal iron in the oligotrophic North
 Atlantic and North Pacific. *Science* 293:847–849.
- [12] Fung IY, et al. (2000) Iron supply and demand in the upper ocean. *Global Biogeochem. Cycles* 14:281–
 295.
- [13] Moore JK, Doney SC, Glover DM, Fung IY (2001) Iron cycling and nutrient-limitation patterns in
 surface waters of the World Ocean. *Deep Sea Res. Part II* 49:463–507.
- [14] Radic A, Lacan F, Murray JW (2011) Iron isotopes in the seawater of the equatorial Pacific Ocean:
 New constraints for the oceanic iron cycle. *Earth Planet. Sci. Lett.* 306:1–10.
- [15] John SG, Adkins J (2012) The vertical distribution of iron stable isotopes in the North Atlantic near Bermuda. *Global Biogeochem. Cycles* 26.
- [16] Conway TM, John SG (2014) Quantification of dissolved iron sources to the North Atlantic Ocean.
 Nature 511:212–215.
- [17] Lacan F, et al. (2008) Measurement of the isotopic composition of dissolved iron in the open ocean.
 Geophys. Res. Lett. 35.

- [18] Owens JD, et al. (2012) Iron isotope and trace metal records of iron cycling in the proto-North
 Atlantic during the Cenomanian-Turonian oceanic anoxic event (OAE-2). *Paleoceanography* 27.
- [19] Frank M, O'Nions RK, Hein JR, Banakar VK (1999) 60 Myr records of major elements and Pb–Nd
 isotopes from hydrogenous ferromanganese crusts: Reconstruction of seawater paleochemistry.
 Geochim. Cosmochim. Acta 63:1689–1708.
- [20] Hein JR, *et al.* (1987) *Farnella Cruise* F7-86-HW, Cobalt-rich Ferromanganese Crust Data Report for
 Karin Ridge and Johnston Island, Central Pacific., (US Geol. Surv.), Open File Rep. 87-663.
- [21] Hein J, et al. (2000) in *Handbook of marine mineral deposits*, ed Cronan DS (CRC Press, Boca Raton,
 FL), pp 239–279.
- [22] Koschinsky A, Hein JR (2003) Uptake of elements from seawater by ferromanganese crusts: solid phase associations and seawater speciation. *Mar. Geol.* 198:331–351.
- [23] Nielsen SG, et al. (2009) Thallium isotope evidence for a permanent increase in marine organic
 carbon export in the early Eocene. *Earth Planet. Sci. Lett.* 278:297–307.
- [24] Bergquist BA, Wu J, Boyle EA (2007) Variability in oceanic dissolved iron is dominated by the
 colloidal fraction. *Geochim. Cosmochim. Acta* 71:2960–2974.
- [25] Broecker WS, Peng TH (1982) *Tracers in the Sea* (Lamont-Doherty Geological Observatory, Columbia University), p 690.
- [26] Henderson GM, Burton KW (1999) Using (²³⁴U/²³⁸U) to assess diffusion rates of isotope tracers in ferromanganese crusts. *Earth Planet. Sci. Lett.* 170:169–179.
- [27] Hein JR, et al. (1993) Two major Cenozoic episodes of phosphogenesis recorded in equatorial
 Pacific seamount deposits. *Paleoceanography* 8:293–311.
- [28] Chu NC, et al. (2006) Evidence for hydrothermal venting in Fe isotope compositions of the deep
 Pacific Ocean through time. *Earth Planet Sci. Lett.* 245:202–217.
- [29] Janecek TR (1985) in *Deep Sea Drilling Project Leg 86, Western North Pacific*, Init. Repts. DSDP, ed
 Turner KL (U.S. Govt. Printing Office, Washington, D.C.) No. 19, pp 589–603.
- [30] Janecek TR, Rea DK (1983) Eolian deposition in the northeast Pacific Ocean: Cenozoic history of
 atmospheric circulation. *Geol. Soc. Am. Bull.* 94:730–738.
- [31] Schlitzer R (2011) Ocean Data View. (http://odv.awi.de/).
- [32] Johnson CM, Beard BL, Albarède F (2004) *Geochemistry of non-traditional stable isotopes*, Rev. Min eral. Geochem. (Mineralogical Society of America, Washington, DC) Vol. 55.
- [33] Levasseur S, Frank M, Hein JR, Halliday AN (2004) The global variation in the iron isotope composition of marine hydrogenetic ferromanganese deposits: implications for seawater chemistry?
 Earth Planet. Sci. Lett. 224:91–105.
- [34] Johnson CM, Beard BL, Albarede F (2004) in *Geochemistry of non-traditional stable isotopes*, Rev.
 Mineral. Geochem., eds Johnson CM, Beard BL, Albarede F (Mineralogical Society of America)
 Vol. 55, pp 1–24.

- [35] van den Berg CMG (1995) Evidence for organic complexation of iron in seawater. *Mar. Chem.* 50:139–157.
- [36] Kraemer SM (2004) Iron oxide dissolution and solubility in the presence of siderophores. *Aquat. Sci.* 66:3–18.
- [37] Dideriksen K, Baker J, Stipp SLS (2008) Fe isotope fractionation between inorganic aqueous Fe (III)
 and a Fe siderophore complex. *Min. Mag.* 72:313–316.
- [38] Dideriksen K, Baker JA, Stipp SLS (2008) Equilibrium Fe isotope fractionation between inorganic
 aqueous Fe (III) and the siderophore complex, Fe (III)-desferrioxamine B. *Earth Planet. Sci. Lett.* 269:280–290.
- [39] Morgan JLL, Wasylenki LE, Nuester J, Anbar AD (2010) Fe Isotope Fractionation during Equilibration of Fe-Organic Complexes. *Environ. Sci. Technol.* 44:6095–6101.
- [40] Ryan BM, Kirby JK, Degryse F, Scheiderich K, McLaughlin MJ (2014) Copper Isotope Fractionation
 during Equilibration with Natural and Synthetic Ligands. *Environ. Sci. Technol.* 48:8620–8626.
- [41] Little SH, Sherman DM, Vance D, Hein JR (2014) Molecular controls on Cu and Zn isotopic fractionation in Fe–Mn crusts. *Earth Planet. Sci. Lett.* 396:213–222.
- [42] Barbeau K, Rue E, Bruland K, Butler A (2001) Photochemical cycling of iron in the surface ocean
 mediated by microbial iron (III)-binding ligands. *Nature* 413:409–413.
- [43] Mawji E, et al. (2008) Hydroxamate siderophores: occurrence and importance in the Atlantic
 Ocean. *Environ. Sci. Technol.* 42:8675–8680.
- [44] Rue EL, Bruland KW (1995) Complexation of iron (III) by natural organic ligands in the Central North Pacific as determined by a new competitive ligand equilibration/adsorptive cathodic stripping voltammetric method. *Mar. Chem.* 50:117–138.
- [45] Cohen KM, Finney SC, Gibbard PL, Fan JX (2013) The ICS International Chronostratigraphic Chart.
 Episodes 36:199–204.
- [46] Boyle EA (1997) What controls dissolved iron concentrations in the world ocean?—A comment.
 Mar. Chem. 57:163–167.
- [47] Bennett SA, et al. (2008) The distribution and stabilisation of dissolved Fe in deep-sea hydrothermal plumes. *Earth Planet. Sci. Lett.* 270:157–167.
- [48] Toner BM, et al. (2009) Preservation of iron(II) by carbon-rich matrices in a hydrothermal plume.
 Nat. Geosci. 2:197–201.
- [49] Waeles M, Baker AR, Jickells T, Hoogewerff J (2007) Global dust teleconnections: aerosol iron solubility and stable isotope composition. *Environ. Chem.* 4:233–237.
- [50] Beard BL, et al. (2003) Application of Fe isotopes to tracing the geochemical and biological cycling
 of Fe. *Chem. Geol.* 195:87–117.
- [51] Severmann S, et al. (2004) The effect of plume processes on the Fe isotope composition of hy drothermally derived Fe in the deep ocean as inferred from the Rainbow vent site, Mid-Atlantic
 Ridge, 36°14'N. *Earth Planet. Sci. Lett.* 225:63–76.

- [52] Rouxel O, Shanks III WC, Bach W, Edwards KJ (2008) Integrated Fe-and S-isotope study of seafloor
 hydrothermal vents at East Pacific Rise 9–10 N. *Chem. Geol.* 252:214–227.
- [53] Wiederhold JG, et al. (2006) Iron isotope fractionation during proton-promoted, ligand-controlled,
 and reductive dissolution of goethite. *Environ. Sci. Technol.* 40:3787–3793.
- ⁷⁰⁵ [54] Butler IB, Archer C, Vance D, Oldroyd A, Rickard D (2005) Fe isotope fractionation on FeS forma-⁷⁰⁶ tion in ambient aqueous solution. *Earth Planet. Sci. Lett.* 236:430–442.
- [55] Severmann S, Johnson CM, Beard BL, McManus J (2006) The effect of early diagenesis on the
 Fe isotope compositions of porewaters and authigenic minerals in continental margin sediments.
 Geochim. Cosmochim. Acta 70:2006–2022.
- [56] Guilbaud R, Butler IB, Ellam RM (2011) Abiotic pyrite formation produces a large Fe isotope
 fractionation. *Science* 332:1548–1551.
- [57] Bennett SA, et al. (2009) Iron isotope fractionation in a buoyant hydrothermal plume, 5 S Mid Atlantic Ridge. *Geochim. Acta* 73:5619–5634.
- [58] Aquilina A, et al. (2013) Diagenetic Mobilisation of Fe and Mn in Hydrothermal Sediments. *Min. Mag.* 77:604.
- [59] Klemm V, Reynolds B, Frank M, Pettke T, Halliday AN (2007) Cenozoic changes in atmospheric
 lead recorded in central Pacific ferromanganese crusts. *Earth Planet. Sci. Lett.* 253:57–66.
- [60] Van Andel TH, Heath GR, Moore TC (1975) Cenozoic History and Paleoceanography of the Central
 Equatorial Pacific Ocean A Regional Synthesis of Deep Sea Drilling Project Data. *Geol. Soc. Am. Mem.* 143:1–223.
- [61] Cogné JP, Humler E (2006) Trends and rhythms in global seafloor generation rate. *Geochem. Geophys. Geosyst.* 7:Q03011.
- [62] Von Damm KL, et al. (1985) Chemistry of submarine hydrothermal solutions at 21 °N, East Pacific
 Rise. *Geochim. Cosmochim. Acta* 49:2197–2220.
- [63] Wu J, Wells ML, Rember R (2011) Dissolved iron anomaly in the deep tropical–subtropical Pacific:
 Evidence for long-range transport of hydrothermal iron. *Geochim. Acta* 75:460–468.
- [64] Fitzsimmons JN, Boyle EA, Jenkins WJ (2014) Distal transport of dissolved hydrothermal iron in
 the deep South Pacific Ocean. *Proc. Natl. Acad. Sci.* p 201418778.
- [65] Lupton J (1998) Hydrothermal helium plumes in the Pacific Ocean. J. Geophys. Res. 103:15853–
 15868.
- [66] Reid JL (1997) On the total geostrophic circulation of the Pacific Ocean: Flow patterns, tracers, and
 transports. *Prog. Oceanogr.* 39:263–352.
- [67] John SG, Mendez J, Moffett J, Adkins J (2012) The flux of iron and iron isotopes from San Pedro
 Basin sediments. *Geochim. Acta* 93:14–29.
- [68] Lam PJ, Bishop JK (2008) The continental margin is a key source of iron to the HNLC North Pacific
 Ocean. *Geophys. Res. Lett.* 35.

- [69] Bruland KW, Rue EL, Smith GJ, DiTullio GR (2005) Iron, macronutrients and diatom blooms in the
 Peru upwelling regime: brown and blue waters of Peru. *Mar. Chem.* 93:81–103.
- [70] Stramma L, Schmidtko S, Levin LA, Johnson GC (2010) Ocean oxygen minima expansions and their biological impacts. *Deep Sea Res. Part I* 57:587–595.
- [71] Davis AS, Gray LB, Clague DA, Hein JR (2002) The Line Islands revisited: New ⁴⁰Ar/³⁹Ar
 geochronologic evidence for episodes of volcanism due to lithospheric extension. *Geochem. Geophys. Geosyst.* 3:1–28.
- [72] Hein JR, *et al.* (1990) Geological, geochemical, geophysical, and oceanographic data and interpre tations of seamounts and Co-rich ferromanganese crusts from the Marshall Islands, KORDI-USGS
 RV *Farnella* cruise F10-89-CP., (US Geol. Surv.), Open File Rep. 90-407.
- [73] Parsons B, Sclater JG (1977) An analysis of the variation of ocean floor bathymetry and heat flow with age. *J. Geophys. Res.* 82:803–827.
- [74] [74] Klemm V, Levasseur S, Frank M, Hein JR, Halliday AN (2005) Osmium isotope stratigraphy of a marine ferromanganese crust. *Earth Planet. Sci. Lett.* 238:42–48.
- [75] [75] Williams HM, et al. (2005) Systematic iron isotope variations in mantle rocks and minerals: the
 effects of partial melting and oxygen fugacity. *Earth Planet. Sci. Lett.* 235:435–452.
- [753] [76] Williams HM, et al. (2009) Fractionation of oxygen and iron isotopes by partial melting processes:
 implications for the interpretation of stable isotope signatures in mafic rocks. *Earth Planet. Sci. Lett.* 283:156–166.
- [77] Mikutta C, et al. (2009) Iron isotope fractionation and atom exchange during sorption of ferrous
 iron to mineral surfaces. *Geochim. Cosmochim. Acta* 73:1795–1812.
- [78] Dideriksen K, Baker JA, Stipp SLS (2006) Iron isotopes in natural carbonate minerals determined
 by MC-ICP-MS with a ⁵⁸Fe-⁵⁴Fe double spike. *Geochim. Cosmochim. Acta* 70:118–132.
- [79] Weyer S, et al. (2005) Iron isotope fractionation during planetary differentiation. *Earth Planet. Sci. Lett.* 240:251–264.
- [80] David K, Frank M, O'Nions RK, Belshaw NS, Arden JW (2001) The Hf isotope composition of
 global seawater and the evolution of Hf isotopes in the deep Pacific Ocean from Fe–Mn crusts.
 Chem. Geol. 178:23–42.
- [81] Johnson KS, Gordon RM, Coale KH (1997) What controls dissolved iron concentrations in the
 world ocean? *Mar. Chem.* 57:137–161.
- [82] Burrell M, Hanfrey CC, Kinch LN, Elliott KA, Michael AJ (2012) Evolution of a novel lysine decar boxylase in siderophore biosynthesis. *Mol. Microbiol.* 86:485–499.
- [83] Ehmann DE, Shaw-Reid CA, Losey HC, Walsh CT (2000) The EntF and EntE adenylation domains
 of *Escherichia coli* enterobactin synthetase: Sequestration and selectivity in acyl-AMP transfers to
 thiolation domain cosubstrates. *Proc. Natl. Acad. Sci.* 97:2509–2514.
- [84] Yamanaka K, et al. (2005) Desferrioxamine E produced by *Streptomyces griseus* stimulates growth
 and development of *Streptomyces tanashiensis*. *Microbiol*. 151:2899–2905.

- [85] Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ (1990) Basic local alignment search tool. J.
 Mol. Biol. 215:403–410.
- [86] Feng DF, Doolittle RF (1997) Converting amino acid alignment scores into measures of evolutionary time: a simulation study of various relationships. *J. Mol. Evo.* 44:361–370.
- [87] Grishin NV (1995) Estimation of the number of amino acid substitutions per site when the substitution rate varies among sites. *J. Mol. Evo.* 41:675–679.
- [88] Feng DF, Cho G, Doolittle RF (1997) Determining divergence times with a protein clock: Update
 and reevaluation. *Proc. Natl. Acad. Sci.* 94:13028–13033.
- [89] Ochman H, Elwyn S, Moran NA (1999) "Calibrating bacterial evolution". Proc. Natl. Acad. Sci.
 96:12638–12643.
- [90] Benjamin WH, Turnbough CL, Posey BS, Briles DE (1985) The ability of *Salmonella typhimurium* to produce the siderophore enterobactin is not a virulence factor in mouse typhoid. *Infect. Immun.* 50:392–397.
- [91] Christensen JN, Halliday AN, Godfrey LV, Hein JR, Rea DK (1997) Climate and ocean dynamics
 and the lead isotopic records in Pacific ferromanganese crusts. *Science* 277:913.
- [92] Pälike H, *et al.* (2012) A Cenozoic record of the equatorial Pacific carbonate compensation depth.
 Nature 488:609–614.
- [93] Zachos J, Pagani M, Sloan L, Thomas E, Billups K (2001) Trends, rhythms, and aberrations in global
 climate 65 Ma to present. *Science* 292:686–693.
- [94] Pagani M, et al. (2011) The role of carbon dioxide during the onset of Antarctic glaciation. *Science* 334:1261–1264.
- [95] Boyle EA, Bergquist BA, Kayser RA, Mahowald N (2005) Iron, manganese, and lead at Hawaii
 Ocean Time-series station ALOHA: Temporal variability and an intermediate water hydrothermal
 plume. *Geochim. Cosmochim. Acta* 69:933–952.
- [96] Tagliabue A, Aumont O, Bopp L (2014) The impact of different external sources of iron on the
 global carbon cycle. *Geophys. Res. Lett.* 41:920–926.
- [97] Douville E, et al. (2002) The rainbow vent fluids (36 14? n, mar): the influence of ultramafic rocks
 and phase separation on trace metal content in mid-atlantic ridge hydrothermal fluids. *Chem. Geol.* 184:37–48.
- [98] Mottl MJ, McConachy TF (1990) Chemical processes in buoyant hydrothermal plumes on the east
 pacific rise near 21 n. *Geochim. Cosmochim. Acta* 54:1911–1927.
- [99] Fitzsimmons J, Conway T, John SG, Boyle EA (2013) Iron Isotopes in Seawater from the Southeast
 Pacific and North Atlantic Oceans. *Min. Mag.* 77:1092.
- ⁸⁰⁷ [100] Lupton JE, Craig H (1981) A Major Helium-3 Source at 15°S on the East Pacific Rise. *Science* ⁸⁰⁸ 214:13–18.
- [101] Rijkenberg MJ, et al. (2014) The Distribution of Dissolved Iron in the West Atlantic Ocean. *PloS One* 9:e101323.

- [102] Nishioka J, Obata H, Tsumune D (2013) Evidence of an extensive spread of hydrothermal dissolved
 iron in the Indian Ocean. *Earth Planet. Sci. Lett.* 361:26–33.
- [103] Toner BM, Marcus MA, Edwards KJ, Rouxel O, German CR (2012) Measuring the form of iron in
 hydrothermal plume particles. *Oceanogr.* 25:209–212.











