Iron minerals as archives of Earth’s redox and biogeochemical evolution

CAROLINE L. PEACOCK¹, STEFAN V. LALONDE² and KURT O. KONHAUSER³

¹ University of Leeds, School of Earth and Environment, Leeds, LS2 9JT, UK, e-mail: C.L.Peacock@leeds.ac.uk
² European Institute for Marine Studies, CNRS-UMR6538 Laboratoire Domaines Océaniques UMR 6538, Brest, France, e-mail: Stefan.Lalonde@univ-brest.fr
³ University of Alberta, Department of Earth and Atmospheric Sciences, Edmonton T6G 2E3 Canada, e-mail: KurtK@ualberta.ca

Abstract
Iron minerals provide sedimentary repositories of chemical information pertaining to Earth’s redox and biogeochemical evolution, from before the Great Oxidation Event some 2.5 billion years ago, to more recent events occurring up to and into the Cenozoic Era. The most powerful chemical information recorded in iron minerals comes in the form of trace-element signatures, most notably their concentrations and stable isotope compositions. Here we provide an introduction to iron mineralogy and the processes responsible for the accumulation and preservation of trace-element signatures in iron minerals, focusing on the deposition of iron minerals in three key ancient sedimentary archives: banded iron formations, ferromanganese crusts and black shales. We introduce the theory and practical use of non-traditional trace-element stable-isotope systems in redox and biogeochemical research, focusing on the recent use of iron, molybdenum and chromium stable isotopes to shed light on the redox and biogeochemical information stored in iron-rich sediments. By analysing both trace-element concentrations and stable-isotope compositions recorded in iron minerals, iron-rich sedimentary archives are providing a unique window into the past, where changes in trace-element signatures shed light on major transitions in Earth’s redox and biogeochemical evolution.

Introduction

This chapter provides a review of the use of iron minerals as repositories of chemical information pertaining to Earth’s redox and biogeochemical evolution, from before the Great Oxidation Event some 2.5 billion years ago, to more recent events occurring up to and into the Cenozoic Era. The most powerful chemical information recorded in iron minerals comes in the form of trace element signatures, most notably their concentrations and stable isotope compositions. We begin our review by providing an introduction to iron mineralogy and the processes responsible for the accumulation and preservation of trace element signatures in iron minerals, focusing on the deposition of these minerals in three key ancient sedimentary archives: banded iron
formations (BIF), ferromanganese crusts and black shales. We then review the theory and practical use of non-traditional trace element stable isotope geochemistry, and in particular the use of emerging non-traditional stable isotope systems, in palaeo redox and biogeochemical research. Finally we focus on how different trace element signatures, recorded in those ancient sedimentary archives, have helped provide a window into the past, where changes in trace element signatures can be interpreted to reflect transitions in Earth’s redox and biogeochemical evolution.

1. Iron minerals as archives of chemical information

Iron (Fe) minerals exist over a very wide range of redox conditions. They are extremely efficient scavengers of other elements, and effectively archive chemical information about the fluids from which they form. In the marine realm, Fe minerals form chemical sediments and cements under a variety of aqueous conditions and geological settings; these range from rapid water-column precipitation in Fe-rich hydrothermal plumes upon mixing with bulk seawater to the slow formation of concretions and cements at redox boundaries in sediments, nodules, and hydrothermal chimneys. The ubiquity of Fe oxyhydroxides in the marine environment, both modern and ancient, and the fact that their composition is intimately linked to the chemistry of the waters from which they precipitate, make them especially promising recorders of the geochemical history of their formation waters. Specifically, the elemental and isotopic enrichments recorded in Fe minerals provide information on the redox state of the contemporaneous environment and the biogeochemical cycling of bio-essential metals, both of which are linked to primary productivity, carbon cycling, and ultimately the evolution of Earth’s surface environments through time.

1.1. Iron mineralogy as a function of redox

Iron is the fourth most abundant element in the Earth’s crust, where it exists predominantly as ferrous iron, Fe(II), in Fe-bearing silicate minerals (e.g. pyroxenes, amphiboles). As these minerals are weathered, Fe(II) is released into solution, largely oxidized to ferric iron, Fe(III), and then delivered to the oceans. Transport involves a continuum of different size fractions, defined as aqueous (<0.02 μm, mainly organically-bound Fe), nanoparticulate (0.02 to 1 μm, mainly Fe(III) oxyhydroxide minerals that are also closely associated with organic species), and larger particles (>1 μm); for further explanation of Fe size fractions see Raiswell and Canfield (2012). There is a myriad of different processes delivering Fe to the oceans, and the relative magnitudes of the associated Fe fluxes vary both spatially and temporally (recently reviewed by Boyd and Ellwood, 2010 and Raiswell and Canfield, 2012). In order of importance, processes supplying Fe to the global ocean include deposition of windborne dust (e.g. Jickells et al., 2005), melting of glaciers and icebergs (e.g. Raiswell et al., 2008), remobilization of Fe from submarine shallow-shelf systems (e.g. Poulton and Raiswell, 2002), Fe transport via rivers (e.g. MacKenzie et al., 1979), and submarine hydrothermal emissions (e.g. Tagliabue et al., 2010; Table 1). A significant
proportion of the riverine Fe flux is probably removed via biogeochemical processes in estuaries, while the remainder is deposited on the continental shelf (e.g. Boyle et al., 1977; Boyd and Ellwood, 2010).

Once Fe reaches the oceans, and depending on prevailing geochemical conditions, it can be (re)precipitated via a number of different mechanisms as a variety of hydrogeneous and typically nanoparticulate Fe(II/III) minerals (for a review see Raiswell and Canfield, 2012). Because Fe is redox active, these minerals can form under almost all water column redox conditions. In addition, many of these minerals have considerable surface area with an amphoteratic surface charge (i.e. can be either positively- or negatively-charged depending on aqueous pH), meaning that they are highly reactive towards both dissolved cations and anions. As such, hydrogeneous Fe minerals make ideal sedimentary archives of trace element concentrations and stable isotope compositions in contemporaneous seawater, recording elemental signatures over a wide range of redox regimes.

Under fully oxic water column conditions, when the concentration of dissolved O₂ is greater than ~10⁻⁶ molar, Fe(II) is oxidized to Fe(III), which precipitates primarily as the nanoparticulate and poorly crystalline Fe(III) oxyhydroxide, ferrihydrite [Fe₃⁺⁻⁵(OH,O)₁₂]. Ferrihydrite is metastable and ages to the more crystalline minerals goethite [α-FeOOH] or hematite [Fe₂O₃] (Cornell and Schwertmann, 2003). Under anoxic conditions, when the concentration of dissolved O₂ is below ~10⁻⁶ molar, a variety of different Fe(II/III) minerals are formed. In non-sulfidic anoxic conditions (total dissolved sulfide < 10⁻⁶ molar) a variety of non-sulfidized Fe minerals can precipitate, including ferrihydrite and/or ferrous carbonates (e.g. siderite [FeCO₃]) and ferrous phosphates (e.g. vivianite [Fe₃(PO₄)₂·8H₂O]). In addition, a mixed Fe(II/III) layered double hydroxide can form, termed green rust (general formula \( \text{Fe}^{II}_{1-x}\text{Fe}^{III}_x\text{OH})_2\] \(\text{H}^{+}\)[(x/n)\(A_n^{-}\)·(m/n)\(H_2O\)]\(\text{H}^{+}\) where \(A_n^{-}\) = CO\(_3\)\(^{-}\), SO\(_4\)\(^{2-}\), Cl\(^{-}\), OH\(^{-}\). While relatively rare today, recent research suggests that green rust might be an important reactive Fe phase in anoxic water columns that are devoid of free sulfide and rich in dissolved Fe(II) (termed ferruginous) (e.g. Zegeye et al., 2012). Under sulfidic

### Table 1. Fe sources and estimated fluxes to the global ocean, compiled and discussed extensively by Raiswell and Canfield (2012).

<table>
<thead>
<tr>
<th>Fe sources to the global ocean</th>
<th>Fe flux (Tg/yr)</th>
</tr>
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<tbody>
<tr>
<td>Aeolian</td>
<td>1.53–3.03</td>
</tr>
<tr>
<td>Icebergs</td>
<td>0.9–1.38</td>
</tr>
<tr>
<td>Sediment Recycling</td>
<td>0.05–0.25</td>
</tr>
<tr>
<td>Riverine</td>
<td>0.14</td>
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<tr>
<td>Hydrothermal</td>
<td>0.05</td>
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<tr>
<td><strong>Total</strong></td>
<td><strong>2.62–4.8</strong></td>
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anoxic conditions where free sulfide is present (total dissolved sulfide $\geq 10^{-6}$ molar; termed euxinic) reactive Fe is precipitated as Fe sulfides, first as metastable makinawite (FeS), which then ages to pyrite (FeS$_2$). The classification of oxic and anoxic environments, in terms of dissolved O$_2$ and sulfide, is taken from Berner (1981) who provides an extended discussion of Fe (and other) minerals that are thermodynamically stable in these geochemical regimes.

In ocean sediments Fe(II/III) minerals are subject to diagenesis, during which they can be reductively dissolved and reprecipitated. Reductive dissolution proceeds via two competing pathways (e.g. Canfield et al., 1993a,b; Van Cappellen and Wang, 1996; Raiswell and Canfield, 1998; Haese, 2000), the first involving microbial reduction (dissimilatory Fe reduction, DIR) and the second involving abiotic reduction by dissolved sulfide, which diffuses upwards through the sediment column from a deeper zone of microbial sulfate reduction (e.g. Berner, 1964; Staubwasser et al., 2006). If Fe(III) oxyhydroxides are reduced by DIR, reactive Fe can go through a repeated cycle of reductive dissolution, upward diffusion, oxidation and reprecipitation as Fe(III) oxyhydroxides or, with partial oxidation, reprecipitation as a myriad of other Fe(II/III) minerals as described above (e.g. Canfield, 1989; Staubwasser et al., 2006). Ultimately, reduction of Fe(III) oxyhydroxides by dissolved sulfide removes Fe from the diagenetic cycle (e.g. Raiswell and Canfield, 1998). The diagenetic formation of Fe sulfide minerals constitutes a major sulfur and Fe exit channel that occurs at anoxic depths in the sedimentary pile where both reduction of Fe(III) and sulfate occur.

Use of Fe-rich sediments as archives of palaeo-seawater composition assumes that the Fe minerals provide a faithful record of seawater trace element concentrations and/or stable isotope compositions, despite their diagenetic history. In some sedimentary systems, where Fe(III) concentrations exceed organic carbon, Fe(III) oxyhydroxides can be preserved in the sediments without reductive dissolution (e.g. Konhauser et al., 2005). Under different scenarios, where reductive dissolution is likely to have occurred, it is important to demonstrate that the diagenetic processes have not significantly affected the original trace element concentrations and/or stable isotope compositions at the bulk scale, i.e. that there has been minimal loss of Fe and elements of interest from the bulk archive over time, and minimal redistribution of the isotopes between the bulk archive and sedimentary pore-waters (e.g. Poulton and Canfield, 2006; Friedich et al., 2011; Robbins et al., 2015). Indeed, in the case of some geochemical proxies, the environmental signal recorded by Fe minerals may be largely a diagenetic one (e.g. the case of Fe isotope compositions; section 2.4.1).

1.2. Uptake of trace elements by iron minerals

1.2.1. Iron oxyhydroxides

Iron oxyhydroxides are highly effective sorbents, a property born out of the functional group and electrostatic charging characteristics of their surfaces. Some minerals (e.g. clays) possess permanent surface charge as the result of structural defects or isomorphous substitution. Others, including most hydrated oxyhydroxide minerals (whether they be Fe oxyhydroxides, silicon oxides, titanium oxides, etc.), develop
surface charge as the result of pH-dependent surface functional group protonation/ deprotonation reactions. These minerals develop a positive surface charge at low pH as surface sites are neutralized or rendered positively charged with the surface sorption of protons and a negative charge at high pH as progressive deprotonation occurs (i.e. they are amphoteric). Adopting a 1pK formalism for the representation of surface sites, then surface sites of Fe oxyhydroxides (including hydrous ferric oxide, HFO, which is synonymous with ferrihydrite) develop charge through the following protonation/deprotonation reaction:

\[ >\text{FeOH}^{-0.5} + H^+ \leftrightarrow >\text{FeOH}_2^{+0.5} \quad \text{pK}_a = \sim 8 \]  

(1)

where \( >\text{FeOH} \) denotes surface Fe hydroxyl functional groups (log acidity (pK_a) value for ferrihydrite given as an example and taken from e.g. Davis and Leckie (1978)). At pH values above and below ~8, surface sites, and thus the net charge on the mineral surface, will be predominately negatively or positively charged as \( >\text{FeOH}^{-0.5} \) or \( >\text{FeOH}_2^{+0.5} \), respectively. At pH ~8, surface sites of opposite charge are present in equal amounts and this is deemed the point of zero charge (PZC). Crucially, ferrihydrite, and other pure Fe oxyhydroxide surfaces in general, are characterized by PZC values around 8, and thus possess abundant negatively and positively charged sites that are strongly reactive under many natural conditions (e.g. Sverjensky and Sahai, 1996). Considering their ubiquity and propensity to form nanometre-scale aggregates with large surface areas (in some cases upwards of 600 m² per dry gram, Roden et al., 2003), Fe oxyhydroxides provide the most prevalent positively charged reactive mineral surface found in nature (Cornell and Schwertmann, 2003). In contrast, the surfaces of silicate minerals generally possess PZC values around 2 (e.g. Ihler, 1979), and are thus strongly negatively charged under most natural conditions.

The surface chemistry of Fe oxyhydroxides has long been studied in relation to the sorption of contaminants in natural waters. Equilibrium surface complexation models (SCM), employing experimentally- and/or theoretically-derived parameters (e.g. stability constants for (de)protonation of the surface sites, surface area, surface site density, surface capacitance), have been developed to account for Fe oxyhydroxide sorption behaviour under a wide variety of pH values, ionic strengths, and sorbent–sorbate ratios (for an introduction to SCM see Davies and Kent, 1990). Among the early works, Dzombak and Morel (1990) provide a comprehensive database of thermodynamic parameters that describe the experimentally observed sorption of a variety of anions and cations to HFO. These surface sorption reactions may proceed to complete surface coverage, and in some cases, sorption that exceeds surface site saturation can lead to surface precipitation (e.g. Ler and Stanforth, 2003).

More recently, research on the sorption of trace elements to Fe oxyhydroxides has focused on determining the speciation and precise molecular coordination environment (e.g. mono- vs. bi-dentate, mono- vs. bi-nuclear) of sorbed trace elements using synchrotron-based spectroscopic techniques (for a review see Brown and Parks, 2001). Following the application of synchrotron spectroscopy to metal sorption systems, a new class of SCM have been developed by studies that first determine the molecular mechanisms of trace element sorption, then invoke this directly observed surface
complex to describe experimental trace element sorption data (e.g. Peacock and Sherman, 2004a,b; Moon and Peacock, 2011, 2012, 2013). These SCM are better able to predict trace element partitioning between solution and solid compared to those based solely on macroscopic sorption experiments. By properly defining the stoichiometry of the surface complex (i.e. the necessary input reaction that represents the formation of the surface complex in the model), and thus its complex-specific dependence on pH, aqueous trace element concentration and surface site density, it is possible to obtain an accurate description of its formation as a function of these same parameters. In order to apply model predictions to both modern and, in particular, ancient natural systems, where pH, trace element concentrations and surface site density (i.e. mineral crystallinity) vary over time and space, it is increasingly desirable to constrain SCM with molecular-level information whenever possible (e.g. for micronutrient trace element concentrations the modern oceans, Peacock and Sherman, 2007; Sherman and Peacock, 2010).

Significant progress has been made in testing predictive models using natural Fe oxyhydroxides (e.g. Tessier et al., 1996; Kennedy et al., 2003; Lalonde et al., 2007). In some cases, highly parameterized SCM may not be practical for the description of natural systems; critical information such as mineral surface area and surface site density may be difficult to determine. In these cases, lumped-process distribution coefficients, based on the Freundlich isotherm approach, are often applied. The distribution coefficient approach, which employs a linear or exponential relationship to describe the distribution of a trace element between solid and solution, benefits from simplicity and ease of execution, but, because it does not determine the speciation or coordination environment of the sorbed trace element, it cannot properly describe the formation of the surface complex and thus the sorption of the trace element as a function of the important environmental parameters (pH, trace element concentration, etc.). Partition coefficients derived via this approach, either experimentally or by measuring trace element concentrations in contemporaneous solids and solutions, are therefore, limited to the particular aqueous conditions (fixed pH, trace element concentration and ionic strength) present in the experiments or at the time of trace element uptake (Langmuir, 1997). Nonetheless, as we shall see below, distribution coefficient models have proven effective for the description of trace element sequestration by natural Fe oxyhydroxides, especially in marine systems (e.g. Feely et al., 1998; Edmonds and German, 2004), and with consideration of these limitations, they have been successfully used to relate trace element concentrations recorded in Fe mineral archives to contemporaneous seawater chemistry (e.g. Konhauser et al., 2009).

1.2.2. Iron sulfides
Like their oxide counterparts, Fe sulfide minerals are generally insoluble, and in anoxic waters and sediment porewaters where Fe$^{2+}$ and S$^{2-}$ (predominately as HS$^{-}$ at marine pH) are present, they precipitate rapidly. Taking the form Fe$_x$S$_y$, seven stable minerals composed entirely of Fe and S have been described, of which four may be found in low-temperature aqueous and sedimentary environments: mackinawite (meta-stable,
amorphous to tetragonal FeS), pyrite (cubic FeS₂), marcasite (orthorhombic FeS₂), and greigite (cubic Fe₃S₄). Precipitation of amorphous mackinawite proceeds faster than the other phases, such that it is often considered a precursor for its more crystalline counterparts. However, mackinawite is not a strict precursor in the sense that it does not appear to experience solid-state transformation during diagenesis, but rather undergoes dissolution and re-precipitation (cf. Rickard and Luther, 2007). Pyrite is the most abundant Fe sulfide mineral that is stable at Earth’s surface, and like Fe oxides, it’s tendency to concentrate trace elements from its formation waters makes pyrite-rich sediments both an important marine trace element sink and a rich palaeo-environmental chemical archive (e.g. Large et al., 2014).

Similar to Fe oxides, Fe sulfide minerals present a reactive surface upon which significant trace element sorption may occur. In the case of disordered mackinawite, the reactive surface appears composed of strongly acidic mono-coordinated sulfur sites (\(>\text{FeSH}^0\)) and weakly acidic tri-coordinated sulfur sites (\(>\text{Fe}_2\text{SH}_0\)), with a specific surface area of \(\sim 350 \text{ m}^2 \text{ g}^{-1}\) (Wolthers et al., 2005). Few thermodynamic surface stability constants exist for trace elements at Fe sulfide mineral surfaces, however, partition coefficients for the adsorption of metal-sulfide forming elements to the mackinawite surface appear to scale with the solubility products of their respective metal sulfides (Morse and Arakaki, 1993). Despite mackinawite being an unstable phase, the linear free-energy relationship that characterizes its trace element uptake appears reflected in the trace element compositions of sedimentary pyrites.

In pyrite-bearing marine sediments, the degree of trace element pyritization (DTMP), defined for any given trace element as the percent of that metal in a sediment that is found in pyrite (Raiswell and Canfield, 1998), also appears to scale with the respective metal sulfide solubility products (Morse and Luther, 1999; Fig. 1). In other words, the more pyrite a sediment contains, the greater the proportion of the total trace element load will be hosted in pyrite. However, not all metals display this

![Figure 1](image-url). The degree to which different trace elements in anoxic sediments are concentrated in pyrite (% DTMP) vs. their own solubility as a metal sulfide (relative to FeS; \(\log K_{\text{MeS}}/K_{\text{FeS}}\)). This relationship speaks to the affinity of different metals for sulfide as well as the kinetics of MeS vs. FeS precipitation. Some metals precipitate slowly as sulfides and thus are more likely to be found substituted into pyrite in anoxic sediments (metals on the pyritization trend), whereas others rapidly form their own metal sulfides and fall off this trend (Pb, Zn, Cd). Redrawn from Morse and Luther (1999).
thermodynamically-controlled behaviour; for those which may rapidly precipitate as their own amorphous metal sulfide, notably, lead (Pb), zinc (Zn), and cadmium (Cd), the kinetics of water exchange are faster than that of Fe$^{2+}$, and these elements tend to form their own metal sulfide phases in pyritised sediments, and DTMP for these elements remains low (Morse and Lurther, 1999). As a consequence, sedimentary pyrite is generally considered an important sink for arsenic (As), mercury (Hg), and molybdenum (Mo), moderately important for cobalt (Co), copper (Cu), manganese (Mn), and nickel (Ni), but generally unimportant for Pb, Zn, and Cd (Huerta-Diaz and Morse, 1992).

While predictive techniques, such as partition coefficient and SCM model calculations, are relatively undeveloped compared to those advanced for Fe oxide minerals, trace element concentrations in sedimentary pyrite nonetheless appear to scale with aqueous concentrations in their local environment, such that pyrite-rich sediments (e.g. black shales) represent a rich archive of information on Earth’s surface geochemical evolution (section 3.3). Indeed, focusing on just pyrite grains from shales, Large et al. (2014) documented the temporal variability of several trace elements, including Mo, Co, Ni, As and Se, throughout geological time. Those authors suggested that in the same way that hydrothermal pyrite tracks the chemistry of ore-forming fluids (Large et al., 2009), sedimentary pyrite can be used to track changes in seawater composition, and ultimately the redox state of Earth’s surface environment. This view has recently found support in the work by Swanner et al. (2014) who provided pyrite, shale, and BIF trace element data that all suggested an expansion of the palaeo-marine Co reservoir between 2.48 and 1.84 Ga, coincident with anoxic marine conditions and expanded hydrothermal activity. Similarly, Gallagher et al. (2015) recently reported a suite of trace element data from Precambrian to Ordovician carbonate-hosted pyrites centred around several key geological events such as the Great Oxidation Event (GOE) ~2.45 Gyr (billion years ago), that were in most cases argued to be largely consistent with previous pyrite and shale records.

1.2.3. Exemplar molybdenum

Molybdenum is the most abundant transition metal in seawater (~105 nmol/kg; Anbar and Knoll, 2002) and can exist in a variety of different oxidation states depending on ocean redox. In oxic conditions, Mo is present as molybdate (Mo(VI)O$_4^{2-}$) and its concentration in seawater is thought to be controlled by an equilibrium reaction in which molybdate sorbs to Fe and manganese (Mn) oxyhydroxide minerals present in ocean sediments (Barling et al., 2004; Goldberg et al., 2009, 2012). Under anoxic non-sulfidic conditions, where dissolved Fe is commonly present in the water column (termed ferruginous), Mo is likely drawn down in association with other minerals, such as green rust (Zegeye et al., 2012). In the presence of dissolved sulfide, molybdate is converted into a series of thiomolybdate complexes, starting with oxythiomolybdates [Mo(VI)O$_4$$_{4-x}$S$_2^{2-}$] (when H$_2$S ~0.1–11 µM), which progressively dominate Mo speciation and possibly culminate in tetrathiomolybdate [Mo(VI)S$_4^{2-}$] (when H$_2$S >11 µM) (Helz et al., 1996; Erickson and Helz, 2000). These thiomolybdates are
extremely reactive and Mo might, therefore, be drawn down via the scavenging of these complexes onto Fe minerals and sulfidized organic particles (likely at low concentrations of dissolved sulfide), or perhaps via the precipitation of Mo-sulfide phases (at higher concentrations of dissolved sulfide (~50–250 μM) and presumably Mo) (Helz et al., 1996; Zheng et al., 2000). On the other hand, these complexes might react with zero-valent sulfur present in natural sulfidic waters to produce Mo(IV)-polysulfides, which might similarly be scavenged by Fe minerals and/or organic matter (Dahl et al., 2010). As such, the concentration and speciation of Mo recorded in Fe mineral archives can be used to infer prevailing ocean redox conditions at the time of sediment deposition.

Understanding how the amount of oxygen in seawater has changed through time is key to elucidating links between the evolution of ocean redox chemistry, climate and life. Oxygen provides an important control on seawater chemistry because many elements present in seawater are critically sensitive to redox conditions - existing in oxic waters either in solution as aqueous ions/organic carbon complexes or as solid forms that may be (depending on the precise chemical nature of the redox environment) sequestered into the sediments under reducing conditions. This drawdown of redox-sensitive elements, such as Mo, provides a means of reconstructing past ocean redox via careful analyses of the rock record. Indeed, the determination of Mo concentrations in marine shales is one of the most commonly used means to identify the most extreme water column redox condition (highly sulfidic, termed euxinic) throughout Earth history (Scott et al., 2008).

The availability of Mo in seawater also has important implications for the biosphere because it is essential for various cellular metabolisms. For example, it is required for nitrate assimilation by eukaryotes and some prokaryotes, and its enhanced drawdown under widespread euxinic conditions in the mid-Proterozoic ocean (~1.8 Gyr) has been widely postulated as an explanation for the maintenance of low levels of atmospheric oxygen and the protracted pace of biological evolution during Earth’s ‘middle age’ (Anbar and Knoll, 2002). As such, intimate links exist between ocean redox, biological activity, atmospheric chemistry and climate, and these can be identified and evaluated by detailed understanding of the environmental behaviour of important redox-sensitive species such as Fe and Mo.

1.3. Iron mineral archives

The most commonly utilized Fe mineral archives are those found in BIF and marine ferromanganese crusts, in which the concentrations and/or stable isotope compositions of trace elements present in the constituent Fe (and Mn) minerals can be interpreted to reflect contemporaneous seawater trace element signatures. In addition to these Fe-rich archives, scavenging of trace elements onto discrete Fe minerals, which are then deposited into marine sediments as relatively dispersed Fe phases, might also provide repositories for contemporaneous trace element seawater signatures. Indeed, trace element enrichment in black shales, which provide an important archive for redox-sensitive trace elements (e.g. Arnold et al., 2004; Scott et al., 2008), probably results
from the scavenging of particle reactive trace element species by Fe minerals (e.g. Helz et al., 1996; Zheng et al., 2000; Dahl et al., 2013).

1.3.1. Banded iron formations

Banded iron formations represent vast quantities of Fe-rich sediments that were precipitated in the oceans between ~3.8 and 1.8 Ga. Trace element signatures recorded in BIF can therefore provide information on contemporaneous seawater chemistry during early Earth evolution. The name BIF stems from the appearance of these enigmatic deposits, composed of alternating bands of Fe-rich (~20–40% Fe) and silica (Si)-rich (~40–50% SiO₂) layers. Banding can often be observed on a wide range of scales, from coarse macrobands (metres in thickness) to mesobands (centimetre-thick units) to millimetre and submillimetre layers. Among the latter is the wide variety of varve-like repetitive laminae, known as microbands (Trendall and Blockley, 1970).

The mineralogy of BIF from the best preserved successions is remarkably uniform, comprising mostly quartz (in the form of chert), magnetite, hematite, Fe-rich silicate minerals (stilpnomelane, minnesotaite, greenalite and riebeckite), carbonate minerals (siderite, ankerite, calcite and dolomite), and minor sulfides (pyrite and pyrrhotite); the presence of both ferric and ferrous minerals give BIF an average oxidation state of Fe²⁺ (Klein and Beukes, 1992). However, it is generally agreed that none of the minerals in BIF are primary in origin. Instead, the minerals reflect significant post-depositional alteration under diagenetic and metamorphic conditions (including, in some cases, post-depositional fluid flow). The effect of increasing temperature and pressure is manifested by the progressive change in mineralogy through replacement and recrystallization, increase in crystal size and obliteration of primary textures (Klein, 2005). For instance, the alternating layers of magnetite and hematite are interpreted to have formed from an initial Fe oxyhydroxide phase, e.g. ferrihydrite, that precipitated in the photic zone when dissolved Fe²⁺ was oxidised and hydrolysed to insoluble ferric Fe. The Fe oxyhydroxide particles then sank through the water column and were deposited on the seafloor where they eventually formed (1) magnetite or Fe carbonates when organic remineralisation was coupled with Fe(III) reduction; (2) hematite, when organic material was lacking; or (3) Fe silicates, possibly in the form of a precursor mineral such as greenalite, when Si-sorbed ferric oxyhydroxides reacted with other cationic species in the sediment pore waters (Morris, 1993). The chert is widely considered to have precipitated from the water column as colloidal Si co-precipitated with Fe-rich particles (Fischer and Knoll, 2009) given that the Archaean ocean had significantly elevated concentrations of dissolved Si, at least as high as at saturation with cristobalite (0.67 mM at 40°C in seawater), and possibly even amorphous Si (2.20 mM) (Siever, 1992; Maliva et al., 2005; Konhauser et al., 2007).

There is still considerable effort towards understanding how BIF formed but almost all researchers now agree that BIF were precipitated from a predominantly anoxic water column (e.g. Holland, 1984). The duration of BIF deposition spans major evolutionary changes in the Earth’s surface composition, from an early anoxic atmosphere dominated by CO₂ and CH₄ to an atmosphere that became partially oxygenated.
Therefore, it is likely that BIF formed via different mechanisms throughout the Precambrian. The traditional model of BIF precipitation assumes the oxidation of dissolved Fe(II) via abiotic oxidation by cyanobacterially-produced O₂ (Cloud, 1973; Klein and Beukes, 1989) and/or biotic oxidation by chemolithotrophic bacteria (Holm, 1989; Planavsky et al., 2009). Both models suggest the presence of free molecular oxygen in the Precambrian ocean and, hence, require the presence of oxygenic photosynthesis at that time of Earth history. Given the recent evidence for oxidative processes having already existed at 3.0 Gyr (Crowe et al., 2013; Planavsky et al., 2014), and perhaps even at 3.2 Gyr (Satkowski et al., 2015), it might be plausible that O₂-driven reactions in the photic zone led to BIF deposition since the Mesoarchaean. An alternative biological mechanism, and one that might explain BIF deposition prior to the Mesoarchaean, involves Fe(II) oxidation via anoxygenic photosynthesis (Czaja et al., 2013; Jones et al., 2015; Pecoits et al., 2015). Phototrophic Fe(II)-oxidizing bacteria were discovered two decades ago (e.g. Widdel et al., 1993), and their metabolism involves light-energy fuelled CO₂ fixation coupled to the microbial oxidation of Fe²⁺. The attractiveness of this concept is that it explains BIF deposition in the absence of molecular oxygen using the abundant availability of Fe²⁺, light and CO₂ at that time (Garrels et al., 1973; Hartman, 1984; Konhauser et al., 2002). It has even been demonstrated by eco-physiological lab experiments in combination with modelling that these phototropic bacteria would have been capable of oxidizing enough Fe(II) to explain the large expansion of BIF deposits (Kappler et al., 2005). There is in addition one abiological mechanism, that being ultraviolet photo-oxidation of Fe²⁺ in the surface waters (e.g. Braterman et al., 1983). However, this view has been challenged on the basis that those early experiments were not conducted in complex solutions mimicking seawater (Konhauser et al., 2007). Whatever the mechanism for their precipitation, BIF today have undergone probably repeated episodes of diagenesis and metamorphism, and their Fe mineralogy is now dominated by magnetite and hematite (for reviews on BIF deposition and mineralogy see Klein, 2005 and Bekker et al., 2010).

1.3.2. Ferromanganese crusts

Marine ferromanganese crusts are found throughout the world’s oceans and are formed of subequal amounts of Fe(III) oxyhydroxide and Mn(III,IV) minerals that have precipitated from seawater and deposited as pavements and coatings on submarine rock outcrops and topographic highs (e.g. Glasby, 1977). Crusts are arguably the slowest growing geological precipitates, with only a few mm deposited per million years (Ma) (e.g. Hein et al., 2000). Accordingly, crust formations of ~10 cm thickness can provide trace element archives stretching over the last ~65 Ma of Earth history. Because ferromanganese crusts precipitate directly from ambient seawater (termed “hydrogenetic”) these archives can be interpreted to reflect contemporaneous seawater chemistry throughout the Cenozoic. Crusts are restricted to submarine rock outcrops and topographic highs because these environments are often kept sediment free, via, for example, local currents generated by obstructional upwelling along seamount flanks.
These currents also promote enhanced turbulent mixing and upwelling, leading to increased primary productivity and development of an oxygen-minimum zone (OMZ) that increases the concentrations of Fe$^{2+}$ and Mn$^{2+}$ which can be scavenged into crusts during oxic conditions resulting from turbulence (Muinos et al., 2013). As a result of these conditions, Fe and Mn oxyhydroxides precipitated from seawater can deposit onto the hard rock surfaces and build up in layers over time, as opposed to elsewhere in the oceans where the same precipitates are deposited and dispersed into seafloor sediments. Coupled sorption and redox reactions occurring between dissolved trace elements present in contemporaneous seawater and the mineral precipitates then sequester metals as the minerals precipitate (via co-precipitation and structural incorporation) and during their transit through the marine water column (via adsorption onto mineral surfaces) (for a general model of crust formation see Koschinsky and Halbach, 1995). Rare and critical trace elements in particular are often concentrated to economically valuable concentrations in marine ferromanganese crusts, making the deep-sea mining of crusts and nodules a potentially attractive endeavour (e.g. Hein et al., 2013).

The Fe mineralogy of crusts is dominated by ferrihydrite, with older crust layers sometimes featuring small (<10%) amounts of goethite, typically attributed to aging of the authigenic ferrihydrite (e.g. Hein et al., 2000), or a switch in precipitation from authigenic ferrihydrite to goethite during a temporary change in palaeo-seawater conditions (e.g. Koschinsky et al., 1997). Manganese mineralogy is dominated by poorly crystalline phyllomanganates, namely $\delta$-MnO$_2$ (often termed vernadite) and birnessite (e.g. Burns and Burns, 1977). Minor quartz, feldspar and other detrital minerals, along with sometimes significant (~20%) carbonate fluorapatite in older crusts subject to diagenesis and/or precipitated during palaeo-seawater transition, are also present (e.g. Hein et al., 1993).

Marine ferromanganese nodules are also prevalent in the deep oceans, and, despite their similarly slow growth rates, they do accumulate and comprise a significant component of some marine and freshwater sediment. For instance, in the Pacific Ocean, it has been estimated that $10^{12}$ tons of nodules exist, predominantly in pelagic sediments, with an annual rate of formation of $6 \times 10^6$ tons (Mero, 1962), while in Oneida Lake, New York, for example, within a 20 km$^2$ area of lake, $10^6$ tons of nodules, that average 15 cm in diameter, exist at the bottom sediment (Dean and Greeson, 1979). Diagenetic nodule formation involves a series of microbially-catalysed reactions. The process begins with cyanobacterial and algal plankton concentrating Fe$^{2+}$ and Mn$^{2+}$ from solution. Upon their death the microbial biomass and those bound metals are transported to the bottom sediment where anaerobic respiratory processes in the suboxic layers liberates Fe$^{2+}$ and Mn$^{2+}$ back into the sediment pore waters. Concurrently, reduction of particulate ferric oxyhydroxides and Mn(III,IV) oxides occurs. Upward diffusion to the sediment–water interface facilitates microbial re-oxidation and incorporation of metals onto some form of nucleus, which can be any solid mineral or organic substrate. With continued accretion of Fe and Mn, a nodule, that may or may not display concentric laminations, forms. Plankton contributes to
metal oxidation by producing high-pH, oxygenated surface waters that are conducive to the re-oxidation (and hydrolysis) reactions (e.g. Richardson et al., 1988). Because nodules grow at the sediment–water interface (termed ‘diagenetic’) it is difficult to determine to what extent their trace element signatures reflect contemporaneous seawater chemistry and their use as archives of seawater chemical information is somewhat limited.

### 1.3.3. Black shales

Shales are an example of marine sediments that, although not technically classed as Fe mineral archives, can never-the-less record contemporaneous trace element seawater signatures, likely as a result of trace element drawdown in association with Fe (and sulfide and organic) phases (e.g. for Mo, Helz et al., 1996; Zheng et al., 2000; Dahl et al., 2013). They comprise fine-grained clastic sediments predominantly composed of clay minerals (e.g. kaolinite, montmorillonite and illite) and silt-sized fragments of other minerals (e.g. quartz and calcite). Shales are typically grey in colour, but addition of variable amounts of minor constituents can significantly alter their appearance; most notably perhaps in the case of black shales which contain > 1% unoxidized carbonaceous material, and display an Fe mineralogy dominated by amorphous Fe sulfides and pyrite (for a summary of the geochemistry of black shale deposits see Vine and Tourtelot, 1970). This relatively high unoxidized carbon content and the presence of Fe sulfides suggests that black shales were deposited under an anoxic water column, where organic carbon was not subject to aerobic oxidative degradation and ferrous Fe was thermodynamically stable. The presence of sedimentary fabrics also indicates a lack of physical disruption by burrowing animals (at least since the Neoproterozoic). This, combined with depositional occurrences that span most of the geological record (e.g. Tourtelot, 1979), means that trace element signatures recorded in black shale archives can represent seawater chemistry during episodes of ocean anoxia occurring throughout Earth history.

### 2. Stable isotope fractionation of trace elements

Subtle physicochemical discrimination between the different stable isotopes of any given element may impart a robust geochemical fingerprint of the processes that the element has experienced, making stable isotope geochemistry a powerful tool for palaeo-environmental reconstruction. This has been recognized and exploited by geochemists for over 60 years, with early studies naturally focusing on the stable isotopes of lighter elements (H, C, N, O and S) whose greater relative mass differences imparted stronger, more easily measured fractionations. These ‘traditional’ stable isotope systems have since been supplemented by ‘non-traditional’ ones whose smaller natural fractionations have now become accessible due to advances in mass spectrometry technology. Natural stable isotope variability has now been described and quantified for an increasing number of non-traditional elements, including Li, B, Mg, Si, Cl, Ca, V, Cr, Fe, Ni, Cu, Zn, Ge, Se, Mo, Cd, Sn, Ce and Eu, and the list continues to grow (see Johnson et al., 2004 dedicated to the subject).
2.1. Stable isotope nomenclature and conventions

Most stable isotope data are reported in delta notation where the isotope composition of a sample is defined as the relative deviation of the sample isotope ratio to the same isotope ratio in a known reference material (e.g. O’Neil, 1986):

$$\delta^{ij}_E = 1000 \left( \frac{R_{ij}^{\text{Sample}}}{R_{ij}^{\text{Reference}}} - 1 \right)$$  \hspace{1cm} (2)$$

where $i$ and $j$ are the particular isotopes used in ratio $R$ of element $E$. Units for $\delta^{ij}_E$ are in parts per thousand, i.e. per mil (%). For the traditional stable isotope systems (H, C, N, O and S), the isotope ratio $R^{ij}$ is expressed as the abundance of the minor isotope over the abundance of the major isotope, for example, D/H and $^{13}$C/$^{12}$C. This convention has ensured a consistent point of reference for traditional stable isotope enrichments and depletions, where a positive or negative value of $\delta^{ij}_E$ indicates that a sample is relatively enriched or depleted in the heavy isotope, respectively, compared to the standard. For several non-traditional stable isotope systems, however, $R^{ij}$ expressed as rare over major isotope does not satisfy the heavy over light convention, because the minor isotope is the isotopically light. This is the case for the Fe and Mo stable isotope systems, where, for example, $^{54}$Fe and $^{95}$Mo are more rare than $^{56}$Fe and $^{98}$Mo. In order for stable isotope enrichments and depletions to be expressed in the same direction (heavy enrichments as positive delta values, light enrichments as negative delta values) between traditional and non-traditional systems, in non-traditional systems the convention heavy over light is usually preferred. Considering $^{54}$Fe and $^{56}$Fe and $^{95}$Mo and $^{98}$Mo, for the Fe and Mo stable isotope systems, respectively, $R^{ij}$ is typically expressed as $^{56}$Fe/$^{54}$Fe and $^{98}$Mo/$^{95}$Mo, with corresponding delta notation of $\delta^{56}$Fe and $\delta^{98}$Mo.

For traditional stable isotope systems the choice of reference material for each isotope system has been internationally standardized, where, for example, the reference material used for reporting the isotope compositions of H and O is Standard Mean Ocean Water (SMOW), and Air for N. For some of the non-traditional systems the reference material is yet to be agreed upon and is therefore not standardized across the field. In more recent work, Fe stable isotope measurements are usually expressed with reference to the isotope composition of the international standard IRMM-14 (Belshaw et al., 2000), however, for the Mo system an international standard has only recently been proposed (NIST SRM 3134; Wen et al., 2010; Greber et al., 2012), and the Mo isotope community has employed various in-house Mo isotope standards over the years to normalize results (for an inter-comparison see Goldberg et al., 2013). In many non-traditional stable isotope systems there is also a choice of isotope ratios that can be measured, because several of the non-traditional elements have multiple naturally occurring stable isotopes. For example, Fe has four naturally occurring stable isotopes $^{54}$Fe (5.84%), $^{56}$Fe (91.7%), $^{57}$Fe (2.12%) and $^{58}$Fe (0.28%), where stable isotope data is commonly reported as $^{56}$Fe/$^{54}$Fe, while Mo has seven naturally occurring stable isotopes $^{92}$Mo (14.84%), $^{94}$Mo (9.25%), $^{95}$Mo (15.92%), $^{96}$Mo (16.68%), $^{97}$Mo (9.55%), $^{98}$Mo (24.13%) and $^{100}$Mo (9.63%), and stable isotope data has been reported
for both $^{97}\text{Mo}/^{95}\text{Mo}$ and $^{98}\text{Mo}/^{95}\text{Mo}$. As both the reference material, REF, and the stable isotopes measured in $R^{\beta}$ can be different for the same non-traditional stable isotope system, care is required when comparing $\delta^{\beta}E$ values between different studies. For an extended review of stable isotope nomenclature and conventions the reader is directed to Johnson et al. (2004).

2.2. Stable isotope fractionation theory

2.2.1. Mass-dependent stable isotope fractionation

Mass-dependent stable isotope fractionation refers to any physical or chemical process that acts to partially separate stable isotopes of the same element between two different reservoirs, where the amount of separation scales in proportion with the difference in the masses of the isotopes. For example, using the O stable isotope system, the mass difference between $^{18}\text{O}$ and $^{16}\text{O}$ is $+2$, while the mass difference between $^{18}\text{O}$ and $^{17}\text{O}$ is $+1$, so considering the partitioning of O isotopes between two different reservoirs, $\delta^{18}\text{O}$ will be approximately twice that of $\delta^{17}\text{O}$ as the result of mass-dependent fractionation. Separation of stable isotopes of the same element between two different reservoirs, and thus two separate substances, occurs because chemical bond strengths are mass-dependent. In the case of the two most abundant isotopes of O, $^{18}\text{O}$ and $^{16}\text{O}$, in an $\text{H}_2\text{O}$ molecule, there is a difference in bond strength between $^{18}\text{O}/\text{C}_0\text{H}$ and $^{16}\text{O}/\text{C}_0\text{H}$. Specifically, $^{18}\text{O}/\text{C}_0\text{H}$ has a lower vibrational frequency, and hence a lower ground state energy (or zero point energy, ZPE), than $^{16}\text{O}/\text{H}$. This means that the bond between $^{18}\text{O}–\text{H}$ is stronger than that between $^{16}\text{O}–\text{H}$. Differences in chemical bond strength lead to differences in reaction rate constants, where species with the stronger bonds will react more slowly than species with the weaker bonds. This effect on reaction rate constants is the origin of kinetic isotope fractionation, which acts to separate isotopes during unidirectional or incomplete processes, perhaps best illustrated during evaporation of water where light water molecules (with $^{16}\text{O}$) will evaporate more rapidly than heavier water molecules (with $^{18}\text{O}$). Differences in chemical bond strength also lead to differences in equilibrium constants for reactions that are otherwise identical. This is the origin of equilibrium isotope fractionation, involving the partial separation of isotopes between two substances that are in chemical equilibrium. The theory of stable isotope fractionation pre-dates the advent of multi-collector inductively coupled plasma mass spectrometry (MC-ICP-MS) and the subsequent discovery of stable isotope fractionation in the heavy trace elements, and includes a few very early studies (Lindemann and Aston, 1919; Lindemann, 1919; Urey and Greiff, 1935). The modern theoretical basis for calculating mass-dependent fractionation was established over 60 years ago in a series of seminal studies by Harold Urey, Jacob Bigeleisen and Maria Mayer (e.g. Urey, 1947; Bigeleisen and Mayer, 1947). A rigorous treatment of mass-dependent stable isotope theory can be found in several texts, including a more recent review of fractionation fundamentals by Criss (1999).

Equilibrium stable isotope fractionation describes the partial separation of isotopes between two or more substances that are in, or are approaching, chemical equilibrium. There are a number of general qualitative chemical rules governing equilibrium stable
isotope fractionations which can be used to predict the magnitude and direction of isotope fractionation between two separate substances (for a complete review see Schauble, 2004). In the first instance, equilibrium isotope fractionations usually decrease as temperature increases, and, all else being equal, fractionations are largest for light elements and for isotopes with very different masses (e.g. Criss, 1999; Schauble, 2004). In addition to these rules, a further consideration of equilibrium thermodynamics provides a means to predict which substances will be enriched in heavy isotopes in a given geochemical system. In equilibrium thermodynamics, all systems strive to achieve their lowest energy state. Because chemical bonds involving heavier isotopes have a lower ground state energy than otherwise identical bonds involving lighter isotopes of the same element, then during equilibrium isotope fractionation the heavy isotope will tend to concentrate in the substance that confers the lowest ground state energy for that particular bond, i.e. in the substance that confers the strongest bonding environment. This process is well demonstrated in the isotope exchange of $^{18}$O and $^{16}$O between liquid water and water vapour:

$$\text{H}_2^{16}\text{O} (l) + \text{H}_2^{18}\text{O} (g) = \text{H}_2^{18}\text{O} (l) + \text{H}_2^{16}\text{O} (g)$$

where at, or approaching, chemical equilibrium, $^{18}$O will concentrate in the stronger O–H bonds of water relative to the weaker O–H bonds of steam. The formation of strong bonding environments, i.e. strong chemical bonds, correlates with several physicochemical properties, including high oxidation state in the element of interest and low coordination number (for a full list see Schauble, 2004). Thus with a choice of bonding environment between substance $A$ with element $E$ in low oxidation state, and substance $B$ with element $E$ in high oxidation state, the heavier isotopes will tend to concentrate in substance $B$, all else being equal. Similarly, with a choice of bonding environment between substance $A$ with element $E$ in high coordination, and substance $B$ with element $E$ in low coordination, the heavier isotopes will again tend to concentrate in substance $B$, all else being equal. Overall, the general rules for equilibrium isotope fractionation predict that large equilibrium fractionations are most likely to occur at low temperature between substances with significantly different oxidation states, coordination numbers, bonding partners and/or electronic configurations (Schauble, 2004). Furthermore, these rules suggest that stable isotope signatures of redox-active trace elements, preserved in the sedimentary record, can provide a record of the palaeo redox conditions present in the environment during the time of sediment deposition. Specifically, the extent of the measured fractionation in the sedimentary record should reflect the extent of oxidation/reduction of that redox-active species in the depositional system, which is itself related to the overall redox state of the contemporaneous environment. In essence, this is the basis for utilizing redox-active trace element stable isotopes recorded in sedimentary archives as tracers of past redox and biogeochemical conditions.

2.2.2. Fractionation factors

The fractionation factor is a convenient quantification of the contrast in isotope compositions between two substances and is defined as (e.g. O’Neil, 1986):
\[ \alpha_{A-B} = \frac{R_{i/j}A}{R_{i/j}B} \]

where \( A \) and \( B \) are two distinct substances, and \( R_{i/j}A \) and \( R_{i/j}B \) are the isotope ratios of the particular element of interest in substance \( A \) and \( B \), respectively. The fractionation factor may also be cast in terms of \( \delta^{i/E} \) values:

\[ \alpha_{A-B} = \frac{1000 + \delta^{i/E}A}{1000 + \delta^{i/E}B} \]

A fractionation factor of 1 indicates that the isotopes of element \( E \) are distributed evenly between substances \( A \) and \( B \) and there is no isotope fractionation. A fractionation factor > 1 indicates isotope \(^1E\) (for non traditional isotope systems \(^1E\) is usually the heavy isotope) is concentrated in substance \( A \), while a fractionation factor of < 1 indicates isotope \(^1E\) is concentrated in substance \( B \). The fractionation factor does not allow differentiation between the physical processes that lead to different isotope compositions in different substances, and thus may reflect equilibrium or non-equilibrium isotope partitioning.

### 2.2.3. Processes producing isotopically distinct reservoirs

There are generally two types of process that can lead to the partial separation of isotopes between two reservoirs, namely, closed-system equilibrium and Rayleigh fractionation. Closed-system equilibrium is best illustrated by considering the slow reaction of two substances \( A \) and \( B \), where \( A \) and \( B \) remain open to complete isotope exchange throughout the reaction. In this process the isotope composition of \( A \) and \( B \) is a function of the fraction of \( A \) remaining and the fraction of \( B \) produced. As the reaction proceeds, the shifting mass balances of \( A \) and \( B \) requires that their \( \delta^{i/E} \) values shift in order to maintain a constant offset in fractionation between \( A \) and \( B \). As such, \( \delta^{i/E}A \) and \( \delta^{i/E}B \) will progress along a straight line as a function of the fraction of \( B \) produced (Fig. 2).

During Rayleigh fractionation the products of a reaction do not continue to exchange with other phases in the system, because, for example, they are constantly isolated or removed. So as the reaction proceeds, product \( B \) is in thermodynamic and isotope equilibrium with reactant \( A \), but is then instantaneously removed from the system. In this case the isotope compositions of \( A \) and \( B \) evolve along an exponential trajectory, and because the product is progressively isolated, large changes in \( \delta^{i/E} \) values in the remaining components can occur (Fig. 2). An example would be the Fe isotope fractionations associated with ferrihydrite precipitating from seawater during the Precambrian, such that the BIF which formed in the water column and then settled out to the seafloor became progressively heavy in \(^{56}\)Fe, while the residual Fe(II) pool in solution became isotopically more depleted (e.g. Rouxel et al., 2005).

In addition to changes in the redox conditions of the contemporaneous environment and changes in the biogeochemical processes that ultimately control the source and sink fluxes of trace elements to and from the oceans, there are a number of independent
physicochemical processes that occur in seawater and which, according to the general rules governing equilibrium stable isotope fractionation (section 2.2.1), should be expected to cause stable isotope fractionation of trace elements and thus affect the isotope signal recorded in oceanic sedimentary archives. The first concerns the fact that almost all dissolved trace elements in seawater have varied speciation chemistry involving both inorganic and organic ligands. Physicochemical differences between the different dissolved aqueous species of element $E$ in seawater, including differences in redox state, coordination number, bonding partners and electronic configurations, are expected to induce stable isotope fractionation of $E$ between the relevant aqueous reservoirs (section 2.2.1). Isotope fractionation between different aqueous species have been studied for several trace element stable isotope systems, including Fe and Mo (e.g. Anbar et al., 2000; Roe et al., 2003; Welch et al., 2003; Anbar et al., 2005; Tossell, 2005; Weeks et al., 2007). Secondly, because there can be differences in coordination number and bonding partners (i.e. bonding environment) between the dissolved aqueous species of $E$ in seawater and the inorganic and organic particulate phases that sequester $E$ into the sediments, more often than not, the act of sequestration itself can induce an isotope fractionation. In particular the stable isotope fractionation of trace elements upon sorption to Fe and Mn oxyhydroxides has been measured for Mo (Barling and Anbar, 2004; Kashiwabara et al., 2011; Wasylenki et al., 2011), for Ni (Wasylenki et al., 2015), for Cu and Zn (Pokrovsky et al., 2005; Balistrieri et al., 2008; Pokrovsky et al., 2008) and for Cd (Wasylenki et al., 2014). In addition, for Fe and Mn oxyhydroxides in particular, sorption of element $E$ can be coupled with its oxidation and the concomitant reductive dissolution of the sorbent mineral phase (e.g. for chromium (Cr), Manceau and Charlet, 1992; for Tl, Peacock and Moon, 2012). According to the rules governing equilibrium stable isotope fractionation, this redox
reaction can also be expected (e.g. for Cr, Schauble et al., 2004), or in select systems has in fact been shown (e.g. for Tl, Peacock and Moon, 2012; Nielsen et al., 2013), to induce stable isotope fractionation of \( E \) during sequestration to the sediments. Despite a number of studies now documenting trace element stable isotope fractionation occurring during sorption to sedimentary mineral phases, relatively few seek to determine the molecular mechanism(s) responsible for the fractionation, and thus provide a fundamental framework for understanding, interpreting and predicting fractionations between palaeo-seawater and contemporaneous trace element sedimentary archives (notable exceptions include, for Mo, Wasylenki et al., 2011; for Ni, Wasylenki et al., 2015; for Cu and Zn, Juliot et al., 2008; Pokrovsky et al., 2008; Little et al., 2014; for Tl, Peacock and Moon, 2012). These studies are crucial for understanding the isotope signals measured in ancient sedimentary archives and subsequently how these archives relate to contemporaneous seawater. In particular, the isotope fractionation of trace elements during sequestration can be the dominant isotope effect in the modern oceans; for example, ferromanganese-rich sediments are known to be the main sedimentary sink for Mo in modern seawater, and during sorption to ferromanganese minerals, light Mo is preferentially sequestered leaving seawater with a heavy isotope signature of 2.3\%\; ; this sequestration process controls the modern global Mo isotope seawater composition (e.g. Barling et al., 2001; Siebert et al., 2003; Barling and Anbar, 2004). Similar sequestration processes are also likely to be important in controlling the global seawater isotope compositions of a range of other trace elements, which are now emerging as potentially important tracers of past biogeochemical processes (e.g. Cu and Zn, Little et al., 2014).

Recent research now indicates that the magnitude and direction of isotope fractionation that occurs during sequestration of trace elements to Fe (and Mn) oxyhydroxide minerals is controlled by the precise molecular mechanism by which the metals are sorbed from solution (e.g. for Ni, Wasylenki et al., 2015). This, in turn, depends on an often complex combination of geochemical, physicochemical and mineralogical factors (e.g. for Ni; Peacock and Sherman, 2007; Peacock, 2009; Peacock et al., 2010). Ultimately, isotope fractionation that occurs as a result of physicochemical processes in seawater must be considered when using sedimentary trace element isotope signals as tracers of contemporaneous seawater isotope composition and, ultimately as proxies for contemporaneous environmental conditions and processes.

### 2.3. Predicted and measured fractionations of trace elements in geological systems

Stable isotope fractionation of the trace elements can be predicted from the theoretical basis for mass-dependent fractionation, established by Urey (1947) and Bigeleisen and Mayer (1947). These calculations are important for the new trace element systems because they provide limits to the fractionations that we might expect to measure in nature, and are particularly useful for systems that cannot be reproduced experimentally (e.g. Johnson et al., 2004). Several of the trace element stable isotope systems have been explored theoretically, including Fe (e.g. Hill and Schauble,
2008), Mo (e.g. Weeks et al., 2007, 2008; Wasylenki et al., 2008, 2011), Cr (e.g. Schauble et al., 2004), Tl (e.g. Schauble, 2007), and Ce (e.g. Nakada et al., 2013). Ideally, calculated fractionations should be verified experimentally, and this is an active research area where several groups are working with both theory and experiment to shed light on measured natural isotope values and isotope variations between reservoirs (e.g. for Mo, Wasylenki et al., 2011; for Cu and Zn, Little et al., 2014).

The Fe and Mo stable isotope systems have perhaps received the most attention in terms of theoretical studies that predict fractionation between different Fe-Fe and Mo-Mo species and experimental studies quantitatively constraining that fractionation in different Fe and Mo systems. As detailed below, both elements experience significant change in their geochemical behaviour with changes in environmental redox conditions, both are important bio-essential metals linked to modern and ancient biogeochemical cycling, and both have revealed a rich sedimentary record of evolving conditions at Earth’s surface.

2.4. Emerging non-traditional stable isotope systems

2.4.1. Iron

Compared to the lower-mass traditional stable isotopes (H, C, N, O and S), fractionations for higher-mass elements are not expected to exceed 10 per mil at temperatures equal or below 100°C (Johnson et al., 2004). Unlike other more volatile isotopic systems (e.g. C and O), Fe isotopes are geochemically robust, retaining their environmental signature even in the face of extreme metamorphic conditions (e.g. Frost et al., 2007; Dauphas et al., 2007), making them a useful tool in deciphering the biogeochemical history of highly deformed and metamorphosed rocks.

Iron has four naturally occurring stable isotopes $^{54}$Fe (5.84%), $^{56}$Fe (91.7%), $^{57}$Fe (2.12%) and $^{58}$Fe (0.28%), where stable isotope data are commonly reported as $^{56}$Fe/$^{54}$Fe. Many natural materials tend to have distinctively narrow $\delta^{56}$Fe ranges, where most igneous rocks fall within the 0.00±0.05 % range, as do riverine and marine sediments, shales, and modern wind-blown sediments (Beard et al., 2003; Fig. 3). In contrast, Fe-rich sediments have characteristically large variations in $\delta^{56}$Fe (Fig. 3). This large variation is interpreted to have been caused by a combination of factors, including mineral-specific equilibrium isotope fractionation, variations in the original Fe isotope composition of the fluids from which these minerals precipitated, and bacterial metabolic processes which might have influenced their formation (Fig. 4; Johnson et al., 2003; 2008b). Metabolically-processed Fe may also retain a ‘vital effect’ due to the redox-related changes driving Fe isotope fractionation during microbially-influenced Fe mineral precipitation (Fig. 4; Johnson et al., 2002).

While Fe isotope systematics has gained widespread attention in applications ranging from chemical weathering, soil formation, the formation of ore deposits, and modern marine metal cycling, their principal application to date in terms of Earth’s redox and biogeochemical evolution has focused on the Precambrian record of Fe isotopes archived in BIF (section 3.1.2).
Figure 3. Iron isotope composition of natural materials. Adapted from Beard et al. (2003).
2.4.2. Molybdenum

The Mo isotope composition of modern and ancient Fe mineral archives has received significant attention due to the strong potential for palaeo-redox reconstruction (Barling et al., 2001; McManus et al., 2002; Siebert et al., 2003; Arnold et al., 2004; Barling and Anbar, 2004; Archer and Vance, 2008; Voegelin et al., 2009; 2010; Planavsky et al., 2014). Molybdenum has seven naturally occurring stable isotopes $^{92}\text{Mo}$ (14.84%), $^{94}\text{Mo}$ (9.25%), $^{95}\text{Mo}$ (15.92%), $^{96}\text{Mo}$ (16.68%), $^{97}\text{Mo}$ (9.55%), $^{98}\text{Mo}$ (24.13%) and $^{100}\text{Mo}$ (9.63%) and Mo isotopic compositions represented in the conventional $\delta$ notation have been primarily reported in two different ways in the literature. The most common reporting is $\delta^{98/95}\text{Mo}$ with respect to the isotope standard "Bern-Mo" (Johnson Matthey ICP standard, lot 602332B; McManus et al., 2002; Siebert et al., 2003; Voegelin et al., 2009, 2010). The second most common reporting is $\delta^{97/95}\text{Mo}$ with respect to Roch-Mo2 (Johnson Matthey Specpure Molybdenum Plasma Standard, lot 7024991; Barling et al., 2001; Barling and Anbar, 2004). Despite this apparent complexity, it is relatively easy to convert from $\delta^{98/95}\text{Mo}$ to $\delta^{97/95}\text{Mo}$ with a simple multiplication factor of 2/3, corresponding to the relative mass difference between these notations. The NIST SRM 3134 Mo standard has since been proposed by Greber et al. (2012) in order to enable easier data comparisons between studies. Goldberg et al. (2013) calibrated the NIST SRM 3134 standard relative to Roch-Mo2 and Bern-Mo as follows: $\delta^{98/95}\text{Mo}_{\text{SRM 3134}} = \delta^{98/95}\text{Mo}_{\text{Roch-Mo2}} + 0.34 \pm 0.05 \%$ and $\delta^{98/95}\text{Mo}_{\text{SRM 3134}} = \delta^{98/95}\text{Mo}_{\text{Bern-Mo}} + 0.27 \pm 0.06\%$.

Figure 4. Pathways of iron cycling and associated isotopic fractionations during the deposition and early diagenesis of iron formation. Adapted from Johnson et al., 2007.
Siebert et al. (2003) were the first to constrain mean ocean $\delta^{98/95}{Mo}$, reporting an isotopically heavy value of $2.63\%\pm 0.1$ (2SD). An even heavier isotopic composition was reported by McManus et al. (2002) for deep marine sediment pore waters, where isotopically light Mo (relative to seawater) accumulates in reducing sediments (the most important marine Mo sink), leaving a low concentration of heavy residual Mo in associated pore waters.

All oceanic Mo sources and sinks have a lighter isotopic composition than seawater (Fig. 5). The isotopic offset between oxic and anoxic sediments is fixed by the isotopic composition of the most important Mo source, river water, which may vary in dissolved Mo isotope composition between 0 and 2.3\% and averages around 0.7\% (Archer and Vance, 2008). The heavy Mo isotopic composition of seawater is largely determined by the areal extent of sediments underlying an anoxic water column, which have a light isotopic composition and are the most important marine sink of Mo (McManus et al., 2002; Siebert et al., 2003). The second most important sink, Fe and Mn oxide-rich sediments, represents an even lighter Mo isotope exit channel, but is less important today than the anoxic sedimentary sink (Arnold et al., 2004). Thus isotopically heavier

![Figure 5](image-url). Mo isotope compositions of marine sediments deposited under diverse water column redox conditions. Adapted from Anbar (2004).
seawater values reflect a rigorous oxic sink removing light Mo isotopes, and isotopically lighter seawater values reflect a more important role for anoxic and euxinic sinks that sequester the heavier isotopes. Uptake of Mo into carbonates is assumed to have an insignificant effect on the Mo isotopic composition of seawater because they contain very little Mo and are thus relatively unimportant from a mass-balance perspective compared to the major sources and sinks identified.

2.4.3. Chromium

The mobile chromium(VI) anion (HCrO$_4^-$/CrO$_4^{2-}$) is the most thermodynamically stable form of Cr in equilibrium with modern atmospheric oxygen. Chromium(III) is the most stable form under reduced conditions, and its mobilization is only prevalent under strongly acidic conditions. The oxidation to Cr(VI) depends upon the solubilisation of primary Cr(III)-bearing mineral phases, such as chromite (FeCr$_2$O$_4$), and the subsequent reaction of dissolved Cr(III) with Mn(IV) oxides (reaction 6). The latter is the only naturally occurring oxidant of Cr(III) at pH<9 (Eary and Rai, 1987), but because it requires the presence of elevated O$_2$ levels to oxidize Mn(II) to Mn(IV), Cr mobility is ultimately tied to environmental oxygenation. Chromium(VI) can be reduced to Cr(III) by microbes (Sikora et al., 2008) and by Fe$^{2+}$ (reaction 7) or Fe(II)-bearing minerals (Ellis et al., 2002). The oxidation of dissolved Fe$^{2+}$ by Cr(VI) proceeds on the order of minutes, and is much faster than with oxygen, even under well aerated, high-pH conditions (Eary and Rai, 1989). This means that in the presence of Fe(II), Cr(IV) is efficiently reduced to Cr(III). The Cr(III) is subsequently and effectively scavenged into Fe(III)-Cr(III) oxyhydroxides owing to the very low solubility of (Fe,Cr)(OH)$_3$ (Sass and Rai, 1987). Moreover, the formation of a solid phase greatly reduces the potential for transformation back to Cr(VI) (Fendorf, 1995).

\[ \text{Cr}^{3+} + 1.5\text{MnO}_2 + \text{H}_2\text{O} \rightarrow \text{HCrO}_4^- + 1.5\text{Mn}^{2+} + \text{H}^+ \]  \hspace{1cm} (6)

\[ 3\text{HCrO}_4^- + 9\text{Fe}^{2+} + 15\text{H}_2\text{O} \rightarrow 3(\text{Fe,Cr})(\text{OH})_3 + 6\text{Fe(OH)}_3 + 15\text{H}^+ \]  \hspace{1cm} (7)

Chromium has four stable isotopes of masses 50 (4.35%), 52 (83.8%), 53 (9.50%) and 54 (2.37%). After Cr(III) oxidation, the Cr(VI) anion is enriched by up to 7% in $^{53}$Cr compared to coexisting compounds containing Cr(III), leaving the aqueous phase with positive $\delta^{53}$Cr values and the residual soils and weathered rock with negative $\delta^{53}$Cr values (Schauble et al., 2004; Zink et al., 2010). This positive Cr(VI) signal should be transferred to the oceans because subsequent sorption of Cr(VI) onto soil and sediment particles apparently produces no isotope effect (Ellis et al., 2004). The microbial reduction of Cr(VI) further generates isotopic shifts of up to 4.1‰, comparable to those produced during abiotic reduction (Ellis et al., 2002). The overall effects of Cr redox reactions are, therefore, to enrich the heavier isotope in the remaining dissolved Cr(VI). Chemical sediments may then capture the signal of oxidative continental weathering if the Cr(VI) pool is large enough to alter the Cr isotope composition of seawater.
3. Iron-based records of Earth’s redox and biogeochemical evolution

3.1. Banded iron formations

One of the unique sedimentological features of the Archaean oceans was the deposition of banded iron formation (BIF). As reviewed below, several aspects of the trace element signatures recorded in these enigmatic deposits are employed to reconstruct Earth’s ancient redox and biogeochemical evolution.

3.1.1. Rare earth elements

Rare earth element (REE) studies in BIF focus on two central facets, namely understanding the sources of Fe to the ancient oceans, and the ancient water column redox conditions. At the heart of REE studies in BIF is the assumption that there is minimal fractionation of REE during precipitation of the primary ferric oxyhydroxides. Iron formations are, therefore, inferred to have trapped a REE signature of seawater at the site of Fe precipitation.

For over half of Earth’s history, seawater was characterized by its high Fe concentrations, leading to the notion that the Archaean oceans (4.0–2.5 Ga) were “ferruginous” throughout much of the anoxic water column. Early studies invoked a continental source of Fe, since Fe(II) was much more mobile during weathering in the absence of atmospheric O\textsubscript{2} (e.g. James, 1954; Lepp and Goldich, 1964) and potentially since continents had a more mafic composition than today (Condie, 1993). Holland (1973) later suggested that Fe was instead sourced from deep marine waters and supplied to the depositional settings via upwelling. With the discovery of modern seafloor-hydrothermal systems (e.g. Corliss et al., 1978) and the subsequent recognition that modern hydrothermal systems may contribute up to 75% of dissolved Fe to the Fe budget in the deep-oceans (Carazzo et al., 2013), submarine venting has become the generally agreed upon source for seawater Fe.

Perhaps the strongest support for a submarine volcanic source for BIF comes from its europium (Eu) enrichment, which indicates a strong influence of high-temperature hydrothermal fluids on the seawater dissolved REE load (e.g. Klinkhammer et al., 1983; Derry and Jacobsen, 1988, 1990). The disparate behaviour of Eu from neighbouring REE in hydrothermal fluids is linked with Eu(III) reduction at high-temperature and low Eh conditions (Klinkhammer et al., 1983; Sverjensky, 1984). It is generally assumed that Fe and REE will not be fractionated during transport from spreading ridges or other exhalative centres, and, therefore, a strong positive Eu anomaly indicates that the Fe in the precursor sediment was hydrothermally derived (e.g. Slack et al., 2007). In this regard, secular trends in the magnitude of Eu anomalies in BIF has historically been assumed to indicate variations in hydrothermal flux, with a long-term decrease in hydrothermal activity from the Eoarchean to Palaeoproterozoic (e.g. Huston and Logan, 2004; Sreenivas and Murakami, 2005). This REE trend is likely linked with an overall decline in the delivery of reductants from Earth’s interior.

Neodymium (Nd) isotopes have also been shown to be highly valuable in tracing the source input of Fe to BIF. For example, a number of BIF record mixing of $^{143}\text{Nd}/^{144}\text{Nd}$ from primarily two sources: (1) anoxic deep water with generally positive initial $\varepsilon\text{Nd}$
(εNd(i)) values reflecting submarine mid-ocean ridge basalt (MORB)-like depleted sources, and (2) shallower waters with generally lower εNd(i), reflecting riverine input from evolved continental sources (e.g. Alibert and McCulloch, 1993; Bau et al., 1997; Frei and Polat, 2007). Newly compiled $^{143}\text{Nd}/^{144}\text{Nd}$ data by Alexander et al. (2009) of BIF older than 2.7 Ga suggest that bulk anoxic Archaean seawater was dominated by high-temperature hydrothermal alteration of mantle-derived oceanic crust, presumably also responsible for delivering the Fe to the BIF. However, deviation from this mantle Nd signal has been recorded in the 2.9 Ga-old Pongola BIF in South Africa, which was deposited in a near-shore, shallow-water environment where continental fluxes of Fe and other solutes contributed greatly to the unit’s composition (Alexander et al., 2009). Similarly, Haugaard et al. (2013) reported positive εNd values in the Si-rich bands of the 2.9 Ga Itilliarsuk BIF of West Greenland, suggesting that those REE were controlled by a local, depleted continental crust, whereas the negative εNd(i) values found in the Fe-rich layers were hydrothermally-sourced. Nevertheless, from Archaean to Proterozoic, there is a gradual decrease in hydrothermal activities as the Earth cooled, and a stronger continental contribution of Nd to the ocean (Miller and O’Nions, 1985; Jacobsen and Pimentel-Klose, 1988).

Anomalies in cerium (Ce) abundance in BIF have been used to constrain water column redox conditions. In general, oxygenated marine settings display a strong negative Ce anomaly when normalized to shale composite ($\text{Ce}_{(\text{SN})}$), while suboxic and anoxic waters lack significant negative $\text{Ce}_{(\text{SN})}$ anomalies and can even show positive anomalies (e.g. German and Elderfield, 1990; Byrne and Sholkovitz, 1996). Oxidation of Ce(III) to Ce(IV) greatly reduces Ce solubility, resulting in its preferential removal onto Mn(IV)-Fe(III)-oxyhydroxides, organic matter and clay particles. In contrast, suboxic and anoxic waters lack significant negative $\text{Ce}_{(\text{SN})}$ anomalies due to reductive dissolution of settling Mn(IV)-Fe(III)-oxyhydroxide particles. Similarly, light REE depletion develops in oxygenated waters due to preferential removal of light vs. heavy REE onto Mn(IV)-Fe(III)-oxyhydroxides and other particle-reactive surfaces. As a result, the ratio of light-to-heavy REE markedly increases across redox boundaries due to reductive dissolution of Mn(IV)-Fe(III)-oxyhydroxides (German et al., 1991; Byrne and Sholkovitz, 1996). In many Archaean and early Palaeoproterozoic BIF there are no Ce anomalies, and thus no deviation from trivalent Ce behaviour (e.g. Alexander et al., 2008; Frei et al., 2008), suggesting that the water column from which ferric oxyhydroxides precipitated was anoxic (Bau and Dulski, 1996). In support of this model, a recent survey of 18 different Palaeoproterozoic and Archean BIF did not display significant Ce anomalies until after atmospheric oxygenation at ~2.5–2.4 Ga (Planavsky et al., 2010a).

There also appear to be differences in trivalent REE behaviour in BIF before and after the permanent rise of atmospheric oxygen. Late Palaeoproterozoic BIF show significant ranges in light-to-heavy REE (Pr/Yb$_{(\text{SN})}$) ratios, both below and above the shale composite value (Planavsky et al., 2010a). This range of light-to-heavy REE and Y/Ho ratios in late Palaeoproterozoic BIF likely reflects variable fractionation of REE + Y by Mn(IV)-Fe(III)-oxyhydroxide precipitation and dissolution. This interpretation implies deposition of late Palaeoproterozoic BIF at ~1.9 Ga in basins with varying
redox conditions and a strong redoxcline separating an upper oxic water column from deeper waters that were suboxic to anoxic (Planavsky et al., 2009). These ranges are also similar to those seen in modern anoxic basins. In contrast, most Archaean and early Palaeoproterozoic BIF deposited before the rise of atmospheric oxygen are characterized by consistent light REE depletion (Planavsky et al., 2010a). This consistent depletion in light REE suggests the lack of a discrete redoxcline and points toward O$_2$-independent Fe(II) oxidation mechanisms.

3.1.2. Trace element isotope compositions

In addition to REE signatures recorded in BIF, Fe isotope signatures archived in these deposits are also used to shed light on ancient redox conditions. The primary Fe minerals that ultimately sedimented to yield BIF were probably formed in the water column as the result of oxidation of Fe(II)$_{aq}$, where overall fractionation appears to be independent of the oxidative pathway, whether by molecular oxygen, anaerobic photoferrotrophy or UV photo-oxidation (Fig. 4; e.g. Bullen et al., 2001; Croal et al., 2004; Balci et al., 2006; Staton et al., 2006). While complete oxidation of an Fe(II)$_{aq}$ pool will result in ferric oxyhydroxides with a $\delta^{56}$Fe composition equal to that of the initial pool, partial oxidation will result in ferric oxyhydroxides that are up to +4‰ higher (e.g. Wu et al., 2012). Reactions between these ferric oxyhydroxide precipitates and various Fe(II)$_{aq}$ pools as they descend through the water column and their diagenesis upon burial result in their conversion to important BIF forming minerals (e.g. hematite, magnetite, siderite, ankerite).

In Precambrian BIF, positive $\delta^{56}$Fe signatures are thought to occur via the interaction of ferric oxyhydroxides formed by partial oxidation of hydrothermally-sourced Fe(II)$_{aq}$, with more pristine pools of hydrothermally sourced Fe(II)$_{aq}$ contributing near-zero $\delta^{56}$Fe values. Such a scenario is easily envisioned for oxidative mechanisms restricted to the upper water column, where isotopically heavy precipitates formed in a zone of partial oxidation rain into a deeper, isotopically pristine hydrothermal Fe(II)$_{aq}$ pool. Alternatively, positive signatures may be generated via the partial or near-complete reduction of ferric oxyhydroxides of nearly any initial isotope composition. Dissimilatory Fe reduction (DIR) by Fe reducing bacteria releases isotopically light Fe(II)$_{aq}$ and leaves the residual oxide isotopically heavy (Fig. 4). This appears reflected in the Fe isotope composition of mineral separates from Precambrian BIF, where hematite is systematically heavier than magnetite, the latter likely formed by reaction of primary Fe oxyhydroxides with isotopically light Fe released by DIR (Johnson et al., 2008a). With few electron acceptors available for anaerobic respiration in marine sediments on the O$_2$-poor early Earth, DIR would have probably constituted a major remineralization pathway during sedimentary diagenesis (Konhauser et al., 2005). The Fe isotope record in both sedimentary pyrites and Precambrian BIF (Johnson et al., 2008b) demonstrates a profound excursion to extremely negative Fe isotope values ~2.7 Ga that would appear to reflect increasingly important DIR in the run-up to Earth surface oxygenation (Fig. 6). In this model, extremely light Fe isotope compositions reflects a period in Earth’s history where reduction of increasingly abundant
Figure 6. Iron isotope compositions of sedimentary sulfide minerals (a), iron formations (b) and whole-rock data from shales (c Authors, correct?) through the Archaean to Mesoproterozoic. Adapted from Johnson et al. (2008) and Planavsky et al. (2012).

sedimentary Fe oxyhydroxides was one of the most important organic carbon remineralisation pathways in deep marine settings on the O2- and SO4-poor early Earth.
The apparently sensitive character of Mo isotope redox proxies has also been exploited in Precambrian Fe formations in the search for the oldest traces of free O$_2$ on Earth. Examining a suite of Precambrian Fe formations of various ages, Planavsky et al. (2014) revealed a trend between Mo isotope compositions and Fe/Mn ratios (Fig. 7), with isotopically lighter Mo isotope compositions occurring as Fe/Mn decreases (and thus Mn increases). This is interpreted as a redox array where under more reducing conditions, when conditions are prohibitive to Mn(II) oxidation and Mn(III,IV) oxide deposition, Mo isotope compositions of BIF tend towards values expected for equilibrium isotope fractionation between Fe oxyhydroxides and seawater (with Mo in Fe oxyhydroxides being $\sim$1‰ lighter than in coeval seawater); when Mn(III,IV) oxides are deposited under more oxidizing conditions, the greater equilibrium Mo isotope fractionation between Mn oxides and seawater is expressed (with Mo in Mn oxides being $\sim$2‰ lighter than coeval seawater). Molybdenum isotope compositions as light as $-0.9‰$ in the 2.95 Ga Sinqeni Fe formation (Pongola Supergroup, S. Africa) could be generated only by Mo adsorption onto Mn(III,IV) oxides during deposition of this deposit; in the study of Planavsky et al. (2014), the Mo isotope data appear to be signalling the ancient presence of Mn(III,IV) oxides, and thus significant O$_2$, in the

![Figure 7](image-url)

*Figure. 7.* Mo isotope compositions of diverse Precambrian iron formations whose trend as a function of Fe/Mn ratios indicate adsorption of Mo onto Mn oxides, and by consequence the presence of appreciable free O$_2$, as far back as 2.96 Ga. Adapted from Planavsky et al. (2014).
environment despite the fact that no Mn(III,IV) oxides are currently found in the deposit (having likely been the subject of dissimilatory Mn(III,IV) reduction, leaving the Fe formation enriched only in Mn(II)). These data are amongst Earth’s earliest evidence for free O$_2$ in the environment, and place important constraints on the timing of the evolution of oxygenic photosynthesis. Mo isotope proxies have also been widely applied in the study of evolving marine palaeo-redox using the more extensive shale record (section 3.3.2).

3.1.3. Trace element concentrations

In natural systems where trace element sequestration by authigenic ferric oxyhydroxides results from a continuum of adsorption and co-precipitation reactions, lumped-process distribution coefficient models can be used to relate the concentration of an element in the Fe oxide to the dissolved concentration present at the time of precipitation, i.e. they reflect contemporaneous seawater trace element signatures. This predictive aspect of metal sorption reactions has been exploited to better understand the BIF record with respect to ancient seawater composition and nutrient limitations on Precambrian primary productivity.

The phosphorus (P) content of BIF has received significant attention as ferric oxyhydroxides possess a strong affinity for dissolved phosphate, and phosphate is a bio-essential nutrient whose availability can limit phytoplankton productivity in modern oceans. Today, particles of ferric oxyhydroxide that precipitate out of some marine hydrothermal vent fluids upon mixing with oxygenated seawater have been noted to strongly adsorb phosphate from seawater, to the point where the resulting particles attain molar P/Fe(III) ratios of over 0.2 (Feely et al., 1990, 1998). Seawater phosphate concentrations at hydrothermally active spreading centres may be strongly affected by adsorption onto ferric oxyhydroxides; at the Juan de Fuca ridge for example, significant negative phosphate anomalies have been noted at depth, with phosphate declining most appreciably (by approximately 60 nM) nearest to the ridge axis, coincident with increased sediment Fe and P concentrations (Feely et al., 1990). Later work by Wheat et al. (1996) indicated that marine hydrothermal systems annually remove about 50% of the preindustrial dissolved riverine phosphate flux, and that the co-precipitation of phosphate with ferric oxyhydroxides in seawater accounts for 36–66% of that removal rate. The extent to which phosphate is incorporated into the particles depends on the available phosphate concentration, in a manner that can be described using a distribution coefficient ($K_D$):

$$K_D = \frac{P/Fe(III)}{[P]}$$

where $P/Fe(III)$ represents the particle’s solid phase molar ratio, and $[P]$ represents the dissolved phosphate concentration at the time of particle precipitation. Both experimental studies (Konhauser et al., 2007) and field data (Feely et al., 1998) demonstrate that this relationship is well described at pH ~8 by a $K_D$ value of approximately 0.06–0.075 mM$^{-1}$. 

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Assuming that the ferric oxyhydroxides precipitates originally constituting Precambrian BIF behaved similarly, Bjerrum and Canfield (2002) applied a $K_D$ value of 0.06 mM\(^{-1}\) to extrapolate Precambrian ocean phosphate concentrations from the concentrations of Fe(III) and P retained in BIF. Their results indicated that ocean phosphate concentrations might have been low enough (10 to 25% of present day values) to significantly limit biological productivity prior to 1.8 Ga.

Konhauser et al. (2007) re-examined the BIF palaeo-proxy for P by evaluating the role of Si as a ferric oxyhydroxide co-precipitate and phosphate adsorption competitor. Throughout the Precambrian, the oceans had a significantly higher concentration of dissolved Si, at least as high as at saturation with cristobalite (0.67 mM at 40°C in seawater), and possibly even amorphous Si (2.20 mM) (Maliva et al., 2005). Silica concentrations close to or at saturation with respect to amorphous Si likely persisted until the early Cretaceous, when the disappearance of VMS-associated hematitic jasper deposits indicate that diatoms or other Si-secreting organisms began actively scavenging Si (Grenne and Slack, 2003), eventually to reduce its concentration to a modern average that is less than 0.10 mM (Tréguer et al., 1995). Crucially, Si strongly adsorbs to ferric oxyhydroxides and effectively competes with phosphate for available adsorption sites. This was demonstrated experimentally by Konhauser et al. (2007), who used a revised $K_D$ approach accounting for Fe-Si co-precipitation indicates that with sufficiently high Si concentrations, the scavenging of phosphate by Fe-rich hydrothermal fluid precipitates is minimal (Fig. 8). The BIF P record, interpreted in this light, might indicate that phosphate was more abundant in Precambrian oceans than today. Additionally, high levels of carbonate saturation during the Precambrian (e.g. Grotzinger, 1990) would have inhibited carbonate fluorapatite (CFA) formation, which today is the largest burial flux for phosphorous in modern oceans (Ruttenberg and Berner, 1993). Combined, these two factors suggest higher, rather than lower dissolved Archean phosphorous concentrations. A subsequent increase in P content of BIF in the Neoproterozoic, following Snowball Earth deglaciations, may then have led to enhanced cyanobacterial photosynthesis, which in turn, produced enough oxygen to facilitate the evolution of animal life (Planavsky et al., 2010b; Fig. 9a). Jones et al. (2015) recently re-evaluated the partitioning of P in solutions even more closely approximating seawater, and found that beyond Si, Ca and Mg can have equally important effects on P partitioning to the surfaces of Fe oxyhydroxide minerals, and the subject of calibration of adsorption-dependent Fe oxyhydroxide proxies remains an area of active investigation.

The Ni content of BIF has also received significant attention because it has been shown that Ni concentrations in BIF have changed dramatically over time, and that a drop in Ni availability in the oceans around 2.7 Ga would have had profound consequences for microorganisms that depended on it, that being methane-producing bacteria called methanogens (Konhauser et al., 2009, 2015; Fig. 9b). These bacteria have a unique Ni requirement for their methane-producing enzymes, and crucially, these bacteria have been implicated in controlling oxygen levels on the ancient Earth as the methane they produced was reactive with oxygen and kept atmospheric oxygen...
levels low. It is possible that a Ni famine eventually led to a cascade of events that began with reduced methane production, the expansion of cyanobacteria into shallow-water settings previously occupied by methanogens, and ultimately increased oxygenic photosynthesis that tipped the atmospheric balance in favour of oxygen, the so-called GOE at 2.45 Ga. Perhaps it is no coincidence that at this time stromatolites expanded and diversified into shallow-water settings (McNamara and Awramik, 1992), and with increased consumption of CO₂, indiscriminate sedimentation of massive amounts of calcium carbonate took place in situ on the sea floor, leading to decimetre- to metre-thick beds that extended over thousands of square kilometres (Grotzinger and Knoll, 1999).

A recent compilation of Cr enrichment in BIF shows a profound enrichment coincident with the GOE around 2.45 Ga (Konhauser et al., 2011; Fig. 9c). After that Cr enrichments started to increase in shallow-water BIF and peaked synchronous with the permanent loss at ~2.32 Ga of mass-independent fractionation of sulfur isotopes that defines the GOE (Bekker et al., 2004, Guo et al., 2009). Given the insolubility of Cr minerals, its mobilization and incorporation into BIF indicates enhanced chemical weathering at that time, most likely associated with the evolution of aerobic continental pyrite oxidation. As the GOE commenced, aerobic chemolithoautotrophic weathering

Figure 8. Experimentally-determined partitioning coefficients ($K_D$; black lines and bold text) for partitioning of phosphate between simulated seawater and ferrihydrite under different ambient dissolved silica conditions (red squares, orange triangles, and green circles represent 0, 0.67, and 2.2 mM dissolved Si, respectively). Also shown are data for modern hydrothermal plumes worldwide (blue circles). This experiment demonstrates that suppressed partitioning of P into iron oxyhydroxides under Precambrian marine Si concentrations may be responsible for the low P/Fe ratio of BIF (area shaded grey). Adapted from Konhauser et al. (2007).
of pyrite on land produced significant acidity, which would have increased sulfate and nutrient fluxes to seawater (Reinhard et al., 2009; Konhauser et al., 2011; Bekker and Holland, 2012). This profound shift in weathering regimes constitutes Earth’s first acid continental drainage system, and accounts for independent evidence of increased supply of sulfate (Bekker et al., 2004) and sulfide-hosted trace elements to the oceans at that time (Scott et al., 2008). With increased sulfate supply to the oceans, marine sulfate-reducing bacteria would have begun to dominate the anoxic layers of the water column, leading to the further marginalization of methanogens to the sulfate-poor bottom sediments, as is the case today. The end result of the progressive decline in methane production (a potent greenhouse gas) was a cooling climate, eventually culminating in the onset of major Palaeoproterozoic glaciations (Zahnle et al., 2006).

During the past few years, Cr isotopes in BIF have also been used to track the progressive oxygenation of Earth’s surface environment during the Archaean and Palaeoproterozoic (Fig. 9d). Frei et al. (2009) measured slight positive Cr
fractionations (+0.04 to +0.29‰) in samples deposited between 2.8 and 2.6 Ga, which they suggested was a first-order proxy for the presence of Cr(VI) in seawater at that time. Because of the efficient reduction of Cr(VI) by Fe^{2+} (which was abundant in the highly ferruginous Archaean oceans), and the near instantaneous precipitation of (Fe,Cr)(OH)₃ at seawater pH, the stable Cr isotope signatures of BIF should thus mirror the initial reaction of Cr(III) from mineral dissolution on land with MnO₂. Interestingly, BIF deposited between 2.5 and 1.9 Ga show minimal positive fractionations in Cr isotopes, suggesting that oxygen levels then dropped in the aftermath of the GOE. This is an unusual finding in that it was previously believed that atmospheric O₂ levels progressively increased post GOE (Kump, 2008), but the Cr isotope record in BIF instead documents transient fluctuations in atmospheric oxygenation. Most recently, Crowe et al. (2013) reported $^{53}$Cr-depletion in 2.96 Ga palaeosols (ancient soil horizons) and $^{53}$Cr-enrichment in overlying 2.96 Ga BIF. These near-contemporaneous units thus seem to have captured the Cr(VI) pool originating from oxidative continental weathering to chemical sedimentation some 500 Ma before the GOE. Based on the assumption that export of Cr(VI) from the weathering environment to the oceans necessitate minimal encounter with Fe(II) in rivers or groundwater, those authors further calculated that oxygen concentration were possibly as high as $3 \times 10^{-4}$ present atmospheric levels.

Frei et al. (2009, 2012) reported further strong positive fractionations ($\delta^{53}$Cr up to +4.9‰) in the late Neoproterozoic, suggestive of increased surface oxygenation at that time. Importantly, this work provided supporting evidence that linked a significant rise in global oxygenation to the emergence of early animals (e.g. Planavsky et al., 2010b). If positive $\delta^{53}$Cr are indicative of oxidative weathering, then the lack of such fractionations suggests minimal O₂ (to produce the MnO₂ catalyst). Indeed, in a recent study of granular Fe formations (reworked BIF) that were deposited between 1.7 and 0.9 Ga, Planavsky et al. (2014) observed $\delta^{53}$Cr of −0.12‰. They hypothesized that the lack of Cr redox cycling must reflect atmospheric O₂ levels <0.1% present atmospheric levels, values below the theoretical estimates for the minimum O₂ requirements of the earliest animals. In other words, after the GOE oxygen levels plummeted, and it was not until the Neoproterozoic that O₂ levels increased sufficiently to facilitate animal evolution.

Cobalt is a bio-essential trace element and co-limiting (e.g. with Fe) nutrient in some regions of the modern oceans. It has been proposed that Co was more abundant in poorly ventilated Precambrian oceans based on the greater utilization of Co by anaerobic microbes relative to plants and animals. Based on the sedimentary record of Co enrichments and a model of marine Co sources and sinks (Fig. 9e), Swanner et al. (2014) proposed that before ~1.8 Ga the large volume of hydrothermal fluids circulating through abundant submarine ultramafic rocks and the predominance of anoxic sediments with a negligible capacity for Co burial resulted in a large marine Co reservoir. A decrease in marine Co concentrations in the Middle Proterozoic resulted from waning hydrothermal Co sources and the expansion of oxic and euxinic sediments that buried Co with Mn(III,IV) oxides or pyrite, respectively. It is interesting to note that the initial diversification of eukaryotes, and thus eukaryotic Co-utilizing genes,
occurred in the middle to late Proterozoic (e.g. Knoll et al., 2007), at a time when marine Co concentrations were declining. Along similar lines, marine cyanobacteria likely utilize Co-binding ligands as a strategy for Co acquisition (Saito and Moffett, 2001; Saito et al., 2005), which may help to explain their adaptation to low-Co environments despite high physiological demand. Perhaps this suggests that the number of Co-utilizing genes encoded by an organism’s genome do not always mirror the metal’s availability.

The lack of a clear link between metabolic metal demands and environmental concentrations is most clearly demonstrated by Zn. The relatively late proliferation of Zn metallomes by eukaryotes had previously been linked to marine Zn biolimitation (Williams and da Silva, 1996; Saito et al., 2003; Dupont et al., 2010). However, a recent examination of the BIF and shale records (Robbins et al., 2013; Scott et al., 2012, respectively; Fig. 9f) indicate that Zn may have been near modern abundances and was likely bioavailable to eukaryotes throughout the Precambrian, casting doubt on the coupled geochemical and eukaryotic evolutions with respect to Zn utilization. A novel possibility stemming from this work is that the late proliferation of Zn metalloenzymes in eukaryotes could have been a biologically intrinsic process related to the regulation of increasingly complex genomes, rather than solely dependent on the dramatic changes in the marine bioavailability of aqueous Zn species.

3.2. Ferromanganese crusts

Ferromanganese crusts have received significant attention due their ability to sorb certain trace elements from seawater, in particular Mo and Co. Molybdenum is supplied to the oceans primarily by the riverine flux of molybdate (MoO$_4^{2-}$) derived from the oxidative weathering of sulfide minerals (e.g. Archer and Vance, 2008). The molybdate anion is highly conservative, with a long residence time, in today’s well mixed oceans. As a result, Mo is the most abundant transition element in the modern oxygenated oceans, at an average concentration of ~100 nM (Bertine and Turekian, 1973). The only significant sink for molybdate in well oxygenated environments is believed to be co-precipitation and burial with Mn oxyhydroxides, but this is a very slow process that removes only minor amounts of Mo from the ocean today (Bertine and Turekian, 1973). The adsorption of Co(II) and Co(III) to surface sites on ferric oxyhydroxides and Mn(III,IV) oxides, respectively, is an important pathway for scavenging of Co under oxic conditions (Koschinsky and Hein, 2003; Takahashi et al., 2007; Stockdale et al., 2010). By contrast, soluble Co maxima occur below the O$_2$-H$_2$S chemocline in modern euxinic basins in conjunction with both the soluble Fe and Mn peaks (Viollier et al., 1995), which indicates that Co is released via the reductive dissolution of both Fe(III) oxyhydroxides and Mn(III,IV) oxides.

3.3. Black shales

3.3.1. Trace element concentrations

In modern sedimentary environments, black shales are used to examine palaeo-productivity based on trace element enrichments that correspond to increased organic
carbon burial (e.g. Tribovillard et al., 2006) or the concentrations of trace elements relative to total organic carbon (Me/TOC) (e.g. Alego and Rowe, 2012). This linkage is mainly due to trace elements with multiple redox states being precipitated and enriched during periods of reducing conditions, e.g. uranium (U) and Mo under sulfidic conditions or Cr under ferruginous conditions (Arnold et al., 2004; Dahl et al., 2013; Partin et al., 2013; Reinhard et al., 2013) or incorporated into microbial biomass, e.g. Ni, Cu, Zn and Cd (Twining and Baines, 2013). Indeed, the utility of shales as palaeo-marine proxies has grown in recent years as an indicator of palaeo-environmental conditions on the ocean floor (i.e. oxic, ferruginous or euxinic) with the advent of Fe speciation analyses that assess the degree of pyritisation by comparing the ratio of pyrite Fe to reactive Fe (e.g. Raiswell and Canfield, 1998; Poulton and Canfield, 2005).

Black shales represent an important fraction of all ancient marine sediments preserved, and because of this, their trace element and isotope compositions have been used to track major biogeochemical events in Earth’s history (e.g. Lyons et al., 2009). This includes, amongst others: (1) the rise of atmospheric oxygen, (2) the onset of oxidative continental weathering, (3) the establishment of aerobic marine ecosystems, (4) the post-GOE drop in atmospheric O₂, and (5) the extent of Proterozoic marine euxinia. Arguably the smoking gun for the rise of atmospheric O₂ (i.e. the GOE) comes from the loss of sulfur isotope mass-independent fractionation (S-MIF) (Farquhar et al., 2000). The processes that lead to S-MIF are restricted to the photochemical dissociation of SO₂ in the upper atmosphere into elemental and water-soluble S species. They then get rained out and incorporated into diagenetic pyrites, such as those associated with black shales. The only way to preserve the S-MIF signal is under low atmospheric O₂ levels because O₂ erases the unique signatures of these reactions and the presence of ozone prevents the penetration of ultraviolet radiation that generates the signals. The last appearance of S-MIF can be dated to pyrites in 2.32 Ga-old black shales of the Rooihoogte/Duitschland and Timeball Hills formations in South Africa, marking this point in Earth’s history as a time of atmospheric oxygenation (Bekker et al., 2004; Guo et al., 2009).

In the past decade, several geochemical studies have been suggestive of oxidative continental weathering having occurred prior to the GOE. For example, increases in Mo and rhenium (Re) concentrations, as well as the first trend towards loss of S-MIF, comes from the 2.5 Ga Mount McRae Shale of Western Australia (Anbar et al., 2007; Kaufman et al., 2007). The trace element enrichments were interpreted as reflecting oxidative weathering of terrestrial sulfide minerals (but at much lower levels than at present), their transport to the oceans as dissolved oxyanion forms (e.g. MoO₄²⁻ and ReO₄⁻), and then their being scavenged into black shales. The shales lack U enrichment, which, because it is predominantly hosted in silicates, would be less affected by rising atmospheric oxygen levels. Interestingly, Mo and Re enrichments were not observed in the upper sections of the shales, suggesting a subsequent drop in atmospheric O₂, or as coined by Anbar et al. (2007), the transient O₂ increase reflected a “whiff” of oxygen. Similar patterns of these metals had been reported in the 2.64–2.50 Ga Ghaap Group in South Africa (Wille et al., 2007). Most recently, Stueken et al. (2012) suggested that an
increase in the total sulfur and Mo supply to marine shales at 2.8 Ga was best explained by the biological oxidation of crustal sulfide minerals.

Evidence in support of transient rises in atmospheric O\textsubscript{2} prior to the GOE implies the evolution and establishment of cyanobacteria in the ocean-water column. The presence of dissolved O\textsubscript{2} in seawater as early as 2.7 Ga is inferred from (i) black shales based on shifts toward heavier $\delta^{15}$N values in kerogen consistent with the onset of an oxic nitrogen cycle (Godfrey and Falkowski, 2009), (ii) C isotope data in kerogens indicating methanotrophy (Eigenbrode and Freeman, 2006; Thomazo et al., 2013), and (iii) coupled Fe and Mo isotopes that indicate the formation of ferric oxyhydroxides in the upper water column which were likely linked to cyanobacteria rather than photoferrotrophs (Czaja et al., 2012): by 2.56 Ga, O\textsubscript{2} levels may have already reached several hundred metres on the continental slope (Kendall et al., 2010). Based on the presence of 10 wt.% organic content in the 3.2 Ga Gorge Creek Group in north-western Australia, and the lack of Fe and sulfur, it is even possible that oxygenic photosynthesis may have evolved then. As Buick (2008) suggested, it is difficult to explain another mode of primary productivity that would have had enough available substrates over such wide an area for such lengths of time.

Until a few years ago, most researchers were of the opinion that once O\textsubscript{2} accumulated in the oceans and atmosphere it would have continued to rise until modern values were reached, sometime in the late Neoproterozoic (e.g. Kump, 2008). However, a study focusing on the temporal trends of U enrichment in black shale demonstrated that the initial rise of atmospheric O\textsubscript{2} at $\sim$2.45 Ga was followed by a dramatic decline to less oxidizing conditions only several hundred million years after the GOE itself (Partin et al., 2013). Somewhat similar trends were observed in the Mo and Mo/TOC records in black shales, but with evidence for a large oceanic Mo inventory occurring only at 2.2 Ga and then dropping to 10–20% of modern ocean levels until the Phanerozoic (Scott et al., 2008). The smaller seawater Mo inventory, in the immediate aftermath of the GOE, was interpreted to reflect the Mo distribution in the continental crust, as has been argued for Re by Hannah et al. (2004). The present continental flux of Mo, like that of Re, is largely derived from organic matter-rich sulfidic shales; at the time of the GOE, the pre-GOE organic matter-rich sulfidic shales were not significantly enriched in Mo above crustal levels (Yamaguchi, 2002; Scott et al., 2008). The predominant crustal reservoir of Mo that oxidized to form molybdate at this time would have been crystalline crustal rocks with generally low Mo content. This would result in a small supply of Mo to the oceans in the immediate aftermath of the GOE. The time for ingrowth of a large reservoir of Mo-enriched shales may have been comparable to the crustal residence time (200 Myr), and the process would ultimately lead to higher oceanic Mo contents as its concentration increased in the source rocks (e.g. black shales) that were being weathered.

The solubility and burial of U and Mo also depends on the sulfide concentration because both are chalcophilic. Therefore, in modern euxinic environments, such as the Black Sea, Mo sinks outpace Mo supply. Low Mo concentrations develop in the water column, and the magnitude of Mo enrichment and Mo/TOC ratios in sediments drop
precipitously (Algeo and Lyons 2006). This feature of Mo geochemistry forms the basis for the ‘bio-inorganic bridge’ hypothesis, whereby changes in the chemistry of the early oceans directly affected the bioavailability of trace elements, and in turn, the evolution of life (Anbar and Knoll, 2002). Those authors suggested that the drawdown of Mo under euxinic conditions limited primary photosynthetic productivity in the photic zone of the oceans for over a billion years between the late Palaeoproterozoic and Neoproterozoic (the so-called ‘boring billion’; Buick et al., 1995). The cause for the global euxinia was attributed to the progressive rise in atmospheric O$_2$ after the GOE that increased pyrite oxidation on land, which in turn led to a greater supply of dissolved sulfate to the oceans, enhanced sulfate reduction in the oceans, and the generation of sufficient dissolved sulfide that the total dissolved Fe flux in the oceans was titrated out by 1.8 Ga (Canfield, 1998). It was not until the Neoproterozoic, in association with a second large oxidation of the Earth’s surface, that the deep oceans became oxygenated. However, the observed Mo drawdown and complementary Mo isotope data during that time interval are inconsistent with ocean-wide euxinia (e.g. Arnold et al., 2004), and a recent investigation of sedimentary Mo and Cr enrichments in black shales, using mass-balance box models, demonstrated that only $\sim$1–10% of the modern seafloor area needed to be anoxic at that time (Reinhard et al., 2013).

### 3.3.2. Molybdenum isotopes in shales

When Mo removal from seawater is quantitative, the Mo isotope composition of reducing sediments may be directly linked to the isotopic composition of contemporaneous seawater, and in turn, to the relative importance of reducing and oxic areas of the seafloor. This property of the Mo isotope system has been extensively exploited to reconstruct palaeo-redox from black shales, whose organic-rich nature testify to depletion of available electron acceptors in the sedimentary pile and the establishment of reducing conditions (either anoxic or euxinic) amenable to recording the Mo isotope composition of contemporaneous seawater (section 2.4.2). Using a conservative assumption of 70% Mo removal by euxinic black shales in the mid-Proterozoic, Arnold et al. (2004) used simple isotope mass balance to calculate the Mo isotopic composition of mid-Proterozoic seawater to be around 0.8‰. This isotopically light value for seawater was interpreted to reflect expanded euxinia and a suppressed oxic sink. Assuming a riverine source near 0‰, Arnold et al. (2004) initially estimated a ratio between the fractional importance of oxic ($f_{ox}$) and euxinic ($f_{eux}$) sinks as being 10 times lower than today ($f_{ox}/f_{eux} < 0.4$ in the mid-Proterozoic vs. $\sim 3$ today). Using a newly-determined modern riverine source value of $\sim 0.7‰$, Archer and Vance (2008) re-interpreted the low isotopic composition of mid-Proterozoic black shales (relative to modern seawater) as reflecting near total drawdown of the marine Mo reservoir as the result of expanded euxinia, with sedimentary Mo isotope compositions tending towards the source composition (mostly river water). In other words, Mo isotope data from shales reveal that mid-Proterozoic oceans were characterized by important euxinic Mo drawdown and little evidence for any important role for oxic sedimentary Mo exit channels. Similarly light (relative to modern seawater) Mo sedimentary isotope
compositions indicating near total absence of oxic Mo exit channels have been reported for black shales from the 2.05 Ga Palaeoproterozoic Shunga carbon burial event in the aftermath of the GOE (Asael et al., 2013), during the mid-Neoproterozoic in the run-up to the appearance of the Ediacaran fauna (Dahl et al., 2011), and during Phanerozoic ocean anoxic events (OAE) such as the Toarcian (early Jurassic) OAE (Pearce et al., 2008). Using a mass-balance model with spatially variable metal burial rates and by also considering Cr (whose marine exit channels appear insensitive to the distinction between anoxia and euxinia), Reinhard et al. (2013) refined sedimentary Mo exit channels under low-O$_2$ (anoxic and euxinic) conditions, finding that a relatively small spatial extent of euxinia (1–10% of the seafloor, vs. 0.1% today) would be sufficient to crash the marine dissolved Mo reservoir, and by inference, draw seawater Mo isotope compositions towards riverine sources. Early models indicating widespread global euxinia throughout mid-Proterozoic oceans (e.g. Canfield et al., 1998) are thus better characterized by expanded but spatially-restricted euxinia combined with widespread anoxia.

Molybdenum isotope-based redox proxies appear to be very sensitive to even the smallest amounts of free oxygen, and are now being used to examine trace amounts of dissolved Mo that may have been released prior to the ~2.4 Ga GOE and sustained oxidation of crustal Mo sources. Wille et al. (2007) demonstrated small but non-negligible Mo isotope deviations from crustal values in 2.6–2.5 Ga black shales from the Transvaal Supergroup, South Africa, indicating the production of some free O$_2$ at that time. In only slightly younger (2.50 Ga) black shales in the Hamersley basin, Western Australia, work by Anbar and colleagues (Anbar et al., 2007; Duan et al., 2010) has revealed coupled Mo abundance and isotope excursions occurring while the atmosphere remained anoxic, re-enforcing the idea that small, local oxygen-production events (i.e. “whiffs”) likely preceded global oxygenation of the atmosphere. Kurzweil et al. (2015) extended the Hamersley Mo isotope dataset back by nearly 100 Ma to reveal a long-term trend of increasingly heavy Mo isotope compositions upwards in the basin. This would indicate a gradual increase in the prevalence of oxic Mo exit channels for seawater Mo prior to the GOE, consistent with the early occurrence of non-negligible Mo isotope fractionation as reported by Wille et al. (2007) for the contemporaneous Griqualand West basin, S. Africa.

4. Conclusions and future directions

By nature of their physicochemical surface properties, redox active Fe minerals are extremely efficient scavengers of trace elements from natural waters and so can be used to provide a rich archive of trace element concentrations and stable isotope compositions in terrestrial and marine environments over the entire history of their formation. These trace element signals can then be interpreted to shed light on environmental conditions in these environments at the time of mineral precipitation. Three Fe mineral archives are currently employed to help understand the evolution of Earth’s redox and biogeochemical conditions, namely banded iron formations (BIF),
ferromanganese crusts and black shales, where although shales are not technically classed as Fe-rich sediments, it is likely that many of their trace elements are sequestered in association with Fe minerals. In these sediments Fe oxyhydroxides, including ferrihydrite, goethite, hematite and magnetite, and Fe sulfides, including pyrite, provide the primary Fe-rich phases for trace element sorption.

To date Fe mineral archives have provided a wealth of trace element concentration and stable isotope data which has been interpreted to reflect environmental conditions during Earth’s earliest oceans and atmosphere, during the Archaean in the run up to and events after the Great Oxidation Event (GOE), through subsequent Ocean Anoxic Events (OAE), and into the relatively recent geological history of the Cenozoic. In light of these studies, it is certain that these enigmatic deposits have revolutionised our understanding of the Earth system. To continue forward with their investigation and bring new insights into Earth’s redox and biogeochemical evolution, it is apparent that we must continue to improve our understanding of both the Fe mineral archives and the behaviour of their trace element impurities, both at the small-scale within the archives and at the larger-scale within the environment as a whole. By continuing to investigate the precipitation, deposition and diagenesis of Fe minerals, and both the macro- and microscopic uptake of trace elements to these minerals, we can be increasingly confident in the fidelity of trace element signals recorded in Fe mineral archives, and better able to relate these signals to the contemporaneous environment. Similarly, by continuing to investigate the role and function of trace elements in the environment, we can continue to make links between trace element signals recorded in Fe mineral archives and biogeochemical processes that reflect contemporaneous environmental conditions.

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