Invited review

Trace elements at the intersection of marine biological and geochemical evolution


Abstract

Life requires a wide variety of bioessential trace elements to act as structural components and reactive centers in metalloenzymes. These requirements differ between organisms and have evolved over geological time, likely guided in some part by environmental conditions. Until recently, most of what was understood regarding trace element concentrations in the Precambrian oceans was inferred by extrapolation, geochemical modeling, and/or genomic studies. However, in the past decade, the increasing availability of trace element and isotopic data for sedimentary rocks of all ages has yielded new, and potentially more direct, insights into secular changes in seawater composition—and ultimately the evolution of the marine biosphere. Compiled records of many bioessential trace elements (including Ni, Mo, P, Zn, Co, Cr, Se, and I) provide new insight into how trace element abundance in Earth’s ancient oceans may have been linked to biological evolution. Several of these trace elements display redox-sensitive behavior, while others are redox-sensitive but not bioessential (e.g., Cr, U). Their temporal trends in sedimentary archives provide useful constraints on changes in atmosphere-ocean redox conditions that are linked to biological evolution, for example, the activity of oxygen-producing, photosynthetic cyanobacteria. In this review, we summarize available Precambrian trace element proxy data, and discuss how temporal trends in the seawater concentrations of specific trace elements may be linked to the evolution of both simple and complex life. We also examine several biologically relevant and/or redox-sensitive trace elements that have yet to be fully examined in the sedimentary rock record (e.g., Cu, Cd, W) and suggest several directions for future studies.

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1. Introduction

The trace elements utilized in metalloenzymes today are commonly thought to reflect, to some degree, the availability of trace elements in ancient seawater when those metalloenzymes first evolved. It was first realized in 1988 that trace metal availability exerted significant control over phytoplankton productivity (Martin and Fitzwater, 1988). In the mid 1990s, as part of the first edition of their seminal book, Frausto da Silva and Williams (2001) suggested that a cell’s trace element inventory was directly related to the conditions under which the host organism evolved. This profound suggestion, along with increasing evidence for absolute Co requirements in cyanobacteria, led, to some degree, the availability of trace elements in ancient seawater when those metalloenzymes first evolved. It was first realized in 1988 that trace metal availability exerted significant control over phytoplankton productivity (Martin and Fitzwater, 1988). In the mid 1990s, as part of the first edition of their seminal book, Frausto da Silva and Williams (2001) suggested that a cell’s trace element inventory was directly related to the conditions under which the host organism evolved. This profound suggestion, along with increasing recognition that trace element availability exerted control over productivity and other metabolic activities, stimulated a new generation of studies examining trace element use and limitation in marine microorganisms (e.g., Sunda and Huntsman, 1995; Saito et al., 2002). Indeed, Sunda and Huntsman (1995) arguably provide the first experimental evidence for absolute Co requirements in cyanobacteria.

The idea that an organism’s trace element requirements are dependent on the environment in which it evolved stems from the simple observation that the biogeochemical cycling of many bioessential elements can vary dramatically under different aqueous conditions (see also Williams and Rickaby, 2012). In this light, some trace element limitations observed in the modern ocean might be thought of as an evolutionary legacy of earlier life adapted to more replete conditions. This idea is easily illustrated with iron (Fe), which is the most common metal co-factor and a limiting micronutrient in large regions of the oceans today, especially high-nutrient low chlorophyll (HNLC) regions (for reviews see Zahariev et al., 2008; Moore et al., 2013). By contrast, Fe would have been much more abundant in surface waters on a more reducing Earth when basic microbial metabolic machinery was being established (e.g., Poulton and Canfield, 2011; David and Alm, 2011). Thus, modern iron demand can be thought of as an evolutionary relict, stemming from the emergence of lineages under ancient environmental conditions of relative Fe abundance. The leading alternative to this model is that organismal elemental requirements are driven almost entirely by utility, i.e., cellular function, with shifts in biological requirements decoupled from corresponding environmental abundances (Scott et al., 2013; Robbins et al., 2013; Stüeken et al., 2015). Some continuum likely exists between these two scenarios.

If biological trace element requirements are related to changes in their paleo-seawater concentrations, then comparative microbial phylogenomics should provide some insight into paleomarine chemistry. Zerkle et al. (2005) surveyed multiple microbial genomes to track the distribution of metalloenzymes in prokaryotes over geological time and evaluated biogeochemical signatures from which inferences about paleomarine trace element concentrations could be made. Further, Zerkle et al. (2005) proposed an evolutionary trajectory for the use of several metals in metalloenzymes, highlighting instances that matched inferred seawater chemistry and several that did not. Using an alternative approach, Dupont et al. (2006) examined the diversification of structural domains in metal-binding proteins across modern proteomes, and similarly suggested that their trace element evolutionary pathways track to some degree changes in paleomarine geochemistry. Dupont et al. (2010) furthered this idea by linking increased atmosphere-ocean oxygenation during the late Neoproterozoic to increased reliance on certain bioessential trace metals, such as Zn, Cu, and Mo. This transition, in turn, may have been a contributing factor to the evolution and diversification of eukaryotes at that time. In at least one case, careful examination of the rock record reveals that biological innovation, rather than evolving marine trace element concentrations, may have guided biological dependency (Zn, c.f. Section 4.4; Scott et al., 2013; Robbins et al., 2013). As both these studies indicate a relatively constant marine reservoir of Zn, the rapid proliferation observed in Zn metalloenzymes in the Neoproterozoic (e.g., Dupont...
et al., 2010) may instead be attributable to the utility of Zn in eukaryotic metalloenzymes rather than evolving environmental conditions (see also Section 4.4).

Regardless of which factors have shaped the elemental stoichiometry of microorganisms, constraining trace element abundances in seawater through time is of paramount importance for understanding the evolution of marine biogeochemistry on a mechanistic level. The potential for evolving marine geochemistry to affect the bioavailability of biologically critical trace elements was popularized, at least from an Earth Sciences perspective, by Anbar and Knoll (2002). These authors proposed that Proterozoic ocean chemistry and biological evolution may be viewed as linked via a “bioinorganic bridge”, whereby high concentrations of dissolved sulfide in the marine environment during much of the Proterozoic limited the bioavailability of critical trace elements in seawater, such as Mo. This not only had a negative impact on global primary productivity, but it delayed the evolution of early eukaryotes. Not surprisingly, there has been an increased focus in recent years on exploring Precambrian sedimentary rocks for direct proxies of paleomarine chemistry and redox conditions. Much of this work has been directed towards the iron formation (IF), black shale, and pyrite records of the marine paleoenvironment; although carbonates also provide important information, their complex diageneric histories and susceptibility to overprinting complicates their use.

As iron- and silica-rich chemical sedimentary deposits, IF have proven useful because the trace metal content in the least metamorphosed units is almost entirely derived from seawater (see Bjerrum and Canfield, 2002; Konhauser et al., 2007, 2009, 2015; Robbins et al., 2013). This is further supported by intense filtering of the IF record to remove samples showing indications of detrital inputs (e.g., >1% Al₂O₃ or >0.1% TiO₂; see Konhauser et al., 2009, 2011; Robbins et al., 2013; Partin et al., 2013a; Swanner et al., 2014 for further discussion). Therefore, it is thought that their chemical composition directly reflects availability in the water column at the time of mineral precipitation and deposition.

Another powerful source of information is the shale record, especially organic matter-rich, fine-grained, siliciclastics (with >0.5 wt% total organic carbon, TOC), which are attractive because (1) they have modern analogues in anoxic basins (e.g., Black Sea, Cariaco Basin), (2) they provide a more continuous temporal record since shale is relatively common in the geologic record, (3) several trace elements (e.g., Mo, U) are known to scale with organic carbon during deposition and burial in euxinic water columns (Algeo and Lyons, 2006), and (4) a direct relationship between concentrations in organic-rich sediments and dissolved concentrations in overlying anoxic and sulfidic bottom waters has been demonstrated for some trace elements (e.g., Mo, Zn; Algeo and Lyons, 2006; Scott et al., 2013).

Recently, several other sedimentary rock types are receiving increased attention for their trace element proxy potential, including chert (Baldwin et al., 2011) and diagenetic pyrite (e.g., Swanner et al., 2013, 2014; Large et al., 2014; Gallagher et al., 2015). Each of these archives has the potential to provide unique insights into past marine trace element concentrations, but as outlined below, they also have obvious limitations.

The IF and shale trace element records published to date generally support the idea that redox chemistry played a central role in the evolution of marine elemental cycling, with two particular events standing out: the Great Oxidation Event (GOE) ~2.4 billion years ago (Ga), and the Neoproterozoic oxygenation event (NOE) ~0.7 Ga. The GOE represents the permanent rise of oxygen to above 10^-5 of present atmospheric levels [PAL], an upper limit for the production of large isotopic signatures by sulfur mass-independent fractionation (S-MIF) (Pavlov and Kasting, 2002) that effectively disappeared from the sedimentary record between 2.45 and 2.32 Ga (e.g., Bekker et al., 2004; Farquhar et al., 2000, 2011). Recent studies based on the trace element proxies of Cr and U, as indicators for continental oxidative weathering, have pushed the onset of the GOE back to 2.48–2.47 Ga (Konhauser et al., 2011 and Partin et al., 2013a), respectively. The period surrounding the GOE is likely better thought of as a long-lived dynamic transition rather than a discrete event (Lyons et al., 2014a). However, for the purpose of this review, we will refer to the age of the GOE as ~2.4 Ga.

In this review, we first provide a brief description of the geochemical modeling (Fig. 1) and genomic work done thus far, as well as the IF and black shale records, highlighting the features that make trace elements useful as paleomarine proxies. Subsequently, we discuss several bioessential trace elements in the order in which they were first explored in the literature. Finally, we identify several trace elements that have yet to be investigated in detail and, in this light, outline several opportunities for future work. Although the primary purpose of this paper is to review our knowledge of the records of trace element evolution in seawater, we will also highlight trace metal evidence for Earth’s protracted redox evolution. Given the control exerted on many metals by the prevailing redox condition and the requirement of oxygen for complex life to evolve, these trace metal redox signals speak directly to the activity of the biosphere, and more specifically that of photosynthetic cyanobacteria.

1.1. Evidence for early life

It is now generally accepted that life evolved relatively early in Earth history, with putative evidence pointing to the existence of a biosphere as early as 4.1 Ga (see Bell et al., 2015) and more convincingly by 3.7 Ga.
The evidence for early life comes in a variety of forms that include $^{13}$C-depleted organic carbon residues (e.g., Mojsis et al., 1996; Rosing, 1999; Rosing and Frei, 2004; McKeegan et al., 2007), microfossils (e.g., Schopf and Packer, 1987; Schopf, 1993, 2006), purported ichnofossils (Furnes et al., 2004; Banerjee et al., 2006, 2007), purported ichnofossils (Furnes et al., 2004; Banerjee et al., 2006, 2007), stromatolites or other microbially induced sedimentary structures (MISS) (Walter et al., 1980; Hofmann et al., 1999; Grotzinger and Knoll, 1999; Allwood et al., 2006; Van Kranendonk et al., 2003, 2008; Heubeck, 2009; Noffke, 2009; Noffke et al., 2013; Nutman et al., 2016), and molecular biomarkers (Brocks et al., 1999, 2003a,b,c; Eigenbrode and Freeman, 2006; Waldbauer et al., 2009). Each of these indicators for early life has been challenged, generally by invoking abiogenic mechanisms or contamination by younger material to explain their origins (e.g., carbon isotopes: van Zuilen et al., 2002; Lepland et al., 2011; microfossils: Brasier et al., 2002; Garcia-Ruiz et al., 2003; ichnofossils: Grosch and McLoughlin, 2014; stromatolites: Lowe, 1994; biomarkers: Rasmussen et al., 2008; French et al., 2015). Despite these challenges, however, there is a growing consensus that favors an origin of life on Earth before ~3.5 Ga, and perhaps much earlier.

For reasons still debated, complex animal life did not evolve until almost three billion years later in the Neoproterozoic (e.g., Pecoits et al., 2012). Following the evolution of stem group eukaryotes in the early mid-Proterozoic (Butterfield, 2000; Knoll, 2014), there is a relatively static period in eukaryotic diversity lasting from 1.8 to 0.8 Ga. This relative evolutionary stasis was punctuated by two critical events that occurred in relatively short order: first, the evolution of metazoans in the Cryogenian (e.g., Love et al., 2009; Erwin et al., 2011), and second, the rapid diversification of complex animal life in the Ediacaran and into the Cambrian – although this event also likely has roots in the Cryogenian (Fedonkin, 2003; Love et al., 2009; Erwin et al., 2011). This billion-year stagnation in eukaryotic diversification is often attributed to the late rise of atmospheric oxygen (e.g., Nursall, 1959; Knoll and Carroll, 1999; Fedonkin, 2003; Planavsky et al., 2014a) to levels required by metazoans (i.e. 0.5% to 4% PAL; Mills et al., 2014). Recent work by Mills et al. (2014), however, suggests that primitive metazoans, such as sponges, may have needed very little oxygen in the water column in order to thrive. Additionally, changes in the availability of critical trace elements (Figs. 1 and 2) – themselves linked to the evolving redox state of the Earth – have been suggested to have influenced eukaryotic diversification (e.g., Anbar and Knoll, 2002; Williams and Frausto da Silva, 2003; Saito et al., 2003; Dupont et al., 2010; Williams and Rickaby, 2012). The slow rate of eukaryotic evolution, limited variation in the carbon isotope record (e.g., Brasier and Lindsay, 1998), and a paucity of evidence for glaciation on Earth during this time period (Eyles and Young, 1994) have ultimately led to the mid-Proterozoic (1.8 to 0.8 Ga) being termed the ‘boring billion’ (e.g., Brasier and Lindsay, 1998).

As life likely emerged at the end of the Hadean or early Eoarchean (4.1–3.7 Ga), it would have been subjected to geochemical conditions in the oceans that were fundamentally different to those present today. Earth’s history has been marked by the advent of plate tectonics, growth of the continental crust, and protracted oxygenation of the atmosphere-ocean system—amongst other fundamental transitions. All these events had major impacts on seawater chemistry. To

![Fig. 2. Updated version of a classic figure from Williams and Frausto da Silva (2003; their Fig. 4), highlighting the concentration of selected elements in the modern ocean (Elderfield and Schultz, 1996; Williams and Frausto da Silva, 2003), the primitive ocean (as per Williams and Frausto da Silva, 2003), hydrothermal fluids (Trefry and Metz, 1989; Elderfield and Schultz, 1996; Douville et al., 2002; Kishida et al., 2004), and values based on the proxy record for ~2.7 Ga (Jones et al., 2015; Scott et al., 2008, 2013; Konhauser et al., 2009; Robbins et al., 2013). Clear differences exist between the predicted primitive ocean values of Williams and Frausto da Silva (2003), who based their estimates off of sulfide mineral solubility, and those indicated by the proxy record. Some proxy records exhibit values more typical of hydrothermal fluids, possibly indicating a greater relative contribution from hydrothermal sources to the early oceans (e.g., Robbins et al., 2013).](image)
understand the interplay between ocean chemistry and the emergence and diversification of life, it is necessary to examine both predictive models for ancient ocean chemistry and the chemical record of ancient sedimentary rocks. Accordingly, such efforts can be divided into the two broad areas emphasized in the review: (1) modeling, cultures, and modern observations, and (2) the sedimentary proxy records.

2. Modeling and experimental views on trace element limitations

2.1. Geochemical and biological modeling approaches

The notion that changes in seawater composition (Figs. 1 and 2) drove evolution was initially championed in the works of Frausto da Silva and Williams (e.g., Frausto da Silva and Williams, 2001; Williams and Frausto da Silva, 2002, 2003, 2004). In their 2001 book *The Biological Chemistry of the Elements*, Frausto da Silva and Williams suggested that trace element concentrations could have been significantly different on the early Earth (Fig. 2) and that trace element bioavailability may have been controlled by changes in the paleomarine concentration of dissolved sulfide. They pointed to the biological utilization of Ni and Co as examples, suggesting that prokaryotes evolved in a reducing environment where these trace elements could have acted as catalysts for early metabolisms, consistent with their use in methanogenesis and by hyperthermophiles. The idea that evolution could be chemically constrained was furthered by Williams and Frausto da Silva (2003) who suggested that changes in mineral solubility may have driven increasing availability of trace elements, such as V, Cu, and Zn (amongst others), as the Earth became more oxidizing (Fig. 2). Williams and Frausto da Silva (2004) classified the genome, proteome, and environmental chemistry as “the trinity of life,” inextricably linking these three factors to biological evolution. A more recent view of this work has been provided by Williams and Rickaby (2012), where they further suggested that organisms may be grouped into chemotypes depending on similarities in their trace metal requirements. Saiito et al. (2003) modeled the solubility of a suite of biologically essential trace elements (Fe, Mn, Zn, Ni, Cu, Cd, and Co) to evaluate plausible concentrations in ancient seawater under ferruginous (anoxic and euxinic) conditions relative to Cd, Cu, and Zn. Following increasing oxygenation in late Neoproterozoic (Canfield et al., 2007; Scott et al., 2008; Sahoo et al., 2012, 2016; Lyons et al., 2014a; Planavsky et al., 2014a), these sulfide complexes would have become less abundant, resulting in greater availability of Cd, Cu, and Zn. Under more oxic conditions first observed in the Cryogenian and continuing to develop into the Cambrian–Ordovician (Large et al., 2014; Sperling et al., 2015; Sahoo et al., 2016), the models of Saiito et al. (2003) indicated that Fe and Mn concentrations in seawater would have been drawn down to modern levels as the result of oxide mineral precipitation. This would have presented a challenge for many microbial clades, as Fe is the most common metal co-factor for both prokaryotes and eukaryotes (Dupont et al., 2010). The models of Saiito et al. (2003) support the hypothesis that modern cyanobacterial trace element nutrient requirements may be viewed as the direct result of their early evolution in ancient oceans limited in certain trace elements (Cu, Cd, Zn) and enriched in others (Fe, Mn, Co, Ni). Inverse modeling based on the utilization of trace elements in metalloenzymes (Zerkle et al., 2005) further supports a strong linkage between evolving ocean chemistry and biological trace element dependency. This work suggests that the utilization of trace metals in biology generally follows the pattern of Fe ≫ Zn > Mn ≫ Mo, Co, Cu ≫ Ni > W, V.

Links between the chemical evolution of the early oceans and the trace element complement of organisms are also informed by the emergence or disappearance of metal-binding protein fold families (FF) or fold super families (FSF). Fold families are groups of proteins that are related by structure, function, and sequence, and are considered to have a common evolutionary origin (Dupont et al., 2006). For families whose proteins contain metal-binding domains, it is possible to predict which metals occupy these domains, such that evolutionary

![The periodic table of elements showing biologically essential elements as identified by Frausto da Silva and Williams (2001), and highlighting those that have been investigated in the sedimentary record to date. Blue indicates major bioessential elements, yellow – bioessential trace elements that have not been investigated in the proxy record, purple – major elements with biological importance whose general seawater geochemical behavior is fairly well known, and red – trace elements investigated in at least one of the sedimentary proxy records, such as IF, black shales, sedimentary pyrite, and/or carbonates.](image-url)
relationships between protein structures may be used to reveal trends in metal acquisition in biological systems. The loss or gain of these FF or FSF can be viewed as key evolutionary events that in many cases appear to track fundamental shifts in paleomarine redox chemistry (Dupont et al., 2006, 2010).

Much of the current work focused on trace element proxy records (Fig. 3) has been inspired by the idea that trace metal biolimitation may have been significantly different in the deep past (Javaux et al., 2001; Anbar and Knoll, 2002). In Anbar and Knoll’s (2002) ‘bio-inorganic bridge’, changes in early ocean redox chemistry directly affected the bioavailability of trace elements and, in turn, the evolutionary trajectory of life. Inspired by arguments for widespread marine euxinia during the Mesoproterozoic (1.8–1.0 Ga) (Canfield, 1998), they suggested that such conditions would have limited the seawater Mo reservoir. As nitrogenases containing a Mo-Fe metal cofactor are more efficient than alternative Fe-Fe and V-Fe versions (Eady, 1996), low Mo concentrations would have negatively impacted the ability of primary producers to fix N₂. Consequently, Mo limitation could have stifled primary productivity and perhaps even eukaryote evolution (Anbar and Knoll, 2002).

Indeed, the genomic record for several trace metals has been linked to evolving marine geochemistry. Molybdenum utilization in organisms may have developed in tandem with increasing Mo availability following the early stages of biospheric oxygenation, at which point it became critical to nitrogen fixation (Williams and Frausto da Silva, 2002, 2004; Zerkle et al., 2005; Boyd et al., 2011); although Zerkle et al. (2006) and Glass et al. (2009) have both shown that very low Mo concentrations are capable of supporting near modern levels of nitrogen fixation. Similarly, based on eukaryotic cellular requirements, geochemical modeling, and the late emergence of eukaryotic Zn metalloenzymes, it was believed that Zn concentrations in the early oceans would have been a possible barrier to eukaryotic evolution (Williams and Frausto da Silva, 2002, 2003; Saito et al., 2003; Dupont et al., 2006, 2010; Williams and Rickaby, 2012). However, recent studies have suggested that the link between trace element availability, utilization, and metallome requirements may have been more complex. For instance, Stüeken et al. (2015a) presented nitrogen isotope data from ~3.0 Ga sedimentary units suggesting the activity of Mo–Fe nitrogenase, which might indicate that Mo was present at low but physiologically-sufficient levels in the early ocean. This is a scenario supported by the culture studies of Zerkle et al. (2006) and Glass et al. (2009). In the case of Zn, the records of enrichments in both black shales and IFS suggest that the size of the oceanic Zn reservoir has been relatively constant since the Archean (Scott et al., 2013; Robbins et al., 2013). Thus, in some cases in the deep past, trace elements may have been employed in biological systems despite strongly limiting seawater concentrations (the case of Mo), while in others, sedimentary records are at odds with marine trace metal histories suggested by geochemical models and protein structural phylogeny (the case of Zn).

Saito et al. (2003) emphasized that their projections of Precambrian seawater metal concentrations (Fig. 1) are based on thermodynamic models of mineral solubility and speciation in simulated seawater and are thus inherently limited. For many elements, seawater abundances are subject to kinetic controls (e.g., Broecker, 1971) where the dissolved reservoir scales with input/output fluxes without approaching solubility limits. A further limitation is that many of these models assume a homogeneous ocean, which is at odds with observations of the modern oceans. In today’s oceans, trace element abundances vary both laterally and with depth, and these may vary over several orders of magnitude (Bruland and Lohan, 2003). As well, Moore et al. (2013) have highlighted spatial heterogeneity on a global scale in surface waters with regards to both major limiting nutrients such as nitrate or phosphate, as well as trace metals. Further, studies of ancient environments have already shown that there can be basin scale differences in water column chemistry as the result of stratification with depth or proximity to shoreline (e.g., Poulton et al., 2010). Given our knowledge regarding spatial variation both in modern oceans and as recorded by the sedimentary record, the assumption of a homogenous ocean is certainly incorrect. However, many aspects of modeling approaches have yet to be fully explored for trace elements in the modern ocean, much less under ancient ocean conditions with dramatically different chemical and redox regimes.

2.2. Culture experiments and modern environmental observations

A number of studies have utilized either pure cultures, industrial samples, or natural environments to test and examine the limiting effects of trace metals on biology: for a comprehensive example we draw the reader’s attention to Glass and Orphan (2012) who discuss the trace metal limitations of methanogens and methanotrophs. In modern marine environments, Morel and Price (2003) have demonstrated how plankton are able to utilize very low levels of trace metals via complexing ligands that likely evolved to be highly efficient at extracting low levels of these micronutrients from seawater. As well, Morel (2008) related the elemental stoichiometry of modern phytoplankton to the cycling of trace elements in the oceans, and further suggested that the trace metal cycles may have been affected by the advent of strong biological recycling in order to fulfill microbial needs.

Several critical ideas may be gleaned from these reviews and the works that they are based on. First, although trace metals may be biolimiting, levels vary between different organisms, and a universally biolimiting concentration for trace metals may not exist. Second, most prokaryotes and single celled eukaryotes seem to favor trace metal concentrations on the order of nM to mM levels. Indeed, Glass and Orphan (2012) discuss how the production of methane can be stimulated by the addition of 0.2–2 μM of trace metals such as Fe, Cu, or Mo to some methanogen communities. In laboratory cultures with freshwater cyanobacteria, Zerkle et al. (2006) suggest that nitrogen assimilation through molybdenum-nitrogenase can occur at Mo levels as low as 5 nM, suggesting that these enzymes can be active over a broad range of concentrations. Third, trace metal concentrations do have an upper limit, after which they become toxic as opposed to being beneficial for the organism. Finally, given the low levels of trace metals in modern oceans, organisms have had to develop strategies for dealing with the possibility of micronutrient limitation (e.g., Morel and Price, 2003; Morel, 2008). This may include the development of siderophores to assist in scavenging any Fe present, or even the exclusion of certain trace metals traditionally used in a metalloenzymes in favor of a metal free variety. Such strategies lead to the question of whether they are a more recent development or a hold-over from evolution in an ocean with similarly low levels of trace metals, effectively comparable to modern? Ongoing research in these areas will be fundamental to understanding microbial evolution and our interpretation of the sedimentary record.

3. Proxy records

3.1. Traditional proxy records for redox conditions

Two more traditional and key proxy records that have informed our understanding of the Earth’s redox evolution include S-MIF and Fe speciation. Increasingly, these are being augmented with metal stable isotopes, such as Mo and Cr (see Sections 4.2 and 4.6). Here we will provide a brief overview of these records, and the redox constraints they have placed on the evolution of the Earth. This will provide a basic framework for the reader to interpret the newer, and often more controversial, trace metal isotope and proxy records for evolving oxygen levels. We have included these redox considerations for two reasons. First, trace metal isotopes are becoming increasingly used in the field of geochemistry and new datasets are being rapidly generated. Second, and perhaps more important for the purposes of this review, even if the metals themselves do not have a direct
3.1.1. Sulfur mass-independent fractionations

The disappearance of S-MIF is perhaps the most accepted evidence for the onset of widespread and permanent oxygenation of the Earth’s surface to levels above $10^{-5}$ times present atmospheric levels (PAL) (see Pavlov and Kasting, 2002; Bekker et al., 2004; Farquhar et al., 2000, 2011). Below this threshold, photolytic reactions between ultraviolet rays and sulfur gases in the atmosphere produce isotopic anomalies that deviate from mass dependent behavior and are subsequently recorded in marine sedimentary rocks. However, once oxygenic photosynthesis leads to an accumulation of appreciable oxygen, above $10^{-5}$ PAL, this allows an ozone layer to form, shielding the Earth from harmful UV rays and suppressing S-MIF. Although modeling has suggested that the S-MIF signal may be carried forward for 10–100 Ma through sedimentary recycling (Reinhardt et al., 2013a), it remains a definitive marker for the first major rise in atmospheric oxygen and a fundamental change in the redox state of the Earth.

3.1.2. Iron speciation

Iron speciation, a technique developed by Berner, Canfield, and colleagues (e.g., Berner, 1970; Canfield, 1989) and refined by Poulton and Canfield (2005) for application to ancient sediments, has offered many new insights into the evolving redox state of the early Earth. Iron speciation is predicated on determining the amount of Fe in a given sample that has been partitioned into various phases—carbonates, ferric oxyhydroxides, magnetite, and sulfide—relative to the total amount in a sample; see Poulton and Canfield (2011) for a brief review. The ratio of Fe in highly reactive phases (the sum of the four aforementioned phases) to total Fe effectively diagnoses sediment deposition from anoxic water column settings. Combining this further with the ratio of pyrite extractable Fe to highly reactive Fe allows samples deemed anoxic to be further categorized as ferruginous or euxinic. Fe speciation analyses have shed new light on the spatial complexity of water column redoxlines (e.g., Poulton et al., 2010), and have shown that the Proterozoic was likely dominated by ferruginous conditions (Planavsky et al., 2011; Poulton and Canfield, 2011). Poulton and Canfield (2011) highlight several fundamental shifts based on a number of Fe-speciation analyses. They suggest that the oceans were dominantly ferruginous until the late Archean after which a surface oxic layer likely formed. During the Paleoproterozoic to Neoproterozoic, this surface oxic layer probably persisted, but with a euxinic wedge on the shelf with underlying ferruginous waters (Li et al., 2010). A recent assessment of a compilation of Fe speciation data suggests that bottom waters may have remained ferruginous well into the Paleozoic (Sperling et al., 2015).

3.2. Proxy records for trace element evolution

3.2.1. Iron formations

Iron formations (IF) are iron-rich (15–40 wt%) and siliceous (40–60 wt%) sedimentary deposits that precipitated from seawater throughout much of the Archean and Paleoproterozoic (3.75–1.85 Ga) (James, 1954; Trendall, 2002; Klein, 2005). Deposition of IF appears tied to periods of enhanced magmatic and hydrothermal activity (associated with large igneous province emplacement; Isley and Abbott, 1999) that delivered large amounts of ferrous iron to anoxic deep oceans (Bekker et al., 2010, 2014). Low concentrations of $Al_2O_3$ (<1 wt%) and incompatible elements (Ti, Zr, Th, Hf and Sc < 20 ppm) are commonly observed in IF, indicating minimal detrital input during deposition, although this is not universal for all iron formations.

Iron formations may be divided into two petrographic affinities: banded iron formation (BIF) and granular iron formation (GIF). BIF are characterized by distinctive layering of variable thickness, from macrobrecks (meters in thickness), to the more characteristic mesobands (centimeter-thick units) from which they draw their name, to millimeter and submillimeter microbands (e.g., Trendall and Blockley, 1970; Morris, 1993; Krapež et al., 2003). GIF typically lack banding and consist of granules of chert or other silicates and iron oxides with early diagenetic chert cement filling pore spaces (e.g., Simonson, 1985). GIF dominate the Archean and are more important in terms of total IF tonnage (Bekker et al., 2010). GIF first appear in the rock record at ca. 2.4 Ga (Simonson and Goode, 1989) and are the most common type of iron formation in the Paleooproterozoic, reaching their peak abundance ca. 1.88 Ga. After a 1.88 Ga pulse of IF deposition, which appears to have been globally synchronous (Rasmussen et al., 2012), they effectively disappear in the middle Proterozoic, returning in the Neooproterozoic in association with widespread “Snowball Earth” glaciation (Hoffman et al., 1998). While the Phanerozoic is devoid of the IF resembling those of the Precambrian, the iron oxide-rich sedimentary record is continued into the Phanerozoic in the form of ironstones (see Mücke and Farshad, 2005 for review) and exhalative deposits (see Lyons et al., 2006 for review).

Iron formation deposition spans several major redox changes in Earth’s surface composition—from an early anoxic atmosphere to an atmosphere that became at least partially oxygenated (e.g., Klein, 2005; Bekker et al., 2010). Therefore, it is likely that IF formed via different mechanisms throughout the Precambrian. A number of recent reviews detail IF occurrence, mineralogy, mechanisms of formation, depositional environments, and diagenetic history (see Klein, 2005; Bekker et al., 2010, 2014; Posth et al., 2014). For the purpose of this review, the importance of IF is its ability to record marine signatures, and specifically archive trace element concentrations in the Precambrian.

Evidence supporting the idea that IFs record autogenic marine signatures includes marine-like rare earth element and yttrium (REE + Y) patterns and small-scale chemical variations that argue for the preservation of environmental signals (e.g., Bau and Möller, 1993; Bau and Dülk, 1996; Bolhar et al., 2004; Alexander et al., 2008; Pecoits et al., 2009; Planavsky et al., 2010a; Haukård et al., 2013, 2016). A concern potentially compromising the IF record is the possibility of post-depositional mobilization of trace elements, which can overprint or even eradicate autogenic marine signatures. However, limited post-depositional mobilization or addition of trace elements in IF is indicated by small-scale REE and Fe isotope variations, both within and between Fe-rich mesobands despite experiencing diagenetic and metamorphic conditions up to amphibolite facies (e.g., Bau, 1993; Frost et al., 2007; Whitehouse and Fedo, 2007; Steininhoefel et al., 2010). Trace element mobilization efforts for IF have often limited their scope to samples falling at greenschist facies or below in an effort to provide the most robust estimates possible of trace element abundances.

Recently, the use of IF as paleoenvironmental proxies for trace element abundances has been questioned because laboratory studies of Fe(II)-redox driven recrystallization suggest that this process may overprint autogenic trace element records (Friedrich et al., 2011; Friedrich and Catalano, 2012). These studies focused largely on the initial sorption of trace metals and subsequent mobilization that occurs upon further interaction with Fe(II)-rich fluids. They also specifically highlighted the potential for the mobilization of Ni (Friedrich et al., 2011) and Zn (Friedrich and Catalano, 2012). Generally, losses were < 10% when ferrihydrite doped with Ni or Zn was placed in a Fe(II) solution that contained no Ni or Zn. However, further experimental work (Friedrich et al., 2012) has shown that when impurities of Al, Cr, or Sn are present in the ferrihydrite, Ni and Zn mobilization is attenuated. Such a scenario is likely more comparable to natural iron oxyhydroxides formed in the Precambrian oceans that were the precursors to the minerals presently found in IFs, as they would have incorporated various trace metals and other minor impurities from the water column. Ultimately,
the studies of Friedich et al. (2011) and Friedich and Catalano (2012) are based on systems inherently at disequilibrium and that are unlikely to be truly representative of the formation of IF particles in equilibrium with contemporaneous ferruginous seawater. Further, none of these studies have assessed the potential for the mobility of trace elements during later diagenetic mineral phase transitions. In this regard, recent experimental work using diagenetic capsule experiments (Robbins et al., 2015) demonstrated limited mobility for Ni and Zn during simulated diagenetic treatments at high temperature and pressure that capture the transformation from ferrhydrate to hematite. Overall, when all lines of evidence are considered, it is reasonable to conclude that IF do indeed preserve authigenic signatures and thus record the abundances of biologically critical trace elements in ancient oceans with high fidelity.

Recent attempts to connect the record of trace metals in IF to coeval trace element abundances in seawater may be hampered by the empirical sorption models (e.g., linear partitioning, or KD, models) typically employed to determine metal partitioning between seawater, microbes, mineral colloids, and organic ligands in the water column. These partitioning models are only applicable at the experimental conditions tested and say nothing about the chemical mechanisms of trace metal uptake (Sposito, 1982; Goldberg and Criscenti, 2007). Erel and Stolper (1993), improving on earlier KD studies such as those of Byrne and Kim (1990) and Koeppenkastrop and De Carlo (1992), proposed a semi-empirical model that linked the binding of marine REE to microbes and particulate matter to their first hydroxide thermodynamic binding constants (e.g., constants for hydrox ferric oxides found in Dzombak and Morel, 1990). Aqueous REE concentrations were successfully modeled by employing REE-carbonate complexation mass action constants. While admitting that their model could not account for factors including pH and binding site concentrations, Erel and Stolper (1993) found that the model was able to predict REE concentrations in modern seawater and approximate REE patterns observed in Archean BIF.

More recently the surface complexation modeling (SCM) approach, grounded in equilibrium thermodynamics through mass action expressions, has been extended beyond minerals to successfully model trace metal adsorption onto microbial surfaces (Fein et al., 1997, 2005), including marine cyanobacteria (Li et al., 2015) and anoxicogenic photosynthesizers (Martinez et al., 2016). Although data-intensive, the SCM approach allows us to predict the impacts of the aqueous speciation of metals, redox transitions, pH, the precipitation and dissolution of solid phases, competition of multiple metals for specific types of surface binding sites, varying metal-to-sorbent ratios, and ionic strength on the final distribution of trace metals in a system (Davis et al., 1998; Koretsky, 2000)—without conducting additional experiments. For example, SCM studies allow for systematic investigation of the impacts of paleoseawater salinity (Sanford et al., 2013), pH (Pearson and Palmer, 1999; Ohnemueller et al., 2014) and competition on the uptake of trace metals to particles in the photic zone, for instance, Fe-Mn oxyhydroxides and planktonic microbes. Binding constants from disparate studies in the literature could be combined, unlike in empirical approaches such as the KD model. Ultimately, the further application of SCM promises to better connect paleoseawater geochemistry and microbiology to trace metals trends observed in the rock record.

A final limitation on the use of the IF record is the lack of direct modern analogues. However, several studies have used Phanerzoic ironstones and hydrothermal exhaltes to extend the record from the Precambrian to the modern (e.g., Konhauser et al., 2009, 2011, 2015; Partin et al., 2013a; Robbins et al., 2013; Swanner et al., 2014). Such hydrothermal deposits provide an opportunity to test experimental and hypothesized partitioning scenarios for trace elements onto IF—but only to a limited extent. As these partitioning scenarios are sensitive to matrix effects, such as different Si concentrations (e.g., Konhauser et al., 2007, 2009) and the presence of additional cations (Jones et al., 2015), see also Section 4.1), they cannot be directly equated. Although the KD value for P adsorbing to Fe in modern hydrothermal particles (Bjerrum and Canfield, 2002) is very close to the experimentally-derived value for Si-free seawater developed by Konhauser et al. (2007), further refinement of these partitioning scenarios is needed, and SCM may be a useful tool for future work. Additionally, the broad scaling between trace metals and Fe in IF suggest that first order partitioning trends are largely preserved (e.g., Robbins et al., 2013; Konhauser et al., 2015). The observed scaling between Zn and Fe in modern hydrothermal deposits falls just above that for IF Zn/Fe ratios (Robbins et al., 2013). Collectively, these considerations suggest that the use of modern hydrothermal exhaltes to extend the IF record is justified.

3.2.2. Shales

The shale record is another powerful source of information, especially organic matter-rich mudrocks, also known as black shales (with >0.5 wt% TOC). Shales are fine-grained sedimentary rocks containing variable amounts of organic matter that are typically deposited in low energy environments and can provide key information regarding local bottom-water redox conditions and the extent of primary paleo-productivity. The latter may be inferred from trace element enrichments that are intimately associated with organic carbon burial fluxes, which are in turn favored under anoxic depositional conditions (e.g., Ni, Cu, Zn, Cd; Tribovillard et al., 2006; Algeo and Rowe, 2012). Local bottom water redox conditions are indicated by enrichments in redox sensitive trace metals. Such trace metals are generally soluble in oxygenated seawater but are removed in anoxic seawater or sediment pore waters through authigenic mineral formation and/or uptake by organic matter (e.g., U, V, Mo, Re, Cr, and Co).

Organic-poor shales deposited from well-oxygenated bottom waters typically have low metal burial rates (with possible exceptions, such as Mn and Co) and thus they have generally muted metal enrichments. By contrast, organic-rich black shales deposited under anoxic and sulfidic pore waters may become enriched in redox-sensitive trace metals, especially Mo (Algeo and Lyons, 2006; Märs et al., 2008; Scott and Lyons, 2012). It is possible for black shales to form beneath oxygenated bottom waters if sedimentation rates are sufficiently high for organic matter to be buried rapidly and escape oxidation or if productivity rates are sufficiently high (e.g., Sageman et al., 2003). Elevated Re and U enrichments without Mo enrichment in black shales are particularly useful indicators for mill bottom water oxygenation and limited oxygen penetration below the sediment-water interface (Crusius et al., 1996; Morford and Emerson, 1999; Morford et al., 2005; Algeo and Tribovillard, 2009). Deposition from anoxic and non-sulfidic bottom waters can also be inferred from trace metal contents, in particular mild Mo enrichments that indicate dissolved sulfide was confined to sediment pore waters (Scott and Lyons, 2012).

The utility of black shales as paleomarine proxies has grown in recent years. In addition to trace metal contents, sedimentary Fe speciation analysis (Poulton and Canfield, 2005) serves as an indicator of local paleoredox conditions on the ocean floor (i.e., oxic; anoxic, ferruginous; or euxinic) (e.g., Poulton et al., 2004, 2010; Canfield et al., 2007, 2008; Lyons et al., 2009; Reinhard et al., 2009; Planavsky et al., 2011; Asael et al., 2013). The trace element concentrations of anoxic and euxinic black shales have been used to track corresponding elemental concentrations in the oceans through time (e.g., Scott et al., 2008, 2013; Partin et al., 2013b; Reinhard et al., 2013b). If black shales can independently be determined to have been deposited under specific redox conditions (by Fe speciation), the degree of metal enrichment (Mo, U, Re, V, Cr) can then be used to track first order shifts in metal concentrations and the global marine redox state (Emerson and Heusted, 1991; Lyons et al., 2009). This idea builds from two key principles (explored below): (1) the dominant marine redox condition is the primary control on the size of the dissolved ocean inventory of redox sensitive elements (Emerson and Heusted, 1991) and (2) the marine metal reservoir exerts a first order control on enrichments in euxinic and anoxic
sediments (Algeo and Lyons, 2006; Lyons et al., 2009; Scott et al., 2008; Reinhard et al., 2013b).

Trace element enrichments in the organic-rich sediments of modern anoxic basins provide the foundation for interpreting ancient seawater metal inventories from organic-rich black shales. Algeo and Lyons (2006) showed that the dissolved Mo concentration in euxinic bottom waters is proportional to the observed Mo/TOC ratios in the underlying organic-rich sediments. A similar relationship was demonstrated for Zn (Scott et al., 2013) and likely exists for most trace metals (e.g., Large et al., 2014). Such observations were derived by comparison of data from modern anoxic basins with various degrees of water mass restriction from the open ocean. Isolated anoxic basins typically have lower dissolved metal concentrations because of slow recharge of trace metals from ocean circulation, and a largely anoxic ocean would similarly be expected to have low concentrations due to widespread trace metal draw down (Emerson and Heusted, 1991). A temporal compilation of metal enrichments from black shales deposited in sedimentary basins with relatively unrestricted connection to the open ocean should therefore reveal the evolution of seawater metal reservoirs through time.

For some trace elements, the black shale record provides valuable independent information about the evolution of seawater trace element reservoirs, such as Zn, complementing the IF record (c.f. Scott et al., 2013 and Robbins et al., 2013). A significant advantage of the black shale record over the IF archives is that examples from modern settings, such as the Cariaco Basin and the Black Sea (e.g., Lyons and Berner, 1992; Lyons et al., 2003; Algeo and Lyons, 2006; Scott et al., 2008, 2013; Reinhard et al., 2013b), can be used as analogues to understand the processes underpinning the formation and the pathways of trace metal uptake for their ancient counterparts. Furthermore, the distribution of black shales in the rock record is more continuous compared to IFs and ironstones, whose deposition was limited to certain periods of the geologic record.

One of the additional advantages of black shales is that they have low permeability, which contributes to the retention of primary depositional signatures despite the potential for metals to be remobilized during diagenetic processes. While the possibility for diagenetic mobilization does exist, several lines of geochemical evidence can be used to indicate a primary depositional signature. For instance, Tribovillard et al. (2006) indicated that unless black shales are exposed to an influx of oxidizing agents after deposition, a number of trace elements commonly associated with sulfides should remain relatively immobile during diagenesis; these include Mo, Zn, Ni, and Co, amongst others. Further, the Re-Os radioactive isotope system can provide precise and accurate depositional ages for black shales and, at the same time, confirm that redox-sensitive trace elements have not been significantly affected by post-depositional processes (e.g., Kendall et al., 2009).

3.2.3. Sedimentary to early diagenetic pyrite

The sedimentary pyrite record can also prove useful for tracking trace element abundances. Pyrite (FeS2) can form in sediments contemporaneous with deposition or during early diagenesis, and it may therefore incorporate trace element signatures characteristic of seawater (see Large et al., 2014). Focusing on in situ analysis of pyrite grains from black shales, Large et al. (2014) documented the temporal variability of several trace elements, including Mo, Co, Ni, As, and Se, through geological time. Those authors suggested that in the same way that hydrothermal pyrite can track the chemistry of ore-forming fluids (Large et al., 2009), syngentic to early diagenetic pyrite can track changes in seawater composition. Large et al. (2014) also provided an assessment of the importance of pyrite versus matrix trace element incorporation, sulfide recrystallization, and the location in the water column or sediment where pyrite forms, which suggest the record is favorable to recording seawater signatures; a result further supported by several additional studies (Gregory et al., 2014; Large et al., 2015; Mukherjee and Large, 2016). Indeed, pyrite frambooids formed in the water column may prove to be most useful, but this possibility remains to be tested further. The use of the pyrite records has recently found support in the work of Gallagher et al. (2015) who reported a suite of trace element data (Mo, Ni, As, Co, Zn) from Precambrian to Ordovician carbonate-hosted pyrite deposited in shallow marine environments. Their data were centered on the Archean-Proterozoic and Proterozoic-Phanerozoic transitions and were largely consistent with previous assertions from the shale-hosted pyrite, IF, and black shale records.

Large et al. (2015) have also used laser ablation analysis of sedimentary pyrite in shales to identify cyclical variations in several key trace elements, including Mo, Se, and Cd, in the late Precambrian through the Phanerozoic. The cyclical pattern in trace element abundances are ascribed to changes in continental uplift and weathering fluxes, and it appears that mass extinction events seem to coincide with periods of anoxia and oceanic nutrient depletion. However, such large changes in input fluxes could be compensated for by relatively minor increases in the extent of anoxia, which would have contributed to the extinction (e.g., Sahoo et al., 2012; Reinhard et al., 2013b). As with the black shale and IF records, the greatest value of the pyrite record likely lies in its ability to produce broad first-order trends.

3.2.4. Carbonates

Carbonate rocks can also capture and preserve records of ancient marine chemistry. For example, carbonate-associated sulfate (CAS) has been used to study the oxygenation of the early Earth by tracking seawater sulfate levels and their isotopic properties (e.g., Kah et al., 2004; Gellaty and Lyons, 2005; Gill et al., 2007, 2011; Guo et al., 2009; Planavsky et al., 2012; Guibald et al., 2015). In terms of redox sensitive trace elements, a recent study has examined δ13C/δ18O ratios (Hardisty et al., 2014) in order to track surface oxidation of the ocean. The authors showed evidence for an increase in iodate, the oxidized version of iodine, following the GOE and attribute this to the development of an aerobic iodine cycle.

In general, the most important hurdle for any carbonate-specific redox proxy is a thorough understanding of the behavior of the proxy during marine, meteoric, and burial diagenesis. These processes can alter carbonate δ34S and δ18O, as well as the trace element compositions (e.g., Schrag et al., 2013; Swart and Kennedy, 2012), limiting their use for paleoenvironmental proxies. For instance, CAS concentrations have been shown to decrease by orders of magnitude during meteoric argonite-to-calcite alteration, but δ34S CAS values are preserved during the same process (Gill et al., 2008). A similar study found that seawater δ34S is also preserved even after extensive authigenic carbonate precipitation in pore waters with active sulfate reduction (Lyons et al., 2004), again indicating that original marine isotope ratios are retained through this process.

3.2.5. Chert as a possible trace element archive

To date IF and black shales have been the dominant rock types, with sedimentary pyrite being increasingly used to infer evolution of paleomarine trace element reservoirs and redox conditions. A further potential archive that has yet to receive much attention is the Precambrian chert record, with Baldwin et al. (2011) recently discussing the potential for Precambrian cherts to record paleoceanic signatures. Abiogenic cherts are predominantly formed in the Precambrian due to high marine silica concentrations in the absence of a biological sink (Siever, 1992; Maliva et al., 2005). Several mechanisms have been proposed for the primary formation of abiogenic Precambrian cherts, including direct precipitation of amorphous silica from seawater (Siever, 1992; Maliva et al., 2005) and more recently sedimentation of sand-sized silica grains (Stefurak et al., 2014, 2015). Thus far, Precambrian cherts have mainly been used to examine paleoceanic temperatures from their oxygen and silicon isotope compositions (e.g., Knauth and Lowe, 2003; Knauth, 2005; Robert and Chausidon, 2006; Marin-Carbone et al., 2014). However, questions remain as to whether these are truly primary marine signals. Marin-Carbone et al. (2014)
outlined a set of criteria, both petrographic and geochemical, that can be used to help identify pristine sedimentary cherts. Trace element abundances themselves may also help resolve primary versus secondary signals preserved in cherts (Baldwin et al., 2011; Marin-Carbonne et al., 2014). If appropriate samples are identified, the trace element record in cherts could be a powerful complement, both spatially and temporally, to existing trace element datasets.

4. Bioessential trace elements, their records, and implications for changes in seawater chemistry and prevailing redox conditions

4.1. Phosphorus

Phosphorus (P) is key for all life and fills a variety of biological roles, including the formation of phospholipids, cellular membranes, and nucleic acids. Due to strong biological scavenging and sorption to Fe oxyhydroxides, P is typically present at fairly low concentrations in seawater (0.001–3.5 μM; Bruland and Lohan (2003)). In modern oceans P shows a strong nutrient-type profile, and can often be found below the average marine concentration of ~2.3 μM in surface waters (Bruland, 1980; Bruland and Lohan, 2003).

Phosphorus is typically considered to be the limiting nutrient for marine productivity on geologic time scales (Tyrell, 1999). Based on P/Fe ratios in IF (Fig. 4), Bjerrum and Canfield (2002) suggested that prior to 1.9 Ga, P sorption to precipitating iron oxyhydroxides would have drawn down the marine P reservoir to levels of around 0.15 to 0.6 μM, thus limiting primary productivity in the surface oceans. Those authors argued that IF acted as a major sink for P, and consequently, their deposition resulted in a P crisis. Their argument has its roots in the observed drawdown of P by Fe precipitates near modern hydrothermal vents and the fact that the IF record is characterized by P/Fe ratios significantly lower than those observed in iron-rich plume particles today (e.g., Feely et al., 1991, 1998).

However, silica concentrations in the Precambrian oceans were likely much higher (up to 2.2 mM, effectively saturated with respect to amorphous silica; c.f. Siever, 1992; Maliva et al., 2005) than in modern oceans (up to ~0.1 mM; Bruland and Lohan, 2003). The presence of aqueous silica can greatly affect the partitioning of P onto Fe oxyhydroxides. Konhauser et al. (2007) demonstrated that a high silica ocean would have attenuated phosphate adsorption to precipitating Fe oxyhydroxides, thereby keeping P in solution while still accounting for low P/Fe ratios in IF (Fig. 4) — and casting doubt on the idea of an Archean P crisis. Subsequently, Planavsky et al. (2010b) combined the partitioning coefficients derived by Konhauser et al. (2007) with P/Fe ratios in the IF record and suggested that P levels in the Precambrian oceans were at least similar to modern seawater, and perhaps even higher, although this remains debated. Furthermore, they identified a large influx of P into the oceans coincident with the end of global Neoproterozoic glaciations. More recently, it has been argued that the Neoproterozoic influx of P to the oceans may instead be directly attributed to the weathering of large igneous provinces (LIPs) (Horton, 2015). This idea is based on compiled P concentrations in LIPs and average mid-ocean ridge basalts, as well as the emplacement of several LIPs that occurred just prior to Neoproterozoic global glaciations and the NOE.

It should be pointed out, however, that there is renewed debate on the partitioning of P in the early oceans and its implications for the early biosphere. For instance, Jones et al. (2015) suggested that divalent cations, such as Ca2+ and Mg2+, may also play a strong role in governing the partitioning of P in ancient silica-rich oceans, renewing the possibility of an Archean, and possibly Proterozoic phosphate crisis. By contrast, it has been suggested that if the ferrihydrite precipitates were not a major flux of sedimenting trace elements, perhaps it was the biomass itself that contributed (via intracellular assimilation) to the P inventory of BIF (Li et al., 2011).

Reconstructing the ancient P biogeochemical cycle from the shale record is generally restricted to the Phanerozoic. For instance, März et al. (2008) investigated the P and redox sensitive metal content of a Cretaceous black shale interval from the ca. 86 Ma old Coniacian-Santonian ocean anoxic event (OAE). These authors found that P concentrations were relatively low during deposition from euxinic waters, unlike redox sensitive trace metals, such as Zn, V, or Cd. By contrast, high P concentrations were found in black shales deposited from anoxic and non-sulfidic waters where P burial was coupled to Fe oxyhydroxide formation. This strong coupling of P and Fe during burial suggests that the black shale record in deep time can be used to distinguish anoxic/non-sulfidic versus euxinic deposition. This relationship offers the potential to shed light not only on paleomarine P abundances but also in resolving the ongoing debate regarding the interpretation of P/Fe ratios in IF. Additionally, Large et al. (2015) measured P in a suite of fifty Phanerozoic black shales and found a cyclical variation in P that seemed to coincide with changes in other trace elements, including Se and Mo. Large et al. (2015) further correlated these nutrient peaks to periods of rapid evolutionary change, such as the Cambrian explosion and the rise of tetrapods, whereas periods of nutrient depletion in the Phanerozoic record seem to coincide with mass extinctions. Overall, the Phanerozoic shale studies of P highlight the potential power of the black shale record in elucidating temporal trends in P availability.

4.2. Molybdenum

Molybdenum (Mo) is important in a number of vital enzymes, including nitrogenase (used for nitrogen fixation), nitrate reductase (reduction of nitrate to nitrite), and a eukaryotic enzyme for nitrate assimilation (e.g., Kisker et al., 1997; Williams and Frausto da Silva, 2002). Because dissolved Mo is efficiently removed from sulfidic seawater, it was suggested that Mo, amongst other elements (e.g., Fe, Cu), would have been bio-limiting in the Proterozoic (Javaux et al., 2001; Anbar and Knoll, 2002) if the seafloor was covered by euxinic waters, as originally hypothesized by Canfield (1998). Anbar and Knoll (2002) referred to this relationship as a ‘bio-inorganic bridge’ through which the geochemistry of the early oceans is linked to the evolution of the biosphere. Molybdenum was considered by those authors to be the best example of how a ‘bio-inorganic bridge’ could be reflected in the sedimentary record and was the first of many metals examined from this perspective (Scott et al., 2008).

Molybdenum is sourced to the oceans primarily though oxidative weathering of the continents. Under oxic conditions Mo is largely conservative, and in modern oceans it is the most abundant transition metal in seawater (~105 nM; Collier, 1985). However, in the presence of abundant (10−2–10−4 M) free hydrogen sulfide, Mo is converted to a series of particle-reactive thiomolybdates and is efficiently removed.

![Fig. 4. Molar P to Fe ratios in iron formations through time. Reproduced from Planavsky et al. (2010b).](image-url)
from seawater (the so-called Mo “geochemical switch”; Helz et al., 1996). On a global scale, most Mo is removed to anoxic sediments where sulfide is restricted to the pore waters. However, the rate of Mo burial is an order of magnitude higher under euxinic conditions where hydrogen sulfide is present in the bottom water column. Algeo and Lyons (2006) demonstrated that the Mo/TOC ratio of sediments in modern euxinic settings is positively correlated with the concentration of dissolved Mo in the water column, which scales with local controls and/or the global extent of euxinia. For these reasons, the concentration of Mo in sediments deposited under well-constrained water column redox conditions can be used to track the oxygenation of the oceans through time. For instance, following the GOE, when the Mo weathering flux is assumed to have been robust, concentrations of Mo in euxinic sediments can provide a useful constraint on the spatial extent of euxinia on a global scale, with higher euxinic sediment Mo concentrations expected in oceans that were otherwise well-oxygenated and Mo replete (e.g., Scott et al., 2008; Dahl et al., 2010; Scott and Lyons, 2012; Kendall et al., 2015a).

Scott et al. (2008) identified two distinct first-order increases in the Mo concentrations of black shales related to stepwise increases in atmospheric oxygenation. The first is a diffuse boundary at ~2 Ga that corresponds to an initial rise in Mo/TOC ratios following the GOE. This likely resulted from an increased riverine Mo flux coupled to surface ocean oxygenation, but with subsequently suppressed enrichments related to widespread euxinia (Fig. 5). The second, sharper boundary at ~551 Ma is coincident with increased ocean oxygenation in the late Neoproterozoic. A temporal trend roughly similar to that of the black shale record (Fig. 5) was also observed in a compilation of Mo concentrations in synsedimentary to early diagenetic pyrite through time (Large et al., 2014). In the pyrite record, Large et al. (2014, 2015) identify the Mo increase in the Neoproterozoic at ~660 Ma. This boundary has since been pushed back to ca. 800 Ma following the discovery of high Mo concentrations in earlier Neoproterozoic black shales (Sahoo et al., 2012, 2016; Chen et al., 2015; Thomson et al., 2015). However, this earlier pulse in Mo appears to be transient, with Mo enrichments showing a systematic stratigraphic drop (Sahoo et al., 2016).

Scott et al. (2008) suggested that the abundance of Mo and Mo/TOC ratios in Proterozoic black shales are consistent with a paleomarine Mo reservoir that was only 10–20% of the modern ocean. Such a reduced reservoir is close to concentrations that are biomimicking for nitrogen-fixing cyanobacteria (~5% of modern seawater; Zerkle et al., 2006). Importantly, Scott et al. (2008) pointed out that the lower Mo levels indicated for the Proterozoic did not necessarily require widespread euxinic conditions. Expansion of less Mo-reactive, non-euxinic sediments (where sulfide is present but restricted to the pore waters; Scott and Lyons, 2012) to just 10% of the seafloor could accomplish the observed drawdown. Recent work emphasizing an updated mass-balance model for the Mo geochemical cycle coupled with other geochemical data (e.g., sedimentary Fe speciation; Cr concentrations) point to a redox-stratified Proterozoic ocean with a greater extent of water column euxinia (up to 1–10% of the seafloor) compared to today—but also containing a wide expanse of anoxic and ferruginous deep waters where Mo burial rates were likely lower than those under euxinic conditions (Planavsky et al., 2011; Reinhard et al., 2013b).

Molybdenum (Fig. 5) has also been used to track early stages of photosynthetic O2 production since its primary source to the oceans is through oxidative weathering of the continents (Siebert et al., 2005; Anbar et al., 2007; Wille et al., 2007; Kendall et al., 2010). Following river transport of Mo to the oceans, Mo accumulates in surface waters if they have been mildly oxygenated. In deeper anoxic waters, Mo reacts with sulfide to form oxythiomolybdate complexes (e.g., MoO4 − 2S2 −), which are scavenged from solution by organic matter and sulfide minerals (Erickson and Helz, 2000; Algeo and Lyons, 2006; Helz et al., 2011; Chappaz et al., 2014). Therefore, the availability of Mo in Archean black shales, similar to rhenium (Re), can be used as evidence for mild surface ocean oxygenation and possibly brief “whiffs” of atmospheric oxygen (Anbar et al., 2007). Similar logic was applied to mild Mo enrichments observed in ca. 2.7–2.5 Ga black shales from other sedimentary sections, such as in the Griqualand West Basin, South Africa (Siebert et al., 2005; Wille et al., 2007). An alternative explanation is provided by Lalonde and Konhauser (2015) who suggested that the oxidative weathering responsible for these brief mobilizations of Mo may be due to local oxidation of crustal material by microbial mats on land, a scenario not reliant on atmospheric oxygenation. This remains an area of intense interest and ongoing research.

Kendall et al. (2010) used the coupled geochemical behavior of Mo and Re to show that mild oxygenation, possibly spanning hundreds of meters of the upper water column, occurred on the slope of the 2.6–2.5 Ga Campbell-Malmani carbonate platform. This interpretation stems from the similar ease by which Mo and Re are mobilized from the readily oxidized crustal sulfide minerals, yet they exhibit different burial rates when deposited in sediments where oxygen penetration and dissolved sulfide concentrations in pore waters are low (Crusius et al., 1996; Morford and Emerson, 1999; Morford et al., 2005). Such conditions result in authigenic Re enrichment in sediments without Mo because Mo sequestration requires free sulfide, either in the water column or sediment pore waters.

Molybdenum isotopes have also been used to reconstruct the oxygenation of the oceans through time. Duan et al. (2010) used Mo isotope data from the ca. 2.5 Ga Mt. McRae shale to infer that small amounts of oxygen mobilized Mo from crustal sulfide minerals and that some Mo was subsequently adsorbed to oxide mineral surfaces on land or in the surface oceans ~50 Ma before the GOE. Similarly, Czaja et al. (2012) used coupled Mo and Fe isotope data from 2.68–2.50 Ga carbonates and black shales deposited on the slope of the Campbellrand-Malmani platform to confirm the presence of free dissolved O2 in the water column at the time of deposition. This relationship suggests that Fe oxyhydroxides formed on the Campbellrand-Malmani carbonate platform were likely the result of Fe(II) oxidation by dissolved O2 rather than by photoferrotrophic, and importantly, that photosynthetic O2 production by cyanobacteria was initiated by ~2.7 Ga. However, an alternative plausible explanation for the observed isotopic variations of Czaja et al. (2012) is the precipitation of isotopically heavy Fe oxyhydroxides as the result of anoxogenic photosynthesis and carrying isotopically

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**Fig. 5.** Molybdenum in black shales through time. The Mo record in black shales shows a relatively systematic increase through time, with a minor spike near the GOE, followed by a decrease in the Paleoproterozoic and Mesoproterozoic, and an increase in the Neoproterozoic to modern. This can be seen in the range of values recorded (blue) as well as formation average (black). This plot is a variation of data presented in Reinhard et al. (2013b). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
light Mo sorbed to the mineral surface—thereby not definitively indicating the process of oxygenic photosynthesis.

Recently, Planavsky et al. (2014b) examined the Mo isotope record of Mn-rich IF of the ca. 2.95 Ga Sinqeni Formation, Pongola Supergroup (South Africa) and found evidence for small amounts of oxygen that likely would have existed in the form of a transient oxygen oasis at the ocean’s surface (as per Olson et al., 2013). Planavsky et al. (2014b) further argued that the Mo isotope data of the Sinqeni Formation preserves the expression of Mo isotope fractionation during adsorption of Mo to Mn(IV)-oxides in shallow marine settings. The inferred presence of Mn(IV) oxides requires that free oxygen was available in shallow marine waters because O$_2$ is the only oxidizing agent other than H$_2$O$_2$ strong enough for oxidation of Mn(II) to Mn(IV); however, see Johnson et al. (2013) for an alternative view. Hence, the Mo enrichment and isotope records suggest that oxygenic photosynthesis likely evolved by >500 million years prior to the GOE.

Molybdenum isotope data has also been used in conjunction with U and Fe isotope data and sedimentary Fe speciation to demonstrate that a dramatic decline in ocean oxygenation occurred following the GOE (Asael et al., 2013; Partin et al., 2013b). Asael et al. (2013) calculated a dramatic decline in ocean oxygenation occurred following the GOE (Asael et al., 2013; Partin et al., 2013b). Asael et al. (2013) calculated the seawater Mo isotope composition (δ$^{98}$Mo) to be 0.85 ± 0.20‰, which is significantly lighter than modern seawater (2.3‰) and thus consistent with an appreciable extent of euxinic waters in the Proterozoic ocean. The interpretation of euxinia was further supported by the iron speciation analysis conducted on their samples. A similar conclusion was drawn for the late Paleoproterozoic and early Mesoproterozoic ocean based on sedimentary Fe speciation and Mo concentration and isotope data from the ca. 1.8 Ga Rove Formation (Canada) and ca. 1.4 Ga Velkerri Formation (Australia) (Arnold et al., 2004; Kendall et al., 2009, 2011). Mo isotope signatures for expanded ocean euxinia have also been documented from Neoproterozoic black shales, including the ~750 Ma Walcott Member of the Chuar Group, Grand Canyon (Dahl et al., 2011) and the ca. 640 Ma Black River Dolomite (Kendall et al., 2015a). It is not until ca. 560 Ma that seawater Mo isotope compositions similar to modern seawater are first inferred from the sedimentary rock record (see Sahoo et al., 2016).

A recent compilation of Mo isotope data for euxinic black shales also reveals that significant oscillations in ocean redox conditions, and hence the seawater Mo inventory, may have occurred across the Precambrian-Phanerozoic transition before the advent of permanent widespread ocean oxygenation (Dahl et al., 2010; Kendall et al., 2015a; Sahoo et al., 2016). This view adds a new dimension to existing evidence for predominantly ferruginous deep ocean conditions in the Proterozoic (e.g., Canfield et al., 2007, 2008; Poulton et al., 2010; Poulton and Canfield, 2011; Johnston et al., 2010; Planavsky et al., 2011; Reinhard et al., 2013b; Li et al., 2015; Guibald et al., 2015), whereby transient ocean oxygenation events (OOEs) may have punctuated periods of relatively redox stasis (Sahoo et al., 2016). However, some caution must be exercised when viewing the Mo isotope record in black shales as a consistent tracer of seawater signatures. This is due to variability observed in the modern sediments of the euxinic Black Sea, where Mo isotope fractionations vary as a function of sulfide concentration (Neubert et al., 2008). Essentially, in order to record a seawater signature, a critical level of at least 11 μmol L$^{-1}$ H$_2$S$_{(aq)}$ is required. Under more weakly euxinic conditions, Mo removal may not be quantitative, which can result in variable isotopic fractionation between the sediment and overlying water column. This isotopic fractionation is due, at least in part, to the fractionation of Mo isotopes amongst various aqueous species (Tossell, 2005), and accordingly a critical amount of sulfide is required to ensure that all Mo is in the tetrathiomolybdate phase and that Mo is quantitatively scavenged. Under such a scenario, the Mo isotope composition of sediment deposited under non- to weakly-euxinic conditions would depart from that of the source (i.e., contemporaneous seawater), thereby affecting the ability of euxinic sediments to track the Mo isotope composition of the early oceans. Accordingly, in order to track the Mo isotope composition of the early oceans, black shales need to be deposited under conditions with persistent and appreciable levels of dissolved sulfide (Neubert et al., 2008; Gordon et al., 2009; Arnold et al., 2012).

The paleomarine Mo record (Fig. 5) also has important implications for biological evolution in the Neoproterozoic. In line with related suggestions by Anbar and Knoll (2002), Boyd et al. (2011) suggested that the emergence of Mo-Fe nitrogenases appears to correlate with, at least transiently, increasing seawater Mo concentrations around the GOE. However, this timing has recently been challenged on geochemical evidence. Stüeken et al. (2015a) report a suite of fluvial to marine sedimentary rocks at 3.2 Ga that have an average nitrogen isotopic composition (δ$^{15}$N) of 0.0 ± 1.2‰. They argue that the only way to record such a signal is through a biological pathway utilizing Mo-nitrogenase and that any abiological pathway of nitrogen fixation or an alternative nitrogenase would inherently impart a substantially different N isotope fractionation. These studies suggest not only a relatively early origin for Mo-nitrogenase but also have implications for the bioavailability of Mo at very low dissolved concentrations in the Mesoarchean. This would be physiologically consistent with observations of Mo-nitrogenase activity at very low Mo levels in culture studies (Zerkle et al., 2006; Glass et al., 2009). By contrast, Sánchez-Baracaldo et al. (2014) used phylogenetic reconstruction to suggest the Cryogenian (0.85 to 0.635 Ga) as the origin of N-fixing cyanobacteria. However, the history of Mo as recorded by the chemical sedimentary record is more consistent with an early origin for Mo-nitrogenase, or at least its precursor, and a subsequently consistent limitation of N-fixation by a Mo-depleted Proterozoic ocean (i.e., Lyons et al., 2014b).

4.3. Nickel

Nickel (Ni) is critical for many prokaryotic metalloenzymes. It is utilized in carbon reduction by both acetogenic and methanogenic bacteria, as well as the cofactors methyl-coenzyme M reductase and acetyl-CoA synthase, both of which are required for methane production (Hausinger, 1987; Zerkle et al., 2005, and references therein). Furthermore, Ni is used in hydrogenases, carbon monoxide dehydrogenase, and catalyzes the reduction of CO$_2$ to CO and the resultant production of acetyl-CoA (e.g., Ragsdale and Kumar, 1996). In non-methanogens, Ni may be used in urease and in a recently evolved superoxide dismutase found in many marine organisms (Frausto da Silva and Williams, 2001; Dupont et al., 2008).

Initial estimates of Ni concentrations in the early ocean were derived from geochemical modeling (Saito et al., 2003) and microbial genomics...
Studies in the ferruginous ocean analogue Lake Matano, Indonesia, have shown that green rust may play an important role in scavenging Ni from ferruginous water columns (Zegeye et al., 2012). Further, production of trace hydrogen sulfide through microbial sulfate reduction may play a strong role in governing aqueous Ni concentrations, even under ferruginous conditions (Crowe et al., 2008; Zegeye et al., 2012). In Lake Matano, Ni and Co exhibit divergent behavior implying that different biogeochemical process govern concentrations of each metal (Crowe et al., 2008).

4.4. Zinc

Zinc (Zn) is amongst the most biologically important trace metals, particularly for eukaryotes, and is a component in a wide variety of metallo-peptides and polymerases (see Lipscomb and Sträter, 1996 for a review on Zn enzymology). Many Zn metalloenzymes are used in processes involving DNA or RNA synthesis (e.g., Lipscomb and Sträter, 1996; Berg and Shi, 1996). In eukaryotes, Zn is also used in Zn-fingers, small protein structural motifs which act as signaling agents in processes centralized in the nucleus and are thought to have emerged relatively late (e.g., Berg and Shi, 1996; Dupont et al., 2006, 2010).

Prior to compilations of Zn data from the sedimentary proxy record, the only estimates for Zn concentrations and bioavailability came from geochemical modeling (e.g., Saito et al., 2003) (Fig. 1). This modeling work was consistent with the emergence of eukaryotic metalloenzymes and rapid diversification of eukaryotes in the Neoproterozoic following a transition from a widely anoxic ocean with expanded euxinia to a well-oxygenated ocean (Dupont et al., 2010). This view provides a possible explanation for the delay in eukaryotic diversification.

Recently, however, two studies have reevaluated paleomarine concentrations through time, and thereby Zn bioavailability, using the sedimentary rock record. Scott et al. (2013) focused on black shales, building from the observation that Zn/Al ratios in sediments from modern eucinian basins are positively correlated with dissolved Zn concentrations in bottom waters. Scott et al. (2013) found there was no evidence in the Precambrian black shale record to infer a depleted paleomarine Zn reservoir—because the average Zn concentration in Precambrian and Phanerozoic eucinian shales are not significantly different. Accordingly, they suggested that seawater Zn levels remained broadly uniform throughout time at near modern levels and several orders of magnitude above concentrations that would be biolimiting. This finding was bolstered by Robbins et al. (2013) who examined Zn concentrations in iron formation through time (produced from Robbins et al., 2013). This version includes all available IF data and is plotted without the removal of samples indicating detrital contamination (i.e., <1% Al2O3 or >0.1% TiO2). Even when the all samples are included, Zn/Fe ratios in IF trend to fall in a field that spans 2–3 orders of magnitude and is relatively consistent through time. The only notable deviation from this long term trend is the most modern samples, which may be reflective of local Zn enrichment in near-vent hydrothermal exhalative sediments. See Robbins et al. (2013) for further discussion.

![Fig. 7. Concentrations of Zn in iron formation through time (produced from Robbins et al., 2013). This version includes all available IF data and is plotted without the removal of samples indicating detrital contamination (i.e., <1% Al2O3 or >0.1% TiO2). Even when the all samples are included, Zn/Fe ratios in IF trend to fall in a field that spans 2–3 orders of magnitude and is relatively consistent through time. The only notable deviation from this long term trend is the most modern samples, which may be reflective of local Zn enrichment in near-vent hydrothermal exhalative sediments. See Robbins et al. (2013) for further discussion.](https://example.com/fig7)
and Zn/Fe ratios in IF (Fig. 7) and found generally constant Zn enrichments through time. When viewed alongside updated geochemical models for Zn speciation, and considering hypothesized partitioning scenarios for Zn and Fe co-precipitation, Robbins et al. (2013) estimated a paleomarine Zn reservoir on the order of 10 nM. This value is several orders of magnitude above the ~10^{-13} M concentration considered as biolimiting (Brand et al., 1983) and in excellent agreement with the black shale record (Scott et al., 2013). However, Zn displays nutrient-type behavior in modern seawater, and it is possible that Zn drawdown also occurred in ancient surface waters; sedimentary proxy records are generally considered to reflect the overall size of the paleomarine reservoir without addressing the finer details of its vertical structure.

The updated view of a relatively static paleomarine Zn reservoir (Fig. 7) contradicts the findings of earlier geochemical models that linked (1) the low Zn requirements of cyanobacteria to the predicted low levels of total Zn in the Precambrian oceans, and also (2) the proliferation of eukaryotes to an increase in total Zn during the Phanerozoic (e.g., Saito et al., 2003; Dupont et al., 2010). This disparity can be linked to an overestimation of the effects of Zn aqueous complexation by sulfides during the Precambrian by early geochemical models (e.g. Saito et al., 2003). Additionally, this model directly contradicts the behavior of Zn in some modern anoxic aquatic systems. For instance, in some conditions where a strong redoxcline exists, such as in Jellyfish Lake, Palau, total dissolved Zn concentrations increase with depth due to the formation of aqueous sulfide complexes (Landing et al., 1991; Dierssen et al., 2001). Alternatively, Robbins et al. (2013) proposed that an increased proportion of hydrothermal fluxes relative to riverine fluxes and the transport of hydrothermal Fe- and Zn-rich fluids to more distal environments in anoxic Archean and Paleoproterozoic oceans could have helped maintain a large marine Zn reservoir.

This view of severe Zn limitation (e.g., Saito et al., 2003) was also based the assumption that its availability for biological use was limited to free dissolved Zn^{2+}. In the modern oceans, the majority of Zn in surface waters is complexed with organic ligands (90–98%). This organic ligand pool has traditionally been viewed as being a non-bioavailable component of the total Zn pool (Bruland, 1989). This framework was based on culture work that has either not applied synthetic ligands to culture or introduced EDTA (a strongly binding organic ligand), neither of which are an accurate representative of natural waters. In contrast, it was noted more recently by Lohan et al. (2005) that marine phytoplankton appear to be much more tolerant of low dissolved Zn concentrations than in culture studies. The authors then went on to propose that Zn binding organic ligands may be produced by phytoplankton in order to facilitate biological uptake, and to regulate ambient Zn concentrations. Consistent with this idea is recent culture work by Xu et al. (2012) and Aristilde et al. (2012), where it was demonstrated that weakly binding organic ligands can in fact increase Zn uptake rates in phytoplankton. This work highlights a gap in our understanding of Zn forms available for biological use, and complicates predictions on its past bioavailability. As such, debate of Zn bioavailability calls for new methods and data so as to directly test this in future work.

There is also potential for further inferences to be drawn from the stable isotope record of Zn in chemical sediments, such as carbonates. A study on a Neoproterozoic (Marinoan) cap carbonate sequence shows variations in δ66Zn that are suggested to record an increase in surface runoff and continental weathering coincident with the onset of deglaciation and a subsequent increase in primary productivity caused by development of nutrient-rich surface waters (Kunzmann et al., 2013). A single study on Zn isotopes in IF (Pons et al., 2013) found some of the most enriched δ66Zn values between 2.7 Ga and 1.8 Ga. Variations were considered to be rapid and governed by two factors: preferential incorporation of isotopically heavy Zn into carbonates and a significant increase in the amount of marine sediments exposed to weathering around 2.7 Ga as the result of increased continental land mass. These studies highlight the potential of the Zn stable isotope record for understanding links between trace metal in past environments availability and biological productivity, as well as examining source-sink relationships.

4.5. Cobalt

The most recognized biochemical role of cobalt (Co) is its participation as a cofactor in cobalamin (vitamin B12), which is essential for a number of metalloenzymes, including methionine synthase and ribonucleotide reductase (e.g., Marsh, 1999; Frausto da Silva and Williams, 2001). In previous modeling work, researchers have pointed to the antiquity of the cobalamin cofactor, suggesting that its origin somewhere between 3.5 and 2.7 Ga is consistent with the evolution of cyanobacteria in a Co-rich ocean (Saito et al., 2003). Additional biochemical functions of Co exist (Kobayashi and Shimizu, 1999), including in cambialistic carbonic anhydrase enzymes (Morel et al., 1994; Roberts et al., 1997), and perhaps more will be discovered with emerging metalloproteomic techniques (e.g., Waldron et al., 2007; Aguirre et al., 2013). Modeling efforts have indicated that seawater Co concentrations could have been relatively high in the Archean, decreased in the Proterozoic, and then further decreased following Neoproterozoic ocean oxygenation (Saito et al., 2003). Despite the predicted decreases in the seawater concentration of Co over time, the biological utilization of Neoproterozoic ocean oxygenation (Saito et al., 2003). This may be related to the occurrence of Co-binding ligands in Co-limited waters, which exerts strong control over the speciation of Co in the modern ocean (Saito and Moffett, 2001; Saito et al., 2005). Indeed, a high degree of Co complexation may favor cyanobacteria over eukaryotic phytoplankton in modern settings containing picomolar levels of dissolved Co, for example in the Costa Rica upwelling dome (Saito et al., 2005; Ahlgren et al., 2014).

An examination of Co concentrations in IF and pyrite through time suggests an expansion of the paleomarine Co reservoir between 2.8 and 1.8 Ga (Swanner et al., 2014). This conclusion is indicated by a large increase in Co/Ti ratios in IF (Fig. 8), euxinic shales, and pyrite relative to the evolving continental crust, with a peak in Co/Ti values observed at ~2.4 Ga. The expansion of the Co reservoir between 2.8 and 1.8 Ga (Fig. 8) may be coincident with increased mantle plume activity and associated hydrothermal inputs (Swanner et al., 2014). This expansion is also broadly consistent with an increase in genes that bind Co around this time (David and Alm, 2011). Such conditions simultaneously allow for the establishment of the ferruginous conditions necessary for IF precipitation and an increase in the amount of Co introduced to the ocean. Pervasive anoxic conditions would help keep Co in solution, and as such the residence time of Co would be higher in more anoxic oceans compared to modern, well-oxygenated oceans (c.f., Swanner et al., 2014; also Saito and Moffett, 2002). This is consistent with the
relatively high Co concentrations in the anoxic ferruginous waters of Lake Matano compared to its oxic surface waters (Crowe et al., 2008). However, the syngentic and early diagenetic pyrite record for Co presented by Large et al. (2014) indicates a steadier decline in Co levels from 3 Ga to present. Resolving the discrepancy between the IF and pyrite records would be of great interest, and is an area where a more complete black shale record may prove useful in resolving the temporal trends in marine Co from the Archean through to the modern. If the pyrite record for Co proves correct in predicting the overall trajectory, it will be quite telling, as this would then mirror the general trajectories for Fe and Mn as well, and may offer support for trace element variations as correlative to large igneous events.

4.6. Chromium

Although chromium (Cr) has been identified as a biological component in some higher-level organisms, it is generally regarded as a toxin (Frausto da Silva and Williams, 2001). As such, temporal trends in seawater are unlikely to mirror direct evolutionary controls. Rather, investigations in the sedimentary record have focused on Cr abundances and isotope compositions for tracking oxygenation as well as the impact of anaerobes in the surface environment. This utility is due to reduced Cr in the form of Cr(III) being effectively immobile at neutral to alkaline pH; Cr becomes mobile when it is oxidized to Cr(VI). Furthermore, Cr is subject to strong stable isotope fractionation during redox reactions, especially reduction (Ellis et al., 2002). As such, variations in Cr abundances and isotopic compositions are likely to represent changes in the redox state of the oceans or atmosphere, as well as associated changes in the mechanisms of Cr mobility and sediment provenance.

Reinhard et al. (2013b) investigated Cr and Mo enrichments in anoxic and euxinic shales, coupling these two elemental systems in order to constrain the extent of anoxia versus euxinia in the early oceans. For Cr, there are no significant authigenic enrichments in middle Proterozoic black shales and this was taken to indicate pervasive anoxia (Fig. 9). Conversely, Mo is enriched in middle Proterozoic black shales to an intermediate degree relative to Archean and Phanerozoic samples (Fig. 5), which indicates the presence of euxinic conditions that were relatively limited on the seafloor—although still far greater than the extents observed today. This distinction is the result of differences in the geochemistry of Cr and Mo. Efficient burial of Cr can occur in sediments deposited under anoxic and ferruginous waters. By contrast, Mo burial, as discussed above, is most efficient in sediments deposited from euxinic waters. Together, these two elements can constrain the relative extent of ferruginous and euxinic conditions. In this regard, Reinhard et al. (2013b) estimated at least 30–40% of the middle Proterozoic seafloor was anoxic, and possibly much more, with only 1–10% of the seafloor being euxinic. Large et al. (2014) also investigated Cr and Mo concentrations in sedimentary pyrite and found patterns consistent with the dominance of anoxic settings in the Proterozoic from 2.15 to 0.7 Ga, with euxinic settings more prevalent between 1.2 and 0.8 Ga. Large et al. (2014) attributed a decrease in Cr concentrations, from the Archean to the Proterozoic (Fig. 9), as reflective of a change in source availability. They argued that Cr should be sourced from ultramafic rocks similar to Ni and Co, and that this decrease may reflect a decrease in the abundance of the source (Fig. 9; also see their Fig. 8c).

Chromium isotope enrichments have also been used to track the oxygenation of the atmosphere and oceans. For instance, small variations in Cr isotope composition in 2.8–2.6 Ga IF were first suggested to document brief pulses in atmospheric oxygen, which mobilized Cr via oxidative weathering and led to Cr sequestration in IF prior to the GOE (Frei et al., 2009). Those authors argued that increases in $\delta^{53}$Cr of +0.04 to +0.29‰ in IFs at 2.7, and again at 1.8 Ga, are the direct result of Cr(III) being oxidized to Cr(VI) and that oxidation of Cr must have been catalyzed by oxidized Mn$^{4+}$ in the form of MnO$_2$. Subsequently, Konhauser et al. (2011) examined temporal trends in the degree of Cr enrichment in IF and found a peak at 2.48–2.32 Ga (Fig. 10). This peak was suggested to indicate the onset of acidic weathering triggered by microaerophilic iron-oxidizing bacteria which accelerated the weathering of pyrite as soon as some atmospheric oxygen became available. Muted $\delta^{53}$Cr variations (−0.3 to +0.3‰) at this time were taken as being indicative of Cr cycling in reduced form (Konhauser et al., 2011), in contrast with highly variable values coincident with ocean oxygenation in the Neoproterozoic (up to +4.9‰; Frei et al., 2009).

Chromium isotope data were also acquired from rocks from the Pongola Supergroup, South Africa, in an effort to seek evidence for earlier signs of photosynthetic oxygen production. Crowe et al. (2013) found that the $\delta^{53}$Cr isotope compositions of 2.96 Ga paleosols were fractionated relative to the crustal values, indicating oxidative mobilization of Cr and the presence of low levels of atmospheric oxygen (<10$^{-4}$ PAL; the level required to prevent reduction of Cr(VI) by Fe(III) during transport to the ocean). This conclusion would suggest the presence and activity of oxygenic photosynthesis about ~600 million years prior to S-MIF disappearance during the GOE. The idea of early O$_2$ production is supported by the Mo isotope composition of IF also obtained from within the Pongola Supergroup, where scaling between Mo isotope composition and Mn/Fe ratio indicate the presence of Mn(IV) oxide exit channel for Mo at that time (Planavsky et al., 2014b). However, it is also plausible that the Cr isotope fractionation observed by Crowe et al. (2013) in the 2.96 Ga paleosols is due to the localized production of massive oxidation of pyrite by aerobic bacteria on land around the time of the GOE. The data, normalized to the upper continental restoration model of Condie (1993), has been replotted from Konhauser et al. (2011).
of oxygen by cyanobacteria within a microbial mat (see Lalonde and Konhauser, 2015). In either case, the sedimentary Cr record, in terms of abundance and especially in isotopic composition, appears to provide important clues regarding oxygen production due to photosynthesis either in the early oceans, or on land, and corresponding oxidative weathering on the early continents. Investigations into the isotopic record of Cr in black shales and IF will continue to be a useful tracer for paleoredox conditions in the Precambrian oceans.

Recently, Planavsky et al. (2014a) presented δ53Cr data from Precambrian and Phanerozoic ironstones, as well as Neoproterozoic to Phanerozoic black shales and mudstones. Collectively, this record shows that δ53Cr values in Precambrian black shales are similar in composition to crustal Cr until the Neoproterozoic. Beginning approximately 0.8 Ga, relatively large positive δ53Cr values are observed in black shales and mudstones, as well as in some Phanerozoic ironstones. Those authors interpreted these trends in δ53Cr to suggest that atmospheric oxygen levels were <0.1% PAL (below this level, Cr[III] oxidation is limited by the lack of Mn oxides) prior to the Neoproterozoic. This concentration would have been limiting for metazoans, potentially explaining the delay in their diversification (although see Zhang et al. (2016) for an alternative opinion, as well as a comment on the opposing view by Planavsky et al. (2016)). The capacity of δ53Cr from shales and ironstones to resolve changes in oceanic oxygenation state depends on our ability to discriminate authigenic from detrital Cr, and this stands as a key challenge and opportunity. Nevertheless, the Cr record links biological and geochemical evolution indirectly via the effect that O2 has on both.

In an effort to establish a more rigorous calibration of the Cr isotope redox proxy in the modern, Reinhard et al. (2014) examined Cr isotope fractionations in the recent sediments of the Cariaco Basin, Venezuela. A slight increase in δ53Cr coupled with elevated Cr enrichments is coincident with the onset of deep-water euxinic conditions in the basin. The authors argued that this observation suggests that such sediments, and their ancient equivalents, can be used to track the Cr isotope composition of the oceans over time—because the δ53Cr signature of the overlying water column is approximately recorded by sediments deposited under an anoxic water column. Similar evidence of the reliability of black shales in tracking δ53Cr signatures is presented by Gueguen et al. (2016) from the upwelling zone of the Peru Margin, comparable to modern deepwater sediments. However, Scheiderich et al. (2015) recommended a more cautious approach to Cr isotope proxies based on important variability in Cr concentrations and isotopic compositions in the Arctic ocean and between the Arctic, Atlantic, and Pacific Oceans, emphasizing the need to better understand modern marine Cr cycling.

Most recently, Cole et al. (2016) have expanded the record of black shale δ53Cr values throughout Earth history, illustrating a base-level shift at ~850 Ma from largely unfractionated δ53Cr values similar to the composition of continental crust to highly fractionated δ53Cr values. The authors have used this record to suggest a lack of significant terrestrial oxidative weathering of Cr prior to 850 Ma indicative of oxygen levels below ~0.1% PAL (based on the pO2 estimate from Planavsky et al., 2014a). Again, capacity to constrain authigenic versus detrital Cr is key to these interpretations.

4.7. Iodine

The history of iodine (I) cycling through time is biologically relevant, as its redox transformations occur at oxygen levels similar to proposed minimal requirements of early animals. Under anoxic conditions iodine occurs in the form of iodide (I−), which is excluded from the crystal lattice of carbonates during their precipitation. When dissolved oxygen concentrations are ≥1 μM, iodate (IO3−) is formed (Farrenkopf and Luther, 2002; Luther and Campbell, 1991; Rue et al., 1997) and is incorporated into carbonate minerals (Lu et al., 2010). This relationship suggests that the simple presence of iodine in carbonate indicates the presence of O2 ≥1 μM at the site of carbonate precipitation, i.e., in surface waters (Hardisty et al., 2014). This level of dissolved O2 in surface waters is generally consistent with the 0.1–1% PAL proposed as necessary for early animals (Mills et al., 2014; Sperling et al., 2013). However, the level of oxygen at depth remains unconstrained by the I record, and may still have posed a challenge to complex life.

A study of iodine concentrations in the carbonate record from the Neoarchean through Paleoproterozoic, indicate oxidation of the surface coincident with the GOE. This is reflected by a switch from the absence to presence of carbonate-bound iodine (Hardisty et al., 2014). Such a record indicates the potential for habitable marine environments, at least in terms of minimum O2 thresholds, hundreds of millions of years before the first evidence for animal life or eukaryotic diversification (Cingras et al., 2011; Knoll, 2014). However, records of iodine time are currently limited, with measurements lacking through most of the Proterozoic and Paleozoic.

Beyond redox chemistry at biologically critical O2 levels, iodine is a bioessential element (Frausto da Silva and Williams, 2001), specifically within marine algae that accumulate iodine primarily in the form of iodide to reach total iodine concentrations up to several hundred ppm (Elderfield and Truesdale, 1980; Fuse et al., 1989; Muramatsu and Hans Wedepohl, 1998). These values are far greater than those found in limestones (a few ppm) (Lu et al., 2010; Muramatsu and Hans Wedepohl, 1998; Zhou et al., 2014, 2015). This organically bound iodine can ultimately end up as a component of organic-rich rocks, particularly black shales, where iodine-to-carbon ratios are slightly elevated relative to those found in algae: −1.5 × 10−4 (Elderfield and Truesdale, 1980). However, most of the original organic-bound iodine is lost through diagenetic carbon remineralization (Kennedy and Elderfield, 1987). The reason for such high concentrations in marine algae is still unclear, but at least one primary role is as an antioxidant (Küpper et al., 1988, Venturi et al., 2000). It has also been shown that brown algae assimilate iodine as iodide, which is excreted to react with oxygen and other reactive oxygen species, providing protection against oxidative stress (Küpper et al., 1988). Regardless of this and other roles, the end result is that biological assimilation and burial act as the main marine iodine sink, with no known quantitatively important mineralogical sinks. Diagenesis of organic matter is thought to release ~99% of bound iodine back to seawater, such that internal cycling dominates the marine iodine budget; contributions from continental weathering and hydrothermalism appear negligible today (Lu et al., 2010).

Though the redox requirements for the accumulation of marine iodate act as the primary control for I/(Ca+Mg) ratios, the potential for carbonates to track changes in the marine iodine reservoir as well as the dominant iodine species in shallow ocean settings could provide a critical constraint on the evolution and abundance of marine algae through time. In oceanian form, the halogens Br, I, and Cl are all relatively soluble, and one might predict that marine iodine concentrations relative to the bulk silicate Earth would be similar to that of Br and Cl. However, normalized seawater is depleted in iodine by a factor of ~40 when compared to these other halogens (Sharp and Draper, 2013). This discrepancy can be explained by uptake and burial in association with marine algae, whose affinity for iodine is high relative to Br and Cl (Elderfield and Truesdale, 1980; Sharp and Draper, 2013), but this possibility requires that marine iodine concentrations have decreased through time. Perhaps decreases in marine iodine abundance through time occurred in pulses following the emergence and dominance of various algal clades? This pathway could be particularly relevant for the Proterozoic, where filamentous microfossils suggest red algae rhodophytes in carbonates up to 1200 Ma old (Butterfield, 2000; Knoll et al., 2013) and potential green algae chlorophytes as old as 1800 Ma (Knoll, 2014; Moczydlowska et al., 2011).

Recent studies have tested the idea that organic carbon burial provides a secondary control on I/(Ca+Mg) ratios on shorter timescales by evaluating I/(Ca+Mg) in association with positive δ13C,carb excursions during Phanerozoic Oceanic Anoxic Events (OAEs), particularly OAE-2 (Lu et al., 2010; Zhou et al., 2015). The expectation is that if

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tracking the evolution of the iodine reservoir, $I/(Ca+Mg)$ ratios will system-
atically decrease to similar minimum values in global sections in step with contemporaneous $\delta^{13}C_{\text{carb}}$ excursions. Though variable and particularly low $I/(Ca+Mg)$ ratios are observed in association with OAEs, these studies found that patterns and decreases in $I/(Ca+Mg)$ varied from section to section, and in most cases decrease prior to the $\delta^{13}C$ excursion—the opposite of what is expected given that the residence time of iodine exceeds that of dissolved inorganic carbon by an order of magnitude (Broecker et al., 1982). Instead, the $I/(Ca+Mg)$ ra-
tios may indicate a shift to low oxygen marine conditions prior to the OAE, as defined by $\delta^{13}C$ and black shale deposition. In other words, if changes in the total marine iodine reservoir coincide with anoxia, either locally or globally, this reservoir change may be unobservable because as $I/(Ca+Mg)$ exclusively tracks iodate (Lu et al., 2010; Zhou et al., 2015).

4.8. Selenium

In biological systems, selenium (Se) is found predominately in the alternative amino-acid selenocysteine, a cysteine analogue where the sulfur is replaced by selenium (Hatfield and Gladyshev, 2002). Selenocysteine is used in a variety of oxidoreductases involved in regu-
loating oxidative stress, including methionine sulfoxide reductase and glutathione peroxidase. The proliferation of selenoproteins in select lin-
eages, including the smallest free living eukaryote, Ostreococcus, has been suggested to be due to the selenocysteine-containing proteins being more efficient than the corresponding cysteine version (Palenik et al., 2007). Recently, this possibility was shown to be true for a thioredoxin, with the selenocysteine version of the protein being 10-
fold more catalytically active (Kim et al., 2015). Given these examples of Se utilization in biological systems, it is likely that Se plays a key role as a catalyst in intercellular processes.

The potential of Se and its stable isotopes as a paleo-redox tracer was first assessed by Johnson (2004) and Johnson and Bullen (2004), who calculated that significant stable isotope fractionation should occur dur-
ing redox transformations between its four common valence states (VI, IV, 0, –II). Mitchell et al. (2012) examined Se abundances and $\delta^{82/76}Se$ from a series of Cambrian to Holocene marine sediments and found sig-
nificant variability in Se concentrations, but a narrow range of Se iso-
topes values (between −1 to 1‰). Since then, more complete compilations of Se through time have been presented from the pyrite record (Large et al., 2014), as well as marine and terrestrial mudrocks from the Archean through to the modern (Stüeken et al., 2015b). In their analyses of syngenetic and early diagenetic pyrite, Large et al. (2014) showed a temporal trend in Se with several small peaks at 3.0 and 2.4 Ga, along with a Paleoproterozoic to Mesoproterozoic perturba-
tion between 1800 and 1300 Ma, and finally a rise in the Neoproterozoic to Phanerozoic. Large et al. (2014) further suggested that this trend in Se may be used as a paleo-proxy for tracking oxygenation. Those same authors also suggested that the peak in Se at ~3.0 Ga in the pyrite record is consistent with Cr and Mo isotope data at that time (Crowe et al., 2013; Planavsky et al., 2014b). Accordingly, the pyrite record would then indicate low levels of $O_2$ in the Proterozoic, although with some variability, before the onset of widespread ocean oxygenation in the Neoproterozoic.

Stüeken et al.’s examination of the Se cycle and its redox dynamic shed light on a number of aspects related to the oxygenation of the Earth, and we highlight a few key findings here. First, Stüeken et al. (2015b) suggested that increasing marine Se and S levels, and differ-
ences between terrestrial and marine settings between 2.8 and 2.7 Ga, imply that there was terrestrial oxidative weathering at that time. In 2.8–2.7 Ga non-marine samples, Stüeken et al. (2015b) found negative-
ly fractionated $\delta^{73/72}Se$ values, as opposed to coeval marine samples which are generally positive, and this negative fractionation was attrib-
uted to oxidative weathering in terrestrial settings. Second, despite hav-
ing several redox states, there is no evidence for mass-independent fractionation (MIF) of Se prior to the GOE. A general absence of Se-MIF is consistent with the findings of Pogge von Strandmann et al. (2014) who examined a small suite of 2.65–2.5 Ga black shales from the Ghaap Group in South Africa which had previously shown large S-MIF signals and should have been favorable for recording a Se-MIF signal, if one existed. Third, there is a decrease in sedimentary $\delta^{82/78}Se$ values during the period between the middle Proterozoic and the mid-
Paleozoic, which Stüeken et al. (2015b) attributed to the oxygenation of the deep ocean in the Neoproterozoic. Pogge von Strandmann et al. (2015) also examined the Se isotope record in the Neoproterozoic and found declining $\delta^{82/76}Se$ values in black shales that they attributed to a pro-
tracted oxygenation of the ocean–atmosphere system between −770 Ma and the early Cambrian.

Se isotope analyses have also been used to infer the presence of free $O_2$ in the Neoarchean (Stüeken et al., 2015c). Selenium isotopes were measured on samples from the 2.5 Ga Mt. McRae shale where Anbar et al. (2007) first reported a pre-GOE ‘whiff’ of oxygen. A peak in $\delta^{82/76}Se$ and Se enrichment factors led Stüeken et al. (2015c) to sug-
gest oxidative mobilization and fractionation of Se from terrestrial sources, similar to the conclusions of Anbar et al. (2007), Reinhard et al. (2009), Duan et al. (2010), and Kendall et al. (2015b), based on S-Mo-Re-Os systematics in the same horizon.

Selenium has also garnered interest in more recent sediments for its possible implications regarding biological activity in the Phanerozoic, especially its chemistry coincident to mass extinctions. Several global mass extinctions in the Phanerzoic have been correlated to severe Se depletion in the pyrite record, including the end-Ordovician, −Devonian and −Triassic events (Long et al., 2016). Coincident with these extinctions the pyrite record shows a dramatic decrease in Se, al-
most two orders of magnitude, suggesting that organisms relying on Se for biological functions may have become stressed, and that this may have been a contributing factor in the extinctions. Selenium has also provided clues for other Phanerozoic mass extinctions. For instance, Stüeken et al. (2015d) interpreted a profile from an outcrop in southern Alberta, Canada to indicate a brief period of euxinia, followed by a highly productive oxic period before the end-Permian mass extinction. They observed a large negative $\delta^{82/76}Se$ excursion and interpreted it to reflect a large export of organic Se to the sediments immediately prior to the extinction, suggesting that productivity was high and nutrient limita-
tion was likely not a cause of the extinction, at least locally. However, Stüeken et al. (2015d) do suggest that nutrient limitation may have played a role in the recovery from the mass extinction. In these ways we can see how Se evidence in black shales and pyrite may shed light on more recent evolutionary events in marine geochemical and biolog-
ical evolution.

4.9. Uranium

The redox-sensitive element uranium (U), though not bioessential, provides constraints on the dynamics of the evolving oxygenation of the Earth. Some bacteria catalyze the reduction of $U(VI)$ to $U(IV)$ in order to obtain energy, including a few dissipatory Fe(III)- and sulfate-reducing bacteria (e.g., Lovley et al., 1991; Behrends and Van Cappellen, 2005). Microbial $U(IV)$ reduction, in addition to abiotic mechanisms of reduction such as sorption to organic matter or co-
precipitation with iron oxides, sequesters marine U into sediments. These processes provide a means to use U concentration in the sedi-
mentary record as a proxy for changes in the oxygen content of the atmosphere-ocean system. In terms of the sedimentary record, U abund-
ances have been investigated in both the IF and black shale records (Partin et al. (2013a,b), respectively, while U isotopes have been used in some studies to better illuminate the oxygenation of the oceans (Asael et al., 2013; Kendall et al., 2013, 2015a; Dahl et al., 2014). More recent work has also demonstrated that U isotopes can potentially be used to track the biotic reduction of $U(VI)$, providing the potential for a novel redox biosignature proxy in the rock record (Stylo et al., 2015).
in modern euxinic settings (Asael et al., 2013). This low redox state is consistent with a decrease in oxygenation of the atmosphere-ocean system following the GOE, as inferred for the late Paleoproterozoic from the black shale U abundance record (Partin et al., 2013b). Similarly, Kendall et al. (2015a) used coupled U and Mo isotopes from the Ediacaran Doushantuo Formation to examine ocean oxygenation dynamics around the time of metazoan diversification. High $\delta^{238}$U values indicate pervasive ocean oxygenation between 560 and 551 Ma. Collectively, studies such as these have demonstrated the utility in using U, a non-bioessential element, for tracking the redox evolution of the Earth’s atmosphere and oceans. This, in turn, has profound effects for the evolution of the biosphere, and perhaps even for tracking the biotic signature of U reduction in the rock record (Stylo et al., 2015).

5. Future work

5.1. General directions

The study of trace element concentrations in sedimentary archives through geological time is a relatively young field, and only a handful of bioessential trace elements have been investigated in the sedimentary record – IF, black shale, and pyrite (Figs. 3 and 12). Additionally, other sedimentary rock types that might serve as proxies for ancient ocean trace element concentrations—chert and carbonates—demand further study, including potential overprints during diagenesis and later burial. Chert and carbonate records have yet to be extensively explored, and are poised to provide further insights on paleomarine trace element abundances. When viewed through a genomic lens in terms of metalloenzyme evolution and diversification, the constraints imposed by geochemical modeling and the paleomarine proxy record should significantly improve our understanding of the links between evolving marine geochemistry, Earth surface redox conditions, and early biological evolution (Fig. 12).

5.2. Metals remaining to be investigated

Specific avenues for future work that would be of immediate interest include examining the sedimentary proxy records for temporal trends in bioessential elements that have yet to be fully explored. Two in particular, copper (Cu) and vanadium (V), stand out in this respect. Cu is used in ‘blue’ copper proteins for electron transfer, energy capture, and in other oxidative enzymes (e.g., Solomon et al., 1996). The emergence of Cu-metalloproteins is thought to have been quite late (Dupont et al., 2010). Similar to Zn, thermodynamic modeling indicates that the seawater Cu reservoir should have reached a low in Proterozoic oceans as the result of expanded water column euxinia (e.g., Frausto da Silva and Williams, 2001; Saito et al., 2003). However, a recent examination of authigenic Cu enrichments in BIF and black shales, normalized to detrital input using Ti, are remarkably static throughout the Precambrian (Chi-Fru et al., 2016). Despite the static abundance record, there is significant variation in black shale Cu isotope compositions over time that appears to be due to variations in Cu sinks and sources (Chi-Fru et al., 2016). The preferential sequestration of $^{65}$Cu by iron oxides (e.g., Ballistrieri et al., 2008) likely enriched seawater in residual $^{63}$Cu, which would have been incorporated into planktonic biomass and ultimately deposited into black shales depleted in $^{65}$Cu. After the GOE, oxidative continental weathering of sulfides should have increased the supply of dissolved Cu(II) and delivered more $^{65}$Cu-rich runoff to the oceans, while at the same time the isotopically light sink associated with iron oxides waned. This evolution towards heavy $^{63}$Cu values coincides with a shift to negative sedimentary $\delta^{56}$Fe values and increased marine sulfate after the GOE, and is traceable through Phanerzoic black shales to modern marine settings, where marine dissolved and sedimentary $\delta^{56}$Cu values are universally positive.
Zerkle et al. (2005) suggested that since the onset of widespread ocean oxygenation in the Neoproterozoic, the biological utilization of Cu enzymes doubled, likely as a result of increased Cu availability. In the modern ocean, Cu-metalloproteins are involved in the first step of nitrification, last step of denitrification, and during ammonia oxidation; the net result is a substantial bacterial Cu requirement (Amin et al., 2013). Low seawater Cu concentrations and bioavailability may also have had significant implications for the Precambrian atmosphere. In this regard, Buick (2007) proposed that in a Cu-limited ocean, denitrification would be incomplete since Cu is an essential component of the enzymes involved in that metabolism. The effect could have been a build-up of N₂O with significant climatic implications because N₂O is a...
potent greenhouse gas. The results of Chi-Fru et al. (2016) would seem to indicate that such a scenario was unlikely, considering the relatively static sedimentary Cu abundance record they present. Additional work is required to understand how Cu abundances and stable isotopes respond to changes in ocean and atmosphere redox—and how the biological history of copper utilization fits into the evolving redox landscape.

Vanadium is also an important bioessential trace element for the early nitrogen cycle. Both Anbar and Knoll (2002) and Zhang et al. (2014) suggested that the V-Fe nitrogenase varieties might have contributed to nitrogen fixation during the Paleoproterozoic, although the V variety is less efficient (Eady, 1996). Correspondingly, Zhang et al. (2014) indicated that low δ15N values in Archean cherts may be more consistent with alternative nitrogenases using Fe or V instead of Mo, suggesting a more important role for these variants in deep time—an interpretation challenged by the recent work of Stüeken et al. (2015a) (see Section 3.2). A preliminary trend for V in euxinic black shales through time was presented in Sahoo et al. (2012) (see their Fig. 3b). It was argued that seawater V concentrations were low in the mid-Proterozoic, and that V enrichments in the Doushantuo Formation (~635–630 Ma) were consistent with a well-oxygenated ocean. Overall, it appears that the black shale record for V mirrors that of Mo, and is relatively consistent with Neoproterozoic enrichments seen in the pyrite record (Large et al., 2014). However, the IF archives have not yet been investigated, and an extensive temporal examination of V in the context of its biological importance is lacking. Further, and perhaps most importantly, the cellular stoichiometry and specific requirements for V remain unclear.

Collectively, the temporal trends of Mo, V, and potentially Cu, suggest a scenario where N-fixing cyanobacteria in the early oceans may have been affected by biolimiting levels of multiple trace metals, especially prior to the GOE. As such, an understanding of Cu and V will be essential in order to obtain a complete picture of the N cycle prior to Neoproterozoic ocean oxygenation. The evolving picture for widespread nitrogen fixation on the early Earth is becoming more sophisticated and complex, and ultimately may have been influenced by the intricate interplay of several key trace metals in the early oceans.

Additional biologically relevant elements whose concentrations in sedimentary archives have not yet been investigated on billion-year timescales (Fig. 3) include cadmium (Cd) and tungsten (W). In the case of Cd, the black shale and sedimentary pyrite records may be of special interest because Cd-sulfides have recently been identified as an important sink from oxygen-deficient water columns (Janssen et al., 2014). Cadmium has been shown to stimulate Thalassiosira weissflogii under Zn limiting conditions because it is used in an alternative form of carbonic anhydrase, the enzyme that facilitates the interconversion of dissolved bicarbonate to CO2, and vice versa (Lane and Morel, 2000). Tungsten is used by primary anaerobic prokaryotes and mostly filled roles now played by Mo (Williams and Frausto da Silva, 2002; Pushie et al., 2014). Hence, W may also play a key role in the biogeochemical cycling of N prior to the Neoproterozoic. In modern bacteria and archaea, W is generally important for hyperthermophiles, suggesting that W bioavailability may have been critical for early life given the ancient origin of these organisms. In modern extant lineages, it may also fill biological roles by substituting for Mo (e.g., Frausto da Silva and Williams, 2001; L’vov et al., 2002).

Finally, explaining the continued utilization of Fe (and Mn) in biology despite decreasing seawater concentrations and bioavailability from the Archean to the Phanerozoic (Zerkle et al., 2005) is of great interest to biogeochemists. There are several pathways that have evolved to utilize alternative trace metals in place of Fe (see below). Dissolved Fe concentrations in the Archean and Paleoproterozoic oceans would have had to have been high in order to permit the formation of extensive IF via Fe(II) oxidation by anoxicogenic and oxyanoxic photosynthesis (Bekker et al., 2010). However, Anbar and Knoll (2002) suggested that declining seawater Fe concentrations after the Paleoproterozoic, triggered by the increased precipitation of Fe-oxides and sulfides (Canfield, 1998), may have suppressed the activity of nitrogenase (whose different versions all depend on Fe), and in turn, potentially limited Mesoproterozoic primary productivity via a dearth of fixed N.

High dissolved concentrations of Fe in Archean seawater may have also directly impacted the activity of cyanobacteria. Swanner et al. (2015) showed that for cyanobacteria cultured under high-Fe conditions, there is an increase in destructive reactive oxygen species (ROS) produced in the cell; indeed, they reported a two-fold increase in ROS at 100 µM Fe(II) and a five-fold increase at 1000 µM Fe(II). These ROS would have negatively affected the survival of early cyanobacteria, potentially delaying the proliferation of oxyangetic photosynthesis prior to the GOE. However, despite declining Fe concentrations following the final Paleoproterozoic episode of widespread IF deposition at 1.88 Ga (Rasmussen et al., 2012), Fe utilization in biological systems remains high. Several recent molecular innovations are employed by organisms in the modern environment to reduce Fe requirements—but always with a tradeoff. These tradeoffs include the replacement of a Fe-containing superoxide dismutase with a Ni-containing version (Dupont et al., 2012), exchanging an Fe protein with a Cu protein in the electron transport chains in photosynthesis (Peers and Price, 2006) and ammonia oxidation (Santoro et al., 2015), and using B2 instead of Fe for ribonucleotide reductase. In all cases, the exchange necessitates the acquisition of a different micronutrient. Declining Fe concentrations may also have stimulated the evolutionary development of organic ligand complexes, such as siderophores, that target elements such as Fe for facilitated acquisition (Hider, 1984; Neilands, 1989; Kendall et al., 2012).

6. Conclusions

There has been significant progress made in recent years in the examination of sedimentary proxy records (black shales, IF, pyrite) for the paleomarine availability of biologically critical trace elements, including P, Mo, Ni, Zn, Cr, and Co (Fig. 12). In several instances, fundamental differences between thermodynamic based solubility models and the sedimentary proxy record highlight the need for a multidisciplinary approach when evaluating paleomarine trace element abundances. As such, updated geochemical models should accompany future studies of the sedimentary proxy record in order to provide parsimonious explanations for paleomarine concentrations and resultant biological implications. It is critical in studies of the temporal trends of biocritical trace elements and organismal stoichiometries to consider both thermodynamic and kinetic controls on the composition of seawater. Collectively, studies on the proxy record thus far indicate that there is a complex and dynamic interplay between biological and geochemical processes that have not only affected the composition of Earth’s surface through time, but the evolution of life as well.

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