Cobalt and marine redox evolution

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1. Introduction

The availability of bio-essential trace elements such as Fe, Mo, Zn, Co, Ni, and Cu underpins the emergence, long-term evolution, and activity of life on our planet. The record of trace element utilization imprinted in modern organisms is commonly thought to reflect metal availability in seawater when key metalloproteins evolved (Dupont et al., 2006; Fraústo da Silva and Williams, 2001; Zerkle et al., 2005). The availability and removal of trace elements within aqueous habitats for life reflects the compositional evolution of the Earth's crust, but is also controlled by redox changes driven by metabolic innovation (Anbar, 2008). Temporal patterns in the concentrations and isotopic variations of trace elements in ancient sediments can serve as proxies for major changes in redox conditions in the oceans and atmosphere over geologic timescales (Anbar, 2008; Konhauser et al., 2009; Scott et al., 2008, 2012), and can provide a means to test the idea that environmental availability controlled the evolutionary history of metal utilization. However, empirical records of metal variation through time have...
Cobalt is a bio-essential metal for life, forming amongst others, the central cobalt-corrin complex of cobalamin (vitamin B₁₂). Eukaryotes use Co primarily as cobalamin in, for example, methionine synthesis. Bacteria and archaea additionally use cobalamin in enzymes for anaerobic metabolisms, including fermentation, dehalogenation, and one-carbon compound electron transfers (Banerjee et al., 2007). Genomes of karyotes use Co primarily as cobalamin in methionine synthesis. Bacteria and archaea additionally use cobalamin in enzymes for anaerobic metabolisms, including fermentation, dehalogenation, and one-carbon compound electron transfers (Banerjee et al., 2007). Genomic reconstructions support the early biological utilization of Co (David and Alm, 2011; Dupont et al., 2006), perhaps reflecting an ancient abundance of dissolved marine Co relative to the modern oceans. However, this model has not yet been tested against the geological record.

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Cobalt concentrations in modern seawater vary from 3 to 120 pM (Saito and Moffett, 2002), with variations dependent on interactions with other metals, biota and organic matter. Cobalt shows nutrient-like behavior with surface minimum concentrations due to biological uptake by phytoplankton (Saito et al., 2010; Saito and Moffett, 2002). Yet strong Co ligands complex nearly all of the dissolved Co in some oligotrophic waters (Noble et al., 2008; Saito et al., 2004; Saito and Moffett, 2001; Saito et al., 2005), although some portion of this complexed Co pool is likely bioavailable (Saito et al., 2004; Saito and Moffett, 2001). In coastal and deep waters, Co behaves as a scavenged-type element (Moffett and Ho, 1996; Saito et al., 2004), becoming oxidized and adsorbed to Mn(III,IV) oxides as they precipitate (Murray and Dillard, 1979). Co scavenging is thought to be catalyzed by Mn(II)-oxidizing bacteria (Moffett and Ho, 1996; Murray et al., 2007).

Previous estimates of marine Co concentrations through Earth’s history were based on thermodynamic considerations and assumptions regarding evolving marine redox and chemical composition (Saito et al., 2003), and did not consider how some key sources and sinks changed through time. Furthermore, equilibrium mineral precipitation models neglect kinetic control over precipitation. In this study, we use the sedimentary record of Co to track first-order changes in the marine Co reservoir through time, along with estimated magnitudes of the modern sources and sinks to infer the causes for Co reservoir change. We also suggest that the contents of Co of authigenic marine pyrite can be used as a proxy for marine Co concentrations.

2. Behavior of Co in marine environments

The formation of authigenic marine phases is subject to thermodynamic control, based on the abundance and speciation of ions, redox, and pH conditions. However, kinetically-driven scavenging reactions also influence the composition of marine precipitates. As a basis for the discussion below, Eh–pH diagrams detailing Co speciation were generated with the Act2 module of Geochemists’ Workbench using the Minteq thermodynamic database from 2005. Cobalt(II) was set to 100 nM, Fe(II) and Mn(II) to 50 μM, bicarbonate to 5 mM, silica to 2.2 mM (saturation with amorphous silica; Konhauser et al., 2007) in seawater ionic strength; physical constraints were 25 °C and 1 atmosphere ambient pressure. Cobalt does not form carbonate compounds, andCo(II) precipitates are only possible at very high Eh–pH conditions (Fig. 1). Cobalt(II) is oxidized to Co(III) in the same Eh–pH space where Mn(II) oxidation occurs, accounting for the oxidation of Co(II) and adsorption of Co(III) to precipitated Mn(III,IV) oxides (Murray and Dillard, 1979; Takahashi et al., 2007). Iron(II) is oxi-
Co-precipitation of Co with Fe (oxyhydr)oxides can be inferred based on a correlation coefficient of 0.763 from previously published data from hydrothermal sediments at the Endeavor Segment, Juan de Fuca Ridge (Hrishceva and Scott, 2007).

Adsorption of Co(II) and Co(III) to surface sites on Fe(III) (oxyhydr)oxides (Musić et al., 1979) and Mn(III,IV) oxides (Takahashi et al., 2007), respectively, is an important pathway for scavenging of Co under oxic conditions (Koschinsky and Hein, 2003; Stockdale et al., 2010; Takahashi et al., 2007). Soluble Co maxima occur below the O$_2$–H$_2$S chemocline in modern euxinic basins in conjunction with both the soluble Fe and Mn peaks (Dryssen and Kremling, 1990; Öztürk, 1995; Viollier et al., 1995), which indicates that Co is released by reductive dissolution of both Mn(III,IV) oxides and Fe(III) (oxyhydr)oxides. Both poorly crystalline and crystalline Fe(III) (oxyhydr)oxide surfaces efficiently scavenge Co(II) in waters with pH above ~7 (Gunnarsson et al., 2000; Musić et al., 1979; Dzombak and Morel, 1990), likely as bidentate inner-sphere complexes (Reak et al., 2011), while Co(III) is substituted for Mn in Mn(III/IV) oxides (Manceau et al., 1997). Correlation between Co and Fe in hydrothermal sediments collected from Endeavor Segment, Juan de Fuca (Fig. 2; data from Hrishceva and Scott, 2007) implies a common delivery path for both metals and is consistent with an Fe(III) (oxyhydr)oxide Co shuttle.

Sorption of Co(II) to ferrihydrite, likely the dominant Co scavenging pathway in Fe(II)-rich seawater (e.g. Konhauser et al., 2009), was investigated with MINTEQ using constants and site densities previously determined for ferrihydrite (Dzombak and Morel, 1990). Activities of dissolved components were corrected using the Davies equation, and modeling utilized 100 pM initial Co(II), 0.56M NaCl electrolyte to simulate seawater, and 1 g/L free ferrihydrite mineral (a constant concentration of surface sites during steady-state production of ferrihydrite). We compared Co sorption with and without Co ligands at modern (ca. 40 pM; Saito et al., 2004; Saito and Moffett, 2001) and high (400 pM) concentrations to investigate whether Co ligands prevent sorption of Co(II) to surface sites of ferrihydrite. The amount of Co(II) sorbed was unaffected at marine pH with 40 pM ligands, but Co(II) sorption to ferrihydrite was negligible when ligand concentrations were 400 pM (Fig. 3). We note that this model utilizes data for ligands specific to Co(II), but Co(III)-ligand complexes are likely extremely inert (Saito et al., 2005), and should further decrease the pool of Co available for metal sorption. We also did not examine the effects of inorganic species (e.g. Si), which can compete with metals for surface binding sites on ferrihydrite (Konhauser et al., 2007, 2009). As illustrated by those studies, the presence of Si should lower the amount of metal bound for any given dissolved Co concentration.

In oligotrophic surface waters, Co concentrations are controlled by phytoplankton uptake and binding to organic ligands (Saito and Moffett, 2001, 2002) rather than by the scavenging reactions with Mn that occur in coastal and deep waters (Moffett and Ho, 1996). Cobalt concentrations in phytoplankton are similar to abundances of Cd and Cu and enriched by as much as 10× over Mo (Ho et al., 2003). Organic material can also sorb trace elements and transfer them to sediments (Broecker and Peng, 1982; Krauskopf, 1956), although this process is not as quantitatively significant for Co as...
it is for other metals (e.g., Cd, Zn; Algeo and Maynard, 2004; Yee and Fein, 2003).

The formation of sulfide minerals governs the concentrations of Co in anoxic and sulfidic waters. Cobalt sulfide is more soluble than sulfides of some other biologically important elements (e.g., Cu, Zn; Saito et al., 2003), and dissolved Co concentrations below the chemocline of sulfidic waters can exceed average ocean concentrations by several orders of magnitude (Dryssen et al., 1990). Previous calculations suggested that the formation of Co-sulfide should scavenge dissolved Co from sulfidic waters (Dryssen and Kremling, 1990; Voilierr et al., 1995). The exchange of the bisulfide ion with water molecules hydrating dissolved Co(II) is slower than with those hydrating Fe(II), likely contributing to the persistence of dissolved Co(II) in sulfidic waters (Morse and Luther, 1999). For most sulfidic marine systems, this means that while FeS is more soluble than CoS from a thermodynamic point of view; e.g., Saito et al., 2003), FeS formation is kinetically favored, and Co is incorporated into FeS rather than precipitating as CoS (Huerta-Diaz and Morse, 1992; Morse and Arakaki, 1993).

The concentration of metals, including Co, in sulfidic waters is also buffered by the formation of soluble metal-sulfide complexes (Daskalakis and Helz, 1992), although this seems to be more important for metals such as Cd, Zn, and Cu that form stronger sulfide complexes. However, above 1 μM total sulfide, soluble Co-sulfide complexes will reduce the dissolved Co(II) pool (Saito et al., 2003). Regardless of the initial phase of aqueous precipitate, Co ultimately substitutes into pyrite during sedimentary diagenesis (Huerta-Diaz and Morse, 1992; Stockdale et al., 2010).

3. Marine Co sources and sinks

The concentration of elements in seawater reflects a balance between delivery and removal of elements via precipitation and adsorption processes (e.g., Broecker, 1971; Krauskopf, 1956). To quantitatively interpret changes in the marine Co reservoir, as recorded by sediments deposited under different redox conditions, we detail below estimates of the fluxes of dissolved Co to seawater, and mass accumulation rates (MAR) for sediments deposited under oxic, anoxic and euxinic conditions (Table 1).

3.1. Marine Co sources

The amount of Co present in the crust ultimately governs the amount of Co delivered to the oceans through fluid–rock interaction. Minerals with higher Fe, Mg, and Cr contents are also enriched in Co (Carr and Turekian, 1961), specifically olivine and pyroxene present in ultramafic rocks (Glassley and Piper, 1978). Dissolved Co is delivered to seawater via rivers and hydrothermal fluids that source Co predominantly from mafic and ultramafic rocks. The estimated riverine flux of Co is 5.5 × 10^{12} g kyr^{-1} (Table 1; Gaillardet et al., 2003). Cobalt fluxes from the continents were likely higher prior to 2.5 Ga due to their more mafic compositions, after which average Co concentrations in the continental crust dropped from 22 to 15 ppm (Condie, 1993). Dust can add dissolved Co to surface waters, but this process is likely to be minor and geographically and seasonally restricted (Shelley et al., 2012).

Although recent work has highlighted the role of hydrothermal fluids in supplying scavenged-type elements to the ocean reservoir (e.g., Fe; Saito et al., 2013; Tagliabue et al., 2010), initial measurements have found little evidence for Co fluxes from hydrothermal systems to seawater (Noble et al., 2012, 2008). However, hydrothermal fluids contain Co in concentrations often several orders of magnitude above average seawater (Metz and Trefry, 2000), implying that delivery of hydrothermal Co to open oceans is limited by efficient, near-field scavenging reactions in oxic seawater (German et al., 1991). Using the fluid flow volume through high- and low-temperature hydrothermal systems as estimated from the oceanic Mg budget by Elderfield and Schultz (1996), we calculated Co fluxes out of both types of systems as described by Reinhard et al. (2013). The Co anomaly for high-temperature systems utilized the difference between concentrations within the Plume vent on the Juan de Fuca Ridge (200 nmol kg^{-1} at 246°C) and bottom waters (0.02 nmol kg^{-1}; Metz and Trefry, 2000). The Co anomaly for low-temperature systems was based on Co data from site 1027 on the Juan de Fuca Ridge (0.7 nmol kg^{-1} at 64°C) relative to bottom waters (0.03 nmol kg^{-1}; Wheat et al., 2003). We estimate a total hydrothermal Co flux of 1.2 × 10^{11} g Co kyr^{-1}; 2.4% of the total Co flux (Table 1). We recognize that the net fluxes of most hydrothermally derived metals to the open ocean remain poorly constrained; metals released from hydrothermal vents are readily incorporated into sulfide or oxide precipitates within plumes, thereby diminishing dispersion. However, nanoparticulate sulfides or organic ligands may stabilize and transport trace elements away from vents (Sander and Koschinsky, 2011; Toner et al., 2009; Yee et al., 2011), increasing trace element, and probably Co, fluxes to the global ocean. We therefore anticipate that global fluxes of Co from hydrothermal systems will be refined in the near future (e.g. GEOTRACES).

3.2. Oxic sedimentary Co sinks

In sediments where oxygen penetrates at least 1 cm, authigenic Fe(III) (oxyhydr)oxides, Mn(III/IV) oxides and associated Co are permanently buried (Brumsack, 1989; Froelich et al., 1979). Any Fe(II) and Mn(II) released during dissimilatory microbial reduction is likely to be reoxidized and immobilized before diffusing out of sediments. Deposition of phytoplankton biomass may also add Co to oxic sediments (Saito et al., 2004). Regardless of the pathway for authigenic Co delivery to sediments, Co released during early diagenesis is immobilized by scavenging with Fe(III) (oxyhydr)oxides and Mn(III/IV) oxides. Cobalt fluxes to oxic continental margin and hydrothermal sediments are higher than those in the deep sea (Douglas and Adeney, 2000; Koschinsky and Hein, 2003), but these fluxes are not well-constrained. We use an estimate of 2.3–5 μg Co cm^{-2} kyr^{-1} (Krishnaswami, 1976) from deep-sea pelagic sediments for an average authigenic oxic Co MAR.

3.3. Euxinic sedimentary Co sinks

Modern sulfidic (sulfide in porewaters) or euxinic (sulfide in bottom waters) environments include basins where authigenic Fe(III) (oxyhydr)oxides and Mn(III/IV) oxides are reductively dissolved below the O2–H2S transition zone, releasing associated Co
In anoxic marine sediments lacking dissolved sulfate, Fe(III) (oxyhydr)oxides and Mn(III/IV) oxides are subject to dissimilatory microbial reduction. Fluxes of Co(I) and Mn(II) have been observed out of sediments underlying oxygen- and sulfide-poor bottom water (Brumsack, 1989; Saito et al., 2004), and plumes of dissolved Co mobilized from anoxic sediments have been observed over oxygen minimum zones (OMZs; Noble et al., 2012), demonstrating the mobility of Co under anoxic conditions. While remobilization under anoxic conditions is an important source of Co in some coastal environments, this process does not represent an exogenous supply of Co, and so we have not included reducing sediments as a source in flux estimates. Because Co is readily mobilized from anoxic sediments, there is no significant flux of Co to anoxic sediments, and Co in anoxic sediments solely reflects what is added with detrital minerals (Brumsack, 1989; van der Weijden et al., 2006). Thus, a decrease in the extent of anoxic sediments at the expense of oxic or euxinic sediments will decrease the size of dissolved Co reservoir, while expansion of anoxic sediments should increase the size of the reservoir.

Generally speaking, sediments deposited under an anoxic but non-sulfidic water column, which are often referred to as ferruginous sediments, are also unlikely to permanently remove Co. Rare in the modern but common in the Precambrian, ferruginous sediments are characterized by overlying water column with anoxic conditions with Fe(II) as the main redox buffer (Planavsky et al., 2011). These redox conditions also allowed for the deposition of iron formations (IF), which are likely to be a significant Co sink (Fig. 4). However, IFs reflect stabilization of Fe(III) (oxyhydr)oxides under anoxic conditions where there was anomalously high local Fe(III) fluxes. Furthermore, most oxide facies IF are found in deep water settings, and oxides are preserved due to a lack of sufficient organic carbon to drive complete Fe(III) reduction (Konhauser et al., 2005). Therefore, IF likely have higher capacities for Co burial (with authigenic Fe(III) (oxyhydr)oxides) than modern anoxic sediments, but IFs represent a small fraction of overall marine ferruginous settings.

4. Archives of marine Co concentrations

We propose that sedimentary Co concentrations can serve as an archive of marine Co concentrations. We use this premise to evaluate how key sources and sinks for Co have varied through time. This compilation also informs the evolution of marine redox conditions. We focus on two types of sedimentary archives: those in which Co was sequestered by authigenic Fe(II) (oxyhydr)oxides (IF database), or by iron sulfides (sedimentary pyrite and euxinic shale databases).

4.1. Iron formations as a Co archive

Iron(III) (oxyhydr)oxides, generally thought to be the precursor phases to most IF, have extremely large and reactive surfaces that extensively adsorb cations, including Co(II), at marine pH (Dzombak and Morel, 1990). Therefore, we propose that the record of authigenic Co enrichment in IF can be used to evaluate large-scale changes in marine Co concentrations through time. Similar chemical principles justified using Ni, P, and Zn enrichments in IF to track changes in seawater concentrations of these elements (Konhauser et al., 2009; Planavsky et al., 2010; Robbins et al., 2013). Precambrian IF are chemical precipitates with minor detrital input. Authigenic, but diagnostically altered, Fe-rich (hematite, magnetite, siderite) and Si-rich (quartz) phases dominate the mineralogy of IF, and so their trace element composition is often inferred to reflect input of these elements from authigenic vs. detrital phases. Post-depositional alteration of the primary trace element seawater signatures is generally minimal unless IF have experienced hydrothermal alteration or near surface weathering (Bau and Moeller, 1993).
Historically, Precambrian IF are divided into Algoma-type, which have a spatially limited extent and formed in proximity to volcanic and hydrothermal settings, and Superior-type, which are more extensive and where deposited under marine conditions on a continental shelf or an isolated basin. Both deposits are represented in our dataset and we assign each IF as being Algoma- or Superior-type for simplicity; in reality there is a gradation between these IF types (see Bekker et al., 2010). Large, basin-scale IF deposition experienced a hiatus in the Middle Proterozoic, and a return to IF deposition in the Neoproterozoic was followed by permanent cessation of IF deposition. The lack of IF in intervening intervals is evident in our compilation. We further utilize exhalite deposits and oolitic ironstones to extend the record of Co burial with authigenic Fe oxide facies into the Phanerozoic.

We utilize an expanded dataset of Konhauser et al. (2011, 2009), comprising published values as well as new data acquired by bulk analysis and in situ LA-ICP-MS. Supplementary Table 1 reports Co, Al, Ti, Fe, Mn, and S concentrations (where available) for the 1353 Co data points used in this study. The supplementary information also includes references for the published data and the descriptions of iron formations analyzed in this study.

### 4.2. Pyrite as a Co archive

The partitioning of Co into iron sulfide phases is dependent on Co and Fe concentrations, but importantly for our purposes, it appears to be largely independent of the amount of hydrogen sulfide (Morse and Arakaki, 1993). In most sulfidic environments, dissolved Fe$^{2+}$ concentrations are near levels predicted from equilibrium with the amorphous Fe–S phase mackinawite (e.g. Helz et al., 2011). Finally, silicate phases react with sulfide 108$\times$ more slowly than Fe(III) (oxyhydr)oxide minerals (Canfield et al., 1992), so Co in detrital minerals should not contribute to Co concentrations in pyrite. Therefore, it is reasonable to assume that the degree of authigenic Co enrichment in sediments deposited below euxinic waters should, to a first order, reflect the dissolved Co concentrations in the water column.

In order to use trace element concentrations within pyrite as proxies for seawater concentrations during sediment deposition, it is essential that (1) pyrites formed within the sediments that host them, and (2) the metal inventory of pyrite was not overprinted by secondary alteration processes. In the absence of free oxygen in the Archean atmosphere, detrital pyrite was delivered to marine sediments (Rasmussen and Buick, 1999). Therefore, only sulfides with early diagenetic textures (e.g. nodules or disseminated grains) from shales with abundant organic carbon and sulfur and were included in this study (Rouxel et al., 2005), although not all of the sediments included in this study have been definitively demonstrated to have been deposited under euxinic conditions. Post-depositional disturbances to metal content were screened by selecting samples with no obvious sign of alteration (Rouxel et al., 2005, 2006).

Individual pyrite grains from black shales were digested and trace element data were acquired by Thermo Element2 HR-ICP-MS at Woods Hole Oceanographic Institution as described by Rouxel et al. (2005). A basic description of each sample, as well as the geological setting and age constrains for the host rock, are reported by Rouxel et al. (2005). Previously unreported samples include Devonian-age black shales from the Illinois basin, pyrite nodules from the.ca. 1.8 Ga Gunflint Formation of Kakabeka Falls in Ontario, Canada, and pyrite from the ca. 2.7 Ga Manjeri Formation of the Belingwe Belt, Zimbabwe. Cobalt concentrations are reported in Supplementary Table 2. Trace element compositions of pyrite from modern anoxic sediments are taken from Huerta-Diaz and Morse (1992) and reference cited therein.

### Table 2

<table>
<thead>
<tr>
<th>Age bin</th>
<th>Mean Co/Ti $(\mu g g^{-1})$ ± SD</th>
<th>p-value*</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\geq 2.80$ Ga</td>
<td>79.85 ± 1.72</td>
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<tr>
<td>2.75 to 1.88 Ga</td>
<td>150.57 ± 2.67</td>
<td>$&lt;0.0001$</td>
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<tr>
<td>$&lt; 1.72$ Ga</td>
<td>62.25 ± 1.84</td>
<td>$&lt;0.0001$</td>
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### 4.3. Bulk euxinic shales as a Co archive

Cobalt is enriched in sediments deposited under euxinic conditions (Algeo and Maynard, 2004), and the shales used in this study have Mo contents $> 25$ ppm, consistent with euxinia (Scott and Lyons, 2012). Authigenic Co was distinguished from detrital Co via normalization to a conservative element, in this case Al. This approach is justified as the Co/Al ratio of fine-grained material derived from continents has varied little through time (Condie, 1993), and is similar amongst several compilations (Condie, 1993; Kamber et al., 2005; Wedephol, 1971). The Co and Al values for euxinic sediments used in this study have all been previously published (see Supplementary references). Because it is not always straightforward to distinguish detrital from authigenic trace element signatures in shales (e.g. Van der Weijden, 2002), we emphasize that the IF and pyrite datasets are likely to be the most robust indicators of the marine reservoir.

### 4.4. Statistical analysis of Co datasets

When grouped by ages that correspond to global events (described below), Co concentrations were log-normally distributed. The statistical differences between average time-binned values were compared using an unpaired t-test of log concentration values. The $p$ values for these comparisons are reported in Table 2. The mean and one standard deviation (SD) of log concentration values were then back-transformed to concentration values, and these are reported as mean $(\mu g g^{-1})$ ± SD in accordance with the multiplicative nature of a log-normal distribution (Limpert et al., 2001).

### 5. Evolution of the marine Co reservoir

#### 5.1. Reconstructing changes in Co sources and sinks through Earth history

The datasets of Co in IF, pyrite and shale reveal time-resolved patterns in the delivery of Co to marine sediments, and thereby indicate first-order changes to the marine Co reservoir through time (Fig. 4). The average Co/Ti $(\mu g g^{-1})$ of IF $> 2.80$ Ga (79.85 ± 1.72) are significantly lower than IF deposited between 2.75 to 1.88 Ga (150.57 ± 2.67). Average values then significantly drop in IF, exhalite, and oolitic ironstones deposited between 1.72 Ga and modern times (62.25 ± 1.84; Table 2), despite that exhalites are prime
records of locally derived hydrothermal Co. There is also a significant difference between the average concentrations of Co (ppm) in pyrites from shales deposited between 2.8 and 1.84 Ga (7.34 - 1.77) and those deposited from 1.8 to 0.3 Ga (3.87 - 1.96; Table 2), signifying a concordance of the IF and shale pyrite Co records, and likely global-scale trends in marine Co concentrations.

Higher seawater Co concentrations in the interval between 2.8 and 1.84 Ga reflect pervasive anoxia and a higher hydrothermal Co flux to the marine reservoir during the emplacement of oceanic crust, likely due in part to several mantle plume events at this time (Barley et al., 2005; Rasmussen et al., 2012). These events also likely supplied the Fe for major IF deposited from 2.5 to 2.4 Ga, and again between 1.9 and 1.8 Ga (Barley et al., 1997; Rasmussen et al., 2012). In hydrothermal systems, the supply of Co tracks that of Fe (Douville et al., 2002) because the solubility of both elements is enhanced at higher temperature and Cl concentrations (Metz and Trefry, 2000). Archean-aged hydrothermal systems are thought to have experienced higher heat flow, enhancing the supply of Fe (Isley, 1995), and likely Co to seawater. Higher Co concentrations from Archean hydrothermal fluids are also more likely due to the prevalence of ultramafic oceanic crust (Arndt, 1983). Compositional control on Co concentrations in hydrothermal fluids is indicated in the ultramafic Rainbow vent field on the Mid-Atlantic Ridge, where Co concentrations reach up to 13 μM Co (Douville et al., 2002), an enrichment of at least 10^5 above seawater concentrations. Persistent anoxia would have allowed the dispersion of dissolved Co plumes without trapping near source by oxidative scavenging (Noble et al., 2012). These factors indicate an increased proportion of hydrothermal Co fluxes relative to continental Co fluxes in comparison to modern Co inputs (Table 1).

The Co concentrations in younger than 1.84 Ga IF are comparable to those prior to 2.8 Ga (Table 2), potentially indicating that weathering of mafic to ultramafic Neoarchean continental crust is not an essential aspect of the large enrichments. This is in contrast to Ni, whose supply to oceans from weathering of emergent oceanic plateaus waned after 2.7–2.6 Ga (Kamber, 2010; Konhauser et al., 2009). Although Co and Ni have generally similar low-temperature geochemical behavior, they are decoupled during high-temperature hydrothermal alteration. Nickel is not as efficiently leached as Co, resulting in low hydrothermal Ni fluxes (Douville et al., 2002). Furthermore, Co-chloride complexes are more stable than Cu- and, probably, Ni-chloride complexes, resulting in higher Co solubility at low temperatures (Metz and Trefry, 2000). The increase in marine sediment Co concentrations after 2.8 Ga may be driven by changes in the riverine flux of Co from the continents due to increased, permanent, subaerial exposure. Other authors have suggested the emergence of continents between 2.9 and 2.7 Ga (e.g. Pons et al., 2013 and references within). However, the crustal growth rate slows and Co concentrations drop at 2.5 Ga, from 22–25 ppm to 15 ppm (Condie, 1993), in the midst of the highest sedimentary Co concentrations (Figs. 4 and 5).

We propose that the drop in Co concentrations in marine sediments that occurs after 1.84 Ga reflects a decrease in intense, plume-related hydrothermal activity (Rasmussen et al., 2012). As supply of hydrothermal Fe and deposition of massive IF waned in the late Paleoproterozoic, so too did the hydrothermal Co flux, shifting toward modern conditions in which continentally-derived Co dominates Co influx (Table 1). Hydrothermal supply to the oceans reached modern levels by 0.7–0.8 Ga (Derry and Jacobsen, 1988), with episodic larger mantle inputs re-occurring throughout the Proterozoic and Phanerozoic (Peng et al., 2011; Veizer et al., 1983). Additional Middle Proterozoic samples would be needed to test whether later mantle plume events resulted in a return to globally high marine Co concentrations.

The drop in seawater Co concentrations after 1.88 Ga occurs at a time when the extent of euxinic environments in the oceans increased (Poulton et al., 2004) at the expense of ferruginous sediments. Without well-constrained depositional rates for IF, it is impossible to estimate Co MAR from ferruginous settings, although our IF dataset attests to the fact that Co is effectively buried under ferruginous conditions. Nevertheless, given that IFs are rare marine sediments, the contribution of ferruginous sediments to overall Co removal in the oceans was likely low. Further, as stressed above, the Co sink associated with ferruginous settings will be lower than foroxic sediments; although Fe(III) (oxyhydr)oxides sorb Co, Mn(III/IV) oxides, which form at higher Eh, are much more effective at scavenging Co from seawater (Stockdale et al., 2010). Therefore, expansion of euxinic settings to less than 10% of seafloor area in the Middle Proterozoic (Reinhard et al., 2013), is unlikely to explain the drop in the Co reservoir size (cf. Saito et al., 2003) without invoking a waning hydrothermal Co source. Significant areal expansion of oxic and euxinic sediments might explain the drop in the Co reservoir, but oxic conditions suitable for Co scavenging were likely confined to Middle Proterozoic surface waters (Poulton et al., 2004), and sediments deposited under oxic conditions were of limited extent (Reinhard et al., 2013).
The logarithmic range of Co concentrations in sediments, even those from the same formation, is a phenomenon that has been observed for other trace elements (e.g. Zn; Robbins et al., 2013; Scott et al., 2012). This variability is likely primary, based on similar phenomena in modern authigenic sediments (Figs. 2 and 4). For Co, this may reflect temporal variability in marine Co concentrations, which is expected because Co is a non-conservative element within the oceans and has an extremely short residence time in seawater, 280 yr by our estimate (Table 1; 40–120 yr; Saito and Moffett, 2002). Due to its short residence time, marine Co concentrations respond quickly to perturbations in sources or sinks, such as the development of OMZ that fluctuate on decadal time scales (Noble et al., 2012; Stramma et al., 2008). The highest IF Co/Ti (nearly 160,000 μg g−1) and pyrite Co (nearly 2000 ppm) are found in the 2.32 Ga Timeball Hill Formation and the underlying Rooihoogte formations. Post-depositional hydrothermal overprint is likely not responsible for elevated Co concentrations because they do not correspond to high Pb or Cu, concentrations (data not shown), which would suggest mineralization. Some of this variation may be primary, and representative of temporally or spatially variable Co concentrations within the basin. However, some variability could be tied to re-distribution between Fe-phases during diagenesis. Despite these variations, the robust statistical differences in time-binned averages (Table 2) validate that shifts in sedimentary Co map onto global events, and hence, indicate reservoir changes.

Although there are no significant changes in the average Co concentrations from Middle Proterozoic to Phanerozoic in any of the sedimentary records (Table 2), there is significant variability in Phanerozoic euxinic shales (Fig. 5), many of which were deposited during ocean anoxic events (OAE), which reflect transient rather than pervasive anoxic conditions. The average Co/Al ratios from euxinic black shales deposited during the Cretaceous OAE-2A at Demerara Rise are ~2× larger than euxinic sediments deposited before or after (their Fig. 9; Hetzel et al., 2009). This is dramatically different from other redox-sensitive metals such as Mo and V, whose restricted supply was exhausted during the OAE, leading to depletions of Mo and V during the peak of the OAE relative to sediments deposited before and after. Cobalt is mobilized from anoxic sediments (Noble et al., 2012), and therefore increased Co burial in euxinic sediments might reflect this greater reservoir during anoxic events, and in fact be a proxy for low-oxygen conditions (Saito et al., 2010). OAE Co enrichments may also be linked to, or augmented by a hydrothermal Co pulse (e.g. Brumsack, 2006). Importantly, there is independent evidence for increased hydrothermal input from a sharp shift toward less radiogenic (hydrothermally-derived) initial O isotope values during the onset of the OAE (Turgeon and Creaser, 2008). Furthermore, other authors have also documented sedimentary Co increases during OAE intervals linked to hydrothermal activity (Orth et al., 1993; Snow et al., 2005). Therefore, the OAE Co records are consistent with the notions developed above that anoxia and hydrothermal activity play a critical role in the global marine Co cycle, and that marine Co concentrations respond on shorter timescales than conservative elements such as Mo.

Surprisingly, we find little evidence for a change in the marine Co reservoir with the deep ocean oxygenation, potentially beginning as early as 635 Ma (Sahoo et al., 2012). The average Co/Al (μg g−1) for Proterozoic euxinic shales (11.51 – 1.73) is not significantly different from Phanerozoic-aged shales (9.33 – 1.97), although we emphasize that Phanerozoic Co is dominated by OAE samples, and likely reflect transient anoxia. There is also no statistical support for changes in IF or shale pyrite Co concentrations during similar time intervals (data not shown). We note however, that a paucity of datapoints from the Middle Proterozoic for all three databases hinders interpretation of changes to the Co reservoir with deep ocean oxygenation. The highest Phanerozoic Co concentrations occur in exhalites, which demonstrate that near-source scavenging reactions prevent hydrothermal Co dispersal under anoxic conditions, and therefore Phanerozoic exhalites likely reflect only local Co concentrations. Non-OAE euxinic shale pyrites would likely best track changes in the Co reservoir associated with ocean oxygenation. Because expanding oxic sediments at the expense of anoxic should enhance Co removal, and Neoproterozoic hydrothermal activity is likely similar to modern (Derry and Jacobsen, 1988), ocean ventilation should have further decreased the Co reservoir size.

5.2. Marine cobalt reservoir size and biological evolution

Our empirical record of the marine Co reservoir affords an opportunity to discuss hypotheses of Co utilization in biology, and to consider how Co-utilization in aquatic organisms fit to these hypotheses. The biological utilization of metalloproteins may reflect the availability of elements in the environment in which the protein first appeared (e.g. Nisbet and Fowler, 1996). Genomic analyses of all three domains of life indicate that Co-binding proteins originated after 3.3 Ga (David and Alm, 2011). Given the uncertainties in this age stated by the authors (>250 My), it is feasible that origin of many Co-binding proteins coincides with the enhanced marine Co availability we observe after 2.8 Ga, generally supporting the availability hypothesis. However, Co is chemically suited to catalyze reactions involving hydrogen rather than oxygen (Fraisto da Silva and Williams, 2001), and any preferential Co utilization by early organisms may simply reflect the abundance of reduced energy sources such as methane, carbon monoxide, and hydrogen on the early Earth (Zerkle et al., 2005). Because higher Co availability is linked to periods in Earth's history when the oceans were anoxic, had on average lower Co burial rates, higher Co mobility, and probably longer Co residence times, environmental availability and catalytic suitability may both be reflected in the utilization of Co in organisms or early-evolved proteins.

Consistent with a greater utilization of Co by early organisms, bacteria and archaea have a greater number of genes that encode for Co-binding proteins (Zhang and Gladyshev, 2010), and their genomes encode a larger proportion of Co-binding proteins than eukarya (Dupont et al., 2006). In contrast, a lack of encoded Co-binding proteins in eukarya should indicate evolution predominate- nitely after 1.8 Ga. This is broadly consistent with initial diversification of eukaryotes in the middle to late Proterozoic (Knoll et al., 2007). The persistence of the cobalamin-requiring gene metH in eukaryotic phytoplankton, involved in biosynthesis of the essential amino acid methionine, suggests that lower marine Co concentrations were not a sufficient selection pressure to drive loss of the metH gene in favor of a cobalamin-independent but less efficient gene (Bertrand et al., 2013), or that key Co-binding proteins were maintained and few new Co-binding proteins were acquired during genome expansion. Alternately, the persistence of metH in eukaryotic phytoplankton may reflect a later acquisition (Croft et al., 2005), and it often occurs in eukaryotic phytoplankton that already contained the cobalamin-independent gene (Helliwell et al., 2011). Thus, eukaryotic metH persistence may be related to its efficiency rather than Co availability.

Enzymes that directly bind Co rather than cobalamin also offer insight into how metal availability regulates enzyme utilization. Both diatoms and eukaryotic algae can substitute Co (or Cd) for Zn in the carboxy anhydrase enzyme that interconverts CO2 and bicarbonate when Zn concentrations are limiting (Morel et al., 1994; Saito and Goepfert, 2008). Such substitution may be a strategy for coastal algae to deal with intense metal drawdown during algal blooms (Saito and Goepfert, 2008), but also indicates that expression of metal-binding proteins encoded at the genomic level are
affected by temporally and spatially variable metal concentrations. The trends in Co sedimentary records presented here reflect geological control over element delivery and burial; short-term and spatial variability will therefore not be resolvable with our dataset. Furthermore, the amount of metal-binding protein expressed (i.e. the metalloenzyme) should fluctuate with physical and chemical conditions and the physiological state of the cell (Bertrand et al., 2013; Saito et al., 2011). Proteomics is therefore an important complement to genomic investigation of why utilization of metal-binding proteins persist, are lost or acquired as metal concentrations change.

Finally, although genomic utilization may reflect availability of metals during evolution, it can also indicate later adaptations to changing metal abundances. For instance marine cyanobacteria have an absolute requirement for Co that cannot be met by other metals (Saito et al., 2002; Sunda and Huntsman, 1995), which may reflect evolution in largely anoxic or sulfidic oceans with greater Co availability (Saito et al., 2003), yet thrive in theoxic oceans where Co concentrations are often less than 40 pM. Cobalt utilization may have persisted in cyanobacteria as Co availability declined due to the acquisition of Co-binding ligands (Saito and Moffett, 2001; Saito et al., 2005). The hypothesis that availability dictates utilization implies that the earliest life lacked strategies for acquisition of metals (Nisbet and Fowler, 1996). However, it is clear that modern organisms possess strategies to deal with limitation. Thus, the evolutionary history of metal-acquisition genes may be an important consideration when comparing metal availability to biological utilization, and many of these proteins are still being identified (Zhang and Gladyshev, 2010).

6. Conclusions

Trace elements are proxies for tracking marine redox evolution, but most studies have focused on metals that partition strongly into anoxic or sulfidic sediments (e.g., Mo). For Co, and Fe and Mn, which have short residence times in the modern ocean, marine concentrations respond dynamically to changes in delivery and removal. High seawater Co concentrations from ~2.8–1.84 Ga, recorded by IF and authigenic pyrites, resulted from widespread anoxia and enhanced hydrothermal activity and are probably linked to Fe fluxes that resulted in the deposition of IF. The marine Co reservoir decreased after ~1.84 Ga due to waning hydrothermal Co delivery. The expansion of euxinic sediments at the expense of anoxic sediments, which are a negligible sink for Co, may have also contributed to the Middle Proterozoic Co reservoir decrease. Variability of Phanerozoic IF and euxinic shale Co concentrations are linked to localized hydrothermal activity and/or transient anoxic conditions, and as such obscure any global changes in the marine Co reservoir associated with deep-water oxidation. Our study of the concentrations of Co in marine sediments through time reveals a more nuanced view of the marine Co reservoir through Earth’s history than is possible with theoretical models based on thermodynamic equilibrium (e.g. Saito et al., 2003). This emerging view of the evolution of the marine Co reservoir through time provides a framework for interpreting Co availability influenced the acquisition and utilization of Co in biology.

Acknowledgements

EDS acknowledges support of the National Science Foundation (NSF) IRFP and a Deutsche Forschungsgemeinschaft grant (KA 1736/24-1). NJP acknowledges support of the NSF EAR-PF and NSF ELT program. SVL gratefully acknowledges postdoctoral fellowship support from NSERC and LabExMER. Discovery Grants and CGS-M from NSERC supported KOK, AB, and LJR, respectively. OR received support from Europero Mer and IFREMER. MAS was supported by the Gordon and Betty Moore Foundation and the NSF Chemical Oceanography program. SJM was supported by the NASA Exobiology Program, NASA’s Astrobiology Institute Fund for International Cooperation, the University of Colorado Center for Astrobiology, J.W. Fulbright Foundation, University of Colorado’s Office of the Vice Chancellor for Research, and a sabbatical stay at the Centre de Recherches Pétrographiques et Géochimiques (CRPG-Nancy). This manuscript benefited from helpful discussions with Chris Reinhard and Martin Wille.

Appendix A. Supplementary material

Supplementary material related to this article can be found online at http://dx.doi.org/10.1016/j.epsl.2014.01.001.

References


