Proterozoic ocean redox and biogeochemical stasis

Christopher T. Reinhard^{a,b,1}, Noah J. Planavsky^{a,b}, Leslie J. Robbins^c, Camille A. Partin^d, Benjamin C. Gill^e, Stefan V. Lalonde^f, Andrey Bekker^d, Kurt O. Konhauser^c, and Timothy W. Lyons^b

^aDivision of Geological and Planetary Sciences, California Institute of Technology, Pasadena, CA 91125; ^bDepartment of Earth Sciences, University of California, Riverside, CA 92521; ^cDepartment of Earth and Atmospheric Sciences, University of Alberta, Edmonton, AB, Canada T6G 2E3; ^dDepartment of Geological Sciences, University of Manitoba, Winnipeg, MB, Canada R3T 2N2; ^eDepartment of Geosciences, Virginia Institute of Technology, Blacksburg, VA 24061; and ^fUMR6539 Domaines Océaniques, Institut Universitaire Européen de la Mer, Université de Bretagne Occidentale, 29280 Plouzane, France

Edited by Mark H. Thiemens, University of California at San Diego, La Jolla, CA, and approved February 15, 2013 (received for review May 22, 2012)

The partial pressure of oxygen in Earth's atmosphere has increased dramatically through time, and this increase is thought to have occurred in two rapid steps at both ends of the Proterozoic Eon (~2.5-0.543 Ga). However, the trajectory and mechanisms of Earth's oxygenation are still poorly constrained, and little is known regarding attendant changes in ocean ventilation and seafloor redox. We have a particularly poor understanding of ocean chemistry during the mid-Proterozoic (~1.8-0.8 Ga). Given the coupling between redoxsensitive trace element cycles and planktonic productivity, various models for mid-Proterozoic ocean chemistry imply different effects on the biogeochemical cycling of major and trace nutrients, with potential ecological constraints on emerging eukaryotic life. Here, we exploit the differing redox behavior of molybdenum and chromium to provide constraints on seafloor redox evolution by coupling a large database of sedimentary metal enrichments to a mass balance model that includes spatially variant metal burial rates. We find that the metal enrichment record implies a Proterozoic deep ocean characterized by pervasive anoxia relative to the Phanerozoic (at least ~30–40% of modern seafloor area) but a relatively small extent of euxinic (anoxic and sulfidic) seafloor (less than ~1-10% of modern seafloor area). Our model suggests that the oceanic Mo reservoir is extremely sensitive to perturbations in the extent of sulfidic seafloor and that the record of Mo and chromium enrichments through time is consistent with the possibility of a Mo-N colimited marine biosphere during many periods of Earth's history.

paleoceanography | geobiology

he chemical composition of the oceans has changed dramatically with the oxidation of Earth's surface (1), and this process has profoundly influenced and been influenced by the evolutionary and ecological history of life (2). The early Earth was characterized by a reducing ocean-atmosphere system, whereas the Phanerozoic Eon (< 0.543 Ga) is known for a stably oxygenated biosphere conducive to the radiation of large, metabolically demanding animal body plans and the development of complex ecosystems (3). Although a rise in atmospheric O_2 is constrained to have occurred near the Archean-Proterozoic boundary (~2.4 Ga), the redox characteristics of surface environments during Earth's middle age (1.8-0.543 Ga) are less well known. The ocean was historically envisaged to have become ventilated at ~1.8 Ga based on the disappearance of economic iron deposits (banded iron formations; 2), but over the past decade it has been commonly assumed that the mid-Proterozoic Earth was home to a globally euxinic ocean, a model derived from theory (4) and supported by evidence for at least local sulfidic conditions in Proterozoic marine systems (5-8). However, the record of sedimentary molybdenum (Mo) enrichment through Earth's history has been interpreted to suggest that euxinia on a global scale was unlikely (9).

More recently, it has been proposed that the deep ocean remained anoxic until the close of the Proterozoic, but that euxinia was limited to marginal settings with high organic matter loading (10–13). In anoxic settings with low dissolved sulfide levels, ferrous iron will accumulate—thus these anoxic but nonsulfidic settings have been termed "ferruginous" (10). This model has also found

support in at least local evidence for ferruginous marine conditions during the mid-Proterozoic (12, 13). However, it has been notoriously difficult to estimate the extent of this redox state on a global scale, even in the much more recent ocean—largely because most of the ancient deep seafloor has been subducted.

In principle, trace metal enrichments in anoxic shales can record information about seafloor redox on a global scale. Following the establishment of pervasive oxidative weathering after the initial rise of atmospheric oxygen at ~2.4 Ga (14), the concentration of many redox-sensitive elements in the ocean has been primarily controlled by marine redox conditions. For example, in today's well-oxygenated oceans, Mo is the most abundant transition metal in seawater (~107 nM; ref. 15), despite its very low crustal abundance (~1-2 ppm; ref. 16). Under sulfidic marine conditions the burial fluxes of Mo exceed those in oxygenated settings by several orders of magnitude (17). Hence, it follows that when sulfidic conditions are more widespread than today, global seawater concentrations of Mo will be much lower. Because the enrichment of Mo in sulfidic shales scales with dissolved seawater Mo concentrations (18). Mo enrichments in marine shales (independently elucidated as being deposited under euxinic conditions with a strong connection to the open ocean) can be used to track the global extent of sulfidic conditions (9). Substantial Mo enrichment in an ancient euxinic marine shale, such as occurs in modern euxinic marine sediments, implies that sulfidic bottom waters represent a very small extent of the global seafloor.

In principle, a similar approach can be used with other metals such as Cr, which, importantly, will also be reduced and buried in sediments under anoxic conditions but without the requirement of free sulfide. Chromium is readily immobilized as $(Fe,Cr)(OH)_3$ under ferruginous conditions (19, 20) and will be reduced and rendered insoluble by reaction with a wide range of other reductants under sulfidic or even denitrifying conditions (21–23). Thus, comparing Mo enrichments in independently constrained euxinic shales and Cr enrichments in independently constrained anoxic shales can offer a unique and complementary perspective on the global redox landscape of the ocean.

A better understanding of the marine Mo cycle in Proterozoic oceans may also illuminate key controlling factors in biological evolution and ecosystem development during the emergence of eukaryotic life. The biogeochemical cycles of marine trace elements form a crucial link between the inorganic chemistry of seawater and the biological modulation of atmospheric composition. The availability of iron, for example, has been invoked as a primary control on local carbon export fluxes and atmospheric

Author contributions: C.T.R. and N.J.P. designed research; C.T.R., N.J.P., L.J.R., C.A.P., B.C.G., S.V.L., A.B., K.O.K., and T.W.L. performed research; C.T.R., N.J.P., L.J.R., C.A.P., B.C.G., and S.V.L. analyzed data; and C.T.R., N.J.P., L.J.R., C.A.P., B.C.G., S.V.L., A.B., K.O.K., and T.W.L. wrote the paper.

The authors declare no conflict of interest.

This article is a PNAS Direct Submission.

¹To whom correspondence should be addressed. E-mail: reinhard@caltech.edu.

This article contains supporting information online at www.pnas.org/lookup/suppl/doi:10. 1073/pnas.1208622110/-/DCSupplemental.

 pCO_2 on glacial-interglacial timescales (24, 25). However, the leverage exerted by Fe on recent oceanic carbon fixation is most fundamentally driven by the sparing solubility of Fe in an ocean that is well-ventilated by an oxygen-rich atmosphere. By analogy, on a more reducing Earth surface Mo is likely to be a key colimiting trace nutrient given its importance in biological nitrogen fixation, assimilatory/dissimilatory nitrate reduction, and a number of other metabolically significant electron transfer processes (26–28).

To move forward in our understanding of Proterozoic redox evolution, we present a unique view of Cr and Mo enrichments in anoxic shales and a complementary modeling approach to interpret these data. From this vantage point, we present evidence that anoxic conditions were a globally important feature in the mid-Proterozoic ocean. In our analysis, we take anoxic environments to include those that are euxinic (anoxic and H₂S-rich), ferruginous (anoxic and Fe²⁺-rich), and NO₃⁻-buffered (i.e., anoxic but with low concentrations of both H₂S and Fe²⁺). We note, however, that the latter environments are likely to be spatially and temporally limited given the relatively low concentration (and thus redox buffering capacity) of NO₃⁻ in seawater, particularly during the Proterozoic (29).

Despite evidence for pervasive marine anoxia during the Proterozoic, we highlight that euxinia covered only a small portion of the seafloor. On this basis, we present a framework for linking Mo enrichments to seawater Mo concentrations that points toward Mo–N colimitation in the Proterozoic ocean. Therefore, despite a more limited extent of euxinia than previously envisaged, life in the Proterozoic ocean was heavily influenced by a prevalence of sulfide in the water column that far exceeded the small amounts of euxinia that characterize the modern ocean.

Mid-Proterozoic Geochemical Record

We present a record of Cr and Mo enrichments in anoxic and euxinic black shales through time (Fig. 1). Samples for this study (n > 3,000) come from our analytical efforts and a literature survey (Fig. S1 and Table S1). Our own data include results from over 300 Precambrian samples and modern anoxic systems. Samples were filtered for basic lithology (fine-grained siliciclastics) using a combination of basic sedimentary petrology and major-element thresholds. We relied on well-established paleoproxies rooted in Fe-S systematics to infer the redox state of the water column overlying the site of shale deposition. Importantly, these paleoredox proxies are calibrated to delineate anoxic settings (where Cr will be reduced and buried) and euxinic settings (where both Cr and Mo will be reduced/sulfidized and buried). The Fe–S paleoproxies have recently been reviewed in detail (30, 31), and full information on sample filters is provided in SI *Discussion*. We emphasize the noncircularity of our approach to analyzing the shale record—specifically, we constrained local paleoredox to have been anoxic or euxinic with no appeal to sedimentary trace metal enrichments as fingerprints of those conditions. This approach allows us to use the metal enrichments themselves as proxies for broader ocean redox state and its control on the ocean-wide inventories of those metals.

There are significant Mo enrichments in mid-Proterozoic (~1.8–0.6 Ga) euxinic shales (Fig. 1*B*). However, these enrichments are significantly lower than those observed in late Proterozoic and Phanerozoic euxinic equivalents (9). Proterozoic enrichments range from less than 10 to greater than 100 ppm, compared with concentrations on the order of ~1–2 ppm in average upper crust (16). The total mean for temporally binned mid-Proterozoic euxinic shale data (2.0–0.74 Gya) is 40.5 ppm (± 22.5 at the 95% confidence level) compared with the Phanerozoic where the total mean is 170.2 ppm (± 33.4 ppm at the 95% confidence level).

In strong contrast with the Mo record, there are no discernible Cr enrichments in mid-Proterozoic anoxic shales. We report Cr enrichments by normalizing to Ti content, as detrital inputs of Cr to marine sediments can be substantial and are greatly in excess of those for Mo. The total mean for Cr/Ti values for mid-Proterozoic anoxic shales is 1.69×10^{-2} (ppm/ppm), and the 95% confidence interval $(1.45 \times 10^{-2} \text{ to } 1.93 \times 10^{-2})$ is indistinguishable from post-Archean average upper crust (32–34) (Fig. 14). There is a marked increase in Cr/Ti values indicating enrichments of tens to hundreds of ppm. This pattern is mirrored in the sharp rise in Mo enrichments through the same interval.

We hypothesize that the enrichment trends for both metals reflect the progressive expansion of marine anoxia between 2.0 and 1.8 Ga and persistent oceanic Cr drawdown during the mid-Proterozoic. Because of the different conditions required for the reduction, immobilization, and accumulation in sediments for Cr (anoxic) and Mo (euxinic), we suggest that a relatively small proportion of oceanic anoxia was represented by euxinic conditions, which allowed a moderate although muted seawater Mo reservoir to coexist with a strongly depleted Cr reservoir. We note also that the much greater density of Cr data once the datasets have been filtered for local redox suggests qualitatively that anoxic shales were much more common than euxinic shales during the Proterozoic.

In contrast, the Phanerozoic record is generally characterized by elevated enrichments of both elements, suggesting that for most of the Phanerozoic both anoxic and euxinic conditions were less spatially and/or temporally widespread. Comparatively short-lived Phanerozoic oceanic anoxic events (OAEs), such as those famously expressed in the Mesozoic, are a notable exception. We suggest that combining enrichment records for elements that respond to



Fig. 1. Sedimentary Cr (A) and Mo (B) enrichments in anoxic and euxinic black shales through time. Because of the relatively high Cr content of typical detrital material, Cr enrichments are expressed as Cr/Ti ratios. Gray diamonds represent all filtered data, whereas black circles represent temporally binned averages. Blue boxes show the total mean (±95% confidence interval) of temporally binned averages for the mid-Proterozoic and Phanerozoic (*SI Discussion*). (*Insets*) Cumulative frequency distribution of enrichments for the mid-Proterozoic (gray curve) and the Phanerozoic (black curve). Green boxes show the composition of average post-Archean upper crust (31–33) used to approximate the detrital input. Note the log scale.

the presence of free HS^- in anoxic marine environments (Mo, Zn, etc.) with elements that respond to anoxia more generally (Cr, Re, V, etc.) may allow us to place more detailed constraints on the fabric of seafloor redox and bioinorganic feedbacks throughout Earth's history. We can then expand this approach by interpreting such data within a global mass balance framework.

Interpreting the Enrichment Record: Model for Global Mass Balance and Burial in Marine Sediments

Our quantitative model begins with a conventional mass balance formulation (35), in which the ocean is treated as a single well-mixed reservoir (Fig. S2)—a reasonable assumption given the relatively long residence times of the elements of interest (*SI Discussion*). The globally averaged concentration of a metal in the ocean evolves as

$$\frac{d}{dt} \int_{v} [\text{Me}] \ dv = F_{in} - F_{out},$$

where [Me] represents the seawater concentration of a given metal, integrated over ocean volume v. The terms F_{in} and F_{out} represent input and output fluxes, respectively. In both cases, input fluxes associated with riverine delivery and/or seawaterbasalt interaction are grouped into a single input term (F_{in} ; SI Discussion and Fig. S3), whereas output fluxes (F_{out}) are broken into burial terms specific to each metal cycle (Fig. S2 and Tables S2–S6). Riverine input dominates the overall input flux for both metals (SI Discussion), and this flux is unlikely to have varied significantly (relative to variations in the removal fluxes) after Earth's initial oxygenation. Sink fluxes (burial in sediments) are a function of the characteristic burial rate and areal extent of a given sink environment (*i*):

$$F_i = k \int\limits_{A_i} b_i^{\rm ini} \, dA_i,$$

where A_i represents the seafloor area of each sink environment (oxic, ferruginous, sulfidic, etc.), and b_i^{ini} represents the globally averaged initial burial rate characteristic of that environment. In this equation, k is a reaction coefficient that relates the burial flux to the seawater concentration. For a strictly first-order model, $k = [\text{Me}]_{/}[\text{Me}]_{\text{M}}$, where $[\text{Me}]_{l}$ is the mean oceanic concentration of a given metal at time t, and $[\text{Me}]_{\text{M}}$ is the modern seawater concentration. As previously noted (36), this kind of first-order mass balance approach to specifying removal fluxes is a specific variant of the more generalized case:

$$F_i = k^{\alpha_i} \int\limits_{A_i} b_i^{\rm ini} \, dA_i,$$

where $\alpha = 1.0$.

Combining the above terms yields an expression for each removal flux:

$$F_i = A_i b_i^{\text{ini}} \left[\frac{[\text{Me}]_t}{[\text{Me}]_{\text{M}}} \right]^{\alpha_i}$$

Following previous approaches, we first assume that $\alpha = 1.0$ (i.e., first-order mass balance). This approach is grounded in the notion that the burial rate of a metal in a given sink environment will scale in a roughly linear fashion with the ambient seawater reservoir size (18, 35). After substitution and rearrangement of the above equations, we arrive at a generalized mass balance equation for both metals:

$$\frac{d}{dt} \int_{v} [\text{Me}] dv = F_{in} - \sum A_i b_i^{ini} \frac{[\text{Me}]_t}{[\text{Me}]_M}$$

Because we are mainly interested in broad (~10⁶ y) shifts in deep ocean redox, we assume steady-state conditions for both metal systems. Assuming steady state (i.e., d[Me]/dt = 0) yields an expression for the average oceanic concentration of a given metal:

$$[\mathrm{Me}]_t = [\mathrm{Me}]_{\mathrm{M}} \frac{F_{in}}{\Sigma A_i b_i^{ini}}.$$

An important component of our model is the specification of spatially variant metal burial rates. Most past treatments of oceanic metal mass balance suffer from the physically unrealistic assumption that the metal burial rates characteristic of modern environments, typically encountered in restricted or marginal settings such as the Black Sea and Cariaco Basin, where overall sediment and carbon fluxes are high, can be scaled to very large regions of the abyssal seafloor where bulk sediment delivery and organic carbon fluxes are typically much lower (37, 38). We have attempted to avoid the same oversimplification by coupling an algorithm that addresses carbon flux to the seafloor as a function of depth (39) to a polynomial function fitted to bathymetric data (40) and by tuning an imposed burial ratio parameter ($R_{Me/C}$) to reproduce the modern globally averaged burial rate for each metal (*SI Discussion* and Fig. S4).

The essential assumption here is that a given region of the seafloor will have a characteristic "burial capacity" for Mo and Cr, regulated to first order by carbon flux to the sediment, and that this capacity will only be realized if the environment is anoxic (in the case of Cr) or euxinic (in the case of Mo). From a mechanistic perspective, this approach builds from clear evidence that the burial in sediments of many redox sensitive metals in anoxic settings scales strongly with carbon flux to the sediments (18, 41, 42). We hold that this approach allows for a more realistic depiction of perturbations to seawater metal inventories as a function of seafloor redox dynamics by smoothly decreasing globally averaged burial rates as larger regions of the seafloor become anoxic (Cr) or sulfidic (Mo).

Interpreting the Enrichment Record: Model Results

Our approach assumes, by definition, that the burial rate of a given metal in an authigenically active environment (i.e., environments that remove Cr and/or Mo from seawater and sequester them within the sediment column) scales with the ambient concentration in seawater:

$$b_i = b_i^{\text{ini}} \left[\frac{[\text{Me}]_t}{[\text{Me}]_{\text{M}}} \right]^{\alpha_i}$$

The seawater reservoir is controlled largely by how this relationship is expressed on a global scale, but this relationship will also apply to individual settings or regions of the seafloor. As a result, we can envision a generalized authigenically active setting and estimate sedimentary metal enrichments as a function of seawater concentration, in turn controlled by the relative areas of different redox environments on a global scale.

The results of such an exercise are shown in Fig. 2. Here, we have used as our starting point burial rates and overall sediment mass accumulation rates from one of the best-characterized perennially euxinic basins on the modern Earth, the Cariaco Basin in Venezuela. Our purpose here is to depict a generalized setting accumulating Cr and Mo within sediments that has a relatively open connection to the seawater metal reservoir(s), and for that reason we have chosen the Cariaco Basin over the highly

Fig. 2. Estimated authigenic sedimentary enrichments ([X]_{auth}) for Cr (A) and Mo (B) in a generalized anoxic or euxinic setting, respectively, as a function of anoxic and sulfidic seafloor area (A_{anox}, A_{sulf}). Black curves represent a bulk mass accumulation rate of 1.0×10^{-2} g·cm⁻²·y⁻¹, whereas gray dotted curves represent a factor of 1.5 above and below this value (SI Discussion; Fig. S5). The blue box in A represents the approximate area of seafloor anoxia required to drop below an enrichment threshold of 5 μ g·g⁻¹, a conservative value for our purposes given the negligible enrichments recorded by mid-Proterozoic anoxic shales. The red box in B shows the approximate sulfidic seafloor area consistent with the range of mid-Proterozoic Mo enrichments, and is scaled relative to the y axis according to the 95% confidence interval of temporally binned averages



shown in Fig. 1. Seafloor areas are shown as a percentage relative to modern seafloor area (%) and in terms of raw area (km²).

restricted Black Sea, which shows clear local reservoir effects for Mo (18). In essence, we pose the question, "How would an anoxic or euxinic continental margin or epicontinental environment, such as that represented in the marine black shale record, respond to a particular perturbation to seafloor redox state?" We can then scale this relationship to spatially varying organic carbon burial and bulk sedimentation to inform metal uptake away from the continental margin.

A striking pattern emerges when we consider the magnitude of enrichment that can be achieved in an authigenically active environment under different oceanic redox conditions (Fig. 2). If our model is correct, the negligible sedimentary Cr enrichments characteristic of the entire mid-Proterozoic would require extremely pervasive anoxic conditions. Our approach (which is likely conservative; *SI Discussion*) suggests that at least $\sim 30-40\%$ of the seafloor must have underlain anoxic deep waters to drive Cr enrichments to crustal values for sustained periods. We stress that this is a minimum estimate, and that our results are also fully consistent with virtually complete seafloor anoxia.

The Mo enrichment record, however, tells a very different story. Enrichments in euxinic environments during the mid-Proterozoic are muted relative to the Phanerozoic, a pattern that emerges as a consequence of more widespread sulfidic deposition relative to most of the Phanerozoic and is reinforced when Mo enrichments are normalized to total organic carbon (TOC; ref. 9). However, Mo enrichments in Proterozoic euxinic environments that are mostly well above crustal values are inconsistent with pervasive, ocean-scale euxinia (Fig. 2B). Instead, our model results point to roughly ~1-10% of the seafloor as having been euxinic during the mid-Proterozoic, although there is likely to have been dynamic expansion/contraction of the area of euxinic seafloor area within and occasionally beyond this range-as related, for example, to spatiotemporal patterns of primary production along ocean margins. A similar range of euxinic seafloor is implied for some brief periods of the Phanerozoic (43, 44), but the record of appreciable Cr enrichment during this latter phase of Earth history indicates much more spatially and temporally restricted anoxia overall.

Marine Euxinia: Global Effects of Regional Ocean Chemistry

The record of Cr and Mo enrichment, when interpreted in light of our model results, necessitates that euxinia covered a relatively small fraction of overall seafloor area during the mid-Proterozoic despite pervasive anoxic conditions on a global scale. Such a result adds to growing evidence that Proterozoic deep ocean chemistry was dominated by anoxic but nonsulfidic (ferruginous) conditions (10, 12, 13), in contrast with most modern anoxic marine settings that tend toward euxinia. Nevertheless, euxinia in the mid-Proterozoic ocean was likely orders of magnitude more widespread than

5360 | www.pnas.org/cgi/doi/10.1073/pnas.1208622110

today's estimate of $\sim 0.1\%$ of the seafloor, and the deleterious impacts on nutrient availability could have been enough to significantly alter global biogeochemical processes. In effect, the sensitivity of the oceanic Mo reservoir to small perturbations in the extent of sulfidic seafloor suggests that a strong distinction should be made between a sulfidic ocean in an oceanographic sense, in which a large proportion of oceanic volume and basin- or global-scale areas of the seafloor (much more than our estimate of $\sim 1-10\%$) are characterized by sulfidic waters, and an ocean that is sulfidic from a nutrient or biological perspective. From a biological perspective, trace nutrient colimitation of marine primary producers will be strongly controlled by the extent of euxinic conditions despite euxinia not being persistent throughout most of the ocean. These conditions are, of course, not mutually exclusivea globally sulfidic ocean will almost certainly result in trace element colimitation. However, the distinction is important as it highlights the leverage that relatively small regions of the ocean can exert on the biogeochemical cycling of certain biolimiting trace elements (9, 26).

To further explore this concept, we use the model to estimate globally averaged seawater [Mo] under variable scaling between the ambient seawater concentration and burial rate within sediments. The most common approach, as discussed above, is to assume strictly first-order (i.e., linear) scaling ($\alpha = 1.0$). Although data are somewhat limited and it is difficult to establish precisely what the form of this relationship should be, data from the most well-characterized perennially euxinic settings on the modern Earth suggest that this relationship may in fact be nonlinear (Fig. 3A). The effect of this parameter on steady-state globally averaged seawater [Mo] is shown in Fig. 3B. Because lowering the value for α allows for a higher burial rate (and thus removal flux from the ocean) at a given value for seawater [Mo], the concentration is ultimately drawn down to much lower steady-state values for a given perturbation.

Although the term α regulates the scaling between ambient [Mo] and Mo burial rate on a global scale in an integrated sense across diverse redox settings, it will also do so within individual authigenically active environments. Importantly, this means that the sedimentary enrichments predicted by the model for a given extent of euxinic seafloor do not vary as a function of α . Changes in this parameter are reflected by the steady-state concentration of Mo in seawater (the same applies for Cr). It is clear from this exercise (Fig. 3) that even relatively small areas of the seafloor overlain by euxinic water masses (a fraction of modern continental shelf area) are sufficient to draw the ocean's average Mo concentration to ~10 nM (an order of magnitude below that seen in the modern ocean), even with strictly linear scaling between ambient [Mo] and burial flux.



Fig. 3. Effects of deviating from a strictly first-order model. (*A*) Mo burial rates as a function of ambient dissolved Mo concentration (shown as a proportion of modern seawater [Mo]/[Mo]_M) for a range of α -values between 1.0 (strict first-order) and 0.25. Curves are calculated assuming a modern globally averaged euxinic burial rate of 1.53 µg·cm⁻²·y⁻¹. Black circles represent values for well-characterized perennially euxinic marine basins on the modern Earth [Black Sea (BS), Framvaren Fjord (FF), and the Cariaco Trench (CT)]. (*B*) Steady-state globally averaged seawater Mo concentrations as a function of sulfidic seafloor area (A_{sulf}) for different values of α . The shaded box depicts values below 10 nM.

Further work is needed to better pinpoint the levels of seawater Mo that should be considered biologically limiting, but available evidence is consistent with biolimiting concentrations in mid-Proterozoic oceans. Culturing experiments with modern strains of diazotrophic (nitrogen-fixing) cyanobacteria generally indicate that rates of nitrogen fixation and overall growth become impacted by Mo availability once concentrations fall to within the \sim 1–10-nM range (45–49). Some strains seem to show resilience to Mo scarcity until concentrations fall below ~5 nM (48), but in general there seems to be a sharp change in overall growth rates, cell-specific nitrogen fixation rates, and stoichiometric growth status within the 1-10-nM range. Even small changes in relative rates of diazotrophy, if expressed globally and on protracted timescales, can be expected to have large effects on carbon, nitrogen, and oxygen cycling as well as ecosystem structure in the surface ocean.

Although Mo enrichments in anoxic shales deposited during the mid-Proterozoic do not approach those characteristic of Phanerozoic settings, enrichment levels are nonetheless maintained well above crustal values. Thus, Mo enrichments in mid-Proterozoic euxinic marine settings seem poised within a very sensitive region of parameter space. We stress that such a relationship implies some kind of stabilizing feedback controlled by Mo-N colimitation (e.g., ref. 9). In this scenario, widespread euxinic conditions would deplete the Mo reservoir, thereby limiting primary productivity and carbon export flux. This in turn would reduce the amount of biomass oxidized via microbial sulfate reduction (which produces HS⁻), limiting sulfide accumulation in marine settings. The ultimate result would be for Mo concentrations to rebound (a negative feedback; ref. 9). However, it would be difficult to transition from a Mo-N colimited system to a strongly oxidizing, Mo-replete ocean. Such a shift would need to be driven ultimately by a longterm increase in sedimentary burial of organic matter, but this burial would lead to a corresponding increase in Mo burial fluxes pushing the system back to Mo-N colimitation. The link between primary production and Mo removal from the ocean would again be the microbial production of hydrogen sulfide needed for efficient Mo burial. The response time of Mo in a Mo-depleted ocean is likely to be short enough (relative to the residence time of oxygen in the ocean/atmosphere system) to induce a rapid and efficient stabilizing feedback on redox conditions. Iron will be orders of magnitude more soluble under any form of anoxia (euxinic or ferruginous) than it is in the modern ocean. In this light, in a reducing ocean, the coupled C-S-Fe-Mo biogeochemical cycles form an attractor-driving the marine system toward persistent trace metal-macronutrient colimitation. This relationship is similar, in essence, to the control exerted by limited Fe solubility in an oxidizing and well-ventilated ocean, but we expect that the

stabilizing feedbacks and sensitivity responses will be very different between the two systems.

Conclusions

Exploration of the Cr and Mo enrichment record in anoxic marine shales during the last ~2.0 Ga within a mass balance framework reveals that the mid-Proterozoic ocean was characterized by pervasive anoxic conditions, as manifested by negligible Cr enrichments in anoxic shales, but limited euxinia, as reflected in nontrivial Mo enrichments in euxinic shales that are nonetheless guite muted relative to most Phanerozoic equivalents. The Phanerozoic ocean appears to have been marked by more circumscribed anoxia on the whole, with anoxic shales typically showing substantial Cr enrichments. As a result, a potentially much larger *relative* fraction of this anoxia may have been represented by euxinic conditions, in particular during Cretaceous OAEs and periods of anomalously widespread anoxia during the Paleozoic (44). It remains to be explored if these episodes represent a fundamentally different mode of anoxic marine conditions or whether they can be viewed as temporary reversions to mid-Proterozoic conditions.

In addition, our model points toward a view in which the chemistry of small and dynamic regions of the seafloor exerts fundamental control on biological carbon and oxygen cycling through bioinorganic feedbacks related to trace element availability (9, 26), in much the same way that carbon cycling and export in large regions of the modern well-ventilated ocean are controlled by the availability of Fe. Moving forward, it will be important to explore in detail, and with a wide range of organisms, the thresholds at which diazotrophs are strongly impacted by Mo availability. It will also be important to develop explicit ecological models aimed at delineating the constraints and feedbacks associated with Mo-N colimited planktonic ecosystems. For example, elevated growth rates and doubling times due to greater overall Fe availability (as the solubility of Fe in any anoxic state will be orders of magnitude above that seen in oxic systems) may be able to compensate for lower cell-specific rates of nitrogen fixation within the context of ecosystem nitrogen supply. Further, although there is some understanding of the Mo requirements for assimilatory nitrate uptake (e.g., ref. 50), little is known regarding the effects of Mo availability on dissimilatory nitrate reductase. Finally, it is clear that some diazotrophs show biochemical idiosyncrasies aimed at dealing with Mo scarcity (49), and recent work on the exquisite adaptation of some diazotrophic organisms to Fe limitation in the modern oceans (51) begs for a more thorough exploration of the biochemistry of Mo-limited diazotrophy.

In any case, our results provide strong independent evidence for an emerging first-order model of Proterozoic ocean redox structure. In this model, the surface ocean is well-ventilated through air-sea gas exchange and local biological O2 production, but the deep ocean is anoxic largely as a result of equilibration with atmospheric pO_2 at least 1–2 orders of magnitude below the modern value in regions of deep-water formation (4). The increased mobility and transport of dissolved Fe(II) under reducing conditions, combined with spatially heterogeneous carbon fluxes through marine systems (as constrained by the intensity of vertical exchange through upwelling and eddy diffusion), yielded an ocean that was pervasively anoxic (i.e., redoxbuffered by Fe²⁺ or NO₃⁻) with localized regions of euxinia in marginal settings (12, 13). We note that although our model is, in principle, somewhat sensitive to variations in the sourcing of Cr to the ocean (SI Discussion), it is also supported by data from other elements whose sourcing to the ocean should only be weakly dependent on atmospheric pO_2 (52). This emerging model provides a backdrop for the early evolution and ecological expansion of eukaryotic organisms (26, 53) and the biogeochemical

- 1. Holland HD (1984) The Chemical Evolution of the Atmosphere and Oceans (Princeton Univ Press, Princeton).
- 2. Cloud P (1972) A working model of the primitive Earth. Am J Sci 272(6):537-548.
- Payne JL, et al. (2009) Two-phase increase in the maximum size of life over 3.5 billion years reflects biological innovation and environmental opportunity. Proc Natl Acad Sci USA 106(1):24–27.
- Canfield DE (1998) A new model for Proterozoic ocean chemistry. Nature 396(6692): 450–453.
- Shen Y, Canfield DE, Knoll AH (2002) Middle Proterozoic ocean chemistry: Evidence from the McArthur Basin, Northern Australia. Am J Sci 302(2):81–109.
- Shen Y, Knoll AH, Walter MR (2003) Evidence for low sulphate and anoxia in a mid-Proterozoic marine basin. Nature 423(6940):632–635.
- Poulton SW, Fralick PW, Canfield DE (2004) The transition to a sulphidic ocean approximately 1.84 billion years ago. Nature 431(7005):173–177.
- Brocks JJ, et al. (2005) Biomarker evidence for green and purple sulphur bacteria in a stratified Palaeoproterozoic sea. Nature 437(7060):866–870.
- 9. Scott C, et al. (2008) Tracing the stepwise oxygenation of the Proterozoic ocean. *Nature* 452(7186):456–459.
- Canfield DE, et al. (2008) Ferruginous conditions dominated later Neoproterozoic deep-water chemistry. Science 321(5891):949–952.
- 11. Lyons TW, Reinhard CT, Scott C (2009) Redox redux. Geobiology 7(5):489-494.
- Poulton SW, Fralick PW, Canfield DE (2010) Spatial variability in oceanic redox structure 1.8 billion years ago. Nat Geosci 3(7):486–490.
- Planavsky NJ, et al. (2011) Widespread iron-rich conditions in the mid-Proterozoic ocean. Nature 477(7365):448–451.
- Bekker A, et al. (2004) Dating the rise of atmospheric oxygen. Nature 427(6970): 117–120.
- Collier RW (1985) Molybdenum in the Northeast Pacific Ocean. Limnol Oceanogr 30(6):1351–1354.
- Taylor SR, McLennan SM (1995) The geochemical evolution of the continental crust. *Rev Geophys* 33(2):241–265.
- 17. Emerson SR, Huested SS (1991) Ocean anoxia and the concentrations of molybdenum and vanadium in seawater. *Mar Chem* 34(3-4):177–196.
- Algeo TJ, Lyons TW (2006) Mo-total organic carbon covariation in modern anoxic marine environments: Implications for analysis of paleoredox and paleohydrographic conditions. *Paleoceanography*, 10.1029/2004PA001112.
- Eary LE, Rai D (1989) Kinetics of chromate reduction by ferrous ions derived from hematite and biotite at 25°C. Am J Sci 289(2):180–213.
- Fendorf SE, Li G (1996) Kinetics of chromate reduction by ferrous iron. Environ Sci Technol 30(5):1614–1617.
- Pettine M, Millero FJ, Passino R (1994) Reduction of chromium (VI) with hydrogen sulfide in NaCl media. Mar Chem 46(4):335–344.
- Graham AM, Bouwer EJ (2010) Rates of hexavalent chromium reduction in anoxic estuarine sediments: pH effects and the role of acid volatile sulfides. *Environ Sci Technol* 44(1):136–142.
- Rue EL, Smith GJ, Cutter GA, Bruland KW (1997) The response of trace element redox couples to suboxic conditions in the water column. *Deep Sea Res Part I Oceanogr Res Pap* 44(1):113–134.
- Martin JH (1990) Glacial-interglacial CO₂ change: The iron hypothesis. Paleoceanography 5(1):1–13.
- Behrenfeld MJ, Kolber ZS (1999) Widespread iron limitation of phytoplankton in the South Pacific Ocean. Science 283(5403):840–843.
- Anbar AD, Knoll AH (2002) Proterozoic ocean chemistry and evolution: A bioinorganic bridge? *Science* 297(5584):1137–1142.
- Schwarz G, Mendel RR, Ribbe MW (2009) Molybdenum cofactors, enzymes and pathways. Nature 460(7257):839–847.
- Glass JB, Wolfe-Simon F, Anbar AD (2009) Coevolution of metal availability and nitrogen assimilation in cyanobacteria and algae. *Geobiology* 7(2):100–123.
- Fennel K, Follows M, Falkowski PG (2005) The co-evolution of the nitrogen, carbon and oxygen cycles in the Proterozoic ocean. Am J Sci 305(6-8):526–545.

feedbacks controlled by the progressive restructuring of primary producing communities (54). Finally, the sensitivity of the oceanic Mo reservoir to perturbation, combined with the relative uniformity of the Mo enrichment record in Proterozoic euxinic shales, implies that this redox structure may have been stable on long timescales as a function of Mo–N colimitation in the surface ocean. This hypothesis can be tested through the generation of more Proterozoic shale data, whereas further modeling might constrain how robust such a feedback could be and what conditions would have been required to overcome it during the later Proterozoic oxygenation of the deep ocean and subsequent evolution of macroscopic life.

ACKNOWLEDGMENTS. This research was supported by a National Aeronautics and Space Administration Exobiology grant (to T.W.L.), a National Science Foundation graduate fellowship (to N.J.P.), and a Natural Sciences and Engineering Research Council (Canada) Discovery Grant (to A.B.).

- Lyons TW, et al. (2009) Tracking euxinia in the ancient ocean: A multiproxy perspective and Proterozoic case study. Annu Rev Earth Planet Sci 37:507–534.
- Poulton SW, Canfield DE (2011) Ferruginous conditions: A dominant feature of the ocean through Earth's history. *Elements* 7(2):107–112.
- Condie KC (1993) Chemical composition and evolution of the upper continental crust: Contrasting results from surface samples and shales. *Chem Geol* 104(1-4):1–37.
- McLennan SM (2001) Relationships between the trace element composition of sedimentary rocks and upper continental crust. *Geochem Geophys Geosyst*, 2000GC000109.
- Rudnick RL, Gao S (2003) Composition of the continental crust, *Treatise on Geochemistry*, eds Holland HD, Turekian KK, Vol 3, pp 1–64.
- Rosenthal Y, Boyle EA, Labeyrie L, Oppo D (1995) Glacial enrichments of authigenic Cd and U in Subantarctic sediments: A climatic control on the elements' oceanic budget? *Paleoceanography* 10(3):395–413.
- Dahl TW, et al. (2011) Molybdenum evidence for expansive sulfidic water masses in ~750 Ma oceans. Earth Planet Sci Lett 311(3-4):264–274.
- Jahnke RA (1996) The global ocean flux of particulate organic carbon: Areal distribution and magnitude. Global Biogeochem Cycles 10(1):71–88.
- Dunne JP, Sarmiento JL, Gnanadesikan A (2007) A synthesis of global particle export from the surface ocean and cycling through the ocean interior and on the seafloor. *Global Biogeochem Cycles*, 10.1029/2006GB002907.
- Middelburg JJ, Soetaert K, Herman PMJ, Heip CHR (1996) Denitrification in marine sediments: A model study. *Global Biogeochem Cycles* 10(4):661–673.
- Menard HW, Smith SM (1966) Hypsometry of ocean basin provinces. J Geophys Res 71(18):4305–4325.
- Algeo TJ, Maynard JB (2004) Trace-element behavior and redox facies in core shales of Upper Pennsylvanian Kansas-type cyclothems. Chem Geol 206(3-4):289–318.
- 42. Dean WE (2007) Sediment geochemical records of productivity and oxygen depletion along the margin of western North America during the past 60,000 years: Teleconnections with Greenland Ice and the Cariaco Basin. Q Sci Rev 26(1-2):98–114.
- Pearce CR, Cohen AS, Coe AL, Burton KW (2008) Molybdenum isotope evidence for global ocean anoxia coupled with perturbations to the carbon cycle during the Early Jurassic. *Geology* 36(3):231–234.
- Gill BC, et al. (2011) Geochemical evidence for widespread euxinia in the later Cambrian ocean. Nature 469(7328):80–83.
- Fay P, de Vasconcelos L (1974) Nitrogen metabolism and ultrastructure in Anabaena cylindrica. Arch Microbiol 99(1):221–230.
- Jacobs R, Lind O (1977) The combined relationship of temperature and molybdenum concentration to nitrogen fixation by Anabaena Cylindrica. Microb Ecol 3(3):205–217.
- Zahalak M, Pratte B, Werth KJ, Thiel T (2004) Molybdate transport and its effect on nitrogen utilization in the cyanobacterium *Anabaena variabilis* ATCC 29413. *Mol Microbiol* 51(2):539–549.
- Zerkle AL, House CH, Cox RP, Canfield DE (2006) Metal limitation of cyanobacterial N₂ fixation and implications for the Precambrian nitrogen cycle. *Geobiology* 4(4): 285–297.
- Glass JB, Wolfe-Simon F, Elser JJ, Anbar AD (2010) Molybdenum-nitrogen co-limitation in freshwater and coastal heterocystous cyanobacteria. *Limnol Oceanogr* 55(2):667–676.
- Glass JB, Axler RP, Chandra S, Goldman CR (2012) Molybdenum limitation of microbial nitrogen assimilation in aquatic ecosystems and pure cultures. *Front Microbiol*, 10.3389/fmicb.2012.00331.
- Saito MA, et al. (2011) Iron conservation by reduction of metalloenzyme inventories in the marine diazotroph Crocosphaera watsonii. Proc Natl Acad Sci USA 108(6): 2184–2189.
- Partin C, et al. (2012) Fluctuations in Precambrian atmospheric and oceanic oxygen levels: A new Precambrian paradigm emerging? *Mineralogical Magazine* 76(6):2207.
- Javaux EJ, Knoll AH, Walter MR (2001) Morphological and ecological complexity in early eukaryotic ecosystems. *Nature* 412(6842):66–69.
- Johnston DT, Wolfe-Simon F, Pearson A, Knoll AH (2009) Anoxygenic photosynthesis modulated Proterozoic oxygen and sustained Earth's middle age. Proc Natl Acad Sci USA 106(40):16925–16929.

Supporting Information

Reinhard et al. 10.1073/pnas.1208622110

SI Discussion

Metal Enrichment Database and Filtering Protocols. Through our own analytical efforts and a literature survey, we have assembled a database of molybdenum (Mo) and chromium (Cr) concentrations for over 3,000 samples. Data sources are shown in Table S1. Samples were initially filtered to represent solely fine-grained siliciclastic sediments, using basic petrographic observation and major element thresholds. Samples were required to contain weight percent (wt %) levels of iron (Fe) and aluminum (Al). Samples containing less than 1.0 wt % total organic carbon were also removed.

The information contained within a particular degree of authigenic enrichment of Cr or Mo depends on local depositional redox. Thus, samples were further filtered such that Cr data were only analyzed from anoxic shales, and Mo data were only analyzed from euxinic shales. Anoxic shales were delineated as having Fe_T/Al > 0.5 (1) and/or Fe_{HR}/Fe_T > 0.38 (2–4), where Fe_{HR} designates "highly reactive" Fe (Fe that is reactive to dissolved H₂S on syngenetic or diagenetic timescales; ref. 5) and Fe_T represents total Fe. Euxinic settings were delineated by combining the above thresholds for anoxia with either Fe_{PY}/Fe_{HR} > 0.7 (4, 6) or with elevated values for degree of pyritization (DOP > 0.6; ref. 7), defined as (8)

$$DOP = \frac{Fe_{PY}}{Fe_{PY} + Fe_{HCL}}$$

where Fe_{HCl} is Fe soluble in a 1-min boiling concentrated HCl leach and Fe_{PY} denotes pyrite Fe. Because elevated DOP has also been shown to require enhanced Fe mobility and transport (1, 7), and because it is an extremely robust analytical measurement, DOP > 0.8 supersedes all other redox filters in the designation of euxinia. In some cases, total sulfur content is used to calculate DOP_T, according to

$$DOP_T = \frac{Fe_S}{Fe_S + Fe_T}$$

where Fe_S denotes the inferred amount of sulfur-bound Fe assuming that total sulfur represents pyrite sulfur (9). Modern systems (such as the Black Sea and Cariaco Basin) were additionally filtered by site location, for extreme silicilastic dilution, and the presence of bioturbation, and units with fewer than ten filtered samples were not included in overall statistical calculations.

Our focus here is on the contrast in enrichment records between the mid-Proterozoic and the Phanerozoic. However, as noted in the main text, there appear to be substantial Cr enrichments in late Paleoproterozoic anoxic shales. Recent work has suggested an increase and subsequent decrease in Earth surface oxidation during the Paleoproterozoic (10, 11), and we suggest in the main text that the Cr enrichment pattern is possibly related to this, with the subsequent drawdown of Cr enrichments representing the progressive expansion of anoxic marine environments. Shown in Fig. S1 is the full enrichment record, including Archean and Paleoproterozoic data. The Mo enrichment record in euxinic shales (Fig. S1B) is similar to previous work (12), showing minimal enrichment during the Archean and appreciable although muted enrichments (relative to the Phanerozoic) during most of the Proterozoic. The Cr enrichment record shows some very high enrichments during the Archean and Paleoproterozoic (Fig. S1A). Some of this trend is almost certainly because Archean upper crust was more mafic in composition, and much of these enrichments could be linked to very Cr-rich source terrains. The Paleoproterozoic enrichments, on the other hand, may well be authigenic enrichments. Future isotopic and petrographic work is required to distinguish between these two models.

Modern Mo Mass Balance. We begin by assuming steady state, wherein a single input flux (F_{in}) is balanced by removal via authigenic burial into three main sedimentary sinks: an oxic sink (F_{ox}) , a reducing sediment sink (F_{red}) , and a sulfidic sink (F_{sulf}) . Our balanced modern Mo budget is shown in Fig. S2 and Table S2, and individual removal terms are discussed below. The weathering flux of Mo on the modern Earth is dominated by the mobilization of Mo from sulfide mineral phases or organics in sedimentary and igneous rocks and transport as dissolved MoO₄²⁻ and we set as F_{in} a recently obtained modern riverine flux of dissolved MoO₄² to the ocean (13). This flux is somewhat larger than those conventionally used, but is derived from the most extensive riverine database generated to date, representing 38 rivers across 5 continents and including 11 of 19 large-scale drainage areas. However, regardless of our choice of estimate for the riverine Mo flux, sensitivity analysis (Fig. S3) indicates that our conclusions are weakly sensitive to the assumed value of the input flux over a wide range. We neglect hydrothermal fluxes of Mo to/from the ocean, as these are either poorly established or likely to be quantitatively small (see below).

Oxic settings are defined as those in which Mn is permanently removed from the ocean as an oxide phase (with associated adsorbed Mo). In many oxic deep-sea settings, dissolved O₂ penetrates to the sediment-basalt interface (14, 15) and this Mn (and Mo) will effectively be buried permanently. In other settings, dissimilatory microbial Mn reduction deeper in the sediment column can remobilize Mn (and, presumably, associated Mo). However, when O₂ penetration depths are large (multiple centimeters or more) upward-diffusing Mn will be quantitatively oxidized at a steady-state oxidation front (16-18), effectively removing Mn and Mo from the ocean on a timescale characteristic of tectonic recycling of seafloor sediments (on the order of $\sim 10^8$ y). Morford and Emerson (19) suggest that once O₂ penetration falls below ~1 cm, Mn and Mo will be recycled and released from shallow sediments. We therefore characterize oxic seafloor as being the areal extent of sediments in which O₂ penetration exceeds 1 cm. This is estimated using global diagenetic models (20, 21) to be $\sim 3 \times 10^8$ km², or roughly 84% of modern seafloor area. We stress that there are fairly large regions of the seafloor that are essentially Mo neutral (see below), such that the total seafloor area for the entire budget need not sum to 100%. This area is then combined with a burial rate of $2.75 \times 10^{-3} \,\mu \text{g} \cdot \text{cm}^{-2} \cdot \text{y}^{-1}$, estimated by compiling Mo burial rates in oxic settings (22) and by combining Mn burial rates in oxic pelagic sediments (23) with a characteristic Mo/Mn ratio of 2×10^{-3} (24). The combined sink is shown in Table S2.

Sulfidic settings are defined as environments in which dissolved H_2S accumulates at or above the sediment-water interface. This includes traditional euxinic settings (Black Sea, Cariaco Basin), but is also meant to include small areas of the seafloor below regions of intense upwelling (Peru margin, Namibian shelf), where dissolved H_2S is present at high levels essentially at the sediment-water interface and occasionally breaches into the water column (25, 26). Our sulfidic sink is calculated by combining estimates of seafloor area, authigenic enrichment, and bulk mass accumulation rate (MAR) for modern sulfidic settings (2, 27–39). The globally averaged sulfidic burial calculated

through such an approach will be biased low—given that the modern extent of euxinic seafloor, on an areal basis, is dominated by the Black Sea, and this setting is characterized by low burial rates due to restricted exchange over the Bosporus sill and an evolved Mo reservoir (36). As a result, the global sulfidic burial rates implemented in the model are referenced to a modern globally averaged sulfidic burial rate that neglects the influence of the Black Sea. This is done in an effort to represent the burial capacity of marine settings with unfettered access to the seawater Mo reservoir (12).

The final sink is reducing sediments. This sink represents environments that have been referred to by the rather ambiguous term "suboxic." We follow ref. 12 in designating these environments as those in which dissolved H₂S accumulation is restricted to pore waters, but further point out that reducing sediments in which O_2 penetration is less than ~1 cm and H_2S accumulation occurs more deeply in the sediments do not effectively bury Mo (19). From a mechanistic perspective, these reducing sediment environments are typically associated with relatively low bottomwater O2, but the effectiveness of Mo sequestration in these settings is most likely a more complex function of Mn flux to sediments (and, thus, bottom-water O₂), sedimentation rate, and labile organic carbon flux to the sediment-water interface. In any case, we use a somewhat moderate burial rate for reducing sediments of 0.27 μ g·cm⁻²·y⁻¹ (37, 40–43), and use the remaining parameters of the budget to solve for the seafloor area represented by this sink (Table S2). Although we present a revised approach for estimating the global Mo removal fluxes, our result is similar to previous estimates based on consideration of bulk burial rates (12) and isotope mass balance (40).

Modern Cr Mass Balance. As for Mo, we begin by assuming steady state, with a single input flux (F_{in}) balanced by three authigenic burial fluxes: an oxic sink (F_{ox}) , a reducing sediment sink (F_{red}) , and an anoxic sink (F_{anox}). In our modeling analysis, we take anoxic environments to include those that are euxinic (anoxic and H₂S-rich), ferruginous (anoxic and Fe²⁺-rich), and $NO_3^$ buffered (i.e., anoxic but with low concentrations of both H_2S and Fe^{2+}). We note, however, that the latter environments are likely to be spatially and temporally limited, given the relatively low concentration (and thus redox buffering capacity) of NO₃⁻ in seawater. Potential hydrothermal fluxes to/from the ocean are neglected in our treatment of the modern Cr cycle, as currently available data suggest that these fluxes are quantitatively insignificant (see below). The Cr mass balance is rather poorly constrained-compared with that for Mo. However, we suggest that although our mass balance is likely to be revised as better estimates of fluxes and reservoirs become available, this is very unlikely to change our fundamental conclusions.

Our input flux is calculated following the method of ref. 13. In brief, we compiled a database of dissolved [Cr] values and annual discharge rates for rivers (Table S3), and used this to calculate discharge-weighted dissolved [Cr] values for individual largescale drainage regions according to the available data. The Cr flux from each large-scale drainage region was then computed by combining the discharge-weighted [Cr] value with the total discharge flux from each region. These regional fluxes were combined and used to estimate a global discharge-weighted dissolved [Cr] value of 14.82 nM. This value was then combined with an estimated global exorheic flux of 3.9×10^4 km³·y⁻¹ (44), yielding an estimate of the discharge-weighted flux of dissolved Cr to the modern ocean. Although dissolved Cr data are less common than dissolved major ion data for rivers, the large-scale drainage regions in our database account for $\sim 70\%$ of the overall exorheic flux to the oceans. Assuming minimal estuarine removal (45, 46), and combining this estimate with an average seawater concentration of 4 nM (47) and an ocean volume of 1.37×10^{21} L, this yields a residence time for Cr in the modern ocean of ~9,500 y

(i.e., over a factor of 9 greater than characteristic timescales of ocean mixing). To our knowledge, estimates of this sort are few, but ours is well in line with previous attempts (e.g., ref. 48). As for Mo, sensitivity analysis (Fig. S3) indicates that our conclusions are not likely to be fundamentally altered unless input fluxes to the ocean become extremely low.

Dissolved Cr(VI) species should become adsorbed onto the surface of metal- and Al-oxide phases (49-51). We therefore expect some nontrivial burial flux of Cr in oxic settings, although we note that sorption to Al-oxide phases decreases sharply when approaching circumneutral pH (52). The Cr content of pelagic red clays, although often elevated above crustal values with respect to Cr/Ti ratios, is rather variable. We use in our budget a relatively low oxic Cr burial rate of $1.0 \times 10^{-3} \,\mu\text{g}\,\text{cm}^{-2}\,\text{y}^{-1}$, of the same order as our much better constrained Mo burial flux. , of This corresponds to a sediment with a Cr/Ti ratio of 1.87×10^{-2} , consistent with typical values from pelagic red clays (53-55), accumulating at a burial rate of 1.0×10^{-3} g cm⁻² y⁻¹. Because the burial of Cr in oxic settings should depend on the efficiency of metal oxide burial, this burial rate is then combined with the same areal extent of oxic seafloor (defined by sediment O_2 penetration depth) discussed above.

The anoxic sink for Cr is defined in a similar manner to the sulfidic sink for Mo, a natural result of the fact that on the modern Earth the relative mobility and transport of S and Fe are such that anoxic settings tend to become euxinic (anoxic and sulfidic). We use Cr/Ti ratios from the Cariaco Basin (34, 56, 57) to obtain a modern anoxic burial rate of ~0.5 μ g·cm⁻²·y⁻¹, and scale this to the seafloor area of anoxic environments as discussed above for the modern Mo budget. This burial rate is roughly of the same order as that for Mo in euxinic settings, although we acknowledge that these estimates will improve with further generation and analysis of Cr data in anoxic marine systems. However, Cr will be reduced and immobilized as Cr(III) via a wide range of reductants-dissolved H₂S is not necessary (58-60). Indeed, Cr(VI) reduction to Cr(III) has been shown to take place in the openwater column of the eastern tropical Pacific, coincident with the onset of microbial denitrification (61). This provides a crucial distinction with the behavior of Mo, in that effective Mo capture requires the additional presence of free dissolved sulfide, and forms the centerpiece of our analysis. The reducing sediment sink is again solved for using the other parameters of the budget. We assume an authigenic burial rate of 0.15 µg·cm⁻²·y⁻¹, derived from combining Cr/Ti ratios in the Gulf of California (62) with the requisite bulk MAR (63). This aspect of the budget is not well constrained, but we consider it unlikely that such settings will authigenically bury Cr at rates much higher than this. In other words, we use what we consider to be a relatively high burial rate to avoid underestimating the magnitude of this sink relative to the anoxic sink, rendering this portion of the budget conservative for the conclusions presented here. Parameters for our modern balanced Cr budget are shown in Fig. S2 and Table S4.

Hydrothermal Cycling of Mo and Cr. The systematics of Mo and Cr in hydrothermal systems and the effects of hydrothermal processes on the Earth surface cycles of Mo and Cr have not been explored in detail, but we can place some basic constraints on the possible effects of high- and low-temperature seawater–basalt interaction on the mass balances of Mo and Cr in the ocean. The water flux through a high-temperature hydrothermal system $(F_{(ht)}; \text{ in kg } \cdot y^{-1})$ can be estimated as (64)

$$F_{(ht)} = \frac{Q_{(ht)}}{\Delta T_{(ht)}c_p},$$

where $Q_{(ht)}$ is the hydrothermal heat flux, $\Delta T_{(ht)}$ is the seawater temperature anomaly, and c_p is the specific heat of seawater (at

seafloor pressure and vent fluid temperature). We can combine this with a concentration anomaly for a given metal (Δ [Me] = [Me]_{*SW*} – [Me]_{*fluid*}) to estimate a high-temperature hydrothermal flux to/from seawater (F_{hvd}) as

$$F_{(ht)} = \frac{Q_{(ht)}}{\Delta T_{(ht)}c_p} \Delta[\text{Me}],$$

Results of this calculation for both Mo and Cr are shown in Table S5. These calculations suggest that high-temperature seawaterbasalt interaction represents a removal flux of both Mo and Cr that is very small relative to the riverine flux of either element. We note that such estimates are inherently imprecise, given uncertainties in the magnitude of on-axis heat flow (ref. 64 and references therein) and analytical difficulties associated with obtaining unadulterated fluid chemistry. In the case of hightemperature fluids, it is most likely that these concentrations have been perturbed by mixing with seawater Cr and/or Mo, which would cause us to underestimate the magnitude of these sink terms. However, this should have a negligible effect on our result, given that reported concentration anomalies indicate near-complete removal of both elements during high-temperature seawater-basalt interaction (Table S5). Assuming complete removal of seawater Mo from the circulating fluid (i.e., Δ [Mo] = 107 nM) would increase our estimated high-temperature removal flux from 0.85 to 0.91% of the total input flux. Making the same assumption for Cr (i.e., Δ [Cr] = 4 nM), would have a trivial effect on the estimated high-temperature removal flux.

Low-temperature, off-axis hydrothermal systems are a much more difficult problem to address. Pristine vent fluid composition is not well constrained for many settings, but, more importantly, the global water flux through such systems is very poorly constrained. Given that the temperature anomaly is probably small, a much larger water flux would be necessary to dissipate the requisite heat flow. As a result, even a very small concentration anomaly may result in a significant flux to/from seawater on a global scale. Magnesium (Mg²⁺) substitutes readily for calcium (Ca^{2+}) during seafloor basalt alteration (65), and is removed from seawater during hydrothermal fluid evolution at both high and low temperature (66-68). Using the above method of calculation, the high-temperature removal flux of Mg^{2+} from sea-water can be estimated as ~1.3 × 10¹² mol·y⁻¹. By combining the global discharge rate used above with a global average riverine Mg²⁺ concentration of 128 μ mol kg⁻¹ (69), we derive a global riverine Mg²⁺ flux of ~5.0 × 10¹² mol y⁻¹. If we assume that the balance between the riverine flux and removal during hightemperature seawater-basalt interaction is made up by lowtemperature flow, we can use the Mg²⁺ concentration anomaly $(\Delta[Mg^{2+}] = [Mg^{2+}]_{SW} - [Mg^{2+}]_{fluid})$ of well-constrained diffuse flow systems such as that along the Juan de Fuca Ridge to calculate an approximate water flux through such systems of 9.5 \times $10^{13} \text{ kg} \cdot \text{y}^{-1}$

Combining this estimated water flux with available chemical anomalies for Mo and Cr allows us to place rough limits on the magnitude of the low-temperature fluxes of these elements to seawater (Table S6). These may be upper limits given available constraints, as the calculations assume no other removal fluxes of Mg^{2+} from seawater [i.e., uptake during carbonate burial or clay mineral alteration during "reverse weathering" reactions (70)]. We suggest that although low-temperature fluxes are likely to be somewhat larger than those that occur during on-axis fluid flow, they are still a relatively small fraction of the corresponding riverine fluxes. Given the framework outlined above, it is highly unlikely that the flux of either element will exceed ~10% of their respective riverine inputs. Furthermore, as stated above, sensitivity analysis indicates that our results are not strongly affected by reasonable changes in Cr and/or Mo input fluxes (Fig. S2). Offshore Scaling of Metal Burial Rates in the Model. Our modeling approach essentially involves balancing the modern steady-state cycles of both Cr and Mo and applying a continuous range of perturbations to this balanced cycle to explore the new steady state attained under different oceanic redox regimes. In doing so, we begin with a conventional first-order mass balance formulation. This class of model, often used to explore the dynamics of various chemical tracers in the ocean and their isotope systems, makes the implicit assumption that the burial fluxes characteristic of some particular environment (typically organic-rich continental margin sediments or marginal restricted basins) can be universally applied to extremely large regions of the seafloor. In other words, it is assumed that a burial rate characteristic of, say, the Peru margin can be applied to the abyssal realm of the ocean. This is almost certainly physically unrealistic, as open ocean settings are characterized by much lower bulk sediment fluxes, and, in particular, organic carbon fluxes (71-74). As a result, if a particular region of the abyssal ocean becomes authigenically active for some chemical constituent of seawater, it can be expected that the removal rates of that constituent into the sediment column will be much lower than those seen in more marginal settings. The net result will be a system that is overly sensitive to perturbation, as burial fluxes in large regions of the deep sea will be overestimated. This dilemma, inherent in conventional first-order mass balance analysis, has been noted by some previous work (75, 76) but has not been explored in detail. This problem is particularly acute for redox-sensitive transition metals, such as Mo and Cr, given that the organic matter flux is typically thought to be directly involved in metal sequestration (e.g., ref. 36).

We have attempted to alleviate this problem by adding a "pseudospatial" dimension to the conventional one-box ocean mass balance approach. We take an algorithm used in global diagenetic models (77) for organic carbon flux to the seafloor as a function of depth, which is then coupled to a polynomial function fitted to bathymetric data for the modern ocean (78). We then use a burial flux ratio ($R_{Me/C}$, where Me refers to Mo or Cr) for each element, a tuned parameter resulting in a relationship that encodes a decrease in local (and globally averaged) metal burial rates as larger regions of the seafloor become authigenically active. Values for $R_{Me/C}$ are tuned to reproduce the modern condition (i.e., the modern globally averaged burial rates at $\sim 0.1\%$ seafloor anoxia; Fig. S4). The essential concept here is that a given region of the seafloor has a characteristic burial capacity for either Cr or Mo, regulated to first order by the relative carbon flux through the water column and to the sediments, and that this burial capacity will only be reached when a region of the ocean achieves the requisite redox characteristics for each metal.

We stress that because the metal burial rates are derived by using a tunable ratio, this pattern is not explicitly dependent on the absolute value of the carbon flux to the seafloor at a given depth-rather, it hinges on the observation that carbon fluxes to the seafloor will decrease as one moves out into the deep sea, with the first-order topology depicted in Fig. S3. This is important, as dramatically different redox structures within the ocean, extreme differences in the composition of primary producing communities, mineral ballasting, etc., might be expected to result in significant differences in the absolute value of the carbon flux to the seafloor within different regions of the ocean. However, we consider it unlikely that the basic pattern of an offshore decrease in carbon fluxes has changed much throughout Earth's history on a global scale. In addition, although the basic bathymetry of the ocean has doubtless changed throughout Earth's history, we consider the modern depth-area curve to represent a reasonable first approximation. This approach must ultimately be refined if used in efforts to delineate more subtle changes in ocean redox, or if applied to periods during which continental

configuration and/or bathymetry are better constrained, but we contend that it provides a much more realistic depiction of the sensitivity of Cr and Mo mass balance to perturbation than previous model treatments. Further work should focus on the development and implementation of more spatially explicit approaches for dealing with the effects of seafloor redox perturbation on biogeochemical cycling and isotope systematics, for example coupling efficient models of benthic diagenesis that can be forced by gridded domains (79) to Earth system models of intermediate complexity (e.g., GENIE; ref. 80).

Prescribed Perturbations in the Model and the Role of Reducing Sediments. As discussed above, our model analysis involves balancing the modern steady-state cycles of Cr and Mo, applying a continuous range of perturbations to seafloor redox state, and establishing the ultimate steady-state conditions and local burial rates attained by the model system. Because our model includes a representation of offshore decreases in authigenic burial rates, essentially a spatial component, we must make some explicit assumptions about the basic seafloor environments in which perturbations begin and expand.

We assume first that $\sim 5\%$ of the shallow seafloor remains essentially authigenically neutral unless it becomes absolutely necessary to encroach upon this area (i.e., above 95% seafloor anoxia or euxinia). This assumption is meant to encompass coastal sediments deposited within the well-oxygenated mixed layer of the ocean. In addition, we assume that if atmospheric oxygen levels are low enough such that large portions of the oceanic mixed layer are anoxic, then the vast majority of the ocean will almost certainly be anoxic as well, effectively rendering the exercise moot and making our conclusions with respect to anoxia rather self-evident without changing our conclusions with respect to euxinia. Perturbations are then applied by expanding a given redox state (anoxic or euxinic) from the shallow shelf out into the deep sea. An important corollary of this approach is that during a given perturbation the first environments to become authigenically active are characterized by the highest metal burial rates. We view this as generally justifiable on mechanistic grounds, as elevated carbon fluxes through the water column and water column oxidant depletion are most commonly seen along ocean margins.

However, it is important to entertain the possibility that the nature of perturbations to seafloor redox may not be the same for Cr and Mo. For example, given that deep-sea anoxia during the mid-Proterozoic was most likely caused by gas-exchange constraints expressed in deep-water formation regions (81), rather than local reductant (i.e., carbon) input, it may be argued that large regions of the deep sea would first become anoxic and authigenically active for Cr, whereas euxinic environments driven by the combined effects of more reducing source waters and local carbon flux would be limited to marginal environments. In effect, this would result in less efficient Cr removal and similar Mo removal compared with the results presented here, which would in turn require a larger area of marine anoxia for a given Cr reservoir change. Thus, to remain conservative we prescribe that perturbations to both models begin and expand from settings where metal accumulation rates are highest. The reverse scenario, in which euxinia develops first in the abyssal realm of the ocean but anoxia is confined to the shelves, is difficult to imagine simply because of regional variability in carbon flux.

Another important assumption that is made in our modeling exercise is that the seafloor area of reducing sediments is fixed at a constant value (\sim 1.9% for the Mo model and \sim 4.9% for the Cr model). However, it is reasonable to expect that if the ocean becomes less oxygenated on a global scale there should be a firstorder expansion of reducing sediment environments. With respect to our basic conclusions, it is clear that this is a much larger concern for Cr than for Mo. Expanding the reducing sediment sink in the Mo model would only serve to decrease the extent of euxinic seafloor inferred for a given calculated enrichment. In other words, our interpretation that the mid-Proterozoic Mo enrichment record in euxinic shales implies relatively limited euxinic seafloor is rendered conservative by neglecting the expansion of reducing sediments in the model, and our conclusion that euxinia represents a small relative fraction of overall anoxia will remain unchanged. In the case of Cr, it might be argued that expansion of the reducing sediment sink together with expanding anoxia could result in our model significantly overestimating the amount of anoxic seafloor necessary to drive shifts in the seawater Cr reservoir. We consider this a problem that is somewhat scale-dependent, and that it is unlikely to change our fundamental conclusions for two reasons. First, in our model experiments we have artificially enhanced the impact of the reducing sediment sink, by choosing a relatively high Cr burial rate and by specifying that this burial rate applies to $\sim 5\%$ of the seafloor in addition to the expansion of anoxia in shelf environments. When we consider that the offshore scaling of metal burial rates (Fig. S4) should equally well apply to reducing sediment environments, this essentially amounts to "double counting" ~5% of the seafloor as being both reducing sediments and anoxic, with the highest metal burial rates specified for both. Alternatively, we can envision this as representing the exclusive expansion of anoxia in marginal settings, whereas $\sim 5\%$ of the seafloor offshore is covered by reducing sediments with unrealistically high burial rates. In either case, alternative approaches would need to either expand the reducing sediment sink at the expense of anoxic environments on the shelf, or expand it within the deep sea where metal burial capacity decreases sharply (Fig. S4). Both approaches would yield a comparable (and in some cases smaller) overall removal flux into reducing sediment environments.

Lastly, it is unlikely for very large regions of the seafloor to be characterized by this type of chemical environment at steady-state timescales. Such systems can be driven by sharp redox gradients, as often occurs in modern continental margin settings, but these environments require a rather unique combination of extremely high organic matter (i.e., reductant) loading and a degree of bottom-water ventilation such that the system does not become truly anoxic. Such sharp redox gradients will be difficult to maintain across large regions of the deep sea as a result of generally attenuated local organic carbon flux to offshore sediments. Alternatively, such a system can be driven by oxidant limitation. However, it is difficult to imagine this kind of system persisting on an extremely large scale, as it is poorly redox-buffered-small changes to circulation or carbon flux will result in the development of either true anoxia or increased bottom water O₂ such that the environment becomes effectively oxic with respect to metal burial.

Using the Model to Calculate Authigenic Metal Enrichments. From a qualitative perspective, it is difficult to avoid the conclusion that the coupled enrichment records require much more pervasive anoxia than that implied by equivalent Phanerozoic settings, but also that the relative fraction of anoxia represented by sulfidic deposition was not large. Our attempt to place more quantitative constraints on this conceptual interpretation involves using the scaling between seawater reservoir size and metal burial rates (inherent in a first-order mass balance model) to estimate sedimentary enrichments by assuming a bulk MAR in a hypothetical siliciclastic-dominated continental margin setting.

These two parameters (authigenic metal burial rate and bulk sediment MAR) should not be arbitrarily decoupled. This is true arithmetically, as metal burial rates in modern settings are in fact derived from bulk MARs. It is also expected on mechanistic grounds, as higher bulk MARs result in more rapid delivery of reactive mineral surfaces and organic carbon, and more rapid and efficient burial of authigenically sequestered elements. Indeed, there is good evidence from a range of modern (82) and ancient (83) settings that metal burial rates will scale in a general sense with bulk sediment MARs—in other words, metal enrichments will not simply scale linearly with changes in MAR. This issue is similar to that discussed above, in that an arbitrary decoupling of metal burial rates from bulk MARs is akin to applying a metal burial rate from a continental margin setting to the abyssal realm of the ocean, an approach we consider physically unrealistic.

Because this scaling between metal burial rate and bulk MAR is a somewhat broad relationship, we use well-constrained recent Cariaco Basin sediment data as a guide. We separate the range of Cr and Mo burial rates and bulk MAR values constrained for the Cariaco Basin over the last ~20,000 y, and sequentially combine them to explore the effect of a reasonable decoupling between these two parameters (Fig. S5). In an effort to render our estimates conservative, we choose combinations of metal burial rate

- 1. Lyons TW, Severmann S (2006) A critical look at iron paleoredox proxies: New insights from modern euxinic marine basins. *Geochim Cosmochim Acta* 70:5698–5722.
- Poulton SW, Fralick PW, Canfield DE (2004) The transition to a sulphidic ocean approximately 1.84 billion years ago. Nature 431(7005):173–177.
- Poulton SW, Canfield DE (2005) Development of a sequential extraction procedure for iron: implications for iron partitioning in continentally derived particulates. *Chem Geol* 214:209–221.
- Poulton SW, Canfield DE (2011) Ferruginous conditions: A dominant feature of the ocean through Earth's history. *Elements* 7:107–112.
- Canfield DE (1989) Reactive iron in marine sediments. Geochim Cosmochim Acta 53: 619–632.
- März C, et al. (2008) Redox sensitivity of P cycling during marine black shale formation: Dynamics of sulfidic and anoxic, non-sulfidic bottom waters. *Geochim Cosmochim Acta* 72:3703–3717.
- Lyons TW, et al. (2009) Tracking euxinia in the ancient ocean: A multiproxy perspective and Proterozoic case study. Ann Rev Earth Planet Sci Lett 37:507–534.
- Raiswell R, Buckley R, Berner RA, Anderson TF (1988) Degree of pyritization of iron as a paleoenvironmental indicator of bottom-water oxygenation. J Sed Petr 58:812–819.
- Algeo TJ, Maynard JB (2008) Trace-metal covariation as a guide to water-mass conditions in ancient anoxic marine environments. *Geosphere* 4:872–887.
- Kump LR, et al. (2011) Isotopic evidence for massive oxidation of organic matter following the great oxidation event. *Science* 334(6063):1694–1696.
- Planavsky NJ, Bekker A, Hofmann A, Owens JD, Lyons TW (2012) Sulfur record of rising and falling marine oxygen and sulfate levels during the Lomagundi event. Proc Natl Acad Sci USA 109(45):18300–18305.
- Scott C, et al. (2008) Tracing the stepwise oxygenation of the Proterozoic ocean. Nature 452(7186):456–459.
- Miller CA, Peucker-Ehrenbrink B, Walker BD, Marcantonio F (2011) Re-assessing the surface cycling of molybdenum and rhenium. *Geochim Cosmochim Acta* 75:7146–7179.
- D'Hondt S, et al. (2009) Subseafloor sedimentary life in the South Pacific Gyre. Proc Natl Acad Sci USA 106(28):11651–11656.
- Fischer JP, Ferdelman TG, D'Hondt S, Røy H, Wenzhöfer F (2009) Oxygen penetration deep into the sediment of the South Pacific gyre. *Biogeosciences* 6:1467–1478.
- Froelich PN, et al. (1979) Early oxidation of organic matter in pelagic sediments of the eastern equatorial Atlantic: Suboxic diagenesis. *Geochim Cosmochim Acta* 43: 1075–1090.
- Finney BP, Lyle MW, Heath GR (1988) Sedimentation at MANOP Site H (Eastern Equatorial Pacific) over the past 400,000 years: Climatically induced redox variations and their effects on transition metal cycling. *Paleoceanography* 3:169–189.
- Burdige DJ (1993) The biogeochemistry of manganese and iron reduction in marine sediments. *Earth Sci Rev* 35:249–284.
- Morford JL, Emerson S (1999) The geochemistry of redox sensitive trace metals in sediments. Geochim Cosmochim Acta 63:1735–1750.
- Meile C, Van Cappellen P (2003) Global estimates of enhanced solute transport in marine sediments. *Limnol Oceanogr* 48:777–786.
- Thullner M, Dale AW, Regnier P (2009) Global-scale quantification of mineralization pathways in marine sediments: A reaction-transport modeling approach. *Geochem Geophys Geosyst*, 10.1029/2009GC002484.
- 22. Bertine KK, Turekian KK (1973) Molybdenum in marine deposits. *Geochim Cosmochim Acta* 37:1415–1434.
- Bender ML, Ku T-L, Broecker WS (1970) Accumulation rates of manganese in pelagic sediments and nodules. *Earth Planet Sci Lett* 8:143–148.
- Shimmield GB, Price NB (1986) The behaviour of molybdenum and manganese during early sediment diagenesis – offshore Baja California, Mexico. Mar Chem 19:261–280.
- Brüchert V, et al. (2003) Regulation of bacterial sulfate reduction and hydrogen sulfide fluxes in the central Namibian coastal upwelling zone. *Geochim Cosmochim Acta* 67:4505–4518.
- Brüchert V, Currie B, Peard KR (2009) Hydrogen sulphide and methane emissions on the central Namibian shelf. Prog Oceanogr 83:169–179.
- 27. Piper DZ, Dean WE (2002) Trace element deposition in the Cariaco Basin, Venezuela shelf, under sulfate-reducing conditions a history of the local hydrography and global climate, 20 ka to the present, USGS Professional Paper, Vol 1670, 45 pp.
- Francois R (1988) A study on the regulation of the concentrations of some trace metals (Rb, Sr, Zn, Pb, Cu, V, Cr, Ni, Mn and Mo) in Saanich Inlet sediments, British Columbia, Canada. Mar Geol 83:285–308.

and bulk MAR that result in relatively low enrichments for Cr and relatively high enrichments for Mo, and these are presented in the main text (Fig. S5).

No single combination of parameters will be adequate to describe the entire shale record, but we view this broad range as sufficient to encompass the vast majority of environments represented in our database. It is apparent from this exercise that a very strong decoupling between metal burial rate and bulk MAR values, which we consider unrealistic, is necessary to invalidate our basic conclusions. Furthermore, this condition would need to pertain to every mid-Proterozoic anoxic shale in our database over ~1.5 Ga, whereas fortuitously being alleviated during the early Phanerozoic, a combination of circumstances that that would be improbable.

- Ravizza G, Turekian KK, Hay BJ (1991) The geochemistry of rhenium and osmium in recent sediments from the Black Sea. *Geochim Cosmochim Acta* 55:3741–3752.
- Algeo TJ (2004) Can marine anoxic events draw down the trace element inventory of seawater? Geology 32(12):1057–1060.
- Buesseler KO, Benitez CR (1994) Determination of mass accumulation rates and sediment radionuclide inventories in the deep Black Sea. Deep Sea Res Part I Oceanogr Res Pap 41:1605–1615.
- Dean WE, Piper DZ, Peterson LC (1999) Molybdenum accumulation in Cariaco basin sediment over the past 24 k.y.: A record of water-column anoxia and climate. *Geology* 27:507–510.
- Giraudeau J, Meyers PA, Christensen BA (2002) Accumulation of organic and inorganic carbon in Pliocene-Pleistocene sediments along the SW African margin. *Mar Geol* 180: 49–69.
- Lyons TW, Werne JP, Hollander DJ, Murray RW (2003) Contrasting sulfur geochemistry and Fe/Al and Mo/Al ratios across the last oxic-to-anoxic transition in the Cariaco Basin, Venezuela. Chem Geol 195:131–157.
- Borchers SL, Schnetger B, Böning P, Brumsack H-J (2005) Geochemical signatures of the Namibian diatom belt: Perennial upwelling and intermittent anoxia. Geochem Geophys Geosyst, 10.1029/2004GC000886.
- Algeo TJ, Lyons TW (2006) Mo-total organic carbon covariation in modern anoxic marine environments: Implications for analysis of paleoredox and paleohydrographic conditions. *Paleoceanography*, 10.1029/2004PA001112.
- McManus J, et al. (2006) Molybdenum and uranium geochemistry in continental margin sediments: Paleoproxy potential. *Geochim Cosmochim Acta* 70:4643–4662.
- Sifeddine A, et al. (2008) Laminated sediments from the central Peruvian continental slope: A 500 year record of upwelling system productivity, terrestrial runoff and redox conditions. Prog Oceanogr 79:190–197.
- Gutierrez D, et al. (2009) Rapid reorganization in ocean biogeochemistry off Peru towards the end of the Little Ice Age. *Biogeosciences* 6:835–848.
- Poulson Brucker RL, McManus J, Severmann S, Berelson WM (2009) Molybdenum behavior during early diagenesis: Insights from Mo isotopes. *Geochem Geophys Geosyst*, 10.1029/2008GC002180.
- Brumsack H-J (1989) Geochemistry of recent TOC-rich sediments from the Gulf of California and the Black Sea. Geol Rundsch 78:851–882.
- Zheng Y, Anderson RF, van Geen A, Kuwabara J (2000) Authigenic molybdenum formation in marine sediments: A link to pore water sulfide in the Santa Barbara Basin. *Geochim Cosmochim Acta* 64:4165–4178.
- Poulson RL, Siebert C, McManus J, Berelson WM (2006) Authigenic molybdenum isotope signatures in marine sediments. *Geology* 34:617–620.
- Peuker-Ehrenbrink B (2009) Land2Sea database of river drainage basin sizes, annual water discharges, and suspended sediment fluxes. *Geochem Geophys Geosyst*, 10.1029/2008GC002356.
- Cranston RE, Murray JW (1980) Chromium species in the Columbia River and estuary. Limnol Oceanogr 25:1104–1112.
- Abu-Saba KE, Flegal AR (1995) Chromium in San Francisco Bay: Superposition of geochemical processes causes complex spatial distributions of redox species. Mar Chem 49:189–199.
- Jeandel C, Minster JF (1987) Chromium behavior in the ocean: Global versus regional processes. Global Biogeochem Cycles 1:131–154.
- van der Weijden CH, Reith M (1982) Chromium(III) chromium(VI) interconversions in seawater. Mar Chem 11:565–572.
- Ellis AS, Johnson TM, Bullen TD (2004) Using chromium stable isotope ratios to quantify Cr(VI) reduction: Lack of sorption effects. *Environ Sci Technol* 38(13): 3604–3607.
- Smith E, Ghiassi K (2006) Chromate removal by an iron sorbent: Mechanism and modeling. Water Environ Res 78(1):84–93.
- Zachara JM, Girvin DC, Schmidt RL, Resch CT (1987) Chromate adsorption on amorphous iron oxyhydroxide in the presence of major groundwater ions. *Environ Sci Technol* 21(6):589–594.
- Zachara JM, Cowan CE, Schmidt RL, Ainsworth CC (1988) Chromate adsorption by kaolinite. *Clays Clay Miner* 36:317–326.
- 53. Turekian KK, Imbrie J (1966) The distribution of trace elements in deep-sea sediments of the Atlantic Ocean. *Earth Planet Sci Lett* 1:161–168.
- Chester R, Hughes MJ (1969) The trace element geochemistry of a North Pacific pelagic clay core. *Deep-Sea Res* 16:639–654.

- Cronan DS (1969) Average abundances of Mn, Fe, Ni, Co, Cu, Pb, Mo, V, Cr, Ti and P in Pacific pelagic clays. Geochim Cosmochim Acta 33:1562–1565.
- Piper DZ, Dean WE (2002) Trace-element deposition in the Cariaco Basin under sulfate reducing conditions—a history of the local hydrography and global climate, 20 ka to present. USGS Professional Paper (US Geological Survey, Denver), Vol 1670, 41 pp.
- 57. Dean WE (2007) Sediment geochemical records of productivity and oxygen depletion along the margin of western North America during the past 60,000 years: Teleconnections with Greenland Ice and the Cariaco Basin. Q Sci Rev 26:98–114.
- Eary LE, Rai D (1989) Kinetics of chromate reduction by ferrous ions derived from hematite and biotite at 25°C. Am J Sci 289:180–213.
- Fendorf SE, Li G (1996) Kinetics of chromate reduction by ferrous iron. Environ Sci Technol 30:1614–1617.
- Bond DL, Fendorf S (2003) Kinetics and structural constraints of chromate reduction by green rusts. *Environ Sci Technol* 37(12):2750–2757.
- Rue EL, Smith GJ, Cutter GA, Bruland KW (1997) The response of trace element redox couples to suboxic conditions in the water column. *Deep Sea Res Part I Oceanogr Res Pap* 44:113–134.
- Brumsack H-J, Gieskes JM (1983) Interstitial water trace-metal chemistry of laminated sediments from the Gulf of California, Mexico. Mar Chem 14:89–106.
- 63. Calvert SE (1966) Accumulation of diatomaceous silica in sediments of the Gulf of California. *Bull Geol Soc Amer* 77:569–596.
- Elderfield H, Schultz A (1996) Mid-ocean ridge hydrothermal fluxes and the chemical composition of the ocean. Annu Rev Earth Planet Sci 24:191–224.
- Berner RA (2004) A model for calcium, magnesium and sulfate in seawater over Phanerozoic time. Am J Sci 304:438–453.
- Seyfried WE, Jr., Mottle MJ (1982) Hydrothermal alteration of basalt by seawater under seawater-dominated conditions. *Geochim Cosmochim Acta* 46:985–1002.
- Von Damm KL, Edmond JM, Measures CI, Grant B (1985) Chemistry of submarine hydrothermal solutions at Guymas Basin, Gulf of California. *Geochim Cosmochim* Acta 49:2221–2237.
- Wheat CG, Mottl MJ, Rudnicki M (2002) Trace element and REE composition of a low-temperature ridge-flank hydrothermal spring. *Geochim Cosmochim Acta* 66: 3693–3705.
- Emerson S, Hedges J (2008) Chemical Oceanography and the Marine Carbon Cycle (Cambridge Univ Press, Cambridge, UK).

- Michalopoulos P, Aller RC (1995) Rapid clay mineral formation in Amazon Delta sediments: Reverse weathering and oceanic elemental cycles. Science 270:614–617.
- Jahnke RA (1996) The global ocean flux of particulate organic carbon: Areal distribution and magnitude. *Global Biogeochem Cycles* 10:71–88.
- Middelburg JJ, Soetaert K, Herman PMJ (1997) Empirical relationships for use in global diagenetic models. Deep Sea Res Part I Oceanogr Res Pap 44:327–344.
- Sarmiento JL, Gruber N (2006) Ocean Biogeochemical Dynamics (Princeton Univ Press, Princeton).
- Dunne JP, Sarmiento JL, Gnanadesikan A (2007) A synthesis of global particle export from the surface ocean and cycling through the ocean interior and on the seafloor. *Global Biogeochem Cycles* 21:10.1029/2006GB002907.
- Dahl TW, et al. (2010) Devonian rise in atmospheric oxygen correlated to the radiations of terrestrial plants and large predatory fish. Proc Natl Acad Sci USA 107(42):17911–17915.
- Dahl TW, et al. (2011) Molybdenum evidence for expansive sulfidic water masses in ~750 Ma oceans. Earth Planet Sci Lett 311:264–274.
- Middelburg JJ, Soetaert K, Herman PMJ, Heip CHR (1996) Denitrification in marine sediments: A model study. Global Biogeochem Cycles 10:661–673.
- Menard HW, Smith SM (1966) Hypsometry of ocean basin provinces. J Geophys Res 71: 4305–4325.
- Archer DE, Morford JL, Emerson SR (2002) A model of suboxic sedimentary diagenesis suitable for automatic tuning and gridded global domains. *Global Biogeochem Cycles* 16:10.1029/2000GB001288.
- Ridgwell A, et al. (2007) Marine geochemical data assimilation in an efficient Earth System Model of global biogeochemical cycling. *Biogeosciences* 4:87–104.
- Canfield DE (1998) A new model for Proterozoic ocean chemistry. Nature 396: 450–453.
- Meyers SR, Sageman BB, Lyons TW (2005) Organic carbon burial rate and the molybdenum proxy: Theoretical framework and application to Cenomanian-Turonian oceanic anoxic event 2. *Paleoceanography* 20:10.1029/2004PA001068.
- Hetzel A, Böttcher ME, Wortmann UG, Brumsack H-J (2009) Paleo-redox conditions during OAE 2 reflected in Demerara Rise sediment geochemistry (ODP Leg 207). *Palaeogeogr Palaeoclimatol Palaeoecol* 273:302–328.



Fig. S1. Complete database for Cr (A) and Mo (B) enrichments through time, filtered according to protocols discussed in the text. Shaded boxes denote the range of upper modern crustal concentrations.



Fig. S2. Schematic of the global Cr (*A*) and Mo (*B*) mass balance models, showing the modern balanced state. Average modern seawater concentrations ($[Cr]_M$ and $[Mo]_M$), reservoir sizes (R_{Cr} and R_{Mo}), and residence times (τ_{res}) are shown. Dominant authigenic removal processes are depicted schematically for each sink [oxic (ox), anoxic (*anox*), reducing (*red*), and sulfidic (*sulf*) sinks]. C_{org} represents organic carbon, whereas terms shown as $S\{x-y\}$ denote sorption processes. Hydrothermal fluxes are neglected in our treatment here (see *SI Discussion* for further details). Fluxes are in 10⁷ mol·y⁻¹.



Fig. S3. Effect of changes in the assumed input flux on model results. A-C apply to the Cr model. Each curve shows the areal extent of seafloor anoxia (A_{anox}) at which authigenic Cr enrichments decrease below 5 μ g·g⁻¹. Each panel represents a different bulk MAR (increasing from left to right, with a range of a factor of 1.5 around 1.0×10^{-2} g·cm⁻²·y⁻¹, the approximate value for deep sediments of the modern Cariaco Basin), whereas the curves represent low (dashed gray; 0.5μ g·cm⁻²·y⁻¹), medium (black; 0.75μ g·cm⁻²·y⁻¹) and high (dashed gray; 1.0μ g·cm⁻²·y⁻¹) authigenic Cr burial rates. D-F apply to the Mo model, with each curve showing the areal extent of sulfidic (euxinic) seafloor (A_{sulf}) at which authigenic Cr enrichments decrease below 40 μ g·g⁻¹ (the mid-Proterozoic total mean). The range of bulk MAR values is the same as in A-C, with the various curves in each panel representing low (dashed gray; 1.0μ g·cm⁻²·y⁻¹), medium (black; 1.5μ g·cm⁻²·y⁻¹), and high (dashed gray; 2.0μ g·cm⁻²·y⁻¹) authigenic Mo burial rates.



Fig. S4. Parametrization of offshore metal burial rate scaling in the model. Globally averaged authigenic Cr burial rates are shown in *A* as a function of anoxic seafloor area (A_{anox}). The blue filled circle represents the modern state, with the blue dotted line depicting a constant burial rate decoupled from the extent of anoxic seafloor area. Globally averaged authigenic Mo burial rates are shown in *B* as a function of sulfidic (euxinic) seafloor area (A_{sulf}). The red filled circle represents the modern state, with the red dotted line depicting a constant burial rate decoupled from the extent of sulfidic (euxinic) seafloor area (A_{sulf}). The red filled circle represents the modern state, with the red dotted line depicting a constant burial rate decoupled from the extent of euxinic seafloor area (essentially the models used in refs. 1–4).

- 1. Scott C, et al. (2008) Tracing the stepwise oxygenation of the Proterozoic ocean. Nature 452(7186):456-459.
- Dahl TW, et al. (2010) Devonian rise in atmospheric oxygen correlated to the radiations of terrestrial plants and large predatory fish. Proc Natl Acad Sci USA 107(42):17911–17915.
 Dahl TW, et al. (2011) Molybdenum evidence for expansive sulfidic water masses in ~750 Ma oceans. Earth Planet Sci Lett 311:264–274.
- 4. Kendall B, Creaser RA, Gordon GW, Anbar AD (2009) Re-Os and Mo isotope systematics of black shales from the Middle Proterozoic Velkerri and Wollogorang Formations, McArthur Basin, northern Australia. Geochim Cosmochim Acta 73:2534–2558.



Fig. S5. Range of metal burial rates and bulk sediment MAR explored in the model. *A*, *C*, and *E* depict estimated authigenic Cr enrichments as a function of anoxic seafloor area at a range of plausible metal burial rates and bulk sediment MAR. The blue box represents our conservative threshold for Cr enrichment as constrained by the shale record. *B*, *D*, and *F* depict estimated authigenic Mo enrichments as a function of sulfidic (euxinic) seafloor area at a range of plausible metal burial rates and bulk sediment MAR. The blue box represents our conservative threshold for Cr enrichment as constrained by the shale record. *B*, *D*, and *F* depict estimated authigenic Mo enrichments as a function of sulfidic (euxinic) seafloor area at a range of plausible metal burial rates and bulk sediment MAR. The red box represents the 95% confidence interval around the overall mean for the mid-Proterozoic shale enrichment record. Each panel represents a different bulk MAR (increasing from top to bottom, and depicting a range of a factor of 1.5 around $1.0 \times 10^{-2} \text{ g-cm}^{-2} \text{ y}^{-1}$, the approximate value for deep sediments of the modern Cariaco Basin) and the contours are labeled by authigenic metal burial rate (in $\mu \text{g-cm}^{-2} \text{ y}^{-1}$). The solid black curves are those depicted in the main text.

Other Supporting Information Files

Table S1 (DOCX) Table S2 (DOCX) Table S3 (DOCX) Table S4 (DOCX) Table S5 (DOCX) Table S6 (DOCX)