Strong evidence for a weakly oxygenated ocean–atmosphere system during the Proterozoic

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Earth’s surface has undergone a protracted oxygenation, which is commonly assumed to have profoundly affected the biosphere. However, basic aspects of this history are still debated—foremost oxygen (O₂) levels in the oceans and atmosphere during the billion years leading up to the rise of algae and animals. Here we use isotopic ratios of iron (Fe) in ironstones—Fe-rich sedimentary rocks deposited in nearshore marine settings—as a proxy for O₂ levels in shallow seawater. We show that partial oxidation of dissolved Fe(II) was characteristic of Proterozoic shallow marine environments, whereas younger ironstones formed via complete oxidation of Fe(II). Regardless of the Fe(II) source, partial Fe(II) oxidation requires low O₂ in the shallow oceans, settings crucial to eukaryotic evolution. Low O₂ in surface waters can be linked to markedly low atmospheric O₂—likely requiring less than 1% of modern levels. Based on our records, these conditions persisted (at least periodically) until a shift toward higher surface O₂ levels between ca. 900 and 750 Ma, coincident with an apparent rise in eukaryotic ecosystem complexity. This supports the case that a first-order shift in surface O₂ levels during this interval may have selected for life modes adapted to more oxygenated environments.

Iron (Fe) in seawater is soluble under anoxic conditions [as Fe(II)] and poorly soluble under oxic conditions [precipitating as Fe(III) oxyhydroxides]. Although oxidation of Fe(II) can also occur under anoxic conditions (via photo-oxidation or anoxicogenic photosynthesis; ref. 13), the rate and extent of Fe(II) oxidation in aqueous environments—either abiogenic or biologically mediated—will depend greatly on the abundance of O₂. Therefore, secular records of marine Fe(II) oxidation can potentially be used to track O₂ levels in seawater. At the low surface O₂ levels characteristic of early Earth, marine Fe(II) availability would have been comparatively high (sourced from hydrothermal inputs and the redox cycling of continental Fe), and partial oxidation of this seawater Fe(II) reservoir would have been common (14, 15). At higher O₂ levels characteristic of the modern oceans and atmosphere, any Fe(II) inputs to surface waters are quantitatively oxidized (16).

Based upon the kinetics of Fe(II) oxidation and the associated Fe isotope systematics (16–18), it is possible to estimate the abundance of O₂ in seawater based upon the Fe isotope irontone |

Significance

Earth’s transition from anoxic oceans and atmosphere to a well-oxygenated state led to major changes in nearly every surficial system. However, estimates of surface oxygen levels in the billion years preceding this shift span two orders of magnitude, suggesting a poor understanding of the evolution of the oxygen cycle. We use the isotopic record of iron oxides deposited in ancient shallow marine environments to show that oxygen remained at extremely low levels in the ocean–atmosphere system for most of Earth’s history, and that a rise in oxygen occurred in step with the expansion of complex, eukaryotic ecosystems. These results indicate that Earth is capable of stabilizing at low atmospheric oxygen levels, with important implications for exploration of exoplanet biosignatures.

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composition of chemical sediments (14). The partial oxidation of dissolved Fe(II) can lead to equilibrium fractionation whereby the produced Fe(III) (oxyhydr)oxides are enriched in $^{56}\text{Fe}$ by up to 1.0 to 3.2$\%$ (reported as the ratio of $^{56}\text{Fe}/^{54}\text{Fe}$ relative to the IRMM-014 standard reference material in parts per thousand; $\delta^{56}\text{Fe}$) (17, 18). More rapid oxidation effectively lowers this overall fractionation (14, 18), and quantitative oxidation produces Fe(III) oxyhydroxides with an Fe isotope composition that mirrors that of the Fe(II) source. Biologically mediated Fe(II) oxidation (facilitated by anoxic or anaerobic photosynthetic bacteria under anoxic conditions or microaerophilic chemotrophs under suboxic conditions) results in similar Fe isotope fractionations to that of abiotic Fe(II) oxidation (18) (SI Appendix). Compared to paleoredox proxies based on trace element geochemistry, the Fe is isotopic composition of Fe-rich chemical sedimentary rocks is expected to be strongly rock-buffered, less likely to be affected by detrital contamination, and resistant to post depositional alteration. Therefore, Fe-rich rocks can serve as robust Fe isotope archives in deep time (14, 18, 19) and can be used to constrain the redox state of ancient seawater.

**The Ironstone Record of Shallow Marine Redox Conditions**

The Proterozoic oceans were likely to have been characterized by a strong redox depth gradient, with anoxic and Fe(II)-bearing (ferruginous) deep ocean waters overlain by O$_2$-bearing surface waters in contact with an oxygenated atmosphere, with anoxic and H$_2$S-bearing (euxinic) zones variably developed at mid-depths along productive continental margins (2). Constraining the O$_2$ concentrations ($[O_2]$) of this oxygenated surface layer is essential to understanding the evolution of early aerobic eukaryotes. Iron isotope studies of Fe-rich sedimentary rocks can offer important insights into ancient marine redox conditions. However, most of these studies have focused on banded iron formations, which are generally interpreted to have been deposited in deeper (below wave base) marine settings that are temporaneous local euxinia (19, 23, 24) (SI Appendix). The Fe isotope composition of hydrothermal Fe(II) and igneous rocks covers a limited range around $0_{\%}$, (typically $-0.5 < \delta^{56}\text{Fe} < 0.3_{\%}$), whereas Fe(II) sourced from Fe(III) reduction is generally characterized by negative $\delta^{56}\text{Fe}$ values (18, 28). Fractionation beyond these input values can therefore implicate genetic processes. The studied Phanerozoic ironstones and Holocene ferruginous shallow marine sediments that formed under a well-oxygenated ocean–atmosphere system.

**Ironstone Iron Isotope Compositions**

Ironstones comprise authigenic precipitates that require an abundant supply of mobile Fe(II) for their genesis, which can be supplied by anoxic seawater enriched in Fe(II) from distal hydrothermal sources (19, 21, 26) or a benthic Fe(II) flux sourced from the reduction of continental Fe(III) (e.g., ref. 27). The Fe isotopic composition of hydrothermal Fe(II) and igneous rocks covers a limited range around $0_{\%}$, which are indistinguishable from those of hydrothermal and igneous Fe(II) sources (Fig. 2). The results show a statistically significant difference between the Proterozoic and Phanerozoic ironstones. In contrast to the younger equivalents, nearly all of the studied Phanerozoic ironstones have highly variable and mostly positive $\delta^{56}\text{Fe}$ values (Fig. 2). The exception to this rule is the youngest of the Proterozoic ironstones in our dataset—the ca. 750 Ma Galeros Formation in Arizona—which displays a notably limited range in $\delta^{56}\text{Fe}$ values (Fig. 2) that are not statistically different from those of Phanerozoic ironstones.

**An Isotopic Fingerprint of Marine Partial Fe(II) Oxidation**

The Fe isotopic composition of Fe(III) oxide-bearing sediments is determined by several factors, including the Fe(II) source, the rates of Fe(II) supply and oxidation, and the relative proportion of authigenic Fe(III) (oxyhydr)oxides compared to the total Fe contents. In well-oxygenated surface waters, the oxidation of seawater Fe(II) goes to completion, resulting in a narrow range in $\delta^{56}\text{Fe}$ values that reflects that of the Fe(II) source (likely close to $0_{\%}$) (16, 17). The much greater spread in Fe isotope compositions in the Proterozoic ironstones (relative to the Phanerozoic ironstones), including positive and negative $\delta^{56}\text{Fe}$ values, is best explained by distillation processes involving partial Fe(II) oxidation.

Although negative $\delta^{56}\text{Fe}$ values in ironstones could indicate that the local seawater was enriched in isotopically light Fe(II) produced via distillation Fe(III) reduction (17, 18, 28), this
signature could also result from the ongoing removal of isotopically heavy Fe via the progressive partial oxidation of a seawater Fe(II) reservoir (i.e., isotopic distillation) (15, 18). When Fe(II) oxidation is extensive, the $\delta^{56}$Fe values of the resultant Fe(III) (oxyhydr)oxides can follow a predictable distribution that is weighted toward higher $\delta^{56}$Fe values (approximating the instantaneous product of Fe(II) oxidation; SI Appendix), with a gradual trend of decreasing relative frequency toward lower $\delta^{56}$Fe values. This trend is most clearly evident in the data from the ca. 1.650 Ma Chuanlinggou Formation ironstone (North China), for which $\delta^{56}$Fe values are most commonly between 0.6 and 1.0‰, with more negative values becoming progressively less common (Fig. 3A). Assuming an initial seawater Fe(II) with $\delta^{56}$Fe values close to 0‰, this may suggest an instantaneous fractionation of $-0.8$‰ associated with Fe(II) oxidation and the precipitation of ironstone Fe(III) (oxyhydr)oxides in the low $O_2$ Proterozoic surface waters. The interpretation that the Fe isotope signature of the Chuanlinggou ironstone is controlled by partial oxidation is strongly supported by a negative correlation between $\delta^{56}$Fe values and Mn enrichment (Mn/Fe ratios; Fig. 3B), which are typically considered a proxy for marine oxidation (29).

Most of the other Proterozoic ironstones do not show this full distribution ranging from positive to negative $\delta^{56}$Fe values and generally show a spread in $\delta^{56}$Fe values from $\sim$0 to 1‰ (Fig. 2). This may suggest that Fe(II) oxidation in the waters sourcing these ironstones was not as extensive during their formation (Fig. 3). Local and regional variation in the $\delta^{56}$Fe values of individual ironstone samples is an expected consequence when partial oxidation dominates, as there can be temporal and spatial variability in the extent of oxidation of different ferruginous water masses. Under a low $pO_2$ atmosphere, $[O_2]$ in shallow seawater can be locally elevated due to local primary production (30) but is unlikely to be depleted relative to gas-exchange equilibrium with the overlying atmosphere (SI Appendix, Fig. S2).

Further, upwelling rates and initial dissolved Fe(II) concentrations can be variable, leading to different extents of Fe(II) oxidation at low $O_2$ levels. The spread in the isotopic composition of these ironstones (including negative and positive $\delta^{56}$Fe values) is consistent with partial oxidation of an Fe(II) source with $\delta^{56}$Fe values close to 0‰ (18).

These Fe isotope trends are suggestive of a high dissolved Fe(II):$O_2$ ratio of the contemporaneous shallow seawater. However, in order to attempt to leverage these Fe isotope data to obtain estimates for surface $O_2$ levels, it is important to consider other possible influences on the ironstone Fe isotope signatures such as variation in the isotopic composition of the Fe(II) source. In modern, oxic river waters and surface seawater, the stabilization in solution of otherwise highly insoluble Fe(III) by complexation with organic ligands can lead to dissolved Fe(III) with positive $\delta^{56}$Fe values (e.g., refs. 31 and 32). However, this process has not been shown to result in isotopically heavy Fe in the marginal sediments (31, 33), and the bulk isotopic composition of Fe in rivers and estuarine sediments is typically close to (or slightly lighter than) average crustal values (31, 34). Given that organic ligands stabilize Fe(III) in solution, we suggest that ligand-bound Fe(III) did not play a major role in the formation of ironstones.

Kinetic isotope effects during the precipitation of Fe(II) sulfides from dissolved Fe(II) can involve negative fractionation, and this is expressed most strongly during partial pyrite formation when $H_2S$ availability is limited (35). Iron isotopic fractionation related to pyrite formation has been shown to be an important process in modern euxinic basins and may have been active in localized euxinic water masses during the Proterozoic (36). However, the possible effects of this process on the isotopic composition of marine Fe(II) would have depended upon the relative availability of Fe(II) and $H_2S$ in the mid-depth euxinic zone, where developed. If ferruginous seawater interacted with euxinic water masses in the middle Proterozoic, $H_2S$
would likely have been in excess and seawater Fe(II) would have been depleted due to quantitatively pyrite formation, producing sedimentary pyrite that largely matches the δ56Fe values of the original Fe(II) source (36). In this scenario, isotopically heavy Fe(II) is not supplied to shallow waters due to quantitatively drawdown near the redoxcline, and therefore this process would not affect the isotopic composition of nearshore sediments. These Fe–S systematics are demonstrated by data from modern euxinic basins where sedimentary pyrite is not strongly fractionated (28) and seawater Fe(II) near or above the redoxcline is not 56Fe-enriched (36).

For sedimentary pyrite deposited after the Great Oxidation Event, δ56Fe values are typically unfractionated or positive (15, 17) (SI Appendix, Fig. S7). This may suggest that fractionation and isotopic distillation due to partial pyrite formation was not an important process in the middle Proterozoic (SI Appendix). More data from this interval are needed, and investigations into the triple Fe isotope composition of Proterozoic Fe(II) sulfides and Fe(III) oxides may help to better elucidate the importance of the sulfide Fe sink during this time (37). In sum, the sedimentary Fe record is more consistent with a partial oxidation control or local Fe cycling, the positive δ56Fe values in our data can only be explained by the partial oxidation of Fe(II) due to a high Fe(II)/O2 ratio (e.g., refs. 14, 17, 18).

Low Surface O2 during the Middle Proterozoic

In modern redox-stratified lakes with anoxic and Fe(II)-rich bottom waters, considered an analog for the low-O2 oceans of the early Earth, surface waters are oxygenated due to gas exchange with the atmosphere. In these settings, upwelling Fe(II) from the ferruginous water column is completely and rapidly oxidized, and the resultant Fe(III) oxohydroxides are isotopically unfractionated relative to the dissolved Fe(II) pool (16). Therefore, partial Fe(II) oxidation in the upper water column requires lower [O2] than present levels. By combining our Fe isotope data with a model for Fe(II) oxidation kinetics, we can use ironstones to estimate the ancient shallow seawater [O2].
Ironstone samples with $\delta^{56}$Fe values that fall within the range of igneous and hydrothermal Fe(II) sources could be consistent with quantitative Fe(II) oxidation. For ironstone samples that are fractionated beyond the relatively narrow range of $\delta^{56}$Fe values of possible input values, we can use a distillation model to invert each ironstone $\delta^{56}$Fe value and estimate the fraction of the Fe(II) reservoir that was oxidized ($f_{\text{ox}}$; Fig. 4A and SI Appendix). This process is repeated using different values for the assumed combined isotope fractionation effect for Fe(II) oxidation and precipitation in shallow seawater. Following this, we can compare these results for the extent of Fe(II) oxidation to those obtained using a probabilistic model of Fe(II) oxidation kinetics that takes into account a wide range of possible marine pH values, surface water temperatures, and variable residence times of water masses within the shallow ocean (Table 1 and SI Appendix, Fig. S1). We run this model at a range of seawater $[O_2]$ and utilize the results from a Lagrangian model embedded within a general circulation model (12) to approximate a surface water residence time. This produces a statistical distribution of the values for $f_{\text{ox}}$ obtained for each $\delta^{56}$Fe data point; these results are shown in Fig. 4. Based on the results of this modeling approach, we find that the signatures observed in Proterozoic ironstones are most compatible with shallow seawater $[O_2]$ below $\sim 5 \mu$mol/kg (Fig. 4B). At these low $O_2$ levels, some proportion of the Fe(II) oxidation is likely to have been mediated by microaerophilic chemotrophic bacteria, which would lead to isotopic fractionation similar to that of abiotic oxidation (18). Given that microbial rates of Fe(II) oxidation at low $[O_2]$ are higher than abiotic oxidation rates (38), our $[O_2]$ estimate of $< 5 \mu$mol/kg is likely to be conservatively high. This is evident when comparing our model constraints to the much lower estimates for shallow water $[O_2]$ that would be obtained using a dispersion–reaction modeling approach that has been used in other Fe isotope studies (14, 37). Such low surface water $[O_2]$—as indicated by the maximum constraints provided by our model—would be expected to have exerted a powerful control on the ecology of even the shallowest marine environments during this time (39, 40).

Surface seawater $[O_2]$ is linked to atmospheric $pO_2$ via air–sea gas exchange. The Fe(III) (oxy)hydroxide precursor minerals of the studied ironstones were precipitated from shallow seawater (above storm wave base). In modern marine environments, such settings are well mixed and unlikely to be deficient in dissolved $O_2$ relative to equilibrium with the overlying oxygenated atmosphere, except in cases of extreme eutrophy (41). In the mid-Proterozoic, the flux of $O_2$ supplied by air–sea gas exchange at $pO_2 \geq 1\%$ of the present atmospheric level (PAL) is unlikely to have been exhausted by upwelling ferruginous seawater, given estimates of dissolved Fe(II) concentrations and assuming reasonable upwelling rates (SI Appendix, Fig. S2; calculations in SI Appendix). Therefore, the predominance of Fe isotopic evidence for low surface water $[O_2]$ from ironstones, deposited in a range of paleocontinents and time periods in the mid-Proterozoic, is extremely difficult to explain unless atmospheric $pO_2$ was low during for at least intervals of the Proterozoic. Assuming gas-exchange equilibrium, with solubility corrections for temperature and salinity (42), our results strongly suggest that atmospheric $pO_2$ reached levels below 1% PAL (Fig. 4B), at least during the sampled intervals throughout the mid-Proterozoic. These results are supported by other geological evidence, such as mid-Proterozoic paleosols that suggest Fe loss during weathering, which is also consistent with a low-$O_2$ atmosphere (43). Regardless of whether low shallow water $[O_2]$
might have been toxic to the earliest animal life, low $O_2$ levels in the ocean–atmosphere system would have exerted an important limitation on marine nutrient levels, primary productivity, and the potential for oceans to sustain complex, eukaryote-rich ecosystems (44). As such, new, robust evidence for the presence of low surface $O_2$ levels bolsters the case that oxygenation may have played a significant role in constraining the structure and energetic scope of complex life in the Proterozoic biosphere (3, 45).

Late Proterozoic Oxygenation and the Rise of Eukaryotes

The hypothesis that there was a mechanistic link between a rise in atmospheric $O_2$ and the emergence of animals toward the end of the Proterozoic (e.g., ref. 3) remains debated. Specifically, an $O_2$ control on eukaryotic diversity has recently been challenged on the grounds that the $O_2$ requirements of primitive animals were exceeded well before their evolution (7–9). Much of the debate stems from uncertainties associated with the assumptions used in various paleoredox approaches to translate geochemical data from sedimentary rocks into $O_2$ estimates (5, 9, 10), leading to estimates that span over an order of magnitude (e.g., refs. 4, 5, 7, 8, 43).

For instance, the chromium (Cr) isotope proxy system has been used to argue for a low atmospheric $O_2$ (i.e., 0.1 to 1.0% PAL) (4, 46) as well as higher levels (>1 to 10% PAL) (7, 10). This proxy links the Cr isotope composition of marine sediments to atmospheric $O_2$, whereby the oxidative weathering of terrestrial rocks supplies fractionated Cr to the marine realm (e.g., ref. 46). Nonredox-dependent and marine processes have also been shown to fractionate Cr, and the interpretation of atmospheric $O_2$ based upon Cr isotope data may yield multiple, nonunique solutions (47, 48). Therefore, the Cr sedimentary isotope record tracks a complex history of Cr weathering, transport, and burial and compared to the Fe isotope proxy is less likely to represent changes in shallow $O_2$ critical to eukaryotic evolution. Further, the Cr isotope $O_2$ proxy relies on the preservation of a trace element which may be overprinted during diagenesis and alteration (49, 50).

Another approach that has been used to estimate mid-Proterozoic paleoredox conditions involves the abundance of cerium (Ce) in marine carbonate rocks (5, 6, 12), as Ce is preferentially scavenged from seawater underoxic conditions (51). However, poorly constrained Ce oxidation kinetics and debate over the oxidative pathway (e.g., refs. 51 and 52) present a challenge for attempts to quantitatively link Ce anomaly records to $O_2$ levels, and different modeling approaches have led to discrepancies in estimates for mid-Proterozoic atmospheric $O_2$ (5, 12).

In comparison, the kinetics of Fe(II) oxidation—and the associated Fe isotope systematics—are relatively well-constrained, building on the foundation of research spanning over two decades and summarized in recent literature reviews (17, 18). Our approach is based on a major element, and compared to some other proxies ironstone Fe isotope signals are more likely to be preserved during the postdepositional history of the rocks. Perhaps most importantly, many mid-Proterozoic redox proxy data come from sedimentary rocks deposited in comparatively deeper marine settings (such as shales). The ironstone Fe isotope proxy is more directly related to $O_2$ levels in shallow marine environments that hosted the evolution of early aerobic eukaryotes, and our kinetic model for Fe(II) oxidation is able to provide critical quantitative constraints on relevant shallow seawater $O_2$.

Geochemical evidence (largely from the shale record) has been used to argue for the oxygenation of the deep oceans in the late Proterozoic, possibly during the Ediacaran Period (635 to 541 Ma) (e.g., refs. 45 and 53). However, estimates for the timing of hypothesized late Proterozoic oxygenation span hundreds of millions of years, and the magnitude of this hypothesized rise in $O_2$ lacks consensus, making it difficult to test hypotheses regarding the cause-and-effect relationships between oxygenation and eukaryotic evolution (e.g., see ref. 11). Our ironstone dataset suggests an earlier oxygenation event, which led to more $O_2$-rich shallow waters in the Tonian Period (1,000 to 720 Ma), as evidenced by positive $^{56}Fe$ values in the ca. 900 Ma Nelson Head and McClure ironstones of Canada, yet negligible fractionation in the ca. 750 Ma Galeros Formation ironstone (Fig. 2). The absence of a correlation between $^{56}Fe$ values and detrital proxies (SI Appendix, Fig. S6), and the similar petrography of the Galeros Formation ironstone to other Proterozoic ironstones, suggests that these observed trends are not driven by mineralogical controls or detrital contribution. To our knowledge, there are no other shallow-marine ironstones reported from the late Proterozoic; positive $^{56}Fe$ values in iron formations associated with the ca. 720 Ma Snowball Earth event were deposited under ice cover in disequilibrium with the atmosphere (54) and were not considered in this study. Therefore, although they provide only a limited temporal snapshot, our new ironstone data suggest a change in surface $O_2$ levels in the Tonian Period and narrow down the timing of this oxygenation to between ca. 900 and 750 Ma (SI Appendix for geochronological details). This oxygenation is broadly coincident with independent evidence for redox-driven biogeochemical change during this interval, including the reorganization of the phosphate (55) and sulfate (e.g., ref. 56) cycles, as recorded by marine sedimentary rocks.

Implications for Global Oxygen Cycles and Eukaryotic Evolution

The ironstone record of low atmospheric $O_2$ (less than ~1% PAL) during the mid-Proterozoic highlights gaps in our mechanistic understanding of the $O_2$ cycle, given that some models have questioned the stability of low-$O_2$ worlds (e.g., see ref. 57). This possibility suggests that following the Great Oxidation Event, atmospheric $O_2$ was high enough to form an oxygen layer that shielded near-surface $O_2$ from photolysis (58), yet the oxygen cycle included sufficient feedbacks that could maintain $O_2$ below ~1% PAL. The subsequent oxygenation of surface environments toward the end of the Proterozoic also coincides with biomarker evidence for the rise of eukaryote-dominated ecosystems (59, 60) and an apparent rise in eukaryotic fossil diversity (61, 62). Evidence for drill holes and biomineralized scales in late Tonian fossils (63, 64) may indicate greater eukaryotic predation during this time. As the ingestion of larger cells is an $O_2$-intensive activity (61), and eukaryotic predation is thought to be near-absent in modern anoxic habitats (40), low marine $[O_2]$ may have restricted this behavior in earlier ecosystems. Further, redox-related nutrient limitations on primary productivity prior to the middle Tonian may have exerted a strong control on ecosystem structure and eukaryotic predation (44). A mid-to-late-Tonian oxygenation event could therefore have provided both the requisite $O_2$ and nutrient supply for the proliferation of eukaryotic predation via a concomitant reorganization of the phosphorus cycle (44). Some molecular clock analyses estimate that animals may have first diverged during this time (65) (Fig. 24), and tentative fossil evidence for primitive metazoa (sponges) is reported from mid-Tonian strata (66). The selective pressure of expanded eukaryotic predation may have been a controlling factor in the timing of the emergence of animal multicellularity during this time (61, 62). Therefore, our evidence from the ironstone record provides critical support for the idea that surface $O_2$ levels were changing in step with eukaryotic evolution in the Proterozoic.

Materials and Methods

Ironstone samples were collected from surface outcrop and supplemented by samples from the University of Cincinnati Museum and the Yale Peabody Museum collections. These samples were prepared as polished thin sections.
and analyzed using reflected light microscopy, Raman spectroscopy, and scanning electron microscopy. Well-preserved ironstone samples were crushed in an agate mill to a fine powder, and sample powders (n = 201) were digested for bulk rock geochemical analyses. Major and trace element compositions were determined using an inductively coupled plasma mass-spectrometer (ICP-MS). For Fe isotope analysis, Fe was purified from sample solutions using ion-exchange chromatography and measured using a multicollector ICP-MS. Refer to SI Appendix for detailed methods.

**Data Availability.** All study data are included in the article and/or supporting information. All samples used in this study will be archived in the Yale Peabody Museum upon publication of this work.

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**SI APPENDIX**

**APPENDIX A.**

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