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Post-depositional REE mobility in a Paleoarchean banded iron formation revealed by La-Ce geochronology: A cautionary tale for signals of ancient oxygenation



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ABSTRACT

Precambrian banded iron formations (BIF) are chemical sedimentary deposits whose trace element signatures have been widely used to interrogate the chemical composition and redox state of ancient seawater. Here we investigated trace element signatures in BIF of the 3.22 Ga Moodies Group, Barberton Greenstone Belt (South Africa), which are interbedded with near-shore siliciclastic sedimentary rocks and represent one of the oldest known shallow-water occurrences of BIF. Unusual rare earth element (REE) signatures, notably with pronounced negative Ce anomalies in shale-normalized spectra, have been previously reported for chemical sediments of the Moodies Group, which we confirm here through an expanded dataset for Moodies BIF spanning three different localities. We find negative Ce anomalies as low as 0.2 Ce/Ce* that are associated with unusual enrichment of LREE relative to HREE in the sample set. While total REE abundances and certain REE features appear strongly related to the concentration of detrital indicators (e.g., Zr), and are likely primary, other features, notably LREE enrichment, cannot be explained as a primary feature of the sediment. This is better explained by later addition of REE from a LREE-enriched but Ce-depleted fluid that generated the significant negative Ce anomalies observed in surface samples of Moodies Group BIF. This REE addition event influenced both Sm-Nd and La-Ce isotope systematics, the latter yielding an isochron of 60 \pm 32 Ma, thus constraining the timing of emplacement of the negative Ce anomalies to the past 100 Ma, possibly upon surface exposure of the Barberton Greenstone Belt to wetter conditions during the Cenozoic. Our findings constitute a cautionary tale in that even the most immobile elemental redox proxies may be more sensitive to post-depositional modification than previously thought, and demonstrate the clear advantage offered by paleoredox proxies coupled to radiometric geochronometers to enable the direct dating of ancient signals of Earth surface oxygenation.

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1. Introduction

Banded iron formations (BIF) are chemical sediments deposited throughout much of the Precambrian. They consist primarily of iron oxide minerals (hematite, magnetite) and cryptocrystalline quartz (chert), along with lesser amounts of carbonates, iron-rich clays, and other secondary alteration minerals, depending on the metamorphic grade (Bekker et al., 2010; Konhauser et al., 2017). The precursor chemical minerals thought to have initially formed BIF (amorphous Fe(II)- and Fe(III)-hydroxides, carbonates, and silicates) are unstable, and have universally recrystallized to a more stable metamorphic assemblage of dehydrated minerals. Nonetheless, BIF have been widely considered a useful proxy for the chemical composition of ancient seawater and porewaters in which they formed (see Konhauser et al., 2017, for review). Being highly nonporous and impermeable, BIF are conventionally thought to be relatively robust against significant disturbance during metamorphism or metasomatism (e.g., Robbins et al., 2019). Indeed, mil-

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Fig. 1. Location and stratigraphy of Moodies Group BIF in the Barberton Greenstone Belt. (A) Map showing sampling locations. (B) Stratigraphic columns of the Stolzburg Syncline, Moodies Hills Block, Dycedale Syncline and Eureka Syncline. Abbreviations of stratigraphic units follow Anhaeusser (1976). (For interpretation of the colors in the figure(s), the reader is referred to the web version of this article.)

limeter scale variations in trace elements (e.g., Bau, 1993; Bau and Dulski, 1996) and in iron isotope compositions in BIF (e.g. Frost et al., 2007), along with experiments simulating BIF diagenesis (e.g. Posth et al., 2013), have been used to argue for general elemental immobility, even for samples that have experienced complete recrystallization during amphibolite-facies metamorphism.

Rare earth element (REE) patterns, and in particular anomalies in the abundance of cerium (Ce) relative to its neighbors, are commonly used to interrogate redox conditions of ancient seawater. Cerium is particularly prone to fractionation from other REE because oxidation of Ce(III) to Ce(IV) under mildly oxidizing conditions greatly reduces Ce solubility, resulting in its preferential removal from solution onto Mn(IV)-Fe(III)-oxyhydroxides, organic matter, and clay particles (Byrne and Sholkovitz, 1996). Under oxidizing conditions, seawater is thus characterized by a strong negative Ce anomaly. Accordingly, the presence of negative cerium anomalies in chemical sediments has often been used to infer oxidative conditions during deposition. Despite the potential for significant insight into the primary REE chemistry of ancient seawater, the interpretation of cerium anomalies in ancient sedimentary rocks requires caution because post-depositional REE modifications have shown that cerium may behave differently from other REE (Mongelli, 1993; Cotten et al., 1995). Hayashi et al. (2004) were the first to use 138 La- 138 Ce isotope systematics ($T_{1/2} = 292.5$ Ga) in an attempt to constrain the timing of emplacement of a cerium anomaly in ancient sedimentary rocks. The first La-Ce isotope measurements obtained by Hayashi et al. (2004) on samples from the Barberton Greenstone Belt (BGB) show that the presence of cerium anomalies must be interpreted with caution. They attributed Ce anomalies in the BGB, including a single BIF sample from the Moodies Group with $Ce/Ce^* = 0.93$ (calculated by Hayashi et al. in the absence of Pr data as the chondrite-normalized projection between La and Nd), to post-depositional remobilization of REE prior to ca. 1.1 Ga (Havashi et al., 2004).

In this paper we present major and trace element measurements in 48 BIF outcrop samples collected at three localities in the Moodies Group, dated at 3.22 Ga (Heubeck et al., 2013). Measurements of the La-Ce and Sm-Nd isotope systematics are presented in samples characterized by a large range of cerium anomalies. This dataset permits; (1) direct dating of the formation of the cerium anomaly, and (2) resolution of whether REE spectra represent primary features or were later modified during secondary events, such as surficial weathering of exposed outcrop. Here we demonstrate that the negative cerium anomalies in Moodies BIF were produced by relatively recent fluid addition and that this signature cannot be used to discuss redox conditions during their deposition.

2. Geology of Moodies Group BIF

The Moodies Group of the BGB is comprised of approximately 3.5 km of sand- and siltstone, subordinate conglomerate and volcanics, and minor ferruginous sediment that were mostly deposited in shallow-marine and/or terrestrial settings (Heubeck, 2019). Onset of Moodies deposition postdates the ages of dacitic volcanics conformably underlying basal Moodies strata (3225 \pm 3 Ma, Kröner et al., 1991; 3223 \pm 1 Ma Heubeck et al., 2013). Dacitic ash-fall tuffs in the central BGB, approximately mid-section in the stratigraphic column (near BIF unit MdI2, Fig. 1), have been dated at 3219 \pm 3 Ma (Heubeck et al., 2013), similar to the age of a porphyritic dike crosscutting topmost Moodies units dated at 3219 \pm 9 Ma (Heubeck et al., 2013). Thus, Moodies deposition occurred approximatively between 3223 and 3219 Ma, and likely within a few Ma of 3222 Ma ago (Heubeck et al., 2013). Regional metamorphic grade is lower greenschist facies (Tice et al., 2004), with evidence for metamorphic overprint during at least three events: (1) emplacement of the Kaap Valley Pluton at 3214 \pm 4 Ma, (2) during late granite plutonism at ca. 3100 Ma (Toulkeridis et al., 1994), and (3) related to fluid circulation associated with Ventersdorp Supergroup volcanism or the Limpopo orogeny ca. 2650-2700 Ma ago (Toulkeridis et al., 1998).

In the north-central part of BGB, up to four regionally mappable ferruginous units, 2 to 6 m thick and comprised of jaspillite and BIF, are interbedded with fine-grained sand- and siltstones

(Heubeck, 2019). Geological mapping and interpretation of sedimentary structures suggest that deposition of ferruginous units occurred largely in prodelta- and protected bay / lagoonal settings. The BIF samples examined in this study were collected from surface outcrops at three localities in the Moodies Group (Fig. 1, supplementary materials): (1) unit MdI2 in the Clutha Creek section of the Eureka Syncline (samples CC; 25°41′52.81″S, 31°5'10.89"E), (2) unit MdI2 from the Devil's Staircase Road section of the Moodies Hills Block (samples DSR; 25°49'55.82"S, 31°0'50.49"E), and (3) unit MdI1 from the Stolzburg Syncline (samples SB; 25°54'8.04"S, 30°50'44.90"E). Samples were collected using hammers, chisels, and a sledge, avoiding clear signs of alteration, and then trimmed for fresh surfaces; fractured pieces were discarded. All samples are distal from hydrothermally overprinted brittle-ductile shear zones that are common in the northern BGB. In all three sections, BIF occur in thick, conformable sections of thinly bedded, small-scale-slumped siltstones and veryfine-grained sandstone interbedded with rare laminated shale. See supplementary material for detailed descriptions of each sample site and a lithological column of the DSR site, placing the samples in a stratigraphic context.

3. Methods

All samples were prepared using trace-element clean methods (crushing and powdering in pure tungsten carbide and agate, respectively) and all digestions and column separations were performed in clean laboratories at the European Institute for Marine Studies in Brest and at the Laboratoire Magmas et Volcans in Clermont-Ferrand, France. Full details of the analytical methods are provided in the supplementary materials. Briefly, for analysis of major elements, samples were digested at 90 °C in PFA vials using concentrated HNO₃ and HF, followed by immediate neutralization with 20 gL⁻¹ H₃BO₃ to retain Si, as per Cotten et al. (1995). After dilution by a factor of 400, samples were analyzed using an Yvon Horiba Ultima 2 Inductively Coupled Plasma Optical Emission Spectrometer (ICP-OES). For analysis of trace elements and Nd isotopes, between 80 and 120 mg of powder was digested sequentially in PFA vials at 90 °C using concentrated HF-HNO₃, aqua regia, and 6M HCl. For trace element analysis, aliquots were resuspended in 2% HNO3 with indium as an internal standard and analyzed using a Thermo Scientific Element2 High-Resolution Inductively Coupled Plasma Mass Spectrometer (ICP-MS) at the Pôle Spectrométrie Océan in Brest, France. To assure complete digestion and confirm the REE data acquired after HF-HNO₃-Aqua Regia digestion in PFA beakers, a selected set of sample powders were re-analyzed by HR-ICP-MS after Na₂O₂-NaOH alkaline fusion in glassy carbon crucibles at 650 °C in the presence of a Tm spike, following the protocol and data reduction procedure of Bayon et al. (2009).

For Nd isotopes, aliquots were evaporated and re-dissolved in 2.5M HCl for chemical separation of REE using cationic BioRad[®] AG50X8 200-400 mesh columns, followed by further purification on LnSpec Eichrom resin. Nd isotopic measurements were performed during two analytical sessions at the Pôle de Spectrométrie Océan in Brest using a Thermo Scientific Neptune multi-collector ICP-MS and a Thermo Scientific Triton Thermal Ionization Mass Spectrometer (TIMS, Triton).

For Ce isotopes, 100-150 mg of rock powder was digested using a HF-HNO₃ dissolution technique. The chemical separation used to separate the Ce fraction involved four steps of column chemistry. Isotopic measurements were performed on a Thermo Scientific Thermal Ionization Mass Spectrometer (TIMS) at the Laboratoire Magmas et Volcans as previously described by Bonnand et al. (2019). Together with the samples, Ce_{AMES} and Ce_{LMV} reference materials were analyzed and gave ¹³⁸Ce/¹⁴²Ce values of 0.0225743 \pm 0.0000007 (2 s.d.; n = 26) and 0.0225705 \pm 0.0000006 (2 s.d.; n = 48), respectively.

4. Results

4.1. Major and trace elements

The Moodies Group BIF are composed of chert, magnetite, and hematite, with minor phyllosilicates (likely stilpnomelane and chamosite), goethite, and rare ankerite (see Table S2 for X-ray diffraction method and data). Samples showed a large range in the concentration of detrital indicators, with Al₂O₃ ranging from 8.5 wt.% to <0.1%, and with correlated concentrations of Sc, Zr, Th, and Hf that reached sub-ppm values in the least contaminated samples (see Table S3 in the supplementary information for complete elemental dataset). Iron concentrations ranged from 73.8 wt.% Fe2O3Total to 5.8 wt.%; 43 out of 54 samples meet the definition of iron formation of >15 wt.% Fe (James, 1954), including two samples with Al₂O₃ between 6 and 8 wt.% (Fig. S1). Phosphorus concentrations averaged 0.11 wt.% P₂O₅ and no apatite or other P-rich phases were detected by either X-ray diffraction or µXRF scanning (data not shown). Redox-sensitive trace elements, such as U and Mo, are present at sub-ppm levels. Transition metals (e.g., V, Cr, Mn, Co, Ni, Cu and Zn) show a variability of approximately one order of magnitude in absolute abundances, similar to other larger BIF datasets (Fig. S2; e.g., Konhauser et al., 2011).

Shale-normalized (SN) spectra for rare earth elements + yttrium (REY_{SN}; simplified as REE throughout this manuscript) span a wide range in abundances relative to Post-Archean Australian Shale (PAAS; updated values from McLennan, 1989), ranging from below 0.01 to close to 1 (Fig. 2). Total REE concentrations show strong correlation with detrital indicators, such as Zr concentrations (Fig. 3). REE spectra are highly diverse, with some showing light REE (LREE) enrichment relative to heavy REE (HREE) enrichment, while others show the reverse. Nearly all samples show suprachondritic Y/Ho ratios, ranging from 20.5 to 45.5, with an average of 35.0, as well as significant positive La and Eu anomalies, and mild positive Gd anomalies. A striking feature of the spectra are highly variable Ce anomalies, with Ce/Ce* reaching values as low as 0.19. A subset of the samples digested by alkaline fusion and analyzed for REE using a Tm spike (Bayon et al., 2009) confirm the large range in La/Ce and Ce/Ce* that is observed for samples digested using concentrated acids in PFA vials (Table S4). The cerium anomaly is calculated here using the equation derived by Lawrence et al. (2006) with $Ce/Ce^* = Ce_{SN}/(Pr_{SN} * Pr_{SN}/Nd_{SN})$, where La is not considered in the calculation as to avoid false negative Ce anomalies that may arise simply as the result of La enrichment.

4.2. Nd and Ce isotopes

Nd isotope compositions (¹⁴³Nd/¹⁴⁴Nd) of 22 selected samples from all three localities range from 0.510836 to 0.511459, while ¹⁴⁷Sm/¹⁴⁴Nd ratios range from 0.0911 to 0.1453. On a ¹⁴⁷Sm/¹⁴⁴Nd – ¹⁴³Nd/¹⁴⁴Nd plot, samples from the SB locality do not show any correlation. Samples from the DSR locality are tightly clustered. With such a small range of Sm/Nd ratios, it is not possible to calculate a robust isochron. Samples from the CC locality show a positive correlation that yields an errorchron with an apparent age of 2770 \pm 530 Ma (MSWD = 16).

 138 La/ 142 Ce ratios for the five DSR samples analyzed range from 0.004 to 0.013, while the SB sample analyzed showed a value of 0.004 (Fig. 4b). Cerium isotope compositions (138 Ce/ 142 Ce) ranged from 0.02257053 to 0.02257522 (Table S5). On a 138 La/ 142 Ce vs. 138 Ce/ 142 Ce plot, the five analyzed DSR samples define an isochron



Fig. 2. PAAS-normalized REE patterns of Moodies BIF analyzed in this study (DSR = Devil's Staircase Road of the Moodies Hills block, CC = Clutha Creek section of the Eureka Syncline, SB = eastern Stolzburg Syncline).



Fig. 3. Total REE content versus ${\rm Zr}$ content in the Moodies BIF. Abbreviations as in Fig. 2.

that yields an apparent age of 60 ± 32 Ma. The single analyzed SB sample overlaps the range defined by the DSR samples.

5. Discussion

5.1. Unusual REE signatures in Moodies Group BIF

Samples of the shallow-water BIF of the Moodies Group analyzed here show some rather unusual characteristics in their REE systematics. Most striking is the presence of strong Ce anomalies, ranging down to Ce/Ce^{*} = 0.19. Because cerium is readily oxidized from Ce(III) to Ce(IV) in the presence of free oxygen, this process leaves oxic waters depleted in Ce, creating characteristic negative Ce anomalies in oxygenated seawater (shale-normalized Ce anomalies, Ce/Ce^{*}, of about 0.3 – 0.1; e.g., German et al., 1995). BIF deposited before the GOE generally lack Ce anomalies, a feature attributed to the prevalence of anoxic water column conditions (e.g., Bau and Dulski, 1996). Moodies Group BIF samples show significant Ce anomalies at all three sample locations (Ce/Ce^{*} = 0.2 – 0.95; Fig. 2), with samples from DSR and SB localities showing the most pronounced anomalies. The magnitude of Ce anomalies observed in our dataset reaches more extreme values than those

observed by Hayashi et al. (2004) in three samples from the Moodies and underlying Fig Tree Groups. One might be tempted to equate these anomalies with oxic Ce cycling at the time of deposition; however, Hayashi et al. (2004) suggested that such anomalies may instead represent post-depositional mobilization of REE with a maximum age of 1.1 Ga.

Another unusual feature of the REE spectra is the highly variable degree of LREE vs. HREE enrichment. Modern seawater, and most detritus-free ancient chemical sediments (both BIF and carbonates; e.g., Bolhar et al., 2004), generally shows HREE enrichment relative to LREE due to lanthanide contraction and the decrease in ionic radii going from LREE to HREE as the result of an increasingly filled f-electron shell. This contraction results in a more important extent of aqueous carbonate complexation for HREE relative to LREE (Lee and Byrne, 1992), leaving a greater proportion of LREE available for adsorption to reactive particle surfaces, thus decreasing their concentration in seawater relative to the more strongly carbonate-complexed HREE. Another consequence of the lanthanide contraction is that LREE are more efficiently mobilized by alteration fluids bearing important REE ligands such as fluoride, phosphate, and carbonate (e.g., Wood, 1990).

Both syn-depositional and post-depositional processes may influence REE systematics. BIF, however, have generally been considered robust against post-depositional alteration of their REE signatures (Bau, 1993), even by protracted high-grade metamorphism (Bolhar et al., 2004). Nonetheless, the highly atypical nature of the REE signatures observed in this study, and the fact that all samples were obtained from surface outcrop, lead us to examine these assumptions more closely. We first consider controls over REE systematics that can be confidently linked to syn-depositional processes, such as the seawater precipitation of iron and silica chemical components and the admixture of detrital contaminants, before examining alternative explanations for chemical features that are difficult to explain as primary.

5.2. Syn-depositional control over REE signatures: chemical sedimentation vs. detrital contamination

For the original protolith, prior to any post-depositional alteration, it is reasonable to assume that REE were derived from two contrasting sources; those (1) scavenged directly from seawater onto the surface of the original chemical precipitates, and



Fig. 4. (a) Sm-Nd isotope composition of Moodies Group BIF analyzed in this study. The age and the MSDW were calculated using DSR and CC samples. (b) La-Ce isotope composition of selected samples from the SB and DSR localities.

(2) delivered physically in association with detrital components. Distinguishing these two sources can be difficult as both primary depositional processes and post-depositional alteration processes have the potential to generate correlations that do not imply causation. For example, total REE concentrations in this dataset show mild but notable correlations with major elements which are typically associated with a predominately chemical sedimentary origin (e.g., Fe, Si, Mn, P; Pearson's correlation coefficients of 0.21, -0.41, 0.28, and 0.59, respectively). However, even stronger correlations are observed between total REE concentrations and the concentrations of detrital indicators (e.g., Al, Ti, Zr, Hf, Th; Pearson's correlation coefficients of 0.91, 0.86, 0.94, 0.91, and 0.95, respectively). A variety of processes may induce such correlations. These include early depositional and diagenetic processes such as REE adsorption to iron oxide precipitates or sequestration of REE into early diagenetic phosphate minerals. However, multiple features of the REE spectra show no relation to the inferred chemical sedimentary component. For instance, there exists no good correlation between Ce/Ce* nor LREE/HREE (evaluated as Nd/Yb, for example) and the concentrations of Fe, Si, Mn, or P.

On the other hand, some features of the REE spectra can be explained by the admixture of detrital components. In order to assess the influence of detrital contributions on REE spectra of the Moodies BIF, we modeled the addition of detrital components to a detritus-poor sample. The chemical composition of the Archean continental crust, used as a detrital component in our model, is highly debated (e.g. Martin et al., 2005). We chose to use an average composition of TTG and Sanukitoids presented in Halla et al. (2017) as representative of the Archean crust and as a material most likely to increase the LREE/HREE slope. The models are presented in Fig. 5 and the chemical composition of the detrital component is given in Table S6. As shown in Fig. 5a, Moodies samples are characterized by large variations in La/Yb ratios. The addition of detrital material to a detritus-poor sample results in an increase in the La/Yb ratio as shown in Fig. 5a. The model succeeds in reproducing the general trends observed in CC and SB samples but not the high La/Yb ratios in detritus-poor DSR samples. The model also explains variations observed in Yb and Th concentrations (Fig. 5b). This suggests that the increase in Yb in the Moodies samples is primarily controlled by the addition of detrital material. During the addition of detrital material to a detrituspoor sample, the chemical signatures of the REE spectra are modified (Fig. 5c). For example, the LREE depletion, characteristic of detritus-poor samples, is erased by the addition of about 10-15% detrital material. The addition of detrital material also results in the suppression of the positive Y anomaly (Fig. 5c). However, during the addition of detrital material, the positive Eu anomaly is not affected and remains a characteristic feature of detrital-rich samples (Fig. 2). This indicates that the chemical composition of the crustal component is characterized by a positive Eu anomaly, evidence for which is also apparent in Barberton TTG rocks (Moyen et al., 2019, their supplementary data). During the addition of detrital components, the La/Ce and Sm/Nd ratios both decrease slightly.

Detrital contributions can explain large variations in total REE concentrations and also the correlation between Yb and Zr, but fail to explain certain features in the dataset, notably the high La/Yb ratios seen in DSR samples at low Zr abundances, as well as the presence of Ce anomalies in general. As discussed in the next section, these are better explained by post-depositional alteration processes. Detrital contributions appeared to have modified the sensitivity of both of these features (La/Yb and Ce anomalies) to post-depositional alteration.

5.3. Fluid addition

REE spectra have been clearly affected by a secondary process that resulted in differential REE mobilization. Two distinct mechanisms of REE mobilization could be proposed to explain such variations; (1) removal of REE during chemical weathering, and/or (2) addition of REE by fluid circulation. The REE concentrations in aqueous solutions are affected by several parameters, including the original composition of the protolith, the chemical composition of the fluid, and physicochemical conditions during the fluid-rock interaction. For example, during chemical weathering and leaching of REE in basalts, it has been shown that LREE are preferentially mobilized compared to HREE (Cotten et al., 1995). However, it's important to note that the relative enrichment of LREE and HREE can be highly variable as a function of protolith, fluid chemistry, degree of alteration, and biological effects (Laveuf and Cornu, 2009). Importantly, during chemical weathering and fluid-rock interaction in tropical environments, Ce^{3+} is oxidized to Ce^{4+} and becomes insoluble, leaving the fluid with a strong negative anomaly (Cotten et al., 1995). The removal of REE by chemical weathering would thus result in depletion of the total REE concentration of the protolith, leaving a strong LREE depletion in the residual altered rock and a strong positive Ce anomaly, which is the opposite of what is observed in Moodies BIF.

We propose instead that fluid addition is largely responsible for generating the unusual REE spectra in the samples. As discussed below, the initial REE content of the iron formations themselves also appears to have played a role in the modification of REE spectra features during fluid addition.

To further explore the possible impact of fluid addition, we have developed two fluid addition models (Fig. 6). The fluid used in these models bears deliberately simple REE spectra and shows



Fig. 5. (a) La/Yb ratio versus Zr concentrations for Moodies Group BIF. (b) Yb concentration versus Th concentration of the Moodies Group BIF. (c) PAAS-normalized REE patterns for a detritus-poor sample (CC-14-52) together with models of admixture of detrital input. Dashed lines in (a) and (b) represent the admixing model of detrital material. Values for Archean Crust in (c) are from Halla et al. (2017) (Table S6). The values given in (c) show the amount of detrital material added to the detritus-poor sample. The three samples in dashed grey lines in (c) are sandstones interbedded in the studied sections (data in table S3). See text for details.

two main features; (1) a positive linear enrichment from Lu to La, and (2) a strong negative Ce anomaly (Ce/Ce^{*} = 0.27). The REE concentrations in the model fluid presented in Fig. 6 are listed in Table S6. The two models presented in Fig. 6 represent the admixing of REE from fluids of the same composition to rocks of differing initial compositions. Two end-member cases are presented where the starting compositions were either detritus-poor or detritus-rich, respectively. During fluid addition, the total REE concentration increases, the REE spectra become progressively en-



Fig. 6. Ce anomaly versus shale-normalized Ce/Yb ratio in Moodies Group BIF. The dashed line represents the admixing model; arrow indicates increasing admixture of detrital material. The dotted and solid lines show fluid addition models with detritus-rich and detritus-poor starting composition, respectively. See text for details.

riched in LREE, and negative Ce/Ce* values are produced (Fig. 6). Importantly, the Ce/Yb ratio will either increase or decrease, depending on the composition of the protolith (i.e., detritus-rich or -poor). Similarly, in the case of La/Yb (Fig. 5a), detritally contaminated samples (high Zr) do not reach La/Yb ratios above crustal values, whereas detritus-free samples (low Zr) show both high and low La/Yb values. Samples showing low La/Yb values appear to retain the light REE depletion expected for seawater precipitates. However, samples with high La/Yb ratios cannot be explained by a seawater origin nor by detrital contamination; the best explanation for samples found above crustal La/Yb values is late LREE addition.

The REE concentration in the fluid is unknown which prevents us from determining the amount of fluid addition necessary to explain the observed variations. Nevertheless, it is clear that the amount of inherited REE required to alter spectra will be greater for a detritus-rich sample than for a detritus-poor sample. The model we present illustrates the late generation of multiple features of the unusual REE spectra using relatively simple and plausible assumptions, notably a preferential mobility of LREE compared to HREE. It is possible that the source of the mobilized REE is the host rock itself, in which case the fluid would be expected to carry pre-existing anomalies, such as a positive Eu anomaly. However, when considering fluids of different compositions and relative REE mobilities, it quickly becomes apparent that the system is poorly constrained and that diverse fluid compositions may be evoked to explain any given alteration pattern, regardless of the protolith from which REE in the fluids derived. What is apparent is that these fluids were enriched in LREE while at the same time depleted in Ce.

Groundwaters today are characterized by REE concentrations that are generally low, in the nanomolar to picomolar range, including deeper (\sim 200 m) groundwaters draining REE-rich lithologies such as granite (Janssen and Verweij, 2003; Noack et al., 2014; Munemoto et al., 2015). Mean iron formation Ce and Pr concentrations observed in our dataset are 10 and 1.3 ppm, respectively. Mass balance indicates that to supply an equivalent amount of REE from groundwater, volumes ranging from as little as 200 L per kg of iron formation, to as much as ten million liters, are implied. Acidic geothermal waters may carry higher REE loads, up to \sim 0.1 ppm (Wood, 2006), and could supply the necessary REE in as little as 100 L per kg of iron formation. Crucially, the BGB already



Fig. 7. (a) ¹³⁸Ce/¹⁴²Ce versus shale-normalized Ce/Yb ratio. (b) ¹⁴⁷Sm/¹⁴⁴Nd versus shale-normalized Ce/Yb ratio in the Moodies Group BIF analyzed in this study. The dotted and solid lines show fluid addition models with detritus-rich and detritus-poor starting composition, respectively. See text for details.

contains evidence of significant REE mobility during the formation of "ironstone pods" as the result of iron dissolution and reprecipitation in shallow groundwater systems (Lowe and Byerly, 2007). Contrary to the samples studied herein, these pods show a range of features that are inconsistent with an Archean origin, namely a lack of deformation, stratification aligned with modern topographic slopes, abundant cavities, and goethite drip features. Importantly, these ironstone pods bear both positive and negative Ce anomalies, reaching as low as 0.2 Ce/Ce*, and REE concentrations that are similar to those reported here for the Moodies BIF (Hren et al., 2006). They thus offer an attractive analogue for how significant quantities of REE may have been mobilized under near-surface conditions and with sufficient oxygen exposure to generate large Ce anomalies in markedly older rocks.

5.4. Applicability of Sm-Nd and La-Ce geochronometers for dating fluid addition

As discussed above, fluid addition likely had a strong influence on REE elemental ratios. In Fig. 7, we present the impact of our fluid addition model on the La/Ce and Sm/Nd ratios. The La/Ce ratio is most strongly affected because the added fluid is heavily depleted in Ce. This results in an increase in the La/Ce ratio of the altered samples. In the models presented in Fig. 7a, the ¹³⁸La/¹⁴²Ce ratio evolves from 0.004 to 0.015. During fluid addition, the Sm/Nd ratios are also affected and slightly decrease from 0.13 to 0.10 (Fig. 7b). The modification of parent/daughter elemental ratios during fluid addition has strong implications for the time-dependent evolution of Ce and Nd isotopic compositions because the relatively large variations in La/Ce produced during alteration means that it is possible to use Ce isotopes to constrain the timing of fluid addition.

A large spread in La/Ce ratios, if ancient, should give rise to a corresponding large spread in the ¹³⁸Ce/¹⁴²Ce ratio. However, if the fractionation of La and Ce is more recent, ingrowth of ¹³⁸Ce will be small, considering the long half-life of ¹³⁸La, and samples will show a restricted range in ¹³⁸Ce/¹⁴²Ce despite a large apparent range in La/Ce. We have measured the Ce isotopic compositions of six samples with varying ¹³⁸La/¹⁴²Ce ratios (from 0.004 to 0.012), and a plot of ¹³⁸Ce/¹⁴²Ce versus ¹³⁸La/¹⁴²Ce gives an isochron age of 60 \pm 32 Ma (Fig. 4b). This suggests that the La-Ce elemental fractionation occurred considerably more recently than deposition of the Moodies BIF. In Fig. 8a, we report the theoretical isochrons for two fractionation ages (the known 3.2 Ga U/Pb zircon-constrained depositional age, and the La/Ce fractionation age of 1.1 Ga suggested as a maximum age by Hayashi et al., 2004), anchored for simplicity at the same initial 138 Ce/ 142 Ce value as indicated by our data. As is clearly evident, the data reported in this study does not show the expected ¹³⁸Ce ingrowth predicted by a 3.2 Ga isochron, but rather shows a restricted range in ¹³⁸Ce/¹⁴²Ce, consistent with a suite of sediments that possessed similar La/Ce ratios for most of their history. This confirms that the La/Ce fractionation is not a primary feature of the Moodies BIF, and crucially, the negative Ce anomalies were generated postdepositionally. Our data also show a less inclined slope than the 1.1 Ga isochron, which contradicts the La/Ce fractionation ages proposed by Hayashi et al. (2004). This maximum age was essentially proposed to explain the radiogenic Ce isotopic composition of a banded ferruginous chert from the Fig Tree Group ($\varepsilon Ce = 9.2$). All our samples, on the contrary, plot on the terrestrial Ce-Nd array defined by Israel et al. (2019) with present day ε Ce and ε Nd mean values for the less-perturbed DSR samples of 3.9 and -33.1, respectively.

The fractionation of La from Ce, and thus the generation of negative Ce anomalies, is more recent than proposed by Hayashi et al. (2004). Nonetheless, considering the long half-life of ¹³⁸La, we emphasize that the geochronological resolution of our dataset is limited. It is safe to assume that La/Ce fractionation likely occurred in the last 100 Ma. This is consistent with the findings of Lowe and Byerly (2007), who inferred that the "ironstone pods" found in multiple localities in the Barberton Greenstone Belt formed during the Cenozoic, most likely during the Pleistocene. These ironstone pods are composed largely of goethite that would not have survived metamorphic temperatures exceeding ~100°C without recrystallization to hematite. The BIF sampled in this study also contain a small, but non-negligible, quantity of goethite in some samples (Table S2). This similarly indicates that some hydrological modification must have occurred at near-surface conditions. There appears to be no correlation between the magnitude of the Ce anomaly and any feature of sample mineralogy that would indicate a consistent mineralogical pattern of post-depositional alteration (Table S2). While the Moodies BIF show multiple features consistent with an Archean origin (deformation, strike and dip consistent with adjacent siliciclastic units, mineral assemblages consistent with metamorphosed BIF), it is likely that these units were also subject to hydrological modification during wetter periods of the Cenozoic.

Fluid addition also played a role in the Sm/Nd variations measured in the Moodies BIF, with consequences for Sm-Nd geochronology. In a plot of 143 Nd/ 144 Nd versus 147 Sm/ 144 Nd, the samples from the CC locality lie on an isochron with an age of 2770 \pm 530 Ma, whereas the other localities either plot outside of



Fig. 8. (a) La-Ce isotope composition of selected samples from the SB and DSR localities; (b) Sm-Nd isotope composition of Moodies Group BIF analyzed in this study. In (a), the black and red lines are theoritical Sm-Nd isochrons, anchored at the same initial ¹³⁸Ce/¹⁴²Ce value, at 1.1 and 3.2 Ga, respectively. In (b), the blue line is the isochron calculated using samples from the CC locality only;the black line is a theoretical isochron calculated using the U/Pb-constrained depositional age. The arrow in (b) indicates the modification induced by fluid addition.



Fig. 9. Degree of Sm/Nd-ratio deviation from the 3.2 Ga isochron (in %) versus magnitude of the Ce anomaly of Moodies Group BIF. See text for details.

the isochron (most SB samples) or show little variation (DSR). In order to test the influence of fluid addition on the Sm/Nd isochron, we calculated the theoretical 3.2 Ga isochron anchored by the sample with the highest Sm/Nd ratio because this sample was least affected by detrital input or fluid addition, which both act to decrease the Sm/Nd ratio (Fig. 8b). It can be clearly seen that the majority of points lie to the left of the theoretical isochron. This is consistent with the models presented where fluid addition is shown to decrease the Sm/Nd ratio and move points to the left of the isochron. The deviation from the 3.2 Ga isochron can be calculated (as the difference in Sm/Nd ratio expressed in %) and plotted as a function of the magnitude of the cerium anomaly (Fig. 9). This deviation scales positively with the magnitudes of negative Ce anomalies in the same samples and is consistent with Sm/Nd ratios that have been affected by fluid addition. However, this fluid delivery did not change the Nd isotope composition. SB samples that are the most affected by the suspected fluid addition show 143 Nd/ 144 Nd ratios in a narrow range (<2 ε), showing that the fluid may have been internal to the examined BIF units.

It is important to note that Sm/Nd ratios are less susceptible to fractionation relative to La/Ce ratios because neither undergo redox transformation under surface or upper-mantle conditions. The shifts in Sm/Nd produced by fluid addition are thus small, and while perceptible on an isochron plot, may still give an isochron age that is within error of the known depositional age (which

is the case of the isochron coinciding with the CC samples in Fig. 4). Ultimately, while the influence of fluid addition on Sm-Nd geochronology may be apparent, it is unsuitable for resolving the timing of fluid addition due to the muted effects on the Sm/Nd ratio. Sm-Nd geochronology should thus not be used to constrain the timing of emplacement of Ce anomalies in ancient rocks, metased-imentary or otherwise.

5.5. Implications for geochemical proxy reconstruction of Archean redox conditions

Redox-sensitive geochemical proxies such as Ce/Ce^{*} are widely applied to reconstruct the history of Earth surface oxygenation. In Archean black shales, enrichments in redox-sensitive elements (S, Mo, Re, Os), as well as Fe, Mo and C isotopes, all suggest that O_2 -rich niches already existed ca. 2.6 to 3.0 Ga (e.g., Wille et al., 2007; Czaja et al., 2012; Stüeken et al., 2012). In diverse metasediments of the Pongola Supergroup, isotope compositions of S, Fe, Cr, Mo, and U have all similarly been used to indicate the presence of some free O_2 ca. 2.96 Ga ago (e.g. Crowe et al., 2013; Eickmann et al., 2018). Sedimentary Fe isotope composition and U enrichment have also been used to argue for oxygenated surface waters as far back as 3.2 Ga ago (Satkoski et al., 2015).

Nevertheless, a Paleo- to Mesoarchean origin for oxygenic photosynthesis remains controversial (e.g., Ward et al., 2016). Part of this controversy is related to alternative interpretations of geochemical records suggested as proxies for O₂ in deep time, especially emerging metal isotope proxies (e.g., Konhauser et al., 2011; Kaufman, 2014) and to uncertainty in the robustness and syngenicity of these signals (Kirschvink et al., 2012; Kaufman, 2014). Indeed, post-depositional oxidation of originally reduced Archean metasediments has been widely demonstrated (e.g., Li et al., 2012; Albut et al., 2018). For instance, Hoashi et al. (2009) hypothesized that hematite in the 3.46 Ga Marble Bar chert (Pilbara Supergroup, Australia) occurred as a primary precipitate formed in the presence of seawater, yet this was later discounted on petrographic grounds (Rasmussen et al., 2014). Similarly, Cr isotope fractionation in the 2.96 Ga Singeni IF (Pongola Supergroup, South Africa) that was used to suggest transient atmospheric oxygenation ca. 3.0 Ga (Crowe et al., 2013) was later shown to be present in surface samples but not in fresh drill core (Albut et al., 2018), and since confirmed using U-series isotopic disequilibrium to represent modern weathering (Albut et al., 2019). Due to the uncertainties surrounding the timing of oxidation, it appears that geochemical proxy evidence indicating the former presence of O₂ in water may

not alone be sufficient to resolve the debate surrounding the origin of oxygenic photosynthesis.

Several geochemical proxies for O₂ form radiogenic systems that are amenable to dating, and radiogenic approaches for constraining the timing of oxidation have now been successfully applied using Re-Os (Kendall et al., 2015) and U-Th-Pb (Li et al., 2012; Satkoski et al., 2015). The latter approach provides the only direct geochronological support to date for an oxidative proxy signal dating as far back as 3.2 Ga ago. Our study underlines the necessity of such an approach. Ce/Ce* is considered one of the most robust redox proxies due to the general insolubility of REE at Earth surface conditions, by their immobility during greenschistand amphibolite-facies metamorphism, and by the fact that Ce becomes even more immobile when oxidized. Indeed, with the exception of the Hayashi et al. (2004) study and this work, we are unaware of any reports of confirmed late emplacement of negative Ce anomalies in Archean-aged metasedimentary rocks. However, several published datasets in Archean rocks warrant reexamination of the inferred syn-depositional origin of negative Ce anomalies. For example, an Archean paleosol from the Singhbhum Craton displays strongly negative Ce anomalies that were used to suggest transient atmospheric oxygenation sometime prior to 3.02 Ga, yet iron loss and U immobility within the paleosol, as well as the presence of redox-sensitive detrital pyrite and uraninite in overlying sediments, indicate reducing conditions (Mukhopadhyay et al., 2014). Furthermore, the sample showing the most pronounced Ce anomaly conspicuously shows pronounced LREE enrichment but comparable Ce abundance relative to other samples. In a study compiling REE data from multiple studies, Kato et al. (2006) highlighted that several different BIF occurrences deposited prior to 2.5 Ga are characterized by negative Ce anomalies. It would be prudent to re-examine these samples for telltale signs of REE mobilization and the generation of Ce anomalies by fluid addition, which may be indicated by LREE enrichment. While late emplacement of a negative Ce anomaly by an alteration fluid simply requires oxic conditions at some point during the fluid's trajectory, it is important to note that this anomaly could persist if conditions became anoxic further along the flow path. Contrary to some redox indicators, such as uranium that are rapidly scrubbed out of solution at reductive redox boundaries (c.f. Abdelouas et al., 2000, and reference therein), only dilution by non-fractionated REE, or possibly reductive dissolution of minerals that already bore a positive Ce anomaly, would erase a negative Ce anomaly from an alteration fluid. Such a scenario may lead to the emplacement of negative Ce anomalies in rocks that simultaneously bear indicators of anoxia, such as U immobility.

This work constitutes a cautionary tale in that even the most robust (e.g., least easily reset) elemental and isotopic proxies for ancient oxygenation, such as REE and their isotopes, are susceptible to post-depositional alteration. In the case of REE-based proxies, as described here, there are a variety of supplemental indicators that can be used to reveal such processes, such as modifications to the REE spectra and perturbation to Nd and Ce isotope systematics. This is not the case for all elemental and isotopic redox proxies. Therefore, when inferring the ancient presence of O_2 from elemental and isotopic redox proxies, additional confirmatory information regarding the syn-depositional nature of the signal is required.

6. Conclusion

Some BIF of the Moodies Group show unusual features in their shale-normalized REE spectra, most notably pronounced negative Ce anomalies and variable LREE vs. HREE enrichment, with some samples possessing LREE that are enriched compared to HREE. We examined the origin of these features with respect to poten-

tial syn- and post-depositional processes that may have influenced the REE spectra. While total REE abundances scale weakly with the concentrations of elements that were likely derived from the chemical components of the sediment (Fe, Si, Mn, P), there appears to be little relation between their concentrations and the features of the REE spectra themselves. By contrast, total REE abundances scaled strongly in a positive fashion with the concentrations of indicators of detrital contamination, notably Al, Ti, Hf, Th, and Zr. Detrital contamination clearly exerted control over REE features in these samples but cannot explain the presence of significant negative Ce anomalies nor extreme LREE enrichments relative to HREE, which we attribute to post-depositional mobilization of REE and addition of REE by alteration fluids. A simple model for the late addition of REE can explain these unusual spectral features for both detritus-rich and detritus-poor samples. The influence of this late REE addition is also evident in the Ce and Nd isotope systematics of these samples. La/Ce ratios were significantly altered by fluid addition, and the samples form a La-Ce isochron that constrains the timing of fluid addition to within the last 100 Ma. This is consistent with previous suggestions that negative Ce anomalies in ferruginous sediments of the Barberton Greenstone Belt are not primary features and confirms the utility of the La-Ce isotope approach for constraining the timing of oxidation as expressed by the Ce anomaly redox proxy. Sm/Nd ratios were less perturbed by fluid addition; nevertheless, the effect of fluid addition is evident in the Nd isotopic compositions of the samples. The deviation of the Sm-Nd isochron is proportional to the importance of fluid addition, as inferred by the magnitude of the negative Ce anomaly. This work highlights the need for careful consideration of the potential for post-depositional alteration to modify elemental and isotopic redox proxies. Caution is warranted when inferring the ancient presence of O₂ in the absence of direct geochronological constraints on the age of the proxy signal.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary material

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