An anoxic, Fe(II)-rich, U-poor ocean 3.46 billion years ago

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Abstract

The oxidation state of the atmosphere and oceans on the early Earth remains controversial. Although it is accepted by many workers that the Archean atmosphere and ocean were anoxic, hematite in the 3.46 billion-year-old (Ga) Marble Bar Chert (MBC) from Pilbara Craton, NW Australia has figured prominently in arguments that the Paleoarchean atmosphere and ocean was fully oxygenated. In this study, we report the Fe isotope compositions and U concentrations of the MBC, and show that the samples have extreme heavy Fe isotope enrichment, where $\delta^{56}$Fe values range between +1.5\% and +2.6\%, the highest $\delta^{56}$Fe values for bulk samples yet reported. The high $\delta^{56}$Fe values of the MBC require very low levels of oxidation and, in addition, point to a Paleoarchean ocean that had high aqueous Fe(II) contents. A dispersion/reaction model indicates that $O_2$ contents in the photic zone of the ocean were less than $10^{-3}$ \textmu M, which suggests that the ocean was essentially anoxic. An independent test of anoxic conditions is provided by U–Th–Pb isotope systematics, which show that U contents in the Paleoarchean ocean were likely below 0.02 ppb, two orders-of-magnitude lower than the modern ocean. Collectively, the Fe and U data indicate a reduced, Fe(II)-rich, U-poor environment in the Archean oceans at 3.46 billion years ago. Given the evidence for photosynthetic communities provided by broadly coeval stromatolites, these results suggest that an important photosynthetic pathway in the Paleoarchean oceans may have been anoxygenic photosynthetic Fe(II) oxidation.

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1. INTRODUCTION

The evolution of photosynthesis was a critical evolutionary step in the history of life on Earth (e.g., Canfield, 2005). It has been proposed that anoxygenic photosynthesis preceded oxygenic photosynthesis (e.g., Hohmann-Marriott and Blankenship, 2011), in part because the Archean atmosphere and oceans of Earth are generally considered to have been anoxic, and also because anoxygenic photosynthesis is deeply rooted in the 16S rRNA tree of life (Xiong et al., 2000). The first rise in atmospheric oxygen, termed the “Great Oxidation Event” (GOE), has been proposed to have occurred between $\sim$2.45 and 2.2 Ga (Holland, 1984, 2006). The GOE model was developed based on geologic occurrences of detrital pyrite, siderite, and uraninite in sediments deposited before the GOE, as well as retention of Fe in paleosols after the GOE (Holland, 1999), amongst other geological observations, although some workers have argued for a much earlier oxygenation of the atmosphere (e.g., Hoashi et al., 2009). Support for the GOE at $\sim$2.4 Ga comes from the disappearance of mass-independent fractionation of S isotope after the GOE (Farquhar et al., 2000). Recent geochemical studies, however, increasingly provide evidence for a more complex evolution of
atmospheric \( \text{O}_2 \) levels prior to the GOE as compared to a simple “step-function” (Anbar et al., 2007; Kaufman et al., 2007; Wille et al., 2007; Garvin et al., 2009; Godfrey and Falkowski, 2009; Ono et al., 2009; Duan et al., 2010; Kendall et al., 2010; Voegelin et al., 2010; Czaja et al., 2012; Reinhardt et al., 2013).

Attempts to constrain the evolution of oxygenic photosynthesis have been controversial. For example, the timing of the evolution of oxygenic photosynthesis has been partially constrained to \( \approx 2.7 \) Ga, based on molecular biomarkers (Brocks et al., 1999; Eigenbrode and Freeman, 2006; Eigenbrode et al., 2008; Waldbauer et al., 2009), although this line of research has been highly controversial (e.g. Rasmussen et al., 2008). Large microfossils of 3.4–3.1 Ga age have been interpreted to be possible eukaryotes or possible cyanobacteria, which, if confirmed, would suggest an even earlier origin for oxygenic photosynthesis (Sugitani et al., 2007, 2010; Javaux et al., 2010). Other researchers point to geological evidence that may indicate stellarlter oxygenation of the atmosphere, and therefore, a very early appearance of oxygenic photosynthesis. For example, hematite in the 3.46 billion-year-old (Ga) Marble Bar Chert Member (MBC), and in the stratigraphically younger Apex Basalt, from Western Australia, has been interpreted to reflect an \( \text{O}_2 \)-bearing Archean ocean and atmosphere in the Paleoarchean (Hoashi et al., 2009; Kato et al., 2009). Other workers have disagreed with an early timing of oxidation for the hematite-bearing Apex Basalt (Li et al., 2012).

In this study, we focus on the origin and paleo-environmental significance of hematite from the 3.46 Ga Marble Bar Chert Member (MBC) of the Duffer Formation, from the Pilbara Craton, Western Australia (Van Kranendonk et al., 2007b). Hoashi et al. (2009) interpreted hematite in the MBC to have precipitated from a fully-oxygenated Archean ocean at the time of deposition at 3.46 Ga, which in turn would suggest that oxygenic photosynthesis had evolved prior to that time. Alternatively, Van Kranendonk et al. (2008) suggested that hematite in jaspilite chert from the conformably underlying ca. 3.48 Ga Dresser Formation formed as a result of alteration of primary siderite during circulation of high pH hydrothermal fluids. Here, we combine Fe isotopes, which can constrain the extent of oxidation, with U–Th–Pb isotopes, which provide an estimate of seawater U contents as an independent measure of oxygen abundance, and as a test for post-depositional alteration of the host rocks. Our results indicate that hematite in the MBC was precipitated by a very small extent of oxidation from a Fe(II)-rich, U-poor ocean at 3.46 Ga. We conclude that hematite in the 3.46 Ga MBC cannot be used to infer an origin for oxygenic photosynthesis prior to 3.5 Ga.

2. GEOLOGICAL BACKGROUNDS AND SAMPLES

The Marble Bar Chert Member (MBC) member of the ca. 3.46 Ga Duffer Formation is the most prominent chert unit in the Warrawoona Group, which is the lowest stratigraphic part of the well-preserved Pilbara Supergroup in the Paleoarchean Pilbara Craton in northwestern Australia (Van Kranendonk et al., 2002, 2007a) (Fig. 1). Thick successions of pillow basalts and finely bedded cherts within the Warrawoona Group indicate predominantly deep marine deposition (Van Kranendonk et al., 2007a). The MBC is generally 100–200 m thick, and extends north–south for over 120 km, west of the town of Marble Bar (Fig. 1). The depositional age of the MBC is well constrained to 3.46 Ga by U–Pb zircon geochronology on conformably underlying felsic volcanic rocks of the Duffer Formation (Fig. 1; Van Kranendonk et al., 2007b). A bedding-parallel shear zone that developed locally along the contact between the MBC and the overlying Apex Basalt (Kato et al., 2009) reflects the effects of tilting during tectonic deformation between 3.3 Ga and 2.9 Ga (Van Kranendonk et al., 2007a). The MBC and Apex Basalt at Marble Bar were buried to at least 3 km depth prior to 2.78 Ga, at which time they were uplifted, eroded, and overlain by rocks of the Fortescue Group. Phanerozoic uplift has once more exposed these rocks to the effects of recent weathering (Thorne, 2001; Van Kranendonk et al., 2007b).

Hematite-bearing chert layers (jasper) are abundant in outcrops of the MBC (Appendix 1; Van Kranendonk, 2006, 2010). Van Kranendonk (2006) interpreted the jasper layers to predate white chert within the MBC, as the latter occur as cross-cutting veins or in situ replacement of the hematite-bearing chert. However, the Australian continent has been exposed to deep weathering since the Triassic (Anand, 2005), which has produced weathering profiles that are commonly 50–100 m deep in Western Australia (Anand and Paine, 2002; Anand, 2005). A critical question, therefore, is the age of hematite formation in the MBC – is it early Archean, or could it be Neoarchean or even Phanerozoic?

The Archean Biosphere Drilling Project (ABDP) was initiated with the goal of obtaining samples below surface weathering zones, and the site for the first diamond drill core from this program (ABDP-1) is located about 4 km southwest of the town of Marble Bar (Fig. 1). The 260 m deep drill core of ABDP-1 intersects basalts of the 3.474–3.463 Ga Duffer Formation, the MBC, and the 3.46 Ga Apex Basalt (Fig. 1). There are abundant hematite-bearing bands in the drill core samples, even at depths greater than 100 m (Van Kranendonk, 2010).

The MBC samples analyzed in this study come from drill core depths of 169 m and 176 m in ABDP-1 (Figs. 1–3), and were sampled at the Geological Survey of Western Australia, Perth. These samples were analyzed for U–Th–Pb and Fe isotopes. In addition, we report new Fe isotope data for the Apex Basalt samples analyzed for U–Th–Pb isotopes by Li et al. (2012) from depths of 190–262 m, which have been variably oxidized. A non-oxidized basalt sample from the Duffer Formation at 42.6 m depth was also analyzed (Fig. 1).

3. ANALYTICAL METHODS

3.1. Sample preparation

Sampling of chert was undertaken using a thin (0.2 mm) diamond saw blade. For Fe isotope analysis, a small wedge-shaped piece 1 × 1 mm in size was cut from a quarter drill
core for each hematite-bearing red layer of the MBC (Figs. 2 and 3). For U–Th–Pb isotope analysis, a larger slab (cm size, >0.15 g) was cut because of the very low U and Th concentrations in chert samples. Both hematite-rich red layers and hematite-poor white layers of the MBC were sampled. Samples were examined under a binocular microscope before and after cutting in order to ensure that cracks, veins, and other secondary features were avoided. Samples were cleaned using acetone, 0.2 M HCl, and 18.2 MΩ H₂O in an ultrasonic bath for more than 10 min each to remove surface contamination, before being dried and weighed. This procedure ensured complete removal of any surface Fe, Pb and U contamination that may have been introduced during coring or sampling. Powder
samples of Apex Basalt and Duffer Formation basalt that were analyzed for Fe isotope compositions were the same as those studied by Li et al. (2012), and sample preparation details may be found in that study.

Following the sample cleaning procedure noted above, sample digestion and ion-exchange chromatography were carried out in a clean room with HEPA-filtered air. Samples were digested using a mixture of double-distilled HF and Optima grade HNO₃ in capped Savillex beakers that were heated overnight at 130 °C. Dissolved samples were converted to chloride form using double-distilled 8 M HCl and solutions were checked under a binocular microscope to ensure that the entire sample had dissolved and that no fluorides were present.

3.2. Iron isotope measurements

Prior to Fe isotope analysis, a small aliquot of each dissolved sample was taken for total Fe concentration measurement using the Ferrozine method (Stooley, 1970).

Fig. 2. Variation of Fe and U–Th–Pb isotope data along ABDP-1 drill core at a depth of 169.6–169.8 m. Blue triangles on core photo denote samples for Fe isotope analysis, and the adjacent small-font numbers denote depth in mm. Blue rectangles on core photo denote samples for U–Th–Pb isotope analysis. A, B, and C denote different sides of the quarter drill core. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

Fig. 3. Variation of Fe and U–Th–Pb isotope data along ABDP-1 drill core at a core depth of 176.9–177.0 m. Blue triangles on core photo denote samples for Fe isotope analysis, and the adjacent small-font numbers denote depth in mm. Blue rectangles on core photo denote samples for U–Th–Pb isotope analysis. A, B, and C denote different sides of the quarter drill core. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
Approximately 100 μg of Fe from each sample was purified using anion-exchange resin (BioRad AG 1X4 200–400 mesh resin) and HCl (e.g., Beard et al., 2003). Iron solutions were diluted to 600 ppb and isotopic measurements were conducted using a Micromass IsoProbe MC-ICP-MS and an Aridus desolvating nebulizer that aspirated at ~50 μL/min. Mass spectrometry followed the procedures reported by Beard et al. (2003). Isotopic data are reported as $^{56}$Fe/$^{54}$Fe and $^{57}$Fe/$^{54}$Fe ratios in standard delta (δ) notation, in units of per mil ($\delta$), and the average of igneous rocks as the standard reference reservoir (Beard et al., 2003):

$$\delta^{56}\text{Fe} = \left[\frac{(^{56}\text{Fe}/^{54}\text{Fe})_{\text{sample}}}{(^{56}\text{Fe}/^{54}\text{Fe})_{\text{standard}}} - 1\right] \times 1000$$

(1)

$$\delta^{57}\text{Fe} = \left[\frac{(^{57}\text{Fe}/^{54}\text{Fe})_{\text{sample}}}{(^{57}\text{Fe}/^{54}\text{Fe})_{\text{standard}}} - 1\right] \times 1000$$

(2)

Relative to the average of igneous rocks, the international Fe isotope standard IRMM-014 has a δ$^{56}$Fe value of −0.09‰ on this scale (Beard et al., 2003). The external long-term reproducibility (2-SD) for δ$^{56}$Fe measurements using this method is ±0.08‰, as determined from analysis of multiple in-house Fe standard solutions, and synthetic samples (Fe standard solutions doped with matrix elements) that were processed through the ion-exchange procedure together with drill core samples (Appendix 2).

3.3. U–Th–Pb isotope measurements

For U–Th–Pb isotope analysis, each digested sample was divided into two aliquots; one aliquot was used for Pb isotope analysis and the other was spiked with a mixed $^{235}$U–$^{238}$Th. $^{206}$Pb tracer to determine U, Th, and Pb concentrations by isotope-dilution mass spectrometry. Solutions were chemically purified by anion-exchange chromatography, where initial cation separation was done using 0.6 M HBr, followed by extraction of Pb using 6 M HCl (Li et al., 2012). The U and Th in the 0.6 M HBr wash was purified using a second anion-exchange column using HNO$_3$, followed by HCl to separate a combined U and Th cut from the rest of the sample (Li et al., 2012). Total procedural blanks for Pb chemistry was 50–80 pg, which was less than 0.1% of the amount of Pb in the samples. Total procedural blanks for Th and U were <50 pg, which is less than 10% of the amount of Th and U in the samples. Isotope-dilution analysis of Pb, U, and Th produced concentration uncertainties of <0.03%, <0.23%, and <0.08%, respectively.

Lead isotope ratios were determined on a VG Instruments Sector 54 thermal ionization mass spectrometer using a static multi-collector routine and a $^{208}$Pb ion signal of 1.7–2.0 $\times 10^{-11}$ A. Lead was loaded onto single Re filaments and run using the Si–gel H$_3$PO$_4$ technique. Lead isotope ratios were corrected for mass fractionation by +0.125% per amu, as constrained by long-term measurement of the $^{207}$Pb/$^{206}$Pb ratio of NIST SRM-981 and the $^{206}$Pb/$^{207}$Pb ratio of SRM-982. External precision of Pb isotope analysis was ±0.052% per amu (2 SD, n = 30), based on repeat measurement of SRM-981 and SRM-982 during the analytical session.

Isotopic ratios of $^{235}$U/$^{238}$U and $^{229}$Th/$^{232}$Th were simultaneously measured using a Micromass IsoProbe MC-ICP-MS and a 50 μl/min self-aspirating, concentric-flow nebulizer and an Aridus$^{\circ}$ desolvating nebulization system. Instrumental mass bias was corrected using a sample-standard bracketing technique, using mixtures of Ames Th mixed with NBL-114 natural U ($^{238}$U/$^{235}$U = 137.88) as bracketing standards. The samples were diluted to match the total ion intensity of the standards. There were no memory effects during analysis, as on-peak zero measurements of U and Th remained unchanged after washout between samples. Tailing effects were negligible, and correction for abundance sensitivity did not shift the results outside analytical error.

4. RESULTS

Iron isotope compositions and concentrations for MBC and basaltic samples, and Pb isotope compositions and U–Th–Pb concentrations for MBC samples, are given in Appendix 2. Iron and U–Th–Pb isotope data for MBC samples are correlated to drill cores in Figs. 2 and 3. Iron isotope compositions of the basaltic samples are plotted against Fe content, Fe($^{III}$/Fe$_{\text{Total}}$) and Fe/Th ratios in Fig. 4.

4.1. Iron isotope results

The MBC samples have very high δ$^{56}$Fe values, ranging between +1.53‰ and +2.63‰ (Figs. 2 and 3; Appendix 2), that define the upper limit of measured δ$^{56}$Fe values for bulk terrestrial samples. The δ$^{56}$Fe values of samples from a drill core depth of 169.6–169.8 m vary between +1.98‰ and +2.63‰ (Fig. 2). These are systematically higher than δ$^{56}$Fe values of samples from a drill core depth of 176.90–170.00 m (Fig. 3). There is a steadily decreasing trend in δ$^{56}$Fe values from +2.63‰ at 169.61 m to +2.03‰ at 169.65 m (Fig. 2), and an increase in δ$^{56}$Fe values from +1.53‰ at 176.97 m to +1.88‰ at 177 m (Fig. 3). Other than these relations, there is no significant variation in δ$^{56}$Fe values between layers (Table EA-1). There is no correlation between δ$^{56}$Fe values and Fe concentrations, nor between isotopic composition and color (red/black) of the hematite-bearing layers. It should be noted that the Fe isotope compositions of hematite-poor white layers are consistent with those of the hematite-rich (red) layers (Figs. 2 and 3), indicating that δ$^{56}$Fe values do not correlate with Fe contents.

In contrast to the MBC samples, basaltic samples from the ABDP-I drill core have much lower δ$^{56}$Fe values, ranging between −0.40‰ and +0.66‰ (Fig. 4). The majority of these samples cluster around 0‰, and the weighted average δ$^{56}$Fe value for all basaltic samples is 0.00‰ equal to the average of igneous rocks (Beard et al., 2003). The δ$^{56}$Fe values of basaltic samples only become significantly variable as Fe($^{III}$/Fe$_{\text{Total}}$) ratios increase to >0.9 (Fig. 4). There is no correlation between the δ$^{56}$Fe values and total Fe concen-
trations in the basalts, nor between $\delta^{56}$Fe values and Fe/Th ratios (Fig. 4).

### 4.2. U–Th–Pb isotope results

Lead concentrations in the MBC samples vary between 0.37 and 12.95 ppm, U between 3 and 60 ppb, and Th between 4.5 and 42.23 ppb (Appendix 2). Concentrations of Pb, U, and Th in the MBC samples are systematically lower than those of the basaltic samples from the same drill core, which are 1–58 ppm for Pb, 80–1040 ppb for U, and 233–694 ppb for Th (Li et al., 2012). Importantly, U contents are much lower than those of modern oceanic sediments (0.306–4.889 ppm, global average 1.68 ppm; Plank and Langmuir, 1998). The MBC samples have very non-radiogenic Pb isotope compositions (low $\delta^{206}$Pb/$\delta^{204}$Pb, $\delta^{207}$Pb/$\delta^{204}$Pb and $\delta^{208}$Pb/$\delta^{204}$Pb ratios relative to average crust) that, in general, overlap those of basaltic samples from the ABDP-1 drill core. The $\delta^{206}$Pb/$\delta^{204}$Pb ratios of samples from a drill core depth of 169.6–169.8 m vary between 13.699 and 13.891 (Fig. 2), which are systematically higher than the $\delta^{206}$Pb/$\delta^{204}$Pb ratios of samples from a drill core depth of 176.90–170.00 m that vary between 12.684 and 13.153 (Figs. 2 and 3).

### 5. DISCUSSION

#### 5.1. Contrasting origin for hematite in the Marble Bar Chert and Apex Basalt

The hematite-bearing bands of the MBC have the highest $\delta^{56}$Fe values ever reported from natural bulk rocks (Fig. 5). These contrast sharply with the igneous and near-igneous values (average $\delta^{56}$Fe ~0‰; Beard et al., 2003) of variably oxidized samples of the Apex Basalt (Fig. 4). The large contrast in Fe isotope compositions between hematite in the MBC and that in the oxidized portions (hematite and goethite; Kato et al., 2009) of the Apex Basalt indicates distinct processes of formation.

The small variation in $\delta^{56}$Fe values of the Apex Basalt (Fig. 4 and Table S1), averaging 0‰, suggests that Fe in Apex Basalt is primarily igneous in origin. The limited range in $\delta^{56}$Fe values, which occurs only in the most oxidized samples, most likely reflects small extents of internal redistribution of Fe during oxidation. The fact that the two samples that have low $\delta^{56}$Fe values also have high Fe/Th ratios, and that the samples with the highest $\delta^{56}$Fe values also have low Fe/Th ratios (Fig. 4) implies that some samples might have released isotopically light Fe into fluids during weathering. This isotopically light Fe was then oxidized and re-precipitated, causing enrichment of isotopically light Fe oxides. Iron mobility could have been promoted by weathering of sulfides that were formed at 2.76 Ga at Marble Bar (Kato et al., 2009), which produced a locally acidic and redox-active environment that might enable small-scale Fe mobility and Fe isotope fractionation. Although Kato et al. (2009) suggest that oxidation occurred in the Archean, Li et al. (2012) documented Phanerozoic U mobility that correlated with the extent of oxidation,
indicating that oxidation of the Apex Basalt most likely occurred via channelled groundwater flow during deep Phanerozoic weathering. Under such conditions, large-scale Fe mobility would not be expected, and this is supported by the limited range in $\delta^{56}$Fe values and the fact that the average $\delta^{56}$Fe value lies at the value for igneous rocks.

In the MBC, the very high $\delta^{56}$Fe values rule out the possibility that Fe was transported from the Apex Basalt. Moreover, the very high $\delta^{56}$Fe values of the MBC are inconsistent with hematite formation by in situ oxidation of Fe(II)-bearing minerals such as siderite, as has been proposed for some jaspers in the underlying Dresser Formation (Van Kranendonk et al., 2008). Siderite that precipitated from an Archean ocean should have had a $\delta^{56}$Fe value below $-0.5_{\text{oo}}$ (Polyakov and Mineev, 2000; Wiesli et al., 2004; Johnson et al., 2008; Rustad et al., 2010), and in situ alteration to hematite should retain the negative $\delta^{56}$Fe values. Furthermore, the low solubility of Fe(III) oxides and hydroxides at circum-neutral pH (e.g., Kuma et al., 1996, and references therein) makes it unlikely that significant quantities of Fe could be leached from the MBC under oxidized conditions. We therefore conclude that the hematite in the MBC was not produced by the Phanerozoic oxidation event that oxidized the Apex Basalt at Marble Bar. This interpretation is consistent with the fracture and vein patterns in the drill core (Fig. 1C), that suggest groundwater movement was likely along channelled fracture systems.

5.2. Isotopic constraints of oxidation of Fe(II) in the Archean ocean

The likely source of aqueous Fe(II) that was oxidized to form hematite in the MBC was hydrothermal fluids, which should have had a $\delta^{56}$Fe value of around $0_{\text{oo}}$, or slightly negative (Yamaguchi et al., 2005; Johnson et al., 2008). Oxidation of aqueous Fe(II) in modern hydrothermal systems produces precipitates that have slightly negative $\delta^{56}$Fe values (Fig. 5), reflecting essentially quantitative oxidation. The very high $\delta^{56}$Fe values of hematite in the MBC, therefore, do not support a fully oxidized Archean ocean, as proposed by Hoashi et al. (2009).

Rather, the very high $\delta^{56}$Fe values can only be explained by partial oxidation of aqueous Fe(II), given the $\sim3-4_{\text{oo}}$ fractionation in $^{56}$Fe/$^{54}$Fe between oxides and aqueous Fe(II) (Wu et al., 2012).

Constraints on the extent of oxidation of aqueous Fe(II) in an Archean ocean can be made using a one-dimensional dispersion/reaction model, which assumes aqueous Fe(II) released from hydrothermal vents diffuses upwards to the photic zone, followed by oxidation of Fe(II)$_{aq}$ to Fe(III) hydroxides (Czaja et al., 2012, 2013). The limitation of the dispersion/reaction model is that it may not accurately describe upwelling of hydrothermal Fe(II), as the solution of the model relies on a steady-state condition of the ocean. Nevertheless, the dispersion/reaction model is superior compared with a simple Rayleigh fractionation model that is commonly used in geochemical studies, which is not an appropriate model for interpreting Fe isotope fractionation during oxidation in the photic zone because it is a closed-system model and does not account for a continual influx of Fe(II)$_{aq}$ and outflux of Fe(OH)$_3$ precipitates. Using the dispersion/reaction model of Czaja et al. (2012, 2013), we produced vertical profiles for concentrations and isotope compositions of Fe(II)$_{aq}$ and Fe(OH)$_3$, at different rates of Fe(II)$_{aq}$ oxidation (Appendix 3). In the model, oxidation could occur either anaerobically or aerobically, which simulates oxidation by either anoxygenic Fe(II)-oxidizing photosynthetic bacteria, or by free oxygen generated by oxygenic photosynthetic bacteria, respectively; in the latter case, our model can provide constraints on the amount of free oxygen that may have been present.

Dispersion/reaction model of Czaja et al. (2012, 2013) produces broadly similar profiles of concentrations and isotopic compositions of Fe(II)$_{aq}$ and Fe(OH)$_3$ over a wide range of input parameters (Appendix 3). Although many input parameters may be varied in the model, the most important factor that affects the results is the Fe(II) oxidation rate ($f_{\text{Fe-oxidation}}$). At a high rate of Fe(II) oxidation, which may be produced either by high O$_2$ from photosynthesis or by high activity of anoxygenic Fe(II)-oxidation, a low-Fe(II)$_{aq}$ zone is developed in the uppermost part of the photic zone. At such high rates of oxidation, the weighted average $\delta^{56}$Fe value for Fe(OH)$_3$ is close to $0_{\text{oo}}$ (Fig. 6; Appendix 3), reflecting quantitative Fe(II) oxidation of the Fe(II) hydrothermal input, assumed to have $\delta^{56}$Fe = $0_{\text{oo}}$. At lower Fe(II) oxidation rates, however, high Fe(II)$_{aq}$ concentrations develop in the photic zone, which in turn decreases the upwards Fe(II)$_{aq}$ flux ($f_{\text{Fe-transport}}$), as required to maintain a steady-state condition in the dispersion/reaction model ($f_{\text{Fe-transport}} = f_{\text{Fe-oxidation}}$). This relation occurs because the Fe(II)$_{aq}$ flux is a function of the Fe(II)$_{aq}$ concentration gradient in the water column and the eddy dispersion coefficient. Under these conditions, Fe(II)$_{aq}$ in the photic zone is partially oxidized, and the oxidation product, Fe(OH)$_3$ precipitates, has positive $\delta^{56}$Fe values (Fig. 6; Appendix 3).

Dispersion/reaction modeling shows a rough linear relation between $\delta^{56}$Fe values of Fe(III) hydroxides and the oxidation flux ($f_{\text{Fe-oxidation}}$) between zero and $f_{\text{ref}}$, a parameter defined as the maximum diffusive Fe(II) flux from the Fe(II) source to the photic zone ($f_{\text{ref}} = D \times \Delta C_{\text{ref}}/\Delta z_{\text{ref}}$; Kappler et al., 2005) (Fig. 6). D is the eddy dispersion coefficient, $\Delta C_{\text{ref}}$ is the Fe concentration difference between Fe(II) source and Fe-depleted zone (C$_{\text{(Fe(II)-source=0)}}$), and $\Delta z_{\text{ref}}$ is the distance between the Fe(II) source and the bottom of the photic zone. The slope of the rough linear relation is dependent on the Fe isotope fractionation factor between aqueous Fe(II) and Fe(III) hydroxides (Fig. 6). At high rates of photosynthesis, complete depletion of Fe(II) develops in the top of the photic zone. In the case of anoxygenic photosynthesis, microbes are concentrated in the bottom of the photic zone and are limited by access to Fe(II). This is a scenario that has been proposed and discussed by Kappler et al. (2005) for anoxygenic photosynthetic oxidation of Fe(II). In the case of high oxygenic photosynthesis, complete depletion of Fe(II) also develops in the top of the photic zone at high rates of oxygen production, accompanied by a build-up of free O$_2$, which is transported downward in the water column to react with Fe(II) below the photic zone. It should be noted that $f_{\text{Fe-oxidation}}$ can be greater than...
and, in this case, free O$_2$ begins to build up and the oxygenated water column expands downwards to below the photic zone, which decreases the diffusion distance of aqueous Fe(II) at a given depth. In both cases (anoxygenic or oxygenic photosynthesis), close-to-zero $\delta^{56}$Fe values for Fe(III) hydroxides are predicted, as Fe(II) is quantitatively oxidized.

The modeling results indicate that the $\delta^{56}$Fe values of the ferric hydroxides/oxides may be used as a proxy for Fe(II)$_{aq}$ abundance in the photic zone. The high $\delta^{56}$Fe values of the MBC iron oxides require a reduced, Fe(II)$_{aq}$-rich, photic zone. This, in turn, places constraints on permissible oxygen contents, which vary as a function of $\delta^{56}$Fe values of Fe(OH)$_3$ precipitates (Fig. 7). Important variables are the $^{56}$Fe/$^{54}$Fe fractionation factor between Fe(OH)$_3$ and Fe(II)$_{aq}$, and the depth of the basin, the latter of which exerts an influence on the rate of Fe(OH)$_3$ precipitation relative to Fe(II)$_{aq}$ influx. The small changes in $\delta^{56}$Fe values in the MBC core may reflect subtle changes in extent of Fe(II) oxidation. The fact that the $\delta^{56}$Fe values in a particular section of core do not correlate with Fe-rich or Fe-poor banding (Figs. 2 and 3) suggests that the variable Fe concentrations in MBC bands reflect variations in Fe(III) oxide accumulation rates (or SiO$_2$ precipitation rate) at the time of deposition, rather than changes in the extent of oxidation. In all cases, however, the O$_2$ contents estimated from the Fe(II) oxidation rate increases, aqueous Fe(II) contents in the photic zone decrease, and eventually aqueous Fe(II) becomes depleted in the top of photic zone. With increasing activity of oxygenic photosynthesis, free O$_2$ begins to build up and the O$_2$-bearing water column expands downwards to below the photic zone, which increases the diffusion distance of aqueous Fe(II) at a given depth from the Fe(II) source. The relations plotted in Fig. 6 were specifically chosen because they are insensitive to parameters and boundary conditions of the dispersion/reaction model. The most important parameters that determine the $\delta^{56}$Fe values of Fe(III) hydroxide precipitates is the extent of oxidation and the Fe(OH)$_3$–Fe(II)$_{aq}$ Fe isotope fractionation factor.

$f_{\text{ref}}$ and, in this case, free O$_2$ builds up and the oxygenated zone expands downwards to below photic zone, which decreases the diffusion distance of aqueous Fe(II) (Fig. 6B).

Fig. 6. Summary results of dispersion/reaction modeling of the Fe(II) oxidation at 3.46 Ga Marble Bar assuming Fe(II) oxidation by anoxygenic photosynthetic Fe(II) oxidizers (A), or by O$_2$ produced by oxygenic photosynthesis (B). $\Delta^{56}$Fe$_{\text{Fe(OH)3-Fe(II)aq}}$ is the Fe isotope fractionation factor between ferric hydroxides and aqueous Fe(II). Fe isotope composition of Fe(OH)$_3$ and Fe(II) concentrations at the top and bottom of photic zone are plotted against the $f_{\text{Fe-oxidation}}/f_{\text{ref}}$ Ratio. $f_{\text{Fe-oxidation}}$ is the rate of Fe oxidation expressed as the flux of oxidized Fe that is settling as Fe(OH)$_3$,$f_{\text{ref}}$ is the maximum flux of Fe(II) from the source to the bottom of photic zone by diffusion (for details, see Section 5.2). At low rates of Fe(II) oxidation, such as $f_{\text{Fe-oxidation}} < 0.8f_{\text{ref}}$, the reducing capacity of Fe(II) provided by diffusion from the hydrothermal Fe(II) source dominates the photic zone, and even the top of photic zone contains significant amounts of aqueous Fe(II). As the Fe(II) oxidation rate increases, aqueous Fe(II) contents in the photic zone decrease, and eventually aqueous Fe(II) becomes depleted in the top of photic zone. The fact that the $\delta^{56}$Fe values of Fe(III) hydroxide precipitates are insensitive to parameters and boundary conditions of the dispersion/reaction model. The most important parameters that determine the $\delta^{56}$Fe values of Fe(III) hydroxide precipitates is the extent of oxidation and the Fe(OH)$_3$–Fe(II)$_{aq}$ Fe isotope fractionation factor.

Fig. 7. Summary results of dispersion/reaction modeling showing the relation between average photic zone O$_2$ concentrations and the weighted average $\delta^{56}$Fe values of the Fe(OH)$_3$ produced, assuming Fe(II) oxidation by O$_2$ produced by oxygenic photosynthesis. The curves represent results for Fe isotope fractionation factors ($\delta^{56}$Fe$_{\text{Fe(OH)3-Fe(II)aq}}$) of 4.0‰ and 2.5‰ in a 200 m basin and a 500 m basin. The shaded area represents the $\delta^{56}$Fe value range measured in the MBC samples.
mated for the photic zone are less than $10^{-3}$ μM to produce the high measured δ$^{56}$Fe values, which is less than 0.0003% of modern O$_2$ contents in the photic zone. This conclusion is robust and is insensitive to changes in input parameters, including choice of Fe(OH)$_3$–Fe(II)$_{aq}$ Fe isotope fractionation factors, although the later parameter would be important if attempting to distinguish between O$_2$ contents of $10^{-3}$ to $10^{-6}$ μM (Fig. 7). For all practical purposes, however, all calculated O$_2$ contents of $<10^{-3}$ μM can be considered to indicate anoxic conditions in the photic zone, and this conclusion contrast greatly with the proposal of Hoashi et al. (2009) that the Fe oxides in the MBC reflect deposition in an oxygenated ocean.

5.3. U–Th–Pb isotope constraints on Archean ocean U contents

Uranium abundances in seawater will broadly correlate with oxygen contents because of the high solubility of U(VI) species relative to reduced species (Langmuir, 1978). Below we estimate the maximum seawater U contents of the Paleoarchean seawater from which the MBC precipitated, which provides an independent estimate for seawater oxygen contents relative to those based on Fe isotopes. Our approach is to estimate U contents using U–Th–Pb geochronology, which provides insights into changes due to radioactive decay and post-formation alteration. These issues are critical to assess in such ancient rocks because they bear on the confidence with which U can be used as a paleo-environmental proxy.

5.3.1. Post-depositional mobility of U and Pb in the MBC

It is possible that the low measured U contents in the MBC reflect post-depositional U loss through leaching by oxygenated groundwater circulation, but this can be critically evaluated using U–Th–Pb isotopes. The 206$^{Pb}$/204$^{Pb}$–208$^{Pb}$/204$^{Pb}$–207$^{Pb}$/204$^{Pb}$ variations of the MBC indicate non-radiogenic isotope compositions, plotting along Pb–Pb arrays that correspond to low-$^{238}$U/$^{204}$Pb ratios (μ) between 0 and 4 (Fig. 8), indicating long-term low U/Pb ratios. The MBC samples have Pb isotope compositions that overlap the basaltic samples in the ABDF-1 drill core, and lie between two Pb isotope end members (i.e., the non-radiogenic "ore lead" and the radiogenic "basalt lead") that were previously reported from the Warrawoona Group (Fig. 8; Li et al., 2012). On a 206$^{Pb}$/204$^{Pb}$–207$^{Pb}$/204$^{Pb}$ plot, the MBC samples scatter about a line that has a slope equivalent to an age of 3435 ± 140 Ma and a MSWD of 2.9 (insert of Fig. 8B; best-fit line calculated using Isoplot v2.49 (Ludwig, 1999), where σ errors for Pb isotope ratios are ±0.052%/amu). This age matches the depositional age of the MBC that is constrained by zircon U–Pb geochronology (Fig. 1). It is important to note, however, that all Pb reservoirs in the region liealong the isochron (Fig. 8), and therefore mixing relations will produce the same array (see discussion in Li et al., 2012). The MBC samples do not, however, plot on isochrons that correspond to a 3.46 Ga age on plots of 206$^{Pb}$/204$^{Pb}$–238$^{U}$/204$^{Pb}$ and 206$^{Pb}$/204$^{Pb}$–232$^{Th}$/204$^{Pb}$ (Fig. 9), indicating open-system behavior for the MBC samples.

![Fig. 8.](image-url) On a 208$^{Pb}$/204$^{Pb}$–232$^{Th}$/204$^{Pb}$ diagram (Fig. 9B), all of the MBC chert samples plot to the left side of the reference isochron. Because Th should be relatively immobile, the very low 232$^{Th}$/208$^{Pb}$ ratios likely reflect Pb addition, and the range of 208$^{Pb}$/204$^{Pb}$ ratios can be explained as a mixture of low-206$^{Pb}$/204$^{Pb}$–208$^{Pb}$/204$^{Pb}$ "ore Pb" and high–208$^{Pb}$/204$^{Pb}$–207$^{Pb}$/204$^{Pb}$ "basalt Pb" components that have been added to the chert. The data relations on a 206$^{Pb}$/204$^{Pb}$–238$^{U}$/204$^{Pb}$ diagram (Fig. 9A) are also consistent with Pb addition, although mobility of U is also possible, and is evaluated in Section 5.3.2 below. The isochron diagrams in Fig. 9A suggest that both the MBC and Apex
Basalt have been influenced by the same event(s), as Pb addition has also been reported in the variably oxidized Apex Basalt (Li et al., 2012). Addition of "ore Pb" could have occurred at any time since 3.46 Ga because "ore Pb" defines the starting point of the $^{206}\text{Pb}/^{204}\text{Pb}$–$^{207}\text{Pb}/^{204}\text{Pb}$ "isochron" (see discussion in Li et al., 2012), whereas addition of "basalt lead" likely occurred recently, because the slope of the mixing trend on the $^{206}\text{Pb}/^{204}\text{Pb}$–$^{207}\text{Pb}/^{204}\text{Pb}$ diagram (Fig. 8B) represents an age that is consistent with the depositional age.

A key question is the effect of Pb addition to the U/Pb ratios, which is important to understand before U contents of the MBC at the time of deposition can be estimated. To account for the influence of Pb addition to the MBC, we evaluate the U/Pb ratios in the MBC by modeling the radiogenic decay of the primary depositional U component and compare the model results to measured results (Fig. 10). We assume a reference that represents samples that contained 5 ppm Pb with variable Pb isotope compositions ($\mu = 0.1–3$, $\kappa = 0.25$; for definitions of $\mu$ and $\kappa$, see Faure 1986), to approximate the measured Pb isotope compositions of the MBC samples, and these reference values define a family of lines in a $^{208}\text{Pb}/^{204}\text{Pb}$–$^{206}\text{Pb}/^{204}\text{Pb}$ plot (Fig. 10). The importance of incorporating $^{208}\text{Pb}/^{204}\text{Pb}$ ratios lies in the relative immobility of $^{232}\text{Th}$, the parent to $^{208}\text{Pb}$. We define $U^*/\text{Pb}$ as the deviation in measured U/Pb ratios relative to those required to produce the least radiogenic Pb isotope ratios of the MBC. $U^*$ could be viewed as "excess U" in the MBC at 3.46 Ga ago and it is further described mathematically in the following section (Section 5.3.2). Modeling results in Fig. 10 indicate that the Pb isotope deviations define a narrow range between 0 and $+0.8\%$ for $U^*/\text{Pb}$ ratios. In contrast, if we assume that 3.46 Ga U contents were 100 or 300 ppb higher ($U^*/\text{Pb} = +2\%$ and $+6\%$, respectively), but had recently decreased, the measured $^{206}\text{Pb}/^{204}\text{Pb}$–$^{208}\text{Pb}/^{204}\text{Pb}$ variations would have been highly skewed toward high $^{206}\text{Pb}/^{204}\text{Pb}$ ratios (Fig. 10), which is not observed. This conclusion is robust and independent of issues related to Pb addition, as U/Pb ratios are used in the model (Fig. 10). We cannot, however, rule out the possibility of U addition, a Phanerozoic event observed in the Apex Basalt adjacent to the MBC (Li et al., 2012). Therefore, the measured U contents in MBC samples ultimately will only constrain the upper limit of U contents in the MBC at the time of deposition at 3.46 Ga.

5.3.2. Estimating Archean seawater U concentration

A complete assessment of seawater U contents at the time of deposition of the MBC requires a mass-balance relation that accounts for four U components in the MBC samples: (1) U adsorbed by Fe(III) hydroxides during deposition in an Archean ocean ($U_{\text{adsorbed}}$) at 3.46 Ga, (2) an Archean detrital U component ($U_{\text{detrital}}$), (3) possible U added by a Phanerozoic event ($U_{\text{Phanerozoic}}$), and (4) the decrease in U contents of the Archean components due to $^{238}\text{U}$ and $^{235}\text{U}$ decay. The possibility of U addition in the Phanerozoic is based on the evidence presented by Li et al. (2012) for the Apex Basalt. Collectively, these terms may be described by the equation:

$$U_{\text{measured}} = U_{\text{adsorbed}} + U_{\text{detrital}} + U_{\text{Phanerozoic}} - U_{\text{decay}} \quad (3)$$

The detrital component of U can be constrained using the measured Th contents and a U/Th ratio of 0.25 that approximates that of average crust (Wignall and Myers, 1988; Chung and Chang, 1996; Fisher and Wignall, 2001). The amount of U decrease due to radioactive decay is calculated to be $47\%$, based on decay constants of $^{235}\text{U}$ and $^{235}\text{U}$, present-day $^{238}\text{U}/^{235}\text{U}$ ratio, and an age of 3.46 Ga. The amount of U added during the Phanerozoic is difficult to constrain. An upper limit on U adsorbed, nevertheless, may be constrained by assuming $U_{\text{Phanerozoic}}$ is zero, although we postulate that the estimated $U_{\text{adsorbed}}$ content could be an order of magnitude lower than the upper.

![Fig. 9](https://example.com/image-url)
bounds calculated here, given that Li et al. (2012) documented increases of up to 890% in the U/Th ratios of the ABDP-1 basalts during the Phanerozoic.

Uranium contents of seawater may be constrained using the \( k_d \) values discussed above, the aqueous oxides/hydroxides, adsorption coefficients for \( U \) are very high for Fe(III) oxides and hydroxides, more than three-orders-of-magnitude higher than those for \( U \) adsorption to silicates, including SiO2 phases (Waite et al., 1992), indicating that Fe(III) oxides/hydroxides should have controlled the \( U \) budget of the MBC. Partition coefficients \( (k_d) \) for \( U \) between Fe(III) hydroxide and aqueous solution are pH and pCO2 dependent (Wazne et al., 2003; Davis et al., 2004). There is a general consensus that the pH value for Archean seawater was below that of the modern ocean, perhaps 7.5 or lower, based on expected higher atmospheric CO2 contents (Walker, 1983; Grozinger and Kasting, 1993; Ohmoto et al., 2004). Under these conditions, the \( k_d \) between Fe(III) hydroxide and aqueous solution varies between 106 and 108 (Wazne et al., 2003; Davis et al., 2004). Using a conservative \( k_d \) of 106 and the \( U^{*}/Fe_2O_3 \) ratios of the MBC \( (U^* = U^{\text{measured}} + U^{\text{decay}} - U^{\text{detr}}) \), upper limits for Archean seawater \( U \) contents during precipitation of the MBC are estimated at between 0.02 and 0.75 ppb (Fig. 11). Given the possibility of Phanerozoic \( U \) addition discussed above, such estimates are upper bounds and the seawater value at the time of deposition could be lower than 0.02 ppb \( U \). We note that the highest inferred \( U^{*}/Fe_2O_3 \) values are those measured in samples that have the lowest Fe contents (Fig. 11), which suggests that the highest \( U \) contents inferred for Archean seawater are those samples that could be most susceptible to Phanerozoic \( U \) addition, given their low Fe contents. We therefore conclude that the \( U \) concentration of 3.46 Ga seawater was at least two orders of magnitude lower than that of modern seawater (3 ppb; see refs in Barnes and Cochran, 1990). Although there may be refinements that could be made to \( k_d \) estimates with new experimental data, we have chosen a conservative value. Uncertainties in the detrital component are less than 5%, given the narrow range in crustal \( U/Th \) ratios (e.g., Li et al., 2012).

Uranium occurs in primary igneous rocks as insoluble \( U(IV) \) compounds that, when oxidized, form highly soluble \( U(VI) \) species (Langmuir, 1978). The low redox potential of the \( U(VI)/U(IV) \) couple (0.27 V, Langmuir, 1978) makes \( U \) an element that is very sensitive to continental oxidation. The low \( U \) content in Archean seawater inferred here therefore complements geological evidence of detrital uraninite preservation in the Archean (Rasmussen and Buick, 1999), which attests to an anoxic Archean atmospheric and ocean conditions at 3.46 Ga. Such results are consistent with the implications of the high \( \delta^{18}Fe \) values discussed above. Our results do not agree with the study by Rosing and Frei (2004), who argue for \( U \)-rich Archean seawater based on complex and partially reset \( Pb \) isotope compositions of 3.8 Ga metasedimentary rocks of the Isua Supracrustal Belt of southern West Greenland.

5.4. Implications for photosynthesis in the Archean ocean

The Fe isotope compositions of the MBC are consistent with the suggestion by Hoashi et al. (2009) that hematite in the MBC formed via oxidation of \( Fe(II) \) in the Archean ocean. Iron isotope data, however, rule out an oxygenated ocean as the cause of this \( Fe(II) \) oxidation, as proposed by these authors. Importantly, our modeling demonstrates that the occurrence of \( Fe(III) \) minerals cannot be taken as evidence for oxygenation of the atmosphere. There are several possible mechanisms for \( Fe(II) \) oxidation, including UV photo-oxidation, anaerobic photosynthetic \( Fe(II)-oxi-
dizing bacteria, or free oxygen produced by oxygenic photosynthesis. Although UV photo-oxidation of Fe(II)$_{aq}$ has been proposed as a mechanism for Fe(III) oxide precipitation in the Archean (Brateman et al., 1983), this is not supported by the experimental work of Konhauser et al. (2007), which demonstrated that UV photo-oxidation occurs at a rate much slower than the rate of precipitation of Fe(II) minerals such as greenschist and siderite under Archean seawater conditions (high dissolved silica, Fe(II), and high pCO$_2$). Although additional experimental work exploring a range of conditions may be required to fully rule out UV photo-oxidation, the evidence at hand suggests this is an unlikely process. Precipitation experiments indicate that siderite is isotopically lighter than aqueous Fe(II) by ca. 0.5‰ in $\delta^{56}$Fe values (Wiesli et al., 2004), and greenschist probably also has low $\delta^{56}$Fe values based on field studies (Frost et al., 2007). It is thus unlikely that hematite formed from these precursor minerals by oxidation or pH change (Garrels and Christ, 1965). Both reaction kinetics and Fe isotope compositions, therefore, rule out UV photo-oxidation as the origin of hematite in the MBC.

Remaining possible mechanisms for Fe(II) oxidation include oxidation by free oxygen, or by anaerobic photosynthetic Fe(II)-oxidizing bacteria. Under an anoxic Archean atmosphere, as suggested by a low seawater U concentration (this study) and occurrence of mass-independent fractionation of S isotopes at this time (Farquhar et al., 2007), any free oxygen would have been confined to localized “oxygen oases”, such as might be associated with blooms of oxygenic photosynthesizing bacteria. Alternatively, anaerobic photoautotrophic microorganisms could have used Fe(II) rather than H$_2$O as an electron donor, producing Fe(III) rather than O$_2$ (Widdel et al., 1993). Experiments in systems that only contain Fe show that anaerobic photoautotrophic microorganisms produce hydrous ferric oxide as metabolic products that are enriched in heavy Fe isotopes by $\sim$1.5 ± 0.2‰ in $\delta^{56}$Fe values relative to Fe(II) solutions (Croal et al., 2004), and Fe(II)-oxidizing nitrate-reducing bacteria may produce even larger fractionations of $\sim$3‰ (Kappler et al., 2010). Co-precipitation of iron oxides with Si, as would be expected during formation of the MBC, produces further enrichment of heavy Fe isotopes due to binding changes (Wu et al., 2012), where the Fe–Si hydroxide-Fe(II)$_{aq}$ fractionation factor can be as high as 4‰ in $^{56}$Fe/$^{54}$Fe. The Fe isotope compositions of the MBC, therefore, match well with these experimentally determined Fe isotope fractionations, and it can be reasonably concluded that anaerobic photosynthesis, in a Si-bearing system, could produce the measured $\delta^{56}$Fe values for Fe(II) to Fe(III) oxides in the MBC.

The very high $\delta^{56}$Fe values require that the photic zone of Paleoarchean ocean was rich in Fe(II), the most important electron donor that could have supported anoxygenic photosynthesis (Canfield, 2005). We suggest that the combination of high-$\delta^{56}$Fe values and low U contents could be a signature for anaerobic photosynthetic Fe(II)-oxidizing bacteria. Because anoxygenic photosynthesis is likely to be a deeply rooted metabolism (Blankenship, 1992; Hoffmann-Marriott and Blankenship, 2011), and phylogenetic evidence suggests that oxygenic photosynthesis evolved after anoxygenic photosynthesis (Xiong et al., 2000), our results suggest that anoxygenic photosynthesis can be traced back to 3.46 Ga ago. Recent work by Czaja et al. (2013) on 3.8 Ga BIFs from the Isua Supracrustal Belt suggests that this metabolism may have been present even earlier, although the range in $\delta^{56}$Fe values measured at Isua is less than that measured in the MBC.

The MBC samples contain ppb-levels of U and Th that are orders-of-magnitude lower than that of Phanerozoic cherts. The low Th contents indicate low detrital components in the MBC samples. U–Th–Pb isotope systematics indicate that the MBC has undergone post-depositional alteration that added external Pb components to the MBC, which is consistent with results obtained from the Apex Basalt (Li et al., 2012). Modeling of U–Pb isotope decay, however, indicates that the low U contents of the MBC cannot be explained by recent U loss, despite the evidence...
for Pb mobility. Based on the U–Th–Pb isotope data, we constrain the upper limit of U concentration that was adsorbed by Fe(III) hydroxides during deposition of the MBC, from which we calculate an upper limit of U concentration in 3.46 Ga seawater at ~0.02 ppb, at least two orders of magnitude lower than the modern seawater value. Because U is mobile only when oxidized, the low U concentration in the MBC samples is an independent geochemical proxy that indicates an anoxic atmosphere in the Archean at 3.46 Ga. Our U–Th–Pb results also highlight the importance of determining the “age” of U in ancient sedimentary rocks when using U as a paleo-redox proxy, either through element abundances or mass-dependent isotope fractionations.

Collectively, the extreme enrichment in heavy Fe isotopes and low-U contents rule out the hypothesis that the Archean ocean was oxic (Hoashi et al., 2009). The MBC is laterally extensive (~120 km) and was precipitated in a deep marine environment, suggesting that the Fe(II)-rich, U-poor nature inferred for seawater at 3.46 Ga existed at a basinal scale. Our results provide additional support for a growing body of evidence for anoxic conditions at ~3.4 Ga, and additionally point to anoxygenic photosynthetic Fe(II) oxidation as a likely phototrophic pathway in the Paleoarchean. The results presented here do not support the presence of oxygenic photosynthesis as the mechanism of oxidation of Fe(II) in the MBC.

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APPENDIX A. SUPPLEMENTARY DATA

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