Environmental niches and metabolic diversity in Neoarchean lakes

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Abstract

The diversification of macro-organisms over the last 500 million years often coincided with the development of new environmental niches. Microbial diversification over the last 4 billion years likely followed similar patterns. However, linkages between environmental settings and microbial ecology have so far not been described from the ancient rock record. In this study we investigated carbon, nitrogen and molybdenum isotopes and iron speciation in five non-marine stratigraphic units of the Neoarchean Fortescue Group, Western Australia, that are similar in age (2.78-2.72 Ga) but differ in their hydro-geologic setting. Our data suggest that the felsic-dominated and hydrologically open lakes of the Bellary and Hardey formations were probably dominated by methanogenesis ($\delta^{13}C_{org} = -38.7 \pm 4.2\%$) and biological N$_2$ fixation ($\delta^{15}N_{bulk} = -0.6 \pm 1.0\%$), whereas the Mt. Roe, Tumbiana and Kylene Formations, with more mafic siliciclastic sediments, preserve evidence of methanotrophy ($\delta^{13}C_{org}$ as low as -57.4%, $\delta^{13}C_{carb}$ as low as -9.2%) and NH$_3$ loss under alkaline conditions. Evidence of oxygenic photosynthesis is recorded only in the closed evaporitic Tumbiana lakes marked by abundant stromatolites, limited evidence of Fe and S cycling, fractionated Mo isotopes ($\delta^{98/95}Mo = +0.4 \pm 0.4\%$) and the widest range in $\delta^{13}C_{org}$ (-57% to -15%), suggesting oxidative processes and multiple carbon fixation pathways. Methanotrophy in the three mafic settings was probably coupled to a combination of oxidants, including O$_2$ and SO$_4^{2-}$. Overall, our results may indicate that early microbial evolution on the Precambrian Earth was in part influenced by geological parameters. We speculate that expanding habitats, such as those linked to continental growth, may have been an important factor in the evolution of life.

1. Introduction

The paleontological record of plants and animals has demonstrated that diversification of macro-organisms was often facilitated by the provision of new environmental niches (Benton, 1995). A famous example includes the colonization of continental habitats, driven by evolutionary adaptations to higher solar radiation, desiccation and highly variable water chemistry (Little, 1990; Kenrick & Crane, 1997). Since then, continents have been important localities for the origination of new species, at least partly driven by the immense environmental heterogeneity of land surfaces (Stein et al., 2014). With the advent of next-generation sequencing, it has become increasingly clear that similar environmental selection effects also operate on Bacteria and Archaea and shape
microbial diversity on spatial scales ranging from millimeters to hundreds of kilometers (Horner-Devine et al., 2004; Smith et al., 2005; Martiny et al., 2006). Experimental work has confirmed that Bacteria and Archaea can adapt and speciate rapidly in response to environmental gradients (Rainey & Travisano, 1998), and similar to plants and animals, they appear to be dispersal-limited and may be constrained by geographic barriers (Whitaker et al., 2003). Hence, it is conceivable that the earliest diversification of microbial life in the Archean (e.g., David & Alm, 2011) was also driven to some extent by the diversity of environmental niches.

To test this hypothesis, we turned to the Neoarchean lower Fortescue Group (2.78-2.72 Ga) in Western Australia (Fig. 1), which comprises a sequence of igneous and sedimentary rock units that are typically interpreted as being deposited in non-marine environments and that capture several distinct sets of environmental conditions (Blake et al., 2004). The mineralogy of the detrital sediment is highly variable, with a mostly felsic source for the Bellary and Hardey formations and a mafic source for the Mt. Roe, Kylena and Tumbiana formations. While the Kylena and Tumbiana formations are carbonate-rich and likely represent large stagnant evaporative lakes, the Bellary and Hardey formations represent open fluvo-lacustrine systems (Blake, 1993), and the Mt. Roe formation has only small shallow ephemeral ponds (Rye & Holland, 2000). This set of environmental niches is not unique to the Neoarchean, but it creates a window into diverse microbial habitats a few hundred million years before the Paleoproterozoic rise of atmospheric oxygen (Lyons et al., 2014).

We hypothesize that this diversity of environmental settings resulted in biogeographic patterns in microbial ecology. We addressed this question with a range of geochemical tools including carbon and nitrogen isotopes, molybdenum isotopes, iron speciation and trace element abundances to test for variation in metabolic strategies, nutrient supply and redox settings. Many of these stratigraphic units have previously been investigated in separate geochemical studies (Buick, 1992; Rasmussen & Buick, 1999; Rye & Holland, 2000; Eigenbrode & Freeman, 2006; Bolhar & van Kranendonk, 2007; Thomazo et al., 2009; Thomazo et al., 2011; Coffey et al., 2013; Wille et al., 2013; Flannery et al., 2014; Stüeken et al., 2015a; Stüeken et al., 2015b; Slotznick & Fischer, 2016) but have so far not been compared within an environmental context. We provide previously unrecognized correlations among microbial metabolism, redox conditions, hydrological settings and geological substrate on Neoarchean land surfaces.

2. Geological Setting

The Fortescue Group rests unconformably on the Pilbara craton in the northwest of Western Australia (Blake & Barley, 1992; Blake et al., 2004). The lower part, which was the focus of this study, comprises three subaerial flood basalt sequences (Mt. Roe, Kylena, Maddina) that are separated by terrigenous sedimentary units (Fig. 2). Accommodation space for the sediments was probably created by WNW–ESE extension of the craton, which progressively opened up a series of half-grabens. The flood basalts are thought to be analogous to Phanerozoic flood basalts associated with continental rifting (Barley et al., 1992). The metamorphic grade of these rocks is relatively low—sub-greenschist facies specifically—and thus well suited to our study (Smith et al., 1982).

2.1. Bellary Formation

The basal unit of the Fortescue Group is the Bellary Formation, which remains undated and is composed of siliciclastic sediments, including conglomerates, diamictites and thin mudstones. It rests unconformably on granitoids and gneisses, which provide a maximum age of
2.85 Ga (Hall, 2005). Although basalt overlies the unconformity in many areas, the source terrain of the sediments is dominated by granite and other felsic lithologies of the Pilbara craton, as evidenced by the felsic composition of conglomeratic clasts (Fig. 3a). We did not observe basalt close to our sampling sites. The setting of Bellary Formation has been interpreted as fluvio-lacustrine with local alluvial fans (Barley et al., 1992; Blake, 1993) filling paleo-valleys cut into the unconformity surface with relief of up to several hundred meters and a lateral extent of several kilometers (Blake, 1993).

2.2. Mt. Roe Formation

The overlying Mt. Roe basalt contains discontinuous (less than a few kilometers along strike) siliciclastic and volcaniclastic sedimentary beds that are few meters thick and separate individual basalt flows. Silty mudrocks are typically graded and moderately sorted (Fig. 3c), suggesting pulsed, perhaps storm-controlled deposition into standing water. The sediments contain abundant opaque particles that likely represent devitrified volcanic glass eroded from basaltic flow tops (Fig. 3b). Paleosols on some of the basalt flows further support the non-marine depositional setting (e.g., Macfarlane et al., 1994; Rye & Holland, 2000). Leaching of ferrous iron from these paleosols indicates anoxic conditions at ~2.77 Ga (Yang et al., 2002). Deposition apparently occurred in shallow, short-lived (ephemeral) ponds rather than deep perennial lakes (Rye & Holland, 2000) with sedimentation occurring in distinct pulses from a very local source.

2.3. Hardey Formation

The Hardey Formation overlies the Mt. Roe Formation and is dominated by sandstone with subordinate conglomerates and mudrocks. It is interrupted by a felsic porphyry unit. Detrital grains in the Hardey Formation are predominantly composed of quartz and feldspar (Fig. 3d), reflecting a felsic provenance (Blake, 1993). The depositional environment is interpreted to be an open, braided fluvial system, based on the moderate degree of sorting and unidirectional current ripples and trough cross-bedding in sandstones, indicating one predominant direction of water flow—as in a river (Blake, 1993) (Fig. 3f). As for the Mt. Roe, mudrocks of the Hardey Fm are commonly graded (Fig. 3e) and probably represent overbank deposits or ephemeral lakes, indicated by their symmetrical and truncated wave ripples (Fig. 3f). The mudrock units are mostly less than 10 meters thick and extend laterally for no more than a few kilometers. The conglomeratic units contain deposits of heavy minerals, including pyrite, siderite and uraninite, which attest to an oxygen-depleted atmosphere at the time of deposition around 2.75-2.76 Ga (Rasmussen & Buick, 1999; Hall, 2005).

2.4. Kylena basalt, Mopoke Member

The Kylena basalt above the Hardey Formation contains the Mopoke Member, a thin (~10-meter-thick) stromatolitic carbonate unit that crops out discontinuously (Fig. 3g). The continuity along strike for many segments is less than 10 kilometers. The carbonate is largely calcite with minor dolomite (Flannery et al., 2014, this study). The carbonates of the Mopoke Member are interlaminated with shale that includes volcaniclastic material, likely derived from the Kylena basalt vents. Patterns for rare earth elements in the carbonates suggest a non-marine depositional setting (Bolhar & van Kranendonk, 2007), whereas occurrences of mud-drapes over symmetric wave ripples and relatively heavy \( \delta^{13}C_{org} \) values compared to the lacustrine Tumbiana Formation have been interpreted as evidence of either a marine influence or a distinct lake (Flannery et al., 2014). As further investigation shows that carbon and nitrogen isotope data are indeed similar to
those of the Tumbiana Formation (discussed below) (Stüeken et al., 2015b), we tend toward the lacustrine interpretation.

2.5. Tumbiana Formation

The Tumbiana Formation is a thick sequence (several hundred meters in many places) of tuffaceous wacke, mudrock and carbonate that extends for hundreds of kilometers across the Pilbara Craton. This continuity is broken up by NNE-trending growth faults, which fragment the sites of deposition into separate sub-basins many tens of kilometers across. The carbonate consists mostly of calcite (Buick, 1992, this study) and is commonly stromatolitic (Fig. 3h). Siliciclastic sediments with felsic mineralogies are rare but present, with trough cross-bedding suggesting a fluvial source in this case. Multiple lines of evidence indicate a lacustrine setting, including the abundance of desiccation cracks and teepees (Fig. 3i), rapid lateral and vertical facies changes (Buick, 1992; Awramik & Buchheim, 2009), the absence of tidal indicators (cf. Sakurai et al., 2005; Awramik & Buchheim, 2009), the scarcity of dolomite and the absence of gypsum despite the presence of halite (Buick, 1992), elevated and variable $^{87}\text{Sr}/^{86}\text{Sr}$ ratios (Awramik & Buchheim, 2009) and the absence of Y and Gd anomalies and the presence of variable slopes in REE patterns (Bolhar & van Kranendonk, 2007; Coffey et al., 2013). The microbial mats and stromatolites were likely produced by cyanobacteria, as indicated by the morphology of tufted mats (Flannery & Walter, 2012) and the scarcity of sulfides and other authigenic iron minerals suggesting that H$_2$S and ferrous iron were not available as electron donors for anoxygenic phototrophs (Buick, 1992). The lack of Ce anomalies suggests that oxygen did not build up in the water column (Coffey et al., 2013). Individual stromatolites may have a synoptic relief of up to 2 meters, indicating that water was at least that deep for prolonged periods.

3. Methods

In addition to compiling data from the literature, we made new measurements of samples from the Hardey, Mt. Roe, Kylena and Tumbiana formations. Geochemical measurements followed standard protocols and therefore will only be described briefly.

3.1. Sample collection and preparation

Samples of the Bellary and Kylena formations and some samples of the Tumbiana and Hardey formations were collected from outcrop in the field (Stüeken et al., 2015b). We obtained additional samples of the Mt. Roe Formation, the Hardey Formation and the Tumbiana Formation from drill cores obtained as part of the NASA Astrobiology Drilling Project (cores ABDP-6, ABDP-3, ABDP-10). We also included Tumbiana samples from core SV-1 and WRL-1 (Stüeken et al., 2015b). Weathering rinds and the outer surfaces of drill-core samples were removed with a rock saw. The rocks were then crushed into cm-sized chips, cleaned with alcohol, 2N HCl and DI-H$_2$O (18MΩ) and pulverized with a Al$_2$O$_3$-ceramic puck mill. Powders were stored in scintillation vials that had been baked out in the muffle furnace at 500°C overnight.

3.2. Geochemical analyses

3.2.1 Organic carbon and nitrogen isotopes

Carbon and nitrogen isotope measurements were performed in the University of Washington Isolab following protocols described by Stüeken et al. (2013; 2015a; 2015b). For organic carbon and bulk nitrogen, powders were decarbonated with 6N HCl (reagent grade, applied three times at 60°C overnight), rinsed three times with DI-H$_2$O (18MΩ) and dried in a closed oven.
for two days. Dry powders were analyzed by flash-combustion with an elemental analyzer (Costech) coupled to a continuous-flow IRMS (Thermo MAT253). Data were corrected by two-point normalization with calibrated in-house standards (Paul et al., 2007). Analytical precision was 0.18‰ for δ13Corg and 0.50‰ for δ15N, as determined by the standard deviation of replicate samples. Relative precisions of C and N concentrations were 1% and 4%, respectively. Long-term isotopic precision over several weeks was monitored with our in-house rock standard UW-McRae, for which we obtained a value of 5.22 ± 0.20 ‰ for δ15N and -37.83 ± 0.10‰ for δ13Corg—in good agreement with previous studies (5.51±0.19‰ and -37.89 ± 0.21‰, Stüeken et al., 2015a). Carbon and nitrogen isotope data are reported in standard delta notation, referenced to VPDB for C and to atmospheric N2 for N.

3.2.2. Carbonate carbon and oxygen isotopes

For carbonate-bound carbon and oxygen isotopes, untreated powders were weighed into glass vials and reacted with concentrated phosphoric acid at 80°C in a Kiel III Carbonate-Device. The resulting CO2 gas was purified cryogenically and analyzed with a dual-inlet IRMS (Thermo Finnigan Delta Plus). As for carbon and nitrogen isotopes, we used a two-point calibration to correct the raw data. The average precision was 0.04‰ for both δ13Ccarb and δ18Ocarb. We assumed calcite mineralogy for the oxygen isotope measurements, as confirmed by high molar Ca/Mg ratios measured in Na-acetate extracts (range 4 to 283, median 40, excluding samples with less than 2% carbonate; see below). The CO2 gas pressure was calibrated for carbonate quantity (relative precision 6.4‰). Carbon and oxygen isotope data are reported in standard delta notation, referenced to VPDB for C and O.

3.2.3. Elemental abundances

Elemental abundances and iron speciation were measured at the University of California, Riverside (Reinhard et al., 2013). Aliquots of untreated rock powder were first ashed in ceramic crucibles (800°C overnight) and then digested in Teflon beakers with 4ml HNO3 and 1ml HF at 120°C overnight. Fluoride precipitates were removed with 1-2 treatments with aqua regia (3ml HCl + 1ml HNO3). All acids were concentrated and trace-metal grade. Sample residues were stored in 5% (v/v) HNO3. Elemental analyses were performed by ICP-MS (Agilent 7500ce). Average precision for all elements was 6%. Measured concentrations of the USGS rock standard SCo-1 agreed to with 7% with nominal values. Enrichment factors (EF) for trace elements (X) relative to continental crust were calculated as EFx = ([Xsample]/[Alsample])/(Xcrust)/[Alcrust]), where concentrations in average upper continental crust were taken from Rudnick & Gao (2014).

3.2.4. Iron speciation

For the extraction of different iron phases, around 100mg of powder were treated sequentially with sodium acetate (1M, 10ml, buffered to pH 4.5 with acetic acid, 48h), sodium dithionite (50g/L, 10ml, buffered to pH 4.8 with 0.35M acetic acid and 0.2M sodium citrate, 2h) and ammonium oxalate (0.2M, 10ml, buffered to pH 3.2 with 0.17M NH4OH, 6h) to extract iron from carbonates, ferric oxides and magnetite, respectively (Poulton & Canfield, 2005; Reinhard et al., 2009; Reinhard et al., 2013). For each treatment, the samples were placed on a shaker table with the reagent at room temperature and then centrifuged. An aliquot of the supernatant was saved for analyses of iron contents by ICP-MS (Agilent 7500ce). Pyrite-bound iron was determined using the Cr-reduction method with a separate powder aliquot (Canfield et al., 1986). Around 0.5-2g of powder were reacted with hot CrCl3 (1M in 0.5M HCl, 35ml) and concentrated HCl (15ml) in one
step under N$_2$ gas for 2h, and the resulting H$_2$S gas was trapped with zinc acetate. The abundance of sulfur was determined by iodometric titration (Canfield et al., 1986). We assumed an FeS$_2$ stoichiometry to calculate the concentration of pyrite Fe from measured values of pyrite S. The precision was 7% on average. The sum of all extracts in each sample, including pyrite-bound iron (Fe$_{PY}$) is defined as the total amount of ‘highly reactive iron’ (Fe$_{HR}$).

3.2.5. Molybdenum isotopes

Molybdenum isotopes were measured at Yale University following established protocols (Asael et al., 2013; Planavsky et al., 2014). An aliquot of the bulk digest was doped with a $^{97}$Mo-$^{100}$Mo double spike, targeting a spike/sample molybdenum ratio of 1. The molybdenum was purified from matrix elements with two ion-exchange columns (AG-MP-1M and AG50W-X8) and analyzed by multi-collector ICP-MS (Thermo Neptune) at Mo concentrations of 50ppb. Isotopic ratios are expressed in $\delta^{98/95}$Mo (‰) = 1000 · [(Mo/95Mo)$_{\text{sample}}$/(Mo/95Mo)$_{\text{NIST}3134}$ - 1], where NIST 3134 was set to +0.25 ‰ as proposed by Nägler et al. (2014). We obtained a value of -0.41 ± 0.03‰ (1σ) for the USGS reference standard NOD-1, in good agreement with previous work (-0.42 ± 0.02‰, Asael et al., 2013). The average standard deviation of sample replicates was 0.03‰ (1σ).

4. Results

Analytical results are tabulated in the Appendix.

4.1. Carbon isotopes

New organic and inorganic $\delta^{13}$C values for the Tumbiana Formation range from -57.4‰ to -36.3‰ and -6.8‰ to +1.0‰, respectively, similar to previous results (-57.1‰ to -15.0‰ and -9.2‰ to +1.9‰, Eigenbrode & Freeman, 2006; Thomazo et al., 2009; Stüeken et al., 2015b). Low $\delta^{13}$C$_{\text{carb}}$ values tend to occur primarily in mudrocks with less than 40% carbonate and associated with light $\delta^{13}$C$_{\text{org}}$ values (Fig 3). Inorganic $\delta^{13}$C in the Kylena Formation (-1.9‰ to +2.9‰) is broadly similar to that of the Tumbiana Formation and in good agreement with measurements by Flannery et al. (2014) (-5.7‰ to +2.1‰). In the Hardey and Bellary formations, where carbonate is rare, $\delta^{13}$C$_{\text{org}}$ values (-42.3‰ to -24.9‰, Watanabe et al., 2004; Stüeken et al., 2015a) fall toward the heavier end of the range seen in the Tumbiana Formation. The Mt. Roe Formation has light $\delta^{13}$C$_{\text{carb}}$ values (-55.3‰ to -50.1‰) similar to those of the Tumbiana and Kylena formations (Flannery et al., 2014; Stüeken et al., 2015b, this study), albeit with much less scatter.

4.2. Nitrogen isotopes

New nitrogen isotope data from the ABDP-3 drill core from the Hardey Formation ($\delta^{15}$N = -2.3‰ to +1.8‰) agree well with those previously obtained from outcrop samples (-2.2‰ to +0.7‰, Stüeken et al., 2015a). Nitrogen isotope data from the Mt. Roe ( $\delta^{15}$N = +7.3‰ to +15.8‰) fall between those of the Hardey and Tumbiana formations (Thomazo et al., 2011; Stüeken et al., 2015b). The Tumbiana Formation has been noted for the highest nitrogen isotope ratios found to date in the rock record, with $\delta^{15}$N values that extend up to 50‰ (Thomazo et al., 2011; Stieken et al., 2015b), paired with unusually high ratios of organic carbon to total nitrogen (119-1719, Thomazo et al., 2011; Stüeken et al., 2015b). In the Mt Roe Formation, TOC/TN ratios (85-914) are also elevated compared to the Hardey and Bellary formations (5-271, Stüeken et al., 2015b, this study).

4.3. Molybdenum isotopes and metal enrichments
Molybdenum isotope data ($\delta^{98/95}$Mo) from sedimentary rocks of the Tumbiana Formation range from -0.1‰ to +1.8‰, with a mean of 0.4 ± 0.4‰. This range is larger than that reported from the Mt. Roe and Hardey formations (+0.2‰ to +0.7‰, Wille et al., 2013). Molybdenum is not significantly enriched in any of the formations relative to modern average upper continental crust ($EF_{Mo} = 2.5 \pm 1.5$, with a few outliers with < 1% Al). Enrichments in other transition metals are similarly small or absent.

4.4. Iron speciation

Ratios of total iron to aluminum are elevated relative to average crust (0.5, Rudnick & Gao, 2014) in the Tumbiana (0.7-3.7) and Kylena formations (0.9-2.9) and in some samples of the Mt. Roe Formation (0.03-0.9)—but not in the Hardey Formation (0.2-0.6). Ratios of highly reactive iron ($Fe_{HR}$) to total iron ($Fe_T$) fall mostly between 0.1 and 0.4 in the Tumbiana, Kylena and Mt. Roe formations; three samples in the Mt. Roe Formation are higher (up to 0.6). In the Hardey Formation, $Fe_{HR}/Fe_T$ falls mostly below 0.1. Ratios of pyrite-bound iron ($Fe_{Py}$) to $Fe_{HR}$ range from 0 to 0.5; four samples of the Mt. Roe Formation plot between 0.6 and 0.9.

5. Interpretation

The various proxies used in this study reveal distinct environmental and ecological properties in each depositional setting. Summaries of those patterns follow.

5.1. Iron speciation

The iron speciation proxy for paleoredox has been calibrated for marine environments with relatively low sedimentation rates (Raiswell & Canfield, 1998; Anderson & Raiswell, 2004). In marine and non-marine settings, however, high background rates of siliciclastic sedimentation in combination with relatively low abundances of reactive iron ($Fe_{HR}$) may decrease the ratio of $Fe_{HR}$ to total iron ($Fe_T$) and thus dilute potential enrichment patterns (e.g., Lyons & Severmann, 2006). This possibility is particularly likely when the sediments are both mafic and tuffaceous, as in the Tumbiana, Kylena and Mt. Roe formations, because (a) mafic minerals are iron-rich, and (b) airfall tuff can be associated with rapid sedimentation rates. Furthermore, (c) the solubility of Fe$^{2+}$ decreases markedly with increasing pH (Morgan & Lahav, 2007). The mobility of reactive iron in lakes and rivers could thus be higher or lower compared to seawater, depending on pH, which would also affect the $Fe_{HR}/Fe_T$ ratio irrespective of the redox state. For these three reasons, standard definitions of oxic, ferruginous and euxinic fields in terms of $Fe_{Py}/Fe_{HR}$ versus $Fe_{HR}/Fe_T$ crossplots (Fig. 5, e.g., Poulton & Canfield, 2011) are therefore not necessarily applicable to non-marine settings. Nevertheless, our iron data carry some environmental information about pH and sulfide availability.

5.1.1. Tumbiana and Kylena formations

The Tumbiana and Kylena formations have notably high $Fe_T/Al$ ratios between 0.7 and 3.7 (mean 1.7 ± 0.7), consistent with abundant mafic tuff and basaltic sediment sources. Although iron-rich mafic minerals and tuff are generally susceptible to weathering, the $Fe_{HR}/Fe_T$ ratios of these samples are notably low (< 0.3). It is unlikely that the lack of enrichment in $Fe_{HR}/Fe_T$ resulted from the lack of an appropriate carrier phase, because dissolved Fe$^{2+}$, if present, could have been captured by the abundant carbonate. Furthermore, the Tumbiana lakes were at least partly closed basins with evidence of halite precipitation (Buick, 1992), making it unlikely that reactive Fe$^{2+}$ was lost by fluvial export. The relative lack of $Fe_{HR}$ is thus consistent with suppressed iron
solubility irrespective or redox under alkaline conditions in these lakes, as previously proposed (Stüeken et al., 2015b) (discussed below). Little ferrous iron would have been released into solution under such conditions. Instead it would have been retained in clays and zeolite minerals that formed during alteration of volcanic ash. This would explain the scarcity of authigenic iron minerals such as carbonates, oxides and sulfides.

5.1.2. Hardey Formation

High pH has not previously been invoked for the Hardey Formation, and we find no evidence for it in our data. Ratios of total iron to aluminum cluster tightly around a mean of 0.5 ± 0.1, equivalent to average upper continental crust (0.5, Rudnick & Gao, 2014). Ratios of FeHR to FeT of < 0.2 overlap with ‘oxic’ marine sediments that experienced minimal addition of authigenic iron (Poulton & Raiswell, 2002). However, oxic conditions are incompatible with the record of detrital pyrite, uraninite and siderite in these sediments (Rasmussen & Buick, 1999), unless the sedimentation rate was very high. It is perhaps more likely that detrital iron-silicate minerals diluted any authigenic iron minerals, and/or the formation of authigenic iron minerals was simply not favored in the absence of significant amounts of oxygen, carbonate or sulfide. It is therefore not possible to infer a redox setting for the Hardey Formation from the iron speciation data.

5.1.3. Mt. Roe Formation

In the Mt. Roe Formation, FeT/Al ratios vary between 0.03 and 0.91, although given the mafic provenance we might predict values as high as those seen in the Tumbiana Formation. Paleosols in the Mt. Roe basalt have been noted for iron loss as a result of anoxic, acidic weathering (Macfarlane et al., 1994), and it is likely that clay particles in the lake deposits that we sampled were sourced from those exposed and leached horizons. Unlike the other units, the Mt. Roe Formation shows some FeP/FeHR ratios above 0.5, suggesting that sulfur was more readily available. Sulfate may have been supplied by localized oxidative weathering (Stüeken et al., 2012; Lalonde & Konhauser, 2015) or by volcanic outgassing of SO2 and subsequent biological reduction. In the other formations, where FeP/FeHR ratios are generally lower (Fig. 5), sulfur input seems to have been relatively less.

In conclusion, the iron speciation data cannot resolve whether any of these fluvio-lacustrine water bodies in the Fortescue Group were oxic or anoxic because the iron-based proxies are presently limited in their relevance to non-marine environments. However, the low FeHR/FeT ratios in the Tumbiana and Kylena formations are consistent with the interpretation of high pH in those settings (Stüeken et al., 2015b), and they indicate that none of these water bodies developed euxinia, as FeP/FeHR are mostly below 0.7 (Poulton & Canfield, 2011). Slightly elevated FeP/FeHR ratios may indicate the presence of sulfide in some sedimentary pore waters.

5.2. Trace elements and molybdenum isotopes

The concentrations of trace elements appear to be primarily controlled by bedrock composition with no detectable overprint by redox processes or hydrothermal activity. Mild enrichments in V, Ni and Cr (EF > 2) compared to average modern continental crust (Rudnick & Gao, 2014), paired with mild depletions in U (mostly EF < 1), are consistent with a mafic sediment provenance and a generally more mafic Archean crust (Taylor & McLennan, 1995). The absence of significant enrichments in Cu (EF ~ 1-5), Zn (EF ~ 1-2) and As (EF ~ 1-3) argues against input from hydrothermal fluids, either during or after deposition. Furthermore, the absence of marked
depletions in first-row transition metals argues against metasomatic alteration, as observed in other geographic regions in the Fortescue Group (White et al., 2014a; White et al., 2014b).

High sedimentation rates, either through fluvial particle transport or rainout of volcanic ash, could easily have masked small elemental enrichments in the sediments. The molybdenum isotope record, however, appears to be more sensitive to Mo mobility than elemental abundances. The wide range of δ⁹⁸/⁹⁵Mo values in the Tumbiana Formation (-0.1‰ to +1.8‰, Fig. 6) exceeds those of basalts (-0.2‰ to +0.3‰) and granites (+0.1‰ to +0.6‰) (Voegelin et al., 2014, this study), which implies some degree of isotopic fractionation in the aqueous phase. The data thus strongly suggest that Mo was present in aqueous solution in the Tumbiana lakes. One possible source of dissolved Mo is alteration of volcanic glass, which can liberate Mo even in the absence of oxygen (Greaney et al., 2016). However, if this process were important, one would expect high levels of dissolved Mo and thus the potential for a wide range of fractionations in the Mt. Roe Formation where volcanic ash is abundant. This is not observed (Fig. 6). Thus, it is perhaps more likely that the Mo in the Tumbiana Formation was liberated by low levels of localized oxidative weathering (Stüeken et al., 2012; Lalonde & Konhauser, 2015). Isotopic fractionations may have been induced as the dissolved Mo was partially sequestered into sulfide minerals (Poulson et al., 2006; Siebert et al., 2006; Neubert et al., 2008; Kendall et al., 2017) or assimilated into biomass (Zerkle et al., 2011). Regardless of the mechanism that fractionated the Mo isotopes, the isotopic range is evidence that Mo was present in solution, possibly as a result of mildly oxic conditions and consistent with claims of oxygentic photosynthesis based on the tufted mats and stromatolites in the Tumbiana Formation (Buick, 1992; Flannery & Walter, 2012).

The comparatively small ranges of δ⁹⁸/⁹⁵Mo in the Hardey Formation (+0.4‰ to +0.5‰) and Mt. Roe Formation (+0.2‰ to +0.6‰) (Wille et al., 2013) may suggest that Mo was less mobile in these settings compared to the Tumbiana lakes; however, we cannot rule out that small amounts of authigenic, isotopically fractionated Mo precipitates were swamped by detrital Mo.

5.3. Nitrogen fixation in the felsic sedimentary systems

Total nitrogen and organic carbon are well correlated in our new samples from the Hardey Formation (r² = 0.8), suggesting that the nitrogen was derived from biomass. A similar though weaker trend (r² = 0.4) was observed in previous analyses of outcrop samples from the Hardey and Bellary formations (Stüeken et al., 2015a). It is possible that these samples contain some fraction of detrital silicate-bound nitrogen. However, the tight N isotope range in both the Bellary and Hardey formations (Fig. 7) suggest that any potential isotopic effects from the admixture of a detrital N phase are minor. We can also rule out a significant metamorphic overprint, because such effects only become significant (> 1‰) for greenschist facies and above (reviewed by Thomazo & Papineau, 2013). Hence, the δ¹⁵N values in these samples, falling around the atmospheric composition of 0‰, are most parsimoniously explained by biological N₂ fixation using Mo-based nitrogenase (Stüeken et al., 2015a). The isotopic composition of biomass from organisms using this enzyme lies between -2‰ and +1‰ (reviewed by Casciotti, 2009), and 26 out of 30 data points from these two formations plot within this range. Alternative nitrogenases using Fe or V instead of Mo can be reasonably ruled out because they would produce larger fractionations between -6‰ and -8‰ relative to the atmosphere (Zhang et al., 2014). Similarly, abiotic fixation products such as HCN from photochemistry or NO₃ from lightning would likely have been more fractionated (< -5‰, Ingerson, 1953; Moore, 1977; Kuga et al., 2014), and additional isotopic fractionation would have been imparted on such abiotic NO₃ species during reduction to N₂ or NH₄⁺ (McCreary et al., 1983; Sigman et al., 2009). In short, the tight cluster of δ¹⁵N values around a mean of -0.6 ± 1.0‰
is inconsistent with secondary processing of a fractionated NO$_3^-$ reservoir and instead is best explained by a single dominant metabolism, in this case biological N$_2$ fixation.

As noted above, Mo isotopes and concentrations in the Hardey Formation argue against a large Mo reservoir (Wille et al., 2013). At concentrations below 1-5 nM (100-20 times less than in the modern ocean), Mo can become limiting for modern nitrogen-fixing organisms (Fay & de Vasconcelos, 1974; Attridge & Rowell, 1997; Zerkle et al., 2006; Glass et al., 2010). In some modern lakes, Mo levels can be < 5 nM and thus severely limit but not preclude biological productivity (Glass et al., 2012). It is conceivable, therefore, that ancient N$_2$ fixers were adapted to low Mo concentrations through enzymatic pathways that enhanced the efficiency of Mo utilization (Glass et al., 2013).

5.4. NH$_3$ volatilization from alkaline lakes

As noted previously (Thomazo et al., 2011; Stüeken et al., 2015b), the Tumbiana Formation stands out for its unusually high nitrogen isotope ratios of up to +50‰ (mean +31 ± 9‰, 1 s.d., Fig. 7). The same is true for the Kylena Formation (up to +30‰, mean +24 ± 7‰). These data are most reasonably interpreted as evidence for high pH (>9), which favors $^{14}$N loss via NH$_3$ volatilization (Stüeken et al., 2015b). Aqueous NH$_4^+$ released from dead biomass during degradation and diagenesis transforms into volatile NH$_3$ with a pKa of 9.25, and this process is associated with a fractionation of 45‰ at room temperature and 1 bar atmospheric pressure (Li et al., 2012). With a starting composition around 0‰ from N$_2$ fixation and some degree of Rayleigh distillation, NH$_3$ volatilization alone could have generated $\delta^{15}$N values up to +50‰. This interpretation is supported by the high C/N ratios of these rocks (458 ± 326), despite their low metamorphic grade (Stüeken et al., 2015b). Such extreme C/N ratios cannot be reconciled with a biological nitrogen removal process and instead require a post-mortem mechanism that preferentially removes N over C. Hence the high $\delta^{15}$N values and C/N ratios were probably both consequences of alkaline NH$_3$ devolatilization. It is possible that additional redox processes were operative in the Tumbiana lakes (Thomazo et al., 2011), as is the case in surface waters of modern alkaline lakes where NH$_3$ builds up at depth and is released during seasonal water-column overturn (Jellison et al., 1993). If NO$_3^-$ was a nitrogen source for some organisms, it could have elevated the average starting composition of biomass to > 0‰, as in the modern ocean. In such a case, values of up to +50‰ could then be explained with less extreme degrees of NH$_3$ volatilization.

The Mt. Roe Formation, which is also mafic in its mineralogy but was probably hydrologically ephemeral (Rye & Holland, 2000), is not as enriched in $\delta^{15}$N (mean +12 ± 3‰) compared to the Kylena and Tumbiana formations, but it is similarly depleted in nitrogen (C/N = 561 ± 269). The total spread in $\delta^{15}$N over 9‰ (from +7‰ to +16‰) is much wider than the range observed for the Bellary and Hardey formations and consistent with some form of secondary processing of nitrogen in the environment. In the modern ocean, for example, $\delta^{15}$N values of up to +15‰ can be generated by partial denitrification of NO$_3^-$ to N$_2$ and uptake of the residual heavy nitrate (Sigman et al., 2009; Tesdal et al., 2013), but in the absence of significant Mo isotope fractionations (Wille et al., 2013) or stromatolitic indications of O$_2$ production, we see no evidence suggesting appreciable NO$_3^-$ production. It is thus likely that N loss and associated isotopic fractionation were caused by NH$_3$ volatilization from alkaline waters, similar to the Tumbiana Formation. The reason for the disparity between the Mt. Roe and Tumbiana formations could be a higher degree of redox cycling in oxic surface waters of the Tumbiana lakes, which, as noted above, may have elevated the $\delta^{15}$N starting composition of biomass. Furthermore, differences in water depth could affect the overall degree of N loss from a system. In the relatively ephemeral
system of the Mt Roe, where high pH conditions perhaps only developed transiently in sedimentary pore waters in contact with tuffaceous material, NH₃ loss may be relatively less severe compared to long-lived evaporitic settings.

Claims of alkaline conditions in the Tumbiana, Kylena and Mt. Roe formations are consistent with the mafic substrate. Volcanic glass, in particular basaltic glass, is known to act as a strong base that consumes H⁺ and releases cations into solution during aqueous alteration (Gislason & Eugster, 1987; Gysi & Stefansson, 2008). Modern alkaline lakes are therefore commonly associated with areas of active volcanism, such as the East African Rift (Cerling, 1994; Kempe & Kazmierczak, 2011).

A previous interpretation of high pH in the Tumbiana lakes (Stüeken et al., 2015b) was recently contested by Hahn et al. (2015) based on rare earth element (REE) patterns. Modern alkaline lake waters can be highly depleted in light (LREE) over heavy REE, with (Pr/Yb)₉₉₈₈ < 0.01 (Moeller & Bau, 1993), and it was argued that similar depletions should be observed in carbonate precipitates (Hahn et al., 2015). However, there are no known examples of chemical sediments from bona fide alkaline lakes that capture the extreme LREE depletion seen in the water column. For example, cherts from modern alkaline Lake Magadi ((Pr/Yb)₉₉₈₈ = 0.1-0.7) (Kerrich et al., 2002) and carbonates from alkaline Lake Abhé ((Pr/Yb)₉₉₈₈ = 0.2-1.5) (Dekov et al., 2014) have REE patterns that are significantly flatter than those from alkaline waters, and the REE pattern of one carbonate sample from the Green River Formation ((Pr/Yb)₉₉₈₈ = 0.7), a well-known Eocene alkaline lake, falls within the range of Tumbiana carbonates ((Pr/Yb)₉₉₈₈ = 0.4-1.5) (Bolhar & van Kranendonk, 2007; Coffey et al., 2013). Hence REE patterns may not be a straightforward proxy for alkaline conditions in the rock record.

Another argument against high pH has been raised based on a relatively small spread of δ⁴⁴/⁴⁰Ca values in Tumbiana carbonates compared to waters of modern alkaline Mono Lake (Blättler et al., 2017), where high [HCO₃⁻]/[Ca²⁺] ratios induce a Rayleigh effect during carbonate precipitation, leading to a larger spread in δ⁴⁴/⁴⁰Ca (Nielsen & DePaolo, 2013). However, the calcium cycle in Mono Lake is not comparable to the Tumbiana Formation for several reasons. First, Ca²⁺ in Mono Lake is sourced primarily from small rivers and local springs with variable isotopic composition (Nielsen & DePaolo, 2013), whereas in the Tumbiana Formation, Ca²⁺ would most likely have been leached from airfall tuff during intra-basinal water-rock interactions. Fluvial deposits are rare in the Tumbiana Formation, and spring deposits are absent, whereas tuff is ubiquitous and abundant (Buell, 1992). Hence Ca²⁺ recharge was likely more uniform in spatial extent and isotopic composition compared to Mono Lake. Possible differences in reservoir size could also play an important role. Second, waters of Mono Lake are enriched in phosphate compared to modern seawater by more than a factor of 20, which inhibits the precipitation of CaCO₃ and imparts strong kinetic effects on Ca isotope fractionations (Nielsen & DePaolo, 2013). It is unlikely that dissolved phosphate concentrations were similarly high in the Tumbiana lakes, because our samples show P/Al ratios (P/Al *1000 = 14.1 ± 5.8, n = 61, unpublished data) that are only about twice as high as in Neoarchean marine shales (P/Al *1000 = 6.7 ± 5.8, n = 41, Yamaguchi, 2002) and average continental crust (P/Al *1000 = 8, Rudnick & Gao, 2014). Hence, Mono Lake is not a suitable analog for the Tumbiana Formation in terms of Ca chemistry. Most importantly, carbonate precipitates in Mono Lake actually show a very narrow spread in δ⁴⁴/⁴⁰Ca—much narrower than the isotopic spread in dissolved Ca (Nielsen & DePaolo, 2013) and roughly identical to that reported from Tumbiana carbonates (Blättler et al., 2017). Hence, carbonates may simply not capture the Rayleigh effect observed in the aqueous phase, which makes the δ⁴⁴/⁴⁰Ca proxy problematic for tracking alkalinity in deep time.
5.5. Carbon cycling

The Neoarchean (2.8-2.5 Ga) was a time of exceptionally light organic carbon isotope ratios (Krissansen-Totton et al., 2015). Marine sediments commonly fall between -30‰ to -45‰ compared to a range of -20‰ to -30‰ for the remainder of Earth history. This pattern of extreme $^{13}$C depletion is usually attributed to widespread production of isotopically light methane that was partially reincorporated into biomass via methanotrophy (Hayes, 1994; Hinrichs, 2002; Eigenbrode & Freeman, 2006; Fischer et al., 2009; Czaja et al., 2010; Zerkle et al., 2012; Izon et al., 2015). Slotznick & Fischer (2016) recently proposed that light organic $\delta^{13}$C values in the Tumbiana Formation were instead the result of acetogenesis and reassimilation of light acetate, which would not necessitate any oxidant, unlike methanotrophy. We will revisit this hypothesis below.

5.5.1. Preservation of carbon isotope signatures

As noted above, nitrogen isotope ratios in the three mafic settings in the Fortescue Group have likely been affected by NH$_3$ loss under alkaline conditions. There are no known effects of high pH on carbon isotopes. In general, diagenesis can lower organic $\delta^{13}$C values by 1-2‰ through selective preservation of heavier compounds (Lehman et al., 2002) and even more so through the addition of isotopically light biomass from methanogenic and methanotrophic organisms. The latter can be significant in samples where $\delta^{13}$C$_{org}$ is less than -35‰ and will be further discussed below.

Metamorphic alteration can lead to a re-equilibration between organic matter and carbonate such that $\delta^{13}$C$_{org}$ increases while $\delta^{13}$C$_{carb}$ decreases. However, this effect is minor (<3‰) below greenschist facies metamorphism (McKirdy & Powell, 1974; Hoefs & Frey, 1976; Hayes et al., 1983; Schidlowsk, 1987). Furthermore, as noted above (Fig. 4), we tend to observe light $\delta^{13}$C$_{carb}$ values in Tumbiana samples with light $\delta^{13}$C$_{org}$, which is opposite to the expected metamorphic trend and more likely an artifact of diagenesis. In these samples, amount of carbonate-bound carbon is in nearly all cases at least equal or greater than the amount of organic-bound carbon. If the observed trend were the result of metamorphic overprinting rather than diagenesis, then organic $\delta^{13}$C$_{org}$ should be more positive in samples with light $\delta^{13}$C$_{carb}$, which is not observed. A significant degree of metamorphic alteration of primary signals is therefore unlikely.

5.5.2. The Calvin cycle and oxygenic phototrophy in the Tumbiana Formation

The organic carbon isotope data from the Hardey and Bellary formations (mean -38.7 ± 4.2‰) (Fig. 7) fall within the Neoarchean marine range, whereas the Tumbiana, Kylena and Mt. Roe formations (mean -45.9 ± 7.8‰) are systematically lighter. Values heavier than -35‰ may be explained by primary production through the Calvin-Benson-Bassham (or reductive pentose phosphate) cycle (-15‰ to -35‰, Schidlowski, 1987; Zerkle et al., 2005). This pathway is used by a range of organisms, including cyanobacteria, some anoxygenic phototrophs and some chemotrophs, and it may have been the dominant mode of CO$_2$ fixation over most of Earth history (Nisbet et al., 2007). In the case of the Tumbiana and Kylena formations, the presence of stromatolites, the distinctive morphology of the tufted mats and the inferred scarcity of ferrous iron and sulfide as electron donors for anoxygenic phototrophs have been used to suggest that cyanobacterial productivity was at least in part responsible for primary production (Buick, 1992; Flannery & Walter, 2012). The most parsimonious explanation for the heavier organic $\delta^{13}$C values as high as -15‰ may thus be a cyanobacterial contribution. In the Hardey, Bellary and Mt. Roe
forms, by contrast, the Calvin cycle was probably not a dominant mode of productivity, because δ\(^{13}\)C\(_{\text{org}}\) values heavier than -35‰ are rare at these sites.

5.5.3. Autotrophic methanogenesis in the Hardey and Bellary formations

Organic carbon isotope values down to around -42‰ could reflect CO\(_2\) fixation by the acetyl-CoA pathway, which is used by many anaerobic organisms, including autotrophic methanogens, acetogens and some sulfate reducers (Schidlowski, 1987; Zerkle et al., 2005). Values significantly lighter than -42‰ are incompatible with any known CO\(_2\) fixation pathway that uses atmospheric CO\(_2\) with a composition of -7‰ (Schidlowski, 1987; Zerkle et al., 2005) and could instead reflect incorporation of isotopically light CH\(_4\) into biomass (Hayes, 1994; Hinrichs, 2002; Eigenbrode & Freeman, 2006). Two alternatives to autotrophic methanogenesis, acetogenesis and methanotrophy for δ\(^{13}\)C\(_{\text{org}}\) values below -35‰ are possible, but not likely for the Hardey and Bellary formations. First, such light values could have resulted from CO\(_2\) fixation by the Calvin cycle, if the CO\(_2\) source was lighter than atmospheric CO\(_2\). However, open fluvio-lacustrine settings such as the Hardey and Bellary were likely equilibrated with the CO\(_2\)-rich Archean atmosphere through dynamic mixing, as is true today (Mook, 2001). Atmospheric CO\(_2\) is set to a value of -7‰ today because it is buffered by the much larger marine DIC pool through an equilibrium fractionation of roughly 7-8‰ between gaseous CO\(_2\) and aqueous HCO\(_3^-\) (Hayes et al., 1999). The carbon isotope composition of the Neoarchean atmosphere would probably have been about the same as today, as suggested by the nearly stable δ\(^{13}\)C\(_{\text{carb}}\) record through time in phases that capture marine DIC (Krispanssen-Totton et al., 2015).

The second alternative possibility to autotrophic methanogenesis, acetogenesis and methanotrophy is heterotrophic methanogenesis using isotopically light organic substrates for conversion to methane and biomass (Londry et al., 2008). The biomass of such heterotrophic methanogens can be up to 30‰ depleted compared to their carbon source (Londry et al., 2008) and could thus explain organic carbon isotope values down to -60‰ if the starting substrate were produced by the Calvin cycle or another CO\(_2\) fixation pathway. However, we would expect that a signature of those organisms would result in a wide spread of δ\(^{13}\)C\(_{\text{org}}\), ranging from the composition of the primary producers to that of the heterotrophic consumers. In the Hardey and Bellary formations, the strong mode around -40‰ (minimum -42.5‰, Fig. 7) suggests that a single process was dominant and is thus inconsistent with heterotrophic methanogenesis.

We are left with autotrophic methanogenesis, acetogenesis (the acetyl-CoA carbon fixation pathway) and methanotrophy. Methanotrophy was probably insignificant for lack of an oxidant. The scarcity of pyrite in these sediments suggests that sulfate reducers were minor, and as noted above, the presence of redox-sensitive detrital minerals, the scarcity of δ\(^{13}\)C\(_{\text{org}}\) values within the range of cyanobacteria, as well as the nitrogen isotope data argue against appreciable O\(_2\). Collectively, these observations suggest that methanotrophy was not a major metabolisms in the Hardey and Bellary settings. Hence, autotrophic methanogenesis and/or acetogenesis using atmospheric H\(_2\) and CO\(_2\) are perhaps the most plausible metabolisms in these fluvio-lacustrine settings. Importantly, autotrophic methanogens require a minimum atmospheric H\(_2\) mixing ratio of around 40 ppm (Kral et al., 1998), which is at the bottom end of values inferred from climate models of the Archean (e.g. Kharecha et al., 2005; Claire et al., 2014) and may provide a useful lower limit.

5.5.4. Methanotrophy in the Tumbiana, Kylena and Mt. Roe formations
The light but highly variable $\delta^{13}C_{\text{org}}$ data from the more mafic Tumbiana, Kylena and Mt. Roe formations imply some contribution from either heterotrophic methanogenesis or some form of methane or acetate assimilation. Slotznick & Fischer (2016) tried to rule out methanotrophy in the Tumbiana Formation on the basis of inorganic $\delta^{13}C_{\text{carb}}$ data from stromatolites. Methanotrophy generates isotopically light CO$_2$, which mixes with other dissolved inorganic carbon (DIC) and can lead to isotopically light carbonate precipitates. The two stromatolites analyzed by Slotznick & Fischer (2016) showed a tight cluster of $\delta^{13}C_{\text{carb}}$ from -1‰ to +1‰, which they argued was inconsistent with methanotrophic CO$_2$ production. However, a compilation of 150 $\delta^{13}C_{\text{carb}}$ data points for the Tumbiana from our study and previous investigations (Thomazo et al., 2009; Coffey et al., 2013) shows a wide range of $\delta^{13}C_{\text{carb}}$ values from -9.2‰ to +1.9‰, with a mean of -0.8 ± 2.0‰ (Fig. 4a). In the Kylena Formation, $\delta^{13}C_{\text{carb}}$ values range from -5.7‰ to +2.1‰ (mean +0.1 ± 2.2‰, n=19, Flannery et al., 2014, this study). The more negative values tend to occur in samples with less than 40% carbonate and are plausibly explained by mixing of methanotrophic CO$_2$ with a larger dissolved DIC reservoir. As discussed above, the Tumbiana lakes may have been alkaline with a pH > 9, and hence DIC levels may have exceeded 1000 mM—using the framework of Slotznick & Fischer (2016, their Fig. 4a). At such high DIC concentrations, a contribution of isotopically light CO$_2$ to DIC via methanotrophy would be strongly diluted. Figure 4b of Slotznick & Fischer (2016) suggests the expected $\delta^{13}C_{\text{carb}}$ values at high pH would be heavier than -2.5‰, making a methanotrophic contribution difficult to detect – except from samples with lower carbonate abundances where the methanotrophic signature becomes more obvious (Fig. 4a). Acetate-based anaerobic metabolisms may still have been present, but the occurrence of numerous light $\delta^{13}C_{\text{carb}}$ values supports previous arguments for methanotrophy during deposition of the Tumbiana and Kylena formations (Hayes, 1994; Hinrichs, 2002; Eigenbrode & Freeman, 2006). Light $\delta^{13}C_{\text{carb}}$ values paired with light $\delta^{13}C_{\text{org}}$ values also argue against heterotrophic methanogenesis as the sole cause of the light $\delta^{13}C_{\text{org}}$ data, because that metabolism does not generate isotopically light CO$_2$ or HCO$_3^-$.

5.5.5. Variable electron acceptors for methanotrophs

The likelihood of methanotrophy leaves open the question of which substrate was used as an electron acceptor for the oxidation of methane. We suspect that both aerobic and anaerobic processes may have played a role. In the Mt. Roe Formation, the moderate amounts of pyrite (Fig. 5) may be a result of sulfate reduction, which can be coupled to methane oxidation even at low sulfate concentrations (Beal et al., 2011). We see no evidence of O$_2$ production in the Mt. Roe Formation, which would also rule out nitrate as an electron acceptor. There is no evidence for extensive iron redox cycling in any of these non-marine settings, which is also supported by iron isotope data (Czaja et al., 2010). Hence, sulfate would be the most plausible oxidant for methane oxidation in the Mt. Roe Formation. In the Tumbiana and Kylena formations, where stromatolites (likely cyanobacterial) are common and sulfide minerals scarce (Buick, 1992, this study), aerobic methanotrophy is likely to have played a more important role. However, small but significant fractionations in sulfur isotopes suggest that some microbial sulfate reduction also occurred at this site (Thomazo et al., 2009). Light $\delta^{13}C_{\text{carb}}$ values in relatively carbonate-poor mudrocks (Fig. 4a) are consistent with a diagenetic process that oxidized methane below the oxic surface layer of the Tumbiana lakes (i.e., sulfate reduction linked to anaerobic oxidation of methane). We therefore favor a combination of aerobic and anaerobic metabolisms for methanotrophy.

6. Environmental patterns
In summary, our data reveal patterns in microbial metabolism in the Fortescue Group that appear to have been driven by physicochemical characteristics of the environments (Fig. 8, Table 1), consistent with studies of modern microbial biogeography (Martiny et al., 2006). The more felsic Hardey and Bellary formations were deposited in open fluvio-lacustrine settings hosted N2-fixers and perhaps autotrophic methanogens using atmospheric H2 and CO2, but we see no clear evidence for photosynthetic activity, sulfate reduction or methanotrophy. These metabolisms may have been present at low levels, but the tightly clustered δ13Corg data around -40‰ suggest that they were not producing significant amounts of biomass. Nitrogen isotopes are consistent with nitrogen-limited conditions and molybdenum-based fixation of atmospheric N2. In the mafic, evaporitic Tumbiana and Kylena formations, which represent closed systems, our iron-speciation data suggest that ferrous iron was insoluble and that sulfide was not abundant. This conclusion supports the inference of alkaline conditions (Stüeken et al., 2015b). It also supports the interpretation that the abundant stromatolites in these stratigraphic units were built by oxygenic phototrophs, because alternative electron donors other than H2O were scarce (Buick, 1992). The light δ13Corg values down to -57‰ are most likely the result of methanotrophy, using a combination of O2 and SO4²⁻ as oxidants. This conclusion is supported by light δ13Ccarb values ranging down to -9‰ (cf. Slotznick & Fischer, 2016) and slightly fractionated sulfur isotopes in pyrite (Thomazo et al., 2009). In the mafic but shallower ephemeral ponds of the Mt. Roe Formation, similarly light δ13Corg values are also interpreted as evidence of methanotrophy, but in this case we have seen no indication of photosynthetic activity that could have provided O2. Instead, SO4²⁻ may have been the dominant oxidant of CH4, as indicated by the moderate amounts of pyrite in some of our Mt. Roe samples.

Overall, methanotrophy appears to have been favored in mafic systems, and oxygenic photosynthetic activity was favored in long-lived evaporitic mafic systems that evolved to alkaline conditions (Fig. 8). Importantly, modern cyanobacteria are most productive and diverse in alkaline lakes relative to other lakes and the open ocean (Zavarzin, 1993), and hence mafic volcanic terrains on Archean continents may have been ideal habitats for them to thrive. We speculate that the propensity for methanotrophy in mafic environments may be related to a higher supply of oxidants in the form of biogenic O2 from thriving cyanobacterial colonies and/or biogenic or volcanogenic SO4²⁻.

7. Ecological and evolutionary implications

Our results support the idea that environmental diversity on the Archean Earth fostered microbial diversity. A range of microbial metabolisms along environmental gradients has previously been described from marine environments of Archean and Proterozoic age (e.g. Eigenbrode & Freeman, 2006; Tice & Lowe, 2006; Guy et al., 2012; Stüeken, 2013; Luo et al., 2014; Koehler et al., 2017). In those marine settings, environmental gradation is usually ascribed to differing redox settings between shallow and deep habitats. Our data suggest that biological diversity was at least partly controlled by geological processes such as volcanism and sediment composition and partly by geographical and hydrological parameters. Such environmental controls are commonly seen today in studies of microbial biogeography (Martiny et al., 2006), but they have not previously been recognized in ancient microbial ecosystems. By analogy to the extreme diversification seen in macroscopic multicellular eukaryotes in terrestrial environments (Little, 1990; Kenrick & Crane, 1997), it is conceivable that geological, geographical and hydrological factors such as those described here contributed to the evolution of early life, in addition to stochastic biological mutations and innovations. Varying physico-chemical conditions like those
manifest across the Fortescue water bodies could well have acted as drivers for biochemical evolution in earlier times by emphasizing different metabolic substrates at differing bio-availabilities in distinct habitats.

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**Figure captions**

**Figure 1: Geographic location of the Fortescue Group.** The map was modified from Buick (1992). Red stars mark sampling sites for this and previous studies that produced data used in this manuscript. BC = Beabea Creek (Tumbiana). WRL-1 (Tumbiana), SV-1 (Tumbiana), ABDP-3 (Hardey), ABDP-6 (Mt. Roe) and ABDP-10 (Tumbiana) are drill cores.
Figure 2: Stratigraphy of the Neoarchean Fortescue Group. The stratigraphic sequence and ages are adopted from Blake et al. 2004.
Figure 3: Field photos and photomicrographs. (a) Felsic conglomerate in the Bellary Formation. Lens cap in the top left corner is ~7cm wide. (b) Silt lamina in the Mt. Roe Formation. Opaque silt grains are devitrified mafic volcanic fragments. Scale bar = 0.3mm. (c) Graded lamination in shale of the Mt. Roe Formation. Scale bar = 0.3mm. (d) Silt to very fine sand lamina in the Hardey Formation, crossed polars. Grains are mostly quartz, feldspar and muscovite. Scale bar = 0.3mm. (e) Graded shale in the Hardey Formation. Scale bar = 0.3mm. (f) Wave ripples (lower surface) and current ripples (upper surface) in the Hardey Formation. Dashed lines on the upper surface mark catenary ripple crests. Width of hammer stem ~4cm. (g) Stromatolitic carbonate in the Kylena Formation. (h) Stromatolites in the Tumbiana Formation. (i) Teepee structure in the Tumbiana Formation as evidence of desiccation. Hammer is ~40cm.
Figure 4: Carbon isotope trends in the Tumbiana Formation. (a) Inorganic carbon isotopes versus carbonate content. Dashed lines mark marine range (Eigenbrode & Freeman, 2006). (b) inorganic versus organic carbon isotopes.

Figure 5: Iron speciation. Boundaries between oxic, ferruginous and euxinic conditions are taken from Poulton & Canfield (2011) calibrated for marine sediments. The threshold between oxic and anoxic settings as inferred from Fe\textsubscript{HR}/Fe\textsubscript{T} does not apply in lacustrine and fluvial settings, because this ratio can be affected by high sedimentation rates and pH effects on Fe solubility (see text). The threshold for euxinia based on Fe\textsubscript{PY}/Fe\textsubscript{HR} remains valid. Our data thus indicate that none of these sites were euxinic, except for perhaps parts of the Mt Roe Formation.
**Figure 6: Molybdenum isotopes.** Mt. Roe and Hardey data are taken from Wille et al. (2013). The basalt range is that of basalts underlying the Tumbiana Formation. It encompasses the range of modern basalts (-0.2‰ to +0.3‰, Voegelin et al., 2014). The wide range of values in sediments from the Tumbiana lakes indicates that Mo was mobile in this setting, possibly as a result of mildly oxic conditions proximal to cyanobacterial mats (Buick, 1992; Flannery & Walter, 2012).

![Molybdenum isotope plot](image)

**Figure 7: Carbon and nitrogen isotopes.** The mafic settings show evidence of NH$_3$ volatilization at high pH (>9), consistent with the development of alkaline conditions in modern volcanic terrains (Kempe & Kazmierczak, 2011). The felsic settings are dominated by biological N$_2$ fixation. Evidence of methanotrophy occurs in the mafic sites but not in the felsic environment. The heavier end of the δ$^{13}$C$_{org}$ spectrum in the Tumbiana Formation may reflect photosynthetic organisms as invoked by Buick (1992) and Flannery & Walter (2012). All data from the Mt Roe Formation and most data from the Hardey Formation are from this study. Additional data are taken from Thomazo et al. (2011), Stücken et al. (2015a) and Stücken et al. (2015b).
Figure 8: Environmental and metabolic patterns in the non-marine Fortescue Group. In the more felsic and hydrologically open settings (Bellary and Hardey formations), carbon isotopes suggest the presence of autotrophic methanogens. Other metabolisms were likely minor, if present. The mostly mafic and hydrologically open Mt Roe setting shows evidence of methanotrophy, possibly coupled to microbial sulfate reduction. The mafic and evaporitic Tumbiana and Kylena lakes likely hosted oxygenic photosynthesizers, as well as methanotrophs and minor sulfate reducers. Hence environmental factors evidently affected the relative abundances of microbial groups.
### Table 1: Environmental characteristics

<table>
<thead>
<tr>
<th>Formation</th>
<th>Depth (substrate topography and sed. thickness)</th>
<th>Extent (strike length)</th>
<th>proportion of exogenous siliciclastic sediment</th>
<th>evaporation</th>
<th>tuff input</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tumbiana</td>
<td>a few 10s of meters</td>
<td>many 10s of kilometers</td>
<td>little (mostly closed)</td>
<td>much (halite)</td>
<td>high</td>
</tr>
<tr>
<td>Kylena</td>
<td>~10 meters</td>
<td>~10 kilometers</td>
<td>none (mostly closed)</td>
<td>some (carbonate)</td>
<td>high</td>
</tr>
<tr>
<td>Hardey</td>
<td>a few 10s of meters</td>
<td>a few kilometers</td>
<td>much (open)</td>
<td>none evident</td>
<td>none</td>
</tr>
<tr>
<td>Mt Roe</td>
<td>a few meters</td>
<td>&lt;1 kilometer</td>
<td>moderate (pulsed)</td>
<td>none evident</td>
<td>moderate</td>
</tr>
<tr>
<td>Bellary</td>
<td>many 10s of meters</td>
<td>a few kilometers</td>
<td>some (open)</td>
<td>none evident</td>
<td>none</td>
</tr>
</tbody>
</table>
References


Watanabe Y, Bevacqua DC, Ohmoto H (2004) Microbial ecosystem in the oldest freshwater lake revealed from a drill core of the 2.76 Ga Hardey Formation, Pilbara District, Western Australia. AGU Fall Meeting Abstracts, 1, 0996.


