Abstract

A persistently aerobic marine nitrogen cycle featuring the biologically mediated oxidation of ammonium to nitrate has likely been in place since the Great Oxidation Event (GOE) some 2.3 billion years ago. Although nitrogen isotope data from some Neoarchaean sediments suggests transient nitrate availability prior to the GOE, these data are open to other interpretations. This is especially so as these data come from deep-water environments that were spatially divorced from shallow-water settings that were the most likely sites for the accumulation of oxygen and the generation of nitrate. Here we present the first nitrogen isotope data from contemporaneous shallow-water sediments to constrain the nitrogen cycle in shallow Late Archean settings. The BH-1 Sacha core through the Campbellrand-Malmani carbonate platform records a transition from a shallow siliciclastic/carbonate ramp to a rimmed carbonate shelf with the potential for reduced communication with the open ocean. In these settings nitrogen isotope ($\delta^{15}$N) data from sub-
peri-tidal and lagoonal settings are close to 0‰, indicating diazotrophy or the complete utilization of
remineralised ammonium with an isotopic composition of near 0‰. Our dataset also includes
negative δ^{15}N values that suggest the presence of an ammonium pool of concentrations sufficient to
have allowed for non-quantitative assimilation. We suggest that this condition may have been the
result of upwelling of phosphorus-rich deep waters into the photic zone, stimulating primary
productivity and creating an enhanced flux of organic matter that was subsequently remineralised
and persisted in the dominantly anoxic Neoarchaean marine environment. Notably, we find only
limited evidence of coupled nitrification/denitrification, even in these shallow water environments,
calling into question previous suggestions that the Late Archean nitrogen cycle was characterized by
widespread aerobic nitrogen cycling. Rather, aerobic nitrogen cycling was likely spatially
heterogeneous and tied to loci of high oxygen production while zones of shallow water anoxia
persisted.

Keywords

• Nitrogen Isotopes
• Carbon Isotopes
• Neoarchean
1. Introduction

Nitrogen (N) is an essential nutrient for the construction of all biomolecules. Despite the biological importance of N\textsubscript{2} and its abundance in the atmosphere, only diazotrophic (N\textsubscript{2}-fixing) microbes can directly assimilate di-nitrogen into biomass, which ultimately provides the primary source of nitrogen to the biosphere. In the absence of oxygen, the uptake of ammonium (NH\textsubscript{4}\textsuperscript{+}) from remineralised diazotrophic biomass is the primary source of nitrogen for non-diazotrophs. In oxygen-rich waters, NH\textsubscript{4}\textsuperscript{+} is rapidly recycled or oxidised by nitrifying microbes to nitrite (NO\textsubscript{2}\textsuperscript{-}) and nitrate (NO\textsubscript{3}\textsuperscript{-}). Nitrate is an important component of dissolved inorganic nitrogen (DIN) in oxygenated waters, and serves as the dominant nitrogen source for primary productivity in modern surface oceans. In such conditions diazotrophic activity will be reduced due to competition with nitrate assimilators for other nutrients (e.g. Sigman et al., 2009).

In low-oxygen settings, such as modern oxygen minimum zones (OMZs), NO\textsubscript{2}\textsuperscript{-} and NO\textsubscript{3}\textsuperscript{-} can also be utilised as electron acceptors in chemotrophic metabolisms. For example, NO\textsubscript{3}\textsuperscript{-} can be used in heterotrophic denitrification or during dissimilatory nitrate reduction to ammonium (DNRA) (Granger et al., 2008), whilst NO\textsubscript{2}\textsuperscript{-} is utilised in the anaerobic oxidation of ammonium (anammox).

Importantly, both annamox and denitrification remove bioavailable nitrogen from the oceanic reservoir and return it to the atmosphere (Cline and Kaplan, 1975), which in extreme cases can lead to nitrate limitation. When bioavailable nitrogen is scarce an expansion of diazotrophy can occur due to reduced competition for nutrients, if other nutrients like Fe or bioactive trace elements are not limiting (Weber and Deutsch, 2013; Sigman et al., 2009).

Due to the redox sensitive nature of these N cycling processes, the global N cycle is believed to have evolved over geologic time alongside changes in the redox state of Earth’s oceans and atmosphere (e.g., Stüeken et al., 2016). The 2.33 Ga Great Oxidation Event (GOE) (Luo et al., 2016) marked the time when atmospheric oxygen (O\textsubscript{2}) first exceeded 10\textsuperscript{-5} times present atmospheric levels (PAL), as constrained by the disappearance of mass-independent fractionation (MIF) of sulphur isotopes.
This change in surface redox conditions seemingly coincided with the widespread expansion of aerobic nitrogen cycling in the world’s oceans (e.g., Zerkle et al., 2017; Luo et al., 2018). However, this narrative could be overly simplistic. In particular, small increases of around 2‰ in δ¹⁵N values are preserved in the 2.7-2.5 Ga sediments of the Campbellrand-Malmani carbonate platform from South Africa (Godfrey and Falkowski, 2009), a setting that is spatially and temporally correlated to the sediments analysed in this study, albeit further offshore and a deeper depositional setting. Other positive excursions in δ¹⁵N are reported from studies of Australian Neoarchaean sediments (Garvin et al., 2009; Busigny et al., 2013) and in all settings these nitrogen isotope values have been explained by transient or localized aerobic nitrogen cycling. Alternatively, these small positive shifts in δ¹⁵N values could represent uptake of a residual pool of ¹⁵N-enriched ammonium produced by partial nitrification or partial assimilation, or by nitrogen redox cycling independent of environmental oxygenation (e.g., Thomazo et al., 2011; Busigny et al., 2013; Ader et al., 2016). However, there are currently no δ¹⁵N records of contemporaneous shallow-water sediments available to distinguish between these alternatives.

Here, we investigate δ¹⁵N values preserved in shallow-water sediments from the 2.7-2.5 Ga Campbellrand-Malmani carbonate platform. These sediments, from the BH-1 Sacha core, represent the shallow-water equivalent to previously investigated deeper-water sediments (GKP01 core; Fischer et al., 2009), in which δ¹⁵N values rose by about 2‰ and were interpreted to represent transient aerobic nitrogen cycling (Godfrey and Falkowski, 2009). If aerobic nitrogen cycling was even intermittently pervasive in this basin, we would expect sediments from these shallower water settings to record similar δ¹⁵N values to deeper-water sediments, particularly given their potential proximity to surface ocean oxygen oases, as shown by the presence of microbial sedimentary structures (Altermann and Siegfried, 1997).
2. Tracking the nitrogen cycle through time

Transformations in the marine nitrogen cycle can modify nitrogen isotope signals \( \delta^{15}N (\%) \)
\[
\delta^{15}N = \left( \frac{^{15}N/^{14}N}_{\text{sample}} / {^{15}N/^{14}N}_{\text{air}} - 1 \right) \times 1000
\]
leading to fractionation effects between the reactants and products (Casciotti, 2009; Sigman et al., 2009; Zerkle et al., 2008; Brunner et al., 2013; Möbius, 2013; Zhang et al., 2014; McCready et al., 1983; Ader et al., 2016). However, in modern settings, most nitrogen assimilation or redox transformations utilise all available substrates, leading to no apparent fractionation effects (Sigman et al., 2009). The exceptions are diazotrophy and nitrate/nitrite-reduction processes, including denitrification, DNRA, and anammox. Diazotrophy generally produces biomass with \( \delta^{15}N \) values between \( \sim +1 \) and \( \sim -1\% \) (Zhang et al., 2014; Bauersachs et al., 2009), although extreme values as low as \( \sim -4\% \) have been experimentally demonstrated (e.g., as reviewed in Zerkle et al., 2008). The contribution of diazotrophy to total biomass in modern ocean settings is limited except under oligotrophic conditions, as \( \text{NO}_3^- \) and \( \text{NH}_4^+ \) assimilators compete more successfully for other nutrients (Weber and Deutsch, 2013). However, in most oxygen-rich settings, \( \text{NH}_4^+ \) sourced from remineralised biomass is rapidly oxidized, such that nitrate is the main form of nutrient N. In the modern ocean incomplete denitrification in the water column renders this pool of bioavailable \( ^{15}N \)-enriched, leading to marine organic matter (OM) with an average \( \delta^{15}N \) of \( +5 \% \) to \( +6\% \) (Peters et al., 1978; Sigman et al., 2003; Galbraith et al., 2013).

The \( \delta^{15}N \) of marine biomass thus records the dominant forms of bioavailable nitrogen utilised for primary productivity, particularly on a local scale. These signals are ultimately archived in sedimentary rocks, either directly in OM or retained as mineral-associated nitrogen derived from degraded OM (e.g., Stüeken et al., 2016; Freudenthal et al., 1999). Variations in the \( \delta^{15}N \) of sedimentary rocks can thus be utilized to track changes in the marine nitrogen cycle through geological time.

3. Geological setting
The Campbellrand-Malmani Platform in South Africa (Fig. 1A, B and C) is part of the Transvaal Supergroup. A ~3,700 m section of these rocks was recovered in the BH1-Sacha core and logged in detail by Altermann and Siegfried (1997). The lowermost recovered strata are a succession of interbedded doloarenites, quartzites, and shales with microbial laminites from the top of the Vryburg Formation (Fm) (Altermann and Siegfried, 1997). Thick (> 10 m) shale beds in the lower part of the core likely represent deeper shelf conditions when marine transgressions mark the beginning of several deepening and shallowing cycles, with carbonates likely representative of the shallowest regressive conditions. Thin (< 10 m) shales interbedded with carbonates throughout the core are mostly likely evidence of slightly deeper lagoonal conditions during generally regressive periods (Altermann and Siegfried, 1997). A major flooding surface defines the base of the overlying Boomplaas Fm, but its top is defined by an oolite bed indicating shallow, wave-agitated conditions. The Lokammona Fm is marked by several, variably developed, shoaling cycles similar to those in the Vryburg Fm (Sumner and Beukes, 2006). These three formations comprise the Schmidtsdrift Subgroup, and are overlain by the Monteville Fm of the lowermost Campbellrand Subgroup, which shows cycles similar to the Lokammona Fm (Sumner and Beukes, 2006). Together, these four formations are interpreted as representing deposition of sediments on an evolving ramp structure (Fig. 1B).

From the Reivilo Fm upwards, depositional conditions are considered to have been generally shallow, with subtidal, intertidal or lagoonal conditions dominating as inferred from the presence of domal, columnar, elongate and small, bifurcating, finger-like stromatolites, birds-eye structures, flat-pebble breccias and chert beds (Altermann and Siegfried, 1997; Sumner and Beukes, 2006; Fischer and Knoll, 2009). These facies are considered to represent the replacement of ramp-like conditions with a rimmed-, carbonate-dominated shallow-shelf, platform-top (Sumner and Grotzinger, 2004; Sumner and Beukes 2006). A maximum metamorphic grade of sub-greenschist is reported for the section of the Campbellrand-Malmani Platform studied here (Button, 1973; Miyano and Beukes, 1984; Fischer et al., 2009), consistent with minimal alteration of stable isotope values as indicated by
our data (see discussion below). A U-Pb zircon age of 2,714 ± 8 Ma from the underlying Ventersdorp
Supergroup provides a maximum age for the Campbellrand-Malmani Platform (Armstrong et al.,
1991) and tuffs within the Schmidtsdrift and Campbellrand subgroups have yielded stratigraphically
coherent Neoarchaean ages (Barton et al., 1995; Walraven and Martini, 1995; Fischer et al., 2009;
Knoll and Beukes, 2009; ranges of published dates are in the supplementary material).

4. Methods

Samples were collected from the National Core Repository at Donkerhoek (Pretoria, South Africa).
Sampling was predominantly focussed on black shales, which were anticipated to have high-TOC
content. Samples were ground into homogenous powders in agate ball mills, and carbonate was
removed via two 24 hour digestions in 10 % (vol/vol) HCl. After each digestion the samples were
centrifuged and the supernatant was discarded and replaced with fresh 10% HCl. Sample residues
were washed until pH neutral using ultrapure water (18.2 MΩ•cm) and dried at < 40 °C. Carbonate
content was calculated gravimetrically from dry sample residues which were then homogenised
using an agate pestle and mortar and stored in glass vials. Several samples of mid-grey carbonate
rocks, assumed to contain some OM, were also decarbonated to assess the possibility of extracting
kerogen. This method of carbonate abundance determination is prone to inaccuracy. However, here
it is used only to provide a broad assessment of lithological characteristics, for example between
rocks with high carbonate content (e.g. > 85%), and those with low carbonate concentrations (e.g.
<10%). Kerogen was extracted at the University of St Andrews by digestion of decarbonated residues
in HF/HCl following established protocols (e.g., Zerkle et al., 2017).

Nitrogen isotope values for kerogen (δ\(^{15}\)N\(_{\text{org}}\)) and decarbonated sediments (δ\(^{15}\)N\(_{\text{bulk}}\)) were
determined by nano-EA-IRMS in the GAPP Lab at Syracuse University, USA. Prior to analyses,
samples and standards were placed into a vacuum chamber overnight to remove atmospheric
nitrogen and flooded with Argon (Ar) prior to analysis. Upon introduction into the EA, tin capsule
sealed samples were further purged with helium (He) for 45 seconds prior to combustion in an
Elementar Vario Isotope Cube coupled to a Trace Gas analyser. Oxidation and reduction reactor temperatures were 1100 °C and 650 °C, respectively. Helium carrier gas flow was 150 ml/min; and the O₂ pulse was set for 90 seconds. Resultant sample gas was trapped in a liquid-N silica gel filled cryotrap, before release to an Elementar Isoprime 100 IRMS via an Agilent CarboBond capillary column (25 m x 0.53 mm x 5 um) with a He flow rate of ~2 cm³ min⁻¹ (e.g., Polissar et al., 2008; Zerkle et al., 2017; Luo et al., 2018).

Data accuracy for δ¹⁵Nₐᵣg was assessed using the IAEA N1 (Ammonium Sulphate, (NH₄)₂ SO₄) standard which provided a mean δ¹⁵N value of 0.11± 0.50‰ (1σ; n=30) versus a certified value of +0.40 ± 0.2‰ (1σ).

The extremely sensitive nature of the nano-EA-IRMS method generates relatively large peak height sizes for the blank, in comparison to standard IRMS techniques. This, coupled with relatively nitrogen-poor kerogen samples, led to blank-sample ratios that could introduce variability in δ¹⁵Nₐᵣg measurements. Blank-corrected δ¹⁵Nₐᵣg were calculated by blank extraction using the following equation:

$$\delta^{15}N_{\text{org}} = \frac{(\text{Peak Area}_{\text{sample}} \times \delta^{15}N_{\text{sample}}) - (\text{Peak Area}_{\text{blank}} \times \delta^{15}N_{\text{blank}})}{(\text{Peak Area}_{\text{sample}} - \text{Peak Area}_{\text{blank}})}$$

The size (Peak Areablank) and isotopic composition of the blank (δ¹⁵Nblank) contribution to each sample measurement was calculated by measuring a blank (without tin cup - as we have found no difference in blank size and isotopic value whether a tin cup was included or not) before and after each five samples and calculating incremental change for both values between the two blanks.

Duplicates of sample BH1-330, corrected to IAEA N1 indicate reproducibility of δ¹⁵Nₐᵣg was ± 0.02‰ (1σ), whilst triplicates of samples BH1-1963 and BH1-3144.33 produced values ± 0.49‰ (1σ) and ± 0.32‰ (1σ), respectively. Duplicates of samples with large blank contributions resulting from exceptionally low Nₐᵣg produced higher variability, resulting in δ¹⁵Nₐᵣg measurements on samples below 0.05 wt. % N being discarded. Error bars for δ¹⁵Nₐᵣg plots were calculated using standard
deviation of triplicate (BH1-1963.6; 0.49‰) and standard deviation of standards (0.50‰; n=30),
where error bar = √(error of standards² + largest error of corrected sample²)).

Decarbonated sediments were also analysed to measure δ¹⁵Nbulk values. Again, nitrogen abundances in these decarbonated residues (TNbulk) were exceptionally low, ranging from 0.0018 wt. % to 0.017 wt. %, so samples were run in triplicate and δ¹⁵Nbulk values determined by Keeling plots. These samples were corrected to the reference material NIST1547 – Peach Leaves, which returned a δ¹⁵N value of +1.76‰ (n=8, determined by Keeling plots) against a certified value of +1.98‰ for measurement of triplicate samples and +2.10‰ (n=5) for sample 3364 reruns (Table S4). Error bars for δ¹⁵Nbulk reflect the combined standard deviation of standards and the intercept errors of keeling plots for each individual sample triplicate using the same formula. These individual errors are shown in supplementary table four.

Organic carbon isotope values (δ¹³Corg) for this study were measured at the University of St Andrews via flash-combustion of decarbonated residues using a Costech 4010 EA equipped with a zero blank autosampler and interfaced with a Thermo Finnigan Delta Plus XP IRMS in continuous flow mode. Data accuracy was verified using an internal standard (n=5), and returned values of -25.07 ± 0.18‰ (1σ) and -25.05 ± 0.11‰ (1σ) against an accepted value of -25.04‰.

Nitrogen and carbon isotope values are reported using the standard delta notation showing per mil deviations from V-PDB for δ¹³C, and relative to Nair for δ¹⁵N.

The abundance of nitrogen in kerogen (Norg) and in decarbonated residues (TNbulk) and the total organic carbon content (TOC) of whole rock powders were calculated by comparing peak areas generated during isotope analyses with those of standards with known abundances (Table S3 and S4). To calculate TOC abundances, the carbon yield was adjusted according to the mass lost during decarbonation.
Potassium (K) and iron (Fe) abundances for samples 2709 and 3364 were determined by XRF at Department of Earth and Environment, Franklin and Marshall College, USA. Standards returned 0.51 wt. % ±0.0014 (1σ) against a certified value of 0.52 wt. % for K₂O and 12.42 wt. % ±0.0135 (1σ) against a certified value of 12.30 wt. % for Fe₂O₃. Data for the remaining four samples were analysed at University of St Andrews using standard XRF providing a standard deviation of 1σ of 0.02 wt. % for K₂O.

5. Results

Nitrogen isotope values from kerogen (δ¹⁵Norg) for BH1-Sacha (Fig. 2; Table 1) range from −2.73‰ to +3.18‰ (mean −0.23 ± 1.59‰ here and elsewhere, 1σ, n=21). With the exception of samples BH1-3113.32 and BH1-3144.33, all δ¹⁵Norg values of greater than 0‰ are from samples with carbonate abundance in excess of 85% (Fig. 2; Table S1). Bulk nitrogen isotope values (δ¹⁵Nbulk) range from −3.30‰ to +2.94‰ (mean +0.58 ± 2.14‰, n=6). Organic carbon isotope values (δ¹³Corg) for BH1-Sacha (Fig. 2; Table S1, including a subset from Izon et al. [2015]) range from −46.09‰ to −27.36‰ (mean −36.69 ± 3.76‰, n=26).

For data binned by depositional setting, the ramp-top settings of the Schmidsdrift Subgroup and overlying Monteville Fm have mean δ¹⁵Nbulk values of +0.11 ± 2.01‰ (n=5) and δ¹⁵Norg values of +0.22 ± 1.96‰, n=7 (Fig. 2; Table S1). The highest δ¹⁵Norg values measured are from two carbonate-rich facies: +2.48‰ at 3306.3 m core depth, and +3.18‰ at 3114.5 m. The highest and lowest δ¹⁵Norg values from siliciclastic facies in this section are +0.53‰ and −2.03‰, whilst δ¹⁵Nbulk values range from −3.30‰ to +2.02‰. Mean δ¹³Corg values from these ramp depositional settings are −34.27 ± 3.63‰, n=11.

In the platform-top/rimmed-shelf facies from the base of the Reivilo Fm through the Gamohaan Fm, the mean δ¹⁵Norg values are −0.52± 1.36‰ (n=15) with a single δ¹⁵Nbulk value of +2.94‰ (Fig. 2; Table S1). Again, the three carbonate samples had significantly higher δ¹⁵N values, up to +3.17‰. Mean δ¹³Corg values for these facies are −38.13 ± 3.26‰ (n=15).
6. Discussion

6.1. Preservation of Primary Isotopic Signals

Diagenetic and metamorphic processes can produce changes in primary carbon and nitrogen isotope values, generally driving δ^{13}C_{org} and δ^{15}N values heavier and/or causing a divergence between bulk rock and kerogen δ^{15}N. At the sub-greenschist facies conditions of the Campbellrand-Malmani Platform (Button, 1973; Miyano and Beukes, 1984; Fischer et al., 2009) such effects should be minimal, and not exceed 1 – 2‰ (Stüeken et al., 2017). However, we examined trends in stable isotope values alongside elemental abundances to evaluate possible contributions from post-depositional alteration.

Thermal maturation during diagenetic or metamorphic processes can preferentially remove ^{12}C and ^{14}N from sediments, leading to diagnostic positive correlations between δ^{15}N_{bulk} and δ^{13}C_{org}, and negative correlations between the abundances of TOC and δ^{13}C_{org}, the abundances of TN_{org} and δ^{15}N_{org}, and the abundances of TN_{bulk} and δ^{15}N_{bulk}. We note a weak positive correlation between TN_{org} and δ^{15}N_{org} (R^2 = 0.28; Fig. 3A) and a moderately positive correlation between TN_{bulk} and δ^{15}N_{bulk} (R^2 = 0.51; Fig. 3E). We also note a weak positive correlation between δ^{15}N_{org} and δ^{13}C_{org} (R^2 = 0.24; Fig. 3B) and a weak negative correlation between TOC/TN_{org} and δ^{15}N_{org} (R^2 = 0.37, Fig. 3C). We note no correlation between TOC and δ^{13}C_{org} (R^2 = 0.10; Fig. 3D), and no correlation between δ^{15}N_{bulk} and δ^{13}C_{org} (R^2 = 0.05; Fig. 3F). Therefore, our data show little evidence for significant post-depositional alteration of δ^{13}C_{org} or δ^{15}N, consistent with the low metamorphic grade of these sediments. We also examine the potential influence of nitrogen adsorbed onto detrital clays by comparing potassium (K) content with TN_{bulk} and δ^{15}N_{bulk} (Figs. 3G and H).

We do note that there is a moderate correlation (R^2 = 0.69) between K and δ^{15}N_{bulk} suggesting that higher concentrations of clay-bound ammonium could be associated with more-positive δ^{15}N_{bulk} values, although, this correlation is somewhat influenced by the one sample in this small data set.
The sample from 3364m is unusual in that negative (-3.3‰) $\delta^{15}N_{\text{bulk}}$ values are associated with very low K abundances (0.97%) in comparison to the remaining samples (Table S6). When this sample is removed from the analyses the correlation between K and $\delta^{15}N_{\text{bulk}}$ is more modest ($R^2 = 0.32$) which more closely reflects the very limited correlation ($R^2 = 0.17$) between $TN_{\text{bulk}}$ and K abundances.

6.2. Nitrogen cycling in a marginal marine environment

The majority of samples analysed in this study are from relatively thin shale units that were intercalated with stromatolitic carbonates. These thin shales are interpreted to have been deposited during small marine transgressions, overprinted upon longer-term, relatively shallow, potentially lagoonal conditions in platform and ramp-top settings (Altermann and Siegfried, 1997; Sumner and Beukes, 2006; Ergolu et al., 2017). These thin shales are distinct from thicker shale units that likely represent more open and deeper marine settings. Given the shallow nature of this setting and the presence of microbial mats within the photic zone, such conditions should be prime candidates for pre-GOE oxygen oases. Although the persistence of sulphur MIF in the BH-1 Sacha record indicates that atmospheric oxygen concentrations must have remained less than 1 ppm during this time (Izon et al., 2015), oxygen concentrations could have been sufficiently high in the water column or locally associated with benthic microbial mats to allow for aerobic biogeochemical cycling. Indeed, trace element and iron speciation analyses from Neoarchaean sediments of the Campbellrand-Malmani Platform suggest a stratified ocean with mildly oxygenated surface oxygen oases (Kendall et al., 2010; Czaja et al., 2012; Ergolu et al., 2015) and localized oxygen production in microbial mats (Zerkle et al., 2012). Previous studies of trends in $\delta^{15}N$ from deeper water sediments from this basin have also been interpreted to represent transient aerobic N cycling (Godfrey and Falkowski, 2009).

We find that $\delta^{15}N_{\text{org}}$ values from siliciclastic sediments deposited in these near-shore settings are inconsistent with widespread coupled nitrification/denitrification. In particular, $\delta^{15}N$ values close to
0‰ are more consistent with a nitrogen cycle dominated by diazotrophy and the recycling of NH$_4^+$, and suggest that any NO$_3^-$ generated from localized oxidation of ammonium was quantitatively and rapidly removed by denitrification and anammox. Even if NO$_3^-$ was present, it likely provided a secondary nutrient N source given that ammonium could have been persistently available under widely anoxic water column conditions. In modern settings (e.g., Higgins et al., 2012) cyanobacteria can assimilate both NH$_4^+$ and NO$_3^-$ as a source of nitrogen, and it has been noted that genes controlling the assimilation of oxidised nitrogen sources can be repressed in the presence of ammonium (Flores et al., 2005).

The lower part of the BH1-Sacha core (e.g., the Schmidsdrift Subgroup and Monteville Fm) represents depositional conditions that were likely open to the ocean. Both δ$^{15}$N$_{\text{bulk}}$ and δ$^{15}$N$_{\text{org}}$ values in these ramp facies (with the exception of samples at 3114.5m and 3306.3m, discussed below) are again consistent with anaerobic nitrogen cycling. Intriguingly, one sample (3364m) provided a δ$^{15}$N$_{\text{bulk}}$ value of –3.3‰, and another sample (3125.05m) a δ$^{15}$N$_{\text{org}}$ value of -2.0‰, showing greater $^{15}$N depletion than biomass generally produced by modern marine diazotrophs. This is especially the case when considering that the long-term diagenetic or metamorphic effects could possibly have elevated δ$^{15}$N by a few per mille in comparison to the δ$^{15}$N of primary organic matter. We suggest that these strongly negative δ$^{15}$N values indicate the generation of biomass from partial assimilation of NH$_4^+$, as recently suggested for ~2.7 Ga sediments (Yang et al., 2019). Ammonium could have accumulated in deep waters under upwelling zones that supported high rates of diazotrophic productivity and organic matter export (Fig. 4A, B). The remineralisation of this organic matter at depth under anoxic conditions could have led to the formation of a deep-water NH$_4^+$ pool. In turn, these ammonium-rich waters could have been upwelled, leading to partial NH$_4^+$ assimilation by primary producers, thus forming $^{15}$N-depleted organic matter. Such settings could have included the ramp front during early platform formation (e.g., thicker shale units at 3364m) and later the mature platform front (the latter a depositional setting not recorded in the BH1-Sacha core), both of which represented relatively deeper conditions in contrast to shallow lagoonal depositional settings.
Other negative δ\textsuperscript{15}N values (e.g., those associated with thin shales intercalated with carbonates) could represent times when communication between the lagoonal setting and the open ocean was more vigorous and allowed NH\textsubscript{4}+-rich deep waters to temporarily inundate the lagoon.

Sediments at 3364m have high Fe concentrations (14.8%; Table S6) that approach the lower boundary of Fe abundances in banded iron formations, despite being deposited in open-ramp conditions. It is possible that in these Fe-rich conditions, diazotrophs utilizing the FeMo nitrogenase enzyme produced biomass with δ\textsuperscript{15}N values down to -4‰, similar to those reported in Zerkle et al. (2008). In addition, diazotrophs using alternative nitrogenase enzymes (e.g., Fe-Fe or Fe-V) have been shown to produce biomass with δ\textsuperscript{15}N values down to -7‰ when Mo is limited (Zhang et al., 2014). However, large variations in δ\textsuperscript{98}Mo have been reported for the lower Campbellrand-Malmani Platform, which provide evidence for a sizeable Mo reservoir (Eroglu et al., 2015). It is therefore unlikely that diazotrophs would have relied on these less efficient alternative enzymes in this environment.

Such low δ\textsuperscript{15}N values are unusual, but not unreported in modern oceans and in the more recent geologic past (e.g. Higgins et al., 2012). For example, δ\textsuperscript{15}N values of ~–5‰ have been reported for particulate nitrogen, and attributed to the partial utilization of ammonium (Rau et al., 1991) and nitrate (Altabet and Francois, 1994). In the latter case, these low values are associated with incomplete utilisation of nitrate in seasonally upwelling waters. However, this degree of 15N-depletion is not recorded in subsequent sedimentary δ\textsuperscript{15}N values due to the integration of multiple seasonal signals representing different rates of nitrate utilisation. The fact that some low δ\textsuperscript{15}N values are recorded in the Neoarchaean sediments studied here suggests that ammonium had accumulated to significant levels to remain under-utilised and that nitrogen limitation was not a significant factor in the wider oceans.

Intriguingly, widespread partial assimilation of ammonium could have left the residual pool of NH\textsubscript{4}+-relatively 15N-enriched (e.g., Yang et al., 2019). If transported offshore, uptake from a pool of heavier
NH$_4^+$ could generate $^{15}$N-enriched organic matter in more distal ocean settings (Fig. 4A, B; e.g., Busigny et al., 2013). This was illustrated numerically by Yang et al. (2019) using a simple Rayleigh model based on the experimental calibrations of isotope effects during ammonium uptake (Hoch et al., 1992). This scenario provides an alternative explanation for the presence of positive $\delta^{15}$N values recorded in relatively distal Neoarchaean settings in the Griqualand West Basin (e.g., Godfrey and Falkowski, 2009).

The majority of our $\delta^{15}$N data point to a largely anaerobic marine N cycle, driven by N$_2$ fixation and varying degrees of ammonium uptake and recycling. However, a handful of carbonate-rich facies show positive $\delta^{15}$N$_{org}$ values outside the range of typical values for N$_2$ fixation (up to +3.2‰; Fig. 4, Table S1). Elevated $\delta^{15}$N values could indicate syn-depositional oxidative degradation (Freudenthal et al., 2001; Möbius et al., 2010). However, these $\delta^{15}$N$_{org}$ values could hint at the potential influence of nitrification and incomplete denitrification in localised, mildly-oxygenated settings during the deposition of carbonates, potentially periods when waters were likely shallower than when thin muds were deposited. Ergolu et al. (2017) have argued that large differences in $\delta^{13}$C$_{org}$ values between carbonates and mudstones in the shallower parts of the Campbellrand-Malmani Platform were due to different consortia of microbes that comprise microbial mats in the two settings. They suggested that during the deposition of carbonates, microbial mats could have had larger proportions of oxygenic photoautotroph than during the deposition of mudrocks, with anaerobic microbes more prevalent during the deposition of the latter. This scenario could support shallow microbial mat communities as epicentres of highly localised nitrification-denitrification, without a major role for aerobic N cycling in the Late Archean.

6. Conclusions

Here we describe the Neoarchean nitrogen cycle in a shallow marginal marine setting, during the evolution of a well-defined carbonate platform into a relatively isolated platform- top/ rimmed-shelf lagoonal environment. Nitrogen isotope values range from -3.3 to +3.2‰ and are consistent with a
predominantly anaerobic nitrogen cycle, dominated by N₂ fixation and the assimilation and recycling of upwelling NH₄⁺. Evidence for the presence of oxidised nitrogen species is limited, suggesting that any NO₃⁻ generated during periods of transient oxygen availability was quantitatively removed by anaerobic respiration. However, rare δ¹⁵N values > 0 ‰ recorded in carbonate-rich sediments hint at highly localized areas of nitrification and partial denitrification. In addition, the inclusion of some very light δ¹⁵N values suggest partial assimilation of an upwelling pool of ammonium, leaving a residual pool of ¹⁵N-enriched NH₄⁺ that could potentially explain small increases in δ¹⁵N from contemporaneous open-ocean settings. We thus conclude that N cycling in the shore-proximal marine environment of the Griqualand West basin was controlled by anaerobic processes and recycling of bioavailable nitrogen, and that nitrification/denitrification was unlikely to have been widespread prior to the end of the Neoarchaean.

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References


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Figure Legends

**Fig. 1.** (A) Simplified stratigraphy of BH1-Sacha core (Altermann and Siegfried, 1997). (B) Simplified structure of Campbellrand-Malmani platform-top and ramp showing placement of BH1-Sacha core (Sumner and Beukes, 2006). (C) Position of BH1-Sacha drill site (star symbol) in South Africa (Sumner and Beukes, 2006). Dates in Figs. A and B are from a review in Sumner and Beukes (2009). More dates and references from correlative stratigraphy are presented in the supplementary information which confirm a date of 2588 to 2549 Ma for the Monteville Fm.

**Fig. 2.** Geochemical data for BH1-Sacha core. Nitrogen isotope data $\delta^{15}$N$_{org}$ (‰) are shown as black dots for silicates and triangles for carbonates, whilst $\delta^{15}$N$_{bulk}$ (‰) are shown as open pentagons. Carbonate abundance data (wt. %) are shown as dots. TOC (wt. %) and $\delta^{13}$C$_{org}$ (%) data from this study are shown as dots, whilst data from Izon et al. (2015) are shown as open squares. Symbols for lithologies in core diagram as per Fig. 1A. Error bars for $\delta^{15}$N$_{org}$ reflect the combined standard deviation of N1 standards and triplicates of sample BH1-1963.6 using the equation: error bar = $\sqrt{\text{ equation }}$.

**Fig. 3.** Cross plots of geochemical data. Filled circles represent data from this study. Open squares include TOC and $\delta^{13}$C$_{org}$ data from Izon et al. (2015). Data shown: (A) TN$_{org}$ and $\delta^{15}$N$_{org}$, (B) $\delta^{15}$N$_{org}$ and $\delta^{13}$C$_{org}$, (C) TOC/TN$_{org}$ and $\delta^{15}$N$_{org}$ (D) TOC and $\delta^{13}$C$_{org}$, (E) TN$_{bulk}$ and $\delta^{15}$N$_{bulk}$, (F) $\delta^{15}$N$_{bulk}$ and $\delta^{13}$C$_{org}$, (G) potassium and $\delta^{15}$N$_{bulk}$, and (H) TN$_{bulk}$ and $\delta^{15}$N$_{bulk}$.

**Fig. 4.** Proposed model for facies-dependent nitrogen cycling at the ocean margin. This depositional setting is a well-defined marine marginal ramp and platform-top that could have reduced communication with the open ocean and led to somewhat isolated shallow depositional settings. Platform/ramp bathymetry and stromatolite assemblages are based upon idealised Proterozoic ramp and platform settings by Walter et al. (1992). Shown here are: (A) a shallow ramp top setting, representing the Schmidsdrift Subgroup and Monteville Fm; and (B) a lagoonal depositional setting, representing the Campbellrand Subgroup overlying the Monteville Fm. The proposed mechanisms include: (1) transport of diazotrophic biomass to the seafloor; (2) remineralization of OM to NH$_4^+$ and shoreward transport; (3) NH$_4^+$ assimilation, producing OM with -ve $\delta^{15}$N values; (4) transport of a residual pool of NH$_4^+$ (possibly with +ve $\delta^{15}$N values); (5) open ocean ammonium assimilation, producing OM with +ve $\delta^{15}$N values; and, (6) potential for restricted open-marine influence but (7) potential for variability in communication as sea levels fluctuated.
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