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Ocean redox structure across the Late Neoproterozoic Oxygenation Event: A

nitrogen	isotope	perspective
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27 Abstract

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The end of the Neoproterozoic Era (1000 to 541 Ma) is widely believed to have seen the transition from a dominantly anoxic to an oxygenated deep ocean. This purported redox transition appears to be closely linked temporally with metazoan radiation and extraordinary perturbations to the global carbon cycle. However, the geochemical record of this transition is not straightforward, and individual data sets have been variably interpreted to indicate full oxygenation by the early Ediacaran Period (635 to 541 Ma) and deep ocean anoxia persevering as late as the early Cambrian. Because any change in marine redox structure would have profoundly impacted nitrogen nutrient cycling in the global ocean, the N isotope signature of sedimentary rocks ($\delta^{15}N_{sed}$) should reflect the Neoproterozoic deep-ocean redox transition. We present new N isotope data from Amazonia, northwest Canada, northeast Svalbard, and South China that span the Cryogenian glaciations (~750 to 580 Ma). These and previously published data reveal a N-isotope distribution that closely resembles modern marine sediments, with a mode in δ^{15} N close to 4‰ and range from -4 and +11‰. No apparent change is seen between the Cryogenian and Ediacarian. Data from earlier Proterozoic samples show a similar distribution, but shifted slightly towards more negative $\delta^{15}N$ values and with a wider range. The most parsimonious explanation for the similarity of these N-isotope distribution is that as in the modern ocean, nitrate (and hence O₂) was stable in most of the middle-late Neoproterozoic ocean, and possibly much of Proterozoic Eon. However, nitrate would likely have been depleted in partially restricted basins and oxygen minimum zones (OMZs), which may have been more widespread than in the modern ocean.

1. Introduction

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It is widely argued that the appearance of animals in the geological record is linked to a major step in the oxygenation of Earth's surface environment, known as the Neoproterozoic Oxygenation Event (NOE) (Canfield et al., 2007, 2008; Och and Shields-Zhou, 2012; Scott et al., 2008). The NOE is more difficult to document in the geological record than its early Paleoproterozoic counterpart, the Great Oxidation Event (GOE), whose onset is well established based on the sedimentary record of redox-sensitive detrital minerals and mass independent fractionation of sulfur isotopes (e.g. Farguhar et al., 2000; Lyons et al., 2014). Although the NOE has not been formally defined, one criterion is the timing of ventilation of the deep ocean with sufficient dissolved oxygen to accommodate the earliest animals (Och and Shields-Zhou, 2012). Whereas the cause-and-effect relationship between oxygenation and animal evolution and the oxygen threshold required for early animals are still debated (e.g., Butterfield, 2009; Sperling et al., 2013), eukaryotic diversification and the oxygenation of the environment must have been closely coupled through a complex network of biogeochemical feedbacks linked through the origin, export, decay and burial of organic matter (Butterfield, 2009; Close et al., 2011; Logan et al., 1995; Pawlowska et al., 2013). Iron speciation data from Neoproterozoic mudstones suggest a transition from dominantly anoxic and ferruginous deep oceans with prevalent sulfidic (euxinic) conditions in continental margin settings (Planavsky et al., 2011; Poulton and Canfield, 2011) to dominantly anoxic and ferruginous deep oceans with only occasional evidence for euxinic conditions in the earlymiddle Neoproterozoic (Canfield et al., 2008), followed by deep-ocean ventilation in the late Ediacaran (Canfield et al., 2007). However, these data only provide information about local depositional redox conditions and do not bear directly on the state of the deep ocean. Trace metal

abundances (Mo, U, V, and Cr) in euxinic sediments, although sensitive to local conditions, are also strongly dependent on the volume of oxygenated water masses, hence they bear on global redox conditions in the oceans (Algeo and Rowe, 2012). A series of separate studies and compilations of trace metal data have been published recently. All show a dramatic increase in abundances in the Ediacaran first interpreted to record deep ocean ventilation (635–541 Ma; Scott et al., 2008; Och and Shields-Zhou, 2012; Sahoo et al., 2012; Partin et al., 2013), broadly consistent with iron speciation data. Yet, data from some individual basins variably indicate persistently anoxic (Li et al., 2010; Johnston et al., 2013) versus oxygenated (Johnston et al., 2012; Sansjofre et al., 2014) bottom-waters throughout most of the Ediacaran Period. Other data imply widespread oxygen-deficient conditions in the earliest Cambrian (e.g. Schroder and Grotzinger, 2007; Cremonese et al., 2013) despite the appearance of complexly burrowing bilaterians. An emerging model is that anoxic conditions on continental platforms was locally controlled (Kah and Bartley, 2011; Shen et al., 2011), the deep ocean being pervasively oxygenated, but with a low oxygen concentration preventing the stabilization of oxic conditions in platforms throughout the Ediacaran Period (Johnston et al., 2012, 2013). Recent modeling coupled with trace metal compilations further suggests that no more than 40% of the pre-NOE seafloor could have been anoxic and no more than 10% euxinic (Partin et al., 2013; Reinhard et al., 2013). These models and existing data thus allow for the possibility that much of the global ocean may have been oxygenated prior to the Ediacaran Period.

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We have applied the nitrogen isotope system to the question of the NOE because the N-isotopic composition of marine sediments ($\delta^{15}N_{sed}$; where the standard delta notation and normalization to atmospheric N_2 is used; Mariotti et al., 1981) is strongly dependent on the nitrogen biogeochemical cycle in the water column, itself controlled by the ocean redox structure (Boyle

et al., 2013; Canfield et al., 2010; Fennel et al., 2005; Quan and Falkowski, 2009). Although the present day nitrogen cycle is highly complex and remains poorly understood in detail (e.g. Lam et al., 2011; Sigman et al., 2009a), the current understanding of its basic features and of its isotopic imprint in the sedimentary record (Galbraith et al., 2008; Sigman et al., 2009b) have already motivated the application of $\delta^{15}N_{sed}$ to elucidate nitrogen biogeochemical cycling in past oceans. Nitrogen isotopes have been applied to the Phanerozoic (e.g. Algeo et al., 2008; Higgins et al., 2012; Junium and Arthur, 2007; Quan et al., 2008) and Precambrian records (Beaumont and Robert, 1999; Busigny et al., 2013; Garvin et al., 2009; Godfrey and Falkowski, 2009; Godfrey et al., 2013; Kump et al., 2011; Papineau et al., 2009, 2013; Stücken, 2013; Thomazo et al., 2009, 2011) as a proxy for both ocean redox and nitrogen biogeochemistry. Here we add a new nitrogen isotope dataset from ~750 to 580 Ma strata from multiple cratons to a growing Neoproterozoic database of both bulk sedimentary ($\delta^{15}N_{sed}$) and kerogen ($\delta^{15}N_{ker}$) nitrogen isotope data (Cremonese et al., 2013; Kikumoto et al., 2014; Spangenberg et al., 2014) to test whether the NOE is recorded in nitrogen isotope signatures. These results show that the $\delta^{15}N_{sed}$ distributions for the Cryogenian and the Ediacaran resemble one another and that documented in the modern ocean.

1.1. Isotope Biogeochemistry of the Nitrogen Oceanic Cycle

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Nitrogen-isotopic compositions of nitrate, particulate organic matter and surface sediments have been extensively studied to characterize the nitrogen cycle in present oceans and its transcription into the sedimentary record (Fig. 1a, e.g. Robinson et al., 2012; Somes et al., 2010; Tesdal et al., 2013; Thunell et al., 2004). Below we summarize the basic processes and pathways and associated isotope fractionation (expressed using the enrichment factor notation in ‰, $\epsilon_{a-b} \approx \delta^{15} N_a - \delta^{15} N_b$) in the oceanic nitrogen cycle.

The initial source of all bioavailable-N to the oceans is the fixation of atmospheric N₂ (N₂fixation) by aerobic or anaerobic autotrophs, such as cyanobacteria, which transform molecular N_2 into organic matter (via NH_4^+) with a small isotope fractionation ($\varepsilon_{org-N2} \sim -4$ to 0%; e.g. Zerkle et al., 2008). Other primary photosynthetic organisms assimilate remineralized forms of nitrogen mostly as NO₃ or NH₄⁺. The mineralization of organic matter produces NH₄⁺ (ammonification) with no net isotope fractionation owing to its efficiency (Möbius, 2013). If NH₄⁺ remains stable it can then be assimilated with an isotope effect that increases strongly with its availability ($\epsilon_{org-NH4} \sim 0\%$ to -27%) such that the organic matter can be markedly ^{15}N depleted where NH₄⁺ is abundant (Pennock et al., 1996). The NH₄⁺ may also be oxidized. If this oxidation is not complete, the residual NH₄⁺ will be enriched in ¹⁵N, because the first oxidation step to nitrite (NO₂⁻) involves a strong fractionation ($\varepsilon_{NO2-NH4} \sim -41\%$ to -13%; Casciotti et al., 2003; Mariotti et al., 1981; Santoro and Casciotti, 2011). In oxic environments, this oxidation is accomplished by nitrification (i.e. sequential oxidation to NO₂ and NO₃) and goes to completion, unless ammonium is concurrently assimilated as a nutrient. Subsequent NO₃⁻ assimilation entails fractionation with $\epsilon_{org-NO3}$ between 0 and -8‰ in NO_3 -limited and NO_3 replete conditions, respectively (Pennock et al., 1996; Somes et al., 2010). Under dysoxic conditions (i.e. typically in OMZs or within redox transition zones), NO₃ as well as available NH₄⁺ and NO₂⁻ are converted into gaseous species (NO₂ and/or N₂) by an incompletely understood combination of metabolic pathways that include heterotrophic denitrification and anammox (anaerobic oxidation of NH₄⁺ by NO₂⁻), nitrification, nitrate reduction to NH₄⁺, chemolithotrophic sulfide-dependent denitrification, co-oxidation of methane and ammonium, and most probably other metabolisms yet to be indentified (Lam et al., 2009, 2011; Lavik et al., 2009; Mandernack et al., 2009; Wenk et al., 2013). In present day OMZs where nitrate

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consumption does not reach completion, the net result of these processes is a pronounced 15 N-enrichment in the residual NO_3^- pool (Voss et al., 2001) similar to that of the experimentally determined for heterotrophic denitrification ($\epsilon_{NO3-\ N2} \sim +15$ to +30%) (Granger et al., 2008; Mariotti et al., 1981). When NO_3^- consumption proceeds to completion, as in anoxic porewaters, hardly any isotope effect is expressed (Lehmann et al., 2007). Finally, within stable redox transition zones, nitrate (from above) and ammonium (from below) are also near-quantitatively converted into N_2 or N_2O resulting in a discrete zone of 15 N-enrichment in nitrate and ammonium ($\epsilon_{NO3-N2} \sim +10$ %; $\epsilon_{NH4-N2} \sim +10$ %) tied to decreasing concentrations (Fuchsman et al., 2008; Thunell et al., 2004; Wenk et al., 2014).

The ocean's redox structure thus tightly controls the nitrogen cycle through the complex network of metabolic pathways by which nitrogen flows between reservoirs. The local redox conditions determine the dissolved inorganic nitrogen speciation, nitrate being stable in oxic waters and ammonium in anoxic waters. The nature (i.e. OMZs versus redox transition zones), location (shallow versus deep), and spatial extent of dysoxic zones determine the rate of nitrate and ammonium conversion to N_2 or N_2O , as well as their ^{15}N enrichment.

1.2. Expected $\delta^{15}N_{sed}$ signature under different ocean redox structures

On continental margins and in anoxic basins, $\delta^{15}N_{sed}$ mostly records the isotopic composition of primary producers, with only minor modification of (e.g. Robinson et al., 2012; Tesdal et al., 2013; Thunell et al., 2004). The $\delta^{15}N$ of the primary producers depends on the mass and isotope balance between the two main sources of nitrogen supporting the new primary productivity, i.e. N_2 -fixation and upwelled nitrate and/or ammonium reaching the photic zone (e.g. Somes et al., 2010). Because both this balance and the nitrate isotope composition are ultimately linked to the

redox structure of the global ocean, $\delta^{15}N_{sed}$ should reflect this redox structure, even though its isotopic signature is acquired in surface waters.

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We review conceptual scenarios for the nitrogen cycle and $\delta^{15}N_{sed}$ expression based on two ocean redox structures envisaged for the Neoproterozoic: a dominantly oxygenated ocean and a redox-stratified ocean. In a dominantly oxygenated ocean (Fig. 1a), nitrate is the main form of bioavailable nitrogen and it is generated by nitrification of the ammonium released by remineralized organic matter. In the surface mixed layer, which broadly corresponds to the euphotic zone, ammonium and nitrate are usually quantitatively reassimilated to support primary productivity, without net isotope fractionation. Deeper, regenerated nitrate accumulates and eventually returns to the surface ocean in upwelling zones. When the upwelling passes through dysoxic or anoxic OMZs, a fraction of this nitrate pool is denitrified, increasing the $\delta^{15}N$ of residual nitrate ultimately delivered to the surface. The nitrogen isotope composition of nitrate $(\delta^{15}N_{NO3})$ in surface waters can thus strongly vary regionally (between +1 and +15%) depending on ocean circulation patterns and on deep and intermediate water redox condition (Somes et al., 2010). The $\delta^{15}N_{sed}$ distribution of present-day ocean surface sediments captures this range of variation, with a mode at 5-6% (Fig. 2a; Tesdal et al., 2013), a small negative tail approaching 1‰ and a large positive tail to +15‰ corresponding to OMZs.

In a redox-stratified ocean (Fig 1b and c), the inorganic nitrogen generated by organic matter mineralization will be in the form of nitrate above the redox transition zone and ammonium below. The redox transition is unlikely to be stable in the surface mixed layer, owing to its rapid mixing time. Thus in most locations the redox transition zone should stabilize below the surface mixed layer, except possibly in up-welling zone where it may reach the surface (Kump et al., 2005)(Fig. 1c). In a stable situation, nitrate is depleted by assimilation in the surface mixed layer

and accumulates below until it reaches the redox transition zone, where it is denitrified. Below the redox transition zone, ammonium accumulates. In this configuration, ammonium does not reach the surface. It is quantitatively converted to N₂ or N₂O within the redox transition zone by coupled nitrification-heterotrophic denitrification and anammox, as observed today in the Cariaco basin and the Black sea (Fuchsman et al., 2008; Meckler et al., 2007; Thunell et al., 2004). Overall this scenario results in extensive loss of bioavailable nitrogen (nitrate and ammonium) compared to the oxic ocean scenario. Only nitrate accumulated in between the surface mixed layer and the redox transition zone may ultimately remix into the surface ocean in upwelling zones (Fig. 1b). But this accumulation is strongly limited by down-welling zones where denitrification will occur as the nitrate crosses the redox transition zone. The amount of nitrate available for recharging the surface ocean is thus strongly controlled by the depth of the redox transition zone. The closer the redox transition zone is to the base of the surface mixed layer, the smaller the nitrate reservoir is. Resulting nitrate limitation is compensated by N₂. fixation, which drives surface nitrate δ^{15} N towards 0% (Quan and Falkowski, 2009), as observed in Mediterranean sapropels and sedimentary rocks deposited during ocean anoxic events (OAEs; Fig. 2b and references therein). Locally, upwelling currents may be vigorous enough for some of the ammonium to traverse the redox transition zone before being fully converted to N₂ or N₂O. In this case, the ¹⁵N-enriched residual ammonium will be assimilated either directly or after having been oxidized to nitrate, producing positive $\delta^{15}N_{\text{sed}}$ signatures similar to those found in present-day OMZs (Fig. 1b). This process has recently been proposed to explain positive $\delta^{15}N$ values on a late Paleoproterozoic platform (Godfrey et al., 2013). If the redox transition zone was shallow enough (Fig. 1c), it may impinge on the surface mixed layer in upwelling zones possibly reaching the surface (Kump et al., 2005). In this case, ammonium would be competitively

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assimilated by primary producers and converted to N_2 or N_2O by coupled nitrification–heterotrophic denitrification and anammox. Strong upwelling would favor ammonium assimilation and negative $\delta^{15}N_{sed}$, as has been proposed to explain values as low as -4‰ during OAE2 (Higgins et al., 2012), but where conversion to N_2 or N_2O dominates, $\delta^{15}N_{sed}$ values would trend positive (e.g., Papineau et al., 2009). Given the instability of the redox transition zone in the surface mixed layer, $\delta^{15}N_{sed}$ values should be highly variable both temporally and spatially, as observed in the c.a. 2.0 Ga Lower Aravalli Group (Papineau et al., 2009)(Fig. 2g). The $\delta^{15}N_{sed}$ distribution of a stratified redox scenario should reflect it with high variability, ranging from negative to positive values, and with a mode close to 0‰.

This brief overview highlights the difficulty of predicting $\delta^{15}N_{sed}$ signatures under non-actualistic scenarios. It also demonstrates the difficulty in extrapolating results from a single basin to interpreting the global nitrogen cycle given the intrinsic complexity and heterogeneity of the nitrogen cycle. However, it raises the possibility that distributions of $\delta^{15}N_{sed}$ from several globally distributed locations hold key information on the global nitrogen cycle. Specifically, the $\delta^{15}N_{sed}$ distribution in redox-stratified scenarios are expected to show a mode closer to 0% (Quan and Falkowski, 2009) with marked tails toward negative and positive $\delta^{15}N_{sed}$ values in upwelling regions. For this reason we have chosen to integrate data from multiple basins to produce a global picture of $\delta^{15}N_{sed}$ spanning the purported NOE, i.e. from 750 to 580 Ma.

2. Samples

We present new $\delta^{15}N_{sed}$ data from four late Neoproterozoic basins that straddled the equator during the Marinoan glaciation (Fig. 3): (i) the Datangpo and Doushantuo formations sampled from the Yangjiaping section (~720-580 Ma), South China; (ii) the Mirasol d'Oeste and Guia

formations sampled from 4 sections spanning a cross-section of the Araras platform, Brazil; (iii) the Elbobreen and Dracoisen formations sampled from a composite section (~750–600 Ma) in northeast Svalbard; (iv) and the Twitya and Sheepbed formations from a composite section (~660–600 Ma) in northwest Canada. Most samples are Ediacaran (635–541 Ma) in age, such that this data set spans the first fossil evidence of bilaterians (~575 Ma; Narbonne and Gehling, 2003) and record what is commonly regarded as a transition from dominantly anoxic to oxygenated oceans (Fike et al., 2006; Sahoo et al., 2012). A small subset of older samples extends the record beyond the Cryogenian (c.a. 720–635 Ma) glaciations. A description of geological backgrounds for the sampled sections is included in the supplementary material. We only summarize below the depositional depth and redox conditions inferred for each of these sections.

In the Yangtze platform (South China) sections, iron speciation, trace metal contents, sulfur and carbon isotopes, and pyrite morphologies indicate that the water column was redox stratified during the deposition of both the Datangpo and Doushantuo formations (Ader et al., 2009; McFadden et al., 2007; Li et al., 2010; Li et al., 2012; Wang et al., 2012). However, enrichment of Mo in Doushantuo shales indicates that a significant percentage of the open ocean must have been oxygenated at the time (Sahoo et al., 2012). At Yangjiaping, both formations were probably deposited in oxic shallow waters (Ader et al., 2009; Li et al., 2012; Wang et al., 2012), while at the Wuhe section, for which previous $\delta^{15}N_{sed}$ results are available (Kikumoto et al., 2014), the Doushantuo Formation was deposited in a deep anoxic inner-shelf environment.

On the Araras platform, the four studied sections span a platform transect (Fig. S1) in an overall deepening upward succession: the lower Mirassol d'Oeste Formation was deposited in shallow water and overlying Guia Formation was deposited in increasingly deeper environments, with

the upper part of the unit deposited in a slope setting. The trace metal contents in the same samples for which $\delta^{15}N_{sed}$ have been obtained indicate a fully oxygenated water column, with possible episodes of porewater anoxia approaching the sediment-water interface on the slope (Sansjofre et al., 2014).

Shales from Syalbard and Canada were all deposited below storm wave base. For the Elbobreen

and Dracoisen formations (Northeast Svalbard) Fe/Al data lie close to the local detrital baseline of 0.58 with a few values above but most lying below (Table S1 in supplementary material). These results are not conclusive, but are consistent with liberation of iron from ferrouginous OMZs (Fe/Al < 0.58) and subsequent addition to more oxygenated parts of the water column, as occurs in modern OMZs (Scholz et al., 2014). We have no direct constraints on the depositional redox conditions of our samples from the Twitya and Sheepbed formations (Northwest Canada). However, iron speciation data are available for correlative sections of the lower Sheepbed Formation (Johnston et al., 2013; Shen et al., 2008). These data were interpreted to indicate anoxic and ferruginous deep waters with variations in the redox transition zone depth, such that the depositional environment may have been oxic in some locations. We note, however, that FeHR/FeT data lie close to the 0.38 threshold used to discriminate anoxic from oxic environments and the FeT/Al ratio are close to both the generic detrital ratio of 0.53 (Johnston et al., 2013), which means the data do not conclusively demonstrate anoxic conditions. As for Svalbard, it is possible that ferruginous conditions were confined to an OMZ in an otherwise oxygenated ocean.

3. Nitrogen stable isotope analyses

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Samples were ground in an agate mortar and sieved to ensure a grain size <140 μm . To concentrate nitrogen in the insoluble residue, most carbonate-rich samples were first decarbonated in HCl 6 N overnight at room temperature, followed by 2 h at 80°C. Residues were washed with distilled water, centrifuged and dried at 50°C. When enough material was available and N-content was above 200 ppm, samples were prepared by conventional sealed tube combustion to convert total nitrogen to N_2 (Dumas combustion). This method is hereafter referred to as the conventional method (see Ader et al. (1998) for a detailed description). The N_2 was then measured on a dual inlet Thermo Finnigan Delta+XP mass spectrometer allowing analyses on as low as 2 micromoles N_2 with a precision of \pm 0.2% (1 σ). Nitrogen blanks were approximately 0.15 micromole, thus representing less than 10% of the measured nitrogen on the least N-rich samples. The reproducibility of the $\delta^{15}N_{sed}$ measurements was \pm 0.2% (1 σ).

For samples containing a limited amount of material, N-isotope analyses were performed using sealed-tube combustion and on-line N_2 purification on an ultra-high vacuum line with direct introduction to a static mass spectrometer (hereafter referred to as the static method) following an established protocol (Ader et al., 2006; Busigny et al., 2003, 2005; Thomazo et al., 2011). Powdered samples (between 5 and 30 mg) were embedded in a platinum parcel, then evacuated under vacuum at 150°C before being sealed in an evacuated and pre-purified quartz tube with an excess of CuO and Cu wire and CaO granules. The sample was combusted at 950 °C for 6 h and the resulting N_2 purified and quantified on an ultra-high vacuum line directly coupled to our static mass spectrometer. Precision for $\delta^{15}N$ via this method is \pm 0.3% (1 σ) based on repeat analyses. Blanks yielded $\delta^{15}N$ values of \pm 1.5 \pm 3% and higher nitrogen amounts (\pm 30 ng) than expected based on prior work in our lab (Ader et al., 2006; Busigny et al., 2005; Thomazo et al., 2011), which is likely a result of the lower sample degassing temperature set at 150°C to avoid

nitrogen devolatilisation, the thermal maturation experienced by our samples being lower than greenschist facies. In most cases this blank content accounts for less than 3% of the total nitrogen, but reached 10% in some of the most nitrogen-poor samples. All results are blank-corrected and given in standard δ notation, calibrated to Air (Mariotti et al., 1981).

In order to ascertain whether bulk and decarbonated samples, as well as conventional and static methods, yield comparable results, selected samples were analyzed by multiple methods (Table S2). $\delta^{15}N_{sed}$ results obtained on bulk and decarbonated samples using the conventional or the static method compare well given the reproducibility of each method. When all data are considered together, the $\delta^{15}N_{sed}$ reproducibility (1 σ) is usually better than $\pm 0.35\%$, and always better than $\pm 0.63\%$. N-contents also compare quite well between methods, except for pure carbonate samples, for which bulk N-contents measured on the decarbonated residue are often lower than those measured directly on bulk samples. It is likely that a significant fraction of the very small residue was lost during the decarbonation step, probably as a supernatant during rinsing and centrifugation. Therefore, N-contents determined using the decarbonated residue of carbonate samples are probably slightly underestimated.

4. Results

Our $\delta^{15}N_{sed}$ results are listed in the supplementary information Table S3 and presented as chemostratigraphic profiles (together with available $\delta^{13}C_{org}$ and $\delta^{13}C_{carb}$ data) for each of the studied sections in Figure 4. For the Araras platform (n=41), $\delta^{15}N_{sed}$ data have been obtained from 4 sections spanning the platform (Fig. 4a-c). $\delta^{15}N_{sed}$ data for the shallow-water cap dolostone of the Mirassol d'Oeste Formation (directly overlying the Marinoan glacial deposits) are only available in the Terconi-Camil composite section. They show the largest distribution

with $\delta^{15}N_{sed}$ values ranging from +2 to +10‰ (Fig. 4a). $\delta^{15}N_{sed}$ data for the overlying Guia Formation are available for all three sections (Fig. 4a-c). They lie between +2 and +6‰, and no systematic variation is apparent either temporally or between sections.

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For South China, all samples come from the Yangikiaping section (Fig. 4d). In the Datangpo Formation, $\delta^{15}N_{sed}$ data cluster around +4%. In the Doushantuo Formation, $\delta^{15}N_{sed}$ values increase within the 4 meter-thick cap carbonate from +3% up to +5.5%, at which they remain through the subsequent 40 m. Above, $\delta^{15}N_{sed}$ values are more variable, but overall decrease to +2.5% in the lower Dengying Formation. Recently published $\delta^{15}N_{sed}$ data from the Doushantuo Formation from a drill-core in the Three Gorges area that penetrates a deep inner-shelf setting on the Yangtze platform (Kikumoto et al., 2014) show similar $\delta^{15}N_{sed}$ values than the Yangjiaping section, which correspond to the shelf margin (Fig. 2c). Both sections also display similar $\delta^{15}N_{sed}$ stratigraphic variations (Fig. 4d, this study and Fig. 2, Kikumoto et al., 2014) in spite of the different depositional setting, lithologies and $\delta^{13}C_{org}$ and $\delta^{13}C_{carb}$ stratigraphic variations. Samples from Svalbard (n=18) present very homogeneous $\delta^{15}N_{sed}$ values ranging from +4.1 to +6.3% (Fig. 4e), although data from the post-Marinoan Dracoisen Formation show a distinct positive upward trend from +4 to +5.5% (Fig. 4e). All samples from northwest Canada (n=8) fall between +2 and 4% (Fig. 4f). No obvious correlation with $\delta^{13}C_{org}$ and $\delta^{13}C_{carb}$ is evident in any of the sections (Fig. 4 and Fig. 5d).

These results are synthesized in histograms in Figure 2 together with $\delta^{15}N_{sed}$ data previously published for the Doushantuo and Dengying formations, South China (Cremonese et al., 2013;) and $\delta^{15}N_{kerogen}$ data from the Tamengo Formation, Brazil (Kikumoto et al., 2014; Spangenberg et al., 2014). $\delta^{15}N_{sed}$ values for South China range between +2.2 and +8.5% (Fig. 2c); for the Amazonian craton between +2.2 and +10.5%; for the Late Ediacaran Tamengo Formation

between -3.3 and +3.1‰, defining the lower end of the distribution (Fig. 2d); for northwestern Canada between +2 and +4‰ and for Svalbard between +4 and +7‰ (Fig. 2e). Overall, the $\delta^{15}N$ values (n = 158) range from -3.3 to +10.5‰, with roughly 40% of the data falling between +3 and +6‰ (Fig. 2f). No time-based evolution is evident from the comparison of the Cryogenian values to those of the Ediacaran (Fig. 2f).

5. Discussion

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5.1. Preservation of the $\delta^{15}N_{sed}$ signature

In sediments from continental margins and anoxic depositional settings, $\delta^{15}N$ of primary producers is recorded with only minor modification in $\delta^{15}N_{sed}$ in spite of the early diagenetic remineralization of organic matter and ammonium fixation in clay minerals (e.g. reviews in Robinson et al., 2012; Thomazo et al., 2011). The conservation of primary signatures is illustrated in sediments from the eastern Mediterranean sapropels and the OAE2 event in the proto-North Atlantic by the similarity between $\delta^{15}N_{\text{sed}}$, $\delta^{15}N_{\text{kerogen}}$ and $\delta^{15}N$ of primary producers as reconstructed from porphyrin $\delta^{15}N$ (Higgins et al., 2010; 2012). However, most Precambrian sedimentary rocks have experienced some degree of post-depositional $\delta^{15}N_{sed}$ modification. Devolatilization of fixed ammonium during metamorphism has been identified as the main culprit. At metamorphic grades higher than greenschist facies, a decrease in N content is often coupled to an increase in $\delta^{15}N_{sed}$ as ^{15}N -depleted fixed ammonium is preferentially volatilized (Bebout and Fogel, 1992; Busigny et al., 2003; Jia and Kerrich 2004; Jia, 2006; Mingram and Brauer 2001; Pinti et al., 2009; Yui et al., 2009). The reliability of $\delta^{15}N_{sed}$ in Precambrian sedimentary rocks as a tracer for initial $\delta^{15}N$ of primary producers thus remains controversial. This problem has been addressed by comparing $\delta^{15}N_{sed}$ and $\delta^{15}N_{kerogen}$ data from the same

successions (Godfrey et al., 2009, 2013; Kump et al., 2011), on the basis that $\delta^{15}N_{kerogen}$ should not be modified by metamorphism (Ader et al., 1998, 2006; Boudou et al., 2008). So far, these comparisons have shown mixed results: two successions show $\delta^{15}N_{sed}$ to be higher than $\delta^{15}N_{kerogen}$ by up to +6% (Kump et al., 2011) and +12% (Godfrey et al., 2013), whereas others successions yield comparable $\delta^{15}N_{sed}$ and $\delta^{15}N_{kerogen}$ values (Godfrey et al., 2009, 2013). The two successions with higher $\delta^{15}N_{sed}$ than $\delta^{15}N_{kerogen}$ are recognized as having undergone lower greenschist facies (Kump et al., 2011) or higher (Godfrey et al., 2013) peak metamorphic conditions. Peak metamorphic temperatures for the other successions are not reported. These mixed results raise the specter of overprinting of $\delta^{15}N_{sed}$ by devolatilization and/or of overprinting of $\delta^{15}N_{kerogen}$ by N-contamination during the HF extraction of N-poor kerogen (Ader et al., 2006) in rocks metamorphosed to greenschist or higher facies. They nevertheless suggest that $\delta^{15}N_{sed}$ may be preserved in rocks that have experienced minimal metamorphism.

In any case, for this study the maximum metamorphic conditions for the studied locations were below greenschist facies (Supplementary material), minimizing the risk of post-depositional modification of the $\delta^{15}N_{sed}$ signal. N/C ratios strongly vary (Fig. 5b), but the N and TOC contents of decarbonated samples show a rough correlation (Fig. 5c), suggesting N originates from the organic carbon matter. The fact that $\delta^{15}N_{sed}$ does not correlate with either N_{decarb} content (Fig. 5a) or C/N ratio (Fig. 5b) suggests that even if some nitrogen was redistributed between organic matter, clay minerals and fluid phases during diagenesis and low-grade metamorphism, bulk $\delta^{15}N_{sed}$ values should not have been significantly modified. Finally, it would be very unlikely for burial and metamorphic processes to have resulted in the observed similarities in $\delta^{15}N_{sed}$ values across all studied locations, as well as across transects of the Araras and Yangtze platforms, in

spite of the variable lithologies and thermal histories. We thus proceed on the assumption that $\delta^{15}N_{sed}$ values reflect primary biomass signatures.

5.2 Significance of $\delta^{15}N_{sed}$ in the context of global ocean redox structure

The distribution of $\delta^{15}N_{sed}$ in analyzed Neoproterozoic rocks (mode of ~ +4‰, Fig. 2c) is clearly more like that of modern sediments (mode of ~ +5‰, Fig. 2a) than to that of OAEs (mode of ~ -1‰, Fig. 2b). The fact that most $\delta^{15}N_{sed}$ values do not cluster in the -4 to + 2‰ range typical of OAEs, which have been consistently interpreted as representing episodes of redox stratification, indicates that during most of the studied time interval, the nitrogen cycle and ocean redox structure differed from that of a redox-stratified ocean. The Neoproterozoic $\delta^{15}N_{sed}$ distribution with a mode close to +4‰ would be compatible with redox-stratified ocean only if all studied sections (except that of the Rio de la Plata craton with $\delta^{15}N_{ker}$ close to 0‰) were located in the vicinity of upwelling zones, which would be an extraordinary coincidence. Consequently, a globally redox-stratified ocean is an unlikely scenario to explain the Cryogenian-Ediacaran $\delta^{15}N_{sed}$ distribution.

In contrast, the $\delta^{15}N$ distribution for the ~750 to 580 Ma period, while being markedly different from that of OAEs, is strikingly similar to that of modern sediments (Fig. 2a). The most parsimonious hypothesis is thus that it reflects a nitrogen cycle similar to the one operating in the modern ocean, where nitrate was stable throughout the ocean, except in anoxic OMZs and locally restricted anoxic basins. It implies that most of the late Neoproterozoic oceans contained enough dissolved O_2 to stabilize nitrate, with the probable exception of transient OAEs. This interpretation is compatible with the existing knowledge of the redox depositional conditions for the studied samples. Samples from the Araras platform (Amazonian craton) show oxic

sections are compatible with deposition under oxygenated conditions or within anoxic OMZs in an otherwise oxygenated ocean. The Datangpo and Doushantuo formations (South China) were both deposited from a redox-stratified water column (Li et al. 2010; Li et a., 2012), yet their δ¹⁵N_{sed} values in both shelf margin and inner-shelf basin settings are positive and lie in the same range as the other locations. These patterns are fully compatible with nitrate having being supplied to these areas through surface currents in a scenario analogous to the present-day anoxic Cariaco Basin, where $\delta^{15}N_{sed}$ values of ~+3.5% are maintained (Thunell et al., 2004). The $\delta^{15}N_{sed}$ distribution compiled from these locations is thus highly likely to record the $\delta^{15}N$ distribution of the global ocean nitrate inventory, albeit with minor regional influences. As to the negative tail of the distribution defined by samples from the Tamengo Formation deposited in an anoxic basin (Spangenberg et al., 2014), it records the regional signal of N₂-fixation and/or ammonium assimilation expected in the case of redox stratification (Junium and Arthur 2007; Higgins et al., 2012). In detail, the δ^{15} N distribution for the ~750 to 580 Ma period shows minor differences with the modern distribution. While these differences may simply reflect sampling bias, they may also speak to important differences in the global ocean redox structure, which we explore here. Compared to the modern $\delta^{15}N$ distribution, that of the ~750 to 580 Ma is slightly shifted towards lower $\delta^{15}N_{sed}$ values, its positive tail is less prominent, and its negative tail extends to lower values (as low as -3.3%; Figs. 2, 7). Its mode is also 1% lower than that of modern sediments. This shift to lower $\delta^{15}N_{\text{sed}}$ values is compatible with the hypothesis of an oxygenated ocean, but one containing less dissolved oxygen than the modern ocean. Restricted anoxic basins would be more common, increasing the proportion of $\delta^{15}N_{sed}$ values close to 0‰. Similarly, OMZs would

depositional conditions (Sansjofre et al., 2014), while available data for Svalbard and Canada

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be more widespread and severe, possibly reaching euxinia, which would have resulted in more common quantitative denitrification, hence driving N_2 -fixation. O_2 penetration in sediments would also have been less compared to the modern ocean, increasing the surface of sediments experiencing conditions favorable to quantitative sedimentary denitrification. Combined, these effects would have decreased the nitrate inventory of the ocean without significantly increasing residual nitrate $\delta^{15}N$ values, driving N_2 -fixation to compensate for the nitrate loss and hence decreasing $\delta^{15}N$ of the global nitrate inventory. Importantly, the absence of marked change in $\delta^{15}N$ signatures during the latter half of the Neoproterozoic (Fig. 4 and Fig. 2f) implies no fundamental and irreversible change in the nitrogen cycle during this interval. These data imply no first order change in oceans redox structure across the NOE and by extension, suggest that pervasive oxygenation of the oceans occurred before 750 Ma.

Few $\delta^{15}N$ data are available for the early Neoproterozoic and late Mesoproterozoic, but significant data have recently been published for the early Mesoproterozoic and Paleoproterozoic (~2.1 to 1.4 Ma; Busigny et al., 2013; Kump et al., 2011; Papineau et al., 2009, 2013; Stüeken, 2013). These data are synthetized in Figure 2g. The mode of the 2.1–1.4 Ga $\delta^{15}N$ distribution is closer to +2‰ compared with +4‰ for the ~750 to 580 Ma period, but is clearly distinct from the -1‰ mode characteristic of the Phanerozoic OAEs. The offset to lower $\delta^{15}N$ values can be interpreted along the same lines of reasoning as above—increased area of restricted anoxic basins and euxinic OMZs, along with less pervasive O_2 penetration in sediments overlain by an oxygenated water column. Redox-stratified restricted basins and OMZs on continental platforms were likely more numerous than during the Neoproterozoic. The extremely large range of $\delta^{15}N$ (from -4 up to +24‰) identified in the Arivilli group (India) may indeed correspond to a restricted anoxic basin or an euxinic OMZ, where the positive $\delta^{15}N$ reflects impingement of the

redox transition zone on the surface mixed layer (Papineau et al., 2009, 2013). This interpretation implies a similar N-cycle than in the modern ocean with nitrate being stable in most of the ocean. It is at odds with most previous interpretations of Proterozoic N-cycling which are predicated on the assumption of a redox-stratified ocean. However, recent interpretations of trace metal compilations suggest that a minimum of 60% of the seafloor was overlain by oxygenated waters in the Proterozoic (Partin et al., 2013; Reinhard et al., 2013). This significant revision allows for the possibility that the $\delta^{15}N$ distribution for the 2.1–1.4 Ga period may reflect a modern style N-cycle in pervasively oxygenated oceans.

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It is difficult to place a quantitative constraint on the dissolved O₂ concentration of Proterozoic oceans given our poor knowledge of the O2 abundance required to stabilize nitrate. Measurements performed using the recently developed highly sensitive STOX oxygen sensors show that the accumulation of NO₂, the intermediate compound associated with N-loss in OMZs, is restricted to O2 concentrations lower than 50 nM (Thamdrup et al., 2012). A conservative lower estimate for O₂ concentration in a significant part of the oceans between 740 and 580 Ma can thus be fixed at 50 nM. However, this is probably underestimated by at least two orders of magnitude, considering that the activity of metabolic pathways controlling N-losses in OMZs have been detected at O₂ concentrations as high as 20 μM (Kalvelage et al., 2011), which prevent NO₂ accumulation. In any case, the widespread stability of nitrate implies that the Cryogenian and Ediacaran oceans were neither dominantly ferruginous nor sulfidic, because nitrate would have been reduced by ferrous iron or sulfide microbial oxidation, respectively (Canfield et al., 2010; Lavik et al., 2009; Pantke et al., 2012; Weber et al., 2006). Apart from possible episodic OAEs, anoxic waters (ferruginous or sulfidic) must therefore have been restricted to intracratonic and silled basins, zones of widespread upwelling on open continental

shelves, and probably to a thin layer of bottom water overlying the sediment-water interface (Fig. 1d).

5.3. Reconciling Redox Proxies

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A host of redox proxy datasets have been published recently for the Neoproterozoic and early Cambrian with seemingly contradictory implications. Compilations of trace metal data from euxinic black shales through time broadly indicate a major oxygenation event sometime in the late Neoproterozoic: the NOE (Fig. 6; Partin et al., 2013; Och and Shields-Zhou, 2012; Scott et al., 2008). Canfield et al. (2007) argued more specifically that deep ocean oxygenation occurred in the middle Ediacaran Period based on a compilation of iron speciation data from the Ediacaran succession of southeast Newfoundland. On the other hand, high-resolution trace metal abundance datasets from the organic-rich Doushantuo Formation in from South China show a prominent spike in Mo, V, and U in the early Ediacaran Period, which imply ocean oxygenation occurred shortly after the Marinoan (end-Cryogenian) glaciation (Sahoo et al., 2012). Other data sets from individual basins variably imply oxic, anoxic, and euxinic continental shelves during the Ediacaran (Johnston et al., 2012, 2013; Li et al., 2010; Sansjofre et al., 2014), while studies of the Precambrian-Cambrian boundary interval indicate ocean anoxia in the late Ediacaran early Cambrian (Kimura et al., 1997; Schröder and Grotzinger, 2007). Do these seemingly contradictory ocean redox datasets imply that the samples or proxies are unreliable or can these data be reconciled with a single model for Neoproterozoic oxygenation? We argue for the latter based on the $\delta^{15}N_{sed}$ record. The most parsimonious explanation for the

We argue for the latter based on the $\delta^{15}N_{sed}$ record. The most parsimonious explanation for the similarity of the $\delta^{15}N_{sed}$ distribution for the 750–580 Ma period to that of modern sediments is that nitrate was largely stable in the Neoproterozoic ocean by ~750 Ma, and hence that the

oceans were at least partially oxygenated by this time, albeit likely with low total dissolved oxygen concentrations. If correct this interpretation implies that deep-ocean ventilation occurred prior to the NOE. However, lower ocean O₂ concentrations than in the modern ocean may have allowed for persistent and broad OMZs that produced widespread anoxic conditions on continental shelves, which may even have reached the wind-mixed layer in some occasions as suggested by one reported occurrence of photic zone anoxia biomarkers (Olcott et al., 2005). The extent of anoxia and the occurrence of euxinia, in turn, were dependent on carbon export from the surface ocean, and hence sensitive to local nutrient variability, allowing for variable redox conditions within single basins and between basins. Oxic conditions likely prevailed not only in the surface ocean but also in the deeper ocean and away from highly productive continental margins. However, the sedimentary record of deep Neoproterozoic ocean basins is largely absent from the geological record, accounting for a strong bias in available local redox proxy data towards water column anoxia.

Our interpretation is compatible with the latest reinterpretation of trace metal data, recognized proxies for the global ocean redox (Algeo and Rowe, 2011). Reinhard et al. (2013) used a simple mass balance model to show that the low trace metal abundances in black shales prior to 630 Ma allow for a minimum of 60% of the seafloor to have been oxygenated throughout the Proterozoic. Given that the degree to which anoxia and/or euxinia extended into the water column from the seafloor remains completely unconstrained, anoxia may even have been restricted to microbial matts covering the sediment-water interface (Pawlowska et al., 2013) or to a thin layer of bottom water, leaving most of the water column oxygenated enough to allow nitrate accumulation (Fig. 1d). Therefore the unambiguous shift in trace metal abundance at 630 Ma (Och and Shields-Zhou, 2012; Partin et al., 2013; Reinhard et al., 2013; Sahoo et al., 2012),

rather than heralding deep ocean oxygenation, may instead only record an increase in atmospheric O_2 , and hence dissolved O_2 , beyond a critical threshold, thereby increasing the penetration depth of O_2 within seafloor sediments and hence total surface area of oxygenated seafloor on continental shelves. This may have been the threshold that permitted the earliest animals to evolve or develop more energy intensive metabolisms, such as burrowing and predation (Sperling et al., 2013).

6. Conclusions

The $\delta^{15}N_{sed}$ datasets presented here (Fig. 2, 4 and 6) suggests that the global nitrogen cycle operated similarly in the second half of the Neoproterozoic Era to in the modern ocean. If true, then it implies that nitrate was stable in the latter Neoproterozoic ocean, which, by extension, was at least mildly oxygenated. This conclusion is compatible with recent modeling of trace element data that suggest that most of the global Proterozoic ocean was sufficiently oxygenated to also accumulate trace metals (Partin et al., 2013; Reinhard et al., 2013). Although this hypothesis does not discount a possible rise in atmospheric and oceanic oxygen levels during this period or significant fluctuations associated with snowball glaciations, it does suggest that the major transition from a dominantly anoxic to oxygenated deep-ocean did not happen between \sim 750 and 580 Ma but rather sometime prior to 750 Ma, and perhaps much earlier in the Proterozoic.

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Figure legends

Figure 1: Schematic representation of the nitrogen cycle and resulting nitrogen-isotopic imprint on marine sediments (a) in a modern-like oxic ocean in which oxygen penetrates into the sediment except in OMZs where denitrification does not reach completion, (b) in a redox-stratified ocean with a relatively deep redox transition zone, where nitrate and ammonium are quantitatively converted to N_2/N_2O at the redox transition zone, except at vigorous upwelling locations where ammonium is nitrified concurrently with conversion to gaseous species, (c) in a redox stratified ocean with a shallow redox transition zone stabilizing below the surface mixed layer, except in upwelling zones where anoxic waters may reach the surface, allowing ammonium to be assimilated at the same time as being converted to N_2 and N_2O , c) in our model for the 750 to 580 Ma ocean with extended and severely OMZz and limited oxygen penetration in surface sediments. The red color indicates anoxic waters and the blue color oxygenated waters. $\varepsilon_{ap} \approx 0$ indicates cases where the reaction is complete so that the apparent isotope fractionation (ε_{ap}) is close to nil.

Figure 2: Comparison of $\delta^{15}N_{sed}$ distribution in modern oceanic sediments, in Phanerozoic anoxic events and in Neoproterozoic sediments. (a) Modern sediment $\delta^{15}N_{sed}$, modified after Tesdal et al. (2013). (b) $\delta^{15}N_{sed}$ from anoxic events in the Phanerozoic: (i) in black, the Cretaceous ocean anoxic events OAEs 1a, 1b, 1d and 2 (Arnaboldi and Meyers, 2006; Dumitrescu and Brassel, 2006; Jenkyns et al., 2007; Junium and Arthur, 2007; Kuypers et al., 2002, 2004; Meyers 2006); (ii) in dark grey, early Toarcian OAE (T-OAE) (Jenkyns et al., 2001); (iii) in light grey, the Quaternary and Cenozoic Mediterranean sapropels (Higgins et al., 2010; Meyers and Bernasconi, 2005; Sachs et al., 1999); (iv) in white, other Phanerozoic anoxic events, of unknown spatial extent, which correspond to the Triassic-Jurassic transition (Quan et al., 2008), the Permian-Triassic transition (Cao et al., 2009; Luo et al., 2011), the late Devonian (Calvert et al., 1996; Levman and Bitter, 2002; Sliwinski et al., 2011), the late Ordovician (LaPorte et al., 2009) and the early Cambrian (Cremonese et al., 2013). (c) Histogram of the δ¹⁵N_{sed} data for the Yangtze platform (South China craton): in grey, data from the Yangjiaping section representative of the shelf margin (this study), in white, previously published data from the top of the Dengying Formation in the Xiaotan section (Cremonese et al., 2013) and the Doushantuo and Dengying formations in the Wuhe section, representative of the inner shelf basin (Kikumoto et al., 2014). (d) $\delta^{15}N_{sed}$ distributions for Brazil: in black, data from the Amazon craton (this study); in white, $\delta^{15}N_{kerogen}$ values reported for the Tamengo Formation, Rio de la Plata craton (Spangenberg et al., 2014). (e) $\delta^{15}N_{sed}$ values for northwestern Canada (white) and Svalbard (grey). (f) Histogram of all the $\delta^{15}N_{sed}$ (and $\delta^{15}N_{kerogen}$) data obtained in this study compiled with the previous late Neoproterozoic data (Cremonese et al., 2013; Kikumoto et al., 2014; Spangenberg et al., 2014). (g) Histogram of published $\delta^{15}N_{sed}$ and $\delta^{15}N_{kerogen}$ data for Paleoproterozoic and Mesoproterozoic successions: 1.4–1.5 Ga Belt Basin, Montana ($\delta^{15}N_{sed}$

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from Stüeken et al., 2013); 1.8–1.9 Ga Animikie Basin, North America ($\delta^{15}N_{sed}$ from Godfrey et al., 2013); 2.0–2.1 Ga Onega Basin, NW Russia ($\delta^{15}N_{kerogen}$ from Kump et al., 2011) and 1.9–2.1 Ga Aravilli group, India ($\delta^{15}N_{sed}$ from Papineau et al., 2009, 2013).

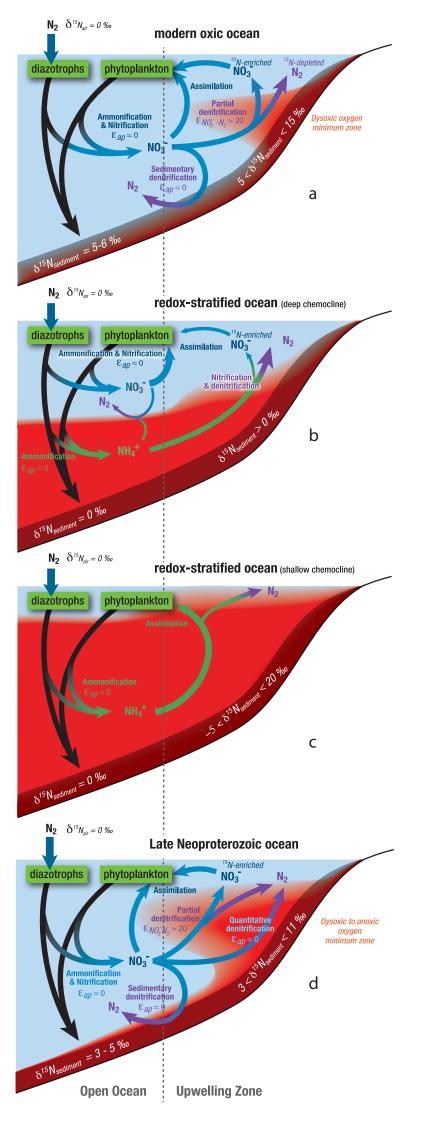
Figure 3: Paleogeographic reconstruction for 630 Ma (Modified from Li et al., 2013). Red empty stars indicate the locations of the four sample sets (South China, Amazon, NW Canada and NE Svalbard).

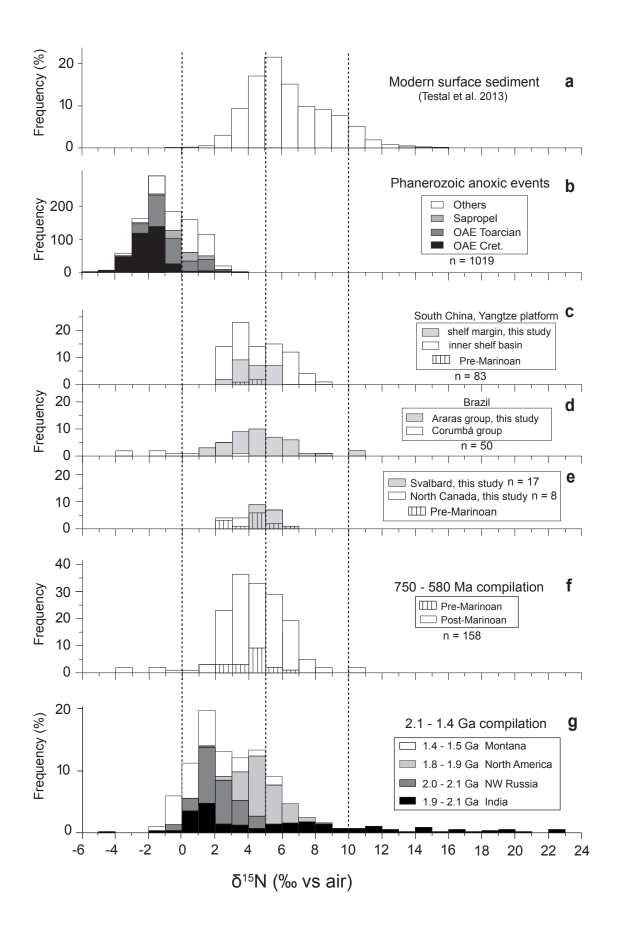
Figure 4: Chemostratigraphic representation of the present study $\delta^{15}N_{sed}$ results together with available $\delta^{13}C_{carb}$ and $\delta^{13}C_{org}$ data. (a–c) Sections spanning the Araras platform on the southeastern margin of the Amazonian craton (Fig. S1): (a) Composite Terconi-Camil section; (b) Carmelo section; (c) Copacel section. For Camil, Carmelo and Copacel sections, $\delta^{13}C_{carb}$ and $\delta^{13}C_{org}$ are from Sansjofre (2011) and for Terconi section from Sansjofre et al. (2011). (d) Yangjiaping section, Shelf margin of the Yangtze platform, South China craton. $\delta^{13}C_{carb}$ and $\delta^{13}C_{org}$ are from Macouin et al. (2004) and Ader et al. (2009). (e) Composite northeast Svalbard section. Previously published $\delta^{13}C_{carb}$ and $\delta^{13}C_{org}$ are from Halverson (2011). (f) Composite northwest Canada section. $\delta^{13}C_{carb}$ are from Hoffman and Schrag (2002).

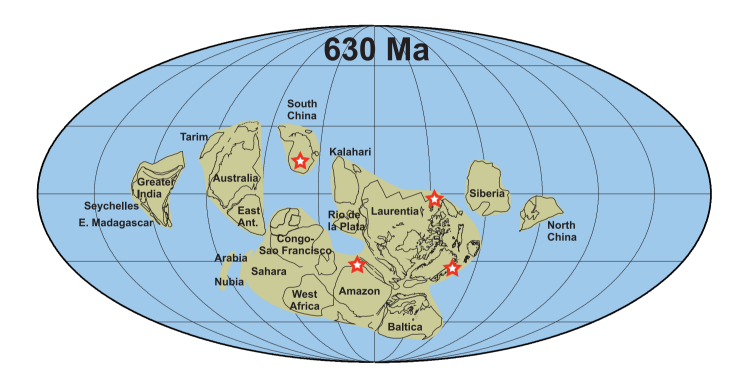
Figure 5: N and $\delta^{15}N_{sed}$ data analyses. (a) Cross-plots of $\delta^{15}N_{sed}$ values with the nitrogen content in ppm in the decarbonated residue (N_{decarb} ppm). (b) Cross-plots of $\delta^{15}N_{sed}$ values with C/N. (c) Cross-plots of N_{decarb} ppm with TOC_{decarb} ppm in log scales showing a broad correlation passing

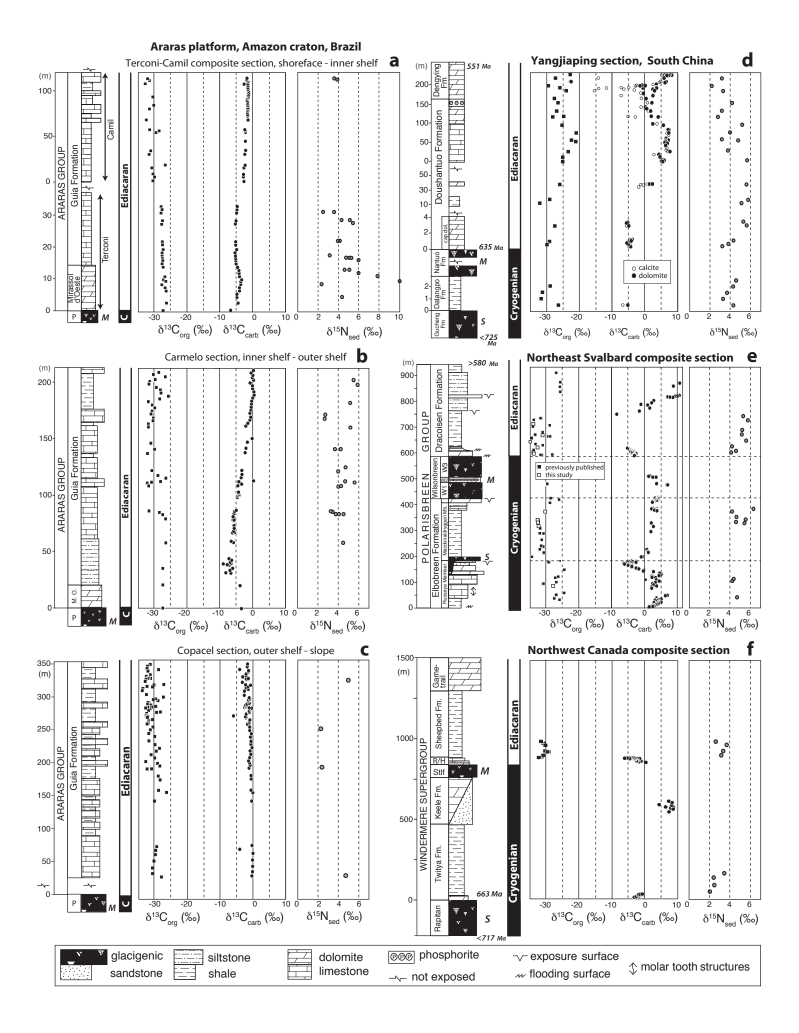
through the origin, as expected in the case of an organic origin for the sedimentary nitrogen. (d) Cross-plots of $\delta^{15}N_{sed}$ values with $\delta^{13}C_{org}$ showing no correlation.

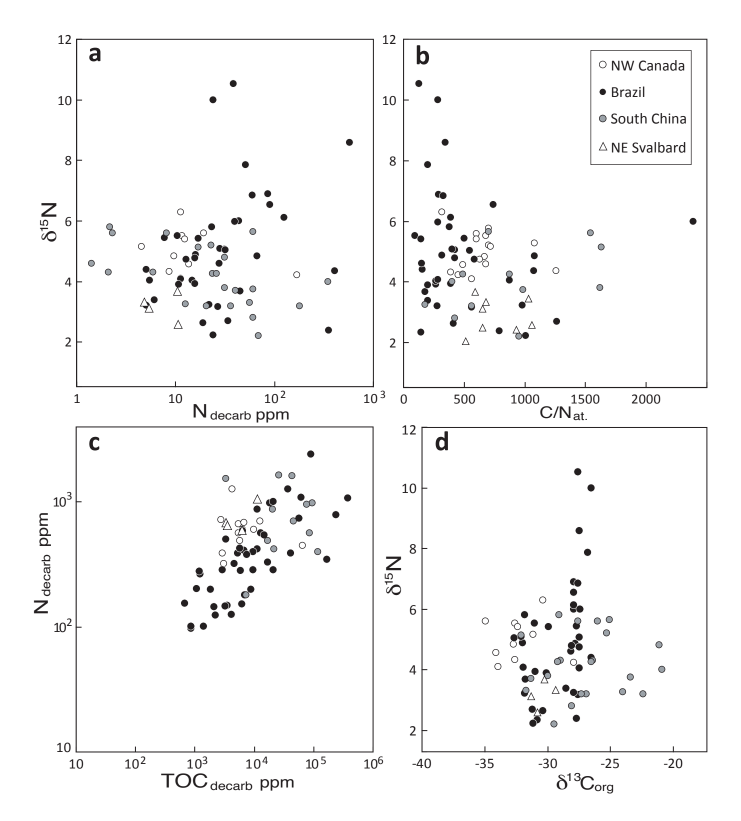
Figure 6: Summary figure showing the major features of biospheric and paleoenvironmental evolution from the early Neoproterozoic to the early Cambrian. Ranges of eukaryotes summarized from Macdonald et al. (2010). Composite carbonate δ^{13} C profile modified from Halverson and Shields-Zhou (2011). Mo concentrations in euxinic black shales from Sahoo et al. (2012). Nitrogen isotope data from this contribution, Cremonese et al. (2013), Kikumoto et al. (2014), and Spangenberg et al. (2014).











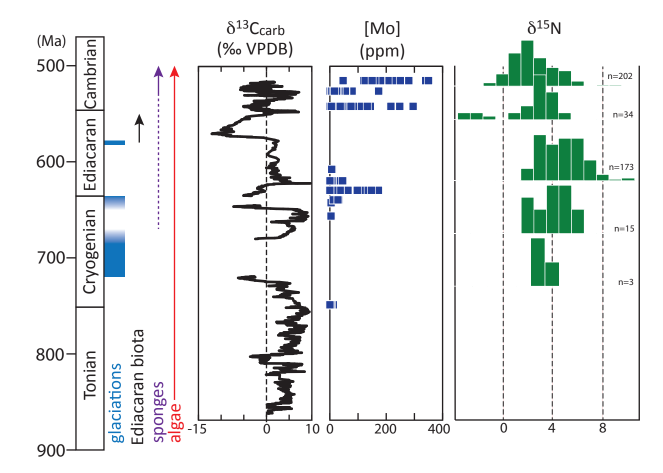


Table S1: FeT/Al ratios for Svalbard shales from the three units samples for nitrogen isotope analyses.

Sample	Stratigraphic unit	FeT/Al
C426 120	1	0.27
G436.138	lower Dracoisen Fm	0.37
G436.120	lower Dracoisen Fm	0.51
G436.105	lower Dracoisen Fm	0.71
G436.101.5	lower Dracoisen Fm	0.42
G436.95.5	lower Dracoisen Fm	0.45
G436.88	lower Dracoisen Fm	0.42
G436.64	lower Dracoisen Fm	0.44
G436.42	lower Dracoisen Fm	0.40
G436.10.5	lower Dracoisen Fm	0.49
G435.79.5	MacDonaldryggen Mbr	0.61
G435.76	MacDonaldryggen Mbr	0.62
G435.66	MacDonaldryggen Mbr	0.57
G435.56	MacDonaldryggen Mbr	0.58
G435.45	MacDonaldryggen Mbr	0.55
G435.35	MacDonaldryggen Mbr	0.48
G435.25	MacDonaldryggen Mbr	0.43
G435.18	MacDonaldryggen Mbr	0.32
G521.46	lower Russøya Mbr	0.37
G521.44.4	lower Russøya Mbr	0.57
G521.44.5 G521.40.5	lower Russøya Mbr	0.97
G521.40.3 G521.36	lower Russøya Mbr	0.57
G521.33	lower Russøya Mbr	1.16
G521.30	lower Russøya Mbr	0.27
G521.26.4	lower Russøya Mbr	0.27
G521.23.4	lower Russøya Mbr	0.66
G521.19.6	lower Russøya Mbr	0.45
G521.19.0 G521.18.9	lower Russøya Mbr	0.19
G521.13.2	lower Russøya Mbr	0.19
G521.13.2 G521.6.2	lower Russøya Mbr	0.24
G521.3.7	lower Russøya Mbr	0.24
0.521.5.7	10 woi Russwya wioi	0.54
Detrital baseline		
G526f	Wilsonbreen Fm	0.55
G526j	Wilsonbreen Fm	0.61

Table S2: Comparison of nitrogen content and $\delta^{15}N$ results obtained for bulk and decarbonated samples, measured by either static or by conventional isotope ratio mass

spectrometry.

	_		Bulk sa	mples		D				
Samples	decarb a	Sta	atic	Conv	entional	St	atic	Con	ventional	Average
name	wt %	$N_{bulk} \\$	$\delta^{15}N$	$N_{bulk} \\$	$\delta^{15}N$	N_{bulk}	$\delta^{15}N$	$N_{bulk} \\$	$\delta^{15}N$	$\delta^{15}N$ ‰
		ppm	±0.3‰	ppm	±0.15‰	ppm	±0.3‰	ppm	±0.15‰	
TeS 46	93	60	3.6			63	2.5	71	3.6	3.2±0.63
TeS 30	80	97	5.3			81	5.1	81	4.8	5.1 ± 0.25
CRS 16	60					301	5.1	300	4.6	4.8 ± 0.30
δXM01	10	770	4.1					530	3.3	3.7 ± 0.55
XM0107	11	945	4.2	970	4.7			980	4.0	4.3 ± 0.35
do1008	93	70	4.4			61	4.1			4.3 ± 0.21
do1013	59	685	5.4					630	5.8	5.6 ± 0.28
HU 99	5	1664	5.4	1640	4.9			1550	5.1	5.1 ± 0.28
HU 110	84	140	5.6					110	5.7	5.7 ± 0.07
do0603	98	32	3.5			17	4.0			3.8 ± 0.35
do0329	74	200	4.1					130	4.4	4.2 ± 0.21
do0351	97	19	3.4			5	3.1			3.3 ± 0.21
HU 180	89	140	2.5			101	1.9			2.2 ± 0.42

^a decarb: weight percent of the initial mass remaining after HCl attack

Table S3: δ^{15} N, δ^{13} C_{carb}, C_{org}, δ^{13} C_{org}, N content and N/C results for late Neoproterozoic samples from Brazil, NW Canada, NE Svalbard and South China. δ^{13} C_{carb}, δ^{13} C_{org} and C_{org} are from other studies for South China (Ader et al., 2009) and Brazil (Sansjofre et al., 2011; Sansjofre, 2011).

Formation	Stratigr.	Sample name	decarb		δ ¹⁵ N ±	0.1‰		[N] p	pm de	carb ¹	[N] ppm	Corg	ı %	δ ¹³ C _{carb}	δ ¹³ C _{org}	C/N
Formation	height (m)	Sample name	(wt%)	1 st	2 nd	3 rd	Avg	1 st	2 nd	Avg	bulk ²	decarb	Bulk	± 0.1‰	± 0.1‰	at
Yangjiaping section	ı, Yangtze platf	orm, South Chin	a ³													
Datangpo	-93	XM0101	54	4.3 ^{Sb}							210 ^{Sb}	0.23	0.11	-5.2	-26.4	5.9
Datangpo	-92.5	XM0103	12	3.3 ^{Sb}							1010 Sb	5.6	4.89		-31.7	56
Datangpo	-92	δXM01	10	4.1 ^{Sb}	3.3 ^b		3.7				770 ^b	3.0	2.69		-31.3	41
Datangpo	-91.5	XM0107	11	4.2 ^{Sb}	4.7 ^b	4.0	4.3	1100			945 ^{Sb}	0.19	0.17		-29.0	2.1
Datangpo	-91	XM0109	9	4.6 ^b							970 ^b	0.13	0.12	-1.8	-26.2	1.4
Doushantuo	0.4	do1002		3.2 ^{Sb}							70 ^{Sb}			-4.3		
Doushantuo	0.7	do1004	94	3.8				1620			97*	4.4	0.26	-5.0	-30.0	32
Doushantuo	1.1	do1008	93	4.4 ^{Sb}	4.1 ^S		4.3	875 ^S			70 ^{Sb}	2.0	0.14	-4.5	-29.2	27
Doushantuo	3.3	do1013	59	5.4 ^{Sb}	5.8		5.6	1540			685 ^{Sb}	0.33	0.14	-4.9	-27.6	2.3
Doushantuo	3.8	do1015		5.3 ^{Sb}							590 ^{Sb}			-5.2		
Doushantuo	11.7	HU 99	5	5.4 ^{Sb}	4.9 ^b	5.1	5.1	1630			1664 ^{Sb}	2.6	2.44		-32.1	17
Doushantuo	15.6	δdo 0406	6	5.8 ^b							1110 ^b	0.22	0.21		-29.1	2.2
Doushantuo	38.6	HU 106	52	5.6 ^{Sb}							336 ^b	0.49	0.24	2.3	-26.0	8.2
Doushantuo	0.7	HU 110	84	5.6 ^{Sb}	5.7		5.7	700			140 ^{Sb}	4.6	0.73	5.5	-25.1	61
Doushantuo	26.6	do 0603	98	3.5 ^{Sb}	4.0 ^S		3.8	983 ^S			32 ^{Sb}	9.6	0.17	7.7	-23.4	61
Doushantuo	52.9	HU 125	96	4.8 ^{Sb}							60 ^{Sb}	3.9	0.16	6.9	-21.1	31
Doushantuo	56.0	do 0625	97	3.2 ^S				560 ^S			15.9*	8.6	0.24	7.2	-22.4	179
Doushantuo	74.5	do 0635	98	4.0 ^S				400 ^S			9.5*	11.9	0.28	7.3	-20.9	346
Doushantuo	94.4	do 0301	75	5.2 ^{Sb}							310 ^{Sb}	2.5	0.61	5.7	-25.3	23
Doushantuo	115.9	do 0309	99	2.8				421			4,1*	2.2	0.02	4.1	-28.1	60
Doushantuo	131.0	do 0319	59	3.2 ^{Sb}							510 ^{Sb}	2.2	0.90	1.8	-26.9	20
Doushantuo	152.1	do 0329	74	4.1 ^{Sb}	4.4		4.3	490			200 ^{Sb}	1.7	0.44	2.2	-26.5	26
Dengying	183.3	do 0351	97	3.4 ^{Sb}	3.1 ^S		3.3	180 ^S			19 ^{Sb}	0.73	0.02	3.0	-24.0	12
Dengying	196.0	HU 180	89	2.5 ^{Sb}	1.9 ^S		2.2	950 ^S			140 ^{Sb}	7.7	0.83	-1.4	-29.5	69
Dengying	216.0	do1105	99.8	3.2 ^{Sb}							20 ^{Sb}	37	0.06	5.9	-27.3	36

Table S1 continued

Formation	Stratigr.	Sample name	decarb		δ^{15} N ±	0.1‰		[N]	pm de	carb ¹	[N] ppm	Corg	%	$\delta^{13} C_{carb}$		C/N
Formation	height (m)	Sample name	(%)	1 st	2 nd	3 rd	Avg	1 st	2 nd	Avg	bulk ²	decarb	Bulk	± 0.1‰		at
Terconi section, Bra	ızil ⁴															
Mirassol d'Oeste	4.2	TeS 5	99.5	4.4 ^S				1073 ^S			5*	38	0.18	-4.7		409
Mirassol d'Oeste	8.2	TeS 13	99.8	2.4 ^S				788 ^S			1*	24	0.05	-3.5	-27.7	352
Mirassol d'Oeste	9.2	TeS 15	95	10.0 ^S				283 ^S			14*	0.58	0.03	-3.5	-26.5	24
Mirassol d'Oeste	10.7	TeS 18	93	7.9				223			14*	0.88	0.06	-3.9	-26.8	51
Mirassol d'Oeste	11.7	TeS 20	93	6.0 ^S				2391 ^S			168*	9.0	0.63	-4	-27.9	44
Mirassol d'Oeste	12.7	TeS 22	92	4.6 ^S	5.1 ^S		4.9	1101 ^s	1061 ^S	1081	91*	6.2	0.52	-4.1	-27.8	66
Guia	15.7	TeS 28	89	6.0				286			32*	1.0	0.11	-5.1	-27.4	39
Guia	16.4	TeS 30	80	5.1 ^S	4.8	5.3 ^{Sb}	5.1	399 ^S	444	422	97 ^{Sb}	1.1	0.23	-5.1	-27.5	31
Guia	17.2	TeS 32	89	3.2				563			63*	1.3	0.15	-5.2	-27.6	27
Guia	21.7	TeS 40	92	4.0	4.1		4.0	875	637		71*	1.1	0.09	-5.1	-27.5	15
Guia	27.2	TeS 42	85	5.4				504			75*	0.34	0.05	-5.2	-27.7	7.8
Guia	28.2	TeS 44	88	4.3	5.1		4.7	532	636	584	73*	0.64	0.08	-5	-27.5	13
Guia	30.7	TeS 46	93	2.5 ^S	3.6	3.6^{Sb}	3.2	889 ^S	1072	980	60 ^{Sb}	1.8	0.13	-4.8	-27.9	22
Mirassol d'Oeste	12.8	TR+2.80	93	6.9				287			20*	2.1	0.15	-4.9	-27.9	86
Mirassol d'Oeste	10.6	TR +0.60	91	6.8	6.8		6.8	406	249	328	30*	1.7	0.15	-8.8	-27.6	60
Mirassol d'Oeste	10.2	TR +0.20	91	6.1				388			35*	4.2	0.38	-9.5	-27.9	125
Mirassol d'Oeste	10	TR -0.3	93	10.2	10.8		10.5	126	125	126	9*	0.41	0.03	-4.1	-27.6	38
Mirassol d'Oeste	8.3	TR-1.7	96	6.2	6.9		6.5	772	710	741	30*	5.7	0.23	-4.1	-27.9	90
Mirassol d'Oeste	7.3	TR-2.7	98	8.6				345			7*	17	0.34	-3.9	-27.5	575
Camil section, Brazi	l ⁴															
Guia	112.7	CAS 1	90	3.7				181			18*	0.69	0.07	-1.8	-31.8	45
Guia	111.7	CAS 2	90	3.9				267			27*			-2.0		
Carmelo section, Br	azil ⁴															
Guia	57.6	CRS 0	31	4.4				155			108	0.07	0.05	-5.3	-26.5	5.1
Guia	82.9	CRS 2n	24	4.4	4.0	3.7	4.0	260	270	265	201	0.12	0.09	-5.1	-27.5	5.5
Guia	84.8	CRS 3	63	3.4				203			75	0.11	0.04	-4.9	-28.5	6.1
Guia	85.6	CRS 4	20	3.2				278			224	0.12	0.10	-4.5	-31.8	5.0
Guia	107.3	CRS 5	64	4.2	3.9		4.1	284	299	292	108	0.29	0.10	-4.1	-31.9	11

Table S1 continued

Formation	Stratigr.	Comple nome	decarb		δ ¹⁵ N ±	0.1‰		[N] p	pm de	carb ¹	[N] ppm	Corg	%	δ ¹³ C _{carb}	$\delta^{13}C_{org}$	C/N
Formation	height (m)	Sample name	(%)	1 st	2 nd	3 rd	Avg	1 st	2 nd	Avg	bulk ²	decarb	Bulk	± 0.1‰		at
Guia	110.8	CRS 7	69	5.5				97			30	0.09	0.03	-3.9	-31.0	10
Guia	111.6	CRS 8	98	4.6 ^S				149 ^S			4*	0.36	0.01	0.0	-28.2	28
Guia	120.6	CRS 11	97	3.9				390			10*	0.53	0.01	-3.1	-31.0	16
Guia	140.1	CRS 13	81	4.2	3.6		3.9	203	196	199	38	0.19	0.04	-2.0	-30.1	11
Guia	159.2	CRS 16	60	5.1 ^S	4.7		4.9	336 ^S	301	319	134	0.46	0.18	-2.0	-32.0	16
Guia	167.1	CRS 19	96	2.6				410			16*	0.67	0.03	-0.7	-30.4	19
Guia	170.6	CRS 20	98	2.7				1264			21*	3.7	0.06	-1.0	-31.2	34
Guia	178.6	CRS 23	93					151			10*	0.63	0.04	-0.5	-30.8	48
Guia	181.2	CRS 24	83	5.1				369			68	0.96	0.16	-0.5	-32.1	28
Guia	191.8	CRS 28	96					124			5*	0.22	0.01	-0.3	-28.2	21
Guia	197.1	CRS 30	20	5.8 ^S				380 ^S			304	0.76	0.60	-0.9	-31.8	23
Guia	201.3	CRS 32	76	5.4				145			35	0.21	0.05	-0.3	-29.9	17
Guia	207.7	CRS 35	90					98			10*	0.14	0.01	-0.0	-28.7	16
Copacel section, A	raras platform,	Brazil ⁴														
Guia	35.1	NOS 13	98	4.8				425			7*	0.57	0.01	-0.2	-28.1	16
Guia	76.2	NOS 8	68					101			32	0.09	0.03	-3.9	-29.7	10
Guia	201.3	CP 1	89	2.3				146				0.32	0.03	-1.0	-30.8	
Guia	259.3	CP13	95	2.2				1009			48*	2.1	0.10	-0.8	-31.2	24
Guia	333.8	CP 51	75	5.0				545			138	1.5	0.38	-3.0	-32.6	32
Northeast Svalbard	d composite sec	ction⁵														
Dracoisen	19.4	G136-19.4		4.1				540	580	560		0.54			-33.9	11
Dracoisen	24.1	G136-24.1		4.7	4.5		4.6	490	490	490		0.57			-34.1	14
Dracoisen	105	G436-105		5.2				700								
Dracoisen	88	G436-88		5.2				720				0.28			-31.2	4.5
Dracoisen	64	G436-64		5.6				600				0.99			-34.9	19
Dracoisen	42	G436-42		4.3				390				0.29			-32.6	8.7
Dracoisen	159	G306-159		5.3				1080								
Dracoisen	143	G306-143		5.8				700				1.26				

Table S1 continued

Formation Stratigr. Sam height (m)	Sample name	decarb		$\delta^{15}N$ ±	0.1‰		[N] p	pm de	carb ¹	[N] ppm	Corg %		$\delta^{13}C_{carb}$	δ ¹³ C _{org}	C/N	
	Sample name	(%)	1 st	2 nd	3 rd	Avg	1 st	2 nd	Avg	bulk ²	decarb	Bulk	± 0.1‰	± 0.1‰	at	
Macdonaldryggen	-25	G435-25		6.3				320				0.31			-30.4	11
Macdonaldryggen	-35	G435-35		4.2												
Macdonaldryggen	-56	G435-56		4.8				670				0.55			-32.7	9.6
Macdonaldryggen	-66	G435-66		5.5				680				0.68			-32.6	12
Macdonaldryggen	-76	G435-76		4.6				680								
Macdonaldryggen	-79.5	G435-79.5		5.5	5.3		5.4	610	590	600		0.64			-32.4	12
Russøya Mb.	114	G406-35		4.2	4.5		4.4	1280	1240	1260		0.42				
Russøya Mb.	106.5	G406-42		4.2				450				6.56			-27.9	170
Russøya Mb.	43.5	P5014-43.5		4.7				630								
Northwest Canada c	omposite sect	tion ⁶														
Sheepbed	94	G4-94		2.6				1060				1.13			-30.8	11
Sheepbed	73	G4-93		3.7				590				0.62			-30.3	10
Sheepbed	36	G4-36		3.3				680				0.33			-29.4	4.9
Sheepbed	8	G4-8		3.1				650				0.35			-31.3	5.4
Twitya	163	F849.163		3.5 ^b							1030 ^b					
Twitya	135	F849.135		2.4 ^b							930 ^b					
Twitya	87	F849.87		2.5 ^b							650 ^b					
Twitya	46	F849.46		2.1 ^b							510 ^b					

Most N content and δ^{15} N measurements were performed on decarbonated samples by conventional mass spectrometry. N content and δ^{15} N measurements performed on bulk samples are indicated by 5 and measurements performed using the static mass spectrometry are indicated by 5 .

¹ [N] ppm decarb: values correspond to the direct measurement of the nitrogen content in the decarbonated residu.

² [N] ppm bulk: values correspond either to direct measurement of the nitrogen content in bulk samples or in the recalculated nitrogen content in the bulk samples from the nitrogen content in the decarbonated residu and the mass pourcent of mass loss after decarbonatation (decarb %).

^{*} indicates most probably underestimated N content in bulk rocks, owing to possible mass loss during the decarbonatation procedure (see method section in the main text and Table S2)

³ For the South China data, the 0 datum correspond to the contact between the cap carbonate and the Nantuo Diamictite.

⁴ For the Brazilian data, the datums are as follows, 0 = contact with the Marinoan Diamictites of the Puga Formation for Terconi, Carmelo and Copacel sections. For Camil section, 0 = firt outcrop of the Guia Formation observed in Camil quarry, which is 114 m below the Guia-Serra Do Quilombo transition.

⁵ For the Svalbard data, the datums are as follows: for the Dracoisen formation, 0 = based of the formation; for the Macdonaldryggen Member, 0 = the top of the member; for the Russøya Member, 0 = the base of the member

⁶ For the NW Canada data, the 0 m datums are the base of each formation.