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## REPORT



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# Iron-mediated anaerobic ammonium oxidation recorded in the early Archean ferruginous ocean

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## Abstract

The nitrogen isotopic composition of organic matter is controlled by metabolic activity and redox speciation and has therefore largely been used to uncover the early evolution of life and ocean oxygenation. Specifically, positive  $\delta^{15}N$  values found in well-preserved sedimentary rocks are often interpreted as reflecting the stability of a nitrate pool sustained by water column partial oxygenation. This study adds much-needed data to the sparse Paleoarchean record, providing carbon and nitrogen concentrations and isotopic compositions for more than fifty samples from the 3.4 Ga Buck Reef Chert sedimentary deposit (BRC, Barberton Greenstone Belt). In the overall anoxic and ferruginous conditions of the BRC depositional environment, these samples yield positive  $\delta^{15}$ N values up to +6.1‰. We argue that without a stable pool of nitrates, these values are best explained by non-quantitative oxidation of ammonium via the Feammox pathway, a metabolic co-cycling between iron and nitrogen through the oxidation of ammonium in the presence of iron oxides. Our data contribute to the understanding of how the nitrogen cycle operated under reducing, anoxic, and ferruginous conditions, which are relevant to most of the Archean. Most importantly, they invite to carefully consider the meaning of positive  $\delta^{15}$ N signatures in Archean sediments.

## KEYWORDS

ammonium oxidation, biogeochemistry, Buck Reef Chert, Feammox, ferruginous ocean, nitrogen isotopes, Paleoarchean

## 1 | INTRODUCTION

Nitrogen is an essential nutrient for all living organisms, and its availability strongly influences biological productivity. As water column redox speciation controls active metabolic pathways of the biogeochemical nitrogen cycle, the nitrogen isotopic composition of sedimentary rocks can record oceanic redox changes in the early oceans. On the modern Earth, the atmosphere is the main surface nitrogen reservoir, where N is present mostly as gaseous dinitrogen  $N_2$  (Ward, 2012,  $\delta^{15}N = 0\%$ ). However, nitrogen in its  $N_2$  form can only be assimilated by diazotrophs, prokaryotic organisms who possess the nitrogenase enzyme and are thus capable of biological  $N_2$  fixation (Raymond et al., 2004). Bioavailable "fixed" forms of nitrogen are therefore almost exclusively provided to the biosphere by  $N_2$ -fixating

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diazotrophs and their mineralization in the water column or in the sediment, referred to as ammonification (Sigman et al., 2009). Fractionations imparted by biological N<sub>2</sub> fixation with classical Mobased nitrogenase range from -2‰ to +2‰, whereas alternative nitrogenases using Fe or V as cofactors can impart negative fractionations as large as -8‰ (Zhang et al., 2014). In the modern ocean, transfers between the different oceanic nitrogen reservoirs (N2,  $NH_{4}^{+}$ ,  $NO_{3}^{-}$  and dissolved and particulate organic nitrogen DON and PON) are mainly controlled by biological processes involving redox reactions. The isotopic composition of these nitrogen sources determines the isotopic composition of primary producers and therefore the sedimentary  $\delta^{15}$ N. Ammonium release during organic matter remineralization (ammonification) does not impart significant isotope fractionation (Möbius, 2013), and ammonium is rapidly assimilated into the biomass, as it is the preferred inorganic nitrogen source for many organisms. Consequently, even if ammonium assimilation preferentially incorporates <sup>14</sup>N, its fractionation is almost never expressed. In oxic conditions prevailing during the Phanerozoic, ammonium can also be successively nitrified to nitrites NO2<sup>-</sup> and nitrates NO<sub>3</sub><sup>-</sup>, which are either assimilated by photosynthetic organisms or biologically reduced through denitrification or anaerobic ammonium oxidation (anammox) in dysoxic and anoxic conditions (Dalsgaard & Thamdrup, 2002; Sigman et al., 2009). Denitrification and anammox release N<sub>2</sub>O or N<sub>2</sub>, which are subsequently lost to the atmosphere, making them the major oceanic sinks of fixed nitrogen. As denitrification imparts a large nitrogen isotope fractionation of ≈30‰ (Sigman et al., 2009), modern and by extension Phanerozoic sedimentary  $\delta^{15}$ N values around +5‰ result from the balance between N<sub>2</sub> fixation inputs and denitrification/anammox outputs.

The Archean biogeochemical nitrogen cycle is thought to have been very different from the Phanerozoic, as the redox states of the ocean and the atmosphere have evolved together with rising oxygen levels, impacting environmental conditions and biological evolution. The analysis of 3.5 Ga fluid inclusions containing N<sub>2</sub> (Nishizawa et al., 2007) points toward a stable isotopic composition of atmospheric nitrogen ( $\delta^{15}N_{N2} = 0$ %) at least since the Paleoarchean. Without oxygen levels significant enough to allow nitrification, nitrites and nitrates were likely scarce or absent. Therefore, ammonium is often assumed to have been the dominant form of fixed N prior to the rise of oxygen levels (Beaumont & Robert, 1999; Canfield et al., 2010). Potential abiotic sources of ammonium, including hydrothermal reduction, lightning, and volcanic eruptions, have been reviewed in Stüeken et al. (2016) and their magnitude was deemed too small to have sustained the Archean biosphere. The range of Paleoarchean sedimentary  $\delta^{15}$ N values (reviewed in Ader et al., 2016) centers around 0‰, which is compatible with a primitive nitrogen cycle dominated by biological N<sub>2</sub> fixation, ammonification, and ammonium assimilation. The rise of oceanic oxidants preceding the Great Oxidation Event (GOE) might have initiated nitrification/ denitrification during the Neoarchean, when positive  $\delta^{15}N$  values have been observed (Garvin et al., 2009; Godfrey & Falkowski, 2009; Stüeken et al., 2015; Thomazo et al., 2011), paving the way towards an aerobic nitrogen cycle similar to modern environments.

The first nitrogen isotope Precambrian time series was published by Beaumont and Robert (1999). Since then a lot of data have been acquired but only a few for the Paleoarchean (reviewed in Ader et al., 2016; Mettam & Zerkle, 2021; Stüeken et al., 2016). This is due to the scarcity of well-preserved Paleoarchean geological formations and the difficulty of analyzing nitrogen-poor samples. Therefore, the exceptionally well-preserved and continuous sedimentary record provided by the 900 m BARB3 drill core through the 3.4 Ga Buck Reef Chert offers a unique opportunity to gain some insight into Paleoarchean environments in which an early biosphere could have developed. Moreover, it opens a deep-time window to explore early nitrogen cycle mechanisms such as the onset of N<sub>2</sub> fixation and redox reactions.

#### 2 **GEOLOGICAL CONTEXT**

The 3416±5 Ma (Krüner et al., 1991) Buck Reef Chert (BRC) is the largest chert deposit of the Kromberg Formation in the Onverwacht Group, located in the Barberton Greenstone Belt, Kaapvaal craton, South Africa (Lowe & Worrell, 1999). The ICDP-sponsored BARB3 drill core (Hofmann et al., 2013) through the BRC sedimentary deposit records a transition between two sedimentary units: a basal chert-dominated unit from 900 to 600 m depth and a BIF-dominated unit from 600 to 400m depth. Microscopic observations on the studied BARB3 samples matches detailed facies studies performed on outcrop samples by Tice and Lowe (2006a), except for the evaporitic facies that has not been intersected by the drill core.

The chert-dominated unit consists of a succession of black and white banded cherts with sand-sized carbonaceous grains and laminae, granular cherts, chert breccia and ferruginous cherts with intercalated siderite layers (Figure 1; Figure S1).

The BIF-dominated unit is mainly composed of thick layers of iron-rich carbonates alternating with a smaller proportion of chert beds. Siderite occurs as densely packed rhombic and spherulitic crystals, typical of early diagenetic siderite.

A distinctive feature of the BIF-dominated unit is the occurrence of thin layers ≈1 cm thick of microcrystalline quartz and carbonate characterized by subspherical domains (25 µm) containing micron to sub-micron-sized inclusions of red iron oxides. Single quartz crystals 50µm in diameter enclose the spheres. These jasper layers are located in the depth intervals of 580-560, 545-500, and 480-400 m (Figure 1, Figure S1). Euhedral iron-rich carbonate crystals in jasper occur in varying amounts (ca. 5%-10%) and are either disseminated or in lenticular clusters, suggesting their early diagenetic nature. The jasper layers are unrelated to any zones where oxic groundwaters could have circulated at depth, which suggests they are primary.

The transition between the chert- and BIF-dominated units is thought to record a temporal and environmental change from a wave-dominated shallow platform to a deeper low-energy basinal depositional environment (Tice & Lowe, 2004) and could also reflect chemical variability linked to an increased supply of dissolved iron through hydrothermal pulses. The origin of the BRC silicified



FIGURE 1 Stratigraphic evolution of carbonate content (wt. %), TOC (wt. %), TN (ppm), TOC/TN,  $\delta^{13}C_{org}$  (‰ vs. PDB), and  $\delta^{15}N$  (‰ vs. air) along the BARB3 drill core.

laminations is variously interpreted in the literature, and four main hypotheses stand out: (i) water column chemical precipitation (Ledevin et al., 2019), (ii) secondary and late silicification of a laminated sedimentary or volcano-sedimentary precursor (de Vries et al., 2006), (iii) segregation from an initial homogenous mix of silica and other components during early diagenesis (Tice & Lowe, 2006a), (iv) pulses of hydrothermal silicification (Geilert et al., 2014). The origin of siderite in the BIF-dominated unit is also debated, as ironrich carbonates can either precipitate directly in a stratified water column rich in dissolved iron (Klein & Beukes, 1989) or form in the sediment close to the sediment-water interface during diagenesis (Heimann et al., 2010). A hydrothermal origin for siderite has also been suggested for some Archean BIFs (Bolhar et al., 2005).

Regional metamorphism does not exceed the greenschist facies (Tice et al., 2004), with peak temperatures of  $310\pm50^{\circ}$ C, according to Raman geothermometry of carbonaceous matter (Alleon et al., 2021).

## 3 | METHODS

## 3.1 | Sampling

Fifty-one samples were chosen along the drill core according to their organic matter content (TOC>0.1% in the digestion residue) that mostly reflects their potential in containing enough nitrogen

to analyze. Those samples are composed of various proportions of bands of chert and siderite. Only samples CT6, CT7, and CT18 contain jasper layers. Accordingly, the amount of carbonates in the samples has been used to define two lithological endmembers: "chert" (when carbonates range from 0 to 50 wt. %) and "siderite" (when carbonates range from 50 to 100 wt. %).

## 3.2 | Mineralogy

Mineralogy was investigated before and after chemical treatment using X-Ray Diffraction (XRD) at the Biogéosciences Laboratory (Université de Bourgogne Franche-Comté, France). Diffractograms were obtained with a Bruker D8 Endeavor diffractometer with CuK $\alpha$ radiation, LynxEye XE-T detector and Ni filter, under 40kV voltage and 25 mA intensity. The goniometer scanned from 2° to 65° 2 $\theta$  for each run. Identification of crystalline phases was based on the position and area of their respective mean basal reflections.

## 3.3 | Chemical treatment

Samples were first crushed into powder using a ring and puck mill at the Biogéosciences Laboratory (Université de Bourgogne Franche-Comté) in order to obtain sample powder smaller than  $60\mu m$ . Carbonate-free residues were obtained by mixing sample powders

with 6 N HCl for 24 h, followed by a second step of digestion at 80°C for 4 h. The powder was then rinsed with deionized distilled water to a neutral pH and oven-dried at 40°C for 48 h. Carbonate content, expressed in weight percent (wt. %), was evaluated through gravimetric mass balance after HCl digestion. Some kerogen extracts were commercially produced by Global Geolab Ltd, Alberta, Canada. Powders were digested using HCl and HF, and kerogens were separated out by heavy liquid separation with zinc bromide.

#### C isotopic analyses 3.4

The decarbonated residues were poured into tin capsules (20 to 80mg of powder) and weighted using a Sartorius M2P ultra-balance before TOC and  $\delta^{13}C_{org}$  measurements were performed using a Thermo Fisher Scientific Flash Smart elemental analyzer, coupled to a Thermo Fisher Scientific Delta V isotope ratio mass spectrometer (EA-IRMS) via a Conflo IV interface at the Biogéosciences laboratory. Certified USGS40 ( $\delta^{13}C_{org} = -26.2\%$ , TOC = 40.82 wt. %) and caffeine IAEA-600 ( $\delta^{13}C_{org} = -27.77\%$ ) reference materials were used for the calibration. Total organic carbon (TOC) contents are expressed in dry weight percentage (wt. %) of the non-decarbonated bulk powder and isotope results are reported in delta-notation relative to V-PDB. Each measurement session included three to four standard measurements at the beginning and at the end, as well as one standard measurement every 10 samples. The mean  $\delta^{13}C_{ore}$ precision for standards is 0.8‰ and the mean accuracy 0.5‰. The number of replicates for each sample is shown in Table 1. The mean external reproducibility ( $2\sigma$ ), based on sample replicate analyses and including powder resampling and reprocessing via chemical treatment, is  $\pm 0.06$  wt. % for the TOC content and  $\pm 0.86$ % for the  $\delta^{13}C_{org}$ . The  $\delta^{13}C_{org}$  of 10 kerogen extracts (Table 1) has also been measured following the same procedure mentioned above: their average deviation from corresponding bulk  $\delta^{13}\mathsf{C}_{_{\mathrm{org}}}$  measurements is on average  $0.63 \pm 0.68$ %.

#### 3.5 N isotopic analyses

As most samples contain less than 200 ppm N (Table 1) and the proportion of silicate-bound nitrogen is unknown, the above-described EA-IRMS method is not sensitive enough for reliable bulk nitrogen isotopic analyses (Ader et al., 2016; Boocock et al., 2020). Samples were therefore analyzed using the "classical method" developed at IPGP described in Ader et al. (2014, 2016), in which N<sub>2</sub> is produced offline through sealed-tube Dumas combustion and cryogenically purified in a vacuum line. Up to 200 mg of decarbonated residual powder was put into a quartz tube with CuO and Cu wires, purified beforehand at 900°C for 2 h in a muffle furnace to prevent contamination. Samples were degassed for 12h at 150°C under vacuum to remove adsorbed atmospheric N<sub>2</sub> and organics. Quartz tubes were then sealed directly under vacuum and combusted in a muffle furnace at 950°C for 6 h under oxidizing conditions by oxygen liberated

from the CuO wires, then cooled at 600°C for 2 h, allowing residual oxygen to combine with cupric oxide and nitrous oxide to be reduced by copper, and finally cooled to ambient temperature (Beaumont & Robert, 1999; Kendall and Grim, 1990). This allows the chemical separation of gaseous products. Busigny et al. (2005) have shown that the extraction yield for this protocol is 100% for both organic and mineral nitrogen, including ammonium in minerals such as phyllosilicates. The content of each quartz tube is liberated in the vacuum line with a tube cracker, where  $CO_2$  and  $H_2O$  are trapped cryogenically to avoid any isobaric interferences. The purified incondensable N<sub>a</sub> gas is concentrated into a calibrated volume for quantification using a Toepler pump (Hg manometer). Standard analytical procedures for nitrogen usually include CaO in the reagents to trap gaseous CO<sub>2</sub> and H<sub>2</sub>O liberated from samples (Kendall and Grim, 1990), but given that Busigny et al. (2005) have shown that it significantly contributes to analytical blanks, we performed a few tests on samples from the BRC which show that the addition of CaO does not yield significant  $\delta^{15}N$  differences. Most samples were consequently analyzed without the addition of CaO, except for the two kerogen extracts. Analytical blanks for the entire procedure are <0.05 micromoles N, which represents less than 10% of the gas for smaller samples and less than 1% for more concentrated samples. Purified N<sub>2</sub> is analyzed by dual-inlet mass spectrometry using a ThermoFinnigan DeltaPlus XP IRMS. Possible air contamination and isobaric interferences due to CO are monitored by scanning of m/z 12 (C from CO<sub>2</sub>, CO, CH<sub>4</sub>, or organic compounds), 18 (H<sub>2</sub>O), 30 (C<sup>18</sup>O), 32 (O<sub>2</sub>), 40 (atmospheric Ar), and 44 (CO<sub>2</sub>). External  $\delta^{15}$ N reproducibility falls between 0.10 and 1.02‰ with a mean of  $0.44 \pm 0.29\%$  (n = 14). Samples that were replicated are reported in Table 1. Samples with no external reproducibility are reported with an error of +0.63‰, which is the external reproducibility of sample 75A replicated 10 times at different concentrations (Table S1). Kerogen extracts of two samples (CT16 and CT18U) yield TN contents and  $\delta^{15}N$  values identical within external reproducibility to those measured on their decarbonated counterparts (Table S2).

### RESULTS 4

Iron-rich carbonate throughout the BRC is identified mostly as siderite, with a few samples containing ankerite, based on XRD analyses (Figures S2 and S3; Table S3), which is consistent with previous investigations (Tice et al., 2004; Tice & Lowe, 2006a; Tice & Lowe, 2006b). Carbonate content obtained through HCI digestion is highly variable (2% to 99%) depending on the relative proportions of bands of chert versus siderite. Total organic carbon (TOC) contents vary between 0.01% and 5.01% with a mean value of  $0.66 \pm 1.01\%$ (n = 52). Total nitrogen (TN) content ranges from 11 to 259 ppm with a mean of  $55 \pm 53$  ppm (n = 55), excluding two N-rich outliers (samples 75A and 73A showing TN contents of 1823 and 705 ppm, respectively). TOC/TN ratios are below 150, which is relatively low for Archean sedimentary rocks (Yamaguchi, 2002); it ranges from 3 to 141 with a mean TOC/TN of  $72 \pm 24$  (n = 51).

viation (SD) an	d number of rep	licates, for eac	th sample and	alyzed along t	the BARB	3 drill core.										:RIN I
		Carbonate														ET AL.
ample	Depth (m)	content (wt. %)	TOC (wt. %)	SD	Nb repl.	TN (wt. ppm)	SD	Nb repl.	TOC/TN	δ <sup>13</sup> C <sub>org</sub> (‰ vs. PDB)	SD	Nb repl.	δ <sup>15</sup> N (‰ vs. air)	SD	Nb repl.	
7A <sup>a</sup>	421.37	96	0.01		1	41		1	ю	-27.96	0.84	1	-0.3	0.63	1	
CT19U <sup>a</sup>	433.18	93	0.40	0.14	2	45		1	87	-26.68	0.84	1	0.2	0.63	1	
CT19L <sup>a</sup>	433.24	54	0.21	0.02	e	26		1	81	-29.58	1.73	с	0.2	0.63	1	
CT18U <sup>a</sup>	437.27	83	0.69	0.21	2	114	e	2	60	-28.13	0.35	2	-0.3	0.10	2	
CT18L <sup>a</sup>	437.36	65	0.28	0.02	т	35		1	78	-29.37	2.77	ო	-0.1	0.63	1	
CT17 <sup>a</sup>	448.64	86	0.42	0.07	с	58	ю	с	73	-28.59	1.23	2	0.6	0.22	ო	
CT16 <sup>a</sup>	459.03	70	0.55	0.15	2	06	5	ო	61	-29.78	0.39	2	1.1	0.18	ო	
CT15 <sup>a</sup>	466.33	35	0.41	0.01	С	50		1	81	-30.23	1.93	ო	0.8	0.34	2	
CT14U <sup>a</sup>	470.82	60	0.11	0.02	т	18		1	60	-28.81	0.39	2	3.3	0.63	1	
CT14M <sup>a</sup>	470.94	81	0.10	0.01	с	18	1	2	57	-26.82	1.11	ო	1.5	0.25	2	
CT14L <sup>a</sup>	471.05	73	0.17	0.02	с	24		1	69	-27.84	0.40	т	2.9	0.63	1	
CT13 <sup>a</sup>	483.39	36	0.30	0.15	ო	36	ო	2	83	-32.45	2.69	ო	0.7	0.64	2	
75A	483.81	30	3.77		1	1823	96	10	21	-36.11	0.84	1	4.0	0.63	10	
CT5U_U	491.66	39	0.14	0.04	т	29		1	48	-31.42	0.29	ო	6.1	0.63	1	
CT5U_M	491.74	75	0.45	0.09	с	82		1	55	-30.43	0.83	ო	4.5	0.63	1	
CT5U_L	491.82	47	0.11	0.01	т	22		1	49	-29.17	0.21	ო	5.0	0.63	1	
CT5L_U	491.9	76	0.18	0.02	б	28		1	66	-29.14	0.17	ო	4.9	0.63	1	
CT5L_M	491.98	49	0.16	0.03	С	32		1	49	-29.54	0.22	ო	3.4	0.63	1	
CT5L_L	492.04	96	0.37	0.25	2	61		1	61	-28.93	0.71	2	4.4	0.63	1	- 9
73A	498.09	60	5.01		1	705	34	ო	71	-31.53	0.84	1	2.8	0.31	ო	<b>gek</b>
CT6U <sup>a</sup>	504.68	71	0.16	0.01	e	20		1	82	-27.51	0.36	ო	1.6	0.63	1	biol
CT6L <sup>a</sup>	504.8	89	0.26	0.08	с	40	2	2	65	-28.13	0.36	с	2.0	0.19	7	og
72B <sup>a</sup>	511.05	66				23		1		-28.97	0.84	1	2.9	0.63	1	У
21A	517.53	92				16		1		-27.91	0.84	1				0
CT7 <sup>a</sup>	520.05	75	0.22	0.01	e	28		1	77	-28.44	0.61	e	1.9	0.63	1	3
20A <sup>a</sup>	538.26	06				13		1		-27.17	0.84	1	3.8	0.63	1	
CT12U	548.08	98	0.37	0.28	2	60		1	61	-29.80	0.84	1	4.3	0.63	1	W
CT12L	548.15	89	0.38	0.03	ю	40		1	97	-32.08	0.19	ю	5.2	0.63	1	IL
CT11U	558.62	70	0.16	0.01	ი	17		7	95	-27.93	0.50	ю	5.3	0.63	~	ΕY
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		Carbonate content	TOC (wt.		qN	TN (wt.		qN		δ <sup>13</sup> C <sub>ore</sub> (‰		qN	δ <sup>15</sup> Ν (‰		qN
Sample	Depth (m)	(wt. %)	(%	SD	repl.	(mdd	SD	repl.	TOC/TN	vs. PDB)	SD	repl.	vs. air)	SD	repl.
CT11M	558.68	92	0.31	0.04	ო	40		Ļ	77	-29.01	0.25	ო	5.0	0.63	1
CT11L	558.75	71	0.09	0.02	ო	13		7	73	-28.68	0.15	с	5.4	0.63	1
CT10	562.96	98	0.23	0.08	с	32		1	70	-29.82	0.01	2	3.3	0.63	1
CT9	573.11	54	0.14	0.00	ო	17		1	84	-29.62	2.69	ო	3.4	0.63	1
CT8U_U	581.76	61	0.17	0.05	ო	23	4	2	76	-31.16	1.42	ო	4.1	0.63	1
CT8U_L	581.88	49	0.14	0.01	ო	20	2	e	72	-27.48	0.20	с	5.0	0.12	2
CT8L_U	581.93	98	0.27	0.05	ო	32		Ļ	85	-28.43	0.84	1	4.3	0.63	1
CT8L_M	581.96	75	0.18	0.01	ო	21		7	85	-28.92	0.28	с	4.3	0.63	4
CT8L_L	582	56	0.23	0.02	ო	30		Ļ	75	-29.31	0.14	ო	3.9	0.63	1
18A	583.03	96				12		7		-28.10	0.84	1	4.5	0.63	1
17A	616.77	13	1.41		1	180	7	e	78	-40.35	0.84	1	4.3	0.57	ო
16A	662.32	68	0.96		1	66		1	98	-35.91	0.84	1	2.7	0.63	1
61A	682	39	3.65		1	259		7	141	-39.40	0.84	1	3.4	0.63	1
58A	747.59	59	1.83		1	137		2	134	-31.72	0.84	1	1.9	1.02	2
CT4U	770.65	19	0.11	0.04	2	11		1	66	-30.18	2.47	2			
CT4L_U	770.8	6	0.37	0.05	2	41		1	89	-30.01	0.33	7	2.0	0.63	1
CT4L_L	770.95	2	0.28	0.00	ю	42		1	66	-30.68	0.03	2	3.6	0.63	1
9A	794.12	23	1.89		1	159	2	2	118	-39.39	0.84	1	2.8	0.75	2
52A	813.05	23	2.61		1					-36.77	0.84	1			
CT3U	837.32	4	0.78	0.03	2	140		1	56	-33.01	0.32	7	5.8	0.63	1
CT3L	837.51	С	0.62	0.06	2	181		1	34	-32.15	0.45	7	5.6	0.63	1
49A	865.76	29	0.05		1	27	18	2	17	-29.37	0.84	1			
CT2U	874.65	2	0.47	0.06	2	71		1	66	-30.52	0.99	2	3.3	0.63	1
CT2L	874.87	16	0.32	0.01	2	33	4	ю	96	-30.24	1.28	7	1.4	0.82	7
CT1U	883.47	9	0.78	0.05	2	94		1	83	-30.99	3.43	7	1.3	0.63	1
CT1L	883.65	ю	0.08	0.01	2	16		7	49	-29.25	0.04	2			
1A	896	13	0.88		1	128		Ţ	69	-38.49	0.84	1	3.9	0.63	1
Samples containing jas	iper layers.														

Note: Green shade indicates Cherts (carbonates <50wt. %); Grey shade indicates Siderites (carbonates >50 wt. %).

Along the drill core,  $\delta^{15}$ N values (n = 51) range between -0.3%and + 6.1% with a mean  $\delta^{15}$ N of  $3.1 \pm 1.8\%$ .  $\delta^{15}$ N variations are not correlated with either organic carbon ( $R^2 = .0308$ ) or nitrogen concentrations ( $R^2 = .0043$ ) (Figure 3). The average  $\delta^{15}$ N value in the chert-dominated unit does not significantly differ from the average  $\delta^{15}$ N value in the BIF-dominated unit, nor do  $\delta^{15}$ N values in chert samples significantly differ from  $\delta^{15}$ N values in siderite samples (Figure 2, p > .05). However,  $\delta^{15}$ N values (n = 17) from the BIF-dominated unit with jasper layers are significantly different (Figure 2, p < .05) from  $\delta^{15}$ N values (n = 34) in core intervals lacking jasper, with a mean  $\delta^{15}$ N of 1.3‰ and 4.0‰ respectively.  $\delta^{15}$ N values display stratigraphic variations along the BARB3 drill core, with markedly positive values in most of the core, and values decreasing toward 0‰ in the upper part, between depths of 400 and 500 m.

 $\delta^{13}\mathrm{C}_{\mathrm{org}}$  in the BIF-dominated unit and in the chert-dominated unit show close mean values of -29.3  $\pm$  1.8% (n = 39) and - 33.4  $\pm$  4.0% (n = 17) respectively, but the latter displays a much larger range of variations (from -40.4% to -29.3%), whereas values in the BIF-dominated unit are much more homogenous (from -32.5% to -26.7% leaving out one outlier). The same  $\delta^{13}\mathrm{C}_{\mathrm{org}}$  variations hold true for chert and siderite samples (Figure 2).

## 5 | DISCUSSION

The potential impact of post-depositional processes on the  $\delta^{15}N$  is discussed in Supplementary Information. All indicators point to a good preservation of the BRC samples primary nitrogen isotopic signature. Besides, even if metamorphic processes in the greenschist

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facies may have slightly shifted the  $\delta^{15}N$  (<2‰), they cannot account for the large range of values observed in the BRC and their rapid stratigraphic variations.

## 5.1 | A biotic organic source of N

Nitrogen-bearing organic molecules produced in organic hazes such as in Titan's atmosphere (tholins) are mentioned as an abiotic alternative to organic matter production during the Archean (Izon et al., 2015; Trainer et al., 2006). However, organic molecules generated through experimental aerosols production setups in plasma or photochemical flow reactors are depleted in <sup>15</sup>N (from 0.8‰ to -25.5‰) compared with the initial N<sub>2</sub> gas from which they precipitated (Kuga et al., 2014; Sebree et al., 2016). If the  $\delta^{15}N_{N2}$  value of the Archean atmosphere was similar to that of the present-day atmosphere ( $\delta^{15}N_{N2} = 0$ %, Nishizawa et al., 2007), then the range of  $\delta^{15}$ N values observed in the BRC is not consistent with a significant contribution of <sup>15</sup>N-depleted abiotic organic matter. The relatively stable TN along the drill core would also mean a sustained abiotic production of these organic molecules, which seems unlikely given the suggested rapid oscillations between hazy and haze-free states evidenced for the Neoarchean atmosphere (Izon et al., 2015; Zerkle et al., 2012). Indeed, oscillations that have only been observed over stratigraphic intervals of a few dozens of meters are hard to reconcile with the 350 m-thick BRC, even if the sedimentation rate remains poorly constrained.

While hydrothermal activity may have been widespread on Archean oceans seafloor and fluid circulations have likely



FIGURE 2 Boxplots showing the  $\delta^{15}$ N and  $\delta^{13}$ C<sub>org</sub> range of values and means between (1) the chert-dominated unit and the BIF-dominated unit, (2) chert samples and siderite samples and (3) areas of the drill core with or without jasper layers.







chert siderite

o areas with small jasper layers



siderite chert o areas with small jasper layers

FIGURE 3 Crossplots: TOC (wt. %) vs. TN (wt. ppm);  $\delta^{15}$ N (‰ vs. air) vs. TN (wt. ppm);  $\delta^{15}$ N (‰ vs. air) vs. TOC/TN.

influenced the availability of nutrients like dissolved iron (Poulton & Canfield, 2011), it seems unlikely that hydrothermal circulations either made a strong contribution of abiotic organic nitrogen to the BRC or that it significantly shifted its bulk  $\delta^{15}$ N signal. Indeed, the contribution of mantellic N to hydrothermal systems seems low compared to hydrothermal remobilization of previously sedimented nitrogen (Stüeken et al., 2021). Neither typical mantellic  $\delta^{15}$ N values around -5‰ (Cartigny & Marty, 2013) nor  $\delta^{15}$ N values measured for altered oceanic crust (ranging from -12% to +8‰, Li et al., 2007) have been found in the BRC samples. Instead, the BRC lowest  $\delta^{15}$ N values are consistent with the range of fractionation displayed by Mo-based biological fixation of N<sub>2</sub> by diazotrophs (Nishizawa et al., 2014; Sigman et al., 2009; Zhang et al., 2014).

Abiotic graphite formation has been identified in highly metamorphosed, siderite-bearing sedimentary rocks of the Eoarchean Isua Supracrustal Belt (van Zuilen et al., 2003). In the BRC, organic matter is not preferentially associated with siderite layers, which suggests that it was not formed by thermal decomposition of siderite.

Therefore, the most probable hypothesis is that nitrogen found in the BRC samples has a biotic source, namely biological N<sub>2</sub> fixation by diazotrophs in the photic zone, followed by organic matter mineralization in the water column or the sediment.

### 5.2 Anoxic and reducing paleoenvironmental conditions

The lithologies of the BRC (mainly chert and siderite), the lack of Ce anomalies (Tice & Lowe, 2006b), and the presence of massindependent fractionation of sulfur isotopes in sulfides (S-MIF, Galić, 2015) are consistent with overall reducing and anoxic conditions. The thin jasper layers within the BIF-dominated unit indicate that iron oxidation may nonetheless have taken place punctually, possibly due to anoxygenic photosynthesis (Tice & Lowe, 2006b) or UV photolysis (Konhauser et al., 2007).

The Archean ocean was likely saturated in silica, which either could have precipitated in the water column or formed silica gels at the sediment surface (Ledevin et al., 2019). The main iron source in Archean oceans is likely Fe(II) provided by hydrothermalism (Poulton & Canfield, 2011). The siderite enrichment in the BIF-dominated unit could either indicate deepening of the water column such that the depositional setting was overlain by Fe(II)-rich bottom waters in a stratified ocean (Tice & Lowe, 2004), or periods of increased hydrothermal activity releasing large quantities of dissolved Fe, indicating a chemical shift from a silica-rich to a carbonate-rich fluid.

## 5.3 | Nitrogen cycling in the Paleoarchean BRC

In the anoxic and ferruginous environmental context of the BRC, nitrites and nitrates would not have been thermodynamically stable. Therefore, the water column nitrogen pool was most probably fixed ammonium (Beaumont & Robert, 1999; Canfield et al., 2010), either released during the degradation of organic matter or released from lithified sediments through fluid circulations (Stüeken et al., 2021).

Nevertheless, nitrification in a stratified ocean with an oxic surface layer has been proposed for some Neoarchean and Paleoproterozoic settings (Garvin et al., 2009; Godfrey & Falkowski, 2009; Kipp et al., 2018; Koehler et al., 2018). While the production of small amounts of nitrates cannot be fully ruled out, it is unlikely that nitrification could have been sustained in the reducing and anoxic conditions of the depositional environment of the Paleoarchean BRC. Indeed, an anoxic atmosphere evidenced by the S-MIF data (Galić, 2015) seems incompatible with an oceanic surface layer hosting a stable nitrate reservoir. In addition, the small range of pyrites  $\delta^{34}$ S values (Galić, 2015) indicate that sulfates were scarce, further supporting the absence of significant concentrations of oxidized anions in the BRC water column. Therefore, without a stable pool of nitrates that could explain the enrichment in <sup>15</sup>N of a residual nitrate pool through denitrification and its subsequent assimilation by primary producers, a mechanism that can fractionate  $NH_4^+$  before its assimilation is required to explain the observed positive nitrogen isotopic signatures.

Partial biological assimilation of NH<sub>4</sub><sup>+</sup> can enrich residual organic matter in <sup>14</sup>N ( $\varepsilon \approx -4\%$  to -27%, Hoch et al., 1992), if the pool of ammonium is not quantitatively consumed, which might be possible in upwelling regions. However, the expected distribution of  $\delta^{15}$ N values should be centered around 0%, displaying both the upwelled <sup>15</sup>N-depleted and the sinking <sup>15</sup>N-enriched pools of ammonium. While this mechanism has been proposed to explain the negative and near zero  $\delta^{15}$ N values found in the Paleoarchean (Ader et al., 2016; Beaumont & Robert, 1999),  $\delta^{15}$ N values in the BRC are centered around 3.3‰ with no negative values. This means that in the BRC depositional environment, an additional mechanism enriching residual ammonium in <sup>15</sup>N was operating at a regional scale.

The hypothesis of positive ammonium  $\delta^{15}N$  values resulting from NH<sub>3</sub> degassing at high pH (Stüeken et al., 2015) (pH>9.2) is compatible with the precipitation of iron-rich carbonates but seems unlikely in the context of the BRC given that positive  $\delta^{15}N$  values are also found in chert layers (Figure 1), which require decreasing pH values (Wright, 2022).

The oxidation of ammonium is therefore the only remaining mechanism capable of producing a  ${}^{15}NH_4^{+}$ -enriched pool that could have been recorded in the BRC organic N after assimilation.

Most metabolic pathways oxidizing  $NH_4^+$  need free  $O_2$ , and even classic anaerobic ammonium oxidation (anammox) requires the presence of nitrites, which cannot be produced without  $O_2$  (Grotzinger & Kasting, 1993). However, in anoxic and ferruginous conditions,  $NH_4^+$  can be oxidized to  $N_2$  in the presence of iron oxides through the Feammox reaction (Stüeken et al., 2016):

$$6 \operatorname{Fe}(OH)_3 + 2 \operatorname{NH}_4^+ + 10 \operatorname{H}^+ \rightarrow 6 \operatorname{Fe}^{2+} + \operatorname{N}_2 + 18 \operatorname{H}_2O$$

This reaction has only recently been described in modern environments, namely in iron-rich conditions in wetland soils and marine sediments (Rios-Del Toro et al., 2018; Yang et al., 2012). Such a metabolism is compatible with Eh-pH conditions and iron speciation during the Archean period, as it is thermodynamically favored when Fe<sup>2+</sup> concentrations are below 100 $\mu$ M and NH<sub>4</sub><sup>+</sup> concentrations are above 1  $\mu$ M, for pH between 6 and 8 (Stüeken et al., 2016). These conditions were probably achieved in the Paleoarchean seafloor, given the available constraints on its pH values around 6.5 to 7 (Halevy & Bachan, 2017). Ammonium and iron concentrations measured in modern anoxic and ferruginous environments (Petrash et al., 2022) and estimated for ancient environments (Canfield, 2005; Tosca et al., 2019) are also in accordance with this hypothesis.

Jasper layers observed in the BRC suggest that iron redox cycling (from anoxygenic photosynthesis or UV photolysis) was occurring and could have provided iron oxides to the Feammox reaction. Ammonium released from biomass degradation during diagenesis could have been in direct contact with iron oxides in the sediment, allowing Feammox to operate while residual <sup>15</sup>N-enriched ammonium could have exchanged isotopically with organic matter or been assimilated by benthic organisms. Feammox is also capable of producing N<sub>2</sub> at rates that can outcompete denitrification in iron-rich tropical soils (Yang et al., 2012). It therefore provides a return nitrogen flux to the atmosphere before the spread of denitrification, which is consistent with a stable atmospheric N<sub>2</sub> reservoir from 3.5 Ga (Nishizawa et al., 2007).

While the fractionation factor for Feammox remains to be determined, all known ammonium oxidation pathways enrich residual  $NH_4^+$  in <sup>15</sup>N. In particular, anammox fractionation can equal that of denitrification (Brunner et al., 2013). The assimilation of the residual <sup>15</sup>N-enriched ammonium into the biomass will then lead to positive sedimentary  $\delta^{15}N$  values (Figure 4).

Such positive values in the BRC are observed where jasper layers are absent, indicating limited iron oxide availability and/or reduction in those iron oxides by reactions with organic matter or sulfide during diagenesis. By contrast, the lowest  $\delta^{15}$ N values are observed in the intervals of the BIF-dominated unit that contain jasper layers, that is, in environments where iron oxides were either abundant or well preserved. The return to a diazotrophic environment could either mean that the Feammox reaction did not take place in settings where waters were saturated in Fe<sup>2+</sup> to concentrations capable of inhibiting this metabolic pathway or that iron oxides were not scavenged by sulfides and abundant enough to allow NH<sub>4</sub><sup>+</sup> to be entirely consumed by the Feammox reaction, leaving no detectable fractionation (Figure 4).

As to organic carbon isotopic signatures,  $\delta^{13}C_{org}$  values in the BIF-dominated unit are centered around -30‰, which is consistent with carbon fixation through the Calvin cycle, a pathway that exists in both chemoautotrophs and photosynthesizers, either anoxygenic or oxygenic. Such values fall in the typical range for photosynthesis,



FIGURE 4 Schematic model illustrating how the iron and nitrogen cycles could have interacted in an anoxic and ferruginous Archean ocean, as recorded in the BRC formation. Following the sedimentary model described by Tice and Lowe (2004), the chert-dominated unit is related to platform settings and gives way to the BIF-dominated unit as depth increases toward basin settings. The water column is chemostratified, with a lower  $Fe^{2+}$ -rich layer supplied by hydrothermal iron fluxes. Siderite precipitates either at the chemocline or in the sediment and accumulates on the seafloor, forming the BIF-dominated unit. The abundance of siderite follows a gradient away from the hydrothermal discharge zones. In the absence of free O<sub>2</sub> in the water column, iron oxides observed in the BRC jasper layers can be produced through photoferrotrophy when dissolved  $Fe^{2+}$  reaches the photic zone. Organic nitrogen is sourced to the water column by diazotrophic N<sub>2</sub> fixation and is either consumed by organisms or ammonified as NH<sub>4</sub><sup>+</sup>. In the chert-dominated unit, iron oxides are scarce and therefore quantitatively consumed by Fearmox. The residual ammonium becomes enriched in <sup>15</sup> N and can carry this enrichment to organic matter when reassimilated, which would produce positive sedimentary  $\delta^{15}N$  values. In the iron-rich settings of the BIF-dominated unit,  $\delta^{15}N$  values are close to 0‰, recording only biological N<sub>2</sub> fixation by diazotrophs. This could either result from quantitative consumption of NH<sub>4</sub><sup>+</sup> by Fearmox due to an excess of iron oxides or to an inhibition of Fearmox by dissolved Fe<sup>2+</sup> concentrations exceeding 100 µM.

likely anoxygenic given other oxidation indicators. The chertdominated unit displays more variable  $\delta^{13}C_{org}$  values, which points toward a larger diversity of metabolisms in shallow depositional environments, possibly within benthic communities of biomats (Tice & Lowe, 2004). Some of the strongly negative values, down to -40‰, are consistent with methanotrophy/methanogenesis, which is all the more interesting because iron oxides can also be used to oxidize methane (Lepot, 2020). These strongly negative  $\delta^{13}C_{org}$  values are thus consistent with Fe(III)-driven oxidation of organic products, including methane and ammonium.

## 6 | CONCLUSION

Carbon and nitrogen isotopic compositions in the 3.4 Ga Buck Reef Chert display a large range of variations. However, while the sedimentary  $\delta^{13}C_{org}$  signal seems to change with the depositional environment or chemical conditions, the bulk sedimentary  $\delta^{15}N$  signal seems to rather primarily reflect iron oxides availability.  $\delta^{15}N$  values are consistently positive and reach up to +6.1‰ in both jasper-free cherts and siderites of the two main sedimentary units, whereas they display a distinctive diazotrophic Mo-based N<sub>2</sub> fixation signature close to 0‰ in the jasper-bearing facies of the upper BIFdominated unit. Positive  $\delta^{15}N$  values observed in sediments that are more scarce in iron could have resulted from the partial oxidation of NH4<sup>+</sup> to N2 through the Feammox reaction, whereas sediments deposited in an environment saturated in Fe<sup>2+</sup> would show no isotopic nitrogen fractionation, either from direct Feammox inhibition or from complete consumption of the ammonium pool. Accordingly, these findings suggest that iron availability could have controlled the nitrogen isotopic signature recorded in sediments through an ammonium oxidative pathway. More importantly, these results call for careful consideration of the meaning of positive  $\delta^{15}N$  signatures in Archean ferruginous sediments. Indeed, while <sup>15</sup>N-enriched sedimentary organic matter has often been used as a redox indicator of a stable pool of nitrates in "oxygen oases," this study shows that in anoxic settings where Fe(III) minerals are present, the oxidation of ammonium can lead to similar positive  $\delta^{15}N$  signatures. As anoxic and ferruginous conditions are thought to have been widespread in the Archean ocean, it might be necessary to carefully consider iron abundance and speciation when interpreting the nitrogen isotopic signatures and to rely on complementary redox indicators.

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## DATA AVAILABILITY STATEMENT

The data that supports the findings of this study are available in the supplementary material of this article.

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## SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.

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