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4 **Title: Limited oxygen production in the Mesoarchean ocean**

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25 performed research; F.OO., A.H., J.E.S., E.E.S., S.W.P., M.B. and A.B. acquired and  
26 analyzed data; F.OO. wrote the paper with significant input from all authors.

27 **Abstract:** The Archean Eon was a time of predominantly anoxic Earth surface conditions,  
28 where anaerobic processes controlled bio-essential element cycles. In contrast to oxygen oases  
29 well documented for the Neoproterozoic (2.8-2.5 billion years ago; Ga), the magnitude, spatial  
30 extent, and underlying causes of possible Mesoproterozoic (3.2-2.8 Ga) surface ocean oxygenation  
31 remain controversial. Here, we report  $\delta^{15}\text{N}$  and  $\delta^{13}\text{C}$  values coupled with local seawater redox  
32 data for Mesoproterozoic shales of the Mozaan Group (Pongola Supergroup, South Africa), which  
33 were deposited during an episode of enhanced Mn (oxyhydr)oxide precipitation between ~2.95  
34 and 2.85 Ga. Iron and Mn redox systematics are consistent with an oxygen oasis in the  
35 Mesoproterozoic anoxic ocean, but  $\delta^{15}\text{N}$  data indicate a Mo-based diazotrophic biosphere with no  
36 compelling evidence for a significant aerobic nitrogen cycle. We propose that in contrast to the  
37 Neoproterozoic, dissolved  $\text{O}_2$  levels were either too low or too limited in extent to develop a large  
38 and stable nitrate reservoir in the Mesoproterozoic ocean. Since biological  $\text{N}_2$  fixation was  
39 evidently active in this environment, the growth and proliferation of  $\text{O}_2$ -producing organisms  
40 were likely suppressed by nutrients other than nitrogen (e.g., phosphorus), which would have  
41 limited the expansion of oxygenated conditions during the Mesoproterozoic.

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43 **Keywords:** oxygen oasis; nitrogen isotopes; nutrient limitation; oxygenic photosynthesis;  
44 Mesoproterozoic

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46 **Significance Statement:** Episodic development of “oxygen oases” during the Archean Eon  
47 characterizes the hundreds of millions of years transition to permanent oxygenation in the  
48 atmosphere-hydrosphere system at the Great Oxidation Event (~2.4-2.3 Ga). One of these well-  
49 characterized “oxygen oases” is recorded in Mesoproterozoic (~2.95-2.85 Ga) sediments of the  
50 Pongola Supergroup. We show that in contrast to the Neoproterozoic (2.8-2.5 Ga), biological  
51 oxygen production in a shallow ocean having Mo-based nitrogen fixation was not sufficient to  
52 result in a dissolved nitrogen reservoir that would carry the isotopic effects of an aerobic  
53 nitrogen cycle. Nevertheless, it appears that low concentrations of bioavailable phosphorus,  
54 rather than nitrogen, suppressed the growth and expansion of oxygenic photosynthesizers, and  
55 may explain why pervasive and permanent oxygenation was delayed during the Archean Eon.

56  
57 **body**

58 A dramatic rise in atmospheric oxygen level during the Great Oxidation Event (GOE) at ~2.4  
59 Ga is marked by the disappearance of mass-independent fractionation of sulfur isotopes,  
60 oxidation of detrital pyrite and uraninite, and the appearance of red beds, reflecting the  
61 irreversible transition from an anoxic to an oxic world (1-2). While it is widely accepted that  
62 oxygenic photosynthesis was a first-order control on the GOE (3), Archean shallow-marine  
63 “oxygen oases” and “whiffs” of atmospheric oxygen ( $\text{O}_2$ ) have been proposed to have occurred  
64 up to several hundred million years prior to the GOE (4-18). However, while processes that  
65 drove oxygen production during transient and localized oxygenation events in the Neoproterozoic  
66 (2.8-2.5 Ga) are supported by a wide range of geochemical proxies (e.g., 4-6, 13-18), those  
67 from the Mesoproterozoic (3.2-2.8 Ga) are constrained by only a limited number of studies (7-12).

68 The nitrogen (N) cycle from the early Archean up to ~2.7 Ga is widely considered to have been  
69 dominated by bioavailable ammonia ( $\text{NH}_4^+$ ) under anoxic water column conditions (15, 16).  
70 Oxidation of  $\text{NH}_4^+$  would have been suppressed in an early Archean ocean characterized by  
71 extremely low  $\text{O}_2$  concentrations (15-17). Free  $\text{O}_2$  is produced through oxygenic  
72 photosynthesis, the rate of which is mainly controlled by the concentrations of bioavailable N  
73 and phosphorus (P) (19-24). While the sedimentary  $\delta^{15}\text{N}$  record suggests that N was  
74 bioavailable and that diazotrophic Mo-based nitrogenase dominated  $\text{N}_2$  fixation in the  
75 Mesoarchean, the record also places a robust minimum age for the occurrence of aerobic N  
76 cycling at ~2.72 Ga in the Neoproterozoic (e.g., 15, 16 and references therein). Indeed, prominent  
77 N isotope excursions in the Neoproterozoic provide evidence for temporary  $\text{NH}_4^+$  oxidation, and  
78 thus the  $\delta^{15}\text{N}$  record has been used to infer the development of locally oxygenated surface  
79 ocean environments after ~2.7 Ga (5, 15, 17, 18, 25).

80 Independently, stable isotope systematics of redox-sensitive elements such as Fe, Mo, U and  
81 S, as well as locally enhanced Mn (oxyhydr)oxide precipitation, support an earlier emergence  
82 of oxygenic photosynthesis and episodic development of “oxygen oases” in the Mesoarchean  
83 surface ocean (7-9, 11), well before currently accepted evidence for oxidative nitrogen cycling.  
84 Furthermore, phylogenomic estimates based on molecular clocks also suggest that  
85 cyanobacterial stems capable of oxygenic photosynthesis might find their roots in the Archean,  
86 with a development and progressive diversification starting as early as ~3.5 Ga (26-28).  
87 However, the factors that caused a delay in pervasive oxygenation of the atmosphere-  
88 hydrosphere system after the establishment of oxygenic photosynthesis in the early Archean  
89 remain poorly constrained, particularly with regard to the role of their two main bio-limiting  
90 nutrients, N and P (19-23). Modelling studies have demonstrated that low dissolved P  
91 concentrations would severely suppress the rate of oxygenic photosynthesis and ultimately the  
92 spatial extent of Archean oxygen oases (29). However, there is currently no consensus on  
93 dissolved P concentrations in the Archean ocean (21-23, 30-32).

94 In order to assess controls on the spatial development and intensity of Earth’s first oxygen  
95 oases, we measured nitrogen ( $\delta^{15}\text{N}$ ) and organic carbon ( $\delta^{13}\text{C}_{\text{org}}$ ) isotopes, local water column  
96 redox proxies (Fe speciation and Mn concentrations), and elemental data for shales of the  
97 ~2.95-2.85 Ga Mozaan Group, Pongola Supergroup, South Africa (see Supplementary  
98 Information for geologic setting and all data). Our aim is to clarify the factors that controlled  
99 the nature and development of oxygen oases in the Mesoarchean.

100

## 101 **Results and discussion**

102 **Water column redox reconstruction.** Our samples span a shallow-marine (above wave base)  
103 depositional setting in the White Mfolozi Inlier, to a deeper-water (below wave base)  
104 equivalent in the Nongoma area, and comprise three sequences deposited at different water  
105 depths (Figs. 1, S1, S2; Table S1). In the White Mfolozi Inlier, sequence I, deposited in the  
106 most proximal, intertidal to shallow subtidal setting, is characterized by high Mn contents and  
107 Mn/Fe ratios compared to average values for shales of the Pongola Supergroup (33; Table S1),  
108 mostly high ratios of highly-reactive Fe to total Fe (FeHR/FeT) and high Fe/Al ratios (see  
109 Methods for detailed analytical techniques). Sequence II was deposited in a deep subtidal, but  
110 above fair-weather wave base setting, and shows a progressive decrease in Mn, Mn/Fe,  
111 FeHR/FeT and Fe/Al, while Mn and Fe contents are higher than in average Pongola shales (33;  
112 Table S1). The uppermost sequence III represents deepening to between fair-weather and storm  
113 wave base, and is characterized by persistently low Mn, Mn/Fe, FeHR/FeT and Fe/Al, with Mn  
114 and Fe contents similar to those in average Pongola shales (33; Table S1). In the more distal,  
115 deeper-water setting of the Nongoma area, where distinct compositional trends were not  
116 observed, Fe/Al ratios tend to be high, but Mn contents remain low and Mn/Fe ratios shift to  
117 values lower than the average for shales of the Pongola Supergroup (33; Table S1).

118 To explain these data, we invoke upwelling of anoxic waters that were rich in Fe<sup>2+</sup> and Mn<sup>2+</sup>  
119 into oxic shallow waters. Precipitation of Fe as (oxyhydr)oxide minerals may have started  
120 under low oxygen or anoxic conditions, potentially via photoferrotrophy in shallower waters  
121 directly overlying deeper anoxic waters (34), and this likely explains the observed Fe  
122 enrichments in the distal Nongoma setting (Fig. 1b). Fe(II) oxidation would have been  
123 progressive during upwelling, leading to increased FeHR/FeT and Fe/Al enrichments (Fig. 1a)  
124 as water depth shallowed through a redoxcline (as captured by sequence II in the White Mfolozi  
125 Inlier) into the shallow and locally oxygenated waters of sequence I, where Mn(IV)  
126 (oxyhydr)oxides precipitated (Fig. 1a; Fig 2). Increased Mn/Fe ratios in shallower waters thus  
127 reflect progressive removal of dissolved Fe(II) and/or enhanced precipitation of Mn  
128 (oxyhydr)oxides as upwelling anoxic waters reached the redox threshold for Mn(II) oxidation.  
129 However, sequence III in the White Mfolozi Inlier has Mn/Fe ratios similar to the average value  
130 for shales of the Pongola Supergroup, with no evidence for FeHR enrichment, likely reflecting  
131 deeper anoxic waters where there was limited oxidant availability to promote Fe- or Mn-

132 (oxyhydr)oxide precipitation in the water column. At first glance, Fe enrichments in shallower  
133 waters and their absence in deeper waters of the White Mfolozi Inlier may appear contradictory,  
134 since Fe enrichments are commonly taken to denote water column anoxia (35). However, our  
135 data are entirely consistent with current understanding of how Fe enrichment may be enhanced  
136 under anoxic ferruginous conditions, whereby one prominent pathway for developing high  
137 FeHR/FeT and Fe/Al ratios is via upwelling of deep, anoxic waters into shallower oxic settings  
138 (35, 36).

139 Once Mn and Fe (oxyhydr)oxides had formed and became deposited, they were then largely  
140 converted to carbonate minerals through microbial respiration during early diagenesis, as  
141 indicated by high Fe<sub>carb</sub> concentrations in sediments of sequence I and II (Table S1). In support  
142 of this, highly negative  $\delta^{13}\text{C}$  (between  $-22$  and  $-13\%$ , VPDB) and  $\delta^{18}\text{O}$  values (between  $-21$   
143 and  $-8\%$ , VPDB) indicate carbonate precipitation through organic carbon remineralization  
144 during diagenesis (7, 37). This happened below the sediment-water interface in sediments  
145 deposited below a water column characterized by relatively high rates of organic carbon (OC)  
146 burial (high productivity) (7). In contrast, Fe<sub>carb</sub> is scarce in the deeper water sequence III and  
147 the more distal Nongoma setting (Table S1), where instead most of the Fe is associated with  
148 chlorite and stilpnomelane (7). It is thus likely that Fe (oxyhydr)oxides were converted to Fe-  
149 rich clay minerals during diagenesis in this setting, probably via reverse weathering (38).  
150 Another possibility involves conversion of Fe (oxyhydr)oxides into mixed ferrous/ferric phases  
151 such as green rust during settling (39, 40), before their final transformation to stilpnomelane  
152 and chlorite during diagenesis and metamorphism. Thermodynamic estimates based on the  
153 chemical composition of Fe-chlorite showed that Fe- and Mn-rich clay minerals of the Mozaan  
154 Group formed during diagenesis and metamorphism (37). Regardless of the precise nature of  
155 precursor Fe minerals, Fe/Al ratios much higher than those in average Pongola shales (33;  
156 Table S1) indicate that their precipitation gave rise to significant Fe enrichments in the deep-  
157 water sediments, and during upwelling of deep ferruginous waters into shallower oxic settings.  
158 The  $\delta^{13}\text{C}_{\text{org}}$  values average  $-27.6\%$  in the shallow-water sequence I samples, reflecting isotopic  
159 fractionations expected during autotrophic  $\text{CO}_2$  fixation (41). During deposition of sequence  
160 II,  $\delta^{13}\text{C}_{\text{org}}$  values progressively decrease to the average value of  $-30\%$ , and down to  $-38\%$  in  
161 the deep-water settings of the White Mfolozi Inlier and Nongoma areas (Fig. 1). The highly  
162 negative  $\delta^{13}\text{C}_{\text{org}}$  values in these deeper-water, ferruginous settings likely reflect biological  
163 carbon cycling with a significant contribution from methanogens and methanotrophs (42). The  
164 variability in biological processes with water-depth might be linked to the water column redox

165 gradient, where (1) high Mn/Fe ratios and Mn(II) oxidation (which requires free O<sub>2</sub>) are  
166 consistent with photoautotrophic CO<sub>2</sub> fixation and oxygenic photosynthesis in the shallow-  
167 water settings, and (2) Fe enrichments without Mn(II) oxidation (Mn/Fe ratios lower than the  
168 average Pongola shale values) are consistent with methanotrophs utilizing dissolved O<sub>2</sub> or  
169 Fe(III) compounds to oxidize methane at the redoxcline or chemocline, respectively, under  
170 hypoxic or anoxic conditions in deeper-water settings. In view of this, the water column  
171 appears to have been both redox and ecologically stratified.

172

173 **N isotope systematics and preservation of primary isotopic signals.** Our geochemical data  
174 suggest a shallow-water oxygen oasis in the Mesoarchean Pongola sea at ~2.9 Ga. If these  
175 conditions were stable and extensive enough to support oxic nitrogen metabolism, then this  
176 should be reflected in nitrogen isotope systematics, as observed in younger Neoproterozoic  
177 sedimentary successions (5, 15, 17, 18, 25). Large N isotope heterogeneity revealed by the  
178 NanoSIMS technique in isolated microfossils from the ~3.0 Ga Farrel Quartzite (Western  
179 Australia) has been linked to biological aerobic nitrification (43), indicating the emergence of  
180 this metabolic pathway even before deposition of the Mozaan Group. In ancient marine  
181 sediments,  $\delta^{15}\text{N}$  values between  $-4$  and  $+2\text{‰}$  (Air-N<sub>2</sub>) are usually attributed to isotopic  
182 fractionation imparted during biological N<sub>2</sub> fixation using the Mo-nitrogenase enzyme (Nif; 5,  
183 15-18, 25). The use of V- (Vnf) and Fe-based (Anf) alternative nitrogenase enzymes produces  
184 more depleted  $\delta^{15}\text{N}$  values, between  $-6$  and  $-8\text{‰}$  (15, 44).  $\delta^{15}\text{N}$  values above  $+4\text{‰}$  would  
185 provide compelling evidence for an aerobic N cycle coupling nitrification and  
186 denitrification/anammox processes (*e.g.*, 5, 15, 17, 18, 25). Nitrogen isotope values for 18 out  
187 of 22 samples fall in the range of  $-5$  to  $+3\text{‰}$  (Air-N<sub>2</sub>), and reflect isotopic fractionation driven  
188 by Mo-based diazotrophy (Fig. 1; Table S2). Positive values above  $+4\text{‰}$  are limited to 4  
189 samples, including 2 from the White Mfolozi Inlier and 2 others from the more distal Nongoma  
190 area.

191 Here, we exclude abiotic sources for bioavailable nitrogen, because they were probably too  
192 small in magnitude and should have otherwise dominated the early Precambrian  $\delta^{15}\text{N}$  record,  
193 counter to what is observed (15). However, several mechanisms can alter the original  $\delta^{15}\text{N}$   
194 values of marine biomass, ranging from early diagenesis to the thermal degradation of organic  
195 matter (OM) during deeper burial diagenesis and metamorphism (15, 45-48). The redox state  
196 of the water column, sedimentation rate, and OM accumulation can also impart different N

197 isotope fractionations between sinking organic particles and surficial marine sediments.  $\text{NH}_4^+$   
198 release during OM remineralization below the sediment-water interface and partial oxidation  
199 in pore waters can increase bulk sediment  $\delta^{15}\text{N}$  ( $\delta^{15}\text{N}_{\text{bulk}}$ ) values by  $\sim 4\%$  under oxic diagenetic  
200 conditions, while this effect tends to be minimal during anoxic diagenesis, with an isotopic  
201 fractionation  $< 1\%$  (15, 45). The predominance of Fe- and Mn-carbonate minerals derived from  
202 the reduction of Fe- and Mn-(oxyhydr)oxides indicate anoxic diagenetic conditions (7, 37) and  
203 thus likely a minimal effect of early diagenetic processes on primary  $\delta^{15}\text{N}$  values. Importantly,  
204 oxic diagenesis would result in isotopic compositions that reflect an aerobic nitrogen cycle,  
205 which is not seen in our dataset.

206 Organic-bound  $\text{NH}_4^+$  can also be released through thermal devolatilization of organic matter  
207 during burial diagenesis and metamorphism, resulting in a maximum increase in  $\delta^{15}\text{N}_{\text{bulk}}$  values  
208 of 1–2‰ at greenschist facies, 3–4‰ at lower amphibolite facies, and up to 6–12‰ at upper  
209 amphibolite facies; even larger offsets can occur in sedimentary rocks affected by circulating  
210 fluids (46). The Mozaan Group experienced lower greenschist facies metamorphism (37),  
211 suggesting a maximum increase in  $\delta^{15}\text{N}_{\text{bulk}}$  values of less than 2‰. In order to alleviate  
212 potential effects caused by mechanisms described above on  $\delta^{15}\text{N}$  values, N isotope data were  
213 also measured on extracted kerogen ( $\delta^{15}\text{N}_{\text{ker}}$ ) to compare with bulk sample data ( $\delta^{15}\text{N}_{\text{bulk}}$ ). The  
214 offset between  $\delta^{15}\text{N}_{\text{bulk}}$  and  $\delta^{15}\text{N}_{\text{ker}}$  values allows evaluation of the extent of preservation of the  
215 N isotope signature imparted by the initially deposited biomass. Two samples from the  
216 Nongoma area showing evidence for hydrothermal processes are characterized by large offsets  
217 between  $\delta^{15}\text{N}_{\text{bulk}}$  and  $\delta^{15}\text{N}_{\text{ker}}$  values and very positive N isotope values (Fig. 1; Table S2), which  
218 likely supports post-depositional alteration by circulating fluids. Therefore, these 2 samples  
219 will not be further considered in this study. In contrast, the minimal offset in most of the studied  
220 samples supports good preservation of the primary isotopic signature (Fig. 1; Table S2). The  
221 samples with minimal offset between  $\delta^{15}\text{N}_{\text{bulk}}$  and  $\delta^{15}\text{N}_{\text{ker}}$  values also lack evidence of  
222 secondary alteration by later circulating fluids or hydrothermal processes (37). Furthermore, a  
223 minimal effect of post-depositional processes on the isotope composition of biomass is also  
224 indicated by the absence of co-variation among  $\delta^{15}\text{N}$  and TN,  $\delta^{15}\text{N}$  and C/N,  $\delta^{15}\text{N}$  and  $\delta^{13}\text{C}$ , as  
225 well as between  $\delta^{13}\text{C}_{\text{TOC}}$  and TOC for bulk sediments (Fig. 3). A weak negative co-variation  
226 between  $\Delta^{15}\text{N}_{\text{ker-bulk}}$  (the difference between  $\delta^{15}\text{N}_{\text{ker}}$  and  $\delta^{15}\text{N}_{\text{bulk}}$ ) and total K further supports  
227 a minimal contribution of ammoniated phyllosilicates (*e.g.*,  $\text{NH}_4^+$  substituted for  $\text{K}^+$ ) with a  
228 distinct isotopic composition (see Fig. S4).

229

230 **A lack of evidence for aerobic nitrogen cycling in the Mesoproterozoic Pongola basin oxygen**  
231 **oasis.** In the modern ocean, where the main processes intrinsic to the aerobic N cycle, including  
232 N<sub>2</sub> fixation, nitrification, and denitrification/anammox, are at play,  $\delta^{15}\text{N}$  values of +5 to +7‰  
233 in sedimentary organic matter reflect <sup>14</sup>N loss to the atmosphere through denitrification and  
234 anammox in oxygen-minimum zones (48). Buried biomass is an indirect archive of this  
235 process, because organisms assimilate the isotopically heavy nitrate as a nutrient. Therefore,  
236  $\delta^{15}\text{N}$  values  $\geq +4\text{‰}$  found in Neoproterozoic marine sediments are interpreted to reflect temporary  
237 aerobic N cycling (*e.g.*, 5, 15, 17, 18, 25). In contrast, the nitrogen isotope data of the Mozaan  
238 Group are inconsistent with the establishment of a significant aerobic nitrogen cycle. Assuming  
239 that Mn(II) oxidation occurs at higher redox potential than NH<sub>4</sub><sup>+</sup> oxidation, redox conditions  
240 may have been sufficient for evolved nitrifying bacteria in the Pongola basin, because high Mn  
241 concentrations in sequence I indicate the precipitation of Mn(IV) oxyhydroxide minerals (7),  
242 which required O<sub>2</sub>. Photochemical oxidation of Mn is inhibited under Fe-saturated conditions  
243 (49) and, unlike Fe(II), significant oxidation of Mn requires oxygen and a catalyst (50). It is  
244 thus likely that free O<sub>2</sub> was locally available in the water column during deposition of sequence  
245 I (7) and that the subtle increase in  $\delta^{15}\text{N}_{\text{ker}}$  values (up to +5.2‰) measured in 2 samples from  
246 the upper part of sequence I (deposited at the redoxcline) may reflect  
247 nitrification/denitrification and uptake of residual nitrate in the water column (Figs. 2).  
248 However, the absence of a more compelling isotopic shift in  $\delta^{15}\text{N}$  values over the extended  
249 stratigraphic interval indicates that NO<sub>3</sub><sup>-</sup> (*i.e.* the residual nitrogen species that carries the  
250 isotopic information in the modern ocean) did not build up to high enough concentrations to be  
251 an important nitrogen source to the biosphere.

252 While the iron and manganese proxies have the capacity to promptly respond to redox  
253 perturbations on a local scale, the nitrogen isotope proxy requires the build-up of a dissolved  
254 nitrogen reservoir that carries the isotopic effects of redox reactions at the ecosystem scale. A  
255 good modern analog illustrating such a discrepancy between Mn (higher redox potential) and  
256 N (lower redox potential) cycles can be found in the Black Sea. Here, rapid oxygen-dependent  
257 microbial Mn(II) oxidation is observed at low micromolar (< 3-5  $\mu\text{M}$ ) dissolved oxygen  
258 contents in the suboxic zone of the Black Sea (51), where nitrification produces a maximum  
259 nitrate concentration of only 3.5  $\mu\text{M}$  (52). However, this maximum nitrate level appears to be  
260 too low to leave an isotopic signature of aerobic N cycling (53) (in contrast to open ocean  
261 nitrate concentrations of up to ~35  $\mu\text{M}$ , which leave an average  $\delta^{15}\text{N}$  signal of around +5‰;  
262 *ref 15 and references therein*). Estimates of O<sub>2</sub> content based on  $\delta^{56}\text{Fe}$  variations in



263 Mesoarchean oxygen oases suggest a maximum concentration of 10  $\mu\text{M}$  (7, 11). Such dissolved  
264  $\text{O}_2$  levels are thus consistent with the potential activity of both Mn(II) oxidizing and nitrifying  
265 bacteria during deposition of the Mozaan Group.

266 It appears that the geographical extent of oxygen oases was likely too restricted in the  
267 Mesoarchean ocean to develop a nitrate reservoir that was large enough to leave an isotopic  
268 signature, which contrasts with their Neoproterozoic equivalents (assuming water-column  $\text{O}_2$   
269 concentrations were similar in the Mesoarchean and Neoproterozoic oxygen oases; 7, 11, 29).  
270 Overwhelming supply of reducing inputs (e.g., Fe(II) and Mn(II)) from submarine volcanism  
271 to the Mesoarchean ocean could have suppressed more widespread oxygenation and thus  
272 limited nitrification. However, iron formation (IF) secular records (e.g., 54, 55) indicate that  
273 volumetrically, the Neoproterozoic IFs are much larger than their Mesoarchean analogs, and yet  
274 the Neoproterozoic to early Paleoproterozoic IFs are characterized by compelling evidence for  
275 aerobic N cycling (5, 25, 56). It therefore appears that the reducing sinks from submarine  
276 volcanism were not the main driving factor that suppressed the expression of aerobic N cycle  
277 in the Mesoarchean ocean.

278

279 **Implications for oxygenic photosynthesis in the Mesoarchean ocean.** Overall, our data  
280 reveal an ecosystem that was dominated by Mo-based diazotrophy, in an oxygen oasis where  
281 a combination of the restricted spatial extent and low dissolved  $\text{O}_2$  concentration likely limited  
282 the build-up of a sufficient nitrate reservoir to impart an isotopic expression of aerobic N  
283 cycling. In the modern ocean, cyanobacteria are the main  $\text{N}_2$  fixers (57), and our data suggest  
284 that this relationship may extend back to the Mesoarchean. Mo-based diazotrophy requires  
285 soluble  $\text{MoO}_4^{2-}$  availability in the ocean. In the modern ocean, Mo is mainly delivered via  
286 riverine inputs following oxidative continental weathering, with a minor contribution from  
287 submarine hydrothermal systems (see ref. 16 and references therein). The mild Mo enrichment  
288 (relative to average concentration for the upper crust) recorded by the Mozaan Group (Table  
289 S1) suggests that dissolved Mo was available in this oxygen oasis (8). However, it has been  
290 shown that a very low Mo content (down to 1 nM, which is ca. 1% of modern seawater  
291 concentrations) can sustain Mo-based diazotrophy in modern environments (58). In view of  
292 this, submarine hydrothermal Mo inputs could have been sufficient to sustain Mo-based  $\text{N}_2$   
293 fixation in the Archean ocean (see ref. 16 and references therein). Moreover, it has also been  
294 demonstrated that continental Mo could have been mobilized and delivered to the ocean even

295 under Archean anoxic atmospheric conditions ( $O_2$  concentration  $< 10^{-5}$  PAL; 59, 60 and  
296 references therein).

297 Free  $O_2$  produced by oxygenic photosynthesis had an impact on the water-column redox and  
298 ecological gradients of the Pongola basin, as observed in stable isotope data confirming aerobic  
299 Fe and Mn cycling in shallow-water settings (7). Bioavailable N and P are the main nutrients  
300 that control marine productivity over time (*e.g.*, 19-23). Their scarcity may have limited  
301 biological  $O_2$  production, resulting in delayed pervasive and permanent oxygenation of the  
302 atmosphere-hydrosphere system after the emergence of oxygenic photosynthesis in the  
303 Mesoarchean. Since our  $\delta^{15}N$  data indicate that N was bioavailable in the Mesoarchean marine  
304 oasis, P scarcity could have been the main limiting factor in biological  $O_2$  production,  
305 consistent with previous biogeochemical modelling (29). Indeed, we observe very low P  
306 contents in the Pongola sediments (Fig. 1; Table S1), which would be entirely consistent with  
307 the suggestion of widespread P limitation under global ferruginous conditions (21), prior to  
308 more extensive anoxic P recycling linked to the build-up of seawater sulfate following more  
309 expansive environmental oxygenation (22).

310 It is possible that positive  $\delta^{15}N$  values in the ca. 3.2 Ga riverine deposits of the Moodies Group,  
311 South Africa, interpreted as evidence for denitrification (12), and a weakly oxidizing U cycle  
312 in the Mesoarchean ocean (11, 61), may reflect episodes of mild, local oxidizing conditions in  
313 the atmosphere-hydrosphere system (ref. 13 and references therein). However, estimates based  
314 on preserved Mesoarchean detrital uraninite in the Witwatersrand Supergroup (South Africa)  
315 deposited contemporaneous with the Pongola Supergroup suggest that atmospheric  $O_2$   
316 concentrations were lower than  $3.2 \times 10^{-5}$  atm (62). Furthermore, the general absence of  $\delta^{15}N$   
317 values above +4‰ in most Mesoarchean marine and continental deposits around the world (15,  
318 16, 63; this study), is consistent with the view that low rates of biological  $O_2$  production limited  
319 the geographical extent of oases and, ultimately, controlled the size of the seawater nitrate  
320 reservoir, which did not reach the level necessary to leave a more widespread and persistent  
321 isotopic signature of an aerobic N cycle. Regardless of the mechanism/s that controlled  
322 dissolved P concentrations in Archean oceans, oxygenation of Earth's early biosphere was  
323 apparently limited by a low supply of bio-available P, rather than N, under anoxic to very low-  
324 oxygen surface conditions.

325

326 **Methods**

327 **Major and trace elements.** Powdered samples were analysed for major element  
328 concentrations by X-ray fluorescence spectroscopy. Analysis was carried out on fusion beads,  
329 using a PANalytical MagiX Pro PW2540 spectrometer at the University of Johannesburg.  
330 Accuracy was checked with certified reference materials and was better than 1%. Elemental  
331 concentrations are reported in wt.% with a detection limit of 0.04 wt.%. Trace elements were  
332 measured at the Isotope Geochemistry Lab, University of Tuebingen (Germany) according to  
333 the analytical procedure previously described (10, 64). Around 30 mg of ashed powdered  
334 samples (heated to 600°C for 12 hours to ash organic compounds) were dissolved using a mix  
335 of concentrated and distilled HF (2 mL) and HNO<sub>3</sub> (0.3 mL) in screw-top 15 mL Savillex®  
336 PFA beakers at 120°C for 4 days. After evaporation at 80°C, samples were taken up in 1.5 mL  
337 6 M HCl and re-dissolved in closed beakers at 130°C for 1 day. The samples were evaporated  
338 to incipient dryness at 90°C and reacted twice with 0.3 mL aliquots of concentrated HNO<sub>3</sub> with  
339 evaporation at 90°C in between to remove excess F and Cl. An aliquot of 5 M HNO<sub>3</sub> (1 mL)  
340 was added to the sample residues and heated at 80°C for ~1 hour for dissolving the samples.  
341 Analyses were performed in the iCap-Qc ICP-MS coupled to an ESI SC-2 DX auto-sampler  
342 with an ESI Fast uptake system equipped with a 4 mL sample loop. For analysis, solution  
343 samples in 1 mL 5 M HNO<sub>3</sub> were diluted twice; first in MQ water (dilution factor 1000) and  
344 second in an internal standard solution made of 0.3 M HNO<sub>3</sub> (dilution factor 10,000) containing  
345 a spike mixture of <sup>6</sup>Li (~3 ppb), In (~1 ppb), Re (~1 ppb), and Bi (~1 ppb). Analytical accuracy,  
346 estimated from the 1 r.s.d. of the mean, varied between 3 and 15% and was monitored by  
347 repeated measurements of reference materials OU-6, QS-1, W-2a, and AGV-2. Enrichment  
348 factors were calculated as  $(\text{element}/\text{Al})_{\text{sample}}/(\text{element}/\text{Al})_{\text{reference}}$  using the average  
349 concentrations for the upper crust as reference (65).

350 **Iron speciation analysis.** Iron speciation analysis was performed at the University of Leeds,  
351 UK using a calibrated sequential extraction protocol followed by Fe analysis via AAS (66).  
352 This method is designed to quantify four different pools of Fe considered to be highly-reactive  
353 (FeHR) towards H<sub>2</sub>S in surface and near-surface environments: (1) pyrite S extracted via Cr-  
354 reduction followed by trapping as Ag<sub>2</sub>S, with Fe calculated assuming an FeS<sub>2</sub> stoichiometry  
355 (Fe<sub>Py</sub>); (2) carbonate-associated iron extracted with a sodium acetate solution (Fe<sub>Carb</sub>); (3) ferric  
356 oxides extracted with a dithionite solution (Fe<sub>Ox</sub>); and (4) mixed-valence iron oxides,  
357 principally magnetite, extracted using ammonium oxalate (Fe<sub>Mag</sub>). The total Fe content in  
358 ancient marine shales (Fe<sub>T</sub>) represents the sum of FeHR and Fe bound in silicates (66, 67). It  
359 has been established that marine shales deposited under oxic water column conditions are

360 characterized by low ratios of FeHR/FeT < 0.22 driven by the lack of Fe mineral precipitation  
361 following transport of Fe(II) under anoxic water column conditions. Under anoxic water  
362 column conditions, FeHR/FeT ratios tend to be higher (above 0.38), whereas values between  
363 0.22 and 0.38 are considered equivocal due to additional processes (*e.g.*, rapid sedimentation)  
364 that may obscure Fe enrichments under anoxic water column conditions (35). Furthermore, for  
365 anoxic water column conditions with FeHR/FeT > 0.38, FePy/FeHR < 0.7 represents  
366 ferruginous conditions, whereas FePy/FeHR values above 0.7–0.8 characterize euxinic  
367 conditions (35, 67). In some cases, FeHR can be converted to poorly reactive silicates (FePRS)  
368 during diagenesis resulting in apparently low FeHR/FeT ratios, which can be wrongly  
369 interpreted to reflect oxic water column conditions (35, 67). Therefore, it is required to pair  
370 FeT/Al ratios and FePRS concentrations in order to distinguish samples deposited under oxic  
371 from those deposited under anoxic water column conditions, where FeT/Al ≤ 0.55 ± 0.11 would  
372 indicate oxic water column conditions (67). FeT/Al > 0.66 would reflect local Fe enrichments  
373 either under anoxic water column conditions, or due to input of anoxic hydrothermal fluids into  
374 oxic seawater (67).

375 **Carbon and nitrogen isotope analyses.** The nitrogen isotope composition of bulk rock  
376 ( $\delta^{15}\text{N}_{\text{bulk}}$ ) and the carbon and nitrogen isotope compositions of the extracted kerogen ( $\delta^{13}\text{C}_{\text{org}}$ ,  
377  $\delta^{15}\text{N}_{\text{ker}}$ ) were determined by elemental analysis/isotope ratio mass spectrometry (EA/IRMS) at  
378 the Institute of Earth Surface Dynamics of the University of Lausanne (Switzerland), using a  
379 Carlo Erba 1108 (Fisons Instruments, Milan, Italy) elemental analyzer connected to a Delta V  
380 Plus isotope ratio mass spectrometer via a ConFlo III open split interface (both of Thermo  
381 Fisher Scientific, Bremen, Germany) operated under continuous helium flow (68, 69). The  
382 kerogen, the fraction of organic matter insoluble in organic solvents, non-oxidizing acids and  
383 bases, dispersed in lithified sediments, was isolated through several steps. These steps are based  
384 on the dichloromethane-extraction procedure followed by HF–HCl treatment at the Institute of  
385 Earth Surface Dynamics, University of Lausanne. The procedure modified from Durand and  
386 Nicaise (70) involved Soxhlet extraction with a mixture of methanol and dichloromethane for  
387 removal of the soluble organic fraction (bitumen), removal of carbonates (as well as sulfides,  
388 sulfates, oxides and hydroxides) by treatment with 6N HCl, removal of silicates by treatment  
389 with a mixture of 40% HF and 6N HCl. The acid treatments were done at 65–70°C while stirring  
390 with a PTFE coated magnetic stirrer. The solid residue was thoroughly washed with warm  
391 deionized water and water purified with a Direct-Q UV3 Millipore® System, and then dried at  
392 40°C. Mineralogical analysis of the dried solid residues was conducted at the Institute of Earth

393 Sciences of the University of Lausanne using a Thermo Scientific ARL X-TRA Diffractometer,  
394 and confirmed the complete elimination of silicates (mainly clay minerals), which constituted  
395 up to 35% of the bulk rock samples before the HCl-HF treatment.

396 The stable isotope compositions are reported in the delta ( $\delta$ ) notation defined as

$$397 \quad \delta^i E_{sample/standard} = \frac{R(^i E / ^j E)_{sample}}{R(^i E / ^j E)_{standard}} - 1$$

398 where  $R$  is the molar ratio of the heavy ( $^i E$ ) to light ( $^j E$ ) isotope of chemical element  $E$  (*e.g.*,  
399  $^{13}\text{C}/^{12}\text{C}$  or  $^{15}\text{N}/^{14}\text{N}$ ). The standard for carbon isotope ratios ( $\delta^{13}\text{C}$ ) is Vienna Pee Dee  
400 Belemnite limestone (VPDB), for nitrogen isotope ratios ( $\delta^{15}\text{N}$ ) is molecular nitrogen in air  
401 (Air- $\text{N}_2$ ). For calibration and normalization of the measured isotopic ratios to the  
402 international scales, a 3- or 4-point calibration was used with international reference materials  
403 (RMs) and UNIL (University of Lausanne) in-house standards. The  $\delta^{13}\text{C}$  and  $\delta^{15}\text{N}$  values of  
404 the in-house standards were normalized with the RMs USGS64, USGS65 and USGS66. The  
405 RMs used together with the UNIL-standards for normalization and quality control of the  
406 measured  $\delta^{13}\text{C}$  and  $\delta^{15}\text{N}$  values were USGS-24, USGS-40, USGS-41 and IAEA-600.  
407 Average  $\delta^{13}\text{C}$  value obtained for USGS-40 was  $-26.4 \pm 0.1\%$  ( $n = 6$ ), which is in good  
408 agreement with accepted value of  $-26.39\%$  (71). The accuracy of the analyses was checked  
409 periodically through the analysis of international reference materials. For  $\delta^{15}\text{N}$ , we obtained –  
410  $4.5 \pm 0.2\%$  ( $n = 6$ ) for USGS-40 and  $+1.0 \pm 0.2\%$  for IAEA-600, also in good agreement  
411 with accepted values of  $-4.52\%$  and  $+1.0\%$ , respectively (71). The carbon and nitrogen  
412 concentrations (TOC and TN) were determined from the peak areas of the major isotopes  
413 using the calibrations for  $\delta^{13}\text{C}$  and  $\delta^{15}\text{N}$  values. The repeatability was better than 0.2 wt.% for  
414 carbon and nitrogen contents.

415 Additional nitrogen isotope analyses of bulk rock samples were carried out at the University  
416 of St. Andrews with an EA Isolink coupled via a Conflo IV to a MAT 253 IRMS (Thermo  
417 Fisher Scientific, Bremen, Germany). Untreated rock powders were weighed into tin capsules  
418 and combusted in a helium stream at 1020°C with a 5-sec pulse of  $\text{O}_2$  gas (with a flow rate of  
419 250 ml/min). Chromium oxide granules were used as an additional combustion aid in the  
420 reactor.  $\text{SO}_2$  was trapped with silvered cobaltic cobaltous oxide in the reactor, while  $\text{CO}_2$  and  
421  $\text{H}_2\text{O}$  were trapped at room temperature with carbosorb and magnesium perchlorate,  
422 respectively. Nitrogen oxides were reduced to  $\text{N}_2$  with a Cu column at 650°C, which also  
423 trapped excess of  $\text{O}_2$  gas.  $\text{N}_2$  was further purified with a GC column at 50°C. A blank was run

424 after each sample, and a set of standards was included before and after every set of five samples.  
425 USGS40 and USGS41 were used for isotopic calibration while SGR-1 was used as a quality  
426 control standard. We obtained a value of  $+17.4 \pm 0.5\%$  ( $n = 8$ ) for SGR-1, which is in good  
427 agreement with previous measurements ( $+17.4 \pm 0.4\%$ , ref. 72).

428

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442

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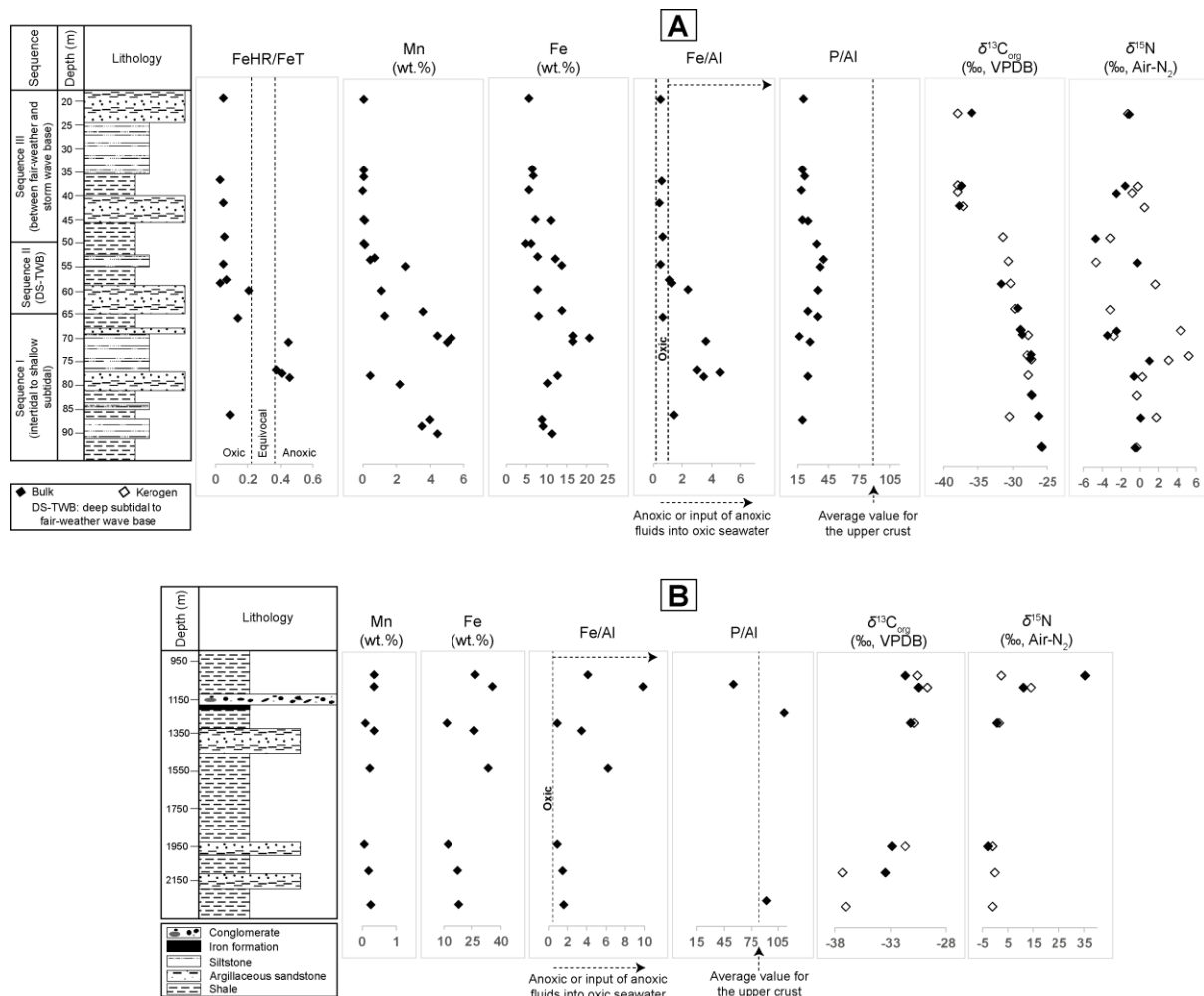
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622 **Figures and Legends**

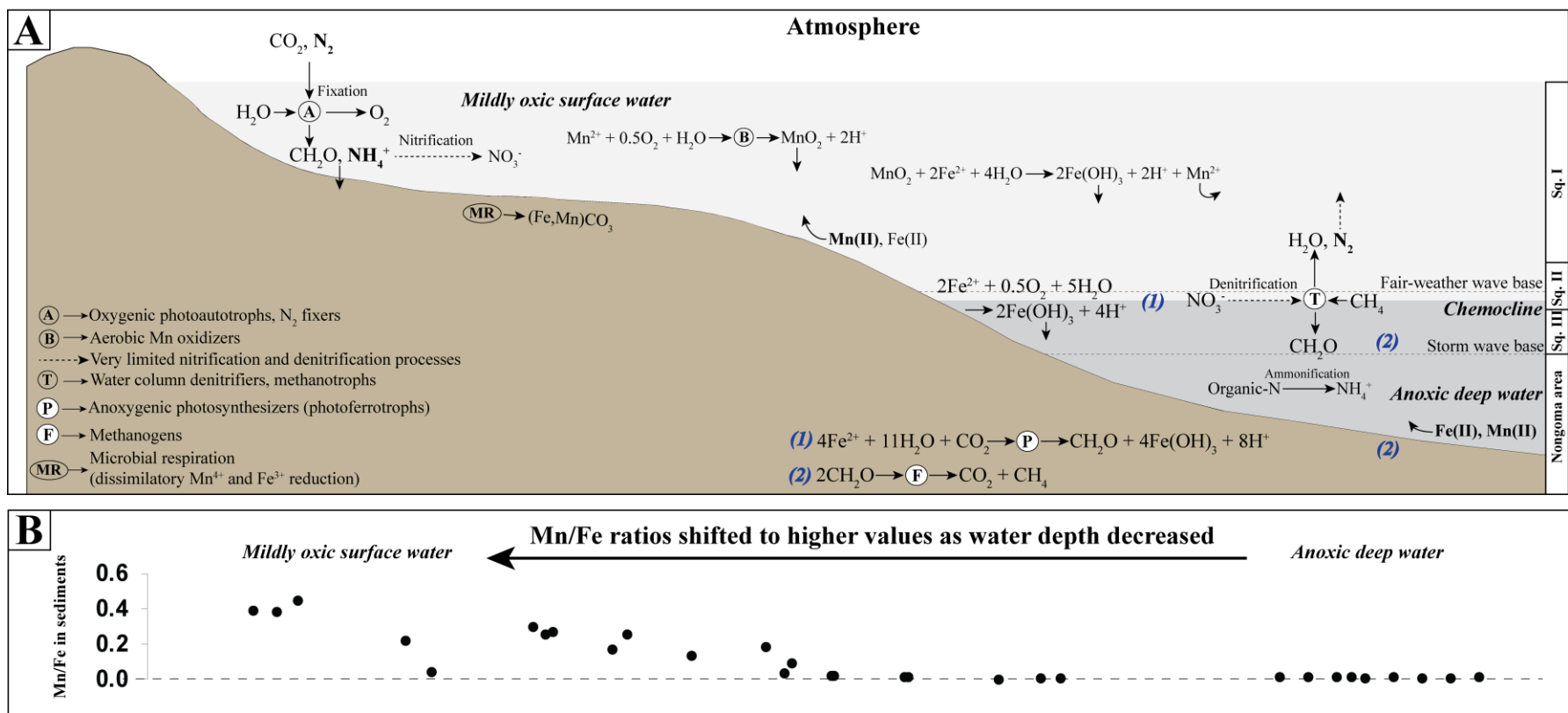


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625 **Fig. 1.** Geochemical data for shale samples plotted along the lithostratigraphic columns of the  
 626 studied sections of the Mozaan Group (**A**) from the shallow part of the Ntombe Formation in  
 627 the White Mfolozi Inlier (Pongola basin), and (**B**) its deeper-water equivalent in the  
 628 Nongoma area (see Fig. S2 for details). Sequences are defined based on water depth  
 629 indicators and chemostratigraphic data. Vertical lines and horizontal arrows on Fe/Al plots  
 630 are based on the description provided in analytical methods (ref. 67), whereas the average  
 631 value for the upper crust on P/Al plots is from ref. 65.

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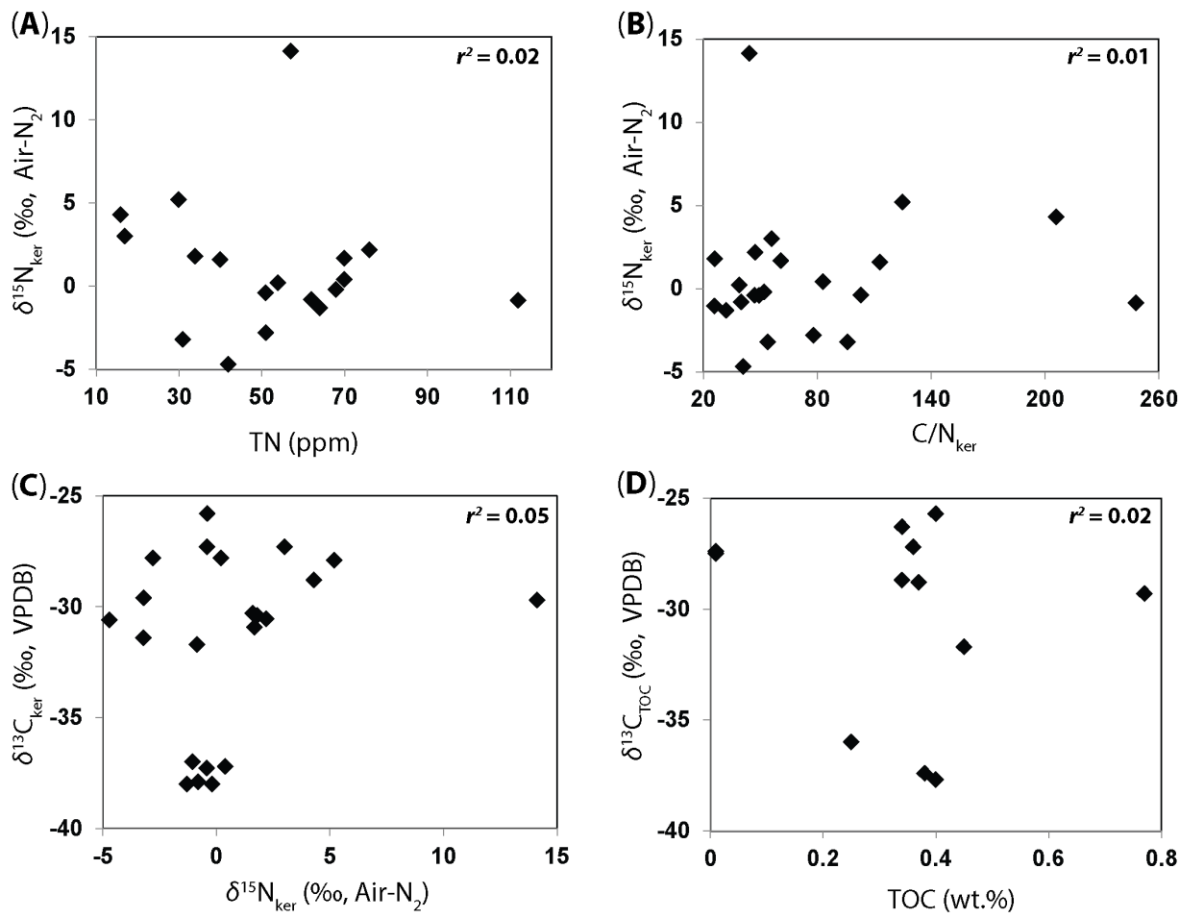
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**Fig. 2.** Proposed paleoenvironmental reconstruction of the Mesoarchean Pongola basin during deposition of the Ntombe Formation, Mozaan Group (modified from ref. 7). **(A)** Water column chemistry and biogeochemical cycles developed in the localized oxygenated surface waters (recorded by the sequences I and II), overlying anoxic deep-waters (recorded by the sequence III and the sedimentary succession in the Nongoma area). Low biological  $\text{O}_2$  production in shallow-marine environments likely limited expression of nitrification and denitrification signals in sediments deposited in the Pongola basin **(B)** Mn/Fe ratios in sediments reflective of seawater redox increase towards the shoreline as ferruginous waters upwelled from anoxic, deep settings to mildly oxygenated, shallow-marine environments. Sq. I (sequence I); Sq. II (sequence II); Sq. III (sequence III).



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643 **Fig. 3.** Cross-plots showing relationships among C and N elemental and isotopic data, with  
 644 no obvious co-variation among these parameters suggesting a minimal impact of post-  
 645 depositional processes on the original C and N isotopic composition of the marine biomass.  
 646 (A) TN vs.  $\delta^{15}\text{N}$  of kerogen ( $\delta^{15}\text{N}_{\text{ker}}$ ); (B) Atomic C/N ratios in kerogen ( $\text{C}/\text{N}_{\text{ker}}$ ) vs.  $\delta^{15}\text{N}_{\text{ker}}$ ;  
 647 (C)  $\delta^{15}\text{N}_{\text{ker}}$  vs.  $\delta^{13}\text{C}_{\text{org}}$  of kerogen; (D) TOC vs.  $\delta^{13}\text{C}_{\text{org}}$  of bulk sediment.