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Uncovering the Ediacaran phosphorus cycle

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Abstract

Phosphorus is a limiting nutrient that is thought to control oceanic oxygen levels to a large extent¹⁻³. A possible increase in marine phosphorus concentrations during the Ediacaran Period (~635-539 Myr) has been proposed as a driver for increasing oxygen levels⁴⁻⁶. Little is known, however, about the nature and evolution of phosphorus cycling during this time⁴. Here we use carbonate-associated phosphate (CAP) from six globally-distributed sections to reconstruct oceanic phosphorous concentrations during a large negative carbon isotope excursion – the Shuram Excursion (SE), which co-occurred with global oceanic oxygenation⁷⁻⁹. Our data suggest pulsed increases in oceanic phosphorus concentrations during the falling and rising limbs of the SE. Using a quantitative biogeochemical model, we propose that this observation could be explained by carbon dioxide and phosphorus release from marine organic matter oxidation primarily by sulphate, with additional phosphorus release from carbon dioxide-driven weathering on land. Collectively, this may have resulted in elevated organic-pyrite burial and ocean oxygenation. Our CAP data also seem to suggest equivalent ocean phosphorus concentrations under maximum and minimum extents of ocean anoxia across the SE. This observation may reflect decoupled phosphorus and ocean anoxia cycles, as opposed to their coupled nature in the modern ocean. Our findings point to external stimuli such as sulphate weathering rather than internal oceanic phosphorus-oxygen cycling alone as a possible control on oceanic oxygenation in the Ediacaran, and in turn this may help explain the prolonged rise of atmospheric oxygen levels.

42 Phosphorus (P) is considered the primary limiting nutrient for marine productivity on geological
43 timescales ¹. Consequently, it has been proposed that the coupled cycling of P and oxygen (O₂)
44 maintains oxygen availability over million-year timescales, preventing runaway ocean anoxia in the
45 Phanerozoic (<539 Myr ago) ¹⁻³ and, whereby sustaining oxygen-demanding eukaryotic life. This
46 relationship exists because dissolved P and O₂ follow negative feedback cycles, where higher
47 oceanic P concentrations under ocean anoxia enhance photosynthetic production of oxygen, which
48 oxygenates previously anoxic water masses ^{1,2}. This feedback arises due to release of P from both
49 the decomposition of organic matter and iron (Fe) (oxyhydr)oxides in anoxic sediments ¹⁰. Under
50 oxygenated bottom waters, this released P generally becomes trapped in the sediment via a variety
51 of pathways, including uptake by Fe (oxyhydr)oxides, biological sequestration of polyphosphates
52 ^{2,10}, or formation of authigenic phosphate minerals, all of which prevent the return of P to the oceans
53 ¹⁰. Under anoxic bottom waters, a proportion of the mobilised P can diffuse out of the sediment ¹¹,
54 however, enhanced organic matter preservation ¹¹ and iron mineral formation under ferruginous
55 (anoxic and Fe²⁺-containing) conditions ¹² can aid P retention in the sediments. Consequently,
56 euxinic conditions (anoxic and sulphidic) are particularly conducive to sedimentary P release
57 because of uptake of dissolved iron as sulphide minerals, thereby limiting P sequestration by iron
58 minerals ¹².

59 The coupled cycling of P and O₂ may have supported high levels of ocean P in the largely anoxic
60 Precambrian oceans (>539 Myr ago) ¹³, as interpreted from P/Fe ratios in iron-formations ¹⁴.
61 Conversely, alternative interpretations of this P/Fe record suggest low oceanic P ^{15,16}, due to
62 effective drawdown of P by iron minerals and sequestration under ferruginous conditions ^{15,17}.
63 Alternatively, low P Precambrian oceans may have arisen due to a lack of oxidants, such as oxygen

64 and sulphate, which suppressed P release from organic matter ^{4,5}. Some researchers have
65 speculated that these low marine P levels substantially increased during the Ediacaran in
66 conjunction with extensive deep ocean oxygenation, giving rise to modern-like oceanic P levels ⁴⁻⁶.
67 The Ediacaran P cycle is thus key to any understanding of how the interaction of P and O₂ cycles
68 evolved from the Precambrian to the modern world and its potential role in the co-evolution of
69 atmospheric O₂ and complex life. However, little is known about the nature and evolution of the
70 Ediacaran P cycle. To shed light on the Ediacaran P cycle, we use the newly developed Carbonate
71 Associated Phosphate (CAP) proxy to directly document variability in oceanic phosphate
72 concentrations during a major Ediacaran ocean oxygenation event ^{7-9,18}. This episode witnessed
73 major diversification of multicellular eukaryotic and metazoan life ^{18,19} and the largest known
74 negative carbonate-carbon isotope ($\delta^{13}\text{C}_{\text{carb}}$) excursion in Earth's history, referred to here as the
75 Shuram Excursion (SE).

76 We analysed six globally distributed and well-preserved carbonate sections (Fig. 1a; see
77 Methods) to capture oceanic phosphate variability during the SE: the Jiulongwan and Sishang
78 sections of the Doushantuo and Dengying formations from South China, the Mochia-Khutuk section
79 of the Shuiquan formation in North-western China, the Cerro Rajón section of the Clemente, Pitiquito
80 and Gamuza formations of northern Mexico, the Death Valley section of the Johnnie Formation of
81 South-western USA, and the Parachilna Gorge section of the Wonoka Formation of South Australia.
82 All sections preserve >10‰ shifts in $\delta^{13}\text{C}_{\text{carb}}$ values characteristic of the SE (Fig. 1b-g) ²⁰⁻²⁷. The six
83 sections reported here are taken to represent global expressions of the SE and are temporally
84 correlated via a combination of chemostratigraphy, fossil assemblages, lithological correlation and
85 palaeomagnetism (see Supplementary Information 2).

86 Among all the measured sections, CAP shows a similar “M-shape” pattern, which is broadly
87 coincident with four stages in the temporal evolution of the $\delta^{13}\text{C}_{\text{carb}}$ trend at each section (Stage I-
88 IV; Fig. 1b-g). Note, that the Death Valley and Parachilna Gorge sections do not preserve the final
89 recovery and initial drop in $\delta^{13}\text{C}_{\text{carb}}$ of the SE, respectively, and the Sishang section is faulted in the
90 middle (Fig. 1d-e, g); however, the sections capture three out of the four CAP shifts identified in
91 other global sections. In Stage I (falling limb of the SE), there is a large increase in CAP (“P-1st” in
92 Fig. 1) at the onset of declining $\delta^{13}\text{C}_{\text{carb}}$ values, with CAP values reaching between 0.39 and 1.33
93 mmol/mol [CAP/(Ca+Mg)]. This increase is followed by an equally large decrease in CAP to a
94 minimum during the mid-SE (Stage II; the plateau in SE $\delta^{13}\text{C}_{\text{carb}}$ values), with CAP values ranging
95 between 0.03 and 0.24 mmol/mol. A rebound in CAP begins in the mid-SE (Stage II) and continues
96 to increase as $\delta^{13}\text{C}_{\text{carb}}$ values increase toward the termination of the SE, reaching values of 0.32 to
97 2.18 mmol/mol (Stage III; rising limb of the SE; “P-2nd” in Fig. 1)), which is then followed by a gradual
98 decline in CAP values to between 0.07 and 0.20 mmol/mol following the termination of the SE (Stage
99 IV; a return to pre-SE $\delta^{13}\text{C}_{\text{carb}}$ values). Given that the CAP trends in all sections are connected to
100 the $\delta^{13}\text{C}_{\text{carb}}$ excursion, and since the SE is approximately synchronous ²⁸, a broadly
101 contemporaneous shift in global oceanic P levels is implied.

102 CAP can be used to produce direct estimates of relative changes in palaeoceanic phosphate
103 levels at the time of carbonate precipitation, if contamination, diagenetic alteration and other
104 potential changes in oceanic chemistry can be evaluated ²⁹. Our CAP extractions show no signs of
105 contamination by other sources of sedimentary P, as there is little to no correlation between CAP
106 and bulk rock P, total organic carbon (TOC) or La (an element used for tracking phosphate mineral
107 dissolution ^{29,30}) (Extended Data Table 1; see Supplementary Information 3). Encouragingly, an

108 independent methodology using in-situ analysis of CAP in carbonate preserving primary textures
109 gives trends consistent with our bulk CAP data from the Dengying Fm. in the Jiulongwan section of
110 South China (Fig. 1b) ³⁰.

111 While the global occurrence of our CAP trends is difficult to reconcile with diagenetic alteration,
112 globally synched diagenetic alteration of carbonate has been purported to result after marine
113 regression exposes platforms to meteoric waters ³¹. The SE, however, is recognised as a marine
114 transgression event, which is unlikely to expose carbonate sediments to meteoric alteration ³².
115 Nevertheless, in order to assess the potential effects of diagenetic alteration on CAP we utilise an
116 open-system diagenetic model to evaluate chemical signals of alteration (Extended Data Fig. 1).
117 Our diagenetic model shows that CAP is altered at similar or greater fluid-to-rock ratios than a
118 number of carbonate proxies, such as $\delta^{13}\text{C}_{\text{carb}}$, $\delta^{18}\text{O}$, $\delta^{44/40}\text{Ca}$ or Mn/Sr. Therefore, CAP could
119 correlate with these proxies across the SE if altered, but this is not observed in our samples
120 (Extended Data Table 1; Fig. 1) ^{29,30}. Furthermore, coexisting $\delta^{34}\text{S}$ values in Carbonate-Associated
121 Sulphate ($\delta^{34}\text{S}_{\text{CAS}}$) decline throughout the SE ^{20-22,24} (Fig. 1b-d), which is inconsistent with known
122 diagenetic pore water chemistry and the increasingly elevated $\delta^{34}\text{S}_{\text{CAS}}$ values that commonly occur
123 in diagenetic carbonates due to microbial sulphate reduction ³³. In addition, local redox proxies (e.g.,
124 $\text{I}/(\text{Ca}+\text{Mg})$, Ce/Ce^* , Fe-speciation) measured among globally distributed SE sections record differing
125 local redox conditions ³⁴⁻³⁶, which is also inconsistent with a global diagenetic pathway (see
126 Supplementary Information 3). Finally, $^{87}\text{Sr}/^{86}\text{Sr}$ values and trends in the Cerro Rajón and Death
127 Valley sections reflect local diagenetic alteration (see Supplementary Information 2) yet preserve
128 similar CAP trends to other SE sections (Fig. 1).

129 Importantly, calcite and dolomite minerals in the study sections have undergone different

130 diagenetic pathways and may have differences in distribution coefficients for CAP²⁸. Despite this,
131 CAP trends remain consistent among sections regardless of whether they are hosted in calcite or
132 dolomite minerals (Fig. 1b-g). This strongly argues against a dominant diagenetic and mineralogical
133 control on the CAP trends, and therefore we conclude that the CAP trends observed in this study
134 most likely represent primary changes in seawater chemistry although diagenetic alteration of local
135 CAP values may have exerted a minor influence. However, we note that in addition to diagenetic
136 processes, changes in ocean pH, alkalinity, temperature and carbonate precipitation can all impact
137 CAP values without concomitant changes in oceanic phosphate concentrations. We consider all of
138 these potentially impacting factors on our CAP data in Supplementary Information 3, 4 and 9, and
139 find that changes in oceanic phosphate concentration is the most plausible overarching control on
140 the global CAP trends that we observed, albeit with possible minor overprints from changes in
141 seawater chemistry.

142 Given the lower-than-mantle $\delta^{13}\text{C}_{\text{carb}}$ values of the SE, some models which report the SE as a
143 primary-seawater-signal have centred on oxidation of an organic matter reservoir, such as dissolved
144 organic matter (DOM)^{37,38}, fossil kerogen²³, or methane³⁹, all of which contain ¹³C-depleted carbon,
145 and with the exception of methane, also contain P. Assuming oxidant supply exceeds demand, these
146 models are supported by many redox studies which suggest a transient global oceanic oxidation
147 event during the SE, as supported by potential global-scale redox proxies such as $\delta^{238}\text{U}_{\text{carb}}$ (Fig. 1b-
148 d). Other models have invoked a diagenetic origin or air-sea exchange of CO₂³² for the low $\delta^{13}\text{C}_{\text{carb}}$
149 values of the SE^{31,40}, but it is difficult to envisage how these models could reconcile the co-evolving
150 trends of our new CAP data with the $\delta^{13}\text{C}_{\text{carb}}$ and redox proxy trends of the SE (see Supplementary
151 Information 3 and 7).

152 Here we entertain a number of primary seawater signal hypotheses for the SE (see
153 Supplementary Information 7 and Extended Data Fig. 2 for details). As discussed below, we find
154 that a best-fit solution is achieved when rising oceanic sulphate levels (driven by enhanced
155 weathering) stimulates sulphate-limited microbial sulphate reduction and oxidation of a DOM
156 reservoir ³⁷ (Fig. 2). This mechanism is supported by higher values of Carbonate-Associated
157 Sulphate (CAS) concentration and lower values of $\delta^{34}\text{S}_{\text{CAS}}$ during the SE (Fig. 1b-d) ²⁰⁻²⁴, which are
158 indicative of a growing oceanic sulphate reservoir ³⁷. Enhanced weathering (and accordingly
159 sulphate supply) during the SE is supported by elevated $^{87}\text{Sr}/^{86}\text{Sr}$ ⁴¹ (Fig. 1b-e) and $\epsilon\text{Nd}(t)$ records
160 ⁴². We quantitatively explore this scenario and its impact on ocean P using the COPSE
161 biogeochemical model (Fig. 3; Extended Data Figs. 2-7).

162 It is well documented that the Ediacaran oceans were dominated by deeper anoxic waters ^{43,44},
163 and therefore free oxygen would have been in contact with a small fraction of the deep-ocean DOM
164 pool. Consequently, sulphate is the primary oxidant in our model. Indeed, if a DOM reservoir was
165 oxidised by free oxygen, our modeling indicates that ocean anoxia would not significantly decrease
166 during the SE, which would allow for sustained high ocean P levels resulting from elevated
167 weathering and DOM oxidation (Extended Data Fig. 3). However, this contrasts with coexisting
168 $\delta^{238}\text{U}_{\text{carb}}$ data showing a substantial decrease in ocean anoxia, as well as the CAP records showing
169 a drop in oceanic P levels during the SE ^{8,35} (Fig. 1b-g). Similarly, models for DOM oxidation by a
170 combination of sulphate and free oxygen are also inconsistent with the observed $\delta^{238}\text{U}_{\text{carb}}$ -CAP
171 records (Extended Data Fig. 3).

172 The sizes of the modelled DOM reservoir and sulphate pulse to the oceans were adjusted to
173 produce a SE with a duration of 15 Myr (see Supplementary Information 5), while also reproducing

174 the two observed CAP peaks (Extended Data Figs. 4-6; P-1st and P-2nd in Fig. 1). Varying the
175 magnitude of the additional sulphate pulse to the oceans has little effect on modelled CAP trends,
176 while burying all additional sulphate as pyrite creates a pO_2 trend inconsistent with redox proxies
177 from the SE (Extended Data Fig. 5). Similarly, if gypsum is assumed not to covary with seawater
178 sulphate levels, increasing sulphate levels prevent a return to anoxia after the SE, which is
179 inconsistent with coexisting $\delta^{238}U_{carb}$ and $\delta^{34}S_{CAS}$ records (Extended Data Fig. 5). Finally, to best fit
180 our CAP data, we suppressed oceanic P levels in COSPE under anoxia by assuming that P is also
181 buried in association with Fe^{2+} , rather than only with Fe^{3+} (Fig. 3; also see Extended Data Fig. 7 for
182 model results with no Fe^{2+} -associated P burial). This choice is based on the predominantly
183 ferruginous ocean conditions in the late Neoproterozoic and previous suggestions that ferruginous
184 oceans may have drawn down ocean P through uptake via a variety of Fe mineral precipitates^{6,15,17}.
185 The relative magnitudes of the modelled ocean P peaks were varied by changing the onset and
186 waning of Fe^{2+} -P burial and whether HCO_3^- is released alongside CO_2 during sulphate reduction of
187 DOM (Fig. 3). Details and rationale for the model choices are provided in the Methods section and
188 Supplementary Information 6-8.

189 Our quantitative modelling indicates that the SE CAP profiles may have recorded feedbacks in
190 weathering, organic matter burial/oxidation, and the extent of marine anoxia that bridges previously
191 elusive $\delta^{13}C$ and palaeoredox proxy relationships. We detail the scenario below: (1) Additional
192 sulphate input to the oceans from weathering triggers the oxidation of a DOM reservoir by sulphate
193 at the beginning of the SE (Fig. 2a). This released organic-bound C and P, which led to declining
194 oceanic $\delta^{13}C$ and increased oceanic P as well as atmospheric pCO_2 , thereby elevating P input to
195 the oceans via silicate weathering (Stage I and P-1st in Figs. 1b-g, 3b). CO_2 -driven weathering also

196 resulted in an elevated $^{87}\text{Sr}/^{86}\text{Sr}$ curve across the SE (Figs. 1b-e, 3d)⁴¹. (2) Greater P availability
197 would then elevate primary productivity and concomitant organic matter and pyrite burial, which
198 increased $p\text{O}_2$ (Fig. 2a) and decreased the extent of ocean anoxia (Fig. 2b), as suggested by the
199 observed global shift to more positive $\delta^{238}\text{U}$ values (Figs. 1b-d; 3c). (3) A decrease in ocean anoxia
200 would increase P burial and lead to a decrease in the oceanic P reservoir (Fig. 2b), consistent with
201 declining CAP values, and coincident maximum $\delta^{238}\text{U}$ values (Stage II in Figs. 1b-g, 3b-c). (4) As
202 the size of the DOM reservoir dwindled, oceanic $\delta^{13}\text{C}$ would start to return to pre-SE values (Stage
203 III in Figs. 1b-g, 3a). Simultaneously, decreasing organic and pyrite burial drove atmospheric oxygen
204 lower, which supported by a reduced P reservoir and consequently lower rates of oxygenic
205 photosynthesis, resulted in lower $p\text{O}_2$ levels and increasing marine anoxia (Fig. 2c), which may be
206 recorded by a transition to increasingly negative $\delta^{238}\text{U}$ values (Stage III in Figs. 1b-d and 3c)^{8,45}.
207 Other independent carbonate- and siliclastic-hosted redox proxies (e.g., Tl and Mo isotopes^{46,47},
208 carbonate-bound iodate³⁵ and Fe-speciation³⁶), corroborate the increase after a prior decrease in
209 ocean anoxia across the SE. Increasing ocean anoxia would decrease P burial, which, when
210 coupled to continued P release from DOM and weathering, would lead to another increase in the P
211 reservoir (Fig. 2c), as observed in the CAP data (P-2nd in Figs. 1b-g, 3b). (5) Finally, cessation of
212 massive DOM oxidation and reduced continental weathering led to a waning P input from DOM and
213 continental weathering, which in turn decreased organic and pyrite burial supporting a return of
214 elevated anoxia, as evidenced by negative $\delta^{238}\text{U}$ values during the terminal SE in multiple global
215 carbonate sections^{7,9} and a return to pre-SE $\delta^{13}\text{C}_{\text{carb}}$ values and coexisting low CAP values (Stage
216 IV in Figs. 1b-g and 3a-c).

217 An alternative to DOM oxidation could be methane oxidation²³ by sulphate. However, anaerobic

218 oxidation of methane is less likely because methane oxidation will not release P directly and is
219 unlikely to lead to $p\text{CO}_2$ increases (and thus elevated P weathering) —favouring alkalinity increases
220 instead. The possibility of small temporal differences in our $\delta^{13}\text{C}_{\text{carb}}$ and CAP trends among globally
221 distributed sections (Fig. 1) may have resulted from heterogeneous redox conditions in the
222 Ediacaran which led to variable spatiotemporal rates of DOM oxidation, continental weathering and
223 element cycling²². Consequently, we conclude that our CAP data in cohort with coexisting C, S, and
224 U isotopic data, as well as other redox proxies, are best explained by the oxidation of an organic
225 matter reservoir (DOM or fossil organic matter) by sulphate and resulting organic and pyrite burial,
226 which facilitated ocean oxygenation and perhaps the coeval diversification of complex life in the
227 Ediacaran (Fig. 3e)^{18,19}.

228 The CAP proxy predominantly records local seawater P levels, reflecting P fluxes from
229 surrounding sediments, riverine input, ocean upwelling etc. Whether local seawater P levels are
230 controlled by local or global processes will depend on the magnitude of fluxes involved at different
231 timescales. Observations of long-term oceanic P cycling suggest that upwelling of deep ocean P
232 onto continental shelves is quantitatively larger than local continental shelf P fluxes¹⁰. This can be
233 demonstrated using a quantitative four-box ocean P cycle model which predicts that continental
234 shelf P concentrations, aside from the most proximal areas and mixed layer, track deep ocean P
235 levels over geological timescales, due to the supply from upwelling^{3,13} (Extended Data Fig. 8). Given
236 the relatively homogenous concentration of deep ocean P, and a residence time greater than the
237 oceanic mixing time¹⁰, upwelling of deep ocean P is expected to result in continental shelf P
238 concentrations that capture overall P levels in the global ocean when viewed over geological
239 timescales. Consequently, our CAP trends from continental shelves are expected to represent global

240 changes in the average ocean P concentration over geological timescales, and their global
241 relationships are thus preserved despite the potential for local overprints that include redox controls
242 on continental shelf P concentrations (see Supplementary Information 8). This view is supported by
243 similar CAP trends among our global sections, despite having been deposited under differing local
244 redox conditions ^{34,35}.

245 Given that CAP and $\delta^{238}\text{U}_{\text{carb}}$ trends may reflect relative changes in the average global ocean P
246 concentration and extent of ocean floor anoxia respectively, our study provides a unique opportunity
247 to explore dynamics in the Ediacaran P and O₂ cycles. As predicted by observations of modern
248 aquatic environments, expansion of oxygenated bottom waters during the SE (evidenced by
249 $\delta^{238}\text{U}_{\text{carb}}$) should be accompanied by declines in marine P concentration (as seen in CAP).
250 Surprisingly, however, minimal extents of ocean floor anoxia (i.e., the most positive $\delta^{238}\text{U}_{\text{carb}}$ values
251 in Stage II of the SE) are associated with CAP values of equal magnitude to those CAP values
252 during periods of maximum ocean floor anoxia before/ after the SE (i.e., the most negative $\delta^{238}\text{U}_{\text{carb}}$
253 values in the early Stage I and Stage IV) at all studied sections (Fig. 1b-d). This observation is in
254 stark contrast to modern P and O₂ cycling where widespread ocean floor anoxia supports higher
255 ocean P levels relative to periods with limited ocean floor anoxia. Similarly, $\delta^{238}\text{U}_{\text{carb}}$ and CAP co-
256 vary, increasing and decreasing at the start and end of the SE, respectively (Fig. 1b-e), which also
257 contrasts with canonical models of modern P and O₂ cycles. This difference indicates that the
258 Ediacaran P and O₂ cycles may have been fundamentally differed to the modern (Fig. 3b-c;
259 Extended Data Fig. 7) ^{1-3,11}. Note that, this conclusion is based on the comparison of CAP values
260 from discrete intervals when $\delta^{238}\text{U}_{\text{carb}}$ data are at their maximum and minimum, and therefore a
261 precise temporal correlation of CAP trends among the studied sections does not affect the

262 conclusions. Moreover, CAP and $\delta^{238}\text{U}_{\text{carb}}$ were measured in identical sample powders and minerals,
263 ruling out effects of temporal or mineralogical differences.

264 One may argue that P fluxes from continental weathering and organic matter oxidation could
265 have maintained high ocean P levels during the SE, effectively balancing the increased P sinks
266 under oxygenated bottom waters. However, current quantitative models of P and O_2 cycling, indicate
267 that even under the most extreme P input fluxes, oceanic P levels are still expected to be lower
268 during minimal oceanic anoxia during the SE, relative to before or after the SE when the extent of
269 seafloor anoxia was greater (see Extended Data Fig. 4 and Supplementary Information 9).
270 Additionally, modeling and experimental results suggest that changes in pH, alkalinity or carbonate
271 precipitation rate resulting from shifts in surface weathering, $p\text{CO}_2$ and organic remineralisation are
272 unlikely to have decoupled the CAP and $\delta^{238}\text{U}_{\text{carb}}$ values (see Extended Data Fig. 9 and
273 Supplementary Information 4 and 9).

274 The roughly equivalent oceanic P levels under maximum and minimum extents of ocean floor
275 anoxia across the SE provides the first direct observation for a decoupling of the modern-style
276 negative feedback cycles between oceanic P and O_2 . Currently there are no other coupled datasets
277 of CAP and $\delta^{238}\text{U}_{\text{carb}}$ which capture a modern-style P and O_2 cycle that we can compare our dataset
278 to. However, chemical traits of redox-dependant phosphorus burial are frequently observed in
279 Phanerozoic sediments ^{2,11,12}, but largely absent in Ediacaran sediments deposited under
280 ferruginous conditions ⁴⁸, which may support our conclusion of decoupled P and O_2 cycles in
281 Ediacaran oceans. For example, $\text{C}_{\text{org}}/\text{P}_{\text{Tot}}$ (total sediment organic carbon/ total sediment P) values
282 are roughly invariant among siliciclastic sediments deposited under variable redox conditions in the

283 Ediacaran Doushantuo Formation ⁴⁸, whereas C_{org}/P_{Tot} commonly covaries with local redox
284 conditions in Phanerozoic sediments ^{2,11,12}.

285 While we favour a primary seawater signal as described above, and cannot entirely rule out an
286 early diagenetic interpretation of the CAP trends, such a scenario is unlikely to change our
287 conclusion of decoupled P and O₂ cycling in the Ediacaran Ocean. This is because in modern marine
288 sediments, increasingly ¹³C-depleted pore waters carry increasingly higher P concentrations
289 following the remineralisation of organic matter. Therefore, if carbonate deposited during the SE
290 indeed captured diagenetic porewater chemistry, the more ¹³C-depleted fluids (down to -12‰)
291 should have carried higher solution P concentrations relative to fluids surrounding carbonate
292 deposited before or after the SE with high $\delta^{13}C_{carb}$ values (up to +5‰). However, carbonates with
293 these endmember $\delta^{13}C_{carb}$ values across the SE carry equable CAP values (Fig. 1b-g). According
294 to our early diagenetic model (Supplementary Information 3), in order to achieve a $\delta^{13}C_{carb}$ shift from
295 +5 to -12‰ in diagenetic carbonates, CAP is expected to increase by at least a factor of four yet
296 plots of our $\delta^{13}C_{carb}$ and CAP data show no such trend (Extended Data Fig. 1d). This decoupling
297 between $\delta^{13}C_{carb}$ and CAP required substantial sequestration of P released from diagenetic organic
298 remineralisation in reducing Ediacaran pore waters (directly evidenced by bottom-water redox
299 proxies, e.g., Fe speciation data across the interval of highly depleted $\delta^{13}C$ values in the Jiulongwan
300 section ⁴³). This P sequestration in anoxic porewaters (and Ediacaran water columns) ^{43,44} is also in
301 stark contrast to modern P and O₂ cycling, thus supporting a decoupled P and O₂ cycle in the
302 Ediacaran (see more detailed discussion in Supplementary Information 3). Moreover, purported
303 models for the SE which achieve low $\delta^{13}C$ in seawater via non-DOM-oxidation mechanisms ³² still
304 advocate for carbonate deposited during the SE to have captured seawater chemistry. Consequently,

305 the observed CAP and $\delta^{238}\text{U}_{\text{carb}}$ trends would be faithful recorders of seawater chemistry and
306 therefore support the conclusion of decoupled P and O₂ cycles in the Ediacaran.

307 The decoupling of ocean P recycling and anoxia observed here likely results from persistent P
308 removal by organic or iron mineral phases under widespread oxidant-depleted, ferruginous
309 conditions in Ediacaran Oceans¹⁶. Given similar redox conditions and potentially a small P reservoir
310 in Precambrian oceans preceding the Ediacaran⁴⁹, it is reasonable to infer that such a decoupling
311 of the P and O₂ cycles may have existed throughout most of the Precambrian⁵⁰. Importantly, a
312 weakening of the oceanic P and O₂ feedbacks as suggested here would have locked the
313 Precambrian oceans and atmosphere into a largely anoxic state, because ocean anoxia would not
314 have resulted in elevated marine P availability, oxygenic photosynthesis and organic matter burial
315 relative to periods of lower marine anoxia. Consequently, Precambrian oceanic oxygenation events,
316 such as the SE and maybe others (e.g., the Great Oxygenation Event⁵⁰), may have relied on
317 external stimuli such as sulphate weathering inputs rather than internal oceanic P-O₂ cycling alone
318^{4,5}. This may explain the prolonged rise of oxygen and complex life over Earth history until the
319 removal of widespread ferruginous conditions or increases in sulphate/oxidant availability in the
320 Phanerozoic which shifted the Ediacaran P and O₂ cycles into a modern-style, negative feedback
321 cycle¹. In turn this transition may explain the prolonged oxygenation and metazoan habitation of the
322 Phanerozoic oceans¹.

323
324

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334 **Author Contributions**

335 C.L led the research. C.L and M.S.D designed the research. M.S.D, Z.Z, M.C and H.G performed
336 analyses. W.S and B.J.W.M conducted modelling work. C.L, T.W.L, D.S.H, S.J.L, M.W.W, A.vS.H,
337 K.L, M.C, H.G provided samples and assistance in the field. S.W.P provided analytical assistance.
338 M.S.D, C.L, and W.S wrote the manuscript with important discussion and contributions from all
339 authors.

340

341 **Data Availability Statement**

342 All data generated or analysed during this study are available at
343 https://figshare.com/articles/dataset/Dodd_et_al_2023_xlsx/22274293 and included with the
344 published article (and its supplementary information files).

345 **Materials Availability**

346 All samples were collected and exported in a responsible manner and in accordance with relevant
347 permits and local laws. Global coordinates and/ or location information and drill core names are
348 given for all samples collected in the supplementary information files. Requests for materials
349 should be addressed to C.L.

350

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489

490 **Methods**

491 **Bulk-rock Phosphorus analysis**

492 Although altered material could not be completely avoided, fresh pieces of only fine-grained
493 carbonate rock were selected and crushed to a powder. Phosphorus measurements were conducted
494 by dissolving ~50 mg of sample powder using a standard HNO_3 –HF digestion protocol, in which
495 HNO_3 –HF (1:1) and HNO_3 were sequentially added to a 15-mL Teflon bomb at 190°C until complete

496 digestion was achieved. Distilled HNO₃ and trace metal-grade HF reagents were used for all
497 samples. The leachates were then measured by ICP-MS using an Agilent 7700x inductively coupled
498 plasma mass spectrometer (ICP-MS). Analytical errors were better than ±7% for phosphorus based
499 on duplicate analyses of two USGS standards (BCR-2, BHVO-2) and two Chinese national
500 standards (GSR1, GSR5).

501

502 **Carbon analysis**

503 Total carbon (TC) and total organic carbon (TOC) were measured using a CS-900 Carbon
504 Analyser. TC was measured directly by weighing out ~0.1g of rock powder with iron shavings and
505 tungsten pellets to aid combustion in the analyser. A calcium carbonate carbon standard (AR1034)
506 was analysed after every 10 samples to check reproducibility (± 0.1 wt% C; n = 30). To measure
507 TOC, an aliquot of sample powder (~1 g) was reacted with 6 M HCl to decarbonate the sample,
508 rinsed with deionised water to neutral pH, then centrifuged and dried, and finally measured in the
509 analyser as for TC. Total inorganic carbon (TIC) was then calculated by the difference between TC
510 and TOC.

511

512 **Carbonate carbon isotope analysis**

513 About 60 to 300 µg of sample powder was loaded into a vial after drying at 70°C for 24 hr in an
514 argon atmosphere. The samples were then reacted with 100% phosphoric acid under a vacuum at
515 70°C for 220 s using a Kiel IV device. The resulting CO₂ was subsequently introduced into a MAT
516 253 isotope ratio mass spectrometer (IRMS) for isotopic measurements. Delta values were
517 calibrated relative to international reference standard NBS-19 ($\delta^{13}\text{C} = +1.95\text{‰}$; $\delta^{18}\text{O} = -2.20\text{‰}$)

518 and Chinese national standard GBW04416 ($\delta^{13}\text{C} = +1.61 \pm 0.03\text{‰}$; $\delta^{18}\text{O} = 1.59 \pm 0.11\text{‰}$).

519 Carbon and oxygen isotope data for carbonates are reported relative to Vienna Pee Dee

520 Belemnite (VPDB) with a precision of better than $\pm 0.1\text{‰}$ based on duplicate analyses of

521 GBW04416 and the study samples.

522

523 **Carbonate-Associated Phosphate (CAP) analysis**

524 CAP was measured following previously established protocols ²⁹. Only samples

525 comprising >50% carbonate were chosen for analysis. Based on sample TIC and carbonate

526 mineralogy, sufficient sample powder was weighed out to yield ~1 mmol of carbonate. An

527 appropriate amount of 2% vol/vol acetic acid was then added to the powder to dissolve up to 70%

528 of the carbonate. After 30 minutes the sample was centrifuged, and the leachate was extracted. An

529 aliquot of leachate was taken for major and trace element analyses and measured using an Expec

530 6000 Inductively Coupled Plasma Optical Emission Spectrometer (ICP-OES) for Ca and Mg and an

531 Agilent 7700x ICP-MS for trace elements. Analytical errors were better than $\pm 2\%$ for all studied

532 elements based on duplicate analyses of two USGS standards (BCR-2, BHVO-2) and two Chinese

533 national standards (GSR1, GSR5). Another aliquot was taken and mixed with H_2SO_4 to achieve a 1

534 M H_2SO_4 concentration and left for 1 hour to remove Ca as CaSO_4 . Then P was measured by

535 spectrophotometry using the molybdenum-blue method at a wavelength of 880 nm, with a relative

536 standard deviation of better than $\pm 5\%$. The sample residue was then washed with 50ml of 10% NaCl

537 solution buffered to pH 8 with NaHCO_3 for 24 hours, and P was measured by the molybdenum-blue

538 method. This sequence was repeated until P in the wash was below detection limit (0.05 ppm). The

539 P in the leach and washes were then summed together to give CAP, which was then normalised to

540 the Ca and Mg concentrations in the leach. Three Ediacaran dolomite sediments of varying TIC and
541 TOC were run alongside unknown samples to check reproducibility for quality assurance, repeat
542 CAP measurements were within $\pm 5\%$.

543

544 **Strontium isotope analysis**

545 The extraction method for strontium isotope analysis was based on previous works⁵¹⁻⁵³. Around
546 0.1g of sample was weighed out and then washed for 24 hours in 10ml of 1M ammonium acetate
547 buffered to pH 8 in order to remove exchangeable strontium. The sample was then centrifuged, and
548 the wash discarded and the sample rinsed with Milli-Q water. Subsequently the sample was partially
549 leached in 2% acetic acid to dissolve 30% of the sample carbonate and left to react overnight. The
550 sample was then centrifuged, and the acetic leach discarded and the sample rinsed with Milli-Q
551 water. Following this a further aliquot of 2% acetic acid was added to dissolve 40% of the sample
552 carbonate and left to react for 30 minutes. The sample was then centrifuged, and the leachate
553 collected for Sr isotope analysis.

554 The leachate collected was then centrifuged to remove insoluble residues and then decanted,
555 dried, and subsequently dissolved in HNO₃. Strontium separation by cation exchange was carried
556 out using automatic column chromatography separation device, see Zhou et al., (2021)⁵⁵ for details.
557 Approximately 200–300 ng of the dried sample was transferred onto a degassed and pre-baked
558 high purity Re filament with Ta₂O₅ activator. The prepared filaments were measured using the VG
559 Sector 54 thermal ionisation mass spectrometer in the Thermal Ionisation Mass Spectrometry facility
560 of China University of Geoscience (Wuhan). Filaments were transferred to a sample carousel,
561 heated under vacuum to a temperature between 1450 and 1650 °C, and analysed when a stable

562 signal was detected on the mass 88 ion beam. Approximately 100 $^{87}\text{Sr}/^{86}\text{Sr}$ ratio data points were
563 collected for each sample. Final data have been corrected for fractionation using the standard value
564 $^{86}\text{Sr}/^{88}\text{Sr} = 0.1194$. The fraction of ^{87}Sr resulting from in situ decay from ^{87}Rb was removed by
565 measurement of rubidium abundance at mass 85. Repeated analysis of the BCR-2 and BHVO-2
566 standards yields an average value of $^{87}\text{Sr}/^{86}\text{Sr} = 0.70498$ and 0.70347 ± 0.00001 (2σ) during the
567 analytical window.

568

569 **Carbonate precipitation experiments**

570 The precipitation of carbonate and uptake of CAP from seawater solutions followed previous
571 methodologies²⁹. In brief, a peristaltic pump was used to add solutions of 1M CaCl_2 and 1M Na_2CO_3
572 with KH_2PO_4 to 1L solutions of synthetic seawater made to emulate modern day seawater chemistry.
573 Solutions were constantly stirred, and an air pump was used to bubble air through the solution in
574 order to maintain a stable pH. For precipitation rate experiments variable amounts of carbonate
575 seeds were used to vary the precipitation rate in conjunction with varying the pump rates of CaCl_2
576 and Na_2CO_3 . Solution chemistry was checked periodically for pH, alkalinity, calcium and phosphorus
577 concentrations. A Thermofisher pH electrode was used to check pH with an accuracy better than \pm
578 0.1 units, alkalinity was measured by titration of seawater aliquots with 0.1M HCl, phosphorus and
579 calcium were measured as outlined above in CAP analysis. At the end of the experimental run,
580 carbonate was collected by filtration through 0.2 μm PES filters and the carbonate washed with 1M
581 NaHCO_3 until phosphorus concentrations in the washes were below detection. Once the carbonate
582 samples were clean of adsorbed phosphorus, 0.1g was completely dissolved in 2M HCl and

583 solutions subsequently analysed for P, Ca, and Mg as outlined above in CAP analysis. Carbonate
584 precipitation rates were calculated as mole/m²/s following Barkan et al. (2020)⁵⁴.

585

586 **COPSE biogeochemical modelling**

587 The COPSE model was used to calculate biogeochemical feedbacks between P cycle and DOM
588 oxidation during the Shuram Excursion (SE). This model follows the work of ref.⁵⁵ and is solved in
589 MATLAB using the Ordinary Differential Equation (ODE) suite. Here, we provide a key description
590 of our model work, and the full description of our model work is given in the Supplementary
591 Information.

592 **Steady-state calculation for low background phosphorus concentration under high ocean**

593 **anoxia in Ediacaran oceans.** Based on the observed CAP and redox proxy trends across the SE

594 we can infer that a mechanism(s) was maintaining P at lower levels during the early Stage I and

595 Stage IV of the SE under high ocean anoxia. While it is ambiguous what mechanism(s) could

596 maintain a lower P reservoir under anoxia during the Ediacaran^{4-6,15}, we chose to modify the iron-

597 bound P burial (Equation 1) to maintain a low oceanic P reservoir during high ocean anoxia because

598 Fe²⁺-P burial is being recently recognized as an important P sink in Precambrian ferruginous-

599 dominated oceans (see Supplementary Information 6 for rationale)^{11, 14-16}, but also allow for an

300 increase in oceanic P burial with decreasing ocean anoxia as modern P cycle. We stress that this

301 modification of the COPSE model is a qualitative change, as there are no current quantitative

302 relationships for Fe²⁺-P burial. Importantly, this modification does not significantly change the model

303 results between model runs with and without Fe²⁺-P burial; except for lowering ocean P levels under

304 high ocean anoxia (Extended Data Fig. 7). Therefore, this modification is appropriate for this study,

305 however future modelling work will require field relationships linking enhanced P burial under anoxia
 306 to produce more meaningful results. Thus, we modified the flux of iron-bound P burial in the standard
 307 COPSE model and include a ferrous iron scavenging flux that has a positive relationship with the
 308 degree of anoxia (ANOX) as below:

$$309 \quad \text{fepb}_{sws} = \frac{k_1}{k_2} \cdot \frac{P_t}{P_0} \quad (1)$$

$$310 \quad \text{fepb}_{fe3} = \text{fepb}_{sws} \cdot (1 - ANOX) = \frac{k_1}{k_2} \cdot \frac{P_t}{P_0} \cdot (1 - ANOX) \quad (2)$$

$$311 \quad \text{fepb}_{fe2} = (\text{fepb}_{sws} - \text{fepb}_{fe3}) \cdot ANOX = \frac{k_1}{k_2} \cdot \frac{P_t}{P_0} \cdot ANOX^2 \quad (3)$$

$$312 \quad \text{fepb}_{tot} = \text{fepb}_{fe3} + \text{fepb}_{fe2} = \frac{k_1}{k_2} \cdot \frac{P_t}{P_0} \cdot (ANOX^2 - ANOX + 1) \quad (4)$$

313 Here fepb_{sws} is the flux of iron-bound P burial reaching the seawater-sediment interface, and fepb_{fe3}
 314 is the final flux of P that is buried with Fe^{3+} , while fepb_{fe2} is the Fe^{2+} scavenging flux including
 315 vivianite burial, and fepb_{tot} is the total flux of iron-bound P burial. P_t and P_0 are the mass of the
 316 marine P reservoir at time t and present day, respectively. $k_1 = 10^9 \text{ mol P yr}^{-1}$ and $k_2 = 0.997527$
 317 are constants, and $0 < ANOX < 0.997527$. This modification changes the original definition of the
 318 total iron-bound P burial which is negatively correlated with anoxia in the standard COPSE model.
 319 In practice, this allows for a low ocean P concentration in anoxic-ferruginous Precambrian oceans
 320 given a $p\text{O}_2 = 0.01 \text{ PAL}$ (see Supplementary Information for details of Fe-scavenging). We chose
 321 this modelling approach because the simplistic separation of Fe-P burial into Fe^{2+} - and Fe^{3+} -P burial
 322 allowed us to increase P burial under high ANOX (per the CAP and redox data) while also keeping
 323 the original COPSE model P and O_2 cycle in effect.

324

325 **Weathering sulphate input.** We run the COPSE model to steady state, forcing the model conditions
 326 to represent the expected geochemical conditions for Ediacaran period. We included a DOM

327 reservoir that responds to a weathering sulphate pulse ($sulfate_{pulse}$), which uses a simple step-
 328 forcing that follows previous work³⁷. The time-dependent forcing function for the sulphate pulse is:

$$329 \quad sulfate_{pulse} = interp1([-10 \ 0 \ 1 \ 15 \ 16 \ 30], [0 \ 0 \ x \ x \ 0 \ 0])$$

330 where the first vector is time (m.y.) and the second vector is the additional sulphate input relative to
 331 the background weathering flux ($x = 3, 4, \text{ or } 5$). The sulphate pulse is set to last for ~15 million years,
 332 which follows the geological $^{87}\text{Sr}/^{86}\text{Sr}$ and $\epsilon\text{Nd}(t)$ records for elevated weathering input during the
 333 SE⁴³, as well as the rough time-taken for the duration of the SE (see Supplementary Information 5
 334 for details on the age framework of the SE). For the model runs in the manuscript, both normal
 335 steady-state ‘background’ flux (subscript ‘background’) and additional weathering pulse input
 336 (subscript ‘pulse’) are working together and respond to changing geochemical conditions:

$$337 \quad gypw_{background} = k_{gypw} \cdot U \cdot R \cdot \frac{carb_w}{k_{carb_w}} \quad (5)$$

$$338 \quad gypw_{pulse} = gypw_{background} \cdot sulfate_{pulse} \quad (6)$$

$$339 \quad pyrw_{background} = k_{pyrw} \cdot U \cdot R \cdot \frac{carb_w}{k_{carb_w}} \quad (7)$$

$$340 \quad pyrw_{pulse} = pyrw_{background} \cdot sulfate_{pulse} \quad (8)$$

341 Here the k_{gypw} and k_{pyrw} are the present-day weathering rates of gypsum and pyrite, U and R are
 342 the normalised uplift and river runoff forcing, and the $\frac{carb_w}{k_{carb_w}}$ is a climate dependence representing
 343 the effect of $p\text{CO}_2$ on global carbonate weathering. For this study, the test case of $x = 5$ is used for
 344 our idealised (‘best fit’) scenario (Extended Data Fig. 6), which maintains a negative $\delta^{13}\text{C}_{carb}$
 345 excursion of ca. -10 ‰ for million years.

346

347 **DOM oxidation.** The size of the DOM reservoir (model term DOC) is set at the beginning of the
 348 model run. This reservoir has an output flux via DOM oxidation, which is driven by sulphate reduction

349 37:

$$350 \quad \frac{dDOC}{dt} = \begin{cases} 0, & DOC < 10^{12} \text{ mol} \\ 2 \cdot (gypw_{pulse} + pyr_{pulse}) \cdot \frac{S_t}{S_0} \cdot \frac{DOC_t}{DOC_0}, & DOC \geq 10^{12} \text{ mol} \end{cases} \quad (9)$$

351 Here S_t and S_0 are the mass of the marine sulphate reservoirs at time t and present day, DOC_t and
352 DOC_0 are the mass of DOM reservoir at t time and pre-SE, respectively. The $DOC_0 = 30$ times the
353 mass of the modern DIC reservoir following previous estimation, in order to maintain a $\delta^{13}C_{carb}$
354 excursion of ~15 million years and reaching values of -10 ‰. We consider sulphate as the only
355 direct oxidant for DOM oxidation, and its additional pulses set the rate of DOM consumption. The
356 normal background sulphate fluxes are set not to be involved in DOM oxidation, but instead balance
357 the oxidation of fresh organic carbon. In nature, free oxygen can also oxidise DOM once its local
358 concentration crosses over a certain threshold (e.g., ~4 μM ⁵⁸. We also test this possibility of DOM
359 oxidation by free oxygen using two output fluxes via both sulphate and oxygen, which is given in the
360 Extended Data Fig. 3 and Supplementary Information 7. We follow previous models that set DOM
361 oxidation to terminate when the DOM reservoir becomes smaller than 10^{12} moles, rather than zero,
362 to prevent system instability³⁷.

363

364 **Sulphate burial.** The COPSE model assumes that marine sulphate is removed via the burial of
365 gypsum and pyrite, which are linearly proportional to the total oceanic sulphate concentration⁵⁵. In
366 order to balance the additional sulphate input, we add additional fluxes of both gypsum
367 ($mgsb_{additional}$) and pyrite ($mpsb_{additional}$) burial, among which the pyrite burial is equal to half the
368 DOM oxidation rate. The remaining sulphate pulse that is not consumed by DOM is assumed to be
369 buried as gypsum:

$$370 \quad mgsb_{additional} = (gypw_{pulse} + pyr_{pulse} - \frac{dDOC}{2 \cdot dt}) \cdot \frac{S_t}{S_0} \quad (10)$$

371
$$mpsb_{additional} = \frac{dDOC}{2 \cdot dt} \quad (11)$$

372

373 **Phosphorus mass balance.** In this study, elevated sulphate supply to the Ediacaran Ocean,
374 following previous proposals ²⁴, is proposed to create a large long-term oxidant source. This input
375 of oxidising power oxidised a large marine DOM pool, releasing DIC and P. The P released by DOM
376 oxidation (*pdoc*) is assumed to be a net source for the marine P reservoir, which is calculated using
377 the rate of DOM oxidation ($\frac{dDOC}{dt}$) and the C/P_{DOC} ratio. The new P mass is estimated with the
378 following equations, adapted from ref. ⁵⁵:

379
$$\frac{dP}{dt} = psea + pdoc - mopb - capb - fepb_{tot} \quad (12)$$

380 Here *psea* is the weathering flux of reactive P to the ocean, *mopb* and *capb* is P burial with organic
381 carbon and calcium, respectively. We chose a fixed ratio of $C/P_{DOC} = 1000$ as the default, but also
382 tested the influence of changing C/P_{DOC} (250, 1000, 3500, and no P release) on the marine P
383 concentration (see Extend Data Fig. 4).

384 **Full model description.** See Supplementary Information for full modified COPSE model equations
385 and parameters.

386

387 **Code availability.**

388 MATLAB code for COPSE is freely available at <https://github.com/sjdaines/COPSE/releases>.

389 **Methods References**

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- 707

708 Main text figure captions

709 **Figure 1. Carbonate-Associated Phosphate (CAP; given as CAP/(Ca+Mg)) and accompanying**
710 **key geochemical records in six globally distributed sections which recorded the Shuram**
711 **Excursion (SE), the largest known negative C-isotope excursion in carbonate ($\delta^{13}\text{C}_{\text{carb}}$) from**
712 **the geological record.** a) Global palaeogeographic map showing the locations of study regions
713 (red stars) at ~560 Ma, modified after ref. ²⁵. b) Jiulongwan section (South China); c) Cerro Rajón
714 section (Mexico), inset graph shows entire Ediacaran section; d) Death Valley section (USA), e)
715 Parachilna Gorge section (Australia). f) Mochia-Khutuk section (North-western China), g) Sishang
716 section (South China). Geochemical data sources: CAP and ⁸⁷Sr/ ⁸⁶Sr data are from this study, also
717 $\delta^{13}\text{C}_{\text{carb}}$ of the Sishang and Parachilna Gorge sections are data from this study; other C-, U- and S-
718 isotope data from refs. ^{7,8,21,22,23,24,26,27}. P-1st = first CAP peak; P-2nd = second CAP peak. PQ Fm.,
719 Pitiquito Formation; Gm Fm., Gamuza Formation; DY Fm., Dengying Formation; VPDB, Vienna Pee
720 Dee Belemnite; VCDT, Vienna Canyon Diablo Troilite. Stratigraphic heights are meters above the
721 first exposed outcrops of Ediacaran strata, except in the Death Valley, Parachilna and Mochia-
722 Khutuk sections which are meters above the lower boundary of the measured geological formation
723 and Sishang metres below Dengying Formation. Light blue circles are CAP in calcite and dark blue
724 circles CAP in dolomite. White circles are $\delta^{13}\text{C}_{\text{carb}}$ and CAP outliers and error bars show a 5% error

725 margin. Black line through CAP data = LOESS smoothing. Stages I to IV are defined as follows:
726 Stage I (falling limb of $\delta^{13}\text{C}_{\text{carb}}$), Stage II (plateau of $\delta^{13}\text{C}_{\text{carb}}$ values), Stage III (rising limb of $\delta^{13}\text{C}_{\text{carb}}$)
727 and Stage IV (return to pre-SE $\delta^{13}\text{C}_{\text{carb}}$ values). See Supplementary Information 2 for correlation
728 between sections and published data.

729

730 **Figure 2. Conceptual model illustrating the co-evolution of the phosphorus cycle and**
731 **dissolved organic matter (DOM) oxidation during the Shuram Excursion (SE) (see text for**
732 **detailed description of the panels).** a) Increasing weathering sulphate oxidises oceanic DOM
733 increasing oceanic P and atmospheric O_2 and CO_2 ; b) elevated atmospheric O_2 decreases ocean
734 anoxia and increases sedimentary P burial; c) cessation of DOM oxidation and pyrite burial leads to
735 a decrease in atmospheric O_2 and the return of ocean anoxia. Stage I-IV refers to SE stages I-IV in
736 Figure 1. Arrows from Fe^{2+} and Fe^{3+} signify the burial of P with Fe under anoxic and oxic conditions,
737 respectively.

738

739 **Figure 3. Biogeochemical model results which best replicate the observed Carbonate-**
740 **Associated Phosphate (CAP) and C- and U-isotope records in Figure 1.** a) Ocean carbonate
741 C-isotopic composition ($\delta^{13}\text{C}_{\text{carb}}$); b) ocean P concentration ($[\text{P}]_{\text{sw}}$); c) ocean carbonate uranium
742 isotopic composition ($\delta^{238}\text{U}_{\text{carb}}$); d) ocean sulphate sulphur isotopic composition ($\delta^{34}\text{S}_{\text{CAS}}$); e) ocean
743 carbonate strontium isotopic composition ($^{87}\text{Sr}/^{86}\text{Sr}$); f) Ediacaran fossil record (adopted after ref ¹⁹).
744 Note the different time axes of panels a-e from panel f. Stage I-IV and P-1st and P-2nd refer to
745 Shuram Excursion (SE) $\delta^{13}\text{C}_{\text{carb}}$ profile and CAP peaks in Figure 1, respectively. Turquoise lines
746 represent model run with continuous Fe^{2+} -P burial and no HCO_3^- release from DOM oxidation; blue

747 lines represent Fe^{2+} -P burial stops during SE and no HCO_3^- release from DOM oxidation; dark blue
748 lines represent Fe^{2+} -P burial stops during SE and HCO_3^- released during DOM oxidation, which
749 halves CO_2 release from DOM (see Supplementary Information 4 and 6 for details).

750

751 **Extended Data Table and Figure Captions**

752

753 **Extended Data Table 1. Statistics for Carbonate-Associated Phosphate (CAP) extraction and**

754 **diagenetic measures.** The strong, statistically significant correlation between bulk rock La and bulk

755 rock P in the Jiulongwan section derives from the dissolution of phosphate minerals. Yet, no statically

756 significant correlations are seen between La and CAP in the partial leach, implying minimal

757 contamination from phosphate minerals for bulk rock P. Unless otherwise stated all elemental data

758 is derived from CAP partial leach extraction. Isotopic data is obtained for the bulk rock. Note, when

759 removing < 10% of data points as outliers, significant levels $[P(\alpha)]$ all increase above 0.01 for all

760 sections, indicating statistically insignificant correlations. All trendlines are linear fits. TOC = Total

761 Organic Carbon.

762

763 **Extended Data Figure 1. Open system diagenetic evolution fluid-rock interaction model. (a-c)**

764 Fluid -rock alteration models showing the relative order of alteration for CAP, CAS, Fe, Mn, IO_3 , $\delta^{13}\text{C}$,

765 $\delta^{34}\text{S}_{\text{CAS}}$, $\delta^{238}\text{U}$, $\delta^{44/40}\text{Ca}$, $^{87}\text{Sr}/^{86}\text{Sr}$. Multiple curves are presented for $\delta^{13}\text{C}$ and $\delta^{34}\text{S}_{\text{CAS}}$ under varying

766 dissolved inorganic carbon (DIC) and sulphate concentrations in the diagenetic fluid. (d) Fluid-rock

767 alteration model showing the predicted trends between CAP and $\delta^{13}\text{C}$, grey points are CAP and

768 $\delta^{13}\text{C}$ data from all study sections. Solid and dashed lines represent different pore water DIC

769 concentration and $\delta^{13}\text{C}$ compositions. Dotted line is trendline through datapoints with R value. See
770 Supplementary Information for model description. Yellow stars mark the point at which 50% of the
771 CAP value has been altered, the red/ grey stars mark the point at which 50% of the element of
772 interest has been altered. CAP = Carbonate-Associated Phosphate, CAS = Carbonate-Associated
773 Sulphate.

774

775 **Extended Data Figure 2. COPSE model results comparing different hypotheses (dissolved**
776 **organic matter (DOM) oxidation by sulphate only, elevated organic matter recycling, elevated**
777 **weathering by uplift, elevated weathering by volcanism) for the observed changes in ocean**
778 **P during the Shuram Excursion. a, A, I, i.** Relative increase in sulphate addition versus
779 background flux. **b, B, II, ii.** Phosphorus concentration in seawater ($[\text{P}]_{\text{sw}}$). **c, C, III, iii.** Relative
780 atmospheric oxygen concentration ($p\text{O}_2$). **d, D, IV, iv.** Degree of ocean anoxia (Anoxia). **e, E, V, v.**
781 Modelled marine carbonate carbon isotope composition ($\delta^{13}\text{C}_{\text{carb}}$). Note in panel E, the $\delta^{13}\text{C}_{\text{carb}}$
782 reflects the $\delta^{13}\text{C}$ of porewater dissolved inorganic carbon (DIC), not oceanic $\delta^{13}\text{C}$. **f, F, VI, vi.**
783 Modelled marine sulphate sulphur isotope composition ($\delta^{34}\text{S}_{\text{sulphate}}$). **g, G, VII, vii.** Modelled marine
784 carbonate uranium isotope composition ($\delta^{238}\text{U}_{\text{carb}}$). For the DOM oxidation hypothesis, we run the
785 COPSE model with a DOM reservoir 30 times the size of the modern marine DIC reservoir and the
786 C:P of the DOM reservoir is 1000, while the C:P of organic matter in the organic matter recycle
787 model is 250 (see Supplementary Information 6 for more details). PAL = Present Atmospheric Level.
788

789 **Extended Data Figure 3. COPSE model results comparing the oxidation of a dissolve organic**
790 **matter (DOM) reservoir using sulphate, free oxygen (Shields et al., 2019) and sulphate + free**

791 **oxygen, respectively. a, A, I** Relative increase in sulphate addition versus background flux (the
792 varying colour shades of the model lines reflect the varying magnitude of the sulphate pulse for
793 different model runs). **b, B, II.** Phosphorus concentration in seawater ($[P]_{sw}$). **c, C, III.** Relative
794 atmospheric oxygen concentration (pO_2). **d, D, IV.** Degree of ocean anoxia (Anoxia). **e, E, V.**
795 Modelled marine carbonate carbon isotope composition ($\delta^{13}C_{carb}$). **f, F, VI.** Modelled marine sulphate
796 sulphur isotope composition ($\delta^{34}S_{sulphate}$). The C:P of the DOM reservoir is set to 1000 in all model
797 runs. The magnitude of the sulphate pulses for each model is variable because higher additional
798 sulphate fluxes are required for models where DOM is oxidised by O_2 resulting from pyrite burial, in
799 comparison to models where DOM is oxidised only by sulphate (see Supplementary Information 7
300 for details).

301

302 **Extended Data Figure 4. COPSE model results with varying sizes (sub-figures A-F) and**
303 **variable P content (a-d) of an initial dissolved organic matter (DOM) reservoir and higher**
304 **initial steady-state pO_2 of 20% present atmospheric level (PAL), with C:P of DOM = 250 (sub-**
305 **figures a-d) (no Fe^{2+} -P burial) (sub-figures I-IV). A.** Size of DOM reservoir in moles of carbon. **B.**
306 Phosphorus concentration in seawater ($[P]_{sw}$). **C.** Relative atmospheric oxygen concentration (pO_2).
307 **D.** Degree of ocean anoxia (Anoxia). **E.** Modelled marine carbonate carbon isotope composition
308 ($\delta^{13}C_{carb}$). **F.** Modelled marine carbonate uranium isotope composition ($\delta^{238}U_{carb}$). In panels A-F, we
309 choose an additional sulphate pulse of 4 times that of the background flux and the C:P of the DOM
310 reservoir is 1000. **a, I.** Size of DOM reservoir in moles of carbon. **b, II.** Phosphorus concentration in
311 seawater ($[P]_{sw}$). **c, III.** Relative atmospheric oxygen concentration (pO_2). **d, IV.** Degree of ocean
312 anoxia. In panels a-d, we choose a sulphate input of 4 times the background flux and the size of the

313 DOM reservoir is 30 times that of the size of the modern marine dissolved inorganic carbon (DIC)
314 reservoir. For panels I-IV, the DOM reservoir is 30 times that of the size of the modern marine
315 dissolved inorganic carbon (DIC). Higher steady-state pO_2 was achieved by adjusting the model
316 terrestrial P input flux and gypsum burial.

317

318 **Extended Data Figure 5. COPSE model results varying the magnitude of an additional**
319 **sulphate pulse for dissolved organic matter (DOM) oxidation by sulphate only (no Fe^{2+} -P**
320 **burial) (a-h), burying all additional sulphate pulse as pyrite (I-VIII) and setting gypsum burial**
321 **to a constant rate (i-viii). a.** Size of DOM reservoir in moles of carbon. **b.** Modelled marine
322 carbonate carbon isotope composition ($\delta^{13}C_{carb}$). **c.** Phosphorus concentration in seawater ($[P]_{sw}$).
323 **d.** Modelled marine carbonate uranium isotope composition ($\delta^{238}U_{carb}$). **e.** Relative atmospheric
324 oxygen concentration (pO_2). **f.** Degree of ocean anoxia. **g.** Modelled marine sulphate sulphur isotope
325 composition ($\delta^{34}S_{sulphate}$). **h.** Modelled marine carbonate strontium isotope composition ($^{87}Sr/^{86}Sr$).
326 The blue line, the grey line, and the dashed grey line are additional sulphate pulses of 3, 4 and 5
327 times the background flux respectively. **I, i.** Relative atmospheric oxygen concentration (pO_2). **II, ii.**
328 Phosphorus concentration in seawater ($[P]_{sw}$). **III, iii.** Modelled marine carbonate carbon isotope
329 composition ($\delta^{13}C_{carb}$). **VI, vi.** evolution of ocean anoxia. **V, v.** Modelled marine carbonate uranium
330 isotope composition ($\delta^{238}U_{carb}$). **VI, vi.** Modelled marine sulphate sulphur isotope composition
331 ($\delta^{34}S_{sulphate}$). **VII, vii.** Sulphate concentration in seawater ($[SO_4]_{sw}$). **VIII, viii.** Oceanic gypsum burial
332 rate (mgsb). Here we used a DOM reservoir size that is 30 times that of the size of the modern
333 marine dissolved inorganic carbon (DIC) reservoir. PAL = present atmospheric level, POL = present
334 oceanic level.

335

336 **Extended Data Figure 6. Full COPSE model outputs for dissolved organic matter (DOM)**
337 **oxidation by sulphate with constant (Fe^{2+}) -bound phosphorus burial, using a sulphate pulse**
338 **of 4 times background flux, DOM C:P of 1000, and the size of the DOM reservoir is 30 times**
339 **the size of the modern marine dissolved inorganic carbon (DIC) reservoir. a.** Weathering
340 sulphate pulse versus background flux. **b.** DOM oxidation flux (DOM_{ox}) in moles of carbon per year.
341 **c.** DOM reservoir (DOM_{pool}) in moles of carbon. **d.** P concentration in sea water ($[\text{P}]_{\text{sw}}$). **e.** Relative
342 atmospheric oxygen concentration ($p\text{O}_2$) to present atmospheric level (PAL). **f.** Degree of marine
343 anoxia (Anoxia). **g.** Modelled marine carbonate carbon isotope composition ($\delta^{13}\text{C}_{\text{carb}}$). **h.** Modelled
344 marine sulphate sulphur isotope composition ($\delta^{34}\text{S}_{\text{sulphate}}$). **i.** Silicate weathering flux (silw) in moles
345 of carbon per year. **j.** Modelled marine carbonate strontium isotope composition ($^{87}\text{Sr}/^{86}\text{Sr}$). **k.**
346 Relative marine sulphate concentration ($[\text{SO}_4^{2-}]_{\text{sw}}$) to present oceanic level (POL). **l.** Relative marine
347 new primary productivity (newp) to POL. **m.** Relative atmospheric carbon dioxide concentration
348 ($p\text{CO}_2$) to PAL. **n.** Average global temperature (Temp) in $^{\circ}\text{C}$. **o.** Organic carbon weathering flux
349 (oxidw) in moles of carbon per year. **p.** Marine organic carbon burial flux (mocb) in moles of carbon.
350 **q.** Gypsum sulfur weathering flux (gypw) in moles of sulfur. **r.** Pyrite sulfur weathering flux (pyrw) in
351 moles of sulfur. **s.** Marine pyrite sulphur burial flux (mpsb) in moles of sulphur. **t.** Marine gypsum
352 sulphur burial flux (mgsb) in moles of sulphur. **u.** Phosphorus releasing flux from DOM oxidation
353 ($\text{DOM}_{\text{ox_P}}$) in moles of phosphorus. **v.** Flux of weathered phosphorus reaching the sea (psea) in
354 moles of phosphorus. **w.** Total iron-bound phosphorus burial flux (fepb) in moles of phosphorus. **x.**
355 Carbonate-bound phosphorus burial flux (capb) in moles of phosphorus. **y.** Marine organic
356 phosphorus burial flux (mopb) in moles of phosphorus. **z.** Ferric iron (Fe^{3+}) -bound phosphorus burial

357 [fepb(Fe³⁺)] in moles of phosphorus. **aa.** Ferrous iron (Fe²⁺)-bound phosphorus burial [fepb(Fe²⁺)]
358 in moles of phosphorus.

359

360 **Extended Data Figure 7. Comparison of COPSE model results for ocean P cycling with and**
361 **without P burial by Fe²⁺ scavenging.** **a.** Ocean inorganic carbon isotopic composition ($\delta^{13}\text{C}_{\text{carb}}$). **b.**
362 Ocean P concentration ($[\text{P}]_{\text{sw}}$); red shaded areas represent uncertainty windows in Fig. 3. **c.** Ocean
363 uranium isotopic composition recorded in carbonates ($\delta^{238}\text{U}_{\text{carb}}$). **d.** Ocean sulphur isotopic
364 composition recorded in carbonate-associated sulphate ($\delta^{34}\text{S}_{\text{CAS}}$). **e.** Ocean strontium isotopic
365 composition ($^{87}\text{Sr}/^{86}\text{Sr}$). **f.** Ediacaran fossil record adapted after Darroch et al. (2018). Stages I to IV
366 are defined as the SE intervals of falling limb, plateau, rising limb and post-SE, respectively, as in
367 Figure 1 which are matched with modelled ocean P reservoir shifts. Model parameters for outputs
368 are the same as detailed in Fig. 3 and Extended Data Fig. 6 except for the red line which excludes
369 P burial by Fe²⁺-scavenging (i.e., a modern-style P and O₂ cycle). P-1st and P-2nd refers to carbonate-
370 associated phosphate (CAP) peaks in Figure 1.

371

372 **Extended Data Figure 8. Model output of a quantitative 4-box ocean P cycle model.** Output
373 from Figures 4a and 5a in Alcott et al. (2019) with the P concentrations in the respective boxes
374 (proximal shelf, distal shelf, deep ocean) plotted. This shows the relative concentration of soluble
375 reactive phosphorus in each ocean box during a model solution in which P levels are oscillating on
376 a large scale. These results show that even under substantial changes in P concentration, the distal
377 shelf (i.e., the area of the shelf that is not dominated by riverine input) is expected to be strongly
378 linked to the deep ocean P concentration. See Alcott et al. (2019) for full model details.

379

380 **Extended data Figure 9. Experimental constraints on the effects of alkalinity (a) and**
381 **precipitation rate (b) on carbonate-associated phosphate (CAP) values in carbonate. a. CAP**
382 uptake increases with progressively lower $[\text{CO}_3^{2-}]$ and alkalinity concentrations. **b. CAP uptake**
383 decreases with increasing precipitation rate. The changes in CAP over the observed ranges in
384 alkalinity and precipitation rate are small compared to the effects of phosphate concentration and
385 solution pH (Dodd et al., 2021). All trendlines are linear fits. Error bars are $\pm 5\%$ for CAP and ± 0.1
386 for Ca/ ALK.

387





