

This is a repository copy of Unique Neoproterozoic carbon isotope excursions sustained by coupled evaporite dissolution and pyrite burial.

White Rose Research Online URL for this paper: <u>https://eprints.whiterose.ac.uk/149033/</u>

Version: Accepted Version

Article:

Shields, GA, Mills, BJW orcid.org/0000-0002-9141-0931, Zhu, M et al. (3 more authors) (2019) Unique Neoproterozoic carbon isotope excursions sustained by coupled evaporite dissolution and pyrite burial. Nature Geoscience, 12 (10). pp. 823-827. ISSN 1752-0894

https://doi.org/10.1038/s41561-019-0434-3

© 2019, The Author(s), under exclusive licence to Springer Nature Limited. This is an author produced version of an article published in Nature Geoscience. Uploaded in accordance with the publisher's self-archiving policy.

Reuse

Items deposited in White Rose Research Online are protected by copyright, with all rights reserved unless indicated otherwise. They may be downloaded and/or printed for private study, or other acts as permitted by national copyright laws. The publisher or other rights holders may allow further reproduction and re-use of the full text version. This is indicated by the licence information on the White Rose Research Online record for the item.

Takedown

If you consider content in White Rose Research Online to be in breach of UK law, please notify us by emailing eprints@whiterose.ac.uk including the URL of the record and the reason for the withdrawal request.



1	Unique Neoproterozoic carbon isotope excursions sustained
2	by coupled evaporite dissolution and pyrite burial
3	
4	Graham A. Shields ¹ , Benjamin J.W. Mills ² , Maoyan Zhu ^{3,4} , Timothy D. Raub ⁵ Stuart
6	Danies & Timotry W. Lenton
7 8	¹ Department of Earth Sciences, University College London, Gower Place, London, WC1E 6BT, UK
9	² School of Earth and Environment, University of Leeds, Leeds, LS2 9JT, UK
10 11 12	³ State Key Laboratory of Palaeobiology and Stratigraphy & Center for Excellence in Life and Paleoenvironment, Nanjing Institute of Geology and Palaeontology, Chinese Academy of Sciences, Nanjing, 210008, China
13 14	⁴ College of Earth Sciences, University of Chinese Academy of Sciences, Beijing, 100049, China
15	⁵ School of Earth and Environmental Sciences, University of St. Andrews, KY16 9AL, UK
16	⁶ Global Systems Institute, University of Exeter, Exeter, EX4 4QE, UK
17	
18	Correspondence and requests for materials should be addressed to G.S.
19	(g.shields@ucl.ac.uk), B.M (b.mills@leeds.ac.uk) or M.Z. (myzhu@nigpas.ac.cn)
20	
21	

23 The Neoproterozoic Era witnessed a succession of biological innovations that culminated 24 in a wide range of animal body plans and behaviours during the Ediacaran-Cambrian 25 radiations. Intriguingly, this interval is also marked by perturbations to the global carbon 26 cycle, as evidenced by extreme fluctuations in climate and carbon isotopes. The 27 Neoproterozoic isotope record has defied parsimonious explanation because sustained ¹²C-enrichment (low δ^{13} C) in seawater seems to imply that substantially more oxygen was 28 consumed by organic carbon oxidation than could possibly have been available. Here we 29 30 propose a solution to this problem, in which carbon and oxygen cycles can maintain dynamic equilibrium during negative δ^{13} C excursions when surplus oxidant is generated 31 through bacterial reduction of sulfate that originates from evaporite weathering. 32 33 Coupling of evaporite dissolution with pyrite burial drives a positive feedback loop 34 whereby net oxidation of marine organic carbon can sustain greenhouse forcing of 35 chemical weathering, nutrient input and ocean margin euxinia. Our proposed model is 36 particularly applicable to the late Ediacaran 'Shuram' isotope excursion that directly 37 preceded the emergence of energetic metazoan metabolisms during the Ediacaran-38 Cambrian transition. Non-steady state sulfate dynamics are likely to have contributed to 39 climate change, episodic ocean oxygenation and opportunistic radiations of aerobic life 40 forms during the Neoproterozoic Era.

41

The Neoproterozoic Era (1000 – c.540 Ma) marks a turning point in Earth history when groups of morphologically complex multicellular eukaryotes, including algae and animals, attained ecological dominance, irrevocably changing Earth System dynamics¹. These biological radiations took place amid fluctuating climate, including two prolonged episodes of global glaciation during the Cryogenian Period (c.715 – c.660 and c.650 – c.635 Ma) and short-lived, regional ice ages during the Ediacaran Period (e.g. c.580 Ma), interspersed with warmer 48 intervals. The world's oceans also became episodically more oxygenated during the Neoproterozoic with the extent of oxygenated seafloor reaching near-modern levels at times 49 during the early Cambrian². Both climate and oxygenation are regulated by Earth's long-term 50 51 carbon cycle, and so perhaps unsurprisingly this interval is characterised by extreme carbon isotope instability³ (Fig. 1). Since their discovery over 30 years ago^{4-7} , the uniquely high 52 amplitudes of Neoproterozoic δ^{13} C excursions have defied conventional interpretation^{3,8–10}. 53 54 Here we relate the largest of these anomalies to the transfer of oxidant from the evaporite rock 55 reservoir to the surface environment via the coupling of sulfate weathering and pyrite burial. 56 Such pulses of evaporite weathering are predicted to have occurred during the Ediacaran Period 57 (see detailed account in SI 4), in particular, as Rodinia's passive margins underwent tectonic 58 uplift during the amalgamation of Gondwanaland¹¹.

59 Conventional carbon isotope mass balance is based on the principle that the isotopic composition of carbon input via outgassing and weathering, and that of sedimentary carbon 60 61 outputs are equal on time scales of >10⁵ years. This δ^{13} C value is considered to be unchanging at -5‰, which is taken to be the average composition of crustal carbon. Because organic matter 62 is depleted in ¹³C, and carbonate rocks precipitate in isotopic equilibrium with ambient 63 dissolved inorganic carbon (DIC), the mean δ^{13} C value of carbonate rocks and fossils can be 64 used to determine the proportion that sedimentary organic matter makes up of the total 65 sedimentary carbon sink. This proportion is generally referred to as f_{org} , which has varied over 66 Earth history between 0.1 and 0.3^{12} . During the Neoproterozoic Era, globally correlative 67 marine carbonate rocks from at least three intervals (~720 Ma Garvellach¹³, ~650 Ma 68 Trezona^{14,15}, ~560 Ma Shuram^{5,16}/Wonoka⁷/DOUNCE¹⁷ anomalies) are characterised by $\delta^{13}C$ 69 70 values below -5% (Fig. 1), which can, using a conventional mass balance approach, only be 71 explained by negative rates of organic burial. This is particularly true of the late Ediacaran Shuram excursion, during which δ^{13} C remained below -8‰ for at least ~10 Myr^{18,19}. In order 72

73 to address this quandary, it was proposed that the pool of dissolved organic matter (DOM) in 74 the Proterozoic ocean was much larger than today, and that negative excursions represent nonsteady-state remineralisation of that pool³. However, later numerical treatments of this 75 model^{20,21} pointed out that the Earth system cannot remain out of oxygen (and carbon) balance 76 77 for such a long period of time. In other words, there is insufficient oxidant even in the modern atmosphere and oceans to remineralise enough organic matter to drive a -8% δ^{13} C excursion 78 79 for several million years. As a result, many authors have interpreted extreme negative anomalies as diagenetic alteration²², authigenic cements¹⁰ or as regional phenomena²³. 80 81 However, such arguments appeal to an inexplicable sampling bias, whereby globally 82 correlative isotopic signatures are presumed to be unrepresentative of the global carbonate sink. 83 Here we take a different approach to the problem of negative δ^{13} C excursions by viewing them 84 in terms of a linked carbon-sulfur-oxygen system, whereby changes in oxidant dynamics 85 caused an excess of organic carbon oxidation over burial, resulting in a smaller DOM reservoir. 86 For steady state to be maintained throughout a negative δ^{13} C excursion, shrinkage of the DOM pool would need to match surplus oxidant production for the duration of the anomaly. If we 87 consider plausible δ^{13} C values of -10‰¹⁷ and -35‰ for deposited carbonates and kerogen 88 (globally averaged carbon sinks), respectively, and $-30\%^{24}$ and -5% for the DOM reservoir 89 90 and crustal carbon (globally averaged carbon sources), respectively, then organic carbon 91 oxidation would need to increase over the background rate by approximately a factor of three 92 (Methods 1). This oxidant imbalance then requires three times as much oxygen as could have 93 been supplied by organic burial alone. Therefore, it can only plausibly be sustained via the 94 other major net source of oxygen to the Earth system: pyrite burial.

95 Although bacterial sulfate reduction coupled with pyrite burial releases on a mole-for-mole 96 basis almost twice as much oxygen as organic burial²⁵, it is generally assumed that the oxygen 97 released by pyrite burial is approximately matched by the oxygen consumed during pyrite 98 weathering. However, most riverine sulfate derives from the weathering of evaporites²⁶, rates 99 of which due to the sporadic and regional nature of evaporite deposits, will vary considerably 100 over time²⁷. During parts of the Proterozoic, when oceans were both iron-rich and anoxic²⁸, 101 and so prone to sulfate reducing conditions (euxinia) at productive margins, one might 102 realistically suppose that the amount of oxidising power transferred from rock sulfate to the 103 surface environment would also have varied considerably, particularly during times when no 104 basin-scale evaporite deposits were forming.

105 By writing a simple steady state mass-balance for the surface carbon cycle (Methods 1) we can 106 map the broad relationship between the evaporite sulfate input rate, net DOM oxidation and 107 δ^{13} C composition of the ocean-atmosphere system (Fig. 2). These calculations show that an evaporite weathering flux of around 1.1×10^{13} mol S yr⁻¹ (around 10 times the modern flux, 108 as proposed for the dissolution event during the early $Cenozoic^{27}$) could sustain a carbon 109 110 isotope excursion of between -10‰ and -15‰, depending on the proportion of the riverine 111 sulfate flux that is eventually buried as pyrite. Thus, the amount of oxidant required to achieve 112 a deep negative carbon isotope excursion through net organic carbon oxidation may reasonably 113 result from basin-scale evaporite dissolution.

114 Although plausible, we acknowledge that the above steady-state approximation is highly 115 idealised and does not capture the true dynamics of the expected events, which include both positive and negative feedbacks (Fig. 3). For a negative δ^{13} C excursion to be generated, oxidant 116 117 input needs to be coupled to shrinkage of a marine DOM reservoir, rather than accumulating 118 as an atmospheric oxygen increase. The rate of DOM oxidation is controlled by the deep ocean 119 redox state, which itself is largely controlled by the abundance of atmospheric oxygen. 120 Therefore, the process of DOM oxidation must be self-limiting and it should not be possible to 121 deplete the surface oxygen reservoir beyond the level that causes the deep ocean to become 122 entirely anoxic. Net oxidation of organic carbon should cause a substantial rise in atmospheric

123 CO_2 concentration and hence additional climate feedbacks. Rising CO_2 and temperature would 124 support enhanced continental weathering, with the potential to drive further evaporite 125 dissolution and therefore sustain oxidant delivery.

126 The network of long-term biogeochemical feedbacks between the sulfur, carbon and oxygen cycles is adequately represented by the COPSE biogeochemical box model²⁹, and in order to 127 explore how changes in the evaporite sulfate weathering flux might affect the oxidant balance 128 129 in a Proterozoic marine environment, we modify COPSE to include a dynamic reservoir of 130 deep-ocean DOM (modelled as DOC) and an extra input flux of sulfate from weathering 131 (Methods 2 and SI). The model is first set up for an 'Ediacaran like' steady state ($pO_2 = 0.05$ 132 PAL, $pCO_2 = 13$ PAL, SO₄ = 0.1 of present ocean level and a mostly anoxic deep ocean). This 133 is achieved by fixing background tectonic parameters (uplift, degassing) at assumed values for 134 600 Ma, while lowering the phosphorus input rate by $50\%^{30}$.

135 We then perturb the system by adding a weathering pulse of sulfate from evaporite dissolution, 136 and a smaller pulse of sulfate from pyrite oxidation, assuming that pyrite-bearing sediments 137 would also be exposed during uplift. These pulses follow simple stepwise increases and decreases²⁷, but also include a dependence on climate through runoff as summarized in figure 138 139 3 (see methods for details). Figure 4a shows the overall amount of S delivery from evaporite dissolution ($\sim 50 - \sim 100 \times 10^{18}$ mol), which is chosen to be similar to that proposed for 140 basin-scale evaporite dissolution in the Cenozoic ($\sim 1.1 \times 10^{13}$ mol S yr⁻¹ for ~ 5 Myrs)²⁷, 141 142 although we use a longer timeframe to compare to the long duration of the Shuram excursion. The control model run with no DOC reservoir is shown in grey in figure 4. As shown 143 144 previously²⁷, this level of sulfate input raises ocean sulfate concentration considerably. Increased burial of pyrite leads to a substantial increase in pO_2 and less prevalence of anoxia 145 (shown as fraction of anoxic seafloor). Oxidative weathering of fossil organic carbon increases 146 as O₂ rises, causing a small reduction in δ^{13} C and increase in pCO₂. Seawater δ^{34} S decreases 147

during the evaporite dissolution event as the combined δ^{34} S value of inputted evaporite and pyrite sulfur is lower than the initial seawater value (we set δ^{34} S_{pyr} = 0‰ and δ^{34} S_{evap} = 15‰ for the weathered material cf.³¹). The increase in pyrite burial buffers against this change by driving δ^{34} S to more positive values (manifest as a 'hump' in the model δ^{34} S curve), but is insufficient to reverse the overall trend.

153 We view net oxidation of deep ocean DOC as a feedback process driven by other model 154 variables, and model it accordingly. Model runs include a DOC reservoir of size 20 times (light 155 blue) and 30 times (dark blue) the modern DIC reservoir, respectively. The reservoir is assumed 156 to have a carbon isotopic composition of -30% and is allowed to be oxidized when the degree 157 of anoxia (ANOX) is reduced below a threshold value (see Methods 2). In both runs, oxidation 158 of DOC begins as the deep ocean becomes more widely oxygenated, however, as DOC 159 oxidation is an oxygen sink, ANOX remains at the threshold value until the DOC reservoir 160 nears depletion. In this period, the model is in a quasi-steady state wherein the transition to an 161 oxygenated deep ocean is prevented by the net oxidation of DOC. During this state, ocean δ^{13} C 162 is around -9‰, close to the value suggested by the simple calculations shown in Figure 2 for 163 this level of sulfate input. Persistent anoxia, nutrient delivery, and the related increase in pyrite burial rates drive δ^{34} S values higher than the control run, but the model still produces a negative 164 excursion in δ^{34} S, consistent with available observations^{16,32}. On depletion of the DOC 165 166 reservoir, the deep ocean can be oxygenated, and the model gradually returns to follow the 167 control experiment.

The duration of the carbon isotope anomaly in our COPSE model reconstruction depends on the size of the DOM pool and on being able to maintain high rates of pyrite burial, but crucially does not depend on (or deplete) the atmospheric O_2 pool, which is predicted to increase during the course of the excursion. A high rate of pyrite burial would be maintained in part due to the effect of organic remineralisation on raising pCO_2 and global temperatures, which produces a 173 positive feedback loop (Fig. 3), whereby high chemical weathering rates and nutrient input can 174 sustain euxinic ocean margins until either the DOM pool or the evaporite weathering flux 175 decreases below a certain threshold value. Our treatment of these aspects is necessarily simple: 176 we assume in the model run shown that 80% of the inputted sulfate is buried as pyrite by a near-shore biota that is sensitive to river inputs³³. We also assume that gypsum, pyrite and 177 178 organic carbon weathering fluxes are related to the model global rate of runoff (in addition to 179 following a prescribed stepwise increase and decrease for pyrite and gypsum). We show further 180 model runs in the SI in which the climate-weathering effect is not considered, and in which 181 only evaporite inputs are considered, without any weathering of pyrite. We also show runs 182 where differing amounts of P release from DOM oxidation fuels additional primary productivity. Sustained, highly-negative δ^{13} C excursions remain possible in all of these cases, 183 but a closely corresponding negative δ^{34} S excursion can only be achieved when a smaller pulse 184 of pyrite weathering accompanies the evaporite weathering pulse. While pyrite weathering is 185 important in setting seawater δ^{34} S values, pyrite weathering and deposition form an O₂-neutral 186 187 cycle over long timescales so other model processes are relatively unaffected.

188 This evaporite dissolution / DOM oxidation scenario appears to be the most parsimonious 189 solution to the Shuram C-isotope conundrum in that it predicts the extent of oxic seafloor to 190 increase towards the end of the excursion, while maintaining high sulfate concentrations, which is in line with geochemical studies^{34,35}. Furthermore, the Shuram anomaly coincided with 191 orogenic uplift relating to the formation of Gondwanaland³⁶, and in particular the tectonic 192 193 inversion of all major basin-scale evaporite sulfate deposits of Tonian age (see SI 4). Although 194 our model fits best the late Ediacaran Shuram anomaly, coupled evaporite dissolution and 195 pyrite burial may have also played a role in other extreme negative carbon isotope excursions 196 of the Neoproterozoic, which all occurred after one of the largest evaporite depositional events in Earth history between c.830 Ma and c.770 Ma³⁷. The succeeding interval of major carbon 197

198 cycle disruption from c.770 Ma until c.550 Ma was a time of little or no basin-scale evaporite 199 deposition, suggesting that, as in the Cenozoic⁵, the sulfate weathering-deposition cycle was 200 not in steady-state. However, unlike the Cenozoic, the low atmospheric oxygen and anoxic 201 deep ocean of the Neoproterozoic allowed evaporite-derived oxidizing power to be effectively 202 transmitted into a negative δ^{13} C signal.

203 The existence of a series of negative carbon isotope excursions during much of the 204 Neoproterozoic Era indicates that the DOM pool underwent dynamic size changes throughout 205 this time, and served as a buffer against oxygenation and climate change, but only when the 206 pool was sufficiently large. Exhaustion of the DOM pool may have occurred during the Shuram 207 anomaly, suggesting that the expansion of aerobic Ediacaran fauna at that time was an 208 opportunistic radiation in response to a transient oxidant surplus. But it is possible that a greatly-reduced DOM pool may have continued to wax and wane until well into the Cambrian 209 210 Period, during which time geochemical evidence suggests that redox conditions on the seafloor 211 reached modern proportions for the first time².

212

213 Acknowledgements

This work was supported by the NERC-NSFC programme 'Biosphere Evolution, Transitions and Resilience' through grant NE/P013643/1 to G.A.S. and M.Z. and NE/P013651/1 to T.M.L., by grant NE/R010129/1 to G.A.S. and B.J.W.M., a University of Leeds Academic Fellowship to B.J.W.M., and by the National Natural Science Foundation of China (41661134048) and Strategic Priority Research Program (B) of the Chinese Academy of Sciences (XDB18000000) to M.Z.

220 Author contributions

G.A.S. and B.J.W.M. conceived the project. B.J.W.M. created the model, which was revised from previous versions created by T.M.L., B.J.W.M. and S.D. All authors contributed to data interpretation and the writing of the manuscript.

224



Figure 1. Carbonate carbon isotope record (reproduced from ³⁸). Grey area indicates values below the average continental crust and mantle value of between -5‰ and -6‰ with three major excursions at ~720 (Garvellach anomaly¹³), ~650 (Trezona anomaly¹⁴) and ~560 Ma (Shuram anomaly^{5,16}). Smaller post-glacial excursions occur after Cryogenian (Sturtian and Marinoan) low latitude glacial events (blue bars). Black dashed line indicates progressive deep ocean oxygenation between ~580 and ~520 Ma³⁹⁻⁴¹. Note that excursions to below the mantle value occur before and during deep-ocean oxygenation and climate events.



234

Figure 2. Negative carbon isotope excursion driven by net oxidation of a dissolved organic
carbon reservoir via coupled evaporite weathering and pyrite burial. The magnitude of the

- $\delta^{13}C$ anomaly depends on the O₂ production rate from pyrite burial, which results from the
- 238 evaporite weathering flux and the fraction of this flux that is buried as pyrite.









Figure 4. COPSE Model forced with sulfate input and including differently sized DOC reservoirs. A. Input rates of sulfate from gypsum and pyrite weathering. B. Size of DOC reservoir in moles of carbon. C. A. Relative atmospheric oxygen concentration. D. Degree of ocean anoxia. E. Relative atmospheric carbon dioxide concentration. F. Relative ocean sulfate concentration. G. Calculated $\delta^{13}C$ of new carbonate. H. Calculated seawater $\delta^{34}S$.

259

261 Methods

1. Steady state mass balance calculations

263 In order to estimate the surplus oxygen flux needed to sustain the late Ediacaran Shuram anomaly, we apply standard isotope mass balance, and assume δ^{13} C values of -10%¹⁷ and -264 35‰ for deposited carbonates ($\delta^{13}C_{carb}$) and kerogen ($\delta^{13}C_{org}$), respectively, and -30‰²⁴ and -265 266 5‰ for the integrated carbon sources ($\delta^{13}C_{input}$) of DOM oxidation and crustal carbon, respectively. Following standard C-isotope mass balance, $\delta^{13}C_{carb} = (\Delta^{13}C_{carb-org}) \cdot f_{org} +$ 267 δ^{13} C_{input} at steady state, and so during the excursion when $f_{org} = 0.1^{12}$, then δ^{13} C_{input} = -12.5%. 268 269 The proportion that DOM oxidation contributed to the global carbon cycle, i.e. $f_{\text{DOM}} = 0.3$, whereby $\delta^{13}C_{\text{excursion input}} = -12.5\% = \delta^{13}C_{\text{pre-excursion input}}(1-f_{\text{DOM}}) + \delta^{13}C_{\text{DOM}}f_{\text{DOM}}$. A 270 conservative estimate for the requisite surplus oxygen flux (f_{DOM}) would therefore be about 271 272 three times greater than that supplied by organic burial (f_{org}) alone, thus requiring a

273 contribution from other sources, most likely pyrite burial.

In order to explore the potential for evaporite dissolution to drive surface system oxygenation and negative carbon isotope excursions, we first solve a simple isotope mass balance calculation for the δ^{13} C composition of the total combined atmosphere and ocean carbon pool (A). Variations in A over time follow the formulation:

278
$$\frac{dA}{dt} = F_{oxidw} + F_{ocdeg} + F_{carbw} + F_{ccdeg} - F_{ocb} - F_{mccb} - F_{sfw} + F_{DOCox}$$
(1)

279 Where F_{oxidw} is oxidative weathering, F_{ocdeg} is organic carbon metamorphism and 280 degassing, F_{carbw} is carbonate weathering, F_{ccdeg} is carbonate degassing, F_{ocb} is organic 281 carbon burial, F_{mccb} is marine carbonate burial and F_{sfw} is seafloor weathering 282 (following^{29,21}). We add F_{DOCox} to represent the oxidation of a deep ocean reservoir of 283 dissolved organic carbon. As a general approximation to the expected Ediacaran carbon cycle 284 we take $F_{oxidw} = 2.5 \times 10^{12} \text{ mol yr}^{-1}$, $F_{ocdeg} = 0.5 \times 10^{12} \text{ mol yr}^{-1}$, $F_{carbw} = 8 \times 10^{12} \text{ mol}$ 285 yr^{-1} , $F_{ccdeg} = 6 \times 10^{12}$ mol yr^{-1} , $F_{ocb} = 3 \times 10^{12}$ mol yr^{-1} , $F_{mccb} = 12 \times 10^{12}$ mol yr^{-1} , and 286 $F_{sfw} = 2 \times 10^{12}$ mol yr^{-1} . Here $\frac{dA}{dt} = 0$, thus the carbon cycle is in steady state, and the 287 fraction of carbon buried in organic form (f_{org}) is 0.2.

288 We assume that the sulfur cycle begins at steady state, and allow an addition of sulfate 289 from evaporite dissolution, F_{evap} (in moles S). To maintain long-term steady state this 290 evaporite must leave the system in oxidised (e.g. gypsum) or reduced (e.g. pyrite) forms. We denote f_{py} as the fraction of the evaporite input that exits the system as pyrite and experiment 291 292 with values of 0.5 and 1. Assuming that DOC oxidation is driven solely by excess oxygen produced by burial of pyrite, we set the flux of DOC oxidation as $F_{DOCox} = 2 \times f_{py}F_{evap}$. To 293 294 maintain long term steady state in the carbon cycle, the flux of additional CO₂ from DOC 295 oxidation must be balanced by burial of carbonates and organic carbon. We assume this occurs at the initial ratio of 4:1 in favour of carbonates (e.g. $f_{org} = 0.2$). Thus the equation 296 297 for long term carbon isotopic mass balance is:

298
$$F_{oxidw}\delta_G + F_{ocdeg}\delta_G + F_{carbw}\delta_C + F_{ccdeg}\delta_C - F_{ocb}(\delta_A - \Delta B) - F_{mccb}\delta_A - F_{sfw}\delta_A + F_{ccdeg}\delta_C + F_{ccdeg}\delta_C - F_{ocb}(\delta_A - \Delta B) - F_{mccb}\delta_A - F_{sfw}\delta_A + F_{ccdeg}\delta_C + F_{ccdeg}\delta_C + F_{ccdeg}\delta_C - F_{ocb}(\delta_A - \Delta B) - F_{mccb}\delta_A - F_{sfw}\delta_A + F_{ccdeg}\delta_C + F_{ccdeg}\delta_C + F_{ccdeg}\delta_C - F_{ocb}(\delta_A - \Delta B) - F_{mccb}\delta_A - F_{sfw}\delta_A + F_{ccdeg}\delta_C + F_{ccdeg}\delta_A - F_{sfw}\delta_A + F_{ccdeg}\delta_C + F_{cc$$

$$F_{DOCox}\delta_{DOC} - 0.2F_{DOCox}(\delta_A - \Delta B) - 0.8F_{DOCox}\delta_A = 0$$
(2)

where δ_A is the isotopic composition of atmosphere and ocean carbon, δ_C is the compositions of buried carbonates, δ_G is the composition of buried organic carbon and δ_{DOC} is the composition of the oceanic DOC reservoir. We take $\delta_C = 0\%$, $\delta_G = -25\%$, $\Delta B = 25\%$, $\delta_{DOC} = -30\%$, and solve for δ_A under varying values for F_{evap} . Results are shown in figure 2.

305

306 2. COPSE model reconstructions

We run the COPSE model²¹ to steady state under assumed Ediacaran forcings, add a deep ocean reservoir of dissolved organic carbon that responds to the degree of ocean anoxia, then impose an evaporite dissolution and pyrite burial event.

310 Ediacaran steady state

The 'Ediacaran' steady state is achieved by fixing all model parameters at the assumed values for 600 Ma. In all but two cases, these values are assumed to be the same as at the beginning of the Phanerozoic, but we lower the rate of phosphorus input by 50% to reduce atmospheric oxygen (e.g. as in Daines et al. 2017^{22}) and use a static gypsum burial rate of 50% of the present day to reduce ocean sulfate. The steady state has $pO_2 = ~0.05$ PAL, pCO_2 = ~13 PAL, SO₄ = ~0.1 of present ocean level and a mostly anoxic deep ocean (*ANOX* ≈ 0.9).

317 Dissolved organic carbon reservoir

The size of the model DOC reservoir is set at the beginning of the model run. The reservoir has a single output flux via DOC oxidation, which depends on the degree of ocean anoxia (*ANOX*):

321
$$\frac{dDOC}{dt} = \begin{cases} 0, DOC < 1 \times 10^{12} \ mol \\ -\frac{k}{1+e^{-a(1-ANOX-c)}} \left(\frac{DOC}{DOC_0}\right), DOC \ge 1 \times 10^{12} \ mol \end{cases}$$
(3)

here a = 300, c = 0.35 and $k = 1 \times 10^{14}$ mol yr⁻¹ are scaling parameters for the sigmoid function, which define the anoxia level at which DOC oxidation begins and the rapidity of the transition. In practice, this function allows for geologically-rapid oxidation of DOC when ANOX < 0.7. The threshold here is chosen to be slightly below the model steady state so that the DOC reservoir is stable under the COPSE Ediacaran setup. DOC oxidation is terminated when the reservoir becomes smaller than 10^{12} moles, rather than zero, to prevent system instability. The rate of DOC oxidation is controlled by O₂ supply and never reaches the value 329 of k. This is shown in the SI for different values of c. We assume that DOC is oxidised

330 directly by O₂, although oxidation via microbial sulfate reduction is also possible.

331 Sulfate input

An uplift and evaporite dissolution event is prescribed in the model using a simple step-forcing that follows previous work on evaporite dissolution⁵. The time-dependent forcing function for sulfate input is:

335
$$S_{input} = [0\ 1\ 10\ 16], [0\ 7\ 7\ 0]$$
 (4)

where the first bracket is time in million years and the second bracket is the additional sulfate
input in terms of the background weathering fluxes. For the model run in the manuscript,
steady state 'background' and additional 'pulse' inputs of both pyrite and gypsum are
considered:

342
$$pyrw_{background} = k_{pyrw} \cdot g_{runoff}$$
 (7)

343
$$pyrw_{pulse} = k_{pyrw} \cdot g_{runoff} \cdot S_{input} \tag{8}$$

here k_{gyp} and k_{pyr} are the present day weathering rates of gypsum and pyrite, and g_{runoff} is a climate-dependence representing the effect of global runoff on weathering fluxes, e.g. Berner (1994)⁴²:

347
$$g_{runoff} = 1 + 0.087(T - T_0)$$
(9)

348 where T is global average surface temperature and T_0 is the present day value.

349 Sulfate burial

	The COPSE model assumes that sulfate burial fluxes are linearly proportional to the	
351	total oceanic sulfate concentration, which means that concentration would have to rise to	
352	many times the present day level in order to bury large amounts of pyrite. We add an	
353	additional flux of pyrite burial (mpsbadditional) that is directly related to the pulsed weathering	
354	input of sulfate, so pyrite burial is more clearly related to the sulfate supply to high-	
355	productivity near-shore environments. A partitioning constant f_{py} is used to determine what	
356	fraction of the pulsed sulfate input is buried as pyrite. This is set at 0.8 in the plot shown in	
357	the ms. Additional sulfate that is not buried as pyrite is assumed to be buried as gypsum:	
358	$mpsb_{additional} = fpy \cdot (pyrw_{pulse} + gypw_{pulse}) $ (10)	
359	$mgsb_{additional} = (1 - fpy) \cdot (pyrw_{pulse} + gypw_{pulse}) $ (11)	
360		
361	Other model alterations	
362	The following alterations are made to COPSE to make the model more applicable to	
363	the scenario being tested:	
	-	
364		
364 365	1. COPSE uses a sigmoid function to calculate the degree of ocean anoxia, <i>ANOX</i> . A	
364 365 366	 COPSE uses a sigmoid function to calculate the degree of ocean anoxia, ANOX. A modified version of the function was previously presented³⁰, based on the anoxia 	
364 365 366 367	 COPSE uses a sigmoid function to calculate the degree of ocean anoxia, ANOX. A modified version of the function was previously presented³⁰, based on the anoxia response of 3D ocean models. Whilst the functions are similar and the choice 	
 364 365 366 367 368 	 COPSE uses a sigmoid function to calculate the degree of ocean anoxia, <i>ANOX</i>. A modified version of the function was previously presented³⁰, based on the anoxia response of 3D ocean models. Whilst the functions are similar and the choice makes little difference in the Phanerozoic version of COPSE, the Watson et al. 	
 364 365 366 367 368 369 	 COPSE uses a sigmoid function to calculate the degree of ocean anoxia, <i>ANOX</i>. A modified version of the function was previously presented³⁰, based on the anoxia response of 3D ocean models. Whilst the functions are similar and the choice makes little difference in the Phanerozoic version of COPSE, the Watson et al. version of the function has a more gradual transition to anoxia and allows the 	
 364 365 366 367 368 369 370 	 COPSE uses a sigmoid function to calculate the degree of ocean anoxia, <i>ANOX</i>. A modified version of the function was previously presented³⁰, based on the anoxia response of 3D ocean models. Whilst the functions are similar and the choice makes little difference in the Phanerozoic version of COPSE, the Watson et al. version of the function has a more gradual transition to anoxia and allows the model to more easily assume an 'Ediacaran-like' steady state under minimal 	
 364 365 366 367 368 369 370 371 	 COPSE uses a sigmoid function to calculate the degree of ocean anoxia, ANOX. A modified version of the function was previously presented³⁰, based on the anoxia response of 3D ocean models. Whilst the functions are similar and the choice makes little difference in the Phanerozoic version of COPSE, the Watson et al. version of the function has a more gradual transition to anoxia and allows the model to more easily assume an 'Ediacaran-like' steady state under minimal alteration of other parameters and is therefore used here. 	
 364 365 366 367 368 369 370 371 372 	 COPSE uses a sigmoid function to calculate the degree of ocean anoxia, <i>ANOX</i>. A modified version of the function was previously presented³⁰, based on the anoxia response of 3D ocean models. Whilst the functions are similar and the choice makes little difference in the Phanerozoic version of COPSE, the Watson et al. version of the function has a more gradual transition to anoxia and allows the model to more easily assume an 'Ediacaran-like' steady state under minimal alteration of other parameters and is therefore used here. COPSE predicts low δ¹³C values for the Ediacaran steady state (~-2‰). In order 	
 364 365 366 367 368 369 370 371 372 373 	 COPSE uses a sigmoid function to calculate the degree of ocean anoxia, <i>ANOX</i>. A modified version of the function was previously presented³⁰, based on the anoxia response of 3D ocean models. Whilst the functions are similar and the choice makes little difference in the Phanerozoic version of COPSE, the Watson et al. version of the function has a more gradual transition to anoxia and allows the model to more easily assume an 'Ediacaran-like' steady state under minimal alteration of other parameters and is therefore used here. COPSE predicts low δ¹³C values for the Ediacaran steady state (~-2‰). In order to clearly test the size of the evaporite-induced excursion, we set the overall 	

375		carbon inputs and makes newly formed $\delta^{13}C_{carb}$ is ~0‰. This alteration merely
376		shifts the baseline of δ^{13} C.
377	3.	We take crustal values of $\delta^{34}S_{pyrite} = 0\%$ and $\delta^{34}S_{gypsum} = 30\%$ to reproduce
378		Ediacaran pre-excursion baseline values ¹⁶ , and assume that the pulse of evaporite
379		weathering has δ^{34} S _{gypsum} = 15‰.
380	4.	COPSE has a relatively high rate of gypsum weathering and burial at present day,
381		we alter the present day reference rate of gypsum weathering to 1×10^{12} mol S yr ⁻
382		¹ , to better represent the background rate used in ref 5, for which our evaporite
383		dissolution scenario is based.
384	3. Additi	onal model experiments
385	Se	ee supplementary information 1 for additional model runs where we assume no
386	climate de	ependence for sulfur weathering fluxes, and consider the role of pyrite versus
387	gypsum v	veathering, and test uncertainty in the DOC oxidation function.
388	4. Full m	odel description
389	Se	e supplementary information 2 for full modified COPSE model equations and
390	descriptio	on. MATLAB code for COPSE is freely available at
391	https://git	hub.com/sjdaines/COPSE/releases
392		
393	Reference	es
394	1. Le	nton, T. M., Boyle, R. A., Poulton, S. W., Shields-Zhou, G. A. & Butterfield, N. J.
395	Co	-evolution of eukaryotes and ocean oxygenation in the Neoproterozoic era. Nat.
396	Ge	osci. 7, (2014).
397	2. Ch	en, X. et al. Rise to modern levels of ocean oxygenation coincided with the
398	Ca	mbrian radiation of animals. Nat. Commun. 6, 1–7 (2015).

399	3.	Rothman, D. H., Hayes, J. M. & Summons, R. E. Dynamics of the Neoproterozoic
400		carbon cycle. Proc. Natl. Acad. Sci. USA 100, 8124-8129 (2003).
401	4.	Knoll, A. H., Hayes, J. M., Kaufman, a J., Swett, K. & Lambert, I. B. Secular
402		variation in carbon isotope ratios from Upper Proterozoic successions of Svalbard and
403		East Greenland. Nature 321, 832–838 (1986).
404	5.	Burns, S. J. & Matter, A. Carbon isotopic record of the latest Proterozoic from Oman.
405		<i>Eclogae Geol. Helv.</i> 86 , 595–607 (1993).
406	6.	Kaufman, A. J., Knoll, A. H. & Narbonne, G. M. Isotopes, ice ages, and terminal
407		Proterozoic earth history. Proc. Natl. Acad. Sci. U. S. A. 94, 6600–6605 (1997).
408	7.	Calver, C. R. Isotope stratigraphy of the Ediacarian (Neoproterozoic III) of the
409		Adelaide Rift Complex, Australia, and the overprint of water column stratification.
410		Precambrian Res. 100, 121–150 (2000).
411	8.	Melezhik, V., Fallick, A. E. & Pokrovsky, B. G. Enigmatic nature of thick sedimentary
412		carbonates depleted in 13C beyond the canonical mantle value: The challenges to our
413		understanding of the terrestrial carbon cycle. Precambrian Res. 137, 131-165 (2005).
414	9.	Grotzinger, J. P., Fike, D. a. & Fischer, W. W. Enigmatic origin of the largest-known
415		carbon isotope excursion in Earth's history. Nat. Geosci. 4, 285–292 (2011).
416	10.	Schrag, D. P., Higgins, J. A., Macdonald, F. A. & Johnston, D. T. Authigenic
417		carbonate and the history of the global carbon cycle. Science (80). 339, 540-3
418		(2013).
419	11.	Li, Z. X. et al. Assembly, configuration, and break-up history of Rodinia: A synthesis.
420		Precambrian Res. (2008). doi:10.1016/j.precamres.2007.04.021
421	12.	Krissansen-Totton, J., Buick, R. & Catling, D. C. A statistical analysis of the carbon

- 422 isotope record from the Archean to Phanerozoic and implications for the rise of
- 423 oxygen. Am. J. Sci. **315**, 275–316 (2015).
- 424 13. Fairchild, I. J. *et al.* Tonian-Cryogenian boundary sections of Argyll, Scotland.
 425 *Precambrian Res.* 1–28 (2017). doi:10.1016/j.precamres.2017.09.020
- 426 14. McKirdy, D. M. et al. A chemostratigraphic overview of the late Cryogenian
- 427 interglacial sequence in the Adelaide Fold-Thrust Belt, South Australia. *Precambrian*428 *Research* 106, 149–186 (2001).
- 429 15. Rose, C. V. *et al.* Constraints on the origin and relative timing of the Trezona ?? 13C
 430 anomaly below the end-Cryogenian glaciation. *Earth Planet. Sci. Lett.* **319–320**, 241–
- 431 250 (2012).
- 432 16. Fike, D. A, Grotzinger, J. P., Pratt, L. M. & Summons, R. E. Oxidation of the
 433 Ediacaran ocean. *Nature* 444, 744–7 (2006).
- 434 17. Lu, M. *et al.* The DOUNCE event at the top of the Ediacaran Doushantuo Formation,
 435 South China: Broad stratigraphic occurrence and non-diagenetic origin. *Precambrian*436 *Res.* 225, (2013).
- 437 18. Condon, D. *et al.* U-Pb ages from the Neoproterozoic Doushantuo Formation, China.
 438 *Science* 308, 95–98 (2005).
- 439 19. Gong, Z., Kodama, K. P. & Li, Y. X. Rock magnetic cyclostratigraphy of the
- 440 Doushantuo Formation, South China and its implications for the duration of the
- 441 Shuram carbon isotope excursion. *Precambrian Res.* **289**, 62–74 (2017).
- 442 20. Bristow, T. F. & Kennedy, M. J. Carbon isotope excursions and the oxidant budget of
 443 the Ediacaran atmosphere and ocean. *Geology* 36, 863–866 (2008).
- 444 21. Bjerrum, C. J. & Canfield, D. E. Towards a quantitative understanding of the late

445		Neoproterozoic carbon cycle. Proc. Natl. Acad. Sci. U. S. A. 108, 5542–5547 (2011).
446	22.	Derry, L. A. A burial diagenesis origin for the Ediacaran Shuram-Wonoka carbon
447		isotope anomaly. Earth Planet. Sci. Lett. 294, 152-162 (2010).
448	23.	Li, C. et al. Uncovering the spatial heterogeneity of Ediacaran carbon cycling.
449		<i>Geobiology</i> 15 , 211–224 (2017).
450	24.	Lee, C., Love, G. D., Fischer, W. W., Grotzinger, J. P. & Halverson, G. P. Marine
451		organic matter cycling during the Ediacaran Shuram excursion. Geology 43, 1103-
452		1106 (2015).
453	25.	Garrels, R. M. & Lerman, A. Coupling of the sedimentary sulfur and carbon cycles -
454		an improved model. Am. J. Sci. 989–1007 (1984).
455	26.	Burke, A. et al. Sulfur isotopes in rivers: Insights into global weathering budgets,
456		pyrite oxidation, and the modern sulfur cycle. Earth Planet. Sci. Lett. (2018).
457		doi:10.1016/j.epsl.2018.05.022
458	27.	Wortmann, U. G. & Paytan, A. Rapid variability of seawater chemistry over the past
459		130 million years. Science (80). 337, 334–336 (2012).
460	28.	Guilbaud, R., Poulton, S. W., Butterfield, N. J., Zhu, M. & Shields-Zhou, G. A. A
461		global transition to ferruginous conditions in the early Neoproterozoic oceans. Nat.
462		<i>Geosci.</i> 8 , (2015).
463	29.	Lenton, T. M., Daines, S. J. & Mills, B. J. W. COPSE reloaded: An improved model of
464		biogeochemical cycling over Phanerozoic time. <i>Earth-Science Rev.</i> 178, 1–28 (2017).
465	30.	Daines, S. J., Mills, B. J. W. & Lenton, T. M. Atmospheric oxygen regulation at low
466		Proterozoic levels by incomplete oxidative weathering of sedimentary organic carbon.
467		Nat. Commun. 8, (2017).

468	31.	Canfield, D. E. & Farquhar, J. Animal evolution, bioturbation, and the sulfate
469		concentration of the oceans. Proc. Natl. Acad. Sci. U. S. A. 106, 8123-8127 (2009).
470	32.	Osburn, M. R., Owens, J., Bergmann, K. D., Lyons, T. W. & Grotzinger, J. P.
471		Dynamic changes in sulfate sulfur isotopes preceding the Ediacaran Shuram
472		Excursion. Geochim. Cosmochim. Acta 170, 204–224 (2015).
473	33.	Laakso, T. A. & Schrag, D. P. A small marine biosphere in the Proterozoic.
474		Geobiology (2019). doi:10.1111/gbi.12323
475	34.	Kendall, B. et al. Uranium and molybdenum isotope evidence for an episode of
476		widespread ocean oxygenation during the late ediacaran period. Geochim. Cosmochim.
477		<i>Acta</i> 156 , 173–193 (2015).
478	35.	Shi, W. et al. Sulfur isotope evidence for transient marine-shelf oxidation during the
479		Ediacaran Shuram Excursion. Geology 46, 267–270 (2018).
480	36.	Campbell, I. H. & Squire, R. J. The mountains that triggered the Late Neoproterozoic
481		increase in oxygen: The Second Great Oxidation Event. Geochim. Cosmochim. Acta
482		74 , 4187–4206 (2010).
483	37.	Prince, J. K. G., Rainbird, R. H. & Wing, B. A. Evaporite deposition in the mid-
484		Neoproterozoic as a driver for changes in seawater chemistry and the biogeochemical
485		cycle of sulfur. Geology (2019). doi:10.1130/g45464.1
486	38.	Saltzman, M. R. & Thomas, E. Chapter 11 - Carbon Isotope Stratigraphy. in The
487		Geologic Time Scale 207-232 (2012). doi:http://dx.doi.org/10.1016/B978-0-444-
488		59425-9.00011-1
489	39.	Canfield, D. E., Poulton, S. W. & Narbonne, G. M. Late-Neoproterozoic Deep-Ocean
490		Oxygenation and the Rise of Animal Life. Science (80). 315, 92-95 (2007).

491	40.	Chen, X. et al. Rise to modern levels of ocean oxygenation coincided with the
492		Cambrian radiation of animals. Nat. Commun. 6, (2015).
493	41.	Sahoo, S. K. et al. Oceanic oxygenation events in the anoxic Ediacaran ocean.
494		<i>Geobiology</i> 14 , (2016).
495	42.	Berner, R. A. GEOCARB II: a revised model of atmospheric CO2over Phanerozoic
496		time. Am. J. Sci. (1994). doi:10.2475/ajs.294.1.56