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Ferruginous oceans during OAE1a and collapse of the marine sulfate pool

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79 **Highlights**

80 • Fe-speciation and redox sensitive trace element enrichment patterns in
81 sediments from OAE1a (~120 Ma) reveal deposition under anoxic, ferruginous
82 conditions in both the paleo-Tethys and paleo-Pacific oceans

83

84 • Widespread ferruginous ocean conditions during OAE1a are only possible with
85 low seawater sulfate concentrations, which must have dropped well below 300
86 μM , and possibly below 100 μM .

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88 • Mass balance of the sulfur cycle predicts that biomass associated organic S (bio-
89 sulfur) may have played an important role as a global S-sink during OAE1a.

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109 **Abstract**

110 Seawater sulfate is one of the largest oxidant pools at Earth's surface today and
111 its concentration in the oceans is generally assumed to have varied between 5 and
112 28 mM since the early Phanerozoic Eon. Intermittent and potentially global
113 Oceanic Anoxic Events (OAEs) are accompanied by changes in seawater sulfate
114 concentrations and signal perturbations in the Earth system associated with major
115 climatic anomalies and biological crises. Ferruginous (Fe-rich) ocean conditions
116 developed transiently during multiple OAEs, implying strong variability in seawater
117 chemistry and global biogeochemical cycles. The precise evolution of seawater
118 sulfate concentrations during OAEs, however, is uncertain and thus models that
119 aim to mechanistically link oceanic anoxia to broad-scale disruptions in the Earth
120 system remain equivocal. Here, we use analyses of Fe-speciation and redox
121 sensitive trace metals in slope sediments deposited in the Tethys and Pacific
122 oceans to constrain seawater sulfate concentrations and underlying dynamics in
123 marine chemistry during OAE1a, ~120 Ma. We find that large parts of the global
124 oceans were anoxic and ferruginous for more than 1 million years. Calculations
125 show that the development of ferruginous conditions requires that seawater sulfate
126 concentrations drop below 300 μM and possibly below 100 μM , which is an order
127 of magnitude lower than previous minimum estimates. Such a collapse of the
128 seawater sulfate pool over a time scale of only one-hundred thousand years is a
129 key and previously unrecognized feature of Phanerozoic Earth surface redox
130 budgets. Critically, this unprecedented sensitivity has potential to dramatically alter
131 global biogeochemical cycles, marine biology, and climate on remarkably short
132 timescales.

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136 **1.0 Introduction**

137 Seawater chemistry is generally thought to have evolved to its current well-
138 oxygenated, sulfate-rich state between 540 and 420 million years ago (Ma)
139 (Stolper and Keller, 2018). Throughout much of the preceding 3.5 billion years, the
140 oceans were largely anoxic, predominantly Fe-rich (ferruginous), and punctuated
141 by intervals of widespread hydrogen sulfide-rich conditions (euxinic) (Poulton and
142 Canfield, 2011). These conditions waned in the early Phanerozoic, and thus, for
143 much of the last 500 Myrs, marine and global biogeochemical cycles were thought
144 to have operated much as they do today. Widespread oceanic anoxia, however,
145 re-emerged intermittently in the Phanerozoic Eon and was particularly prevalent
146 during warm periods such as the Cretaceous (Jenkyns, 2010). The oceans
147 developed euxinia during some of these Oceanic Anoxic Events (OAEs) (Marz et
148 al., 2008; Poulton et al., 2015) when pelagic microbial respiration was channelled
149 through sulfate reduction producing hydrogen sulfide (H₂S) that accumulated in
150 poorly ventilated water masses. Emerging evidence, however, also suggests that
151 ferruginous conditions developed during several OAEs (OAE2, OAE3) (Marz et al.,
152 2008; Poulton et al., 2015). Since development of ferruginous conditions hinges
153 on the balance between Fe- and S-delivery and removal from the oceans, temporal
154 shifts between euxinic and ferruginous conditions imply large-scale variability in
155 ocean chemistry and the S-cycle.

156 The sulfur isotope record and analyses of fluid inclusions from the
157 Cretaceous Period reveal that background seawater sulfate concentrations were
158 much lower (3-10 mM) than the modern oceans (28 mM) (Gomes et al., 2016; Mills
159 et al., 2017; Timofeeff et al., 2006). Episodic evaporite deposition drew seawater
160 sulfate concentrations down even further, possibly to as low as 1 mM, during the
161 Early Cretaceous (Wortmann and Chernyavsky, 2007) and this evaporite
162 deposition likely took place intermittently both before (Davison, 2007), and after
163 (Chaboureau et al., 2013), Aptian OAE1a (~120 Ma). Whereas some stratigraphic
164 reconstructions imply evaporite deposition contemporaneous with OAE1a
165 (Tedeschi et al., 2017), this timing is not well supported by independent
166 chronostratigraphic data. Regardless, low seawater sulfate concentrations in the

167 Aptian oceans could have strongly influenced global biogeochemical cycling
168 across the OAE1a interval. For example, seawater sulfate concentrations are an
169 important control on marine methane (CH₄) budgets, with super-millimolar sulfate
170 concentrations attenuating the release of CH₄ from modern marine sediments to
171 the atmosphere through microbial anaerobic methane oxidation (Reeburgh, 2007).
172 Sub-millimolar sulfate concentrations, in contrast, can lead to large-scale oceanic
173 CH₄ efflux with corresponding implications for climate (Olson et al., 2016).

174 Paleomarine sulfate concentrations have mostly been estimated through
175 two approaches; 1) the rate method, and 2) the microbial sulfate reduction (MSR)
176 trend method (e.g., Algeo et al. (2015), Supplementary Information), but these,
177 importantly, do not resolve seawater sulfate concentrations below about 1 mM.
178 The rate and MSR methods are based on the magnitude of S-isotope fractionation
179 between cogenetic sulfate and sulfide ($\Delta^{34}\text{S}_{\text{CAS-Pyrite}}$), the rate of change in
180 seawater sulfate $\delta^{34}\text{S}$ (based on $\delta^{34}\text{S}$ in carbonate associated sulfate (CAS) or
181 barite minerals), empirical relationships between $\Delta^{34}\text{S}_{\text{CAS-Pyrite}}$, and sulfate
182 concentrations calibrated in modern environments (Algeo et al., 2015). When
183 applied to the early Aptian oceans using existing S-isotope data (Gomes et al.,
184 2016; Kristall et al., 2018; Mills et al., 2017), these methods yield estimates for
185 seawater sulfate that are less than 1-3 mM (Fig. 1). There are, however, critical
186 limitations to these estimates. For example, the rate method only provides
187 maximum estimates for seawater sulfate concentrations because the measured
188 rate of change in $\delta^{34}\text{S}_{\text{Seawater}}$ in a given sedimentary unit is generally smaller than
189 the calculated theoretical maximum, which is mathematically obtained when the
190 source flux of sulfate to the oceans is zero (Algeo et al., 2015) (Supplementary
191 Information) (Fig. 1). Since this scenario does not occur in nature, estimates
192 provided by the rate method tend to be larger than actual seawater sulfate
193 concentrations (Algeo et al., 2015). On the other hand, while there is a relationship
194 between sulfate concentrations and the magnitude of S-isotope fractionations
195 (Algeo et al., 2015), the controls on microbial sulfate reduction and associated
196 isotope fractionation are complex and incompletely known (Canfield and Farquhar,
197 2012), confounding precise application of the MSR method to the geologic record.

198 Both methods are thus incapable of resolving seawater sulfate concentrations
199 across the Aptian beyond coarse maximum estimates of 1-3 mM (Fig. 1). The
200 sensitivity of biogeochemical cycles below this range, however, is important and
201 highlights the need for new data and models that provide more accurate and
202 precise reconstructions of seawater sulfate concentrations during the Aptian Age.

203 Recognizing the limitations of the rate and MSR methods, we took a
204 different approach to reconstructing seawater sulfate concentrations in the Aptian
205 oceans. We studied sedimentary rocks from the Tethys (Cismon drill core) and
206 Pacific (Deep Sea Drilling Project (DSDP) Site 463) oceans that capture OAE1a
207 (Fig. S1). OAE1a is delineated by organic matter (OM)-rich black shale containing
208 units that were deposited over more than a million years (Erba et al., 2010;
209 Malinverno et al., 2010). We conducted a suite of geochemical and mineralogical
210 analyses, which collectively reveal that OAE1a sediments were deposited under
211 anoxic Fe-rich (ferruginous) water column conditions that extended from the
212 Tethys to the Pacific oceans and persisted for more than 1 million years. Our
213 modeling reveals that development of these ferruginous conditions was a response
214 to collapse of the seawater sulfate pool associated with expanded oceanic anoxia.

215

216 **2.0 Methods**

217 *2.1 Sediment digestions and Fe-speciation analyses*

218 We took care to work with well-preserved sedimentary rock samples from
219 both the Cismon and DSDP Site 463 drill cores. These rocks were powdered first
220 using an agate mill and then by hand using an agate mortar and pestle. Sample
221 splits (200 mg) were entirely digested in a lithium metaborate fusion, using a
222 sample to LiO₂ flux ratio of 1:1. Sample splits of 500 mg of sediment were
223 subjected to the Fe-speciation sequential extraction scheme (Poulton and
224 Canfield, 2005). We also performed a revised extraction scheme that included an
225 organic matter leach, as indicated in (Table S1). For the revised scheme our
226 “highly reactive, Fe_{HR}” pool is defined as carbonate-associated Fe (Fe_{carb}, 0.5 N
227 HCl extractable Fe), organic matter associated Fe (Fe_{OM}), ferric (oxyhydr)oxides
228 including magnetite (Fe_{Oxides}, sum of dithionite and oxalate extractable Fe, Fe_{Goe}

229 and $\text{Fe}_{\text{Magnetite}}$, Table S1), and pyrite (Fe_{Pyr}) ($\text{Fe}_{\text{HR}} = \text{Fe}_{\text{Carb}} + \text{Fe}_{\text{OM}} + \text{Fe}_{\text{Oxides}} +$
230 Fe_{Pyr}). The Fe_{Tot} pool is the sum of all Fe_{HR} pools and Fe contained in silicate
231 minerals (Fe_{Sil}).

232

233 *2.2 Elemental concentrations*

234 Extract Fe concentrations were measured by both flame atomic adsorption
235 spectroscopy (Flame AAS, Varian 875) and inductively coupled optical emission
236 mass spectroscopy (ICP-OES, Varian 725ES). Extract major and trace element
237 concentrations were measured by quadrupole inductively coupled plasma mass
238 spectroscopy (Q-ICP-MS, Perkin Elmer NexION 300D), while major elemental
239 concentrations were determined by inductively coupled optical emission mass
240 spectroscopy (ICP-OES, Varian 725ES). For flame AAS measurements, precision
241 on triplicate measurements was 1.2% (1 RSD) and our limit of detection in solution
242 was $80 \mu\text{g L}^{-1}$, or roughly $35 \mu\text{g g}^{-1}$ sediment Fe, based on dilutions. For our ICP-
243 OES measurements precision on triplicate measurements for Fe was 2.2% (1
244 RSD) and our limit of detection in solution was $6 \mu\text{g L}^{-1}$, or roughly $30 \mu\text{g g}^{-1}$
245 sediment, based on dilutions. For Al analysis via ICP-OES, precision on triplicate
246 measurements was 1.2% (1 RSD) and our limit of detection in solution was $6 \mu\text{g}$
247 L^{-1} , or roughly $33 \mu\text{g g}^{-1}$, based on dilutions. Our extractions dissolved >92% of the
248 Fe from the PACS-2 international reference standard. Errors on Fe concentrations
249 in the different leachates based on triplicate extractions of the PACS-2
250 international reference standard are as follows (reported as 1 sigma RSD); $\text{Fe}_{\text{Aca}} \pm 6.0\%$,
251 $\text{Fe}_{0.5\text{NHCl}} \pm 5.8\%$, $\text{Fe}_{\text{Dith}} \pm 2.8\%$, $\text{Fe}_{\text{Oxa}} \pm 10.1\%$, $\text{Fe}_{\text{Sil}} \pm 3.4\%$.

252 For our Q-ICP-MS measurements, precision on Cr was <1% (1 RSD) and
253 our limit of detection in solution was $0.03 \mu\text{g L}^{-1}$, or roughly $0.7 \mu\text{g g}^{-1}$ sediment Cr,
254 based on dilutions. For Cr analysis via ICP-OES, we achieved an RSD of <1% and
255 our limit of detection was roughly $5.3 \mu\text{g g}^{-1}$ sediment, based on dilutions. For V
256 analysis via ICP-OES, we achieved an RSD of <1% and our limit of detection was
257 $0.26 \mu\text{g g}^{-1}$ sediment, based on dilutions. For U and Mo analysis via Q-ICP-MS,
258 we achieved RSDs of <1% and 3.5% respectively, and our limits of detection were
259 roughly 0.10 and 0.40, respectively $\mu\text{g g}^{-1}$ sediment, based on dilutions. Our fusion

260 digestions dissolved ~100%, 97% and 100% of the Cr in the BHVO-2, MESS-3,
261 and PACS-2 international reference standards, respectively.

262

263 *2.3 Microscopy and XRD*

264 Polished thin sections of the Cismon and DSDP Site 463 rocks were imaged
265 on a Hitachi S-4800 field emission scanning electron microscope (Hitachi S-4800
266 FEG SEM) equipped with field emission gun. Elemental concentrations were used
267 to infer mineralogy, and these were determined by energy-dispersive X-ray
268 spectroscopy (EDS, Oxford Instruments X-Max 80 Detector) based on X-ray
269 fluorescence at the relevant emission energies for Fe, C, S, P, Ca and O. Sediment
270 mineralogy was determined by powder X-ray diffraction (XRD). Rock powders
271 were mounted on non-diffracting silica plates. Continuous-scan X-ray diffraction
272 data were collected over a range 3-90°2θ with CoKα radiation on a Rigaku Miniflex
273 diffractometer. We analyzed the X-ray diffractograms using the International
274 Centre for Diffraction Database PDF-4, RRUFF database, and Search-Match
275 software by JADE. The XRD data is plotted in Figure. S2.

276

277 *2.4 1D water column reactive transport model*

278 We developed a reaction transport model to explore how surface ocean
279 sulfate concentrations influence water column sulfate drawdown and rates of pyrite
280 deposition. This model was developed and described in detail in Crowe et al.
281 (2014) with additional details provided in the Supplementary Information. The
282 model predicts seawater sulfate distributions under steady-state conditions by
283 describing changes in sulfate concentration with depth as a function of vertical
284 transport and rates of sulfate reduction to sulfide as:

285

$$286 \quad K_z \frac{\partial^2 [SO_4^{2-}]}{\partial x^2} - R_{SR} = 0 \quad (1)$$

287

288 where K_z is the eddy diffusivity governing vertical mixing ($m^2 d^{-1}$), x is the depth in
289 the water column (m), and R_{SR} is the sulfate reduction rate ($\mu mol m^{-3} d^{-1}$). For K_z ,
290 we implemented a range of values ($0.01 - 1 m^2 d^{-1}$), of which the highest value is

291 similar to globally averaged open ocean vertical transport rates (Munk and
292 Wunsch, 1998), and with lower values representing more strongly stratified seas
293 (Table S2). Sulfate reduction rates were calculated with a Michaelis-Menten like
294 kinetic description:

295

$$296 \quad R_{SR} = \frac{V_{max} [SO_4^{2-}]}{K_m + [SO_4^{2-}]} \quad (2)$$

297

298 where K_m is the half-saturation constant (μM). For K_m , we considered a range of
299 values from 3.6 to 450 μM , which are characteristic of organisms from modern
300 environments with both high and low seawater sulfate concentrations (Table S2).
301 V_{max} is the maximum rate of sulfate reduction ($\mu\text{mol m}^{-3} \text{d}^{-1}$), when sulfate supply
302 is unlimited, and thus corresponds to the scenario when sulfate reduction is limited
303 by organic matter availability. V_{max} can, therefore, be estimated based on models
304 of organic matter degradation rates in modern anoxic marine systems. We thus
305 parameterized V_{max} according to carbon degradation rates in the modern ocean
306 under both high ($V_{max} = 15 \mu\text{M yr}^{-1}$) and low productivity ($V_{max} = 1 \mu\text{M yr}^{-1}$) scenarios
307 (Crowe et al., 2014; Hartnett and Devol, 2003) (Table S2).

308 We set two conditions that tether our model reconstructions for seawater
309 sulfate to the rock record: 1) pyrite burial fluxes were used to place tight limits on
310 total sulfide production fluxes, which cannot be greater than the pyrite burial fluxes
311 recorded in OAE1a sediments assuming all H_2S is quantitatively converted to
312 pyrite; and 2) under ferruginous conditions, seawater sulfate concentrations must
313 be quantitatively drawn down to preserve non-pyritized Fe_{HR} in the presence of
314 abundant organic matter. Pyrite burial fluxes were calculated by combining
315 sedimentation rates (Malinverno et al., 2010) with pyrite concentrations in the
316 Cison and DSDP Site 463 sediments. Together, these two conditions place
317 upper limits on the maximum sulfate flux that can be converted to pyrite through
318 microbial sulfate reduction and reaction of the sulfide produced with Fe_{HR} . We thus
319 use our model to identify a parameter space of possible upper bounds for seawater
320 sulfate concentrations.

321

322 2.5 Sulfur mass balance model

323 To connect the results of our reactive transport modeling to the requisite
324 dynamics in the global sulfur cycle we developed a box model following published
325 studies (Gomes et al., 2016; Mills et al., 2017). Through a system of coupled
326 evolution equations, we track the mass and isotopic composition of marine sulfate
327 as a function of sources and sinks of sulfur to and from the ocean using the
328 following equations;

329

$$330 \quad \frac{\partial M_s}{\partial t} = F_w + F_h - (F_{pyr} + F_{evap} + F_{OM} + F_{CAS}) \quad (3)$$

331

$$332 \quad \frac{\partial \delta^{34}S_{sulphate}}{\partial t} = ((F_w \delta_w + F_h \delta_h) - \delta^{34}S_{sulphate} (F_w + F_h) - F_{pyr} \Delta^{34}S) / M_s \quad (4)$$

333

334 Here, M_s is the mass of sulfur in the ocean; F_h , F_w , are the hydrothermal (including
335 volcanic), and weathering input fluxes of S, respectively; F_{pyr} , F_{OM} , F_{CAS}
336 and F_{evap} are the burial fluxes of pyrite, biomass associated organic S (bio-sulfur),
337 carbonate associated sulfate, and evaporites, respectively. $\delta^{34}S_{sulphate}$ is the S-
338 isotope composition of seawater sulfate. δ_w and δ_h are the S-isotope composition
339 of the weathering (5.2‰) and hydrothermal (3.2 ‰) inputs respectively, $\Delta^{34}S$ is the
340 average isotope fractionation factor associated with pyrite deposition (36.0‰). To
341 obtain an initial value for M_s , we assume an ocean volume of $1.38 \times 10^{18} \text{ m}^3$ and
342 an initial sulfate concentration of 4 mM, in accordance with Cretaceous estimates
343 based on the chemical composition of fluid inclusions and like previous modeling
344 work (Gomes et al., 2016; Timofeeff et al., 2006; Wortmann and Chernyavsky,
345 2007). We take $\delta^{34}S_{sulphate} = 19.5\text{‰}$ as an initial value, which is within error of
346 measurements of pre-OAE1a $\delta^{34}S_{CAS}$. From these initial conditions, we establish
347 a pre-OAE1a steady-state condition (Table S4). Additional details can be found in
348 the Supplementary Information.

349

350 3.0 Results

351 3.1 Fe-speciation

352 Fe-speciation analyses reveal enrichments of pyritizable Fe (Fe_{HR}) across
353 the OM-rich shale intervals that define OAE1a in both Cismon and DSDP Site 463
354 rocks (Fig. 2), relative to rocks stratigraphically above and below. Ratios of
355 $Fe_{HR}/Fe_{Tot} > 0.38$ imply sediment deposition beneath anoxic waters if ratios of
356 Fe_{Tot}/Al are also > 0.5 and C_{org} contents are > 0.5 wt% (Raiswell et al., 2018).
357 Fe_{HR}/Fe_{Tot} values recorded in the Cismon core during OAE1a are consistently
358 above 0.38 and have $Fe_{Tot}/Al > 0.5$ as well as $C_{org} > 0.5$ wt%, diagnostic of
359 deposition below an anoxic water column (Fig. 3). Fe_{HR}/Fe_{Tot} ratios in rocks that
360 bound OAE1a have $Fe_{HR}/Fe_{Tot} < 0.38$ (Fig. 3). Similarly, Fe_{HR}/Fe_{Tot} ratios in rocks
361 deposited at DSDP Site 463 also capture intervals with values > 0.38 and Fe/Al
362 > 0.5 and $C_{org} > 0.5$ wt%. However, some Fe_{HR}/Fe_{Tot} values in sediments deposited
363 during OAE1a at DSDP Site 463, are below the $Fe_{HR}/Fe_{Tot} > 0.38$ threshold (Fig. 3).
364 The sediments deposited during OAE1a, at both sites studied, also preserve
365 appreciable non-pyritized Fe_{HR} (Fig. 2). Fe_{Pyr}/Fe_{HR} ratios are a direct measure of
366 the degree of pyritization of the highly reactive Fe_{HR} pool and ratios < 0.7 indicate
367 deposition under a ferruginous water column with excess highly reactive Fe
368 (Raiswell et al., 2018). All rocks deposited at the Cismon and DSDP 463 sites
369 during the OAE1a interval have $Fe_{Pyr}/Fe_{HR} < 0.7$ (Fig. 3).

370 The 0.5 N HCl leach selectively dissolves poorly crystalline Fe (oxyhydr)-
371 oxides, reactive Fe-carbonates, and acid volatile sulfide (AVS). Determination of
372 both Fe(II) and Fe(III) in the 0.5 N HCl leach, therefore, provides a means of further
373 speciating Fe between highly reactive Fe(II) and Fe(III) phases, as well as
374 determining the association of other elements, like P, with these phases. We find
375 that there was detectable Fe(III) in most of the 0.5 N HCl extracts (Fig. S3), but
376 that this Fe(III) represented a very small component of the total highly reactive Fe
377 in the sediments. For Cismon sediments this translates to an average of $\sim 0.5\%$ of
378 the Fe_{HR} pool. For DSDP Site 463 sediments this translates to an average of $\sim 2.0\%$
379 of the Fe_{HR} pool (Fig. S3). Furthermore, we observe that on average only 7% and
380 5% of the total P is associated with poorly crystalline Fe-(oxyhydr)-oxide phases
381 in the Cismon and DSDP Site 463 sediments, respectively (Fig. S3).

382

383 *3.2 Redox sensitive trace elements (RSTEs)*

384 Multiple redox sensitive trace elements (RSTEs) display strong enrichments
385 relative to the Post Archean Average Shale (PAAS) reference material (McLennan,
386 2001). Rhenium is highly enriched in OAE1a sediments (Fig. 4), with average Re
387 concentrations of 100 and 2.0 ng g⁻¹ in the Cismon and DSDP Site 463 cores,
388 respectively. Average Re concentrations in rocks that bound OAE1a are 1.0 and
389 1.0 ng g⁻¹ in the Cismon and DSDP Site 463 cores, respectively (Fig. 3). During
390 OAE1a average shale normalized Re enrichment factors (Re_{EF}) are 746 and 18 in
391 the Cismon and DSDP Site 463 cores, respectively. Average Re_{EF} in rocks that
392 bound OAE1a are 15 and 21 in the Cismon and DSDP Site 463 cores, respectively
393 (Fig. 4). During OAE1a Cr enrichment factors (Cr_{EF}) are 5.0 and 4.0 in the Cismon
394 and DSDP Site 463 cores, respectively. Average Cr_{EF} in rocks that bound OAE1a
395 are 3.9 and 1.1 in the Cismon and DSDP Site 463 cores, respectively (Fig. 4).
396 During OAE1a average V enrichment factors (V_{EF}) are 2.5 and 1.7 in the Cismon
397 and DSDP Site 463 cores, respectively. Average V_{EF} in rocks that bound OAE1a
398 are 1.1 and 1.3 in the Cismon and DSDP Site 463 cores, respectively (Fig. 4). In
399 addition to our new RSTE data, and in contrast to Re, V, and Cr, a compilation of
400 Mo concentrations in sediments deposited across OAE1a (Charbonnier et al.,
401 2018; Follmi, 2012; Westermann et al., 2013) shows that only 2 out of 196 samples
402 analysed have Mo concentrations greater than the 25 µg g⁻¹, a threshold that
403 implies deposition under euxinic conditions (Lyons and Severmann, 2006) (Fig.
404 4e).

405

406 *3.3 Microscopy and XRD*

407 In both the Cismon and DSDP Site 463 sediments, the presence of Fe_{HR} is
408 demonstrable through X-ray diffraction analyses (Fig. S2). An appreciable fraction
409 of this Fe_{HR} is preserved as Fe_{Carb} (Supplementary Data), which operationally
410 reflects the mineral siderite (Fig. S2). In thin section, pyrite crystals in the Cismon
411 and DSDP Site 463 rocks are well-preserved, lacking evidence of oxidation rims,
412 dissolution pitting, or pervasive cracking (Fig. 5). A collection of images of
413 consistently well-preserved pyrite (including framboids) from multiple thin sections

414 are broadly representative of the OAE1a sediments (Fig. 5). Additionally, electron
415 micrographs and corresponding SEM-EDS analyses reveal pristine Ca-P-rich
416 minerals (apatite) and demonstrate a much stronger association of P with Ca than
417 Fe (Fig. 5).

418

419 *3.4 1D water column reaction transport modeling*

420 Results from the water-column reactive transport modeling illustrate how
421 sulfate drawdown and pyrite burial fluxes scale as a function of surface ocean
422 sulfate concentrations. These calculations yield estimates for seawater sulfate that
423 must be less than $\sim 300 \mu\text{M}$, and less than $100 \mu\text{M}$ under most scenarios (Fig. 6).
424 A sensitivity analysis of our model and its relevant parameters (V_{max} , K_m and K_z)
425 provides constraints on allowable seawater sulfate concentrations that fulfill
426 conditions that sulfate concentrations be drawn down to negligible values
427 (condition 1, Methods section 3.4) without exceeding the pyrite/S-burial rates
428 recorded in the OAE1a sediments (condition 2, Methods section 3.4). Higher
429 values of K_m result in lower rates of sulfate reduction and pyrite burial. Higher
430 values for V_{max} result in high rates of sulfate reduction and pyrite burial. The K_z
431 values used are conservative with respect to sulfate concentrations as higher
432 values for K_z lead to lower possible surface seawater sulfate concentrations (Fig.
433 6) (Table S2). In general, the scenario which leads to the maximum predicted
434 seawater sulfate concentration ($\sim 300 \mu\text{M}$) corresponds to V_{max} values similar to
435 respiration rates in modern anoxic water columns ($\sim 7 \mu\text{M yr}^{-1}$), high K_m ($450 \mu\text{M}$),
436 and slightly more sluggish vertical mixing (K_z , $0.1 \text{ m}^{-2} \text{ d}^{-1}$) relative to the modern
437 open oceans (Fig. 6) (Table S2). All other scenarios that satisfy model conditions
438 1 and 2 (Section 2.4) yield lower seawater sulfate concentrations.

439

440 *3.5 Sulfur mass-balance model*

441 Global sulfur mass balance modeling reveals that low μM seawater sulfate
442 concentrations during OAE1a are supported by existing constraints on global S-
443 budgets and S-isotope data. A perturbation to the global S-cycle lasting roughly
444 0.5 Myrs led to enhanced hydrothermal and riverine S-inputs (Mills et al., 2017) of

445 $\sim 2.3 \pm 0.6 \text{ Tmol yr}^{-1}$ and $\sim 3.1 \pm 0.8 \text{ Tmol yr}^{-1}$, respectively (Fig. 8), assuming a
446 conservative $\sim 25\%$ uncertainty for the different S-fluxes (Canfield and Farquhar,
447 2012). To both balance enhanced S-inputs and draw seawater sulfate down to
448 concentrations that satisfy the 1D modeling results (Fig. 6), requires S-burial fluxes
449 of $5.4 \pm 1.7 \text{ Tmol yr}^{-1}$ (95% confidence interval, Fig. 8). The Aptian S-isotope
450 record, furthermore, is satisfied, within the uncertainties of Cretaceous S-fluxes
451 (Table S4), when oceanic S-removal fluxes are appropriately distributed between
452 pathways that are strongly isotopically fractionated from seawater (i.e., pyrite), and
453 those that are not (i.e., CAS, barite, and bio-sulfur, Fig. 8). Following OAE1a, the
454 seawater sulfate pool can rebound to mM sulfate concentrations on timescales of
455 $\sim 1 \text{ Myrs}$ through a reduction in S-removal fluxes and the continued supply of S
456 through volcanism and weathering that is largely unchanged over this time scale
457 (Jellinek et al., 2020) (Fig. 8).

458

459 **4.0 Discussion**

460 *4.1 Sample fidelity*

461 The OAE1a rocks we analyzed from both Cismon and DSDP 463 appear
462 pristine with little evidence for post-depositional oxidation. Previous work observed
463 extensive pyrite oxidation during storage of OAE sediments (OAE2, Kraal et al.,
464 2009). We thus assessed potential post-depositional oxidation through detailed
465 optical and electron microscopy, focusing on observations of pyrite crystal
466 morphology and texture along cracks, or near sample edges where pyrite grains
467 are most exposed to the atmosphere. Visually, all pyrite grains observed in both
468 cores appear well-preserved (Fig. 5) with no evidence of post-depositional
469 oxidation. We also observe pristine Ca-P-rich minerals and limited association of
470 P with Fe (Fig. 5). Pyrite oxidation can cause dissolution of Ca-P minerals and a
471 redistribution of P into newly formed Fe-(oxyhydr)oxide minerals (Kraal et al.,
472 2009). Thus, the preservation of Ca-P-rich minerals in our samples and lack of Fe-
473 associated P further implies negligible oxidation artifacts (Supplementary
474 Information).

475 Tests for more subtle effects of post depositional alteration based on
476 geochemical information support microscopic observations and demonstrate
477 negligible oxidation of sediment Fe-species. Key Fe- and S-bearing minerals
478 sensitive to oxidation include acid volatile S-minerals (AVS), pyrite, and/or siderite.
479 Recognizing that the initial product of acid volatile S (AVS), pyrite, and/or siderite
480 oxidation is poorly crystalline Fe (oxyhydr)-oxides (Luther III et al., 1982), such
481 oxidation would be observed as 0.5 N HCl leachable Fe(III). Most Fe leached in
482 0.5 N HCl was Fe(II) and the concentration of 0.5 N HCl leachable Fe(III) in all
483 rocks analyzed was a negligible (<2%) component of the total Fe_{HR} (Fig. S3).
484 Importantly, 0.5 N HCl leachable Fe(III) is not expected to be preserved in anoxic,
485 OM-rich sediments, and the Fe(III) we measured in this fraction may thus be the
486 product of very limited post depositional oxidation. Critically, however, such a small
487 amount of post depositional oxidation has a negligible effect on our Fe-speciation
488 results and their interpretation (Fig. S4), as we show in detail below.

489 The conversion of both pyrite and siderite to Fe (oxyhydr)-oxides during
490 sample oxidation can cause a redistribution of Fe from Fe_{Pyr} and Fe_{Carb} to Fe_{Ox} .
491 Because these pools are summed in the Fe_{HR} pool, oxidation has little potential to
492 influence the Fe_{HR}/Fe_{Tot} or Fe/Al ratios, that are used to discriminate between oxic
493 versus anoxic conditions. However, Fe-speciation discriminates between
494 ferruginous and euxinic conditions based on Fe_{Pyr}/Fe_{HR} , with ferruginous conditions
495 indicated at conservative ratios <0.7. Given that pyrite oxidation decreases this
496 ratio, it has potential to obscure signals for euxinia. We thus tested our capacity to
497 accurately discriminate between ferruginous and euxinic conditions based on
498 Fe_{Pyr}/Fe_{HR} ratios measured in our samples by fully unpacking the wealth of
499 information in Fe-speciation analyses (Fig. S4). As one example, we summed
500 Fe_{Ox} , the product of Fe_{Pyr} oxidation, with Fe_{Pyr} to come up with a maximum possible
501 'pre-oxidation' ratio, Fe'_{Pyr}/Fe_{HR} . Fe'_{Pyr}/Fe_{HR} ratios in both the Cismon and DSDP
502 Site 463 sediments are nearly all below the conservative <0.7 threshold for the
503 delineation of ferruginous conditions (Fig. 3). As a further example, we assumed
504 all the sulfur present in the sediment was originally pyrite and calculate
505 conservative pyritized Fe values (S_{Tot}/Fe_{HR}). We find that these values are also

506 below the conservative threshold of 0.7 in both the Cismon and DSDP Site 463
507 sediments (Fig. S4). We employed an additional 3 tests (Supplementary
508 Information) and together these all demonstrate that our Fe-speciation analyses
509 are robust, and that sample oxidation had negligible, if any, effect on our ability to
510 discriminate between euxinic and ferruginous conditions (Fig. S4).

511

512 *4.2 Anoxic ferruginous conditions*

513 Fe-speciation in sediments deposited at both the Cismon and DSDP Site
514 463 sites records deposition under anoxic, ferruginous water column conditions
515 (Fig. 3). Ratios of $Fe_{HR}/Fe_{Tot} > 0.38$ in all the Cismon sediments deposited during
516 OAE1a are diagnostic of anoxic conditions in the Tethys Ocean. Some Fe_{HR}/Fe_{Tot}
517 values in sediments deposited during OAE1a at DSDP Site 463, however, are
518 below the threshold (>0.38) used to diagnose anoxia and are thus ambiguous to
519 depositional redox state based on Fe-speciation alone. We note, however, that Fe-
520 speciation analyses cannot diagnose sediment deposition under oxic conditions.
521 This is because of Fe mass-balance, which dictates that enrichment of Fe_{HR} at one
522 depositional location by necessity requires its depletion in another. Some
523 sediments must, therefore, act as a source of Fe_{HR} . It follows that Fe_{HR}/Fe_{Tot} ratios
524 < 0.38 can also result from deposition beneath an anoxic water column and such
525 ratios signify that the strength of the Fe-shuttle, and Fe-delivery to the open Pacific
526 Ocean (DSDP Site 463), likely waxed and waned throughout OAE1a. In contrast,
527 continental slope sediments, like those deposited at the Tethys site (Cismon core),
528 likely served as more persistent and effective oceanic sinks for Fe_{HR} . Sediments
529 deposited during OAE1a at both sites preserve appreciable non-pyritized Fe_{HR} ,
530 which thus signals deposition and burial in a sulfide poor, ferruginous setting.
531 Values < 0.7 are due to preservation of non-pyritized Fe_{HR} and, by definition, such
532 values require insufficient sulfide supply to pyritize the available Fe_{HR} , and this thus
533 also precludes accumulation of free sulfide and the development of euxinia. Values
534 of $Fe_{Pyr}/Fe_{HR} < 0.7$ preserved in sediments deposited under anoxic conditions
535 ($Fe_{HR}/Fe_{Tot} > 0.38$), therefore, either reflect sulfate depletion and deposition from a

536 sulfate and sulfide poor water column, or lack of sulfide production from available
537 sulfate, which we revisit below.

538 The distribution of RSTEs in sediments from both the Cismon and DSDP
539 463 sites also reveals deposition under anoxic, ferruginous conditions. Rhenium is
540 highly enriched in OAE1a sediments, and this confirms deposition under anoxic
541 conditions at both sites given tendency for Re enrichment under both ferruginous,
542 and to a lesser degree euxinic conditions (Tribovillard et al., 2006) (Fig. 3).
543 Deposition under euxinic conditions is often accompanied by strong Mo
544 enrichments and deposition under euxinic conditions can be conditionally inferred
545 when sedimentary Mo concentrations are $>25 \mu\text{g g}^{-1}$ (Lyons and Severmann,
546 2006). However, most OAE1a samples analysed here have a conspicuous lack of
547 Mo enrichment with correspondingly strong enrichments in other RSTEs (V, Cr,
548 Re) (Fig. 4). This supports the Fe-speciation data and diagnoses the deposition of
549 sediments at the Cismon and DSDP 463 sites under ferruginous, rather than
550 euxinic conditions.

551 Our conclusion for deposition of OAE1a sediments under ferruginous
552 conditions is underscored by comparisons of RSTE distributions in OAE1a
553 sediments and those deposited in euxinic regions in the modern oceans. Strikingly,
554 the general pattern of RSTE enrichments (Cr, V, Re) with lack of corresponding
555 Mo enrichment in the OAE1a sediments, strongly contrasts observations from
556 modern euxinic sediments (Bennett and Canfield, 2020) (Fig. 7). This can be seen
557 when enrichment factors (EF) of these RSTEs are normalized to Mo_{EF} in the
558 OAE1a sediments and compared to ratios of $\text{RSTE}_{\text{EF}}/\text{Mo}_{\text{EF}}$ in example modern
559 euxinic basins (Fig. 7). Most strikingly, Cr, V, and Re are on average 150, 50, and
560 10 times more enriched in the OAE1a sediments than in modern euxinic
561 sediments, relative to Mo, respectively (Fig. 7). Furthermore, concentrations of Mo
562 $>25 \mu\text{g g}^{-1}$, sporadically observed in some OAE1a sediments from other sites (Fig.
563 4e), need not reflect deposition from euxinic waters. Instead, such enrichments
564 can be achieved through a variety of processes and given the Fe-speciation and
565 OM contents of OAE1a sediments, likely reflect Fe-oxide and OM-Mo shuttling to
566 the sediment (Hardisty et al., 2018). We note that development of global euxinia

567 for a brief interval at the initiation of OAE1a, and/or hydrological restriction and
568 reservoir effects, have potential to induce widespread drawdown in seawater
569 RSTE inventories (Tribovillard et al., 2006), possibly contributing to muted Mo_{EF} .
570 Such drawdown, however, doesn't explain the broader pattern of RSTE
571 distributions given that several non-Mo RSTEs are strongly enriched. Instead, the
572 RSTE enrichment patterns of OAE1a sediments must reflect pervasive anoxic
573 conditions throughout OAE1a and given known mechanisms for differential RSTE
574 enrichments, and our Fe-speciation results, the enrichments in Cr, V, and Re are
575 best explained by deposition from ferruginous oceans.

576

577 *4.3 Surface-ocean sulfate scarcity during OAE1a*

578 In organic matter (OM)-rich settings, like those that supported deposition of
579 OAE1a sediments, residual sulfate is incompatible with $Fe_{Pyr}/Fe_{HR} < 0.7$. This is
580 because under anoxic conditions, available OM supports sulfate reduction and
581 corresponding sulfide production until either sulfate or OM is exhausted.
582 $Fe_{Pyr}/Fe_{HR} < 0.7$ can develop in scenarios where there is insufficient OM to fuel
583 sulfate reduction, as is the case in many modern hydrothermal sediments (e.g.,
584 Southeast Pacific Rise (SEPR)), which are different than the sediments deposited
585 during OAE1a. The two scenarios can be easily distinguished by both the sediment
586 OM content (e.g., OAE1a > 8 wt%, SEPR < 0.01 wt%), and by the speciation of non-
587 pyritized Fe_{HR} . Appreciable OM tends to fuel reduction of Fe_{HR} and conversion of
588 ferric (oxyhydr)-oxide phases to minerals like siderite and magnetite, whereas lack
589 of OM tends to preserve ferric (oxyhydr)-oxides as minerals like goethite or
590 hematite, as in the goethite dominated sediments of the SEPR (Poulton and
591 Canfield, 2006). Sulfate reduction and resulting pyritization of Fe_{HR} during
592 deposition of OAE1a sediments was, therefore, not OM-limited, and instead must
593 have been sulfate limited. This conclusion is underscored by the appreciable
594 fraction of Fe_{HR} preserved as Fe_{Carb} (Supplementary Data), which operationally
595 reflects the mineral siderite, the presence of which was also unambiguously
596 demonstrated through X-ray diffraction analyses (Fig. S2). Notably, the
597 preservation of siderite also requires depletion of sulfate since siderite is rapidly

598 ($t_{1/2} = 22$ minutes) converted to pyrite when exposed to sulfide (Berner, 1981;
599 McAnena, 2011). All rocks deposited at the Cismon and DSDP 463 sites during
600 the OAE1a interval have $Fe_{Pyr}/Fe_{HR} < 0.7$, and host appreciable $Fe(II)_{HR}$ phases.
601 The concurrent preservation of abundant sedimentary OM and non-pyritized
602 $Fe(II)_{HR}$ thus demonstrates that pyritization was limited by sulfate availability and
603 implies sulfate drawdown to negligible concentrations.

604 Surface ocean seawater sulfate concentrations were below $300 \mu\text{M}$ and
605 potentially less than $100 \mu\text{M}$ during the deposition of OAE1a sediments. Using
606 rates of microbial sulfate reduction and physical transport parameterized based on
607 modern marine organisms and ecosystems (Table S2), 1D water-column modeling
608 across scenarios that capture the entire possible parameter space yields estimates
609 for seawater sulfate that were less than $\sim 300 \mu\text{M}$ in all scenarios, and less than
610 $100 \mu\text{M}$ under most reasonable scenarios (Fig. 6). Importantly, imposing higher
611 sulfate concentrations in this model with realistic transport across oceanic
612 pycnoclines and sediment accumulation rates, yields pyrite burial fluxes much
613 higher than those recorded in OAE1a sediments (conservative maximum = 10
614 $\text{mmol m}^{-2} \text{yr}^{-1}$) (Fig. 6). Other parameter values at higher sulfate concentrations
615 and appropriately low pyrite burial rates leave residual sulfate in the water column
616 and are thus incompatible with the preservation of non-pyritized $Fe(II)_{HR}$, in OM-
617 rich sediments, and sediment deposition under ferruginous conditions (Fig. 6).

618

619 *4.4 Seawater sulfate dynamics in the Aptian age*

620 Our observation of ferruginous conditions during OAE1a requires a decline
621 of the seawater sulfate pool from low mM to hundreds of μM concentrations over
622 an interval on the order of 50 kyrs, commensurate with the rapid onset of
623 ferruginous conditions during OAE1a, as delineated by the Fe-speciation and C-
624 isotope records (Fig. 2). Seawater sulfate concentrations were drawn down to as
625 low as 1 mM preceding OAE1a as the result of evaporite deposition (Wortmann
626 and Chernyavsky, 2007). Evaporite deposition effectively draws sulfate down to 1
627 mM , but when sulfate concentrations drop below 1 mM , seawater saturation with
628 respect to sulfate-minerals (gypsum) during evaporation requires unrealistically

629 high Ca^{2+} concentrations that are inconsistent with Aptian seawater chemistry
630 (Timofeeff et al., 2006) (Fig. S5). Our observation of ferruginous conditions and
631 sub-mM surface ocean seawater sulfate concentrations thus effectively rule out
632 the deposition of OM-rich shales at the same time as the formation of evaporites
633 containing abundant gypsum. This requires that evaporitic gypsum deposition
634 likely took place before the OAE1a interval, in the Late Barremian to early Aptian,
635 as previously considered (Davison, 2007). Seawater sulfate drawdown to <300
636 μM , instead, requires a second phase of sulfate sequestration through microbial
637 sulfate reduction and pyrite deposition and burial. Expansion of the pyrite S-sink
638 during OAE1a provides a plausible mechanism to lower seawater sulfate
639 concentrations and drive subsequent development of ferruginous conditions.

640 Enhanced pyrite burial associated with expansion of oceanic anoxia offsets
641 strong hydrothermal and weathering inputs of S to the oceans during OAE1a and
642 contributes to sulfate drawdown. Sulfur mass balance modeling reveals that an
643 increase in global pyrite deposition rates from 0.66 to a maximum of 2.5 ± 0.6 Tmol
644 yr^{-1} (Fig. 8), partly offsets enhanced S-inputs and contributes to sulfate drawdown,
645 while remaining consistent with the S-isotope record (Fig. 8). We note that pyrite
646 deposition rates are higher under anoxic conditions and indeed pyrite burial rates
647 are elevated by up to a factor of ~ 10 during OAE1a (Fig. 2). Such enhanced pyrite
648 burial could be achieved by expanding the global extent of water column anoxia.
649 For example, if water column anoxia expanded from 0.1% , its extent in the modern
650 ocean (Martin et al., 1987), to up to $\sim 10\%$, the increase from 0.66 to 2.5 ± 0.6 Tmol
651 yr^{-1} could be achieved with area specific pyrite deposition rates of 0.07 ± 0.02 mol
652 $\text{S m}^{-2} \text{ yr}^{-1}$ in regions of ocean anoxia, which is similar to rates of pyrite burial in
653 sediments underlying modern OMZs (Fig. S6). While enhanced pyrite burial during
654 OAE1a can remove some S (Fig. 8), the potentially strong input fluxes from
655 hydrothermalism and weathering require an additional S-sink to draw seawater
656 sulfate concentrations down to levels that support ferruginous conditions.

657 Bio-sulfur is an additional, yet often overlooked, sulfur sink that when
658 combined with pyrite burial, is sufficiently large to draw seawater sulfate down to
659 <300 μM . Sinks such as bio-sulfur, CAS, or barite associated with OM, are all

660 known to operate in low sulfate modern and ancient environments (Horner et al.,
661 2017; Paris et al., 2014). While bio-sulfur is a major pathway for S burial in
662 lacustrine environments, it is often neglected in the marine S cycle. Marine
663 organisms have molar C/S ratios between ~30 and ~50 (Chen et al., 1996;
664 Fagerbakke et al., 1996), and thus the burial flux of organic S is likely on the order
665 of 5 to 10 wt% that of organic C. Marine organisms, furthermore, assimilate S with
666 a $\delta^{34}\text{S}$ composition nearly identical to seawater (Werne et al., 2003), and indeed,
667 OAE1a S-isotope mass balance requires non-pyrite sinks that carry near seawater
668 $\delta^{34}\text{S}$ values (Fig. 8). Given the total S and OM contents of the Cismon sediments
669 deposited during OAE1a (Bottini et al., 2012), and assuming all non-pyrite S is
670 buried as bio-sulfur, we calculate C:S molar ratios between 5 and 49 (average of
671 24), revealing that OM buried during OAE1a has a similar C:S composition to that
672 of modern bio-sulfur. OM burial during OAE1a increased by an average factor of
673 ~10 (range = 1 – 40x) at the Cismon and DSDP 463 sites (Fig. 2). If this OM was
674 ~5 wt% S, a conservative 10-fold increase in OM burial during OAE1a ($\sim 3.5 \pm 0.9$
675 Tmol S yr^{-1}) would be, together with pyrite burial ($2.5 \pm 0.6 \text{ Tmol S yr}^{-1}$), sufficient
676 to draw the marine sulfate reservoir down to $550 \pm 290 \mu\text{M}$ during OAE1a (Fig. 8),
677 within error of estimates from our 1D model (Fig. 6).

678 Although uncertainties in Cretaceous S-budgets and fluxes are large, our
679 modeling estimates plausible seawater sulfate concentrations within available
680 constraints. Importantly, our model results reproduce $\delta^{34}\text{S}_{\text{CAS}}$ records and S burial
681 fluxes in OAE1a sediments at seawater sulfate concentrations that are consistent
682 with ferruginous conditions. The development of ferruginous conditions can,
683 therefore, be attributed to widespread oceanic anoxia and ensuing sulfate
684 drawdown through pelagic sulfate reduction and enhanced burial of bio-sulfur, all
685 against the backdrop of low Cretaceous seawater sulfate concentrations, strong
686 hydrothermalism, and weathering. Furthermore, at a seawater sulfate
687 concentration of $\sim 250 \mu\text{M}$ and peak S-fluxes in our mass balance model, we
688 calculate a residence time for seawater sulfate of ~ 55 kyrs, which is an order of
689 magnitude longer than the modern ocean mixing time (~ 3 kyrs). Relative

690 homogeneity in Aptian $\delta^{34}\text{S}_{\text{CAS}}$ records is thus expected, even with seawater
691 sulfate concentrations well below 1 mM.

692

693 **5.0 Implications and conclusions**

694 At 28 mM, seawater sulfate is an oxidant pool twice the size of modern
695 atmospheric O_2 . A decline to below 300 μM seawater sulfate concentrations
696 consequently indicates a large-scale reorganization of global oxidant pools during
697 OAE1a with implications for marine ecology, biogeochemical cycling, and climate.
698 Water column anoxia, for example, may have extended at least transiently into the
699 photic zone during OAE1a with potential to influence photosynthetic ecology.
700 Indeed, biomarkers indicative of green S-bacteria (phylum Chlorobi) have been
701 recovered in sediments deposited during OAE1a (van Breugel et al., 2007).
702 Preservation of biomarkers from green S-bacteria in Aptian ferruginous sediments
703 could signal the return of photoferotrophy to the Phanerozoic oceans, as they are
704 known to grow on ferrous Fe (photoferotrophy) (Crowe et al., 2008). Low-sulfate,
705 ferruginous oceans would also have channelled OM degradation from sulfate
706 reduction to methanogenesis and limited the consumption of this methane through
707 anaerobic oxidation with sulfate (Reeburgh, 2007). Seawater sulfate
708 concentrations $<300 \mu\text{M}$ could thus promote marine methane efflux to the
709 atmosphere with potential greenhouse warming.

710 Development of ferruginous conditions during OAE1a reveals large-scale
711 dynamics in Earth's biogeochemical cycles over intervals of <100 kyrs. The
712 development of ferruginous ocean conditions during multiple OAEs may thus
713 signify a general instability in Earth surface redox budgets and the recurrent
714 reorganization of major oxidant pools at Earth's surface, like seawater sulfate,
715 during the Phanerozoic Eon. In particular, emerging reconstructions of seawater
716 sulfate concentrations during multiple Phanerozoic OAEs, suggest repeated
717 collapse of the seawater sulfate reservoir to below 1 mM, possibly below 100 μM ,
718 over short timescales (<50 kyrs) (He et al., 2020). The mechanisms driving this
719 reorganization remain uncertain but could be addressed through better constraints
720 on global S-budgets and the drivers of ocean deoxygenation, as well as Earth

721 system modeling that resolves such biogeochemical dynamics over relatively short
722 time scales.

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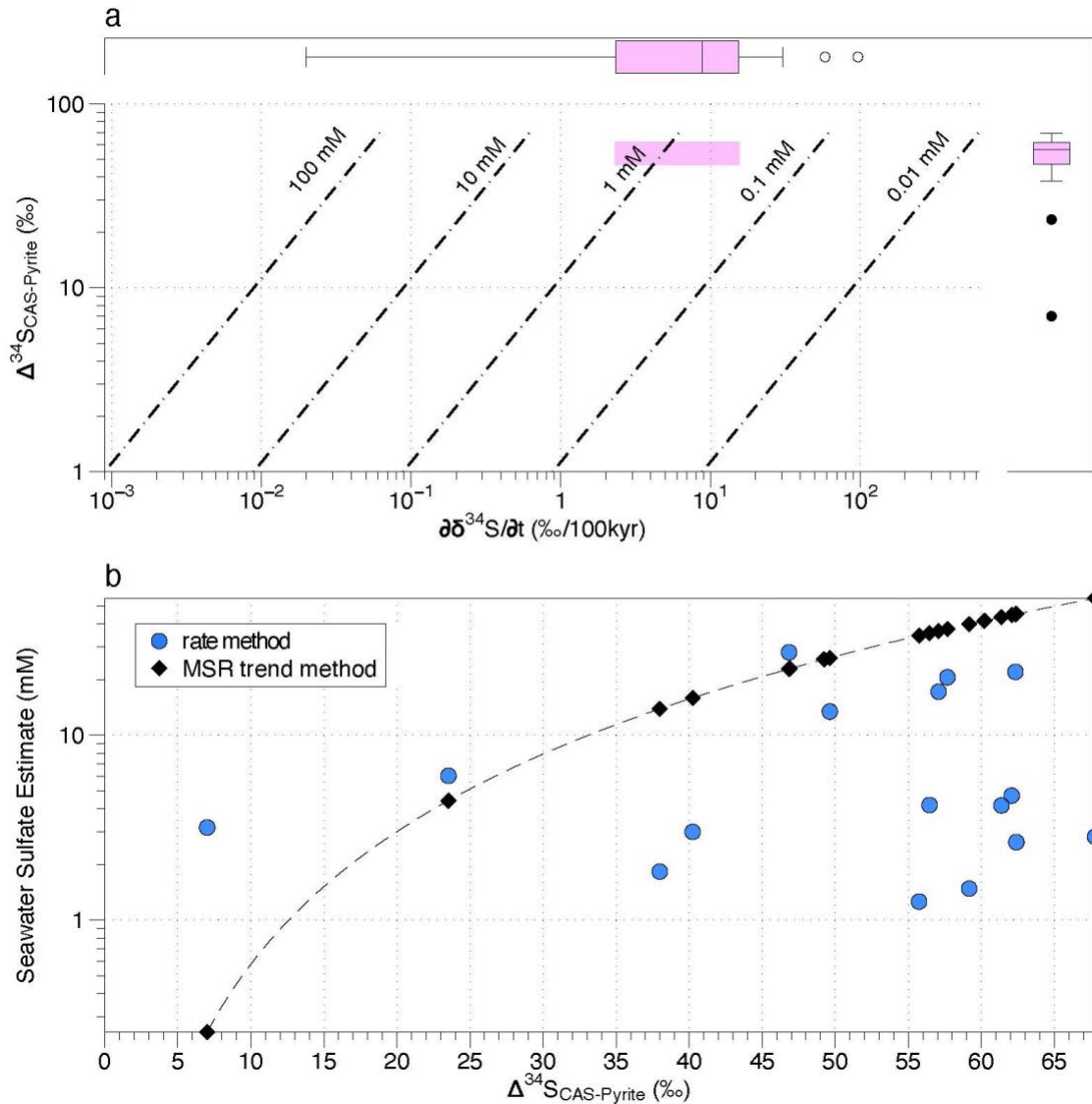
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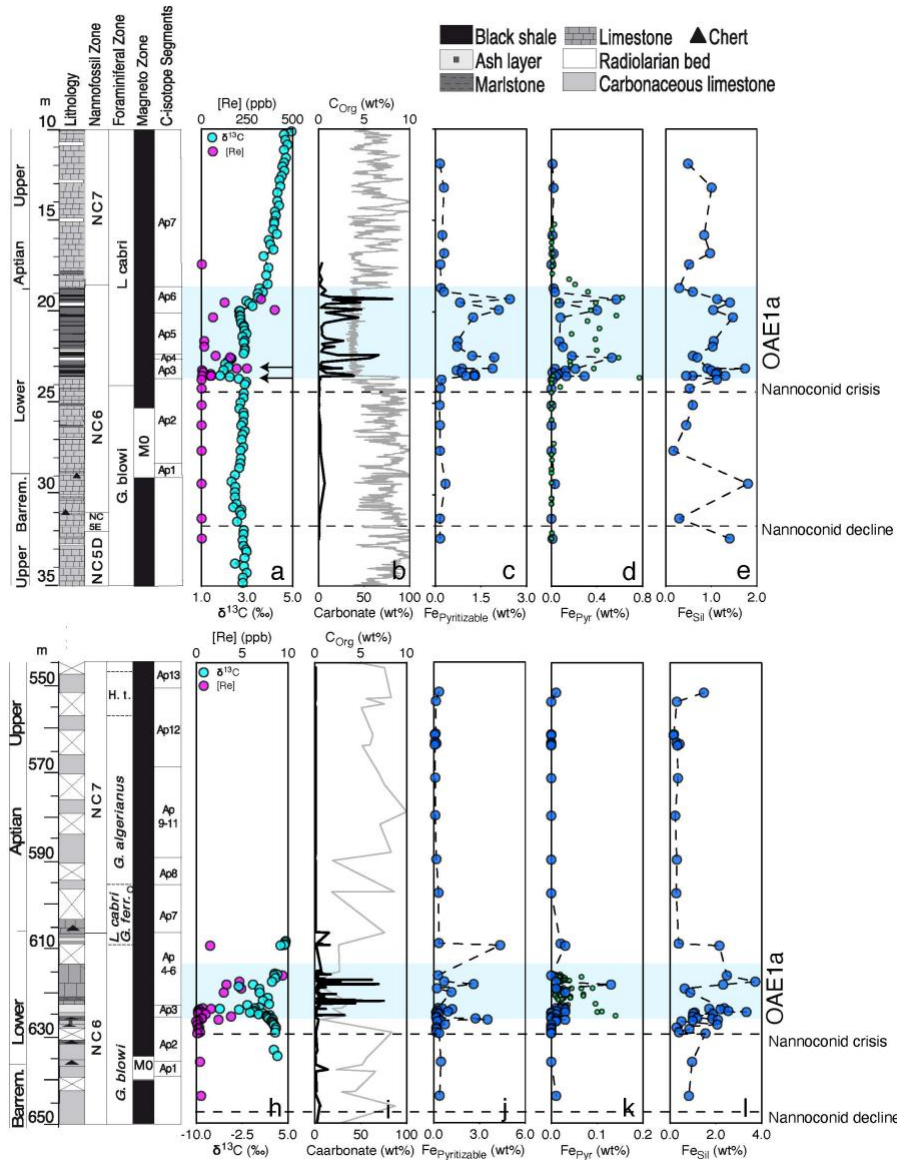
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734 **Figures and captions**



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Figure 1. (a) Rate method estimates for early Aptian seawater sulfate concentrations using the approach presented in (Algeo et al., 2015). Seawater sulfate concentrations were calculated using a S-pyrite burial flux of $1.3 \times 10^{12} \text{ mol yr}^{-1}$. The x-axis box and whisker plot represent $\partial \delta^{34}\text{S}_{\text{CAS}} / \partial t$ calculated using Aptian S-isotope data from (Gomes et al., 2016; Kristall et al., 2018; Mills et al., 2017). The y-axis box and whisker plot represents $\Delta^{34}\text{S}_{\text{CAS-Pyrite}}$ from (Gomes et al., 2016; Kristall et al., 2018; Mills et al., 2017) The pink shaded region represents the parameter space where the ranges between the 25th and 75th percentiles of the $\Delta^{34}\text{S}_{\text{CAS}}$ and $\partial \delta^{34}\text{S}_{\text{CAS}} / \partial t$ distributions overlap, corresponding to a likely range of range of seawater sulfate estimates. **(b)** All data points represent paired cogenetic S-isotope data for Aptian sulfate ($\delta^{34}\text{S}_{\text{CAS}}$) and sulfide ($\delta^{34}\text{S}_{\text{Pyrite}}$) (Gomes et al., 2016). For the rate method, seawater sulfate concentrations were calculated using a S-pyrite burial flux of $1.3 \times 10^{12} \text{ mol yr}^{-1}$. For the MSR trend method, the values were calculated using the relationship presented in (Algeo et al., 2015) (Supplementary Information).



751
 752 **Figure 2.** Fe-speciation and carbon isotope records for Cison and DSDP Site 463 cores.
 753 Integrated stratigraphy of the Cison and DSDP Site 463 after (Bottini et al., 2015; Erba
 754 et al., 2010). The grey shaded region (OAE1a) represents ~1.1 Myrs (Malinverno et al.,
 755 2010), C-isotope stages C2–C7 (Bottini et al., 2015). Panels (a–e) are Cison data and
 756 panels (f–j) are DSDP Site 463 data. **(a)** Cison C-isotope data from (Bottini et al., 2015).
 757 Rhenium concentration data from (Bottini et al., 2012). **(b)** Carbonate C, and organic
 758 matter C data after (Bottini et al., 2012; Bottini et al., 2015). **(c)** $Fe_{Pyritizable}$; sum of all
 759 pyritizable Fe_{HR} pools (Fe_{Carb} , Fe_{OM} , Fe_{Ox}) **(d)** Fe_{Sil} ; silicate Fe **(e)** Fe_{Pyr} ; pyrite Fe. Green
 760 data represent pyrite concentration data from (Gomes et al., 2016). **(f)** DSDP Site 463 C-
 761 isotope data from (Bottini et al., 2012). Rhenium concentration data from (Bottini et al.,
 762 2012). **(g)** Carbonate C data from (Bottini et al., 2015), organic C data from (Bottini et al.,
 763 2015; van Breugel et al., 2007). **(h)** $Fe_{Pyritizable}$; sum of all pyritizable Fe_{HR} pools (Fe_{Carb} ,
 764 Fe_{OM} , Fe_{Ox}) **(i)** Fe_{Sil} ; silicate Fe. **(j)** Fe_{Pyr} ; pyrite Fe. Green data represent pyrite
 765 concentration estimates from (Mélières et al., 1981). The start of the nannoconid decline
 766 and beginning of the nannoconid crisis are marked with dashed lines (Erba et al., 2010).

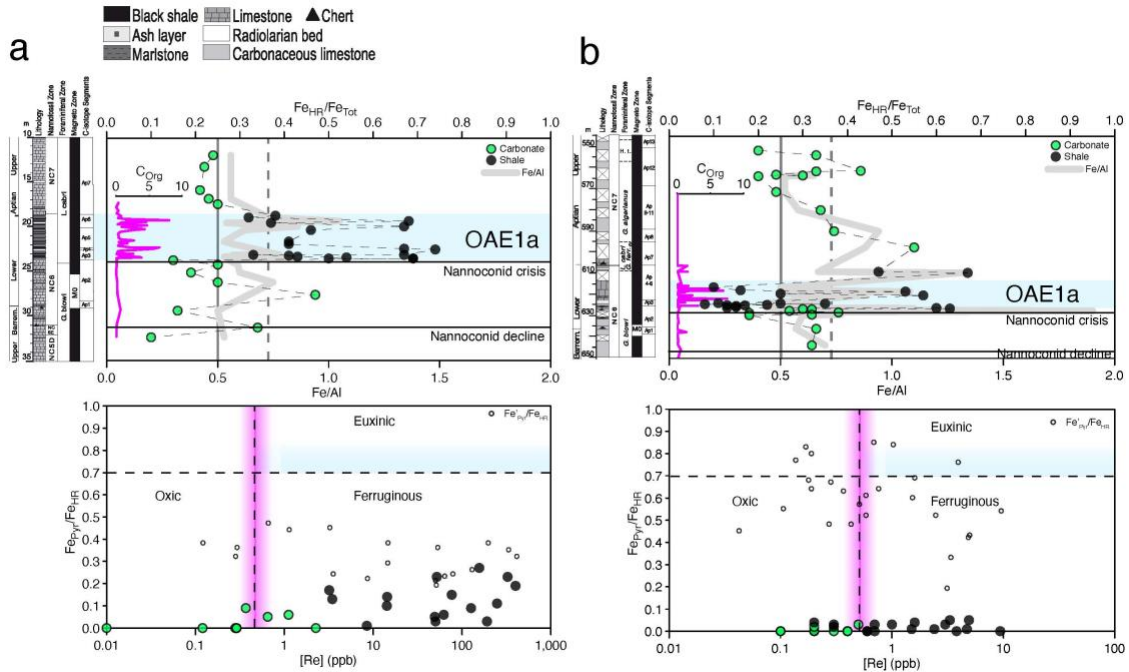
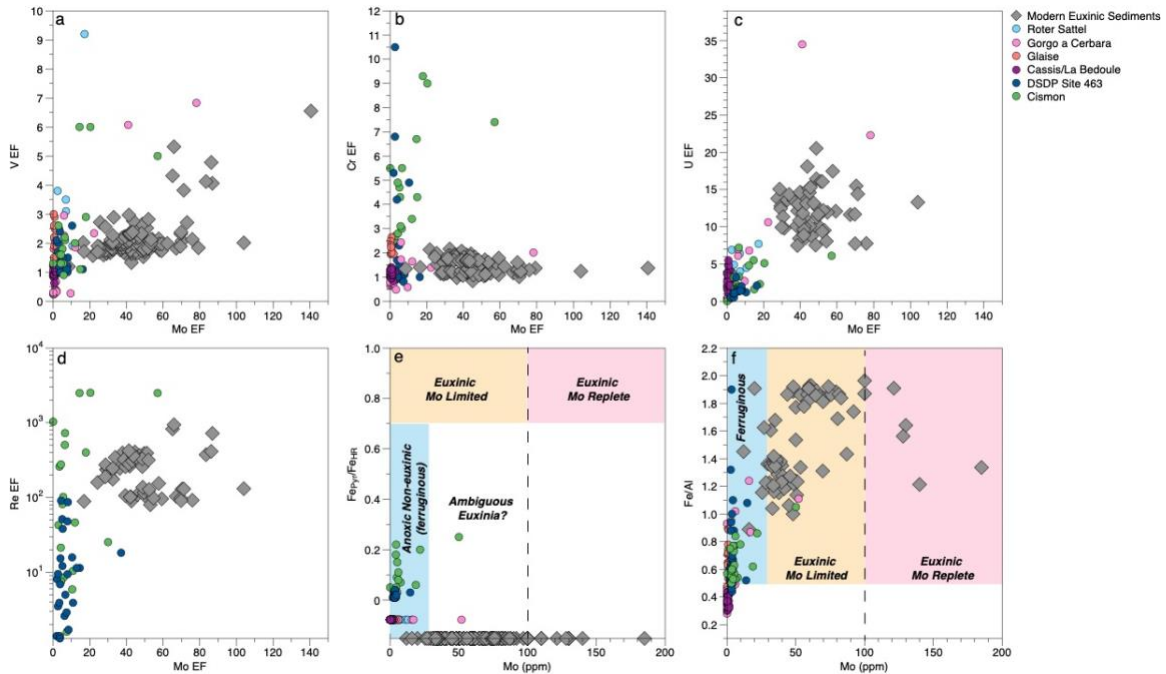


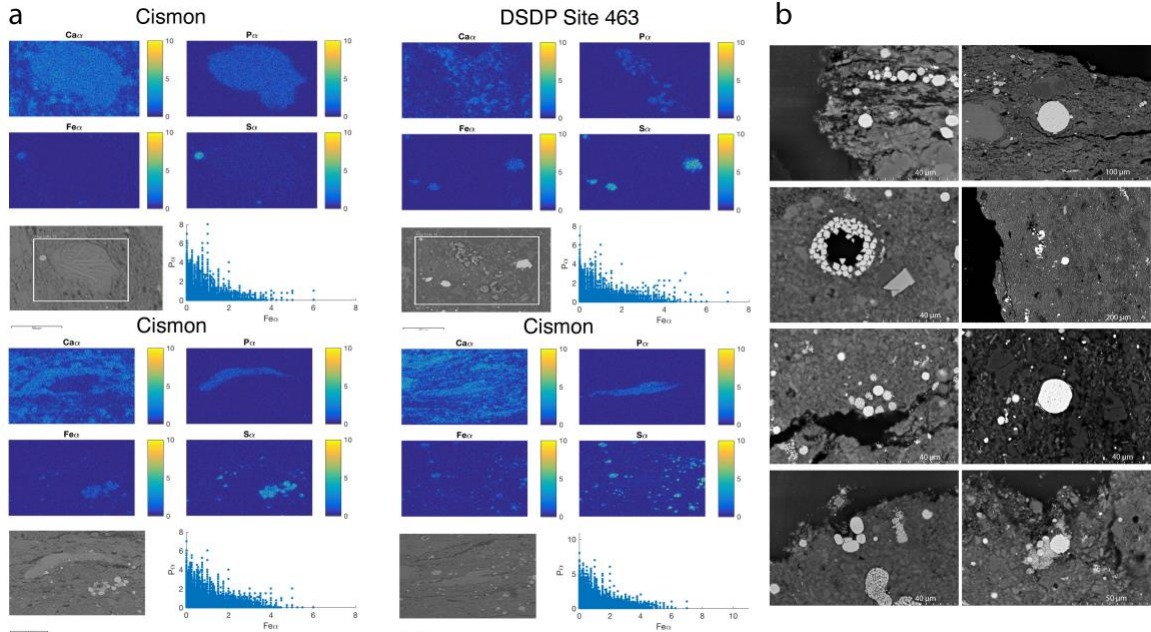
Figure 3. (a) Fe-speciation and Fe/Al records of the Cismon and DSDP Site 463 **(b)**. The vertical and horizontal dotted lines refer to the oxic-anoxic threshold ($Fe_{HR}/Fe_{Tot} = 0.38$) and a conservative ferruginous-euxinic threshold ($Fe_{Pyr}/Fe_{HR} = 0.70$), respectively. The open circles represent Fe_{Pyr}^1/Fe_{HR} , the amount of pyrite present in the samples assuming the unlikely scenario where the entire Fe_{Ox} pool is a result of Fe_{Pyr} oxidation. The solid vertical line in the top panels refers to the Fe/Al ratio of 0.5. The solid vertical line in the bottom panels refers to the average rhenium concentration of the PAAS, with the purple shading representing a 2 sigma uncertainty on this value (McLennan, 2001). Litho-, bio- and magneto-stratigraphy is the same as Figure 1.

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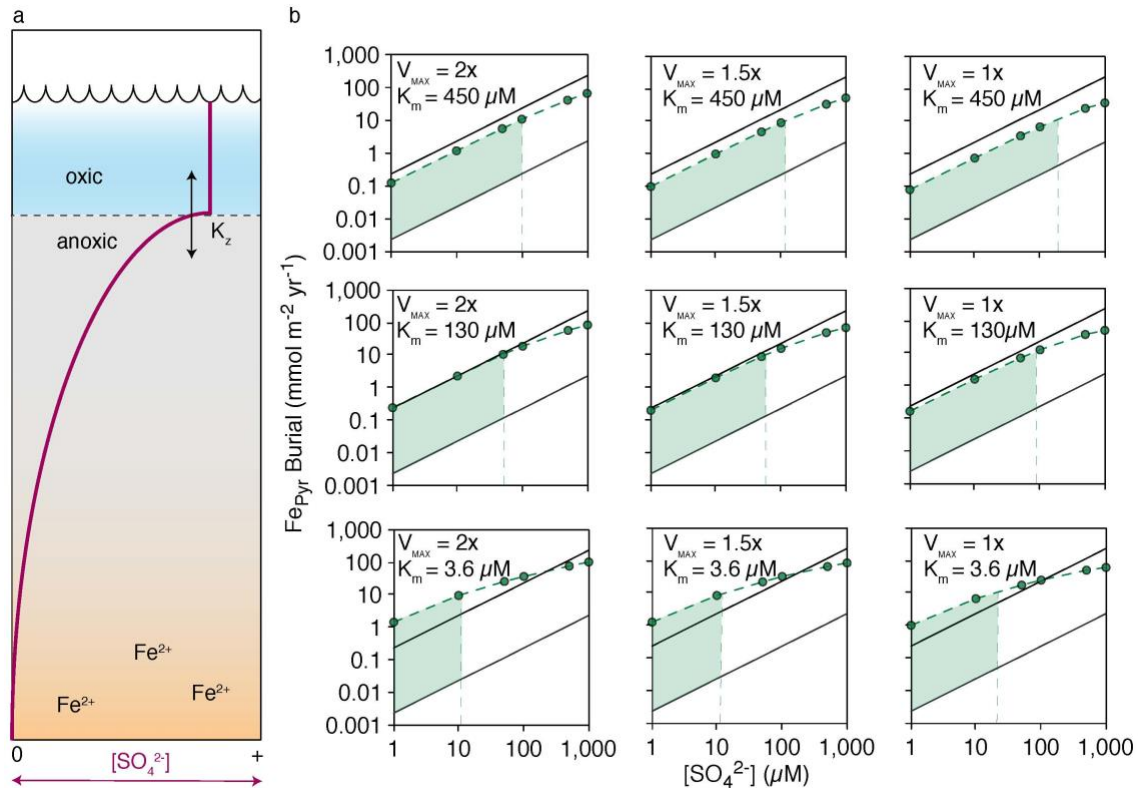
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Figure 4. RSTE enrichment factors (EFs) and molybdenum concentrations. EFs were calculated by normalizing to the PAAS. Data from other OAE1a sedimentary sections as follows; Gorgo a Cerbara, Glaise, and Cassis la Bédoule (Follmi, 2012; Westermann et al., 2013), Roter Sattel (Charbonnier et al., 2018). All modern environment RSTE data come from hydrologically restricted euxinic basins (Bennett and Canfield, 2020). **(a)** V EFs vs Mo EFs. **(b)** Cr EFs vs. Mo EFs. **(c)** U EFs vs. Mo EFs. **(d)** Re EFs vs. Mo EFs. **(e)** Fe_{Pyr}/Fe_{HR} vs. Mo concentrations. The data for other OAE1a sections and modern environments do not have corresponding Fe_{Pyr} values and thus these data plot on the x-axis. All OAE1a data points (2 analyses excepted) exhibit Mo concentrations that unambiguously demonstrate non-euxinic conditions. **(f)** Fe/Al vs. Mo concentrations. Fe/Al ratios >0.5 imply anoxic conditions (Lyons and Severmann, 2006). All OAE1a data points that plot above this threshold (2 analyses excepted) exhibit Mo concentrations that unambiguously demonstrate ferruginous (non-euxinic) conditions.



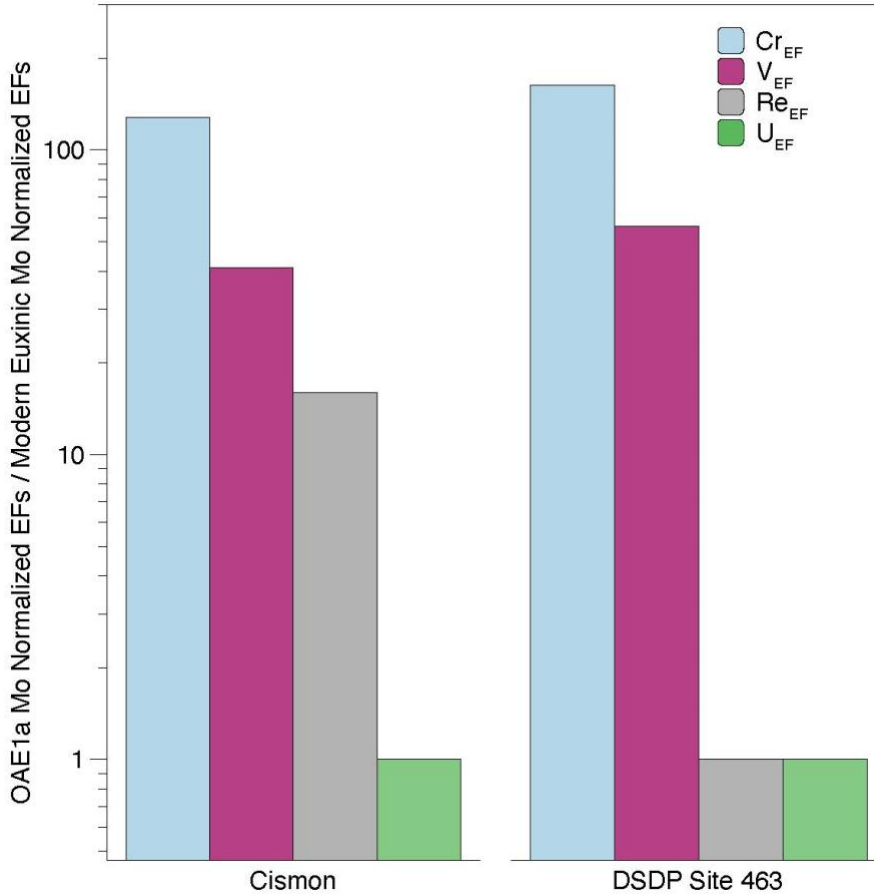
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 821 **Figure 5. (a)** SEM-EDS analyses of Ca-P bearing minerals (apatite) in the Cismon and
 822 DSDP Site 463 sediments. We observe a tight coupling between the distribution of Ca and
 823 P in the OAE1a samples, and no relationship between the distribution of Fe and P,
 824 implying no redistribution of P into freshly formed poorly crystalline Fe-(oxyhydr)oxide
 825 phases and a lack of the pyrite oxidation mechanism observed by Kraal et al. (2009). **(b)**
 826 Compilation of back scatter electron (BSE) SEM images obtained on thin sections of
 827 OAE1a samples from both the Cismon and DSDP Site 463 sediments. We observe no
 828 strong evidence for pyrite oxidation, even on crystals located near thin section boundaries
 829 where oxidation would be most concentrated.

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 852 **Figure 6.** 1D water column reaction transport model sensitivity analyses. **(a)** Schematic
 853 model representation of a stratified Cretaceous water column. The model is designed such
 854 that seawater sulfate must be exhausted by the bottom of the domain (sediment-water
 855 interface) under anoxic conditions via sulfate reduction promoting the development of
 856 ferruginous conditions. **(b)** Water column depth integrated sulfate reduction rates
 857 (equivalent to pyrite burial). The data points represent different model outputs for the
 858 indicated scenarios. The solid black lines (equation = $[SO_4^{2-}]/\partial z * K_z$) mark the pyrite
 859 burial rates needed to support complete drawdown of sulfate at a given surface seawater
 860 sulfate concentration. The lower line corresponds to a $K_z = 0.01 \text{ m}^2 \text{ d}^{-1}$ and the upper line
 861 corresponds to a K_z of $1 \text{ m}^2 \text{ d}^{-1}$. The only allowable model runs thus need to plot above
 862 these lines. The green shading corresponds to the allowable parameter space that
 863 satisfies both the requirement for complete sulfate drawdown and the conservative
 864 condition for local pyrite burial rates to be less than $10 \text{ mmol m}^{-2} \text{ yr}^{-1}$ (Fig. S6). Maximum
 865 allowable sulfate concentrations come from the intersection of the green shading
 866 (extended green dashed lines) with the x-axis.

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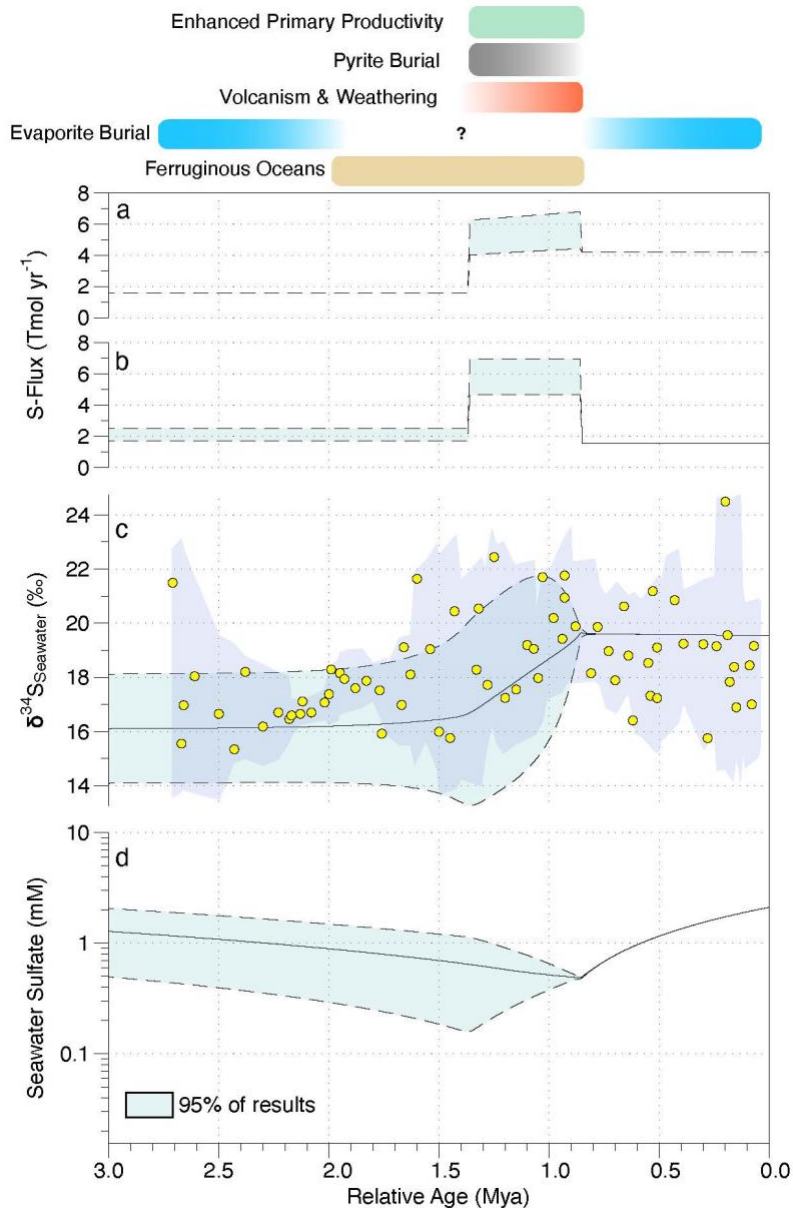


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Figure 7. OAE1a RSTE enrichment factors (EFs) compared to modern euxinic sediments. Displayed are EFs of RSTEs normalized to Mo enrichment factor in the OAE1a sediments compared to RSTE EFs normalized to Mo in example modern euxinic basins;

$\frac{(OAE1a \frac{RSTE_{EF}}{Mo_{EF}})}{(Modern\ Euxinic \frac{RSTE_{EF}}{Mo_{EF}})}$. We find that the RSTE enrichment factors of Cr, V, and Re, metals that

do not require sulfide to be buried under anoxic conditions, are dramatically enriched in both the Cison and DSDP 463 sediments, relative to modern euxinic basins. Modern data is from (Bennett and Canfield, 2020).



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Figure 8. Evolution of the Cretaceous seawater sulfate reservoir. The coloured bars represent the conceptual model and proposed timing of early Cretaceous events associated with the development of OAE1a. **(a)** Modelled S-input and **(b)** burial fluxes. Solid lines represent the mean results of the stochastic modeling (Supplementary Information), whereas the shaded region between the dashed lines encompasses 95% of the results. **(c)** Modelled $\delta^{34}\text{S}_{\text{sulfate}}$. Solid lines represent the mean results of the stochastic modeling (Supplementary Information), whereas the shaded region between the dashed lines encompasses 95% of the results. The yellow data points represent Aptian $\delta^{34}\text{S}_{\text{CAS}}$ data from (Gomes et al., 2016; Kristall et al., 2018; Mills et al., 2017). The purple shading represents a 5-point $\delta^{34}\text{S}_{\text{CAS}}$ moving average of the S-isotope data using a 2s.d. error envelope. **(d)** Modeled seawater sulfate concentrations. Solid lines represent the mean results of the stochastic modeling (Supplementary Information), whereas the shaded region between the dashed lines encompasses 95% of the results.

Acknowledgements

915

916 This work was funded through NSERC Discovery Grants to Sean A. Crowe (0487)
917 and Roger Francois, the Canadian Foundation for Innovation, the Canada
918 Research Chairs Program, and a UBC 4-Year Fellowship to Kohen W. Bauer. EE
919 and CB were funded by PRIN 2017RX9XXY to EE. We thank the International
920 Ocean Discovery Program (IODP) for providing the DSDP Site 463 samples. We
921 thank John Greenough for help with the milling of rock samples. We also
922 acknowledge Steve Calvert for participating in helpful discussions on rhenium
923 geochemistry.

924

925 **Data availability statement**

926

927 The datasets and models generated during and/or analysed during the current
928 study are available from the corresponding author on reasonable request.

929

930 **CRedit author statement**

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932 **KWB**: validation, formal analysis, investigation, data curation, writing,
933 visualization. **CB**: resources, data curation, visualization, validation, writing. **SK**:
934 validation, software, methodology, formal analysis, writing. **MJ**: validation, writing.
935 **RF**: conceptualization, validation, supervision, funding acquisition, writing. **EE**:
936 resources, visualization, validation, writing. **SAC**: conceptualization, validation,
937 supervision, project administration, funding acquisition, writing.

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