

VARIABILITY IN SULFUR ISOTOPE RECORDS OF PHANEROZOIC SEAWATER SULFATE

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Key Points

- 6710 measurements of $\delta^{34}\text{S}$ of sulfate in Phanerozoic sedimentary rocks were compiled and systematically updated to a consistent time scale
- Records derived from evaporites, barite, and carbonate-associated sulfate are similar, but also contain dramatic short-term discrepancies
- Variation created by diagenetic and depositional processes increases with age in all records, obscuring temporal trends in marine sulfate

Plain Language Summary

Sedimentary rocks deposited in ancient marine basins preserve a record of seawater composition. We compare the sulfur isotopic composition of three sedimentary materials that contain sulfate—a major ion in seawater important for carbon and oxygen cycling. Evaporite salts, the mineral barite, and trace sulfate in limestone each reveal the same first-order trends over the last 541 million years, but also display substantial shorter order discrepancies that reflect how the materials capture and store paleoceanographic information. These discrepancies partially obscure understanding of the relationship between life, ocean chemistry, and climate.

Keywords

Phanerozoic seawater composition; marine sulfate proxies; carbonate-associated sulfate; evaporites; barite; sulfur isotope ratios; kriging

25 Abstract

26 The $\delta^{34}\text{S}$ of seawater sulfate reflects processes operating at the nexus of sulfur, carbon, and
27 oxygen cycles. However, knowledge of past seawater sulfate $\delta^{34}\text{S}$ values must be derived
28 from proxy materials that are impacted differently by depositional and post-depositional
29 processes. We produced new timeseries estimates for the $\delta^{34}\text{S}$ value of seawater sulfate by
30 combining 6710 published data from three sedimentary archives—marine barite, evaporites,
31 and carbonate-associated sulfate—with updated age constraints on the deposits. Robust
32 features in multiple records capture temporal trends in the $\delta^{34}\text{S}$ value of seawater and its
33 interplay with other Phanerozoic geochemical and stratigraphic trends. However, high-
34 frequency discordances indicate that each record is differentially prone to depositional biases
35 and diagenetic overprints. The amount of noise, quantified from the variograms of each
36 record, increases with age for all $\delta^{34}\text{S}$ proxies, indicating that post-depositional processes
37 obscure detailed knowledge of seawater sulfate's $\delta^{34}\text{S}$ value deeper in time.

38 1 Introduction

39 Seawater sulfate acts as a major oxidant of organic carbon, controlling the cadence of its
40 burial in sediments and connecting the carbon, sulfur, and oxygen cycles (Bowles et al.,
41 2014; Jørgensen, 1982). Microbial sulfate reduction (MSR), reoxidation of sulfide, and the
42 burial and oxidation of pyrite govern sedimentary inorganic carbon and alkalinity fluxes
43 (Ben-Yaakov, 1973; Froelich et al., 1979). Pyrite in sedimentary rocks may be exposed and
44 oxidized during uplift, erosion, and weathering—impacting Earth's dioxygen and carbon
45 dioxide budgets on tectonic timescales (Burke et al., 2018; Kump & Garrels, 1986; M. A.
46 Torres et al., 2014). Over Phanerozoic time (the past 541 Myr), the burial of sulfide and
47 disulfide minerals must have balanced the acid produced and dioxygen consumed during
48 terrestrial pyrite weathering. Therefore, tracking ancient sulfate fluxes related to these
49 processes illuminates when, how, and where the Earth system achieves this balance, and
50 what happens during intervals of unsteadiness.

51 Thode et al. (1953) first recognized that a record of ancient marine sulfur isotopic
52 compositions ($\delta^{34}\text{S}$) could constrain changes to Earth's biogeochemical cycles, and Ault and
53 Kulp (1959) applied mass balance assumptions in an early effort to quantify important sulfur
54 fluxes. Isotope fractionations during MSR preferentially enrich the residual sulfate in ^{34}S by
55 several percent (Bradley et al., 2016; Harrison & Thode, 1958; Sim et al., 2011). When more
56 sulfate is reduced and fixed into pyrite, removing more light sulfur isotopes from the oceans,
57 the remaining sulfate in seawater becomes enriched in the heavy, rare isotopes. Holland
58 (1973) first attempted to calculate changes in dioxygen fluxes from $\delta^{34}\text{S}$ data. Holser (1977)
59 further recognized that rapid changes in the $\delta^{34}\text{S}$ value of seawater coincide with intervals of
60 biotic crises and dramatic reorganizations of Earth's climate and biosphere. The subsequent
61 forty years have seen many efforts to derive an accurate and precise record of how the $\delta^{34}\text{S}$
62 value of seawater sulfate has changed over Earth history.

63 Three sedimentary materials constitute proxy archives of Phanerozoic seawater sulfate $\delta^{34}\text{S}$
64 values: (1) marine evaporites, which include sulfate salts precipitated from evaporated
65 seawater in marginal marine basins; (2) marine barite, which forms from a suite of
66 biogeochemical processes associated with sinking particles in pelagic waters; and (3)
67 carbonate-associated sulfate (CAS), which is minor sulfate incorporated into the crystal
68 lattice of biogenic and abiogenic calcite, aragonite, and dolomite phases that accumulate in
69 sedimentary rocks.

70 Important reviews (Bottrell & Newton, 2006; Claypool et al., 1980; Holser et al., 1989;
71 Strauss, 1997; Veizer et al., 1980) on the evolution of the Phanerozoic sulfur cycle have
72 assumed that these proxies more-or-less accurately preserve the isotopic composition of
73 ancient seawater sulfate. This assumption is reasonable because Phanerozoic seawater likely
74 contained abundant sulfate as a conservative, well-mixed anion. Modern seawater has 28
75 mmol/kg sulfate, which has an approximate residence time of more than 10 Myr—much
76 longer than the mixing time of the oceans (Bottrell & Newton, 2006; Walker, 1986).
77 Supergiant gypsum and anhydrite deposits in the sedimentary record indicate that sulfate has
78 been a major constituent in ancient seawater, as well. These deposits, which represent long-
79 lived intervals of basin recharge and evaporation of seawater (Warren, 2010), formed
80 episodically from Mesoproterozoic through Phanerozoic time (Grotzinger & Kasting, 1993;
81 Pope & Grotzinger, 2003). The composition of fluid inclusions in halite from evaporite
82 deposits further suggested that sulfate maintained at least millimolar concentrations
83 throughout Phanerozoic time (Lowenstein et al., 2003).

84 Important features in the $\delta^{34}\text{S}$ age curves were observed in multiple datasets on both long
85 and short timescales. All archives exhibited high $\delta^{34}\text{S}$ values in early Paleozoic time, fell to
86 minima in the late Paleozoic, and increased to modern values ($\sim 21\text{‰}$) over Mesozoic and
87 Cenozoic time. This pattern was originally noted in the evaporite record by Ault and Kulp
88 (1959) and reaffirmed by more extensive evaporite compilations (Claypool et al., 1980;
89 Holser et al., 1989; Holser & Kaplan, 1966; Strauss, 1997). Burdett et al. (1989) produced
90 the first continuous biogenic CAS dataset for the Neogene Period and demonstrated that it
91 agreed with the evaporite $\delta^{34}\text{S}$ record. Kampschulte et al. (2001) and Kampschulte and
92 Strauss (2004) then demonstrated that biogenic CAS captured the first-order features of the
93 Phanerozoic evaporite record, and could be correlated with higher resolution and confidence
94 than evaporites to the carbonate carbon isotope record. The $\delta^{34}\text{S}$ pattern covaries with many
95 other geochemical records of changing seawater composition (Hannisdal & Peters, 2011;
96 Prokoph et al., 2008), and so has been interpreted to reflect long-term changes related to the
97 assembly and breakup of Pangea (Turchyn & DePaolo, 2019).

98 In addition to long-term trends, Holser (1977) identified shorter fluctuations (5–50 Myr) in
99 the Upper Devonian and lower Triassic evaporite record; these excursions are recorded by
100 CAS as well (Kampschulte & Strauss, 2004). Increased temporal resolution from barite and
101 CAS found additional rapid excursions, notably associated with Jurassic and Cretaceous
102 intervals of widespread organic-rich shale deposition (Gill, Lyons, & Jenkyns, 2011; Paytan
103 et al., 2004) and Paleogene carbon cycle perturbation (Paytan et al., 1998; Rennie et al.,

104 2018). In addition, some $\delta^{34}\text{S}$ records with high temporal resolution, especially derived from
105 CAS, have rapid variability (Kah et al., 2016; Kampschulte et al., 2001), and data from
106 multiple locations containing similar-age strata have $\delta^{34}\text{S}$ heterogeneity (Gill, Lyons, Young,
107 et al., 2011; Present et al., 2015).

108 Although seawater sulfate was likely well-mixed for much of Phanerozoic time, these rapidly
109 varying datasets indicated that short periods of sulfate drawdown may have been expressed
110 as high spatial and temporal $\delta^{34}\text{S}$ gradients (Holser, 1977; Kah et al., 2004, 2016). If these
111 gradients represent globally relevant budgets of carbon, nutrients, and oxidizing capacity,
112 then the residence time of sulfate in ancient oceans must have been much shorter than today.
113 An analogy to the carbon cycle is illustrative. Isotopic fractionations between oxidized and
114 reduced species are comparable for carbon and sulfur. The biological pump—
115 remineralization of sinking organic matter that is fractionated by tens of permille from
116 dissolved inorganic carbon—is only capable of creating inorganic carbon isotopic gradients
117 of less than 3‰ given Pliocene-age to present nutrient inventories and *ca.* 2 mmol/kg
118 bicarbonate (Toggweiler & Sarmiento, 1985). Therefore, even small gradients in the $\delta^{34}\text{S}$ of
119 marine sulfate, of similar magnitude to carbon isotope gradients driven by the biological
120 pump, would have required both a higher proportion of anaerobic organic carbon
121 remineralization and more than an order of magnitude smaller sulfate inventory.

122 However, the implicit assumption that proxies for seawater $\delta^{34}\text{S}$ values are suitably accurate
123 and precise to demonstrate rapid changes in seawater's composition has not been tested. The
124 processes by which the proxy materials form and incorporate sulfate from seawater may
125 affect their $\delta^{34}\text{S}$ value, complicating the reconstruction of Phanerozoic seawater's
126 composition but providing nuance on biogeochemical sulfur cycling and its imprint on the
127 rock record.

128 We produced a new timeseries to estimate the Phanerozoic history of the $\delta^{34}\text{S}$ value of
129 seawater sulfate by synthesizing published geochemical data with updated geochronology
130 and stratigraphic correlations. We attribute some of the differences between archives to
131 mechanics of how sulfate is incorporated into and preserved in sedimentary rocks. This
132 approach tests the assumption that each archive samples the same history of seawater $\delta^{34}\text{S}$
133 values, quantifies uncertainty in proxy archives, and reveals that some major sources of
134 variance are themselves produced by biogeochemical processes that may have varied through
135 Phanerozoic time.

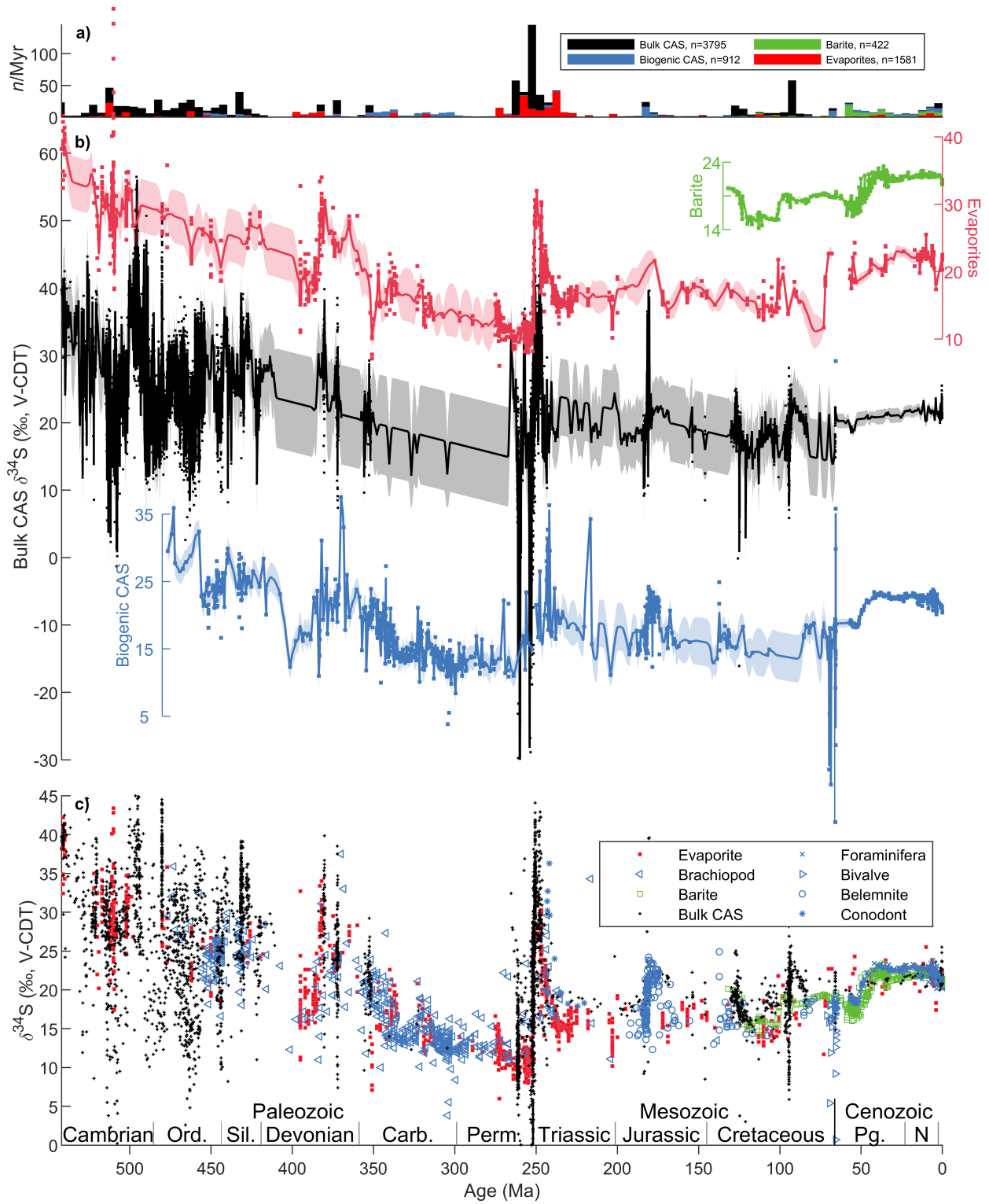
136 **2 Synthesis of Phanerozoic seawater sulfate $\delta^{34}\text{S}$ proxy data**

137 We compiled 6710 measurements from 108 references that reported $\delta^{34}\text{S}$ values in
138 Phanerozoic marine evaporites, bulk rock CAS, biogenic CAS, or marine barite. Each $\delta^{34}\text{S}$
139 value was assigned an age using the International Commission on Stratigraphy 2016/04 time
140 scale (Cohen et al., 2013; updated) (Figure 1). The Supporting Information enumerates the
141 $\delta^{34}\text{S}$ data, assigned age, data type, data source, and method and literature used for each age
142 assignment.

Each proxy material has different, irregularly spaced temporal distributions (Figure 1a). To estimate Phanerozoic $\delta^{34}\text{S}$ trends, each proxy record was interpolated at 50 kyr resolution (Figure 1b). Kriging—a geostatistical approach using autocorrelation to quantify stochastic components in spatiotemporal data—was used to weight data for interpolation and estimate confidence intervals (Gebbers, 2010). Because kriging uses the empirical autocorrelation structure of the data to produce weights, it is well suited for irregularly spaced data. Autocorrelation varies between two endmembers of linearly detrended variance: at the maximum is the variance of all points in that geologic interval, and at the minimum is the unresolved chatter between data closely spaced in time. The kriged uncertainty on the interpolations reflects this increase in variance, such that interpolated values further from data have larger uncertainties up to the population variance according to the observed range of autocorrelation. Kriging was done on each geologic material, partitioned by era, by modelling variograms—functions describing how the variance per point (semivariance) of pairs of linearly detrended data varies with their average separation distance in time (Supporting Information). Paleogeography was not considered, so spatial variability was collapsed into the temporally unresolved chatter within each era.

159

Figure 1 [next page]. Records of Phanerozoic seawater sulfate $\delta^{34}\text{S}$ generated from proxy materials. **(a)** The average number of $\delta^{34}\text{S}$ analyses of each proxy per Myr, in 5 Myr bins, illustrates the temporal bias in the sampling of each material through Phanerozoic time. **(b)** Interpolated proxy records of the $\delta^{34}\text{S}$ composition of sulfate over Phanerozoic time. Shading indicates the kriged 1σ confidence intervals. **(c)** All compiled proxy data for the $\delta^{34}\text{S}$ of Phanerozoic seawater.



167 **3 Discussion**

168 *3.1 Distribution of $\delta^{34}\text{S}$ in proxies*

169 During seventy years of effort to determine a history of Phanerozoic seawater sulfate $\delta^{34}\text{S}$
170 from different geologic materials, it has implicitly been assumed that each proxy samples the
171 same primary population of seawater $\delta^{34}\text{S}$ compositions through space and time. However,
172 comparison of all Phanerozoic $\delta^{34}\text{S}$ data for each proxy indicates that the four datasets do not
173 come from the same distribution (Supporting Information, non-parametric Kruskal-Wallis
174 one-way analysis of variance, $\chi^2[3,6709] = 684.54$, $p \ll 0.001$). Therefore, each proxy likely
175 has different temporally or spatially variable sampling biases or reflects different
176 biogeochemical processes that contribute to variance in the time-series of ancient sulfate's
177 $\delta^{34}\text{S}$.

178 Major $\delta^{34}\text{S}$ trends and excursions in Cenozoic, Mesozoic, and late Paleozoic records are
179 exhibited in multiple archives, but significant discrepancies and gaps are apparent in records
180 from Cambrian to Devonian time (Figure 1c). In early- to mid-Paleozoic strata, biogenic
181 carbonates are sparse, marine barite is absent, and bulk CAS $\delta^{34}\text{S}$ values diverge from
182 evaporites by greater than 10‰ (Figure 2a). Additionally, Paleozoic variance is highest for
183 all records (Figure 2b).

184 The evaporite record, being comprised of massive amounts of sulfate but limited in spatial
185 and temporal extent, likely captures long-term $\delta^{34}\text{S}$ trends. The bulk CAS, biogenic CAS,
186 and barite records have higher temporal resolution than the evaporite record for much of the
187 Phanerozoic, potentially capturing shorter $\delta^{34}\text{S}$ excursions.

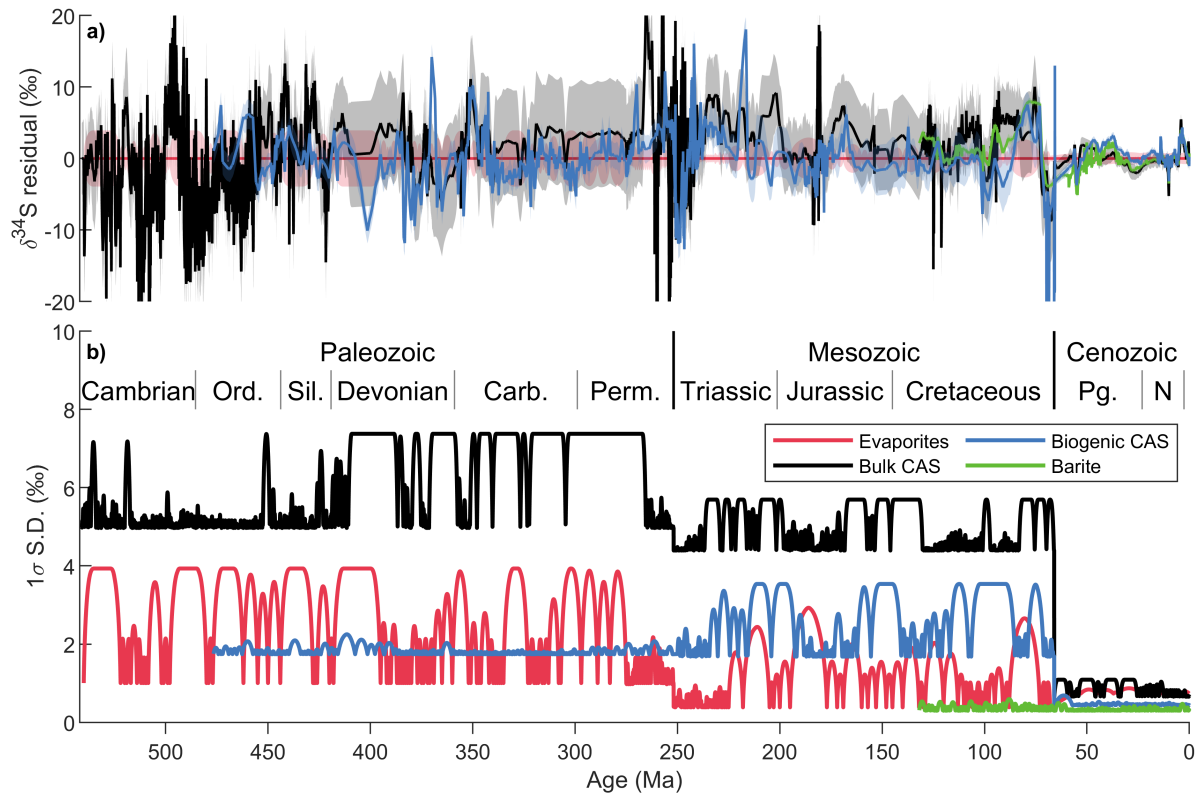


Figure 2. Comparison of $\delta^{34}\text{S}$ values and variance generated from proxy materials. **(a)** Residuals between the evaporite record and each other record shaded with kriged 1σ confidence intervals. **(b)** Confidence intervals produced from kriging data from each proxy in each era. Where data is sparse, the confidence intervals approach the standard deviation of linearly detrended data in each geologic era, excluding 1st and 99th percentile outliers. Where there is data, the confidence interval is the uncorrelated chatter determined from the semivariance of data temporally closer than the mean minimum time between data pairs.

3.2 Sources of $\delta^{34}\text{S}$ variance

The $\delta^{34}\text{S}$ variability for each proxy is plotted in Figure 2b. The maxima in each era on each curve represents the standard deviation of detrended $\delta^{34}\text{S}$ data over each geologic era. For example, the standard deviation of linearly detrended Paleozoic bulk CAS data is 7.4‰ (excluding 1st and 99th percentile outliers), while that of all Cenozoic barite data is 1.3‰. These standard deviations can be interpreted as a naive description of expected variability where data is sparse, and reflect the combination of local spatiotemporal trends in the proxy record plus an uncorrelated random component. The uncorrelated, random component is estimated by the semivariance of pairs of data that are closer together than the mean

206 minimum time between all pairs of data (Gebbers, 2010). The uncorrelated variances for
207 each proxy are plotted as the minima in each era on each curve in Figure 2b.

208 Uncorrelated variance is a metric that convolves multiple sources of uncertainty. Sources of
209 variance of geologic interest include temporally unresolved variability in seawater $\delta^{34}\text{S}$
210 values and temporally incoherent variability in how the sedimentary archives were formed
211 or altered. These sources of variance may be temporally unresolved due either to spatial
212 variability of seawater sulfate's $\delta^{34}\text{S}$ at a given time, or to temporal variability more rapid
213 than the resolution of the record. In addition, the uncorrelated variance captures analytical
214 uncertainty related to making $\delta^{34}\text{S}$ measurements in each archive, and non-systematic error
215 in age assignments of proxy materials. While the relative contributions of each of these
216 sources of uncertainty may differ between proxies or with age, the uncorrelated variance
217 metric—like the population variance—describes the data's structure and how predictive a
218 given $\delta^{34}\text{S}$ measurement is of other nearby values.

219 This analysis produced two key results: the uncorrelated variance is different for each
220 archive, and for all archives it increases with age. Cenozoic and Mesozoic CAS data have
221 uncorrelated variance larger than that of evaporites and barite. Uncorrelated Paleozoic bulk
222 rock CAS data have a standard deviation more than twice that of biogenic CAS and
223 evaporites. Differences between multiple proxies of the same age indicate that the
224 uncorrelated variance is likely caused, in part, by variability inherent to how $\delta^{34}\text{S}$ is
225 preserved, rather than just inadequate sampling of primary spatial and temporal variability
226 of seawater sulfate.

227 The remaining analysis considers the sources of variance in each archive that may have
228 contributed to the uncorrelated variance. Importantly, trends statistically distinguishable
229 from the uncorrelated variance need not represent true trends in the $\delta^{34}\text{S}$ of Phanerozoic
230 seawater. The same sources of variance controlling the uncorrelated data may themselves
231 have spatial or temporal components that lead to biased estimates of Phanerozoic seawater's
232 composition in the proxy records. The uncorrelated variance, in part, quantifies the
233 disagreement between contemporaneous records from different localities. Trends in the data
234 smaller than the uncorrelated variance are indistinguishable from random noise. This is true
235 even for individual records from stratigraphic successions with coherent $\delta^{34}\text{S}$ trends: a given
236 stratigraphic succession may clearly resolve a trend in the $\delta^{34}\text{S}$ of the proxy but fail to
237 statistically resolve a global trend in the $\delta^{34}\text{S}$ of seawater sulfate.

238 3.2.1 Evaporites

239 Deposits of carbonate, sulfate, and halide salts form as seawater evaporates in restricted
240 basins. Throughout Phanerozoic time, bedded marine evaporites formed subaqueously, in
241 salinas (hypersaline lagoons) and salt pans, and subaerially, in supratidal sabkha
242 environments. Extremely thick (>100s of meters) evaporite deposits have also formed in
243 deeper-water environments. Deposition and preservation of evaporites require favorable
244 climatic and tectonic conditions where restricted basins experience net evaporation (Warren,

245 2010). Therefore, the evaporite record has limited spatial and temporal continuity (Claypool
246 et al., 1980; Strauss, 1997).

247 Because evaporites are massive products of seawater sulfate, they are largely expected to
248 provide an accurate proxy for the $\delta^{34}\text{S}$ of ancient seawater sulfate. However, because they
249 form in marginal marine environments often with biologically adverse salinities, it can be
250 difficult to constrain their geologic age with biostratigraphy. In many deposits, it is also
251 challenging to discern depositional environment or deconvolve marine and non-marine
252 geochemical signatures (Hardie, 1984; Kendall & Harwood, 1989; Lu & Meyers, 2003). The
253 restricted, marginal marine settings in which many evaporites form are prone to changes in
254 fluid source or depositional environment with minor base-level changes (Playà et al., 2007).
255 Basins rich in evaporites also often form diapirs that drive salt tectonics, which complicates
256 a deposit's internal stratigraphy (Nielsen, 1989).

257 Evaporites can have a $\delta^{34}\text{S}$ range of 1‰ to 6‰ within a formation (Thode & Monster, 1965).
258 This variability cannot be attributed to fractionation during gypsum precipitation, which
259 produces sulfate salt prior to halite saturation that has a $\delta^{34}\text{S}$ composition 1‰ to 2‰ higher
260 than the unevaporated seawater (Raab & Spiro, 1991). Salinity stratification in evaporating
261 basins can promote water-column anoxia and allows MSR to distill sulfate to higher $\delta^{34}\text{S}$
262 compositions than the original seawater; in some cases, evaporite $\delta^{34}\text{S}$ compositions are
263 higher than other proxies from the same depositional basin (Fike & Grotzinger, 2010).
264 Consequently, early workers hypothesized that the isotopic composition of ancient seawater
265 was best reflected by the lowest $\delta^{34}\text{S}$ value in an evaporite succession (Ault & Kulp, 1959;
266 Davies & Krouse, 1975; Thode & Monster, 1965). However, evaporite basins in marginal
267 marine environments are recharged not only by unadulterated seawater, but also by
268 groundwater and runoff with $\delta^{34}\text{S}$ compositions biased either higher or lower than seawater
269 from remobilized older evaporite deposits or weathered sedimentary pyrite and organic
270 sulfur (Nielsen & Rieke, 1964; Utrilla et al., 1992). Finally, high organic carbon
271 concentrations in many evaporite deposits can promote isotope fractionation by
272 thermochemical sulfate reduction during burial diagenesis (Vinogradov, 2007).

273 Some of the uncorrelated variance in evaporite isotope ratio data also results from poor
274 stratigraphic control (Supporting Information). Here we used updated stratigraphic
275 information to better constrain the age of evaporite data, but the record can further benefit
276 from higher-resolution sample collection with improved stratigraphic control during
277 intervals where $\delta^{34}\text{S}$ changes appear in other records. Modern stratigraphic models permit
278 correlation of evaporitic strata to better-constrained carbonate and clastic strata. Bernasconi
279 et al. (2017) recently produced a high-resolution evaporite record that resolved the major
280 early Triassic $\delta^{34}\text{S}$ excursions seen in earlier datasets; thus careful correlation and assignment
281 of geologic ages permits tracking changes in the Phanerozoic sulfur cycle with evaporites.
282 Indeed, the stratigraphic control for Mesozoic evaporites provided by Bernasconi et al.
283 (2017) likely drives the low standard deviation of uncorrelated Mesozoic evaporite data to
284 values (0.4‰) comparable to that of the marine barite record (0.3‰).

285 3.2.2 Barite

286 Barite precipitates from hydrothermal fluids, sediment pore fluids, and from particles within
287 the marine water column (Paytan et al., 1993, 2002). Barite is under-saturated in most of the
288 oceans (Chow & Goldberg, 1960; Church & Wolgemuth, 1972). However, barite has been
289 observed in sediment traps in the upper 200 m in the water column, especially in high-
290 productivity regions, and is associated with sulfate enrichment from decaying organic matter
291 (Bishop, 1988). While barite super-saturation is achieved predominately by the addition of
292 sulfate from oxidizing organic sulfur (Horner et al., 2017; Jacquet et al., 2007), marine barite
293 apparently precipitates with $\delta^{34}\text{S}$ values within 0.4‰ of modern seawater (Paytan et al., 1998,
294 2002). Barite is subsequently transported to sediments by fecal pellets and marine snow
295 (Bishop, 1988), and preserved in oxic marine sediments in high-productivity regions where
296 enough barite is delivered to saturate pore fluids (Church & Wolgemuth, 1972). Sulfate
297 reduction in anoxic sediments can cause dissolution of barite, which re-precipitates at the
298 base of the sulfate reduction zone with extremely high $\delta^{34}\text{S}$ compositions (M. E. Torres et
299 al., 1996).

300 Marine barite is considered an accurate proxy for ancient seawater $\delta^{34}\text{S}$ because it
301 precipitates in the open-ocean water column and is texturally distinguishable from diagenetic
302 barite that forms in anoxic sediments at redox fronts (Paytan et al., 1993). However, the
303 marine barite record is limited by the availability of open-marine sediments that deposited in
304 high-productivity regions where both authigenic enrichment of barite occurs and pore fluid
305 sulfate concentrations remain above zero (Paytan et al., 1993). Consequently, the barite $\delta^{34}\text{S}$
306 record is unlikely to be extended much further than the current dataset spanning the last 130
307 Myr. Bedded barite deposits are associated with economically-important disulfide mineral
308 deposits (C. A. Johnson et al., 2009), but contain large $\delta^{34}\text{S}$ variability (>10‰) and do not
309 resolve the ancient seawater record any better than other proxy materials. Additionally, with
310 few exceptions (e.g., Yao et al., 2018), the temporal resolution of the marine barite $\delta^{34}\text{S}$
311 record is unlikely to dramatically improve, especially during biogeochemical events
312 characterized by low marine productivity (such as the Cretaceous-Paleogene boundary) or
313 bottom-water anoxia (such as ocean anoxic events) that would have limited authigenic barite
314 enrichment or preservation.

315 3.2.3 Carbonate-associated sulfate

316 Limestones and dolomites deposited continuously throughout Phanerozoic time,
317 accumulating in marginal marine and open-ocean environments. A minor amount of sulfate
318 is incorporated into biogenic and abiogenic carbonate phases. Biogenic carbonates often
319 contain part-per-thousand sulfate by mass, while inorganic cements typically contain
320 hundreds of parts-per-million (Barkan et al., 2020; Busenberg & Plummer, 1985; Giri &
321 Swart, 2019; Paris, Fehrenbacher, et al., 2014; Staudt & Schoonen, 1995). Recent sediments
322 from various peritidal carbonate platform environments include CAS with an isotopic
323 composition similar to modern seawater (Lyons et al., 2004). CAS, therefore, complements

324 and exceeds the temporal resolution and completeness of the evaporite and barite records
325 (Strauss, 1997).

326 Diagenetic processes may exchange sulfate with the primary carbonate and alter its isotopic
327 composition (Fichtner et al., 2017; Murray et al., 2020; Present et al., 2015, 2019).
328 Kampschulte & Strauss (2004) suggested that the variability of multiple $\delta^{34}\text{S}$ analyses from
329 contemporaneous stratigraphic successions could be used to quantify the effect of diagenesis
330 on the CAS record. However, rapidly-changing and disparate CAS $\delta^{34}\text{S}$ compositions have
331 since been generated and interpreted—especially in Paleozoic studies—as intervals of
332 heterogeneous seawater sulfate $\delta^{34}\text{S}$ reflecting periods of low sulfate concentrations and low
333 marine sulfate residence times (e.g., Gill, Lyons, Young, et al., 2011; Kah et al., 2004).

334 Limestones and dolomites are comprised of mud or grains that precipitated both biologically
335 and abiotically from seawater, with cements binding them together. Each of these
336 components may recrystallize in pore fluids whose chemical composition reflects marine,
337 meteoric, and burial diagenetic processes. A combustion CAS analysis typically requires 10
338 g to 100 g of carbonate (Wotte et al., 2012), and this mass requirement dictates that samples
339 mix components that may have precipitated and/or recrystallized at different times. Further,
340 CAS analyses may be contaminated by sulfur from co-occurring phases, including sulfide
341 and disulfide minerals, sulfur-bearing organic material, and sulfate salts (Edwards et al.,
342 2019; Marengo, Corsetti, Hammond, et al., 2008; Present et al., 2015; Theiling & Coleman,
343 2015; Wotte et al., 2012). Recent application of plasma-source mass spectrometry for sulfur
344 isotope analysis has permitted $\delta^{34}\text{S}$ analyses on less than one-thousandth as much sulfate,
345 corresponding to 5 mg to 50 mg of carbonate (Paris, Adkins, et al., 2014; Paris et al., 2013;
346 Present et al., 2015, 2019; Rennie et al., 2018). Well-preserved biogenic grains,
347 recrystallized grains, matrix, and cements contain CAS with $\delta^{34}\text{S}$ compositions varying by
348 as much as 25‰, spanning most of range of CAS analyses from the entire Phanerozoic
349 (Present et al., 2015, 2019). Therefore, much of the variability of CAS $\delta^{34}\text{S}$ data may not
350 reflect the $\delta^{34}\text{S}$ composition of ancient seawater sulfate. Identifying components that retain
351 the $\delta^{34}\text{S}$ of sulfate incorporated from syndepositional seawater is critical to precisely and
352 accurately exploit the CAS $\delta^{34}\text{S}$ archive.

353 CAS can reflect the $\delta^{34}\text{S}$ of syndepositional seawater sulfate if the carbonate component did
354 not recrystallize after precipitation, if recrystallization and cementation occurred in contact
355 with a low-sulfate fluid, or if the $\delta^{34}\text{S}$ of pore fluid sulfate was not fractionated from seawater
356 (Gill et al., 2008; Lyons et al., 2004; Rennie & Turchyn, 2014). Alteration occurs if the
357 sediments recrystallize above the depth at which sulfate is completely consumed by MSR
358 but deep enough that some distillation of $\delta^{34}\text{S}$ within sediment pore fluid has occurred
359 (Edwards et al., 2019; Fike et al., 2015; Present et al., 2019; Rennie & Turchyn, 2014; Witts
360 et al., 2018). Additionally, some ancient carbonates contain CAS with anomalously low $\delta^{34}\text{S}$
361 interpreted to result from the incorporation of sulfate from sulfide that was reoxidized during
362 diagenesis or weathering (Baldermann et al., 2015; Edwards et al., 2019; Fichtner et al.,
363 2017; Fike et al., 2015; Marengo, Corsetti, Kaufman, et al., 2008; Present et al., 2015, 2019;
364 Rennie & Turchyn, 2014; Riccardi et al., 2006; Yan et al., 2013). Carbonates recrystallizing

365 during burial may also be prone to diagenetic modification of the $\delta^{34}\text{S}$ of CAS if the burial
366 fluids were sulfate rich (Fichtner et al., 2017, 2018; Present et al., 2015). The $\delta^{34}\text{S}$ in burial
367 fluids may be highly variable, and include sulfate from hydrocarbon or organic matter
368 degradation, dissolved evaporites, groundwater modified by MSR, or sulfate released by
369 dissolution of CAS (Dogramaci et al., 2001; Fichtner et al., 2018; Murray et al., 2020; Present
370 et al., 2019; Thode & Monster, 1965, 1970).

371 These diagenetic controls on the $\delta^{34}\text{S}$ of CAS decrease the precision and accuracy of the
372 proxy. This is quantified by its uncorrelated variance, which is much higher than that
373 observed in other seawater sulfate $\delta^{34}\text{S}$ proxies. Uncorrelated Paleozoic CAS data has a
374 standard deviation of 5.0‰, and that of Mesozoic CAS is 4.4‰, which is five to ten times
375 larger than that of Paleozoic and Mesozoic evaporites (1.0‰ and 0.4‰, respectively).
376 Further, diagenesis may have impacted accuracy by systematically biasing the $\delta^{34}\text{S}$ of CAS
377 with respect to the primary composition of seawater sulfate. For example, base level often
378 controls the stratigraphic arrangement of facies in carbonates successions, which can impart
379 biases as large as 10‰ on the $\delta^{34}\text{S}$ of CAS (Present et al., 2019; Richardson, Keating, et al.,
380 2019). Both the random and systematic variability is on the order of well-resolved rapid
381 changes of 3‰ to 6‰ in the $\delta^{34}\text{S}$ of marine barite and biogenic CAS.

382 3.2.4 Biogenic CAS

383 Biogenic CAS may offer a more robust $\delta^{34}\text{S}$ record than bulk CAS because biogenic
384 carbonate can often be readily separated from other limestone components, preservation
385 quality can be assessed, and vital effects appear to be small in most taxa (Kampschulte et al.,
386 2001; Paris, Fehrenbacher, et al., 2014; Present et al., 2015). In modern and cultured
387 biogenic carbonates, the incorporated sulfate has an isotopic composition within 2‰ of the
388 seawater from which it precipitated (Burdett et al., 1989; Kampschulte et al., 2001; Kaplan
389 et al., 1963; Mekhtiyeva, 1974; Paris et al., 2013; Paris, Fehrenbacher, et al., 2014; Present
390 et al., 2015). Recently, Rennie et al. (2018) produced a taxon-specific foraminiferal CAS
391 record with variance and secular trends comparable to the marine barite record.

392 Low-magnesium calcite, precipitated by many brachiopods, belemnites, and planktonic
393 foraminifera, is stable at Earth's surface and shallow burial conditions. The low-magnesium
394 calcite biogenic CAS $\delta^{34}\text{S}$ record has significantly improved the resolution of the
395 Phanerozoic $\delta^{34}\text{S}$ record during two key periods. First, during the Toarcian (Jurassic) Ocean
396 Anoxic Event, belemnite CAS displays a large (6‰) $\delta^{34}\text{S}$ excursion that is not well resolved
397 in the evaporite record (Gill, Lyons, & Jenkyns, 2011; Newton et al., 2011). Second, during
398 Carboniferous time, brachiopods record a prolonged recovery from a $\delta^{34}\text{S}$ maximum in
399 middle Devonian time (D. L. Johnson et al., 2020; Kampschulte et al., 2001; N. Wu et al.,
400 2014). However, aragonite and high-magnesium calcite, precipitated by many bivalves,
401 gastropods, corals, trilobites, echinoderms, bryozoans, and marine algae, dissolves and/or
402 recrystallizes much more readily than low-magnesium calcite (Brand & Veizer, 1980). Few
403 studies have investigated CAS $\delta^{34}\text{S}$ from formerly aragonitic fossils (Mekhtiyeva, 1974;
404 Present et al., 2015; Witts et al., 2018).

405 Unfortunately, well-preserved biogenic carbonate is rare in the rock record, especially during
406 intervals of climatic or biologic crisis (e.g., mass extinction events). Even apparently well-
407 preserved biogenic carbonate can still be susceptible to diagenetic alteration (Fichtner et al.,
408 2018; Witts et al., 2018). Like the marine barite record, a significant expansion of the
409 biogenic CAS $\delta^{34}\text{S}$ proxy record is limited by the availability of suitable sample material.

410 *3.3 Discrepant early Phanerozoic proxy records*

411 While all archives imperfectly estimate ancient seawater's composition, they provide
412 generally indistinguishable estimates considering the sources of uncertainty discussed
413 (Figure 2a). Paleozoic bulk rock CAS data, as a notable exception, commonly exhibit rapid
414 $\delta^{34}\text{S}$ variability (Figure 1b), but other archives with less uncorrelated variance are absent or
415 lack temporal resolution (Figure 1a). Throughout Phanerozoic strata, CAS data consistently
416 display more unresolved variance than other archives, yet they record the same long-term
417 trends (Figure 2), suggesting that some $\delta^{34}\text{S}$ excursions recorded by CAS may not represent
418 changes in the composition of the ocean. The high uncorrelated variance in all early
419 Paleozoic archives may mask $\delta^{34}\text{S}$ excursions on the order of those well-resolved in younger
420 strata by all archives. Spatial and temporal variability in early Paleozoic CAS data may
421 represent short residence times of sulfate in sulfidic oceans (e.g., Gill, Lyons, Young, et al.,
422 2011; Kah et al., 2016), local diagenetic effects on the $\delta^{34}\text{S}$ of carbonate rocks (Present et al.,
423 2015, 2019; Richardson, Keating, et al., 2019; Richardson, Newville, et al., 2019), or both
424 (Edwards et al., 2019; Rose et al., 2019).

425 CAS $\delta^{34}\text{S}$ excursions often correlate with global perturbations evidenced by carbon isotope
426 excursions and trace metal, pyrite sulfur isotope, and bioturbation intensity records (Canfield
427 & Farquhar, 2009; Fike et al., 2015; Gill et al., 2007; Jones & Fike, 2013; Kah et al., 2016;
428 Saltzman et al., 2015). Perhaps some CAS $\delta^{34}\text{S}$ excursions reflect widespread
429 biogeochemical changes at the interface between pore fluid sulfur cycling and carbonate
430 sediment diagenesis, including sulfate, dioxygen, and nutrient availability, organic
431 productivity, or metabolic or oceanographic changes in carbonate mineral saturation (Rennie
432 & Turchyn, 2014). Because part of the $\delta^{34}\text{S}$ variance in all archives derives from early
433 diagenetic processes—such as MSR, pyrite formation, and sulfide reoxidation—
434 consideration of these processes may reveal important temporal changes in carbon cycling
435 in marine pore fluids (Present et al., 2019; Richardson, Keating, et al., 2019; N. Wu et al.,
436 2010).

437 **4 Conclusions**

438 Phanerozoic $\delta^{34}\text{S}$ data were compiled from evaporites, barite, biogenic CAS, and bulk rock
439 CAS and updated to a consistent time scale. The subset of seawater sulfate's $\delta^{34}\text{S}$ history
440 possibly sampled by each proxy varied in space and time, and different suites of depositional
441 and post-depositional processes added variance to each archive. The variance in each record
442 increases with age, but the changing contribution of primary and secondary sources of
443 variability over Phanerozoic time remains unclear.

444 Bulk CAS contains a statistically significant different distribution of $\delta^{34}\text{S}$ compositions than
445 the biogenic CAS, evaporite, or barite records. Early diagenetic overprinting of CAS occurs
446 in depositional environments where carbonate recrystallization and cementation coincides
447 with sulfate-rich pore fluids with modified $\delta^{34}\text{S}$ values. Despite these complications, bulk
448 CAS can be widely applied in ancient sedimentary basins and is the only archive readily able
449 to resolve sulfur cycle changes during rapid biogeochemical events. Extending the breadth
450 and resolution of the $\delta^{34}\text{S}$ record requires developing mechanistic understanding of how
451 biogeochemical perturbations affect the marine diagenesis of carbonate rocks.

452 Acknowledgements

453 No new data were collected for this study. Datasets compiled for this research are tabulated
454 in the Supplemental Material and referenced below. We thank Caltech DocuServe for
455 obtaining many of the publications containing the compiled data, and John Grotzinger and
456 Joe Kirschvink for thoughtful advising and feedback. Constructive reviews by Akshay
457 Mehra and Julia Wilcots were greatly appreciated.

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