Pleistocene sediment offloading and the global sulfur cycle

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Abstract. Quaternary sea level fluctuations have greatly affected the sediment budgets of the continental shelves. Previous studies suggested that this caused a considerable increase in the net loss of shelf sediments. Since sediment accumulation and erosion are closely tied to the formation and re-oxidation of pyrite, we use a high-resolution record of sulfur isotope ratios ($^{34}$S/$^{32}$S) of marine sulfate to evaluate the implications of the so-called “shelf sediment offloading” on the global sulfur cycle. Modeling of our $\delta^{34}$S record suggests that erosion during sea level lowstands was only partly compensated by increased sedimentation during times of rising sea level and sea level highstands. Furthermore, our data suggests that shelf systems reached a new equilibrium state about 700 ka, which considerably slowed or terminated shelf sediment offloading.

1 Introduction

The Pliocene–early Pleistocene was characterized by relatively small (20–50 m) but frequent sea level changes in the precession and obliquity frequency bands (Miller et al., 2011). During the Middle Pleistocene, this pattern changed and large sea level fluctuations in the 100 ky frequency range gradually became dominant. At times, global sea level dropped as low as 130–150 m below present day sea level (Miller et al., 2011), exposing large areas of shelf to weathering and erosion. These sea level changes must have fundamentally altered the balance between sedimentation and erosion on continental shelves. Hay and Southam (1977) proposed that the repeated exposure and inundation of the continental shelves has led to a massive transfer of sediments from continental shelves to the deep ocean. They estimate that as much as $5 \times 10^{21}$ g of detrital sediment may have been removed by this so-called “sediment offloading” (Hay and Southam, 1977).

Although intuitively a convincing hypothesis, a quantitative analysis which includes the rates of sediment delivery to the deep ocean is missing. Hay and Southam (1977) hypothesized that the pattern of sea level falls controls the sediment delivery into the deep ocean. For example, during the first large sea level drop sediment transfer would be exceptionally large and the intensity of sediment erosion will decrease with consequent events, as the sediment reservoir available for erosion will become depleted (Hay and Southam, 1977; Hay, 1998; Hay et al., 2002).

Adding/removing sediments from the shelf is closely coupled to the burial/erosion of pyrite in those sediments. During interglacials, high sea levels result in expanded shelf areas. Coincidentally, the shelf areas are characterized by high pyrite burial rates (Jørgensen, 1982; Berner, 1982). During sea level lowstands, formerly inundated shelf areas are being replaced by low-lying coastal plains transected by rivers. This affects sedimentary sulfur cycling in two ways: (1) pyrite burial is reduced, and (2) fine-grained and unlithified sediments in the exposed shelf (de Haas et al., 2002) are eroded (Gibbs and Kump, 1994; Foster and Vance, 2006) and pyrite and organic sulfur (S) contained in the eroded sediments are oxidized (Turchyn and Schrag, 2004).

Pyrite formation is mediated by microbial sulfate reduction (MSR) and microbial sulfur disproportionation, which produce a large S-isotope ratio difference between pyrite and concomitant seawater sulfate (up to 70 ‰; Canfield and Thamdrup, 1994; Wortmann et al., 2001; Rudnicki et al., 2001; Böttcher et al., 2001, 2005; Brunner and Bernasconi, 2005; Sim et al., 2011). Accordingly, the burial of large amounts of pyrite will result in a more positive sulfur isotope value of seawater sulfate ($\delta^{34}$S), whereas the oxidation...
of large amounts of pyrite will cause a decrease of the seawater sulfate δ34S value. In the following, we take advantage of this relationship and use past changes of seawater sulfate δ34S value to track changes in pyrite burial/oxidation on continental shelves and their relation to changes in global sea level.

The δ34S value of seawater sulfate is uniform throughout the ocean, reflecting the long residence time of marine sulfate (∼10–20 Myr; Jørgensen and Kasten, 2006) compared to the ocean mixing time (∼1600 yr). The evolution of the δ34S value of sulfate thus serves as a proxy for past changes in the sulfur cycle (Paytan et al., 1998, 2004; Wortmann and Chernyavsky, 2007; Wortmann and Paytan, 2012).

Authigenic marine barite crystals continuously record past changes of seawater sulfate δ34S value (Paytan et al., 1998). Here we use a new high-resolution marine barite δ34S record of the last 3 Myr to delineate the onset and duration of these changes, which allows us to validate/test the shelf sediment offloading hypothesis.

2 Geological setting

We use sediment samples from eastern equatorial Pacific sites 849D (0°10.993′N, 110°31.167′W) and 851B (2°46.223′N, 110°34.308′W) obtained by advanced piston coring (APC) during Leg 138 of the Ocean Drilling Program (ODP). Site 849D is located below a highly productive equatorial divergence zone at a depth of 3839 m (Shipboard Scientific party, 1992a). Site 851B is located within the northern limit of western-flowing South Equatorial Current at the depth of 3760 m, within the equatorial high-productivity zone (Shipboard Scientific Party, 1992b).

Sediments at both locations consist of diatom nannofossil ooze (Shipboard Scientific Party, 1992a and b). Sedimentation rates were moderate since the late Pliocene, varying between 25–35 m Myr⁻¹ at Site 849D and 15–20 m Myr⁻¹ at Site 851B (Shipboard Scientific Party, 1992a and b).

Marine barite forms in the water column recording seawater S isotope ratios (Griffith and Paytan, 2012). After burial in the sediment, barite is stable during diagenesis except in environments with high rates of sulfate reduction where sulfate in pore waters is exhausted (e.g., Torres et al., 1996; Griffith and Paytan, 2012). In sulfate-reducing environments, barite is soluble releasing barium to solution. In this barium will diffuse and barite will reprecipitate forming diagenetic barite with typically anomalously high δ34S signatures (Paytan et al., 2002). Sites 849D an 851B are characterized by low organic matter (OM) concentrations and high sulfate concentrations in the interstitial waters (0.2 wt%, OM, 25–28 mM SO₄²⁻; Shipboard Scientific Party, 1992a, b). These conditions suggest that the barite samples in sediments at these sites are not affected by barite dissolution and/or reprecipitation and thus originate from sinking particles in the water column (e.g., marine barite).

Sample ages are estimated using high-resolution (10–50 kyr) orbitally tuned age models by Shackleton et al. (1995) which are based on magnetostratigraphy, biosтратigraphy, gamma ray attenuation porosity measurements (GRAPE) and δ18O records of benthic foraminifera.

3 Methods

Here, we used the sequential dissolution method of Paytan et al. (1996) to extract barite crystals from marine sediments. We have modified the original method to better address concerns about pyrite contamination (DeBond et al., 2012) and to improve the workflow. Unlike the original method organic matter is removed by heating the sample in the furnace at 700 °C instead of oxidizing it with hot bleach overnight. We also changed the order of the extraction steps so that iron and manganese oxyhydroxides are now dissolved with 0.2 N hydroxylamine hydrochloride in 25 % acetic acid at the end of the process. Between steps we centrifuged samples, decanted the supernatant and washed the residue three times with ultrapure deionized water.

In order to prevent oxidation of reduced sulfur during the carbonate leaching process, we added 50 mL of 5 % tin chloride (SnCl₂) solution to 1 L of HCl to maintain reducing conditions during the leaching step (instead of bubbling N₂ gas as in the original procedure). In addition, the HCl is flushed with Argon before the carbonate dissolution. This is the step in which we expected pyrite to be prone to oxidation if present in the sediments.

We examined the purity of the extracted barite with X-ray diffraction. Furthermore, we checked for presence of diagenetic barite using scanning electron microscopy and Energy dispersive X-ray spectroscopy (Paytan et al., 2002). If samples contained residual mineral phases like rutile, we dissolved the extracted barite with sodium carbonate and subsequently reprecipitated pure BaSO₄ (Breit et al., 1985).

3.1 Isotope analysis

Sulfur isotopes are analyzed with a continuous flow isotope ratio mass spectrometer system (CFIRMS) using a Eurovector Elemental Analyzer (EA) coupled via a Finnigan Conflo III open split interface to a Finnigan MAT 253 mass spectrometer. Solid barite samples (200 µg) are mixed in a tin cup with ∼ 600 µg of vanadium pentoxide (V₂O₅) powder and introduced into the EA, where the sulfate from barite (BaSO₄) is converted to sulfur dioxide gas (SO₂) by flash combustion at 1700 °C in an oxygen atmosphere. Measurements are calibrated using international sulfate standards NBS 127, IAEA SO5 and IAEA SO6 (relative to Vienna Canyon Diablo Troilite, hereafter VCDT, +21.1 ‰, +0.49 ‰, and −34.05 ‰, respectively; Copley et al., 2001) and an in-house synthetic BaSO₄ (Sigma-Aldrich) standard (8.6 ‰, VCDT). Repeated measurements of the in-house standard (typically
> 10 measurements per run) and international standards (3–4 measurements per standard per run) yield an average reproducibility of 0.15 ‰ (1 standard deviation – \( \sigma \)).

3.2 Statistical analysis

The isotope data includes errors in sample-assigned ages and uncertainties of how well a single measurement represents the seawater sulfate \( \delta^{34}\text{S} \) value. Note that the latter uncertainty not only includes analytical precision (which can be quantified) but also sample origin, sample handling and sample extraction. We therefore have to assume that each measurement carries an unknown error (or noise).

However, the \( \delta^{34}\text{S} \) value of seawater sulfate at any given time \( t \) depends to a certain degree on the \( \delta^{34}\text{S} \) of sulfate at a given time before \( (t - \Delta t) \). This allows us to apply a “local regression smoothing” technique (LOESS, Cleveland, 1979) to estimate the likely value for the \( \delta^{34}\text{S} \) of sulfate at any time of interest.

We used the default LOESS module provided by the statistical software package R (R Core Team, 2012). The 95 % confidence interval is calculated for each data point from the standard errors returned by the LOESS function.

3.3 Sulfur cycle model

We describe the sulfur cycle using the following mass conservation equation:

\[
\frac{d}{dt} M_{\text{SO}_4}(t) = F_{\text{wp}}(t) - F_{\text{bp}}(t) + F_{\text{we}}(t) - F_{\text{be}} + F_v, \tag{1}
\]

where \( M_{\text{SO}_4} \) denotes the mass of sulfate in the ocean, calculated from the sulfate concentration and the ocean volume; \( F_{\text{wp}} \) and \( F_{\text{we}} \) denote the pyrite and evaporite weathering input, respectively; \( F_v \) denotes the volcanic flux, and \( F_{\text{bp}} \) and \( F_{\text{be}} \) denote the pyrite and evaporite precipitation flux, respectively.

We can formulate a similar mass conservation equation for the respective isotopes of sulfur (\( ^{32}\text{S} \) and \( ^{34}\text{S} \)), as in Eq. (2):

\[
\frac{d}{dt} M_{\text{SO}_4}^{32}(t) = F_{\text{wp}}^{32}(t) - F_{\text{bp}}^{32}(t) + F_{\text{we}}^{32}(t) + F_v^{32}(t) - F_{\text{be}}^{32}, \tag{2}
\]

where \( M_{\text{SO}_4}^{32} \) denotes mass of \( ^{32}\text{S} \) in the ocean calculated from known mass of sulfate and its isotopic composition; \( F_{\text{wp}}^{32} \) and \( F_{\text{bp}}^{32} \) denote \( ^{32}\text{S} \) input from pyrite weathering and \( ^{32}\text{S} \) removal by pyrite burial, respectively; \( F_v^{32} \) denotes the \( ^{32}\text{S} \) input from volcanic flux; \( F_{\text{we}}^{32} \) and \( F_{\text{be}}^{32} \) denote the \( ^{32}\text{S} \) input from evaporite weathering and removal by evaporite precipitation, respectively.

In order to achieve an initial steady state we use modern values for the sulfur isotope composition and volume of the fluxes as boundary conditions (e.g., Berner, 1982; Kump, 1989; Hansen and Wallmann, 2003; Bottrell and Newton, 2006; see Table 1. for additional details). Note that the average isotopic composition of buried pyrite (\( \delta^{34}\text{S}_{\text{pyrite}} \)) is calculated so that other fluxes are in steady state.

From steady state condition,

\[
\frac{d}{dt} M_{\text{SO}_4}(t) = 0, \tag{3}
\]

we can calculate the average \( \delta^{34}\text{S}_{\text{pyrite}} \) value using Eq. (4) and Eq. (5):

\[
F_{\text{bp}}(t) = F_{\text{bp}}^{32}\text{S}(t) + F_{\text{bp}}^{34}\text{S}(t), \tag{4}
\]

\[
F_{\text{bp}}^{34}\text{S}(t) = F_{\text{we}}^{34}\text{S}(t) + F_{\text{wp}}^{34}\text{S}(t) + F_v^{34}\text{S} - F_{\text{be}}^{32}\text{S}. \tag{5}
\]

This yields \( \delta^{34}\text{S}_{\text{pyrite}} \) value of \(-18\) ‰, which is in agreement with previous estimates (Strauss, 1997; Seal, 2006; Leavitt et al., 2013). This implies the average offset (\( \Delta_{\text{pyr}} \)) between \( \delta^{34}\text{S}_{\text{seawater}} \) and \( \delta^{34}\text{S}_{\text{pyrite}} \) of \(-40\) ‰ (VCDT) which is similar to the Cenozoic average (Leavitt et al., 2013).

3.4 Model forcing

The objective of our model is to evaluate the effect of sea level changes on pyrite burial and weathering on the continental shelf and use these changes to track shelf sediment offloading. This requires that we consider two boxes for pyrite burial/erosion. The first box allows for pyrite burial and erosion in the shelf area as a function of the sea level, whereas the second box describes constant pyrite burial in the deep sea and pyrite erosion on continents. Our model follows the “rapid recycling” concept (Berner, 1987; see also Halevy et al., 2012) which separates inputs and outputs, to and from young and old sediment sub-reservoirs. This separation of young vs. old is geologically reasonable, because young sediments deposited in the near-shore environments are more likely to be weathered than older sediments (Berner, 1987). The novelty in our approach is that we scale fluxes into and from shelf sediments in proportion to the shelf area, as calculated from the high-resolution Quaternary sea level record (Miller et al., 2011).

We assume that up to 90 % of the total amount of pyrite is buried in the continental shelf (e.g., Berner, 1982; Canfield et al., 1992; Jørgensen, 1982). In deep water environments, the supply of OM is greatly reduced, and MSR and pyrite burial rates are orders of magnitudes smaller than in the shelf. In a first approximation, we can therefore treat pyrite burial in the deep-water box as constant.

There are, however, caveats to this assumption. Pyrite burial could increase if we (a) increase the delivery of reactive OM to the deep ocean by increasing export production, (b) introduce anoxic conditions, (c) move coastal upwelling zones offshore during glaciations, and (d) increase input of reactive iron to deep water environments through glacial erosion, iceberg sediment delivery, etc. Although some researchers argued for increased productivity (e.g., Murray et al., 1993; Filippelli et al., 2007) this is disputed by others.
Table 1. Model fluxes and sulfur isotope ratios in the steady state.

<table>
<thead>
<tr>
<th>Flux</th>
<th>Initial flux</th>
<th>Isotopic composition [%]</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Evaporite weathering</td>
<td>$2 \times 10^{12}$</td>
<td>21 Kump (1989), Garrels and Lerman (1981), see also Hansen and Wallmann (2003), for $\delta^{34}S_{\text{Evap}}$ see Claypool et al. (1980)</td>
<td></td>
</tr>
<tr>
<td>Volcanic flux</td>
<td>$0.34 \times 10^{12}$</td>
<td>3 Hansen and Wallmann (2003) and references therein</td>
<td></td>
</tr>
</tbody>
</table>

Note: The global sulfur fluxes are not well constrained. All fluxes used to achieve steady state are scaled within the range of previously published estimates (see reference list). The initial sulfate concentration is $27 \text{ mmol L}^{-1}$ which is in the range of estimates based on fluid inclusions in halite for late Miocene/Pliocene (Horita et al., 2002; Zimmermann, 2000; Brennan et al., 2013).

$^a$ Steady state value calculated as a function of other known fluxes (see text).

$^b$ This is used for model initialization. Later on the isotope value of the respective seawater sulfate.

(e.g., Nameroff et al., 2004; Francois et al., 1997; Dean et al., 1997). On the other hand, while redox proxies support decreased oxygen levels in some parts of the deep glacial ocean (François et al., 1997; Thomson et al., 1990; Mangini et al., 2001; Dean et al., 1997), other areas, specifically continental margins, show the opposite trend (i.e., higher oxygen levels, Ganeshram et al., 2002). Overall, Pleistocene trends of deep sea oxygenation are difficult to assess because they are dependent on several factors including circulation patterns, local productivity and temperature which show a high degree of temporal and spatial variability (e.g., Jaccard et al., 2010; Keeling et al., 2010). The offshore migration of upwelling zones during lowstands may increase burial of organic matter and pyrite in the slope and abyssal regions. However, the overall impact of this migration is likely small because upwelling areas have a small areal extent and thus a modest impact on the overall organic carbon burial (< 10% Berner, 1982; Hedges and Keil, 1995; Hu and Cai, 2011). Furthermore, pyrite burial is often limited in the upwelling zones by the lack of reactive iron (e.g., Morse and Emeis, 1990; Mossmann et al., 1991; Schenau et al., 2002; Brüchert et al., 2003; Suits and Arthur, 2000). The glacial inputs of iron by glacier meltwater and icebergs are likely significantly larger than at present, but most of it is in the form of crystalline iron oxides and silicates (e.g., Poulton and Raiswell, 2002; Raiswell et al., 2006) which react with sulfide on 100 kyr–1 Myr timescales (Canfield et al., 1992; Raiswell and Canfield, 1998). On the other hand, the majority of reactive Fe is trapped in fjords/estuaries (85–90%; Raiswell et al. 2006; Schroth et al., 2014) and thus unlikely to significantly contribute to pyrite burial in deep water environments. For the purpose of this model we therefore assume that pyrite burial in abyssal environments can be treated as constant.

Sediment offloading will also introduce pyrite and OM into the abyssal box. However, this redistributed pyrite cannot be counted twice, and thus will not alter the overall pyrite burial. The case for OM is however more complicated, as the additional OM will promote increased MSR. The extent of this OM support of MSR is however less clear as the remobilized OM is dominantly refractory in nature.

If sediments are quickly transferred from shelf to abyssal regions (e.g., gravity flows), OM and pyrite in redistributed sediments are highly vulnerable to oxidation due to greater oxygen penetration in deep water environments (Colley et al., 1984; Cowie et al., 1995; Thomson et al., 1998; Robinson, 2001). The process will continue until the diffusive supply of oxygen is interrupted either through deposition of pelagic sediment or another turbidite layer, which can last from a few thousand years to > 10 kyr (Cowie et al., 1995; Thomson et al., 1998; De Lange, 1998; Robinson, 2001). While we cannot quantify the amount of pyrite oxidized in this manner, we note that it is likely a significant portion of the overall pyrite weathering, particularly during glaciations when turbidite events are common (Weaver et al., 2000; Piper and Normark, 2009; Meiburg and Kneller, 2010).

We used the sea level estimates of Miller et al. (2011) to calculate the size of the global shelf area. Sea level variations (Fig. 1a) are often modified by local signals (e.g., gravity, mantle flow, isostatic rebound). However, the use of a global mean hypsometric curve (ETOPO-5, National Geophysical Data Center, 1988) is likely to average out local signals.

We calculate the shelf area ($A_s$; see Fig. 1b) as a function of sea level at any given point in time using a model cu-
The shelf area is used to force the fluxes affected by pyrite and evaporite weathering and pyrite burial in continental slope and pelagic environments. The second box represents pyrite and evaporite weathering and pyrite burial. First we divide these fluxes into two boxes. The first box corresponds to constant weathering on continents and constant pyrite burial in continental slope and pelagic environments. The second box represents pyrite and evaporite weathering and pyrite burial on the shelf and varies in proportion to calculated shelf area (Eqs. 7, 8). The pyrite weathering flux is calculated as follows:

\[ F_{wp} = F_{wp}^o \times \left[ 1 + \frac{A_{max} - A_s}{A_s} \right]. \]  

(7)

where \( A_{max} \) is the maximum extent of shelf area; \( F_{wp} \) is the calculated pyrite weathering flux corresponding to shelf change \( A_s^o - A_s^* \); \( F_{wp}^o \) is the minimum pyrite weathering flux corresponding to maximum shelf extent \( (A_{max}) \). We assume \( F_{wp}^o \) to be 90% of the steady state value calculated for the modern conditions. This assumption is based on the estimates of maximum shelf flooding area in the past 3 Myr. During times of maximum flooding the sea level may have been up to 10 m higher than the current sea level (Miller et al., 2011), corresponding to a 10% larger shelf area. At present some pyrite weathering takes place on this previously inundated shelf area. Therefore, we assume that during times of maximum extent of shelf inundation, pyrite weathering was lower and only 90% of that today, because pyrite rich shelf sediments were flooded.

Since shelf environments also contain large deposits of evaporites (e.g., Warren, 2006 and 2010), shelf area variations across glacial–interglacial cycles also affect evaporite weathering. Here we assume that evaporite and pyrite weathering covary and as a result of this the sulfur isotopic composition of sulfate input is steady at \( \sim 5.6 \% \) VCDT, which is in line with present day riverine sulfate \( \delta^{34}S \) values and Phanerozoic estimates (on average 3–8 ‰ VCDT; Canfield, 2013, and references therein). Evaporite weathering is thus calculated as follows:

\[ F_{we} = F_{we}^o \times \left[ 1 + \frac{A_{max} - A_s}{A_s} \right] \]  

(8)

where \( A_{max} \) is the maximum extent of shelf area; \( F_{we} \) is the calculated evaporite weathering flux corresponding to shelf change \( A_s^o - A_s^* \); \( F_{we}^o \) is the minimum evaporite weathering flux corresponding to maximum shelf extent \( (A_{max}) \). We assume that \( F_{we}^o \) is 90% of the steady state value.

The pyrite burial flux is calculated as follows Eq. (9):

\[ F_{bp} = F_{bp-abyssal} + F_{bp-shelf} \times \frac{A_s - A_{min}}{A_{max} - A_{min}}, \]  

(9)

where \( F_{bp-abyssal} \) corresponds to the minimum pyrite burial which takes place in slope and abyssal environments at minimum shelf extent, in this case \( 0.6 \times 10^{12} \text{molS yr}^{-1} \); \( F_{bp-shelf} \) is the portion of pyrite that is buried on the shelf at the maximum shelf extent \( (A_{max}) \) assumed to be \( 1.4 \times 10^{12} \text{molS yr}^{-1} \); \( A_{min} \) is the minimum shelf extent. These numbers are based on present-day estimates of sulfate reduction rates and pyrite burial in sediments at different water depth (Jørgensen, 1982; Jørgensen and Kasten, 2006; Thullner et al., 2009).

When considering pyrite burial on the shelf, we distinguish between old pyrite and pyrite which can be remobilized. The former represents the total shelf storage of
pyrite (∼10^{19} \text{ mol S}; \text{Charlson et al., 1992}), while the latter corresponds to the pyrite sub-reservoir in offloaded shelf sediments. Hay and Southam (1977) estimate that 5 × 10^{21} \text{ g} of shelf sediment was offloaded during the Pleistocene. If we take an average concentration of pyrite in pre-Holocene fine grained sediments (0.35 wt%; \text{Berner, 1982}), this corresponds to a pyrite reservoir of 6 × 10^{17} \text{ mol S}. However, this estimate carries a potentially large error. Although modern observations suggest that pyrite is quickly oxidized in reworked shelf sediments (e.g., \text{Amazon shelf; Aller et al., 1986}), it is unclear if all of this eroded pyrite was oxidized. Furthermore, the measured content of pyrite sulfur in shelf sediments is highly variable (0.1–1.5 wt%; \text{Berner, 1982}). Therefore, we suggest that the pyrite sub-reservoir in offloaded sediments could be 6 × 10^{17} ± 3 × 10^{17} \text{ mol S} and we use this range in model runs.

The average isotopic composition of pyrite across glacial–interglacial periods is not known. The sulfur isotope fractionation is negatively correlated with sedimentation rate (e.g., \text{Canfield, 2004; Kampschulte and Strauss, 2004; Bottrell and Newton, 2006}). Estimates for modern pyrite burial vary between 30 and 90% of the total sulfur burial with the remainder being buried as gypsum or anhydrite (e.g., \text{Canfield et al., 2004; Cogné and Humler, 2006; White et al., 2006}). However, recent evidence suggests that these changes were minor (e.g., \text{Foster and Vance, 2006}). Nonetheless, pyrite weathering is not restricted to the continental interiors, but happens each time marine sediments are exposed to erosion.

4 Results and Discussion

Our results show that between 3 and ∼1.5 Ma the seawater δ^{34}S_{SO_4} values fluctuate around ∼22‰ (VCDT) with a standard deviation (1σ) of 0.2‰. In the interval between 1.5 and 0.7 Ma we observe a steady decline from ∼22 to 20.7‰ (VCDT) (Fig. 2). This minimum is followed by an upwards trend from 20.7‰ (VCDT) at 0.7 Ma to 21.1‰ (VCDT) at 0.6 Ma. In the past 0.3 Ma there is a decline from 21.1 to ∼20.7‰ (VCDT) in the most recent sediments (Fig. 2; see Table A1 in Appendix for full list of samples).

4.1 Causes of seawater sulfate δ^{34}S change

Considering the long residence time of sulfate in the ocean (10^{7} \text{ yr}; \text{Jørgensen and Casten, 2006}), a −1‰ shift between 1.5 and 0.7 Ma, implies a massive change in the balance of the sulfur input/output fluxes. Possible explanations include (a) an order of magnitude increase of volcanic and hydrothermal S release, (b) a drastic increase in pyrite weathering, and (c) a massive decrease in pyrite burial. An order of magnitude increase of volcanic S input is incompatible with the geological record which shows no evidence for intensification of volcanic activity in the Pleistocene compared to the earlier periods of the Cenozoic (\text{Kaiho and Saito, 1994; Mason et al., 2004; Cogné and Humler, 2006; White et al., 2006}).

Pyrite weathering could have been affected by changes in continental erosion rates in the past 3 Myr (e.g., \text{Raymo et al., 1988}). However, recent evidence suggests that these changes were minor (e.g., \text{Foster and Vance, 2006}). Nonetheless, pyrite weathering is not restricted to the continental interiors, but happens each time marine sediments are exposed to erosion.

Glacially induced sea level drops will expose large swaths of previously ocean-covered shelf areas to subaerial weathering and erosion. Coincidentally, the shallow shelf is also the location of the highest pyrite burial rates (\text{Jørgensen, 1982}). First-order approximations show that shelf-area-
Increased pyrite weathering during lowstands depletes the shelf pyrite sub-reservoir, which we estimate to be $6 \times 10^{17} \pm 3 \times 10^{17}$ mol S (see Methods). We run a sensitivity analysis to determine how model output changes in this region of shelf pyrite sub-reservoir input parameter uncertainty.

Although the recent work by Leavitt et al. (2013) suggests that $\Delta_{\text{pyr}}$ increases with decreasing shelf area and vice versa on timescales of $10^7$–$10^9$ yr, the glacial–interglacial variations are not constrained. Due to long residence time of sulfate ($10^7$ yr) it is unclear if $\Delta_{\text{pyr}}$ changes on this timescale affect seawater sulfate $\delta^{34}$S. We explore the impact of $\Delta_{\text{pyr}}$ temporal variations by comparing model outputs produced assuming “fixed” $\Delta_{\text{pyr}}$ ($-40\%$) or variable $\Delta_{\text{pyr}}$, calculated in inverse proportion to the shelf area and within the range of Phanerozoic estimates in Leavitt et al. (2013) (see Methods for details and Fig. 4 for variable $\Delta_{\text{pyr}}$).

We start our model at 3 Ma (late Pliocene) and forward the resulting sulfur isotopic composition of seawater sulfate as a function of the sea level estimates published by Miller et al. (2011). Our model captures the shape and magnitude of the $\delta^{34}$S signal quite well (Fig. 5). Specifically, the decline of $\delta^{34}$S values in the past 1.5 Myr is well represented. This suggests that, during this time interval, larger sea level fluctuation of up to $-150$ m (Lisiecki and Raymo 2005; Clark et al., 2006; Miller et al., 2011) drastically increased the transfer of shelf sediments into the deep ocean.

During the interglacial periods, sea level rise creates large accommodation volumes, but Hay and Southam (1977) proposed that the creation of accommodation space outstripped sediment supply, resulting in a net loss of $5 \times 10^{21}$ g of shelf sediment during the Quaternary. This interpretation is supported by our $\delta^{34}$S data, which suggest that the balance between pyrite weathering and pyrite burial shifts in favor of pyrite weathering with increasing sea level variations during the Middle Pleistocene.

The cumulative pyrite weathering is, however, constrained by variations of seawater sulfate $\delta^{34}$S. Interestingly, the steady decline of the $\delta^{34}$S isotope values appears to slow down or to stop around $\sim 700$ ka (Fig. 2). If we accept the premise that the pyrite burial and weathering are linked to sedimentation and subaerial shelf erosion, the stabilization of seawater sulfate $\delta^{34}$S implies that sediment offloading has come to an end or, in other words, shelf sedimentation and erosion dynamics must have reached a new equilibrium adapted to the climate driven 100 kyr sea level cycles.

The model output is affected by the initial size of pyrite shelf sub-reservoir (Fig. 5). The mean estimate for pyrite sub-reservoir parameter ($6 \times 10^{17}$ mol S) produces model output within a 95% confidence interval of $\delta^{34}$S data. Assuming the low estimate of $3 \times 10^{17}$ mol S, the model output undershoots the data because the shelf pyrite sub-reservoir is quickly depleted (Fig. 5). On the other hand, if we use the higher shelf pyrite sub-reservoir estimate ($9 \times 10^{17}$ mol S),
the model output exceeds the measured seawater sulfate $\delta^{34}$S drop (Fig. 5).

The model outputs are almost identical whether we use “fixed” or variable $\Delta_{\text{pyr}}$, which suggests that our model is not sensitive to changes of $\Delta_{\text{pyr}}$ (see Fig. 5). This is corroborated by sensitivity test involving only variable $\Delta_{\text{pyr}}$, which produced constant output with a small positive shift

in the past 1 Myr (Fig. A1). However, we are not arguing that $\Delta_{\text{pyr}}$ was constant throughout the Quaternary. Environmental factors controlling $\Delta_{\text{pyr}}$ (sedimentation rate, areal extent of shelf environments; Leavitt et al., 2013) have changed during this period, which caused both increase and decrease of $\Delta_{\text{pyr}}$. For example, during interglacials $\Delta_{\text{pyr}}$ could have been $-35\%$, while during glacials it could have fallen to $-50\%$ (see Fig. 4). Since $\Delta_{\text{pyr}}$ fluctuated in both directions, the average for the whole Quaternary was similar to our “fixed” value of $-40\%$. While better temporal understanding of $\Delta_{\text{pyr}}$ variations during the Quaternary would certainly improve our interpretation, it is unlikely that $\Delta_{\text{pyr}}$ controlled seawater sulfate $\delta^{34}$S.

In our model, we consider pyrite burial in abyssal regions as constant across glacial–interglacial cycles. However, this may not be the case, as pyrite burial during glaciations can be affected by increased OM burial, emergence of bottom water anoxia, migration of upwelling zones or by higher iron input (see Sect. 3.4 for more details). Higher pyrite burial during glacial periods, for any of these reasons, would have to be compensated by increased pyrite weathering. While this cannot be completely excluded, we note that the change of the seawater sulfate $\delta^{34}$S value due to glacial–interglacial variations of pyrite burial alone is $\sim 30\%$ of the overall $\delta^{34}$S shift (Fig. A2 in Appendix). Since pyrite burial in deep ocean is small ($\sim 10\%$ of the overall pyrite burial; Berner, 1982; Canfield et al., 1992; Jørgensen, 1982) any change of this sink would likely have a minor impact on the seawater sul-
This standpoint is valid for long-term changes of seawater sulfur fluxes (e.g., Kurtz et al., 2003). While fate reservoirs changes slowly in response to the long-term processes, which underlines that large and sluggish seawater sulfate changes are consistent with a traditional view of the sulfur cycle (Fig. A3a, b; for forcing see A4). These damped model outputs show first-order agreement with half of deep ocean DIC storage. Since our \( \delta^{34}S \) record also suggests massive short-term imbalance between sulfur inputs and outputs, we argue that the initial fluxes at half the steady state value, although conceivable, are likely not representative of the Quaternary sulfur cycle.

The model runs assuming initial fluxes at twice the steady state values fall within the 95% confidence interval of \( \delta^{34}S \) data if we assume the shelf pyrite sub-reservoir of \( 9 \times 10^{17} \) mol S and undershoot the data with the shelf pyrite sub-reservoir of \( 3 \times 10^{17} \) mol S (see Fig. A3c and d). The fluxes calculated in these sensitivity runs are at the higher end of previously published values (Fig. A5; for comparison with similar fluxes see estimate of glacial pyrite weathering in Turchyn and Schrag, 2004). Therefore, we posit that they constrain the upper limit of the shelf pyrite sub-reservoir which is approximately \( 9 \times 10^{17} \) mol S. Together with our main model run (see Figs. 3 and 5) this suggests that glacial shelf sediment erosion resulted in oxidation between \( 6 \times 10^{17} \) and \( 9 \times 10^{17} \) mol of pyrite S during the Quaternary.

### 4.2 Pyrite oxidation and the carbon cycle

A massive increase of pyrite weathering implied by our results has important implications on the carbon cycle. Namely, pyrite weathering produces very strong sulfuric acid which dissolves carbonates (e.g., Spence and Telmer, 2005; Calmels et al., 2007). Since continental shelf sediments are rich in carbonates (de Haas et al., 2002) the production of sulfuric acid is likely balanced by carbonate dissolution, which delivers dissolved inorganic carbon (DIC) into the ocean-atmosphere system. Per each mole, of sulfate two moles of DIC are transferred to the ocean (Torres et al., 2014) (Reaction R1).

\[
2\text{CaCO}_3 + \text{H}_2\text{SO}_4 \rightarrow 2\text{Ca}^{2+} + 2\text{HCO}_3^- + \text{SO}_4^{2-} \quad \text{(R1)}
\]

Therefore, pyrite weathering effectively increases DIC storage in the ocean (Torres et al., 2014). If integrated over the entire period of the \( \delta^{34}S \) value shift, pyrite oxidation results in a net transfer of \( 14,000–21,000 \) PgC or about a third to a half of deep ocean DIC storage.

### 5 Conclusions

This study shows that the intensification of Quaternary glaciations in the past 1.5 Myr and concomitant periodic changes in shelf area likely affected the balance of weathering fluxes of sulfate/sulfide and the burial of pyrite. We propose that during glacial periods, pyrite weathering drastically increases as a result of subaerial shelf erosion and that the increased erosion rates are not fully compensated by increased pyrite burial during sea level highstands.

The declining seawater \( \delta^{34}S \) values support the idea that the transition to the climate driven 100 kyr sea level variations resulted in a net reduction of shelf sediment vol-

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**Figure 5.** Model output – seawater sulfate \( \delta^{34}S \) value. Brown solid line – model output with constant offset of \( -40 \% \) between \( \delta^{34}S_{\text{seawater}} \) and \( \delta^{34}S_{\text{pyrite}} \) values. Grey solid line – model output with variable offset between \( \delta^{34}S_{\text{seawater}} \) and \( \delta^{34}S_{\text{pyrite}} \) values (see Fig. 3). Red dashed line – model output with constant \( \Delta_{\text{pyr}} \) offset and \( 50 \% \) larger shelf pyrite sub-reservoir. Green dashed line – model output with constant \( \Delta_{\text{pyr}} \) offset and \( 50 \% \) smaller shelf pyrite sub-reservoir. The shaded area is the 95% confidence interval of a LOESS approximation of the "true" \( \delta^{34}S_{\text{SO}_4^2} \) value.
Our data show that the steady decline in the seawater $\delta^{34}\text{S}$ values stops around 700 ka. We consider it likely that this stabilization indicates the termination of the massive net “sediment offloading” (Hay and Southam, 1977) and heralds a new equilibrium between shelf erosion during sea level lowstands and sediment resupply during sea level highstands.

The resuspension of previously deposited sediments oxidized large amounts of pyrite back to sulfate. Our model results suggest that this would have increased the marine sulfate concentration by $\sim 1–3$ mM in good agreement with sulfate concentration estimates based on fluid inclusions (Brennan et al., 2013) and estimates of the ocean alkalinity based on boron isotopes (Hoenisch et al., 2009). Furthermore, it is likely that shelf sediment offloading may have impacted additional elements that are predominantly buried in the shelf sediments such as phosphorus and carbon (e.g., Berner, 1982; Wollast, 1991; Ruttenberg, 2003) with possible implications to their biogeochemical cycles as well as ocean productivity.
Appendix A: Sample list, model results and forcing of sensitivity runs

Table A1. Sample list with sulfur isotope results.

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Figure A1. Model output using variable $\Delta_{\text{pyr}}$. Black solid line – model output produced assuming constant steady state fluxes and variable $\Delta_{\text{pyr}}$. Green dashed line – model output produced assuming constant fluxes at 200% steady state values and variable $\Delta_{\text{pyr}}$. Brown dashed line – model output produced assuming constant fluxes at 50% steady state values and variable $\Delta_{\text{pyr}}$. The shaded area represents the 95% confidence interval of a LOESS approximation of the “true” $\delta^{34}$S$_{\text{SO}_4}$ value. Note that the size of pyrite reservoir in these model experiments remains the same because input and output fluxes are kept constant.

Figure A2. Model output with variable pyrite burial and fixed weathering fluxes and $\Delta_{\text{pyr}}$. Black solid line – model output produced assuming initial fluxes at steady state values, variable pyrite burial and fixed weathering fluxes and $\Delta_{\text{pyr}}$. Green dashed line – model output produced assuming initial fluxes at 200% steady state values, variable pyrite burial and fixed weathering fluxes and $\Delta_{\text{pyr}}$. Brown dashed line – model output produced assuming initial fluxes at 50% steady state values, variable pyrite burial, fixed weathering fluxes and $\Delta_{\text{pyr}}$. The shaded area represents the 95% confidence interval of a LOESS approximation of the “true” $\delta^{34}$S$_{\text{SO}_4}$ composition.
Figure A3. Sensitivity model tests. (a) Model output using fixed $\Delta_{\text{pyr}}$ and assuming initial fluxes at 50% steady state values. (b) Model output using variable $\Delta_{\text{pyr}}$ and assuming initial fluxes at 50% steady state values. (c) Model output using fixed $\Delta_{\text{pyr}}$ and assuming initial fluxes at 200% steady state values. (d) Model output using variable $\Delta_{\text{pyr}}$ and assuming initial fluxes at 200% steady state values. Note: black solid line – model outputs produced assuming the low estimate for shelf pyrite sub-reservoir. Brown solid line – model outputs produced assuming the high estimate for shelf pyrite sub-reservoir. The shaded area represents the 95% confidence interval of a LOESS approximation of the “true” $\delta^{34}\text{S}_{\text{SO}_4}$ value (for forcing see Figs. A4–A5).
Figure A4. Model forcing with initial fluxes at 50% steady state values. (a) Pyrite weathering ($F_{wp}$). (b) Evaporite weathering ($F_{we}$). (c) Pyrite burial ($F_{bp}$). (d) Pyrite sub-reservoir on the shelf. Note: the pyrite burial flux is not affected by the size of initial pyrite shelf sub-reservoir.

Figure A5. Model forcing with initial fluxes at 200% steady state values. (a) Pyrite weathering ($F_{wp}$). (b) Evaporite weathering ($F_{we}$). (c) Pyrite burial ($F_{bp}$). (d) Pyrite sub-reservoir on the shelf. Note: the pyrite burial flux is not affected by the size of pyrite shelf sub-reservoir.
Author contributions. S. Markovic produced the isotopic data and developed the modeling methodology; all authors assisted in drafting of the manuscript.

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Edited by: T. Treude

References


