Mass-independent isotopic signatures of volcanic sulfate from three supereruption ash deposits in Lake Tecopa, California

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A B S T R A C T

Hundreds to thousands of megatons of sulfur dioxide released by supereruptions can change chemical and physical properties of the atmosphere and thus induce climate perturbations. We present oxygen and sulfur isotope analyses of sulfate in 48 volcanic ash samples, and 26 sediment samples from dry lake beds in the Tecopa basin, California, USA. These ash layers represent three supereruptions, including the 0.64 Ma lava Creek Tuff, 2.04 Ma Huckleberry Ridge Tuff and 0.76 Ma Bishop Tuff. Mass-independent oxygen signatures (Δ17O up to 2.26‰) that are present in these ash units, and not in associated sediments, indicate oxidation of volcanic SO2 by mass-independent ozone and its products. In this study, we consider the formation, deposition, preservation and dilution of mass-independent volcanic sulfate (MIVS). Using the isotopic compositions of the sulfates, we construct a mixing model that demonstrates that the main source of sulfate in Lake Tecopa is mass-dependent sediment-derived sulfate (MDSDS, >77%). However, ash beds still preserve up to 23% of MIVS that initially had undiluted Δ17O value around 8‰, and Δ34S as low as −0.35‰, and Δ33S up to 1.08‰. Therefore, despite potential dilution by MDSDS, the MIVS signatures can be preserved in the geologic record for few million years, if deposited as gypsum in arid environments, alkaline or saline lake. The oxygen and sulfur mass-independent signatures of the volcanic sulfates indicate that photolysis and oxidation of volcanic SO2 has been achieved in the upper atmosphere. Since only supervolcanic eruptions were shown to generate massive amount of mass-independent sulfate, it requires that up to 20–60% of the global ozone layer is consumed as a result of supervolcanic SO2 released. This may occur as a result of a strong physical and chemical degradation of the tropopause; we speculate that the distinction between the high-troposphere and the low-stratosphere, at least locally, could be erased by supereruptions, and recorded by MIVS.

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1. Introduction

Volcanic eruptions of large magnitudes (called supereruptions) release many megatons of volcanic ash and SO2 gas, which upon turning into H2SO4 aerosols, reflect solar radiation and cause Earth’s surface cooling and stratosphere heating (e.g. Robock, 2000 and references therein). Slow oxidation of SO2 in the stratosphere leads to a global climate cool that lasts years to decades (e.g. Blong, 1984; Zielinski et al., 1996; Robock and Oppenheimer, 2003). Both oxygen and sulfur isotopic compositions of volcanic sulfates provide insight into the pathways of SO2 oxidation and residence time in the atmosphere.

Volcanic plumes from large eruptions can reach high altitudes (20–40 km). At such altitude, most oxygen-bearing molecular species (oxyanions) in the Earth’s atmosphere exhibit mass-independent isotope behavior (e.g. Thiemens, 2006 and references therein), with ozone Δ17O values of up +10 to +80‰ (+32‰ on average, e.g. Kranckowsky et al., 2000). The Δ17O-excess1 measured in volcanic sulfate from large eruptions (Bao et al., 2003; Bindeman et al., 2007) could ultimately be inherited from the ozone.

The volcanic sulfate has been isotopically studied after leaching of volcanic ash samples (e.g. Bao et al., 2003; Bindeman et al., 2007), and by extracting volcanic ash from ice cores from Antarctica (e.g. Savarino et al., 2003a,b; Baroni et al., 2007). The ice core sulfates show mass-independent behavior not only of oxygen, but also sulfur isotopes. The sulfur mass-independence require photodissociation of SO2 or SO3 by <310 nm short wave UV radiation (Farquhar et al., 2000; Farquhar et al., 2001) in the upper stratosphere, where the O3 shield is not thick enough to prevent it (e.g. Farquhar and Wing, 2003). Bindeman et al. (2007) showed that sulfates from distal ash deposits of the largest caldera-forming eruptions (>600 km²) for the past 2 Ma possess positive and high Δ17O of 0.7 to 3.5‰, while sulfate from smaller eruptions (<10 km³) show Δ17O < 0.3‰. Bao et al. (2003) measured

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1 The Δ17O-excess, or deviation from the terrestrial mass fractionation line (Δ17O=−0.52×δ18O) is expressed as Δ17O=δ17O−0.52×δ18O; see Miller (2002) for different formulations.
Δ^{17}O < 0.2‰ in sulfate leached from products of modern small eruptions, but reported the record high Δ^{17}O value of +5.84‰ in sulfate from 28 Ma ash layers in western Nebraska from an unidentified, sulfur-rich and large-volume eruption. The high Δ^{17}O in sulfates from large eruptions have been interpreted as reflecting the volcanic SO₂ oxidation by ozone-derived radicals in the troposphere (e.g. "dry fog", Bao et al., 2003) or stratosphere (e.g. Bindeman et al., 2007). Savarino et al. (2003a) additionally reported Δ^{17}O = 4.7‰ in stratospherically-processed sulfate collected in ash deposits from the medium-scale 1991 Mt Pinatubo eruption in the Antarctica ice.

While previous studies created an important framework, they provided insufficient details on the formation, deposition and preservation of mass-independent volcanic sulfates (MIVS). In this paper, we present a detailed study of sulfate extracted from volcanic ash and enclosing sediments in the presently dry Lake Tecopa, near Death Valley in California, USA. This sedimentary basin deposit contains ash layers from the three largest eruptions that occurred in the western USA during the past 2 Ma: two eruptions from Yellowstone: the 0.64 Ma, 1000 km² Lava Creek Tuff (LCT) and the 2.04 Ma, 2500 km², Huckleberry Ridge Tuff (HRT), and one eruption from the Long Valley caldera: the 0.76 Ma, 750 km² Bishop tuff (BT). We performed analyses of δ^{18}O, Δ^{17}O, and δ^{34}S in extracted sulfates and measurement of ash, sulfate and CaCO₃ concentrations. We additionally employ and reinterpret Δ^{33}S and Δ^{36}S data from Bindeman et al. (2007) as being mass-independent. Based on petrographic observations and measured isotopic values, we estimate the proportion of sedimentary and volcanic sulfate sources in Lake Tecopa and discuss the preservation of oxygen mass-independent volcanic sulfates (MIVS) in sedimentary records of intermountain basins. By linear algebra and mass-balance, we estimate the primary volcanic sulfate Δ^{17}O, Δ^{33}S and Δ^{36}S values, and speculate on the quantity of atmospheric ozone consumed in the process of volcanic sulfate formation during these three supereruptions.

2. Geological setting and sampling

The presently dry Lake Tecopa is located in the major intermontane basin to the southeast of Death Valley (Fig. 1). From the Pliocene to middle Pleistocene, Lake Tecopa oscillated between playa to shallow-lake conditions and even during its deepest stages (5–10 m) it remained both saline and alkaline (e.g. Morrison, 1999 and references therein). In this lake, the sedimentary deposits are 100 meters thick and contain at least 15 ash layers (Sheppard and Gude, 1968; Strakey and Blackmon, 1979; Hillhouse, 1987) that were deposited by airfall and redeposited into the lake by fluvial processes. Among these volcanic deposits three are meters-thick and are continuously present across the lake: the 0.64 Ma LCT, 0.76 Ma BT, and 2.04 Ma HRT (Figs. 1 and 2). The LCT ash bed is usually 2–2.5 m thick, reaching up to 3–4 m close to the town of Shoshone; (Fig. 2), the BT ash layer is commonly 1–1.5 m thick, and the affected by K-feldspar diagenesis HRT ash bed is the thinnest (0.5–0.7 m; Fig. 2 and 3). We estimate that these ash layers represent 15–20 vol.% of the Lake Tecopa sedimentary volume studied here (Fig. 3).

We collected 48 samples from LCT, BT and HRT ash beds and 26 samples from enclosing sediments along 18 different profiles. In each sampling location we collected ash every 10–30 cm from the bottom to the top of each layer, and extending into the sediments from 0.5–2 m below and above each ash beds (Table 1 and Fig. 3). The majority of samples were collected in the northern part of the Tecopa basin, where the ash layers are best exposed and not affected by diagenetic processes (e.g. Sheppard and Gude, 1968), as verified by fresh ash shards that are optically isotropic. A few samples were collected in the zeolite and K-feldspar diagenic facies for comparison (Fig. 2).

3. Methods

Sulfate was leached from 40–80 g of samples with HCl-acidified solution (pH = 1–2) and precipitated as barite (BaSO₄) by adding barium chloride (BaCl₂) to the solution while evaporating at T < 80 °C. Barite was purified from potential nitrate contamination by repeated dissolution–reprecipitation with chelating DTPA agent, because adsorbed nitrate contamination was suggested to influence the barite Δ^{17}O values (e.g. Bao, 2006). In this study, like an earlier study by Bindeman et al. (2007), nitrate contamination was found to be minor.

Fig. 1. Map of North America showing the distribution of air-fall deposit from supervolcanic eruptions of Yellowstone: Huckleberry Ridge Tuff (2.04 Ma) and Lava Creek Tuff (0.64 Ma) and Long Valley: Bishop Tuff (0.76 Ma). The location of the dry lake Tecopa is indicated by a white labeled circle.
(<2–3 wt.%) and does not affect $\Delta^{17}$O significantly. Indeed, measured $\Delta^{17}$O values from 16 untreated and treated barite plot less than 0.1‰ from the 1:1 line (Supplementary material).

Oxygen isotopes from 4–7 mg of extracted barite powder were analyzed at the Stable Isotope Laboratory at the University of Oregon by laser fluorination, using methods similar to Bao and Thiemens...
(2000). Extracted O₂ gas was then frozen in a molecular sieve (zeolite 13 Å) and run in a dual inlet mode in a Finnigan MAT253 mass spectrometer. Some samples were frozen in both a single and double molecular sieve (zeolite 13 Å) trap system to check for potential NF⁺ contamination that can affect mass 33 (e.g. Rumble et al., 1997; Pack et al., 2007; Supplementary material). We found such contamination to be negligible.

Sample duplicates, and replicates of barite international standard NBS127 (n = 7) and an in-house standard (IB04-13; n = 12) give a reproducibility (in 2σ) of δ¹⁸O= ±0.6‰, δ¹⁷O= ±0.3‰ and Δ¹⁷O= ±0.05‰. The 35–45% O₂ yield during barite laser fluorination leads to low-δ¹⁷O values, therefore based on the certified δ¹⁷O value of NBS127 (δ¹⁷O = 8.6‰), we applied a correction factor of +8.01‰ for all analyzed samples. Notice that this correction, which is a mass-independent parameter, does not affect Δ¹⁷O values. Day-to-day variations were also monitored by the GMG garnet standard (δ¹⁷O = 5.75‰). Repeated leaching and analysis from the same samples yielded external errors (in 2σ) of the whole method better than δ¹⁸O ± 1.2‰ and Δ¹⁷O ± 0.1‰ (Supplementary material).

Sulfur isotopes were analyzed via an elemental analyzer in the stable isotope laboratory at the University of Maryland. Multiple analyses of NBS127 give δ³⁴S = −8.37‰ ± 0.05 (in 2σ) and based on the certified value (211‰) a correction factor of +29.47‰ was applied for all samples.

In addition to microscopic examination, we investigated sulfate size, position, and morphology at the Smithsonian Institution of Washington DC. SEM imaging and X-ray mapping of one ash (LCT8) and one sediment (2BT+1) sample used the FEI Nova NanoSEM 600 variable pressure ultra high resolution SEM and one sediment (2BT+1) sample used the ThermoNoran Rapid microdiffactrometer.

4. Results

4.1. Petrography of sediment and ash layers

Lake Tecopa sediments mainly consist of lacustrine mudstone including clay, silt and sand (Sheppard and Gude, 1968; Hillhouse, 1987; Fig. 3). Some claystones are interbedded whereas sandstones appear as lenses in the mudstone. X-ray diffraction analyses performed on sediment (sample 2BT+1) led to the identification of calcite, clay minerals (montmorillonite and sepiolite), quartz, albite and micas. Authigenic zeolite and K-feldspar are also present in sediments as well as in ash located close to the center of the Tecopa basin, but absent in the northern and shallowest part of the basin (Fig. 2).

A macroscopic and microscopic petrographic investigation of each ash layer showed that the lower contact of the volcanic deposit is sharp, created by a massive and rapid air fall into the Tecopa basin. By contrast, the upper contact is gradual, showing progressive mixing of ash with the overlying sediments (Fig. 3). Locally, the upper part of the ash layers is seen as being fluvially reworked, and in other places, the volcanic deposit is cemented by carbonates that result in hard beds. The proportion of birefringent lithics across the optically isotropic ash bed was estimated with a petrographic microscope for most samples (Fig. 3), and changes gradually from the bottom to the top of each ash layer. We identified some of these lithics, via X-ray diffraction analyses, as halite and montmorillonite (sample LCT8).

The sediment below the ash layers is always ash-poor (<5%). The bottom of each ash layer consists of almost pure glass (>95% of 100–200 µm glass shards) and a very high ash proportion (>90%) remains across most of the ash layer (Fig. 3). At the top, where there is some cross-bedding and evidence of fluvial reworking of the volcanic deposit, the proportion of ash decreases progressively from 70 to 30%.

The overlying sediment is heavily contaminated with ash (10–30%). In some places, the overlying sediment also contains ash-rich lenses (>50%; Fig. 3).

4.2. Sulfate appearance and concentrations

SEM imaging and X-ray mapping (100 µm × 100 µm) of one ash (LCT8) and one sediment (2BT+1) samples succeeded in identifying gypsum (or anhydrite) particles between 1 and 10 µm, but generally 3–5 µm in size (Fig. 4). Likewise, Bindeman et al. (2007) reported CaSO₄ particles in their samples. High magnification scanning of individual gypsum particles in ash samples, demonstrate that the sulfates occur as reprecipitated clumps (Fig. 4).

In 85% of the samples, sulfate concentration was high and variable: ash layers contained 304 ± 151 (2σ) ppm SO₄ and sediments contained 653 ± 482 (2σ) ppm SO₄ (Table 1 and Fig. 3). It is noteworthy that in a single ash layer, the sulfate content varies irregularly, both vertically and laterally. For instance in the LCT profile 1 (Fig. 2; Table 1), SO₄ concentration ranges between 30 ppm and 365 ppm, and only in two samples (1LCT2 and 1LCT5), sulfate concentration was too low for extraction (<7 ppm; Table 1). Similarly, when comparing LCT profile 1 and profile 2 (Fig. 2 and Table 1), which are a few hundred meters apart, 75% of the samples from profile 1 have significant amount (~7 ppm) of sulfate while this is the case for only 10–15% in the profile 2. Overall, in only 15% of the samples, the sulfate (SO₄) concentration was too low for extraction. However, since the neighboring samples have plentiful sulfate it appears that highly variable sulfate concentration is due to very recent secondary leaching and loss. As an example, we did not find any sulfate in samples from 1 m thick ash layer from the Toba supereruption (74 ka) in Malaysia. It is likely that the wet climate in this region had naturally leached sulfate from ash. Likewise, recent erosion and leaching probably affected the Lake Tecopa sulfate distribution. However, secondary leaching is a mass-dependent process, unable to change the sulfate mass-independent Δ¹⁷O signature. Moreover, it has been demonstrated that leaching and sulfate loss does not significantly change the sulfate δ¹⁸O and δ³⁴S (e.g. Van Stempvoort and Krouse, 1994; Khademi et al., 1997).

4.3. Δ¹⁷O, δ¹⁸O and δ³⁴S values of sulfate

Overall, the sulfates extracted from the ash layers exhibit large ranges in oxygen and sulfur isotopic compositions: the δ¹⁸O ranges from 0.0 to 12.1‰ while the δ³⁴S ranges from 6.1 to 19.8‰. The sediment-extracted sulfates show a similarly large diversity: δ¹⁷O ranges from 2.8 to 10.0‰ (Fig. 3). The highest Δ¹⁷O values, up to 2.26‰ (Fig. 3 and 5), were measured in sulfates from ash whereas sulfate from the sediments has usually Δ¹⁷O <0.5‰. The lower contact between ash layers and underlaying sediment is characterized by a sharp increase of the Δ¹⁷O values, in agreement with the ash proportion (Fig. 3). The maximum Δ¹⁷O were usually measured in the lower part of ash layers, where ash concentration exceeds 95% of the sample (Fig. 3). The maximum Δ¹⁷O value gradually decreases to the top of ash beds and is roughly correlated with the proportion of ash in samples (Fig. 3). Additionally, high Δ¹⁷O values, up to 1.78‰, were measured in fluvially-redeposited ash-rich sediment lenses as observed on top of the LCT ash layers (Fig. 3). The sediment background Δ¹⁷O values are in the 0.15 to 0.5‰ range, and remain approximately constant for 2 Ma in the studied sections. Progressively older and less thick ash layers exhibit a decrease in the maximum measured Δ¹⁷O values: Δ¹⁷O = 2.26‰ in the 0.64 Ma LCT, Δ¹⁷O = 1.25‰ in the 0.76 Ma BT, and Δ¹⁷O = 0.76‰ in the 2.04 Ma HRT (Fig. 3).

It appears in these studied profiles that the main process that affects sulfate isotopic values, and Δ¹⁷O in particular, is sediment-ash sulfate mixing. The pattern of the inferred mutual cross-contamination is not simple since sediments contain on average twice the amount of...
sulfate as ash. The fresh ash samples generally have sulfate concentration lower than 400 ppm and high $\Delta^{17}O$ (>0.5‰), while sediments are usually sulfate-richer (>400 ppm) and have lower $\Delta^{17}O$ values (<0.5‰; Fig. 6). This pattern alone suggests volcanic origin of high $\Delta^{17}O$ in ash. Sulfate from ash, when mixed with sediments yields lower $\Delta^{17}O$ (<0.6‰) than in fresh ash, while the sediments contaminated by ash have higher $\Delta^{17}O$ than the sediment background (0.5‰). While $\Delta^{17}O$ in ash-uncontaminated sediments is systematically lower than 0.5‰, the $\Delta^{17}O$-excesses are still significantly higher than 0.0 ± 0.1‰ as would be expected for sediments derived from mass-dependent sources. There is a possibility that mass-independent signatures are coming from long-term atmospheric aerosol deposition (e.g. Bao et al., 2001), however the time-averaged amount of such precipitation in the Tecopa basin would not generate $\Delta^{17}O$ higher than 0.2‰ (see below).

Other stable isotope parameters and concentrations exhibit variable levels of correlation with lithology: the $\delta^{34}S$ decreases in ash layers (Fig. 3), and anticorrelates with $\Delta^{17}O$ (Fig. 5); less clear correlation is observed with sulfate concentrations and $\delta^{13}C$ (Fig. 3). Carbonate concentration is lower in ash beds than in sediment (Fig. 3), and this is most likely due to the fact that we sampled the freshest ash.

Finally, no systematic correlation between $\Delta^{17}O$ and the degree of diagenesis has been observed (Table 1 and Fig. 2): a negative correlation for LCT samples and slightly positive for BT samples. Diagenesis is a mass-dependent process incapable of changing the $\Delta^{17}O$ values of sulfates.

5. Discussion

The following discussion addresses 1) the sources of sulfate present in the studied sedimentary sections, 2) the mechanisms of incorporation of volcanic sulfate into ash beds, 3) the preservation of oxygen mass-independent volcanic sulfate in a sedimentary basin under arid environment conditions, 4) the sulfur mass-independent signature and 5) atmospheric perturbations due to the release of large amounts of volcanic SO$_2$, as revealed by its $\Delta^{17}O$ signature.

5.1. Sulfate sources

5.1.1. Mass-independent sulfate

In the arid environment of Lake Tecopa, the main source of initial $^{17}O$-excess in sediment is from atmospheric sulfate aerosols. Bao and Reheis (2003) measured maximum $\Delta^{17}O$ values of 0.85–0.44‰ in sulfate collected in dust traps. Long-term accumulation of sulfate is exemplified by desert varnishes, which have $\Delta^{17}O$ values of up to 0.6–0.8‰ (Bao et al., 2001). Sulfate in desert varnishes allow assessment of atmospheric accumulation rates and $\Delta^{17}O$ values during a few hundred thousand years. Using the maximum accumulation rate of 5 µm/1000 years for a desert varnish (Dorn, 1998) that contains 0.6 wt.% of mass-independent SO$_4$ (Bao et al., 2001), the deposition rate of mass-independent atmospheric sulfates is 30 µm/Ma. By contrast, in Lake Tecopa, playa or alluvial sediments had an average accumulation rate of ~40 m/Ma (Morrison, 1999), or ~24 mm/yr of the equivalent sulfate (considering ~600 ppm in average in sediment). If we assume that atmospheric sulfate precipitation is 100 times greater than that recorded in varnishes, then the atmospheric mass-independent sulfate represents 12.5% (75 ppm) of the sediment sulfate. Therefore, considering the maximum $\Delta^{17}O$ value of 1.2‰ (most likely ~0.5‰ on average) for atmospheric aerosols at Death Valley (Bao et al., 2001), the resulting $\Delta^{17}O$ of sediment sulfate is <0.2‰. Thus, the atmospheric mass-independent signature is extensively diluted by non-atmospheric mass-dependent sulfate leading to negligible $\Delta^{17}O$-excess in sediments. This is supported by $\Delta^{17}O$ as low as 0.2‰ in modern day evaporitic sulfate at the surface of Lake Tecopa (Bindeman et al., 2007).

We conclude that long term atmospheric deposition is incapable of explaining the large concentration of high $\Delta^{17}O$ sulfate in ash layers, which must be volcanic in origin. A few hundreds ppm SO$_4$ deposited in a few years into a meter-thick ash layers requires a sulfate accumulation rate ~10$^4$ times greater than the average atmospheric sulfate aerosol accumulation rate estimated above. Indeed, this can be achieved by large scale volcanic eruptions.

In Lake Tecopa, the highest $\Delta^{17}O$ measured in ash sulfates is 2.26‰, but this $\Delta^{17}O$ value was diluted by mass-dependent sedimentary sulfate ($\Delta^{17}O$ ~0‰). How high was the initial undiluted $\Delta^{17}O$ signature of those ash layers? If we assume that the overall amount of sulfate in the lake remained approximately constant through time, and use the estimate that ash represents 15–20 vol.% of sediments in the Lake Tecopa basin, and assume that the mass-independent signature of sulfate in sediments was derived from the ash, we can estimate the following initial range of $\Delta^{17}O$ in the freshly deposited layers of volcanic ash:

\[
\begin{align*}
\Delta^{17}O_{\text{ash}} & \times \Delta^{17}O_{\text{sed}} = \Delta^{17}O_{\text{ash}} + \Delta^{17}O_{\text{sed}} \\
\Delta^{17}O_{\text{ash}} \times \Delta^{17}O_{\text{sed}} & = \Delta^{17}O_{\text{ash}} + \Delta^{17}O_{\text{sed}} \\
\end{align*}
\]

$I$: initial; $F$: final

\[
\begin{align*}
X_{\text{ash}}& = \Delta^{17}O_{\text{ash}} \\
X_{\text{sed}}& = \Delta^{17}O_{\text{sed}} \\
\end{align*}
\]

The application of this simple mass-balance gives an initial $\Delta^{17}O$ for the MIVS between 6.5‰ and 8.5‰. This estimate approaches the theoretical maximum $\Delta^{17}O$ value of 8.0‰ that we discuss below. It is thus apparent that the initially large $\Delta^{17}O$ signal was diluted in ash layers while increasing $\Delta^{17}O$ up to 0.5‰ in sediments.

5.1.2. Mass-dependent sulfates

Sedimentary sulfate derived from weathering and oxidation of igneous sulfides during a few Ma should have $\Delta^{17}O$ value close to 0‰ and variable $\delta^{34}S$ and $\delta^{18}O$ values. In anoxic conditions, microbial activity can significantly affect $\delta^{34}S$ without changing $\Delta^{17}O$ signatures. However, the fact that the Lake Tecopa was a shallow lake (~<5–10 m; e.g. Larsen 2008) as evidenced by the presence of ripple marks and cross-laminations in the sediments, and the fact that no S-bearing minerals such as pyrite are observed in our samples suggest limited (if any) sulfate biogenic reduction effects on sulfate isotopic values.

In order to explain the ~8‰ $\delta^{18}O$ and ~5‰ $\delta^{34}S$ ranges measured in sulfate sediments, we suggest mixing of two sources of mass-dependent sediment-derived sulfates (MDSDS): 1) lacustrine or fluvial sulfates ($\delta^{18}O = -2\%\text{O}; \delta^{34}S = -5\%\text{S}$) and 2) evaporitic sulfates ($\delta^{18}O = -15\%\text{O}; \delta^{34}S = -22\%\text{S}$).

Lacustrine or fluvial sulfates are formed by aqueous abiotic or biological oxidation of S-bearing minerals. During such process the ambient water is the source of 50% to 100% of the sulfate oxygen depending on the pH or Fe$^{3+}$ concentration (Van Stempvoort and Krouse, 1994). The Lake Tecopa water likely had the same composition as the present day Amargosa river water, $\delta^{18}O = -13\%\text{O}; \delta^{34}S = -15\%\text{S}$ (Yang et al., 1997; Larsen et al., 2001) and $\Delta^{17}O = 0$‰. The sulfates in equilibrium with...
### Table 1

Sulfate compositions of Lake Tecopa ash layers and sediments.

<table>
<thead>
<tr>
<th>Samples</th>
<th>Height (m)</th>
<th>SO$_4$ (ppm)</th>
<th>CaCO$_3$</th>
<th>$\Delta^{18}O$</th>
<th>$\delta^{18}O$</th>
<th>$\delta^{34}S$</th>
<th>MIVS</th>
<th>MDSDS 1</th>
<th>MDSDS 2</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>1LCT+4</td>
<td>4.3</td>
<td>211</td>
<td>4</td>
<td>1.95</td>
<td>7.95</td>
<td>0.22</td>
<td>0.65</td>
<td>0.24</td>
<td>1.11</td>
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<tr>
<td>1LCT+4+</td>
<td>4.3</td>
<td>211</td>
<td>4</td>
<td>1.78</td>
<td>5.64</td>
<td>15.06</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1LCT+3</td>
<td>3.6</td>
<td>369</td>
<td>4</td>
<td>0.62</td>
<td>10.01</td>
<td>16.62</td>
<td>0.08</td>
<td>0.20</td>
<td>0.62</td>
<td>0.90</td>
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<tr>
<td>1LCT+2</td>
<td>2.6</td>
<td>38</td>
<td>4</td>
<td>0.96</td>
<td>8.49</td>
<td>16.87</td>
<td>0.17</td>
<td>0.39</td>
<td>0.18</td>
<td>0.74</td>
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<tr>
<td>1LCT+1</td>
<td>2.2</td>
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<td>0</td>
<td>1.55</td>
<td>4.76</td>
<td>14.28</td>
<td>0.19</td>
<td>0.64</td>
<td>0.21</td>
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<td>1LCT8</td>
<td>2.1</td>
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<td>0</td>
<td>0.92</td>
<td>1.97</td>
<td>15.74</td>
<td>0.11</td>
<td>0.86</td>
<td>0.13</td>
<td>1.10</td>
</tr>
<tr>
<td>1LCT8+</td>
<td>365</td>
<td>0</td>
<td>4</td>
<td>0.64</td>
<td>2.22</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1LCT7</td>
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<td>33</td>
<td>0</td>
<td>1.36</td>
<td>4.07</td>
<td>12.93</td>
<td>0.17</td>
<td>0.60</td>
<td>0.18</td>
<td>0.95</td>
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<tr>
<td>1LCT6</td>
<td>1.7</td>
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<td>0</td>
<td>0.78</td>
<td>-0.06</td>
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<td>0.10</td>
<td>0.81</td>
<td>0.01</td>
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<td>0</td>
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<td>4.39</td>
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...
this water would have a low $\delta^{18}O$ (~−2‰), $\delta^{34}S$~15‰ and $\Delta^{17}O$~0–0.2‰ (Fig. 5). Evaporitic sulfate are also generated by aqueous oxidation of S-compounds, however in this case, water evaporation induces isotopic fractionation, leading to the concentration of the heaviest isotopes. This is supported by the measurements performed in salt from Lake Tecopa $\delta^{18}O$~15‰, $\delta^{34}S$~22‰ and $\Delta^{17}O$~0‰ (e.g. Bindeman et al., 2007). Sulfate derived from the dissolution of old evaporitic deposits is also a possibility.

Another mass-dependent sulfate source that can provide long-term contribution is mass-dependent volcanic sulfates (MDVS). They are usually delivered by small eruptions that release SO2 that is oxidized by mass-dependent pathways during atmospheric transport. The MDVS may have a composition close to the tropospheric processed sulfate precipitated in volcanic plumes, close to the eruption center with $\Delta^{17}O$~0‰, $\delta^{18}O$~−6‰ and $\delta^{34}S$~−6–8‰ as measured in different studies (Luhr and Logan, 2002; Bao and Reheis, 2003; Bindeman et al., 2007; Mather et al., 2007).

### 5.1.3. Mixing model

Fig. 5 shows $\Delta^{17}O$ vs. $\delta^{18}O$ and $\Delta^{17}O$ vs. $\delta^{34}S$ correlations, where these sulfate source end-members are plotted. Data is fitted with an ash-

![Fig. 4. SEM images and X-ray mapping (current of 15 kV and 2 nA) of sediment (2BT+1) and ash (LCT8) samples. Ca concentration is in red and S concentration in blue. The arrows show where Ca and S maps overlap indicating the presence of CaCO3 crystals aggregates (gypsum or anhydrite).](image-url)
sediment mixing model, which determines their relative contribution in each of our samples. As the main interest of this study is to trace MIVS signature, the proposed mixing model consists of MIVS and two MDSDS end-members. In order to calibrate the robustness of the linear model, several different combinations of potential isotopic values of end-member sulfate sources were tested. The isotopic composition of the end-members that best fit our data have the following composition:

MIVS: $\delta^{18}O = 1.5 \text{‰}$, $\delta^{34}S = 0 \text{‰}$ and $\Delta^{17}O = 8 \text{‰}$ and

MDSDS: lacustrine or fluvial: $\delta^{18}O = -2 \text{‰}$, $\delta^{34}S = 15 \text{‰}$ and $\Delta^{17}O = 0 \text{‰}$ and evaporitic: $\delta^{18}O = 20 \text{‰}$, $\delta^{34}S = 12 \text{‰}$ and $\Delta^{17}O = 0 \text{‰}$. The relative participation of each end-member in our samples has been evaluated by monitoring the standard deviation of the sums of the three source proportions; such error propagation approach is similar to that proposed by Bao and Marchant (2006). However, if other computed mixing models are performed with $\delta^{18}O$ and $\delta^{34}S$ – 3‰ and $\Delta^{17}O$ 0.2 – 0.3‰, isotopically-different end-members, similar conclusions are drawn from our dataset.

The calculated proportions of MIVS yield 23% in LCT, 18% in BT and 9% in HRT, decreasing with age (Table 1 and Fig. 3). There is no evidence supporting that the far larger (2500 km$^3$), HRT, eruption initially contained less MIVS than the 1000 km$^3$ LCT or the 750 km$^3$ BT. Rather, we interpret its lower MIVS contents as resulting from greater dilution of HRT sulfate by MDSDS with time.

This mixing model demonstrates that the main source (>77%; Table 1) of sulfate in the Lake Tecopa stratigraphic record is mass-dependent sedimentary sulfate (MDSDS). Assuming a more-or-less constant isotopic composition of the sedimentary sulfate sources with time, the fluctuations between “lacustrine or fluvial” and “evaporitic” sources could reveal the oscillating lake’s depth history. The proportion of MDSDS in sediments least affected by ash leachates reaches...
98%, while the ash layer itself is quite dilute and contain between 77% and 93% of MDSDS (Table 1), which represents between 15 and 45 ppm of MIVS remaining in ash layers.

5.1.4. Preservation of $\Delta^{17}$O signal in sulfate

In the case of complete sediment-ash sulfate mixing, one would expect homogeneous sulfate concentrations, and isotopic values, through the sedimentary record. Higher sulfate concentration in sediment than ash (Table 1 and Fig. 6) indicates that sulfates from ash and sediment are not fully mixed. Thus, one major result of the mixing model, is the demonstration that although heavily diluted, the MIVS is preserved in ash beds from Lake Tecopa for hundreds of thousands to few million years. The proportion of MIVS in ash layers (and $\Delta^{17}$O signature) decays with age and/or thickness of the volcanic deposits from LCT through BT to the HRT ash layer (Fig. 3). Consequently, the $\Delta^{17}$O signal decays in the geologic record can be quantified using our three data points (LCT, BT and HRT) with a hypothetical modern “supervolcanic” ash deposit with undiluted 100% MIVS with $\Delta^{17}$O = 8‰. In this quantification, it is assumed that the modern supereruption and ash deposition happened 10 years ago, and was followed by continuous acid precipitations. We observe that the best $\Delta^{17}$O values happened 10 years ago, and was followed by continuous acid precipitations. We observe that the best value is as high as 8‰, precluding an important role played by the H$_2$O$_2$ oxidation pathway (R1). This also leads to the conclusion that the oxidation pathways (R2 and/or R3) play a major role in the formation of MIVS. On average, the atmospheric ozone, which is 90% stratospheric, has an average $\Delta^{17}$O value of 32‰ and a high $\delta^{18}$O value up to 100–150‰ (e.g. Kranckowsky et al., 2000). If OH radicals possess the same $\Delta^{17}$O as their ozone precursors during photodissociation, SO$_2$ oxidation into MIVS inherits 1/4 of the oxygen atoms from the mass-independent O$_2$ (R2) or OH radicals (R3) leading to $\Delta^{17}$O of about 8‰ (e.g. Thiemens, 2006). High $\delta^{18}$O (~15‰) and low $\delta^{34}$S (~0‰) are also expected for MIVS as deduced from the correlation between these isotopic ratios and $\Delta^{17}$O (Fig. 5). It appears from Bao et al. (2003) and our dataset that the processed high $\Delta^{17}$O sulfates are lighter in terms of S-isotopes than the initial volcanic SO$_2$.

5.3. Incorporation of MIVS in ash layers, and surface dynamics of its accumulation in sediment basins

Supereruptions release $\geq 10^8$ km$^3$ of ash and $>10^{9}$ megatons of SO$_2$ into the atmosphere (e.g. Wallace, 2001). Although by air-fall deposition ash blankets wide areas after a few days to a few weeks (e.g. HRT: ~3.106 km$^2$; Toba (74 kyr): $\geq 5.10^6$ km$^2$), ice core records from Antarctica show that the oxidation of SO$_2$ gas to SO$_3$ and H$_2$SO$_4$ aerosols requires months to years (e.g. Zielinski et al., 1996). Thus MIVS can fall as sulfuric acid precipitation: wet (acid rain), or dry (e.g. “dry fogs”). Then, it is incorporated into ash-covered ground and forms gypsum (Fig. 4), months to years after the ash deposition itself. Calculations suggest that MIVS concentration of 50ppm are possible if the majority of sulfuric acid precipitation occurs in ~5000 km radius from the supervolcano, due to atmospheric circulation (e.g. cyclonic current) that is able to concentrate aerosols clouds in the upper troposphere and lower stratosphere as observed with smaller-scale eruptions (e.g. Rose et al., 2001). In order to account for the meter-thick deposits in Lake Tecopa and similar basins elsewhere, local fluvial and/or eolian redistribution of ash (and sulfate) from the watershed area is required (e.g. Rose et al., 2003). Then, as observed for each outcrop, prolonged intra-basin lacustrine and fluvial reworking of sediment and ash (and sulfate) occurs.

5.4. Mass independent $\Delta^{33}$S and $\Delta^{34}$S

Our detailed isotopic dataset for ash in Lake Tecopa permits discussion on rare sulfur isotopes fractionation after large vs. small volcanic eruptions, and comparison with sulfate in Antarctic ice. The latter shows mass-independent $\Delta^{33}$S of $-0.5\%$ to $1.1\%$ and $\Delta^{34}$S of $-2.7\%$ to 1.98‰ (Savario et al., 2003b; Baroni et al., 2008; Fig. 7), but only small $\Delta^{35}$S ($-0.03\%$ to $-0.07\%$) and $\Delta^{36}$S ($+0.03\%$ to $+0.28\%$) values were previously reported in Lake Tecopa LCT and BT ash beds (Bindeman et al., 2007; Fig. 7). Because these values were small, they were earlier interpreted to reflect mass-dependent processing by kinetic gas phase reaction and/or Rayleigh distillation involving a residual Rayleigh reactant (Bindeman et al., 2007). Since significant $\Delta^{33}$S and $\Delta^{34}$S have not been measured in sulfate from small eruption deposits (Bindeman et al., 2007; Fig. 7), we need to
find a process able to explain the significant mass-independent $\Delta^{33}$S and $\Delta^{34}$S signatures characteristic of sulfate from large eruptions.

Here, we discuss a process able to explain these small but significant $\Delta^{33}$S and $\Delta^{34}$S values measured in sulfate from LCT and BT samples from Lake Tecopa. The linear mixing model above suggests that MIVS constitutes 20–28% of LCT and 10% of the BT sulfates in these samples. Assuming that MIVS was diluted by MDSDS with $\Delta^{34}$S = 0‰ and $\Delta^{36}$S = 0‰, we obtain initial undiluted MIVS values of $\Delta^{33}$S = −0.24‰ to −0.35‰ and $\Delta^{34}$S = 0.27‰ to 1.08‰ (Fig. 7). When plotted together with data for the stratospheric sulfate from Antarctic ice, the estimated MIVS compositions from Lake Tecopa ash extends the negative correlation trend, and plot between photolytic wavelength of 220–248 nm (e.g. Farquhar et al., 2001). This suggests a common photolytic mechanism for both, but also suggest that sulfates in ash layers and in Antarctic ice are generated by different timescales and processes. We propose that ash-sulfates are quickly generated and removed from the stratosphere by ash sedimentation and acid precipitations. Pavlov et al. (2005) modeled that volcanic sulfates generated by photolysis of SO$_2$ have initially small negative $\Delta^{34}$S that later becomes positive. However, Baroni et al. (2007) measured in volcanic sulfate deposited in Antarctica large positive $\Delta^{34}$S values that become negative due to mass-balance effects.

We conclude that sulfate in ash layers most likely record isotopic values of the first volcanic sulfate removed from the atmosphere, with small and negative $\Delta^{33}$S as inferred from the Pavlov et al. (2005) model. On the other hand, ice cores record the volcanic sulfate generated after a long journey in the stratosphere, leading to large mass-independent $\Delta^{33}$S signatures due to long UV radiation exposure as observed in Baroni et al. (2007).

5.5. Climatic impact of supereruptions

The chemical and physical state of the atmosphere after super-volcanic eruptions is a matter of considerable debate and extrapolative modeling (e.g. Robock, 2004). The large $^{17}$O-excesses measured in the Lake Tecopa sulfate indicate atmospheric reaction with ozone or O$_3$-derived compounds to make H$_2$SO$_4$ (reactions R2 and R3) and this process is able to consume ozone and water; thus we discuss the possibility of atmosphere hydration or dehydration and ozone depletion.

5.5.1. Stratosphere hydration or dehydration: implications on MIVS formation

The SO$_2$ oxidation by OH$^*$ radical produces highly hydroscopic SO$_4^-$ compounds that quickly react with water to produce H$_2$SO$_4$ (reaction R3). This reaction feeds the discussion on the relative role of volcanic SO$_2$ in drying out the stratosphere (e.g. Bekki, 1995). However, it is not yet clear how much water is injected into the stratosphere by a major volcanic plume. The generalization of stratospheric “watering”, as observed after the 1982 El Chichón eruption, is still under debate (e.g. Shepherd, 2002; Robock, 2004).

The hydration/dehydration balance after a supereruption may be mostly controlled by the physical state of the tropopause. The tropopause is characterized by a sharp decrease of atmospheric temperature, which normally prevents water transfer from the troposphere to the stratosphere. After large volcanic eruptions, the tropopause, contaminated by sulfate aerosol clouds, is heated up by absorption of the incoming solar radiation (e.g. Robock, 2000), a process able to increase the water flux from the troposphere to the stratosphere (e.g. Joshi and Shine, 2003). As a result, an important consequence of the stratosphere hydration after a supereruption is that due to faster reaction rates (Seinfeld and Pandis, 1998), heterogeneous oxidation reactions R1 and/or R2 play a more important role. Therefore, in a “hydrous” stratosphere, the ozone mass-independent signature would be transferred to volcanic sulfate aerosols more efficiently by the direct SO$_2$ oxidation via O$_3^*$ (R2) rather than by OH$^*$ radicals (R3).

5.5.2. Ozone consumption

In either process of stratospheric dehydration or hydration, ozone molecules are consumed during the oxidation of volcanic SO$_2$. In 1991, the 10 km$^3$ eruption of Pinatubo released ~20 Mt SO$_2$ and resulted in
an ozone layer depletion of 3–4%, directly attributable to the eruption (Gleason et al., 1993; McCormick et al., 1995). The ozone layer consumption would be much more dramatic after LCT, BT and HRT supereruptions that released 75 to 250 times more SO₂.

The ozone concentration stays approximately constant due to the balance between formation and destruction processes (e.g. Seinfeld and Pandis, 1998). During and shortly after a supereruption, the atmosphere will not be able to counterbalance the ozone depletion due to volcanic aerosol formation. The oxidation of massive amounts of volcanic SO₂ that leads to MIVS can consume ozone indirectly (R2) or directly (R3), leading to temporal decrease of global ozone concentration. When the volume of volcanic SO₂ released by an eruption and the Δ¹⁷O measured in sulfate from the ash deposit are considered together, it is possible to estimate the mass of ozone consumed indirectly (R2) or directly (R3), leading to temporal decrease of global ozone concentration. When the volume of volcanic SO₂ released by an eruption and the Δ¹⁷O measured in sulfate from the ash deposit are considered together, it is possible to estimate the mass of ozone consumed indirectly (R2) or directly (R3), leading to temporal decrease of global ozone concentration.

As the SO₂ is capable of absorbing short wavelength UV radiation (Finlayson-Pitts and Pitts, 2000), therefore prior its oxidation into sulfate it protects the Earth surface while the ozone production is decreased. Nonetheless, depending on the atmospheric circulation, heterogeneous distribution of SO₂ may lead to local ozone holes and direct exposure of the Earth surface to short wavelength UV radiations; and these consequences would be of prime importance for life on the Earth.

5.5.3. Tropospheric vs. stratospheric sulfate transport during supereruptions

Due to remoteness of Lake Tocop (Fig. 1), and the fact that atmospheric precipitation of MIVS occurred in a ~5000km radius from the eruptive centers, stratospheric jet stream transport of ash and aerosols is obviously required.

Moreover, this study shows that up to 20–60% of global ozone could be consumed after supereruptions. As only 10% of the ozone is tropospheric, consumption of at least 10% of the stratospheric ozone and generation of stratospheric MIVS is required.

Confirmation of the mass-invariant signature of S-isotopes in LCT and BT sulfate indicates a volcanic SO₂ photolysis in the upper stratosphere, unless a local ozone hole created by a supereruption permits SO₂ photolysis at lower altitudes.

The preceding discussion leads to the conclusion that mass-independent signatures of volcanic sulfates from supereruptions have features of both tropospheric and stratospheric transport. This may not necessarily be mutually exclusive and reflect physical degradation by temperature increase and chemical degradation by ozone-depleting reactions of the tropopause. We can speculate that the physical and chemical distinction between high-troposphere and low-stratosphere, at least on a local scale, is erased by supereruptions.

Acknowledgement

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.epsl.2009.03.005.

References


1. Sulfate extraction from samples

In order to extract all soluble sulfates, samples weighing 40-80 g were placed in 1L polypropylene beaker containing diluted HCl (0.1M) with a ratio sample/acid usually of 1/5. The pH of the solution was monitored to be between 1 and 2 and for the carbonate-bearing samples, concentrated HCl was added in order to maintain the required pH. After 24 hours at room temperatures, the beakers were heated up at 50-60°C during 1-2 hours before filtration through filter paper. The filtered solution was collected in glass beakers which were left evaporating at 60-80°C on a hot plate. While the solution was evaporating, barium chloride (BaCl₂) was added to the solution in order to precipitate barite (BaSO₄). Usually, after few hours, a white barite powder precipitated; then an excess of BaCl₂ was added to the remaining solution to scavenge it completely of SO₄²⁻ ions for 1-2 hours. We noticed that the two-steps addition of BaCl₂ results in larger crystals (~7-10 μm) of BaSO₄ than a single stage addition of BaCl₂ to the pre-evaporated solution. The resultant barite powder was rinsed 3 times with deionized water and dried out in oven at 60-80°C. It is important to note that during all this process (~48 hrs), isotopic exchanges between water and sulfate are expected to be negligible (Van Stempvoort and Krouse, 1994), because SO₄²⁻ is not a labile anion at these conditions. The sulfate concentration was estimated from the weight of extracted barite from samples on known weight.

For 16 samples, barite was purified from nitrate contamination by dissolution-reprecipitation method using DTPA, since high Δ¹⁷O measured in barite samples from some studies was inferred as coming from absorbed nitrate contamination (e.g. Bao, 2006). Around 20 mg of barite powder was placed in 30 mL of diethylene triamine penta acetic acid (DTPA) solution (0.05M DTPA in 1 M NaOH). After 1-2 hours at room temperature the barite is totally dissolved. Then, we acidified the solution by adding few drops of concentrated HCl (12M) in order to reprecipitate all the barite within 1-2 hours. Few drops of BaCl₂ were also added to help the BaSO₄ reprecipitation. After decanting, the barite powder was rinsed 3 or 4 times with deionized water and dried out. It is noteworthy that DTPA is an N-bearing compound, therefore it is important to abundantly (at least 3-4 times) rinse the purified barite in order to remove all the DTPA from the solution before drying it out and thus preventing any N-contamination that could be due to the DTPA treatment itself. In such a case, the mass interference of NF⁺ on the mass 33 during the supposed “purified barite” analyses will falsely increased the Δ¹⁷O (see below).

In order to quantify the nitrate contamination we run our 6 sulfate samples dissolved in DTPA solution were also run on an ion chromatograph at the Department of Chemistry at the University of Oregon. The measured NO₃⁻/(NO₃⁻+SO₄²⁻) in ash samples are: 2.8 wt.%, 2.4 wt.%, 0.7 wt.%, 0.7 wt.% and 0 wt.% and in one sediment: 0 wt.%.

Therefore, we performed two consecutive dissolution-reprecipitations of each barite powder to make sure that all nitrate occlusions was removed. However, as we demonstrate above the nitrate contamination is systematically lower than 3 wt.% in our samples and as we discuss below, the Δ¹⁷O measured in purified samples is, in the uncertainties, the same that what we measured in unpurified barite. Therefore we did not attempt to carry DTPA treatment for every sample.

2. Oxygen isotope analyses

Oxygen isotopes were analyzed in the Stable Isotope Laboratory at the University of Oregon. Molecular oxygen was extracted from 4-7 mg of barite powder by a 35W CO₂-laser using BrF₃ as reagent and purified from traces of the reagent cryogenically and through the boiling mercury diffusion pump. Extracted O₂ gas was then frozen on molecular sieve (zeolite
13Å) before being expanded into the bellows of a Finnigan MAT253 mass spectrometer and run in a dual inlet mode for 8 cycles.

Some samples were run in both single and double trap system in which O2 gas was refrozen on the second 13Å sieve before being expanded into the mass spectrometer. In the later, the O2 gas extracted from the sample was trapped in the first molecular sieve that was at liquid nitrogen temperature (-196°C). Then we removed the liquid nitrogen flask from the first trap to the second one. Progressively the O2 gas was transferred from the first to the second trap leaving most impurities in the first molecular sieve. When all O2 gas was trapped in the second molecular sieve, the liquid nitrogen flask was removed and the oxygen expanded into the mass spectrometer leaving most of the impurities left in the second molecular sieve. Between each samples, the two molecular sieves were heated up with a heat gun and left degassing for at least 10 minutes. This technique attempted to prevent any traces of nitrogen fluoride contamination leading to isobaric interference of NF+ ion on the mass 33 affecting 17O18O signal (e.g. Rumble III et al., 1997; Pack et al., 2007). The Background scanning of nitrogen masses was monitored on the mass 52 (NF2+) and gave systematically, when detectable, less than 1-2mV. We determined that atmospheric nitrate contamination was not present in our high-vacuum laser line.

Sample duplicates, and replicates of barite international standard NBS127 (n = 7) and in-house standard (IBO4-13; n = 12) gave a reproducibility (in 2σ) usually better than δ18O ± 0.6‰, δ17O ± 0.3‰ and Δ17O ± 0.05‰. The 35-45% O2 yield during barite laser fluorination leads to low-δ18O values, therefore based on the certified δ18O value of NBS127 (δ18O = 8.6‰), we applied a correction factor of +8.01‰ for all analyzed samples. Notice that this correction does not affect Δ17O values, which is a mass-independent parameter. Day to Day variations were also monitored by the GMG garnet standard.

Sulfate extraction and oxygen isotope analysis was replicated for few samples in order to assess the external errors of the whole method. Repeated extractions from different aliquots of the same sample of ash or sediment give reproducibility (in 2σ) better than δ18O ± 1.2‰, δ17O ± 0.7‰ and Δ17O ± 0.1‰. 

We performed measurements on 16 purified barite powders using the DTPA dissolution-reprecipitation method. The Fig.1 shows that all samples plot very close to the line 1:1, meaning that the differences in Δ17O between the purified and unpurified barite are minimal or insignificant (≤0.1‰ out of the analytical uncertainties in 2σ). It is noteworthy that Δ17O values of the purified barite samples are either identical or sometimes 0.1‰ lower than the unpurified one (Fig. 1). This negative deviation can be due to the removal of minor environmental nitrate contamination (<2-3wt.%).

In order to test the hypothesis of mass interference of NF on the mass 33 and to remove hypothetical NF3 contaminant, we analyzed few (N=3) samples with a double molecular sieve (zeolite 13Å) trap system. The difference in Δ17O between the single and double sieve trap systems is negligible (<0.05‰; on the line 1:1 Fig. 1) for two of the analyzed samples. The sample having the highest difference (see the circled points in the Fig. 1) yielded Δ17O = +0.33‰, which is comparable to what we obtained for the DTPA purified barite (Δ17O = +0.40‰). If the contaminant nitrate was mass-independent, the 17O-excess due to this contamination would be homogenized in the O2 gas during the fluorination and won’t be removed by the double sieve trap system. However, in both methods, we are able to prevent the NF mass 33 interference, meaning that in this experiment the source of the Δ17O increase in the untreated sample come from the NF mass interference and not from the mass-independence of the contaminant nitrate. It is important to note that this nitrate contamination in ash samples can potentially slightly increase the Δ17O of less than 0.1‰ (out of the uncertainties) in a sample having Δ17O more than 10 times higher (1.25‰. Fig. 1). Therefore,
it won’t drastically change our overall interpretations that are mainly based on the $\Delta^{17}\text{O}$ variations of 2.3‰ in the Lake Tecopa stratigraphic section.

**Figure Caption**

Figure 1: Comparison between unpurified and purified barite using the DTPA dissolution-reprecipitation method (Bao et al.2006) or purifying the $\text{O}_2$ gas on the fluorination line with a double molecular sieve trap system. The error bars are ±0.1‰ (overall analytical uncertainties in 2σ) for each point and the 1:1 line. This graph demonstrates that all data points plot on or very close to the 1:1 line, indicating the robustness of analyses of unpurified barite with a single molecular sieve trap.