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An isotopic analysis of ionising radiation as a source of sulphuric acid

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Abstract. Sulphuric acid is an important factor in aerosol nucleation and growth. It has been shown that ions enhance the formation of sulphuric acid aerosols, but the exact mechanism has remained undetermined. Furthermore some studies have found a deficiency in the sulphuric acid budget, suggesting a missing source. In this study the production of sulphuric acid from SO2 through a number of different pathways is investigated. The production methods are standard gas phase oxidation by OH radicals produced by ozone photolysis with UV light, liquid phase oxidation by ozone, and gas phase oxidation initiated by gamma rays. The distributions of stable sulphur isotopes in the products and substrate were measured using isotope ratio mass spectrometry. All methods produced sulphate enriched in 34S and we find an enrichment factor (δ34S) of 8.7 ± 0.4 ‰ (1 standard deviation) for the UV-initiated OH reaction. Only UV light (Hg emission at 253.65 nm) produced a clear non-mass-dependent excess of 33S. The pattern of isotopic enrichment produced by gamma rays is similar, but not equal, to that produced by aqueous oxidation of SO2 by ozone. This, combined with the relative yields of the experiments, suggests a mechanism in which ionising radiation may lead to hydrated ion clusters that serve as nanoreactors for S(IV) to S(VI) conversion.

1 Introduction

Aerosols – small particles suspended in air – are frequently formed by condensation of gas molecules in Earth’s atmosphere. The effect of these particles on the radiative balance of our climate system, through direct and indirect mechanisms, is the greatest uncertainty in the radiative forcing budget (Forster et al., 2007). One issue is that the mechanism behind the formation of aerosols remains unknown, even though several mechanisms have been proposed (Kulmala, 2003; Curtius, 2006; Kurtén et al., 2008) and major advances have recently been made (Kirkby et al., 2011). One of these mechanisms is that of ion-induced nucleation (Enghoff and Svensmark, 2008; Kazil et al., 2008; Hirskko et al., 2011) where ionising radiation promotes the early stages of the aerosol nucleation. The prevailing theory on exactly how the ions affect nucleation is that they lower the energy barrier for condensing molecules caused by the increase in energy required to create the interface between the aerosol phase and the gas phase – the surface tension (Lovejoy et al., 2004). Other factors include the recombination of oppositely charged clusters (Yu and Turco, 2000) and the increased growth rate caused by a charge in a molecular cluster (Yu and Turco, 2001). Ionic cluster formation is estimated to account for 1–30% of total particle formation, based on 7 European
measurement sites (Manninen et al., 2010) and ionisation by galactic cosmic rays may increase binary (sulphuric acid and water) and ternary (sulphuric acid, water, ammonia) nucleation rates of nanometer size particles in the tropospheric boundary layer (Kirkby et al., 2011). Svensmark et al. (2007) proposed a chemical mechanism where sulphuric acid is produced by ion chemistry, but until now there has been no experimental evidence supporting this mechanism.

Sulphuric acid is thought to be one of the main components of aerosols nucleated in the atmosphere (Sipilä et al., 2010) and an extra source might therefore be important for understanding aerosol formation. During a measurement campaign in Hyytiälä, Finland the calculated sulphuric acid concentrations were about 10 percent lower than the measured values (Boy et al., 2005), and a study at the Antarctic also showed a missing sulphuric acid source (Jefferson et al., 1998) stimulating the search for possible additional sources of sulphate. An ion mechanism could additionally operate at night, providing a source of sulphuric acid not depending on sunlight.

The initial steps of a mechanism involving ion chemistry have been investigated by e.g. Fehsenfeld and Ferguson (1974) and more recently by Bork et al. (2011a,b) who used ab inito calculations to show that the gas-phase oxidation of SO$_2$ by ozone is facilitated in ionic clusters, but until now there has been no direct experimental investigation. One of the difficulties of such a study is that nucleation requires a certain amount of condensable gas and thus either a liquid or photochemical source of sulphuric acid is normally used (Berndt et al., 2006). Any effect from an ionising source then occurs in parallel with this other source and is thus hard to distinguish. An early study by Raes et al. (1985) showed an increase in particle number when a gamma source was used in addition to photochemical sulphuric acid production and they speculated that both UV light and gamma rays could produce sulphuric acid from SO$_2$.

An alternative technique for distinguishing in situ processes is isotopic analysis. Fractionating processes alter the isotopic composition of chemical species and are used to constrain atmospheric budgets or interpret geological records and ice cores, e.g. Johnson et al. (2002); Brenninkmeijer et al. (2003). Traditional physical processes and most chemical reactions follow a so-called mass-dependent relationship (MDF, mass-dependent fractionation), where enrichments of different isotopes follow a linear relationship according to the ratio of their masses. A deviation from the linear behavior is sometimes observed, but the origin(s) of non-mass-dependent fractionation, abbreviated NMD, is not fully understood. NMD is, however, known to be produced by a small number of mechanisms occurring in the gas phase including photolysis and recombination reactions, as well as some nuclear processes (Guo et al., 2010; Romero and Thiemens, 2003; Thiemens, 2006). Photolytic reactions can result in NMD due to differences in absorption cross sections caused by isotopic substitution (Danielache et al., 2008; Schmidt et al., 2011).

Sulphur has four stable isotopes ($^{32}$S, $^{33}$S, $^{34}$S, and $^{36}$S) and is widely used for isotopic analysis. It is known that the photolytic fractionation of sulphur compounds is wavelength dependent (Farquhar et al., 2000b) and another possible source for NMD is the oxidation channel of excited SO$_2$, a process very likely to happen in the atmosphere. NMD signatures have been found in atmospheric sulphate aerosols (Baroni et al., 2007; Pavlov et al., 2008) and Archean rock samples (Farquhar et al., 2000a; Ono et al., 2003; Ueno et al., 2008; Bao et al., 2007). In a previous study, Harris et al. (2012) investigated different gas and liquid phase oxidation pathways of SO$_2$ in laboratory experiments, but did not observe clear NMD with respect to $^{33}$S, while the sulphate production processes under consideration were generally distinguishable by their $^{34}$S fractionation factors.

While the isotopic fractionations of SO$_2$ by ionising radiation are not known, NMD is not expected. From the Born-Oppenheimer approximation it is evident that molecular polarizability is not changed by isotopic substitution. In particular the S in SO$_2$ is at a symmetric site and it has previously been shown, for ozone, that NMD effects are primarily associated with assymmetric sites (e.g. the two end atoms in ozone) but not at symmetric sites, such as the central atom in ozone (Gao and Marcus, 2001).

In this study we generate sulphuric acid using UV light, gamma sources, and via ozone oxidation in the aqueous phase. Using multiple sulphur isotopic measurements allows us to distinguish between these reactions and evaluate their relative importance.

2 Experimental setup and procedures

The experiments were performed using two separate systems. Sulphuric acid was generated using a chemical reaction chamber at the National Space Institute in Copenhagen. Samples were collected as BaSO$_4$ by flowing the air leaving the reactor through a series of bubblers containing BaCl$_2$. The isotopic composition of the samples were then analysed with an isotope ratio mass spectrometer (IRMS) at the Tokyo Institute of Technology. The following sections describe each experimental system and the procedures employed.

2.1 Chamber setup (Copenhagen)

A schematic of the setup can be seen in Fig. 1. The reaction volume was contained in a cylindrical 501 reaction chamber (100 cm long with a diameter of 25 cm) made of electropolished stainless steel, previously described by Enghoff et al. (2008). A mixture of pure humidified air (Labline 5.0, Strandmøllen, further cleaned with a Thermo Scientific model 1160 Zero Air Supply) was continuously flowed through the vessel at a rate of 1.01 min$^{-1}$ (standard
temperature and pressure (STP)). Previous to entering the chamber 0.41 min⁻¹ of the purified air went through an ozone generator (using photolysis at 184 nm) resulting in about ~400 ppb ozone in the chamber. The rest of the air (0.61 min⁻¹) was humidified by circulating purified (milli-Q) water through a Gore®Ex tube inserted into the air stream, giving a relative humidity of 40–45 %. In addition 100 µl min⁻¹ (STP) of pure SO₂(g) evaporated from a container (Gerling Holz+Co, ≥ 99.95 % wt.) was sent through the chamber. All air entering the chamber was additionally cleaned with a PALL Kleenpack 3 nm particle filter.

The temperature and absolute pressure were that of the laboratory, typically 22 °C (with UV lamps off) and 30 °C (UV on) and 1 bar, with a 1 mbar overpressure in the reaction vessel. Temperature, absolute and differential pressure, and relative humidity were monitored.

An array of UV (Philips TUV PL-L 18 W mercury discharge) lamps were used to illuminate the chamber with UV-radiation primarily at 253.65 nm (the lamp spectrum is shown in the Supplement). The lamps were placed about 20 cm from the chamber, separated by 2 layers of Teflon foil to maintain an air-tight system but transmit UV. The UV light initiates photochemistry in the gas phase, where ozone is photolysed producing excited oxygen atoms that form OH via reaction with H₂O, initiating the oxidation of SO₂ into sulphuric acid (see Appendix C).

Alternatively the chamber could be exposed to gamma radiation from two 35 MBq Cs-137 sources, placed 27 cm from the side of the chamber.

The outlet air from the chamber was connected to a series of glass gas washing bottles (typically 3) used to collect the sulphuric acid. These bottles contained 400 ml of milli-Q water adjusted to pH 2 using hydrochloric acid with an excess (1.0 g) of barium chloride (BaCl₂(H₂O)₂) compared to the amounts of sulphate formed in the experiments (milligrams). The bottles were magnetically stirred to maximize the air-water interaction. When the sulphuric acid entered the bottles it reacted with the barium ions to form BaSO₄ which has a very low solubility in water. The pH of 2 was chosen to prevent SO₂ from the gas phase from dissolving in water forming the sulphite ion (SO₃²⁻), which could potentially precipitate as BaSO₃. Acidic conditions at pH 2 shift the equilibrium away from SO₃²⁻, instead favouring aqueous SO₂ and HSO₃⁻. We attempted to dissolve a small part of some of the samples with 4 M HCl. It did not dissolve, demonstrating that the product was not BaSO₄.

All experiments ran for one week. The sample was then collected by allowing the BaSO₄ to settle, decanting the excess liquid and thoroughly resuspending the remaining solids in milli-Q water. This procedure was repeated three times for each sample to remove any remaining BaCl₂. The remaining solution was transferred to a smaller container and the water was evaporated in an oven, leaving the BaSO₄ as a powder.

The first column of Table 1 shows the experiments used for further analysis. Five different types of experiments were done, to distinguish oxidation pathways for SO₂. UV light and the gamma source were used separately and together. In addition, a series of experiments were run without UV and gamma source to quantify any uptake and oxidation of SO₂ in the liquid phase. Finally, an experiment was run with no UV, no gamma source, and no ozone, which did not yield any detectable amount of sulphuric acid.

The isotopic composition of the liquid SO₃ had to be determined, as a reference point for the isotopic data. About 5 ml of SO₂ (0.2 mmol) was added to a solution consisting of 500 ml milli-Q water with 15 mmol H₂O₂, as oxidizing agent. The solution was adjusted to pH 11 before SO₂ was added to ensure dissolution. The solution was stirred overnight and BaCl₂(H₂O₂)₂ was added. The resulting BaSO₄ was collected using the method described above. A second reference was prepared in the same fashion, except pH was not adjusted, in order to test for any pH dependence, even though the reaction is supposedly almost pH-independent (Seinfeld and Pandis, 2006, chap. 7). The two references yielded 41.8 and 36.7 mg of BaSO₄, respectively, from the ~5 ml of SO₂. This corresponds to 0.18 and 0.16 mmol. The collected amounts suggest full conversion of the SO₂, considering the uncertainty of determining the volume of SO₂ transferred and losses of BaSO₄ from sample collection. A small sample of the reference could not be dissolved with 4 M HCl, showing that the product was sulphate and not sulphite.

2.2 Isotope fractionation setup (Tokyo)

The collected BaSO₄ was converted to SF₆ for isotopic analysis. The first step was conversion into Ag₂S: the BaSO₄ was held at 300 °C and reduced to H₂S with the modified Kiba reagent (concentrated, dehydrated phosphoric acid containing Sn^{2+}, Sakai et al., 1984). The generated H₂S was then
Table 1. Samples and results. The table displays a sample ID, the mass of BaSO₄ recovered, and the isotopic fractionation data (normalized against the references as defined in Appendix B). The sample ID refers to the method of sulphate production: O₃ is the liquid phase oxidation by ozone, UV is when the UV lamps were turned on, and γ is when the gamma source was used. The number at the end of the ID refers to the position of the bottle in the sampling line (see Fig. 1).

<table>
<thead>
<tr>
<th>Sample ID</th>
<th>BaSO₄ Rec mg</th>
<th>δ³⁴S ‰</th>
<th>Δ³³S ‰</th>
</tr>
</thead>
<tbody>
<tr>
<td>O₃-1</td>
<td>2.8</td>
<td>16.0</td>
<td>-0.170</td>
</tr>
<tr>
<td>O₃-3</td>
<td>392.8</td>
<td>10.4</td>
<td>-0.072</td>
</tr>
<tr>
<td>O₃-UV-1</td>
<td>26.1</td>
<td>8.7</td>
<td>0.296</td>
</tr>
<tr>
<td>O₃-UV-3</td>
<td>63.7</td>
<td>12.6</td>
<td>-0.063</td>
</tr>
<tr>
<td>O₃-γ-1</td>
<td>19.4</td>
<td>13.9</td>
<td>-0.041</td>
</tr>
<tr>
<td>O₃-UV-γ-1</td>
<td>16.8</td>
<td>9.7</td>
<td>0.232</td>
</tr>
</tbody>
</table>

3 Results and discussion

The results of the analysis are presented in Table 1. Columns three and four of the table show δ³⁴S and Δ³³S for all samples. Data for Δ³⁶S was also measured and is presented in Appendix A. The fractionation data of δ³⁴S and Δ³³S is presented graphically in Fig. 2. Definitions for the isotopic designations are given in Appendix B.

3.1 Sulphate production

The sulphate mass obtained from each bottle is listed in the second column of Table 1. Sulphate is generated in the setup in all different operation modes and yields vary widely – for instance more sulphuric acid is formed with UV light present than for the gamma source but the sources combined yield less than each individual source. Seeing sulphate in the samples where UV and gamma radiation was not used indicates a process occurring in the trapping bottles themselves. The experiment without ozone, which showed no precipitation, strongly indicates that this process is liquid phase oxidation of SO₂ by ozone where the reactants become dissolved in the liquid phase depending on their Henry’s Law constants, leading to the conversion of SO₂ to H₂SO₄ and not simply dissolved SO₂ precipitating as BaSO₄. Note that the yields in the experiment without UV and gamma radiation increased for each bottle counting from the chamber and that the isotopic signature of this material is consistent with oxidation of SO₂ by O₃. In the experiment with UV and the experiment with gamma there was more sample in the third bottle than in the second but also more in the first than in the second, indicating that two processes were taking place.

We conclude that while the liquid phase oxidation process takes place in all experiments, the amount of sulphate produced via this process should be small in the first bottle, compared to the amount produced in experiments that had other formation processes present. The effect of the aqueous oxidation process on δ³⁴S in the first bottle will thus be small since the amount produced by the gas phase processes are larger (see Tab. 1 for a comparison of the sample masses recovered). Additionally the concentration of ozone is substantially smaller (about half) in the UV experiments, due to photolysis.

3.2 δ³⁴S

The δ³⁴S values of our sample O₃-UV-1 (8.7 ± 0.4 ‰) can be compared to the recently published values by Harris et al. (2012) for the OH reaction. In the Harris et al. study and this work, UV photolysis was used to generate OH via ozone photolysis, in the first case prior to the reaction chamber but in the second case within the reaction chamber – see Sect. 3.3. Their ³⁴α value of 1.0077 ± 0.0022 at 30°C is easily converted to our δ³⁴S notation by subtracting 1, giving 7.7 ± 2.2 ‰, which appears to agree with our result. The δ³⁴S of the OH reaction has also been estimated by Leung et al. (2001), using RRKM transition-state theory and by Tanaka et al. (1994) using ab initio calculations. They do, however, arrive at very different values: 140 and ∼9.0 ‰, respectively. While the Leung et al. result shows enrichment in δ³⁴S, as in this study, the effect is much larger than what we measure.

Harris et al. (2012) also measured liquid phase oxidation, but it is not obvious that we can make a direct comparison to our experiments without UV and gamma radiation. It is clear that not all sulphuric acid is collected in the first bottle in the line as we get more material in the second and third bottles and thus it is not possible to distinguish the fractionation for the liquid phase process. However we note that the δ³⁴S value for our sample O₃-1 (16.0 ± 0.4 ‰) agrees with the value at 22°C from the Harris et al. (2012) study of 14.8 ± 2.7 ‰. The Harris et al. (2012) results are also shown in Fig. 2. Note that Harris et al. (2012) find a Δ³³S value of ∼8 ‰ for the SO₂ + O₃ reaction, but this “may be a measurement artefact as only two samples were measured” (Harris et al., 2012). Also, their reported uncertainties for the Δ³³S values of the SO₂ + OH reaction are quite large due to counting statistics. Therefore we report their data in Fig. 2 with errors on the Δ³³S scale exceeding the depicted y-axis.
The samples O₃-1 and O₃-3 show a difference in δ³⁴S which can be explained by the isotopic composition of the substrate of the bottles which differs when operated in series, i.e. the first bottle leaves depleted substrate to be trapped in later bottles, so that δ³⁴S values are shifted to lower numbers.

The isotopic mixing line in Fig. 2 shows the basic fractionation present in all samples from the ozone process towards the fractionation for the other processes. Following the mixing line we note that the values of both δ³⁴S and Δ³³S of sample O₃-UV-γ-1 lie between the values for samples O₃-UV-1 and O₃-γ-1, indicating that in this case both processes are happening at the same time. Assuming that sample O₃-UV-γ-1 is a combination of the two processes, we find, using a linear combination of the δ³⁴S values, that 79% of sample O₃-UV-γ-1 originates from the UV process and 21% from the γ process, ignoring ozone oxidation.

3.3 Non-mass dependent fractionation (Δ³³S)

NMD was observed in our samples, especially for the UV experiment (O₃-UV-1). This stands in contrast to the mass dependent signal reported by Harris et al. (2012), however their sensitivity to δ³³S is not very high and their experiment was not directly irradiated with UV. NMD signals in SO₂ mainly arise from differences in the absorption spectra of isotopologues (Danielache et al., 2008) and in this experiment the reaction volume, which contains a high SO₂ concentration (0.01%), is directly exposed to light at 253.65 nm, which was not the case in the Harris et al. experiment. These conditions make a host of new reactions possible (Ueno et al., 2009), the most important ones being

\[
\text{SO}_2 + h\nu \rightarrow \text{SO}_2^* \quad \text{(R1)}
\]

\[
\text{SO}_2^* + \text{O}_2 \rightarrow \text{SO}_3 + \text{O}, \quad k=1 \times 10^{-16} \text{ cm}^3 \text{ s}^{-1} \quad \text{(R2)}
\]

\[
\text{SO}_2^* + \text{SO}_2 \rightarrow \text{SO}_3 + \text{SO}, \quad k=4 \times 10^{-12} \text{ cm}^3 \text{ s}^{-1} \quad \text{(R3)}
\]

\[
\text{SO}_2 \rightarrow \text{SO}_2, \quad k=2.2 \times 10^4 \text{ s}^{-1} \quad \text{(R4)}
\]

\[
\text{SO}_2^* + \text{M} \rightarrow \text{SO}_2 + \text{M}, \quad k=1 \times 10^{-11} \text{ cm}^3 \text{ s}^{-1} \quad \text{(R5)}
\]

Note that SO₂* can denote both a singlet and triplet state, where the singlet can be relaxed into the triplet state. The shown reaction rates are for the singlet state but similar relations occur for the triplet. The absorption cross section for SO₂ at 253.65 nm is \(\sim 1 \times 10^{-19} \text{ cm}^2\) (Danielache et al., 2008), and the concentration of SO₂ is 0.01%. The ozone cross section is \(\sim 1 \times 10^{-17} \text{ cm}^2\) at 253.65 nm and we have \(\sim 400 \text{ ppb} \text{ ozone in the reaction volume. At 40–45 \% RH, 0.2 of each O(\text{I}) from Eq. (R7) (Appendix C) becomes OH (Seinfeld and Pandis, 2006, chap. 6) - the remaining O(\text{I}) is quenched back to the ground state O(\text{I})P. The photon flux and pathlength for the light is the same in both cases so relative to each other we have 1 \times 10^{-19} \text{ cm}^2 \times 1 \times 10^{-4} = 1 \times 10^{-23} \text{ cm}^2\) for the SO₂ excitation pathway and 1 \times 10^{-17} \text{ cm}^2 \times 4 \times 10^{-7} \times 0.2 = 8 \times 10^{-25} \text{ cm}^2\) for ozone. This slightly favours the excitation pathway, starting with Reaction (R1). However, the excited SO₂* can easily be relaxed back to the ground state before it is oxidised to SO₃. With 0.01% SO₂ Reaction (R2) is the quickest pathway to SO₃ of Reactions (R2) and (R3), but the quenching reaction, Reaction (R5), is faster still, meaning that most of the excited SO₂ should be relaxed back to the ground state before reacting further, even at these highly elevated concentrations of SO₂.

Using the high resolution spectra obtained by Danielache et al. (2008) we can make an estimate of the expected NMD signal due to photo-excitation. Using the cross section at 253.65 nm, we predict a Δ³³S of 18.3 to 24.7% for SO₂* compared to SO₂ depending on the linewidth of the excitation source. Using more recent data (Danielache et al., 2012) we get 13.5–36.0%e. Since we only have a small conversion of our SO₂ reservoir this can be compared directly to our measured Δ³³S of 0.30%e. This shows that the measured NMD can be explained by photo-excitation. Our δ³⁴S result is thus obtained from a combination of the OH reaction and the UV excitation reaction. However, since our measured Δ³³S is so small compared to the value expected from the excitation pathway most of the sulphuric acid production in the UV experiments most likely proceeds by the OH pathway and thus justifies comparison with the results of Harris et al. (2012). Furthermore the lack of NMD in sample O₃-UV-3 is due to the liquid phase oxidation dominating.

A different study by Farquhar et al. (2001) focused on NMD signals in SO₂ photolysis reactions and described systems with different light sources. Comparing our data to theirs shows the same trends (our lamp compares best, but not perfectly, to their KrF excimer laser), but one has to keep in mind that minute changes in the irradiance...
may shift isotopic effects drastically (Danielache et al., 2008). Nevertheless, the $\Delta^{38}S$ values reported here (see Appendix A) are pointing towards the negative, while the $\Delta^{32}S$ values show a positive trend, just as in the Farquhar et al. (2001) study.

A small non-zero $\Delta^{33}S$ is seen in samples O$_2$-3, O$_2$-UV-3, and O$_2$-$\gamma$-1, while sample O$_2$-1 has a stronger signal. They are all significant with respect to the error, but we do not have a full understanding of the underlying process. In order to understand this effect a model of the heterogeneous reaction could be constructed similar to the sulphate reduction model of Rees (1973) (see also Johnston, 2011).

### 3.4 Gamma experiments

Ionising radiation gives energy to the electron clouds in molecules often generating ion pairs:

$$M + \gamma \rightarrow M^+ + e^- \quad (R6)$$

In our experiment, as well as the atmosphere, M is typically N$_2$. The resulting electrons eventually attach to molecules, depending on their electron affinity and concentrations. Oxygen and nitrogen exist in the greatest abundance and since oxygen has a positive electron affinity the O$_2$ ion is spontaneously formed. The core ion then attracts water to form hydrated clusters, such as O$_2$(H$_2$O)$_n$ – this happens even at otherwise unsaturated conditions, due to the electric charge. There are many options for further reaction – one relevant scheme is presented by Bork et al. (2011a,b). In the troposphere ionisation is mostly caused by secondary particles (muons and electrons) formed by galactic cosmic rays with energies >1 GeV (Bazilevskaya et al., 2008). Although the energy of the gamma source (662 keV) is somewhat lower than that of cosmic rays penetrating the lower atmosphere the ions produced are the same. Based on the difference in $\delta^{34}S$, the mechanism for sulphate formation in the gamma reaction is clearly different from the OH oxidation pathway. The amount produced is also too high to be explained by this mechanism. The estimated OH production per ionisation is about 2 (Müller and Crutzen, 1993) and the ionisation in our chamber is estimated to be 1000 cm$^{-3}$ s$^{-1}$ (based on a simple model using the activity of the source and distance from the chamber to calculate the energy deposited in the volume), resulting in $6 \times 10^{13}$ OH molecules formed during a run of one week, which would give 23 ng of BaSO$_4$, orders of magnitude less than the 19.4 mg collected, even if a small part of this comes from the liquid phase oxidation process. The gamma induced fractionation pattern resembles that of the experiments without UV and gamma radiation more even if they are some standard deviations apart, but in the gamma experiment we see much more material in the first bottle than in the following, the reverse pattern of what is seen in the experiments without UV and gamma radiation. This suggests that the mechanism of the gamma pathway is gas-phase or heterogenous chemistry in the chamber and by a mechanism that somehow enhances the yield far beyond one sulphate per ionisation. That we see a small NMD effect also suggests a chemical pathway. One possible explanation is the mechanism under investigation in Bork et al. (2011a,b) where hydrated ion clusters (Zatula et al., 2011), such as described above, serve as nanoreactors for sulphur oxidation by a process similar to the oxidation of SO$_2$ by ozone observed in the experiments without UV and gamma radiation.

These experiments clearly suggest that ions may catalyse SO$_2$ oxidation. However, any ion catalysed mechanism will terminate with collision with OH or another radical since most radicals have high electron affinities, i.e. can form stable anions. In other words, the electron becomes chemically immobilized. In this experiment the dominant electron scavenger is OH. The contribution is thus dependent on the ratio of SO$_2$ to OH and the ion production rate and we note that in these experiments, the relative concentration of SO$_2$ to OH was much larger than in the atmosphere. The ion induced contribution should be compared to the oxidation of SO$_2$ by OH. The third-order rate constant for this reaction is about $4.5 \times 10^{-31}$ cm$^6$ s$^{-1}$ (Seinfeld and Pandis, 2006, Appendix B). The gamma-induced pathway scales with $P_{ion}$[SO$_2$]/[OH], where $P_{ion}$ is the ion production rate. The UV pathway scales with [SO$_2$]·[OH]·$M$·$k$, where $k$ is the rate constant. Since both pathways scale with [SO$_2$] the relative contribution of the ion mechanism to the UV mechanism does not depend on [SO$_2$]. On the other hand it depends on [OH]$^2$ and on the ion production rate. Thus the relative contribution can vary greatly and will be larger at night when there is little or no OH available. For a daytime [OH] of e.g. $2 \times 10^6$ cm$^{-3}$ and an ion production rate of 4 cm$^{-3}$ s$^{-1}$ the relative contribution will be 9%. This is of course a very rough estimate and a definitive answer will require further research, such as has been started by Bork et al. (2011b).

### 4 Conclusions

In this study we investigated the production of sulphuric acid from SO$_2$ through a number of different pathways. The sulphuric acid was produced using standard gas phase oxidation by OH radicals from ozone photolysis, liquid phase oxidation by ozone, and oxidation initiated by gamma rays. The distributions of stable sulphur isotopes in the products and substrate were measured using isotope ratio mass spectrometry. All methods produced sulphate enriched in $^{34}$S and we find a $\delta^{34}S$ value of 8.7 $\pm$ 0.4‰ (1 standard deviation) for the UV-initiated OH reaction. Only the experiment with UV light (Hg emission at 253.65 nm) produced a clear non-mass-dependent excess of $^{33}$S of around 0.3‰, resulting from a small contribution from photo-excitation. Our results for UV photolysis and possibly aqueous oxidation by ozone, matched isotope patterns seen in earlier measurements. However, these UV photolysis result do not agree with theoretical
studies (Tanaka et al., 1994; Leung et al., 2001), suggesting that additional theoretical work is needed. The pattern of isotopic enrichment produced by gamma rays was distinctly different from that caused by the UV light and similar, but not equal, to that produced by the aqueous oxidation of SO₂. This, combined with the relative yields of the experiments, suggests a mechanism by which ionizing radiation may lead to hydrated ion clusters that serve as nanoreactors for S(IV) to S(VI) conversion.

Appendix A

Δ³⁶S

In addition to δ³⁴S and Δ³³S we also measured Δ³⁶S, using an improved purification system for most of the samples. The values can be seen in Table A1. We refrain from going into a detailed analysis of this data but note that the linear relation between Δ³³S and Δ³⁶S is consistent with the isotope mixing shown in Fig. 2. Additionally, Table A1 displays further information on the amount of material used and recovered during the isotopic analysis.

Appendix B

Definitions

Sulphur isotopic compositions are reported as δ³⁴S and Δ³³/³⁶S values normalized against the mean value of our two SO₂ references:

\[ \delta^{34}S = \frac{i_{\text{Sample}}}{i_{\text{Reference}}} - 1 \]

where \( R \) denotes the isotope ratios (\(^{34}S / ^{32}S\)) of samples and reference.

\[ \Delta^{33}S = \delta^{33}S - \left( \delta^{34}S + 1 \right)^{0.515} - 1 \]
\[ \Delta^{36}S = \delta^{36}S - \left( \delta^{34}S + 1 \right)^{0.90} - 1 \]

Table A1. Isotopic composition of samples. The table displays the sample ID, Δ³⁶S (normalized against the references as defined in Appendix B), the mass of BaSO₄ used, the mass of Ag₂S recovered, the mass of Ag₂S used, and the amount of SF₆ recovered.

<table>
<thead>
<tr>
<th>Sample ID</th>
<th>Δ³⁶S %o</th>
<th>BaSO₄ Used mg</th>
<th>Ag₂S Rec mg</th>
<th>Ag₂S Used mg</th>
<th>SF₆ µmol</th>
</tr>
</thead>
<tbody>
<tr>
<td>O₁-1</td>
<td>0.5</td>
<td>0.689</td>
<td>0.689</td>
<td>2.85</td>
<td></td>
</tr>
<tr>
<td>O₂-3</td>
<td>0.4</td>
<td>8.99</td>
<td>8.84</td>
<td>2.07</td>
<td>7.91</td>
</tr>
<tr>
<td>O₃-UV-1</td>
<td>-1.3</td>
<td>-</td>
<td>-</td>
<td>2.0</td>
<td>6.98</td>
</tr>
<tr>
<td>O₄-UV-3</td>
<td>0.2</td>
<td>9.39</td>
<td>9.48</td>
<td>2.00</td>
<td>8.87</td>
</tr>
<tr>
<td>O₅-UV-γ-1</td>
<td>0.2</td>
<td>5.51</td>
<td>5.09</td>
<td>2.21</td>
<td>9.45</td>
</tr>
<tr>
<td>O₆-UV-γ-1</td>
<td>-1.0</td>
<td>7.90</td>
<td>7.34</td>
<td>2.13</td>
<td>9.80</td>
</tr>
</tbody>
</table>

* This sample was not measured with the improved purification system.

Alternative, Table A1 displays further information on the amount of material used and recovered during the isotopic analysis.

Appendix C

Gas-phase oxidation of SO₂ by OH

Here we list the reactions relevant to the oxidation of SO₂ by OH in the chamber.

\[ O_3 + hv \rightarrow O(¹D) + O_2 \]  
\[ O(¹D) + H_2O \rightarrow 2OH \]  
\[ SO_2 + OH + M \rightarrow HSO_3 + M \]  
\[ HSO_3 + O_2 \rightarrow SO_3 + HO_2 \]  
\[ SO_3 + H_2O \rightarrow H_2SO_4 \]

Supplementary material related to this article is available online at: http://www.atmos-chem-phys.net/12/5319/2012/acp-12-5319-2012-supplement.pdf.

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