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Isotopic archives of sulphate in speleothems

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Abstract

The hemispheric impact of industrial emissions upon atmospheric sulphur loading is reflected in the sulphur depositional history recorded in cores from ice sheets. However, these do not reveal regional variations. Recently deposited speleothems are used here as archives of regional sulphur depositional history at two locations within the United Kingdom and Ireland. δ^{34} S-so₄ and δ^{18} O-so₄ present within speleothem carbonate are measured for the first time as part of a dual isotope approach to decode the speleothem sulphur record. The largely refractory nature of δ^{34} S-so₄ under oxidising conditions enables source provenance of atmospheric SO₂, whereas the complex cycles of isotopic exchange and fractionation during incorporation of oxygen into sulphate molecules enable δ^{18} O-so₄ signatures to yield insights into ambient environmental conditions and biogeochemical cycling in the ecosystem above the cave. δ^{34} S-so₄ values extracted from speleothem carbonate formed within Browns Folly Mine, UK, range from +3.5 to +5.5‰ and δ^{18} O-so₄ +10.3 to +13.7‰. Both signatures lie within the range expected from sulphate deposition in industrial locations and reflect the transfer of sulphate into speleothem calcite with little fractionation. However, δ^{18} O-so₄ signatures at Crag Cave, western Ireland, are isotopically heavier than expected and approach isotopic equilibrium with δ^{18} O-so₄ on the cove sites where drip water flowpaths into the cave remain oxic, speleothems hold the potential to retain records of atmospheric sulphur loading at the local and regional scale. © 2008 Elsevier Ltd. All rights reserved.

1. INTRODUCTION

Anthropogenic emissions of SO_2 have risen since 1850 due to increased industrial activity (Lefohn et al., 1999), rendering atmospheric sulphate aerosols key agents in forcing recent climate (e.g. Smith et al., 2001). Such increased emissions are documented in ice core records of sulphate deposition, which are currently considered the premier archive of past atmospheric sulphur concentrations (e.g., Patris et al., 2000, 2002; Isaksson et al., 2005). However, ice core localities are frequently distal from sources of

SO₂ emissions and reflect a relatively clean atmospheric boundary layer. Due to the relatively short lifetime of sulphate aerosols in the atmosphere, the impact of anthropogenic sulphur emissions on the climate and biogeochemistry of sulphur cycling is most acute at local and regional scales. Historical quantification of sulphur inputs to terrestrial environments has been demonstrated through sulphur isotope ratios in archived soil and herbage samples from the UK (Zhao et al., 1998), from wood of coniferous trees in Japan (Kawamura et al., 2006) and from lake and peat sediment cores in Europe (Bottrell and Coulson, 2003; Coulson et al., 2005; Novák et al., 2005) and Canada (Mayer et al., 2007).

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Speleothems can yield multi-proxy, high resolution records of the palaeoenvironment, and are formed through

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the carbonation of host carbonate bedrocks and re-precipitation of calcite within cave environments (see review by Fairchild et al., 2006a). The incorporation of trace elements, organic molecules and specific stable isotope signatures in speleothem carbonate provides a suite of geochemical parameters capable of capturing a cave's response to the external environment. Thus, trace amounts of sulphur discovered as sulphate in speleothems suggest that stalagmites may record key aspects of atmospheric variability in sulphate content at local and regional scales (Frisia et al., 2005; Fairchild et al., 2006a). Sulphate is inferred to be present in speleothem calcite as Carbonate Associated Sulphate (CAS) identified in the geological record of marine carbonates (e.g., Bottrell and Newton, 2006). XANES analysis has been used in both marine (Pingitore et al., 1995) and speleothem carbonate (Frisia et al., 2005) to identify the oxidation state of the constituent sulphur as sulphate, implying its presence within the calcite lattice as a structural substitution for carbonate. However, sulphur concentration data alone cannot directly identify the source of speleothem CAS (Wadleigh et al., 1996; Frisia et al., 2005). Atmospheric variability in sulphate content is controlled by natural as well as anthropogenic sources of sulphur and may be modified by biogeochemical cycling and additional sources of sulphur stored within the surrounding bedrock and soil waters (Fig. 1). The sulphate content of cave drip waters and associated stalagmites may thus represent a mixed signal, reflecting not only the local variability in atmospheric sulphate content, but also biogeochemical cycling in the ecosystem above the cave. Sulphur and oxygen isotope systematics of sulphate differ markedly, and in principle should enable the provenance of sulphate within speleothems to be established permitting further deconvolution of the mixed atmospheric/biogeochemical signal archived within speleothems.

2. THEORETICAL BACKGROUND

Fig. 1 summarises the potential sources of sulphate contributing to speleothem CAS with characteristic $\delta^{34}S$ compositions. Primary sea-salt aerosols released into the atmosphere from sea spray are derived from the well-mixed reservoir of marine sulphate and have a $\delta^{34}S$ value identical to that of sea water (~+21‰). Biogenic sources of secondary sulphate aerosol are derived from marine emissions of dimethylsulphide (DMS) and are generally depleted in ³⁴S relative to seawater sulphate, with reported $\delta^{34}S$ values of +15.6 ± 3.1‰ (Calhoun et al., 1991). Continental emissions of sulphate aerosol can be divided into anthropogenic and biogenic emissions. In the northern hemisphere, sulphate in precipitation sourced predominantly from anthropogenic emissions is usually assigned an average $\delta^{34}S$ value between



Fig. 1. Sources and cycling of sulphur contributing to bulk stalagmite sulphate composition with characteristic $\delta^{34}S$ compositions. Values are given for predominantly anthropogenic sulphate in rain in industrialised countries of the northern hemisphere (range -3 to +9%; Mayer, 1998); sulphate aerosol from ocean surface (+21%; Rees et al., 1978); marine biogenic emissions as DMS ($+15.6 \pm 3.1\%$; Calhoun et al., 1991); continental and intertidal biogenic emissions (0 to -30%; Nielson, 1974); volcanic eruptions ($\sim0\%$; Nielson, 1974).

-3 and +9% (Mayer, 1998), despite the δ^{34} S composition of the source materials varying beyond this range (see compilation of source material values in Nielson, 1974). Continental biogenic emissions of sulphur consist predominantly of gaseous organic sulphides from the decomposition of organic matter associated with plants and soils. Such emissions have a wide range of δ^{34} S values depending upon the redox status under which they are formed. Under oxidising conditions, their isotopic composition should be similar to that of local precursor sulphate derived from precipitation, groundwater and lithogenic sources. This is quoted to have an average δ^{34} S value close to 0% (Nielson, 1974), although reducing conditions induce extensive fractionation to the source compounds. This fractionation varies with environmental conditions although when the size of the substrate pool is unlimited, gaseous organic sulphides may be fractionated so they are up to 70% lighter than the precursor sulphate (Brunner and Bernasconi, 2005).

Due to the widely differing source composition of oxygen isotopes involved in the oxidation of sulphur compounds to sulphate, the oxygen isotopic composition of sulphate in speleothems should enable the distinction between sulphate which is sourced directly from the atmosphere and that produced by nutrient cycling in terrestrial ecosystems (e.g.: Mayer et al., 1995a.b; Likens et al., 2002), thus aiding process-identification as well as source provenance. Published values for the $\delta^{18}O$ composition of atmospheric sulphate range from -3 to +41.7% V-SMOW (see compilation of published values in Jamieson and Wadleigh, 1999; Jenkins and Bao, 2006). This wide range reflects the different reaction pathways of gaseous sulphide oxidation in the atmosphere, the oxygen isotopic composition of the oxidants involved and the mixing ratio between aerosols formed at source as sulphate (primary aerosols) and those produced in the atmosphere from gaseous precursors (secondary aerosols) (see review by Holt and Kumar, 1991). Secondary aerosols formed from gaseous precursors do not carry oxygen isotopic signatures indicative of source provenance, but reflect only the pathways of sulphide oxidation in the atmosphere due to a rapid isotopic equilibration between reduced sulphoxy intermediate species and H₂O. Source provenance signatures of δ^{18} O in reduced sulphur compounds emitted into the atmosphere are thus erased during oxidation to secondary aerosol species. Primary sulphate aerosols retain their source δ^{18} O-so₄ composition during transport through the atmosphere. These can be sourced from sea spray (+9.7%; Lloyd, 1967) or from industrial activity with δ^{18} O-so₄ signatures up to +45% (Holt et al., 1982; Jamieson and Wadleigh, 1999). The proportion of primary aerosol present in the atmosphere thus significantly influences the value of bulk δ^{18} O-so₄ aerosol deposition.

Subsequent to sulphate aerosol deposition, biogeochemical cycling resets the oxygen isotopic composition of atmospheric-derived sulphate according to environmental conditions. Where aerobic oxidising conditions prevail in the overlying cave soils, biogeochemical cycling of deposited sulphate may induce assimilatory sulphate reduction to organic-S compounds and mineralisation back into the inorganic phase. This causes only minimal fractionation to ${}^{34}S/{}^{32}S$ ratios compared to precursor sulphate derived from precipitation, groundwater and lithogenic sources. Sulphate ${}^{18}O/{}^{16}O$ ratios however, are a product of incorporating four oxygen atoms obtained in varying proportions from the surrounding soil water and atmospheric oxygen (Eq. (1)).

$$\delta^{18}\text{O-so}_4 = [(\delta^{18}\text{O}_{\text{atm O}_2} - \epsilon \text{ so}_4 - \text{atm}) \times f\text{O}_2 \text{ atm}] + (\delta^{18}\text{O}_{\text{water}} \times 1 - f\text{O}_2 \text{ atm})$$
(1)

where $fO_{2 \text{ atm}}$ and $1 - fO_{2 \text{ atm}}$ represent the fraction of oxygen obtained from the atmosphere and soil water, respectively, and $\delta^{18}O_{\text{atm} O_2}$ and $\delta^{18}O_{\text{water}}$ represent the oxygen isotopic compositions of atmospheric O₂ (+23.88‰; Barkan and Luz, 2005) and water. ε so₄-atm is the isotopic enrichment associated with the incorporation of atmospheric O₂ into sulphate, demonstrated experimentally to have a value of -8.7% (Lloyd, 1968).

Sulphoxy intermediate compounds (SO, SO₂, HSO₃⁻) can be exposed to rapid isotopic equilibration with the surrounding water. The isotopic composition of the constituent oxygen can thus be reset such that up to 75% of the oxygen in the final sulphate molecule has an isotopic composition controlled by meteoric water (Holt et al., 1981). Where mineralisation of organic compounds to sulphate is sufficiently rapid such that sulphoxy intermediate species do not become fully equilibrated with meteoric water, the final sulphate molecule may contain a greater proportion of oxygen sourced from atmospheric O_2 (Bottrell, 2007). The δ^{18} O-cas contained within speleothem calcite is thus sensitive to the relative proportions of atmospheric O_2 and water oxygen incorporated into the sulphate molecule. Isotopic signatures may therefore have a theoretical upper end-member composition of +14.8% where all four oxygen atoms are derived from atmospheric O₂ (Bottrell and Newton, 2006) and range towards the oxygen isotopic composition of the ambient water where equilibration with meteoric water imparts up to 75% of the isotopic composition. Where sulphur is contained within organic matter as ester sulphate compounds and liberated through enzyme hydrolysis (e.g. Fitzgerald, 1976), at least three of the oxygen atoms in the newly mineralised sulphate are sourced directly from the organically bound sulphate molecule. As such, a depletion of only 2-3% is observed in oxygen isotope composition compared to the original sulphate ester compound (Mayer et al., 1995b).

Under reducing conditions, processes of microbial sulphate reduction may cause extensive fractionation of δ^{34} S. Fractionation of δ^{18} O-so₄ however, is manifest as an equilibrium isotopic exchange between the ambient water and the residual sulphate pool such that an enrichment factor of +29‰ at 5 °C may be apparent if equilibrium is approached (Fritz et al., 1989; Brunner et al., 2005; Wortmann et al., 2007). During initial stages of sulphate reduction under closed system conditions, δ^{34} S-so₄ and δ^{18} O-so₄ signatures demonstrate a linear fractionation relationship, typical of a kinetic reaction mechanism. When sulphate reduction reaches an advanced state however, residual δ^{18} O-so₄ signatures approach a constant equilibrium value and δ^{34} S-so₄ residual signatures continue to become enriched in ³⁴S according to a typical Rayleigh fractionation trajectory (Brunner et al., 2005). Where cyclical reduction–re-oxidation of sulphate occurs within the same environment, values of δ^{18} O-so₄ may display an offset between the equilibrium oxygen isotope signature and the measured values. Re-oxidation of sulphite to sulphate involves the incorporation of one additional oxygen from an external source such as water (enrichment factor = +4‰; cf.Taylor et al., 1984). The resulting oxygen isotope composition in the residual sulphate can be approximated as;

$$\delta^{18}\text{O-so}_{4} = [0.75 \times (\delta^{18}\text{O-H}_{2}\text{O} + \varepsilon \text{ so}_{4}\text{-H}_{2}\text{O})] + 0.25 \times (\delta^{18}\text{O-H}_{2}\text{O} + 4^{\circ}_{\infty})]$$
(2)

where ε so₄-H₂O is the equilibrium isotopic enrichment between δ^{18} O in the residual sulphate pool and that in the surrounding water (after Brunner et al., 2005). The influence of re-oxidation upon ³⁴S values however will be to retain a signature similar to the initial products if isotope mass balance is maintained (e.g., Barker et al., 1998). The extraction and analysis of δ^{34} S and δ^{18} O from speleothem CAS thus holds the potential to record the sulphur sources and biogeochemical transformations present along the flow pathways of speleothem-feeding drips.

Traditional methods used for the extraction and analysis of SO₄-S and SO₄-O from carbonates rely on the acid digestion of large quantities of material (typically 100-300 g) and the extraction of sulphate as barium sulphate (e.g., Gellatly and Lyons, 2005). The large volumes of carbonate material required have thus frequently constrained the type of samples analysed for sulphate isotopic composition. Where geological carbonates are present in abundance or contain significant quantities of sulphur, such constraints are not applicable. However, in speleothem carbonate, sulphur content is low and the material available for analysis is necessarily limited. New methods to reduce the sample size of carbonate material required for the extraction and analysis of δ^{34} S-so₄ and δ^{18} O-so₄ have been developed as a part of this study, thus enabling the sulphur isotopic characterisation of speleothem archives. Here we report the first determinations of ${}^{34}\text{S}/{}^{32}\text{S}$ and ${}^{18}\text{O}/{}^{16}\text{O}$ ratios of CAS extracted from speleothems. The samples originate from two cave sites in contrasting locations, demonstrating the power of the dual isotope approach for palaeoenvironmental interpretation. We compare modern time series of a stalagmite collected from within Browns Folly Mine, UK, where values of δ^{34} S-cas illustrate a dominant control by anthropogenic SO₂ emissions, with that of a stalagmite from Crag Cave, S.W. Ireland, yielding δ^{34} S-cas and δ^{18} O-cas signatures representative of sulphate reduction and re-oxidation in the soils above the cave.

3. SITE AND SAMPLE DESCRIPTION

Browns Folly Mine, Bathford, SW England, was developed in the 19th century for the extraction of building stone from the surrounding limestone (the Bath Oolite member of the Jurassic Great Oolite series) (for cross section of mine site see Baldini et al., 2005; Fairchild et al., 2006b). Mining activity commenced in 1836 and continued until the mines were abandoned in 1886, thus constraining the maximum possible age of the associated speleothems. The mines remained closed for approximately 100 years until re-opened by cavers in the 1970s (Baker et al., 1998, 1999). Lithologies of the overlying Middle Jurassic bedrock are mainly oolitic and skeletal limestones, containing a minor sand and clay component and a porosity of 0-40% (Fairchild et al., 2006b), with an absence of pyrite and other sulphide minerals. Vertical fractures dissect the horizontally stratified limestone formations providing a route for rapid fissure flow of groundwater into the mine system. Slower seepage flow is fed by minor fissures and water stored within the porous limestone matrix (Baker et al., 1999; Fairchild et al., 2006b). Vegetation has gradually re-established itself over the study site since the cessation of mining operations in 1886 such that secondary deciduous forest of mixed ash, sycamore and oak is now prevalent (Baldini et al., 2005). The sampled stalagmite (BFM-Boss) was extracted in 1996 from a chamber approximately 300 m from the mine entrance (Baldini et al., 2005) and represents a continuous deposition of calcite in annual couplets of clear and inclusion-rich laminae. Three independent lamina thickness count established 77 ± 3 annual couplets at the central growth axis suggesting that BFM-Boss nucleated no later than 1916. Annual calcite deposition rates remained low until 1975, when couplet thicknesses increase dramatically, indicating an increase in growth rate. The nucleation of BFM-Boss approximately 30 years after the closure of the mines is thought to reflect either the time taken for the development of a stable hydrological system subsequent to the cessation of mining activity, or the slow rate of vegetation establishment above the cave site. Baldini et al. (2005) showed that vegetation development had a strong influence on the δ^{13} C and δ^{18} O record of this sample.

Crag Cave, located in County Kerry, Ireland, is developed in Lower Carboniferous limestone some 20 km from the Atlantic coast of SW Ireland (McDermott et al., 1999) (for site details see Tooth and Fairchild, 2003; Baldini et al., 2006). Discovered in 1981, parts of the relatively shallow cave system (c. 20 m deep) were opened as a tourist cave in 1985, although restricted access to the cave interior has ensured the maintenance of near-pristine conditions, rendering it ideal for hydrochemical studies (Baldini et al., 2006). The surrounding bedrock, belonging to the Lower Carboniferous Cloonagh Formation is dominantly limestone, with small amounts of dolomite and some pyrite (Tooth and Fairchild, 2003). Till deposits overlying the cave are of variable thickness (0.1-2.5 m thickness) belonging to the Munsterian (Marine Isotope Stage 6) glaciation. The low permeability of the fine grained clay matrix encourages saturation and a constant supply of water to the stalagmite-feeding drips, and streaks of iron oxide staining demonstrate the presence of oxidising conditions in isolated zones and hence the existence of redox cycling. The presence of a crack and fissure network allows the till deposits to be regarded as a dual-porosity system, facilitating both rapid water throughput via fracture flow and a more constant matrix seepage flow supplied from the large storage capacity of the micropores (Tooth and Fairchild, 2003). The sampled stalagmite, CC-Bil, was removed from a passage 30 m below the ground surface in 2002. The drip that fed stalagmite CC-Bil formed on the tip of a 1 m long soda straw stalactite and is classed as a 'seepage flow site' under the classification of Smart and Friederich (1986). Drip rates at this site and others in close proximity are low (site CC-Bil = 0.08 ml/min) and show very little variation through time (site CC-Bil drip rate CV = 5.4%) (Tooth and Fairchild, 2003; Baldini et al., 2006). Based on visible calcite deposition on loggers underneath the drip feeding stalagmite CC-Bil, this stalagmite was actively growing when collected in 2002. U-series dating demonstrates a basal mean ICP-MS U-Th age of 1752 AD (\pm 7 years) and an approximately linear growth rate over the past 260 years.

4. METHODOLOGY

4.1. Sulphate extraction

Sulphur concentrations in each of the stalagmites were determined by high resolution inductively coupled plasma mass spectrometric analysis (HR-ICPMS) of calcite powders at Kingston University, UK, drilled in 2 mg aliquots from the central axis of the stalagmite and digested in 8 ml of 2% HNO₃ (Aristar grade). Depending upon the concentration of CAS, larger aliquots of calcite ($\sim 200 \text{ mg}$) were prepared for sulphur and oxygen isotope analysis by sampling in a continuous fashion down the stalagmite central growth axis, each sample aggregating several years of growth. Using the concentration data obtained through HR-ICPMS analysis, the size of each carbonate sample was calculated to yield a minimum of $35 \,\mu g$ of sulphur, representing the quantity of sulphur required for analysis by CF-IRMS (continuous flow isotope ratio mass spectrometry) given the instrumental configuration outlined below. Parallel sampling tracks enabled the integrity of δ^{34} S and δ^{18} O analyses to be assessed through sample replication. The methodological approach to sulphur and oxygen isotope extraction was based upon procedures reported in Newton et al. (2004), albeit simplified as the relative purity of stalagmite carbonate precluded the need for pre-treatment with sodium hypochlorite and bromine water to remove organic contaminants and trace amounts of pyrite, respectively. Given the small quantities of sulphur extracted per sample, the following protocol was designed to minimise sample handling and contamination in all steps of the procedure.

Drilled stalagmite powders were digested overnight in 1 ml of 4M hydrochloric acid (Aristar grade) within polyethylene syringes, capped with Luer tips. Syringe filtration through pre-rinsed 0.2 µm nylon filter capsules (Puradisc, Whatman[®]) enabled sample transfer between vessels with the minimum of product loss. The extraction of sulphate as barium sulphate was undertaken following precipitation using barium chloride solution. Two different techniques of precipitate concentration were used to collect the barium sulphate from solution: filtration and centrifugation. Where filtration was the method employed, 0.5 ml of 1 M barium chloride solution was added to the filtered carbonate digest solution within the housing of a second polyethylene syringe. Samples were left to precipitate and mature for 2-3 days, prior to syringe filtration through quartz microfibre filters using an in-line filter capsule (Swinnex, millipore) modified to hold filter papers 7 mm in diameter. Each filter loaded with 35 μ g of S (~250 μ g BaSO₄) was transferred into a tin capsule, dried at 70 °C and stored in a desiccator prior to mass spectrometric analysis (Fig. 2).

However, based upon the decomposition of quartz above 700 °C and the release of constituent oxygen, the use of quartz microfibre filters as a means of pre-concentrating barium sulphate precluded analysis for δ^{18} O-so₄ by this method. Centrifugation in association with an inert medium was thus employed as the method of pre-concentration for the remainder of the study. Carbonate digest solutions were syringe-filtered into 1.5 ml centrifuge vials and precipitated as barium sulphate following addition of 0.2 ml of 1 M barium chloride solution. Contact time between free sulphate ions in solution with strong acid was minimised to \sim 1 h to prevent potential oxygen isotopic exchange between sulphate and water under acidic conditions (e.g., Hurtgen et al., 2002; Newton et al., 2004; Gellatly and Lyons, 2005). Addition of quartz powder ($\sim 2 \text{ mg}$; EuroVector isotopic grade) served as an inert medium onto which barium sulphate could precipitate, thus increasing the mass of product and enabling ease of handling. Sulphate for δ^{18} O analysis was precipitated onto powdered glassy carbon (~0.5 mg, Euro Vector isotopic grade) as the inert medium. Samples of barium sulphate were left to mature for \sim 72 h prior to repeated centrifuging and washing with de-ionised water to remove excess barium. The resultant pellets of quartz powder with barium sulphate and glassy carbon with barium sulphate were oven dried at 70 °C and transferred to tin or silver capsules for δ^{34} S and δ^{18} O analysis, respectively, (Fig. 2).

To overcome possible problems associated with oxygen release during the decomposition of quartz powder contributing to SO_2 beam intensities and thus influencing $\delta^{34}S$ measurements (e.g. Fry et al., 2002), similar quantities of quartz powder were added to standard materials as well as samples prior to combustion. This effectively supplied a large reservoir of oxygen in the form of quartz powder, buffering the oxygen composition of the SO_2 gas for both samples and standards.

4.2. Mass spectrometric analysis

³⁴S/³²S and ¹⁸O/¹⁶O ratios of product barium sulphate were determined using a EuroVector elemental analyser linked to a GV Isoprime continuous flow mass spectrometer at the University of Birmingham. Combustion of samples within tin capsules in the presence of vanadium pentoxide at 1030 °C vielded SO₂ for determination of δ^{34} S-so₄ and pyrolysis within silver capsules in the presence of nickelised carbon at 1250 °C yielded CO for determination of δ^{18} O-so₄. δ^{34} S values were corrected against CDT using within-run analyses of international standard NBS-127 and SO5 (assuming δ^{34} S values of +20.3% and +0.5%, respectively, (IAEA, 2004)) and δ^{18} O was corrected to V-SMOW using NBS-127 and SO6 (assuming δ^{18} O values of +9.3% and -11.3%, respectively, (IAEA, 2004)). Within-run standard replication (1 SD) was <0.3% for both sulphur and oxygen. Sample replication from two parallel sample tracks drilled in stalagmite BFM-Boss were $\leq 0.4\%$ (1 SD) for δ^{34} S values and <0.5% for δ^{18} O values based on sample replication in stalagmite CC-Bil.

Procedural standard solutions of calcium sulphate and sodium sulphate were used to test the integrity of the methodology for precipitation and analysis of small quantities of barium sulphate for δ^{34} S and δ^{18} O determinations, respectively. Calcium sulphate solution precipitated as barium sulphate under acidified conditions and analysed according to the above methodology using quartz microfibre filters, yielded δ^{34} S-so₄ signatures of +3.2% (0.3% 1 SD, n = 20) compared to δ^{34} S values of +2.9% (0.3% 1 SD, n = 13) for analysis of raw calcium sulphate powder. Where centrifuging with quartz powder was used as the method of pre-concentration, analyses of standard calcium sulphate solution precipitated as barium sulphate, yielded δ^{34} S-so₄ isotopic values of +2.6% (0.3% 1 SD, n = 12). Sodium sulphate solution precipitated as a bulk quantity of barium sulphate yielded δ^{18} O-so₄ isotopic signatures of +12.6% (0.4% 1 SD, n = 8) compared to analyses obtained fol-

Fig. 2. Diagram showing the two approaches (centrifugation and filtration) to extracting and concentrating sulphate from speleothem calcite as barium sulphate prior to δ^{34} S and δ^{18} O analysis.

lowing the centrifuging method of pre-concentration under analogous acidified conditions, of +13.0% (0.7% 1 SD, n = 6). Blank contamination associated with δ^{34} S determinations were

Blank contamination associated with δ^{34} S determinations were zero. A small blank signal associated with oxygen analysis is most likely associated with trace levels of oxides on the glassy carbon and nickelised carbon catalyst, but was too small to determine accurately. To overcome the potential impact of blank contribution upon reported δ^{18} O values, standard materials and samples were analysed using the same quantities of glassy carbon powder and nickelised carbon catalyst.

5. RESULTS

5.1. BFM-Boss

Fig. 3 displays sulphur concentrations and δ^{34} S and δ^{18} O values of carbonate associated sulphate in stalagmite BFM-Boss. Sulphur concentrations are between 150 and 250 ppm (Table 1) and are coincidentally similar to those of the surrounding bedrock (range 156–312 ppm) (Table 2). The δ^{34} S-cAs values associated with BFM-Boss are between +3.5 and +5.5‰ and significantly lower than those contained within the surrounding bedrock (+17.9 to +22.0‰). δ^{18} O-cAs signatures within the speleothem range between +10.3‰ and +13.7‰. Table 3 displays the ionic composition of the feeding dripwater to stalagmite BFM-

Boss. Sulphate concentrations displayed as a percentage sea-salt sulphate (% SSS) are calculated on the assumption that sea spray has a molar SO₄/Cl ratio of 0.0516 and is the source of all Cl⁻ in the drip waters (after Holland, 1978), such that 29% (1 SD = 15%) of all sulphate is calculated to be of inorganic marine provenance. Further, sulphate concentrations within drip waters comprise just 1.4% sulphate sourced from the bedrock based on Ca/SO₄ ratios given in Table 2 (Eq. (3)).

This assumes all calcium in the cave drip waters is sourced from dissolution of the karst bedrock, with negligible contribution from dissolution of calcareous dust particulates in the atmosphere, and assumes ratios are unaffected through chemical evolution associated with limited levels of prior calcite precipitation (Tooth and Fairchild, 2003).

5.2. CC-Bil

Sulphur concentrations within stalagmite CC-Bil are between 100 and 200 ppm and values of δ^{34} S-cas range between +16.8 and +19.1% and δ^{18} O-cas between +19.2

Fig. 3. Sulphur concentration, δ^{34} S-CAS and δ^{18} O-CAS in stalagmite BFM-Boss.

and +23.9‰ (Table 4, Fig. 4). Bedrock concentrations of sulphur are low and range between 51 and 92 ppm (Table 2), and isotopes of δ^{34} S and δ^{18} O within the lower Carboniferous limestone within which Crag Cave is formed are reported to range between +13 and +15‰ for δ^{34} S based on CAS-derived S isotopes and +14 to +19‰ for δ^{18} O based on marine evaporates (Bottrell and Newton, 2006). Drip sites feeding and immediately adjacent to stalagmite CC-Bil display sulphate concentrations which are consistently between 7.9 and 8.5 ppm. % SSS consistently accounts for 36% of the measured drip water sulphate and % bedrock-derived sulphate represents just 1.1% of drip water sulphate composition (Table 4).

6. DISCUSSION

6.1. Browns Folly Mine; BFM-Boss

On the understanding that the chloride content of cave drip waters can be used as an indicator of primary sea spray aerosol content at each site, apportionment of speleothem CAS between the remaining sources is possible using the δ^{34} S end-member compositions and δ^{18} O process information outlined above. Based on the % SSS content and ratio of calcium to sulphate in cave drip waters feeding stalagmite BFM-Boss (Table 3), 29% of the constituent sulphate appears to be of primary marine origin and just 1.4% is sourced from the carbonate bedrock. As the δ^{34} S marine end-member value for primary sea-salt aerosol is +21‰ and the bedrock has an average $\delta^{34}S$ composition of +20%, mass balance suggests the additional source of sulphur contributing to contemporary speleothem CAS must have a δ^{34} S value close to -3°_{00} (actual measured δ^{34} S composition +3.5 to +5.5%). On the basis that anthropogenic emissions of sulphur in precipitation are documented to reside between -3 and +9% in industrialised countries of the northern hemisphere (Mayer, 1998), this would suggest the non sea-salt sulphur within stalagmite BFM-BOSS is derived predominantly from anthropogenic pollution (e.g.,

Zhao et al., 1998) (Fig. 5). Such an interpretation is consistent with the industrial nature of the region surrounding this particular site. Continental biogenic emissions of reduced sulphur compounds may also have δ^{34} S values within the range required to account for the additional source of sulphur contributing to speleothem CAS. However, oxidation of such biogenic emissions within the atmosphere would render the constituent oxygen isotopes too light to account for the source of speleothem sulphate (see Fig. 5). Bedrock-derived CAS δ^{34} S values (+17.8 to +21.9%) are considerably higher than sulphate structurally bound within stalagmite calcite. Release of this heavy bedrock sulphate into groundwaters may contribute an additional source of sulphate to the drip water, although in the most recently deposited speleothem calcite, bedrock sulphate contributes just 1.4% of the total stalagmite CAS composition. δ^{34} S values are stable between 1960 and 1992 at \sim +3.6%, although calcite deposited prior to 1962 hosts δ^{34} S-cas values up to +5.5%. This may reflect lower sulphur emissions from industrial activity during the early 20th C and relatively greater contributions from marine and bedrock-derived sources with heavier sulphur isotopic composition. The lack of co variation between sulphur concentrations and sulphur isotopes post 1960 may reflect the dominant source of anthropogenic sulphur in the atmosphere. The minimal relative contribution from additional sulphur sources may cause little perturbation from pollution end-member compositions.

 δ^{18} O-CAS signatures range from +10.3‰ to +13.7‰, with the heavier isotopic values found during the earlier growth phases of the stalagmite. These signatures fall in a range where interpretations of sulphur cycling in the sediments and biomass above the cave are somewhat ambiguous. The δ^{18} O-CAS compositions are similar to those expected in atmospheric sulphate deposition (Fig. 5) and could represent a direct transfer of atmospheric precipitation into the cave environment via rapid flow routing, having undergone very little biogeochemical transformation. Based on 29% sea-salt sulphate contribution as primary

Sulphur concentration and isotope ratios in stalagmite BFM-Boss							
Distance below top of stalagmite (mm)	Year of deposition	S (ppm)	δ ³⁴ S-SO ₄ vs CDT (Rep 1)	δ ³⁴ S-SO ₄ vs CDT (Rep 2)	δ^{34} S-SO ₄ vs CDT (average)	Standard deviation between replicates	δ ¹⁸ O-SO ₄ vs V-SMOW (‰)
0-4	1995–1990	264	3.6	4.0	3.8	0.24	10.6
4-8	1990–1984	214	3.5	3.5	3.5	0.02	10.5
8-12	1984–1973	160	3.8	3.3	3.5	0.35	10.3
12-16	1973-1958	239	3.7	3.3	3.5	0.22	10.9
16-20	1958-1938	130	_	4.3	4.3	_	13.6
20-24	1938–1916	169	_	5.5	5.5	_	13.7

Table 1 Sulphur concentration and isotope ratios in stalagmite BFM-Boss

marine aerosol (δ^{18} O-so₄ of primary marine aerosol is +9.7% (Lloyd, 1967)), the remaining non sea-salt sulphate (NSS) in the youngest part of the stalagmite is calculated to

 Table 2

 Bedrock sulphur concentrations and isotope ratios

Rock	S concentration (ppm)	Ca/SO ₄ ratio (weight ratio)	δ ³⁴ S-SO ₄ vs CDT
Browns Folly	166	806	22.0
Mine 1			
Browns Folly	212	631	22.0
Mine 2			
Browns Folly	269	497	17.6
Mine 3			
Browns Folly	241	555	18.9
Mine 5			
Browns Folly	191	700	21.3
Mine 6			
Browns Folly	156	858	20.1
Mine 7			
Browns Folly	312	429	17.9
Mine 10			
Crag Cave 1	52	2573	_
Crag Cave 2	51	2623	
Crag Cave 3	92	1454	

have an end-member composition of $\pm 10.9\%$, typical for atmospheric sulphate deposition of mixed primary anthropgenic and secondary aerosol sulphate origin.

To produce drip water sulphate with $^{18}O/^{16}O$ ratios identical to those found in speleothem CAS via biogeochemical mineralisation of organic sulphur would require approximately 80% of the constituent oxygen to be sourced from atmospheric O₂ (assuming fractionation during the

	<i>,</i> ,•	1	. ,	<i>.</i>	1 ' CC D'I
Sulphur	concentration	and	isotope	ratios in	stalagmite CC-Bil
D ' .	a			s34g.g.g.	s180.00

Distance below top of stalagmite (mm)	S concentration (ppm)	δ ³⁴ S-SO ₄ vs CDT (‰)	δ^{18} O-SO ₄ vs V-SMOW (‰)
0–10	142	18.6	23.9
10-20	173	17.6	23.0
20-30	184	17.5	22.7
30-40	107	16.9	20.7
40-50	134	16.8	22.5
50-60	184	18.5	22.8
60-70	178	17.7	19.2
70-80	_	19.1	22.6
80–90	_	19.1	22.5
90–100	—	18.7	22.7

Table 3

Drip site ion chemistry

Cave site	Drip site	n*	Ca ²⁺ (mg/l)	SO ₄ ^{2–} (mg/l)	Cl ⁻ (mg/l)	Ca/SO ₄ ratio	% SSS	% Sulphur from bedrock CAS in drip waters
Browns Folly Mine, UK Midlands (January 1996 – February 1998)	BFM-Boss	14	99.0 (14.5)	11.0 (3.4)	24.2 (16.2)	9.0 (3.9)	28.7 (15.0)	1.4
Crag Cave, S.W.	CC-Bil	1	182.8	7.9	19.8	23.1	35.1	1.1
Ireland. Lower chamber. 2006	Seepage flow							
Crag Cave, S.W.	R	8	113.7 (4.7)	8.2 (0.1)	21.6 (0.2)	13.8 (0.5)	36.7 (0.3)	0.7
Ireland. Lower chamber. August 1997	Seepage flow							
Crag Cave, S.W.	Х	10	137.1 (33.2)	8.1 (0.1)	20.8 (0.3)	17.0 (4.4)	36.2 (0.6)	0.8
Ireland. Lower chamber. August 1997	Seepage flow							
Crag Cave, S.W.	Z	11	133.3 (31.7)	8.4 (0.05)	21.7 (0.1)	15.8 (3.8)	36.0 (0.2)	0.8
Ireland. Lower chamber. August 1997	Seepage flow							

Data from Crag Cave drip sites A-Z obtained from Tooth, 2000.

* n represents the number of field samples. 1 standard deviation (1 SD) is given in parentheses.

Fig. 4. Sulphur concentration, δ^{34} S-CAS and δ^{18} O-CAS in stalagmite CC-Bil.

Fig. 5. Diagram illustrating the range of δ^{34} S-cAs and δ^{18} O-cAs in stalagmite BFM-Boss compared to potential sulphate sources. The measured range of δ^{18} O in BFM-Boss is encompassed within the region of precipitation sulphate. Range of measured δ^{34} S-so₄ in precipitation for industrialised countries of the northern hemisphere obtained from Mayer (1998). Range of published δ^{18} O-so₄ in precipitation for midlatitude sites obtained from Jamieson and Wadleigh, 1999; Jenkins and Bao, 2006.

mineralisation and incorporation of atmospheric oxygen into sulphate to be -8.7_{00}° (Lloyd, 1968)) and an average drip water δ^{18} O-H₂O composition of -5_{00}° . However, given that experimental data have demonstrated biological reaction pathways may produce sulphate containing a maximum of 55% atmospheric O₂ only under exceptional circumstances (Toran and Harris, 1989), the biogeochemical mineralisation of organic sulphur compounds to sulphate is not considered to be apparent in the drip water flow pathway feeding this stalagmite (Fig. 5).

If the main source of sulphate contained within organic matter is present as organic ester sulphate compounds, hydrolysis may produce free sulphate ions within a 2-3% range of the oxygen isotopic composition of the original

sulphate compound (Mayer et al., 1995b). This would give an initial oxygen isotopic composition of approximately $12-13_{00}^{\circ}$, consistent with an atmospheric aerosol source.

There is clear evidence at this cave site for both a longterm hydrological storage component and a very rapid flow route at times of high rainfall (Fairchild et al., 2006b), thus both mechanisms (ester sulphate hydrolysis and rapid flow routing) remain possible contributors to the observed δ^{18} O-cas signatures at this site.

6.2. Crag Cave; CC-Bil

Crag Cave presents a contrasting case in being largely isolated from sources of anthropogenic SO_2 emissions and the large scale combustion of fossil fuels due to a predominantly westerly airflow and its position on the western periphery of Europe (Aherne and Farrell, 2002). Prevailing westerly winds typically provide the main source of atmospheric sulphur transporting aerosol of primary sea-salt origin onto the continental margin (Jordan, 1997; Aherne and Farrell, 2002).

Ratios of sulphate to chloride in atmospheric deposition from the region typically define a sea-salt component of ~80% (Jordan, 1997; Aherne and Farrell, 2002), of which the remaining 20% could be sourced from DMS, distal sources of pollution, or continental biogenic emissions. This contrasts with ratios to chloride in drip waters feeding stalagmite CC-Bil, where just 35% of sulphate is of sea-salt origin. This suggests an additional source of sulphur is obtained from somewhere along the drip water flow path. Values of δ^{34} S-cas in recent speleothem calcite are +18.6% (Table 4), leaving an end-member non sea-salt sulphate composition of +17.2%. This value appears typical of measured atmospheric deposition in the region (Bottrell and Novak, 1997; quoted average value of +17.8%) although could represent any mixture of atmospheric sources stored within the soil profile as organic sulphur, prior to re-mineralisation and release into the drip water flow path. Based on a comparison of Ca/SO₄ ratios in bedrocks and drip waters. carbonation and release of sulphate from the surrounding bedrock appears to contribute just 1.1% of sulphate to the drip water composition and is thus insufficient to account for the additional sulphate source. Signatures of δ^{18} O-cas are used below to try and identify the nature of this additional non sea-salt sulphate.

Values of δ^{18} O-cAs are isotopically enriched in ¹⁸O beyond that expected for values of atmospheric sulphate in an environment distal from extensive inputs of industrial pollution. δ^{18} O-cAs values are also in excess of sulphate oxygen signatures produced through (1) biogeochemical mineralisation of sulphur to sulphate under oxidising conditions; (2) oxidation of continental biogenic emissions released into the atmosphere; (3) oxidation of marine DMS compounds; or (4) the release of primary sea-salt aerosol (Fig. 6). The only feasible mechanism of obtaining oxygen isotopes so enriched in ¹⁸O would entail a degree of sul-

Fig. 6. Diagram illustrating the range of δ^{34} S-cAs and δ^{18} O-cAs in stalagmite CC-Bil compared to potential sulphate sources. The range of measured δ^{34} S-cAs and δ^{18} O-cAs is distinct compared to other sources of sulphate, suggesting additional sulphur cycling must modify the composition of the source materials prior to incorporation into stalagmite calcite. Range of measured δ^{34} S-so₄ in precipitation for industrialised countries of the northern hemisphere obtained from Mayer, 1998. Range of published δ^{18} O-so₄ in precipitation for mid-latitude sites obtained from Jamieson and Wadleigh, 1999; Jenkins and Bao, 2006.

phate reduction. At an advanced stage of sulphate reduction, equilibrium oxygen isotope exchange between reduced sulphur compounds and ambient water enables enrichment of ¹⁸O in residual sulphur compounds dependent upon the temperature and isotopic composition of the surrounding water (enrichment $\sim +28\%$ at cave temperature of 10.4 °C; cf. Fritz et al., 1989). Given that the average δ^{18} O-H₂O composition of the drip water feeding stalagmite CC-Bil has a value of -5.4%, equilibrium isotopic exchange during sulphate reduction would thus impart an oxygen isotopic composition approaching +23% in the residual sulphate, similar to that measured in stalagmite CC-Bil (Table 4). The offset between the lowest measured δ^{18} O-cas composition and the proposed equilibrium value of +23% may indicate a small contribution from the re-oxidation of sulphur compounds and the direct incorporation of an oxygen atom from water during sulphite oxidation. The range in δ^{18} O-CAS signatures through time is thus likely explained through differences in the oxygen isotope exchange rates according to changes in the oxygen isotope composition of ambient water, temperature variations and the degree of re-oxidation (Brunner et al., 2005).

Given the extent of sulphate reduction inferred to occur along the drip water flow path, provenancing the source of additional sulphate using isotopic end-member analysis is not possible. The following two mechanisms may produce drip waters at Crag Cave with sulphate isotopic compositions similar to those measured in CC-Bil. (1) The oxidation of pyrite to sulphate would typically contribute a source of isotopically light sulphur. Subsequent reduction of sulphate may produce residual sulphate enriched in ³⁴S and ¹⁸O. Such a mechanism would have to occur under a system where regions of reduction are spatially isolated from areas of sulphide oxidation, thus allowing kinetic fractionation and enrichment of ³⁴S in the residual sulphate pool. Given the local presence of pyrite within bedrock horizons sampled from elsewhere within the cave (Tooth and Fairchild, 2003), this remains a potential source and cannot be disregarded conclusively. (2) Alternatively, the mineralisation of organic sulphur compounds to sulphate may contribute an additional source of sulphur with similar isotopic composition to that found in atmospheric precipitation. Extensive cyclical sulphate reduction/re-oxidation will enrich oxygen isotopes to a few per mille lighter than the equilibrium value of +23%, whilst sulphur isotopes will undergo very little net fractionation between atmospheric deposition and drip water composition as isotope mass balance is maintained. Net fractionation of sulphur isotopes would be close to unity and reduction/oxidation would have to take place without extensive product loss from the reactant pool. The sulphur isotopic composition of speleothem CAS should thus reflect the isotopic composition of atmospheric sulphur deposition, assuming minimal fractionation during uptake and cycling through vegetation. The soil and vegetation thus acts as a store of organic sulphur which is slowly released into dripwater flowpaths and speleothem calcite (cf. Einsiel and Mayer, 2005).

Thus, based on δ^{34} S and δ^{18} O signatures obtained from speleothem CAS at Crag Cave, the drip water feeding stalagmite CC-Bil appears to be subjected to reducing condi-

tions along the flowpath, mostly likely within the till. The water-saturated nature of the fine grained, clay rich till above the cave provides the necessary conditions for sulphate reduction, whilst patches of iron staining and soil pH as low as 3.5 may be indicative of sulphide oxidation utilising FeIII as an oxidising agent under sub-oxic conditions (Soil profile descriptions obtained from Tooth, 2000). For pyrite oxidation and subsequent reduction to produce isotopic values similar to those observed in speleothem CAS, processes of oxidation and reduction must be spatially separate. Where cyclical sulphate reduction/re-oxidation is apparent, the re-oxidation of reduced sulphur compounds draws the δ^{18} O composition of the residual sulphate away from the equilibrium value and may be extensive enough to reverse the kinetic isotopic enrichment of ³⁴S such that there is very little net fractionation of source sulphur compounds.

7. CONCLUSIONS

There is currently very limited knowledge regarding the historical impacts of increased SO₂ emissions from anthropogenic pollution at the local and regional scale. Here, stalagmites have been demonstrated to contain sulphur in the form of calcite associated sulphate which can be readily extracted and analysed for δ^{34} S-so₄ and δ^{18} O-so₄ signatures. Using the techniques established above, sulphur and oxygen isotope measurements are conducted using approximately 200 mg of calcite powder (equivalent to just 35 µg of S) to obtain a time series record of sulphur inputs to the karst system. Source partitioning of the constituent sulphate using end-member isotopic analysis, sea-salt ratios to chloride and bedrock ratios of Ca/SO₄, has identified the dominant component of sulphur at Browns Folly Mine to be of anthropogenic origin reflecting the industrial nature of the surrounding region. Sulphur and oxygen isotope ratios of speleothem CAS at Crag Cave, W. Ireland, reflect processes of sulphate reduction along drip water flow paths and are therefore unsuitable for directly obtaining atmospheric sulphur records. Thus the potential for speleothems to be used as archives of atmospheric sulphur pollution appears encouraging where sediments overlying the cave system are oxidising throughout, as might be expected where thin brown earth soils are found in many temperate and Mediterranean climate zones. Where stalagmites are fed by drip waters of low redox status, for example at high latitudes where clay rich tills or peat soils are frequently found, their use as archives of atmospheric sulphur pollution appears limited due to the extensive fractionation associated with processes of sulphate reduction. However, use of sulphate sulphur and oxygen isotopes as indicators of reducing status may have important implications in future speleothem studies for the interpretation of additional palaeoclimatic variables. At carefully selected cave sites, speleothems hold the potential to record short term variability in atmospheric sulphur loading and thereby enable an understanding of the local and regional significance of anthropogenic vs natural sulphur aerosols in forcing climatic change.

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