1	Sulphate concentration in cave dripwater and speleothems: long-term trends and overview of its
2	significance as proxy of environmental processes and climate changes
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15	Keywords: Sulphur, speleothems; karst water; volcanic eruptions; anthropogenic sulphur peak.
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18	Abstract
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20	Sulphate concentration in speleothems identifies major volcanic eruptions, provide useful information
21	on soil and aquifer dynamics and, similarity to the ¹⁴ C bomb peak, its Anthropocene peak can be used
22	to date recent cave formations. However, the transmission of S from the atmosphere to cave dripwater
23	and its incorporation in speleothems is subjected to biogeochemical cycling and accurate studies of

each cave site are needed in order to assess how the S atmospheric signal is modified and eventuallyencoded in speleothems.

26 This study investigates the role of biogeochemical cycling and aquifer hydrology by utilising published 27 and new dripwater and speleothem data from Grotta di Ernesto (ER) in northern Italy. Here we provide 28 the first long-term record of sulphate concentration in cave dripwater based on over 20 years of 29 measurements. Fast drip site st-ER1 is characterised by a continuous decrease in SO₄ concentration 30 from a high of 7.5 ± 0.8 mg/l in 1993-1994 to a low of 2.2 ± 0.2 mg/l in 2013-2014, and replicates with a 31 delay of ~15 years the decline in the atmospheric SO₂ emissions. The S-series of slow flow ER78 site 32 is further delayed by ~4.5 years in relation to the S retention in the aquifer matrix. The dripwater data 33 are used to extend the previously published S record (1810 – 1998 AD) of stalagmite ER78 and 34 reconstruct the anthropogenic S-peak: this displays a delay of ~ 20 years with respect to the atmospheric 35 S emission peak due to biogeochemical cycling and aquifer storage. However, sulphur recycling above 36 the cave did not operate with the same degree of efficiency through time, which resulted in a variable 37 time delay between S deposition and incorporation into the stalagmite. In the pre-Anthropocene era, 38 and in particular during the cold Little Ice Age, biogeochemical cycling was far less efficient than 39 today, and the fast transmission of the atmospheric signal allowed capture of S released during major 40 volcanic eruptions by stalagmites.

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43 1. Introduction

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Sulphur concentration in Greenland ice cores provided the first evidence of the impact of
anthropogenic SO₂ emissions on regional climate following industrialisation in the Northern
Hemisphere (Neftel et al., 1985; Mayewski et al., 1986). The detailed chemical investigation of

48	Greenland and Alpine ice cores provided further insight into the influence of natural and anthropogenic
49	sulphur sources on atmospheric SO ₄ load and led to the identification of volcanic eruptions
50	characterised by high explosivity (Wagenbach et al., 1988; Legrand and Mayewski, 1997;
51	Schwikowski et al., 1999; Preunkert et al., 2001; Bigler et al., 2002; Patris et al., 2002). The record of a
52	high-elevation (4450 m asl) ice cap in the Monte Rosa massif indicated that natural SO ₄ sources as sea-
53	spray, mineral dust and quiescent volcanic emissions accounted for the S atmospheric signal during
54	pre-industrial periods, but that their combined contribution diminished progressively below 10-15% in
55	the course of the 20 th century as a consequence of the increasing anthropogenic emissions
56	(Schwikowski et al., 1999). During pre-industrial time, major volcanic eruptions significantly increased
57	the dust and aerosol concentration (including SO ₄) in the atmosphere, causing climate perturbation
58	lasting several years. However, the lack of ice core archives at mid-latitudes and at low elevations
59	limits our understanding of the role of both natural and anthropogenic aerosols in modifying regional
60	climate. In these settings, carbonate speleothems (stalagmites and stalactites) provide complementary
61	information by capturing changes in the atmospheric composition as well as climate signals through a
62	range of isotopic and elemental proxies (McDermott, 2004; Fairchild and Treble, 2009).

63 Stalagmites are known to record atmospheric sulphur loads and ecosystem dynamics in karst 64 regions, thus expanding the inventory of trace element proxies extracted from speleothems to a level 65 commensurate with ice cores (Frisia et al., 2005, 2008; Wynn et al., 2008, 2010, 2013). Sulphur concentration trends in stalagmites were related to anthropogenic emissions starting in the late 19th 66 67 century (Frisia et al., 2005, 2008; Uchida et al., 2013). It was also suggested that discrete S peaks in stalagmites record high explosivity index volcanic eruptions (Frisia et al., 2005, 2008; Badertscher et 68 69 al., 2014). Contrary to ice cores, the transmission of S from the atmospheric deposition to cave 70 dripwater and its incorporation in speleothems is not direct. It can be subjected to biogeochemical 71 cycling in the soil and vegetation above the cave, which is highly efficient in forested catchments (eg.

72 Likens et al., 2002; Shanley et al., 2005; Novak et al., 2007). These processes potentially modify the S 73 concentration and S isotopic signature in stalagmites, as well as delay the transmission of the sulphur 74 atmospheric signal into the epikarst (Wynn et al., 2013). 75 In a review of the literature on sulphur cycling in forested catchments, Wynn et al. (2013) 76 highlighted that biogeochemical cycling may retain sulphur in the soil and vegetation overlying the 77 cave system through processes of inorganic adsorption as well as biologically mediated processes of: i) 78 assimilation, which is the uptake and incorporation of sulphate into soil microorganisms and 79 vegetation; ii) mineralisation, which is the conversion of carbon bonded sulphur compounds released 80 from decaying vegetation and soil organic matter into sulphate; iii) oxidation, which is the 81 incorporation of oxygen into inorganic sulphides; and iv) dissimilatory reduction, which is the removal 82 of oxygen from sulphur compounds to ultimately produce H_2S , occurring under reducing conditions. 83 Fast infiltration events and/or infiltration where soil is thin and vegetation cover is scarce allow quick 84 flow through the soil zone. In this case, preferential flow routing through epikarst results in speleothem 85 capture of a sulphate signal largely unmodified with respect to the atmospheric source. Most 86 commonly, the biogeochemical cycling of sulphur through vegetation and soils and the successive 87 mixing with stored soil and aquifer sources will delay, concentrate and fractionate the S chemical or 88 isotopic signal in dripwater and, eventually, speleothems (Wynn et al., 2008, 2010, 2013). 89 Despite the importance of understanding the timing of atmospheric sulphur transfer into caves, there 90 are very few sulphur data available for karst dripwater (Wynn et al., 2008, 2010; Riechelmann et al., 91 2011; Tremaine and Froelich, 2013; Uchida et al., 2013). Moreover, several data are strongly modified 92 by evapotranspiration (Tremaine and Froelich, 2013) and pyrite oxidation in the host rock

93 (Riechelmann et al., 2011), or were collected in caves located near the sea and affected by a high

proportion of sea-salt derived SO₄ (Uchida et al., 2013). Critically dripwater values were not collected

95 close enough in time to the ca. 1975-1980 peak in anthropogenic emissions (Fig. 1) to document the

96 analogous succeeding peak in dripwater values.

97 This study aims to provide further insight into the speleothem sulphur record by reviewing the 98 published literature and presenting new data of dripwater S concentration over time at the well-studied 99 site of Grotta di Ernesto in Northern Italy. In particular, this paper provides the *first long-term record* 100 of sulphate concentration in cave dripwater based on over 20 years of measurements in the cave. The 101 dripwater data are compared with parallel data from meteoric and soil water collected over a shorter 102 time period (2002 - 2007 AD) and with the sulphur dioxide emissions in Italy and Europe. A long-term 103 record of S concentration is presented for a large karst spring in the region whose catchment is at high 104 altitude and, hence, far less influenced by soil and vegetation dynamics. In this way, we aim to assess 105 the influence of soil efficiency, vegetation cover and aquifer structure on the transmission of the S 106 signal. By calculating an in-cave S-partition coefficient, the published S concentration data in Grotta di 107 Ernesto ER78 stalagmite (1810 – 1998 AD; Frisia et al., 2005a, 2008) are extended to 2014 AD. The 108 extended ER78 time series is then compared to both the S emission curves and the major historic 109 volcanic eruptions in order to assess the transmission of the S signal under the cold temperatures and 110 reduced soil efficiency of the Little Ice Age and under the increasing temperature, soil efficiency and S 111 atmospheric load following industrialization. This becomes the basis for determining the transmissivity 112 of the S signal from the atmosphere to the cave under different environmental conditions. Finally, the 113 observed S peak of the industrial era is proposed as a relatively cheap and quick method for confirming modern calcite deposition in addition to and/or replacement of the ¹⁴C-bomb peak method (Genty and 114 115 Massault, 1997, 1999).

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117 **2.** Sulphur in stalagmites and S cycling in forested catchments

119	The transmission of sulphur from atmosphere to cave dripwater and speleothems has been
120	extensively studied at Grotta di Ernesto (ER), a mid-latitude, mid-altitude cave in the southernmost
121	reaches of the Dolomites (Trento province, NE Italy, Fig. 2) (Fairchild et al., 2000, 2009; Frisia et al.,
122	2005, 2008; Wynn et al., 2010, 2013). The first study of S concentration was carried out on annually
123	laminated ER78 stalagmite and highlighted an increasing S trend from 1810 to 1998 AD interpreted as
124	recording an increase in European S emissions through the industrial era (Frisia et al., 2005, 2008;
125	Fairchild and Frisia, 2014). In most of the record, sulphur displayed a clear annual cyclicity, which was
126	interpreted as related to seasonal pH changes driven by cave ventilation (Frisia et al., 2005, 2011), and
127	independent of any seasonal variation in atmospheric S deposition (Wynn et al., 2014a). However, the
128	20 th century sulphate rise in ER78 lagged the rise in atmospheric S emissions (Fig. 1) by ca. 20 years.
129	A similar delay was observed between peak S emissions status in Europe (1975 – 1980 AD) and the S
130	peak in the stalagmite (1996 AD) (cf. Frisia et al., 2008). This delay was attributed to mineralisation
131	and storage of sulphur within the soil and vegetation (eg. Wynn et al., 2013), as well as due to the
132	hydrology of the drip site which was fed by matrix flow draining the porous portion on the aquifer and
133	characterised by a long residence time (Miorandi et al., 2010). Critically, tree ring records of sulphur
134	dynamics sampled from the trees above the cave demonstrated a sulphur peak which occurred earlier
135	than in the stalagmite, thereby supporting the notion of long term storage of sulphur within the
136	biogeochemical cycle (Fairchild et al., 2009; Wynn et al., 2014b).

137The recent portion (1850 - 1996 AD) of stalagmite ER78 was subsequently analysed for138sulphur and sulphur isotopes by Secondary Ionisation Mass Spectrometry by Wynn et al. (2010). The139increasing trend in S concentration over the past 100 years was found to be related to the progressive140incorporation of S derived from SO2 industrial emissions, as demonstrated by sulphur isotopic141composition from values close to the carbonate bedrock end-member in the pre-industrial portion, to142values isotopically depleted in 34 S (Wynn et al., 2010). In the same study, a similar record of

atmospheric pollution was presented for stalagmite Obi84 from Obir cave in the Austrian Alps. Sulphur
in the Austrian stalagmite peaked in the 1980s and then returned toward pre-industrial levels and ³⁴S
enrichment in circa 10 years (Wynn et al., 2010). This suggested a less pronounced influence of
biogeochemical cycling and aquifer storage with respect to the ER site.

Isotope analysis of δ^{34} S-SO₄ and δ^{18} O-SO₄ were used to trace biogeochemical cycling of 147 148 atmospheric sulphur between 2005 and 2007 AD (Wynn et al., 2013), and highlighted that sulphate in 149 soil and dripwater at the ER site is primarily derived from an atmospheric source, while bedrock 150 contribution is only 2.1%. The weighted mean S concentration in meteoric water (423 ppb) suggested a 151 high input of sulphate derived from anthropogenic emissions, with a progressive increase in S 152 concentration in soil water (788 ppb), and cave dripwater (fast-flow st-ER1 = 1060 ppb; slow-flow st-153 ER78 = 1350 ppb). The increase in S content in soil and dripwater with respect to the meteoric signature was accompanied by a shift to higher ³⁴S values suggesting a two-phase acquisition of 154 sulphur: 1) concentration and ³⁴S-enrichment in the soil zone through storage, assimilation and 155 156 mineralisation; 2) S acquisition in the epikarst and in the aquifer by mixing with long residence time 157 water emplaced during times of high industrial SO_2 emissions (Wynn et al., 2013) as suggested by the 158 S trend recorded in the ER78 stalagmite (Frisia et al., 2005, 2008).

159 On the basis of the published results (Frisia et al., 2005, 2008; Wynn et al., 2010, 2013) and by 160 taking into account the different hydrology of the two drip sites (st-ER1 and st-ER78) extensively 161 studied by Miorandi et al. (2010) a conceptual model for sulphur cycling and transport at Grotta di 162 Ernesto was then developed by Wynn et al. (2013). The model identified three different scenarios (Fig. 163 3 panel A to C) and can be utilised as a reference for karst forested catchment in temperate climate 164 settings: A) under low sulphur atmospheric deposition S is biogeochemically modified by assimilation 165 and stored in the soil by mineralisation prior to entering the epikarst; B) during fast infiltration events S 166 is transmitted directly into the epikarst; C) during times of high SO₂ atmospheric deposition the relative

significance of biogeochemical cycling is diminished and most of the S is transmitted unmodified to the epikarst. In the epikarst and in the aquifer S can be rapidly transmitted into the cave via fissure flow, or stored and be subjected to redox cycling in the matrix (Fig. 3). Each drip site in the cave is characterised by a unique combination of the above conditions and the relative importance of each will vary through time as a response to changing environmental and hydrological conditions.

172 Critically, anthropogenic S peak was found in two stalagmites removed from two nearby caves 173 in southwest Japan, which recorded an increase in S concentration in the course of the 20th century 174 (Uchida et al., 2013). Despite the relatively low resolution of the S record in the two Japanese samples 175 (10 years) a clear increase of S concentration is evident after ca. 1950 AD, with peak values reached 176 between 1990 and 2000 AD. The δ^{34} S values suggested that the S concentration peaks were related to 177 the long-distance transport of emissions from coal burning activity in China, (Uchida et al., 2013).

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179 **3. Sulphur in stalagmites as a record of volcanic eruptions**

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181 The issue of S cycling is crucial for assessing sulphur peaks in stalagmites as markers of 182 volcanic eruptions. In the first study highlighting S in speleothems as a proxy for volcanic eruptions, high resolution Synchrotron Radiation X-Ray Fluorescence (SR-XRF) revealed eight distinct sulphate 183 184 peaks 8 to10 µm in thickness within the 5.06 to 5.19±0.16 kyr interval of stalagmite SV1 from Grotta 185 Savi (Trieste, NE Italy) (Frisia et al., 2005). The SR-XRF map for one of the peaks demonstrated that 186 the sulphate peak coincided with a single layer, consistent with a single seasonal input pulse, similar to 187 the pattern of preservation of sulphate aerosol in recent ice cores (Bigler et al., 2002). Given that Grotta 188 Savi soil is less than 50 cm thick, punctuated by rock outcrops and supporting scanty vegetation (Belli 189 et al., 2013) the biogeochemical cycling of S likely had a minor effect. Therefore, it was hypothesized 190 that the sulphate peaks in Grotta Savi stalagmite recorded multiple volcanic emissions (Frisia et al.,

191 2005). Similarly, SR-XRF analysis of stalagmite ER78 in the "pre-anthropogenic" period from 1810 to 192 1900 AD recorded two distinctive S peaks at circa 1815 and 1884-1888 AD, which were commensurate 193 with the ages of the Tambora (1815 AD) and Krakatau (1883 AD) eruptions (Frisia et al., 2008). Thus, 194 S- peaks in stalagmites appeared to be good candidates to mark explosive volcanic eruptions. 195 Following this assumption, a SR-XRF investigation of stalagmite So-1 (Sofular cave, Turkey) 196 was carried out with the aim of refining the age of the Minoan eruption. S concentration was shown to 197 increase above baseline levels when bromine and molybdenum peaked (at 1621 ± 25 BC and 1617 ± 25 198 BC respectively), whereas S peaked at 1589 \pm 25 BC (age model based on U/Th analyses interpolated 199 by chemical annual laminae counting) (Badertscher et al., 2014). Because Br and Mo are only 200 marginally influenced by soil and vegetation processes, their concentration peak was interpreted as 201 tracking the Minoan eruption, whilst S did not provide an accurate age of the volcanic event because it 202 had a delay of 35-40 years related to biogeochemical cycling and aquifer storage (Badertscher et al.,

203 2014). Nevertheless, the S increase above its baseline concentration is almost synchronous with the 204 peak of the "fast response" Mo and Br tracers, which suggests that S remains a reliable marker for 205 volcanism in stalagmites. The So-1 record allows hypothesizing that the Sofular karst systems was 206 characterized at the time of the eruption by a twofold response to S atmospheric signal: one, which 207 allowed the almost immediate transmission to the cave through fractures and conduits; another where 208 the S signal is delayed by biogeochemical cycling in the soil and vegetation and by aquifer storage and 209 mixing (cf. Wynn et al., 2013).

It is important to observe that both in Savi cave (Frisia et al., 2005, 2008) and in Sofular cave (Badertscher et al., 2014) stalagmites, the peaks of S and the other "volcanogenic" elements are concentrated in continuous layers, 10 to 250 μ m thick. Fluorescence maps indicate that the peaks are not related to micro-particle inclusions that can be incorporated within the crystal boundaries as well as along the stalagmite growth surface (Frisia et al., 2005, 2012). Thus, it is important that the S

215 concentration peak can be related to continuous layers when claiming that the peaks are related to
216 precise abrupt events. In this perspective, XRF microscopy should be recognized as fundamental tool in
217 palaeoclimate research.

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219 4. Sites description data sources and methods

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221 The province of Trento, located within the Italian Southern Alps is characterized by mountain 222 ranges consisting of Mesozoic carbonates, reaching elevations up to 3160 m asl, and by a large, 223 Mesozoic carbonate karst plateau to the SE with elevation between 1000 and 2200 m asl, (Fig. 3). The 224 valley axes cut deep through the mountains, with trends dictated by major geological structures: the N-225 S trending Adige Valley and the W-E trending Valsugana, which have valley bottoms at ca. 200 m asl 226 to ca. 400 m asl respectively. The morphology of the terrain has effects on mesoscale circulation and 227 the development of a thermal boundary layer and accumulation of pollutants dispersed from the valley 228 bottoms during daytime heating (Gabrieli et al. 2011).

Back-trajectory studies indicate that the Trento province is mostly influenced by western Mediterranean cyclogenesis and advection of air masses from the eastern tropical Atlantic during extreme events (Bertò et al., 2004). Lagrangian moisture source diagnostic analysis (Sodemann and Zubler, 2010) revealed that the major sources of precipitation in the Southern Alps during 1995–2001 were from the western Mediterranean Sea (31%), the North Atlantic Ocean (33%), the Arctic, Nordic and Baltic Seas (13%), with the remaining 23% from land sources predominantly in the western Mediterranean, Central Europe, and the eastern North.

Grotta di Ernesto (1167 m asl, 45°58'37''N, 11°39'28''E) is a short and shallow (maximum
rock overburden 30 m) downward gallery cut in Jurassic limestone, overlain by clay-rich calcareous
brown soil (Calcari-Mollic Cambisols, pH 6.5–7.7) up to 100 cm thick. The present-day vegetation

above the cave is a mixed conifer - deciduous forest association, composed primarily of *Fagus*

sylvatica, *Picea abies* and *Abies alba* (Fairchild et al. 2009; Miorandi et al. 2010). Most trees started to
grow ca.80 years ago (Borsato et al., 2007a) following forest clearance carried out to facilitate military
operations before World War 1 (cf. Apolloni, 1996). This ecosystem disturbance is recorded in ER78
stalagmite, which shows organic-rich laminae enriched in heavy elements (Cu, Pb, Y and Zn) in the
period from1900 to 1920 AD, suggesting enhanced leaching of trace elements through the disturbed
soil profile (Borsato et al., 2007a).

246 Present day soil and aquifer dynamics at Grotta di Ernesto have been extensively investigated 247 using hydrochemistry, stable isotopes, pCO₂ and radiocarbon (Fairchild et al., 2000, Huang et al., 2001; 248 Borsato et al., 2007a; Fohlmeister et al., 2010; Frisia et al., 2010; Miorandi et al., 2010; Hartland et al., 249 2012; Johnston et al., 2013; Borsato et al., 2015). The aquifer has a dual-permeability, with a rapid 250 fracture-flow and matrix seepage flow component feeding preferentially slow-dripping stalactites 251 (Miorandi et al., 2010). Drip rates respond to seasonal aquifer recharge, which occurs primarily during 252 the spring snowmelt (April to May) and autumn rainfall (October to November). Homogenisation of the δ^{18} O and δ^{13} C signal in dripwaters contrasts with a marked seasonal signal recorded within the soil 253 254 water (Frisia et al., 2010). This is interpreted as mixing in the karst aquifer below the soil zone, 255 indicating a minimum mixing time of 2 years (Fohlmeister et al., 2010).

Acquasanta spring is one of the major karst springs of the Trento province, and its waters are used for domestic consumption. Consequently, the spring has been monitored for microbiological and chemical quality since 1980 (Borsato et al., 2007b; Borsato, 2010), thus representing the longest record for S concentration in karst water available for the region. The resurgence opens at 477 m asl in the Brenta Dolomites ($46^{\circ}13'00"N$, $11^{\circ}02'04"E$), and the mean altitude of the 25 km² catchment is 2000 m asl. A large part of the catchment is characterised by barren karst with pockets of soil supporting alpine shrubs and grass and receives a mean precipitation of 1255 ± 220 mm/year. Rainwater and snowmelt are 263 captured as both diffuse infiltration and as concentrated runoff sinking into depressions and being 264 quickly transmitted into the karst aquifer developed in a >1500 m thick Mesozoic limestone and 265 dolomite succession. Whilst the flow routes in the massive limestone are typically through fractures 266 and conduits in the dolomitic rock composed of Ca-rich dolomite with variable microstructural 267 characteristics and inter crystalline porosity (Frisia and Wenk, 1993), both matrix and fracture/conduit 268 flow are relevant. Similar to the drip regime at Grotta di Ernesto, Acquasanta spring shows strong 269 seasonal discharge variability, with high discharge recorded during snowmelt from May to June (from 800 to >4000 $1 \cdot s^{-1}$) and low discharge in late winter and early spring (from 100 to 400 $1 \cdot s^{-1}$). During the 270 271 highest seasonal discharge, the water mineralisation is at minimum levels, indicating a fast flow mostly 272 through conduits and large fractures, whereas in winter it displays the highest concentrations as the 273 water is drained predominantly from the smaller fissure network and the rock matrix. The Acquasanta 274 spring water should then be considered as typical for fast-response to surface changes; its high-altitude 275 catchment, above the timberline, ensures that little biogeochemical cycling of sulphur has occurred 276 from input to output of the karst waters. It is, therefore, reasonable to expect that Acquasanta waters 277 respond rapidly to the rise and decline in atmospheric S pollution levels.

278 Four dripwater sites at Grotta di Ernesto representing the two flow pathways (fracture-fed and 279 matrix-fed) were studied from 1993 to 2014. Some of the analyses from these studies have already 280 been published although often as aggregate mean values (Table 1). Drip sites st-ER77 (discharge 0.141 ± 0.046 ml min⁻¹) and st-ER78 (discharge 0.031 ± 0.035 ml min⁻¹), located 30 m below the surface and 281 282 0.4 m away one from each other, are slow drips, mostly fed by matrix flow. Drip sites st-ER76 (discharge 0.216 \pm 0.171 ml min⁻¹) and st-ER1 (discharge 12.4 \pm 11.0 ml min⁻¹), located just 20 m below 283 284 the surface represent intermediate (st-ER76) and fast (st-ER1) seasonal drips preferentially fed by 285 fracture flow and small conduits with a quick discharge response (less than 12 hours) to high 286 infiltration events (Miorandi et al. 2010). The time between two successive drops was measured by

chronometer by the same operator. The average of ten intervals for fast drips and three intervals for
very slow drips was then transformed into millilitres per minute by considering a mean drop volume of
0.12 ml, which results from measurements made on several stalactites at Grotta di Ernesto (Miorandi et
al. 2010).

Soil waters were also collected between 2002 and 2008 from two lysimeters installed above Grotta di Ernesto at 0.7 m depth. From July 2004 to December 2007, bulk monthly aggregate meteoric water samples from above the cave and in the meteorological station of Lavarone (30 km W of the cave on the same plateau and at the same altitude) were collected by using 20 cm diameter polyethylene buckets capturing both wet and dry deposition; sampling protocol and analytical details are reported in Wynn et al. (2013).

Spring waters were collected at Acquasanta spring with different sampling frequencies.
Between 1987 and 1989 chemical analyses were made twice a year, few analyses were made between
1990 and 2002, while between 2002 and 2007 chemical analyses were made twice a year. In 2008 and
2009 complete chemical analyses were made on a monthly basis, whereas from 2010 to 2014 analyses
were made one or twice a year.

302 The dripwater, soil water and spring water samples were collected in HCl-cleaned and milli-Q® 303 water rinsed HDPE bottles. The samples were filtered with Millipore filters 0.45 µm and analysed at 304 the hydrochemistry laboratory of Istituto Agrario di San Michele all'Adige (Grotta di Ernesto) and the 305 hydrochemistry laboratory of Agenzia Provinciale per la Protezione dell'Ambiente of the Autonomous 306 Province of Trento (Acquasanta spring). The S content as well as the major ions were analysed by ion 307 chromatography following the analytical techniques described in detail in Bertoldi et al., (2011). The 308 plausibility of the results was tested against ion balance, which was better than 3% for 98% of the 309 samples.

310	Dripwater, soil and meteoric water at Grotta di Ernesto between 2004 and 2008 were analysed
311	by ion chromatography at the University of Birmingham: sampling and analytical protocols are
312	described in detail in Wynn et al., (2013).
313	The theoretical S-content in stalagmite ER78 for the periods $1995 - 1998$, $2005 - 2008$ and
314	2013 – 2014 AD was reconstructed by using dripwater data, and the S partition coefficient was
315	calculated from the dripwater S/Ca ratio for the period in which the stalagmite and the dripwater series
316	overlapped (1995-1998). For each water analysis, we utilised the mean value of 7 calcite XRF analyses
317	corresponding to the water sampling date ± 1 month; a fixed Ca concentration of 400,000 ppm was
318	used in account for the low concentration in other elements in the stalagmite calcite (cf. Huang et al.,
319	2001, Borsato et al., 2007a). This approach to partitioning is an approximate solution, since Busenberg
320	and Plummer (1985) proposed that SO_4^{2-} substitutes for CO_3^{2-} in the calcite lattice and so carbonate ion
321	should substitute for calcium in the partition coefficient equation. This will be treated in a future
322	publication of experimental work.
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324	5. Results: the response of karst waters to atmospheric sulphur load
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326	Since the beginning of the monitoring program at Grotta di Ernesto in 1993, the SO ₄
327	concentration in dripwater has been constantly decreasing. In fast drip site st-ER1 SO ₄ concentrations
328	diminished exponentially ($R^2 = 0.84$, $p < 0.00001$) from 7.5 ±0.8 mg/l in 1993-1994 to 2.2 ±0.2 mg/l in
329	2013-2014 (Fig. 4). A similar exponential decrease ($R^2 = 0.92$, $p < 0.00001$) from 8.1±0.4 mg/l in
330	1995-1996 to 2.9 ±0.2 mg/l in 2013-2014 is evident in the slow drip sites (st-ER77, st-ER78)
331	characterised by an overall higher SO ₄ concentration with respect to fast drip st-ER1; in intermediate

decreased from 4.75±0.51 mg/l in 2002 to 3.74±1.58 mg/l in 2007, similar to that occurring in fast drip
st-ER1.

335	In Acquasanta spring, water SO ₄ concentration decreased slightly ($R^2 = 0.50$, $p < 0.002$) from
336	1.88 ± 0.33 mg/l in 1987-1989 to 1.14 ± 0.09 mg/l in 2013-2014, with a mean value of 1.46 ± 0.37 mg/l in
337	the years 2005 - 2006, when sulphate concentrations in meteoric water measured at Lavarone
338	(1.35±0.62 mg/l) and Grotta di Ernesto stations (1.26±0.50 mg/l) exhibit similar values (Fig. 5). The
339	spring water SO ₄ values and the measured S content in the Mesozoic dolomites and limestones
340	composing the Acquasanta spring aquifer (S $<$ 270 ppm, Wynn et al., 2013), suggest the bedrock
341	contribution to the SO_4 load is < 5% and, therefore, the S concentration in the spring water is
342	predominantly related to the atmospheric input.
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344	6. Discussion
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346	6.1. Sources of S emissions
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348	Available SO ₄ analyses for other karst spring waters of the same region with catchments above
349	2000 m asl and aquifers developed in similar lithology, show sulphate levels and trends similar to those
350	recorded for Acquasanta spring for the last 10 years (Borsato, 2010). However, dripwater in other caves

in the southern part of the Trento province (cf. Fig. 2) exhibit similar, or higher, SO₄ concentration with

respect to Grotta di Ernesto. During the period 2002-2003, dripwater SO₄ concentration in the mid

altitude Giazzera cave (infiltration elevation 1050 m asl) was 4.9 ± 0.2 mg/l, whereas in the low

altitude Bus del Diaol cave (infiltration elevation 355 m asl) was 9.4 ± 0.8 mg/l (Borsato et al.,

355 submitted). This spatial heterogeneity with respect to the infiltration elevation and latitude is similar to

that observed in the chemistry of surface snow in the same region, where the lowest concentrations in

SO₄ and other trace elements were measured in snow collected above 2000 m asl in the northernmost part of the Trento province, and the highest concentrations in snow at mid and low elevations in the southernmost reaches of the studied area (Gabrielli et al., 2008; Gabrieli et al., 2011). This suggests that, in addition to sea-salt, aeolian dust and bedrock contributions, there are two other factors that need to be accounted for when interpreting S concentration in meteoric and karst water: the geographic location relative to sources of S emissions and the altitude of the boundary layer.

363 The latitudinal trend in cave drips, karst spring water and snow can be ascribed to their relative 364 proximity to the highly industrialized Po Valley and Venetian Plain towards the South and East (Fig. 365 3), the most important regional sources of anthropogenic S pollution. This is confirmed by similar 366 observations made in the southern watershed of the central Alps where a clear South to North 367 diminishing trend in the SO₄ atmospheric deposition was documented (Rogora et al., 2006). 368 On the other hand, the decrease in SO_4 concentration with respect to site elevation is related to the 369 position of the local boundary layer (Gabrielli et al., 2008; Gabrieli, 2011). Balloon data from the 370 meteorological station of Milano-Linate (in the Po Valley, 190 km WSW of Grotta di Ernesto) indicate 371 that, in winter, the boundary layer depth (BLD) is higher than 2000 m asl only for a very limited 372 amount of time (15 to 20%), whereas a well-mixed boundary layer exists below 500 m asl (Gabrieli et 373 al. 2011). Therefore, in winter, the pollutants released in the Po Valley and Venetian Plain, as well as 374 on the western and northern side of the Alps, are confined to the lower elevation belt (i.e. Bus del 375 Diaol). The BLD rises in summer, due to stronger insolation that results in more effective convection, 376 and it is higher than 2000 m asl for 72% of the days from April to September (Gabrieli et al. 2011). 377 This allows the pollutants to be lifted and transported horizontally by the synoptic flow across the Alps 378 (Gabrieli et al., 2011). A similar consideration holds true for the western Alps, where the S atmospheric 379 concentration measured in 1991-1993 increased up to two orders of magnitude from low elevation sites 380 (500 m asl) to the highest glacier site (Col du Dôme, French Alps, 4250 m asl). Accordingly, from

1975 to 1985 AD, the summer S concentration in the Col du Dôme ice core was four times higher than
the winter value (Preunkert et al., 2001).

The variability in altitude of the local boundary layer influences the input of atmospheric S, and explains the greater contribution of anthropogenic sulphate in the southernmost sites (nearer to the local S-emissions) and at lower elevations. It is then reasonable to assume that caves at mid altitude in industrialized regions would record the effects of BLD variability, in addition to a regional atmospheric S signal modified by ecosystem and aquifer processes.

388 In assessing the provenance of anthropogenic emissions, the synoptic setting must be 389 considered. By using the aforementioned contribution to the sources to precipitations in the Southern 390 Alps (Sodemann and Zubler, 2010), the emissions affecting the study area over time were calculated. 391 Tabulated emissions are available for different European countries (Mylona, 1996, 1997; Vestreng et 392 al., 2007) and these were combined in the proportions: Italy (proximal location, 50% contribution), 393 Spain and France (North Atlantic region, 36%), and finally Germany, Poland, Hungary and Czech 394 Republic (northern provenance, 14%). As expected, the resulting emission curve is similar to the total 395 European emission curve, although the regional emission curve has a narrower peak that accounts of 396 the more pronounced contribution from Italy (Fig. 1).

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398 6.2. The industrial era S peak, aquifer storage and dilution effect

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400 SO₂ emissions associated with the industrial era and enhanced in 1950 – 2000 AD (Fig. 1) and 401 their fallout in karst soils can be considered as a global "tracing test" for caves around the world, 402 similar to the effects of the ¹⁴C bomb peak produced by nuclear tests during the 1960s (Genty and 403 Massault, 1999). This is particularly true for caves developed in almost pure carbonate rocks where the 404 S content in the host-rock is very low and the S in dripwater is derived almost entirely from

atmospheric sources (cf. Cap. 1). In these contexts the dynamics of sulphur emissions to the
atmosphere will be replicated within cave drip and karstic groundwater according to the variable extent
of biogeochemical cycling, water residence time and the degree of water source mixing. Waters
emerging at Aquasanta spring, where there has been little to no retention in the biogeochemical cycle
and hydrology is dominated by conduits and fracture flow, display sulphur concentrations closely
aligned to immediate atmospheric SO₂ loading (Figure 5).

411 At Ernesto cave, where the retention in the biogeochemical cycling is relevant, the dripwater 412 composition should be more influenced by past atmospheric composition and, by considering that SO₂ 413 emissions in the atmosphere are constantly decreasing following peak emissions status in 1976 AD 414 (Fig. 1), we can expect an higher S content in dripwater with respect to the coeval atmospheric SO₂ 415 loading (Figure 5). For the same reason, drip sites fed predominantly by matrix flow (st-ER77 and st-416 ER78) support concentrations of sulphur which are greater than those drip sites fed by fracture flow 417 dynamics (st-ER76 and st-ER1) in relation to the higher proportion in matrix fed stalactites of older 418 water with respect to fissure fed sites characterised by a shorter residence time.

419 In the case of drips st-ER76 and st-ER1 fed preferentially by fast fissure flow but with a 420 component of matrix flow (Miorandi et al., 2010) a dilution effect can be expected during higher 421 discharge periods, characterized by a major contribution of fast flow through fissures and conduits and 422 a reduced contribution from older water with high S-content stored in the aquifer. This is illustrated in 423 Fig. 6A, where the SO₄ concentration in fast-drip st-ER1 during specific time intervals (to minimize the influence of the progressive decrease in time of the SO₄ concentration in the aquifer matrix) displays a 424 425 negative exponential correlation with discharge. In particular, the SO₄ concentration varies from a high 426 concentration end-member corresponding to the contribution from the porous matrix to a low 427 concentration end-member reflecting the mean composition in the epikarst fractures, and similar to the 428 mean soil water concentration (Fig. 6A). By contrast, for the slow drip sites st-ER77 and st-ER78,

429 where the fissure flow component is negligible, no correlation between water SO_4 concentration and 430 discharge is evident (Fig. 6B).

Finally, by comparing the trends in the S series we can observe an offset of about 4.5 years between st-ER78 (matrix flow) and st-ER1 (predominant fissure flow) (Fig. 4 and A1). This time span should represent the additional delay related to the S retention in the aquifer matrix, although additional variables such as the different flow path of the two drips, the possible S-concentration related to the sulphate reduction in the aquifer and the secondary contribution from the host-rock must be taken into account.

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438 6.3. Sulphur trend in stalagmite ER78

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440 Annually laminated calcite stalagmite ER78 (Frisia et al., 2003), fed by stalactite st-ER78, was 441 analysed by synchrotron radiation X-ray fluorescence (XRF) at 2µm resolution (corresponding to ~32 442 analyses/year for the last century). The measured intensities (counts per seconds) could not be 443 transformed in actual concentrations because of the lack of standards with similar matrix (Frisia et al., 444 2005). The successive analyses by Secondary Ionisation Mass Spectrometry (SIMS) of the topmost part 445 of the stalagmite (Wynn et al., 2010) allowed quantification of the synchrotron XRF data (Fig. A2). In 446 Fig. 7 the quantified S data and the lamina thickness of ER78 are compared with the surface 447 temperature record, the S emission trend as well as with the SO₄ record in the Alpine ice core at Col du 448 Dôme (Preunker et al., 2001).

When considering the characteristic S emission peak from 1950 to 2010 AD, centred at 1976
AD, a delay of ca. 20 years can be observed in both the rising limb of the S series and the maximum

451 value (1996 AD). As discussed above, stalactite st-ER78 is fed by matrix flow with little contribution

452 from fast fissure flow. Thus, the S concentration in ER78 stalagmite is expected to reflect the S

composition in the epikarst and in the vadose zone (Wynn et al., 2013). Therefore, in the present-day situation of relatively low sulphur atmospheric deposition, S is biogeochemically modified in the soil prior to entering the epikarst (scenario A in Fig.3). The 20-year delay for the transmission of the atmospheric S signal to ER78 stalagmite, thus represents the sum of biogeochemical cycling and aquifer storage. However, the biogeochemical cycling of sulphur above the cave may not always have operated to similar extent, dependent upon vegetation density, soil efficiency and levels of sulphur loading to the overlying ecosystem.

During the cold Little Ice Age in the 19th century, temperatures in Northern Italy were 460 significantly lower than the mean 20th century values (-0.98±0.2°C calculated at Milan between 1800 – 461 1880 with respect to the 1901-1995 reference period; Maugeri and Nanni, 1998, Fig. 7). Cooling was 462 more pronounced than the value calculated for the European Alps and the rest of Europe (-0.33±0.1°C, 463 464 Luterbacher et al., 2004; Casty et al., 2005) and was likely responsible for the reduced annual growth 465 rate of stalagmites in Grotta di Ernesto (Frisia et al., 2003). Stalagmite growth rate (measured as annual 466 lamina thickness) depends on a number of factors including temperature, drip-rate, dripwater Ca 467 content, fluid film thickness, cave air pCO₂ (Dreybrodt, 1999) and the presence of growth inhibitors 468 (Meyer, 1984; Frisia et al., 2012). Of these, variations in drip-water supersaturation due to Ca content 469 or drip water pCO_2 (as a function of degassing in the epikarst or in the cave) are the most likely to vary 470 on a multi-year timescale (Frisia et al., 2003). These factors control parent water supersaturation with 471 respect to calcite, and reflect changes in CO_2 production and storage in the soil and epikarst above the 472 cave. In the studied area, these processes are modulated by temperature (Borsato et al., 2015). A 473 decrease of 1°C in the mean annual temperature would cause a decrease of 0.1 in the calcite saturation 474 index (SI_{CC}) (Borsato et al., 2015) and this, by considering the present-day saturation in ER dripwater 475 $(SI_{CC} = 0.38 \pm 0.15, Miorandi et al., 2010)$ would result in significant reduction in the stalagmite growth 476 rate (Dreybrodt, 1999; Frisia et al., 2003). Cooling also results in a reduction of vegetation cover and

477 soil activity (Frisia et al., 2003; Borsato et al., 2015) by slowing the biogeochemical cycling above the 478 cave (scenario C in Fig. 3). Diminished soil efficiency during the Little Ice Age is also testified by the 479 presence of thin (1-4 µm), brown UV-luminescent, organic and trace element-enriched sub-laminae at 480 the beginning of each hydrological year in autumn (Frisia et al., 2003; Borsato et al., 2007a; Scholz et 481 al., 2012). This suggests that microbial decomposition of organic matter in the soil zone was not as efficient as in the 20th century, when organic-rich sub-laminae almost disappear. Under these 482 483 circumstances, any delay in transmitting an atmospheric S signal to the cave was probably reduced to a 484 few years. This may explain why the volcanic eruptions of 1815 and 1883 AD were recorded as narrow 485 and short-term (4-6 years) S spikes, with apparently little delay or modification of the atmospheric 486 signal (Fig. 7).

487 Starting from 1900 AD, the annual growth rate of ER78 increased, following temperature and 488 solar radiation increase. A high sensitivity of the site to solar radiation is confirmed by spectral density 489 at the 11 year solar cycle recorded by the lamina thickness series (Frisia et al., 2003). Higher 490 temperature resulted in higher soil pCO_2 production (Borsato et al., 2015), more efficient host rock dissolution leading to higher Ca^{2+} content in cave waters (Dreybrodt, 1999) and, ultimately, to faster 491 492 growth rate in Grotta di Ernesto stalagmites (Frisia et al., 2003; Smith et al., 2006). Critically, the 493 increase in solar radiation and surface temperature coincided with industrialization, and a steady rise in 494 S emissions from 1890 to 1940 (Fig. 7). From 1950 there was a steep increase in S emissions, which 495 lasted until 1980, and an equally steep decline after that. These features are reproduced with a variable 496 delay by the S trend in ER78 likely related to the increasing vegetation cover and soil efficiency that 497 enhanced S biogeochemical cycling through the uptake and assimilation of sulphate in plants and 498 soil/root microorganisms, and S mineralisation and storage in the soil zone. In addition, the 499 combination of warming plus increasing pollution levels resulted in variations in BLD, which reached 500 the mid altitudes throughout the warm season, causing an enhanced transport of the S-emissions over

501	the Italian Alps at mid altitudes. A constant, high supply of atmospheric S eventually overcame the S
502	biogeochemical cycling capacity, thus, a significant proportion of atmospheric S was transmitted,
503	unmodified to the epikarst (scenario C in Fig. 3). Following these considerations, it is reasonable to
504	infer that ER78 stalagmite captured atmospheric S emissions with limited delay before the 20 th century
505	warming. This explains why pre-1900 explosive volcanic eruptions are recorded by well-defined, short-
506	lived S peaks in the stalagmite record as highlighted by the detrended S concentration (Fig. 7).
507	In the same detrended series between 1890 and 1970, when both lamina thickness and S content
508	commenced to increase, it is possible to observe a marked annual structure related to seasonal
509	fluctuations of dripwater pCO ₂ driven by cave ventilation (Frisia et al., 2011; Wynn et al., 2014a).
510	Starting from 1970, the S structure becomes noisy, and the annual signal is complicated by secondary
511	peaks, most likely in relation to high S flux from the soil zone, and the mixing in the aquifer of
512	different sources characterised by different S content (Fig. 3).
513	
514	
515	6.4. A future projection of the ER78 sulphur series and the anthropogenic S-peak
516	
517	The S concentrations in stalagmite ER78 and dripwater analyses for the corresponding drip site
518	st-ER78, allow for the calculation of the S partition coefficient (D_S) for the period in which the two
519	series overlapped (1995-1998):
520	S
521	Partition coefficient: $D_S = \frac{\frac{S}{Ca}}{\frac{S}{S}} drinwater$
522	Ca unpwater

523 The calculated D_S for each single water analysis are plotted against their corresponding S/Ca 524 ratio in the inset of Fig. 8, where a linear relationship between the calculated D_S and the dripwater S/Ca 525 ratio can be observed:

526

527
$$D_S = 0.148 \text{ S/Ca (molar)} - 0.01$$
 (R² = 0.82, p < 0.0005);

528

By using the mean calculated D_S value ($D_S = 0.0064 \pm 0.10$; cf. inset in Fig. 8) the theoretical stalagmite S content from the dripwater S/Ca data is reconstructed, and used to complement the sulphur record in stalagmite ER78 (Fig. 9). The exponential decrease evidenced by dripwater S/Ca data ($\mathbb{R}^2 =$ 0.89, *p* < 0.0001, Fig. 8) can then be used to project a theoretical stalagmite S-content up to 2030 AD, when the predicted values should reach pre-industrial values (around 15 ppm).

534 The reconstructed stalagmite series can then be compared with the SO_2 emissions: both series 535 are characterised by a prominent peak lasting 70-80 years and with values up to 10-15 times the pre-536 industrial concentration (Fig. 9). The rising limb and the centre of the S peak in the stalagmite are 537 delayed by ~20 years with respect to the S-emission peak, which is explained by biogeochemical 538 cycling and aquifer storage. By contrast, it is difficult to evaluate the delay in the more recent part of 539 the series because there are no equivalent points identifiable in the falling limb between the emissions 540 and the speleothem time series. However, by considering that the emissions in 2005 decreased to levels 541 similar to the 1920s levels and that the projected speleothem values should reach the corresponding 542 levels around 2025, we can estimate a similar delay of ~20 years. The stalagmite peak in its central 543 part (1990 - 2000) is sharper that the corresponding emissions peak: here we suggest that this can be 544 related to the transmission of part of the atmospheric S load unmodified into the epikarst, following the 545 overload in the soil and ecosystem above the cave (scenario C in Fig. 3).

546 On the basis of these considerations, the delay in sulphur transmission between atmosphere and stalagmite ER78 can be divided into three characteristic periods: i) in the pre-industrial era and the first 547 part of the 20th century the delay was of few years (possibly 4 to 8 years and mostly related to the mean 548 549 residence time in the aquifer) as the biogeochemical cycling was not very efficient; ii) starting from 550 1950 AD following the establishment of the forest above the cave and the enhanced atmospheric S 551 load, the delay progressively increased up to 20 years; iii) the increasing atmospheric S deposition 552 caused an "overload" in the soil and ecosystem above the cave, allowing the transmission of part of the 553 atmospheric S unmodified into the epikarst and sharpening the stalagmite peak in its central part (Fig. 554 9).

The anthropogenic S-peak is a very prominent feature in ER78 stalagmite, as well as in the record from Austrian stalagmite Obi84 (Wynn et al., 2010) and in two Japanese stalagmites (Uchida et al., 2013). Therefore, it can be used in speleothem-based palaeoclimate research to refine modern chronology and acquire complementary information regarding the soil and vegetation biogeocycling efficiency through time. In ideal cases, the S record can be characterized by an annual signal related to seasonal fluctuations of dripwater pCO_2 (Frisia et al., 2011; Wynn et al., 2014a) that can be used to further refine speleothem chronology.

The anthropogenic S-peak is complementary to the ¹⁴C bomb peak, which is used to date the 562 modern portion of speleothems (Genty and Massault, 1999; Mattey et al., 2008). With respect to the S-563 peak, the ¹⁴C bomb peak has the advantage of an initial rise around year 1958, which provides a 564 reliable marker for dating and correlation (Hodge et al., 2014). However, the maximum ¹⁴C peak in 565 566 speleothems can be lagged by up to 20 years with respect to the atmospheric peak centred at circa 567 1965, and the stalagmite signal can be affected by differing degrees of attenuation. Notwithstanding, 568 the major disadvantage is the technical aspect of the analyses, which are destructive, expensive and 569 time consuming. By contrast, recent advances of the sulphur analytical techniques such as SR-XRF,

570	SIMS and high mass resolution ICP mass spectrometry (Jochum et al., 2012) are now opening new
571	avenues of approach and we foresee that high resolution analyses of S concentration in recent
572	speleothems will be routinely carried out in future palaeoclimate reconstructions.
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574	
575	8. Conclusions
576	
577	S concentration variability in stalagmites represents a valuable proxy for volcanic eruptions and
578	anthropogenic emissions, and it may potentially become a highly sensitive proxy for high resolution
579	palaeoclimate and palaeoenvironmental reconstructions.
580	The study of soil processes and aquifer dynamics, as well as the knowledge of the synoptic
581	setting and geographic location with respect to emissions source, is an important pre-requisite for
582	successful interpretation of speleothem S series. Variations in the background levels of sulphur
583	contained within speleothems and the shape and duration of S peaks in both Holocene and pre-
584	Holocene stalagmites, especially when used in conjunctions with other trace elements, are able to
585	provide indications about changes in rates of biogeochemical cycling, as well as independent
586	information on the contribution of volcanic processes to atmospheric load. Moreover, when analysed at
587	high temporal resolution, annual cycles in the S record can be used to complement the dating accuracy
588	and the seasonality information gained from other more traditionally studied elements, such as Mg and
589	Sr (Fairchild and Treble, 2009; Badertscher et al., 2014).
590	Eventually, the potential of sulphur in speleothems as marker of volcanic eruptions (and,
591	consequently, as a correlative tool) depends on ecosystem biogeochemical cycling and mean residence
592	time of the water in the aquifer. Where signals in speleothem calcite are reliant on passing from
593	atmosphere to cave via the sulphur biogeochemical cycle, we suggest the best locations to obtain

594 accurate records of past volcanic eruptions from speleothems are mid altitude, and high to mid latitude 595 caves sites with thin soil cover and highly transmissive aquifers. Tropical and equatorial caves with 596 thick soil cover and long water residence time are the worst candidates. However, speleothems from 597 tropical and equatorial settings formed in caves cut in pure carbonate rocks overlain by scanty soils, or 598 characterized by high-rockiness soils, where aquifers are highly transmissive could yield a reliable 599 volcanic record. S should be used in conjunction with other volcanogenic trace elements such as Br 600 and Mo and high resolution mapping techniques could also be useful in order to distinguish thin 601 continuous layers from discrete grain inclusions.

Industrial anthropogenic SO_2 emissions can be considered a global "tracing test" for dripwater and groundwater in the Northern Hemisphere. In this perspective Grotta di Ernesto is particularly apt to capture the anthropogenic S emissions, being near the Venetian Plain, one of the most industrialized areas in Italy. Despite the delay in the transfer of the atmospheric signal from surface to cave, it is here demonstrated that the anthropogenic S peak is well preserved in both dripwater and speleothems. Thus, it is here proposed the use of the anthropogenic S concentration peak as an additional dating tool for recent speleothems, complementary to the radiocarbon bomb peak.

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Item	Period	References	Notes - Laboratory
	1993-1994	Borsato, 1995	Analyses made by <i>Agenzia Provinciale per la</i> <i>Protezione dell'Ambiente, Provincia Autonoma</i> <i>di Trento</i> . Bimonthly analyses from 06/1993 to 07/1994.
ER dripwater	1995-1997	Fairchild et al., 2000	In the paper only the mean dripwater SO ₄ concentration for all the analyses was reported. Bimonthy analyses from 06/1995 to 12/1997.
	2002-2003 and 2013-2014	this work	<i>Istituto Agrario di San Michele all'Adige,</i> <i>Hydrochemistry laboratory</i> . Monthly analyses from 03/2002 to 05/2003.
	2004-2007	Wynn et al., 2013	Monthly analyses from 07/2004 to 12/2007.
FR soil water	2002-2003	this work	<i>Istituto Agrario di San Michele all'Adige,</i> <i>Hydrochemistry laboratory.</i> Bimonthly analyses from 08/2002 to 12/2003.
	2005-2007	Wynn et al., 2013	Monthly analyses from 10/2005 to 12/2007. The S concentration of 14 bedrock samples is also reported.
ER meteoric water	2005-2007	Wynn et al., 2013	Monthly analyses from 12/2005 to 12/2007.
Lavarone meteoric water	2004-2006	Wynn et al., 2013	Monthly analyses from 07/2004 to 11/2006.
Acquasanta spring water	1987-1989, 1991, 1999, 2002-2014	this work	Servizio Geologico Provincia Autonoma di Trento, and Agenzia Provinciale per la Protezione dell'Ambiente. Spot measurements. The data from 2008 – 2009 (monthly analyses) are also included in the internal report: Borsato, 2010.
ER78	1890 - 1998	Frisia et al., 2005; Wynn et al. 2013	Age model in Frisia et al., 2003; XRF intensity data in Frisia et al., 2005; single points SIMS data in Wynn et al., 2013.
stalagmite	1890 - 1998	this work	Quantification of the XRF data.
	1994 - 2014	this work	Calculated theoretical S concentration on the basis of dripwater Sr/Ca and D _{S.}

799 800 Table 1. Synthesis of the sampling periods, sampling frequencies and data sources.









Fig. 2. Location of Grotta di Ernesto (ER) and Acquasanta karst spring (ACQ) on a digital elevation

- 813 model of the Trento province. AV = Adige Valley; VS = Valsugana; PV = Po Valley; VP = Venetian
- 814 Plain; GL = Garda Lake; DL = Bus del Diaol; GZ = Giazzera Cave.





817 Fig. 3. Conceptual model for S cycling and transport in forested catchment karst inspired by the Grotta 818 di Ernesto system (modified after Wynn et al., 2013). Three different scenarios are illustrated: A) under 819 low sulphur atmospheric deposition S is biogeochemically (BG) modified by assimilation and stored in 820 the soil by mineralisation prior to entering the epikarst; B) during fast infiltration events S is 821 transmitted directly into the epikarst and to the cave via preferential fissure flow. C) during times of 822 high SO₂ atmospheric deposition the relative significance of biogeochemical cycling is diminished and 823 most of the S is transmitted unmodified to the epikarst. In the epikarst and in the aquifer S can be 824 transmitted into the cave via fissure flow or stored and subjected to redox cycling (R) in the matrix. 825 Each drip site in the cave is characterised by a unique combination of the above conditions and the 826 relative importance of each will vary through time as a response to changing environmental and 827 hydrological conditions.



Fig. 4. Sulphate concentration time series for rainfall, soil water and cave dripwater at Grotta di
Ernesto compared with the long trend in sulphate concentration in Acquasanta karst spring and sulphur
dioxide emissions in Europe (cf. Fig. 1). For dripwater, green symbols and lines represent fast (st-ER1)
and intermediate (st-ER76) flow, whereas red symbols and lines represent slow flow (st-ER77 and stER78). For data sources see Tab. 1.



Fig. 5. Mean sulphate concentration and standard deviation in meteoric water, soil water and cave
dripwater at Grotta di Ernesto compared with the mean sulphate concentration of Acquasanta spring for
the period July 2005 – November 2006.



Fig. 6. Sulphate concentration vs. drip rate in stalactites st-ER1 (A), st-ER77 and st-ER78 (B) during
selected time intervals. In (A) the mean sulphate concentrations in soil and meteoric water for 20052006 are also plotted outside the vertical secondary axis.



849 Fig. 7. Mean annual temperature in Milan (grey line: annual data; red line: 5 years running mean) 850 (Maugeri and Nanni, 1998), ER78 lamina thickness (grey line: annual data; black line: 3 years running 851 mean) (Frisia et al., 2003), S concentration (grey line: actual data; black line 2 years Gaussian filter) 852 and detrended S concentration in ER78 stalagmite (data from Frisia et al., 2005, 2008) compared with 853 the modelled sulphur dioxide emissions for the Southern Alps (cf. Fig. 1) and the Col du Dôme ice core 854 summer sulphate record (grey line: actual data; black line: first component of single spectra analysis with a 5 year time window) (Preunker et al., 2001). The dashed vertical lines mark: Tambora (April 855 856 1815), Krakatau (August 1883) and Hekla (March 1947) eruptions, the peak of the S emissions (1976) 857 and the maximum value of the ER78 S peak (1996). The structure of the detrended S concentration 858 identifies two distinctive short-lived peaks attributed to the Tambora and Krakatau eruptions as well as 859 a clear annual structure between 1890 and 1970, also highlighted in the blow-up panel. The red star

- 860 marks the end of the Little Ice Age in the Italian Alps. The cross on the bottom left corner visualises
- 861 the mean error (± 5 years) in the oldest part of the record (1810 1890) based on the original lamina
- 862 counting (Frisia et al., 2003), whereas for the recent part of the record (1970 1996) the estimated
- 863 error is ± 1 year.
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Fig. 8. Measured S concentration in stalagmite ER78 (original data from Frisia et al., 2005, 2008) compared with the dripwater S/Ca ratio (red dots and line). The dashed red line is the exponential regression of the dripwater S/Ca data (R^2 =0.89, *p* < 0.0001). Inset: correlation between sulphur partition coefficient (D_S) and dripwater S/Ca calculated in the overlapping period (1995-1998). The age uncertainty based on lamina counting in this part of the stalagmite is ±1 year (Frisia et al., 2003).



Fig. 9. Sulphur time-series for ER78 stalagmite (grey line actual data, black line 2 years Gaussian filter,
Frisia et al. 2005, 2008) compared with the sulphur dioxide emissions (cf. Fig. 1). For ER78 stalagmite
the theoretical series reconstructed by the measured dripwater S/Ca ratio (black dots and line, cf. Fig.
8) and the projected trend until 2030 (dashed black line) are also shown. The blue dashed lines identify
suggested correlation between the series, while the grey bars at the bottom indicate the reconstructed
hydrological scenario (A to C in Fig. 3) and the corresponding mean delay in the stalagmite S series.



Fig. A1. Comparison of sulphur time-series. The series st-ER77 and st-ER78 have been offset by 4.5
years in order to best match series st-ER1. The offset represents the additional delay related to S
retention in the aquifer matrix with respect to the fast fissure flow. The modelled S emissions (cf. Fig.
and the mean sulphate concentrations in meteoric water for 2005-2006 are also plotted.



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Fig. A2. Calibration of sulphur SR-XRF data (Frisia et al., 2005) on the basis of SIMS analyses (Wynn et al., 2008) for ER78 stalagmite. Measurements were carried out on two physically different slabs and, consequently, we choose three separate parts of the scans in order to avoid the presence of distinctive peaks in the two scans possibly related to grain inclusions. The linear calibration value (XRF cps = 0.187 ± 0.025 ppm) was calculated as the arithmetic mean between values obtained in the three regions highlighted by dashed boxes.