# The Elbrus (Caucasus, Russia) sulfate ice core record: reconstruction of past anthropogenic sulfur emissions in southeastern Europe

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**Abstract.** This study reports on the glaciochemistry of a deep ice core (182 m long) drilled in 2009 at Mount Elbrus in the Caucasus, Russia. Radiocarbon dating of the particulate organic carbon fraction in the ice suggests that the basal ice dates to

- 15  $280 \pm 400$  yr CE (Common Era). Based on chemical stratigraphy, the upper 168.6 m of the core were dated by counting annual layers. The seasonally resolved chemical records cover the years 1774-2009 CE, thus, being useful to reconstruct many aspects of atmospheric pollution in south-eastern Europe from pre-industrial times to present-day. After having examined the extent to which the arrival of large dust plumes originating from Sahara and Middle East modifies the chemical composition of the Elbrus (ELB) snow and ice layers, we focus on the dust-free sulfur pollution. The ELB dust-
- free sulfate levels indicate a six- and seven-fold increase from 1774-1900 to 1980-1995 in winter and summer, respectively. Remaining close to  $55 \pm 10$  ppb during the nineteen century, the annual dust-free sulfate levels started to rise at a mean rate of ~3 ppb per year from 1920 to 1950. The annual increase accelerated between 1950 and 1975 (8 ppb per year), levels reaching a maximum between 1980 and 1990 (376 ± 10 ppb) and subsequently decreasing to  $270 \pm 18$  ppb at the beginning of the twenty first century. Long-term dust-free sulfate trends observed in the ELB ice cores are compared with those
- 25 previously obtained in Alpine and Altai (Siberia) ice, the most important differences consist in a much earlier onset and a more pronounced decrease of the sulfur pollution over the three last decades in western Europe than south-eastern Europe and Siberia.

### **1** Introduction

It is now well recognized that the present climate change is not only related to change of long-lived greenhouses gases but also of short-lived climate forcers, with one of the most important components being aerosol, particularly at regional scales. In this way, it has been suggested that aerosols may have weakened the rate of the global warming during the second part of

- 5 the last century (Andreae et al., 2005). However, uncertainties still exist in quantifying the climatic impact of aerosols. The spatial distribution of aerosols is very heterogeneous and requires therefore numerous observations to make these parameters useful as inputs and constraints for transport and chemistry models. An important gap is also related to the fact that direct atmospheric observations are available only from far later (starting with the appearance of the acid rain phenomena in the late 1960s) than man-made activities started to impact the pre-industrial atmosphere. However, to predict future climate the
- 10 knowledge of atmospheric changes in aerosol load and composition from present-day polluted atmosphere back to preindustrial times is required. Chemical records of species trapped in snow deposited on cold glaciers provide a unique and powerful way to reconstruct past atmospheric chemistry changes including aerosol load and composition (see Legrand and Mayewski (1997) for a review).

In Eurasia, a largely industrialized continent, ice cores were extracted from high-elevation glaciers located at various places,

- 15 including the Alps (Preunkert and Legrand, 2013; Schwikowski et al., 2004), the continental Siberian Altai (Eichler et al., 2009; Eichler et al., 2012; Olivier et al., 2006), and Kamchatka (Kawamura et al., 2012). In the Alps, intimately connected to western European emissions, ice cores have been investigated at Col du Dôme (CDD, Mont Blanc, Preunkert, 2001), Fiescherhorn (Bernese Alps, Jenk, 2006), and Colle Gnifetti (CG) in the Monte Rosa region (Schwikowski, 2006; Wagenbach et al., 2012) in view to examine various aspects of atmospheric pollution. The exceptionally high net snow
- 20 accumulation at the CDD site permitted the extraction of seasonally resolved records of various chemical species over the last 100 years (Preunkert and Legrand, 2013). In older ice layers preservation of winter layers at CDD becomes very limited and summer layers become very thin. Conversely, the low and incomplete net snow accumulation rate at CG, which is controlled by wind erosion, highly limits the preservation of winter ice layers (Wagenbach et al., 2012), but is low enough to provide access to an extended time period, at least over the last millennium. Using the EMEP (European Monitoring and
- 25 Evaluation Programme) regional chemistry-transport model and past emission inventories of SO2 in Europe, observed CDD long-term trends of sulfate were fairly well reproduced, leading Fagerli et al., (2007) to conclude that the seasonal changes seen at the CDD alpine site are associated with geographical changes in source regions impacting the site. This is a strong argument for a separate examination of summer and winter data, extracted from alpine ice cores. However, until now, Alpine ice cores document only the last hundred years (at the best back to 1890, Legrand et al., 2018) on a seasonal basis, whereas
- 30 the early stage of the industrialization time period, which is generally considered to have started around 1850, is missed. An ice core recently extracted from the Elbrus (the highest summit of the Caucasus) indicated excellent preservation of summer and winter layers at least back to 1820 (Mikhalenko et al., 2015). Thus, the ELB ice may contain very valuable information on past atmospheric pollution in south-eastern Europe since the beginning of the industrialization.

Here we report on the glaciochemistry of a deep ice core in 2009 in the Caucasus, Russia. Seasonally resolved chemical records were obtained back to 1774 (i.e., well prior to the onset of the industrial period). Data are discussed in two companion papers of which this one. The present paper examines first of all the impact of large dust plumes, which arrive sporadically from Sahara and Middle East, on the chemical composition of the Elbrus (ELB) snow and ice layers. It then

- 5 focuses on long-term dust-free sulfate trends in relation to growing sulfur pollution. The long-term summer and winter trends of dust-free sulfate are discussed with respect to past SO<sub>2</sub> emissions in south-eastern Europe and compared to those extracted at the Alpine site of CDD and the Siberian Altai (Belukha glacier) in relation to SO<sub>2</sub> emissions from western and eastern Europe, respectively. The second paper focuses on calcium (a dust tracer) long-term trend (Kutuzov et al., this issue), discussing its past changes in relation with natural variability, as well as climatic and land use changes in the dust source
- 10 regions Middle East and North Africa.

#### 2 Methods and Dating

A deep ice core was drilled to bedrock (182.6 m, i.e. 142.1 meter water equivalent (mwe)) in 2009 on the western plateau of Mt. Elbrus (43°21'N, 42°26'E; 5115 m above sea level) in the Caucasus, Russia (Fig. 1). Glaciological settings of the drill site are detailed in Mikhalenko et al. (2015). In brief, the surface of the glacier plateau is about 0.5 km<sup>2</sup>, and the surface snow

- accumulation at the drill site is about 1.5 mwe yr<sup>-1</sup>. Ice-penetrating radar measurements made in 2007 and 2009 revealed a maximum glacier thickness of  $255 \pm 8$  m at the central part of the plateau, and minimum values of ~60 m near the western border of the glacier. Borehole measurements indicated temperatures of -17°C at 10 m depth and -2.4°C at 181.8 m depth. Occasionally melting of surface snow can occur, however, the thickness of the infiltration ice layers, which do not form every year, does not exceed 10 mm. After the overall presentation from Mikhalenko et al. (2015), two other studies of the
- 20 ELB ice core were dedicated to black carbon (Lim et al., 2017) and water stable isotope composition on the 126 m upper layers (Kozachek et al., 2017).

## 2.1 Discrete Subsampling of firn and ice, and Chemical Analysis

- Ice cores were subsampled and decontaminated at -15°C using the electric plane tool methodology described in Preunkert and Legrand (2013). In brief, in a first step ice samples were cut with a band saw. After that, all surfaces of the cut samples were decontaminated by removing ~ 3 mm with a pre-cleaned electric plane tool under a clean air bench. A total of 3724 subsamples were obtained along the upper 168.6 m (131.6 mwe) of the Elbrus core. As expected from glacier ice flow (e.g. Paterson and Waddington, 1984), a decrease of the annual ice thickness with depth is observed (Fig. 2). Annual layer thicknesses decreases from 1.5 mwe (0.8 mwe in summer and 0.7 mwe in winter) near the surface, to 0.46 mwe at ~100 m (i.e. 75 mwe) (0.35 mwe in summer and 0.11 mwe in winter), 0.21 mwe at 155 m (i.e. 120 mwe) (0.12 mwe in summer and
- 0.09 mwe in winter) and 0.18 mwe (0.15 mwe in summer and 0.03 mwe in winter) at 157 m (i.e. 122 mwe) depth. To

minimize the loss of temporal resolution with depth along the core, the sample depth resolution was decreased from 10 cm at the top to 5 cm at 70 m (47 m we), and 2 cm at 157 m (122 mwe) depth and below. In this way, an average of 9 summer samples per year were sampled at 157 m depth (compared to 15 summer samples per year near the surface). The large decrease of the net snow accumulation in winter below 155 m (Fig. 2), likely due to more wind erosion of winter than

- 5 summer snow layers upstream the drill site as already observed at other high altitude glacier sites (e.g. Preunkert et al., 2000), leads to a more pronounced loss of resolution in these winter layers compared to the surface layers (12 samples per winter near the surface and 1-2 samples per winter at 157 m depth).
- For measurements of cations (Na<sup>+</sup>, K<sup>+</sup>, Mg<sup>2+</sup>, Ca<sup>2+</sup>, and NH<sub>4</sub><sup>+</sup>), a Dionex ICS 1000 chromatograph equipped with a CS12 separator column was used. For anions, a Dionex 600 equipped with an AS11 separator column was run with a quaternary gradient of eluents (H<sub>2</sub>O, NaOH at 2.5 and 100 mM, and CH<sub>3</sub>OH). A gradient pump system allows the determination of inorganic species (Cl<sup>-</sup>, NO<sub>3</sub><sup>-</sup>, and SO<sub>4</sub><sup>2-</sup>) as well as short-chain carboxylates. Investigated carboxylates include formate (HCO<sub>2</sub><sup>-</sup>), lactate (CH<sub>3</sub>CHOHCO<sub>2</sub><sup>-</sup>), acetate (CH<sub>3</sub>CO<sub>2</sub><sup>-</sup>), glycolate (CH<sub>2</sub>OHCO<sub>2</sub><sup>-</sup>), and glyoxylate (CHOCO<sub>2</sub><sup>-</sup>), oxalate (C<sub>2</sub>O<sub>4</sub><sup>2-</sup>), malonate (CO<sub>2</sub>CH<sub>2</sub>CO<sub>2</sub><sup>2-</sup>), succinate (CO<sub>2</sub>(CH<sub>2</sub>)<sub>2</sub>CO<sub>2</sub><sup>2-</sup>), and glutarate (CO<sub>2</sub>(CH<sub>2</sub>)<sub>3</sub>CO<sub>2</sub><sup>2-</sup>). Details on working conditions are reported in Legrand et al. (2013). For all investigated ions, blanks of the ice
  decontamination procedure were found to be insignificant with respect to respective levels found in the ice cores.
- During the drill operations, an incident occurred at the depth of 31 m and a fluid (Havoline XLC, Texaco company) was poured in the hole to liberate the drill device. This has led contamination of the firn at 31 m down to the firn-ice transition located at 55.7 m depth. Samples covering the 1983-1997 years were contaminated for sodium ( $124 \pm 87$  ppb compared to  $26 \pm 28$  ppb over the 16 preceding years) and potassium ( $35 \pm 25$  ppb compared to  $16 \pm 15$  ppb over the 1966-1982 years).
- 20 One core section (denoted ELB-140) that covers winter 1875/76, summer 1876, and winter 1876/1877 was not analysed. Finally, a part of the ELB-138 ice core section that covers winter 1877/1878 was of poor quality (splitted ice).

### 2.2 Annual Layer Counting

As discussed by Mikhalenko et al. (2015), dating of the Elbrus ice can be done by annual layer counting on the basis of the stratigraphic ammonium and succinic acid records, both exhibiting well-marked winter minima. As previously seen in alpine

- 25 ice cores, ammonium reveals a well-marked maximum in summer due to a maximum of NH<sub>3</sub> emission together with an efficient upward transport in summer (Maupetit et al., 1995; Preunkert et al., 2000, Eichler et al., 2000). Succinic acid is a light dicarboxylic acid for which a strong summer maximum and a quasi-null winter level can be observed in the present-day atmosphere in Europe (Legrand et al., 2007). The very low winter levels of this organic compound are related to the absence of a winter source of this species, which is mainly photo-chemically produced from biogenic precursors. Mikhalenko et al.
- 30 (2015) assumed a concentration limit of 100 ppb ammonium and 5 ppb of succinate to separate winter and summer in the upper layers down to 75.6 m depth (i.e., 1963). To account for an observed decreasing trend of ammonium concentrations with depth (i.e., due to a post 1950 increase as also seen for species like nitrate and sulfate, see below), the ammonium winter criterion was adjusted to 50 ppb between 75.6 and 86.8 m depth (i.e., 1950-1963) and 30 ppb below. Since no

systematic change of succinate with depth is observed, the succinate concentration limit of 5 ppb was also applied in deeper layers. Requiring that at least one of the criteria is fulfilled and the value of the other is below or close to the limit (< 10% above) the annual counting was found to be very accurate dating (a 1-year uncertainty) over the last hundred years when anchored with the stratigraphy with the Katmai 1912 horizon (Mikhalenko et al., 2015). A good agreement (a 2-year

- 5 discrepancy) was also found when comparing this dating with the chronology achieved by annual layer counting of the water stable <sup>18</sup>O isotope back to 1900 (Kozachek et al., 2017). Though the annual counting becomes less evident prior to 1860, Mikhalenko et al. (2015) reported an ice age of 1825 at 156.6 m depth, what is still consistent with the presence of volcanic horizon at around 1833-1940 such as Coseguina (1835).
- We here extended the annual counting down to 168.5 m depth (i.e., 131.6 mwe) by considering a concentration limit of 30 ppb for ammonium and 5 ppb for succinate. With that, ice dates to 1774 CE at the depth of 168.5 m. In Fig. 3 we report the obtained chronology for three different sequences including the deepest one (1774-1784). It could be seen that in the years prior to 1850, quite often a winter layer is made only of one or two samples, whereas summer layers are made still of more than 6 samples. Below 168.5 m depth, the ice core quality becomes rather bad (numerous small pieces of broken ice) rendering subsampling and ice decontamination not evident. Furthermore, as seen in Fig. 4, in contrary to what is observed
- 15 above 168.5 m, the 30 subsamples obtained along a 1 m long core section at 176.3-177.3 m depth reveal an absence of samples with ammonium and succinate concentrations below the applied winter criterion. This hampered the dating of the basal ice layers of the core by annual counting and therefore another dating approach based on <sup>14</sup>C of particulate organic matter was made for this part of the core.

# 2.3 Basal Ice Dating

- 20 Four ice samples located along the deepest 6 m of the core (bottom at 182.65 m, 142.1 mwe) were analysed for radiocarbon in particulate organic carbon (PO<sup>14</sup>C). The lowest 0.5 m of the core were not analyzed due to a large presence of macro-size inorganic particles. To minimize the time interval covered by each sample, sample lengths were kept as short as possible with respect to the detection limit related to working conditions during sampling and analysis. A typical sample length of 26 to 40 cm (available ice core section of 14-21 cm<sup>2</sup>) was used leading to an initial ice sample mass of 430-580 g. After having
- 25 decontaminated ice sample (using a DOC decontamination method according to Preunkert et al., 2011), melted ice was filtered and combusted at 340°C at the Institut de Géophysique Externe (Grenoble) using the inline filtration-oxidation-unit REFILOX (Reinigungs-Filtrations-Oxidationssystem, Hoffmann et al., 2018). Hereby, the resulting ice mass was reduced to 260-320 g containing 4.4 to 6.5 μgC of POC (Table 1). After cryogenic extraction of the CO<sub>2</sub> content, radiocarbon analyses were done at the accelerator mass spectrometer facility at the Curt-Engelhorn-Center Archaeometry (CEZA) in Mannheim
- equipped with a Gas Interface System (GIS) (Hoffmann et al., 2017). Calibration of the retrieved 14C ages was done using OxCal version 4.3 (Bronk Ramsey, 1995, Bronk Ramsey, 2009).
   To test the reliability of the DOC decontamination method (Preunkert et al., 2011) for POC analysis, a 340 °C REFILOX

mass combustion comparison was made between ultrapure water and decontaminated blank ice (3 samples of 200 to 500

mL). To achieve an impurity-free solid ice the ultra-pure water was slowly frozen in polyethylene (PE) foil (Hoffmann et al., 2018 and references in there). The comparison showed that whatever the blank ice volume, the blank values were in the same range than the ultrapure water POC blanks (~ $0.4 \pm 0.25 \mu gC$ ) which were determined during the course of the ELB radiocarbon sample measurements. Thus, the ice decontamination procedure used for DOC ice measurement is also valid for

5 POC ice decontamination.

Since the CO<sub>2</sub> collection line was recently extended to allow sample pooling, we were now able to directly determine the fraction modern carbon ( $F^{14}C$ ) in blanks done with ultrapure water. A  $F^{14}C$  value of 0.71 ± 0.07 was found, measured on three blank samples in total (each consisting of four pooled samples). This value is in agreement with  $F^{14}C$  blank values found in previous studies (as reviewed and adopted in Hoffmann et al., 2018). In Table 1, we report ice sample data after

- blank correction, including correction of the  $F^{14}C$  value as well as correction of the extracted POC mass with the respective ultrapure water blank determined before each ice sample extraction. Using a mean mass related combustion efficiency of the device of 0.7 (Hoffmann et al., 2018), the mean POC concentrations of the four samples obtained by combustion at 340°C is of 25.4 ± 3.1 ngC g<sup>-1</sup> with highest concentration of 29.0 ngC g<sup>-1</sup> for
- the lowest sample analysed at 141.6 mwe (182.0 m). These values are in good agreement with those observed by Hoffmann
  et al. (2018) in the 340°C POC fraction of the lowest 8 mwe of a CG ice core (37 ± 16 ngC g<sup>-1</sup>). Since a similarity between
  CG and ELB was also observed for their preindustrial black carbon content (Lim et al., 2017), we thus exclude significant age errors due to a POC contamination during <sup>14</sup>C sample preparation and analysis.

As seen in Table 1, the mean age of the ELB-178-03 sample (1530 yr cal BP, i.e. 1530 years before 1950) is older than the mean age of sample ELB-181-01 located 2.3 m below the ELB-178-03 sample. However, given age uncertainties, it is

- 20 difficult to conclude that, as observed previously at other mid-latitude glacier sites such as CG (Hoffmann et al., 2018), the ELB radiocarbon ages do not increase monotonically with depth as would be expected from well-behaved ice flow. PO<sup>14</sup>C measurements suggest that the ELB ice core extends to ~  $1670 \pm 400$  yr cal BP (Table 1). This is younger than basal ice ages found at Alpine sites, i.e. ~5000 yr BP for Col du Dome (Preunkert et al., 2019), ~ 4000 yr BP (Hoffmann et al., 2018) and >10,000 yr BP (Jenk et al., 2009) for two CG ice cores, and ~7000 yr BP for Mt. Ortles (3905 m asl) (Gabrielli et al., 2016).
- From the observed temperature gradient in the borehole ELB site, Mikhalenko et al. (2015) calculated a heat flux at the bottom glacier that is presently 4-5 times larger than the mean value for the Earth's surface, possibly due to a heat magma chamber of the Elbrus volcano, leading to potential basal ice melting when ice thicknesses exceeds 220 m. Though the ice at the drill site and upstream is at present frozen to bedrock, we can not exclude that in the past, assuming a more active heat chamber due to the eruption of  $50 \pm 50$  CE (located 1.6 km away from the Eastern Elbrus plateau), a temporary basal ice
- 30 melting had occurred at the drill site. If so, that may explain the young age of basal ice at the volcanic crater site compared to other non-volcanic mountain glaciers. The age of the basal ELB ice is nevertheless largely greater than expected by ice flow model calculations, estimating a basal ice age of less than 400 years at the drill site (Mikhalenko et al. 2015). Fig. 5 summarizes the extended depth-age relation, including annual layer counting back to 1774 CE, the prominent time horizons, and PO<sup>14</sup>C data. To interpolate data to a continuous age-depth relation, a two-parameter fit (based on a simple analytical

expression for the decrease of the annual layer thickness with depth) was used (Nye, 1963; Jenk et al., 2009; Preunkert et al., 2019).

# 3 The Effect of large Dust events on the Chemistry of ELB Ice

Large dust plumes originating from Middle East and less frequently from Sahara reach the Caucasus (Kutuzov et al., 2013).

- 5 As seen in the Alps, these dust events modify the chemistry of snow deposits, in particular with calcium rich alkaline snow layers (Wagenbach et al., 1996). Depositions of these plumes enhance the level of numerous chemical species in Alpine ice because either they are present in dust at the emission stage or, being acidic, they were uptake by the alkaline dust material during transport (Usher et al., 2003). Preunkert (2001) showed that the arrival of dust plumes at CDD enhanced depositions of several cations (sodium, potassium, magnesium, and sodium) as well as acidic anions (sulfate, nitrate, chloride, fluoride, fluoride,
- and carboxylates). To identify these layers in the ELB snow and ice we have estimated the acidity (or alkalinity) of samples by checking the ionic balance between anions and cations with concentrations expressed in micro-equivalents per liter,  $\mu Eq$  $L^{-1}$ ):

$$[H^{+}] = ([F^{-}] + [Cl^{-}] + [NO_{3}^{-}] + [SO_{4}^{2-}] + [MonoAc^{-}] + [DiAc^{2-}]) - ([Na^{+}] + [K^{+}] + [Mg^{2+}] + [Ca^{2+}] + [NH_{4}^{+}])$$
(1)  
with [MonoAc^{-}] = [HCO\_{2}^{-}] + [CH\_{3}CHOHCO\_{2}^{-}] + [CH\_{3}CO\_{2}^{-}] + [CH\_{2}OHCO\_{2}^{-}] + [CHOCO\_{2}^{-}] (2)

15 and 
$$[DiAc^{2-}] = [C_2O_4^{2-}] + [CO_2CH_2CHOHCO_2^{2-}] + [CO_2CH_2CO_2^{2-}] + [CO_2(CH_2)_2CO_2^{2-}] + [CO_2(CH_2)_3CO_2^{2-}]$$
 (3)

In this work, samples that are below the 25% quartile of a robust spline through the calculated raw acidity profile and contain more than 120 ppb of calcium were considered as impacted by dust events. In this way, 616 (on a total of 2524) summer and 67 (on a total of 1150) winter samples were considered as containing large amount of dust. Note that the results are quite

- similar when changing the calcium concentration criteria from 120 ppb to 100 or 140 ppb. Since the frequency of these events has changed over time (Kutuzov et al., this issue) the significance of their impact on the deposition of chemical species is examined. Figure 6 shows the mean ionic budget of samples considered as contaminated by dust and containing high and moderate content of calcium compared to samples assumed to be free of dust events, over the 1950-1980 period. For the largest events ( $Ca^{2+} > 30 \mu Eq L^{-1}$ , Fig. 6A), the increase of calcium, accompanied by a strong increase of the
- 25 alkalinity, reaches a factor of 7.4 compared to dust-free samples (Fig. 6C). In addition, this calcium enhancement is accompanied by an increase of a factor close to 8 for chloride, sodium, potassium and magnesium, whereas ammonium, nitrate, sulfate, and carboxylates are, at the most, enhanced by a factor of 2. When comparing dust samples containing weaker calcium contents (i.e.  $Ca^{2+} < 30 \mu Eq L^{-1}$ , Fig. 6B) with dust-free samples (Fig. 6C), in addition to the increase of calcium (factor of 2) the most significant changes are seen for magnesium (x1.5), sodium and chloride (x1.6), and potassium

30 (x1.3), respectively.

To evaluate the effect of dust events on the deposition of chemical species, we compare in Table 2 averaged chemical concentrations of all samples with those not impacted by dust (denoted  $X_{red}$ ). Averages were obtained on the base of half-

year summers over the half-decades 1996-2000 and 1974-1978 characterized by high and low dust content, respectively (Kutuzov et al., this issue). Though the main impact of dust is as expected, on cations (except ammonium) and chloride, i.e. the constituents of dust particles, the impact is also significant for acidic species like nitrate and sulfate. For instance the increase of nitrate from 1974-1978 to 1996-2000 AD (144 ppb) in the whole dataset is largely (three quarters) related to the

- 5 increase of dust as indicated by the smaller increase after removal of dust samples (increase of 31 ppb for the  $NO_{3}^{-}$  value). Same is true for sulfate, for which the apparent increase between the two periods (90 ppb) is not due to an increase of pollution but of dust, as indicated by the drop of values when the dust free data set is considered (decrease of 48 ppb for the  $SO_{4}^{-2}$  value). The effect of changing dust inputs over time has to be therefore considered when discussing long-term trends in view to relate them to growing anthropogenic emissions (see Sect. 5 for sulfate in relation with SO<sub>2</sub> emissions). Finally,
- 10 the large effect of dust seen for formate (HCOO<sup>-</sup>) and not for acetate (CH<sub>3</sub>COO<sup>-</sup>) is in agreement with previous observations made by Legrand et al. (2003) in Alpine ice and by (Legrand and De Angelis, 1995) in Greenland ice. These studies showed that the presence of formate and acetate in ice follows the uptake of formic and acetic acid from the atmospheric gas phase, and that the incorporation of these weak acids into hydrometeors is pH dependent with a stronger dependence for formic acid, which is a stronger acid than acetic acid. In conclusion, large dust events significantly influence the chemistry of ELB
- 15 ice for many species, requiring a case by case examination depending on the nature of the dust contribution: primary emissions for sodium and other cations (except ammonium), neutralization of the alkaline material during atmospheric transport for nitrate, and both primary gypsum emissions and neutralization of the alkaline material by acidic sulfur during atmospheric transport for sulfate, for instance.

#### 4 Long-term summer and winter trends of sulfate in the Elbrus ice

- A few outliers of unknown origin were observed in the sulfate raw data set including 22 ppm at 166.65 m depth (summer 1780 AD), 2248 ppb at 146.38 m depth (summer 1862 AD), 1080 ppb at 146.11 m depth (summer 1863 AD), and 864 ppb at 154.73 m depth (winter 1833/34 AD) (Fig. 7). These individual values were removed when calculating the corresponding half-year summer and winter means reported in Fig. 8. In addition, single winter samples with sulfate levels of 815 ppb at 160.62 m depth (winter 1810/11 AD) and 3 data points from 150 to 230 ppb corresponding to winters 1786-87, 1827-28, and 1844-45 were not considered and corresponding half-year winter values were not reported.
- A few ELB snow and ice layers are impacted by known volcanic eruptions. As discussed by Mikhalenko et al. (2015), ice layers dated to 1911 and 1913 were probably impacted by the 1912 AD Katmai eruption, and summer layers of 1836 and 1837 by the 1835 AD Coseguina eruption, respectively. In addition, we suspect the 1854 AD Shiveluch eruption to have impacted summer 1854 ice layer and finally, although less evident since this part of the core is made up of splitted ice (see
- 30 Sect. 2), the Cotopaxi 1877 AD eruption may have influenced the winter 1877/1878 layer. To discuss the long-term trends of sulfate in relation to growing SO<sub>2</sub> emissions, these half-year summer and winters means suspected to be contain volcanic debris, were discarded in Fig. 9. To minimize the effect of year-to-year variability due to meteorological transport conditions

we added the first component of single spectra analysis (SSA) with a five-year time window in Fig. 9.

As discussed in Sect. 3, large dust events significantly enhanced the sulfate level of the ELB ice. Since, as detailed in Kutuzov et al. (this issue), their occurrence have changed over the past with more frequent events after 1950, we have examined to what extent they contribute to the sulfate trend. It is difficult to accurately directly correct sulfate concentration

- 5 from the large dust event contribution since the amount of sulfur trapped by the alkaline material during atmospheric transport towards the site would be very different from event to event. For instance, Kozak et al. (2012) reported non-sea-salt-sulfate to non-sea-salt calcium mass ratios in aerosol collected in the eastern Mediterranean ranging between 0.25 (in case of direct arrival at the site of air mass from the Sahara) to 1.15 (when mineral dust passed through polluted sites located in the Balkans and Turkey before arriving at the site). Instead of corrected sulfate values for the large dust events, we
- 10 therefore have reported in Fig. 8 individual values of total sulfate and sulfate calculated after having removed from the average samples suspected to contain large amount of dust (SO<sub>4</sub><sup>2-</sup><sub>red.</sub> values). The influence of large dust events on the long-term SSA winter trend is rather insignificant and if existing (i.e. effect of < 10 ppb) remaining limited to two decades around 1870 and the recent decade (2000-2010) (Fig. 9). Remaining negligible prior to 1850, the large dust event effect on the summer trend gradually increases after 1950, reaching often 100 ppb after 1960. This change of large dust events results 15 from change in the occurrence of drought in North Africa and Middle East regions (Kutuzov et al., this issue).</p>
- In addition to the enhanced frequency of large dust events after 1950, the calcium background concentrations (i.e.,  $Ca^{2+}_{red.}$ ) also changed over time with an increase from  $68 \pm 21$  ppb prior to 1900 CE to  $194 \pm 61$  ppb after 1960 CE. As discussed by Kutuzov et al. (this issue), this change may result from changes in precipitation and soil moisture content in Levant region (Syria and Iraq). In view to discuss the free-dust sulfate changes with respect to anthropogenic emissions, we make an
- attempt to correct the sulfate record from this increase of the background level of dust. To do so, we examined in Fig. 10 the relationship between calcium and sulfate concentration in individual summer samples corresponding to pre-industrial time (1774-1900 CE). Although being poor ( $R^2=0.32$ ), the correlation suggests a mean slope of the linear SO<sub>4</sub>red-Cared relationship close to 1. The use of this value to correct sulfate from background dust emissions would lead to an overestimation of the sulfate dust contribution. Indeed, as seen in Fig. 10, there are numerous samples that contain more
- sulfate than what is expected with respect to the presence of pure calcium sulfate (gypsum, see the blue line reported in Fig. 10), likely to due the presence of sulfate as ammonium sulfate or sulfuric acid. To correct sulfate from the background dust contribution we here have used a sulfate to calcium ratio close to 0.63 (see the red line drawn as the lower envelope of the relationship in Fig. 10) and subtracted this contribution from the  $SO_4^{2^2}$  red. values by using the  $Ca^{2^+}_{red}$  values. As seen in Fig. 9, the derived mean summer and winter pre-industrial free-dust sulfate levels in ELB ice (taken as the mean value observed
- from 1774 to 1850) is of 70 ppb and 40 ppb, respectively. In summer as in winter, the free-dust sulfate values remained close to the pre-industrial values until 1910-1920 (81 ppb instead of 70 ppb in summer, 47 ppb in winter). After 1920, dust-free sulfate levels increased at a mean rate of 5 ppb per year in summer, and 1 ppb yr<sup>-1</sup> in winter. The sulfate increase then accelerated between 1950 and 1975 (10 ppb yr<sup>-1</sup> in summer, 5 ppb yr<sup>-1</sup> in winter), until a maximum of 530 ppb in summer (255 ppb in winter) was reached at the end of the 80's. After 1990, sulfate levels decreased to 390 ppb in summer (154 ppb
  - 9

in winter) during the first decade of the twenty first century.

#### 5 Comparison between Elbrus, Alpine, and Siberian Altai long-term sulfate trends

# 5.1 The Alpine CDD and Siberian Altai (Belukha, BEL) ice core Sulfate Records

The ELB dust-free sulfate long-term trend is compared with those previously extracted from the Alpine CDD site (ice cores denoted C10 and CDK in Fig. 11). C10 sulfate data were presented in Preunkert et al. (2001), and those form CDK in (Legrand et al., 2013). Since winter data from CDD are more limited (only a few pure winter layers are available between 1890 and 1930, Legrand et al., (2018) we here focus on the comparison of summer levels. The two CDD cores were dated by annual layer counting using the pronounced seasonal variations of ammonium. The two chronologies were in excellent agreement over their overlapping period from 1925-1990 (Legrand et al., 2013; Preunkert et al., 2000). A re-evaluation of

- 10 the C10 chronology based on very recently made continuous measurements of heavy metals, as well as a comparison to a well-dated Greenland ice core record (McConnell and Edwards, 2008), resulted in a revised C10 chronology (Legrand et al., 2018). As for C10, continuous measurements of heavy metals are also available in the lowest part of CDK (Preunkert et al., 2019). It was thus possible to identify the distinct Greenland increases of thallium, lead, and cadmium associated with the widespread start of coal burning at the beginning of the Industrial Revolution in 1890 CE also in the CDK core (at 117.8 m
- 15 (90.5 mwe)). This time marker was then used to constrain a revised annual layer counting in the early 20<sup>th</sup>-century part of the CDK record. The dust-free ELB sulfate long-term trend is also compared with the one previously extracted from the Siberian Altai (Belukha glacier, denoted BEL in Fig. 11) by Eichler et al. (2009 and 2012).

## 5.2 ELB versus CDD and BEL sulfate trends

Since available at all three sites, we compared in Fig. 11 the annual long-term trends of dust-free sulfate from ELB, CDD,

- 20 and BEL. The ELB and CDD annual values were calculated as arithmetic mean from 5 yr-SSA winter and summer records, whereas the BEL annual data refer to 5 year averaged raw data. Examination of the three annual records reveal three major differences between the three sites: (1) an impact of anthropogenic emissions already significant in 1910-1930 at CDD but neither at ELB nor at BEL, (2) a maximum of the anthropogenic perturbation which is reached in 1970-1980 at CDD and later at the two other sites (10 years after at ELB, and a few years after at BEL), and (3) a far less pronounced re-decrease at
- 25 the beginning of the 21<sup>th</sup> century at ELB compared to CDD. The re-decrease of sulfate over the very recent decades is somewhat stronger at BEL than at ELB.

Using data from Smith et al. (2011), available at <a href="http://sedac.ciesin.columbia.edu/data/set/haso2-anthro-sulfur-dioxide-emissions-1850-2005-v2-86">http://sedac.ciesin.columbia.edu/data/set/haso2-anthro-sulfur-dioxide-emissions-1850-2005-v2-86</a>, we report in Fig. 12 emissions of SO<sub>2</sub> from countries located nearby the ELB site: Georgia, Azerbaijan, Syria, Irak, Turkey, Russian, Iran or located further north (Ukraine) and west (Bulgaria). In these countries SO<sub>2</sub> emissions became significant after 1930 and reached maximum in the late 80's or later (for Turkey and Iran). This feature

clearly differs from the situation at CDD where emissions from countries located around the site (France, Italy, Spain,



Switzerland and Germany) were already significant in 1930 and exhibited a maximum between the early 70's and the early 80's (Fig. 10). For BEL, Eichler et al. (2009) demonstrated the importance of emissions from eastern Europe for the dust-free sulfate annual record at that site.

- Finally, we tentatively examine the cause of the recent decrease of sulfate, focusing on the summer season for which the most relevant source regions are limited to countries located nearby the site. As discussed by Kutuzov et al. (this issue), 10 day backward air mass trajectories calculated for the ELB site using the NOAA HYSPLIT-4 model suggest that, in summer, air masses arriving at ELB mainly originate from the nearby Georgia, Azerbaijan, Syria, Irak, and from Turkey, South Russian, and North of Iran. As previously discussed by Fagerli et al. (2007), the CDD site in summer is mainly influenced by emissions from France, western Germany, Italy and Spain. Consistently with SO<sub>2</sub> emission changes, the recent sulfate
- 10 decrease is more pronounced at CDD than ELB with a recovered 2005 level (316 ppb) close to the 1950 one (296 ppb) (Fig. 13). At the ELB site this is not the case, here the 2005 level (380 ppb in 2005) is found to be still almost two times higher than the one of 1950 (227 ppb in 1950). An intermediate pattern is seen at BEL, likely due to a weak impact of countries like Turkey that significantly contribute to the ELB record but not the BEL one, and a more strong contribution of emissions from Russian at BEL (see also Eichler et al., 2009) than at ELB.
- 15

## 6. Conclusions

Based on the ammonium and succinate stratigraphy, the upper 168.6 m of the deep ice core extracted at Mt Elbrus (Caucasus) in 2009 were dated by counting annual layers back to 1774 CE. The derived seasonally resolved chemical records cover the years 1774-2009 making this ice core particularly useful to reconstruct many aspects of atmospheric pollution in south-eastern Europe from pre-industrial times (1850 CE) to present-day. Below 169 m depth the annual

- 20 pollution in south-eastern Europe from pre-industrial times (1850 CE) to present-day. Below 169 m depth the annual counting is not possible but radiocarbon analysis of the particulate organic carbon fraction in the basal ice of the glacier suggests an age of  $\sim$ 1670 ± 400 cal yr BP. We have examined the impact on the chemical composition of the Elbrus ice layers of arrival at the site of large dust plumes originating from Sahara and Middle East. We then report on the dust-free sulfate records. The ELB dust-free sulfate record indicates a six- and seven-fold increase from prior to 1900 to 1980-1995 in
- winter and summer, respectively. Still moderate at the beginning of the  $20^{th}$  century, the sulfate increase accelerated after 1950, dust-free annual levels reaching a maximum in 1980-1990 ( $376 \pm 10$  ppb) and subsequently decreasing to  $270 \pm 18$ ppb at the beginning of the  $21^{th}$  century. These long-term sulfate changes observed in the ELB ice cores are compared with those previously obtained in Alpine and Siberian Altai ice. Consistently with past SO<sub>2</sub> emission inventories, a much earlier onset and a more pronounced decrease of the sulfur pollution over the three last decades are observed in western than south-
- 30 eastern and eastern Europe.

# Data availability

Sulfate and calcium data can be made available for scientific purposes upon request to the authors (contact: suzanne.Preunkert@univ-grenoble-alpes.fr or michel.legrand@univ-grenoble-alpes).

# Author contributions

5 S. Preunkert and M. Legrand performed research, analyzed ice samples and data, and wrote the original manuscript. S. Kutuzov performed research, analyzed data, and commented original manuscript. P. Ginot and V. Mikhalenko performed analysis and commented original manuscript. R. Friedrich analyzed ice samples and commented original manuscript.

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15 manuscript.

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**Table 1.** Overview of masses (corrected for blanks but not for combustion efficiency) and conventional  ${}^{14}$ C ages of the Elbrus ice core samples combusted in the REFILOX system. Calibrated date ranges are shown at 68.2% confidence level and are rounded according to (Millard, 2014).

<sup>14</sup> C sampl e name	Depth	Ice mass [g]	POC mass correcte d [µgC]	<sup>14</sup> C correcte d [F <sup>14</sup> C]	Calibrate d 14C date BCE/CE at 68.2%	Calibrated <sup>14</sup> C-age* range at 68.2% [yr cal BP]	Calibrated <sup>14</sup> C-age* [yr cal BP] mean
ELB- 176- 03	$177.11 \pm 0.22 \text{ m}$ (137.89 ± 0.18 mwe)	295	4.5±0.5	0.914± 0.043	670 CE – 1245 CE	1280 - 705	1040
ELB- 178- 03	$179.19 \pm 0.14 \text{ m}$ (139.59 ± 0.12 mwe)	300	5.6±0.5	$0.955 \pm 0.098$	130 CE – 770 CE	1820 - 1180	1530
ELB- 181- 01	$181.50 \pm 0.13 \text{ m}$ (141.19 ± 0.11 mwe)	260	4.4±0.5	0.932± 0.020	440 CE – 1290 CE	1510 - 660	1110
ELB- 181- 03	$182.02 \pm 0.13 \text{ m}$ (141.62 ± 0.11 mwe)	320	6.5±0.5	0.875± 0.021	90 BCE - 680 CE	2040 - 1270	1670

\* age before 1950 CE

**Table 2.** Mean chemical composition of snow layers deposited in periods characterized by low and high dust inputs (1974-1978 and 1996-2000, respectively). Values in parenthesis are mean values calculated after removal of samples containing dust (denoted red.).  $\Delta$  refer to the difference between total and dust reduced (red.) values (i.e., [X] - [X<sub>red.</sub>] for the X species).

	1974-1978 AD	1996-2000 AD
$Ca^{2+}(Ca^{2+}_{red.})$	286 ppb (169 ppb) $\Delta = 117 \text{ ppb}$	544 ppb (186 ppb) $\Delta = 358$ ppb
$Mg^{2+}(Mg^{2+}_{red.})$	25.0 ppb (19.5 ppb) $\Delta = 5.5 \text{ ppb}$	29.5 ppb (18.0 ppb) $\Delta = 11.5$ ppb
$K^{+}(K^{+}_{red.})$	24 ppb (20 ppb) $\Delta = 4$ ppb	$30 \text{ ppb } (20 \text{ ppb})$ $\Delta = 10 \text{ ppb}$
$Na^+ (Na^+_{red.})$	31 ppb (25 ppb) $\Delta = 6 ppb$	$\begin{array}{l} 40 \text{ ppb } (24 \text{ ppb}) \\ \Delta = 16 \text{ ppb} \end{array}$
Cl <sup>-</sup> (Cl <sup>-</sup> red.)	67 ppb (55 ppb) $\Delta = 12$ ppb	88 ppb (46 ppb) $\Delta = 22 \text{ ppb}$
$\mathrm{NH_4}^+(\mathrm{NH_4}^+_{\mathrm{red.}})$	177 ppb (165 ppb) $\Delta = 12$ ppb	199 ppb (149 ppb) $\Delta = 50 \text{ ppb}$
$NO_3^-$ ( $NO_3^-$ red.)	292 ppb (279 ppb) $\Delta = 13$ ppb	436 ppb (310 ppb) $\Delta = 126$ ppb
$SO_4^{2-}(SO_4^{2-}_{red.})$	649 ppb (598 ppb) $\Delta = 51$ ppb	738 ppb (550 ppb) $\Delta = 188$ ppb
HCOO <sup>-</sup> (HCOO <sup>-</sup> <sub>red.</sub> )	168 ppb (135 ppb) $\Delta = 33$ ppb	184 ppb (113 ppb) $\Delta = 71$ ppb
CH <sub>3</sub> COO <sup>-</sup> (CH <sub>3</sub> COO <sup>-</sup> <sub>red.</sub> )	$\begin{array}{c} 48 \text{ ppb } (37 \text{ ppb}) \\ \Delta = 11 \text{ ppb} \end{array}$	$\begin{array}{l} 49 \text{ ppb } (39 \text{ ppb}) \\ \Delta = 10 \text{ ppb} \end{array}$



**Figure 1.** Map showing the location of the Elbrus site (red star) and dust sources (orange shading, based on Kutuzov et al. (this issue) and references therein).



Figure 2. (a) and (b): Mean summer (a) and winter (b) half-year ice thickness along the Elbrus deep ice core. (c) Mean annual ice thickness and (d) summer to winter ratio of ice thickness.



Figure 3. ELB ice chronology at depth intervals of 166.2 to 168.5 m (top), 154.4 to 156.5 m (mid), and from 99.8 to 107.3 m (bottom), based on the ammonium and succinate stratigraphy. Vertical red lines denote yearly dissection based on identification of winter layers (see Sect. 2.2). For the two oldest time-periods (1773-1782 and 1850-1859), each sample was 2 cm long whereas for the most recent time period (1925-1934) one sample was on average 4 cm long. Note that, though being not coherent with the intra seasonal precipitation distribution (see Kozachek et al., 2017), we here assumed that the accumulation is equally distributed within summer and winter seasons.



**Figure 4.** Distribution of succinate and ammonium concentrations observed in the deepest ice layers for which annual counting was possible (i.e., above 168.5 m depth, top) and 10 m below (bottom). The vertical red dashed bars denote the values of the winter criteria.



**Figure 5.** Depth (in mwe) age-relation of the ELB ice core derived from annual layer counting, prominent time horizons, and mean blank corrected and calibrated  $PO^{14}C$  data with  $1\sigma$  age ranges. To interpolate data to a continuous age-depth relation a two-parameter fit was used following Nye (1963).



**Figure 6**. Mean ionic content of ELB layers deposited between 1950 and 1980. A and B: Mean composition of dust event samples containing more and less than 600 ppb (i.e.,  $30 \mu Eq L^{-1}$ ) of calcium. B and C: Mean composition of dust event samples containing less than 600 ppb of calcium compared to samples free of dust. Abbreviations Monoac. and Diac. stand for C<sub>1</sub>-C<sub>3</sub> monocarboxylates and C<sub>2</sub>-C<sub>5</sub> dicarboxylates, respectively (see Eqs. 2 and 3 in Sect. 3).



Figure 7. Raw sulfate data. The red dots denote the outliers, which were removed prior to calculation of the half-year summer and winter means. We also indicate samples possibly impacted by the volcanic eruptions of Cotopaxi and Shiveluch (see text).



Figure 8. Individual summer (red) and winter (blue) half-year means of sulfate along the ELB ice core. Solid circles refer to values calculated considering all samples. Open circle data (SO4 red.) were calculated after having removed samples considered to be impacted
 by large dust events (see Sect. 3).



Figure 9. ELB half-year summer and winter sulfate trends from 1774 to 2010 (the first SSA component with a five-year time window, see text). The black lines refer to raw sulfate values, the dashed lines to sulfate means calculated after removal of dust samples (SO4<sub>red</sub>.), the solid red and blue lines to sulfate values after having corrected SO4<sub>red</sub> values from the background dust contribution (see Sect. 3). Samples suspected to be impacted by volcanoes were removed.



**Figure 10.** Sulfate versus calcium concentrations in summer samples free of large dust events (Red. values) in ice deposited prior to 1900. The blue line refers to a pure gypsum composition. The red line illustrates a lower envelope of the sulfate to calcium relationship.



5 Figure 11. Dust-free sulfate (annual values in black, summer in red, and winter in blue) trends at the CDD (a), ELB (b), and BEL (c) sites. For CDD we report the records derived from the C10 (dashed lines) and CDK (solid lines) ice cores (see Sect. 5.1). Green areas indicate the decades in which the sulfate levels reached their maxima.



**Figure 12.** SO<sub>2</sub> emissions (from 1850 to 2005) from various countries located around the Caucasus (top) and the Alps (bottom). 10 GEO&AZE&IRK&SYR denotes emissions from Georgia, Azerbaijan, Irak, and Syria. Data are from Smith et al. (2011).



Figure 13. Comparison of the dust-free sulfate summer trends (SO42-red. values) from two CDD ice cores (C10 and CDK) (a) and the
ELB ice core (b) with SO2 emissions from countries suspected to contribute to sulfate depositions at the two sites (c for CDD, d for ELB, see discussions in Sect. 5.2). (a) and (b): solid lines refer to the first SSA component with a five-year time window. Horizontal dashed blue lines refer to the respective pre-industrial sulfate levels. (c): SO2 emissions from France, Italy, Spain, Switzerland and half from Germany. (d): SO2 emissions from Turkey, Georgia and Azerbaijan, half from Russia, a quarter from Ukraine, Bulgaria, and Iran. Green areas indicate the decades in which the sulfate levels reached their maxima.