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Water-soluble organic carbon in snow and ice deposited at Alpine, Greenland, and Antarctic sites: a critical review of available data and their atmospheric relevance

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Abstract

While it is now recognized that organic matter dominates the present-day atmospheric aerosol load over continents, its sources remain poorly known. The studies of organic species or organic fractions trapped in ice cores may help to overcome this lack of knowledge. Available data on the dissolved (or total) organic carbon (DOC or TOC) content of snow and ice often appear largely inconsistent and until now no critical review was conducted to understand the causes of these inconsistencies. To draw a more consistent picture of the organic carbon amount present in solid precipitation that accumulates on cold glaciers, we here review available data and, when needed, complete the data set with analyses of selected samples. The different data sets are then discussed by considering the age (modern versus pre-industrial, Holocene versus last glacial maximum) and type (surface snow, firn, or ice) of investigated samples, the deployed method (DOC, TOC) and the applied contamination control. Finally, the OC levels of Antarctic, Greenland and Alpine ice cores are compared and discussed with respect to natural (biomass burning, vegetation emissions) and anthropogenic source (fossil fuel combustion) contributions to atmospheric OC aerosol.

1 Introduction

Although the carbonaceous matter that includes elemental carbon (EC) and organic carbon (OC) is recognized as a major constituent of present-day atmospheric aerosol over continents (Putaud et al., 2004; Legrand and Puxbaum, 2007; Jimenez et al., 2009), there are still major gaps with respect to its sources. In the present-day continental atmosphere, on a mass carbon basis, the elemental carbon (or black carbon, BC) fraction accounts only for 15% of carbonaceous aerosol (see Pio et al., 2007 for Europe), while the dominant fraction consists of organic carbon (OC). Whereas it is rather well established that secondary productions dominate primary emissions of organic carbon (OC) aerosol, the chemical nature of gaseous precursors

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Antarctic plateau are compared to corresponding data already available from central Greenland or from a high-elevated Alpine site.

2 Experimental section

2.1 Organic carbon analysis

5 The device we used in this study to quantify the organic carbon content in snow and ice samples is based on the commercial Phoenix 8000 system (Teledyne Tekmar company), which was modified to allow precise analysis of small sample volumes of ~5 mL (Preunkert et al., 2011). Given the fact that inconsistencies related to the amount of organic carbon present in snow and ice may partly be induced by differences in the employed method, we here concisely detail our working conditions. The liquid sample is first introduced into an inorganic carbon sparger cell together with phosphoric acid and sparged to drive off any inorganic carbon (dissolved CO₂, HCO₃⁻, and CO₃²⁻). Next, the liquid sample is transferred from the inorganic carbon cell to a UV reactor cell where an UV lamp together with an oxidant oxidizes the organic carbon into CO₂. 10 As discussed by Preunkert et al. (2011), to reach blank values as low as possible, no addition of oxidant solution in the UV reactor is done and instead synthetic air is used as carrier gas from which O₂ increases the oxidation efficiency. The hereby produced CO₂ within the UV reactor is swept together with the carrier gas into a Non-Dispersive Infrared (NDIR) detector. 15 Over the course of a day of sample analysis, 3–5 blank measurements were made showing a typical mean ($\pm 1\sigma$) of 25 ± 1.5 ppbC. The respective daily mean blank value is subtracted from the sample values. Taken as twice the variability of the blanks made over a day, the detection limit is about 3 to 5 ppbC. The typical uncertainty of our method, calculated by considering the uncertainty on the determination of the slope 20 of the calibration curve and the variability of the blank value, is of 4% for samples 25

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containing more than 50 ppbC. Due to the blank variability the uncertainty reaches 35% at the 5 ppbC level.

Under working conditions applied in this study, the efficiency of the oxidation relative to the potassium hydrogen phthalate was found to range from 85 to 125% for different water-soluble organic molecule standards (see details in Preunkert et al., 2011). That is true for organic species present at significant levels in natural ice such as short chain mono- and dicarboxylic acids and formaldehyde, as well as for a few other compounds whose presence in ice is expected but not yet quantified (i.e., methanol, and C2–C3 aldehydes as glyoxal, methyl glyoxal and glycolaldehyde). For long chain polyacids that are also present in snow such as HULIS, the efficiency of the oxidation was found to be slightly lower, i.e. $80 \pm 2\%$. The extent to which this method oxidizes the water insoluble organic material that may be present in snow and ice will be discussed in Sect. 3.2.1. 10

2.2 Ancillary chemical analysis

Most snow and ice samples in which the organic carbon content was measured in this study were also investigated for ions. For cations (Na⁺, K⁺, Mg²⁺, Ca²⁺, and NH₄⁺), a Dionex ICS-1000 chromatograph equipped with CG12 and CS12 separator columns was used. For anions, a Dionex DX-600 equipped with AG11 and AS11 separator columns was run with a quaternary gradient of eluents (H₂O, NaOH at 2.5 and 100 mM, and CH₃OH). A gradient pump system allows the determination of inorganic species 15 (F⁻, Cl⁻, NO₃⁻, and SO₄²⁻) as well as short-chain carboxylates. Investigated carboxylates including formate (HCO₂⁻), lactate (CH₃CHOHCO₂⁻), acetate (CH₃CO₂⁻), glycolate (CH₂OHCO₂⁻), and glyoxylate (CHOCO₂⁻), oxalate (C₂O₄²⁻), malate (CO₂CH₂CHOHCO₂²⁻), malonate (CO₂CH₂CO₂²⁻), succinate (CO₂(CH₂)₂CO₂²⁻), and glutarate (CO₂(CH₂)₃CO₂²⁻). The detection limits for major ions like ammonium, nitrate, and sulfate are close to 1 ppb, those for most carboxylates close to 0.1 ppb. 20

In addition to short chain carboxylates, other potential contributors to the OC content of solid precipitation like short chain aldehydes were investigated in some samples.

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First, HCHO was measured in a few snow pit samples collected at Concordia and in the Alps as well as in a few ice core sections from the Greenland NEEM site. Measurements were done with a fluorimetric method that is based on the liquid phase reaction of formaldehyde with acetyl acetone and a $\text{CH}_3\text{COONH}_4\text{-CH}_3\text{COOH}$ mixture (Hantzsch reagent). The deployed analyzer (Aerolaser AL4021) has a detection limit of 0.1 ppb (Legrand et al., 2007a). A few measurements of some other carbonyl compounds including acetaldehyde, propionaldehyde and acetone were achieved in Alpine surface snow as well as in Alpine ice covering the 1940s (Houdier et al., 2000). Hereby, only acetaldehyde was detected at a significant level (~ 4 ppb, i.e. 2 ppbC).

Water-soluble HULIS were investigated in several snow pit samples collected at Concordia and in the Alps. HULIS were extracted with a DEAE (diethylaminoethyl) resin and the quantification of their organic carbon content was made with a Shimadzu TOC analyzer (Guilhermet et al., 2013).

2.3 Sampling and sub-sampling of snow and ice

The OC content was investigated in various snow and ice samples collected on polar ice sheets (Antarctica and Greenland) as well as at the high-elevated Alpine site of the Col du Dôme (4250 m a.s.l., Mont Blanc Massif, French Alps). It has to be emphasized that, the measurement of OC, which is present at trace levels in precipitation of remote regions, is very sensitive to contamination from numerous organic gases present in the ambient air of laboratories. Thus the DOC (or TOC) content of snow deposits is documented until now only in snow samples collected under stringent conditions nearby the surface or in snow pits but not in firn cores. Note that this contamination problem does not hamper measurements of the water insoluble part of OC (WinOC) retained on quartz filters by filtering melted snow and ice samples (Legrand et al., 2007a).

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2.3.1 Surface snow and snow pits

With only two studies conducted in Antarctica (at the South Pole and coastal areas, Table 1) and two studies performed at Summit in central Greenland (Table 2) the OC content of the surface snow pack remains poorly documented (see Sect. 3). With levels reaching several hundreds of ppbC, the Antarctic snow (both coastal and inland) appears unexpectedly rich in OC with respect to the one of central Greenland. Furthermore, the two central Greenland studies show differences by a factor of 4 in the OC abundance of surface snow (Table 2).

With the aim to enlarge this rather inconsistent data set, surface snow and snow pit samples were collected at Concordia located on the high East Antarctic plateau and at the Col du Dôme in the Mt Blanc massif. At Concordia, surface snow (0–5 and 5–10 cm depth) was collected in January 2010. Furthermore, twenty-four snow samples were collected in a 1 m deep snow pit dug in January 2012 at Concordia (Table 1).

At the Col du Dôme, the surface snow layer corresponding to a 3 day old snow precipitation event (having occurred 14 September 2012) was sampled (4 samples between 0 and 20 cm depth), together with the upper most 1.5 m of the snowpack permitting to document snow deposited during the entire summer 2012 as well as the winter 2011/2012 (18 samples). Finally, 4 surface summer snow samples were collected in June 2011 (Table 2).

All samples were collected in airtight glass bottles (Schott, Mainz, Germany) of different sizes (from 100 mL to 5 L). Indeed, whereas 100 mL bottles are adequate for OC sampling even for samples containing less than 5 ppbC of OC, a volume of 500 mL of melted snow is needed to reach a HULIS detection limit of 1 ppbC as required for Antarctic snow and for Alpine winter snow layers. Glass bottles were cleaned by rinsing them several times with ultrapure water (Purelab Ultra MK2 analytic system commercialized by ELGA LabWater/VWS (UK) Ltd, organic content of less than 2 ppbC) a few hours to days before sampling, and then tightly closed until sampling. At Concordia, snow samplings were conducted in the clean air sector located at a few hundred

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meters upwind from the main station. When needed, the snow sampling was done with a glass scraper that was previously washed with ultrapure ELGA water.

Blank measurements of the glass bottle cleaning and storage procedure were made by analysing the OC content of an ultrapure water volume of about 20 % the volume of bottle (i.e. similar to the minimum sample liquid volume collected by filling completely
5 bottle with snow) poured in the respective bottles. The blank value was found to be close to the detection limit of 3–5 ppbC when blank measurements were performed no later than a few hours after the cleaning. Tests made on blank bottles which were put in sealed plastic bags together with those used to collect snow, revealed however that
10 during the storage time between cleaning, sampling and analysis, contamination took place. A typical bottle blank value of 30–50 ppbC was observed when glass bottles were stored for two days in the lab after cleaning. Such a contamination over time of airtight glass bottles is consistent with observations from Preunkert et al. (2011) who reported a mean contamination of 0.13 and 25 ppbC h⁻¹ in a 4 mL volume of
15 ultrapure water put in a 100 mL airtight glass bottle stored in the clean room and in a “general purpose” room of the lab, respectively. Note that when glass bottles were cleaned well before their use to sample snow, larger blank values were observed. For instance, the glass bottle blank of the flasks designed to sample a snowpit at Vostok (not shown) revealed a value close to 160 ppbC, likely due to the storage over several
20 months under various atmospheres before and after the snow sampling. Therefore, the bottle cleaning of snow samples collected within this study (see Tables 1 and 2) was done only a few hours up to 3 days before sampling and a storage bottle blank was systematically analysed. With these precautions bottle blank values generally remained below 25 ppbC. Since these values are significant with respect to measured snow
25 concentrations, they were subtracted from snow concentrations.

Another problem related to the use of Schott glass bottles to sample snow appears at the very low temperatures encountered in Antarctica (–30 °C). At these temperatures, snow grains that stick on the glass thread do not melt even when compressed in the thread plug of the glass bottle during closure of bottle. Thus the bottle cap does not

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close properly and significant contamination takes place during melting of sample at ambient lab temperature. For instance, on a total of 24 bottles used to collect snow pit samples at Concordia in 2012, nine bottles were found to remain well closed during the melting step, whereas fifteen bottles were not closed tightly showing higher DOC
5 content (82 ± 24 ppbC instead of 30 ± 20 ppbC).

2.3.2 Ice core subsampling

OC investigations were conducted in ice cores extracted at various locations in Antarctica and Greenland, as well as at the Col du Dôme Alpine site. With the aim to investigate change of the OC level with respect to the proximity of the ocean in Antarctica,
10 investigations were made on ice cores extracted at the high plateau (Vostok and Dome C) as well as at sites located closer to the coast (i.e. in the D47 core and at the top of the Caroline ice core drilled at the coast) (Table 1). The ages of the ice samples from the Antarctic ice cores were estimated by Parenin et al. (2004) for Vostok, by Jouzel et al. (1989) for Dome C, and by Ciais et al. (1994) for D47. In the case of the Caroline
15 ice core drilled in the margin area of East Antarctica (Yao et al., 1990), the ice from increasing core depth originates from increasing distance from the coast. In Table 1, based on the study from Lorius and Merlivat (1997), we report therefore an estimate of the altitude origin of the investigated ice core sections corresponding to the Holocene. The Dome C and Caroline OC investigations include, for the first time Antarctic ice
20 samples which correspond to the last glacial maximum (LGM).

Ice cores extracted from Summit (central Greenland) and NEEM (North Greenland) were investigated with the aim to extend the limited number of OC data available from Greenland ice and to examine the impact of boreal forest fire plumes, that were shown to often occur in summer (Legrand et al., 1992; Legrand and De Angelis, 1996;
25 Savarino and Legrand, 1998), on the OC levels. The age of ice from the Summit and NEEM sites were estimated by Clausen et al. (1997) and Rasmussen et al. (2013), respectively.

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To remove contamination present on the outer part of ice samples, we applied the procedure detailed by Preunkert et al. (2011) who developed a special glass device in which ice samples are washed with ultrapure water and melted afterwards. During sample melting an inert gas atmosphere is maintained inside the glass device to prevent contact of the ice sample with the ambient air of the lab. The following procedure was applied in routine: after its storage for 1–2 h at 0 °C, the sample is rinsed with ultrapure water and then put in the receptacle, which is at this point filled with 400 mL ultrapure water. Then the receptacle is emptied in the way that more than 50 % of the initial volume of ice was melted. After the complete melting of the remaining ice, the liquid sample is transferred into the Phoenix 8000.

2.4 Ancillary atmospheric data collected at Concordia

In our further ice core discussions, we will consider site relevant atmospheric information on the levels of organic gases and WSOC aerosol, the two key atmospheric fractions expected to contribute to the organic content of snow and ice. As seen in Table 3, respective information is already available for high-elevated Alpine sites and for Summit but remains limited to some organic gases like HCHO and light carboxylic acids at Concordia. Therefore, we here report WSOC aerosol levels observed in a year-round set of 51 high volume filters (circular quartz filters, Gelman Pallflex Tissuquartz 2500QAT-UP, 15 cm diameter) collected at Concordia from 1 January 2011 and 12 January 2012. Sampling was achieved continuously at weekly resolution with a flow rate of 0.7 m³ STP min⁻¹. To reduce contamination by organic material, quartz filters were pretreated by heating during several hours at 450 °C in laboratory. After that, filters were wrapped in thermally treated and cleaned aluminum foil. A 10 cm² piece of each filter was extracted with 12 mL ELGA ultrapure water in view to measure ions and OC. To remove particulate organic matter, the samples were passed through a hydrophilic PTFE filter (0.45 µm, Millex-LCR from Millipore) prior to analysis. Twelve filter blanks representing the sampling procedure (including the filtration prior the OC analysis) were analyzed. The mean blank value reaches 4 ± 1 ng m⁻³ STP and was subtracted from atmospheric

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sample concentrations. HULIS were measured on 5 selected winter and summer filters using a filter surface of 36 cm² for each sample.

3 Results and discussions

3.1 Presentation of data obtained in this study

OC data of snow and ice samples obtained in this work are summarized in Table 1 for Antarctica and in Table 2 for Greenland and the Alps. Antarctic ice samples were not seasonally resolved, and values reported in Table 1 correspond to annual means. In contrast, all OC values reported for Greenland snow and ice layers from Summit and NEEM are seasonally resolved. As previously discussed in numerous publications (see Legrand and Mayewski (1997) and references therein), the seasonality of major ions in Greenland ice exhibits a sodium maximum during the winter season, a calcium peak in spring, and a maximum of ammonium and nitrate in summer (see the examples reported in Figs. 1 and 2). As seen in Fig. 1, the ice layer located at the top of the Summit ice core section exhibits a large increase of the ammonium level. As for the first time discussed by Legrand et al. (1992) such ammonium events that are accompanied by a large formate increase are related to plumes of boreal forest fires that can reach the Greenland ice cap in summer. Similarly, another forest fire event was detected in a NEEM ice core layer located at 140.15 m depth (see Fig. 2 and discussions in Sect. 6.2).

3.2 A critical discussion on data inconsistencies

As seen in Table 1, OC concentrations widely fluctuate from site to site and from very recent snow to old ice, the range being particular large for Antarctic samples (from less than 5 ppb C up to 900 ppb C, Table 1). Prior our shallow snow study conducted at Concordia, data tended to suggest that OC is more abundant in the recent snow layers than in ice. Since this tendency is even more pronounced in Antarctica than in

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Greenland data, we can clearly rule out the assumption of a trend due to anthropogenic emissions. If correct, these high OC contents of Antarctic and Greenland surface snow with respect to deeper layers would suggest a very strong loss of volatile organic carbon after snow deposition. However, as shown in Table 1, our surface snow study at Concordia conflicts with this assumption.

Regarding existing surface snow Summit data we may ask whether the higher OC values reported by Grannas et al. (2004) (400–580 ppb C, Table 2) compared to those reported by Hagler et al. (2007a) (111 ± 45 ppb C, Table 2) are partly due to the use of a combustion method thought to be far more efficient than any UV oxidation method to oxidize insoluble organic particles possibly present in samples. In addition, Table 1 indicates that the highest values reported for Antarctic surface snow at the South Pole and in margin area of Dronning Maud Land and Princess Elizabeth Land were also obtained using a combustion method (instead of the UV method we deployed for surface snow and snow pit samples collected at Concordia). Therefore we will discuss in the next section in what extent the use of different (UV oxidation versus combustion) methods may have induced some differences in the results. Finally, given the high risk of sample contamination, especially for porous material like snow or firn (see Sect. 2.3.1), we will carefully revisit the respective precautions undertaken in the different studies to minimize contamination during sampling and analysis.

3.2.1 TOC or DOC?

When directly applied to liquid samples, combustion methods are thought to efficiently oxidize water-soluble species as well as insoluble organic particles present in samples, thus OC data obtained with this method (see Table 1 and 2) would refer to the TOC content of the respective samples. In contrast, it is generally accepted that the UV induced oxidation methods give rather low recoveries on particulate organic matter (Booth et al., 1998), especially for samples containing a high particulate organic content. Since the recoveries are however not zero, OC determinations obtained with an UV method are only referred as DOC if a sample filtration through a quartz fibre filter was applied prior

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to OC measurements. Whereas this procedure was applied by Hagler et al. (2007a), all the other studies using UV methods (Federer et al., 2008; Twickler et al., 1986; Preunkert et al., 2011; Legrand et al., 2007a, 2013) do not remove insoluble particles prior OC analysis.

If the water insoluble organic fraction (WinOC) of sample is large with respect to the one of water-soluble organic species, the direct use of an UV method without sample filtering would partly account for the above-discussed discrepancies. However, at the very remote places that we discuss here, insoluble organic carbon is not expected to dominate the organic carbon budget. As an example, the contribution of WinOC retained by filtering the samples through quartz filters has been documented for summer snow samples at Summit (Hagler et al., 2007a), showing on average a WinOC/DOC ratio close to 10% at that site. Given this low WinOC contribution, it is even possible that UV methods applied without sample filtering are able to partly (or totally) oxidize the particulate matter present in this kind of samples.

Federer et al. (2008) compared their UV method with a Sievers 900 TOC analyser and report a mean value of 145 ppbC measured with their UV system compared to 155 ppbC with the Sievers in a melted ice sample from Dye-3 (South Greenland). Given the certified blank for the Sievers vial (10 ppbC) the authors concluded that the difference between the two results is not significant. Using a hydrophilic PTFE filter (0.45 μ m, Millex-LCR from Millipore) to remove particulate organic matter from samples, the authors filtered another melted ice sample from Dye-3. This time the unfiltered sample revealed an OC content of 204 ppbC against 190 ppbC measured in the filtered sample, that lead the authors to suggest that their UV method measures rather TOC than DOC in samples containing a small amount of insoluble organic matter. In addition, their results suggest that particulate organic content of precipitation at remote sites such as Greenland contributes for a minor fraction to the total organic content. Attempts made by Preunkert et al. (2011) to quantify the recovery of their method for particulate carbon of melted ice samples were unsuccessful due to contamination occurring during sample filtering. Preunkert et al. (2011) therefore reported their data as dissolved

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organic (DOC) rather than total organic (TOC) carbon. However, at least for samples from Greenland and Antarctica, the above discussions tend to suggest that UV methods oxidize most of insoluble matter at these sites. Thus, Antarctic and Greenland OC values obtained with a UV method without sample filtering reported in Tables 1 and 2 may correspond to values of TOC. They may represent an overestimation by some 10 % if compared for instance with the Greenland DOC data from Hagler et al. (2007a). Therefore, within an uncertainty of 10 %, Antarctic and Greenland OC values obtained with a UV method without sample filtering can be referred either as TOC or as DOC.

For the high Alpine site of Col du Dôme, Legrand et al. (2007a) reported a mean WinOC/DOC ratio of 20 % instead of the 10 % found at Summit. To get a more quantitative idea of the WinOC recovery of the UV method from Preunkert et al. (2011) for this site, the sample filtering method was improved and one of the 2011 summer snow layers collected at the Col du Dôme site (see Sect. 2.3.1) was measured before and after sample filtering with a hydrophilic PTFE filter (0.45 µm). A filtration blank was analysed in parallel indicating an OC contamination of 84 ppbC. The unfiltered sample revealed an OC content of 664 ppbC versus 718 ppbC measured on the filtered sample. These values suggest that around 30 ppbC of WinOC were detected by the UV method in the unfiltered sample. If a WinOC/DOC ratio of 20 % is assumed for this site (see above), a WinOC content in the range of 120 ppbC would be expected, suggesting that only a quarter of the insoluble matter was oxidized by the UV method of Preunkert et al. (2011) in this alpine sample. Therefore, alpine OC values reported in Table 2 may be considered as DOC rather than TOC values.

3.2.2 Sample contamination

Since the inconsistencies seen on the OC content of samples from Greenland and Antarctica largely exceed 100 % and thus cannot be explained with the presence of insoluble organic matter (see Sect. 3.2.1), we revisit data reported in Table 1 and 2 with regard to potential contamination problems. Focusing the discussion on Antarctica where the inconsistencies are larger than elsewhere, levels reported for the South Pole

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and margin areas of Dronning Maud Land and Princess Elizabeth Land are at least one order of magnitude higher than those obtained in this study for Concordia. The sampling of surface snow carried out at the South Pole was not dedicated to detailed investigation of TOC and snow was collected and stored in Whirl-Pak polyethylene bags (Grannas et al., 2004). As previously discussed by Preunkert et al. (2011), one should avoid plastic devices to collect snow when dealing with very low OC level as a few tens of ppbC or less. Grannas et al. (2004) also reported TOC values covering the preceding year of snow sampling at Summit. Here values can be compared to those obtained in another study more focused on DOC and carried out with more stringent sampling conditions (Hagler et al., 2007a). Indeed in this latter study focused on DOC, snow was sampled with air-tight glass bottles previously cleaned in ultrapure water and baked to 550 °C for 12 h. As seen in Table 2, the snow samples collected at Summit in plastic bags exhibit far higher values than those collected into pre-cleaned glass bottles. Thus it is very likely that the near surface snow collected in plastic bags at the South Pole (Grannas et al., 2004) was also contaminated. The same would be true for surface snow samples collected at margin areas of Dronning Maud Land (DML) and Princess Elizabeth Land since samples were also collected by using a polypropylene scoop and stored into sealed low density polyethylene bags (Anthony et al., 2011). In addition, it has to be emphasized that the dust content observed in these samples (Coulter counter measurements) appears also rather high, reaching for instance 400 to 2000 ppb in the DML transect. Since this values largely exceed dust concentrations observed in the Holocene ice at DML (mean of 30 ppb up to 100 ppb, Ruth et al., 2008) it is legitimate to assume again a contamination for these coastal Antarctic surface snow samples.

Other available Antarctic data reported in Table 1 concern ice. Hereby OC values are consistently in the range of 10 ppbC at all Antarctic sites except those from Talos Dome that indicate values from one to several hundreds ppbC. Unexpectedly, the Talos Dome OC values exceed even those observed in pre-industrial ice from Greenland, a region located far closer to strong emissions of various organic compounds from the continental biosphere. The particularity of the Talos Dome ice samples lies in the fact

that the corresponding ice was extracted by using a drill fluid whereas other data were obtained from ice cores extracted by dry drilling. In addition, OC sampling of the Talos Dome ice was done continuously by applying a melting head on the ice (Federer et al., 2008). From our experience, OC measurements are very sensitive to the presence of cracks in the ice, especially when a fluid was used for the drilling. For instance, one of the three NEEM ice sections (186.6 m depth, Table 2) was damaged with a visible inner crack. Although having cleaned it with the same procedure than the other NEEM ice sections, the 3 pieces of ice from this section clearly indicated a remaining huge contamination (mean value of 570 ± 50 ppbC) and required a subsequent intensive rinsing of the melting glass device over several days to recover typical blank levels. From that it can be concluded that the melting head used for the ice decontamination had either crossed existing micro-cracks or induced the formation of new ones, which allowed penetration of drilling liquid inside the Talos Dome OC samples. It is also interesting to examine OC measurements made on ice from the 905 m long Dome C ice core that contains cracks but was extracted in 1978 using a thermal drill but without fluid. As seen in Table 1, in spite of the presence of cracks, OC values observed in this kind of ice remain fairly close to 10 ppbC suggesting that cracks efficiently re-close rapidly after the core extraction limited any subsequent contamination during storage of ice cores.

20 **4 Towards a more comprehensive picture of the OC content of snow and ice**

Several points emerge from the above discussions on existing inconsistencies in the OC literature data. It has to be noted that, the differences related to the insoluble OC recoveries of the deployed (UV versus combustion) methods cannot account for observed large inconsistencies. Instead, the inconsistencies highlighted in Tables 1 and 2 seem to be often related to contamination problems during sampling and storage of snow, as well as to the ice quality and/or its degradation (i.e. formation of cracks) during the subsampling.

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Discarding values obtained in snow sampled into plastic bags or stored into not properly closed glass bottles, an OC concentration of around 10–15 ppbC seems to be a typical value for Antarctic surface snow (Table 1). With the single exception of Talos Dome, most of OC concentrations observed in good quality Holocene ice range between 5 and 10 ppbC. Such a rather low abundance of organic carbon in Antarctica apparently conflicts with data reported by Calace et al. (2005) who reported from 52 to 92 ppb of dissolved fulvic acid in snowpit samples collected at Concordia. Indeed, if correct, such an abundance of the fulvic acid would imply the presence of at least 26–46 ppbC of DOC in Concordia snow. As shown in Table 3, our HULIS measurements made in several snow samples collected at Concordia indicate however far lower concentrations (from 1 to 3 ppbC). Note also that OC values observed in the Holocene ice indicate values close to 5 ppbC at Vostok and slightly higher ones in Caroline ice (~ 10 ppbC) corresponding to precipitation deposited initially between 1320–1550 m elevation. More data are however needed to increase the significance of such a difference and to discuss a possible contribution of the Antarctic ocean to the budget of organics in Antarctic snow.

As done for Antarctica, the consistency of the various OC data gained in Greenland is examined after having excluded the ones obtained from snow sampled into plastic bags (Table 2). Annual mean values observed within this study in 60 pre-industrial Summit ice samples that were not influenced by biomass burning inputs (see Sect. 3.1) are close to 12 ± 7 ppbC. These values are consistent with the more limited Summit data set (10 samples) reported by Preunkert et al. (2011) (23 ± 11 ppbC). A similar range is observed in the NEEM pre-industrial ice (19 ± 7 ppbC, Table 2). Over the more recent years (2002–2005) at Summit, Hagler et al. (2007a) reported a mean annual DOC value of 40 ppbC (i.e., at least twice higher than the pre-industrial level). As further discussed in Sect. 6.2, this difference may be attributed to man made activities. In addition, the latter study reported also a mean DOC value close to 110 ppbC for the 2006 surface summer snow at Summit (Table 2). As discussed by Anderson et al., (2008), it seems that most of post depositional OC effects at that site are due to formic

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and acetic acid re-emission from the snowpack. Since this surface summer value is twice higher than the mean one observed over the 4 previous summers, significant post depositional effects likely occurred at that site. Recent OC values observed at Summit are in the lower limit of the OC range reported for the 1982–1984 yr at Dye-3 (30 ppbC in winter and 320 ppbC in summer, Twickler et al., 1986). As already discussed by Hagler et al. (2007a), this may be related to the different location (south against central Greenland) and elevation (2486 m against 3240 m) of the two sites.

In the Alps, the different dataset consistently show annual mean values close to 70 ppbC prior to World War II (from 1925 to 1936) and higher ones (145 ppbC) over the recent decades (Table 2). Finally, the study of upper surface snow layers indicates a mean annual value of 140 ppbC suggesting no systematic difference compared to the OC levels observed in ice layers corresponding the late 1980's (i.e. corresponding to the youngest ice layers located just below the close-off).

5 The OC snow chemical speciation

The OC content of snow deposits is related to OC aerosols as well as organic gases present in the atmosphere that have been trapped into precipitation. In order to reconstruct relevant information on past change of atmospheric aerosol, it is needed to evaluate which fraction of OC in ice can be attributed to either OC aerosol or organic gases. Among short-chain organic gases present in the atmosphere, those having a strong polar character like monocarboxylic acids, alcohols, aldehydes and ketones can be incorporated in snow and ice and significantly contribute to the OC ice level.

Using data summarized in Table 3, we compare in Fig. 3 the chemical composition of OC present in recent snow layers deposited at Concordia (annual), Summit (summer), and in the Alps (summer). The choice of annual values for Antarctica and summer values for the two other sites is related to the possibility to compare, at each site, air and snow concentrations over the same seasonal interval. Year-round atmospheric data are available at Concordia but the low snow accumulation rate at that site

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prevents seasonal atmospheric parameters to be well archived in snow. Conversely, seasonally resolved snow records that are available at Summit and in the Alps can only be compared to atmospheric observations for summer since most of atmospheric observations are available for that season. In all three cases, chemical measurements included species related to organic gases (monocarboxylic acids and HCHO). Concerning the aerosol fraction, in addition to dicarboxylic acids and methanesulfonic acid that were documented in snow at the three sites, HULIS were determined in the Alps and in Antarctica but not yet in Greenland.

As discussed by Legrand et al. (2013), given their atmospheric abundance and high effective Henry's law constants, formic and acetic acids and to a lesser extent formaldehyde are organic gases that may be significantly incorporated in ice. In the case of the Alps, the authors concluded that C1–C3 mono-carboxylic acids and HCHO represent the most important water-soluble organic gases contributing to the OC content of ice (mean annual contribution ranging from 34 % over the 1971–1988 years to 43 % over the 1921–1951 years). The remaining OC amount present in ice denoted OC* and calculated as $[OC^*] = [OC] - [Monoacid-C] - [HCHO-C]$ would correspond to OC aerosol. Of this OC* quantity, the authors identified C2–C5 dicarboxylic acids and HULIS but a large fraction remained chemically unidentified (i.e. 30 and 45 % of OC over the 1921–1951 and 1971–1988 time period, respectively). As seen in Fig. 3, the unknown fraction of recent alpine summer snow layers accounts for 60 % of OC instead of 44 % over the 1971–1988 years and 27 % over the 1921–1951 years (Legrand et al., 2013). Though based on a single summer value (2012), the increasing trend of the unknown fraction from prior to after World War II revealed by Legrand et al. (2013) (see further discussions in the Sect. 6.2) seems to continue from the late 80's to the very last years.

The most extended OC speciation available for Greenland summer snow deposits is reported in Fig. 3 but does not yet include HULIS. As a consequence, whereas C1–C3 mono-carboxylic acids and HCHO contribute together for some 20 % of OC, the other chemically identified fraction remains very weak (2 % instead of 17 % in the Alps). Note that, as expected but in contrast to the Alps, a small organic aerosol fraction is related

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to the presence of methanesulfonate in Greenland. Finally, Fig. 3 reports for the first time the OC speciation of Antarctic snow. Compared to the other sites, C1–C3 monocarboxylic acids and HCHO represent a weaker contribution (11 % of OC). In addition, not only MSA but also HULIS represents a significant fraction of WSOC at that very remote site. The conclusion drawn for the case of the Alps that C1–C3 mono-carboxylic acids and HCHO represent the most important water-soluble organic gases contributing to the OC content of ice and that consequently the OC* quantity corresponds to OC aerosol, remains likely also true for Greenland and Antarctica except if oceans surrounding the Greenland and Antarctic ice caps emit other water-soluble organic gases than monocarboxylic acids and/or formaldehyde. One possible candidate is dimethylsulfoxide (DMSO), a water-soluble gas produced by the DMS/OH oxidation via the addition channel that is particularly efficient at low temperatures. Typical concentrations of 3 ng C m^{-3} STP of DMSO, which sporadically reach 20 ng C m^{-3} STP, were observed in summer at coastal Antarctica by Jourdain and Legrand (2001). However, as discussed by Legrand et al. (2001), once trapped in the liquid phase (cloud or hydrated aerosol), DMSO is quickly oxidized into MSA. Therefore the contribution of oxidation by-products from DMS emitted by the ocean at high latitudes, contribute rather to the OC* aerosol fraction (via MSA) than to the fraction related to water-soluble organic gases. In the following we assume therefore that the OC* quantity calculated as $[\text{OC}] - [\text{Monoacid-C}] - [\text{HCHO-C}]$ would correspond to OC aerosol fraction in Greenland and Antarctica snow and ice.

6 Atmospheric implications

In this section we will discuss to what extent the observed spatial variations of the OC ice contents are consistent with our present knowledge of organics in the atmosphere. After that, we will initiate a discussion on the origins and potential atmospheric implications of some observed temporal changes detected in Antarctica, Greenland and

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the Alps over various timescales, i.e. from present-day seasonal variations to changes during the last glacial interglacial period.

6.1 Spatial change: Antarctic versus Greenland and Alps

As concluded above, a comprehensive picture on the OC content of pre-industrial ice reveals a typical annual mean level of $\sim 5\text{--}10$ ppbC in Antarctica, 20 ppbC in Greenland and 70 ppbC in the Alps. To highlight the origins of these spatial variations, air and snow concentrations of organic gases suspected to contribute to the OC content of snow, and of various aerosols including those contributing to the WSOC fraction as well as a few well-known inorganic back-up species like sulfate or ammonium are compared in Table 3. Of course, this comparison only considers snow layers deposited over very recent times during which atmospheric observations are available. Though the OC level in the Alps and in Greenland seems to increase by a factor of 2 over the 20th century (see further discussions in Sect. 6.2), Table 3 still permit identification of the main cause driving the increase of the pre-industrial OC level in ice from Antarctica to Greenland and from Greenland to the Alps.

Table 3 suggests that the increase of the OC content from Antarctic to Greenland (from 5–10 to 20 ppbC) and from Greenland to Alpine (from 20 to 70 ppbC) pre-industrial ice is related to an increase of aerosol and to a lesser extent of water soluble gases. As seen in Table 3, water-soluble organic gases are more abundant in the Alps than in Greenland and in Greenland than in Antarctica, the change being particularly large for monocarboxylates. For instance, in surface snow the amount of monocarboxylates increases from ~ 1.2 ppbC (annual mean) in Antarctica, to 10 ppbC (summer) in Greenland, and to ~ 60 ppbC (summer) in the Alps. Due to the various parameters and complexity of involved processes that control the atmospheric budget of carboxylic acids over snow covered regions, a discussion on these geographical differences is complex and clearly out of the scope of the present paper. First, while measurements made at Col du Dome are likely representative of the free European troposphere, this is not the case in Greenland and Antarctica where stable thin atmospheric boundary

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layers often develop. Second, atmospheric gases like HCOOH and CH₃COOH are weak acids, for which incorporation in precipitation strongly depends on cloud pH and cloud water liquid content. Third, at least in Greenland, snow emissions are suspected to strongly control atmospheric concentrations observed in the very stable lowermost atmosphere lying over this ice cap (Dibb and Arsenault, 2002). One consideration that will be relevant in further discussions of these species is the fact that in the present-day atmosphere they are mainly secondary produced from biogenic precursors mainly emitted by the continental biosphere (Paulot et al., 2011). The weak contribution of anthropogenic sources to the budget of monocarboxylic acids was also supported by the examination of their long-term ice core trends in Greenland (Legrand and De Angelis, 1996) and in the Alps (Legrand et al., 2003).

The main cause leading to the observed increase of the OC content from Antarctic to Greenland and from Greenland to the Alpine pre-industrial ice is related to an increase of the OC* fraction. Using concentrations of monocarboxylic acids and HCHO reported in Table 3, we calculate that in surface snow OC* increases from ~13 ppbC (annual mean) in Antarctica, to 44 ppbC (summer) in Greenland, and ~210 ppbC (summer) in the Alps. To relate these snow values with atmospheric observations we also compare snow and air concentrations of sulfate and ammonium at the three sites. At Concordia, sulfate and ammonium data suggest that 1 gram of snow roughly contains the amount of aerosol present in 1 m³ STP of air. The 13 ppbC of OC* found in Concordia surface snow are consistent with the WSOC level observed in the atmosphere (annual mean of 13 ngC m⁻³ STP). At Summit, sulfate and ammonium data suggest that in summer 1 gram of snow roughly contains the amount of aerosol present in 0.5 ± 0.2 m³ STP of air. In this way, the 44 ppbC of OC* found in Greenland summer snow are consistent with the WSOC level observed in the atmosphere (summer mean of 77 ngC m⁻³ STP). In contrast to the case of Antarctica and Greenland for which surface snow and atmospheric measurements cover roughly the same time periods, for the Alps values reported for atmospheric aerosols were obtained in summer 2004 by Legrand et al. (2007b) and Preunkert et al. (2007) whereas summer snow concentrations cover

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the 2012 summer. Considering the recent decreasing trend of sulfate over Europe and reversely the increasing trend of WSOC (see Preunkert and Legrand, 2013) the preceding exercise done for Greenland and Antarctica may be less accurate in the case of the Alps. Nevertheless, as seen in Table 3, the 209 ppbC of OC* found in Alpine summer snow in 2012 are consistent with the WSOC level observed in the summer atmosphere 2004 (150 ngC m⁻³ STP). It therefore appears that the spatial change of the calculated OC* present in snow consistently follows the increase of WSOC concentration in the present atmosphere seen between central Antarctica, central Greenland, and the Alps.

Such spatial changes are consistent with present-day observations made over continents and oceans. For instance, a two-year-round climatology of organic aerosols was conducted in 6 sites located along a west-east transect extending from the Azores in the mid-Atlantic ocean to K-Puszt (Hungary) in central Europe by Pio et al. (2007). Hereby it was shown that WSOC is more than 10 times less abundant at the oceanic site of Azores (210 ngC m⁻³ STP) than over the European continent (3400 ngC m⁻³ STP at K-Puszt, 2700 ngC m⁻³ STP at Schauinsland in the Black Forest, Germany).

6.2 Temporal changes

Preliminary discussions on the causes of some observed temporal changes including seasonal variations, the possible trend from the pre-industrial to the present-day atmosphere, and the response to natural events such as sporadic arrival of forest fire plumes or climatic variations (last glacial/interglacial transition) are here presented.

Starting with Antarctic data, where the too low snow accumulation prevents a good preservation of the seasonal cycle, year-round atmospheric samplings made at Concordia in 2011 (Fig. 4) reveal a weak maximum of WSOC aerosol levels in summer. Note that this contrasts to the rather strong summer maximum observed for sulfate and methanesulfonate in relation to strengthened marine biogenic DMS emissions from the Antarctic oceans at that season (Preunkert et al., 2008). Also the WSOC seasonality differs from the one of sodium that shows higher levels during the second-half of

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winter 2011 than in summer. The species that seems to mimic the WSOC seasonality the best is the one of mineral dust emitted by continents and hereafter long-range transported towards Antarctica. Mineral dust was found to be twice more abundant in summer than in winter in Antarctica (Wagenbach, 1996). In this way, the seasonality of WSOC in Antarctica is consistent with a large dominance of WSOC aerosol sources over continents rather than oceans. In Greenland ice, the OC level is also characterized by a summer maximum (see Figs. 1 and 2). Not enough data are however available to separate spring and summer, two seasons during which the origin of air masses are different (Arctic basin in spring and North America in summer). Finally, in the Alps the seasonal cycle of OC* characterized by a summer maximum was discussed by Legrand et al. (2013) showing that wood burning may represent the main source in winter whereas vegetation emissions represent the dominant source in summer.

Data available in Greenland and in the Alps indicate a significant increase of OC concentrations from the pre-industrial era to the recent years. At both sites, the OC increase remains close to a factor of 2–3 and is mainly related to the OC* fraction. This change is rather moderate compared to the corresponding ones previously revealed for sulfate aerosol in response to growing SO₂ emissions in the Northern Hemisphere over the twentieth century (see a more detailed discussion in Preunkert and Legrand, this issue). In the case of the Alps, the ¹⁴C signature of the DOC content of summer ice revealed that, in spite of this increase, at least 75 % of DOC is of biogenic origin (May et al., 2013). Furthermore, the summer OC* change was highlighted by examining the time lag of the DO¹⁴C in ice with the ¹⁴CO₂ around the bomb peak of 1965, the authors clearly concluded that the main biogenic source contributing to the DOC in ice in summer is the living biosphere, while biomass burning represents a very weak source. There are many possible causes that may have strengthened either biogenic precursors of atmospheric WSOC (e.g. increasing temperature and or higher atmospheric CO₂ level) or the gas-particle conversion rate (e.g. increasing ozone concentrations). However, since most of the OC* fraction change was found to be related to an increase

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of the chemically unidentified fraction of DOC, it is difficult at this stage to discuss further the causes of these changes.

Our Greenland study documents for the first time the DOC content together with its speciation along forest fire events (Figs. 1 and 2). The pioneer study from Legrand et al. (1992) already revealed an input of formate in such events. Here, we can extend the examination of the organic composition of such perturbation. As seen in Fig. 2, it appears that the main contributors to the increase of DOC along such event are monocarboxylates (77 %) mainly as formate, chemically unidentified species (20 %), and to a lesser extent dicarboxylates (2 %) mainly as oxalate, and HCHO (1 %). We have to point out here that in contrast to background formate levels that are likely present in the form of formic acid, the formate observed along with forest fire events is present in the form of ammonium formate aerosol. As discussed in Sect. 5, the HULIS level is not yet documented in Greenland snow and ice. These species may account for the unknown contribution in forest fire events since they are known to be formed during incomplete breakdown of polymeric plant matter (Mayol-Bracero et al., 2002).

Finally, as seen in Table 1, though being still limited to a few samples, the Antarctic data gained on the Caroline ice core suggest a significant decreasing level of OC during the ice age compared to the Holocene. Note that, even probably slightly contaminated due the presence of cracks (see Sect. 3.2.2), the Dome C data also tend to suggest lower levels during ice age than during the Holocene. One possible cause for this decrease during ice age would be the lowering of continental biogenic emission (vegetation) following colder and drier conditions.

7 Conclusions and perspective

Available data on the dissolved (or total) organic carbon content of snow and ice from Antarctic, Greenland and Alpine ice cores were critically reviewed to understand the causes of existing inconsistencies. Most of large inconsistencies were found to be related to contamination problems during sampling and storage of snow, as well as

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degradation of the ice quality (presence of cracks) rather than to the different recovery of deployed (UV versus combustion) methods in oxidizing insoluble OC material present in samples. Discarding outliers from data, a more consistent picture with respect to our knowledge of OC sources, that are predominantly present over continents, merges. For instance, the typical annual pre-industrial ice content is close to ~5–10 ppbC in Antarctica, 20 ppbC in Greenland and 70 ppbC in the Alps. The investigation of chemical organic species present in snow and ice showed that, whatever the site (Antarctica, Greenland, or Alps), the C1–C3 monocarboxylic acids and HCHO represent the most important water-soluble atmospheric organic gases contributing to the OC content of ice and that the remaining amount of OC denoted OC* ($[OC^*] = [OC] - [Monoacid-C] - [HCHO-C]$) would correspond to atmospheric OC aerosol. C2–C5 dicarboxylic acids, HULIS, and sometimes methanesulfonate were found to contribute to the OC* quantity but a large fraction remained chemically unidentified.

A preliminary discussion was done on the change from pre-industrial to present-day atmosphere that was detected in the Alps and Greenland. In a recent study, the case of the alpine OC trend in summer ice was recently documented by $DO^{14}C$ measurements suggesting that in spite of three times higher levels in the 70s and 80s compared to prior World War II, a major fraction of OC is biogenic in origin. Many environmental parameters may have caused strengthened biogenic emissions of atmospheric WSOC including warmer temperatures, higher atmospheric CO_2 , enhanced oxidative capacity of the atmosphere. For Greenland, it has to be emphasized that the OC increasing trend since pre-industrial era is documented only by the level in pre-industrial ice level and in snow collected over the very recent years in snow-pits. A complete study of the change between the pre-industrial times and the present-day is limited by the fact that the cleaning of the firn material in view of OC measurement remains a difficult work. Finally, this paper reports for the first time a few data on the natural variability of the OC content of ice in relation with sporadic arrival of forest fire plumes over Greenland or response to the last glacial/interglacial transition in Antarctica.

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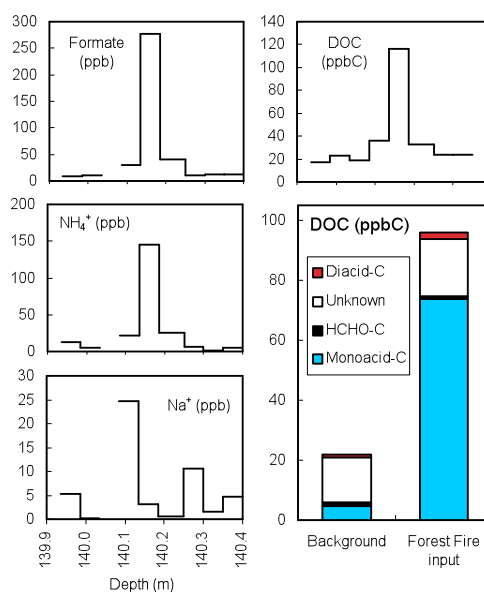


Fig. 2. Chemical signature of a forest fire event recorded in the NEEM ice site (dated to 1465 AD, see Sect. 2.3.2). Characterized by a large amount of ammonium and formate, the event took place in summer as suggested by the sodium minimum of this layer. Comparison of the organic carbon speciation along and outside the event (bottom right) indicates an input of OC mainly in the form of formate, to a lesser extent of dicarboxylates and other unidentified organic species (see Sect. 6.2).

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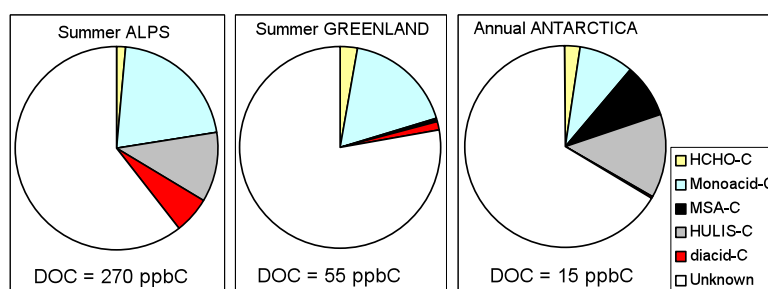


Fig. 3. From left to right: Chemical composition of DOC present in recent summer snow layers at the Col du Dôme and Summit sites, in recent annual snow layers at Concordia (see Sect. 4). The identified organic species include formaldehyde, C1–C3 monocarboxylic (formic, lactic, acetic, glycolic, and glyoxylic) acids, CH_3SO_3^- , C2–C5 dicarboxylic (oxalic, malic, malonic, succinic, and glutaric) acids, HULIS, and the remaining unidentified fraction of OC (see Sect. 4).

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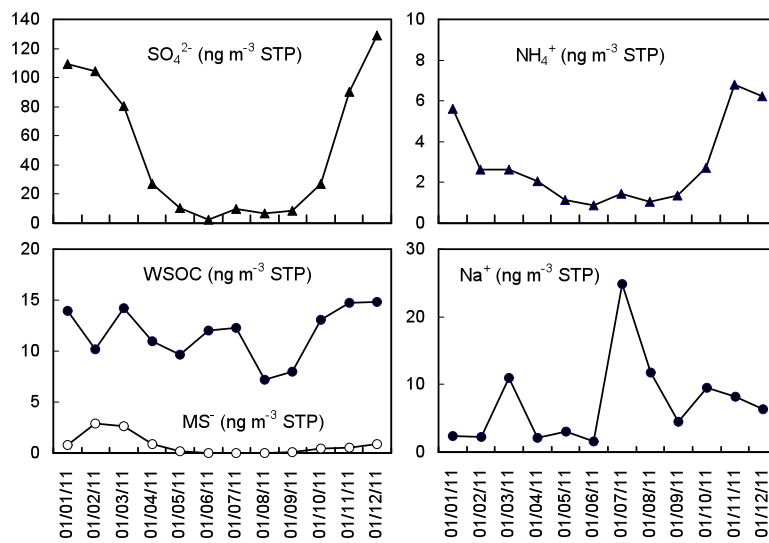


Fig. 4. Year-round bulk aerosol composition observed at Concordia in 2011. Note the higher sodium values in winter than in summer and the well-marked maximum of sulfate and methanesulfonate (CH_3SO_3^-) in summer (see discussions in Sect. 6.2).