Sea ice and pollution-modulated changes in Greenland ice core methanesulfonate and bromine

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Abstract. Reconstruction of past changes in Arctic sea ice extent may be critical for understanding its future evolution. Methanesulfonate (MSA) and bromine concentrations preserved in ice cores have both been proposed as indicators of past sea ice conditions. In this study, two ice cores from central and north-eastern Greenland were analysed at sub-annual resolution for MSA (CH$_3$SO$_3$H) and bromine, covering the time period 1750–2010. We examine correlations between ice core MSA and the HadISST1 ICE sea ice dataset and consult back trajectories to infer the likely source regions. A strong correlation between the low-frequency MSA and bromine records during pre-industrial times indicates that both chemical species are likely linked to processes occurring on or near sea ice in the same source regions. The positive correlation between ice core MSA and bromine persists until the mid-20th century, when the acidity of Greenland ice begins to increase markedly due to increased fossil fuel emissions. After that time, MSA levels decrease as a result of declining sea ice extent but bromine levels increase. We consider several possible explanations and ultimately suggest that increased acidity, specifically nitric acid, of snow on sea ice stimulates the release of reactive Br from sea ice, resulting in increased transport and deposition on the Greenland ice sheet.

1 Introduction

Atmospheric chemistry in the polar regions is strongly modulated by physical, chemical, and biological processes occurring in and around sea ice. These include sea salt aerosol generation, biogenic emissions of sulfur-containing gases and halogenated organics, and the photochemical–heterogeneous reactions leading to release of volatile, reactive bromine species. The resulting chemical signals influence the chemistry of the aerosol deposited on polar ice sheets. For this reason ice core measurements of sea salt ions, methanesulfonate (MSA), and bromine have been examined as potential tracers for sea ice extent (Abram et al., 2013; Spolaor et al., 2013b, 2016; Wolff et al., 2003). The interpretation of such tracers is complicated by the fact that their source functions reflect changes in highly complex systems, and signals are further modified by patterns of atmospheric transport and deposition.

MSA is produced by the atmospheric oxidation of dimethyl sulfide (DMS) ((CH$_3$)$_2$S). DMS is produced throughout the world’s oceans as a breakdown product of the algal metabolite dimethylsulfoniopropionate (DMSP), ((CH$_3$)$_2$S$^-$CH$_2$CH$_2$COO$^-$). DMS emissions are particularly strong in marginal sea ice zones (Sharma et al., 2012), and this source is believed to be a dominant contributor to the MSA signal in polar ice (Curran and Jones, 2000). Ice core MSA records have been used extensively in Antarctica as a proxy for local sea ice dynamics. Although the specifics of the relationship are highly site-dependent (Abram et al., 2013; Curran et al., 2003), MSA has been proven to be a rea-
reasonably good proxy for sea ice conditions (e.g. Curran and Jones, 2000). In the Arctic, the relationship between MSA and sea ice conditions is less straightforward due to the likelihood of multiple source regions with different sea ice conditions contributing to the ice core archived MSA (Abram et al., 2013). Until now, a significant \( r = -0.66 \) relationship between ice core MSA and Arctic sea ice extent (specifically August in the Barents sea) has only been established for a short record from a Svalbard ice core (O’Dwyer et al., 2000).

In this study we analyse the direct correlations between the MSA records from two Greenland ice core sites and the surrounding sea ice conditions in order to demonstrate the utility of MSA as a local sea ice proxy.

In this study, all dissolved or suspended bromine species are measured (including organic bromine) and shall be referred to as “bromine”. The primary source of total inorganic bromine (e.g. \( \text{Br}_2 \), \( \text{Br}^- \), \( \text{HBr} \)) in the marine boundary layer (MBL) is the ocean (Parrella et al., 2012; Sander et al., 2003). At concentrations of less than 0.2 % that of sodium (Na), bromide (\( \text{Br}^- \)) makes a small contribution to ocean salinity. \( \text{Br}^- \) can be concentrated in the high-latitude oceans when the sea water is frozen, since the formation of the ice matrix exudes the sea salts in the form of brine (Abbatt et al., 2012). Small sea-salt aerosol particles blown from the surface of sea ice are typically enriched with bromine (Sander et al., 2003) and satellite imagery has revealed that plumes of bromine (as BrO) are photochemically released from sea ice zones in spring (Nghiem et al., 2012; Schönhardt et al., 2012; Wagner et al., 2001). Recently, studies have begun to link ice core records of bromide enrichment (relative to sea water Na concentrations) preserved in polar ice sheets to that of local sea ice conditions (Spolaor et al., 2013a, b, 2014). Spolaor and co-workers demonstrated the springtime \( \text{Br}^- / \text{Na} \) that is preserved in the ice core is a record of bromine explosion events over adjacent seasonal sea ice. A \( \text{Br}^- / \text{Na} \) enrichment would therefore indicate a larger seasonal sea ice extent or conversely a shorter distance between the ice edge and the ice core site due to decreased multi-year sea ice (Spolaor et al., 2013a). However, like MSA, it is likely that the bromine–sea ice relationship in the Arctic is complicated by the myriad of bromine source regions which influence an ice core record in addition to factors which influence the degree of enrichment of the aerosol as it travels to the ice core site. In this study we compare ice core records of bromine enrichment to those of MSA and other common MBL species in order to determine the influence of sea ice conditions and other factors on bromine concentrations.

Here, we present measurements of MSA, bromine, and elemental tracers of sea salt and crustal input in two Greenland ice cores covering the time period 1750–2010 CE. These ice core records represent the first continuous sub-annual resolution records of bromine in polar ice to extend beyond the satellite era. We examine the relationship between these two sea ice-modulated tracers, their relationship to independent historical estimates of sea ice distribution, and the influence of industrialization on atmospheric and ice core chemistry.

2 Methods

2.1 Ice cores

The 87 m “Summit-2010” ice core was collected in 2010 close to Summit Station, Greenland (72°20’ N 38°17’24” W; Fig. 1). The average snow accumulation at Summit, as determined from the ice core record, is \(~ 0.22 \text{ m year}^{-1} \) water equivalent, with few instances of melt. Due to the relatively high snow accumulation rate, seasonal analysis of the sea salt species concentrations was feasible. The 213 m Tunu core was collected in 2013 (78°2’5.5” N, 33°52’48” W, Fig. 1), approximately 3 km east of the Tunu-N automatic weather station, part of the Greenland Climate Network. The average snow accumulation at Tunu, as determined from the ice core record, is \(~ 0.11 \text{ m year}^{-1} \) water equivalent.

The Summit-2010 and Tunu cores were first dated using volcanic horizons in sulfur (S) from well-dated historic eruptions (e.g. 1815, 1835, 1846, 1854, 1873, 1883, 1912). To aid the volcanic horizon assignment in the Tunu core the non-sea-salt S record was synchronized to the NEEM-2011-S1 volcanic record (Sigl et al., 2015). The dating of both cores was then refined by annual layer counting using a combination of seasonal cycles in Na, Ca, and the ratio of non-sea-salt S/Na for each entire core. In addition, hydrogen peroxide was used as a winter marker in the upper section of the Summit-2010 core. January was defined as the minimum value in the ratio of non-sea-salt S/Na.
Bigler et al. (2002) demonstrated that the input of anthropogenic sulfate during the Arctic haze slightly shifted the seasonal cycle of sulfate preserved in a north-eastern Greenland ice core ice toward spring by, on average, 1 month. The ratio of the Na to S was thus used to define the formal annual layer boundaries because using the ratio of two tracers with opposing seasonality (winter Na maximum and summer S maximum) reduces the sensitivity of the series to any small temporal fluctuations. Comparison with weekly surface snow samples collected from Summit (from 2007–2013; GEOSummit project) confirms the assignment of a non-sea-salt S / Na minimum in December/January. Monthly values were calculated assuming a constant distribution of snowfall within each year. Due to the lower accumulation rate and strong katabatic winds at the Tunu site, constraints from volcanic synchronization played a more important role in developing the depth-age scale for the Tunu core compared with Summit-2010. This dating technique is described in more detail for another Greenland ice core (NEEM-2011-S1) by Sigl et al. (2013, 2015). The annual layer dating for these ice cores resulted in a plutonium record that is consistent with other ice cores from Greenland between 1950 and 1970 and with the emission histories from nuclear weapon testing in the Northern Hemisphere (Arienzo et al., 2016). The error in the dating of the ice core records was estimated as ±0.33 years for the Summit-2010 record and ±1 year for the Tunu record.

2.2 Sampling and analysis

The ice cores were sampled from 33 × 33 mm cross-section sticks using a continuous melter system (McConnell et al., 2002). The silicon carbide melter plate provides three streams from concentric square regions of the ice core sample: an innermost stream (with a cross sectional area of 144 mm²), an intermediate stream (340 mm²), and an outer stream that was discarded along with any contaminants obtained from handling of the ice core. The innermost melt stream was directed to two inductively coupled plasma-mass spectrometers (ICP–MS, Thermo Scientific Element 2 high-resolution with PFA–ST concentric Teflon nebulizer (electrospray ionization – ESI)) run in parallel. All calibrations and runtime standards were run on both instruments and several elements were also measured in duplicate (Na, Ce, Pb) to ensure tracking between both ICP-MSs. In addition, an internal standard of yttrium flowed through the entire analytical system and was used to observe any change in system sensitivity. The instrument measuring bromine was run at medium resolution and was used to define the formal annual layer boundaries because using the ratio of two tracers with opposing seasonality (winter Na maximum and summer S maximum) reduces the sensitivity of the series to any small temporal fluctuations. Comparison with weekly surface snow samples collected from Summit (from 2007–2013; GEOSummit project) confirms the assignment of a non-sea-salt S / Na minimum in December/January. Monthly values were calculated assuming a constant distribution of snowfall within each year. Due to the lower accumulation rate and strong katabatic winds at the Tunu site, constraints from volcanic synchronization played a more important role in developing the depth-age scale for the Tunu core compared with Summit-2010. This dating technique is described in more detail for another Greenland ice core (NEEM-2011-S1) by Sigl et al. (2013, 2015). The annual layer dating for these ice cores resulted in a plutonium record that is consistent with other ice cores from Greenland between 1950 and 1970 and with the emission histories from nuclear weapon testing in the Northern Hemisphere (Arienzo et al., 2016). The error in the dating of the ice core records was estimated as ±0.33 years for the Summit-2010 record and ±1 year for the Tunu record.

2.3 Calculation of anthropogenic Pb, non-sea-salt S, and Br enrichment

The Pb derived from anthropogenic sources (exPb) was calculated as the difference between total lead measure in the ice core, [Pb]_{obs}, and that from dust sources. The Pb from dust was calculated as a fraction of the dust proxy ceurium, ([Ce]_{dust}) as follows:

$$\text{exPb} = [\text{Pb}]_{\text{obs}} - [\text{Ce}]_{\text{obs}} \times \frac{[\text{Pb}]}{[\text{Ce}]}_{\text{dust}},$$

where $[\text{Pb}]_{\text{obs}}$ and $[\text{Ce}]_{\text{obs}}$ are the observed concentrations of Pb and Ce in the ice core, respectively. $[\text{Pb}]/[\text{Ce}]_{\text{dust}}$ is the ratio of Pb to Ce in dust samples.
where the relative amount of Pb in dust, ([Pb]/[Ce]dust), has the constant mass ratio of 0.20588 (Bowen, 1979).

Similarly the amount of non-sea-salt sulfur (nssS) was calculated relative to the sea salt sodium (ssNa):

\[ \text{nssS} = [S]_{\text{obs}} - [\text{ssNa}] \times \left( \frac{\text{SO}_4^{2-}}{[\text{Na}]} \right)_{\text{seawater}}, \]

(2)

where the amount of sulfur relative to Na in sea water, ([SO_4^{2-}]/[Na])_{seawater} has the constant mass ratio of 0.252 (Millero, 1974). The ssNa was calculated by comparison with calcium as both have sea salt and dust origins (Rothlisberger et al., 2002):

\[ \text{ssNa} = \frac{[\text{Na}]_{\text{obs}} \times R_t - [\text{Ca}]_{\text{obs}}}{R_t - R_m}, \]

(3)

where \( R_t \) and \( R_m \) are the Ca/Na mean crustal and mean marine mass ratios of 1.78 and 0.038, respectively (Millero, 1974).

Bromine enrichment factors relative to sea water concentrations were calculated using the following:

\[ \text{enBr}(\text{Na}) = \left( \frac{[\text{Br}]}{[\text{Na}]} \right)_{\text{obs}} \times \left( \frac{[\text{Br}]}{[\text{Na}]} \right)_{\text{seawater}}, \]

(4)

where the ([Br]/[Na])_{seawater} mass ratio is 0.00623 (Millero, 1974).

### 2.4 Air mass back trajectories

To identify the likely sea ice source regions of MSA and Br deposited at the ice core sites, we perform 10-day air mass back trajectories of boundary layer air masses from each ice core site using the GDAS1 archive dataset in the Hysplit4 software (Draxler and Hess, 1998). The starting height of the back trajectories was 500 m to ensure that the monitored air masses travelled close enough to the surface at the ice core site to potentially deposit aerosols. The vertical velocity field was taken from the meteorological data files. Air mass back trajectories were started every 12 h and allowed to travel for 10 days (total number of trajectories hours = 14 400 h per month). The number of hours that the trajectories spent in a 2° × 2° grid was summed over all of the trajectories for that month between the years 2005 and 2013. Previous work showed that the rapid advection of MBL air was the likely source of reactive halogens at Summit (Sjostedt et al., 2007).

### 2.5 Sea ice correlation mapping

In order to assess the relationships between sea ice conditions and ice core chemistry, correlation maps were generated between annual MSA concentrations and monthly sea ice using the HadISST1 ICE dataset at 1° latitude–longitude monthly resolution (Rayner, 2003). Pre-1979 sea ice datasets were interpolated from sea ice extent maps compiled by Walsh (1978) which incorporate a variety of empirical observations. The data were later bias-corrected using modern satellite data (Rayner, 2003). Correlations were performed separately for the satellite period (1979–2012) and for the extended record (1900–2012), excluding the period 1940–1952 when the record has no variability due to scarcity of data (Rayner, 2003). Because strong DMS emissions occur in marginal sea ice zones (Sharma et al., 2012), we considered both sea ice concentration (SIC) and the area of open water in the sea ice pack (OWIP) which represents the size of the marginal sea ice zone. OWIP is defined as the difference between sea ice area (calculated from sea ice concentration over the area of the grid cell) and sea ice extent (NSIDC). A SIC of 15% was used as the threshold for a grid cell to contribute to sea ice extent. The area of OWIP was calculated within the coastal areas as defined by the results of the air mass back trajectories (Sect. 3.4).

Outliers were removed from the MSA time series (see Fig. 2) before the correlations were performed. The outliers were removed using the technique described by Sigl et al. (2013) for identifying volcanic signals using a 25-year running average filter. Correlations were performed on an annual rather than seasonal basis because the seasonality of ice core MSA is distorted due to post-depositional migration of MSA signal at depth in the snowpack (Mulvaney et al., 1992) (Fig. 3 in the article and Fig. S1 in the Supplement).

### 3 Results

#### 3.1 Bromine

Ice core measurements of bromine at Summit and Tunu covering the period 1750–2010 are shown in Fig. 2. Ice core Br levels at each site were stable until ~1820 at Summit and ~1840 at Tunu when they both decreased by ~1 nM, establishing a new baseline that was stable until the mid-1900s. Both ice cores also show a Br peak in the late 20th century. The concentration values and the timing of inflections in concentrations were determined by a three-step linear regression of the dataset. The analysis was performed by simultaneous linear least squares fitting of three straight lines joined by “inflection points” to the dataset. The variables of the fitting procedure were the slopes and intercepts of each line as well as the \( x \) axis locations at which the total function switched from one linear section to the next (the inflection points). Initial guess values were supplied for each variable to help the fitting procedure reach reasonable values. A summary of the regression results can be found in Table S1 in the Supplement.

Sea salt transport onto the Greenland ice sheet occurs predominantly during winter. Historically the wintertime sea salt maximum was believed to be due to increased cyclonic activity over the open oceans (Fischer and Wagenbach, 1996) though more contemporary studies show that blowing snow from the surface of sea ice may be a significant source.
At Summit, a wintertime maximum is observed in the most abundant sea salts, Na and Cl (Fig. 3). Bromine also shows a significant wintertime signal, however the annual maximum appears in midsummer – at concentrations ~70% above winter levels (Fig. 3a). Comparison with Br measured in weekly surface snow samples collected from Summit (from 2007 to 2013; GEOSummit project) confirms that this summer signal is real and not a result of post-depositional modification of seasonality of the bromine signal (Fig. S2). The results from that study confirm that total Br concentrations peak in summer on the ice sheet closely following the Br cycle observed in the Summit-2010 ice core. In addition to the comparison with the GEOSummit data, in the ice cores studied here there are routinely more than 10 measurements made within a yearly layer of snow, giving confidence to the allocation of a summer maximum in bromine at Summit. Analysis of the annual cycle of bromine in the Tunu ice core also shows a summer maximum when averaged over the entire ice core time series but with significantly larger error than observed at Summit. The timing of this peak suggests a predominant summertime deposition of bromine that dwarfs that from winter sea salt sources.

The shape of the annual bromine cycle does change slightly over the course of the Summit record (see Fig. 3). Starting in the early 1900s, the annual bromine cycle slowly becomes broader. A slight shift in the maximum from a solely summer peak in the pre-industrial era towards a broad spring–summer peak by 1970 is observed (Fig. 3 lower panel). Comparison with the sea salt tracer, sodium, which does not undergo the large temporal shift and broadening of its seasonal cycle shows that this change in bromine seasonality is not linked to changes in production or transport of sea salt aerosols or even dating uncertainties in the ice core but perhaps the introduction of an additional, smaller bromine source in the springtime during the industrial era.

Both ice cores show a predominantly positive Br enrichment throughout the year (Figs. S3, S4) relative to both sea salt elements chlorine and sodium. This enrichment reaches a maximum in mid-to-late summer at Summit (Fig. 3). We assume that this enrichment reflects Br enrichment in the aerosol transporting Br to the ice sheet. In a comprehensive review of global aerosol Br measurements, Sander et al. (2003) concluded that in general, aerosols which showed positive Br enrichment factors were of sub-micrometre size. These small aerosols can travel further (lifetimes of around 5–10 days) and due to their larger surface/volume ratio may experience more atmospheric processing than larger aerosols, resulting in the positive enrichment. However, post-depositional reduction of the bromine concentration is a possibility during the summer months due to photolytic processes at the snow surface. This may be the cause of the noisiness of the bromine signal within the lower accumulation Tunu core. However, the increased snow accumulation that occurs during the summer months in both central and northern Greenland (Chen et al., 1997) should act to minimize these bromine-depleting effects driven by increased insolation in summer, and indeed Weller (2004) has shown that accumulation rates of this size are large enough to prevent the post-deposition loss of other species such as nitrate and MSA.

Both sites also show a (small) positive enrichment of chlorine relative to sodium, which is amplified at small sodium concentrations. Chlorine-containing aerosols are expected to undergo similar chemical processing to bromine-containing aerosols but the enrichment factors of bromine (relative to sodium) are much larger, which is likely due to the high solubility of bromine species such as HBr (Sander et al., 2003). Alternatively, the chlorine enrichment could be interpreted as a sodium depletion of the aerosols particularly in those of small diameter where both concentrations are low; this would amplify the bromine enrichment (relative to sodium) but would not explain the bromine enrichment relative to
Figure 3. Upper panels: average seasonal cycle of species in the Summit-2010 ice core. The left-hand y axes are associated with the solid lines and the right-hand y axes associated with the dashed lines. Dashed lines (a–e) average seasonal cycle from depths 43.5–87.3 m (years 1742–1900). Solid lines (a–e) average seasonal cycle from 0 to 43.5 m (years 1900–2010). Error bars indicate the standard error of the monthly value. (a) Total bromine, (b) total sodium, (c) MSA, and (d) nitrate. Units for (a–d) are nM. Note that the seasonal cycle in bromine appears to broaden in the 1900–2010 period (see lower panel). Note also that the MSA maximum shifts from spring in the shallowest part of the ice core (solid line) to winter in the deepest part of the ice core (dashed line) due to post-depositional effects (see Fig. S1). (e) Average seasonal cycle in bromine enrichment (relative to sea salt sodium; see Eq. 3). (f, right) The sea ice extent (SIE × 10^6 km^2) within an area of the East Greenland coast (70–63° N, 15–45° W). (f, left) Area of open water within the sea ice pack (OWIP × 10^6 km^2) for the area defined by SIE. (g, left) Solar insolation at 12:00 GMT at the latitude of Summit (http://eosweb.larc.nasa.gov). (g, right) Annual cycle of the δ^{18}O water signal averaged over 1900–2010 CE. Lower panel: broadening of bromine seasonal cycle in the Summit-2010 ice core. The difference between the summer and spring bromine signal (JJA–MAM) was monitored over the length of the entire ice core. In the pre-industrial era (pre-1850) bromine peaks in summer; realised as positive values of JJA–MAM. After 1900 there is a marked broadening of the seasonal signal towards spring and by ~1970 the seasonal signal maximum is routinely shared between summer and spring realised as an averaged JJA–MAM of approximately zero.

chlorine. It is likely that both halogens undergo some degree of enrichment and the sodium undergoes some depletion in the aerosols though it is difficult to determine this from the data.

A summertime maximum in Br enrichment was also observed by Spolaor et al. (2014) in short segments of Antarctic Law Dome ice core as well as two Arctic ice cores. Spolaor et al. believe that the main source of the inorganic bromine originated from springtime bromine explosion events above sea ice and the summertime maximum could possibly be an indication of lag time between bromine-containing particles becoming airborne and their deposition. Further investigation is needed to definitively establish the seasonality of bromine deposition at the poles. However, the results of the Arctic ice cores studied here suggest that the summer maximum in bromine deposition is indeed real.

In the Tunu ice core, 11% of the monthly bromine enrichment measurements relative to Na were negative (less than the Br/Na seawater ratio, Fig. S3) and 12% were negative relative to Cl. It is possible that the negative enrichment values observed in the Tunu ice core are therefore a result of larger aerosols (> micrometre) reaching the site due to its
proximity to the coast (and thus the likely sea ice aerosol source region) in comparison to Summit.

3.2 MSA

The Summit-2010 MSA record (Fig. 2) replicates that measured by Legrand in 1993 (Legrand et al., 1997) and extends it an additional 17 years (see Fig. S5). The mean Summit-2010 MSA measurements over the period 1984–1992 (2.0 ± 0.7 (1σ) ppb) also compare well with the results of the sub-annually sampled Summit snow pit study performed by Jaffrezo et al. (1994); 2.1 ± 1.8(1σ) ppb. Both the Legrand and Jaffrezo studies measured MSA using ion chromatography of discretely sampled snow and ice. The similarity between the Summit-2010 measurements and the results of these studies demonstrates that the new, continuous technique is able to achieve a comparable accuracy in MSA measured concentrations to the traditional, discrete technique. It also demonstrates that negligible amounts of MSA are being lost by using the continuous melt method.

The Tunu measurements represent the first MSA profile at this location. Replicate measurements of the entire Tunu ice core were performed with the online, continuous technique by melting a secondary stick of ice cut from the original Tunu ice core. The replicate measurements closely followed the original MSA measurements demonstrating the reproducibility, stability, and high precision of the continuous MSA technique (Fig. S6). The Tunu MSA record was also reproduced using discrete samples collected from the CFA system (Fig. S7).

At Summit, MSA concentrations averaged 48 nM in the late 18th century, compared with just 27 nM at Tunu. From 1878 to 1930 MSA concentrations at Summit plateaued at 36 nM after which they began to drop rapidly, at a rate of 0.27 nM year−1, reaching 18 nM by 2000 CE. Large fluctuations in the MSA record after this time make it difficult to assess the most recent trend in Summit MSA concentrations. MSA concentrations in the Tunu core showed a similar temporal variability to those in the Summit record, and until the mid-20th century, were consistently lower in magnitude. MSA concentrations only began to decline consistently at Tunu after 1984, almost 50 years after the rapid decline observed in the Summit record. After 2000 CE, large fluctuations in concentration were again observed, making the modern-day trend in MSA concentration at Tunu difficult to establish.

Comparison with the total sulfur record (Fig. 4) reveals that during the pre-industrial period, MSA contributes to ∼12 and ∼7% of the total sulfur signal at Summit and Tunu, respectively, compared with <2% at the height of the industrial period (1970 CE) at both sites.

The low-frequency, pre-industrial trend in MSA concentrations seen in these ice core records closely follows that of bromine; particularly distinct is the decrease in both MSA and bromine at both sites in the early to mid-1800s (Tables S1 and S2). In the 1900s, however, both sites show a divergence between the MSA and Br records—as MSA begins to decline, Br concentrations increase.

A dramatic shift in the “timing” of the annual MSA maximum in the Summit-2010 ice core is illustrated in Figs. 3c and S1. The signal shifts gradually and continuously along the length of the entire Summit-2010 record from a spring to winter maximum (Fig. S1). This phenomenon has previously been observed in several Antarctic ice cores and has been attributed to post-depositional migration within the ice due to salt gradients (Mulvaney et al., 1992; Weller, 2004). At very low-accumulation ice core sites, post-depositional loss of MSA (and nitrate) must also be considered. Extrapolation of data collected by Weller (2004) from a series of East Antarctic ice cores predicts that sites with annual average accumulations of greater than 105 kg m−1 year−1 (0.105 m year−1) will not show post-depositional loss of MSA (or nitrate). Both ice cores in this study have sufficient average annual accumulation that post-depositional loss of MSA (and nitrate) is predicted to be negligible and so is not discussed further.

3.3 Acidic species

In winter, with the collapse of the polar vortex, polluted air masses enter the Arctic region as the phenomenon known as the Arctic haze (Barrie et al., 1981; Li and Barrie, 1993). SO2 and NOx from the haze are adsorbed onto aerosols or deposited directly on the ice/snow and oxidised to sulphuric (H2SO4) and nitric acid (HNO3). There are also natural sources of SO2 (biomass burning, volcanic eruptions, oceans – Li and Barrie, 1993; McConnell et al., 2007; Sigl et al., 2013) and NOx (microbial activity in soils, biomass burning, lightning discharges – Vestreng et al., 2009) as well as other snow/ice acidifiers including MSA, hydrogen chloride, and organic acids released from biogenic or biomass-burning sources (Pasteris et al., 2012).

The annual cycle for nitrate (NO3−) is shown in Fig. 3d. Before 1900 CE the nitrate shows a seasonal maximum in late summer/early autumn after which the maximum shifts to late spring/early summer. Although there are biological sources of NO3− in the ice core aerosol source regions, in a recent study focused on the NO3− and δ15N-NO3− record in the Summit-2010 ice core, Chellman et al. (2016) concluded that the pre-industrial (1790–1812 CE) NO3− seasonal cycle was driven by biomass-burning emissions. However, in the modern era (1930–2002 CE) oil-burning emissions became the dominant source of NO3− in the snowpack. The change in the dominant NO3− source due to industrialization is the cause of the shift in timing of the seasonal cycle.

Total snow acidity was stable at both sites from 1750 through to ∼1900 CE, except for sporadic, short-lived spikes due to volcanic eruptions. The average pre-industrial acidity was the same at both sites (∼1.8 µM). Both records also show two distinct maxima in acidity centred on 1920 and 1970 CE (Fig. 4), with Tunu displaying higher acid-
Figure 4. Comparison between the measured total sulfur (shown as sulfate) and acidity records from each ice core (top panels). The acidity record is dominated by the influence of the sulfur species until the early 21st century when the NO$_x$ pollution remains elevated whilst anthropogenic sulfur sources are depleted resulting in a slight relative elevation of the total acidity relative to total sulfur concentrations. The large spikes in the acidity and sulfur records are identified as volcanic events. The ice core records cover the period of the 1783 Laki eruption as well as the unknown 1909 eruption and Tambora eruption (Indonesia) in 1815 (Sigl et al., 2013). Comparison between Br/MSA and total acidity (centre panels) and nitrate (NO$_3^-$, bottom panels) measured in the ice cores. The Br/MSA ratio follows the total acidity record closely except where the record is dominated by the sulfur component (e.g. early 1900s). Of the two major acidic species the Br/MSA follows the nitrate most closely at both ice core sites.


The nitrate records from both sites follow the trend in Northern Hemisphere NO$_x$ emissions with a peak in ~1910 and 1990 CE – a result of emissions from increases in both Northern Hemisphere fertilizer usage and biomass and fossil fuel combustion (Felix and Elliott, 2013).

3.4 Air mass back trajectories

Air-mass back-trajectory results demonstrate that air masses reaching the Summit-2010 site between March and July originate primarily from the south/south-east of the ice core site (Fig. 5a). Previous back trajectory analyses by Kahl et al. (1997) also linked individual spikes in their Summit MSA record to air masses that had passed over this same region of coast (SE Greenland) within the previous 1–3 days. Similar back trajectories were calculated for Summit-2010 up to heights of 500 and 10 000 m (total column trajectory, Figs. 5a, S8a) illustrating that air masses that travel in the free troposphere and lower troposphere follow similar back trajectories and likely share the same source regions.

The results for Tunu indicate that air masses arrive primarily from the west coast of Greenland, passing over the Baffin Bay area, but there is also significant contribution from both the SE and NE (in May) coastal areas (Figs. 5b, S8b). Of these two secondary areas it is likely that aerosols transported from the NE would have a greater influence on the ice core concentrations due to proximity to the ice core site.
Figure 5. Air mass back trajectories from the (a) Summit-2010 and (b) Tunu ice core sites over the period 2005–2013 CE. Maps display the fraction of the total number of trajectory hours (ranging between 21,400 and 25,500 h month$^{-1}$) spent at altitudes under 500 m. Back trajectories were allowed to travel for 10 days. New trajectories were started every 12 h. Map grid resolution is $2^\circ \times 2^\circ$. Ice core locations are shown by a pink circle. Maps show that air masses consistently arrive at Summit from the SE Greenland coast with a smaller contribution from the SW coast. Air masses consistently arrive at Tunu from the western Greenland coast with a smaller contribution from the SE and NE coast. The air mass originating from the NE coast is most dominant in May and comparison with the total vertical column profile (Fig. S8) shows it is confined to lower altitudes unlike those from the west coast.
Aerosol deposited at Tunu therefore represents a mixture of source regions, but are likely dominated by the NW Greenland Baffin Bay coastal region.

3.5 MSA–sea ice correlations

Locations which showed a SIC variability greater than 10% (the average estimated range of uncertainty in the satellite measurements) and have a significant correlation to MSA \((t\) test, \(p < 0.05\)) are displayed in Figs. S9 and S10 for the months of March–July. A greater emphasis must be placed on the post-1979 sea ice concentration maps as these were derived from passive microwave satellite data and, where available, operational ice chart data. The likely air mass source regions, as defined by the results of the air mass back trajectories, are indicated by the black bordered regions. Within these areas there is generally a negative correlation between SIC and MSA, particularly in the spring months and only small patches that show large correlation (> 0.4). The large areas of positive correlation along the east coast and in the western Barents Sea are striking for the Summit-2010 record, however, these areas are outside of the defined air mass source region and thus are unlikely to be contributing to the ice core aerosol records. The positive correlation is likely an artefact of the negative autocorrelation between sea ice conditions in this region and the SE coast source region (Fig. S11).

The effect of the estimated error in dating of the MSA records on the SIC correlation maps is explored in Fig. S12. By shifting the dating of the MSA records to either extreme of the dating error estimate and replotting the SIC correlation panels, it is clear the error in the dating of the MSA records does not affect the sign of the correlations displayed on the maps but can have an affect on the magnitude of the correlation found in different locations. This is likely a result of the peaks in the MSA record being shifted in or out of temporal coherence with peaks in SIC at the different locations.

Over the period 1900–2010 CE, highly significant correlation \((t\) test, \(p < 0.001\)) is found between the annual ice core MSA and the amount of OWIP (representing the area of the marginal sea ice zone, Figs. 6a and 7a lower panels) in these aerosol source areas. For both ice cores the source region OWIP trend is followed by the MSA. In the Summit-2010 ice core the highest correlation between annual MSA and monthly OWIP occurs in May \((r = 0.58, p < 0.001\)) though the following months through to July all show highly significant correlations \((July r = 0.53, p < 0.001)\). For comparison, the May SIC correlation map is also shown as the upper panel in Fig. 6a. Figures 3f and S13 demonstrate that this time period (May–July) corresponds to the peak and then rapid decline in the amount of annual OWIP within the Summit-2010 aerosol source area because of the decreasing extent of sea ice. Rapid loss of sea ice reveals areas of biological activity previously capped by the ice, allowing surface–atmosphere exchange of DMS, resulting in the seasonal peak in atmospheric MSA correlation with the peak in the area of OWIP.

At Tunu, the highest correlation over the 1900–2012 CE period is found between annual MSA and annual OWIP \((r = 0.59, p < 0.001)\), though the July OWIP shows the highest monthly correlation and is also highly significant \((r = 0.41, p < 0.002)\). For comparison, the July SIC correlation map is also shown as the upper panel in Fig. 7a. Due to the more northerly location of the Tunu aerosol source region, the sea ice pack in this region is generally less fractured and breakup occurs later in the year, with a sharp peak in OWIP occurring in July (Fig. S13). The higher stability of the ice pack throughout the year compared to that in the Summit-2010 source region is likely reason the Tunu MSA shows highest correlation with the annual average of the OWIP. However, like Summit-2010 the highest monthly OWIP correlation occurs between the annual MSA and the timing of the maximum in annual OWIP (July).

Over the shorter, satellite era (1979–2012 CE) again Tunu shows strongest correlation between annual MSA and annual OWIP though at a much lower significance \((r = 0.32, p < 0.05)\), and the highest monthly correlation occurs in March \((r = 0.2, p < 0.1)\) albeit with low significance. The significance of the Tunu correlation over this period can be dramatically increased (annual OWIP \(r = 0.54, p < 0.001\); March OWIP \(r = 0.63, p < 0.001\)) if the closer, secondary aerosol source region (NE Greenland, 80–73° N, 20–0° W) is assumed to also influence the site in equal proportion. March corresponds to the timing of increased insolation and thus the rapid increase in ice algal production (Leu et al., 2015). The shift from a July to March peak in the correlation of OWIP with annual Tunu MSA may be a result of the reduced overall sea ice extent (SIE) (and thus OWIP), influencing the timing of MSA production. Unfortunately, the post-depositional migration of the MSA signal within the ice cores masks any evidence of true seasonal MSA shifts. Summit-2010 also shows a much less significant monthly OWIP correlation with the annual MSA signal over this time period, with the most significant correlation again occurring in March \((r = 0.4, p < 0.02)\). The greater significance of both the SIC–MSA and OWIP–MSA correlations at both sites over the longer time period is likely a result of the averaging of any MSA production or transport variability as well as the dominance of the low-frequency variability of both time series on the overall correlation.

3.6 MSA–bromine relationship

In an era where climate is driven by only natural forcings, chemical species that share a common source should show broadly consistent variability. This is evident in the pre-industrial section of both ice core records where the relationship between MSA and Br (monitored as Br / MSA) remains constant over the entire period (Fig. 4) despite individual records going through step function changes. Us-
Figure 6. Upper panels: correlation map of monthly sea ice concentration (SIC) derived from the Summit-2010 ice core. The SIC map corresponds to the month which shows the highest OWIP correlation (lower panels) with the annual MSA. Other monthly maps are shown in Fig. S9. (a) HadISST1 ICE dataset from 1900 to 2010 CE correlated with annual records of MSA (with outliers removed). Only locations that showed a SIC variability greater than 10% and have a significant correlation ($t$ test, $p < 0.05$) are displayed. The area of sea ice that is the likely source of MSA (as indicated by the air mass trajectories) are outlined in black (70–63° N, 0–45° W). (b) As for (a) but focused on the satellite period 1979–2010 CE. Lower panels: the correlation between the area of open water within the ice pack (OWIP) calculated within the black outlined areas shown on the upper maps and the annual MSA records (red) with outliers removed (orange) in nM. Summit-2010 MSA shows a significant, positive correlation with the amount of OWIP during spring within the integrated regions over both time periods. The highest correlations were found for March over the 1979–2010 period and May for the 1900–2010 period. In (b) if the MSA source region is enlarged to (70–63° N, 0–60° W) the March OWIP–MSA correlation increases slightly (from 0.38 to 0.4). The Summit-2010 enrBr(Na) and nsiBr records (in nM) are also compared to the same OWIP records. Particularly over the longer time period, there is little correlation between the series.
pre-industrial period, but during the industrial period nsiBr peaks, reaching a broad maximum between 1980 and 2000 CE of ~3.4 and 1.9 nM at Summit and Tunu, respectively.

4 Discussion

The significant correlation between variability of marginal sea ice zone (OWIP) area within the identified source regions and the MSA records suggests that MSA records can be used as a proxy for modern sea ice conditions in these areas. North Atlantic Oscillation proxy records developed in Greenland ice core records (Appenzeller et al., 1998) suggest that although the Northern Hemisphere climate phenomenon has shown variability over the past 200 years, its effect is damped in northern Greenland (Appenzeller et al., 1998; Weißbach et al., 2016) so we can assume that no major changes in atmospheric circulation patterns have occurred to change the source regions for the marine aerosols between the pre-industrial and industrial periods. If this assumption is true, our identification of MSA as a sea ice proxy (specifically a marginal sea ice zone proxy) may be valid for time periods both before and after 1850 at each ice core site.

The MSA records reveal that after 1820 CE a gradual decline in sea ice occurred along the southern Greenland coast (reflected in the Summit-2010 core) and that this decline in sea ice did not extend significantly to the most northern Greenland coastline (reflected in the minimal change in Tunu MSA during this period). It is not unexpected that the Summit-2010 record would show the most dramatic changes in sea ice, since we have demonstrated that the Summit sea ice proxy (MSA) is sourced from the south-east Greenland coast – an area sensitive to climate changes as it is primarily covered by young, fragile sea ice. The timing of the sea ice decline is coincident with the end of the Little Ice Age (LIA), identified from δ18O ice core records as spanning the period 1420–1850 CE in Greenland (Weißbach et al., 2016). The dramatic dip in sea ice reflected in both the Tunu MSA and Br records at 1830 CE (and also seen less dramatically in Summit) also appears in the multi-proxy reconstruction of sea ice extent in the western Nordic seas performed by Macias Fauria et al. (2010). This may be evidence of a 1830 CE sea ice decline event isolated to the East Greenland coast as the ice core records do not replicate the other dramatic, early 20th century fluctuations observed in the latter part of the western Nordic seas reconstruction.

From the ice core records it appears that the greatest decline in Greenland sea ice began in the mid-20th century, dropping to levels that are unprecedented in the last 200 years. This decline is observed along the entirety of the Greenland coast. Sea ice declined first around the southern coast (from 1930 CE, reflected in Summit-2010), followed 54 years later by the more northern coastline (reflected in the Tunu record; see inflection timings in Table S1). This sea ice decline is coincident with the sustained increase in greenhouse gases which has been identified as the major climate forcing and driver of increased global temperatures during the 20th century (Mann et al., 1998) and follows the same general trend in Arctic wide sea ice extent observed by Kinard et al. (2008).

Bromine (more specifically bromine enrichment, Spolaor et al., 2014, and bromine excess, Spolaor et al., 2016) has also been suggested as a possible proxy for sea ice conditions, however the timing of the largest bromine aerosol deposition, in summer, does not coincide with the largest growth or extent of new sea ice. Sea ice begins to increase only at the end of summer as the fractures in the ice cover are relaminated and the ice edge begins to advance southward (see Fig. 3f). Figure S4 compares the record of total bromine and bromine enrichment (calculated relative to sodium, enrBr(Na)) from the Summit-2010 ice core. The major discrepancies between the two records occur when the total sodium signal has sharp maxima causing dips in the enrBr(Na) record in ~1954 and 1990 CE and the magnitude of the low-frequency variability in enrBr(Na) is not as great as in the total bromine record. This is also demonstrated in Figs. 6 and 7 where the enrBr(Na) records are compared with the OWIP records. Whilst both series share high-frequency temporal features, over the longer term (1900–2010) the low-frequency trend is dramatically different. We are not discounting enrBr(Na) as a viable proxy for sea ice conditions, however the use of Na to try and extract the pure sea water component of the Br is complicated by the fact that a lot of Na comes from the sea ice surface as well as from the open ocean. Na itself has been used as a sea ice proxy in several prominent studies (WAIS Divide Project Members, 2013; Wolff et al., 2003) because, like Br, Na is incorporated into the snow on the surface of the sea ice and can be subsequently blown aloft to produce the atmospheric Na signal seen in the ice core. In addition, the Na concentration is fractioned upon the formation of the ice when mirabilite (Na2SO4) is precipitated out of the brine solution at ~8°C (Abbatt et al., 2012).

The calculated, non-sea-ice bromine records (nsiBr) for both ice cores are shown in Figs. 6 and 7. Like the enrBr(Na) records, the nsiBr records share some of the high-frequency features of the OWIP records, however there is no significant correlation between nsiBr and the selected OWIP records over the short time period. This supports the supposition that the nsiBr record is indeed an extraction of the non-sea-ice component of bromine from the total bromine record. Over the longer time period there is a significant negative correlation between OWIP and nsiBr at both sites (Summit-2010: $r = -0.7$, $p < 0.001$, and Tunu: $r = -0.22$, $p < 0.02$). This result is likely an artifact of the positive correlation from the MSA records used to generate the nsiBr records.

So what is the summertime source of bromine? What is the cause of the increase in springtime bromine explosion events in the industrial era (see Fig. 3 lower panel) and why does
Figure 7. Upper panels: correlation maps of monthly sea ice concentration (SIC) derived from the Tunu ice core. (a) HadISST1 ICE dataset from 1900 to 2012 CE correlated with annual records of MSA. The monthly SIC map displayed corresponds to the month which shows the highest OWIP correlation (lower panels) with the annual MSA. Other monthly maps are shown in Fig. S10. Only locations that showed a SIC variability greater than 10 % and have a significant correlation ($t$ test, $p < 0.05$) are displayed. The area of sea ice that is the likely source of MSA (as indicated by the air mass trajectories) are outlined in black ($77–67^\circ$ N, 62–50$^\circ$ W). (b) As for (a) but focused on the satellite period 1979–2012 CE. Lower panels: the correlation between the area of open water within the ice pack (OWIP) calculated within the black outlined areas shown on the upper maps and the annual MSA records (red) with outliers removed (orange). The Tunu enrBr(Na) and nsiBr records (in nM) are also compared to the same OWIP records and show poor correlation, particulary over the longer time period.

the bromine record deviate from the sea ice proxy record (MSA) around the same time? Possible sources of bromine and the factors which may effect the resultant bromine deposition flux are discussed below.

4.1 Alternate sources of bromine

4.1.1 Combustion of coal

Bromine is present in coal (Bowen, 1979; Sturges and Harrison, 1986) and coal burning is therefore a potential source of increased bromine deposition on the Greenland ice sheet over the period 1860–1940 (McConnell and Edwards, 2008). McConnell et al. (2007) demonstrated that pollution from the North American coal-burning era was deposited all over Greenland, leaving as its fingerprint large amounts of black carbon and toxic heavy metals. Sturges (1986) measured the relative concentrations of Br and Pb in particulates emitted from the stacks of coal-fired power stations and found a molar ratio (Br : Pb) ranging between 0.36–0.67 : 1. Figure 8 illustrates that at both Summit and Tunu the exPb (lead not from dust sources) preserved in the ice cores over the coal-burning era (~ 1920) was less than 1nM. This concentration implies that the upper limit to the amount of bromine deposited from coal combustion would be 0.67 nM (assuming no loss of bromine from the particulates during transportaion). This is an insignificant amount compared to the total Br signal preserved in the ice at this time. Coal combustion is not the major cause of the elevated industrial Br concentration.
4.1.2 Leaded gasoline

The largest global, historical, anthropogenic source of bromine is thought to be the combustion of leaded gasoline. Large quantities of 1,2-dibromoethane (DBE) were added to leaded fuel as a scavenger for Pb preventing lead oxide deposition by converting it to volatile lead bromide salts as well as CH$_3$Br (Berg et al., 1983; Nriagu, 1990; Oudijk, 2010). In 1925 CE gasoline had a Br : Pb molar ratio of 2 : 1 in a formulation which is now called “aviation fluid”. The Br : Pb molar ratio was reduced to 1 : 1 in the 1940s except in places such as the Soviet Union which continued to use aviation fluid for motor gasoline (Thomas et al., 1997). Although the consumption of leaded gasoline has been well-documented, particularly in North America, the estimates of the emissions of bromine compounds from the combustion process are still unclear. Estimates of the amount of DBE that is converted into gaseous CH$_3$Br range from 0.1 to 25 % (Bertram and Kolowich, 2000) and direct measurements of exhaust fumes across NW England found a Br : Pb ratio of between 0.65–0.8 : 1 in the airborne particulates (Sturges and Harrison, 1986).

The ratio of Br : Pb in the gasoline formulae can therefore be used only as an upper limit to predict the Br : Pb ratio in gasoline combustion aerosols transported to the ice core sites. Figure 8 shows a comparison between nsiBr and exPb measured in each ice core. Also illustrated is the upper limit of the amount of bromine expected from gasoline sources assuming the 2 : 1 Br : Pb ratio for aviation gasoline over the whole leaded gasoline era. Worldwide leaded gasoline emissions were estimated to have peaked in 1970 CE (Thomas et al., 1997) – an assumption that is supported by the observed timing of the exPb maximum observed in both ice cores. Whilst it is likely that leaded fuel contributed to the increased bromine observed between 1925 and 1970, it is clear that it was not the only contributor to the nsiBr record, particularly after 1970 when the nsiBr record continues to rise despite a worldwide decline in leaded fuel consumption. The disparity between the exPb and nsiBr records suggests the driving force for the enhanced emission of Br was still active and increasing after 1970.

4.1.3 Seasonal salinity changes

Younger sea ice surfaces such as frost flowers, new and first year sea ice have a higher salinity and thus have higher bromine concentrations than older sea ice surfaces (Hunke et al., 2011). The salinity of sea ice is at its maximum at the start of the winter season after which surface salinity slowly diminishes due to gravitational draining (Hunke et al., 2011). As summer approaches, ice continues to undergo desalination due to melting of surface snow which percolates through the ice (Hunke et al., 2011). Satellite observations that the BrO flux from the sea ice declines over summer (despite increasing insolation) is likely due to the combined reduction in young sea ice area and in ice salinity. Ocean surface
salinity decreases in the summer due to the increased meteoric water flux and melting of desalinated sea ice. Salinity increases are therefore unlikely to be the sole cause of the nsiBr flux observed in the ice core records and the observed summer maximum in bromine.

4.1.4 Organic bromine species

Gaseous bromocarbons can be a source of inorganic bromine to the snowpack when they react with OH or to a lesser extent with ·NO₃ or by photolysis (Kerkweg et al., 2008; WMO, 1995) to form the less reactive species HBr, BrNO₃, and HOBr. These species can then be washed out of the atmosphere and deposited on the snow surface due to their high solubility (Fan and Jacob, 1992; Sander et al., 1999; Yung et al., 1980).

The predominant source of gaseous bromine in the atmosphere is methyl bromide, CH₃Br (WMO, 2003). The major modern sources of CH₃Br are fumigation, biomass burning, leaded fuel combustion, coastal marshes, wetlands, rapeseed, and the oceans (WMO, 2003). The ocean is also a major sink for CH₃Br, the temperature-sensitive dissolution occurring through hydrolysis and chloride ion substitution to form bromide (WMO, 1995). Approximately 30 % of CH₃Br was from industrial emissions at the time of the global peak in the CH₃Br mixing ratio (1996–1998) (Montzka and Reimann, 2010). The timing of the massive increases in nsiBr seen at both ice core sites coincides with the timing of maximum anthropogenic emissions of CH₃Br. However, the estimated 2.7 ppt increase in global tropospheric CH₃Br above pre-industrial levels equates to only ~3.7 ppt (0.05 mM) Br incorporated into the snowpack (assuming 100 % conversion efficiency of CH₃Br in soluble Br species). This level is far less than the 2–5 mM increase in nsiBr observed in the ice cores during the industrial period.

Bromoform (CHBr₃) is emitted from vegetation such as marine phytoplankton and seaweed. It has the largest global flux of all the bromocarbons (estimated at almost 5 times that of CH₃Br (Kerkweg et al., 2008). However, it is very short-lived (atmospheric lifetime of ~17 days (Ordóñez et al., 2012) and thus is confined to the marine boundary layer. Inorganic bromine formed from the destruction of CHBr₃ would therefore be representative of only local sources of organic bromine. The biological seasonal cycle maximises the production of CHBr₃ in summer and concentrations are greatly reduced but not negligible in winter (tidal forcing also influences bromocarbon emission by allowing coastal algae to dry out (Kerkweg et al., 2008). The season of Arctic sea ice algae productivity is confined by limitations in available sunlight and nutrients resulting in a mid-to-late spring maximum – depending upon site location (Leu et al., 2015) – as is reflected in the seasonality of the MSA record. Direct transport of bromide enriched aerosols from these algal sources to the ice core sites again cannot explain the summer maximum of bromine observed in the ice. In addition to the incoherence of the seasonality of the bromine ice core signal, to-date biogenic sources have been considered insignificant sources of bromine in the Arctic marine boundary layer compared with the inorganic bromine source from sea salts (Simpson et al., 2007).

4.2 Cause of the springtime increase in bromine flux

4.2.1 Bromine explosion events

Spring is the time of “bromine explosion” events above sea ice. Sea salt aerosols passing through these BrO plumes can become enriched with bromine by adsorbing the gaseous species (Fan and Jacob, 1992; Langendörfer et al., 1999; Lehrer et al., 1997; Moldanová and Ljungström, 2001; Sander et al., 2003). Nghiem (2012) showed that these bromine rich air masses can then be elevated above the planetary boundary layer and transported hundreds of kilometres inland. Increasing the frequency and duration of the bromine explosion events would therefore likely increase the amount of bromine delivered to the ice core sites during spring without influencing the total aerosol flux and thus explain the shift in the bromine seasonal concentrations from a purely summer to a broad spring–summer maxima (Fig. 3).

Springtime field studies at Ny Ålesund, Svalbard have shown positive correlation between atmospheric filterable bromine species and elevated levels of sulfate and nitrate (Langendörfer et al., 1999; Lehrer et al., 1997) suggesting that acidic, anthropogenic pollution may be the driver of the observed increases in annual bromine enrichment during the industrial period and seasonal shift.

4.2.2 Acidity effects on debromination

In remote, relatively clean environments such as the Arctic, even small increases in acidity are thought to affect the cycling of bromine in the snowpack (Finlayson-Pitts, 2003; Pratt et al., 2013; Sander et al., 1999). In the laboratory, increasing the acidity of frozen (Abbatt et al., 2010) and liquid salt solutions (Frinak and Abbott, 2006; George and Anastasio, 2007) increased the yield of gas-phase Br₂ whilst at the same time increasing the solubility of other bromine species, such as HBr. The uptake efficiency of HBr by acidic sulfate aerosols, for example, is estimated at 80 % compared to 30 % for sea salt aerosols (Parrella et al., 2012). Interestingly, Abbatt (1995) demonstrated that HBr is more than 100 times more soluble in supercooled sulfuric acid solutions than HCl. This may explain the cause of bromine enrichment in the aerosol measured in the ice cores relative to the more abundant chloride (Fig. S3). The results of both the laboratory and field studies suggest that increasing snow/ice acidity in the Arctic will likely enhance springtime bromine explosion events above the sea ice whilst the increase in solubility allows the termination products of the explosion to be transported away from the sites on the surface of acidic
aerosols. Increasing springtime bromine aerosol concentrations would increase the average annual bromine concentrations deposited on the ice sheet and could explain the nsiBr records observed in both ice cores.

There are also significant periods over which the calculated nsiBr record shows negative values (e.g. 1815–1870 CE in Summit-2010 and 1860–1940 CE in Tunu). The negative values are a result of the total Br being less than that calculated by interpolation from the smoothed MSA record. Though the sources of Br and MSA are linked – which is what provides the similarities between the general low-frequency trend of the two species – the atmospheric processing, transport and deposition of the two species may be modified by different variables such as changes in atmospheric acidity, for example. These variables cause the short-term differences between the MSA and total Br records preserved in the ice so we believe it is not unreasonable to expect negative values in the calculated non-sea-ice Br record when the MSA and total Br are close (essentially no nsiBr).

Figure 9 illustrates that of the two dominant acidic species preserved in the ice, HNO$_3$ (represented by nitrate) shows the highest correlation to total bromine over sub-decadal time scales at both ice core sites. Records were detrended with an 11-year running average before comparison to isolate the high-frequency components of each record. The bromine–sulfuric acid (represented by sulfate) correlation is not significant. This is primarily because there is no bromine response to the dominant volcanic sulfate spikes throughout the record. The large spikes in sulfate concentrations did not cause a depletion of bromine preserved in the snowpack (Fig. 9). This result might be expected if the increased acidity caused more bromine to volatize. These results suggest that HNO$_3$ is the most influential of the MBL acidic species in the processing and transport of Br on aerosols in the MBL.

4.2.3 NO$_x$ and links to bromine

The snow and atmospheric chemistries of bromine and nitrate (NO$_3^-$) are tightly linked. NO$_3^-$ is one of the main sources of the ·OH radical. The ·OH radical can oxidize bromide salts and cause the release of gas-phase bromine species (Abbatt et al., 2010; Chu and Anastasio, 2005; George and Anastasio, 2007; Jacobi et al., 2014). Morin et al. (2008) observed that the majority of nitrate that is deposited to the snow surface is of the form BrNO$_3$ observed that the majority of nitrate that is deposited to the snow surface is of the form BrNO$_3$ and that the majority of nitrate that is deposited to the snow surface is of the form BrNO$_3$. The reaction

\[ \text{BrNO}_3 + \text{aq} \rightarrow \text{BrO} + \text{HNO}_3 \]

would suppress the emission of BrO from the snowpack and into the interstitial air.

In spring, when the greatest concentrations of BrO are observed over the sea ice the atmospheric concentrations of NO$_x$ species is rising. After 1900 CE there was, on average, a 60% increase in spring NO$_3^-$ concentrations observed in Summit-2010 ice core (Fig. 3d) which, as discussed in Sect. 4.2.2, if reflected in the concentration of acidic aerosols landing on the sea ice (specifically HNO$_3$ concentrations), would enhance the emission of BrO into the MBL. Satellite imagery shows that bromine in the form of BrO is confined primarily to the atmosphere above sea ice (Schönhardt et al., 2012; Wagner et al., 2001) but the presence of measurable bromine concentration hundreds of kilometres inland preserved in the ice cores demonstrates that the bromine must be transported inland, just not in the form of BrO. The reaction

\[ \text{BrNO}_3 + \text{aq} \rightarrow \text{BrO} + \text{HNO}_3 \]
of atmospheric NO$_2$ with BrO can produce the highly soluble BrNO$_3$, which will preserve the bromine in the aerosol allowing it to be transported inland. If there are high NO$_x$ concentrations at the deposition site this will aid in fixing the bromine into the snowpack. This is supported by the observation that NO$_3^-$ snowpack concentrations reach a maximum in summer, coherent with bromine snowpack concentrations even though maximum Br emission from the sea ice occurs in spring. So it appears that NO$_3^-$ in its different forms, as NO$_2$, NO$_3^-$, HNO$_3$ or BrNO$_3$, is intertwined with Br as it cycles between the gas and condensed phases and as it is transported from sea ice source to deposition site. Elevated levels of NO$_3^-$ over the Arctic could thus be the cause of the deviation of the bromine record from the MSA sea ice proxy record.

The high correlation between the pre-industrial (1750–1850 CE) NO$_3^-$ and Br records (Fig. 9) supports this observation of co-transport and sink of Br and NO$_3^-$ into the snowpack, though the natural sources of each are distinctly different. In the industrial era, the low-frequency temporal profile of the total bromine and nitrate records differ considerably, particularly at Summit (Fig. S15), apparently questioning the tight relationship observed before 1850. However, the positive correlation between the nitrate and the Br / MSA (Fig. 4) and nsiBr (Fig. 8) records is striking at both sites. The large relative increase in bromine (compared with MSA) during the era of high NO$_x$ pollution may point to a non-sea-ice source of bromine linked to nitrate emissions or simply an increased springtime emission and summertime deposition of Br from sea ice sources.

Bromine and NO$_x$ species shared a common source in the 20th century through the combustion of leaded gasoline (Sect. 4.1.2). As discussed above, we observe that leaded fuel pollution reaching the Arctic began to decline after 1970 in line with reduced global consumption, but the amount of bromine in excess of natural sources (nsiBr) continued to increase – following the trends in NO$_x$ pollution (Fig. 8a). The continued increase in NO$_x$ despite the decline in leaded fuel combustion is attributed primarily to biomass burning, soil emissions, and unleaded fossil fuel combustion (Lamarque et al., 2013). As the leaded fuel source of bromine began to decline, organic bromine pollutants continued to increase, as was discussed in Sect. 4.1.4. This can only account for a small fraction of the observed Br. The continued correlation between nitrate and nsiBr despite the decoupling of nitrate and bromine anthropogenic sources after 1970 suggests that nitrate pollution is likely influencing the processing of local, natural sources of bromine in the polar MBL, in effect increasing the mobility of the bromine and thus its flux and preservation in the ice sheet.

### 4.2.4 Consequences of nitrate-driven increased bromine mobility in the Arctic

Plumes of BrO emitted from sea ice regions have been linked to mercury deposition events which lead to an increase in the bioavailability of toxic mercury species in polar waters (Parrella et al., 2012). Increased springtime mobilization of bromine from the sea ice induced by anthropogenic nitrate could therefore increase the frequency and duration of these events and thus the mercury toxicity of the oceans. Increased atmospheric bromine concentrations would also increase the frequency of ozone-depletion events (Simpson et al., 2007) thereby altering the oxidative chemistry of the polar MBL.

Whilst several studies have begun to explore bromine records from ice cores as a proxy for past sea ice conditions, the results of this study demonstrate that in an era of massive increases in atmospheric acidity the natural relationship between bromine and sea ice conditions can become distorted, precluding it from being an effective modern-day Arctic sea ice proxy.

### 5 Conclusion

In this study we have shown that high-resolution MSA measurements preserved in ice cores can be used as a proxy for sea ice conditions (specifically the size of the marginal sea ice zone) along specific sections of the Greenland coast. The MSA records show that sea ice began to decline at the end of the LIA and again, more dramatically during the industrial period. Also, unsurprisingly, the changes in sea ice conditions in the northern sites have been less dramatic than along the southern coastline. Comparison between the 260-year records of bromine and MSA presented in this study allow us to show that in the pre-industrial era bromine concentrations preserved in the Greenland ice sheet are also likely linked to the local sea ice conditions. With the decline of sea ice in the modern era and the dramatic increase in acidic pollutants reaching the Arctic the sea ice–bromine connection is distorted, precluding it from being an effective, direct sea ice proxy during the industrial era. The introduction of NO$_x$ pollution, in particular, into the clean Arctic environment promotes mobilization of bromine from the sea ice, which in turn increases the bromine enrichment of the sea salt aerosols, forcing more bromine inland (particularly in spring) than would occur naturally. Nitrate has also been linked with the mechanism for preservation of bromine in the snowpack. The summertime maximum of nitrate may therefore be responsible for the observed summertime bromine maximum preserved in the ice cores. Whilst Northern Hemisphere pollution may prevent bromine from being an effective modern-day sea ice proxy in the Arctic, in Antarctica the anthropogenic flux of nitrate species is thought to be small in comparison with natural sources (Wolff, 2013), leaving room for the possibility that bromine may still be an effective proxy for local Antarctic sea ice conditions and for pre-industrial sea ice reconstructions.
6 Data availability

The raw Tunu data sets are available at the NSF Arctic Data Center website (McConnell, 2016).

The data used in the paper are also supplied in Excel spreadsheet format as a Supplement.

The Supplement related to this article is available online at doi:10.5194/cp-13-39-2017-supplement.

Author contributions. The paper was written and the data analysis performed by Olivia J. Maselli, with expert editing by Eric Saltzman. Ice cores supplied by Joseph R. McConnell. Tunu ice core was collected and processed by Olivia J. Maselli, Joseph R. McConnell, Nathan J. Chellman, Michael Sigl, and Rachael R. Rhodes under the leadership of Beth Bergeron. Ice cores dated by Michael Sigl and Joseph R. McConnell. The ICP–MS and CFA measurements performed by Olivia J. Maselli, Joseph R. McConnell, Nathan J. Chellman, Lawrence Layman, Daniel Pasteris, and Michael Sigl. The MSA measurements were designed and performed by Mackenzie Grieman and Eric Saltzman.

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