Bromine, iodine and sodium in surface snow along the 2013 Talos Dome – GV7 traverse (Northern Victoria Land, East Antarctica)

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Please find in this document:
- The list of relevant changes
- Point-by-point responses to the reviews
- A marked-up manuscript version

List of all relevant changes occurred during discussion and revision
- Minor changes have been addressed as in the two Referee comments.
- Fig. 9 has not been modified, as we think that it results blurred when imported in Word. The figure is shown below in its original resolution. We are happy to change the resolution if it is still considered poor resolution.
Author comment on

Review of Bromine, iodine and sodium in surface snow along the 2013 Talos Dome – GV7 traverse (Northern Victoria Land, East Antarctica)

Referee #2

The manuscript is much improved compared to the initial submission. I recommend publication following some minor corrections outlined here.

We would like to thank the Referee for reading the revised manuscript once again and for providing further comments to the text. We have modified the text following most of the provided comments.

Line 15 amend to: "Halogen chemistry in the polar regions occurs through the release of halogens from sea salt aerosol..."

Due to a similar comment of Michel (Referee3), the sentence has been rephrased: "Halogen chemistry in the polar regions occurs through the release of halogen elements from different sources. Bromine is primarily emitted from sea salt aerosols and other saline condensed phases associated with sea ice surfaces, while iodine is affected by the release of organic compounds from algae colonies living within the sea ice environment."

Line 16 amend to: "saline condensed phases associated with sea ice surfaces..."

Rephrased accordingly.

Line 17 and 18 amend to: "Measurements of halogen species in polar snow samples are limited to a few sites and there is some evidence that they are related to sea ice extent."

The sentence has been modified, but "although" has been left.

Line 48 amend to: "leading to formation of bromine monoxide, BrO, through reaction with ozone:"

Rephrased accordingly.

Lines 56 to 58: The Schoenhardt et al 2012 paper cited does not report tropospheric vertical columns, but total columns. Either amend the statement or find a different reference (although I am not aware that anyone has derived tropospheric columns for IO). "Tropospheric" has been removed.

Line 221: amend to 0.0062

Rephrased accordingly.

Line 230: this is not the dominant sink of Br/source of HBr. Reaction with HCHO has the same rate constant, but HCHO will be present at much higher concentrations than HO2. Actually you don't need to include the reaction at all. R5 has been removed.

Line 243: there is some confusion here. The statement that this parameter (bromine enrichment) is a marker of sea salt aerosol is contrary to what is written in Line 232, which talks of depletion being associated with sea salt aerosol.

We acknowledge the possible misunderstanding. Rephrased as "Such distribution of enrichment supports the theory that this parameter is, in these coastal sites, affected by sea ice signature."

Line 247 amend to: "they are also observed in winter, e.g. in core 8" – there are several examples, and core 8 is just one.

Rephrased accordingly.

Line 253 amend to: "high winter singularities or more extended peaks in cores GV7 and 8 respectively" Rephrased accordingly.
Line 257 amend to: “by multiplying the sea ice concentration”
“Value” has been removed.

Line 273 amend to “with maximum in November”
Rephrased accordingly.

Line 274: While it’s intuitive to think that the combination of sea ice and solar irradiance will generate the seasonality of bromine enrichment, the statement of line 274 “Such comparison suggests that the combined effect of sea ice and insolation drives the seasonality of bromine enrichment” is overstated when looking at Figure 7b. To look at just one example, bromine enrichment in October and November are effectively equal, while %SR.SI are considerably different. The authors need to be clear about the caveats when drawing their conclusions. A better way to present this would be “The combined effect of sea ice and insolation (Fig. 7b, magenta product distribution) shows similar features, with maximum in November, albeit with a much more pronounced springtime increase than seen in the bromine enrichment.” Rephrased accordingly. I don’t think that the statement about sea ice and insolation driving the seasonality of bromine enrichment can be supported by the data. One could find other factors with the same seasonality. For example, the seasonality of short-wave uv radiation alone (~300 nm) gives a November maximum (because of the effect of the ozone hole), so one could equally attribute that to driving seasonality of bromine enrichment. We don’t really agree with this comment, as radiation alone doesn’t give enrichment, as sea ice is needed. Modified as: “Such comparison suggests that the combined effect of sea ice and insolation is related to the seasonality of bromine enrichment.”

Line 317 amend to: “consistent with the homogeneous…”
Rephrased accordingly.

Line 333 again the authors are over-stating what can be drawn from their data. I’d suggest amending to “consistent with a sea ice influence in the area for the extra bromine deposition”. The sea ice influence in this area has been established from previous studies, cited along the text. Our findings are in line with such studies, therefore confirming them.

Lines 335 and 336: in line with my comments above I’d suggest softening the statement to “There is some evidence that the seasonality is linked to the combined effect of sea ice growth and sunlight, which trigger photochemistry above fresh sea ice” We agree with the Referee and the sentence has been modified.

Figure 9 is poor quality (somehow low resolution) and needs to be improved before publishing. We think that the Referee refers to the figure in the word document. The figure resulted blurred because Word doesn’t keep the PDF resolution. We therefore show below the original PDF file, being happy to increase the resolution if required.
Author comment on

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Referee #3 (Michel Legrand)

As already said, data on bromine and iodine in snow and ice are very welcome since they are still rather rare and are potentially interesting for a better understanding of the halogen chemistry at high southern latitudes in the past, for instance.

In the revised version, the authors considered most of my comments and suggestions. I appreciated that the estimates of excess bromine relative to sodium with respect to seawater composition are now shown. I therefore recommend it for publication.

We would like to thank Michel for reading the manuscript once again, and for pointing out the following two comments. We hope that more studies and past records in both the polar regions will improve the knowledge on halogens particularly important in perspective of a warming climate.

I just recommend the following clarification in the abstract.

First sentence of the abstract “Halogen chemistry in the polar regions occurs through the release of sea salt aerosols and other saline condensed phases from sea ice surfaces and organic compounds from algae colonies living within the sea ice environment.” needs to be reworded since algae colonies are thought to influence iodine (not bromine) and conversely saline condensed phases are important for bromine (not iodine).

We agree with the comment. The sentence has been modified considering also Referee2 comment, as the following:

“Halogen chemistry in the polar regions occurs through the release of halogen elements from different sources. Bromine is primarily emitted from sea salt aerosols and other saline condensed phases associated with sea ice surfaces, while iodine is affected by the release of organic compounds from algae colonies living within the sea ice environment.”

I think you can remove the last sentence of the abstract “The flux measurements are consistent with the uniform values of BrO and IO concentrations detected from satellites over the traverse area.” I think that, since BrO is only a small fraction of the bromine family (most of them being water soluble and able to contribute to bromine present in snow), it is quite very dangerous to rely snow deposition of bromine and satellite BrO observations.

We agree with the comment, as it can be interpreted as a measurement of BrO. The sentence has been removed.
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Abstract. Halogen chemistry in the polar regions occurs through the release of halogen elements from different sources. Bromine is primarily emitted from sea salt aerosols and other saline condensed phases associated with sea ice surfaces, while iodine is affected by the release of organic compounds from algae colonies living within the sea ice environment. Measurements of halogen species in polar snow samples are limited to a few sites although there is some evidence that they are shown to be related to sea ice extent. We examine here total bromine, iodine and sodium concentrations in a series of 2 m cores collected during a traverse from Talos Dome (72°48’ S, 159°06’ E) to GV7 (70°41’ S, 158°51’ E), analyzed by Inductively Coupled Plasma Sector Field Mass Spectrometry (ICP-SFMS) at a resolution of 5 cm.

We find a distinct seasonality of the bromine enrichment signal in most of the cores, with maxima during the austral spring. Iodine shows average concentrations of 0.04 ppb with little variability. No distinct seasonality is found for iodine and sodium.

The transect reveals homogeneous air-to-snow fluxes for the three chemical species along the transect, due to competing effects of air masses originating from the Ross Sea and the Southern Ocean. The flux measurements are consistent with the uniform values of BrO and IO concentrations detected from satellites over the traverse area.

Keywords: bromine, iodine, sodium, sea ice, Antarctica, halogens, polar halogen chemistry, Talos Dome.
1. Introduction

Halogen elements play an important role in polar boundary layer chemistry. The release of reactive halogen species from sea ice substrates has been demonstrated to be crucial in the destruction of tropospheric ozone at polar latitudes (so called Ozone Depletion Events) during springtime (Barrie et al., 1988; Simpson et al., 2007; Abbatt et al., 2012).

Although the ocean is the main reservoir of sea salts, various condensed phases of high salinity are found on young sea ice surfaces. During seawater freezing, brine is separated from the frozen water matrix and expulsion processes lead to both upward and downward movement, as temperature decreases (Abbatt et al., 2012). Therefore, high salinity brine, frost flowers and salty blowing snow make newly formed sea ice surfaces a highly efficient substrate for inorganic halides and for their activation and release in the atmosphere (Saiz-Lopez et al., 2012b, Yang et al., 2008). Some studies have also pointed out the role of open-water sea salts as a significant bromine source (Yang et al., 2005; Sander et al., 2003).

Reactive halogen species are involved in cyclic reactions between halogen radicals, their oxides and ozone. Reactions R1-3 show the main reactions for bromine. Atomic bromine radicals result from photolysis of molecular bromine, leading to formation of bromine monoxide oxide, BrO, through the reaction with uptake of ozone:

\[
\begin{align*}
    Br_2 & \rightarrow 2Br \\
    Br + O_3 & \rightarrow BrO + O_2 \\
    BrO + BrO & \rightarrow Br + BrO + O_2
\end{align*}
\]

Self reaction of BrO may form 2 bromine atoms (85%) or a Br₂ molecule (15%) which is readily photolyzed. The mechanism has a catalytic behavior that destroys ozone.

High concentrations of tropospheric vertical columns of BrO and IO have been confirmed by SCIAMACHY (SCanning Imaging Absorption spectroMeter for Atmospheric CartographHY) satellite observations over Antarctic sea ice (Schönhardt et al., 2012).

Bromine can then be recycled and re-emitted from halogen-rich condensed phases (such as sea salt aerosol or other saline solutions) or from sea ice surfaces (Pratt et al., 2013), leading to an exponential increase of bromine in the gas phase (Vogt et al., 1996). Such reactions, known as bromine explosions, lead to enhanced bromine in the atmosphere. A recent 1D chemistry model simulation predicted an increase of bromine deposition on surface snowpack after 24/48 hours of recycling over first year sea ice (Spolaor et al., 2016b). The stability of bromine in the snowpack was investigated at Summit, Greenland (Thomas et al. (2011), to explain the observed mixing ratios of BrO. Measurements in East Antarctica (Legrand et al., 2016) revealed that snowpack cannot account for the observed gas-phase inorganic bromine in the atmosphere.

Bromine enrichment in snow (compared to sodium, relative to sea water) has therefore been recently used to reconstruct sea ice variability from ice cores both in the Antarctic and Arctic regions (Spolaor et al., 2013a, 2016b).

Iodine is emitted by ocean biological colonies and sea ice algae (Saiz-Lopez et al., 2012a; Atkinson et al., 2012) mainly in the form of organic alkyl iodide (R-I) and possibly other compounds. These can be released by wind forced sea spray generation or percolation up to the sea ice surface through brine channels, and are subsequently photolyzed to inorganic species. Plumes of enhanced IO concentrations from satellites and ground based measurements were observed over Antarctic coasts, suggesting a link with biological and chemical sea ice related processes (Schönhardt et al., 2008). Grilli et al. (2013) have shown that ground based IO concentrations in Dumont d’Urville (Indian sector) were more than one order of magnitude lower than in the Atlantic sector (Halley station, Saiz-Lopez et al., 2007), consistent with greater sea ice in the latter. On the other hand, only sporadic events with IO concentrations above detection limits have been observed in the Arctic regions, possibly due to the greater thickness and lower porosity of Arctic sea ice which prevents an efficient release of iodine species in the atmosphere (Mahajan et al., 2010).
Measurements of sea ice related species such as bromine and iodine could therefore allow a sea ice signature to be obtained from ice core records. Until recently, only sodium has been used to qualitatively reconstruct sea ice at glacial-interglacial timescales (e.g. Wolff et al., 2006), but this proxy has limitations at annual and decadal scales, because of the noise input caused by meteorology and open water sources (Abram et al., 2013). Methane sulfonic acid (MSA) is an end product of the oxidation of dimethylsulfide (DMS), which is produced by phytoplankton synthesis of DMSP. MSA deposition has been successfully linked to Antarctic winter sea ice extent (Curran et al., 2003; Abram et al., 2010) and Arctic sea ice conditions (Maselli et al., 2016) on decadal to centennial scales, although some studies reported that the correlation with satellite sea ice observations is strongly site dependent (Abram et al., 2013). Several atmospheric studies reported no evidence of such link (Preunkert et al., 2007; Weller et al., 2011). Post-depositional processes causing loss and migration in the ice layers have also been widely reported to affect MSA, especially at low accumulation sites (Mulvaney et al., 1992; Pastur and Mulvaney, 2000; Delmas et al., 2003; Weller et al., 2004; Isaksson et al., 2005; Abram et al., 2008).

Victoria Land has been intensively studied for the past two decades. The Taylor Dome (Grootes et al., 2001) and Talos Dome (Stenni et al., 2011) deep ice cores respectively provide 150 kyr and 300 kyr climatic records directly influenced by marine airmasses. Sala et al. (2008) pointed out the presence of marine compounds (ikaito) at Talos Dome, typically formed at the early stages of sea ice formation. Their back trajectory calculations also showed that favourable events for air mass advection from the sea ice surface to Talos Dome are rare but likely to occur. An extensive study by Scarchilli et al. (2011) on provenance of air masses has shown that Talos Dome receives 50% of its total precipitation from the west (Indian Ocean), 30% from the east (Ross Sea and Pacific Ocean) and approximately 15% from the interior. Within the framework of the ITASE program (International Trans-Antarctic Scientific Expedition, Mayewski et al., 2005), several traverses were carried out to evaluate the spatial patterns of isotopic values and chemical species linked to marine influence (Magand et al., 2004; Proposito et al., 2002; Becagli et al., 2004, 2005; Benassai et al., 2005).

We present here bromine, iodine and sodium deposition in coastal East Antarctica, by investigating their total concentrations within a series of shallow firn cores, covering the 2010-2013 time period. The cores were drilled during a traverse performed in late December 2013 in Victoria Land (East Antarctica), from Talos Dome (72°48′ S, 159°06′ E) to GV7 (70°41′ S, 158°51′ E). The variability of these species at sub-annual timescales will inform on timing and seasonality as well as spatial patterns of their deposition. Such information is necessary for the interpretation at longer timescales of these elements and possible depositional or post depositional effects. These sub-annual resolution investigations are still limited to the Indian ocean sector (Law Dome - Spolaor et al., 2014) of Antarctica. The only data available on iodine in the Atlantic sector (Neumayer station) have been reported from a snow pit study by Friess et al. (2010). This study will test the regional variability of these tracers, providing measurements from the Ross Sea to the Indian ocean sector that remains otherwise unstudied.

2. Sampling and analyses

2.1 Traverse sampling

The traverse was performed in the northern Victoria Land region of East Antarctica (Fig. 1) from the 20th November 2013 to the 8th January 2014. The starting and ending locations were Talos Dome (72°48′ S, 159°12′ E) and location ‘6’ (see Fig. 1), close to GV7 (70°41′ S, 158°51′ E), for a total distance of about 300 kilometers. Talos Dome (275 km WNW from Mario Zucchelli station) is located approximately 250 km from the Ross Sea and 290 km from the Indian Ocean. GV7 is a peripheral site on the ice divide coming from Talos Dome, located just 95 km from the Indian Ocean.

During the transect, seven shallow cores, labelled hereafter TD (Talos Dome), 10, 9, GV7, 8, 7 and 6 were hand drilled to 2-m depth (except for GV7 which was 2.5 m). The main characteristics of the coring sites are reported in Table 1. Density profiles were obtained from each core immediately after drilling.

The hand auger had a diameter of 10 cm and consisted of an aluminum barrel equipped with fiberglass extensions. The cores were sampled in the cold laboratory at Cà Foscari University of Venice under a class-
100 laminar flow hood. Each core was cut with a commercial hand saw and decontaminated through mechanical chiseling by removing approximately 1 cm of the external layer. Every tool was cleaned each time a piece of sample was decontaminated into three serial baths of ultrapure water, which was changed every 10 washes. The cores were then subsampled at 5 cm resolution (3 cm for the GV7 core) into polyethylene vials previously cleaned with UPW and then kept frozen at -20 °C until analysis.

2.2 Analytical measurements
Total sodium (Na), bromine (Br) and iodine (I) concentrations were determined by Inductively Coupled Plasma - Sector Field Mass Spectrometry (ICP-SFMS Element2, ThermoFischer, Bremen, Germany) at Ca Foscari University of Venice, following the methodology described in Spolaor et al., 2014. The samples were melted one hour before measurements. During this time exposure from direct light was reduced by covering them with aluminum foils, minimizing bromine and iodine photolysis reactions. The introduction system consisted of a cyclonic Peltier-cooled spray chamber (ESI, Omaha, USA). The operational flow rate was kept at 0.4 mL min⁻¹, for an overall sample volume of 5.0 mL. Each sample determination consisted of 5 instrumental detections (less than 2% variations between them). The 5 values were then averaged to provide the final quantification. Each analytical run (10 samples) ended with a HNO₃ (5%) and UPW cleaning session of 3 min to ensure a stable background level throughout the analysis.

The external standards that were used to calibrate the analytes were prepared by gravimetric method by diluting separate stock 1000 ppm IC solution (TraceCERT® purity grade, Sigma-Aldrich, MO, USA) of the three analytes into a primary solution, which was further diluted for into 6 bromine and iodine standards (0.01, 0.05, 0.1, 0.5, 1 and 4 ppb) and 6 sodium standards (0.5, 1, 5, 10, 50 and 100 ppb). The calibration regression lines showed correlation coefficients R²>0.99 (N=6, p=0.05). The detection limits, calculated as three times the standard deviation of the blanks, were 50 and 5 ppt for bromine and iodine respectively and 0.8 ppb for sodium. The reproducibility of the measurements was carried out by repeated measurements of standard samples within the calibration range. The residual standard deviations (RSD) were respectively 5 % (bromine), 3 % (sodium) and 2 % (iodine). Procedural UPW blanks were analyzed periodically to test the cleanliness of the instrument lines.

Stable isotopes of water (¹⁸O and D) measurements were conducted on sub sample aliquots, which were immediately refrozen and shipped to the Center for Ice and Climate (Copenhagen, Denmark). Analyses were carried out using a Cavity Ring-Down Spectrometer (Picarro, Santa Clara, USA) using the method described by Gkinis et al. (2010). Septum-sealed glass vials were used for these measurements to prevent any sample evaporation during the experimental phases.

3. Results and discussion

3.1 Stable water isotopes and snow accumulation

The cores were dated based on the seasonal variations identified in the stable water isotopes (both δ¹⁸O and δD). Midwinters were associated to the relative minima of the isotopic curves (Fig. 2). In case a winter isotopic plateau was found, the center of the plateau was associated to midwinter depth (2011 in core GV7; 2012 and 2011 in core 8; 2010 in core 6). Almost all the cores cover the period between 2010 and late 2013, providing four years of snow deposition. The only exception is represented by core 6, whose upper layer is missing.

The annual deposition signal looks less clear in the two cores that were drilled at the sites with the highest elevation and the closest to the Ross Sea, cores TD and 10, and especially for 2013 in core 10. The two sites are probably the most affected by surface remobilization and isotopic diffusion due to low accumulation. Indeed, non-uniformities in the shallow snow layers such as sastrugi, dunes, wind crusts and other features have been identified as an important aspect of the surface morphology around the Talos Dome area (Frezzotti et al., 2004; 2007).
The annual accumulation rates were calculated by selecting the depth intervals included within consecutive maximum or minimum δ18O values (Table 2). Each snow layer within this interval (i.e. sampling resolution, 5 cm) was multiplied by the density of the snow at that depth, the density curves having the same resolution. The contributions were summed over the annual thickness. Table 2 also includes accumulation rates in Victoria Land reported from previous studies. The GV5 site is located between sites 10 and 9 (Fig. 1).

The accumulation rates found during the traverse are in general agreement with the previous works (Becagli et al., 2004; Frezzotti et al., 2007), except for Talos Dome. The accumulation values calculated from the smoothed isotopic profile in Talos Dome are well above those measured by the stake farm (n=41, C. Scarchilli, personal communication) for the same years. The inconsistency between the accumulation rates derived from the core and those derived from the stake farm and previous measurements suggests that the isotopic assignments of years may be incorrect at this site, and that the profile contains more years than have been assigned. This core therefore is not used in further calculations. The fluxes of deposition of sodium, bromine and iodine in the other cores along the transect are calculated using the accumulation rates from this work.

The accumulation pattern along the transect increases from Talos Dome to the Southern Ocean (GV7, 8, 7, 6), as the previous works have also found (Magand et al., 2004; Frezzotti et al., 2007). Scarchilli et al. (2011) already pointed out how Talos Dome receives 50% of its total precipitation from the north-west (Indian Ocean), 30% from the east (Ross Sea and Pacific Ocean) and approximately 15% from the interior of the plateau. In this picture, our accumulation data show a decrease from the Indian Ocean moving away from the Indian Ocean coasts and approaching Talos Dome.

The sites are located at decreasing altitudes moving from Talos Dome site (highest point) towards the coast facing the Indian sector (site 6). The minimum δ18O value found in each core shows a decreasing trend with altitude, with an elevation gradient of -1.35‰(100m)-1. This super-adiabatic lapse rate is confirmed by the surface snow samples collected taken during the 2001/02 ITASE traverse (Magand et al., 2004).

### 3.2 Sodium, Bromine and Iodine

Sodium shows a mean concentration of 34 ppb, in agreement with published values in this area (Becagli et al., 2004, Bertler et al., 2005, Severi et al., 2009). Among the three elements, sodium shows the highest standard deviation (21 ppb) because of the high variability of sea spray inputs at coastal sites. Singularities up to 200 ppb are probably associated to sea salt rich marine storms. Iodine has an average concentration of 43 ppt, compared to bromine (42%) and sodium (61%).

The bromine enrichment has been calculated as the bromine excess with respect to sea water concentrations, 

\[ \text{Brenr} = \frac{[\text{Br}]}{[\text{Na}]} \]

where [Br] and [Na] are the bromine and sodium concentrations in the sample and 0.0062906 is the bromine-to-sodium concentration ratio in sea water (Millero, 2008). Similarly, non-sea-salt bromine, 

\[ \text{nssBr} = \frac{[\text{Br}]}{[\text{Na}]} - 0.0062 \]

Benassai et al. (2005) have concluded that sea-salt sodium is the dominant fraction (more than 80%) of the total sodium budget in this area. No correction to sodium was therefore applied for this calculation. Despite bromine being a sea salt marker like sodium, it is activated when gas phase HOBr oxidizes bromide over halogen rich sea ice surfaces (i.e. first year sea ice, FYSI) and suspended sea salt aerosol, and exponentially released as Br$_2$ (R4). Following photolysis, atomic bromine radicals can be converted back to HBr (R5): 

\[ \text{HOBrg} + \text{HBr(aq)} \rightarrow \text{Br}_2(g) + \text{H}_2\text{O(aq)} \]  \hspace{1cm} (4)

Following photolysis, atomic bromine radicals can be converted back to HBr. 

\[ \text{Br}_2(g) + \text{HO}_2(aq) \rightarrow \text{HBr(g)} + \text{O}_2(g) \]  \hspace{1cm} (5)

Therefore, sea ice presence should lead to bromine enrichment or depletion, depending whether deposition is dominated by the depleted sea salt aerosol or by the enriched gas phase HBr. Bromine enrichment has already
been linked to sea ice presence in both Arctic and Antarctic coastal sites (Simpson et al., 2005; Spolaor et al., 2013b, 2014, 2016; Valdelongua et al., 2016).

The distributions of bromine enrichment values are reported in Fig. 3, divided into the cores closest to the Ross sea (TD, 10, 9, blue distribution) and to the Indian ocean (GV, 8, 7, 6, red distribution). The first set of cores show on average higher values (5.7 ± 0.3) than the second (4.2 ± 0.2). The variability (rms) is also higher (3.5 ± 0.2) in the first set compared to the ‘Indian ocean’ set (2.5 ± 0.1), because of greater distance covered by the sampling (165 km compared to 40 km). Overall, the values extend from a minimum of 0.5 to 17 with more than 98% of the samples showing values greater than 1 (i.e. sea water value). A detailed insight on the few <1 values revealed that these samples are associated with very high contributions of sodium inputs (>120 ppb), therefore likely associated to strong marine events. Such distribution of enrichment supports the theory that this parameter is, in these coastal sites, affected by sea ice signature, a marker of sea salt aerosol with an extra contribution from sea ice.

The measurements of the chemical species for the different coring sites along the traverse are reported in Fig. 4-5-6 on an age scale (with the exception of Talos Dome which is reported on a depth scale in the supplementary material). Sodium timeseries show great variability: peaks are often found in summer, although they are also observed in winter, e.g., in core 8. These findings confirm that, as previous works pointed out (Curran et al., 1998), in coastal sites storm events carrying open ocean sea salts are more important than sea ice as a sea salt source, although the high level of variability suggests also that meteorology and natural variability play a role (Wagenbach et al., 1998). Bromine and both Br$_{en}$ and nssBr show annual variations, with maximum values in late spring-summer, confirming ice core measurements by Spolaor et al. (2014), Valdelongua et al. (2016), and aerosol measurements by Legrand et al., 2016. Iodine shows a more stable signal throughout the year and high winter singularities or more extended peaks in cores GV7 and 8 respectively.

The timing of the bromine enrichment signal in ice cores relies on the combined effect of sea ice and sunlight, responsible for the photochemical production and release of molecular bromine, Br$_2$ (Pratt et al., 2013). Sea ice area in the 130°E-170°W sector was calculated for the 2010-2013 period using publicly available NSIDC passive microwave sea ice concentration data (Meier et al., 2013), by multiplying the sea concentration value in each grid pixel by the area of the pixel (25 x 25 km$^2$) and integrating over the domain. The longitude sector was decided on the basis of Scarchilli et al. (2011), who concluded that air masses arriving in this area originate from the Ross sea and from the Indian ocean sector, by analyzing 5 day back trajectories from 1980 to 2001. Figure 1 (panel b) shows the minimum and maximum, found in January 2010 and August 2011, respectively. The monthly sea ice areas from 2010 to 2013 were calculated for such sector and plotted in Fig. 7a (blue); each monthly value was normalized to the total annual sea ice area. The minimum sea ice is found in February, while a longer lasting minimum throughout winter and spring is observed, before a rapid decrease from November. Solar radiation values Fig. 7a (red points) were calculated at 71$^\circ$ S, 158$^\circ$ E using the Tropospheric Ultraviolet and Visible Radiation (TUV) Model within the [300,500] nm wavelength interval. Each point represents a daily average of the 15th day of each month of 2012 and it is considered a monthly representation.

The sub annual distribution of bromine enrichment along the transect is shown in Fig. 7b (blue). Each bin contains the cumulative monthly value for every year in every core, normalized by the total value of each year (which may change according to year and location). The histogram is then normalized by the overall sum measured in the transect. The distribution shows a clear sub-annual oscillation with lowest and highest annual contribution in May (autumn) and October-November (late spring), respectively. The combined effect of sea ice and insolation (Fig. 7b, magenta product distribution) shows similar features, with maximum in November, albeit with a much more pronounced springtime increase than seen in the bromine enrichment spring. Such comparison suggests that the combined effect of sea ice and insolation is related to drives the seasonality of bromine enrichment. Monthly sea ice area values are reported in Fig. 7c (blue), together with annual averaged values of bromine enrichment (black) and first year sea ice, FYSI (red), calculated as the difference of maximum and minimum sea ice area. A longer record would be needed to evaluate the correlation between bromine enrichment values and FYSI area and investigate a quantitatively link.

Table 3 shows the average annual iodine concentrations for each location, together with its standard deviation. The mean value (0.043 ppb) is close to the background values found in Antarctic shallow firn cores near the research stations of Neumayer (Frieß et al., 2010) and Casey (Law Dome, Spolaor et al., 2014) respectively.
Unlike previous observations of a clear winter peak of iodine with concentrations up to 0.6 ppb (Neumayer) and 0.3 ppb (Law Dome), no clear seasonality is observed for the transect records, with annual variability around 10-15%. Core 7 (Fig. 6) shows some variability which corresponds to winter peaks. High iodine concentrations are observed in core 8 during the 2012 winter, in association to a strong sea salt (sodium) input, although similar strong winter peaks are observed in 2011 at GV7.

The low background level and low variability of iodine found along the transect reflect a low input of iodine in this area of Antarctica compared to other locations. This picture is confirmed by satellite measurements, which show average IO concentrations close to detection limit over the area of the transect compared to Law Dome, Neumayer, or any other coastal location (Fig. 8, right panel). The high elevation of the traverse area, compared to the others is likely to play a role in preventing efficient iodine transport from the source areas.

Frieß et al. (2010) and Spolaor et al. (2014) have attributed iodine seasonal signal pattern to summertime photochemical recycling of IO from the snowpack, leading to depletion in the summer layers and higher concentrations in winter, when absence of sunlight inhibits photoactivation. The lower variability found across the Northern Victoria Land traverse cores could result from a reduced summer recycling due to low iodine concentrations available the snow.

3.3 Spatial flux variability

Glaciochemistry around Antarctica is very strongly influenced, among several properties, by the distance from the sea and the pathways of the air mass trajectories (Bertler et al., 2005). Atmospheric circulation patterns around the Talos Dome area have been investigated by Scarchilli et al. (2011), who have shown that the main input is represented by the Southern Ocean (Indian sector) with a lower contribution from the Ross Sea.

The spatial variability of sodium, bromine, bromine enrichment and iodine is investigated in Fig. 9. The twelve panels display the annual fluxes of Na, Br, I and integrated annual values of bromine enrichment for each core in relation to its distance from the Indian Ocean. Sodium fluxes show the highest values and variability around the closest locations to the Southern Ocean (GV7, 8, 7, 6), where the accumulation increases. After rapidly decreasing within the first 100 km, the sodium flux becomes stable, as the input from the SO decreases but the one from the Ross sea gradually increases. Bromine exhibits a similar behavior to sodium, with a homogeneous flux within cores 10 and 9 and an increase (up to 3 times) in the last 100 km from the SO. Elevation could partly account for the fractionation of sodium and bromine, having the 180 m of height difference separating GV7,8,7 and 6, and 240 m from GV7 to core 10. The effect of elevation yet is combined to the influence of the distance from the source to resolve the two effects. The pattern of bromine enrichment is linked among other things to the different bromine fractionations during the transport in the gas phase and the aerosol phase, compared to sodium. Unlike sodium and bromine, no decrease is observed for bromine enrichment from our data (Fig. 9, second column), although no clear trend can be inferred. This can be due to the multiple origins of air advection (Ross sea/Indian ocean), to their uneven strength or because the distances are not large enough for any difference to be observed.

A slightly lower fractionation after 100 km from the SO is observed for iodine, consistent with confirming the homogeneous satellite measurements of IO (Fig. 9, right).
4. Conclusions

The 2013/14 Talos Dome – GV7 traverse provided an opportunity to expand the existing sodium dataset in Victoria Land and investigate important features of bromine and iodine temporal and spatial variabilities, so far only available in Antarctica at Law Dome and Neumayer station.

The accumulation rates agree with previous studies, with increasing values from the Ross Sea to the Southern Ocean. Accumulation rates calculated for Talos Dome are higher than previously reported, likely caused by isotopic diffusion and remobilization at this site. Further studies are required at this site in order to access the reproducibility of the climate signal. The locations near the Southern Ocean exhibit high variability due to the higher accumulation.

Sodium and bromine concentrations in the snow samples result in a positive bromine enrichment to seawater, confirming the sea ice influence in the area for the extra bromine deposition. While sodium does not capture clear sub-annual variations associated with sea ice, bromine enrichment shows consistent seasonal variabilities with late spring maxima. There is some evidence that the seasonality is linked to the combined effect of sea ice growth and sunlight, which trigger photochemistry above fresh sea ice. The timing of deposition is coherent among Victoria Land, Law Dome and Dumont d’Urville (Indian sector) and Neumayer (Atlantic sector). Iodine shows an average value of 0.04 ppb, similar to background values observed in the Antarctic coastal locations of Law Dome and Neumayer. Unlike those locations, low iodine annual variability and no consistent seasonality of the signal are observed in the traverse samples.

The spatial variability study reveals homogeneous fluxes of Na, Br, and I over the transect length, with an increase in absolute values and variability at the sites close to the Indian Ocean, due to high accumulation and proximity to the coasts. Uniform satellite values of BrO and IO over Victoria Land are consistent with the snow measurements. A fractionation due to distance of these potential proxies is not found probably due to the combined double input of air masses from the Ross Sea and the Indian Ocean.

A transect covering larger distances and directed towards the interior of the plateau would give an insight on this feature, especially clarifying the spatial pattern of bromine enrichment with respect to differences in gas-phase and aerosol depositions.

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