

RESEARCH ARTICLE

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Key Points:

- Snowmelt onset hinders reactive bromine heterogeneous recycling and ends season of reactive bromine events
- Reactive bromine events occur at subfreezing air temperatures but not at higher temperatures
- Snow appears necessary for reactive bromine heterogeneous recycling, and rainwater can terminate this chemistry

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Snowmelt onset hinders bromine monoxide heterogeneous recycling in the Arctic

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Abstract Reactive bromine radicals (bromine atoms, Br, and bromine monoxide, BrO) deplete ozone and alter tropospheric oxidation chemistry during the Arctic springtime (February–June). As spring transitions to summer (May–June) and snow begins to melt, reactive bromine events cease and BrO becomes low in summer. In this study, we explore the relationship between the end of the reactive bromine season and snowmelt timing. BrO was measured by Multi-AXis Differential Optical Absorption Spectrometer at Utqiagvik (Barrow), AK, from 2012 to 2016 and on drifting buoys deployed in Arctic sea ice from 2011 to 2016, a total of 13 site and year combinations. The BrO seasonal end date (SED) was objectively determined and was compared to surface-air-temperature-derived melt onset date (MOD). The SED was highly correlated with the MOD ($N = 13$, $R^2 = 0.983$, $RMS = 1.9$ days), and BrO is only observed at subfreezing temperatures. In subsets of these sites and years where ancillary data were available, we observed that snowpack depth reduced and rain precipitation occurred within a few days of the SED. These data are consistent with snowpack melting hindering BrO recycling, which is necessary to maintain enhanced BrO concentrations. With a projected warmer Arctic, a shift to earlier snowmelt seasons could alter the timing and role of halogen chemical reactions in the Arctic with impacts on ozone depletion and mercury deposition.

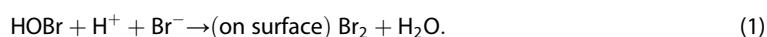
Plain Language Summary Reactive bromine events in the Arctic are common in spring and deplete ozone and cause mercury deposition. These events are affected by snow and ice, which are changing in the Arctic; therefore, we need to understand how environmental conditions affect reactive bromine chemistry. We find that the reactive bromine season ends when snowpack begins to melt. Through these full seasonal observations, we find that reactive bromine events occur to warmer temperatures than previously reported, with 0°C being the observed threshold above which reactive bromine is absent. We also find that snow appears necessary for reactive bromine chemistry and rain stops this chemistry. Earlier snowmelt in a warmer Arctic would end the reactive bromine season earlier, decreasing late springtime ozone depletion and mercury deposition.

1. Introduction

Reactive bromine species (Br and its oxide form, BrO) are strong tropospheric oxidizers [Platt and Hönninger, 2003; Simpson et al., 2007b, 2015; Abbatt et al., 2012]. During Arctic springtime, enhanced BrO mixing ratios have been observed episodically up to ~ 40 pmol mol⁻¹ [Pöhler et al., 2010] as compared to levels found in the marine boundary layer of ~ 1 – 3 pmol mol⁻¹ [Leser et al., 2003] or ~ 1 pmol mol⁻¹ in the free troposphere [Theys et al., 2011]. Ozone normally dominates tropospheric oxidation by providing photooxidants, such as the hydroxyl radical [Jacob, 2000]; however, during springtime reactive bromine rapidly consumes ozone in ozone depletion events [Bottenheim et al., 1986; Oltmans and Komhyr, 1986] and produces BrO [Fan and Jacob, 1992; McConnell et al., 1992; Hausmann and Platt, 1994]. In addition, during ozone depletion events, reactive bromine rapidly oxidizes mercury from Hg⁰ to Hg²⁺, which gets subsequently deposited, becoming more bioavailable [Schroeder et al., 1998; Holmes et al., 2006, 2010; Steffen et al., 2008]. As spring transitions to summer (i.e., late spring, May–June), however, observed BrO levels start decreasing and fewer ozone depletion events are observed [Halfacre et al., 2014].

Snow-covered sea ice and low temperatures dominate springtime conditions above the Arctic Ocean. Also, during springtime, satellite-based spectrometers (e.g., Global Ozone Monitoring Experiment (GOME), Ozone Monitoring Instrument (OMI), and Scanning Imaging Absorption Spectrometer for Atmospheric Chartography (SCIAMACHY)) detect BrO “hot spots” that occur primarily over frozen sea ice [Richter *et al.*, 1998; Wagner and Platt, 1998]. In late spring, the air temperature increases to above freezing and the snowpack starts melting, resulting in a sharp increase in the total snowmelt extent across the Arctic sea ice surface in late May to early June [Nghiem and Neumann, 2007]. At approximately the same time as snowmelt, the satellite-detected BrO hot spots become less common, and they are essentially absent by June [Van Roozendaal *et al.*, 2002]. Ground-based observations of particulate bromine at Barrow, AK, reported by Berg *et al.* [1983] showed enhanced particulate bromine levels (average 82 pptm, parts per trillion by mass) from February to April with decreasing values in May and very low levels (6 pptm) in June. In addition to observing BrO seasonality, ozone depletion events can be used as a proxy for enhanced BrO levels, due to the fact that reactive bromine catalytically destroys ozone. Oltmans *et al.* [1989] measured seasonal surface ozone at Igloodik, Barrow, Mold Bay, and Alert. The average behavior of ozone at Barrow had a winter maximum (November–February), decreased episodically in springtime (March–mid-May), and a secondary peak in June before decaying in summer [Oltmans *et al.*, 1989, 2012].

Current understanding of reactive bromine events in the Arctic [Platt and Hönninger, 2003; Simpson *et al.*, 2007b; Abbatt *et al.*, 2012] requires heterogeneous recycling of BrO to maintain reactive bromine levels. In the absence of recycling, reactive bromine species such as Br atoms would react with species such as formaldehyde (HCHO), producing HBr, which decreases reactive halogen levels, typically with a lifetime of a few hours. As proposed by chemical modeling groups [Fan and Jacob, 1992; McConnell *et al.*, 1992], the key chemical reaction for recycling is



In addition to recycling reactive bromine, reaction (1) is also central to the *bromine explosion* mechanism [Platt and Janssen, 1995; Wennberg, 1999] because it converts one inactive bromide (Br^-) ion to Br_2 , which photolyses, releasing two reactive bromine atoms. Bromide, the inactive species of bromine, is found in sea salt and resides on surfaces such as frozen sea ice, frost flowers (vapor deposited crystalline structures), sea salt aerosols, and snow throughout the Arctic [Abbatt *et al.*, 2012]. Two main requirements for BrO recycling are a high surface area to facilitate the reaction and the presence of bromide on those surfaces.

Laboratory studies have observed reaction (1), on halide-doped frozen surfaces and found that more acidic surfaces were more efficient at producing Br_2 [Huff and Abbatt, 2002; Wren *et al.*, 2013]. This was verified in the field by Pratt *et al.* [2013], who examined reactive halogen production from a snow chamber with snow and ice samples collected near Utqiagvik, AK. They found that only the top 1 cm of the snowpack above both sea ice and tundra produced Br_2 , when exposed to sunlight and ozone, as compared to sea ice and deeper snowpack layers. They also found that the top 1 cm of snow had a higher Br^-/Cl^- ratio and was more acidic, presumably due to atmospheric deposition of gas/particle-phase acids.

For near-surface BrO, snow is likely the predominant site for BrO recycling [Simpson *et al.*, 2007a; Abbatt *et al.*, 2012; Pratt *et al.*, 2013]. Measurements of Arctic snowpack surface area index (m^2 of snow per m^2 of ground) revealed the snowpack to have a very large surface area (up to $3000 \text{ m}^2 \text{ m}^{-2}$ [Dominé *et al.*, 2002]) compared to typical terrestrial soil or oceanic water. Dominé *et al.* [2002] and Domine *et al.* [2011, 2012] classified snowpack layers and measured the surface area of each layer during multiple Arctic field campaigns. Surface area measurements of Arctic snowpack layers revealed the top 1 cm of the snowpack to be generally surface hoar and/or diamond dust with an surface area index ranging from 50 to $120 \text{ m}^2 \text{ m}^{-2}$ and had lower total surface area compared to depth hoar (300 – $500 \text{ m}^2 \text{ m}^{-2}$) and wind-packed layers (1000 – $2500 \text{ m}^2 \text{ m}^{-2}$). The highest total surface area was measured in the wind-packed layers; however, these layers are very dense and less ventilated than surface and depth hoar layers [Albert and Shultz, 2002].

At the onset of snowpack melt, liquid water becomes present in snowpack, and there is a loss of surface area [Dominé *et al.*, 2002; Taillandier *et al.*, 2006; Domine *et al.*, 2007]. Ions elute downward in the snowpack [Johannessen and Henriksen, 1978; Bales *et al.*, 1989], although capillary forces may retain meltwater delaying ionic elution after melt onset [Wever *et al.*, 2014], and potentially redistribution of ions from atmospherically accessible surfaces to grain boundaries. Various studies have examined melt onset of snowpack in the Arctic

[Anderson, 1987; Winebrenner *et al.*, 1994; Kwok *et al.*, 2003; Nghiem and Neumann, 2007; Markus *et al.*, 2009]. Kwok *et al.* [2003] derived melt onset dates by synthetic aperture radar satellite imagery and compared the melt onset dates against individual buoy temperature measurements from the International Arctic Buoy Program, as well as passive microwave brightness temperatures. They reported that melt onset occurred within 1–2 days of the buoy temperature crossing over 0°C; however, the passive microwave brightness temperatures were biased toward a later melt stage. Markus *et al.* [2009] analyzed the minimum and maximum melt season length using the POLar Exchange at the Sea Surface temperature data. The beginning of the maximum melt season, which coincided with the early melt onset date, was the first day the surface temperature increased to above freezing, independent of it staying above freezing. The objective of this study was to understand how environmental conditions associated with the onset of the late spring melt period affect BrO recycling. We used an extensive data set, five spring seasons at Utqiagvik, AK, and eight spring seasons at on-sea-ice sites via O-Buoy measurements [Carlson *et al.*, 2010; Knepp *et al.*, 2010]. Where ancillary data were available, we explored the relationship between BrO abundance and environmental conditions such as temperature, snowpack depth, and precipitation (rain/snow).

2. Data Sources and Methods

2.1. Bromine Monoxide Measurement Sites and Methods

One Multi-AXis Differential Optical Absorption Spectrometer (MAX-DOAS) instrument was located near Utqiagvik, AK, at the Barrow Arctic Research Center (BARC) (156.6679°W, 71.3249°N), where we used 2012–2016 data, which used a consistent observational pattern. MAX-DOAS instruments were also mounted on O-Buoy platforms that were deployed into Arctic Ocean sea ice from 2011 to 2016. Figure 1 shows a map of the O-Buoy locations, and Table 1 shows their coordinates on specific dates. O-Buoy instruments and deployments were a part of the National Science Foundation-funded Arctic Observing Network (AON) project. O-Buoys were designed for year-round measurement of bromine monoxide, ozone, carbon dioxide, and meteorological variables such as temperature and wind [Carlson *et al.*, 2010; Knepp *et al.*, 2010]. O-Buoy deployments started in 2009, and a total of 15 O-Buoys have been deployed. Only a subset of deployed O-Buoy BrO measurements over the time range needed (April–June) is used for this study.

For the most straightforward comparison between these sites and years, we chose to use the directly measured BrO differential slant column density (dSCD), which was observed by MAX-DOAS [Hönninger *et al.*, 2004]. The dSCD is the difference in BrO column between a low elevation view, 1° elevation above the horizon, and the zenith, and is most sensitive to boundary layer BrO. While optimal estimation inverse modeling methods exist to convert dSCD profiles to vertical concentration profiles [Frieß *et al.*, 2011; Peterson *et al.*, 2015], these methods are complex, involve assumptions, and are sometimes numerically unstable. Of particular importance for this work, when observations of dSCD are near zero, the directly measured dSCD may be positive or negative, and the average accurately represents the observations. On the other hand, optimal estimation modeling only allows positive concentrations, so when BrO dSCDs are within noise levels of zero, the optimal estimation inversions contain a positive bias and the average may not correctly represent low BrO levels. Therefore, we chose to use the BrO 1° dSCD as a proxy for the presence or absence of BrO in the boundary layer. Enhanced BrO 1° dSCD measurements clearly indicate that BrO was present. Clouds, aerosol particles, fogs, and blowing snow can reduce visibility, which then reduces the differential pathlength between 1° elevation and zenith views and thus reduces the BrO 1° dSCD. Because reduced visibility periods are typically short-lived (<day), visibility will have little effect on detecting the presence or absence of BrO on multiday seasonal timescales. All BrO observations within $\pm 0.5^\circ$ of the nominal 1° elevation angle were averaged, and the data were averaged over regularized 3 h periods.

2.2. Snow/Rain Precipitation Observations and Snow Depth Measurements

Meteorological routine weather reports (METAR) at Utqiagvik Airport (PABR, 156.7922°W, 71.2826°N) were used to determine snow and/or rain precipitation events. METAR derived from the U.S. National Weather Service Automated Surface Observing System were obtained from the Iowa Environmental Mesonet (<http://mesonet.agron.iastate.edu>). Snow was indicated by the present weather code “SN” in more than one of the METAR observations in the 3-hourly averaging periods, and rain was indicated by the code “RA,” again in more than one METAR line. Snow depth was measured by Ice Mass Balance Buoys (IMB)

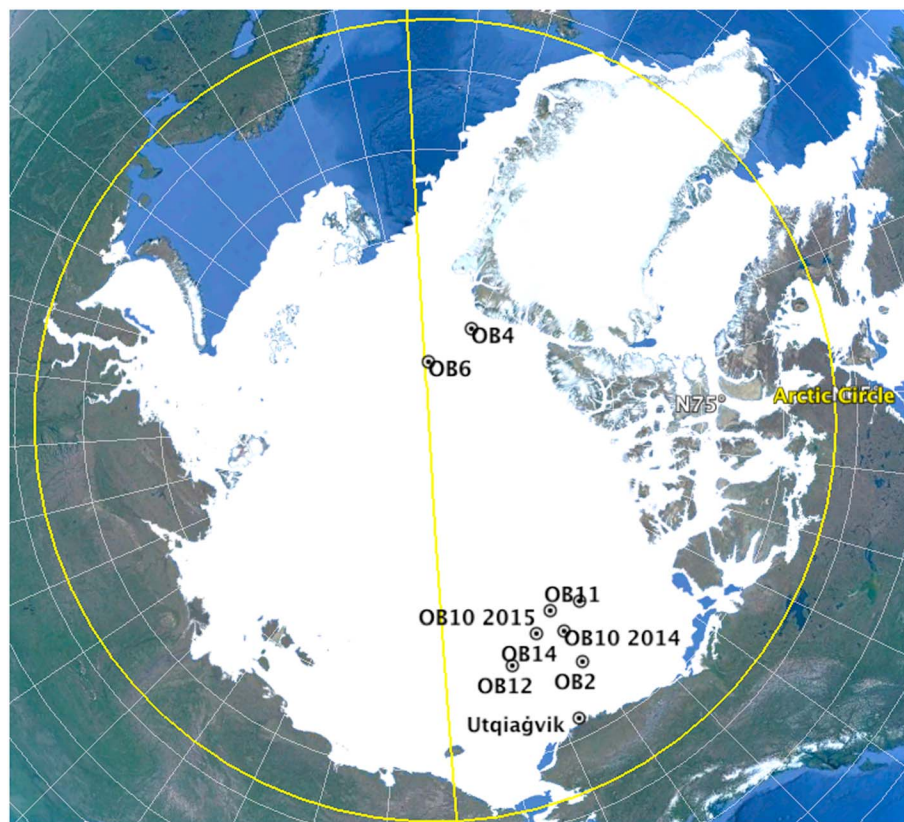


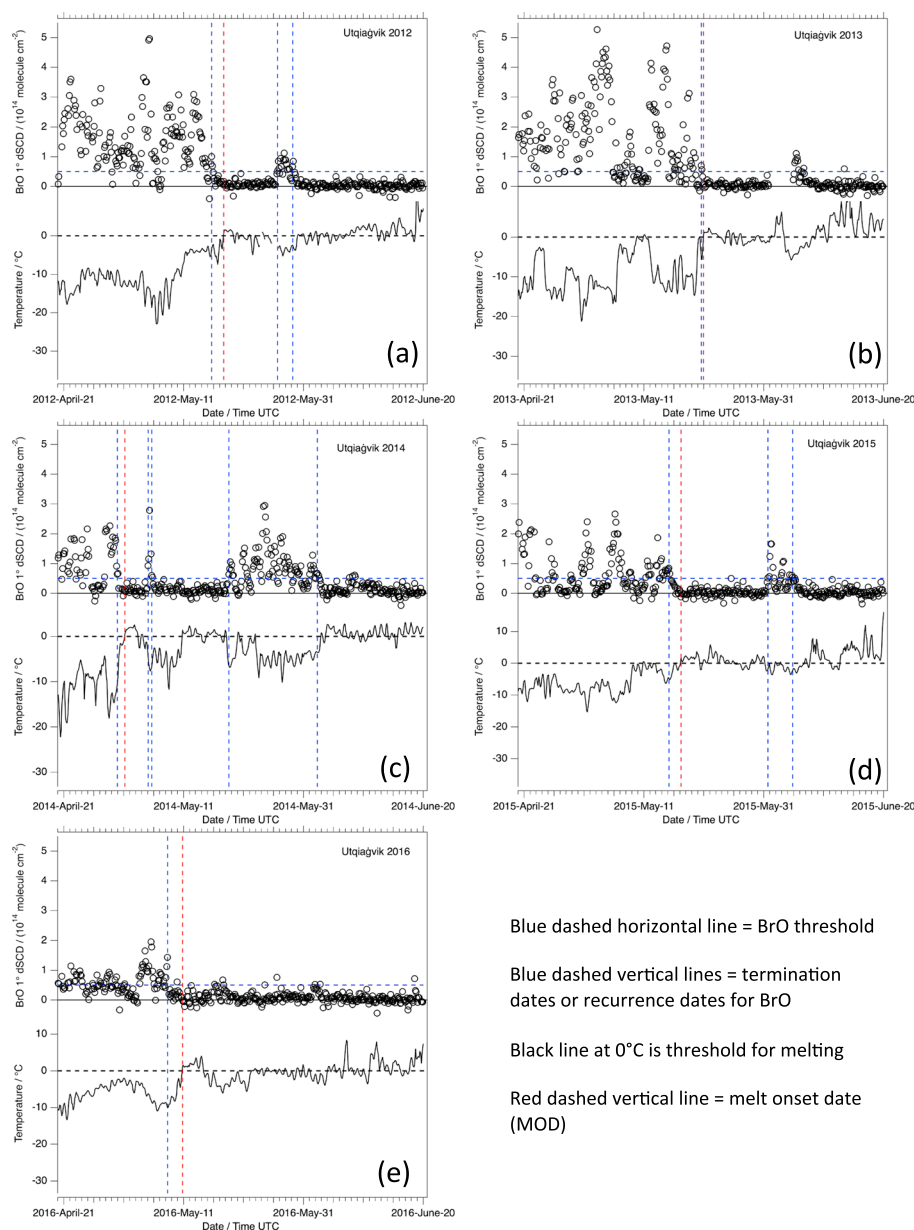
Figure 1. Locations sites used in this study overlaid on May 2015 Multisensor Analyzed Sea Ice Extent - Northern Hemisphere [Fetterer et al., 2010] sea ice extent data visualized in Google Earth. The O-Buoy locations are shown at the location of their seasonal end date, as listed in Table 1. The straight yellow line is 0° and 180° longitude, and the yellow arc is the Arctic Circle.

[Perovich et al., 2015]. An acoustic sounder was mounted on a pole frozen into the ice and positioned above the snow/ice surface. The acoustic sounder measured the distance between the instrument and the snow surface, with an accuracy of 5 mm, every hour, which was subsequently averaged to 3-hourly time resolution. Four O-Buoy-IMB seasonal comparisons were used in this analysis (refer to Table 1 for co-located instruments).

Table 1. Seasonal End Dates (SED) and Melt Onset Dates (MOD) for All Sites and Years^a

Site Year	IMB	Lat	Lon	SED (DOY)	MOD (DOY)	Note
Utqiagvik 2012		71.32	-156.6	15-May-2012 (136)	17-May-2012 (138)	
Utqiagvik 2013		71.32	-156.6	20-May 2013 (140)	20-May 2013 (140)	1
Utqiagvik 2014		71.32	-156.6	29-Apr-2014 (119)	30-Apr-2014 (120)	
Utqiagvik 2015		71.32	-156.6	14-May-2015 (134)	16_May-2015 (136)	2
Utqiagvik 2016		71.32	-156.6	7-May-2016 (128)	10-May-2016 (131)	
O-Buoy 2 2011	2010F	74.24	-150.9	12-Jun-2011 (163)	12-Jun-2011 (163)	3
O-Buoy 4 2012		84.51	-28.9	31-May-2012 (152)	31-May-2012 (152)	
O-Buoy 6 2012		86.80	-4.6	8-Jun-2012 (160)	10-Jun-2012 (162)	4
O-Buoy 10 2014	2013F	76.35	-150.6	11-Jun-2014 (162)	13-Jun-2014 (164)	5
O-Buoy 10 2015	2013F	77.76	-150.7	18-May-2015 (138)	18-May-2015 (138)	
O-Buoy 11 2015	2014I	77.20	-143.4	18-May-2015 (138)	19-May-2015 (139)	
O-Buoy 12 2015		75.83	-164.6	16-May-2015 (136)	18-May-2015 (138)	
O-Buoy 14 2016		77.02	-156.5	15-May-2016 (136)	11-May-2016 (132)	

^aThe IMB column has the name of the ice mass balance buoy. Latitude and longitude are in degrees, N and E being positive. The dates are local date at the time zone of the site, and the numerical day of year (DOY) is listed in parentheses. Note the following: (1) used third MOD for Utqiagvik 2013, (2) did not count oscillations on 11/12 May for Utqiagvik 2015, (3) did not count late May diurnal temperature oscillations for O-Buoy 2 2011, (4) used second MOD for O-Buoy 6 2012, and (5) did not count early may brief warming periods for O-Buoy 10 2014.



Blue dashed horizontal line = BrO threshold

Blue dashed vertical lines = termination dates or recurrence dates for BrO

Black line at 0°C is threshold for melting

Red dashed vertical line = melt onset date (MOD)

Figure 2. (top) BrO 1° elevation dSCD and (bottom) air temperature from April 20 to June 20 at Utqiagvik from 2012 to 2016 averaged in 3 h intervals. The BrO plot includes the threshold used for determining seasonal end date as a blue horizontal dashed line. The blue dashed vertical lines are located at termination and recurrence dates for BrO being above threshold. The horizontal black dashed line at 0°C is the threshold for melt onset. The red dashed vertical line is the melt onset date.

2.3. Objective Determination of BrO Termination and Recurrence Dates

Figure 2 (top) shows BrO 1° dSCD from 20 April to 20 June on five consecutive years at Utqiagvik. During this late season period, we observed that BrO began each season with high variability and values up to $\sim 5 \times 10^{14}$ molecule cm^{-2} . As the season progressed, each BrO record went through a transition where the BrO 1° dSCD became very small and scattered around zero. We call a date on which the BrO goes to zero a termination date. In some years, there is simply one termination date, but some years have a transient recurrence, where BrO is again nonzero, followed by a second termination date. We name the first termination date for a year the seasonal end date (SED).

We develop an objective algorithm to calculate termination and recurrence dates. This algorithm uses two parameters: the threshold below which BrO is considered zero and the duration required for BrO to be

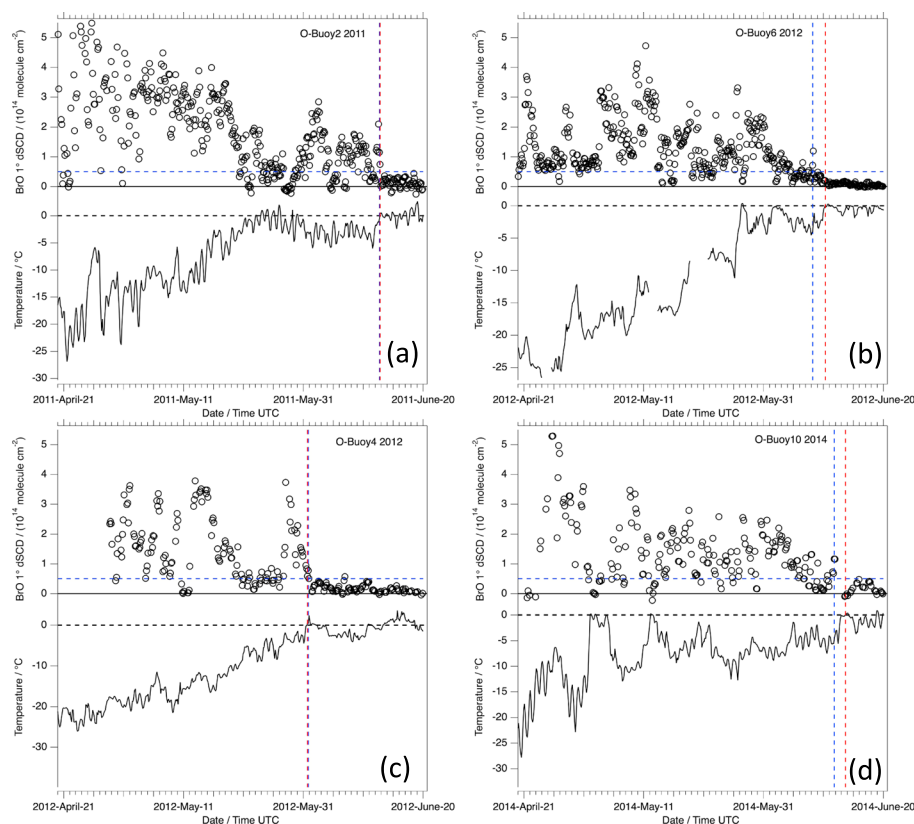


Figure 3. (top) BrO 1° elevation dSCD and (bottom) air temperature from four O-Buoy site/year combinations. The blue and black horizontal dashed lines are BrO and temperature thresholds, respectively. The blue dashed vertical lines are located at BrO termination and recurrence dates, and the red dashed vertical line is the melt onset date.

under this threshold. The threshold is chosen to be 5×10^{13} molecule cm^{-2} , shown as a blue dashed horizontal line on each top plot of Figure 2. The duration is chosen to be 5 days. The algorithm takes a termination date to be the last time when BrO 1° dSCD is above threshold such that not more than one 3 h average is above threshold in the next 5 days. A recurrence is defined as the first point after a termination when BrO is above threshold and remains above threshold for more than one more data point for the next 5 days. If there is a recurrence, there will then be another termination. The blue dashed vertical lines on these figures show the objectively determined termination and recurrence dates. Note that in 2013, there was a data gap that coincided with an apparent BrO recurrence. Because we did not effectively capture this event's timing, we did not report this recurrence.

2.4. Objective Determination of Melt Onset Dates Using Surface Air Temperature

Air temperature at two meters above ground level in Utqiagvik, AK, near the BARC building was obtained from NOAA Earth System Research Laboratory Global Monitoring Division (<http://www.esrl.noaa.gov/gmd/dv/data/>). Air temperature measurements on the O-Buoy instruments were obtained from the arcticdata.io repository (<http://arcticdata.io/>). The O-Buoy temperature sensor was roughly 2–3 m above the surface, depending upon snow depth and ice melt condition. All data are reported as 3 h averages. Based on Markus *et al.* [2009] and Kwok *et al.* [2003], we defined the first date on which the temperature reaches 0°C as the melt onset date (MOD), which is shown as a red dashed vertical line.

3. Results

Figures 2–4, show BrO and temperature records and visually indicate that the BrO termination dates were associated with warming above freezing. Because the temperature after the MOD often oscillates across freezing, we take the first date as the seasonal MOD, which is shown in Table 1 for each site and year. In some instances, we found that there were “early warming periods,” as reported by Winebrenner *et al.* [1994], who

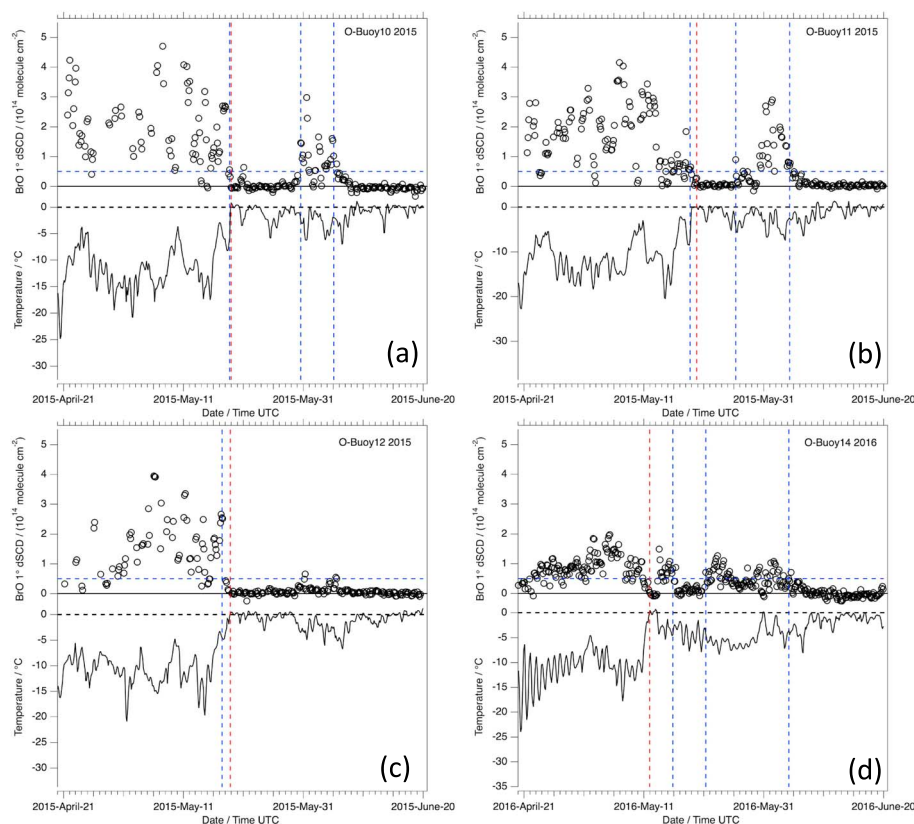


Figure 4. (top) BrO 1° elevation dSCD and (bottom) air temperature from four more O-Buoy site/year combinations. The blue and black horizontal dashed lines are BrO and temperature thresholds, respectively. The blue dashed vertical lines are located at BrO termination and recurrence dates, and the red dashed vertical line is the melt onset date.

detected snowpack melting using backscattering from European Remote Sensing 1 (ERS-1) synthetic aperture radar (SAR) satellite imagery. They reported that almost all melt onsets occurred when the temperature reached 0°C; however, in one case they saw multiple periods where the temperature reached 0°C before the melt onset date. They referred to these periods as “false start” of melting and explained that these short periods “did not lead to irreversible physical changes in those ice and snow properties that control backscattering at 5.3 GHz.” Given that the literature indicates that these early melting periods exist, in five of the 13 cases, we chose to select a later warming above 0°C for the MOD. Notes in Table 1 describe these cases, and they will be discussed in section 4.1. Figure 5a shows that the relationship between MOD and SED is nearly 1:1. A linear regression analysis shows a strong correlation ($R^2 = 0.983$, $N = 13$). The root-mean-square of the difference between the SED and MOD (SED–MOD) for all sites and years was 1.9 days. The average bias of the difference between SED and MOD was -0.8 days. Figure 5b shows the relationship between the BrO 1° dSCD and temperature, showing all valid data between 1 March and 3 July. This figure clearly shows that at subfreezing temperatures, BrO 1° dSCD was highly variable between 0 and 8×10^{14} molecule cm^{-2} , but that upon warming above freezing, BrO 1° dSCD scattered around 0, indicating that BrO was absent above 0°C.

In four sea ice case studies, O-Buoy MAX-DOAS measurements were co-located with Ice-Mass Balance Buoys [Perovich *et al.*, 2015], which allowed for an ancillary study of snow depth in relation to SED. Figure 6 shows that snow depth was fairly constant prior to the SED (first blue dashed vertical line); however, the snowpack height typically began decreasing within a few days of the SED. During recurrence events, which occurred at O-Buoy10 and O-Buoy11 in 2015 (Figures 6c and 6d), we observed that snow depth was stable.

Routine weather observations (METAR) at Utqiagvik, AK, allowed consideration of the weather relationship to the BrO end-of-season behavior. We observed that snow was quite common in this time of year, and rain became more common later in the season. In all years, rain events occurred within a few days of the seasonal end and recurrence dates. Also, we observed that during recurrence events, there was typically snow precipitation and not rain. In 2015, there was a transient rain event on 12 May, before the SED, but this event was

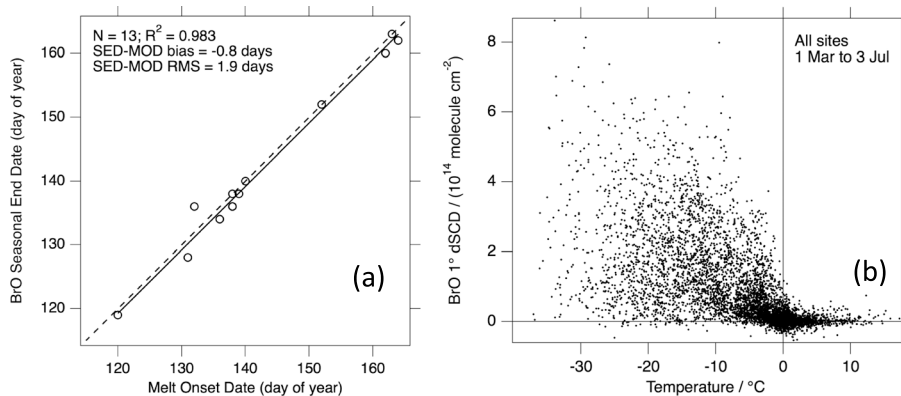


Figure 5. (a) The correlation between the BrO seasonal end date and melt onset date. The black dashed line is the 1:1 line, and the solid line is the orthogonal distance regression (ODR) best fit line. (b) The relationship between BrO 1° dSCD and ambient air temperature.

followed by snow and then a more significant rain period after the SED on 20 and 21 May. In 2016, there were two brief freezing rain events, on 20 and 28 April that preceded the SED, but then a more extended rain period from 10 to 12 May after the SED.

4. Discussion

4.1. The Seasonal End Date Is the Melt Onset Date

Figure 5a shows a strong correlation between SED and MOD. This finding is consistent with midseason studies that indicate that snowpack is a source of reactive halogen species and/or act as surface for recycling

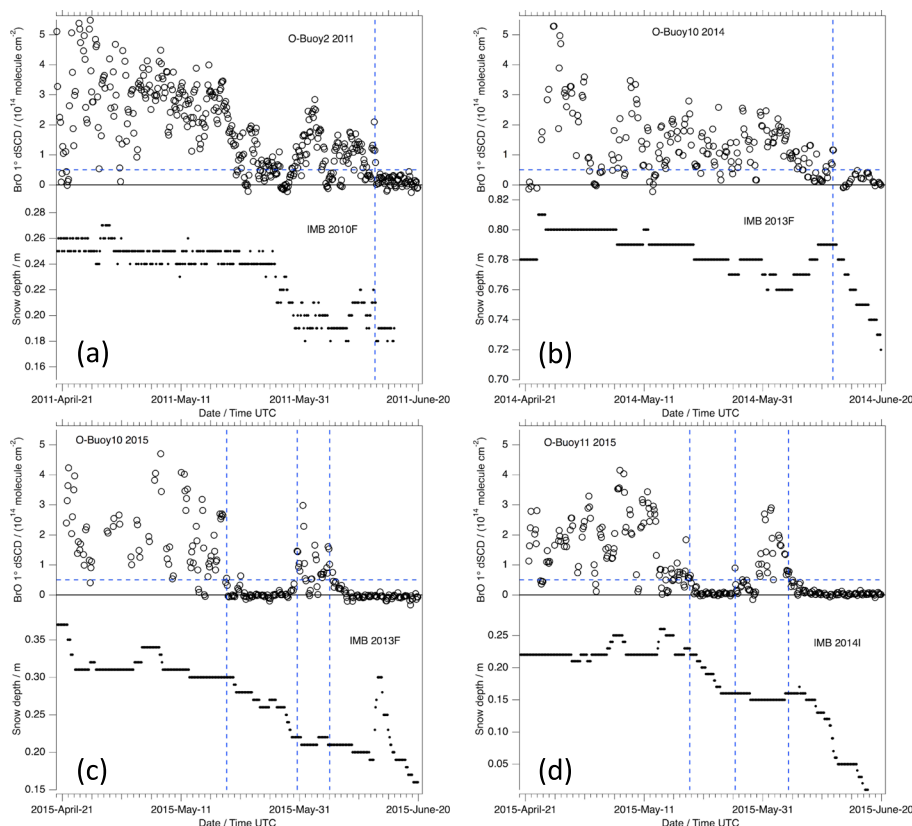


Figure 6. (top) BrO 1° dSCD and (bottom) snow depth for the four O-Buoys that were co-located with ice mass balance buoys (IMBs). The blue dashed lines indicate BrO termination and recurrence dates.

reactive halogens [Simpson *et al.*, 2007a; Pratt *et al.*, 2013]. The need for sufficient surface area is also indicated by studies of BrO aloft [Frieß *et al.*, 2011; Peterson *et al.*, 2017; Simpson *et al.*, 2017] that demonstrate that enhanced aerosol surface area is needed for BrO to be present aloft. The finding $SED \approx MOD$ is useful for predicting the end of the reactive halogen season. This is because satellite-based methods developed for detection of melt onset [Anderson, 1987; Winebrenner *et al.*, 1994; Kwok *et al.*, 2003; Nghiem and Neumann, 2007; Markus *et al.*, 2009] can be now extended to infer an end of the reactive bromine season. The end of BrO season behavior could then be used to validate satellite remote sensing of tropospheric BrO [Theys *et al.*, 2011; Choi *et al.*, 2012; Sihler *et al.*, 2012] on a Pan-Arctic scale.

The goal of the analysis here was to provide an objective means for determining SED and MOD to explore their relationship. However, we invoked literature to argue that some short warming events in five cases were not true melt onset dates. Therefore, it is useful to examine specific cases where either early melt onset was invoked or where the SED may not be perfectly determined by the objective algorithm. Figure 2b shows Utqiaġvik 2013 data, which was a case where we used the third date at which $T > 0^{\circ}\text{C}$ as the MOD. We see that the earlier warming events (10 and 11 May) were associated with subthreshold BrO, but the duration of the low BrO was not 5 days. This short duration caused the objective algorithm to not detect a SED during this warming. Had we manually selected this date as the SED, then there would have been recurrence event starting when the temperature drops and ending at the objectively selected SED. Therefore, the data are consistent with the observation that the reactive halogen season terminates when the temperature goes above freezing. Each of the five cases (Utqiaġvik 2013 and 2015, O-Buoy2 2011, O-Buoy6, 2012, and O-Buoy10 2014) shows the same behavior as just discussed; the early melt event was associated with low BrO 1° dSCD that was either too short in duration (<5 days) or near threshold to be detected as the SED.

Another notable case is that of O-Buoy 14 in 2016, shown in Figure 4d, which shows an early MOD followed by a later SED. The data show that at the time of the MOD, BrO undergoes a typical end-of-season transition to subthreshold levels, but again the duration is too short for the objective algorithm to call this date the SED. If we were to manually adjust these cases, the agreement between MOD and SED could be improved. However, the selection of one of these dates would then be dependent on the other date, so the correlation would be automatic and thus meaningless.

It is also evident from examination of the in-season (before the SED) behavior of BrO that low levels of BrO 1° dSCD can occur for a few days. Therefore, the SED is uncertain by a few days because BrO could have been subthreshold fortuitously, unrelated to snow condition, and we do not attempt to interpret the differences less than a few days. Similarly, past work has shown that MOD is not perfectly determined by temperature metric alone [Winebrenner *et al.*, 1994] but is significantly affected by infrared radiation budget and latent heat fluxes [e.g., Vionnet *et al.*, 2012], leading to the same caveat. Within this view of SED and MOD potentially having a few days uncertainty, we take the high correlation shown in Figure 5a as excellent indication that the melt onset occurs within a couple days of the termination of the reactive bromine season.

4.2. BrO Events Occur up to Freezing, but not at Above-Freezing Temperatures

Figure 5b clearly demonstrates that BrO events occur up to freezing temperatures, and not above freezing. Below freezing, there is a great deal of variability in BrO 1° dSCD, with an increase in the peak dSCD as the temperature decreases from 0°C to approximately -10°C and then a rather flat envelope. However, we note that the dSCD of BrO is an indicator for the detection of BrO, but not directly related to surface concentrations or the lower tropospheric vertical column density of BrO. In the future, work using optimal estimation modeling from these data should assist in quantitatively determining the relationship between temperature and bromine activation. Using data from a subset of these years at Utqiaġvik and two O-Buoys, Peterson *et al.* [2015] found that the lower tropospheric vertical column density of BrO was clearly enhanced at temperatures up to -5°C , in agreement with the analysis here on a more extensive data set. Bottenheim *et al.* [2009] showed that ozone depletion events, which are caused by reactive bromine chemistry, were observed at air temperatures up to -6°C , providing indirect evidence that reactive bromine can occur close to freezing temperatures. On the other hand, Pöhler *et al.* [2010] found surface BrO mixing ratio decayed to zero as temperature increased to -15°C , a much colder threshold than we observe. However, the Pöhler *et al.* [2010] study occurred during the peak reactive bromine season in March–April 2009, and only ever experienced temperatures up to -15°C . The episodic nature of reactive bromine events could have caused low

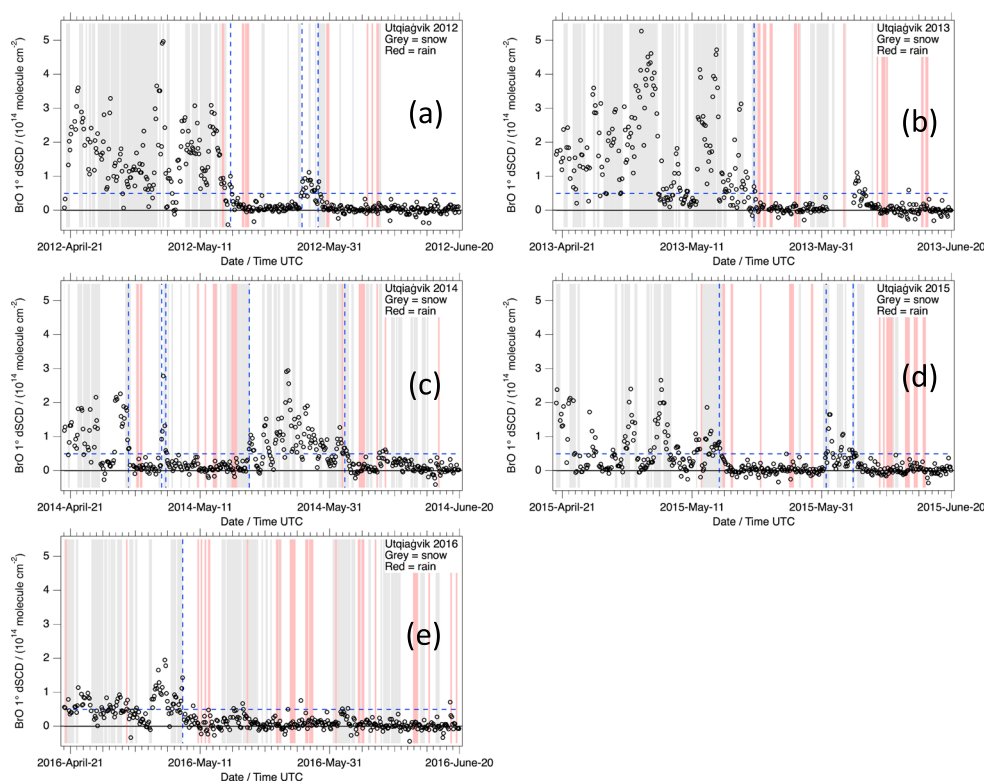


Figure 7. BrO 1° dSCD compared to precipitation observations from the Utqiagvik airport (PABR). During each 3 h interval, the grey background shading indicates >1 METAR code indicating snow, while the red shading shows >1 METAR code indicating rain. The two early season rain events during 2016 were freezing rain events.

BrO mixing ratio in that short-term study, which was interpreted as a threshold. It is clear from the data presented in the current study that BrO is present at temperatures above -15°C , and in fact up to freezing.

4.3. Decaying Snowpack and Rain Precipitation Hinders BrO Recycling

Figure 6 shows that periods of significant decrease in snowpack thickness are associated with subthreshold BrO levels. Figures 6c and 6d particularly show that after all of the termination dates (when BrO went below threshold), snowpack thickness was decreasing. This compaction of snowpack occurs via liquid water presence on snow grains, which causes coalescence and snowpack height decrease [Colbeck, 1982; Brun, 1989]. In one study, end of season compaction decreased the total surface area of a snowpack by an order of magnitude in a few weeks [Taillandier *et al.*, 2006]. In addition, the top of the snowpack may form a melt-freeze crust that has very little surface area [Domine *et al.*, 2007] and may significantly decrease the ventilation of the snowpack [Albert and Perron, 2000]. In addition to warming, rainwater from precipitation would induce snowpack decay due to the increase of liquid water content. Figure 7 shows that rainwater typically occurs within a few days after the BrO termination date. The Utqiagvik 2016 case, shown in Figure 7e, is interesting and unique in having two freezing rain events before the SED. During both of these events BrO is near to below threshold, consistent with either low surface area or decreased through-snow ventilation [Albert and Perron, 2000]. Note by comparing Figure 2e to the prior years (Figure 2, prior panels) that BrO was lower during April 2016 than other years. Potentially, 2016 could show the result of an earlier spring to summer transition [Stone *et al.*, 2002; Markus *et al.*, 2009] as lower BrO during this transitional period due to earlier melting and freezing rain.

4.4. Below Freezing Temperatures and New Surface Area Reinitialize BrO Recycling

All recurrence events were associated with subfreezing temperatures, as evident from Figures 2–4. However, there were periods where the temperature was below freezing, but no recurrence event was observed. Key examples of subfreezing periods that lack BrO are Utqiagvik 7–11 May 2014 (Figure 2c) Utqiagvik 14–21 May

2016 (Figure 2e), O-Buoy12 31 May to 14 June 2015 (Figure 4c), and O-Buoy14 15–20 May 2016 (Figure 4d). These periods indicate that simply being below freezing is not sufficient to regain the capacity to recycle BrO. From the cases at Utqiaġvik where recurrences did occur, 2012, 2014, and 2015, Figures 7a, 7c, and 7d show that snow precipitation preceded the recurrence events. It is likely that the recurrence in early June at Utqiaġvik in 2013 (Figure 7b), which was not well captured due to a data gap, also fits this pattern of snow precipitation before BrO recurrence.

4.5. What Property of Snowpack is Changed at the BrO Season End?

Snowmelt involves multiple coupled phenomena that confound interpretation of why the occurrence of snowmelt is correlated with BrO season termination. When snow begins to melt, the surface goes from solid to liquid, and liquid surfaces have been shown to often have lower heterogeneous reactivity than solid ice [Kahan *et al.*, 2010a, 2010b; Domine *et al.*, 2013]. This diminished reactivity upon liquid surface formation can explain the rapid loss of BrO upon melting. However, we observe a number of cases (see section 4.4) where subfreezing temperatures are experienced after melt onset, but the refrozen snow does not recycle BrO. A possible explanation for these cases is that while in the surface-melted state, bromide (Br^-) ions could redistribute from atmospherically accessible snow surface locations to thermodynamically favored grain boundaries [Baker *et al.*, 2003; Bartels-Rausch *et al.*, 2014] or other locations that are less accessible for heterogeneous chemistry. Refrozen and metamorphosed snowpack would have significantly lower surface area than fresh fallen snow [Domine *et al.*, 2007]. Similarly, a melt-freeze or freezing rain crust on top of snowpack would drastically decrease ventilation throughout the snowpack [Albert and Perron, 2000] and would reduce wind pumping that would allow for Br_2 to be released from the interstitial space of the snowpack [Pratt *et al.*, 2013]. Both the loss of surface area and hindered ventilation would slow BrO recycling, which is a potential cause in the termination behavior when the snowpack begins to melt. Another potential contributor to removal of bromide ions is the “ionic pulse” phenomenon [Johannessen and Henriksen, 1978; Bales *et al.*, 1989], which describes the observation of high ionic impurity concentrations in initial runoff from melting snow. However, actual percolation throughout the snowpack and runoff is likely to be a slower process than top surface melting, which would be inconsistent with the immediacy of BrO termination at MOD. Therefore, it seems more likely that decreased reactivity due to liquid surfaces, loss of surface area, and/or hindered ventilation would be responsible than extensive ion runoff. Which of these factors can be the most important remains to be determined, but all lead to a similar effect—that BrO recycling is hindered upon snowmelt. Laboratory and field manipulation studies that separate these coupled phenomena are encouraged to improve mechanistic understanding of reactive bromine recycling.

5. Conclusion

Melt onset, snowpack depth, and rain/snow precipitation were investigated to understand their effects on BrO recycling. We objectively determined the seasonal end date (SED) to the reactive bromine season, as well as the temperature-based melt onset dates (MOD) for 13 cases from Utqiaġvik, AK, and O-Buoy observations from 2011 to 2016. The strong correlation of the SED occurring within a few days of the MOD indicates these phenomena are strongly coupled. Correlation of BrO abundance with temperature clearly shows that BrO is only present at subfreezing temperatures, and also that BrO is above zero within a few degrees below freezing. From these observations, we find that melt onset hinders BrO recycling and ends the reactive bromine season. Ancillary snowpack depth analysis at sea ice sites revealed that the snow depth decreased within a few days of the SED and MOD, which supports that melt onset of the snowpack inhibits BrO recycling. Rain precipitation data at Utqiaġvik similarly showed that the rain occurred a few days after the SED and suggest that again there is a loss of snowpack surface area due to liquid water, which is hindering BrO recycling. New snow precipitation observations at coastal sites revealed that snow may be necessary to reinitialize BrO recycling but it is not sufficient alone.

The finding that the start of the melt onset season is the end to the BrO season gives us an indication on how BrO may respond to future climate changes. Long-term studies on the summer melt season found that the melt season has lengthened ~8 days per decade for Utqiaġvik, AK [Stone *et al.*, 2002], and ~5 days for the pan-Arctic region [Markus *et al.*, 2009]. These summer melt trends and our results indicate that as the melt season lengthens, the BrO season will get shorter, which would reduce the amount of ozone depletion events and mercury deposition during late spring.

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