Polar nighttime chemistry produces intense reactive bromine events

To cite this version:

HAL Id: insu-01877525
https://hal-insu.archives-ouvertes.fr/insu-01877525
Submitted on 3 Sep 2020

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
Polar nighttime chemistry produces intense reactive bromine events


1Geophysical Institute and Department of Chemistry and Biochemistry, University of Alaska Fairbanks, Fairbanks, AK, USA, 2Institute of Environmental Physics, University of Heidelberg, Heidelberg, Germany, 3LATMOS/IPSL, UPMC Univ. Paris 06 Sorbonne Universités, UVSQ, CNRS, Paris, France

Abstract By examining the origin of airmasses that arrive at Utqiaġvik (formerly Barrow), Alaska, soon after polar sunrise (late January/early February), we identified periods when air arriving at Utqiaġvik had previously resided primarily at higher latitudes in near total darkness. Upon illumination, these airmasses produced high concentrations of reactive bromine, which was detected by differential optical absorption spectroscopy as bromine monoxide (BrO). These observations are consistent with nighttime production of a photolabile reactive bromine precursor (e.g., Br2 or BrCl). A large polar night source of photolabile reactive bromine precursors would contribute seed reactive bromine to daytime reactive bromine events and could export reactive halogens to lower latitudes and the free troposphere.

Plain Language Summary During the spring in the polar regions, unique halogen oxidizers dominate atmospheric chemistry, altering the fate of pollutants such as mercury. The sources of these oxidizers are not well understood, particularly during polar sunrise. Here we report the largest concentration ever detected of one of these species, bromine monoxide (BrO). We find these high concentrations occur when airmasses come out of the polar night, indicating a nocturnal source. Nocturnal production of reactive halogens could act as a seed source for subsequent reactive halogen photochemistry and could export reactive halogens from the polar night all winter.

1. Introduction

Polar springtime reactive halogen events (first reported by Tuckermann et al., 1997) and their impact on ozone depletion (first reported by Barrie et al., 1988) and mercury deposition (first reported by Schroeder et al., 1998) have been known for decades. Satellite observations show that reactive bromine is present in both the Arctic and Antarctic sea ice regions (Chance, 1998; Richter et al., 1998; Wagner & Platt, 1998). However, the mechanism of production and maintenance of high levels of reactive halogens, initiators of these events, and their environmental controls are still elusive (Abbatt et al., 2012; Simpson et al., 2015). A number of field campaigns (Frieß et al., 2011; Peterson et al., 2015; Pöhler et al., 2010; Pratt et al., 2013; Simpson et al., 2017) have been carried out to investigate this phenomenon, but few of those studies covered the full halogen activation season. The Canadian Polar Sunrise Experiments and the ALERT2000 field campaigns were exceptions, observing before and after polar sunrise, which is the time when the sun rises above the horizon for the first time since the prior fall. A key finding from ALERT2000 was that Br2 and BrCl were produced in high concentrations right at the time of polar sunrise (Foster et al., 2001), potentially pointing to a nocturnal source of photolabile halogen species. More recently, studies using autonomous instrumentation and/or long-term monitoring sites have expanded our knowledge of reactive halogen events outside of the traditional peak months of March and April (Burd et al., 2017; Peterson et al., 2015, 2016). Specifically with respect to polar sunrise, Stohl (2006) showed that wintertime polar airmasses can remain for days to weeks in total darkness before being exposed to sunlight, which typically occurs by transport to lower latitudes. Therefore, we measured atmospheric BrO in late January/early February from 2013 to 2018 to quantify reactive bromine levels and couple those measurements with transport modeling to characterize airmass histories prior to sampling.

2. Methods

We calculated 3-day back trajectories using Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT; Stein et al., 2015) and 1° resolution Global Data Assimilation System meteorology that arrive 50 m above sea
level at the Barrow Arctic Research Center (BARC) building (71.3°N 156°W) near Utqiagvik, Alaska, USA, for every hour in late January/February. At each hourly location along this trajectory, we calculated the local solar zenith angle that the parcel would have experienced. Because reactive bromine is produced by photolysis of precursors such as molecular bromine (Br2), we calculated the photolysis rate coefficient for molecular bromine J (Br2) using a clear-sky parameterization that is a function of solar zenith angle (Madronich, 2017; Simpson et al., 2002; see supporting information). The product of J (Br2) times the duration of that time step (hour) is a unitless number describing the average number of photolytic exposures a Br2 molecule would have experienced. We summed the photolysis over the 3-day duration to give a trajectory-integrated clear-sky photoexposure for that airmass at the arrival hour.

HYSPLIT calculations were used for routine hourly calculation of photoexposure throughout this period because they are rapid; however, HYSPLIT does not model the dispersion and mixing of airmasses. Therefore, we also use the FLEXible PARTICle dispersion model (FLEXPART) version 9.02 (Stohl et al., 1998, 2005; Stohl & Thomson, 1999) to study the potential emission source regions influencing measurements at Utqiagvik on specific dates of interest. For this application, FLEXPART is run in backward mode for specific days of interest with particles released at the surface from Utqiagvik (0–20 m above ground level) during a 1-hr period at local noon (release from 20:30 to 21:30 UTC, 11:30 to 12:30 AKST). We release 100,000 particles and follow them back for 10 days to have an airmass history. To drive the FLEXPART model we use the European Centre for Medium-Range Weather Forecasts (ECMWF) ERA-Interim reanalysis meteorology on a 0.75° × 0.75° global grid (60 vertical levels) every 6 hr (00:00, 06:00, 12:00, and 18:00 UTC). We use the plume centroid locations to calculate the photochemical exposure analogous to the HYSPLIT method described above. We construct total column (0–20 km) and surface (0–200 m) potential emissions sensitivities (PESs) for 10 days. The total column PES indicates the region the air resided for the 10 days prior to arriving at Utqiagvik. The surface (0–200 m) PES indicates where air was in contact with the surface and potentially sensitive to surface emissions.

Bromine monoxide was detected by multiple axis differential optical absorption spectroscopy (MAX-DOAS) using an instrument on the roof of the BARC building via methods described previously (Frieß et al., 2011; Peterson et al., 2015; Simpson et al., 2017) to model BrO lower tropospheric vertical column density (LT-VCD) and 0- to 200-m mixing ratio (MR), the results of which are shown in Table 1 and in supporting information Figure S3. The magnitude of these reactive bromine events are high compared to past observations (Morin et al., 2005; Peterson et al., 2015; Simpson et al., 2007), and 90 pmol/mol is double the prior highest observed surface MR (Pöhler et al., 2010). Although it has been noted that MAX-DOAS observations of surface MR can be systematically high during aerosol particle-rich or cloudy conditions (Frieß et al,
five of these six dates were low-aerosol cases with typical aerosol optical depth $< 0.2$ (supporting information Figure S3).

Figure 2 shows a FLEXPART simulation for particles arriving at BARC on 9 February 2017 and demonstrates that this airmass transported south from higher latitudes, where it was still polar night, before arriving at Utqiaġvik. The shaded area shows the location of total polar night at the midpoint during the 10 days of the FLEXPART run, 4 February 2017. A great deal of the potential emission sensitivity of this simulation lies within this region of polar night, particularly the Arctic Ocean and adjacent Russian coastal areas, demonstrating that the air arriving at BARC had large influences from air that spent up to 10 days in polar night.

To examine the relationship between reactive halogen (BrO) events and prior airmass exposure to light, Figure 3 shows these quantities in the period 1–20 February 2017. The years 2013–2016 and 2018 are shown as supplemental figures. Top panel in Figure 3 shows that the minimum of the clear-sky photoexposure occurs on 9 February 2017 (AKST), with the airmass experiencing about 20 photoexposures on the prior part of the trajectory and that the airmass had been in near total darkness.

Figure 3 lower panel shows that BrO is highly elevated on this date, with a peak 1° dSCD of about $17 \times 10^{14}$ molecules per square centimeter. Other BrO events occur after this first event, as is common during springtime, but those airmasses had been exposed to significantly more sunlight (photoexposure) before their arrival at BARC, so they could have had BrO produced by typical springtime photochemistry as is common in March–May in the Arctic (e.g., Frieß et al., 2011; Hönninger & Platt, 2002; Peterson et al., 2016; Pöhler et al., 2010). The first event of the season, however, experienced only about 20 calculated clear-sky photoexposures before arrival at BARC, so somehow produced large amounts of BrO with relatively little prior photochemistry. Examining the other cases of first arrival of a polar night airmass in the years 2013–2016 and 2018 summarized in Table 1 and shown in supporting information Figures S5–S9, we see that the first event of the year commonly occurs with very low ($< 20$ clear-sky photoexposures) prior exposure to photolysis and is associated with high BrO abundance (Table 1). MAX-DOAS analysis also shows that NO$_2$ is low (within error of 0) on these dates, assuring that we are not sampling local pollution from Utqiaġvik.

Late January/February ozone observations (supporting information Figure S10) were typically at background levels (30–40 nmol/mol) in the early part of the month and ozone depletion events (O$_3$ $< 20$ nmol/mol) do not typically occur until the last week of February, indicating that photochemistry is weak at the time of these late January/early February large BrO events. Specifically, the polar night airmass arrival dates (Table 1) showed non-depleted ozone levels (25–37 nmol/mol) at sunrise (see supporting information Figure S3) and ozone decay rates between 0 and $\sim 4$ nmol·mol$^{-1}$·hr$^{-1}$. The largest ozone loss rate occurred on 9 February 2017, the date of highest surface BrO MR. The non-depleted ozone levels on these dates indicates that the airmasses had not been highly processed by reactive halogen chemistry prior to arrival at Utqiaġvik and are consistent with the airmasses being in the dark before arrival.

Table 1. Arrival Dates of First Polar Night Airmasses With Peak BrO dSCD, LT-VCD, and MR and Photoexposures Prior to Arrival Day

<table>
<thead>
<tr>
<th>Date (AKST)</th>
<th>BrO 1° dSCD (molecules per square centimeter)</th>
<th>BrO LT-VCD (molecules per square centimeter)</th>
<th>BrO 0- to 200-m MR (pmol/mol)</th>
<th>Prior photoexposure</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Feb 2013</td>
<td>$9 \times 10^{14}$</td>
<td>$5 \times 10^{13}$</td>
<td>48</td>
<td>$\sim 10$</td>
</tr>
<tr>
<td>7 Feb 2014</td>
<td>$7 \times 10^{14}$</td>
<td>$4 \times 10^{13}$</td>
<td>35</td>
<td>$\sim 5$</td>
</tr>
<tr>
<td>3 Feb 2015</td>
<td>$3 \times 10^{14}$</td>
<td>$3 \times 10^{13}$</td>
<td>20</td>
<td>$\sim 20$</td>
</tr>
<tr>
<td>6 Feb 2016</td>
<td>$5 \times 10^{14}$</td>
<td>$4 \times 10^{13}$</td>
<td>32</td>
<td>$\sim 20$</td>
</tr>
<tr>
<td>9 Feb 2017</td>
<td>$17 \times 10^{14}$</td>
<td>$13 \times 10^{13}$</td>
<td>90</td>
<td>$\sim 20$</td>
</tr>
<tr>
<td>30 Jan 2018</td>
<td>$10 \times 10^{14}$</td>
<td>N/A</td>
<td>55</td>
<td>$\sim 10$</td>
</tr>
</tbody>
</table>

Note. dSCD = differential slant column density; LT-VCD = lower tropospheric vertical column density; MR = mixing ratio.

Figure 1. Spectral confirmation of BrO on 9 February 2017 using a composite of all daytime spectra. The BrO differential slant column density is $1.47 \times 10^{15}$ molecules per square centimeter. See supporting information Figure S4 for fits to other gases.
To consider the degree to which these airmasses resided in the dark, we used the 10-day FLEXPART simulations to calculate photoexposure on longer timescales, where dispersion of the airmass will become more important. Figure 2 shows that the plume center remained for 10 days within the region of total polar night and that a large amount of the surface potential emission sensitivity was also within this circle of darkness. In the supporting information, Figure S11, we calculated integrated photoexposure for the peak BrO day on each year. We find that the exposure is between 4 days (2014) and >10 days (2013 and 2017) for <30 integrated Br₂ photoexposures. The lack of exposure to sunlight on this timescale is also visible in supporting information Figures S12 and S13, which show the FLEXPART simulations for each year.

4. Discussion

Reactions of nocturnal nitrogen oxides (e.g., N₂O₅) with NaBr (sea salt bromide) that produce photolabile bromine precursors (BrNO₂) have been studied and their atmospheric importance has been considered (Finlayson-Pitts et al., 1990). Similar reactions of N₂O₅ with chloride-containing particles that produce photolabile nitryl chloride have been definitively observed in the field through mass spectroscopy (Osthoff et al., 2008; Thornton et al., 2010), providing further evidence that nitrogen oxides can activate halogens through dark reactions (reactions not involving photolysis). Our MAX-DOAS observations episodically identified local NOₓ pollution (coming from Utqiagvik, which lies 5 km southwest of BARC) at the >1-nmol/mol level, but these specific dates of polar night airmass arrival are below NOₓ detection limits (estimated to be ~200 pmol/mol), consistent with the airmasses arriving from the polar ice cap. However, below detection levels of NOₓ, potentially from local pollution or produced by snowpack photochemistry before the airmass entered polar night (Honrath et al., 2002; Jones et al., 2001) or past exposure of these airmasses to NOₓ could
have produced photolabile bromine species via this mechanism. Therefore, dark reactions of nitrogen oxides with sea salt could be a mechanism consistent with our observations of high BrO in airmasses arriving from the polar night.

Oum et al. (1998) proposed that ozone oxidizes bromide (Br\textsuperscript{−}) on ice surfaces to produce Br\textsubscript{2}, providing another potential dark mechanism. Similar chemistry involving ozone oxidation of iodide at the sea water interface has been demonstrated (Carpenter et al., 2013) and provides analogous support for the mechanism of ozone oxidation of halides. Hirokawa et al. (1998) also observed that NaBr reacts with O\textsubscript{3} in the presence of water to produce Br\textsubscript{2}, and Hunt et al. (2004) provided evidence that this reaction may occur via a surface complex. Oum et al. (1998) estimate 0.6-pmol/mol Br\textsubscript{2} could be produced in 10 hr at an ozone exposure of 40 nmol/mol. Therefore, airmasses that remain in the dark for 10 days could produce an estimated 14-pmol/mol Br\textsubscript{2}, which would photolyse and react with ozone to produce ~28-pmol/mol BrO, which is typically less than, but within an order of magnitude of, the observed BrO levels. In addition, polar airmasses may reside in the dark for longer than 10 days due to the efficiency of the Polar Dome in inhibiting transport to lower latitudes, (Klonecki et al., 2003; Stohl, 2006) so even very slow dark reactions may be important. Further laboratory and modeling investigations of dark production mechanisms is clearly necessary, but it appears that these or other unknown mechanisms could produce photolabile bromine precursors that could explain the high BrO concentrations observed here.

Mass-spectroscopic field investigations (Foster et al., 2001) have shown evidence for production of photolabile bromine species at the time of polar sunrise near Alert, Canada. Foster et al. (2001) observed Br\textsubscript{2} MRs of up to 25 pmol/mol and BrCl of up to 35 pmol/mol. Production of either Br\textsubscript{2} or BrCl in the dark could build up a reservoir of photolabile bromine species over time, which then release reactive bromine upon exposure to sunlight through reactions 1 and 2.

Figure 3. BrO observations related to prior exposure of airmasses to photolysis. The top panel shows the clear-sky photo-exposure calculated along a HYSPLIT 3-day back trajectory (red line) and on the day of arrival at BARC (blue dashed line). The bottom panel shows BrO dSCD as a function of view elevation angle versus time. dSCD = differential slant column density.
Photolysis of Br₂ and BrCl at levels observed by Foster et al. (2001) followed by reaction of Br radicals with ozone would lead to 50- and 35-pmol/mol BrO, respectively, in qualitative agreement with the observed levels in this study (Table 1).

Typical springtime reactive bromine release is believed to proceed through reactions 1 and 2 followed by reaction of BrO with radicals (e.g., HO₂) to reform Br₂ in sequence below (Fan & Jacob, 1992; McConnell et al., 1992):

\[
\begin{align*}
\text{Br}_2 + \text{hv} & \rightarrow 2\text{Br} \quad (1) \\
2\text{Br} + \text{O}_3 & \rightarrow \text{BrO} + \text{O}_2 \quad (2)
\end{align*}
\]

\[
\text{BrO} + \text{HO}_2 \rightarrow \text{HOBr} + \text{O}_2 \quad (3)
\]

\[
\text{HOBr} + \text{HBr (on surfaces)} \rightarrow \text{Br}_2 + \text{H}_2\text{O} \quad (4)
\]

This *bromine explosion* (Platt & Janssen, 1995; Wennberg, 1999) sequence only amplifies reactive bromine during daytime, so it would not be responsible for nocturnal formation. It is possible that nocturnally formed reactive bromine precursors could be amplified on the day of detection via faster daytime photochemistry (Nissenson et al., 2014), but Figures 3 and S3 shows that most of these cases do not show an increase in BrO over the detection day. This traditional mechanism is also likely to be slowed under the low solar elevation conditions of late January/early February because HO₂ is produced photochemically and would be expected to be at low concentrations, so reaction 3 is likely to be slow. However, it is possible that there is nocturnal formation of HO₂ precursors that could photolyse to HO₂ radicals, which are then converted to reactive bromine radicals via reactions 4 + 1. To the extent that HOBr may be formed by these reactions or other dark mechanisms, it is likely to be converted to Br₂ by reaction 4, which has been shown to be fast in multiple studies (Fickert et al., 1999; Huff & Abbatt, 2002; Wachsmuth et al., 2002). Recently Nerentorp Mastromonaco et al. (2016) suggested a dark source of bromine could be responsible for observed mercury and ozone depletion in the Antarctic wintertime.

Ozone depletion occurs through reaction 2 followed by reactions of BrO that reform Br₂, but because the sun is very low at this time of year, photolysis of Br₂ (reaction 1) limits the ozone depletion process. For example, on 9 February 2017, Br₂ experiences approximately 50 photoexposures, causing reaction 2 to occur 50 times for each BrO present (~90 pmol/mol), depleting 4.5-nmol/mol O₃, which is consistent with the small ozone depletion observed on this date. Other dates have less reactive bromine and lower photoexposure, which is consistent with the lower observed ozone loss rates. Overall, these considerations indicate that these early season BrO events are different from typical ozone depletion events that occur under higher photoexposure in March–May. For these reasons, it appears that the formation of high BrO abundances in polar night airmasses is due to a dark mechanism rather than traditional bromine release chemistry.

**5. Conclusions**

We observe unprecedented levels of BrO, up to 90 pmol/mol and tropospheric vertical column density of \(1.3 \times 10^{14}\) molecules per square centimeter during February 2017 at Utqiagvik, Alaska. Four of the other five maxima also exceeded the 90th percentile of BrO observations from March–May 2012 to 2016, indicating that late January/early February often has episodes of very high BrO. Back trajectory analysis shows that these peak BrO measurements occurred in air that received minimum prior photolysis of Br₂, indicating a nocturnal source of reactive bromine.

The observation of a nighttime formation mechanism of photolabile bromine species is important because it would imply that the wintertime Arctic Ocean could be producing high concentrations of photolabile bromine gases that would then transport southward to sunlight and release reactive bromine species. Although some of this photolabile bromine reservoir will be trapped under the polar dome, vertical mixing could loft these photolabile species where they could then inject reactive bromine into the lower-latitude free troposphere, potentially enhancing the free-tropospheric BrO loading. Nocturnally produced photolabile bromine would also provide seed reactive bromine for the traditional photochemical bromine release
mechanism on subsequent days. Direct measurement of Br2 and BrCl in nighttime airmasses and determination of the production mechanism and its rate are needed.

References


SIMPSON ET AL. 9993


