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Key Points:

- A combination of factors including snow emissions, vertical mixing, and atmospheric chemistry explain surface Arctic halogen observations
- Snow emissions of halogens impact atmospheric chemistry within a shallow layer near the surface
- Surface HO_x concentrations are increased by up to a factor of 30 due to halogen chemistry

Supporting Information:

Supporting Information may be found in the online version of this article.

Correspondence to:

S. Ahmed and J. L. Thomas, shaddy.ahmed@univ-grenoble-alpes.fr; jennie.thomas@univ-grenoble-alpes.fr

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Author Contributions:

Conceptualization: Shaddy Ahmed, Jennie L. Thomas, Jochen Stutz Data curation: Shaddy Ahmed, Frank Flocke, John J. Orlando, Rebecca S. Hornbrook, Eric C. Apel, Detlev Helmig, Patrick Boylan, L. Gregory Huey, Samuel R. Hall, Kirk Ullmann, Christopher A. Cantrell, Alan Fried Formal analysis: Shaddy Ahmed, Jennie L. Thomas, Katie Tuite, Jochen Stutz Funding acquisition: Jennie L. Thomas

Investigation: Shaddy Ahmed Methodology: Shaddy Ahmed, Jennie L. Thomas, Katie Tuite, Jochen Stutz

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The Role of Snow in Controlling Halogen Chemistry and Boundary Layer Oxidation During Arctic Spring: A 1D Modeling Case Study

Shaddy Ahmed¹, Jennie L. Thomas¹, Katie Tuite², Jochen Stutz², Frank Flocke³, John J. Orlando³, Rebecca S. Hornbrook³, Eric C. Apel³, Louisa K. Emmons³, Detlev Helmig^{4,5}, Patrick Boylan⁴, L. Gregory Huey⁶, Samuel R. Hall³, Kirk Ullmann³, Christopher A. Cantrell^{7,8}, and Alan Fried^{4,9}

¹Université de Grenoble Alpes, CNRS, IRD, Grenoble INP, IGE, Grenoble, France, ²Department of Atmospheric and Oceanic Sciences, University of California, Los Angeles, CA, USA, ³Atmospheric Chemistry Observations & Modeling Laboratory, National Center for Atmospheric Research, Boulder, CO, USA, ⁴Institute of Arctic and Alpine Research, University of Colorado, Boulder, CO, USA, ⁵Boulder A.I.R. LLC, Boulder, CO, USA, ⁶School of Earth and Atmospheric Sciences, Georgia Institute of Technology, Atlanta, GA, USA, ⁷Department of Atmospheric and Oceanic Sciences (ATOC), University of Colorado, Boulder, CO, USA, ⁸Laboratoire Interuniversitaire des Systèmes Atmosphériques (LISA), Creteil, France, ⁹Earth Observing Laboratory, National Center for Atmospheric Research, Boulder, CO, USA

Abstract Reactive chlorine and bromine species emitted from snow and aerosols can significantly alter the oxidative capacity of the polar boundary layer. However, halogen production mechanisms from snow remain highly uncertain, making it difficult for most models to include descriptions of halogen snow emissions and to understand the impact on atmospheric chemistry. We investigate the influence of Arctic halogen emissions from snow on boundary layer oxidation processes using a one-dimensional atmospheric chemistry and transport model (PACT-1D). To understand the combined impact of snow emissions and boundary layer dynamics on atmospheric chemistry, we model Cl₂ and Br₂ primary emissions from snow and include heterogeneous recycling of halogens on both snow and aerosols. We focus on a 2-day case study from the 2009 Ocean-Atmosphere-Sea Ice-Snowpack campaign at Utqiagvik, Alaska. The model reproduces both the diurnal cycle and high quantity of Cl_2 observed, along with the measured concentrations of Br_2 , BrO, and HOBr. Due to the combined effects of emissions, recycling, vertical mixing, and atmospheric chemistry, reactive chlorine is typically confined to the lowest 15 m of the atmosphere, while bromine can impact chemistry up to and above the surface inversion height. Upon including halogen emissions and recycling, the concentration of HO. $(HO_y = OH + HO_y)$ at the surface increases by as much as a factor of 30 at mid-day. The change in HO_y due to halogen chemistry, as well as chlorine atoms derived from snow emissions, significantly reduce volatile organic compound lifetimes within a shallow layer near the surface.

1. Introduction

Halogen chemistry has a large impact on tropospheric chemistry in the polar regions (e.g., Abbatt et al., 2012; Barrie et al., 1988; Oltmans et al., 2012; Simpson et al., 2007, 2015; Steffen et al., 2008, 2013). Recently, new evidence of active Arctic chlorine chemistry has been attributed mainly to photochemical activation of chloride present in surface snow (Custard et al., 2017; Liao et al., 2014). Molecular chlorine (Cl_2) and nitryl chloride ($ClNO_2$), emitted from snow and aerosols, are sources of atomic chlorine (Cl) following their photolysis (McNamara et al., 2019, 2021). The highly reactive nature of Cl atoms makes it important even in trace amounts as Cl atoms react with volatile organic compounds (VOCs) up to three orders of magnitude faster than the more abundant hydroxyl radical (OH) (Atkinson et al., 2006). Active chlorine chemistry occurs simultaneously with reactive bromine chemistry each spring (e.g., Abbatt et al., 2012; Barrie et al., 1988; Simpson et al., 2007, 2015). The latter causes both ozone (O_3) and mercury depletion (Hg^0) in the lowest part of the atmosphere (e.g., Oltmans et al., 2012; Steffen et al., 2008, 2013).

Halogens in the Arctic atmosphere ultimately originate from the ocean as halides (Cl⁻ and Br⁻), which are activated on salty surfaces such as snow on sea ice, continental snow and aerosols (Abbatt et al., 2012). Chlorine and bromine species impact atmospheric chemistry within the polar boundary layer via reactions (R1–R9; where *X*, Y =Cl or Br) (Abbatt et al., 2012; Simpson et al., 2015). Cl₂ photolyzes very quickly during the day (R1), with



Project Administration: Jennie L. Thomas

Resources: Jennie L. Thomas, Eric C. Apel

Software: Shaddy Ahmed, Jennie L. Thomas, Katie Tuite, Jochen Stutz Supervision: Jennie L. Thomas, Jochen Stutz

Validation: Jennie L. Thomas, Jochen Stutz, Frank Flocke, John J. Orlando, Rebecca S. Hornbrook, Detley Helmig. Patrick Boylan, L. Gregory Huey, Samuel R. Hall, Kirk Ullmann, Christopher A. Cantrell, Alan Fried Visualization: Shaddy Ahmed Writing - original draft: Shaddy Ahmed, Jennie L. Thomas Writing - review & editing: Shaddy Ahmed, Jennie L. Thomas, Katie Tuite, Jochen Stutz, Frank Flocke, John J. Orlando, Rebecca S. Hornbrook, Eric C. Apel, Detlev Helmig, Patrick Boylan, L. Gregory Huey, Samuel R. Hall, Kirk Ullmann, Christopher A. Cantrell, Alan

Ulimann, Christopher A. Cantrell, Fried a photolysis lifetime of approximately 10 min, producing Cl atoms that rapidly react with ozone (R2) or VOCs (including methane) ([R3] and [R4]) (Atkinson et al., 2006). Reactions (R3) and (R4) constitute the major reaction pathways of Cl atoms (Platt & Hönninger, 2003). This produces organic peroxy radicals (RO₂), including the methylperoxy radical (CH_3O_2) , which ultimately contribute to hydroperoxyl radical formation (HO_2) . HO₂ production, driven by chlorine chemistry, can impact HO_x chemistry (HO_x = OH + HO₂) by decreasing the OH/HO2 ratio, as well as affecting the reactive bromine cycle (Piot & von Glasow, 2009; Rudolph et al., 1999; Thompson et al., 2015). Molecular bromine (Br_2) is photolyzed very rapidly (photolysis lifetime <1 min) to produce bromine atoms (Br) which can lead to efficient ozone destruction and formation of bromine monoxide (BrO) (Wang et al., 2019). However, Br atoms react appreciably only with a few specific VOCs such as ethyne and the aldehydes and not with methane ([R3] and [R4] only occur for Cl). Br atoms can also react with elemental mercury to deplete near-surface atmospheric Hg⁰ levels and produce more reactive forms of mercury (Hg^{II}) (Steffen et al., 2008, 2013; Wang et al., 2019). Subsequent reaction of BrO with HO₂ forms HOBr (R5), which can be photolyzed to re-form Br and OH (R6). The net result of reactions (R5) and (R6) is the regeneration of a Br atom, which can facilitate further ozone depletion, and the conversion of HO₂ to OH, increasing the oxidative capacity of the atmosphere. At high BrO concentrations, Br2 is also regenerated in the gas phase via self-reaction of BrO (R7).

$$X_2 \xrightarrow{h_\nu} 2X$$
 (R1)

$$X + O_3 \rightarrow XO + O_2 \tag{R2}$$

$$Cl + RH + O_2 \rightarrow RO_2 + HCl$$
 (R3)

$$Cl + CH_4 \rightarrow CH_3 + HCl$$
 (R4)

$$XO + HO_2 \to HOX + O_2 \tag{R5}$$

$$HOX \xrightarrow{hv} X + OH \tag{R6}$$

$$BrO + BrO \to Br_2 + O_2 \tag{R7}$$

Figure 1a highlights the typical diurnal behavior of surface molecular halogen concentrations, snow emissions, solar radiation, and the boundary layer height observed during Arctic spring. Measured diurnal cycles of Cl_2 have shown a double-peaked profile, with peaks in the morning and late afternoon, followed by concentrations dropping below 0.8 parts per trillion by volume (pptv) after midnight (Custard et al., 2016; Liao et al., 2014; McNamara et al., 2019). At sunrise, increased solar radiation drives photochemistry within the snow interstitial air which leads to the release of halogens to the overlying atmosphere via diffusion and wind pumping (Bartels-Rausch et al., 2014; Grannas et al., 2007; Pratt et al., 2013; Thomas et al., 2011; Toyota et al., 2014). Boundary layer mixing modulates surface Cl_2 concentrations, with Cl_2 decreasing during the day due to a combination of its fast photolytic loss (R1) and the effects of boundary layer mixing. Solar heating of the lower atmosphere can cause turbulent mixing of the surface layer, mixing species away from the surface, and increasing the height of the boundary layer (Anderson & Neff, 2008). Low light conditions (i.e., night and early morning) cause a reduction in the photochemical loss of Cl_2 (R1) and a collapse of the boundary layer. This effect has previously been demonstrated to drive evening increases of reactive nitrogen species (NO_x = NO + NO₂) at the surface in both the Arctic and Antarctic (Frey et al., 2015; Honrath et al., 1999, 2002; Thomas et al., 2011).

Figure 1b illustrates some of the key known emission sources of halogens from surface snow in the Arctic. Heterogeneous reactions involving hypohalous acids (e.g., HOCl and HOBr; R8) and halogen nitrates (e.g., $ClONO_2$ and $BrONO_2$; R9), have been recognized as a source of molecular halogens on halide-containing snow and aerosol surfaces (Aguzzi & Rossi, 1999; Deiber et al., 2004; Finlayson-Pitts et al., 1989; Hu et al., 1995; Pratte & Rossi, 2006). Bromine chloride (BrCl) is another product formed via reactions (R8) and (R9) on snow and aerosols, linking the chlorine and bromine chemical cycles (McNamara et al., 2020). BrCl can then be photolyzed, re-forming Br and Cl atoms, as in reaction (R1).

$$HOX + X^{-}/Y^{-} + H^{+} \xrightarrow{(surface)} X_{2}/XY + H_{2}O$$
(R8)

$$XONO_2 + X^-/Y^- \xrightarrow{(surface)} X_2/XY + NO_3^-$$
(R9)





Figure 1. (a) Diurnal evolution of boundary layer height (BLH), solar radiation, chemical snowpack emissions, and surface concentrations of Cl_2 and Br_2 for an "average" day in the Arctic during spring (not drawn to scale). (b) Schematic of key polar halogen emissions from the continental snowpack (X = Cl, Br). Blue arrows represent loss processes and red arrows indicate production. Primary production of halogens is based on ozone and the availability of sunlight. Molecular halogens are also emitted via surface snow and aerosol recycling reactions, dependent on the heterogeneous reactive uptake coefficients γ_s and γ_s , on surface snow and aerosols, respectively.

At present, detailed descriptions of chlorine snow emissions remain absent from most 3D numerical models. Bromine mechanisms are included in some 3D models, but remain under discussion as to the source and recycling mechanisms involving snow (Falk & Sinnhuber, 2018; Fernandez et al., 2019; Herrmann et al., 2021; Marelle et al., 2021; Toyota et al., 2011). Snow is a very complex photochemical medium and the release of halogens is determined by many uncertain processes/variables, including: snow physics; snow/ice chemistry (including photochemistry); gas transport within snow; and impurity concentrations and locations (Bartels-Rausch et al., 2014; Domine et al., 2008; Grannas et al., 2007; McNeill et al., 2012). As a result, modeling snow-covered environments using a first principles approach remains challenging and uncertain (Domine et al., 2013). Zero-dimensional box models are often used to study the effects of halogens on boundary layer chemistry under Arctic conditions (Custard et al., 2015; Liao, Huey, Tanner, et al., 2012; Liao et al., 2014; McNamara et al., 2020; Piot & von Glasow, 2009; Thompson et al., 2015; Wang & Pratt, 2017). An inherent limitation of 0D models, however, is the absence of the vertical dimension necessary for simulating vertical transport and capturing concentration gradients in the atmosphere. Additionally, the physical conditions that characterize the polar regions (low temperatures, limited sunlight during winter, high albedo, etc.) can often create stable low-level temperature inversions resulting in shallow boundary layers (Kahl, 1990). This can greatly impact the vertical distribution of chemical species by acting as a barrier to vertical mixing and transport. One-dimensional models are therefore extremely useful tools which can include these processes to help us better understand the interactions between snow and the atmosphere (Cao et al., 2016; Herrmann et al., 2019; Lehrer et al., 2004; Piot & von Glasow, 2008; Thomas et al., 2012, 2011; Toyota et al., 2014; Wang et al., 2020).

The first observations of high Cl_2 levels within the Arctic boundary layer were reported in spring 2009, during the Ocean-Atmosphere-Sea Ice-Snowpack (OASIS) campaign at Utqiaģvik, Alaska (Liao et al., 2014). Cl_2 mixing ratios of up to 400 pptv were observed and an average noontime Cl atom concentration of 2×10^5 atoms cm⁻³ was estimated from these observations. Daytime Cl_2 mixing ratios were highly correlated with sunlight and surface ozone levels (r^2 value = 0.86), indicating both are key requirements for Cl_2 production. Measurements of VOCs made during the campaign showed a clear impact of chlorine chemistry on VOC oxidation processes (Hornbrook et al., 2016). Measurement-derived estimates of Cl atom concentration suggested the presence of a highly reactive surface layer, which led to an overprediction of VOC production and loss rates compared to the observations



Table 1

Measurements From the OASIS 2009 Campaign Used in This Study

Measurement	Method	Reference	
Meteorology and turbulent fluxes	Ultrasonic anemometers	Boylan et al. (2014)	
Cl ₂ , Br ₂ , BrO, HOBr	Chemical Ionization Mass Spectrometers (CIMS)	Liao et al. (2011, 2014); Liao, Huey, Tanner, et al. (2012)	
OH, HO ₂	Chemical Ionization Mass Spectrometers (CIMS)	Hornbrook et al. (2011); Mauldin III et al. (1998); Tanner et al. (1997)	
NO, NO ₂ , O ₃	Chemiluminescence	Helmig et al. (2012); Villena et al. (2011); Weinheimer et al. (1998)	
НСНО	Difference Frequency Generation Tunable Diode Laser Absorption Spectrometer	Weibring et al. (2007, 2010)	
СО	IR absorption CO analyzer	Parrish et al. (1994)	
VOCs ^a	Trace Organic Gas Analyzer (TOGA)	Hornbrook et al. (2016)	
Aerosol number density and radius	Optical Particle Counter (OPC) and Scanning Mobility Particle Sizers (SMPS)	Woo et al. (2001)	
Actinic flux	CCD Actinic Flux Spectroradiometers (CAFS)	Shetter and Müller (1999)	
^a See Table 1 in Hornbrook et al. (2016).			

(Hornbrook et al., 2016). Interactions between radical chemistry, atmospheric mixing, and snow emissions need to be better understood in order to fully explain these observations and the impacts of halogens on boundary layer oxidation processes.

In this work, we address the following questions:

- 1. What combination of factors, including vertical mixing, land-based snow emissions/recycling, and chemistry explain observations of halogens in the Arctic surface layer?
- 2. How are halogens vertically distributed within the polar boundary layer?
- 3. What is the impact of halogen chemistry on boundary layer oxidation processes as a function of altitude?

We answer these questions using an updated version of the Platform for Atmospheric Chemistry and vertical Transport in 1-dimension (PACT-1D) model (Tuite et al., 2021), which includes descriptions of halogen chemistry, emissions, and recycling. We compare our model with surface measurements of chemical species, including Cl_2 and Br_2 , recorded during the 2009 OASIS campaign at Utqiagvik, Alaska. In Section 2, we introduce the measurements used from the campaign and the new model halogen updates are described in Section 3. The model configuration used in this work is presented in Section 4, followed by the model results and a discussion in Section 5. Finally, the conclusions and perspectives are presented in Section 6.

2. OASIS 2009 Campaign Measurements

In this study, we use measurements taken during the OASIS campaign, which was conducted between March and April 2009 near Utqiaġvik, Alaska. The measurement site was located approximately 5.5 km northeast of the town of Utqiaġvik (71.323 N, 156.663 W; 8 m above sea level) and was chosen for the prevailing northeasterly winds arriving from over the Beaufort Sea. A map of the study site can be found in Boylan et al. (2014) and Hornbrook et al. (2016). Observations from this campaign were chosen due to the extensive chemical and meteorological measurements available, including direct measurements of Cl₂, Br₂, BrO, and a large number of VOCs. A summary of the measurements used in this study is given in Table 1 with the respective instruments/techniques used during OASIS. Meteorological measurements (temperature, winds, relative humidity) were made from two tower stations set up at the field site at several heights. At one of the tower stations, turbulent flux measurements were made using ultrasonic anemometers located at 4 heights (0.6, 1.8, 3.2, and 6.2 m above ground level [AGL]). Inorganic halogens (including Cl₂, Br₂, BrO and HOBr), as well as OH and HO₂, were measured using chemical ionization mass spectrometers (CIMS) at 1.5 m AGL (Hornbrook et al., 2011; Liao et al., 2011, 2014; Liao, Huey, Tanner, et al., 2012; Mauldin III et al., 1998; Tanner et al., 1997). Surface ozone and NO_x measurements were made using



Table 2Heterogeneous Reactions and Reaction Uptake Coefficients on Aerosols (γ_a)				
Reaction	Heterogeneous reactive uptake (γ_a)	Reference		
$\mathrm{HOCl} + \mathrm{Cl}^{\mathrm{(aq)}} \to \mathrm{Cl}_2$	2×10^{-4}	Ammann et al. (2013)		
$HOCl + Br^{-}_{(aq)} \rightarrow BrCl$	2×10^{-4}	Ammann et al. (2013)		
$\text{ClONO}_2 + \text{Cl}_{(aq)}^- \rightarrow \text{Cl}_2 + \text{HNO}_{3(aq)}$	0.03	Aguzzi and Rossi (1999)		
$\text{ClONO}_2 + \text{Br}_{(aq)}^- \rightarrow \text{BrCl} + \text{HNO}_{3(aq)}$	0.05	Aguzzi and Rossi (1999)		
$\text{ClONO}_2 \rightarrow \text{HOCl} + \text{HNO}_{3(\text{aq})}$	0.03	Aguzzi and Rossi (1999)		
$\mathrm{HOBr} + \mathrm{Br}^{\mathrm{(aq)}} \to \mathrm{Br}_2$	0.05	Pratte and Rossi (2006); IUPAC (2009)		
$HOBr + Cl_{(aq)}^{-} \rightarrow BrCl$	0.05	Pratte and Rossi (2006); IUPAC (2009)		
$BrONO_2 + Br_{(aq)}^- \rightarrow Br_2$	0.06	Deiber et al. (2004)		
$BrONO_2 + Cl_{(aq)}^- \rightarrow BrCl$	0.04	Deiber et al. (2004)		
$BrONO_2 \rightarrow HOBr + HNO_{3(aq)}$	0.04	Deiber et al. (2004)		
$\mathrm{N_2O_5} + \mathrm{Cl^{(aq)}} \rightarrow \mathrm{ClNO_2} + \mathrm{HNO_{3(aq)}}$	0.02	Burkholder et al. (2019)		
$\mathrm{N_2O_5} + \mathrm{Br^{(aq)}} \rightarrow \mathrm{BrNO_2} + \mathrm{HNO_{3(aq)}}$	0.011	Seisel et al. (1998)		
$\text{Cl}_2 + \text{Br}^{(\text{aq})} \rightarrow \text{BrCl} + \text{Cl}^{(\text{aq})}$	0.2	Burkholder et al. (2019)		
$\rm OH + HCl \rightarrow 0.5*Cl_2$	0.1	Knipping et al. (2000); Laskin et al. (2006)		

a chemiluminescence instrument on a second tower station, operated by the National Center for Atmospheric Research (NCAR), at 3 different heights (0.6, 1.5, and 5.4 m AGL) (Helmig et al., 2012; Villena et al., 2011; Weinheimer et al., 1998). Additionally, measurements of formaldehyde (HCHO) made by a Difference Frequency Generation Tunable Diode Laser Absorption Spectrometer (Barret et al., 2011; Weibring et al., 2007, 2010), and 18 VOCs measured by a Trace Organic Gas Analyzer (TOGA) (Hornbrook et al., 2016) were made at these same heights. Balloon soundings were also launched during the campaign to record vertical profiles of ozone (Helmig et al., 2012; Oltmans et al., 2012). Carbon monoxide (CO) measurements were made using a CO infrared absorption analyzer (Parrish et al., 1994). Aerosol physical properties (size distribution and number concentration) were measured using an optical particle counter and two scanning mobility particle sizers (Woo et al., 2001). Finally, actinic flux measurements made by CCD Actinic Flux Spectroradiometers (CAFS) were used to calculate photolysis frequencies of 35 different reactions using a modified version of the Tropospheric Ultraviolet and Visible (TUV) radiation model version 4.4 (Madronich & Flocke, 1999; Shetter & Müller, 1999). Data from this campaign are available through the National Science Foundation (NSF) Arctic Data Center at https://arcticdata. io/ (Apel, 2009; Cantrell, 2009; Fried, 2009; Guenther, 2009; Hall, 2009; Smith et al., 2009; Weinheimer, 2009).

3. Description of Halogen Chemistry Within PACT-1D

The Platform for Atmospheric Chemistry and vertical Transport in 1-Dimension (PACT-1D) is the vertical column model used in this work to study Arctic halogen emissions and their impact on oxidation processes during the OASIS campaign. A full description of this model is given in Tuite et al. (2021). Chlorine and bromine gas-phase and heterogeneous reactions are added to this version of the model. Snow emissions and recycling mechanisms of chlorine and bromine have also been implemented and are described in the following sections.

3.1. Gas-Phase and Aerosol Heterogeneous Halogen Chemistry

We update the existing PACT-1D mechanism to include additional chlorine and bromine gas-phase and heterogeneous reactions. The chemical mechanism in PACT-1D is based on the Regional Atmospheric Chemistry Mechanism version 2 (RACM2) (Goliff et al., 2013) using the Kinetic PreProcessor (KPP) (Sandu & Sander, 2006). The additional gas-phase bromine reactions are added following the implementation of Marelle et al. (2021) and are listed in the model chemical mechanism (Ahmed et al., 2022). Reactive and non-reactive heterogeneous uptake reactions of halogens on aerosols are also added to the model (Table 2). We do not include a full description of aerosol aqueous-phase chemistry within the model. We track particulate chloride and bromide as separate species that undergo heterogeneous chemistry to ensure mass balance, which are tracked in the model. We do this by first



initializing the concentration of aerosol-phase chloride and bromide to the chloride and bromide concentrations in fresh sea salt aerosols and then track release and reactive and non reactive uptake. Second-order heterogeneous reactions consuming aerosol-phase halide ions are treated as pseudo first-order reactions, following Marelle et al. (2021), maintaining mass conservation of each species.

3.2. Snow Emission and Recycling of Cl₂ and Br₂

Emissions of molecular halogens from snow have been identified as a key source of Arctic halogen production (Custard et al., 2017; Pratt et al., 2013). Cl_2 and Br_2 production from continental snow have been reported to be a function of both solar radiation and ozone concentration (Custard et al., 2017; Liao et al., 2014; Liu et al., 2017). Halogen species deposited to the snow surface can also undergo recycling mechanisms to re-emit reactive halogens back into the atmosphere (Abbatt et al., 2012; Toyota et al., 2011). We therefore add four parameterizations to describe emissions of Cl_2 and Br_2 in this version of PACT-1D. We include (a) an emission of Cl_2 and Br_2 as a function of the available solar radiation and the surface ozone concentration, and (b) a recycling source of X_2 from the surface conversion of XONO₂ and HOX (where X = Cl, Br) on snow. In both cases, the exact parameterizations are determined by comparing modeled and observed halogen concentrations.

The emission of Cl₂ is parameterized as follows:

$$E_{\text{Cl}_2}^{primary} = F_{(p,\text{Cl})} \times \left(J_{\text{Cl}_2}\right)^{0.5} \times [\text{O}_3]$$
(1)

$$E_{\text{Cl}_2}^{\text{recycling}} = \gamma_{(\text{snow},\text{Cl})} \times \left(D_{\text{CIONO}_2} + D_{\text{HOCI}} \right)$$
(2)

where $E_{Cl_2}^{primary}$ and $E_{Cl_2}^{recycling}$ are the snow emission fluxes of Cl₂, $F_{(p,Cl)}$ is a correction factor which includes a scaling term and the height of the lowest model level (0.01 cm) in units of cm s^{-1/2}, J_{Cl_2} is the calculated photolysis rate of Cl₂, $[O_3]$ is the measured O₃ concentration (in molec cm⁻³), $\gamma_{(snow,Cl)}$ is the probability of heterogeneous conversion on snow to re-form Cl₂ (between 0 and 1), and D_{CIONO_2} and D_{HOCl} are the model-calculated deposition rates of CIONO₂ and HOCl, respectively. In the case of primary Cl₂ emissions (Equation 1), different values of $F_{(p,Cl)}$ were tested in order to reproduce the Cl₂ measurement data in the model (Figure S1 in Supporting Information S1). Observed ambient concentrations of Cl₂ showed a double peaked profile, with an increase in the early morning following sunrise, and a second peak in the late afternoon before falling to almost zero at night. The best-fit primary emission flux for Cl₂ is found to be a function of J_{Cl_2} to the power of 0.5, with $F_{(p,Cl)} = 0.2 \text{ cm} \text{ s}^{-\frac{1}{2}}$. It is also well known that ClONO₂ and HOCl are converted on ice surfaces to re-form Cl₂ (IUPAC, 2009). However, within snow there are a number of complex physical and chemical processes that make these recommendations not directly applicable for snow. We therefore performed a series of sensitivity tests varying $\gamma_{(snow,Cl)}$ between 0 and 1, and found the best fit value of 0.1 for chlorine recycling on snow (Figure S2 in Supporting Information S1).

For Br_2 , the emission sources are described as:

$$E_{\text{Br}_2}^{primary} = F_{(p,\text{Br})} \times J_{\text{Br}_2} \times [\text{O}_3]$$
(3)

$$E_{\rm Br_2}^{recycling} = \gamma_{(snow,\rm Br)} \times \left(D_{\rm BrONO_2} + D_{\rm HOBr} \right) \tag{4}$$

where $E_{Br_2}^{primary}$ and $E_{Br_2}^{recycling}$ are the snow emission fluxes of Br₂, $F_{(p,Br)}$ is a correction factor which includes a scaling term and the height of the lowest model level (0.01 cm) in units of cm, J_{Br_2} is the calculated photolysis rate of Br₂, $\gamma_{(snow,Br)}$ is the heterogeneous conversion efficiency on snow to re-form Br₂ (between 0 and 1), and D_{BrONO_2} and D_{HOBr} are the model-calculated deposition rates of BrONO₂ and HOBr, respectively. For bromine, we found that the observations of bromine species are best described using primary emissions (Equation 3) as a function of J_{Br_2} , with $F_{(p,Br)} = 0.01$ cm (equivalent to the lowest model level height and a scaling factor of 1). The conversion of BrONO₂ and HOBr on ice to re-form Br₂ is known to be more efficient than for chlorine (IUPAC, 2009), which in part facilitates the well known bromine explosion chemistry (Abbatt et al., 2012). We tested a range of possible conversion efficiencies for these reactions and found $\gamma_{(snow,Br)} = 0.6$ best reproduces the observations (Figure S3 in Supporting Information S1). For both Equations 2 and 4, it is assumed that there is an infinite supply of Cl⁻ and Br⁻ in the snow. We do not include conversion of N₂O₅ on snow to form reactive bromine and chlorine due to the low NO_x concentrations compared to pollution influence.



There are large uncertainties in describing both the primary emission flux (Equations 1 and 3) from land-based snow, as well as the recycling of both bromine and chlorine species on snow (Equations 2 and 4), which must be considered in future work that use or further refine these parameterizations. First, there are significant uncertainties in vertical transport near the snow surface and in the lowest portion of the atmosphere (~below 10 m). Therefore, as future work refines our knowledge of these vertical transport processes, we will need to revisit the values used for $F_{(p,Cl)}$, $F_{(p,Br)}$, $\gamma_{(snow,Cl)}$ and $\gamma_{(snow,Br)}$. Second, the main factors driving molecular halogen production from the snowpack are still highly uncertain, with more work needed to improve our understanding. In addition, descriptions of halogen emissions from land-based snow within 3D models remain limited. Bromine emissions triggered from ozone deposition to snow on sea ice is the main process considered by the bromine emissions/ recycling scheme of Toyota et al. (2011). Here, we use ambient ozone concentrations rather than ozone deposition as the trigger for both bromine and chlorine on land-based snow, as suggested from observations. Our equations can be re-formulated as a function of the ozone deposition rate (which is directly dependant on ozone concentration) to be more consistent with equations proposed for snow on sea ice. Finally, production of BrCl from Arctic snow has been measured following irradiation of the snowpack, with multiphase reactions on snow also predicted to be significant contributors of BrCl production (Custard et al., 2017; McNamara et al., 2020). However, flux estimates of BrCl from snow remain uncertain and measurements of BrCl were not available during our selected simulation period (see Section 4.1). We therefore only include BrCl production via heterogeneous reactions on aerosols (Table 2), but, this must be updated in future work to also include BrCl emissions from continental snow.

4. Model Setup

4.1. Selection of OASIS Simulation Period

The model was set up for the dates of 18–19 March 2009 during the campaign; these dates were selected due to the high Cl₂ concentrations recorded and the limited influence from local pollution sources (Figure S4 in Supporting Information S1). The average daytime (06:00–20:00) Cl₂ mixing ratio for the 2 days was 59 pptv and surface ozone levels remained above 10 parts per billion by volume (ppbv), indicating that there was not a major ozone depletion event during this period. Ozonesonde data from profiles launched at the start and just after our modeling case study showed that ozone was well distributed within the lower atmosphere with no significant ozone gradients as a function of altitude (Figure 4c in Oltmans et al. (2012) and Figure 13a in Helmig et al. (2012)). Average background levels of NO_x and CO over the entire campaign were recorded at ~84 pptv and ~160 ppbv, respectively (Villena et al., 2011). Measurements of NO_x and CO between 18 and 19 March do not suggest polluted conditions, with CO levels close to the average background measurements (~160 ppbv), however NO_x levels were above the average background levels (50–500 pptv). Influence from nearby anthropogenic sources was likely to be minimal during this period as winds arriving at the measurement site originated from the Arctic Ocean (north through northeast) for most of 18 and 19 March. Considering these criteria, the period between 18 and 19 March best met the requirements for our modeling case study.

Figures 2a and 2b show the state of sea ice north of Utqiaġvik, on the 18 and 19 March, respectively. A key feature of Figures 2a and 2b is the presence of sea ice leads close to the measurement site. Sea ice leads are visible on both days which are important as they can induce convective mixing of air masses, impacting the concentration of species recorded at the measurement site (Moore et al., 2014). This has been shown to replenish ozone-depleted air masses to near-background concentrations (Moore et al., 2014) as well as influencing the vertical distribution of BrO by mixing it higher in the atmosphere (Simpson et al., 2017).

4.2. Model Configuration

We set up the vertical model grid (Figure 3a) using a total of 112 levels, with a logarithmic spacing for the lowest 1 m of the grid down to a lower boundary of 1×10^{-4} m. The model levels are linearly spaced up to 100 m, by 1-m increments, followed by a non-linear spacing to an upper boundary of 3,000 m. This highly resolved vertical model grid allows us to analyze the impacts of halogen emissions on chemistry very close to the surface.

The 1D model is driven by input data obtained from the measurements (where possible), model output data and calculated explicitly from parameterizations. The atmospheric dynamics (temperature, pressure, relative humidity) are calculated using the 3D Weather Research and Forecasting (WRF) meteorological model (Skamarock et al., 2019) for Utqiagvik, Alaska, and used to drive the 1D model physics in combination with the OASIS ground





Figure 2. (a) Daily Arctic sea ice cover and leads from Willmes and Heinemann (2015) over Northern Alaska on 18 March 2009 and (b) 19 March 2009 with the town of Utqiagivk marked in magenta. Areas shaded in white represent either land cover or data artifacts. (c) 2-m temperature from WRF over Utqiagivk at local noon on 18 March 2009 and (d) 19 March 2009 with 10-m wind speed and wind directions displayed as arrows. (e) FLEXPART-WRF 6-hr backwards surface (0–100 m) potential emission sensitivity (PES) from the measurement site (marked by the green cross) on 18 March 2009–10:00 AKST and (f) 19 March 2009–18:00 AKST. Numbers represent hourly intervals since release.





Figure 3. (a) PACT-1D model grid used with numbers representing the upper model level height in meters above the snow surface. A total of 112 vertical model levels were used. (b) 3D WRF model domain centered on Utqiagvik, Alaska with a 25×25 km horizontal resolution and 50 vertical levels up to a pressure of 50 hPa.

measurements. We use a WRF set up specifically optimized for the Arctic, described in Marelle et al. (2017), with the model domain centered at Utqiaġvik (domain shown in Figure 3b). A horizontal resolution of 25 km \times 25 km is used with a vertical resolution of 50 levels, up to a pressure of 50 hPa. To validate the use of the WRF simulated meteorology, we compare WRF calculated temperatures at Utqiaġvik with surface measurements from OASIS and available vertical temperature profiles in Figure 4. The Integrated Global Radiosonde Archive (IGRA, Durre et al. (2006)) provides radiosonde data twice a day at 00:00 and 12:00 UTC (15:00 and 03:00 AKST, UTC-9, respectively) which we use to compare with our model results. Figure 4 shows that we are able to obtain very good agreement of both the surface and vertical temperature profiles in WRF compared to the observations.

The eddy diffusion coefficients (K_z) in the model are calculated following the parameterization described in Cao et al. (2016) and used in Herrmann et al. (2019). We calculate these values as measurement data of eddy diffusion coefficients during this period were sparse. Vertical K_z profiles are calculated using the measured friction velocities (u_*) at 1.8 m AGL, with the estimated surface inversion height (SIH) derived from the tower turbulent flux measurements. A comparison was made between the calculated K_z values and the available measurement data which showed that calculated values were approximately a factor of 3 greater than the observations. Above the surface inversion layer, we assume a fixed value of $K_z = 1 \text{ cm}^2 \text{ s}^{-1}$, following Cao et al. (2016). In our model runs, we calculate the SIH using a description based on eddy viscosity scaling, following Equation 5 (Zilitinkevich & Baklanov, 2002; Zilitinkevich et al., 2002):

$$SIH = C_s^2 (u_* L/|f|)^{0.5}$$
(5)

where C_s is an empirical constant (estimated as 0.7), u_* is the measured friction velocity, L is the calculated Obukhov length from the measurements and f is the Coriolis parameter (equal to 1.38×10^4 at the latitude of the study site).

Chemical concentrations in the model are initialized using both observations and CAM-chem model data (Buchholz et al., 2019; Emmons et al., 2020). Aerosol surface area and number concentration are fixed to the observations for the duration of the run throughout the boundary layer. To supplement the 35 reactions reported in the CAFS data set, additional photolysis rates were added using the TUV radiation model (version 5.0). Each of these additional rates is scaled to the reported NO₂ photolysis rate (J_{NO_2}) . Chemical emission of NO₂ is also included in the model and is scaled as a function of J_{NO_2} . These emissions are added to the lowest model level, to simulate





Figure 4. Temperature comparison between the WRF model (red) and measurements (black) at Utqiaġvik, Alaska during the simulation period. (a) 2-m temperature from WRF and surface measurements from OASIS. (b–e) Vertical temperature profiles from WRF and NOAA IGRA radiosondes released every 12 hr during the simulation period (dates and times are in Alaska standard time).

photochemical production from snow, and scaled to align with the NO_x levels measured during the simulation period. The 24-hr average NO₂ emission flux we use is 1.71×10^{13} molecules m⁻² s⁻¹, in reasonable agreement with previous Arctic NO_x flux measurements (Honrath et al., 2002). All input data are provided on 15 min time resolution and the model is run using a 20-s time step.

5. Results and Discussion

We study the impact of halogen emissions on oxidation processes during OASIS by performing the following model runs: a reference simulation without halogen emissions from snow (NOSURF); a model run with surface snow emissions and recycling of halogens active (BASE); and several sensitivity runs (FIXO3, AERO, BLD). The model runs are summarized in Table 3 and are discussed in detail in the following section. In all model runs, we include heterogeneous chemistry on aerosols, which participates in active recycling of halogen species in all cases. We present the results and discussion in seven sub-sections. First, we present the meteorological conditions at the measurement site during the modeled period (Section 5.1), followed by an analysis of the NOSURF (Section 5.2) and BASE runs (Section 5.3). We discuss in detail the results of the sensitivity tests performed (Section 5.4), the influence of snow emissions on the vertical extent of halogen concentrations (Section 5.5) and a comparison of the snow emission fluxes with other estimates (Section 5.6). Finally, we analyze the impacts on boundary layer oxidation processes (Section 5.7).



Table 3 Description of the Model Runs Performed in This Study			
Model run	Description		
NOSURF	Run with halogen snowpack emission routines deactivated.		
BASE	Run with halogen snowpack emission and recycling routines active.		
FIXO3	BASE run + O_3 fixed to the observations.		
AERO	BASE run + heterogeneous recycling efficiency on aerosols increased by a factor of 10.		
BLD	BASE run + surface inversion height estimated using expression from Pollard et al. (1973).		

5.1. Meteorological Conditions and Air Mass History

The measurements during OASIS were made approximately 5.5 km northeast of Utqiagvik, Alaska, near the Arctic Ocean (Barret et al., 2011; Boylan et al., 2014; Helmig et al., 2012; Hornbrook et al., 2016; Liao, Huey, Tanner, et al., 2012; Liao et al., 2014; Villena et al., 2011). Figures 2a and 2b show the sea ice cover and lead information over Northern Alaska on 18 and 19 March 2009, respectively. Lead information is obtained from the dataset of Willmes and Heinemann (2015) who used thermal-infrared data retrieved from MODIS and applied a binary segmentation procedure to identify leads (Willmes & Heinemann, 2016). During March, the sample location was snow covered and the surrounding ocean largely covered by sea ice, typically reaching its annual maximum in spring. Sea ice leads are clearly visible during the simulation period north of Utqiagvik, which are known to be important for inducing convective mixing that influences atmospheric chemistry (Moore et al., 2014; Simpson et al., 2017). Meteorological conditions, such as wind speed, wind direction and surface temperature, can also alter surface chemical concentrations via impacts on boundary layer dynamics. Winds on both days were recorded arriving from the northeast, over the Beaufort Sea, carrying clean air masses to the measurement site. During this period, wind speeds were moderate to weak ($<5 \text{ m s}^{-1}$; Figure S4 in Supporting Information S1), lower than much of the campaign period, and surface temperatures were close to the March average. A strong low-level temperature inversion was also observed for the duration of these 2 days, indicating stable boundary layer conditions, which is likely to inhibit vertical mixing of species between the inversion layer top and the overlying atmosphere (Figure S5 in Supporting Information S1).

We use the regional meteorological model WRF (setup described in Section 4.2) to both drive the 1D model atmospheric physics and to understand the regional meteorological conditions during the sampling period. Simulated 2-m temperature and 10-m winds over Northern Alaska are shown in Figures 2c and 2d on 18 and 19 March 2009 (local noon) respectively. The wind direction from WRF on both days captures the northern/northeasterly winds measured at the site, as well as the weaker wind speeds on 19 March. This analysis allows us to next identify the origin for air arriving at the measurement site, using the Lagrangian particle dispersion model, FLEXPART-WRF (Brioude et al., 2013). FLEXPART-WRF is driven by the meteorological conditions simulated by WRF and is run in backward mode to simulate air mass histories for the modeled period. These simulations are performed by releasing a total of 100,000 air parcels at the time when Cl₂ maxima were observed for each day (10:00 a.m. and 6:00 p.m. local time for 18 and 19 March, respectively) and run backwards in time for 6 hr. Figures 2e and 2f, show the calculated surface (0-100 m) potential emission sensitivities (PES) near the measurement site. The PES indicates the air mass origin near the surface and represents the length of time the air mass is sensitive to surface emissions. In Figures 2e and 2f, we show that air masses on these days were unaffected by either the town of Utgiagvik or Prudhoe Bay (southeast of Utgiagvik, not shown on map). Transport of air masses over sea ice, including leads, may impact the halogen concentrations measured at the site. On both days, the PES shows (Figures 2e and 2f) that air passes over leads (Figures 2a and 2b) just before arrival at Utqiagvik. Moore et al. (2014) showed that the rapid recovery of depleted mercury and ozone, which are both tied to the abundance of bromine, can be explained by lead-initiated convection bringing higher concentrations of Hg⁰ and ozone from aloft. This also indicates that high concentrations of bromine in these air masses are likely diluted over leads during the induced convective mixing, as shown by Simpson et al. (2017). Due to the air mass origins and lead locations, and the relatively short lifetimes of Cl_2 and Br_2 , our case is expected to be particularly sensitive to local snow emissions and so we assume that this provides the main source of Cl₂ and Br₂ for our case study (see Section 5.2).





Figure 5. Model comparison with observations at 1.5 m above ground level during 18 and 19 March 2009. NOSURF (blue) and BASE (red) runs are compared with the 10-min averaged measurements (black). NO_x measurements are plotted as an hourly average. Gray shaded areas represent the standard deviation of the average from the instantaneous measurements. Photochemical lifetimes of $Cl_2(\tau_{Cl_2})$ and $Br_2(\tau_{Br_2})$ (orange) and the calculated photolysis rate of $NO_2(J_{NO_2})$ (cyan) are also plotted.

5.2. Model Results Without Snow Emissions or Recycling

A model simulation without halogen emissions from snow or surface recycling (NOSURF) was first performed as a reference simulation. The results from this simulation are compared to measured species at 1.5 m AGL in Figure 5 (blue curve). The halogen species (Cl₂, BrO, and HOBr) in this simulation remain negligible for the duration of the simulation, with the exception of Br_2 which is initialized as described below, showing that additional sources of both chlorine and bromine are required to explain the observations. Surface Br_2 is initialized to the average midnight value (15 pptv) that was recorded during OASIS (Liao, Huey, Tanner, et al., 2012), which



fell rapidly to zero after 08:00 on the first day, indicative of photochemical loss. No significant levels of Br_2 after this period are modeled, suggesting that bromine recycling solely on aerosols is not efficient enough to replenish measured levels of Br_2 and other bromine species. Local snow emissions of Br_2 are therefore necessary to replenish bromine levels during the simulation period.

Surface measurements of both NO and NO₂ were higher than the background average (~82 pptv), reaching daytime peaks of close to 250 pptv and over 400 pptv at night on 19 March. These measurements were likely impacted by both local background NO, emissions from snow (Honrath et al., 1999, 2002) and transient point sources. We filtered out the extreme elevated point sources of pollution (above 500 pptv), and use an hourly average of NO and NO₂ concentrations to smooth out sharp peaks arising from local point sources. The observations (Figures 5e and 5f) show some remaining sharp peaks of NO_x on both days, likely caused by these local emission sources, which are difficult to estimate. In addition, a large increase of NO₂ on the evening of 19 March was recorded, corresponding with a change of wind direction and air mass, bringing air from more polluted regions to the measurement site. Stable conditions and low wind speeds may have also facilitated the build-up of higher NO, concentrations near the surface on these 2 days. The impact of these local point sources, and of advected polluted air masses, are therefore difficult to simulate in the model to represent the true NO_x concentrations observed at the measurement site. Modeled values reach and even exceed the measured daytime peaks, with a large overestimation in NO on day 2, before falling to lower than 100 pptv at night. The low concentration of modeled halogens would certainly contribute to the overestimation of NO_y concentrations via halogen oxide limited reactions with NO and NO₂. Changes in the surface ozone levels over the 2 days are not fully captured by either the NOSURF or BASE simulations. This is possibly due to sea ice leads and convective mixing of ozone down from the free troposphere to the surface. The model currently does not account for the impact of sea ice leads on convective mixing of air masses, and would require further testing to include. Finally, we find a general underestimation of both HCHO and HO₂ levels, indicating missing oxidants and oxidation chemistry, and a predicted midday OH concentration between 0.7×10^6 and 1.5×10^6 molecules cm⁻³ for the 2 days.

5.3. Model Results With Halogen Emissions From Snow and Surface Recycling

When snow and recycling emissions of halogens are active (BASE run), we obtain much better agreement with the measured surface mixing ratios compared to the NOSURF run. Measured Cl₂ levels reached up to 150 and 300 pptv on 18 and 19 March 2009, respectively (10 min average). Figure 5 (red curve) shows the model performs well, capturing both the timing and intensity of the morning and late afternoon Cl₂ peaks on the first day, with some discrepancies on day 2. Early morning increase of Cl₂ was recorded after sunrise, suggesting a photochemical production mechanism, which is captured by the model on both days. Daytime levels of modeled Cl_2 on day 2 are overpredicted, by up to 100 pptv, with the difference possibly explained by weak vertical mixing and a shallow daytime boundary layer. The effects of this on surface chemical concentrations are discussed in more detail in Section 5.5. Nighttime Cl₂ mixing ratios fall to near-zero levels in the model, which is consistent with the measurements on both days. Our model results show that the nighttime (20:00–06:00) reduction of Cl_2 at 1.5 m is largely explained by depositional loss to the ground (see Section 5.5). Together, vertical transport and deposition represent the dominant nighttime loss processes (~94%) for Cl₂ at 1.5 m. Heterogeneous uptake of Cl₂ on aerosols and reaction with bromide has also been suggested as a potential Cl₂ sink (and a source of BrCl) (Hu et al., 1995; McNamara et al., 2020; Wang & Pratt, 2017). We find that the reaction of Cl, with bromide on aerosols accounts for nearly 5% of nighttime removal of Cl₂ at 1.5 m, which comprises the majority (95%) of the nighttime chemical loss for Cl₂.

Modeled bromine species (Br_2 , BrO, and HOBr; Figures 5b–5d, respectively) are also in close agreement with the measurements, with a slight underestimation of BrO and HOBr on day 1. Daytime measurements of Br_2 on these 2 days are missing due to unstable background Br_2 measurements that led to observations below the detection limit (2.0 pptv) (Liao, Huey, Tanner, et al., 2012). We find that modeled daytime levels of Br_2 are close to this 2 pptv detection limit, due its very fast photochemical loss. At night, we find an accumulation of the photolabile Br_2 , via the surface recycling mechanism, which provides reactive bromine for the following day. This is consistent with the average diurnal profile measured for Br_2 during OASIS (Liao, Huey, Tanner, et al., 2012), as well as other Arctic measurement campaigns during spring (McNamara et al., 2020; Wang et al., 2019). Modeled BrO and HOBr diurnal profiles are also in agreement with the observations, with peaks at noon on the first day, indicative of production via Br atoms, and near zero at night. On day 2, a second peak for both BrO and HOBr



is recorded in the late afternoon, coinciding with the evening peak of Cl_2 . This suggests that the second peak in halogen species could possibly be due to a change in the boundary layer meteorology (e.g., collapse of the boundary layer) rather than chemical production.

The model captures the general trend of NO_x (Figures 5e and 5f) and we obtain better agreement with the observations in the BASE run, however, the model does not capture some peaks which may be due to advection of more polluted air masses (e.g., evening of 19 March) or transient point sources. Simulated NO_x levels are highly affected by the presence of halogen emissions, with both NO and NO₂ levels reduced in the BASE run compared to NOSURF. Halogens can react with NO_x to produce halogen nitrites and nitrates (e.g., ClNO₂, ClONO₂, BrONO₂), which act as an important reservoir to sustain reactive halogen chemistry. These species can release halogens back into the atmosphere either directly via photolytic destruction, or by chemical reactions on aerosols and surface snow. O₃ levels in the BASE run also show a steady decline over the 2 days, with O₃ changes dominated by vertical mixing and deposition to the ground in our particular simulation period (Figure 5h). Modeled HCHO (Figure 5g) and HO₂ (Figure 5i) are also in better agreement with the observations following the addition of halogen emissions. We find an increase in the daytime HO_x (HO_x = OH + HO₂) levels by roughly 20–30 times compared to the NOSURF run, indicating much more active HO_x chemistry, which can be attributed to halogen chemistry. Overall, we show that halogen emissions from snow and snow-surface recycling are necessary to reproduce surface concentrations of several key species measured during OASIS, with a considerable impact on HO_x concentrations and oxidative chemistry.

5.4. Model Sensitivity Runs

We investigate the effects of different model uncertainties on surface chemical concentrations by performing three sensitivity tests. The aim of these runs is to explore uncertainties in both the chemical and dynamical mechanisms in our model and their associated impacts on surface concentrations. Specifically, we test whether changes in the modeled ozone concentration, halogen recycling on aerosols, or boundary layer dynamics impact the conclusions drawn from this modeling study. Descriptions of the runs performed are included in Table 3 and are summarized here, followed by a discussion of the results compared with the surface observations (Figure 6).

- 1. **FIXO3**: We first address the impact of ozone on halogen concentrations by fixing the modeled ozone to the measurements within the boundary layer (Figure 6, dashed green curve). Bromine levels in this run are greatly affected by the change in ozone availability due to the reaction with Br atoms (R2). On 18 March, BrO and HOBr are both underestimated in this run compared to the observations, followed by an overestimation on 19 March (when there was more ozone available). Nighttime Br₂ levels are underestimated by ~10 pptv compared to the observations and BASE run, with this difference likely explained by the lower levels of BrO on day 1, resulting in less BrONO₂ formation and recycling to re-form Br₂. The daytime HO₂ concentration on day 2 is approximately 68% lower than the BASE run, due to increased BrO levels and subsequent loss via reaction (R5). The results from this run point to potential inaccuracies in the emission parameterizations of bromine, uncertainties in the downward mixing of ozone in the model from above the boundary layer, or, to the missing treatment of advected air masses, all of which require further exploration and testing
- 2. AERO: To test whether heterogeneous recycling on aerosols could contribute a significant source of halogens, we increase the heterogeneous reactive uptake coefficients for Cl₂, Br₂, and BrCl formation reactions by a factor of 10 (Figure 6, dashed purple curve). This test fails to show a significant increase in the halogen concentrations at the surface, indicating that recycling on aerosols contribute only a minor source of reactive halogens at 1.5 m. Interestingly, nighttime Br₂ levels decreased by up to 5 pptv when compared to the BASE run, caused by lower BrONO₂ levels (~25% reduction) as it was more efficiently recycled on aerosols. We also evaluate the difference in the vertical distribution of halogens between the AERO and BASE run (Figure S6 in Supporting Information S1). The modeled vertical distribution of halogens in the BASE run is discussed in detail in Section 5.5. We find differences of less than 10 pptv between halogens in the AERO run and the BASE run within the lowest 50 m of the model, and see no differences above this height. Specifically, we find an increase in Cl₂ concentrations of several pptv in the AERO run and a small decrease for each modeled bromine species
- 3. **BLD**: We explore uncertainties in the boundary layer dynamics by testing a different expression to calculate the SIH from the meteorological measurements (Figure 6, dashed blue curve). The expression used in this run





Figure 6. Model comparison of sensitivity runs with observations at 1.5 m above ground level during 18 and 19 March 2009. BASE (red), FIXO3 (dashed green), AERO (dashed purple) and BLD (dashed blue) runs are compared with the 10-min averaged measurements (black). NO_x measurements are plotted as an hourly average. Gray shaded areas represent the standard deviation of the average from the instantaneous measurements.

(Equation 6) was originally developed for a stable mixed layer over the ocean by Pollard et al. (1973), and was found to also be applicable to the South Pole by Neff et al. (2008)

$$SIH = 1.2u_*(fN_B)^{-0.5}$$
(6)

$$N_B = \sqrt{\frac{g}{T} \frac{\partial \theta}{\partial z}} \tag{7}$$



where u_* is the measured friction velocity, f is the Coriolis parameter (equal to 1.38×10^4 at the latitude of the study site), N_B is the Brunt-Vaisala frequency, g is the acceleration due to gravity, T is the absolute temperature and $\partial\theta/\partial z$ is the potential temperature gradient. This results in a SIH which is several meters greater than previously used for the BASE run (Figure S5 in Supporting Information S1), leading to some key differences in the modeled chemical species. Most notably, we see a reduction in daytime Cl₂ levels at 1.5 m, by up to 60 pptv, on day 2 compared to the BASE run due to an increased SIH. This shows how sensitive surface concentrations can be to small changes in the boundary layer conditions, with significant uncertainties in vertical transport near the snow surface and lower atmosphere. This is discussed in more detail in Section 5.5.

In summary, the sensitivity runs performed here do not significantly improve model agreement with the observations. However, these sensitivity tests show that modeled surface concentrations are influenced by a number of parameters, which require better understanding of specific processes in order to constrain halogen emissions. Changes in ozone and boundary layer dynamics (vertical mixing) had the largest impacts on halogen concentrations, as well as influencing surface NO_x and HO_x levels. We find that uncertainties in heterogeneous reactions on aerosols do not explain the underestimation of BrO and HOBr on day 1, and represent only a minor source contribution of halogens in our model case. Additional studies designed to investigate these processes and reduce known uncertainties for the Arctic region are needed to further evaluate the source contributions of halogens from snow.

5.5. Vertical Influence of Snow Emissions and Recycling on Halogens

In this section, we use the BASE run to understand the vertical distributions of Cl_2 and Br_2 (Figures 7a and 7b, respectively). No vertically resolved measurements were available for either species, therefore, no direct comparison can be made to the model results. We find that the majority of modeled Cl₂ (approximately 97%) is confined to the lowest 15 m of the atmosphere and rapidly decreases with altitude. This implies highly active chlorine chemistry at the surface. Very little Cl₂ is present above 15 m, indicating a strong vertical gradient in chemical reactivities, with the vertical distribution of Cl₂ influenced by the height of the surface inversion. During the campaign, the surface layer height ranged from as low as a few meters up to several hundreds of meters and was estimated to be very shallow (<50 m) during the simulation period (Boylan et al., 2014). Low-level temperature inversions and shallow boundary layers are a common phenomena in cold polar regions and are frequently characterized by stable conditions and low wind speeds. Typically, solar heating of the surface generates a turbulent well-mixed daytime boundary layer, creating a larger volume in which chemical species can be distributed. This simultaneously increases the vertical transport of species away from the surface and results in decreasing concentrations of chemical species that would otherwise build up near the surface. The diurnal evolution of the surface layer can be seen following this behavior on the first day but not on the second. We are therefore able to capture the daytime reduction in surface Cl₂ levels on day 1, following the morning peak, but overestimate Cl₂ levels on day 2. This high daytime Cl₂ concentration is simulated when the wind speed and estimated SIH were very low (<1 m s⁻¹ and <10 m, respectively), confining Cl₂ to a very shallow layer close to the surface. The BLD sensitivity test also shows similar behavior of the surface layer on the second day, however, estimated a SIH several meters higher during the day, resulting in a reduction of Cl₂ at this time (Figures S5 and S7 in Supporting Information S1). Differences in the SIH estimates between Equations 5 and 6 are discussed in detail by Boylan et al. (2014), but further evaluation is beyond the scope of this study.

The modeled vertical distribution of Br_2 , presented in Figure 7b, shows that Br_2 is present up to and above the SIH during the simulation. During sunlit hours, Br_2 is found only within the first few meters of the model, whereas at night, Br_2 is more abundant (approx. 10 pptv) reaching up to 40 m in height. Previous studies have examined the vertical distribution of bromine in the Arctic via surface-based measurements of BrO (Peterson et al., 2015; Simpson et al., 2017). Peterson et al. (2015) found that the percentage of BrO within the lower troposphere (<200 m) was highly dependent on atmospheric stability. During stable conditions, higher BrO mixing ratios were measured close to the surface, with less BrO distributed within the lower troposphere, compared to unstable conditions which were associated with more vertically distributed BrO events. Simpson et al. (2017) also observed shallow layer events of BrO during stable conditions, and during lead opening events, found that BrO was lofted above the surface via lead-induced convective mixing. Air that had traveled over sea ice, arriving at the measurement site with very low ozone representing ODE conditions and very active bromine chemistry, are not specifically investigated here. However, there are some similarities between what we investigate regarding



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Figure 7. Modeled vertical distributions of (a) Cl_2 and (b) Br_2 during 18 and 19 March 2009 in the BASE run. White trace indicates the model prescribed SIH. Modeled concentration change of (c) Cl_2 and (d) Br_2 due to vertical transport and deposition with respect to time. Black trace indicates the prescribed SIH. Positive values represent upward transport and negative values indicate downward transport.

land-based snow emissions of both bromine and chlorine and what is known about bromine activation from snow on sea ice, where atmospheric boundary layer stability is similar. During the simulation, we find that both BrO and HOBr are present within and above the SIH, reaching heights of up to 40 m during the day (Figure S8 in Supporting Information S1). This is in agreement with the findings of Peterson et al. (2015), as the stable conditions recorded between 18 and 19 March would suggest a shallow layer event of BrO. However, it is also important to note that the air masses arriving at the site could have possibly experienced some lead-induced convective mixing (see Figure 2), which would mix bromine to higher altitudes and this is not considered in the model.

In Figures 7c and 7d, we plot the changes in concentration of Cl_2 and Br_2 due to transport and deposition, respectively. The change in both Cl_2 and Br_2 concentrations, due to vertical transport, is highest during the day following release from snow and transport into the atmosphere. During the night, Cl_2 is mainly transported downward to the surface and lost via deposition to the ground. Deposition in the model is calculated using an approach of molecular collisions with the ground and applying a non-reactive uptake probability (α ; Tuite et al., 2021). This allows us to calculate deposition of different species without prescribing a deposition velocity. For Cl_2 , we set $\alpha = 5 \times 10^{-5}$, following the lower limit recommendation of Burkholder et al. (2019). On 19 March, transport of Cl_2 is clearly limited by the height of the inversion layer, with Cl_2 transport not exceeding more than 15 m altitude, thereby concentrating Cl_2 at the surface. Figure 7d shows that the upward transport of Br_2 is at its maximum during the day on 18 and 19 March. Daytime Br_2 concentrations are confined to the first few meters above ground, indicating that daytime Br_2 is lost via its fast photolytic destruction. Vertically resolved measurements of halogens above the Arctic snow surface are highly desirable for further model evaluation and development.





Figure 8. Snow emission and recycling fluxes of (a) Cl_2 and (b) Br_2 in the BASE run. The total emission flux (black) for each species is plotted together with the primary snow emission flux (orange) and recycling fluxes of XONO₂ (blue) and HOX (red), where X = Cl and Br.

5.6. Modeled Halogen Snow Emission Fluxes Compared to Previous Estimates

Here, we compare the surface emission fluxes of Cl₂ and Br₂ estimated in this work to previous flux estimates. The model emission flux contributions of Cl₂ and Br₂ are shown in Figures 8a and 8b, respectively. Our modeled emissions of both Cl₂ and Br₂ peak at solar noon on each day, coinciding with increased radiation at the snow surface, before falling to zero at night. The peak fluxes are similar to previously reported halogen emission fluxes measured from the Arctic snowpack (Custard et al., 2017), while the exact timing of emissions differs. During February 2014, snowpack flux estimates of Arctic Br₂ and Cl₂ were calculated, based on vertical gradient measurements, for the first time near Utqiagvik. Estimates of these fluxes ranged between $0.7-12 \times 10^8$ and $0.02-1.4 \times 10^9$ molecules cm⁻² s⁻¹ for Br₂ and Cl₂, respectively. We modeled midday fluxes for Cl₂ of 4.3×10^9 and 7.2×10^9 molecules cm⁻² s⁻¹ for 18 and 19 March, respectively, with the primary photochemical snow emission mechanism the main contributor to Cl₂ emissions. This is several times higher than the values reported by Custard et al. (2017). This may in part be explained by increased available sunlight during March compared to February (when the flux measurements were made), enhancing halogen production. Additionally, ambient concentrations of Cl₂ were much lower in February 2014 than those measured in March 2009, with daytime values ranging between 5 and 20 pptv in February 2014 and 59 pptv for the simulation period. Both of these reasons would suggest higher snow emission fluxes of halogens between 18 and 19 March 2009 than estimated previously.

During the OASIS campaign, Liao et al. (2014) estimated an average peak production of Cl_2 of 2.2×10^6 molecules cm⁻³ s⁻¹ using 0D box modeling. Assuming a 15 m mixing layer, this corresponds to an average emission flux of 3.3×10^9 molecules cm⁻² s⁻¹. This is slightly lower than the value we report here, which is likely due to



our focus on an extremely high Cl_2 event during the campaign. Similarly, Wang and Pratt (2017) conducted a 0D box modeling study to simulate halogen snowpack production during March 2012 near Utqiagvik. A peak Cl_2 production rate of $2.9-3.2 \times 10^8$ molecules cm⁻² s⁻¹ was reported (assuming a 10 m effective mixing height), with ambient Cl_2 concentrations ranging between 0 and 20 pptv for the modeled periods. The lower ambient Cl_2 concentrations during March 2012 suggest that solar radiation is not the driving factor for differences between Cl_2 emission fluxes during February and March. Other factors such as atmospheric stability, vertical mixing, and snowpack chemistry are also likely to play a role in controlling molecular halogen emission fluxes from snow. Production of Cl_2 via snow surface recycling of HOCl and ClONO₂ was also minimal over the 2 days, with almost no production of Cl_2 at night, which explains the difference in the nighttime concentrations of Cl_2 and Br_2 . There are uncertainties to the efficiency of this recycling (see Section 3.2), however, sensitivity tests showed no significant increase in Cl_2 when the recycling efficiency of these species was increased (Figure S2 in Supporting Information S1).

Figure 8b shows the modeled Br_2 emission fluxes, with midday values calculated at 4.9×10^8 and 5.0×10^8 molecules cm⁻² s⁻¹ for 18 and 19 March, respectively. These are in close agreement with the range reported by Custard et al. (2017) and are higher than those modeled by Wang and Pratt (2017) of 2.1×10^8 and 3.5×10^6 molecules cm⁻² s⁻¹ for 15 March and 24 March 2012 respectively. Both primary photochemical and snow recycling emissions of Br_2 are important production mechanisms and contribute significantly to the total Br_2 emission flux. Surface recycling of BrONO₂ is the main source of Br_2 on day 1, highlighting the influence of NO_x on bromine chemistry. This mechanism drives the accumulation of Br_2 at night, as this emission source remains significant later into the evening on day 1 compared to the primary snow emission, which falls to zero after sunset. Previous box modeling studies have shown that even under low NO_x levels (<100 pptv), formation of BrONO₂ is significant (Liao, Huey, Tanner, et al., 2012; Thomas et al., 2012; Wang & Pratt, 2017), and under high NO_x levels (<700 pptv) formation of both BrO and HOBr are suppressed, whereas the rate of BrONO₂ formation remained largely unaffected (Custard et al., 2015). Due to the difficulty of measuring BrONO₂, no measurements have yet been reported in the Arctic to the best of our knowledge. Future work remains to compare the partitioning of HOBr and BrONO₂ under different NO_x conditions.

5.7. Boundary Layer VOC Oxidation Processes

We have shown (in Section 5.3) that with the addition of halogen emissions (BASE run), we obtain good agreement with the measured HO₂ concentration at the surface and predict an increase in OH (Figure 5). To further understand the links between halogens, HO_x cycling and oxidative chemistry, we analyze the major HO_x production and loss reactions, as well as VOC chemical lifetimes with respect to OH and Cl. First, we compare the difference in modeled HO_x concentrations between the NOSURF and BASE runs, as well as the change in partitioning of OH/HO₂ between the two runs.

Figure 9a shows the modeled HO_x and Cl atom concentrations in the NOSURF and BASE runs at 1.5 m AGL. We see a clear impact of halogens on surface HO_x concentrations, with up to a 30 times increase at the surface when the halogen snow and recycling emissions are active. This increase is largest within the daytime surface layer, coinciding with the high levels of simulated chlorine atoms, and is shown in Figure 9b as a ratio of HO_x between the BASE and NOSURF runs. Modeled Cl atom concentration at noon is higher than the average concentration predicted during the campaign of 2.0×10^5 atoms cm⁻³ (Liao et al., 2014). We calculate values of 2.9×10^5 and 1.1×10^6 atoms cm⁻³ for 18 and 19 March at noon, respectively. Our higher values can partly be explained by the overestimation of modeled Cl₂ on day 2, as well as the higher Cl₂ levels observed during this period compared to the campaign average Cl₂ levels.

Figure 9c shows the calculated OH/HO₂ ratio at 1.5 m AGL in the NOSURF and BASE model runs. We find a significant shift in the OH/HO₂ ratio toward HO₂ in our BASE run following the addition of halogen emissions compared to the NOSURF run. This difference is largest during the day, within the lowest 40 m of the atmosphere, with up to an order of magnitude difference, as shown in Figure 9d. This shift toward HO₂ in the BASE run can be explained by two main reasons. First, with the chlorine sources active in the BASE run, HO₂ formation via Cl-mediated VOC oxidation is greatly increased, skewing the ratio toward HO₂. This is in support of previous studies, which have suggested that HO₂ can be increased by the presence of chlorine, shifting the OH/HO₂ ratio significantly toward HO₂ (Piot & von Glasow, 2009; Rudolph et al., 1999; Thompson et al., 2015). Second, as the model was not constrained to any observations, the addition of halogen sources had a significant impact





Figure 9. (a) Modeled HO_x concentrations at 1.5 m above ground level in the NOSURF (blue) and BASE (red) runs and Cl-atom concentration (green). (b) HO_x concentration in the BASE run divided by the HO_x concentration in the NOSURF run as a function of altitude. (c) OH/HO₂ ratio at 1.5 m above ground level in the NOSURF (blue) and BASE runs (red). (d) OH/HO₂ ratio in the BASE run divided by the OH/HO₂ ratio in the NOSURF run as a function of altitude.

on the NO_x concentrations. Surface NO_x levels in the NOSURF run were several times greater than the BASE run, which greatly impacted OH formation. Thomas et al. (2012) showed that modeled surface concentrations of OH double with the inclusion of snowpack NO_x sources and bromine chemistry. This was mainly driven by the NO + HO₂ reaction under conditions where the halogen concentrations were significantly lower than those measured at Utqiaġvik during OASIS. In order to further understand HO_x cycling in our model, we analyze the major production and loss reactions of both OH and HO₂.

5.7.1. HO_x Chemical Budget

The main HO_x production and loss reactions at two heights (1.5 and 50.5 m) above the snow are shown in Figure 10. A clear chemical reactivity gradient is shown, with rates at 1.5 m approximately an order of magnitude greater than at 50.5 m, due to increased HO_x and Cl atom concentrations in the lower atmosphere. The principal OH production source in the model is the HO₂ recycling reaction with NO, at both the surface and above the boundary layer at 50.5 m AGL. Halogen-influenced OH production is clearly shown at 1.5 m, accounting for almost a quarter of surface OH production, with photolysis of HOBr (R6) contributing 14% and reactions involving chlorine comprising nearly 10%. This is a significant direct impact of snow-sourced halogens on the OH concentration. Snow emissions of other species, such as nitrous acid (HONO) and hydrogen peroxide (H₂O₂), could also be important sources of OH which may not be fully represented by our simulations due to missing snow emissions of these species in our model runs. At 50.5 m, modeled halogen concentrations are low with limited contribution to OH production at this height. Reaction between ozone and HO₂ is the second most important pathway for OH production at this height and is particularly important as it continues to convert HO₂ to OH





Figure 10. BASE simulation modeled (a) OH production (b) OH loss (c) HO_2 production, and (d) HO_2 loss each at 1.5 and 50.5 m above the snow surface. The five largest contributing reactions are shown for each with the percentage contributions shown as a pie chart.

for several hours after sunset. OH is lost via a multitude of reactions with organics, which can both recycle OH back into HO_2 and act as a source of CH_3O_2 and RO_2 . Mainly, OH loss is dominated by the reaction with CO, accounting for approximately a quarter of OH loss at both heights, which is also an important source of HO_2 .

At 1.5 m, the main HO₂ production reaction is the $CH_3O_2 + NO$ reaction (25%), followed by CO + OH (16%). CH_3O_2 is formed following oxidation of VOCs and methane by OH and Cl, with the rate constant of Cl + CH_4 roughly an order of magnitude greater than OH + CH_4 . Figure 11 shows the major production reactions of CH_3O_2 in our BASE run, with Cl + CH_4 responsible for almost two-thirds (64%) of surface CH_3O_2 production, whereas $OH + CH_4$ accounts for only 5%. At 50.5 m above the surface, production via Cl + CH_4 is negligible due to the low abundance of Cl atoms. Therefore, we can determine that snow emissions of chlorine drive the increase of surface HO_2 levels via CH_3O_2 formation. Consequently, this reaction cycle can also accelerate bromine recycling and ozone depletion, via (R5), linking together the chlorine and bromine chemical cycles. This effect is seen at 1.5 m, with BrO constituting 8% of HO₂ loss, with minimal contribution at 50.5 m.

In summary, it can be clearly seen that halogen emissions from snow make a significant contribution to HO_x production and loss reactions close to the surface. Chlorine and bromine chemical cycles are linked via peroxy radical formation, enhancing HO_x chemistry within the boundary layer, which can significantly impact VOC reactivity and lifetimes.





Figure 11. Modeled production reactions of CH_3O_2 at 1.5 and 50.5 m above the snow surface. The five largest contributing reactions are shown for each with the percentage contributions shown as a pie chart.

5.7.2. VOC Chemical Lifetimes

VOCs were measured during the campaign and the impact of Cl and Br atoms on VOC concentrations are discussed in detail in Hornbrook et al. (2016). The influence of chlorine chemistry on VOCs was determined during the campaign by recording the ratio of isobutane to n-butane (iC_4/C_4), as both alkanes react at similar rates with OH, but n-butane reacts approximately 1.5 times faster with Cl. iC_4/C_4 rose over the course of the campaign, indicating increased Cl-atom processing (and Cl₂ production) with increased availability of sunlight as spring progressed. Table 4 shows the simulated OH and Cl atoms concentrations at 1.5 and 50.5 m AGL used to calculate the chemical lifetimes (τ) of several VOCs presented in Table 5. For computational efficiency, some species within the RACM2 mechanism with similar reactivities are lumped together and treated as one species, such as propane and other similar organic compounds (HC3), as well as for pentane (HC5) and octane (HC8). At the surface, Cl atoms are abundant and rapidly oxidize VOCs, typically on the order of several hours, compared to OH which is generally on the order of days. As previously shown, surface OH concentration increases following the addition of halogen emissions, resulting in a reduction of VOC lifetimes by roughly 43% compared to the NOSURF run. This is a significant increase in the reactivity and processing of VOCs via OH due to the presence of halogens. At 50.5 m above the surface, this difference is minimal as levels of halogen radicals are very

Table 4
OH and Cl Concentrations at 1.5 and 50.5 m, at 12:00 AKST and 48-hr
Average in the NOSURF and BASE Runs

	1.5 m		50.5 m		
	12:00 AKST	00 AKST 48-hr average 12:00 AKST 48-hr		48-hr average	
Species	(molecules cm ⁻³)		(molecules cm ⁻³)		
[OH] _{NOSURF}	6.98×10^5	2.37×10^{5}	3.23×10^5	1.04×10^{5}	
[OH] _{BASE}	1.22×10^{6}	4.57×10^{5}	3.21×10^{5}	1.04×10^{5}	
[Cl] _{BASE}	7.19×10^5	2.65×10^{5}	22	7	

low, demonstrating the impact of chlorine chemistry close to the ground. We also see a clear gradient in chemical lifetimes with height and would expect longer lived VOCs above the boundary layer to act as a reservoir and replenish surface VOC concentrations by downward transport.

Indeed, as shown in Hornbrook et al. (2016), the VOCs sampled indicated more important halogen influence on atmospheric chemistry between the early hours of 18 March and 19 March. These VOC observations are likely a mix of local chemistry that is represented within our 1D model and chemistry that occurred while air masses resided over sea ice prior to sampling. Hornbrook et al. (2016) used ethyne levels to show there was a fairly consistent, but moderately low, Br atom influence atmospheric chemistry on 18 March. At the same time, measured acetaldehyde, propanal, and butanal decreased



Table 5

VOC Lifetimes at 1.5 and 50.5 m With Respect to OH and Cl, at 12:00 AKST and 48-hr Average in the NOSURF and BASE Runs

	12:00 AKST		48-hr average			
Species	$ au_{ m OH}^{ m NOSURF}$	$ au_{ m OH}^{ m BASE}$	$ au_{ m Cl}^{ m BASE}$	$ au_{ m OH}^{ m NOSURF}$	$ au_{ m OH}^{ m BASE}$	$ au_{ m Cl}^{ m BASE}$
1.5 m						
HC3	9.7 days	5.5 days	2.8 hr	28.5 days	14.8 days	7.5 hr
HC5 ^a	4.4 days	2.5 days	1.6 hr	13.1 days	6.8 days	4.4 hr
HC8 ^a	1.8 days	1.0 days	1.1 hr	5.2 days	2.7 days	3.0 hr
Ethane	139 days	79.4 days	7.1 hr	1.1 years	212 days	19.3 hr
Ethene	1.7 days	23.4 hr	2.0 hr	5.0 days	2.6 days	5.3 hr
Acetaldehyde	20.9 hr	12.0 hr	4.8 hr	2.6 days	1.3 days	13.1 hr
Acetone	112 days	64.0 days	11.6 days	329 days	171 days	31.9 days
MEK	15.9 days	9.1 days	9.2 hr	46.9 days	24.3 days	1.0 days
Aldehydes ($\geq C_3$)	15.7 hr	9.0 hr	3.0 hr	1.9 days	1.0 days	8.1 hr
Toluene	2.2 days	1.2 days	6.5 hr	6.4 days	3.3 days	17.8 hr
50.5 m						
HC3	20.8 days	20.9 days	10.3 years	64.6 days	64.6 days	32.4 years
HC5 ^a	9.5 days	9.6 days	6.0 years	29.7 days	29.7 days	18.9 years
HC8 ^a	3.8 days	3.9 days	4.2 years	11.9 days	11.9 days	13.1 years
Ethane	295 days	297 days	26.6 years	2.5 years	2.5 years	83.5 years
Ethene	3.7 days	3.7 days	7.4 years	11.5 days	11.5 days	23.1 years
Acetaldehyde	1.9 days	1.9 days	18.0 years	5.9 days	5.9 days	56.6 years
Acetone	241 days	243 days	1,042 years	2.1 years	2.1 years	3,275 years
MEK	34.4 days	34.6 days	34.2 years	107 days	107 days	108 years
Aldehydes ($\geq C_3$)	1.4 days	1.4 days	11.1 years	4.4 days	4.4 days	34.9 years
Toluene	4.7 days	4.8 days	24.4 years	14.7 days	14.7 days	76.8 years

Note. HC3, HC5, and HC8 represent lumped hydrocarbon species with similar reactivities and average carbon chains of 3, 5, and 8, respectively. We therefore calculate approximate chemical lifetimes using the rate constants of N-pentane and isopentane (in equal proportions) with Cl for HC5 and N-octane and iso-octane (in equal proportions) with Cl for HC8. Rate constants are obtained from Calvert et al. (2015) at 248 K.

^aReactions of HC5 and HC8 with Cl were not included in the chemical mechanism.

by approximately 50%, 75% and 90% respectively (see Figure 14 in Hornbrook et al. (2016)). As well, the butanal observations indicated a gradient between the lowest sampling height, 0.6 m, and the other two sampling heights at 1.5 and 5.4 m, in which the mixing ratio nearest the snow surface reached levels as low as half that at the higher sampling inlets, consistent with Cl atom chemistry near the surface. Overall, our results show that measurements above the Arctic snow surface can be highly influenced by halogen chemistry directly or indirectly via increases in HO_x concentration, resulting in a highly reactive surface layer. Deriving accurate Cl atom concentrations from VOC measurements can therefore be challenging, as surface VOC and Cl atom concentrations are dependent on vertical mixing, surface halogen emissions, and chemistry. In a well-mixed system, downward transport of VOCs can replenish concentrations at the surface, potentially resulting in an underestimation of the derived Cl atom concentration at the ground.

6. Conclusions and Perspectives

In this study, we examined the role of Arctic halogen emissions from snow on boundary layer oxidation processes using an updated version of the PACT-1D model. Snow emissions of Cl_2 and Br_2 were added to the model, including primary production from land-based surface snow and heterogeneous recycling on aerosols and snow.

We compared the model against observations from the 2009 OASIS campaign at Utqiagvik, Alaska, when high atmospheric Cl_2 levels were observed (18–19 March). The modeled halogen concentrations showed excellent agreement with the observations upon the addition of halogen emissions. The main conclusions of our study can be summarized as follows:

- 1. Surface Arctic halogen observations are reproduced by the model when including the combined effects of halogen emissions from snow, vertical mixing and atmospheric chemistry. Primary emissions of Cl₂ from snow, parameterized using solar irradiance and measured surface ozone concentration, can describe surface observations of Arctic Cl₂. Modeled Br₂ levels are in good agreement with observations when using a combination of both primary emissions from snow and heterogeneous surface recycling of BrONO₂ and HOBr. Sensitivity analyses showed that increased heterogeneous recycling of halogens on aerosols could not explain surface observations and only provided a minor source of reactive halogens in our model simulations (AERO simulation)
- 2. Boundary layer dynamics, vertical mixing, chemistry and emissions all strongly impact halogen vertical distribution. During the day, Cl₂ is confined to within the lowest 15 m of the atmosphere on both days of the simulation period. Stable conditions during this period resulted in a shallow surface layer, hindering vertical mixing and impacting surface concentrations. In particular, changes in the model vertical mixing and boundary layer dynamics result in a reduction of up to 60 pptv of Cl₂ at 1.5 m during the day (BLD simulation)
- 3. HO_x radical concentration is increased by up to a factor of 30 with the inclusion of halogen emissions in the model. The increase in OH was primarily driven by elevated HOBr levels and its subsequent photolysis (R6). A significant contributor of HO₂ production is the CH₃O₂ radical formed via the Cl + CH₄ reaction (R4). This also caused a decrease in the modeled OH/HO₂ ratio which is attributable to chlorine chemistry
- 4. Increased HO_x radicals and a high Cl atom concentration near the surface significantly increases chemical reactivity within a shallow layer near the surface. Modeled VOC lifetimes, with respect to OH, are reduced by approximately 43% due to the presence of halogens (BASE run). Cl atoms concentrated near the surface rapidly react with VOCs, but this reactivity becomes much weaker with height and negligible over 15 m above the surface.

We have proposed two model parameterizations for Cl₂ and Br₂ emissions from land-based snow that have been applied to understanding observations during OASIS. Toyota et al. (2011) and recently Marelle et al. (2021) have considered bromine activation, triggered from snow on sea ice, with different efficiency in sunlit compared to dark conditions. In these studies, land-based snow sustains bromine chemistry via conversion of deposited HOBr, BrONO₂, and HBr to form Br_2 via reaction of trace quantities of bromide that is assumed to be present in all land-based snow. This is likely too simplified, but works, due to the fact that activated bromine in the atmosphere is lost via deposition and other processes away from the coasts, which also turns off the land-based snow source of Br₂. Here we propose a slightly different approach, which is consistent with our prior modeling study at Summit, Greenland (Thomas et al., 2011) that showed trace amounts of bromide in snow can be activated via photochemistry without enhanced atmospheric bromine already present. This approach is also consistent with the chamber studies presented in Pratt et al. (2013), which showed that land-based snow can release reactive bromine without the need for any other triggers than ozone and sunlight. Land-based snow as a primary source of activated bromine should be tested in 3D regional models using this proposed parameterization, however we note this may not be important regionally compared to the quantities of bromine that are released from snow on sea ice and sea salt aerosols. Physical changes in the snowpack or other properties tied to temperature also influence bromine release from snow on sea ice. For example, Burd et al. (2017) have shown that over the Arctic Ocean atmospheric bromine chemistry is quickly deactivated upon warming to near freezing temperatures and that bromine chemistry is reactivated upon fresh snowfall. The deactivation of bromine release from snow on sea ice upon reaching near freezing temperatures is already included in some models (Marelle et al., 2021; Toyota et al., 2011). At present, less is known on how temperature changes (including related changes in snow physics, impurity locations, etc.) influence bromine release from land-based snow. The inland snow source of reactive bromine is already partially considered in the Toyota et al. (2011) land-based snow treatment which considers all snow to have reactive bromide available for release upon HBr, BrONO₂, and HOBr deposition. However, the exact description of how this works is not known and hypotheses should be tested in models and further characterized via observations in the future.



Our modeling work suggests that some description of chlorine emissions from snow is needed to capture the atmospheric chemistry occurring near the surface during Arctic spring. Coastal Arctic surface stations where chlorine has been observed are also the same stations where all long term observations of VOCs, aerosols, and ozone are made. Therefore, understanding and modeling how chlorine contributes to atmospheric chemistry at these locations is essential. We have proposed a model parameterization of chlorine release from land-based snow that depends on available sunlight and ozone. This description needs to be tested during other periods of the year and under different conditions. How this chlorine source depends on other parameters, such as snow chloride content, snow properties, etc., is currently not known due to very limited observations of activated chlorine in the Arctic atmosphere. There is also some evidence that chlorine emissions should be considered from snow on sea ice (e.g., Peterson et al. (2019)). The regional and seasonal sensitivity to potential snow emissions of chlorine needs to be explored with model sensitivity studies using reasonable hypotheses and with dedicated observations in the future.

One of the most important conclusions from our study is that the vertical extent of a highly oxidizing layer with active chlorine chemistry may only extend to 15 m above the snow surface. This is much smaller than the vertical resolution of most regional and global models. The effects of this highly oxidizing layer on new particle formation, aerosol processing, VOC chemistry, and other atmospheric chemistry processes need to be explored and eventually included in models. One possibility is to implement sub-grid scale parameterizations of these processes with regional and global chemical models in the future. The influence of bromine chemistry vertically and with respect to the presence of aerosols and mesoscale weather systems has been relatively well characterized (Blechschmidt et al., 2016; Burd et al., 2017; Liao, Huey, Scheuer, et al., 2012; Oltmans et al., 2012; Peterson et al., 2017; Simpson et al., 2017). However, less is known from observations regarding whether there is active chlorine chemistry above the surface. More observations are needed to quantify both bromine and chlorine chemistry above the surface in order to understand how to best include these effects within models.

Data Availability Statement

The code for the model and input files are all publicly available on Zenodo as Ahmed et al. (2022) https://doi. org/10.5281/zenodo.6045999. The model outputs for the BASE run can also be found on Zenodo at https://doi. org/10.5281/zenodo.5654628. The data for this campaign are publicly available through the NSF Arctic Data Center at https://arcticdata.io/.

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