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Key Points:

- Bromine (Br) concentrations increased 3.5-fold from pre-industrial to 1975 and declined 50% by 1999 in a Russian Arctic ice-core
- A robust correlation between ice-core Br and acidity highlights acidity's key role in influencing the atmospheric Br budget
- Model shows acid-catalyzed sea-salt debromination is the largest source of reactive Br and drives ice-core Br trends

Supporting Information:

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

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Anthropogenic Influence on Tropospheric Reactive Bromine Since the Pre-industrial: Implications for Arctic Ice-Core Bromine Trends

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Abstract Tropospheric reactive bromine (Br_y) influences the oxidation capacity of the atmosphere by acting as a sink for ozone and nitrogen oxides. Aerosol acidity plays a crucial role in Br_y abundances through acid-catalyzed debromination from sea-salt-aerosol, the largest global source. Bromine concentrations in a Russian Arctic ice-core, Akademii Nauk, show a 3.5-fold increase from pre-industrial (PI) to the 1970s (peak acidity, PA), and decreased by half to 1999 (present day, PD). Ice-core acidity mirrors this trend, showing robust correlation with bromine, especially after 1940 ($r = 0.9$). Model simulations considering anthropogenic emission changes alone show that atmospheric acidity is the main driver of Br_y changes, consistent with the observed relationship between acidity and bromine. The influence of atmospheric acidity on Br_y should be considered in interpretation of ice-core bromine trends.

Plain Language Summary Reactive bromine in the air impacts major oxidants in our atmosphere, which remove pollutants and greenhouse gases and has changed over time in the Russian Arctic. Ice-core bromine and acidity show a significant increase from pre-industrial to the 1970s followed by a decrease. Our study suggests that human activities caused changes in bromine through the emissions of acidic gases from fossil fuel combustion. Considering relationships between atmospheric acidity and bromine is crucial to interpreting bromine variations in ice cores.

1. Introduction

Tropospheric reactive bromine influences tropospheric oxidation capacity, destroys ozone via catalytic cycles, and perturbs OH to HO_2 ratios toward OH (von Glasow et al., 2004). In coastal polar regions, reactive bromine is responsible for springtime boundary layer ozone depletion events (ODEs) (Barrie et al., 1988; McConnell et al., 1992; Simpson, von Glasow et al., 2007) and gaseous elemental mercury deposition (Steffen et al., 2008; Wang et al., 2019), thus also impacting the transfer of toxic mercury from the atmosphere into the ecosystem.

Natural sources of tropospheric gaseous reactive bromine ($\text{Br}_y = \text{Br} + \text{HBr} + \text{BrNO}_2 + \text{BrNO}_3 + \text{BrO} + \text{HOBr} + 2\text{Br}_2 + \text{IBr} + \text{BrCl}$) include sea-salt debromination (Kerkweg et al., 2008; Yang et al., 2005), organobromines emitted from the marine biosphere (Liang et al., 2010; Quack & Wallace, 2003), saline lakes (Hebestreit et al., 1999; Matveev et al., 2001), and volcanoes (Aiuppa et al., 2005; Bobrowski et al., 2003). Anthropogenic sources of Br_y include methyl bromide (CH_3Br) emissions from agricultural pesticide (Clerbaux et al., 2007), coal burning (Lee et al., 2018), biomass burning, and stratospheric transport from halon degradation. Gaseous CH_3Br and aerosol PbBrCl exhaust emissions from leaded gasoline vehicles have been important since the 1920s, but have diminished since the 1970s because of leaded gasoline abatement in North America and Europe (Habibi, 1973; Lammel et al., 2002; Thomas et al., 1997). The 2-fold increase in summertime ice-core bromine after 1950 observed in the Col du Dome ice-core (French Alps) was largely (70%) attributed to leaded gasoline emissions (Legrand et al., 2021b). Global model simulations showed a 39% increase in global Br_y burden from pre-industrial (PI) to present day (PD) driven by enhanced oceanic iodine production from ozone deposition, increased anthropogenic bromocarbons, and increased bromine flux from stratosphere (Sherwen et al., 2017).

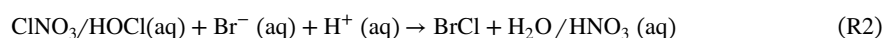
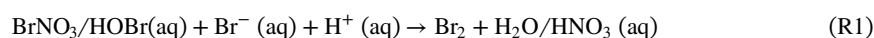
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However, the model did not consider leaded gasoline or sea-salt debromination, the latter being the largest global source (95%) of Br_y (Wang et al., 2021).

Polar regions have additional sources of Br_y from saline, blowing snow-sourced sea-salt-aerosol (Huang et al., 2020; Savelyev et al., 2006; Yang et al., 2008) and direct snowpack emissions (Abbatt et al., 2012; Foster et al., 2001; Pratt et al., 2013; Simpson, Carlson, et al., 2007, Simpson, von, et al., 2007; Stutz et al., 2011). Both sources are most efficient from snow overlying first-year-sea-ice (FYI), which is thinner, more saline (Confer et al., 2023; Frey et al., 2020; Yang et al., 2008), and contains more snow bromine from upward migration of brine from the sea-ice surface (Nandan et al., 2017; Peterson et al., 2019) than snow over multi-year-sea-ice (MYI). The sea-ice source of Br_y has led to the use of Arctic ice-core bromine as a proxy for historical sea-ice extent (Spolaor et al., 2014, 2016; Sturges & Barrie, 1988; Vallelonga et al., 2021). For example, Spolaor et al. (2016) reported a positive correlation ($r = 0.44$) between bromine excess (Br_{exc}) records from a Russian Arctic ice-core, Akademii Nauk (AN), and spring sea-ice area of the Laptev Sea from 1980 to 1998, with both showing a decreasing trend.

Although total sea-ice extent has been decreasing since at least 1979 (Cavalieri & Parkinson, 2012), FYI has been increasing with the decline of MYI (Bougoudis et al., 2020; Confer et al., 2023). Sea-salt concentrations are observed to increase by 9–12% decade⁻¹ during spring and winters in 1980–2017 at Alert, Canada (Confer et al., 2023). Model simulations also show a pan-Arctic increase in surface sea-salt-aerosol concentrations since 1980, mostly due to enhanced blowing snow emissions driven by increased FYI (Confer et al., 2023). With increased FYI in the Arctic since at least 1980, snowpack-emitted bromine may also become a more efficient source (Swanson et al., 2022). Indeed, satellite observations showed increasing trends in tropospheric BrO columns over Arctic sea-ice (+15% decade⁻¹) during polar spring from 1996 to 2017 (Bougoudis et al., 2020).

Although sea-salt release of Br_y is a natural process, the debromination reaction is catalyzed by aerosol acidity (R1–2) (Eigen & Kustin, 1962; Fan & Jacob, 1992), which is influenced by anthropogenic emissions of acidic gases and acidic aerosol precursors such as SO₂ and NO_x(= NO + NO₂).



In the Arctic, snow acidity measured in ice cores began increasing in the 1940s, peaked in the 1970s, and decreased to the present day (e.g., Geng et al., 2014). Therefore, the quality of ice-core bromine as a sea-ice proxy may be compromised by these anthropogenic influences (Maselli et al., 2017). The relative impacts of sea-ice extent and atmospheric acidity on bromine trends since PI remain unclear.

Here, we use a global model to quantify the impact of anthropogenic emissions on tropospheric reactive bromine abundances since the pre-industrial. Since previous work suggests that the Russian Arctic AN ice-core preserves an atmospheric signal while Greenland ice cores may not (Zhai et al., 2023), we compare the model to AN ice-core bromine observations. We use the model-ice core comparison to examine implications of anthropogenic emissions for tropospheric Br_y abundance and interpretation of Arctic ice-core bromine records.

2. Methods

2.1. GEOS-Chem Model and Historical Simulations

We use a global 3D chemical transport model GEOS-Chem (version 11-02d, <https://github.com/geoschem/geoschem/tree/v11-02d-prelim>) for historical simulations. The model is driven by MERRA-2 assimilated meteorological fields from the Goddard Earth Observing System (GEOS) (Gelaro et al., 2017), and contains detailed HO_x-NO_x-VOC-ozone-halogen-aerosol tropospheric chemistry (Wang et al., 2021) and fully coupled stratospheric chemistry (Eastham et al., 2014). Details of the modeled bromine chemistry are shown in Figures S1–S3 in Supporting Information S1. Sea-salt-aerosol debromination occurs in both open ocean (Jaeglé et al., 2011) and blowing snow (Huang & Jaeglé, 2017) sourced sea-salt-aerosol. Following Zhai et al. (2023), ozone dry deposition velocity onto snow and ice is updated to 0.01 cm s⁻¹, consistent with observations (Simpson, Carlson, et al., 2007, Simpson, von Glasow, et al., 2007). Snowpack bromine emissions (Swanson et al., 2022; Zhai et al., 2023) are not included in the model. The magnitude of the snowpack emission source is estimated to be 278 Gg Br/yr during spring in the AN source regions, based on model simulations from Swanson et al. (2022).

Model simulations are performed under three anthropogenic emission scenarios: pre-industrial (PI, CE 1750), peak atmospheric acidity (PA, CE 1975), and present day (PD, CE 1999). Anthropogenic and biomass burning emissions vary between simulations and are from Community Emissions Data System (CEDS, McDuffie et al., 2020) and BB4CMIP6 (van Marle et al., 2017), respectively, from individual years (1750, 1975, 1999). The emitted species include but are not limited to acidic precursors, aerosol, organic compounds, methane, and long- and short-lived organohalogens. Details of the emission setup can be found in Zhai et al. (2021), and trends of anthropogenic emissions of SO₂, NO_x, and NH₃ are in Figure S4 in Supporting Information S1. After a 1-yr spin-up, each simulation is run for 1 yr using 2007 meteorology and sea-ice extent. By using the same meteorology, we aim to isolate changes induced by anthropogenic emissions. All simulations are conducted at 4° × 5° horizontal resolution and 72 vertical levels up to 0.01 hPa.

Additionally, the model considered bromine emissions from coal and leaded gasoline consumption. Bromine emissions from coal-fired power plants in the chemical form HBr are expected to be correlated with SO₂ emissions, assuming similar Br:S ratios in coal and the fact that SO₂ control strategies such as wet flue gas desulphurization also remove bromine from exhaust (Lee et al., 2018; McTigue et al., 2014). To calculate coal burning emitted HBr, we scale coal burning emitted SO₂ with a scale factor of Br_x:SO₂ (where Br_x = BrCl + BrNO₂ + 2×Br₂) = 1.5 × 10⁻⁴ ppb:ppb, which is the median of observed Br_x:SO₂ ratios from Lee et al. (2018). This is likely an underestimate since the major form of reactive bromine emitted from coal burning is HBr, which was not measured.

We also added historical emissions of particle-phase PbBrCl and gas-phase CH₃Br from leaded gasoline for PA. Brominated compounds were added to leaded gasoline as additives since 1923 to prevent lead deposition in engines (Thomas et al., 1997). Leaded gasoline phase-out efforts, starting in the 1970s in North America and 1980s in Europe, effectively eliminated global Pb emissions from this source (Hagner, 2002; Huang et al., 1996; Nriagu, 1990). Annual global bromine usage in gasoline peaked in the early 1970s at 170 ± 20 Gg, decreasing to 100 ± 11 Gg in 1980 (Thomas et al., 1997). Assuming complete Br emission into the atmosphere, we estimated global bromine emissions from leaded gasoline in 1975 (average of the early 1970s and 1980, 135 ± 15 Gg) using the transportation-emitted CO and global mean transportation-CO:Br ratio from a global anthropogenic emission inventory, CEDS (McDuffie et al., 2020). Leaded gasoline-sourced bromine is emitted as 85% PbBrCl and 15% CH₃Br based on exhaust measurements (Habibi, 1973; Harsch & Rasmussen, 1977). PbBrCl is modeled as accumulation-mode aerosol, considering the majority of lead particles emitted are <5 μm in wet diameter (Habibi, 1973). Table S1 in Supporting Information S1 shows global and regional total bromine burden from leaded gasoline and coal combustion.

2.2. Ice-Core Bromine and Acidity Observations

A 724 m ice-core was collected from the Akademii Nauk ice cap (80.5°N, 94.8°E, 750 m a.s.l.) at Severnaya Zemlya in the Russian Arctic from 1999 to 2001 (Fritzsche et al., 2002; Opel et al., 2013). The snow accumulation rate is about 440 kg m⁻² yr⁻¹. According to the updated age model, the ice core covers an estimated time range of year 200 BCE–1999 CE (McConnell et al., 2019), but we only present records since CE 1750. This ice core was chosen because it is expected to preserve an atmospheric signal of bromine due to its high snow accumulation rate and high-latitude location, both of which limit photochemical loss of bromine from the snowpack (Zhai et al., 2023).

Total sodium, total bromine, and acidity records from the AN core are previously published (Zhai et al., 2023) and are briefly described here. Elemental sodium (Na) and bromine (Br) were measured continuously with high resolution inductively coupled plasma mass spectrometry (HR-ICP-MS) with an uncertainty of ±10% using the melter-based ice-core analytical system at the Desert Research Institute (Maselli et al., 2017). Mineral acidity (H⁺) was measured concurrently using continuous flow analysis (Pasteris et al., 2012), with an uncertainty below 5%. Bromine excess (Br_{exc}) is calculated as shown in Equation 1.

$$[\text{Br}_{\text{exc}}] = [\text{Br}]_{\text{ice-core}} - [\text{ssNa}]_{\text{ice-core}} \times ([\text{Br}]/[\text{Na}])_{\text{seawater}} \quad (1)$$

where [X] is mass concentrations in ng × g ice⁻¹, and ([Br]/[Na])_{sea water} is the seawater Br/Na mass ratio of 0.00624 (Millero et al., 2008). [ssNa] is calculated based on ice-core calcium concentrations, as described in Maselli et al. (2017).

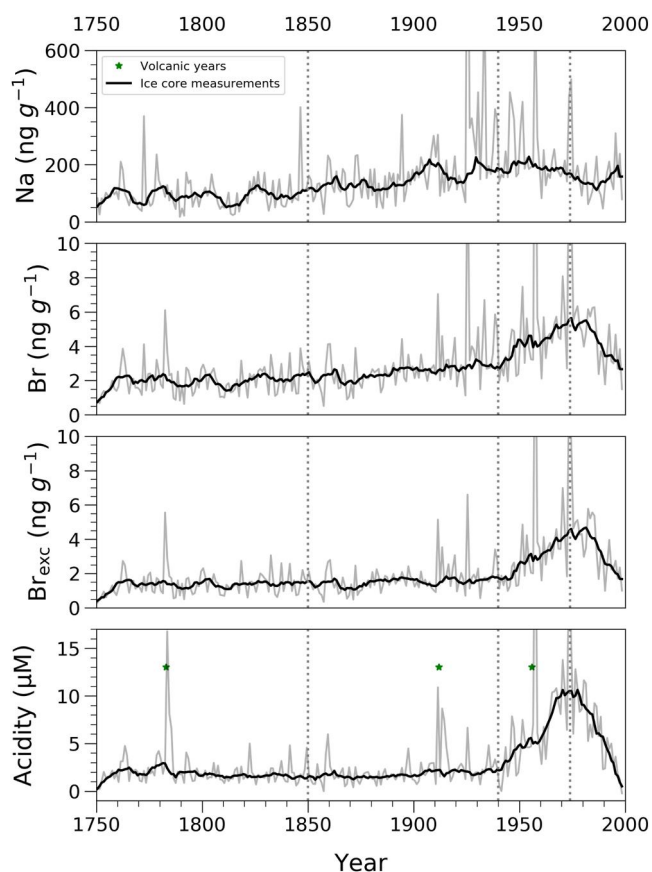


Figure 1. Total sodium, total bromine, Br_{exc} , and acidity concentrations in the Akademii Nauk ice-core. Gray lines are the measured annual ice-core concentrations (total sodium, total bromine, and acidity) or calculations (Br_{exc}), and black lines are the 9-yr running average to reflect long-term variations, with outliers outside of $1.5 \times IQR$ (interquartile range) removed. Green stars mark the large and moderate volcanic years identified in previous study (Opel et al., 2013). Dashed vertical gray lines mark the years 1850, 1940, and 1975.

3. Results

3.1. Ice-Core Bromine Records in the AN Ice-Core

Figure 1 shows total Na, total Br, Br_{exc} , and acidity concentrations in the AN ice core. Change point analysis (Ruggieri, 2013; Zhai et al., 2023) does not identify any change point in the Na record, and there is a slight increasing trend of $0.67 \pm 0.00 \text{ ng g}^{-1} \text{ yr}^{-1}$ in Na throughout the record, resulting in a 77% increase from the pre-1850 average Na concentration to that of post-1970. Total Br, Br_{exc} , and acidity show no trends during 1850–1940, and their large increases start in the 1940s. Total Br has an average concentration of $2.02 \pm 0.88 \text{ ng g}^{-1}$ before 1850, increased by 3.5-fold from pre-1850 to the 1970s, and decreased by 53% after the 1970s to the end of the record (CE 1999). Similar trends are shown in Br_{exc} , with a 4-fold increase from the pre-1850 level to the 1970s, followed by a 59% decrease (Table S2 in Supporting Information S1). Br_{exc}/Br ratio range from 26 to 94%, and are higher ($80\% \pm 6\%$) in the 1970s compared to PI (1750–1850, $69\% \pm 11\%$) and PD (post-1989, $70\% \pm 11\%$). Acidity shows a similar trend as Br, with a background concentration of $2.16 \pm 1.93 \text{ } \mu\text{M}$ before 1850, and an increase of 5.5-fold from pre-1850 to 1975, and decreased by 72% back to PI level by 1999.

Figure 2 shows the relationship between ice-core bromine (Br and Br_{exc}) and acidity. A significant positive correlation ($p < 0.01$) is shown for pre-1850, post-1940, and the full record, with post-1940 showing the highest correlation coefficient ($r = 0.9$). The relationship between ice-core bromine and acidity throughout the record is much stronger than that of bromine and spring sea-ice extent for 1980–1999 reported in Spolaor et al. (2016) ($r = 0.44$), suggesting that it is possible that factors other than sea-ice, such as acidity, may play a role in driving ice-core bromine trends.

3.2. Ice-Core Model Comparison

Figure 3a and Table S3 in Supporting Information S1 show the modeled annual mean tropospheric total bromine burden and its speciation in the AN 5-day back trajectory region (TRJ, Figure S5 in Supporting Information S1) in PI, PA, and PD. Total Br and Br_{exc} are calculated based on Equation 2 and Equation 3, respectively,

$$Br_{total} = Br_y + Br_a^- + Br_c^- \quad (2)$$

$$Br_{exc} = Br_{total} - ssNa \times \left(\frac{[Br]}{[Na]} \right)_{sea\ water} \quad (3)$$

where Br_{total} is total Br, Br_a^- and Br_c^- in Equation 2 are bromide in accumulation-mode and coarse-mode aerosol, respectively, and ssNa sodium in sea-salt-aerosol (both modes).

Modeled Br_y contributes about 95% of total Br in the three time periods, while aerosol bromine is minor (5%); therefore, total Br shows similar distribution and trends as Br_y (Figure S6 in Supporting Information S1). The most abundant Br_y species in PI are Br_2 and BrO, contributing 48% of total Br. Br_2 and BrO burdens remain similar from PI to PA, while HOBr, $BrNO_3$, and BrCl increase by 125%, 392%, and 158%, contributing 24%, 21%, and 11% to total Br in PA, respectively. Increased fractions of $BrNO_3$, HOBr, and BrCl in total Br_y are driven by enhanced NO_x and HO_2 concentrations and more active coupled chlorine-bromine chemistry from PI to PA. Modeled total Br burden and speciation in PD are similar to those of PA. HOBr, $BrNO_2$, and Br_a^- decreased slightly from PA to PD while other species increased.

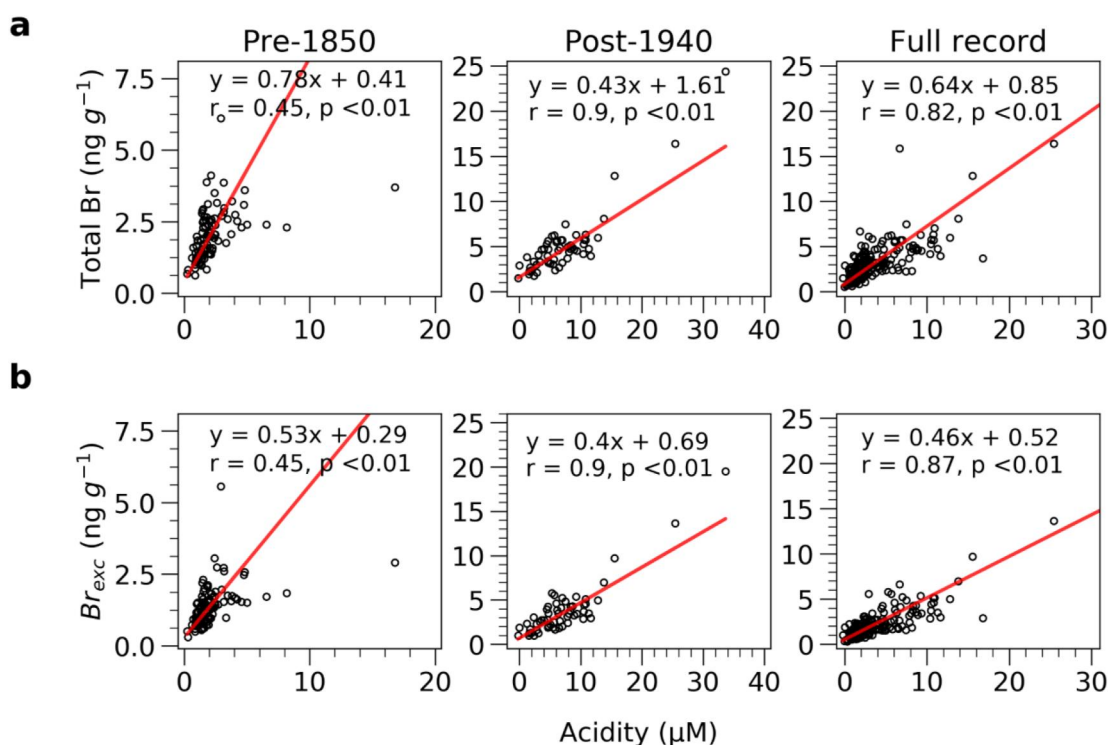


Figure 2. Relationship between concentrations of (a) total Br and acidity, and (b) Br_{exc} and acidity from the AN ice core, for 1750–1850 (pre-1850), 1940–1999 (post-1940), and 1750–1999 (full record). Black circles show the measured Br, Br_{exc} , and acidity concentrations, and red lines represent the reduced major axis regression, with functions shown in each panel.

Br_a^- increased 2.8-fold from PI to PA and decreased 30% from PA to PD (Figures S2 and S3 in Supporting Information S1). In contrast, Br_c^- decreased by 40% from PI to PA and remained low in PD (−0.2% from PA to PD). The opposite trends in Br_a^- and Br_c^- from PI to PA suggest that debromination liberated bromide from coarse-mode aerosol to the gas phase and partially re-partitioned into the accumulation-mode aerosol. $Br_{exc}/total$ Br ratios are 78%, 82%, and 88% for PI, PA, and PD, respectively, consistent with values observed in the AN ice-core (Table S2 in Supporting Information S1).

Figures 3b and 3c compare the percent changes in modeled total bromine and Br_{exc} burdens, respectively, with ice-core total bromine and Br_{exc} concentrations between the 3 time periods. The model predicts a 2-fold increase in total Br and Br_{exc} from PI to PA and only a small decrease (3%) from PA to PD. In comparison, ice-core Br and Br_{exc} show a more symmetrical change centered in 1975 with a 3-fold increase from PI to PA, and 50% decrease from PA to PD. The modeled increase (75% and 96% for total Br and Br_{exc} , respectively) from PI to PD is quantitatively consistent with ice-core observations (64% and 67% for total Br and Br_{exc} , respectively), suggesting that the model may underestimate Br burden in PA (Figure 3b).

3.3. Factors Controlling Modeled Bromine Trends

Figure 4 shows the budget analysis for Br_y , HOBr, $BrNO_3$, and BrCl in the AN TRJ. Sea-salt debromination dominates (>99%) Br_y production in PI, PA, and PD. Organobromines and anthropogenic bromine, including coal combustion and leaded gasoline emissions, contribute less than 1% to the total Br_y sources. Uptake onto sea-salt-aerosol removes >80% of Br_y , while the rest either deposits or is transported out of the TRJ region. Budget analysis shows that HBr uptake on fine-mode-aerosol is the dominant source of Br_a^- , at least one order of magnitude larger than natural emissions from the open ocean and blowing snow on top of sea ice.

Total Br_y production rate increased by 236% from PI to PA. The increase is predominantly driven by enhanced sea-salt-aerosol debromination due to a decrease in aerosol pH from 2.5 to 0.9 in the AN TRJ (Figure S7 in Supporting Information S1). This is consistent with the observed strong correlation ($r = 0.9$, $p < 0.01$) between ice-core Br and acidity after 1940 (Figure 2). From PI to PA, the large increase in Br_y

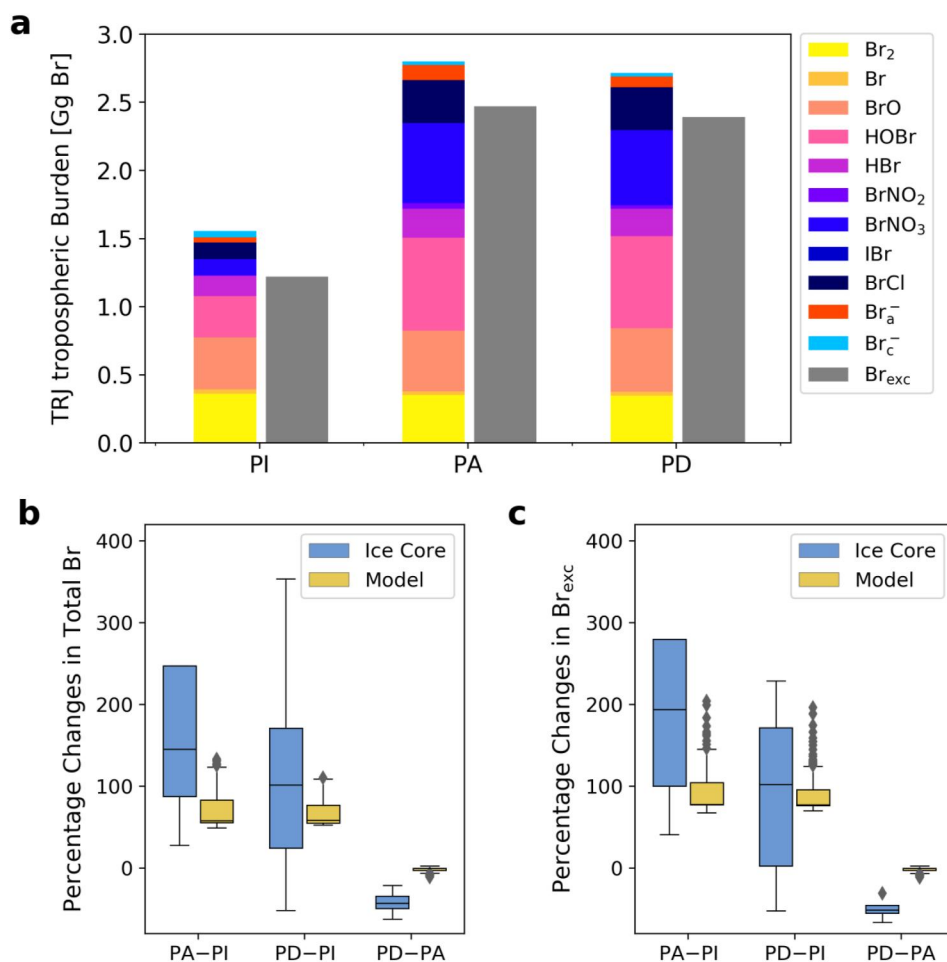


Figure 3. Modeled bromine burden and speciation in the three time periods and model ice-core comparison. (a) Modeled annual mean tropospheric burdens of bromine species (colored bars) and the calculated Br_{exc} burden (gray bars) in the 5-day back trajectory region of AN for PI, PA and PD. Comparison of percentage changes in annual mean tropospheric (b) total Br and (c) Br_{exc} burdens between PI, PA, and PD from AN ice-core records and model simulations. Boxes show the interquartile range (IQR, 25th to 75th percentiles), and the bar in the middle shows the median. Whiskers show $1.5 \times$ IQR range, and data outside of the whiskers are considered outliers.

burden was driven by increases in HOBr, BrNO₃, and BrCl. The only source of BrNO₃ is the reaction of BrO and NO₂, which increased 3.5-fold caused by enhanced NO₂ (Figure S8 in Supporting Information S1) and BrO abundance (+17%). BrCl production rate increased 1.6-fold, and this is mainly driven by more active halogen recycling on accumulation-mode aerosol ($\text{HOBr} + \text{Cl}^- + \text{H}^+ \rightarrow \text{BrCl} + \text{H}_2\text{O}$), caused both by elevated HOBr burden (Figures S1 and S2 in Supporting Information S1) and more acidic aerosol (Figure S7 in Supporting Information S1). HOBr production rate increased 1.2-fold, caused mainly by increased oxidation of BrO by HO₂, which is driven by enhanced BrO abundance (+17%) and higher HO₂ mixing ratios (+33%) from PI to PA.

From PA to PD, Br_y production rate decreased by 21%, due to reduced aerosol acidity (Figure S7 in Supporting Information S1) and less debromination. Interestingly, modeled Br_y tropospheric burden shows minor changes (Figure 3) because the decrease in Br_y production rate is counterbalanced by a decrease in HBr uptake onto sea-salt-aerosol from PA to PD. This decreasing trend in HBr uptake is driven by the lower tropospheric abundance of HBr (−6%) caused by less formation from the reaction between Br radical and acetaldehyde ($\text{Br} + \text{CH}_3\text{CHO} \rightarrow \text{HBr} + \text{CH}_3\text{CO}$), which is the main source of HBr. In other words, the reduction in tropospheric acetaldehyde burden from PA to PD (−29%), resulting from reduced organic precursors (alkanes and alkenes) emitted from coal combustion and vehicles (McDuffie et al., 2020), causes a decrease in Br_y sinks.

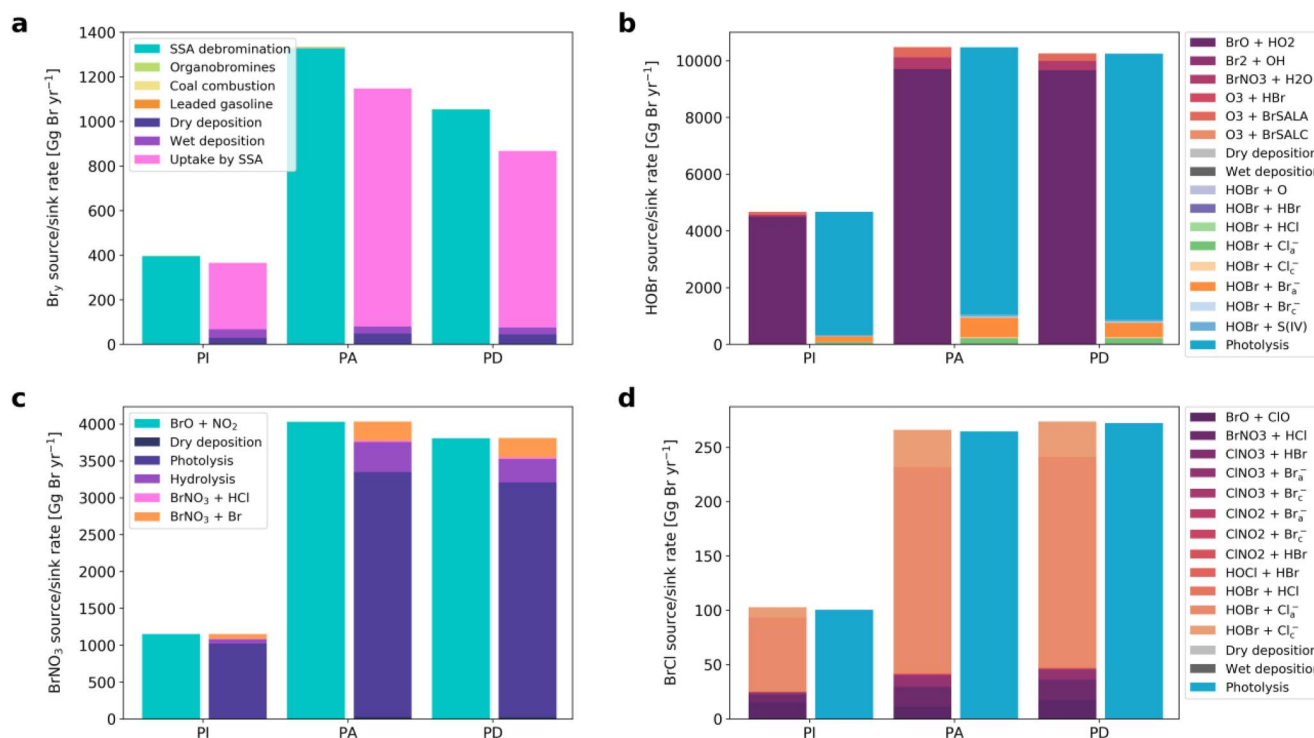


Figure 4. Modeled annual mean sources and sinks of tropospheric (a) Br_y (b) HOBr (c) BrNO_3 and (d) BrCl in the AN TRJ for PI, PA, PD. For each time period, left bar shows the sources and right bar shows the sinks. Differences between total sources and total sinks of Br_y can be explained by Br_y that transported outside of the AN TRJ. X_a^- , X_c^- ($\text{X} = \text{Cl}, \text{Br}$) are chloride and bromide in accumulation- and coarse-mode sea-salt-aerosol, respectively. S(IV) represents SO_3^{2-} and HSO_3^- in cloud droplets.

4. Discussion

The historical simulations with changes in anthropogenic emissions alone can quantitatively capture the AN ice-core bromine changes from PI to PD, but they underestimate the increase in the AN ice-core bromine from PI to PA and do not reproduce the observed -53% decrease in ice-core bromine from PA to PD. No sea-ice changes are simulated, but a model sensitivity study with 1980 meteorology and PA emissions, which has higher total sea-ice extent but lower FYI (-42%) than 2007 (Figure S9 in Supporting Information S1), shows a lower (-6%) tropospheric Br burden due to decreases in blowing snow sourced sea-salt-aerosol (Figure S10 in Supporting Information S1), consistent with the relationship between FYI and blowing snow sourced sea-salt emissions from Confer et al. (2023). The positive relationship between modeled Br_y and FYI is consistent with Bougoudis et al. (2020) and suggests that recent changes in sea ice cannot explain the decline in AN ice-core Br since 1975.

Total bromine measured in the French Alps Col du Dome (CDD) ice-core showed a 2-fold increase in summertime ice-core bromine from the late-1940s to the mid-1970s, followed by a 22% decrease until the end of the record in 2000 (Legrand et al., 2021a). The observed bromine trends were previously interpreted to be driven by leaded gasoline emissions due to the strong correlation between ice-core Br and Pb. Our simulations quantitatively capture the ice-core bromine (Figure S11 in Supporting Information S1) and acidity (Figure S12 in Supporting Information S1) trends. The modeled bromine trends are caused mainly by acid-catalyzed sea-salt debromination, instead of direct emissions from leaded gasoline. Trends in leaded gasoline emissions and anthropogenic acidity are similar.

Acid-catalyzed debromination is the dominant source of Br_y and changes in acidity drive trends in Br_y in the model. The measured acidity in AN ice-core increased 5.5-fold from PI to PA, and decreased back to pre-industrial level in 1999 (Figure 1). However, neither modeled aerosol nor cloud acidity returns to PI levels in PD (Figure S7 in Supporting Information S1). Anthropogenic emissions of acid precursors NO_x and SO_2 in emission inventories declined by 20% and 44%, respectively, from 1975 to 2000 in AN TRJ (Figure S4 in Supporting Information S1), but did not drop back to PI levels. Ammonia emissions declined by 50% (Figure S4 in Supporting Information S1) during the same period, partially counteracting the decrease in SO_2 emissions on

cloud and aerosol pH. The drastic decline in ice-core acidity was not seen in Greenland (Zhai et al., 2023) or Alpine ice-cores (Legrand et al., 2003; Preunkert & Legrand, 2001). The model underestimate in bromine decline from PA to PD is likely due to the underestimated decrease in acidity. The relatively stable modeled Br concentrations from PA to PD is also due to a decline in Br_y loss rate caused by decreased atmospheric acetaldehyde.

The model's limitations also include uncertainties in halogen chemistry, particularly in the HBr/sea-salt Br⁻ partitioning treated as a kinetic reaction with large uncertainties in GEOS-Chem (Ammann et al., 2013; Wang et al., 2021). This results in a model prediction of bromine depletion for accumulation-mode aerosol in the marine boundary layer, in contrast to observed bromine enrichment (Sander et al., 2003). However, our previous work on ice-core chlorine indicates minimal changes in HCl partitioning from PA to PD (Zhai et al., 2021), suggesting similar stable thermodynamic HBr/sea-salt Br⁻ partitioning between these periods.

5. Conclusions

Tropospheric reactive bromine destroys surface ozone and has profound impacts on the tropospheric oxidation capacity (Barrie et al., 1988; von Glasow et al., 2004). Both acid-catalyzed debromination (Wang et al., 2021; Zhu et al., 2019) and sea-ice variations (Spolaor et al., 2016; Vallelonga et al., 2021) can influence ice-core bromine trends. However, the relative importance of these factors on Arctic ice-core bromine trends since the pre-industrial remains unclear. We present bromine records since the pre-industrial from the Russian Arctic AN ice-core since PI, and use GEOS-Chem model simulations to examine the trends. The AN ice core shows a 3.5-fold increase in total Br from pre-1850 to the 1970s, followed by a 53% decrease from the 1970s to 1999. Measured ice-core acidity increased by a factor of 5.5, followed by a 72% decrease after the 1970s and a return to PI levels by the end of the record. Ice-core bromine and acidity are strongly correlated, particularly after 1940 ($r = 0.9$).

The model suggests that sea-salt debromination is the dominant (>99%) source of Br_y in all emission scenarios, and direct emissions of anthropogenic bromine contribute less than 1% to the total Br_y sources. Modeled changes in anthropogenic acidity alone can explain the 64–67% increase in Br and Br_{exc} observed in the AN ice-core between PI and PD. The modeled 2-fold increase in Br_y is smaller than the observed increase in ice-core Br_{exc} (3-fold) from pre-1850 to 1975, and the modeled increase is driven by increases in anthropogenic emissions of acid gas precursors. The modeled relationship between atmospheric Br_y and acidity is consistent with observed relationships between ice-core bromine and acidity from AN (Figure 2) and a mid-latitude alpine ice-core (Legrand et al., 2021a).

The model does not capture the observed decrease in both Br_y (53%) and acidity (72%) after 1975, possibly because of uncertainties in anthropogenic emissions and transport of acidic gas-phase precursors. Since variability in atmospheric Br_y is driven by atmospheric acidity, the modeled underestimate of the decline in acidity since 1975 contributes to the modeled underestimate of the decline in Br_{exc}. Our results suggest that anthropogenic acidity has had a profound influence on the production and abundance of tropospheric reactive bromine (Br_y), with implications for the oxidation capacity of the atmosphere. The use of ice-core bromine as a proxy for past changes in sea-ice extent must consider changes in atmospheric acidity in the interpretation of observed trends, especially when ice-core acidity changes.

Data Availability Statement

Ice-core data for this research can be downloaded from the Arctic Data Center with Creative Commons Attribution (McConnell, 2023). GEOS-Chem is open software and available for download (Yantosca et al., 2021). GEOS-Chem historical simulations output is archived in the Dryad Data Repository (Zhai & Alexander, 2023).

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