Atmos. Chem. Phys. Discuss., 7, 11647–11683, 2007 www.atmos-chem-phys-discuss.net/7/11647/2007/
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Volatile organic compound ratios as probes of halogen atom chemistry in the Arctic

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Received: 3 July 2007 - Accepted: 31 July 2007 - Published: 8 August 2007

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Abstract

Volatile organic compound concentration ratios can be used as indicators of halogen chemistry that occurs during ozone depletion events in the Arctic during spring. Here we use a combination of modeling and measurements of [acetone]/[propanal] as an indicator of bromine chemistry, and [isobutane]/[n-butane] and [methyl ethyl ketone]/[nbutane] are used to study the extent of chlorine chemistry during four ozone depletion events during the Polar Sunrise Experiment of 1995. Using a 0-D photochemistry model in which the input of halogen atoms is controlled and varied, the approximate ratio of [Br]/[CI] can be estimated for each ozone depletion event. It is concluded that there must be an additional source of propanal (likely from the snowpack) to correctly simulate the VOC chemistry of the Arctic, and that the ratio of Br atoms to Cl atoms can vary greatly during ozone depletion events.

Introduction

It is well known that in Arctic regions periods of ozone depletion occur during the late winter and spring, corresponding with Polar Sunrise (Bottenheim et al., 1986, 2002; Oltmans and Komhyr, 1986). This sudden and frequent loss of ozone is highly correlated with elevated concentrations of halogen-containing radicals, the source of which appears to be sea salt aerosols from the Arctic Ocean (Barrie et al., 1988; Simpson et al., 2005), and/or the salt-laden snowpack. While it is widely accepted that bromine chemistry in the gas and aerosol phase creates a catalytic cycle that destroys ozone (Barrie et al., 1988; Fan and Jacob, 1992; Hausmann and Platt, 1994; Vogt et al., 1996), the role of chlorine atoms in ozone destruction, and as important oxidizers, in the Arctic is not as well understood (Jobson et al., 1994; Yokouchi et al., 1994; Shepson et al., 1996; Impey et al., 1997; Tuckermann et al., 1997; Keil and Shepson, 2006). This is mostly due to the analytical difficulty in directly measuring the highly reactive CI, which is thought to exist in very small but kinetically important quantities in the Arc-

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tic boundary layer (on the order of 2×10⁴ molec/cm³ during ozone depletion events) (Jobson et al., 1994; Ramacher et al., 1999; Boudries and Bottenheim, 2000; Keil and Shepson, 2006). Chlorine atom concentrations can be calculated based on decreases in measured alkanes, and relationships between volatile organic compounds (VOCs) can be utilized to further understand the role of Br and Cl (Jobson et al., 1994; Sander et al., 1997; Boudries et al., 2002; Evans et al., 2003).

By examining the relationships between alkanes and their oxidation products, the extent of reaction with various oxidizers in a given parcel of air can be studied. At coastal sites these oxidizers can include OH, Br and Cl radicals (Pszenny et al., 1993; Spicer et al., 1998). An example of this is the oxidation of n-butane to produce a peroxy radical:

$$CH_3CH_2CH_2CH_3 + OH \cdot (or Cl \cdot) + O_2 \rightarrow CH_3CH_2CH(OO \cdot)CH_3 + H_2O (or HCl)$$
 (R1)

This peroxy radical can then go on to react with NO, HO_2 or other peroxy radicals to form methyl ethyl ketone (MEK) and other products:

$$5 \quad \text{CH}_3\text{CH}_2\text{CH}(\text{OO})\text{CH}_3 + \text{NO} \rightarrow \text{CH}_3\text{CH}_2\text{CH}(\text{O})\text{CH}_3 + \text{NO}_2$$
 (R2)

$$CH_3CH_2CH(OO\cdot)CH_3 + HO_2 \rightarrow CH_3CH_2CH(OOH)CH_3 + O_2$$
(R3)

$$CH_3CH_2CH(OO)CH_3 + RO_2 \rightarrow CH_3CH_2C(O)CH_3 + ROH + O_2$$
 (R4)

$$CH_3CH_2CH(OO)CH_3 + RO_2 \rightarrow CH_3CH_2CH(OH)CH_3 + RC(O)R' + O_2$$
(R5)

$$CH_3CH_2CH(OO\cdot)CH_3 + RO_2 \rightarrow CH_3CH_2CH(O\cdot)CH_3 + RO \cdot +O_2$$
(R6)

$$CH_3CH_2CH(O\cdot)CH_3 + O_2 \rightarrow CH_3CH_2C(O)CH_3 + HO_2$$
(R7)

For [CI]/[OH]>0.01, the rate of butane oxidation in Reaction (R1) will be dominated by CI atoms. The rate constants at 245 K (typical of Arctic regions during spring) for the reactions of n-butane and MEK with OH radicals are very

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similar, differing by a factor of 1.4 $(k_{n-butane+OH}=1.64\times10^{-12} \text{ cm}^3/\text{molec s})$ ahue et al., 1998), $k_{MEK+OH}=1.17\times10^{-12} \text{ cm}^3/\text{molec s}$ (Atkinson et al., 2001)), while the rate constants for Cl atom reaction with these species at 245K differ by a factor of 7.8 $(k_{n-butane+Cl}=2.15\times10^{-10} \text{ cm}^3/\text{molecs}$ (Tyndall et al., 1997), $k_{MEK+Cl}=2.75\times10^{-11}\,cm^3/molecs$ (Cuevas et al., 2004)). Due to this, the ratio [MEK]/[n-butane] can be indicative of oxidation by chlorine atoms in the atmosphere under certain conditions of the [CI]/[OH] ratio. If oxidation occurs solely by OH radicals, [MEK]/[n-butane] is expected to change slowly, as the rates of production and destruction can become comparable relatively quickly. However, if chlorine atoms are responsible for the oxidation process, n-butane oxidation to produce MEK outpaces MEK destruction, leading to enhancement in the [MEK]/[n-butane] ratio. Under ozone depletion conditions, when [CI] can be >1×10⁴ molec/cm³, rapid butane oxidation by CI atoms can occur. Bromine atoms are not believed to affect the [MEK]/[n-butane] ratio, because they do not react with small ketones, and only react very slowly with alkanes $(k_{\text{nbutane+Br}} = 2.2 \times 10^{-18} \text{ cm}^3/\text{molec s at } 298 \text{ K (Galiba et al., } 1964)).$

The ratio [isobutane]/[n-butane] is unaffected by OH radical chemistry but is affected by CI atom chemistry, and hence it can also be used as an indicator of chlorine atom chemistry. The rate constants for OH reaction with isobutane and n-butane at 245 K are essentially identical (they differ by a factor of 1.006), at 1.65×10^{-12} and 1.64×10⁻¹² cm³/molecs, respectively (Donahue et al., 1998). However, for Cl reaction with isobutane and n-butane the rate constants at 245 K differ by a factor of 1.65, with values of 1.30×10⁻¹⁰ and 2.15×10⁻¹⁰ cm³/molecs, respectively (Hooshiyar and Niki, 1995; Tyndall et al., 1997). The ratio [isobutane]/[n-butane] is observed to be approximately 0.5 at the point of emission by fossil fuel combustion (Penkett et al., 1993; Jobson et al., 1994; Greenberg et al., 1996; Solberg et al., 1996). However, it has been observed that [isobutane]/[n-butane] is enhanced during ozone depletion events (Jobson et al., 1994; Solberg et al., 1996), which has been ascribed to rapid Cl-atom processing of n-butane. By examining both [MEK]/[n-butane] and [isobutane]/[n-butane], the amount of processing that occurs by both CI and OH radicals can be examined.

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A similar analysis can be used to elucidate the extent of bromine atom chemistry, using the ratio of two different products that result from the oxidation of propane. Similar to n-butane, propane is oxidized to produce peroxy radicals:

$$CH_3CH_2CH_3 + OH \cdot (or Cl \cdot) + O_2 \rightarrow CH_3CH(OO \cdot)CH_3 + H_2O (or HCl)$$
 (R8)

Each of these radicals then proceeds to react with NO, HO₂ or another peroxy radical, similar to Reactions (R2) through (R7). The 2-propyl peroxy radical from Reaction (R8) eventually forms acetone, while the 1-propyl peroxy radical from R9 eventually forms propanal. Using concentrations of 1×10⁸ molec/cm³ for NO, HO₂ and RO₂, the relatively clean Arctic air yields of propanal and acetone are 11% and 14%, respectively, when propane is oxidized by CI, and 6% and 19% when propane is oxidized by OH. It has been shown that bromine atoms react readily with aldehydes via abstraction of the aldehydic H-atom, but not with ketones. The rate constant for reaction of bromine atoms with propanal at 245 K is $9.73 \times 10^{-12} \, \text{cm}^3/\text{molecs}$ (Wallington et al., 1989). Thus when bromine atoms are present the ratio [acetone]/[propanal] will be enhanced due to the preferential removal of propanal. There are some complications in this analysis, in that both acetone and propanal do react with both OH and CI radicals $(k_{acetone+OH@245K}=1.30\times10^{-13} \text{ cm}^3/\text{molec s},$ $k_{\text{acetone+Cl@298\,K}} = 3.50 \times 10^{-12} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \, \text{cm}^3 / \text{molec s}, \, k_{\text{propanal+OH@245\,K}} = 2.61 \times 10^{-11} \,$ $k_{propanal+Cl@298K} = 1.20 \times 10^{-10} \text{ cm}^3/\text{molec s}$) (Wollenhaupt et al., 2000; Atkinson et al., 2001). However, since [Br] is often quite large (>1×10⁷ molec/cm³), relative to [CI] and [OH] in the Arctic springtime boundary layer, and given the rate constant for bromine atom reaction with propanal, the ratio is substantially impacted by [Br]. The contribution of each of these species to [acetone]/[propanal] will be evaluated in the discussion section of this paper.

In this work ambient data that were collected between 17 March and 24 April 1995 at Alert, Nunavut (82.5° N, 62.3° W) are presented and the ratios of the measured VOCs

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are analyzed to further understand the impact and variability of the halogen atoms present during ozone depletion events. Additionally, a 0-D model has been constructed to support the analysis of VOC ratios to evaluate the relative importance of CI, Br and OH radicals.

5 2 Experimental

2.1 Ambient measurements from Polar Sunrise Experiment (PSE), Alert, 1995

Ambient air was scrubbed of ozone by passing it through a potassium iodide trap, and then VOCs were trapped by pumping it through an adsorbent trap consisting of Tenax TA (35/60 mesh, 160 mg, Supelco, Pennsylvania) followed by Carboxen-569 (110 mg, Supelco, Pennsylvania). Typical sample sizes were 1.3 L. This trap was cooled to 5°–10°C while sampling utilizing cold ambient air, and then desorbed under helium flow at 250°C. The sample components were injected into a gas chromatograph (Hewlett Packard 5890) containing a PoraPLOT Q column (10 m×0.32 mm, Varian, California) that was held at 35°C. Once desorption from the trap was complete the GC oven was heated rapidly to 100°C, then increased by 5°C/min to 200°C, and finally ramped quickly to 240°C. A quadrupole mass spectrometer (Hewlett Packard 5972) was used for detection, scanning a mass range of 27–200 a.m.u. at a rate of 24 scans per second. Calibrations were conducted twice per day throughout the study, using a permeation source of acetone. Overall measurement uncertainties for acetone, MEK, n-butane, propanal and isobutane were estimated to be ±10%, 15%, 20%, 30% and 35%, respectively.

The nearby Global Atmospheric Watch (GAW) station, operated by Environment Canada, provided all ozone data for this experiment.

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2.2 0-D model

A 0-D model was created, incorporating much of the known gas-phase chemistry of VOCs and halogens that occurs in the Arctic atmosphere. The VOC chemistry in the model focused primarily on oxidation chemistry of n-butane, isobutane, propane, and their oxidation products. The halogen atom concentrations were varied within the model by creating inputs of Br₂, BrCl and Cl₂, to specifically examine how halogens affect the [MEK]/[n-butane], [isobutane]/[n-butane] and [acetone]/[propanal] ratios, and to use the measured VOC ratios to estimate the relative importance of [Br] and [Cl]. Table 1 shows the 122 gas-phase reactions that were included in the model, as well as their rate constants. When possible, rate constants were calculated for a temperature of 245 K, as this was a representative ambient temperature during the field study.

Table 2 shows the various species that had varying input and/or removal fluxes in the model. These fluxes were represented by using a rate constant for a reaction that either produces the species from a dummy source or removes the species via a first order loss process. Removal rates for species that undergo dry deposition were all set at an equivalent first order rate constant of $1.25 \times 10^{-5} \, \text{s}^{-1}$, corresponding to a dry deposition velocity, V_d , of $0.5 \, \text{cm/s}$ and a boundary layer height of 400 m, based on aircraft observations in the Arctic (Leaitch et al., 1994). The effective deposition rate constants (expressed as negative values) are shown in Table 2.

Surface source fluxes of $\dot{H}ONO$, NO_x and acetone have been measured, and they are believed to be produced photochemically. A zero-order rate constant for each flux was obtained by dividing the measured flux value by the assumed height of the boundary layer (400 m). This rate constant was then scaled by the solar zenith angle calculated within the model. These fluxes were initially entered into the model at the maximum values that have been previously measured for local solar noon (Zhou et al., 2001; Beine et al., 2002; Guimbaud et al., 2002). Some minor adjustment of these values was necessary to obtain concentrations of HONO, NO_x and acetone that corresponded with observations. Although a propanal flux has not been measured in

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the Arctic to date, this flux was found to be necessary to correctly simulate the observed propanal concentrations, as discussed below. In addition, OH values were scaled by the solar zenith angle so that the daytime average OH concentration for each model day was 5×10^5 molec/cm³.

Twenty six photolysis reactions were included in this model, as listed in Table 3. The majority of the photolysis rates were determined using the Tropospheric Ultraviolet-Visible Model (TUV) (Madronich and Flocke, 1998) for 29 March 1995 at Alert. This model was not able to calculate photolysis rates for MEK, BrCl and HOCl. The MEK photolysis rate coefficient was calculated as described in Jenkin et al. (1997), and BrCl and HOCl photolysis rate coefficients were calculated using the following equation:

$$J = \int_{\lambda} \Phi(\lambda) \sigma(\lambda) I_{\lambda} d\lambda \tag{1}$$

where Φ is the quantum yield (assumed to be unity), σ is the absorption cross-section (DeMore et al., 1997) and I is the actinic flux (values are from the TUV model). Photolysis rates within the model varied with the solar zenith angle, as calculated for conditions at Alert on 29 March 1995. The noontime minimum solar zenith angle was 79°.

Table 4 shows the initial concentrations of the species in the model that began with a concentration other than zero. The concentrations of the species in bold were held constant throughout the simulations. The value for water represents 80% relative humidity at the model temperature, the value for ozone is equal to the average ozone concentration at Alert during ambient sampling, and many of the other values are either based on measurements or other models (Michalowski et al., 2000; Grannas et al., 2002).

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3 Results and discussion

3.1 Ambient measurements

Figure 1 shows the measured ambient concentrations of MEK, isobutane, n-butane, propanal, acetone and ozone during the field campaign. In this paper, we focus on four case study ozone depletion events from PSE95, indicated in the figure with the numbers 1 through 4. These events were chosen for further analysis because all of the VOCs of interest were present above their respective detection limits throughout each event. It is clear from this figure that isobutane and n-butane are typically well correlated, which is consistent with their common petroleum fuel source and common radical reaction sinks. Figure 1 also shows a very significant negative correlation between ozone and the VOC oxidation products acetone and MEK. Figure 2 shows the results of the oxidation chemistry more clearly, in which the ratios [isobutane]/[n-butane], [MEK]/[n-butane] and [acetone]/[propanal] are plotted, along with ambient ozone concentrations. When viewed in this manner it can be seen that all three of the VOC ratios appear to be inversely correlated with ozone. As discussed above, this can only imply halogen chemistry, which not only causes ozone depletion, but also has a dramatic effect on the ratios of various VOCs that are present in the Arctic troposphere, and in ways that depend on the ratio [Br]/[CI], as will be shown.

In Fig. 3 the VOC ratios [MEK]/[n-butane] and [isobutane]/[n-butane] are plotted against [acetone]/[propanal] for the four different ozone depletion events of interest. As discussed in the introduction, the x-axes of this figure are indicative of chlorine atom chemistry, while the y-axis is sensitive to bromine atom chemistry. The fact that Event 4 is enhanced in both [MEK]/[n-butane] and [isobutane]/[n-butane] indicates that there is relatively much more chlorine atom chemistry during that event, while Events 1 through 3 all appear to be dominated by bromine chemistry. The data in this figure imply that ozone depletion can occur with highly varying concentrations of Br and Cl atoms, or the ratio [Br]/[CI].

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3.2 Evidence of chlorine atom chemistry

The consecutive reaction scheme that is represented by Reactions (R1) through (R5), and the subsequent removal of MEK by reaction with OH or CI, can be solved mathematically to calculate the impact of oxidizer concentration on the enhancement of [MEK]/[n-butane], using the equation below for consecutive reactions (Apel et al., 2002):

$$\frac{[MEK]}{[n-butane]} = \frac{\alpha \cdot k_1 (1 - e^{(k_1 - k_2)[X]t})}{k_2 - k_1}$$
 (2)

where α is the yield of MEK from n-butane oxidation (α =0.83 when X=OH, and α =0.66 when X=Cl), k_1 is the rate constant for reaction of n-butane with X, k_2 is the rate constant for reaction of MEK with X, and [X] is the average concentration of either OH or Cl over the reaction time, t, in seconds. Using a reaction time of five to ten days (Jobson et al., 1994), [MEK]/[n-butane] can be calculated, and the resulting values range from 0.28 to 0.58 when X is OH ([X]=5×10⁵ molec/cm³), and from 1.57 to 6.23 when X is Cl ([X]=2×10⁴ molec/cm³). When both oxidants are present at the concentrations listed above (i.e. typical concentrations for ozone depletion conditions, Jobson et al., 1994; Boudries and Bottenheim, 2000; Grannas et al., 2002; Ridley et al., 2003), chlorine atoms are responsible for more than 85% of the change in the MEK/n-butane ratio.

3.3 Evidence of bromine atom chemistry

As discussed above, the ratio [acetone]/[propanal] is sensitive to [Br]. However, the dependence is complex. Bromine atom reaction with propane is endothermic by ~46 kJ/mole (Tsang, 1996), so only chlorine and OH radicals can complete the first step (Reaction R10) of the consecutive reaction mechanism shown below. Additionally, only chlorine and OH will oxidize acetone in Reaction (R11). However, in Reaction (R12), propanal can be oxidized by OH, Br and CI.

$$C_3H_8 + (OH \cdot and/or CI \cdot) \rightarrow acetone + propanal$$
 (R10)
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Acetone +
$$(OH \cdot and/or CI) \rightarrow products$$
 (R11)

Propanal + (OH · and/or CI · and/or Br·)
$$\rightarrow$$
 products (R12)

Consider the case where all three oxidizers are present in the atmosphere. By multiplying the rate constants for each reaction (which can be found in Table 1) by typical concentrations of the oxidants during halogen atom chemistry events $([OH]=5\times10^5 \text{ molec/cm}^3, [CI]=2\times10^4 \text{ molec/cm}^3 \text{ and } [Br]=1\times10^7 \text{ molec/cm}^3 \text{ (Jobson)}$ et al., 1994)) the contribution of each oxidizer to the removal rate of acetone and propanal can be calculated. For this condition, 86% of the propanal is destroyed by bromine, while 12% is destroyed by OH and only 2% by Cl. 52% of the acetone destruction is caused by CI, and 48% by OH. However, the removal of acetone by CI and OH occurs much more slowly ($\tau_{acetone} \sim 86$ days) than that of propanal ($\tau_{propanal} \sim 2.5$ h), and indeed is not important on the timescale of the events discussed here. Hence the variation in the ratio [acetone]/[propanal] is mostly determined by Br under these conditions. It is important to note that because there exists a large initial concentration of acetone in the atmosphere before ozone depletion events, acetone production during ozone depletion contributes relatively little to changes in [acetone]/[propanal]. Thus during ozone depletion events propanal destruction will be enhanced and [acetone]/[propanal] will reflect bromine atom chemistry.

Model simulations

Figure 4 shows the results of the model described in Tables 1-4, with no surface flux of propanal present. All concentrations reported from the model represent the average concentration of a given species during the sunlit hours (06:00-20:00 local time). The model was allowed to run for a period of 10 days (Jobson et al., 1994) with various halogen atom concentrations, ranging from no halogens present (all oxidation caused by OH) to a maximum condition of [Br]=4×10⁷ and [Cl]=2×10⁴ molec/cm³ ([Br]/[Cl]=2000). In every case except the "OH only" case, the flux of Cl₂ and/or BrCl

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was adjusted to achieve a Day 9 daytime average [CI] of 2×10⁴ molec/cm³, and the Br₂ flux was varied in order to achieve the [Br]/[CI] ratio desired. The range of [Br]/[CI] examined is representative of the various halogen atom ratio estimates that have been made in the Arctic during ozone depletion events (Jobson et al., 1994; Ramacher et al., 1999; Keil and Shepson, 2006). The solid lines are created from daytime averages of the various species. The individual points on the plot are the same ambient data from PSE95 that are shown in Fig. 3.

While the ambient data and the model results seem to be in the same range when considering [MEK]/[butane] and [isobutane]/[n-butane], e.g. for the case with chlorine atom chemistry only ([Br]/[Cl]=0), the model overestimates [acetone]/[propanal] by orders of magnitude. This is likely due to a missing source of propanal, potentially from the snowpack. Because acetone is so long lived, and the model simulation starts with 600 ppt of acetone, the model does not overestimate [acetone], nor is this ratio sensitive to acetone chemistry. Formaldehyde, acetaldehyde and acetone fluxes have all been measured from the snowpack (Guimbaud et al., 2002; Sumner et al., 2002; Grannas et al., 2004) so it is reasonable to assume that there could be a flux of propanal from the snowpack as well.

Figure 5 shows the results of the model, along with the PSE95 ambient data from Fig. 3, when a radiation-dependent flux of propanal has been added to the model. The magnitude of this flux was adjusted so that the Day 9 average value of [acetone]/[propanal] for the case where [Br]/[Cl]=500 was approximately 80. This flux was then included in the model for all of the other cases shown in the figure, set proportional to calculated radiation. Including this flux greatly increases the agreement between the model and the ambient data, as shown in comparing Figs. 4 and 5. If the height of the Arctic boundary layer is assumed to be 400 m, as discussed earlier, the daytime maximum flux of propanal that is necessary to correctly simulate the VOC ratios measured is calculated to be 2.4×10^9 molec/cm² s. The magnitude of this flux is on the high end compared to the magnitude of measured fluxes for the other oxygenated VOC fluxes that have been measured in the Arctic, namely

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formaldehyde $(4.90\times10^9\,\text{molec/cm}^2\,\text{s})$ (Sumner and Shepson, 1999), acetaldehyde $(4.20\times10^8\,\text{molec/cm}^2\,\text{s})$ and acetone $(6.20\times10^8\,\text{molec/cm}^2\,\text{s})$ (Guimbaud et al., 2002). This likely results from the fact that a volumetric flux corresponding to a 400 m deep boundary layer is being used in this calculation. Given the reactivity of propanal, it likely has a very steep vertical concentration gradient. From the work of Guimbaud et al. (2002), we can calculate the effective mixing height for propanal, given its lifetime. For [Br]/[CI]=500, i.e. [CI]= 2×10^4 , [Br]= 1×10^7 and [OH]= 5×10^5 molec/cm³, the propanal lifetime τ is 2.5 h. The effective mixing height, Z (using K_Z (eddy diffusivity) = $95\,\text{cm}^2/\text{s}$ (Guimbaud et al., 2002)) is then

$$Z_{\text{propanal}} = \sqrt{K_Z \tau} = 9.2 \,\text{m}$$
 (3)

Thus, at solar noon, the appropriate maximum propanal flux (using this value as the effective mixing height), for the purpose of comparing to previously measured carbonyl compound fluxes, is on the order of 5.4×10^7 molec/cm² s. This value is more reasonable, given the relative observed concentrations of acetaldehyde and propanal at Alert.

It can be seen from Fig. 5 that [Br]/[Cl] can vary greatly during ozone depletion events, from ~100 to ~2000. This range of [Br]/[Cl] is consistent with the previous work of Jobson et al. (1994), Impey et al. (1997), Ramacher et al. (1999) and Keil and Shepson (2006). While Events 1–3 appear to be relatively dominated by bromine chemistry, Event 4 is clearly quite different, exhibiting much more chlorine atom chemistry. Comparing the initial slopes of the model output with the data, the ambient [Br]/[Cl] ratios seem to be on the order of 500–800, 800–2000, 800–2000, and 100–200 for Events 1–4, respectively. Evans et al. (2003) discuss that in their model chlorine becomes less active as an oxidant as ozone becomes more depleted, and that chlorine chemistry becomes totally inactive when ozone drops below 10 ppb. This occurs because as the "bromine explosion" chemistry proceeds, aerosol and snowpack salt crystal surfaces become enriched in Br⁻, effectively slowing the relative rate of oxidation of Cl⁻ by HOBr. This is consistent with the measurements of Simpson et al. (2005), who find the snowpack to be relatively enriched in Br⁻ as one moves inland away from the sea

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salt aerosol source, due to the longer lifetime of the fine aerosol Br (and brominecontaining gases such as HBr and HOBr). In other words, deposition of the products of halogen-chemistry processed air tends to enrich the surface in Br⁻. The work of Toom-Sauntry and Barrie (2002) shows clearly that as the spring season progresses, the snowfall at Alert becomes progressively more enriched in Br relative to Cl. Indeed, by May, bromide enrichment factors (relative to fresh sea salt) were as high as ~70. It was also postulated that this results in part from uptake of gas phase bromine species, such as HBr and HOBr. In addition, the reaction of the excess Br with any BrCl that is produced yields Br₂Cl⁻, which can release Cl⁻ to produce gas phase Br₂, inhibiting the release of BrCl into the atmosphere (this assumes that BrCl photolysis is the primary source of Cl atoms). Impey et al. (1997, 1999) observed that photolyzable chlorine became relatively unimportant later in the spring, and under conditions when ozone is completely depleted. This is consistent with the snowfall and snowpack surfaces becoming enriched in Br as spring progresses. While it might be supposed that BrO (and thus HOBr) is required for BrCl production, for Event 4 ozone decreases to ~3 ppb, comparable to that for Event 2, for which we estimate a [Br]/[CI] ratio that is ~10–20 times larger. There are likely several explanations for this variability of [Br]/[Cl]. One plausible explanation is that production of BrCl can occur after ozone is mostly depleted, since it is hypothesized that BrCl is produced from HOBr deposition to the salt-laden snowpack, followed by reaction with Cl⁻ (Adams et al., 2002). If HOBr were mixed throughout the lowest 100 m of the boundary layer, and given a deposition velocity of $0.5 \,\mathrm{cm/s}$, the lifetime of HOBr (H/V_d), and thus the time over which BrCl would still be produced, would be ~6 h. Thus, for a short-term ozone depletion event there could be significant levels of gas phase chlorine atom chemistry, even with ozone concentrations of zero. The relative importance of chlorine and bromine chemistry is more likely a function of the relative concentrations of the two halide ions on the reactive surfaces.

Since Event 4 is different from the other events examined here, it is important to understand what conditions may have led to the exhibited chemistry in that case. It is clear that a potential source of [Br]/[CI] variability is the air mass origin during the event.

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In Fig. 6, 72-h back trajectories are shown for the four ozone depletion events. These trajectories were produced using the HYSPLIT model (http://www.arl.noaa.gov/ready). The air masses that contributed to Events 1 and 2 both arrived having spent most of the previous 72 h over Greenland, while Event 3 began over the ice and then passed over Ellesmere Island before reaching the sampling site. The back trajectory for Event 4, however, is different from the others in that the air mass spent most of the time over the ice pack in the Arctic Ocean, with a relatively longer trajectory. Assuming that the halogens are activated by reaction of HOBr with Br and Cl, the relative concentrations of Br₂ and BrCl (and thus [Br] and [Cl]) will depend on the relative concentrations of Br and Cl in the snowpack, along with other factors such as the relative Henry's Law constants. As discussed by Simpson et al. (2005), there is substantial variation of snowpack [Br⁻]/[Cl⁻], and this tends to increase with distance from the coast. This reflects the idea that with Arctic Ocean trajectories, fresh sea salt aerosol deposits relatively quickly leading to low ratios of [Br-1/[Cl-]] near the coast. It is thus plausible that the conditions leading to Event 4 led to fresh inputs (from air passing over first year sea ice, or polynyas), and an increased probability of activation of Cl⁻. In contrast, air masses that came to the sampling site over areas that are predominantly land (Events 1-3) show evidence of enhanced [Br]/[CI], due to activation within Br enriched snowpacks. Thus, it would seem that the relative activation rates of chlorine and bromine reflect the chemical "age" of the snowpack, and the extent to which there have been fresh inputs of unprocessed sea salt.

4 Conclusions

From the ambient data presented here, and from the model results that are discussed, there are a few conclusions that can be drawn. First, there is reason to believe that there may be a source of propanal that has not yet been measured in the Arctic. This additional source is necessary to correctly simulate the VOC chemistry that occurs during Polar Spring, and it is reasonable to postulate that the missing propanal source

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is oxidation of organic matter in the snowpack (Grannas et al., 2004), given measurements of snowpack photochemical sources of other oxygenated VOCs. It is unlikely that it is an unknown atmospheric source, as this would probably require decomposition of a large alkoxy radical to produce the propyl radical, but such a precursor would be short lived, making substantial concentrations unlikely. Ambient measurements of this propanal flux are needed to verify this conclusion. Second, the ratio of bromine to chlorine atoms appears to be highly variable during ozone depletion events, as seen from the variability in ratios of the alkanes and their oxidation products that are affected by these halogens. Much of this variability for coastal sites may derive from the air mass origin, and the extent of halide processing in the associated snowpacks along the air mass trajectories. Further understanding of the behavior of halogen atoms in the Arctic is very important, as they are thought to be responsible for the springtime ozone and mercury depletion events that have been observed (Schroeder et al., 1998), but the activation mechanisms are not fully understood. For example, Knipping and Dabdub (2002) discuss that Cl⁻ may be activated by OH radicals in saline snowpacks. It is well known that H₂O₂ is an important photolytic precursor for condensed phase OH radicals (Anastasio and Jordan, 2004). Cl₂ can then be produced from the following reactions:

$$OH + CI^{-} \rightarrow CIOH^{-}$$
 (R13)

$$2\text{CIOH}^- \rightarrow \text{Cl}_2 + 2\text{OH}^- \tag{R14}$$

Thus, the high apparent [CI] during Event 4 could be related to poorly understood snowpack photochemical activation of CI⁻. It is clear that further laboratory studies should be conducted to examine the extent to which halogen activation can occur via OH radical chemistry in the condensed phase. This may help explain the existence of low [Br]/[CI] ratios when ozone is low.

Acknowledgements. The authors would like to acknowledge the Jonathan Amy Facility for Chemical Instrumentation at Purdue University for instrumentation support and K. Anlauf for the use of AES/GAW ozone data for PSE95. This work was funded by the National Science

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Foundation, grant no. OPP-0325361. This is publication No. 0621 of the Purdue Climate Change Research Center.

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Table 1. 0-D Model mechanism.

Rxn. #	Reaction	T(K)	Rate constant	Ref.
	First Order Reactions:			
1	PAN→PA+NO ₂	245	4.08×10 ⁻⁸	а
2	$HNO_4 \rightarrow HO_2 + NO_2$	245	1.63×10 ⁻⁴	а
3	$N_2O_5 \rightarrow NO_2 + NO_3$	245	2.67×10 ⁻⁵	
	Second Order Reactions:			
4	OH+CO→HO ₂ +CO ₂	245	2.40×10 ⁻¹³	b
5	$OH+CH_4\rightarrow CH_3O_2$	245	1.47×10 ⁻¹⁵	b
6	$OH+C_2H_6\rightarrow C_2H_5O_2$	245	9.77×10^{-14}	
7	$CI+CH_4\rightarrow HCI+CH_3O_2$	245	3.58×10^{-14}	
8	$CI+C_2H_6\rightarrow HCI+C_2H_5O_2$	245	6.55×10 ⁻¹¹	
9	$CI+C_3H_8\rightarrow HCI+n-C_3H_7O_2$	245	6.02×10^{-11}	С
10	$CI+C_3H_8\rightarrow HCI+i-C_3H_7O_2$	245	7.98×10 ⁻¹¹	С
11	CI+HCHO→HCI+HO ₂ +CO	245	5.80×10^{-11}	
12	CI+CH ₃ CHO→HCI+PA	245	7.20×10^{-11}	
13	CI+propanal→HCI	298	1.20×10 ⁻¹⁰	
14	Cl+acetone→CH ₃ O ₂	298	3.50×10^{-12}	
15	CI+PAN→HCI+HCHO+NO ₃	298	1.00×10^{-14}	
16	$NO+O_3 \rightarrow NO_2+O_2$	245	5.70×10^{-14}	
17	$NO_2 + O_3 \rightarrow NO_3 + O_2$	245	5.15×10 ⁻¹⁸	
18	$NO_2 + NO_3 \rightarrow N_2O_5$	245	1.72×10^{-12}	b
19	$O(^{1}D) + O_{2} \rightarrow O_{3}$	245	2.81×10^{-12}	d
20	PA+NO ₂ →PAN	245	1.25×10 ⁻¹¹	b
21	$PA+NO \rightarrow NO_2+CH_3O_2$	245	2.31×10^{-11}	b

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Table 1. Continued.

Rxn. #	Reaction	T (K)	Rate constant	Ref.
22	$HO_2+NO\rightarrow NO_2+OH$	245	1.08×10 ⁻¹¹	
23	$CH_3O_2+NO\rightarrow NO_2+HO_2+HCHO$	245	8.96×10^{-12}	b
24	$C_2H_5O_2+NO\rightarrow NO_2+CH_3CHO+HO_2$	245	1.16×10 ⁻¹¹	
25	$n-C_3H_7O_2+NO\rightarrow NO_2+propanal$	245	1.21×10 ⁻¹¹	
26	i-C ₃ H ₇ O ₂ +NO→NO ₂ +acetone	245	1.17×10 ⁻¹¹	b
27	$HO_2 + NO_2 \rightarrow HNO_4$	245	6.18×10^{-12}	b
28	$OH+NO_2\rightarrow HNO_3$	245	2.23×10^{-11}	
29	$Br+O_3 \rightarrow BrO+O_2$	245	6.63×10^{-13}	
30	Br+HCHO→HBr+HO ₂ +CO	245	6.63×10^{-13}	
31	Br+CH ₃ CHO→HBr+PA	245	2.99×10^{-12}	
32	Br+propanal→HBr	245	9.73×10^{-12}	е
33	$2HO_2 \rightarrow H_2O_2$	245	2.50×10^{-12}	
34	$CH_3O_2+HO_2\rightarrow CH_3OOH$	245	8.99×10^{-12}	
35	$C_2H_5O_2+HO_2\rightarrow C_2H_5OOH$	245	1.47×10 ⁻¹¹	
36	$CH_3O_2 + CH_3O_2 \rightarrow 2HO_2 + 2HCHO$	245	4.44×10^{-13}	
37	$C_2H_5O_2+CH_3O_2\rightarrow CH_3CHO+HCHO+2HO_2$	298	2.01×10^{-13}	f
38	$C_2H_5O_2+C_2H_5O_2 \rightarrow 2CH_3CHO+2HO_2$	245	6.40×10^{-14}	
39	PA+HO ₂ →PAA	245	2.78×10^{-11}	
40	$CH_3O_2+PA\rightarrow HCHO+CH_3O_2+HO_2$	245	1.51×10 ⁻¹¹	
41	$C_2H_5O_2+PA\rightarrow CH_3CHO+CH_3O_2+HO_2$	245	3.40×10^{-11}	
42	$HO_2+n-C_3H_7O_2 \rightarrow product$	245	3.50×10^{-11}	g
43	$HO_2 + i - C_3 H_7 O_2 \rightarrow product$	245	3.50×10^{-11}	g
44	$CH_3O_2+n-C_3H_7O_2 \rightarrow HCHO+propanal+2HO_2$	245	6.70×10^{-13}	а

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Table 1. Continued.

Rxn. #	Reaction	T (K)	Rate constant	Ref.
45	$CH_3O_2+n-C_3H_7O_2 \rightarrow propanal+CH_3OH+O_2$	245	2.30×10 ⁻¹³	а
46	$CH_3O_2+n-C_3H_7O_2 \rightarrow HCHO+n-C_3H_7OH+O_2$	245	2.30×10^{-13}	a
47	$CH_3O_2+i-C_3H_7O_2\rightarrow HCHO+acetone+2HO_2$	245	1.20×10 ⁻¹⁴	а
48	$CH_3O_2+i-C_3H_7O_2 \rightarrow acetone+CH_3OH+O_2$	245	4.10×10^{-15}	а
49	$CH_3O_2+i-C_3H_7O_2\rightarrow HCHO+i-C_3H_7OH+O_2$	245	4.10×10^{-15}	a
50	n-butane+OH→s-C ₄ H ₉ O ₂	245	1.46×10 ⁻¹²	h
51	n-butane+OH→n-C ₄ H ₉ O ₂	245	1.80×10 ⁻¹³	h
52	n-butane+Cl→s-C ₄ H ₉ O ₂	245	1.53×10 ⁻¹⁰	С
53	n-butane+Cl→n-C ₄ H ₉ O ₂	245	6.24×10^{-11}	С
54	$s-C_4H_9O_2+CH_3O_2 \rightarrow MEK+HCHO+2HO_2$	245	1.20×10 ⁻¹⁴	*(47)
55	$s-C_4H_9O_2+CH_3O_2 \rightarrow MEK+CH_3OH$	245	4.10×10^{-15}	*(48)
56	$s-C_4H_9O_2+CH_3O_2\rightarrow s-C_4H_9OH+HCHO$	245	4.10×10^{-15}	*(49)
57	$s-C_4H_9O_2+NO\rightarrow MEK+NO_2+HO_2$	245	1.17×10 ⁻¹¹	*(26)
58	$n-C_4H_9O_2+CH_3O_2 \rightarrow butanal+HCHO+2HO_2$	245	6.70×10^{-13}	*(44)
59	n-C ₄ H ₉ O ₂ +CH ₃ O ₂ →butanal+CH ₃ OH	245	2.30×10^{-13}	*(45)
60	$n-C_4H_9O_2+CH_3O_2\rightarrow n-C_4H_9OH+HCHO$	245	2.30×10^{-13}	*(46)
61	$n-C_4H_9O_2+NO\rightarrow butanal+NO_2+HO_2$	245	1.21×10 ⁻¹¹	*(25)
62	$s-C_4H_9O_2+HO_2\rightarrow s-C_4H_9OOH+O_2$	245	1.47×10 ⁻¹¹	*(35)
63	$n-C_4H_9O_2+HO_2\rightarrow n-C_4H_9OOH+O_2$	245	1.47×10 ⁻¹¹	*(35)
64	Isobutane+OH→t-C ₄ H ₉ O ₂	245	1.42×10 ⁻¹²	h
65	Isobutane+OH→i-C ₄ H ₉ O ₂	245	2.31×10^{-13}	h
66	Isobutane+Cl→t-C ₄ H ₉ O ₂	245	4.68×10^{-11}	i
67	Isobutane+Cl→i-C ₄ H ₉ O ₂	245	8.32×10^{-11}	i
68	$t-C_4H_9O_2+NO\rightarrow acetone+CH_3O_2+NO_2+HO_2$	245	1.17×10 ⁻¹¹	*(26)

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Table 1. Continued.

Rxn. #	Reaction	T (K)	Rate constant	Ref.
69	$t-C_4H_9O_2+HO_2\rightarrow t-C_4H_9OH+HCHO$	245	1.47×10 ⁻¹¹	*(35)
70	$t-C_4H_9O_2+CH_3O_2 \rightarrow acetone+CH_3O_2+CH_3OH$	245	4.10×10^{-15}	*(48)
71	$t-C_4H_9O_2+CH_3O_2 \rightarrow acetone+CH_3O_2+HCHO+2HO_2$	245	1.20×10 ⁻¹⁴	*(47)
72	$t-C_4H_9O_2+CH_3O_2\rightarrow tC_4H_9OOH+O_2$	245	4.10×10^{-15}	*(49)
73	$i-C_4H_9O_2+NO \rightarrow isobutanal+CH_3O_2+NO_2+HO_2$	245	1.21×10 ⁻¹¹	*(25)
74	$i-C_4H_9O_2+HO_2\rightarrow i-C_4H_9OH+HCHO$	245	1.47×10 ⁻¹¹	*(35)
75	i-C ₄ H ₉ O ₂ +CH ₃ O ₂ →isobutanal+CH ₃ OH	245	2.30×10^{-13}	*(45)
76	i-C ₄ H ₉ O ₂ +CH ₃ O ₂ →isobutanal+HCHO+2HO ₂	245	6.70×10^{-13}	*(44)
77	$i-C_4H_9O_2+CH_3O_2\rightarrow i-C_4H_9OH+HCHO$	245	2.30×10^{-13}	*(46)
78	MEK+OH→products	245	1.17×10 ⁻¹²	
79	MEK+Cl→products	245	2.75×10^{-11}	j
80	Acetone+OH→products	245	1.30×10 ⁻¹³	k
81	Propanal+OH→products	245	2.61×10^{-11}	
82	$O(^{1}D)+H_{2}O\rightarrow 2OH$	245	2.19×10^{-10}	b
83	$O_3+OH\rightarrow HO_2$	245	3.74×10^{-14}	
84	$O_3 + HO_2 \rightarrow OH$	245	1.35×10 ⁻¹⁵	
85	$H_2O_2+OH\rightarrow HO_2$	245	1.51×10 ⁻¹²	
86	$NO+NO_3\rightarrow 2NO_2$	245	2.82×10^{-11}	
87	CH ₃ OOH+OH→HCHO+OH	245	4.47×10^{-12}	b
88	CH ₃ OOH+OH→CH ₃ O ₂	245	4.13×10 ⁻¹²	
89	CH ₃ CHO+OH→PA	245	1.95×10 ⁻¹¹	
90	2PA→2CH ₃ O ₂	245	2.19×10^{-11}	b
91	Butanal+OH→products	245	3.13×10^{-11}	
92	Cl+O ₃ →ClO	245	1.00×10^{-11}	

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Table 1. Continued.

Rxn. #	Reaction	T (K)	Rate constant	Ref.
93	HCI+OH→CI	245	6.76×10^{-13}	
94	CIO+HO ₂ →HOCI	245	8.19×10 ⁻¹²	b
95	CIO+NO→CI+NO ₂	245	2.07×10^{-11}	
96	CIO+NO ₂ →CINO ₃	245	2.18×10 ⁻¹¹	b
97	$CIO+CH_3O_2\rightarrow CI+HCHO+HO_2$	245	2.06×10^{-12}	b
98	$CIO+PA \rightarrow CI+CH_3O_2+CO_2$	245	2.06×10^{-12}	*(97)
99	Br ₂ +Cl→BrCl+Br	245	1.20×10 ⁻¹⁰	1
100	$BrCl+Cl\rightarrow Br+Cl_2$	245	1.50×10 ⁻¹¹	1
101	BrO+ClO→Br+O ₃ +ClO	245	9.29×10^{-12}	1
102	BrO+ClO→Br+Cl	245	7.05×10^{-12}	1
103	BrO+ClO→BrCl	245	1.16×10 ⁻¹²	
104	$Br+C_2H_2 \rightarrow HBr+HO_2+2CO$	298	5.15×10 ⁻¹⁵	е
105	Br+HO ₂ →HBr	245	1.29×10 ⁻¹²	
106	HBr+OH→Br	245	1.10×10 ⁻¹¹	
107	BrO+propanal→HOBr	298	1.50×10 ⁻¹⁴	*(119)
108	BrO+HBr→HOBr+Br	298	2.10×10^{-14}	m
109	BrO+HO ₂ →HOBr	245	4.74×10^{-11}	1
110	$BrO+NO \rightarrow Br+NO_2$	245	2.51×10^{-11}	
111	$BrO+NO_2 \rightarrow BrNO_3$	245	1.22×10 ⁻¹¹	b
112	BrO+CH ₃ O ₂ →HOBr	298	4.00×10^{-12}	n
113	$BrO+CH_3O_2 \rightarrow Br+HCHO+HO_2$	298	1.70×10 ⁻¹²	n
114	2BrO→2Br	245	2.70×10^{-12}	
115	$C_3H_8+OH\rightarrow n-C_3H_7O_2$	245	2.15×10 ⁻¹³	d
116	$C_3H_8 + OH \rightarrow i - C_3H_7O_2$	245	7.12×10^{-13}	
117	BrCl+Br→Br ₂ +Cl	245	3.30×10^{-15}	I

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Table 1. Continued.

Rxn. #	Reaction	T (K)	Rate constant	Ref.
118	Br+Cl ₂ →BrCl+Cl	245	1.10×10 ⁻¹⁵	
119	BrO+HCHO→HOBr	298	1.50×10 ⁻¹⁴	m
120	$HO_2+OH\rightarrow H_2O$	245	1.33×10 ⁻¹⁰	
121	HCHO+OH→HO ₂ +CO	245	9.65×10^{-12}	
122	BrO+CH ₃ CHO→HOBr+PA	245	1.50×10 ⁻¹⁴	*(119)

First- and second-order rate constants are in units of s⁻¹ and cm³ molec⁻¹ s⁻¹, respectively.

All rate constant values from Atkinson et al. (2001) unless otherwise noted: a (Lightfoot et al., 1992), b (DeMore et al., 1997), c (Tyndall et al., 1997), d (Atkinson et al., 1989), e (Wallington et al., 1989), f (Villenave and Lesclaux, 1996), g (Jenkin et al., 1997), h (Donahue et al., 1998) (branching ratios from Atkinson, (1986)), i (Hooshiyar and Niki, 1995) (branching ratios from Cadman, et al., (1976)), j (Cuevas et al., 2004), k (Wollenhaupt et al., 2000), I (Sander and Crutzen, 1996), m (Hansen et al., 1999), n (Aranda et al., 1997)

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^{*} indicates estimated rate constant, with reaction number used for estimate in parenthesis

Table 2. Fluxes included in 0-D model.

Flux number	Species	Radiation dependent	Maximum Value	Source
1	Br ₂	No	Various	*
2	BrCl	No	Various	*
3	Cl ₂	No	Various	*
4	HOBr	No	$-1.25 \times 10^{-5} \mathrm{s}^{-1}$	Michalowski et al. (2000)
5	HOCI	No	$-1.25 \times 10^{-5} \mathrm{s}^{-1}$	Michalowski et al. (2000)
6	HBr	No	$-1.25 \times 10^{-5} \mathrm{s}^{-1}$	Michalowski et al. (2000)
7	HCI	No	$-1.25 \times 10^{-5} \mathrm{s}^{-1}$	Michalowski et al. (2000)
8	N_2O_5	No	$-1.25 \times 10^{-5} \mathrm{s}^{-1}$	Michalowski et al. (2000)
9	HONO	No	$-1.25 \times 10^{-5} \mathrm{s}^{-1}$	Michalowski et al. (2000)
10	HNO_4	No	$-1.25 \times 10^{-5} \mathrm{s}^{-1}$	Michalowski et al. (2000)
11	HONO	Yes	2.95×10^5 molec/cm ³ s	Zhou et al. (2001)
12	NO_x	Yes	1.10×10^6 molec/cm ³ s	Beine et al. (2002)
13	acetone	Yes	1.85×10^4 molec/cm ³ s	Guimbaud et al. (2002)
14	propanal	Yes	5.89×10^4 molec/cm ³ s [#]	*

^{*}Source determined empirically from model results

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^{*}This flux is not always present, as discussed in the text

Table 3. Photolysis reactions included in the 0-D model.

Reaction Number	Reaction	Max. Rate (s ⁻¹)
1	$O_3 \rightarrow O(^1D)$	5.02×10 ⁻⁷
2	$NO_2 \rightarrow NO + O_3$	1.49×10^{-3}
3	$NO_3 \rightarrow NO + O_2$	8.33×10 ⁻³
4	$NO_3 \rightarrow NO_2 + O_3$	5.57×10 ⁻²
5	H ₂ O ₂ →2OH	6.02×10^{-7}
6	HONO→OH+NO	3.04×10^{-4}
7	$HNO_3 \rightarrow OH + NO_2$	2.78×10^{-8}
8	$HNO_4 \rightarrow HO_2 + NO_2$	1.90×10 ⁻⁷
9	HCHO→2HO ₂	1.74×10^{-6}
10	HCHO→CO+H ₂	5.35×10^{-6}
11	$CH_3CHO \rightarrow CH_3O_2 + HO_2$	1.38×10^{-7}
12	Propanal→HO ₂ +C ₂ H ₅ O ₂	5.20×10^{-7}
13	Acetone→CH ₃ O ₂ +PA	4.10×10^{-9}
14	CH ₃ OOH→HCHO+HO ₂ +OH	5.69×10^{-7}
15	Cl ₂ →2 Cl	3.43×10^{-4}
16	HOBr→OH+Br	4.17×10^{-4}
17	$BrO \rightarrow Br + O_3$	4.63×10^{-3}
18	Br ₂ →2Br	8.72×10^{-3}
19	$N_2O_5 \rightarrow NO_2 + NO_3$	3.52×10^{-6}
20	MEK→products	1.10×10 ⁻⁷ *
21	$BrNO_3 \rightarrow BrO + NO_2$	7.86×10^{-5}
22	$BrNO_3 \rightarrow Br + NO_3$	1.67×10^{-4}
23	BrCl→Br+Cl	4.80×10 ⁻⁴ *
24	$CINO_3 \rightarrow CI + NO_3$	5.35×10^{-6}
25	CINO ₃ →CIO+NO ₂	6.36×10^{-7}
26	HOCI→OH+CI	5.69×10 ⁻⁶ *

All photolysis calculations are for Alert, Canada on 29 March 1995 Unless otherwise noted, photolysis rate was calculated using the TUV model, Version 4.3 *Photolysis rate calculated as discussed in the text

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Table 4. Initial model conditions*.

Species	Concentration
	(molec/cm ³)
H ₂ O	2.28×10 ¹⁶
НСНО	6.00×10 ⁹
CH ₃ CHO	3.00×10 ⁹
C_2H_6	4.00×10 ¹⁰
CH ₄	5.66×10 ¹³
O ₃	6.60×10 ¹¹
C_3H_8	1.00×10 ¹⁰
propanal	7.80×10 ⁸
PAN	1.20×10 ¹⁰
n-butane	2.65×10 ⁹
isobutane	1.30×10 ⁹
C_2H_2	2.20×10 ⁹
HONO	1.50×10 ⁸
Acetone	1.40×10 ¹⁰
CO	3.00×10^{12}
HNO_4	1.00×10 ⁸
NO ₂	7.50×10^7
MEK	3.30×10^{8}

^{*}Species in bold are held constant in the model

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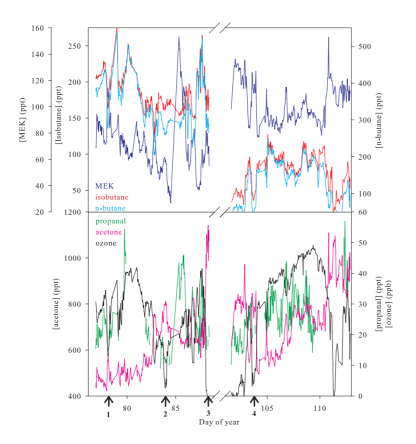
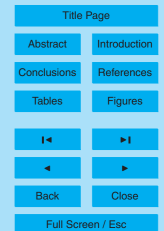


Fig. 1. Ambient concentrations of methyl ethyl ketone, isobutane, n-butane, acetone, propanal and ozone during PSE95. Ozone depletion events of interest, as discussed in the text, are labeled with numbers 1 through 4.

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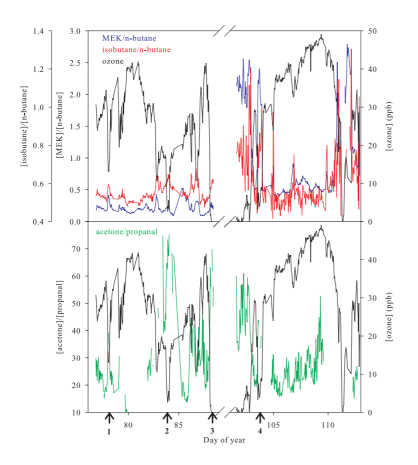
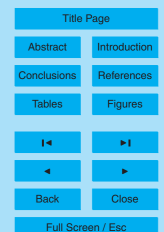


Fig. 2. Ambient measurements of [MEK]/[n-butane], [isobutane]/[n-butane], [acetone]/ [propanal] and [ozone] during PSE95. Ozone depletion events of interest, as discussed in the text, are labeled with numbers 1 through 4.

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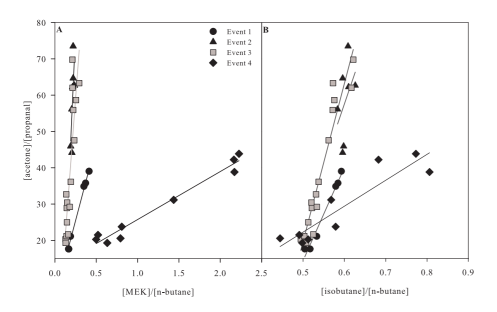
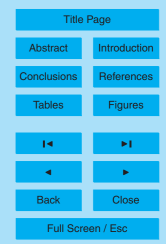


Fig. 3. Ambient [MEK]/[n-butane] **(A)** and [isobutane]/[n-butane] **(B)** vs. [acetone]/[propanal] during PSE95 for the four ozone depletion events indicated in Figs. 1 and 2.

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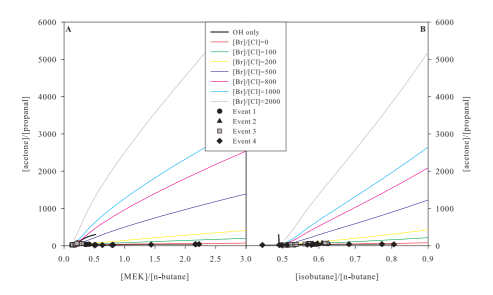


Fig. 4. Model results of [MEK]/[n-butane] **(A)** and [isobutane]/[n-butane] **(B)** vs. [acetone]/ [propanal] for various [Br]/[Cl] when no propanal flux is present are compared to the ambient data shown in Fig. 3. All model results are daytime averages over a time period of ten days, and reported [Br]/[Cl] is the daytime average for model Day 9.

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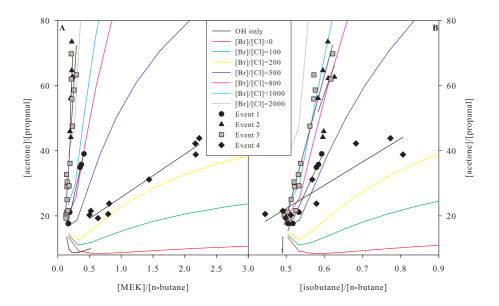


Fig. 5. Model results of [MEK]/[n-butane] **(A)** and [isobutane]/[n-butane] **(B)** vs. [acetone]/ [propanal] for various [Br]/[Cl] when a radiation-dependent propanal flux is present are compared to the ambient data shown in Fig. 3. All model results are daytime averages over a time period of ten days, and reported [Br]/[Cl] is the daytime average for model Day 9.

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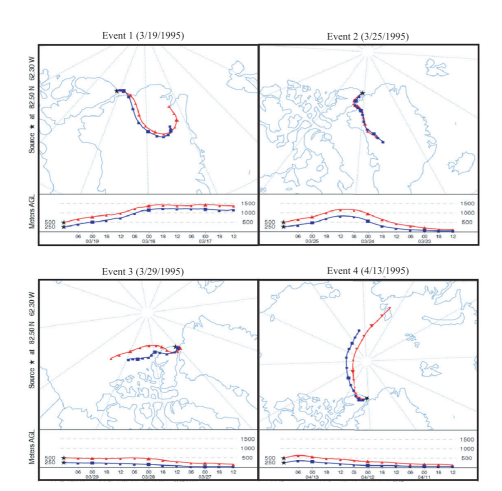
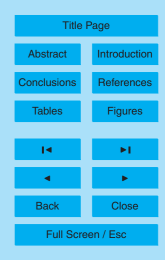


Fig. 6. 72 h HYSPLIT back trajectories are presented for each of the four ozone depletion events of interest during PSE95.

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