An overview of snow photochemistry: Evidence, mechanisms and impacts

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An overview of snow photochemistry: evidence, mechanisms and impacts


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Abstract. It has been shown that sunlit snow and ice plays an important role in processing atmospheric species. Photochemical production of a variety of chemicals has recently been reported to occur in snow/ice and the release of these photochemically generated species may significantly impact the chemistry of the overlying atmosphere. Nitrogen oxide and oxidant precursor fluxes have been measured in a number of snow covered environments, where in some cases the emissions significantly impact the overlying boundary layer. For example, photochemical ozone production (such as that occurring in polluted mid-latitudes) of 3–4 ppbv/day has been observed at South Pole, due to high OH and NO levels present in a relatively shallow boundary layer. Field and laboratory experiments have determined that the origin of the observed NOx flux is the photochemistry of nitrate within the snowpack, however some details of the mechanism have not yet been elucidated. A variety of low molecular weight organic compounds have been shown to be emitted from sunlit snowpacks, the source of which has been proposed to be either direct or indirect photo-oxidation of natural organic materials present in the snow. Although myriad studies have observed active processing of species within irradiated snowpacks, the fundamental chemistry occurring remains poorly understood. Here we consider the nature of snow at a fundamental, physical level; photochemical processes within snow and the caveats needed for comparison to atmospheric photochemistry; our current understanding of nitrogen, oxidant, halogen and organic photochemistry within snow; the current limitations faced by the field and implications for the future.

1 Introduction

It is now widely recognized that the Earth System is tightly interconnected. Changes in one component can strongly affect the state of another; feedbacks between them can have subtle influences that might either amplify or mitigate trends. A connection now receiving growing attention is that between the atmosphere and the cryosphere. The cryosphere forms a large proportion of the Earth’s surface: a seasonal maximum of 40% of land is covered by snow or ice, while several percent of the world’s oceans are covered by sea ice. Traditionally, the cryosphere has been viewed as a “cap”, inhibiting emissions from land and ocean surfaces below and acting itself as a permanent sink of atmospheric species. The snow itself has not been considered beyond its effect on radiative transfer through albedo.

Recent evidence, however, has shown that the polar cryosphere can have a major influence on the overlying atmosphere. Rather than being inert, or simply a sink for impurities, snow is highly photochemically active, with snowpack impurities photolyzed to release reactive trace gases into the boundary layer. Since the initial discoveries of CH3O and NOx production within polar snow (Fuhrer et al., 1996; Sumner and Shepson, 1999; Honrath et al., 1999) evidence for the photochemical production and release of a range of trace gases has been found. These processes appear to be ubiquitous, occurring wherever sunlight shines on snow. The significance of their influence varies according to background concentrations of radicals, and is less important in boundary layers that are anthropogenically perturbed. But in the remote high latitudes, emissions from the snow can dominate boundary layer chemistry. On the Antarctic plateau, for example, some oxidants are as abundant as in the tropical troposphere when viewed in terms of 24 h averages (Mauldin et al., 2004).

The cryosphere, however, is not static. Global snow/ice coverage fluctuations over both seasonal and climatic timescales. In our present interglacial period, snow and ice are not restricted to polar regions but are found at much lower latitudes according to the time of year. Previously, the great ice sheets of the glacial periods covered 25% of the Earth’s surface year-round (as opposed to the present-day 10%) with additionally extensive seasonal snow and sea-ice coverage. Predictions for the future are for considerably less snow coverage than at present. The influence of the cryosphere on atmospheric composition certainly has varied through time and will change in the future.

The science of “snow photochemistry” is relatively young. It is an interdisciplinary subject, drawing on expertise in a wide range of areas. The aim of this paper is to draw this expertise together, and to disseminate information that is relevant for understanding emissions from snow and their influence on atmospheric chemistry. Here we review the detailed chemistry and microphysics of snow itself; explore photochemistry above and within snow; and review observational evidence of the impact snow photochemistry has on the boundary layer and the chemical and physical mechanisms that drive the emissions. Finally we assess current limitations that are impeding progress in understanding, and consider implications for future atmospheres.

2 Unique physical and chemical aspects of snow

2.1 Understanding the location of impurities in snow

Fundamental to the study of snow photochemistry is an appreciation of snow structure, and in particular, the location within snow crystals/grains where impurities reside. It is these impurities that may ultimately undergo reactive processes and generate trace gas products.

Most of the mass of precipitating snow crystals forms by the condensation of water vapor onto an ice-forming nucleus (IFN) or by the freezing of supercooled droplets onto growing ice crystals, a process called riming (Pruppacher and Klett, 1978). IFNs are therefore a source of impurities in
snow crystals, as are the cloud condensation nuclei (CCN) that nucleate supercooled water droplets. Various materials can act as IFN or CCN: plant debris, bacteria, minerals, and the ubiquitous sulfate aerosols (Pruppacher and Klett, 1978; Khvorostyanov and Curry, 2000; Sattler et al., 2001; Targino et al., 2006). Supercooled droplets can also scavenge gases and non-activated aerosols in the cloud. Rimed snow is usually more concentrated in impurities than snow formed solely from the condensation of water vapor (Mitchell and Lamb, 1989; Pouilda et al., 1998). The location of species trapped in rime ice has been little studied. They could form supersaturated solid solutions (a solid-state solution of solutes within ice), or pockets and veins of brine, as observed during the freezing of sea water (Eicken, 1992).

The growth of ice crystals by vapor condensation often takes place in a discontinuous manner, with new layers of water molecules condensing at crystal edges (Nelson and Knight, 1998). Experiments at low temperatures (<190 K) have shown that molecules such as HCl or HNO3 ionize and become solvated on contact with ice surfaces (Horn et al., 1992; Banham et al., 1995). The ions can soon become incorporated in the frozen lattice because of the rapid desorption/adsorption of water from/to the ice surface. Other molecules such as H2O2 and CH2O also become buried in ice structures but little is known of the chemical form that they take (for example as CH2O or H2C(OH)2) or whether they attach to surface hydrogen bonds, dissolve in the quasi-liquid layer (see below) or reside in ice “micropockets”. Larger molecules such as acetone, acetaldehyde and alcohols adsorb on ice by forming hydrogen bonds (Picaud et al., 2000, Sokolov and Abbatt, 2002; Dominé and Rey-Hanot, 2002; Winkler et al., 2002; Hudson et al., 2002; Bartels-Rausch et al., 2004). Semi-volatile organic molecules of low polarity adsorb onto ice through van der Waals interactions (Roth et al., 2004; Goss, 2005), and are considered not to dissolve. These large molecules include polycyclic aromatic hydrocarbons (PAHs) and persistent organic pollutants (POPs) (Jaffrezo et al., 1994; Blais et al., 1998; Daly and Wania, 2004).

The surface and grain-boundaries of ice are disordered (e.g. Petrenko and Withworth, 1999; Döppenschmidt and Butt, 2000; Wei et al., 2001; Girardet and Toubin, 2001; Sadtchenko and Ewing, 2002; Cho et al., 2002) to the extent that near the freezing point, the ice surface layer has been called the quasi-liquid layer (QLL). The lowest temperature where the QLL still exists is uncertain, as it depends on the method used to detect it and on the presence of adsorbed impurities that can lower its temperature range of existence. Wei et al. (2001) show evidence for the existence of the QLL down to −70°C. For ice with very high ionic concentrations, the ions are excluded from the bulk and form a brine layer at the surface or at grain boundaries, with large ionic strengths in that layer (Cho et al., 2002; Carignano et al., 2006). Adsorption of gases on ice surfaces at tropospheric temperatures has sometimes been described as dissolution in the QLL. Molecular dynamics studies confirm that actual solvation and solution of polar molecules within the QLL takes place (Compoint et al., 2002). Therefore, the more soluble and dissociating acids probably even affect the structure of the QLL. For the less soluble gases, most experimental results have been able to describe the adsorption of trace gases on ice by the formalism used for adsorption on solid surfaces (e.g. Sokolov and Abbatt, 2002; Winkler et al., 2002; Dominé and Rauzy, 2004; Ullerstam et al., 2005; Ullerstam and Abbatt, 2005).

During precipitation, falling crystals can scavenge atmospheric aerosols, and gases can adsorb onto the crystal’s surface or even diffuse into its bulk. The composition of a snow crystal arriving at the Earth’s surface is a complex function of the process of formation, the composition of the cloud where it was formed, and the composition of the air mass underlying the cloud (Schwikowski et al., 1998; Franz and Eisenreich, 1998; Lei and Wania, 2004). Following precipitation, the snow will be out of thermodynamic equilibrium with the atmosphere, due to differences in temperature and air composition from that of the originating cloud, thus, gases can be taken up or emitted by the snow (Conkin et al., 1993; McConnell et al., 1998; Hutterli et al., 2002, 2003). Chemical reactions can also take place on the crystal surface, in its volume, or on/in aerosol particles captured by the snow, or scavenged during descent. The availability of species to be emitted or to react will depend on how they were incorporated in the snow.

Adsorbed species react rapidly to changes in thermodynamic conditions and are readily available for reaction with atmospheric gases. Species present on scavenged aerosols located on the surface of snow crystals can be expected to have a reactivity similar to that in the aerosol phase. Species dissolved within ice crystals, or contained in IFN and CCN are completely trapped within ice crystals, and are thus not in contact with atmospheric gases. Their reactivity will be limited to solid phase processes that are extremely slow, and to photolysis, whose efficiency may be severely limited by cage effects. Species forming a solid solution with ice can diffuse within the ice crystalline lattice and reach the surface. The diffusion coefficients, D, measured for HCl, HNO3 and CH2O in ice around −15°C are in the range 10−11 to 10−12 cm2 s−1 (Thibert and Dominé, 1997, 1998; Perrier et al., 2003; see also a critical review of D measurements in Huthwelker et al., 2006). The physical environment of species in rime ice is not clear. However, if they are contained in brine pockets or veins in contact with the atmosphere, their release time will be dictated by their diffusion rate in a liquid phase, with a D value around 10−5 cm2 s−1, much faster than for species in solid solutions.

At present, understanding of the location of impurities in natural snow is limited, so that predicting snow chemical
reactivity based solely on bulk chemical composition is not possible. It is therefore clear that one of the great needs to advance the state of the science is the ability to probe the chemical morphology of snow and ice, in the macro- and microscopic domains (as further discussed in Sect. 6).

2.2 Physical and chemical transformations of snow crystals after deposition

After deposition, snow crystals in the dry snowpack are subjected to temperature gradients that generate water vapor fluxes between crystals. These fluxes cause the sublimation of parts of crystals and condensation on other parts, resulting in changes in snow crystal sizes and shapes, and changes in the physical properties of the snowpack, such as density, porosity, heat conductivity, hardness, specific surface area and albedo. The processes that lead to snowpack metamorphism occur on individual grains, but result in bulk compositional changes to both physical and chemical properties of the snowpack. Snow physics as related to photochemical processes is reviewed by Dominé et al. (2007).

Major changes in snow composition occur after deposition. Examples include the decrease of nitrate in seasonal snowpacks and on ice caps (Mayewski and Legrand, 1990; Nakamura et al., 2000; Röthlisberger et al., 2000, 2002; Beine et al., 2002a), and the increase in sulfate, mineral dust, and sea salt compounds (Harder et al., 2000; Aoki et al., 2000; Dominé et al., 2004). However, relating those changes to a specific process is often difficult. Changes can be caused by physical processes such as dry deposition of gases and aerosols, the formation of surface hoar crystals or the freezing of supercooled droplets during fog events (Bergin et al., 1994, 1995, 1996). Photoreaction of snow species between themselves or with atmospheric gases (Sumner and Shepson, 1999; Spicer et al., 2002; Grannas et al., 2004) may also lead to changes in snow composition, as well as processes directly linked to snow metamorphism such as the release of adsorbed and dissolved species (Hutterli et al., 2004). Sublimation/condensation cycles during metamorphism have the potential to release or trap dissolved species and expose trapped aerosols, changing snow and atmospheric composition and making these species available for gas phase or surface reaction. Solid-state diffusion of species forming solid solutions with ice can also take place. Metamorphism almost always results in the decrease of the specific surface area of snow (Cabanes et al., 2002, 2003), which inevitably results in a decrease in the amount of adsorbed species.

Physical processes and photochemistry can both contribute to decreases in impurity concentrations in snow. For example, release by purely physical processes has been invoked to explain the huge post-depositional decrease in nitrate concentrations in central Antarctic snow (Nakamura et al., 2000; Röthlisberger et al., 2000; Blunier et al., 2005), but nitrate photolysis (Honrath et al., 1999; Ridley et al., 2000; Davis et al., 2001; Jones et al., 2001; Wolff et al., 2002) is also likely to contribute (see further discussion in Sect. 5.3). In addition, the presence of high levels of other impurities in snow affecting the pH influence the preservation of e.g. nitrate and chloride: Elevated levels of sulfate can mobilize nitrate whereas high levels of dust immobilizes chloride in the snow (Röthlisberger et al., 2002, 2003). Fewer processes can affect aerosol species that have long been considered to be irreversibly deposited (Dibb and Jaffrezo, 1997). However, species such as semi-volatile organics, which are partly aerosol bound, can be lost through both physical and photochemical processes. Most observations indicate an increase in aerosol species by dry deposition, a process accelerated by wind, as snow can effectively filter out particles entrained into its pores by wind-driven air motion (e.g., Waddington et al., 1996; Harder et al., 2000; Dominé et al., 2004). Snow metamorphism can also affect the nature and rate of photochemical reactions, largely through changes in snow grain size and specific surface area, hence snow albedo (Dominé et al., 2006) and light penetration depth (Simpson et al., 2002a).

Wet metamorphism can dramatically affect snow composition. Soluble species, including the well-studied acids H$_2$SO$_4$, HNO$_3$ and HCl, have a greater affinity for water than for ice. They partition preferentially to the liquid phase and are readily removed by the first stages of percolation (e.g., Tranter et al., 1986). However, even for inorganic ions there is evidence for significant fractionation driven by preferential elution of some ions (e.g. SO$_4^{2-}$, Ca$^{2+}$, Mg$^{2+}$, K$^+$, Na$^+$) before others (NO$_3^-$, NH$_4^+$, Cl$^−$, F$^−$) (Eichler et al., 2001). Less soluble species such as non-polar organic molecules are also found in meltwaters, but are less easily removed by percolation (Meyer et al., 2006). Particulate material is also entrained by percolating water, but rarely in the early stages and may remain in the snow until the final stages of melting (Hodgkins et al., 1998; Lyons et al., 2003; Meyer et al., 2006).

2.3 Interaction of organics with ice

An understanding of the interfacial interactions between ice/snow and organic molecules comes from various studies, including those of physical and chemical properties of the ice surface (Engquist 1995a, 1995b; Bertilsson et al., 1997, 1999; Wania et al., 1998; Schaff and Roberts, 1999a, 1999b; Girardet and Toubin, 2001; Borodin et al., 2004; Guidi, 2004; Roth et al., 2004; Guzmán et al., 2006a; Heger et al., 2005; Heger and Klán, 2007) and cryogenic chemical behavior of ice contaminants (Sumner and Shepson, 1999; Wania et al., 1999; Dubowski and Hoffmann, 2000; Klán and Holoubek, 2002; Coloussi and Hoffmann, 2003; Klán et al., 2003; Klánová et al., 2003a, 2003b; Grannas et al., 2004; Guzmán et al., 2006b; Heger et al., 2006). Adsorption of various organic molecules on ice surfaces can be described well with a multi-parameter linear free energy relationship, based on the van der Waals and the electron donor/acceptor interactions (such as H-bonding) (Roth et al., 2004). Studies
at low to very low temperatures (80 to 190 K) have found that ice surfaces interact with contaminants via three different important types of surface water molecules, including those with dangling hydrogen or oxygen atoms (Devlin, 1992; Devlin and Buch, 1995). Many organic halocarbon compounds have also been shown to adsorb on water-ices by interactions with the ice surface dangling bonds (Holmes and Sodeau, 1999). Studies of the adsorbed states of some organic molecules, such as acetonitrile, chloroform (Schaaff and Roberts, 1999a, 1999b), acetone (Schaaff and Roberts, 1998), or benzene derivatives (Borodin et al., 2004) have revealed the scope of hydrogen-bonding or dipolar interactions. It is not clear, however, how these low temperature data apply to snow surfaces that exist in the range 200–273 K, where the QLL is present. Most importantly, adsorption, desorption, interaction types, or diffusion of the molecules are known to be temperature and phase-dependent variables (Cho et al., 2002; Abbatt, 2003; Heger et al., 2005).

Some solutes are known to become spontaneously segregated at grain boundaries in the polycrystalline ice during the freezing process (Cohen et al., 1996; Finnegan and Pitter, 1997), however most studies have focused on inorganic ions, with relatively little known about the behavior of neutral organics. Such a solute concentration-enhancing effect (Dash et al., 1995; Takenaka et al., 1996; Cho et al., 2002) may cause solute organic molecules to self-organize (Heger et al., 2005). Many laboratory studies have provided evidence that the ice/snow impurities are located in the QLL on the surface of the single ice crystals (Conklin and Bales, 1993; Dash et al., 1995; Dubowski et al., 2001; Wei et al., 2001; Cho et al., 2002; Dubowski et al., 2002; Chu and Anastasio, 2003; Klánová et al., 2003a; Robinson et al., 2006). Most of these studies, however, used dopant concentrations much higher than natural ones, and this preferential segregation may not always occur in nature. Indeed, species such as CH$_2$O, and possibly others, have a small but non-zero solubility in ice (Burkhart et al., 2002; Perrier et al., 2003), and it is possible that this segregation takes place only once this solubility limit is exceeded, which happens in the laboratory but not in nature. The concentration-enhancing effect in partially frozen aqueous solutions also has been described in connection with the acceleration of some thermal reactions since the 1960s (Grant et al., 1961; Bruice and Butler, 1964; Butler and Bruice, 1964; Fennema, 1975; Takenaka et al., 1992; Takenaka et al., 1996). Recent work has illustrated that the freeze-concentration effect can also have significant impacts on the photochemistry of organics occurring in the QLL on the surface of ice (Bausch et al., 2006).

2.4 Impacts of freezing on snowpack impurities and reactions

Various studies have shown that the action of freezing can affect chemical impurities or reaction processes in snow (Takenaka et al., 1992, 1996; Betterton and Anderson, 2001; O’Driscoll et al., 2006). In many heterogeneous reactions, rate acceleration and/or new product pathways are promoted by natural freeze-thaw cycles which can occur in snow and ice.

For example, it has been shown that both sulfur dioxide and sulfide ions incorporated into ice are oxidized to sulfate ions (Valdez et al., 1989; Finnegan et al., 1991, Betterton and Anderson, 2001). Similarly iodide and bromide ions become oxidized to higher valence species when frozen (Eyal et al., 1964). It was later discovered that the N(III) species, NO$_2^-$ and HONO, can be oxidized by molecular oxygen to nitrate ions upon freezing in aqueous solution at a rate about 10$^5$ faster than that found at room temperature (Takenaka et al., 1992). The mechanism was ascribed to a combination of ion-separation and a “freeze-concentration” pathway in which hydrogen ions were rejected from the ice to solution pockets existing within the overall structure. The reaction was suggested to take place in unfrozen solution “micropockets” surrounded by walls of ice grains. The importance of acidity in driving the reaction was also shown in the experiments, which were performed at pH values between 3.0 and 5.6. To occur within natural snow, therefore, these micro-pockets must be considerably acidic, something that needs further investigation.

3 Introduction to photochemistry in and above snow

The ultraviolet (UV) spectral region is the most critical for photochemistry in the atmosphere and snowpack because those photons possess high enough energy to break chemical bonds, but low enough energy to penetrate the ozone layer and reach the troposphere. Radiation of wavelengths shorter than 290 nm (UV-C region) is completely absorbed by the ozone layer and is thus not important to tropospheric chemistry. Photons in the UV-B region, 290 to 320 nm, pass at least partially through the ozone layer, and are responsible for tropospheric ozone photolysis and production of hydroxyl (OH) radicals through the reaction of O($^1$D) + H$_2$O, although other production mechanisms for OH are likely to be more important at high latitudes. The UV-B levels in the troposphere are highly dependent on the stratospheric ozone abundance and the solar zenith angle, which determines the path of light through the stratosphere. Ozone, bromoform (CHBr$_3$), and nitrate ions in aqueous solution (NO$_3^-$) are UV-B absorbers. The UV-A region, 320 to 400 nm, has low enough energy to pass through the ozone layer with little attenuation, and thus has a lower dependence than UV-B radiation on overhead ozone and solar zenith angle. Important atmospheric chemicals that are photolyzed in the UV-A are NO$_2$, HONO, CH$_2$O, and BrO. The solar zenith angle, ground reflectivity (albedo), and the overhead ozone are critical parameters for considering photochemical rates (Meier et al., 1997). At high latitudes, the sun never gets as high in the sky as it does at lower latitudes. Generally, this effect

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significantly decreases the photolysis rates with increasing latitude. However, the presence of snow at high latitudes, which causes the ground to have a high albedo – often >90% in the UV spectral region – acts to increase atmospheric photolysis rates, sometimes even overcoming the less favorable solar zenith angles. This albedo effect is very significant for UV-A absorbing species, and diurnally-averaged springtime photolysis rates at high latitudes often are comparable to mid-latitude values. Good examples of this effect are seen in the comparability of high and mid-latitude photolysis rates of NO$_2$, BrO, HONO, and CH$_2$O. The same albedo enhancement effect is present in the UV-B spectral region, but the long slant paths of the light through the ozone layer greatly attenuate the UV-B intensity and cause the photolysis rates for UV-B absorbers to be up to an order of magnitude smaller in the high latitudes than at mid latitudes (Simpson et al., 2002b). This effect, which varies with season and latitude, is particularly seen in ozone photolysis resulting in O($^1$D) atoms (Fig. 1) (Lefer et al., 2001). In addition to the aforementioned albedo effect at very high latitudes, there can also be a substantial influence at these latitudes from having 24 h of continuous photolysis and thus continuous photochemistry in summer. Additionally, the loss of stratospheric ozone in both the Antarctic and Arctic will allow for greater penetration of shorter wavelength (and more photochemically reactive) UV radiation to the surface, albeit at a time of year when solar irradiance is reduced compared with the summer.

Photolysis frequencies are quantified by the first-order rate coefficient for a molecule, which is normally termed $J$ (s$^{-1}$), and is given by: (Madronich, 1987; Meier et al., 1997)

$$J = \sigma(\lambda, T)\Phi(\lambda, T)F(\lambda)d\lambda$$

(1)

where $\sigma$ is the absorption cross section and $\Phi$ the quantum yield for the production of the compounds in question. Both $\sigma$ and $\Phi$ are functions of wavelength, $\lambda$, and temperature, $T$. $F$ is the actinic flux (photons cm$^{-2}$ nm$^{-1}$ s$^{-1}$), i.e. the omnidirectional flux of photons of wavelength impinging on the molecule.

In the atmosphere, light rays propagate long distances between scattering events, which allow the actinic flux to be measured directly by using a diffusing optic that collects light from all directions with equal sensitivity (Hofzumahaus et al., 1999). The actinic flux may also be modeled by radiation transfer models, such as the Tropospheric Ultraviolet and Visible model (TUV) (http://cprm.acd.ucar.edu/Models/TUV) (Madronich and Weller, 1990). In the snowpack, it is difficult to place the diffusing optics of an actinic flux spectroradiometer because of their large size and the short scattering length within snow. Therefore, most investigators of photolysis rate coefficients in snow have used measurements of irradiance to constrain radiation transfer models and invert the measurements to actinic fluxes and then photolysis rate coefficients (Simpson et al., 2002a). A complementary method to measure photolysis rate coefficients in snow uses a molecular probe known as a chemical actinometer. The actinometer molecule undergoes a well characterized unimolecular chemical reaction at a rate proportional to the solar actinic flux, and thus the actinic flux in a spectral region can be determined (Qiu et al., 2002; Galbavy et al., 2007a, b). Comparisons of chemical actinometry and spectral radiation measurements have generally shown the methods to agree well (Phillips and Simpson, 2005; Galbavy et al., 2007a, b).

The snowpack is a highly scattering medium with little absorption in the visible and UV region, which makes it appear brilliant white (Wiscombe and Warren, 1980; Grenfell et al., 1981, Warren, 1982; Grenfell et al., 1994). The simplest snowpack radiation transfer models only take into account the scattering coefficient, $S$, which is the probability of a photon scattering per unit length, and the absorption coefficient, $K$, which is the probability of a photon being absorbed per unit length. The scattering coefficient, $S$, is a weak function of wavelength and is most directly related to the snow grain size, or equivalently the specific surface area (SSA), and the interested reader is referred to the companion snow physics review article (Dominé et al., 2007). The absorption coefficient is a strong function of wavelength, and also is very low for pure ice in the visible and near ultraviolet, which makes it very susceptible to large increases due to even trace impurities (Perovich and Govoni, 1991). Thus, in the UV and visible regions, the absorption coefficient of snow is critically dependent on impurity content and chemical nature.

When radiation enters the snowpack from above, the scattering alters its path, converting direct into diffuse radiation. This scattering ultimately redirects the light back upwards and out of the snow, leading to the high albedo of snow in the visible and UV regions. The scattering also enhances the pathlength of the photons in the snowpack and

![Fig. 1. Cloud-free $J[O_3 + h\nu + \rightarrow O_2 + O(1D)]$ from TUV (Tropospheric Ultraviolet and Visible) radiation model for various latitudes and seasons at selected sites where snow photochemistry measurements have been made. Data shown for: South Pole (90° S, 23-December, 2000); Niwot Ridge, Colorado, USA (40° N, 18 April, 2003); Houghton, Michigan, USA (47° N, 14 January, 1999); and Summit, Greenland (74° N, 23 June, 2000).](https://www.atmos-chem-phys.net/7/4329/2007/fig1.png)
thus enhances the absorption probability and photochemical rates for trace absorbers in the snow. The scattering and absorption combine nonlinearly to control the depth to which photons, on average, penetrate into the snowpack. For diffuse radiation and deep and uniform snowpack, the attenuation of light follows the Bouger-Lambert law (Bohren and Barkstrom, 1974), which states that the attenuation of light varies exponentially with depth.

\[ I(d) = I(d') e^{-a_0(\lambda)(d-d')} \]  

In this equation, \( I(d) \) is the irradiance at depth \( d \) and \( a_0(\lambda) \) is the asymptotic flux extinction coefficient. The asymptotic flux extinction coefficient is the inverse of the e-folding depth, \( \varepsilon(\lambda) \), which is the depth over which the intensity of radiation decreases by a factor of e.

\[ \varepsilon(\lambda) = 1/a_0(\lambda) \]  

The e-folding depth of radiation in the UV-B to visible part of the spectrum in snow is typically in the range from 5–25 cm (Grenfell and Maykut, 1977; Grenfell et al., 1981; King and Simpson, 2001; Simpson et al., 2002a; Fisher et al., 2005; Warren et al., 2006; Galbavy et al. 2007a, b). The e-folding depth should be considered to be the characteristic depth of illumination of the snowpack, and the majority of photochemical reactions occur in this region (King and Simpson, 2001).

Penetration of light into snowpack, and thus the amount of photochemistry within the snowpack, is highly dependent on the solar zenith angle (Warren, 1982; Simpson et al., 2002a; Lee-Taylor and Madronich, 2002; Bourgeois et al., 2006). This effect is caused by the fact that snow grains typically act to forward-scatter light that interacts with them. Thus, for glancing incidence radiation that is characteristic of high solar zenith angles, a greater fraction of light is scattered back to space, the albedo is enhanced, and less light enters the snow to drive snowpack photochemistry. Light can impact snowpacks at low solar zenith angles near noon at mid- and low-latitude sites, for example at high altitude snowpacks on mountains. In this low solar zenith angle case, many forward scattering events are required to return a photon to space, and thus more of the illuminating light enters the snowpack and drives photochemistry. Therefore, snowpack photochemistry is highly dependent on the solar zenith angle and should be very rapid for noon-time conditions at low-latitude snowfields and glaciers.

4 Current understanding of snow photochemistry

4.1 Nitrogen oxides

4.1.1 Introduction to nitrogen oxides in Polar regions

Historically, measurements of trace gas chemistry at high latitudes targeted two distinct objectives. One was improved understanding of the chemistry of a clean background atmosphere; tropospheric concentrations of reactive trace gases such as NOx (NO and NO2) were expected to be very low (few parts per trillion by volume (pptv, pmol mol\(^{-1}\)) as in the remote marine boundary layer (Logan, 1983). It was assumed that the dominant sources of total reactive nitrogen oxides (NOx) included downwelling from the stratosphere, or long-range transport of N-species generated at lower latitudes by, e.g., tropical lightning, anthropogenic emissions or biomass burning. A second motivator was to increase our ability to interpret ice core data: nitrate (NO\(_3\)-) is an easy ion to measure from ice cores, so its interpretation in terms of changing atmospheric composition (of NOx or NO\(_3\)-) would be a significant prize. The first polar NO measurements, supported the a priori position. Early measurements at Barrow, an Arctic coastal site, indicated very low NO mixing ratios during most periods analyzed; any enhanced mixing ratios were attributed to local or regional combustion emissions (Honrath and Jaffe, 1992). On the Antarctic Peninsula NO remained below the 5 pptv instrumental detection limit (Jefferson et al., 1998); in retrospect the site was atypical for Antarctica, being surrounded by rock and ocean.

The discovery of elevated NOx mixing ratios within the snowpack interstitial air at Summit, Greenland thus came as a surprise (Honrath et al., 1999). Within the surface snowpack, NOx was a factor of 3 to \( >10 \) times higher than in ambient air and was generally greater than ambient NO\(_y\). Concentrations of NO\(_x\) in interstitial air varied diurnally, indicating that a N-containing reservoir within the snow, most likely nitrate (NO\(_3\)-), was photolyzed to release NO\(_x\) to snowpack interstitial air and potentially to the overlying boundary layer. A proximate source of NO\(_x\) helped to explain anomalous HNO\(_3\) and NO\(_y\) fluxes observed earlier at Summit (Dibb et al., 1998) and confirmed that standard tropospheric chemistry could not be directly applied in the boundary layer above sunlit snow.

4.1.2 Field studies to identify/quantify processes

A number of campaigns were subsequently conducted to look for snowpack NOx production at other locations and to test possible production processes (see Fig. 2 for location of measurement sites mentioned in the text). These studies used surface snow in a variety of ways – in the natural snowpack (Jacobi et al., 2004), as blocks (at Neumayer station, Antarctica, Jones et al., 2000), piles (at Alert, Nunavut, Canada, Beine et al., 2002a) and in flow-through chambers...
The first and final sections are measurements made in ambient air. Middle sections are measurements made within the snowblock, alternatively fully exposed to sunlight and fully shaded to eliminate any photochemical activity. Periods of shading are indicated by cross-hatching.

Several of the early Arctic studies extended measurements to include HONO (see Fig. 4). Certain questions exist about HONO measurements made in locations where mixing ratios are low (Kleffmann et al., 2006), with the data being higher than can be reconciled with model HO\(_x\) and NO\(_x\) chemistry (e.g. Bloss et al., 2006, and see also Sect. 4.2). The high latitude measurements of HONO discussed here should be interpreted with these potential caveats in mind.

A photochemical source of HONO from snow was also indicated, with elevated mixing ratios in snowpack interstitial air that were reduced by shading (Beine et al., 2002a; Dibb et al., 2002). The ratio of photochemical production of HONO compared to NO\(_2\) at Summit ranged from 1:1 to 1:3. Flux studies showed that HONO could also be released into the overlying boundary layer (Zhou et al., 2001; Honrath et al., 2002) (Fig. 4), with an emission ratio of NO\(_x\) (mainly as NO\(_2\)) to HONO of roughly 1:1 measured at Alert (Beine et al., 2002a). Subsequent work at other sites (Ny-Ålesund (Beine et al., 2003; Amoroso et al., 2005) and a high altitude mid-latitude site (Beine et al., 2005)) found that in locations where snow was alkaline, no significant HONO process occurs in most, if not all, sunlit snowpacks across the globe.

The atmospheric significance of the snow photochemistry phenomenon depends on the potential to emit the photoproducts to the overlying boundary layer. A series of flux experiments was conducted at various sites in both polar regions, to detect and quantify NO\(_x\) fluxes out of the snowpack (Jones et al., 2001; Honrath et al., 2002; Beine et al., 2002b; Oncley et al., 2004). In each case, the snowpack was found to be emitting NO\(_x\) into the boundary layer. The flux varied throughout the day, depending on solar intensity, and also changes in turbulence.

Several of the early Arctic studies extended measurements to include HONO (see Fig. 4). Certain questions exist about HONO measurements made in locations where mixing ratios are low (Kleffmann et al., 2006), with the data being higher than can be reconciled with model HO\(_x\) and NO\(_x\) chemistry (e.g. Bloss et al., 2006, and see also Sect. 4.2). The high latitude measurements of HONO discussed here should be interpreted with these potential caveats in mind.
emissions were detected. Furthermore, at Browning Pass, Antarctica, where snow was acidic, surprisingly small emissions of HONO were measured (Beine et al., 2006). This demonstrates the sensitivity of NO\textsubscript{3} emissions to the chemical composition of the snow, not just to physical parameters, as is discussed in detail later (see Sect. 4.1.4).

4.1.3 Field observations of ambient nitrogen oxides

**Seasonal variation of NO and NO\textsubscript{3}:** Figs. 5 and 6 provide an overview of ambient measurements of NO and NO\textsubscript{3} that have been made at high latitudes since the discovery of snowpack nitrogen photochemistry. The data are presented according to latitude and as daily averages, and, except for South Pole, are plotted on the same scale. The original papers show details not apparent in Fig. 5. By considering both NO and NO\textsubscript{3}, it is possible to see whether differences in NO are driven by emissions or by re-partitioning between NO and NO\textsubscript{2}.

Mixing ratios of NO\textsubscript{3} are similar at Summit and Ny-Ålesund, but considerably lower than at Poker Flat. At Alert NO\textsubscript{3} is highly variable, ranging from <5 pptv to over 80 pptv, most likely driven by differing air mass origins, hence source regions. The NO/NO\textsubscript{2} partitioning also varies between sites. For example, NO\textsubscript{3} at Poker Flat is clearly dominated by NO\textsubscript{2}. The data from Ny-Ålesund reflect a more marine signature, with generally low mixing ratios of NO, but relatively high NO\textsubscript{2}. Mixing ratios of NO at Alert clearly follow the increasing intensity of solar radiation through polar sunrise, and achieve higher mixing ratios during April than measured at Ny-Ålesund. At Summit, mixing ratios of both NO and NO\textsubscript{3} show little variability either within a year (spring to summer) or from year to year. Snowpack emissions are a significant source of NO\textsubscript{3} at this remote location on the Greenland plateau (Honrath et al., 2002). The remote coastal Antarctic sites (Neumayer and Halley) have the lowest NO\textsubscript{3} mixing ratios of all the snow-covered sites, reaching maxima on the order of only a few 10 s of pptv or less – nonetheless higher than originally anticipated, and higher than can be achieved in models that neglect snow photochemistry. Neumayer has very low NO throughout the entire year; as expected NO is < 5 pptv (typical instrument detection limits) during the polar night, with a signal appearing during October. The annual maximum appears (in this year) in early December coincident with that of UV-B radiation, rather than with UV-A, suggesting a source driven by photolysis of nitrate in snow rather than photolysis of NO\textsubscript{2} (Weller et al., 2002, see also Sect. 3).

The real surprises regarding N-oxides in polar regions have come from South Pole where summertime mixing ratios of several 100 s pptv of NO have been measured routinely during several campaigns (e.g. ISCAT 98 (Davis et al., 2001); ISCAT 2000 (Slusher et al., 2002; Davis et al., 2004; Huey et al., 2004); ANTCI 2003 (Oltmans et al., 2007)). In 2003, mixing ratios of NO reached an extraordinary 1 ppbv (nmol mol\textsuperscript{-1}). These exceedingly high NO\textsubscript{3} concentrations are highly correlated with atmospheric stability and are believed to result from snowpack emissions across the Antarctic plateau which are concentrated within a very shallow boundary layer (Davis et al., 2004). Mixing ratios of NO\textsubscript{3} at South Pole are high enough for local production of ozone (Crawford et al., 2001), a phenomenon normally associated with polluted atmospheres. Indeed, O\textsubscript{3} enhancements of up to 25 ppbv have been observed (Helmig et al., 2007a). In addition to NO\textsubscript{3}, very substantial concentrations (10’s pptv) of gas phase HNO\textsubscript{3} and H\textsubscript{2}O\textsubscript{2} are measured at this site.

**Diurnal variation in oxidized nitrogen compounds:** Photochemistry occurring within snow and subsequent emissions from the snowpack also drive short timescale variation in boundary layer composition. For example, during the

Polar Sunrise 1998 experiment at Alert, NO\textsubscript{X} showed a clear diurnal variation whose amplitude increased as day length increased (Ridley et al., 2000). Daytime emissions of NO\textsubscript{X} from photolysis of snow NO\textsubscript{3} with formation and fast redeposition of HNO\textsubscript{3} explained this pattern. Surprisingly also, a diurnal variation in NO\textsubscript{Y} was measured at Neumayer (Fig. 7) (Weller et al., 1999). The variation was found to be linked both to exchanges at the snow surface and to changing stability of the boundary layer (Weller et al., 1999). When the boundary layer was shallow and highly stable, NO\textsubscript{Y} mixing ratios were elevated, a relationship that has since also been observed at South Pole, where the highest NO levels were recorded under calm and shallow boundary layer conditions. Davis et al. (2001, 2004) suggested that the low boundary layer height was essential, but not sufficient, to explain the huge enhancements of NO at South Pole compared to other sites.

Fig. 6. Overview of recent NO\textsubscript{X} measurements from high latitude sites. References as for Fig. 5. South Pole NO\textsubscript{X} are based on calculated NO\textsubscript{2} values and measured NO values.
The diurnal variation in NO$_3$ at Neumayer is a composite of diurnal variations in component species. Figure 7 shows that the cycle of NO is clearly well offset from the maximum in NO$_2$ photolysis, evidence of a reduced role for this source at Neumayer. Some interesting new data from Summit show that the evolution of component species is not linear throughout the season from the early spring and into summer, as one might intuitively expect (J. Dibb, personal communication). Summit early spring has large amplitude diurnal variation in NO; no equivalent signal for HONO is evident. By late spring, mixing ratios of HONO have increased and generally track NO and solar elevation. Summertime data collected during the previous year at Summit show NO and HONO varying in tandem. The data suggest that in the early spring, different mechanisms are determining the mixing ratios of NO and HONO, but by the summer, the mixing ratios are being driven by the same (or strongly associated) processes. It is known that HONO evolution out of the snowpack is very different from that of NO, and HONO decomposition leads to NO generation, issues that are further explored below.

4.1.4 Mechanisms for the production of NO, NO$_2$ and HONO

The mechanisms driving the photochemical production of NO$_x$ and HONO in snow have been investigated in controlled studies, mainly in the laboratory but also in the field. For the laboratory studies, a critical point is whether the distribution of NO$_3^-$ in the artificially-generated snow/ice matches that of natural snow, where NO$_3^-$ is probably mostly very close to the ice crystal surface. Another factor to bear in mind is that reactant mixing ratios in some laboratory studies are so much higher than in nature that the relevance of proposed mechanisms in real snow may be questioned. One feature that all the studies reported thus far have in common, however, is that the reactions occur in the QLL or “micropockets” within the ice structure, as discussed in some detail in Sect. 2 of this paper.

The initial studies addressed whether NO$_3^-$ impurities in snow really could produce NO$_x$ when irradiated. Honrath et al. (2000a) made artificial snow by spraying a solution of deionized water doped with NaNO$_3$ into liquid nitrogen. The rapidly frozen “snow” was irradiated and NO$_x$ was indeed detected. Submillimeter ice layers doped with NO$_3^-$ emitted NO$_2$ when continuously irradiated at $\lambda \sim 300$ nm (Dubowski et al., 2001). Laboratory experiments using blocks of Antarctic snow found that production of NO and NO$_2$ ceased at $\lambda > 345$ nm (Cotter et al., 2003). The wavelength dependence of NO$_x$ production in these block studies is consistent with absorption by NO$_3^-$ in aqueous solution (maximum absorption around 300 nm and none above 345 nm). A study to quantify the temperature-dependence of the NO$_3^-$ quantum yield found that the same temperature dependence described results both in solution and in ice, suggesting that photolysis of NO$_3^-$ on ice occurs in the QLL rather than in the bulk ice (Chu and Anastasio, 2003).

Nitrate photolysis in the aqueous phase at wavelengths above 290 nm is classically considered to proceed via two channels:

$$\text{NO}_3^- + h\nu \rightarrow \text{NO}_2 + \text{O}^-$$  \hspace{1cm} (4)

$$\text{NO}_3^- + h\nu \rightarrow \text{NO}_2^- + \text{O}(^3\text{P})$$  \hspace{1cm} (5)

The overall quantum yields for these two channels is roughly 0.01, i.e. only 1% of the photons absorbed lead to products. It appears from two laboratory studies, one studying the aqueous phase (Warneck and Wurzinger, 1988) and the other ice surfaces (Dubowski et al., 2001), that channel 4 exceeds channel 5 by roughly a factor of 8 to 9. A further possible channel in this system results in production of the peroxynitrite ion, OONO$^-$. Although the quantum yield at 254 nm is around 0.1, there is good evidence that the quantum yield decreases significantly with increasing wavelength, and it is unclear whether this channel exists for $\lambda > 280$ nm (see, e.g. Mack and Bolton, 1999). Even if it does exist, any peroxynitrite formed on snow may still not be significant; given that the pKa for HOONO is 6.5. Thus any OONO$^-$ formed will most likely be rapidly protonated to HOONO, whose major fate appears to be very rapid decay to NO$_3^-$ ($\tau \sim 1$ s), so that most OONO$^-$ probably returns back to HNO$_3$.

Channel 5 can be followed by the photolysis of nitrite (NO$_2^-$) via:

$$\text{NO}_2^- + h\nu \rightarrow \text{NO} + \text{O}^-$$  \hspace{1cm} (6)

such that photolysis of NO$_3^-$ can generate NO as a secondary product. Alternatively, NO$_2^-$ can react with oxidants such as ozone or OH:

$$\text{NO}_2^- + \text{OH} \rightarrow \text{NO}_2 + \text{OH}^-$$  \hspace{1cm} (7)
which provides another route for the formation of NO2 (Jacobi and Hilker, 2007; Chu and Anastasio, 2007). Near midday in summer the calculated lifetime of NO2 on polar surface snow is quite short (on the order of several hours), resulting in low estimated snow grain concentrations on the order of 10 nmol kg⁻¹ or less (Chu and Anastasio, 2007).

The dominant product from NO3 photolysis is therefore gaseous NO2, a result that is supported by many field observations which have found NO2 production to noticeably exceed that of NO (e.g. Jones et al., 2000; Dibb et al., 2002). The experiments of Dubowski et al. (2001) suggest, however, that not all of the NO2 is released from the snow, rather only NO2 produced near the ice crystal-air interface is released to the air, possibly then reaching the underlying boundary layer. The rest undergoes secondary chemistry (dark and photochemistry), a result supported by Boxe et al. (2005).

Various mechanisms have been proposed for HONO formation. The pH of melted present day fresh snow is acidic except in regions with strong inputs of dust or sea salt. If we assume that acid/base equilibria known for liquid water can be applied to snow (a hypothesis that is somewhat uncertain), it follows that NO3⁻ in snow can be protonated to produce HONO which will be released into the gas phase:

\[ \text{NO}_3^- + H^+ \rightarrow \text{HONO} \]  

(8)

Under sufficiently acidic conditions, the nitroacidium ion, H2ONO⁺ (pKw=1.7) may also form (Hellebust et al., 2007), which could then react further to produce HONO.

In addition to Reaction (5), another proposed source of NO3⁻ involves the hydrolysis of photo-generated NO2 (Zhou et al., 2001; Boxe et al., 2005), via:

\[ 2\text{NO}_2 + \text{H}_2\text{O} \rightarrow \text{NO}_2^- + \text{NO}_3^- + 2\text{H}^+ \]  

(9)

These authors also suggest the heterogeneous reaction NO + NO2 + H2O → 2 HONO might be significant. McCabe et al. (2005) suggest extensive cage recombination of primary photofragments with the water solvent in the photolysis of NO3⁻, consistent with the proposed mechanisms. However, the concentrations of reactants needed for these reactions are considerably higher than are found in nature so these processes are probably not very likely.

Other mechanisms have also been suggested to produce HONO within snow interstitial air. One example is the reaction of NO2 (produced from NO3⁻ photolysis) with specific photosensitized organics (George et al., 2005; Stemmler et al., 2006). It is not known whether such organic molecules are sufficiently widely found in surface snow to be influential. However, such reactions have been invoked to explain variations in HONO production from snow in coastal Antarctica (Beine et al., 2006), where high concentrations of impurities were found in snow, and where the proximity of the Ross sea polynya could have supplied appreciable amounts of various organic molecules. Certainly humic substances and other plant degradation material are widely found in the Arctic snowpack, as discussed in Sect. 4.4.1. Of note also is that both Reactions (4) and (6) produce O⁻, which will be rapidly protonated to form OH, which may then react with NO to produce HONO:

\[ \text{O}^- + \text{H}^+ \rightarrow \text{OH} \]  

(10)

\[ \text{NO} + \text{OH} \rightarrow \text{HONO} \]  

(11)

However, this pathway is unlikely to be a significant source of HONO since snow grain concentrations of both NO and OH will be quite small. An extensive discussion of HONO formation mechanisms is presented by Cotter et al. (2003) and Jacobi and Hilker (2007). Of particular relevance for the overall discussion here, Jacobi and Hilker (2007) point out that, under natural conditions, the photolysis rates of NO3⁻ in snow are relatively small. As a result, the production rates of the short-lived compounds (such as NO) are also very small, which reduces the likelihood of the possible side and cross-reactions that can be detected under laboratory conditions.

Temperature, pH and ionic content of natural snow will also affect many reactions, and additionally determine whether products are released. Jacobi and Hilker (2007) suggest that direct formation of HONO is highly dependent on the pH of the QLL, with effectively no production at pH\geq5 since the pKa of HONO is 2.8 in solution (Riordan et al., 1995). This is consistent with the field measurements of Beine et al. (2003, 2005) and Amoroso et al. (2005) who found no HONO production in alkaline snow.

4.1.5 Establishing a modeling framework

Irrespective of the mechanism, laboratory and field experiments indicate that NOx production in snow approximates that expected from aqueous photolysis of NO3⁻, extrapolated to subfreezing temperatures (Wolff et al., 2002; Chu and Anastasio 2003; Jacobi and Hilker, 2006). The production rate should be proportional to the concentration of “available” NO3⁻ in snow and the photolysis frequency. The emission of products will be influenced by the microstructural location of NO3⁻, which is influenced by its chemical form (acid or salt) (Beine et al., 2003, 2006). For snow NO3⁻ inventories dominated by HNO3, the NO3⁻ must rapidly reach the surface of the snow crystal, either through initial deposition to the surface or by relatively fast diffusion (Tibert and Domine, 1998), since a very high proportion of it can be lost through physical processes such as volatilization (Röthlisberger et al., 2000). This might not be the case for NO3⁻ trapped as (e.g. Na⁺ or Ca²⁺) salts. This issue is important for sites near the ocean or dust sources, and in other climate regimes such as those prevailing during the last glacial period.

The photolysis frequency can be calculated (Wolff et al., 2002) from the downwelling spectral irradiance at the snow surface, the properties of the snow that determine the actinic flux as a function of depth and wavelength, the absorption
cross-section of aqueous nitrate (Mack and Bolton, 1999), and the quantum yield, which has recently been measured in ice (Chu and Anastasio, 2003) (see Sect. 3 for further details.) Snow temperature is required because the quantum yield is temperature dependent (Chu and Anastasio, 2003). As an indication of the importance of different factors in this calculation, the calculated NO$_3^-$ production (other factors being unchanged) will increase by around a factor of 6 between SZA of 80° and 60°, emphasizing the potential importance of low latitude emissions. The production rate increases by around 1% per 100 m of altitude, and by around 25% at an ozone column of 200 DU compared to 300 DU (i.e. under stratospheric ozone depletion conditions). The quantum yield and production rate will about double at 273 K compared to 253 K. Of course, if the light penetration is doubled, then production rates will also double.

The largest uncertainty is caused by variability in the snow NO$_3^-$ concentration. The map (Fig. 8) shows our estimate of concentrations for important snow-covered regions; where we are aware of a strong seasonality in concentration we use summer values because that is when photolysis occurs.

4.1.6 Sources of snowpack nitrate

**What do isotopic studies tell us of snowpack NO$_3^-$ sources?** The isotopic composition of snowpack NO$_3^-$ should reveal whether photochemical loss is important in the overall budget of snowpack NO$_3^-$ (Freyer et al., 1996) showed that nitrogen isotope composition in Antarctic NO$_3^-$ was closely related to snow accumulation rate, with lower $^{15}$N at higher accumulation sites. This result was later confirmed for Greenland ice (Hastings et al., 2005). For high accumulation sites, Hastings et al. (2004) concluded that $^{15}$N and $^{18}$O of NO$_3^-$ are related to atmospheric sources/processes, in particular NO$_x$ oxidation chemistry, rather than post depositional effects, a result also suggested by other studies (Alexander et al., 2004; Heaton et al., 2004). For low accumulation sites such as Dome C, post-depositional processes profoundly modify the concentration and isotopic composition of snow NO$_3^-$ (Freyer et al., 1996; Blunier et al., 2005). Comparison with the fractionation constant obtained in laboratory photolysis experiments (Blunier et al., 2005) appears to rule out photolysis in the surface snow as the main process leading to changes in NO$_3^-$ isotopic composition, in agreement with calculations that found photolysis could account for up to just 40% (usually less) of observed losses of NO$_3^-$ from Antarctic snow (Wolff et al., 2002). It seems that photochemical production of NO$_x$ from snow NO$_3^-$ is more important for boundary layer chemistry than it is for the budget of NO$_3^-$ in polar snow and ice. Interestingly, in a recent field study combining the collection of year-round aerosols, surface snow, and snow pit samples at South Pole, McCabe et al. (2007) found strong isotopic evidence for a dominant stratospheric source of NO$_3^-$ in winter aerosol and surface snow, but a much stronger tropospheric signature in NO$_3^-$ in the snowpack. They hypothesized that photolysis of the stratospheric NO$_3^-$ produced NO$_x$ which reformed HNO$_3$ (and we note would also likely produce HNO$_2$NO$_2$, Slusher et al., 2002) with tropospheric $^{17}$O signature and redeposited. The recycled (photochemical) NO$_3^-$ was suggested to dominate preserved NO$_3^-$ throughout the 10-year record in the pit, with a larger fraction of recycled NO$_3^-$ seen in years with greater O$_3$ depletion, hence enhanced UV flux in spring and early summer.

**What do NO$_x$ budget studies tell us of snowpack NO$_3^-$ sources?** Various studies have addressed the budget of NO$_x$ at high latitudes. Such studies by definition include numerous measurements, so have been conducted with varying degrees of coverage. Surface snow nitrate exhibits a summertime peak; so, if deposition occurs close to the ground (as opposed to being scavenged by snow aloft and then deposited), there should be a link to the NO$_3^-$ component species listed in Table 1. Uptake would be controlled both by the mixing ratio and the air/snow partitioning of the NO$_x$ constituent, as described in more detail below. There is no consistent story of any one NO$_x$ component dominating over the others across the polar regions where these measurements have been made.

Recent measurements from Halley during the CHABLIS campaign show an interesting contrast between summertime and wintertime NO$_x$ (Jones et al., 2007). During summer (December), the distribution of inorganic (68%) vs organic

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**Fig. 8.** Estimates of snow nitrate concentrations ($\mu$g kg$^{-1}$) for different snow-covered regions. See original references for details. Antarctica and sea ice zone (Mulvaney and Wolff, 1994) (much higher values may be found in the very surface layer in central Antarctic (Röthlisberger et al., 2000) and in coastal regions, where sea salt and mineral aerosols efficiently scavenge nitric acid (Beine et al., 2006)); Greenland and adjacent Arctic islands (Röthlisberger et al., 2002; Koerner et al., 1999); North America: maps at National Atmospheric Deposition Program (NADP) (http://nadp.sws.uiuc.edu/isopleths/annualmaps.asp); Alps (summer concentrations) (Preunkert et al., 2003); rest of Europe: EMEP (http://www.nilu.no/projects/ccc/emepdata.html); Himalayas (Hou et al., 1999); other regions by analogy. The uncertainty on these values due to extrapolation from specific sites is at the very least a factor 2, and this range has to be explored in sensitivity studies.
(32%) NO$_y$ components is quite different than during winter (July) (13% inorganic vs 87% organic). The seasonal variation of NO$_3^-$ concentration in surface snow closely tracks the sum of (HONO + HNO$_3$ + p-NO$_3^-$) in the air and bears no resemblance to the behavior of organic NO$_y$. Which drives what, however, is not yet fully resolved. Some light may be shed by recent observations of oxygen and nitrogen isotopic composition of inorganic aerosol nitrate (p-NO$_3^-$ plus a significant fraction of the inorganic acids) collected on filters (Savarino et al., 2006). Like the oxygen isotopes in NO$_3^-$ at South Pole (McCabe et al., 2007) discussed earlier, these data suggest late winter deposition of NO$_2^-$ from polar stratospheric cloud (PSC) subsidence (in agreement with earlier work by Wagenbach et al., 1998), but a late spring concentration peak in recycled inorganic NO$_3^-$ species driven by snowpack emission of NO$_x$ inland (upwind).

4.1.7 Air-ice partitioning of relevant NO$_y$ species

Assuming photoproducts are created at the ice surface, or in the QLL at the surface, and not in a cage within the ice structure, their release to the firm air depends mainly on their affinity for the ice surface. Much of the published data refer to bulk aqueous solutions, with gas-liquid equilibria described by Henry’s Law. Both NO and NO$_2$ are only weakly soluble in water and interact weakly with ice (e.g. Cheung et al., 2000; Bartels-Rausch et al., 2002). It is likely that they will be lost to firm air before they undergo reaction on the ice surface or in the QLL, as discussed by Jacobi and Hilker (2007).

The acidic gases, HNO$_3$ and HO$_2$NO$_2$, and to a lesser extent HONO, have been shown to be much more strongly adsorbed on ice surfaces (Bartels-Rausch et al., 2002; Huthwelker et al., 2006), so that molecules formed in, or advected to, the firm layer can be adsorbed on ice. Partitioning of the acids between air, ice surface (and/or QLL), and ice matrix is a coupled process of adsorption and bulk diffusion, as described in more detail in the accompanying snow physics paper (Dominé et al., 2007). These processes depend strongly on the presence of other acids, since these affect whether the partitioning species is present as an acidic molecule or an ion (e.g., HNO$_3$ or NO$_3^-$). Finally we note that Henry’s Law coefficients for PAN and methyl nitrate, which constitute an important part of the NO$_y$ family at some locations and times of the year, are only an order of magnitude smaller than for HONO. Some net uptake for these molecules by snow grains might therefore be expected (Ford et al., 2002; Dassau et al., 2004).

4.1.8 Fate of NO$_x$ released to snowpack interstitial air/boundary layer

The production of NO$_x$ within snowpack interstitial air has the potential to influence the chemistry of the overlying atmosphere and also concentrations of NO$_3^-$ (and other compounds) in surface snow and glacial ice. The extent to which this potential is realized depends on the fate of the snowpack NO$_x$. In order for NO$_x$ produced by photochemistry in snow to impact the wider troposphere, it must first escape the snowpack and then escape the near-snow boundary layer. This involves competition between vertical mixing, which is often quite weak over snow covered surfaces (e.g. Munger et al., 1999; Honrath et al., 2002; Oncley et al., 2004; Cohen et al., 2007), and reactions between NO$_3$ and HO$_x$ forming HNO$_3$ and HO$_2$NO$_2$ which redeposit to the snow fairly rapidly. There is abundant evidence supporting significant production and rapid deposition of both acids at South Pole, with lifetimes against deposition on the order of a few hours (Chen et al., 2001; Slusher et al., 2002; Huey et al., 2004; Dibb et al., 2004). In the Arctic little is known about HO$_2$NO$_2$, but it is equally clear that a significant fraction of emitted NO$_x$ reforms HNO$_3$ very close to the snow surface, and much of this is redeposited (Dibb et al., 1998; Munger et al., 1999; Ridley et al., 2000). Of course, any HNO$_3$ and HO$_2$NO$_2$ deposited onto the surface can be photolyzed again, setting up a cycle. The key question is whether this cycle is closed, or leaks some of the NO$_x$ emitted by the snow to higher levels in the atmosphere (note that the NO$_x$ can be exported as NO$_x$ or any of the three acids; HONO, HO$_2$NO$_2$, HNO$_3$, with the first two being rapidly photolyzed in turn to release NO$_x$ again). It has been suggested that even if the NO$_x$ to acid to snow to NO$_x$ cycle is nearly closed, advective transport during the few hours before the acids redeposit could export NO$_x$ emitted from snow off the east Antarctic plateau in the drainage flow (Davis et al., 2006).

Honrath et al. (2002) found that upward fluxes of NO$_x$ plus HONO were larger than the downward fluxes of HNO$_3$ at Summit during summer 2000, suggesting that there is net export of NO$_x$ emitted by snow to the free troposphere over Greenland. On the other hand, investigation of the N and O isotopes of NO$_3^-$ in snow at Summit found diurnal variations consistent with daytime losses due to photolysis, but redeposition of NO$_3^-$ (as HNO$_3$) at night restored the isotopic ratios (Hastings et al., 2004). Over seasonal and annual timescales the net impact of snow photochemistry on the isotopic composition of NO$_3^-$ at Summit was negligible, suggesting that the cycling described above has to be nearly closed. At South Pole it appears quite certain that NO$_x$ from the snow causes enhanced O$_3$ production in the lower several hundred meters of the atmosphere (Crawford et al., 2001; Helming et al., 2007a), suggesting that there has to be some loss of NO$_x$ upward out of the boundary layer. In contrast to the Greenland results, isotopic studies in the Antarctic show that post depositional effects strongly influence the isotopic signature of the remaining snow nitrate (Blunier et al., 2005; McCabe et al. 2007) as well as the isotopic composition of the filterable NO$_3^-$ collected at coastal sites (Wagenbach et al., 1998; Savarino et al., 2006). Observed O and N isotopic fractionations provide strong support for extensive recycling, and appear to be compatible with export of snowpack NO$_x$ from the central plateau to coastal sites.
4.2 Oxidants

4.2.1 Expectations in the absence of snow-atmosphere fluxes

In the troposphere the most important oxidant is the hydroxyl radical (OH). The main source of OH is the reaction of O(1D) + H2O, with photolysis of O3 producing O(1D). As noted in Sect. 2, the global distribution of UV-B radiation results in greatly reduced rates of O3 photolysis at high latitudes compared to the tropics. Combining this with a similarly steep gradient in the abundance of water vapor between the tropics and polar regions leads to the expectation that the production and abundance of OH in the remote troposphere should be greatest in the tropics and least in the polar regions.

This view was consistent with the first Antarctic OH observations conducted during late February 1994 on the coast at Palmer Station (on Anvers Island off the Antarctic Peninsula) as part of the Sulfur Chemistry in the Antarctic Troposphere Experiment (SCATE) (Berresheim and Eisele, 1998).

Using the selected ion chemical ionization mass spectrometry (SICIMS) technique, 24-h and daytime average values for OH were 1.1×105 and 3.0×105 molecule cm−3, respectively (Jefferson et al., 1998). These very low values were attributed to the high average solar zenith angle, extensive cloud cover, and low levels of NO (1–5 pptv). Comparison with models was hampered by uncertainty in the levels of NO, which were below or similar to the instrumental detection limit of 5 pptv. However, by assuming NO levels near this detection limit, modeled and observed OH agreed to within ~30%, with OH production dominated by the reaction O(1D) + H2O, and loss dominated by reaction with CO and CH4. The results are those expected for an unperturbed remote pristine environment at high latitudes, and can be used as a base case in the absence of snowpack emissions.

4.2.2 Recent findings at snow-covered sites: South Pole

Surprisingly, average OH values of 2×105 molecule cm−3 were measured at South Pole (November-December 1998).
Table 2. Mean values of selected parameters measured at South Pole during the 3 recent photochemistry campaigns.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>ISCAT 98 NOV</th>
<th>ISCAT 98 DEC</th>
<th>ISCAT 00 NOV</th>
<th>ISCAT 00 DEC</th>
<th>ANTCI 03 NOV</th>
<th>ANTCI 03 DEC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temp °C</td>
<td>−31.4±4.6</td>
<td>−29.0±2.1</td>
<td>−29.3±2.7</td>
<td>−30.8±1.6</td>
<td>−38.0±4.9</td>
<td>−26.7±2.2</td>
</tr>
<tr>
<td>Dewpoint °C</td>
<td>−34.5±4.9</td>
<td>−31.9±2.1</td>
<td>−33.3±2.9</td>
<td>−30.8±1.6</td>
<td>−38.0±4.9</td>
<td>−26.7±2.2</td>
</tr>
<tr>
<td>W Speed m s⁻¹</td>
<td>6.3±2.4</td>
<td>3.5±1.2</td>
<td>6.1±1.8</td>
<td>4.2±1.6</td>
<td>4.9±1.3</td>
<td>5.2±2.0</td>
</tr>
<tr>
<td>Total O₃ DU</td>
<td>174±24</td>
<td>264±19</td>
<td>282±23</td>
<td>301±16</td>
<td>230±25</td>
<td>287±23</td>
</tr>
<tr>
<td>J(O(¹D)) 10⁻⁶ s⁻¹</td>
<td>8.6±1.3</td>
<td>5.8±1.8</td>
<td>7.6±1.2</td>
<td>12.2±2.2</td>
<td>11.5±1.9</td>
<td></td>
</tr>
<tr>
<td>J(NO₂) 10⁻¹² s⁻¹</td>
<td>0.95±0.01</td>
<td>0.87±0.15</td>
<td>1.03±0.12</td>
<td>1.1±0.1</td>
<td>1.2±0.1</td>
<td></td>
</tr>
<tr>
<td>NO pptv</td>
<td>111±7.4</td>
<td>117±8.6</td>
<td>117±10</td>
<td>107±10</td>
<td>107±10</td>
<td></td>
</tr>
<tr>
<td>O₃ pptv</td>
<td>116±17</td>
<td>116±17</td>
<td>116±17</td>
<td>116±17</td>
<td>116±17</td>
<td></td>
</tr>
<tr>
<td>CO pptv</td>
<td>485±116</td>
<td>393±173</td>
<td>213±10</td>
<td>213±10</td>
<td>213±10</td>
<td></td>
</tr>
<tr>
<td>CH₂O pptv</td>
<td>282±22</td>
<td>200±40</td>
<td>200±40</td>
<td>200±40</td>
<td>200±40</td>
<td></td>
</tr>
<tr>
<td>H₂O₂ pptv</td>
<td>268±111</td>
<td>278±67</td>
<td>278±67</td>
<td>278±67</td>
<td>278±67</td>
<td></td>
</tr>
<tr>
<td>OH 10⁶ molecule cm⁻³</td>
<td>1.8±0.9</td>
<td>2.5±0.6</td>
<td>2.4±1.0</td>
<td>1.5±0.6</td>
<td>1.7±0.9</td>
<td></td>
</tr>
<tr>
<td>HO₂+RO₂ 10⁷ molecule cm⁻³</td>
<td>8.3±2.4</td>
<td>8.3±2.4</td>
<td>8.3±2.4</td>
<td>8.3±2.4</td>
<td>8.3±2.4</td>
<td></td>
</tr>
<tr>
<td>HONO (MC) pptv</td>
<td>27±3</td>
<td>72±21</td>
<td>30±4</td>
<td>30±4</td>
<td>30±4</td>
<td></td>
</tr>
<tr>
<td>HONO (LIF) pptv</td>
<td>32±11</td>
<td>7.4±4.2</td>
<td>5.3±2.5</td>
<td>5.3±2.5</td>
<td>5.3±2.5</td>
<td></td>
</tr>
<tr>
<td>HNO₂ (MC) pptv</td>
<td>24±11</td>
<td>42±15</td>
<td>23±5</td>
<td>23±5</td>
<td>23±5</td>
<td></td>
</tr>
<tr>
<td>HNO₃ (CIMS) pptv</td>
<td>19±11</td>
<td>107±53</td>
<td>86±78</td>
<td>86±78</td>
<td>86±78</td>
<td></td>
</tr>
<tr>
<td>HO₂NO₂ pptv</td>
<td>23±11</td>
<td>47±10</td>
<td>39±1</td>
<td>39±1</td>
<td>39±1</td>
<td></td>
</tr>
</tbody>
</table>

Fig. 9. OH versus NO at South Pole.

during the Investigation of Sulfur Chemistry in the Antarctic Troposphere (ISCAT) campaign (Mauldin et al., 2001). More than an order of magnitude greater than SCATE observations, and equivalent to tropical values, the high OH levels at South Pole mostly result from unexpectedly high NO levels, leading to an intensely oxidizing environment. Diurnal average values for key photolysis frequencies were comparable to equatorial values due to the high albedo (~0.8) and 24 h sunlight conditions (Lefer et al., 2001). In fact, 24 h average values for J(NO₂) (1.1×10⁻² s⁻¹) were 3 times greater than for equatorial conditions while J(O(¹D)) (9.0×10⁻⁶ s⁻¹) was roughly equivalent. Given the dry conditions at South Pole, which reduce the conversion efficiency of O(¹D) from O₃ photolysis into OH, primary production from O₃ photolysis could not explain the observed OH, but ambient NO levels of 225 pptv (median) ranging as high as 600 pptv, provided a strong secondary source of OH (Chen et al., 2001), via the reaction HO₂ + NO → OH + NO₂.

Elevated OH and NO were again observed during the ISCAT 2000 and ANTCI 2003 campaigns, which also occurred in November and December. Measurements from these three field campaigns yield an overall average OH of 2.0(±0.9)×10⁶ molecule cm⁻³ and a median of 1.9×10⁶ molecule cm⁻³ (Mauldin et al., 2001, 2004) (Table 2). The average NO mixing ratio is 187 (±175) pptv and the median value is 122 pptv. While NO values have varied considerably among years (Table 2, Fig. 5), periods with NO levels of several hundred pptv were observed in all years. Average HO₂+RO₂ concentrations measured during more limited periods in 2000 were 8.3 (±2.4)×10⁷ molecule cm⁻³, with a median of 8.1×10⁷ molecule cm⁻³.

Figure 9 displays the relationship between observed OH and NO for all South Pole observations. Peaks in observed OH occur between 70 and 300 pptv of NO. These peak OH values, however, vary by a factor of 3. A similar OH dependence on NO was also seen in the first modeling study of ISCAT 1998 data by Chen et al. (2001). The rapid increase in OH with increasing NO levels on the left of the peak is due to increasing HO₂ to OH conversion by NO, shifting the HO₂ partitioning in favor of OH. Some contribution also comes from enhanced HO₃ production from CH₄ oxidation. The reduction in OH concentration with increasingly high NO levels beyond the peak can mainly be attributed to HO₂ loss via formation of HNO₃ and HO₂NO₂ followed by deposition onto the snow surface. While models including only gas phase chemistry underpredict observed OH, inclusion of HO₂ precursors emitted from the snow during ISCAT 2000 improved model predictions and confirmed that snow
emissions of H$_2$O$_2$ and CH$_2$O are the dominant HO$_x$ source at South Pole (Chen et al., 2004; Hutterli et al., 2004). Mist chamber measurements of HONO ($\sim$30 ppbv average) (Dibb et al., 2004), another important HO$_x$ precursor emitted from the snow, were less encouraging. When these HONO measurements were used in model calculations, predicted values of boundary layer OH were 2–5 times greater than observations. These HONO results were also incompatible with ambient NO$_x$ concentrations given the abundance and very short lifetime of HONO (Chen et al., 2004). As for other polar sites, measurements of HONO are difficult to reconcile with photochemical observations of HO$_x$ and NO$_x$, raising questions about the specificity of the mist chamber measurements (Sjostedt et al., 2005).

Two important and observable consequences result from the intense photochemistry at South Pole. One is the potential for large O$_3$ production rates. Modeling studies of ISCAT 1998 and 2000 showed a net production of $\sim$3–4 ppbv/day. These prompted a reevaluation of historical ozone observations both at the surface and from ozonesondes at South Pole which revealed strong evidence for a surface source of ozone during Austral spring/summer (Crawford et al., 2001; Oltmans et al., 2007). Tethered balloon observations during ANTCI 2003 provided the strongest evidence yet for near-surface ozone production with frequent observations of enhanced ozone (20–25 ppbv) over depths of 200+ m (Helmig et al., 2007a). O$_3$ measurements during the US ITASE traverse between Byrd and South Pole in summer 2002/03 showed up to 2-fold increases of near-surface mixing ratios at sampling locations above 2000 m elevation and indicated that enhanced O$_3$ production is spatially limited to the Antarctic plateau region (Frey et al., 2005). However, ozone can be transported long distances and analysis of surface ozone data from six Antarctic stations gave indications that sites on the exterior of the Antarctic continent are, at least occasionally, influenced by transport of ozone-enriched air from the interior of Antarctica (Helmig et al., 2007b). A consequence is the hypothesized presence of an oxidizing canopy of OH enshrouding the Antarctic plateau (Davis et al., 2004). Observations of NO and NO$_x$ from a Twin Otter aircraft during ANTCI 2003 revealed elevated NO over depths of 500 m and distances of 400 km from South Pole, thus, similar to the conclusions derived from the ozone observations by Frey et al. (2005) demonstrating that the photochemical conditions at South Pole may extend across a large portion of the Antarctic plateau.

4.2.3 Recent findings at snow-covered sites: Halley, Antarctica

The Chemistry of the Antarctic Boundary Layer and the Interaction with Snow (CHABLIS) field campaign took place on the floating Brunt Ice Shelf at Halley and consisted of a year-round study (January 2004–February 2005) and a summer intensive (January–February 2005) (Jones et al., 2005). CHABLIS was the first intensive chemistry field campaign above the snowpack in coastal Antarctica. At the time of writing the work is very recent, with final analysis and modeling still in progress, hence our discussion is brief and qualitative, with most references from published conference proceedings. Peak (daily maxima) OH and HO$_2$ concentrations (measured using laser-induced fluorescence, Heard and Pilling, 2003) varied between 0.9–3.0 $\times$ 10$^6$ and 2.5–9.3 $\times$ 10$^7$ molecule cm$^{-3}$ (1–4 pp), respectively (Bloss et al., 2007). The concentrations of both species declined as the campaign progressed. Despite being at higher latitude, the peak OH concentrations for Halley are considerably higher ($\sim$3–4 times) than observed during SCATE at the Palmer station at a similar time (February) (Jefferson et al., 1998), but are lower than observed at South Pole (in November/December). Airmass back trajectories indicated flow mainly from the Antarctic continent, although on occasion the origin was the Southern Ocean. The site experienced 24 h daylight during the intensive period, however J(O$^1$D) was a factor of 40–50 lower during the “night” compared with the maximum at solar noon. The diurnal profiles for both OH and HO$_2$ are highly distinct, following closely, but not exactly, that of J(O$^1$D). “Night-time” OH was observed above the detection limit (1.5 $\times$ 10$^5$ molecule cm$^{-3}$) on several occasions, in the range $\sim$1.5–2 $\times$ 10$^5$ molecule cm$^{-3}$, and HO$_2$ was always seen “at night” in the range 0.2–1.3 $\times$ 10$^7$ molecule cm$^{-3}$. For the period 2 h either side of local noon, the ratio [HO$_2$]/[OH] was in the range 32–138, and for the entire campaign the average 24 h OH and HO$_2$ concentrations were 3.9 $\times$ 10$^5$ and 2.0 $\times$ 10$^7$ molecule cm$^{-3}$, respectively.

The modeling phase of CHABLIS is at an early stage, and thus we will only discuss here the general factors that control HO$_x$ at Halley to contrast with other Antarctic regions such as South Pole. Typical noon-time concentrations of some of the sources and sinks of HO$_x$ observed during the summer CHABLIS intensive are O$_3$ (10–15 ppbv), NO$_x$ (5–25 pptv), CO (35–40 ppbv), and CH$_2$O (130 pptv). The NO$_x$ levels are considerably higher than at Palmer station (Jefferson et al., 1998) because of snowpack emissions initiated via nitrate ion photolysis (Sect. 4.1), but considerably lower than observed at South Pole due in part to the much deeper boundary layer at Halley. Assuming photolysis of ozone was the only OH source, and reaction with CO and CH$_4$ its only sinks, calculated OH was significantly less than the measured value, even when the upwelling component of actinic flux (with a snow albedo of 0.85) was added to J(O$^1$D). This discrepancy suggests other sources of OH are required, which

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**4.2.4 Recent findings at snow-covered sites: Summit, Greenland**

Snowpack emissions of CH$_2$O, H$_2$O$_2$, and HONO (Dibb et al., 1998, 2002; Honrath et al., 1999; 2002; Hutterli et al., 1999, 2001; Jacobi et al., 2002; Yang et al., 2002) at Summit, Greenland are expected to enhance HO$_x$ levels at this site. In order to directly test the impact of snow emissions on photochemistry at Summit, campaigns were carried out in summer 2003 (July) and spring 2004 (April). Median noontime values of selected parameters are reported for summer, and early and late spring in Table 3 along with predicted values of OH and HO$_2$+RO$_2$ obtained from highly constrained photochemical models (Sjostedt et al., 2007).

During the summer 2003 campaign, high levels of OH were routinely observed (≈1×10$^7$ molecule cm$^{-3}$). These levels were more than a factor of two higher than model predictions constrained to a full set of photochemical precursors. Typical HO$_x$ measurement uncertainties of 40% are reported, so it is unlikely that the disparity is due to measurement error. Conversely, levels of HO$_2$ + RO$_2$ were found to be in excellent agreement with predictions, indicating that peroxy radical sources and sinks were well understood but that the ratio of (RO$_2$ + HO$_2$) to OH was perturbed. The HO$_x$ source for this campaign was found to be dominated by photolysis of O$_3$ and snow-emitted H$_2$O$_2$ with smaller contributions from HONO and CH$_2$O (Chen et al., 2007). The perturbation to the ratio of (RO$_2$ + HO$_2$) to OH was particularly enhanced during an extended period of high winds and blowing snow. Large increases in OH and smaller relative decreases in (RO$_2$ + HO$_2$) characterized these windy periods. Retropulse analysis for this period indicated that marine boundary layer air was rapidly transported (1–2 days) to Summit, suggesting that halogen chemistry can influence observed chemical conditions (Sjostedt et al., 2007). This point is further discussed in Sect. 4.3 below.

The spring 2004 campaign offered an opportunity to observe HO$_x$ chemistry during a period of rapidly increasing temperatures and photolysis frequencies (Sjostedt et al., 2005). OH and HO$_2$+RO$_2$ levels were observed to increase by approximately a factor of five during the month of April. The observations of OH and HO$_2$+RO$_2$ were both found to be in poor agreement with model predictions constrained to all relevant observations (i.e. NO, O$_3$, CH$_2$O, J values, dewpoint, O$_3$, H$_2$O$_2$, CH$_2$O, HONO, etc.). This disagreement was largest early in the campaign when temperatures were the lowest. Conversely, HO$_2$+RO$_2$ was in excellent agreement with model predictions constrained to all relevant observations except for HONO. These results strongly suggest that the measured levels of soluble nitrite (NO$_2^-$) can not be simply interpreted as gas phase HONO, a conclusion already drawn during ISCAT/ANTCI South Pole campaigns (Chen et al., 2004). For this reason, the spring HO$_x$ budget was analyzed assuming HONO was not a significant radical source. This analysis showed that photolysis of H$_2$O$_2$ and CH$_2$O emitted from the snow dominated the radical source early in the spring with O$_3$ photolysis becoming dominant by the end of the campaign. Low ratios of HO$_2$+RO$_2$ to OH were also observed in the spring during periods of high winds and blowing snow. Similar to summer 2003 these periods coincided with transport of marine boundary layer air to Summit.

**Table 3. Median noontime (10:00–15:00 Western Greenland Standard Time) boundary layer values from Summit 2003 and 2004 campaigns.**

<table>
<thead>
<tr>
<th>Species</th>
<th>Early Spring 2004</th>
<th>Late Spring 2004</th>
<th>Summer 2003</th>
</tr>
</thead>
<tbody>
<tr>
<td>meas. HO$_2$+RO$_2$ molecule cm$^{-3}$</td>
<td>6.0×10$^7$</td>
<td>1.9×10$^8$</td>
<td>4.1×10$^8$</td>
</tr>
<tr>
<td>pred. HO$_2$+RO$_2$ molecule cm$^{-3}$</td>
<td>6.3×10$^7$</td>
<td>2.0×10$^8$</td>
<td>4.4×10$^8$</td>
</tr>
<tr>
<td>measured OH molecule cm$^{-3}$</td>
<td>4.1×10$^5$</td>
<td>1.9×10$^6$</td>
<td>8.4×10$^6$</td>
</tr>
<tr>
<td>predicted OH$^a$ molecule cm$^{-3}$</td>
<td>4.0×10$^5$</td>
<td>1.5×10$^6$</td>
<td>3.8×10$^6$</td>
</tr>
<tr>
<td>NO (pptv)</td>
<td>18</td>
<td>19</td>
<td>18</td>
</tr>
<tr>
<td>NO$_2$(pptv)</td>
<td>14</td>
<td>18</td>
<td>23</td>
</tr>
<tr>
<td>H$_2$O$_2$(pptv)</td>
<td>191</td>
<td>622</td>
<td>1800</td>
</tr>
<tr>
<td>CH$_2$O (pptv)</td>
<td>45</td>
<td>111</td>
<td>181</td>
</tr>
<tr>
<td>HONO (pptv)</td>
<td>6</td>
<td>13</td>
<td>13</td>
</tr>
</tbody>
</table>

$^a$Denotes predictions from a photochemical model constrained to measured parameters (NO, O$_3$, H$_2$O$_2$, CH$_2$O, j values, etc.) except for nitrous acid.

$^b$NO$_2$ levels were calculated assuming steady state constrained to measured values of NO, O$_3$, photolysis frequencies, and peroxy radicals.
measurements of major precursors. Yang et al. (2002) used a photostationary state point model to estimate HO₅ for the conditions of the 1999 and 2000 Summit campaigns. Interestingly, estimated OH for summer 1999 was very similar to that observed in 2003. However, estimated (HO₂ + RO₂) values for 1999 were more than 2-fold higher than the 2003 observations. Yang et al. (2002) suggested that the HO₅ source from the photolysis of HONO, CH₂O and H₂O₂ combined was 2-3 times stronger than that from O₃ photolysis, in contrast to 2003 where O₃ was found to be the dominant precursor followed closely by H₂O₂ with only minor production from CH₂O and HONO (Chen et al., 2007).

At Alert, significant concentrations of HONO and CH₂O have been measured, with emissions of both compounds occurring from sunlit snowpack (Zhou et al., 2001; Sumner et al., 2002). Using both snow manipulation experiments and ambient air measurements, Zhou et al. (2001) found that a significant photochemical snowpack source exists for HONO, at times making it the dominant OH source in the boundary layer, implying a much greater role for OH in Arctic springtime chemistry than previously believed. Modeling results of Granas et al. (2002) simulate upwards of 1×10⁶ molecule cm⁻³ OH at Alert at springtime solar noon under typical atmospheric conditions and 7×10⁵ molecule cm⁻³ during ozone depletion events when halogen chemistry is active. In both cases, HONO was hypothesized to be the dominant OH source, followed by CH₂O photolysis. HO₅ removal was dominated by formation of HO₂NO₂ and HNO₃ (81% and 8%) during typical ambient conditions and by HO₂NO₂ and HOBr formation (40% and 45%) during ozone depletion events. Li (1994) also found that calculated OH production rates from HONO dominated both O₃ and CH₂O, with a calculated OH concentration of 3×10⁴ molecule cm⁻³ at noon on 5 April 1992. However, the authors would like to note that these Alert model predictions were not verified by actual HO₅ observations.

Summer observations during the US ITASE ground traverse confirmed snowpack emissions of H₂O₂ and CH₂O across the Antarctic Ice Sheet between 76° S and 90° S (Frey et al., 2005). Model calculations constrained by observed atmospheric hydroperoxides (ROOH), CH₂O and O₃ suggest enhanced NO and OH levels on the East Antarctic plateau, similar to what is seen at South Pole (Frey et al., 2005). However, as the Summit example illustrates, modeling results need to be validated by in situ measurements, particularly of HO₅.

4.2.6 Oxidant production and chemistry in/on snow and ice grains

Much of the past research on snow photochemistry has focused on how these reactions affect the oxidizing capacity of the atmosphere, primarily by emitting NOₓ and precursors of HO₅ from the snowpack into the overlying boundary layer. The chemistry that leads to the release of reactive gases from the snowpack is largely uncharacterized. This condensed phase chemistry includes two general types of reactions: direct photolysis (such as for nitrate) and indirect photoreactions involving a photochemically formed oxidant reacting with a trace contaminant such as an organic compound. In this section we discuss the various oxidants that are present on snow grains and their potential contributions to snowpack chemistry. Of the major oxidants involved in snow photochemistry, the most is probably known about hydrogen peroxide (H₂O₂). H₂O₂ is thermally cycled between the atmosphere and snowpack, with emission occurring during daytime when temperatures increase, and deposition occurring during the cooler night (Sigg et al., 1992; Hutterli et al., 2001). This reversible deposition of H₂O₂ has been described with a physically based atmosphere-to-snow transfer model (McConnell et al., 1997a, b; 1998; Hutterli et al., 2003). Concentrations in fresh snow reflect water-to-H₂O₂ ratios in the cloud, whereas snow exposed sufficiently long to the atmosphere will approach a temperature-dependent partitioning equilibrium analogous to Henry’s Law (Conklin et al., 1993; Bales et al., 1995). Levels of H₂O₂ in fresh snow are altered by subsequent temperature-driven recycling between the snow and the air until the snow is buried below several 10s of cm. In addition to sorption processes, snow metamorphism and wind enhanced ventilation of the top layers of the snowpack can modulate the air-snow exchange, and a net H₂O₂ loss from the snow is expected in the course of a year (Sigg and Neftel, 1988; McConnell et al., 1997a; Hutterli et al., 2001, 2003; Frey et al., 2006). Most measurements on the polar ice sheets suggest net H₂O₂ emission from the snowpack into the boundary layer in summer (Table 4). In one summer study at Summit, Greenland there was a net deposition of H₂O₂ (Jacobi et al., 2002), but this was likely a result of frequent fog events causing significant night-time deposition (Table 4). Estimates of average emission fluxes from the Arctic and Antarctic snowpack in summer vary between 10¹¹ and 10¹³ molecule m⁻² s⁻¹ (Table 4) or 10⁻⁷ and 10⁻⁹ molecule cm⁻³ s⁻¹ for an assumed BL height of 100 m. For comparison, based on one measurement from an illuminated solution of extracted particles, in situ photoformation in airborne particles provides a volumetric H₂O₂ flux of ~10⁴ molecule cm⁻³ s⁻¹ to the boundary layer at Alert, Canada in April (Anastasio and Jordan, 2004). Although the rate is relatively small, this photoformation of H₂O₂ in aerosol particles suggests that deposition of these particles to the snowpack will lead to H₂O₂ photoformation on snow grains as well (Anastasio and Jordan, 2004), which may slow down the loss rate of H₂O₂ from the snow while slightly increasing the emission fluxes. However, this idea of the photochemical formation of H₂O₂ on snow grains is in its infancy (Chu and Anastasio, 2005). Indeed, Jacobi et al. (2006) have identified the photochemical decomposition of H₂O₂ in snow. More work is needed to quantify the snow grain budget of H₂O₂.

Because it is reactive, H₂O₂ likely plays an active role in
Table 4. Average gas phase levels (range in parentheses) of atmospheric radical precursors in air above the snowpack (ambient air) and in the interstitial pore space right below the snow surface (firn air). Positive area flux values indicate net emission, while negative values are equivalent to deposition to the snowpack.\textsuperscript{a}

<table>
<thead>
<tr>
<th>Site</th>
<th>Ambient Air\textsuperscript{b} pptv</th>
<th>Firn Air\textsuperscript{c} pptv</th>
<th>Area Flux molecule m\textsuperscript{-2}s\textsuperscript{-1}</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>H\textsubscript{2}O\textsubscript{2}</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>South Pole</td>
<td>278 (23–773)</td>
<td>884 (375–1515)</td>
<td>9.5x10\textsuperscript{12} up to 2.0x10\textsuperscript{14}\textsuperscript{d}</td>
<td>Hutterli et al., 2004</td>
</tr>
<tr>
<td>90\degree S</td>
<td>5–23 Dec 2000</td>
<td>18–23 Dec 2000, 24 h total</td>
<td>5–23 Dec 2000</td>
<td></td>
</tr>
<tr>
<td>Antarctic Plateau</td>
<td>274 (59–643)</td>
<td>564 (70–3209)</td>
<td>1.4x10\textsuperscript{12} (3.6x10\textsuperscript{11}–4.3x10\textsuperscript{12})\textsuperscript{e}</td>
<td>Frey et al., 2005</td>
</tr>
<tr>
<td>West Antarctica</td>
<td>370 (530–918)</td>
<td>388 (111–692)</td>
<td>3.4x10\textsuperscript{11} (~2.0x10\textsuperscript{11}–8.7x10\textsuperscript{11})\textsuperscript{e}</td>
<td>Frey et al., 2005</td>
</tr>
<tr>
<td>Summit, Greenland</td>
<td>1400 (90–4600)</td>
<td>1630 (1000–2100)</td>
<td>4.9x10\textsuperscript{13} (4.0x10\textsuperscript{13}–5.8x10\textsuperscript{13})\textsuperscript{d}</td>
<td>Hutterli et al., 2001</td>
</tr>
<tr>
<td></td>
<td>650 (–)</td>
<td>5 June–8 July 2000</td>
<td>(–)</td>
<td></td>
</tr>
<tr>
<td><strong>CH\textsubscript{2}O</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>South Pole</td>
<td>103 (27–184)</td>
<td>747 (425–1238)</td>
<td>1.7x10\textsuperscript{12} (9.0x10\textsuperscript{11}–2.9x10\textsuperscript{12})\textsuperscript{d}</td>
<td>Hutterli et al., 2004</td>
</tr>
<tr>
<td>90\degree S</td>
<td>5–23 Dec 2000</td>
<td></td>
<td>5–23 Dec 2000</td>
<td></td>
</tr>
<tr>
<td>Dome C</td>
<td>–</td>
<td></td>
<td>1.5x10\textsuperscript{12} (7.5x10\textsuperscript{11}–3.0x10\textsuperscript{12})\textsuperscript{c}</td>
<td>Hutterli et al., 2002</td>
</tr>
<tr>
<td>75\degree S, 123\degree E</td>
<td>–</td>
<td></td>
<td>3.5x10\textsuperscript{12} (–)\textsuperscript{f}</td>
<td></td>
</tr>
<tr>
<td>Byrd</td>
<td>121 (52–169)</td>
<td>171 (130–209)</td>
<td>5–23 Dec 2000</td>
<td></td>
</tr>
<tr>
<td>Neumayer</td>
<td>(–30–700)</td>
<td></td>
<td>(–)</td>
<td></td>
</tr>
<tr>
<td>70\degree S, 8\degree W</td>
<td>March 1997–Jan 1998</td>
<td></td>
<td>(–1.5x10\textsuperscript{13}–2.5x10\textsuperscript{13})\textsuperscript{j}</td>
<td>Riedel et al., 1999; Riedel et al., 2005</td>
</tr>
<tr>
<td>Summit, Greenland</td>
<td>230 (100–450)</td>
<td>20 Jan–Dec 2000</td>
<td>(–)</td>
<td></td>
</tr>
<tr>
<td>Alert, Canada</td>
<td>(–52–690)</td>
<td>(–120–150)</td>
<td>(–)</td>
<td></td>
</tr>
<tr>
<td>83\degree N, 62\degree W</td>
<td>15 Feb–6 April 1998</td>
<td>9 March 1998</td>
<td>4.9x10\textsuperscript{13} (–)\textsuperscript{f}</td>
<td>Sumner and Shepson, 1999</td>
</tr>
<tr>
<td></td>
<td>(–450–680)</td>
<td>7 April 1998</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>(–650–750)</td>
<td>16 April 1998</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ahmeeck, MI</td>
<td>(–190–310)</td>
<td></td>
<td></td>
<td>Couch et al., 2000</td>
</tr>
<tr>
<td>47\degree N, 88\degree W</td>
<td>Jan 1999, 1 profile</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

snowpack chemistry. Correlations between H\textsubscript{2}O\textsubscript{2} and CO\textsubscript{2} trapped in air bubbles in Greenland ice cores suggest that H\textsubscript{2}O\textsubscript{2} may oxidize organic compounds such as formaldehyde (CH\textsubscript{2}O) to produce CO\textsubscript{2} at depth and possibly in the upper snowpack as well (Tschumi and Stauffer, 2000). The thermal reaction of H\textsubscript{2}O\textsubscript{2} with bromide (or chloride) to form HOB\textsubscript{r} (or HOCl) might also be significant on snow grains as a source of volatile, reactive halogens. While the rates of these reactions are slow (Mohammad and Liebhafsky, 1934), they might be significant in the quasi-liquid layers of snow if the reactants are sufficiently concentrated.

One of the most significant roles of hydrogen peroxide in snow grain chemistry is as a photochemical source of hydroxyl radical (Chu and Anastasio, 2005; Jacobi et al., 2006; France et al., 2007):

\[
\text{H}_2\text{O}_2(\text{snow}) + \text{light} \rightarrow 2\text{OH}
\] (12)
snow grains at midday during summer is typically 300 nmol at Summit indicate that the rate of OH formation on surface zone of Antarctic sea ice (King et al., 2005). Measurements dominant source of OH, as has been suggested for the photic least an order of magnitude), nitrate photolysis will be the much greater than those of hydrogen peroxide (i.e., by at

However, in snow or ice where nitrate concentrations are

Antarctic and Arctic snow grains (Chu and Anastasio, 2005).

Calculations based on laboratory measurements of 12 indicate that H$_2$O$_2$ photolysis is the dominant source of OH on Antarctic and Arctic snow grains (Chu and Anastasio, 2005). However, in snow or ice where nitrate concentrations are much greater than those of hydrogen peroxide (i.e., by at least an order of magnitude), nitrate photolysis will be the dominant source of OH, as has been suggested for the photic zone of Antarctic sea ice (King et al., 2005). Measurements at Summit indicate that the rate of OH formation on surface snow grains at midday during summer is typically 300 nmol L$^{-1}$ h$^{-1}$ and that H$_2$O$_2$ photolysis accounts for nearly all of this production (Anastasio et al., 2007). Based on peak measured concentrations of OH in the interstitial air of surface snow at Summit ($\sim 3 \times 10^8$ molecule cm$^{-3}$; Beyersdorf et al., 2007), the rate of partitioning of gaseous OH to snow grains will be approximately equal to the rate of formation from photolysis of H$_2$O$_2$ on the grains. However, the photolysis of H$_2$O$_2$ on snow grains is not a source of hydroxyl radical to the gas phase, as preliminary results indicate that the OH lifetime on snow grains at Summit during summer is short enough that OH will react before it can evaporate (Anastasio et al., 2007).

Like H$_2$O$_2$, formaldehyde (CH$_2$O) is an important radical precursor (Hutterli et al., 2001; Yang et al., 2002; Chen et al., 2004; Riedel et al., 2005) that is recycled between near-surface snow and the atmosphere (Hutterli et al., 1999, 2002, 2003). Distribution coefficient ($K_d$) values for CH$_2$O, determined between $-5$ and $-35^\circ$C, are about 1% of those reported for H$_2$O$_2$ (Burkhart et al., 2002). Air-snow exchange of CH$_2$O accumulated at or near the ice-air interface of snow grains leads to fast equilibration, and to significant fluxes into the boundary layer (Table 4). Total emission fluxes

### Table 4. Continued.

<table>
<thead>
<tr>
<th>Site</th>
<th>Ambient Air$^a$ pptv</th>
<th>Firn Air$^b$ pptv</th>
<th>Area Flux molecule m$^{-2}$s$^{-1}$</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>CH$_3$CHO</td>
<td>Alert, Canada</td>
<td>90 (70–110)</td>
<td>(220–270)</td>
<td></td>
</tr>
<tr>
<td>83$^\circ$ N, 62$^\circ$ W</td>
<td>22 Apr. 2000, 5 h total</td>
<td>22 Apr. 2000, 5 h total</td>
<td>4.2(±2.1)$\times 10^{12}$ (−)$^d$</td>
<td>Guimbaud et al., 2002</td>
</tr>
<tr>
<td>166 (26–459)</td>
<td>1 Apr.–4 May 2000</td>
<td>−</td>
<td>22 April 2000, 5 h total</td>
<td>Boudries et al., 2002</td>
</tr>
<tr>
<td>1800 (−)</td>
<td>Jan 1999, 1 profile</td>
<td>−(6000–7200)</td>
<td></td>
<td>−</td>
</tr>
<tr>
<td>CH$_3$COOCH$_3$</td>
<td>Alert, Canada</td>
<td>−(380–420)</td>
<td>−(410–580)</td>
<td></td>
</tr>
<tr>
<td>83$^\circ$ N, 62$^\circ$ W</td>
<td>22 Apr. 2000, 5 h total</td>
<td>22 Apr. 2000, 5 h total</td>
<td>−(6.2±4.2)$\times 10^{13}$ (−)$^d$</td>
<td>Guimbaud et al., 2002</td>
</tr>
<tr>
<td>871 (183–1470)</td>
<td>1 Apr.–4 May 2000</td>
<td>−</td>
<td>22 April 2000, 5 h total</td>
<td>Boudries et al., 2002</td>
</tr>
<tr>
<td>600 (−)</td>
<td>Jan 1999, 1 profile</td>
<td>−(2000–4000)</td>
<td></td>
<td>−</td>
</tr>
<tr>
<td>CH$_2$CHO</td>
<td>Ahmeek, MI</td>
<td>−(−)</td>
<td>−</td>
<td>−</td>
</tr>
<tr>
<td>47$^\circ$ N, 88$^\circ$ W</td>
<td>Jan 1999, 1 profile</td>
<td>Jan 1999, 1 profile</td>
<td>−</td>
<td>−</td>
</tr>
</tbody>
</table>

| Alkenes        | Summit, Greenland   | Ethene            | −(up to 163)                       |           |
| 73$^\circ$ N, 39$^\circ$ W | 9 (−)                | June–Aug 1999     | −                                  |           |
| 16 and 19 April 2000 | −(up to 89)         | −                 | −                                  |           |
| 6.2 (−)        | June–Aug 1999       | −(up to 42)       | −                                  |           |
| 3.3            | June–Aug 1999       | −                 | −                                  |           |

$^a$ Studies were included, which report fluxes and/or mixing ratios in ambient and firn air; for a comprehensive compilation of all available atmospheric observations see Sander et al. (available as an electronic supplement to Simpson et al., 2007)

$^b$ Ambient air levels typically at ~1 m above the snow surface

$^c$ Firn air levels within ~0.5 m below the snow surface

$^d$ derived with the gradient method (Monin-Obukhov similarity theory)

$^e$ based on firm-ambient air gradients and effective molecular diffusivities in air

$^f$ derived from concentration changes in the top layer of snow

$^g$ based on simulations with a physical atmosphere-to-snow transfer model

$^h$ Steady-state estimate assuming a BL height of 600 m

$^i$ determined with a quartz chamber

$^j$ inferred from the difference between modeled and observed ambient mixing ratios
of CH₂O (Table 4) originate from both physical and photochemical sources, with the former being dominant. For example, South Pole data suggest that net production of CH₂O within the snow by heterogeneous photochemical processes is likely to contribute not more than 20% to the total measured flux (Hutterli et al., 2004). However, the ratio between these sources will vary at sites with different concentrations of organic precursors in the snow. It will also vary with time as snow temperatures and actinic flux change on diurnal and seasonal time scales. The estimated net fluxes of CH₂O in the Arctic and Antarctic are generally lower than the corresponding H₂O₂ fluxes and range between 10¹⁰ and 10¹² molecule m⁻² s⁻¹, except for the coastal Neumayer, where estimated fluxes of ~2 × 10¹³ molecule m⁻² s⁻¹ were needed to reconcile measured CH₂O mixing ratios (Riedel et al., 2005) (Table 4).

Other radical precursors such as acetaldehyde (CH₃CHO) and acetone (CH₃C(O)CH₃) have been detected in surface snow (Houdier et al., 2002). Firm air levels of these species exceed those in the ambient air, suggesting snowpack emissions (Couch et al., 2000; Boudries et al., 2002; Guimbaud et al., 2002) (see also Sect. 4.4), probably with contributions from both physical and photochemical processes. However, a physical description of the air-snow transfer of these chemical species is still lacking, as well as net flux estimates between snow and atmosphere over longer time periods (Table 4). Modeling results of Grannas et al. (2002) indicate that both a daytime snowpack source (possibly photochemistry) and a nighttime snowpack sink (temperature-dependent uptake) of CH₂O, CH₃CHO, and CH₃C(O)CH₃ were required to correctly model the observed springtime diurnal variation of mixing ratios observed for these species in 2000 at Alert, Canada.

The snowpack chemistry of H₂O₂ and volatile carbonyls (as well as other organic compounds) is likely intimately linked to the hydroxyl radical. Because of its very high reactivity, OH is probably the major oxidant for organic compounds and other contaminants on sunlit snow grains (Anastasio et al., 2007; France et al., 2007), as it is in the atmosphere. Indeed, away from coastal regions with elevated sea-salt levels, the greatest sink for snow grain OH is possibly organic compounds (Anastasio et al., 2007). In this case, most photoformed OH will react with organics and these reactions probably account for a significant portion of the carbon-containing products that are emitted from snow (Dominé and Shepson, 2002; Grannas et al., 2004). In addition to CH₂O and other carbonyls such as those described above (Hutterli et al., 1999; Sumpner and Shepson, 1999; Boudries et al., 2002; Guimbaud et al., 2002; Jacobi et al., 2002; Sumpner et al., 2002; Hutterli et al., 2004), the products formed from OH reactions with organics may also include CO (Haan et al., 2001) and carboxylic acids (Dibb and Arsenault, 2002). It should be noted, however, that other mechanisms – such as direct photoreactions of organic compounds and indirect photoreactions involving other oxidants (Sects. 2.4 and 4.4) are probably also significant in the release of CO and VOCs.

In regions where snow composition is affected by inputs of sea-salt particles, reactions of OH with bromide and chloride on snow grains may also be important. While coastal regions certainly fall in this category, recent observations of boundary layer HO₂ chemistry at Summit suggest that halogen chemistry can also be important in regions 100 s of kilometers from the coast (Sjostedt et al., 2007). Hydroxyl radical oxidizes both bromide and chloride to initially form halogen radical species (such as Br, Br₂⁻, and the analogous chlorine species), which can be further transformed into volatile reactive halogens (such as Br₂ and BrCl) (Simpson et al., 2007). Based on results in solution, a portion of these reactive halogens on snow grains likely reacts with various organic functional groups to form halogenated organic compounds (Anastasio and Matthew, 2006). Such reactions could be responsible for the measured fluxes of alkyl halides out of the snow at Summit (Swanson et al., 2002). The volatile reactive halogens are also important because they will evaporate into the gas phase – both into the interstitial firm air and into the boundary layer – and photolyze to form halogen radicals, which in turn will destroy ozone and react with hydrocarbons, as described by Simpson et al. (2007).

Based on past studies in mid-latitude cloud and fog drops, illuminated polar snow grains likely manufacture a number of other oxidants in addition to OH. These additional oxidants include singlet molecular oxygen (¹O₂*), excited triplet states of organic compounds, and peroxy radicals (Faust, 1994; Anastasio et al., 1997). Initial work has shown that ¹O₂* is indeed formed on illuminated Summit snow and that the corresponding steady-state concentrations are high enough to be a significant sink for electron rich species such as polycyclic aromatic hydrocarbons (McKellar et al., 2005). As far as we are aware, no one has yet attempted to measure the presence of excited triplet states or peroxy radicals on snow grains, but these oxidants probably also affect snowpack chemistry and emissions into the boundary layer.

4.3 Halogens

4.3.1 Introduction

The chemistry of reactive halogens is most widely known through the impacts on ozone, causing dramatic depletions in both the stratosphere and in the polar boundary layer at sunrise. When the relationship between O₃ depletion and bromine was first discovered in the polar boundary layer, gaseous bromocarbons (e.g., CHBr₃) were suggested to be the source of reactive Br. Further study pointed to bromide in sea salt as the ultimate source, though unaltered airborne sea-salt aerosol is not sufficient. The current understanding is that sea salt expelled from the ice lattice during formation of sea ice and to some extent sea-salt aerosol that are deposited on ice is the halogen source. Modifications through interaction with ice surfaces, including brine and frost flowers.
or perhaps just through exposure after deposition onto sea ice and its snow cover during winter play a key role. Current understanding of halogen chemistry and ozone depletion in the polar boundary layer is the subject of a companion paper (Simpson et al., 2007). There is also increasing speculation that halogen chemistry is important throughout much of the troposphere (e.g., Vogt et al., 1996; Platt and Hönninger, 2003; von Glasow and Crutzen, 2006). Direct evidence for the release of photolyzable bromine compounds from the snowpack was first obtained by Foster et al. (2001) at Alert, Canada. Several recent studies investigating snow photochemistry have found evidence that halogen chemistry may be impacting HOX and NOX cycling in coastal Antarctica during the summer (outside of the severe ozone depletion season), and, surprisingly, at 3 km elevation along the crest of the Greenland ice sheet. In this section the evidence that halogen chemistry may be important in unexpected times and places in the polar regions, and the interactions between HOX, NOX and halogen chemistry that appear to result, are briefly introduced.

4.3.2 Direct release of halogen compounds from the snow

Laboratory experiments of Huff and Abbatt (2000, 2002) and Adams et al. (2002) showed that the uptake and reaction of HOB on frozen salt surfaces of variable NaCl/NaBr composition and temperature led to the release of Br2/BrCl to the gas phase. In the case of frozen surfaces no dependence on the pH of the solution that was used to prepare the frozen salt surfaces was observed, whereas in liquid a pH dependence was observed. This result points to a different mechanism in the case of frozen surfaces which so far has not been identified. The mass spectrometer measurements of Foster et al. (2001) and Spicer et al. (2002) showed very high mixing ratios of Br2 and BrCl of up to 27 and 35 pptv, respectively at 1.5 m above the snow surface at Alert, Canada. Enhanced Br2 concentrations within the snowpack (relative to the boundary layer concentrations measured just above the snowpack) were detected at depths down to 20 cm. These measurements provided the first direct evidence of a snowpack source of reactive molecular halogens potentially responsible for the observed tropospheric ozone depletion phenomenon. Comparison of BrO concentrations measured with an in-situ chemical conversion/resonance fluorescence technique and a DOAS system showed a strong vertical gradient of BrO at Alert (Avallone et al., 2003) again pointing to a strong influence from the snowpack. Measurements 1 m and 0.25 m above the snow showed up to 20 pptv of BrO hinting at a possible direct impact of the snowpack on BrO mixing ratios. Modeling results by Piot and von Glasow (pers. comm.) show that the life cycle of an ozone depletion event can only be simulated if re-emission of deposited (mainly particulate) bromine from the snowpack is considered. The exact processes and kinetics behind this remain, however, unknown. For a better understanding of the processes and a more correct inclusion into numerical models it is crucial that the details of halogen cycling in the snow are elucidated.

4.3.3 Atmospheric impact of halogen chemistry

As discussed in previous sections, several polar field campaigns observed OH/HO2 and NO/NO2 ratios that could not satisfactorily be explained with models. It was suggested that this is due to halogen chemistry. In the case of Summit, Greenland (as noted above), the measured OH levels were exceptionally high, and could not be accounted for by standard homogeneous chemistry that did not include halogens (Sjostedt et al., 2007). Since measured peroxy radical levels at Summit are well reproduced by model calculations, ozone production rates (P[O3] = k1[RO2][NO]+k2[HO2][NO]) should be accurately predicted. During the 2003 Summit field campaign average production rates of ∼0.8 ppbv O3 day−1 were calculated for the boundary layer, which is about 2% of the daily average ozone mixing ratio. However, at Summit photochemical production within the surface layer does not appear to have an important influence on surface ozone levels (Helming et al., 2002, 2007c), much in contrast to the findings from South Pole. In fact the boundary layer appears to be slightly depleted in ozone as balloon profiles frequently show higher mixing ratios above the boundary layer (Helming et al., 2002). Firn air measurements of ozone at Summit are lower than ambient, often by a factor of two or more (Peterson and Honrath, 2001). Ozone in the firn air declines with increasing solar irradiance levels, both diurnally and seasonally. While during April, up to 90% of ambient ozone was preserved to depths of 1 m, during mid-summer, only 20–50% remained at the same depth. Diurnal cycles in firn air ozone were clearly evident, with maximum ozone levels observed during the night and minima occurring after solar noon (Helming et al., 2007d). These observations imply that photochemical depletion is the most important removal mechanism of ozone in the interstitial air. The predominantly positive ozone gradients (higher ozone above the surface than in the snowpack) also point towards the snowpack being a sink of boundary layer ozone. This assumption has indeed been confirmed by direct ozone flux measurements at Summit, which show diurnally changing ozone fluxes with maximum deposition rates occurring in the afternoon (Helming, personal communication).

Sjostedt et al. (2007) speculated that the lack of observed ozone production at Summit and the perturbed (HO2+RO2)/OH ratio might be explained by a related chemical mechanism. The presence of halogen radicals could help explain both the slight depletion of boundary layer ozone and the enhancement in OH (Stutz et al., 1999; Foster et al., 2001; von Glasow et al., 2004). It is also possible that this chemistry could be altered by heterogeneous processes due to increased particle surface area (primarily blowing snow) in the high wind events. Alternatively, rapid transport of air that contains halogen compounds and/or sea-salt aerosol from the
Fig. 10. Total vertical column densities of BrO over the Arctic observed from GOME and SCIAMACHY during summer 2003.

marine boundary layer (as suggested by FLEXPART backwards simulations) to Summit might be another mechanism for activating halogen chemistry.

Although there are no measurements of BrO or IO at Summit, there exists ancillary evidence that suggests that both species may be present. For example, both the GOME and SCIAMACHY instruments on the ERS-2/ENVISAT satellites show moderate (approx. $3-5 \times 10^{13}$ molecule cm$^{-2}$) atmospheric columns of BrO during the summertime over the entire Greenland icesheet (see June 2003 composites, Fig. 10) (Richter et al., 1998; Wagner and Platt, 1998). Assuming the satellite BrO retrievals are correct and that the entire column of BrO is in a maximum 1 km boundary layer would result in ambient mixing ratios in the boundary layer of about 20 pptv. Low mixing ratios (1–2 pptv) of soluble gas-phase bromine species (Evans et al., 2003) were consistently detected by the UNH mist chamber during summer 2003. However, on 28 June when OH reached peak values for the 2003 campaign, soluble gas-phase bromide was above 8 pptv from the onset of measurement (10:00 a.m. local time) until 02:30 p.m. and reached a maximum of 11 pptv during this period. Halocarbons such as methyl bromide, ethyl bromide, methyl iodide and ethyl iodide have been observed in the atmosphere at Summit (Swanson et al., 2002). The observed concentrations of all four are enhanced in the firn column indicating active photochemistry involving halogen radicals within the snowpack. Annual changes of isobutane/propane ratios in the firn column at Summit also suggest a seasonal variance of bromine radicals. Consequently, we believe that further study of photochemistry at Summit should address the potential link between HO$_x$ and halogen chemistry in this environment by direct measurement of such species as BrO. A MAXDOAS installed at Summit in summer 2006 will provide preliminary testing of the hypothesis that halogens are impacting chemical cycles at this site. Intensive campaigns to be conducted in 2007 and 2008 will deploy long-path DOAS and in situ instrumentation to provide indirect evidence of halogen chemistry (OH/(HO$_2$ + RO$_2$), NO/NO$_2$, Hg speciation, NMHC) to confirm or refute the hypothesis, and should determine whether snowpack emissions or advection of marine air masses are the dominant source of any tropospheric halogens reaching the crest of the Greenland ice sheet.

In the case of Halley, Antarctica, measurements during the CHABLIS field campaign showed the presence of significant mixing ratios (~5 pptv) of both IO and BrO radicals at noon during the summer, with IO levels being much higher than observed in the Arctic. The levels for both halogen oxides peaked in spring with mixing ratios near 15 pptv (Saiz-Lopez et al., 2007). Under these conditions halogen oxides control the lifetime of NO$_x$ through the formation of halogen nitrates (XONO$_2$) and their subsequent hydrolysis on aerosols (Sander et al., 1999). In addition, the NO$_2$/NO ratio is also affected by the reaction of IO and BrO with NO, and the rate of removal of DMS via reaction with BrO, for which the rate coefficient is an order of magnitude greater than for removal by OH. IO and BrO also affect the oxidizing capacity of the Antarctic boundary layer by reacting with HO$_2$ and hence reducing the HO$_2$/OH ratio.

In order to study the impact of halogens on HO$_x$ concentrations at Halley, a time-resolved photochemical model using the Master Chemical Mechanism v3.1 (Jenkin et al., 1997), which included an explicit non-methane hydrocarbon scheme (11 NMHC species), and an additional halogen scheme, and constrained using measurements of longer-lived sources and sinks, was used to calculate the diurnal variation of OH and HO$_2$ (Bloss et al., 2006). The concentration of photolabile halogen precursors (for example Br$_2$ and I$_2$) was selected to give agreement with [BrO] and [IO] at the DOAS measurement height of ~5 m. When the halogen chemistry was switched off, the model overestimated HO$_2$ by more than a factor of 2 at noon, but when it was included, the agreement
for HO$_2$ was much better, whereas for OH there was a significant model overprediction, suggesting that further sinks are required. Apart from CH$_2$O, there were no measurements during CHABLIS of oxygenated VOCs, a class of compound which may represent a significant unmeasured sink for OH. This was observed, for example, during the North Atlantic Marine Boundary Layer Experiment (NAMBLEX), held at the remote Mace Head station in Ireland, where acetone, methanol and acetaldehyde were significant sinks for OH (Heard et al., 2006). HONO concentrations and J(HONO) were also determined during CHABLIS, but including photolysis of measured HONO as an OH source led to a large overprediction of OH, and this finding requires further investigation. The first and second most important sinks of HO$_2$ were reaction with IO and BrO, respectively, and HOI photolysis dominated OH production. The field and modeling results strongly suggest that the snowpack is a source for the measured halogens (Saiz-Lopez et al., 2007).

4.4 Organics

4.4.1 Organic materials in snow

Organic compounds in the atmosphere exist in particulate matter (solid and liquid) and as gases, originating from both natural (e.g. biomass burning, emission from plants) and anthropogenic (e.g. vehicles, industrial emissions, cooking) sources. They can be incorporated into snow by both wet and dry deposition processes (e.g. Semper and Kawamura, 1994; Franz and Eisenreich, 1998; Wania et al., 1999; Grollert and Puxbaum, 2000; Lei and Wania, 2004; Roth et al., 2004). Section 2 discusses the incorporation of atmospheric constituents into snow. Here we focus on “reactive” organic materials, while we note that other organic compounds (e.g. methane, halogenated organic molecules) are present in snow at significant concentrations, but due to their chemical inertness, their contribution to snow reactivity is considered negligible. Dry deposition is controlled by a number of factors including meteorology, snow surface properties, and, for particulates, the aerosol size distribution. There is a need to improve our understanding of the fundamental processes that deposit specific gas phase and particulate organic compounds to snow.

There are only limited measurements of organic compounds in snow, but these suggest that these compounds are ubiquitous and present at significant concentrations. At remote high latitude sites (Alert, Summit, Greenland, South Pole, see Fig. 2) measurements of total organic carbon, TOC, in snow ranged from <10 to 700 µg C L$^{-1}$ (Twickler et al., 1986; Grannas et al., 2004; Lyons et al., 2007). For comparison, values for sulfate in Greenland snow are in the range 100–300 µg C L$^{-1}$ (e.g., Twickler et al., 1986). By contrast, TOC values at Mount Sonnblick, Austria are higher, ranging from 1000 µg C L$^{-1}$ to 3000 µg C L$^{-1}$ (Haan et al., 2001), probably because of closer proximity to urban sources. Ha-
Organic compounds can be involved in snow photochemistry in a number of ways. They can directly absorb photons, thereby changing the radiation balance within the snowpack, and also create radicals that can react with other compounds (including organics) in snow (Hoffmann, 1996). Measurements on filtered, melted snow samples from Summit and Dome Concordia suggest that these direct photoreactions of organics are significant: unknown, presumably organic, species typically account for approximately half of the sunlight absorbed by these samples, while \( \text{H}_2\text{O}_2 \) and \( \text{NO}_3^- \) account for the other half (Robles and Anastasio, 2007). In addition to direct photoreactions, organic compounds can also react with other radicals generated within the snowpack, such as OH, which has a variety of sources including photodissociation of \( \text{H}_2\text{O}_2 \) (Chu and Anastasio, 2005) and, to a smaller extent, photolysis of nitrate in snow (Zhou et al., 2001; Dibb et al., 2002) (see Sect. 4.2). Finally, organic compounds can react directly with reactive species such as \( \text{H}_2\text{O}_2 \) and \( \text{O}_3 \) that may also be found within snowpack interstitial air.

There have been several reports demonstrating that some organic molecules are subject to post-depositional processing. Jaffrezo et al. (1994) found that the PAH benzo-pyrene decreased by nearly 90% from the surface to the bottom of a 3 m snow pit in Greenland. The results are consistent with measurements by Hagler et al. (2006) which show much higher concentrations of POC in Greenland surface snow than in shallow snowpits. At this time it is not clear whether the observed degradation of particulate organic compounds in snow is dominated by photochemical or physical processes.

A number of studies have observed ambient and/or firn air concentrations of gas phase organic compounds hinting at organic compound-related photochemistry in snow. Dibb and Aresenault (2002) found that the concentrations of formic and acetic acid in firn air were more than ten-fold higher than ambient concentrations. They hypothesize that snow photochemistry, in particular the production of carbonyls such as formaldehyde through oxidation of organics in snow (Summer and Shepson, 1999), are linked with the observed production of organic acids in firn air. Haan et al. (2001) also suggest that carbonyls in snow are linked with photochemical CO production. Similarly, Grannas et al. (2004) found that ice pellets made from melted Arctic and Antarctic snow produced \( \text{CH}_2\text{O} \) and \( \text{CH}_3\text{CHO} \) when irradiated in the laboratory. Production rates were enhanced by addition of nitrate to the melted snow, perhaps due to enhanced OH production. Of particular interest is the finding that a variety of organic compounds characterized in this work, in particular plant vascular material, were hypothesized to be involved in the photochemical production of \( \text{CH}_2\text{O} \) and \( \text{CH}_3\text{CHO} \) (Grannas et al., 2004).

Radical organic compounds generated through snow photochemistry may also play a more general role through subsequent reactions with compounds of environmental importance. Recently Lahoutifard et al. (2003) suggested that organic radicals in snow are responsible for scavenging gas phase mercury. This finding is supported by experiments where the addition of \( \text{H}_2\text{O}_2 \) (an OH radical source) to snow samples caused a dramatic increase in snow mercury concentrations during solar irradiation.

4.4.3 Production rates and snow as a source of boundary-layer gas-phase organics

A number of field and modeling studies indicate that a source of gas-phase organic compounds exists in polar snowpacks, and that the evolution and emission of these species may significantly impact the overlying boundary layer. For example, fluxes of organic compounds such as carbonyls, alkyl halides,
alenes, and organic nitrates out of the snowpack have been measured during various field campaigns (Sumner and Shepson, 1999; Hutterli et al., 1999, 2004; Couch et al., 2000; Grannas et al., 2002, 2004; Guimbaud et al., 2002; Guimbaud et al., 2002; Perrier et al., 2002; Swanson et al., 2002; Dibb et al., 2002; Klán and Holoubek, 2002). Oxygenated hydrocarbons in the atmosphere were measured at Alert from February to May 2000 (Boudries et al., 2002). During sunlight hours, concentrations measured in the snowpack interstitial air were higher than those measured in the overlying boundary layer, implying a positive flux from the snowpack to the Arctic boundary layer, proposed to be related to photochemical production in the snowpack. Guimbaud et al. (2002) found similar behavior for CH$_3$CHO and CH$_3$C(O)CH$_3$ and calculated fluxes from the snowpack of 4.2×10$^7$ and 6.2×10$^8$ molecule cm$^{-2}$ s$^{-1}$, respectively. Hutterli et al. (1999) report fluxes of CH$_3$O$_2$ from sunlit Summit, Greenland snow between 1.4–8.8×10$^6$ molecule cm$^{-2}$ s$^{-1}$, and Jacobi et al. (2002) report 8×10$^7$ molecule cm$^{-2}$ s$^{-1}$. CH$_3$O$_2$ fluxes from sunlit South Pole were on the order of 2×10$^8$ molecule cm$^{-2}$ s$^{-1}$ (Hutterli, 1999; Hutterli et al., 2004). Bottenheim et al. (2002) report a flux of ethene from Alert, Canada snowpack of the order of 1×10$^7$ molecule cm$^{-2}$ s$^{-1}$. Swanson et al. (2002) report production rates of several alkyl halides on the order of 10$^3$–10$^5$ molecule cm$^{-2}$ s$^{-1}$ in irradiated Summit, Greenland snowpack.

Although the aforementioned studies point toward photochemical pathways the emissions of organics are also influenced by physical processing within the snowpack, which in some instances can dominate over photochemical production. In the case of CH$_3$O$_2$, air-snow exchange of CH$_3$O$_2$ accumulated at or near the ice-air interface of snow grains leads to fast equilibration, and to significant fluxes into the boundary layer (see refs in Table 4). In this case, emission is dominated by physical partitioning processes. For example, South Pole data suggest that net photochemical production of CH$_3$O$_2$ within the snow accounts for no more than 20% of the total measured flux (Hutterli et al., 2004). The partitioning of other low molecular weight carbonyls is less well understood. As mentioned above, CH$_3$CHO and CH$_3$C(O)CH$_3$ fluxes were approximated by Guimbaud et al. (2002) and the condensed phase analyses of Houdier et al. (2002) shows seasonal concentration trends in opposition to thermodynamic expectations – implicating photochemical production as a possible explanation for the observations. The modeling work of Grannas et al. (2002) indicates that both snowpack photochemical production and temperature-dependent adsorption/desorption processes play a role in the observed diurnal variations of gas-phase low molecular weight carbonyl compounds in the Arctic boundary layer. To understand the extent to which physical or photochemical processes dominate the measured fluxes of particular organic species will require further fundamental studies of air/snow partitioning and a mechanistic understanding of the processes that may produce these compounds in sunlit snow or ice. It should be noted that fluxes and gradients are related by an exchange velocity and measured concentration gradients may not directly relate to the magnitude of the flux. To quantitatively determine the surface-air flux, and the magnitude of the vertical impact of snowpack emissions/sinks, it is necessary to understand the dynamics above the snow surface, as discussed in Sect. 5.1 and in greater detail in the accompanying boundary layer physics review (Anderson and Neff, 2007).

4.4.4 Processing of anthropogenic organics in snow/ice

Although much focus has been on the natural organic matter present in snow and ice, several laboratory studies have focused on the processing of anthropogenic organic materials in sunlit snow and ice. Laboratory experiments are necessary to elucidate snowpack photochemistry, however, at present the substrates generally used are significantly different from natural snow. Experiments use frozen solutions with high reactant concentrations, and it is possible that reactants are located in highly concentrated pockets of liquid solutions. In natural snow, concentrations are much lower, thus reactants could simply be adsorbed or contained in organic aerosols, and reaction pathways could be different.

Klán and coworkers have reported the unusual photobehavior of halobenzenes (such as chlorobenzene, 2- or 4-dichlorobenzene, bromobenzene, and 1,4-dibromobenzene) in ice (Klán et al., 2000; Klán et al., 2001). Photolysis of relatively concentrated (>10$^{-4}$ mol L$^{-1}$) frozen solutions of chlorobenzene, for example, produced chlorinated biphenyls (PCBs) or terphenyls as the major products because of the reactant concentrations. Similarly, Klánová et al. (2003 a,b) reported that chlorobiphenyldiols were the major products obtained by UV-irradiation of 2- and 4-chlorophenol (>10$^{-7}$ mol L$^{-1}$) in ice during laboratory studies at 253 K. Photolysis of these products (i.e. those resulting from reactions between organic and water molecules) were obtained at temperatures above 263 K and it was suggested that such transformations must occur in a quasi-liquid layer. Subsequent fieldwork confirmed that the same phototransformations as observed in the laboratory can occur in natural sunlight in polar regions, such as Svalbard (Klán et al., 2003). The toxic effect of these photoproducts was determined by Blaha et al. (2004) and found to be greater than for the parent molecules, suggesting that such chemistry occurring in snow and ice could be highly deleterious to the biosphere. Recent work by Sprenkle and Grannas (2006) indicates that certain persistent organic pollutants are also subject to photochemical degradation in snow/ice. Dubowski and Hoffman (2000) investigated the photochemical degradation of 4-nitrophenol in ice pellets and found products similar to what is produced in liquid water. Guzmán et al. (2006b) reported that frozen aqueous solutions of pyruvic acid, irradiated at $\lambda$>300 nm exhibit paramagnetic signals corresponding to distant triplet radical pairs. The photodecarboxylation reaction was shown...
to proceed by the same mechanism with similar quantum efficiencies in water and ice (Guzmán et al., 2006c, 2007).

Oxidation of aromatic and saturated aliphatic hydrocarbons and their derivatives (concentration $= 10^{-3}$–$10^{-5}$ mol L$^{-1}$) by OH, photochemically produced from hydrogen peroxide (concentration $= 10^{-1}$–$10^{-5}$ mol L$^{-1}$), in frozen aqueous solutions was recently investigated by Klán and coworkers (Kláňová et al., 2003a, b; Dolinová et al., 2007). While aromatic molecules (benzene, phenol, or naphthalene) reacted to form the corresponding hydroxy compounds, saturated hydrocarbons (methane, butane, cyclohexane) were oxidized to alcohols or carbonyl compounds. When frozen solutions containing nitrite or nitrate as well as aromatic compounds (such as phenol or 4-methoxyphenol) were photolyzed, the principal chemical processes included nitrilation, hydroxylation and coupling reactions (Matykiewiczova et al., 2007). However, the probability of any bimolecular reaction occurring in the natural environment will ultimately depend on organic contaminant local concentrations and oxidant availability at specific locations of the ice/snow matrix, as well as temperature, wavelength, and photon flux.

Although only a few studies report specific speciation, and we are far from understanding the organic carbon mass balance in snow, it is clear that there are a wide range of compounds existing both as particulates and in the QLL. Evidence suggests that these organic compounds play an important role in the chemistry of snow and the overlying atmosphere. There are still many unknowns including the chemical composition of organic compounds, the processes that deposit them to snow surfaces, their ability to partition between the air and snow, and the fundamental chemical processes that transform them within snow. Currently we are able to predict the course of photoreactions of organic compounds in ice/snow qualitatively but have insufficient data to extrapolate the experimental photolysis rate values to those occurring in the natural environment. Similarly, experiments that duplicate more closely the microphysical characteristics of natural snow are required.

5 Wider impacts of snow photochemistry

From the data presented above, both direct measurements of fluxes, and also the highly unusual and unexpected chemical composition measured in some regions of the polar boundary layer, it is clear that emissions from polar snowpacks influence the overlying boundary layer. Whether trace gas emissions from the snow have influences beyond the local or regional boundary layer is explored in this section.

5.1 The vertical extent of the influence of snowpack photochemistry

To have an atmospheric influence beyond the local boundary layer, trace gas emissions from snow need to reach the higher troposphere. However, extensive snowcover and reduced diurnal radiation cycles at high latitudes result in greatly reduced convective mixing and frequent conditions of shallow boundary layer depths and high atmospheric stability (see Anderson and Neff, 2007). Consequently, snowpack emissions are likely to be “trapped” in a thin air layer above the snow surface. Even differing boundary layer dynamics at various sites can have significant impacts on the vertical extent of snowpack influence. For example, ozone gradients measured during the summer from a tethered balloon at Summit were small and variable, and positive gradients in the lowest few hundred meters (on the order of a few ppbv) were observed more frequently than negative gradients. These data pointed towards a small uptake of ozone to the snow (Helmig et al., 2002). In contrast, much more pronounced, negative ozone gradients were the predominant, summertime condition at South Pole. Ozone near the surface frequently exceeded two times the levels that were observed in the lower free troposphere (Helmig et al., 2007a). Investigations of balloon sonde records further reveal that enhanced ozone concentrations near the surface are a predominant summertime phenomenon at South Pole (Crawford et al., 2001; Oltmans et al., 2007). Due to the lack of diurnal radiation cycles, stable boundary layer conditions with suppressed vertical mixing were noted to be more pronounced and longer-lasting at South Pole than at other polar locations (Cohen et al., 2007; Helmig et al., 2007a). This causes snowpack emissions at South Pole to accumulate to higher mixing ratios than at other sites. For example, mixing ratios of NO in the first few meters above the surface were significantly elevated and dropped to much lower levels at 100 m height above the surface (Fig. 11) (Helmig et al., 2007e).

Under these enhanced NO levels, ozone production occurs at rates reaching $3$–$6$ ppbv day$^{-1}$ (Crawford et al., 2001; Chen et al., 2004). Concentration gradients are directly related to the atmospheric lifetime of the chemical species. Ozone in the polar boundary layer has an estimated lifetime that is about 2 orders of magnitude longer than for NO. This causes ozone concentrations to decrease more slowly with height, with surface enhancements extending to several hundred meters above the surface (Fig. 11). Increased levels of NO in air nearest the surface were also observed during several aircraft flights made over the Antarctic Plateau (Davis et al., 2006). Various flights were made between South Pole and McMurdo station, including a sortie to Vostok and flights to midway (Dome C). These flights showed that over much (and possibly all) of the Plateau, NO levels were substantially higher than those observed along coastal areas. Mixing ratios of HO$_2$ and ozone production rates are expected to be highest not right above the surface, but within a distinct, several 10 s of meters high layer above the surface (Oltmans et al., 2007). Given the non-linearity of HO$_2$–NO$_x$ chemistry and the resulting sharp changes of oxidation rates with height, other chemical reactions are similarly expected to have strong height dependencies.
An earlier set of aircraft profile measurements aimed at investigating tropospheric ozone destruction events (ODEs) were carried out during the 2000 TOPSE campaign, flying from Colorado, through the Hudson Bay area, to north of Alert (Ridley et al., 2003). Several vertical profiles were conducted to as low as 30 m over the surface, including over the Arctic Ocean. At 30 m no evidence was found of an impact of a surface source of NO\textsubscript{x}, while the CH\textsubscript{2}O data were consistent with a significant impact of surface emissions of CH\textsubscript{2}O. The authors also concluded that surface sources of HONO had no impact by 30 m, based on the quite low levels of OH.

On the other hand, the halogens responsible for ODEs can have quite significant mixing ratios up to heights of several kilometers (e.g., Höninger and Platt, 2002; Friess et al., 2004; Höninger et al., 2004), and associated O\textsubscript{3} loss has also been observed to 1 to 2 km (Leaitch et al., 1994; Solberg et al., 1996; Bottenheim et al., 2002; Ridley et al., 2003; Tackett et al., 2007). Please see more discussion on halogen chemistry and ozone in Sect. 4.3 and Simpson et al. (2007).

5.2 Global/regional impacts: modeling assessment and observations

A first study to assess the global impact of the release of NO\textsubscript{x} from the snowpack was carried out by Carver et al. (2004). They used the chemical transport model p-TOMCAT (Savage et al., 2004) which includes a detailed inventory of NO\textsubscript{x} emissions, including for example, lightning and ship emissions. A number of multiannual integrations were carried out with the model to assess the impact of the release of snowpack NO\textsubscript{x}. Different scenarios were used in which the NO\textsubscript{x} emission rate was varied to look at the sensitivity of model results to the emission rate. The NO\textsubscript{x} emission was taken to occur over all snow covered regions around the globe during sunlit hours (solar zenith angle < 90°). Results from the model runs show that the estimated global NO\textsubscript{x} emission from snow is less than 1% of the global total. The polar regions accounted for most of the emissions due to the longer day compared to lower latitudes. Model results for a realistic emission scenario showed very large differences in surface NO\textsubscript{x} for Antarctica, with 10-fold increases in places, whereas in the northern polar regions the impact of snow emissions on surface NO\textsubscript{x} was considerably less. This is due to the much lower background concentration of NO\textsubscript{x} in the southern hemisphere. The study concluded that release of NO\textsubscript{x} from the snowpack makes a significant impact locally over Antarctica, consistent with observations.

Another question is whether the chemistry driving emissions from snow is occurring at other altitudes in the atmosphere. Cirrus clouds, for example, are radiatively important in the atmosphere and also provide surfaces for heterogeneous reactions. Laboratory studies have shown that HNO\textsubscript{3} can be taken up by ice surfaces at tropospheric temperatures (e.g., Hudson et al., 2002) and observations of uptake of HNO\textsubscript{3} by cirrus are accumulating (e.g., Weinheimer et al., 1998; Popp et al., 2004; Ziereis et al., 2004). A long-standing issue within atmospheric chemistry is that numerical models generally overestimate HNO\textsubscript{3}/NO\textsubscript{x} ratios in the upper troposphere (e.g., Chatfield, 1994; Brunner et al., 2005). Various model studies have considered different variables that might affect this ratio, such as lightning and convective transport from the polluted boundary layer (Brunner et al., 2003; Staadt et al., 2003), and heterogeneous reactions on sulfate aerosol (Tie et al., 2003). Calculations indicate that the photolysis of NO\textsubscript{3} on cirrus ice particles is too slow to improve model overestimations of the HNO\textsubscript{3}/NO\textsubscript{x} ratio (Chu and Anastasio, 2003), but there might be other ice chemical processes that are significant in this conversion.

5.3 Implications of snow photochemistry for the ice core community

Ice cores are powerful archives detailing how the Earth’s atmospheric composition and climate have changed over time (e.g., Legrand and Mayewski, 1997; EPICA community members, 2004). Provided there are no significant production or loss processes in the firn column and/or air bubbles, then the composition of air trapped in ice is representative of the overlying air, which, in the case of trace gases with long lifetimes, may also represent an “average” hemispheric
or global concentration. For Antarctic ice, this condition is more or less met, so that ice cores have provided us with detailed records of CH$_4$, CO$_2$, and N$_2$O over timescales up to 800 kyr (Siegenthaler et al., 2005; Spahni et al., 2005). The question, then, is whether photochemistry occurring in the snowpack influences the preserved concentrations of minor impurities, with a consequent effect on our ability to interpret them.

Ice cores hold the potential to tell us about other aspects of atmospheric chemistry, e.g. changes in the oxidative capacity or in NO$_x$ chemistry. The primary molecules and radicals involved, however, are short-lived, so the emphasis has been on studying more stable reaction products, such as CH$_2$O (Staffelbach et al., 1991) and H$_2$O$_2$ (Sigg and Neftel, 1988) for the oxidative capacity, and NO$_3^-$ for NO$_x$ chemistry (Wolff, 1995). Unfortunately there are two issues: (a) the ice record is derived from the polar boundary layer and may be disconnected from the relevant global or regional picture; (b) many of the important chemicals are found in the snow phase and are poorly preserved as the snow is compacted into firn and then ice. Snow photochemistry is relevant to both these issues. For the latter issue, the importance of snow photochemistry depends on the residence time in the photic zone and how this compares with the photolytic lifetime of the impurity. With an e-folding depth of actinic flux on the order of 5–25 cm, snow layers at low accumulation rate sites, such as in central Antarctica, remain under the influence of light for as much as several years and can undergo particularly intense photolytic loss.

For both CH$_2$O and H$_2$O$_2$, it seems now to be well-established that both physical (Hutterli et al., 2002, 2003) and photochemical (Sumner and Shepson, 1999) processes alter the concentrations preserved in polar ice. H$_2$O$_2$ in either the gas or aqueous phase would have a photochemical lifetime of several days in the upper layers of snow; therefore if it can reach the surface of snow grains it can be photolyzed to OH (which itself may then influence the concentrations of other molecules). However, physical exchange of H$_2$O$_2$ between the gas and snow phase is not limited to the photic zone, making it likely that physical processes determine the H$_2$O$_2$ ice core record and no measurable influence from photolysis is expected, since surface snow is buried by subsequent snowfall and moved rapidly below the photic zone. Decadal averages of H$_2$O$_2$ preserved in the West Antarctic Ice Sheet show indeed a strong correlation to local accumulation rate (Frey at al., 2006). For CH$_2$O, production from organic material in the ice (Sumner and Shepson, 1999, Grannas et al., 2004) and photolysis to CO (as one possible product) (Haan et al., 2001) may both occur, but again it seems likely that physical uptake and loss dominates the final preservation of the molecule in the ice sheet (Hutterli et al., 2003, 2004). The implication is that, for these compounds, to understand the relationship between concentrations in the overlying air and those preserved in firn beneath the photic zone, we need to concentrate on models that describe the physical air-firn equilibria, but that we may be able to finesse the photochemical processes in the upper layers. More difficult is the relationship between the chemistry of these compounds in the polar boundary layer and any globally relevant properties. This is certainly heavily influenced both by the confined nature of the boundary layer and by photochemical production and physical emission from snow and ice: for example, at South Pole, concentrations of both molecules are several times higher in the lowest levels of the atmosphere compared to those calculated from models that ignore emissions from the snowpack (Hutterli et al., 2004).

For nitrate in snow, the situation is also complicated. It has been calculated that, for snow accumulation rates and actinic fluxes typical for central Antarctica, as much as 40% of deposited nitrate might be photolyzed in the snowpack (Wolff et al., 2002). Although 40% might seem significant (and recent findings of deeper light penetration into polar snow would suggest even larger photolytic losses, Warren et al., 2006), it turns out that central Antarctic sites undergo huge losses of nitrate (in extreme cases by a factor 100) (Röthlisberger et al., 2000), and these losses continue down to 50 cm or more, where photolytic losses should be small. It therefore seems likely that physical losses of nitric acid dominate over photochemical losses, a suggestion that seems to be confirmed by measurements of $\delta^{15}$N in nitrate in firn (Blunier et al., 2005) (see earlier discussion). In coastal sites with higher snow accumulation rates, smaller losses of nitrate can be expected despite the somewhat higher actinic fluxes. However, we know that snow photochemistry very much dominates the NO$_x$ chemistry of the boundary layer at some sites (Davis et al., 2001), and so there probably is a close relationship between snow nitrate concentrations and local NO$_x$ concentrations, but with the large snow reservoir dominating the system. More interesting at the larger scale is to investigate how the input from outside the combined firn-boundary layer box influences the nitrate concentration preserved in snow and ice, and modeling studies are needed to investigate this further.

A secondary effect of the photochemistry might also be to induce an artificial seasonal cycle, because snow deposited just before the winter is buried before it can be significantly photolized. Alternatively, redeposition of nitric and pernitric acid formed from NO$_x$ generated in the top 10–20 cm of the snowpack could create a summertime surface peak in nitrate that actually represented redistribution of nitrate that accumulated over much longer times (nearly a year at South Pole, or several years at very low accumulation rate sites in central East Antarctica).

One further comment is that under conditions of the last glacial maximum (LGM) we expect compounds such as nitric acid to be “fixed” by reaction with alkaline dust material in the air or the snowpack (Röthlisberger et al., 2002). Under these conditions, physical exchange probably becomes negligible, and photochemical loss may also be reduced (if nitrate no longer sits on the outside of snow crystals). Such changes,
induced ultimately by climate, must also be considered.

In summary, although snow photochemistry is clearly very important for the chemistry of the polar boundary layer, physical exchange seems to be a more important determinant of what is preserved in deeper firn. If, however, we want to learn anything of more than local interest from these concentrations, we need to use modeling to determine which influences from a more regional or global scale can still be discerned in the preserved concentrations, despite the complications of the boundary layer/firn reactor.

6 Current limitations

There are currently a number of major constraints limiting progress in our understanding of snow phase photochemistry and its impacts on atmospheric composition. These are issues for field, laboratory and modeling studies, as discussed here.

6.1 Field studies

6.1.1 Infrastructure and access for field studies

Field studies in polar environments are highly challenging and the difficulties of accessing these regions with appropriate instrumentation are significant limitations in the study of snowpack photochemistry. For example, there are only a limited number of fully supported field laboratories in polar regions, and to allow detailed chemical and physical analysis, they must have access to uncontaminated snow and ice and clean air. Field stations may be carefully located to minimize contamination, but an important area of research is in development of renewable (non-fossil) sources of electric power for research in these extreme environments. This is currently happening at the GeoSummit station Greenland, but similar sites are needed elsewhere in the Arctic and Antarctica. Furthermore, the harsh conditions associated with polar research lead naturally to the majority of research being carried out during the summer months. To really probe the processes at work extended measurement campaigns with year-round duration are necessary.

Assessments of trace gas fluxes out of the snowpack can be used to parameterize numerical model calculations into wider impacts of snow photochemistry. These fluxes are influenced by a number of variables, including atmospheric stability, frequency of fresh snow fall events, snowpack concentrations, and changes in irradiance. More flux measurements carried out during all seasons and addressing a wider range of chemical species are required and these experiments should also aim at differentiating between contributions from physical and photochemical sources. Flux measurements are experimentally very challenging, as they require either a method for the fast and selective measurement of the species of interest for eddy correlation measurement or highly precise and accurate measurements for flux determinations by the tower gradient methods. For many of the gases of interest, such instrumentation is currently not available. Furthermore, micrometeorological flux measurement approaches often fail under the frequently highly stable conditions over snow, therefore improvements in micrometeorological methods for flux measurements under stable conditions are desirable. Finally, much of the previous snow research has been done in polar environments. In order to assess the influence globally, flux measurements are also needed at snow-covered non-polar locations.

To date, few studies have focused on snow photochemistry occurring in mid-latitude regions. It is likely that snowpack photochemistry will be very active at lower latitudes, due to lower solar zenith angle, increased irradiance and higher concentrations of reactive precursors within snow (nitrate, peroxide, organic materials, etc). It could also be expected that the chemistry occurring in the QLL would be more active at mid-latitudes, as the liquid water content of a relatively warm mid-latitude snowpack would be larger than in a much colder, high latitude snowpack. However, the overall impact of snowpack emissions to the overlying boundary layer may be less due to the proximity to anthropogenic influences of e.g. NOx. To assess the influence of mid-latitude snow on atmospheric chemistry will require further field studies in these regions.

6.1.2 Instrumentation for field studies

A crucial question concerns the impact of snowpack photochemistry on the overlying atmosphere, as outlined in Sect. 5. To address this question in the field it is currently necessary to use instrument platforms such as tall towers, tethered balloons, released balloons, blimps, and/or aircraft. For several of these platforms, associated instrumentation must be light weight and low-power, and such instrumentation currently only exists for a very limited suite of molecules. Instrumentation must also be developed for remotely sensing the vertical structure of the atmosphere, including variations in composition, over the snowpack.

Making gas phase measurements within the snowpack interstitial air is difficult, particularly for some molecules at very low mixing ratios. Gas phase sampling rates can be on the order of many liters of air per minute, which leads to efficient artificial ventilation of the snowpack from ambient air above (or interstitial air below) the point of sampling (Albert et al., 2002). This will lead to a potentially mixed signal of both interstitial and boundary layer air and may mask real concentration differences that exist between these two locations. Passive sampling methods could be developed (such as those currently used for persistent organic pollutants, e.g. Farrar et al., 2006), however these often require long exposure times, so high resolution measurements would be prohibited. Additionally, incorporation of such samplers into the snowpack is difficult to achieve without disturbing the very medium they are intended to study.
6.2 Laboratory studies

6.2.1 Laboratory studies on snow surfaces

Laboratory studies of snow surfaces are severely limited by the techniques used to generate the frozen samples studied. Often the analytical detection limits of instrumentation used to monitor processes are well above the typical ambient level of a reactive species present in a natural snowpack. Thus, investigators may find it necessary to use high solute concentrations (relative to what is present in a natural snow sample). If solute concentrations are very high, it is possible that complete freezing will not be achieved and the experimental surface being studied is actually a liquid and not a true disordered QLL. Additionally, samples frozen under laboratory conditions certainly do not represent the true nature of a natural snow formed by e.g., condensation of water vapor onto IFN or riming. Regardless, laboratory studies provide important information about the nature of the surface, reactivity of species in frozen matrices and mechanistic considerations that prove invaluable to field and modeling studies and efforts to improve laboratory techniques are continuing.

6.2.2 Chemical analysis of snow

There is substantial uncertainty about the chemical nature of important snowpack reactants and chromophores, particularly for organic constituents. The chemical analysis of snow for organics is complicated by the fact that much of the organic matter in snow derives both from biota and from bacteria (Sattler et al., 2001; Grannas et al., 2004) and may be in both dissolved and particulate phases. Analysis of meltwater samples will not be representative of the original distribution of organics between the dissolved and particulate phases, thus it is imperative that techniques be developed that can probe the snow chemical environment in situ. This will probably take the form of advanced microscopic or spectroscopic techniques, such as scanning electron microscopy (SEM), nuclear magnetic resonance (NMR) or non-linear spectroscopic methods such as sum frequency generation (SFG) or second harmonic generation (SHG). Similar issues exist for measurement of pH on the surfaces of snow crystals/grains; pH of the intact QLL, for example, is likely to be significantly different to that of the melted crystal. We note also that stable isotope studies on trace species in snow and air is a fast growing field with interesting potentials on key processes taking place in the firn/snow interface. Isotope fractionation factors are a good indicator of kinetic processes, with the possibility of differentiating between physical and chemical processes. Sensitivity is constantly improving and new techniques based on spectroscopy methods (e.g., cavity ring down spectroscopy) are emerging with the advantage of in situ, passive and non-disturbing analysis coupled with light weight and low energy consumption equipment capable of producing high-time resolution records of concentration and isotopic composition.

6.2.3 Nano/micro-scale physical and chemical analysis

A major set of issues for snowpack photochemistry lies within the physical realm of the reactants and photochemistry in the snowpack. As discussed earlier, reactants can be located on snow crystal surfaces and can also be dissolved within the ice crystal lattice, trapped in aerosol particles, at grain boundaries or in pockets of concentrated solutions that could for example be formed during riming. Knowing where reactants reside is fundamental to theoretical studies of snow photochemistry. The issue of the physical environment is a difficult one and has been the subject of only a few studies. Scanning electron microscopy/energy dispersive spectrometry (SEM-EDS) work (Obbard et al., 2003; Barnes and Wolff, 2004) suggests that the location of reactants cannot be explained by considering only the nature of the reactant. Most likely, interactions between different species come into play, as well as the mechanism of formation of the ice. More studies of natural snow using a variety of advanced microscopic and spectroscopic techniques (e.g., SEM-EDS, extended x-ray absorption fine structure (EXAFS), atomic force microscopy (AFM), SFG, SHG, NMR) are in order if we wish to progress on these aspects. The relevance of reactions on deposited aerosol that is incorporated into the snowpack as reactive sites cannot be addressed with current methods. The relative importance of reactions in the snowpack interstitial air, on aerosol particles, in the QLL or other sites is unknown at this point but crucial for our quantitative understanding of the processes and for our ability to eventually include these processes in detail in numerical models.

Furthermore, we currently have a very limited understanding of the kinetics of reactions occurring on/in snow. Indeed, there are numerous unknowns that must be tackled before we can routinely predict the rates of these reactions. For example, the concentrations of OH and other oxidants on snow grains must be measured in order to estimate the reaction rates between oxidants and snow grain contaminants. In addition, the rate constants for these snow grain reactions must be determined. It is currently unclear whether rate constants can be estimated from solution data or whether they are specific for ice at a given temperature and composition. While quantum yields for the direct photolysis of chromophores such as nitrate and hydrogen peroxide behave similarly in solution and ice (e.g., Chu and Anastasio, 2003, 2005), second-order thermal reactions that require collisions between the two reactants are likely to be not as well behaved. Direct photochemical reactions of organic compounds probably also play a significant role in chemistry occurring on snow surfaces, but understanding these processes will require determining the identities of the organics, their light absorption properties and quantum yields for reaction, and the products formed.
6.3 Modeling studies

The real test of our understanding of snowpack photochemistry is whether the first principle based numerical models can simulate observations. The challenges currently presented to us involve not only incomplete, or lack of, understanding of the polar physical and chemical processes (e.g., snow chemistry and air-snow exchanges) but also scarce meteorological observations necessary for model simulations in polar regions. One critical need is for specific, lower dimensional models, based on first principles and including all relevant chemical species, to describe the chemistry of the atmosphere-snow system. Development of modules describing chemistry occurring in the QLL and ice grains is at early stages. These modules will need to be coupled to models of the transport processes that exchange reactants and products between the several condensed phases and the firm air within the snowpack, and between the snowpack and the overlying lowermost part of the atmospheric boundary layer. Heterogeneous processing within this lowermost boundary layer also needs to be simulated because uptake by aerosol, fog and snowflakes contributes to recycling of reactive species emitted from snow; reactions in these atmospheric condensed phases may also transform the emitted species to different chemical forms. Such a unified 1-D model could then link the snowpack and boundary layer to the free troposphere above sunlit snow. A long-term goal is to couple such snowpack models to 3-D atmospheric chemistry transport models to allow calculations of snowpack photochemistry and investigate its impacts over large geographical areas as well as the impact of transport from lower latitudes on snowpack chemical compositions.

7 Conclusions

The capacity for snow on the Earth’s surface to photochemically-generate reactive trace gases and release them into the overlying atmosphere is an important phenomenon that has only recently come to light. Measurements in both polar regions have shown that emissions from snow are fundamental to driving local and regional boundary layer chemistry; early modeling studies are exploring the global effect. The scientific community has made significant progress in understanding snow photochemistry since its initial discovery. Field measurements investigating gas-phase, aerosol-phase and snow-phase chemistry have been undertaken in a variety of locations. Laboratory studies have investigated, both qualitatively and quantitatively, a variety of important factors such as reaction rates and mechanisms, quantum yields, fundamental behavior of molecules at the snow/ice surface, and partitioning of chemicals within snow/ice. There remains a lot to investigate and to learn, but obstacles limiting the progress of our understanding (e.g., the analytical techniques available to study processes at the microscopic and molecular level) are not insignificant.

The Earth’s cryosphere is undergoing significant change. Reductions in sea ice extent in both the Arctic and parts of the Antarctic are already reported, and further reductions as well as decreases in global land snowcover are anticipated. These changes will affect the influence of snowpack photochemistry, adding urgency to our current task of understanding and quantifying relevant processes. For example, a reduction in global snowcover (both over land and ocean) will reduce the net emission of trace gases from snow into the atmosphere while increasing processes that occur on the underlying surfaces. In addition, changes in precipitation rates will affect scavenging processes. The modeling studies carried out to date, as well as field observations, suggest that reductions in snowpack emissions are likely to be regionally important, but the direct effect globally has not yet been addressed. Secondary effects might also be anticipated from a reduced albedo, which will reduce photolysis rates and hence the fate of snowpack products. Similarly, changes in atmospheric stability and mixing, resulting from increased heat inputs will tend to dilute the concentrations of species emitted from the surface, again slowing photochemistry. Further, trace gases emitted from snow or influenced by these emissions (e.g., OH, NO and halogens) are involved in production of CCN from DMS. Reduced snowpack photochemistry would decrease DMS oxidation rates and hence CCN production. Finally, as discussed earlier (and see also Simpson et al., 2007), concentrations of polar boundary layer ozone are also affected by snowpack photochemistry. In polar regions, the radiative impacts of ozone are more important than at lower latitudes due to lower concentrations of water vapor. Hansen et al. (2005) have concluded that tropospheric ozone is an important contributor to warming and sea ice loss in the Arctic. However, in particular through the reactions involving halogens, there is an intriguing positive feedback between sea ice loss, and the contribution of tropospheric ozone to radiative forcing, as loss of sea ice may cause reduced ozone depletion events and higher ozone levels, with increased radiative forcing and warming from tropospheric ozone.

It is thus clear that a warming climate will affect trace gas emissions from snow and all the subsequent processes that these emissions influence. It is essential that we build on our current knowledge in order to develop comprehensive numerical models that can address issues of snow photochemistry and its influence on the regional and global atmosphere both now and in a future warmer world.

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