Quasi-Liquid Layer Formation on Ice under Stratospheric Conditions

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Characterization of the interaction of hydrogen chloride (HCl) with ice is essential to understanding at a molecular level the processes responsible for ozone depletion involving polar stratospheric cloud (PSC) particles. To explain the catalytic role PSC particle surfaces play during chlorine activation, we proposed previously that HCl induces the formation of a disordered region on the ice surface, a 'quasi-liquid layer' (QLL), at stratospheric conditions. The QLL is known to exist in pure ice crystals at temperatures near the melting point, but its existence at stratospheric temperatures (-85°C to -70°C) had not been reported yet. We studied the interaction of HCl with ice under stratospheric conditions using the complementary approach of a) ellipsometry to directly monitor the ice surface, using chemical ionization mass spectrometry (CIMS) to monitor the gas phase species present in the ellipsometry experiments, and b) flow-tube experiments with CIMS detection. Here we show that trace amounts of HCl induce QLL formation at stratospheric temperatures, and that the QLL enhances the chlorine-activation reaction of HCl with chlorine nitrate (ClONO₂), and also enhances acetic acid (CH₃COOH) adsorption.

Surface disorder was observed via ellipsometry on pure single-crystalline hexagonal ice with no HCl present down to approximately -30 °C. This value is consistent with other experimentally determined values of QLL onset temperature for bare ice, which range from a few degrees below the melting point of water to less than -40 °C.

Figure 1 summarizes the results of our investigation of the HCl-ice phase diagram using the ellipsometry-CIMS approach. A change in signal consistent with the formation of a disordered interfacial layer on the ice surface was observed in the range of HCl partial pressures and temperatures in the vicinity of the solid-liquid equilibrium line on the HCl-ice bulk phase diagram. It is important to note that this range of conditions includes those encountered in the polar stratosphere during PSC events. While surface disorder on ice at stratospheric conditions has been predicted theoretically, this is the first report of experimental evidence of HCl-
induced QLL formation at stratospheric temperatures. Assuming the optical constants of this layer to be those of liquid water, we estimate that HCl induces a disordered layer with thicknesses of a few nanometers up to 100 nm depending on the quality of the ice-surface and the location in the phase-diagram.

In the range of HCl partial pressures and temperatures far from the solid-liquid equilibrium line, i.e. in the interior of the ice stability envelope, no surface change was observed. For example, exposure of an ice surface to approximately $10^{-7}$ Torr HCl was observed to induce QLL formation for $T>-35^\circ$C and $T<-65^\circ$C, but in the region $-35^\circ$C$<T<-65^\circ$C we found no evidence of surface change. Indirect support for the observation that HCl-induced surface change is confined to the region of the HCl-ice phase diagram near the solid-liquid equilibrium line is found in the work of Hynes et al., who report that at $10^{-6}$ Torr HCl the uptake coefficient, $\gamma$, of HCl on ice decreased from $\gamma > 0.1$ at 200 K (conditions at which we observe surface change) to $\gamma < 0.01$ upon increasing the temperature above 205 K (conditions at which no surface change was observed in our experiment). Surface disorder is expected to enhance HCl uptake efficiency. Our flow tube-CIMS studies lend further support for the observed trend.

Two types of experiment were performed using the ellipsometer-CIMS system: a) constant temperature experiments in which the ice sample was exposed to a step change in partial pressure of HCl and the ellipsometer signal was monitored over time, and b) constant HCl partial pressures with temperature scanning experiments. Ellipsometer signal time traces for a typical temperature scanning experiment are shown in Figure 2. In the constant temperature experiments, after exposure to HCl, surface change was observed only after an induction time of 1-10 min. An aging effect was observed in that induction times were seen to decrease after the first exposure of a surface to HCl. Both the surface disordering and this aging effect were seen to be reversible.

The reaction of ClONO$_2$ with adsorbed HCl was studied using the flow tube-CIMS technique on zone-refined ice cylinders. As shown in Figure 3, the production of Cl$_2$, and thus the efficiency of the reaction, decreased as we moved from conditions where surface change was observed with the ellipsometer to those where no surface change was observed. At $-77^\circ$C, the reaction proceeds efficiently ($\gamma > 0.1$), and HCl is readily available on the surface for reaction, as has been observed previously. At $-55^\circ$C, with the same reactant concentrations, HCl is not readily available on the surface for reaction, despite the gas-phase presence of HCl. Cl$_2$
evolution consists of an initial 'burst' followed by tailing off of Cl$_2$ release. Kinetic analysis of the 'burst' portion of the Cl$_2$ signal yields a reactive uptake coefficient at least one order of magnitude smaller ($\gamma = 0.014 \pm 0.005$) than that for the same reaction at temperatures where surface change was observed using ellipsometry.

The results of our flow tube-CIMS studies of HCl adsorption on zone refined ice cylinders, as shown in Figure 4, demonstrate that the nature of HCl adsorption on ice differs at the conditions under which surface change was observed with ellipsometry from the HCl-ice interaction at conditions under which no surface change was observed. At -77 °C and 7.4$\times$10$^{-7}$ Torr HCl, almost fully reversible uptake was observed. An initial fast uptake mode ($\sim 10^{14}$ molecule cm$^{-2}$ over 180 sec) was observed, which we attribute to surface adsorption prior to QLL formation. A second adsorption mode was observed consisting of a nearly constant flux of HCl ($\sim 5 \cdot 10^{11}$ molecules cm$^{-2}$ s$^{-1}$) from the surface to the interior of the ice sample, which persisted throughout the time scale of the experiment (1 hour). This loss is observed to be reversible. A memory effect was observed in that this flux increased in magnitude with subsequent exposures of the same ice crystal. These observations of surface-to-bulk loss$^{18,19}$ and memory effect$^{4,17-19}$ are consistent with the findings of other investigators. A model of HCl absorbing into a 100 nm thick film with desorption properties between those of ice and liquid water, with diffusion from the near-surface region into the bulk ice lattice using $D_{\text{HCl}} = 10^{-12}$ cm$^2$ s$^{-1}$ is sufficient to simulate the observed surface-to-bulk loss at long times. In addition, the ClONO$_2$ + HCl reactive uptake experiments indicate that this HCl is readily available for reaction with ClONO$_2$. Therefore, we attribute this loss mainly to absorption of HCl by the disordered surface region.

In flow tube-CIMS studies of HCl uptake at -59 °C and 6.5$\cdot$10$^{-7}$ Torr HCl, conditions under which no surface change was observed with ellipsometry, we observed largely irreversible uptake. We again observed an initial fast mode of HCl adsorption ($\sim 6 \cdot 10^{14}$ molecules cm$^{-2}$ over 500 sec), followed by a slower uptake mode, which leads to saturation within $\sim 3000$ sec. The magnitude of adsorbed HCl decreases with subsequent exposures but adsorption remains largely irreversible. The ClONO$_2$ + HCl reactive uptake experiments indicate that HCl adsorbed under these conditions is for the most part not readily available on the surface for reaction with ClONO$_2$. We attribute this irreversibly adsorbed, inaccessible HCl to diffusion into the bulk and into grain boundaries.
As a third corroboration of these findings, the co-adsorption of HCl and CH₃COOH was studied on zone-refined ice cylinders. CH₃COOH is a weak acid that does not dissociate on bare ice surfaces at -45 °C.²¹ At a constant temperature of -61 °C, at partial pressures of HCl where no surface change was observed with ellipsometry (P₇₅ = 7 · 10⁻⁷ Torr), an uptake of ~ 1.4 · 10¹⁴ molecule cm⁻² was observed. This is an overall enhancement compared to values of acetic acid uptake on bare ice, indicating that the presence of HCl enhances the affinity of acetic acid for the ice surface even when no surface change is observed via ellipsometry. At the same temperature, twofold enhancement in CH₃COOH adsorption (2.9 · 10¹⁴ molecule cm⁻²) was observed at higher HCl concentrations where surface disorder was observed (P₇₅ = 2 · 10⁻⁶ Torr).

In conclusion, we have confirmed that surface disorder induced by the presence of HCl explains the role played by type II PSC particles in catalyzing chlorine activation reactions. This surface disorder also enhances the adsorption of species that do not participate in chlorine activation including weakly polar ones such as CH₃COOH. In the case of warmer temperature atmospheric ice particles, such as those encountered in tropospheric cirrus clouds, surface disorder may play a key role in surface chemistry. The affinity of gas phase adsorbates for these ice surfaces may be greater than previously assumed, with implications for the role of ice particles in the scavenging of semi-volatile gas phase species and the reactive heterogeneous chemistry of the troposphere.²²,²³

METHODS

Ellipsometer-CIMS experiments

The ellipsometer used in these measurements (Beaglehole Instruments) employs a photoelastic birefringence modulator, allowing high sensitivity measurements. The light source is a 50 mW He/Ne laser, focused to a diameter of approximately 1 mm. We observe true melting at 0 °C, and do not see surface disorder far from the solid-liquid equilibrium line in the HCl-ice phase diagram, and therefore we infer that the laser does not melt the ice surface due to local heating. The ice sample was placed in an aluminum sample holder and housed in a vacuum-jacketed flow tube fitted with quartz windows to allow the laser beam to pass through the cooling jacket and flow tube walls. The flow tube was operated in the laminar flow regime (Re < 100) with He as a carrier gas. Ice was placed upstream of the main ice sample to ensure that the
sample was in equilibrium with water vapor, i.e. no net evaporation or condensation took place. The flow tube was interfaced with a CIMS apparatus (Extrel C50) to monitor the gas phase composition via chemical ionization by SF₆⁻ reagent ions. HCl was monitored as SF₅Cl⁻ (162 amu). In an HCl uptake experiment, the ice sample was exposed to a dilute mixture of HCl in He that was introduced to the main He flow through a moveable injector positioned near the centerline of the flow tube. The injector was heated to prevent experimental artifacts due to adsorption of HCl to the injector walls, creating a temperature gradient of < 1°C in the flow tube. Ice samples were prepared from single crystals of ice grown using an adaptation of the Bridgeman technique²⁴ and were observed using crossed polaroids to have grain sizes on the order of centimeters.

Flow tube-CIMS experiments

All flow tube-CIMS experiments employed a 2.5 cm i.d. flow tube operating in the laminar flow regime (Re < 100) that was interfaced with a CIMS apparatus (Extrel C50). Detection occurred via SF₆⁻. HCl was monitored as SF₅Cl⁻ (162 amu), ClONO₂ was monitored as ClONO₂F⁻ (116 amu), Cl₂ was monitored as Cl₂⁻ (70 amu), and CH₃COOH was monitored as CH₃CO₂HF⁻ (79 amu).

Hollow cylindrical ice samples ~ 2 mm thick were prepared using a modification of the Bridgeman technique,²⁴ using a cylindrical mask to form the central flow channel. Ice samples formed in this way were observed using crossed polaroids to have grain sizes on the order of several millimeters.

In an HCl uptake experiment, the ice sample was exposed to HCl via a heated injector as in the ellipsometry-CIMS experiments. The ClONO₂ + HCl experiments and the acetic acid/HCl co-adsorption experiments employed two injectors.

In the ClONO₂ + HCl experiments, first a constant flow of HCl was established. Once the surface was saturated with HCl, the uptake of a constant flow of ClONO₂ and the associated Cl₂ evolution was monitored. ClONO₂ was prepared from ClF and Pb(NO₃)₂ using the method of Schmeisser²⁵ followed by fractional distillation at -77 °C and -95 °C, producing a high-purity sample. Reactive uptake measurements at several exposed ice lengths (and thus reaction times) were converted into kinetic information using the Brown algorithm.²⁶ Values reported here were calculated using a ClONO₂ diffusion coefficient in He of 176 cm²sec⁻¹Torr at 200K, assuming a
dependence. Reported uncertainties reflect a 20% uncertainty in diffusion coefficient and the observed loss rates.

In the acetic acid/HCl co-adsorption experiments, initially a constant flow of HCl was established. Once the surface was saturated with HCl, the uptake of a constant flow of CH$_3$COOH and the associated HCl desorption was monitored. Dimerization of gas-phase acetic acid was accounted for when determining the acetic acid sample concentration and thus the partial pressure of acetic acid in the flow tube. The equilibrium constant for dimerization at 25°C and 1 atm is 0.972.
FIGURES

Figure 1. The phase-diagram of HCl-ice adapted from Molina et al. Thermodynamically stable phases are shown as a function of temperature and HCl partial pressure. “Ice” is the stable phase under polar stratospheric conditions (circled area). “Liquid” refers to a liquid solution, and “Trihydrate” and “Hexahydrate” refer to the crystalline hydrate states. Note that, experimentally, the phase transitions might be observed with a delay due to kinetic barriers. Circles (○) refer to conditions where a change in signal consistent with the formation of a disordered interfacial layer on the ice surface was observed upon exposure of the ice crystal to HCl, whereas filled boxes (■) refer to conditions where no change to the ice surface was observed. Arrows represent experiments involving a scan in temperature during exposure to a constant partial pressure of HCl. Bars represent temperatures at which we observe cease/onset of surface changes. Transition to the hexahydrate phase or the melt is indicated by “A”.

Figure 2. Time study of phase-modulated ellipsometry signal for ice sample exposed to 5 $10^{-7}$ Torr HCl and temperatures ranging from -52.5 °C to -77.2 °C. The traces shown are the ellipsometer signals measured at 50 kHz (x-trace) and 100 kHz (y-trace). The x- and y- traces are related to the real and the imaginary-part of the reflectivity, respectively. The ice sample was held initially at -52.5 °C, cooled to -77.2 °C, and then warmed again to -52.5 °C. As the temperature decreases, there is a discontinuity in the signals corresponding to a surface change at -66.5 °C. When the sample is then warmed to its original temperature, another discontinuity is observed, with the signals returning to their previous levels at -64.5 °C. Experiments such as this one are labeled using arrows in Figure 1.

Figure 3. The reaction of ClONO2 with adsorbed HCl on zone-refined ice cylinders. Experiments shown were conducted at approximately 1-10^{-6} Torr HCl and 5 $10^{-7}$ Torr ClONO2 (pseudo-first order conditions). The left and right panels show studies of the reaction at -56 °C and -77 °C, respectively, and the upper and lower panels show ClONO2 and Cl2 mass spectrometer signals, respectively. The production rate of Cl2 ---and thus the efficiency of the reaction--- decreases as we move from conditions where surface change was observed with the ellipsometer to those where no surface change was observed.
Figure 4. HCl adsorption on zone-refined ice at approximately $7 \cdot 10^{-7}$ Torr HCl. The upper and lower traces are HCl mass spectrometer signal for adsorption at -77 °C and -59 °C, respectively. The nature of the interaction of HCl with the ice samples changes as we move from conditions where surface change was observed with the ellipsometer to those where no surface change was observed.
Figure 1.
Figure 2.

Figure 3.
Figure 4.
REFERENCES


