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Abstract. Recent observations have detected surface active organics in atmospheric aerosols. We have studied the reaction of N$_2$O$_5$ on aqueous natural seawater and NaCl aerosols as a function of sodium dodecyl sulfate (SDS) concentration to test the effect of varying levels of surfactant on gas-aerosol reaction rates. SDS was chosen as a proxy for naturally occurring long chain monocarboxylic acid molecules, such as palmitic or stearic acid, because of its solubility in water and well-characterized surface properties. Experiments were performed using a newly constructed aerosol flow tube coupled to a chemical ionization mass spectrometer for monitoring the gas phase, and a differential mobility analyzer/condensation particle counter for determining aerosol surface area. We find that the presence of $\sim$3.5 wt% SDS in the aerosol, which corresponds to a monolayer surface coverage of $\sim$2 x 10$^{14}$ molecules cm$^{-2}$, suppresses the N$_2$O$_5$ reaction probability, $\gamma_{N_2O_5}$, by approximately a factor of ten, independent of relative humidity. Consistent with this observation is a similar reduction in the rate of ClNO$_2$ product generation measured simultaneously. However, the product yield remains nearly constant under all conditions. The degree of suppression is strongly dependent on SDS content in the aerosol, with no discernable effect at 0.1 wt% SDS, but significant suppression at what we predict to be submonolayer coverages with 0.3–0.6 wt% SDS on NaCl and natural seawater aerosols, respectively.

1 Introduction

Organic material is a major component of natural and anthropogenic atmospheric aerosols (McMurry, 2000). In the past few decades, hundreds of organic species found in atmospheric particles have been identified, although the majority of organic aerosol mass often remains unspeciated (Saxena and Hildemann, 1996; McMurry, 2000; Jacobson et al., 2000). There has been a renewed interest in the sources and fates of surface active organic molecules due to their potential impact on gas-aerosol interfacial chemistry. Surface active films were first extracted from sea salt aerosol by Blanchard (1964) over 40 years ago. Recent studies have shown the presence of fatty acids in marine aerosol sampled over the North Atlantic (O’Dowd et al., 2004; Cavalli et al., 2004) and northern North Pacific (Mochida et al., 2002), with maximum concentrations occurring during periods of high biological productivity. Soft X-ray spectromicroscopy (Russell et al., 2002) and TOF-SIMS (Tervahattu et al., 2002; Petersson and Tyler, 2003) have revealed organic coatings on dried sea salt particles. Tervahattu et al. (2002) used TOF-SIMS to identify the main component of the organic layer as palmitic acid. It has been postulated that marine aerosol may obtain a surfactant component from an organic-rich surface layer that is present on seawater during periods of high biological activity via a bubble-bursting mechanism (Gershey, 1983; Mochida et al., 2002). On aqueous atmospheric aerosol, these surfactant molecules are thought to form an organic surface layer in an “inverted micelle” configuration, possibly affecting the transport of molecules across the air-aerosol interface and surface reactions (Gill et al., 1983; Ellison et al., 1999).

While it is well-known that the presence of a long-chain fatty acid layer inhibits the evaporation of a macroscopic water film (Rideal, 1925; Archer and La Mer, 1955; Rosano and La Mer, 1956), the effect of surface active organics on the interaction of aerosols with gas phase species, particularly in reactive systems, remains poorly characterized. In a recent molecular beam scattering (TOF-MS) study, Lawrence et al. (2005b) found that a butanol coating on supercooled deuterated sulfuric acid films (T=213 K) did not inhibit proton exchange by HCl and HBr, and in fact suggest that the hydrophilic head groups of the butanol may provide additional...
sites for interaction with the adsorbate. Short-chained surfactants such as butanol are known to form less tightly packed films and provide a lower barrier to water evaporation than longer-chained surfactants (Archer and La Mer, 1955; Rosano and La Mer, 1956). Monolayer levels of fatty acids at the air-water interface were found by Mmereki et al. (2004) to inhibit the Langmuir-Hinshelwood surface reaction between ozone and adsorbed anthracene, a polycyclic aromatic hydrocarbon, while a monolayer of 1-octanol was observed to enhance the reaction rate (Mmereki and Donaldson, 2003). Most recently, Thornton and Abbatt (2005) found that the presence of hexanoic acid decreased the N₂O₅ reactive uptake coefficient on artificial seawater (ASW) aerosol by a factor of 3–4 at 70% relative humidity (RH). To our knowledge, these are the only studies of the effect of monolayer organic films on gas-aerosol reactive uptake kinetics, though it has been shown that a multilayer hydrophobic shell (≥15 nm) on aqueous inorganic aerosol causes a significant reduction in the reactive uptake of N₂O₅ (Folkters et al., 2003).

In a continuing effort to elucidate the impact of aerosol organic matter on important heterogeneous reactions, we examined the reaction of N₂O₃ on aqueous natural seawater and NaCl aerosol containing trace amounts (∼10 wt%) of sodium dodecyl sulfate, SDS (C₁₂H₂₅SO₃Na). Our goal was to determine the effect of sub-monolayer surfactant coatings on the kinetics and product yields of this reaction. Our choice of aqueous natural seawater and NaCl aerosol was motivated in part by the fact that the reaction of N₂O₅ with aqueous sea salt aerosols is a sink for nitrogen oxides and a potential source of the chlorine radical in the polluted marine boundary layer. Although discrepancies remain with regard to the product yields of this reaction on aerosols (Behnke et al., 1997; Schweitzer et al., 1998; Thornton and Abbatt, 2005), the reaction is relatively well-characterized (Behnke et al., 1997; Schweitzer et al., 1998; Hoffmann et al., 2003; Stewart et al., 2004; Thornton and Abbatt, 2005) and provides a useful starting point to assess the role of surfactants. The ClNO₂ product, which volatilizes readily from submicron aerosol, provides an additional indicator of the reaction pathways occurring in the aerosol.

Given the lack of direct kinetic studies of gas-aerosol interactions where surfactants are present, SDS was chosen as a proxy for natural organic surfactant molecules because it is readily soluble in water and because the SDS-NaCl-H₂O system has been characterized in bulk solutions (Matijevic and Pethica, 1958; Rehfeld, 1967; Persson et al., 2003; Sorjamaa et al., 2004; Li et al., 1998). SDS is similar to fatty acids in that it contains a long aliphatic tail on a polar head group whose anionic character is likely pH dependent. The aliphatic tail of SDS is analogous to that of lauric acid (C₁₂), which has been identified in both natural marine (Mayol-Bracero et al., 2001) and continental aerosols (Gill et al., 1983; Duce et al., 1983; Rogge et al., 1993). We find that the N₂O₅ reaction probability decreases with increasing SDS mass fraction, and eventually plateaus at 3.5 wt% SDS, beyond which no further decrease is observed. These results suggest that even small quantities of a C₁₂ surfactant on the surface of aqueous atmospheric aerosol have the potential to inhibit the rates of gas-aerosol mass transfer and heterogeneous chemistry.

2 Experimental

Experiments were performed using a recently constructed aerosol flow tube coupled to a custom-built chemical ionization mass spectrometer (CIMS) for monitoring the gas phase, and a differential mobility analyzer/condensation particle counter (Grimm Technologies) for determining aerosol surface area. The experimental setup and procedure closely resembles that described in detail previously (Thornton et al., 2003; Thornton and Abbatt, 2005), but we highlight significant differences below.

2.1 Aerosol generation and characterization

Sub-micron aqueous NaCl aerosol were generated by atomizing a 0.05 M solution of NaCl in deionized water using a commercial constant output atomizer (TSI). We chose to use NaCl in addition to natural seawater due to lower wall losses of N₂O₅ which allow us to better quantify slow aerosol reactions. The presence of Mg²⁺ in seawater aerosol appears to lead to a greater retention of water on the reactor walls upon aerosol impaction (Cziczo and Abbatt, 2000).

To generate SDS-doped NaCl aerosol, small quantities of a stock solution prepared from SDS crystals and deionized water were added to the 0.05 M NaCl solution to create mixtures with 10⁻⁵–10⁻³ M SDS which were then atomized. Similarly, sub-micron natural seawater (NSW) aerosol were generated by atomizing a dilute solution of natural seawater in deionized water (1:7.3). This dilution was chosen to create a 0.05 M solution assuming a NaCl concentration of 0.415 M in natural seawater. The sample was taken from the Caribbean Sea (18°30.09′ N, 64°21.47′ W) in July 2005. SDS-doped NSW aerosol was generated using the same procedure as the SDS-doped NaCl aerosol.

The atomizer output (~1 slpm) was mixed with a ~4 slpm humidified N₂ flow and transported (~10 s) to the kinetics flow tube. The RH of the equilibrated N₂/aerosol stream was monitored at the inlet to the aerosol flow tube with a commercial hygrometer (Fisher Scientific, Vaisala). Reported values for RH are accurate to within 2%. The humidified aerosol stream amounted to >94% of the total flow in the flow tube, and thus the RH inside the reactor is assumed to be that of the aerosol flow. Experiments were performed at 50%–72% RH. We did not perform experiments at RH <50% because NaCl aerosol crystallize at 43% RH (Cziczo and Abbatt, 2000) and our goal was to assess the effect of SDS on aqueous particles.

Because water evaporates from aerosol when the 100% RH atomizer output combines with the drier bulk flow, the
aerosol become greatly concentrated relative to the bulk atomizer solution. Following Tang et al. (1997), at 62% RH, an aqueous NaCl aerosol will be 8.8 M NaCl. Assuming the same proportional increase in concentration (176x), we estimate that 0.001 M SDS in the atomizer solution corresponds to 0.176 M SDS in the aerosol. Based on a a volume of 1.1×10⁻¹⁷ L calculated using the surface area weighted average particle radius, 0.176 M SDS corresponds to an average of 1.2×10⁸ SDS molecules/particle. To avoid confusion concerning bulk-surface partitioning of SDS and the total amount of SDS available per particle, we will refer to SDS concentration as a solute mass fraction (wt%), instead of molarity. Thus, atomizing a solution containing 0.001 M SDS yields aerosol at 62% RH that are 10 wt% SDS.

The aerosol population was continuously sampled at the exit of the kinetics flow tube by a differential mobility analyzer (DMA) coupled to an ultra-fine condensation particle counter (CPC). So that the RH of the DMA sheath flow was similar to the sample flow, the DMA/CPC drew humidified bulk flow for ~1 h prior to starting experiments. The aerosol, with and without SDS present, exhibited a log-normal size distribution with a typical geometric standard deviation of 2.0. NaCl aerosol had an average mean surface area-weighted particle radius, \( r_p \), of 140±10 nm, while for NSW aerosol, \( r_p \) = 160±10 nm. Representative number and surface area-weighted aerosol size distributions for aqueous NaCl aerosol at 61% RH with and without 10 wt% SDS are shown in Fig. 1. Total surface area was typically between 9×10⁻⁴–2.2×10⁻³ cm² cm⁻³ for NaCl and NSW aerosol and between 5×10⁻⁴–1.5×10⁻³ cm² cm⁻³ with SDS present. The lower surface area concentrations were used with the SDS mixtures to minimize foaming in the atomizer outlet. In addition, at the highest surface area concentrations generated with the SDS mixtures, there was a noticeable shift in the \( r_p \) to higher sizes (300–400 nm). As the DMA is unable to accurately size aerosol above ~800 nm, the measured surface area under these conditions may be an underestimate causing our reported reaction probabilities to be biased high by at most 50%. This issue does not impact our conclusions.

2.2 \( \text{N}_2\text{O}_3 \) synthesis and delivery

\( \text{N}_2\text{O}_3 \) was synthesized using NO₂ and O₃, and was collected at ~83°C. O₃ was generated using a pen-ray lamp (UVP). The \( \text{N}_2\text{O}_3 \) sample was stored at ~50°C using a cold finger (NESLAB). During experiments, the sample was kept between ~70°C and ~74°C. The vapor pressure of \( \text{N}_2\text{O}_3 \) at ~70°C is estimated to be 0.006±0.003 Torr (Thornton et al., 2003). \( \text{N}_2\text{O}_3 \) was delivered to the flow tube by flowing 5 sccm dry \( \text{N}_2 \) through the trap. The saturated flow was then combined with a 100 sccm \( \text{N}_2 \) carrier flow and transported through ~3 m of 3 mm OD PFA tubing into the flow tube reactor. Typical initial concentrations in the flow reactor were 5.5×10¹¹ molecules cm⁻³ (22 ppb).

![Fig. 1](image-url) Representative number (top panel) and surface area-weighted (bottom panel) aerosol size distributions for aqueous NaCl aerosol at 61% RH with and without 10 wt% SDS present. Both populations represent ~1.2×10⁻³ cm² cm⁻³ total aerosol surface area.

2.3 Flow tube reactor

The kinetics flow tube reactor consisted of a vertically oriented Pyrex tube, 90 cm long, with an internal radius of 1.5 cm. To minimize wall reactions, the inner wall of the flow tube was coated with halocarbon wax and rinsed with water and methanol between sets of experiments. The RH-equilibrated aerosol stream was introduced at the top of the flow tube through a perpendicular side-arm. \( \text{N}_2\text{O}_3 \) was introduced via an injector consisting of a 3 mm OD PFA tube housed inside a moveable 6 mm stainless steel tube for structural support. The injector position was changed to vary reaction time and thus obtain kinetic information. All experiments were performed at ambient temperature (295 K) and pressure (760 Torr). The flow velocity down the tube was maintained by a critical orifice located at the bottom of the flow tube, in front of the low pressure CIMS region, which drew 1.6 slpm. An additional 0.3 slpm was drawn by the DMA/CPC. Thus the total flow in the flow tube was 1.9 slpm, resulting in a flow velocity of 4.5 cm s⁻¹. In order to study kinetics under well-mixed, fully developed laminar flow conditions, experiments were limited to the region between 30–75 cm.

[1635–1644, 2006]
2.4 CIMS detection of reactants and products

\( \text{N}_2\text{O}_5 \) and \( \text{ClNO}_2 \) were detected via CIMS using \( \Gamma^- \) as a reagent ion. \( \Gamma^- \) was generated by flowing trace amounts of \( \text{CH}_3\text{I} \) in 0.8 slpm \( \text{N}_2 \) through a \( ^{210}\text{Po} \) ionizer (NRD) into the chemical ionization region which consists of a 3.8 cm OD stainless steel tube manifold. The ionizer was positioned axially inside this manifold, with the outlet ~2.5 cm away from the mass spectrometer front aperture. Flow from the kinetics reactor (1.6 slpm) was introduced upstream of the ionizer outlet. The pressure in the CIMS region was ~60 Torr, yielding a reaction time for ion-neutral reactions of ~20 ms.

\( \text{N}_2\text{O}_5 \) was monitored as \( \text{NO}_3^- \) (62 amu), with a sensitivity at 50% \( RH \) of 1.6 cps ppt\(^{-1}\). A higher sensitivity can be achieved, but in order to reduce background signal at 62 amu attributed to \( \text{HNO}_3 \) (Thornton et al., 2003), ion-neutral reaction time was shortened and the electric field inside the collisional dissociation chamber (CDC) was altered from that which provided optimal \( \text{N}_2\text{O}_5 \) sensitivity. \( \text{ClNO}_2 \) was monitored as \( \text{ICl}^- \) (161.9 amu and 163.9 amu), as in Thornton and Abbatt (2005), and also \( \Gamma^-\text{ClNO}_2 \) (207.9 and 209.9 amu), the association reaction product with \( \Gamma^- \). The \( \text{ICl}^- \) signal was a constant fraction of the \( \Gamma^-\text{ClNO}_2 \) signal (207.9 amu: 161.9 amu=1.9:1), although this ratio can be expected to depend on the electric field in the CDC. In a control experiment, we found that HCl vapor reacts with \( \Gamma^- \) to form \( \Gamma^-\text{HCl} \) under our operating conditions. These two observations suggest that the \( \text{ICl}^- \) signal is uniquely attributable to \( \text{ClNO}_2 \) (Thornton and Abbatt, 2005).

3 Results

Shown in Fig. 2a is a set of \( \text{N}_2\text{O}_5 \) decays obtained during experiments at 62% \( RH \) for which 22 ppb \( \text{N}_2\text{O}_5 \) was initially present. Data are shown for \( \text{N}_2\text{O}_5 \) loss in the absence of aerosol, or “wall loss” (triangles), reaction with NaCl aerosol (diamonds), and reaction with NaCl aerosol containing 10 wt% SDS (open squares). For both aerosol decays, aerosol surface area was \( S_a=1.3\times10^{-3} \text{ cm}^2 \text{ cm}^{-3} \) for both aerosol decays, and the wall loss was obtained immediately prior to the SDS-doped aerosol run.

\[
\frac{d^2 S_{\text{ICl}}} {dt^2} = -k_p S_{\text{ICl}}
\]

where \( S_{\text{ICl}}^\infty \) and \( k_p \) are adjustable parameters. For first order loss to multiple products, we can expect \( k_p=k^f \) (Steinfeld et al., 1999). The first order rate constants obtained from \( \text{ClNO}_2 \) growth curves and Eq. (2) are in close agreement with those obtained from \( \text{N}_2\text{O}_5 \) decays, exhibiting similar trends.
in surface area, with slightly more scatter. On average, $k_p$ were within 20% of the $k_f$ determined simultaneously.

The corrected first order rate constant, $k_f$, is related to the reaction probability, $\gamma_{\text{obs}}^{\text{N}_2\text{O}_5}$, according to

$$k_f = \frac{\gamma_{\text{obs}}^{\text{N}_2\text{O}_5} \omega S_a}{4}$$

where $\omega$ is the molecular velocity of $\text{N}_2\text{O}_5$ and $S_a$ is the aerosol surface area per volume. Equation (2) is only an approximation, but effects due to gas-phase diffusion limitations and modification of the collision frequency due to a net reactive flux to the aerosol are negligible for the small aerosol and moderate reactive uptake coefficients determined here. Shown in Fig. 3 are the corrected first order rate constants for reaction of $\text{N}_2\text{O}_5$ with aqueous NaCl aerosol (diamonds) and aqueous NaCl aerosol containing 10 wt% SDS (open squares) as a function of aerosol surface area. Data obtained at various relative humidities are shown together on the same plot, since no relative humidity dependence was observed for $RH$ above 50%, with or without SDS present (see Table 1). Error bars shown represent the 1σ standard deviation in the raw data propagated through the weighted linear least squares fit and the Brown correction. The lines shown were obtained from weighted linear least squares fits to the data. The 1σ confidence limits for the fits are also shown as grey lines. For reaction with pure aqueous NaCl aerosol, the slope of the weighted linear least squares fit corresponds to a reaction probability of 0.015±0.002, and for aqueous NaCl aerosol containing 10 wt% SDS, the slope yields $\gamma_{\text{N}_2\text{O}_5}^{\text{NaCl}}$=0.002±0.001. A similar analysis for NSW aerosol yields a $\gamma_{\text{N}_2\text{O}_5}^{\text{NSW}}$ of 0.034±0.002 at 60% $RH$ in the absence of SDS, roughly twice that for aqueous NaCl aerosol. Table 1 summarizes $\gamma_{\text{N}_2\text{O}_5}$ determined from individual decays and Eq. (2) for the reaction of $\text{N}_2\text{O}_5$ with aqueous NaCl aerosol, NSW aerosol, and these aerosols containing 10 wt% SDS, measured at ~51%, ~60%, and 71% $RH$.

Figure 4a shows $\gamma_{\text{N}_2\text{O}_5}$ determined at ~60% $RH$ for both aqueous NaCl (diamonds) and NSW aerosol (circles) versus SDS concentration, which ranges from 0.1 to 10 wt% (0.00176 M–0.176 M SDS) depending on the SDS concentration used in the atomizer solution. For comparison, NaCl is estimated to be 8.8 M (Tang et al., 1997) under these conditions. For aqueous NaCl aerosol, when 0.1 wt% SDS is present in the aerosol, no effect is observed, i.e. $\gamma_{\text{N}_2\text{O}_5}$ is the same as when no SDS is present. Starting at 0.3 wt% SDS in the aerosol, $\gamma_{\text{N}_2\text{O}_5}$ is reduced by a factor of 3 relative to no SDS present, and decreases further with increasing SDS concentration until ~3.5 wt% SDS, where $\gamma_{\text{N}_2\text{O}_5}$~0.002±0.001. Above 3.5 wt% SDS, no further decrease in $\gamma_{\text{N}_2\text{O}_5}$ is observed. A similar suppression in $\gamma_{\text{N}_2\text{O}_5}$ by SDS is observed for aqueous NSW aerosol: at 5.5 wt% SDS, $\gamma_{\text{N}_2\text{O}_5}$ is reduced by a factor of 3 relative to no SDS present, and ultimately $\gamma_{\text{N}_2\text{O}_5}$~0.001±0.001 at 10 wt% SDS. Figure 4b shows $\gamma_{\text{N}_2\text{O}_5}$ plotted versus estimated overall fractional surface coverage, $\Theta$, which takes into account the distribution in surface coverages present in each experiment as a result of the polydisperse nature of our aerosol. We discuss our approach to calculating $\Theta$ in Sect. 4.2.

![Graph](image.png)

**Fig. 3.** Corrected first order rate constants for reaction of $\text{N}_2\text{O}_5$ with NaCl aerosol (♦) and reaction with NaCl aerosol containing 10 wt% SDS (□) as a function of measured aerosol surface area. Lines shown represent weighted linear least squares fits to the data, with confidence limits of 1σ. For all experiments shown, ~22 ppb $\text{N}_2\text{O}_5$ was present initially. Data obtained at 51%, 62%, and 72% $RH$ are shown.

| Aerosol Type       | $RH$ (%) | $\gamma_{\text{N}_2\text{O}_5}$ |
|--------------------|----------|--------------------------------|
| NaCl               | 51       | 0.016±0.001                    |
| NaCl               | 61       | 0.015±0.001                    |
| NaCl               | 71       | 0.015±0.001                    |
| NaCl/10 wt% SDS    | 50       | 0.00002±0.0001                 |
| NaCl/10 wt% SDS    | 61       | 0.002±0.001                    |
| NaCl/10 wt% SDS    | 71       | 0.005±0.001                    |
| NSW                | 60       | 0.034±0.002                    |
| NSW/10 wt% SDS     | 60       | 0.001±0.001                    |

The ClNO$_2$ product yield, $\gamma_{\text{ClNO}_2}$, was calculated for each decay from the change in $\text{N}_2\text{O}_5$ signal and the change in ClNO$_2$ signal taking into account the combined contributions of the four signals attributable to ClNO$_2$ signal at 162 amu,
That ClNO₂ was observed at all in the presence of monolayer coverages of SDS suggests N₂O₅ does penetrate to the particle bulk and γ\textsuperscript{N₂O₅} \textgreater 0 on the timescales of our experiments.

4 Discussion and atmospheric implications

4.1 Kinetics and product yields on pure aqueous NaCl and NSW aerosols

Our measured reactive uptake coefficients for N₂O₅ on aqueous NaCl aerosols fall in between those reported in other studies, which range from \sim 0.004–0.03 (Behnke et al., 1997; Schweitzer et al., 1998; Stewart et al., 2004). Behnke et al. (1997) and Stewart et al. (2004) also reported γ\textsuperscript{N₂O₅} to be independent of RH above 50% RH on aqueous NaCl aerosol. Thornton and Abbatt (2005) measured γ\textsuperscript{N₂O₅} = 0.03±0.008 on artificial seawater, in close agreement with our measured value of γ\textsuperscript{N₂O₅} = 0.034±0.002 for aqueous NSW aerosol at 60% RH. Our finding that N₂O₅ is more reactive on aqueous NSW aerosol than on aqueous NaCl aerosol is similar to the findings of Stewart et al. (2004), although the absolute values of our measured γ\textsuperscript{N₂O₅} are not in good agreement with theirs. It is somewhat surprising that the reaction probability is larger by nearly a factor of 2 on NSW aerosol than on NaCl aerosol, especially when both systems are aqueous. The liquid water content will be very similar for the two aqueous systems, and thus suggests the additional solutes present in seawater (Mg²⁺, Br⁻, etc.) enhance the reactive uptake rate.

It is not surprising that we did not observe signs of suppression of N₂O₅ uptake due to naturally occurring surfactants on the surface of the NSW aerosol used in this study. The seawater was taken from an area with low biological activity and diluted \sim 10× before atomization. Furthermore, given that the atomizer draws from the bulk of the solution, and that surface active species in seawater are most likely water insoluble (e.g. stearic or palmitic acid), any such surfactants would not be present in the aerosol generated by atomization. We can’t rule out the possibility that we might observe suppression if the aerosol were generated via a bubble-breaking mechanism from undiluted seawater sampled during a period of high biological activity (Gershey, 1983; Mochida et al., 2002).

The following mechanism has been proposed for the reaction of N₂O₅ with aqueous NaCl and sea salt aerosols (Thornton et al., 2003; Thornton and Abbatt, 2005):

\begin{equation}
\text{N}_2\text{O}_5 \ (g) \rightarrow \text{N}_2\text{O}_5 \ (aq) \quad (4)
\end{equation}

\begin{equation}
\text{N}_2\text{O}_5 \ (aq) + \text{H}_2\text{O} \leftrightarrow \text{H}_2\text{ONO}_2^+ \ (aq) + \text{NO}_3^- \ (aq) \quad (5)
\end{equation}

\begin{equation}
\text{H}_2\text{ONO}_2^+ \ (aq) + \text{H}_2\text{O} \rightarrow \text{H}_3\text{O}^+ \ (aq) + \text{HNO}_3 \ (aq) \quad (6)
\end{equation}

\begin{equation}
\text{H}_2\text{ONO}_2^+ \ (aq) + \text{Cl}^- \ (aq) \rightarrow \text{ClNO}_2 \ (aq) + \text{H}_2\text{O} \quad (7)
\end{equation}
Our observed ClNO$_2$ yield of ~90% for N$_2$O$_5$ reaction on aqueous NaCl aerosol is consistent with yields of unity reported by Schweitzer et al. (1998), Hoffman et al. (2003), and Behnke et al. (1997) (for their wetted wall flow tube experiments). Behnke et al. (1997) also reported a yield of 65% for their smog chamber measurements using aqueous NaCl aerosol, and attributed the lower value to ClNO$_2$ loss on the Teflon walls of the chamber. Thornton and Abbatt (2005) reported a yield of 50±10% on artificial seawater aerosol at a single RH, based only on observations of ICl$^-$. We use a very similar approach and detection technique as was used in that study, except that we were able to measure I$^-$/ClNO$_2$ as well. Given that our present yield determinations of 80% are lower limits, it is likely the true yield on sea salt aerosol is greater than 80% (but less than 100%). N$_2$O$_5$ reaction on sea salt aerosol may also produce BrNO$_2$ (Schweitzer et al., 1998). However, ClNO$_2$ was the only halogen product detected in this study when using NSW aerosol. According to the above mechanism, a ClNO$_2$ yield of near unity implies that $K_k$ for the reaction $\gamma N_2O_5 \rightarrow \gamma K_k x [H_2O]$ (Steinfeld et al., 1999), Thornton et al. (2003) determined a second order hydrolysis rate constant of $2.6 \times 10^4$ M$^{-1}$ s$^{-1}$, equivalent to the product $K_k$s in this mechanism. Using the aerosol concentrations of H$_2$O and Cl$^-$ at 70% RH, and assuming a value of 2 M atm$^{-1}$ for the Henry’s law constant of N$_2$O$_5$ (Fried et al., 1994; Jacob, 2000), we estimate that $K_k k_7 \gg 1.8 \times 10^7$ M$^{-1}$ s$^{-1}$, or $k_7/k_6 > 10$. $K_k$ is not known. Such estimates of aqueous phase rate constants are highly uncertain as they carry assumptions about solubility and that mass accommodation is not rate-limiting.

4.2 Effect of aerosol surfactant levels

The presence of SDS leads to a suppression of $\gamma N_2O_5$ for both NaCl and NSW aerosol. We observe the onset of suppression, in terms of SDS concentration, to differ between NaCl and NSW aerosol. While some of this difference can be explained by uncertainty in the atomizer SDS concentration, it is possible that the onset of suppression depends partly on the bulk composition of the aerosol. In both systems, $\gamma N_2O_5$ decreases with increasing SDS mass fraction until about 3.5 wt% SDS, beyond which no further decrease is observed. This behaviour is suggestive of a monolayer of SDS forming at the gas-aerosol interface, beyond which no further suppression is possible. We examine this aspect in more detail below, but note here that the low values of $\gamma N_2O_5$ observed at 3.5–10 wt% SDS are near the limit of detection due to wall loss and the precision in individual decays, so it is possible that $\gamma N_2O_5$ could continue to decrease for SDS >3.5 wt%. The mechanism of suppression could be that SDS inhibits either N$_2$O$_5$ uptake from the gas to the bulk aqueous phase, or reactivity by disrupting the distribution of reactants in the bulk or near-surface regions. The fact that the ClNO$_2$ yield is the same with or without SDS present indicates that both N$_2$O$_5$ hydrolysis (Reactions 5 and 6) and the reaction with Cl$^-$ (Reactions 5 and 7) are inhibited equally. Thus, we conclude that the main effect of the surface organic film is to inhibit the uptake of N$_2$O$_5$ from the gas phase by the aqueous bulk.

Surfactants in aqueous solution will partition to the surface until the surface is saturated (Myers, 1988), and in any model system, surfactant concentration at the surface must be treated separately from the bulk monomer concentration (Myers, 1988; Sorjamaa et al., 2004). For ionic single chain surfactants, like SDS or fatty acids, the head group determines how closely the surfactant molecules can pack at the surface (Myers, 1988). The sulfate head group of SDS is most similar to a carboxylic acid, although surfactants typically present in atmospheric aerosol can also have alcohol, aldehyde, ketone, ester, or amine head groups (Gill et al., 1983; Mochida et al., 2002; Peterson and Tyler, 2003; Tervahattu et al., 2002; Russell et al., 2002). Based on studies of dilute SDS-NaCl-H$_2$O mixtures (Matijevic and Pethica, 1958; Persson et al., 2003), we can assume an SDS footprint of ~50 Å$^2$, corresponding to a saturation surface coverage of $2 \times 10^{14}$ molecules cm$^{-2}$. At 3.5 wt% SDS, the concentration beyond which we did not observe additional suppression, we predict a surface coverage of $\sim 2 \times 10^{14}$ molecules cm$^{-2}$, assuming all SDS partitions to the aerosol surface, and a typical particle surface area of $2.5 \times 10^{14}$ cm$^{-2}$. At the highest SDS concentrations used in this study (10 wt%) we predict a surface coverage of $\sim 5 \times 10^{14}$ molecules cm$^{-2}$ if all the SDS partitioned to the surface. This coverage would require an SDS footprint of $\sim 20$ Å$^2$, which is the tight-packing “Pockels limit” (Pockels, 1891). Whether such a tightly packed film persists in our aerosol or whether SDS loading of the aqueous bulk and possibly micelle formation has begun above 3.5 wt%, we cannot say, but this issue is only relevant for the highest SDS concentration used in this study.

Micelles will form in the solution at bulk surfactant monomer concentrations above what is known as the critical micelle concentration (cmc). Following trends in the cmc of SDS as determined for dilute aqueous NaCl solutions, we expect the cmc to be much less than $10^{-3}$ M SDS in our 9 M NaCl aerosol (Li et al., 1998; Matijevic and Pethica, 1958). But because of the small size and thus high surface area to volume ratio of our droplets, most SDS molecules present in the aerosol are partitioned to the surface, resulting in bulk monomer concentrations well below the cmc for all runs except that using 10 wt% SDS. If micelle formation has begun at this point, it appears to have no effect on the ability of the monolayer film to act as a barrier to N$_2$O$_5$ uptake, as the observed $\gamma N_2O_5$ remains nearly a factor of 10 lower than in runs without SDS.

It is particularly intriguing that we observe significant suppression in $\gamma N_2O_5$ on both NaCl and NSW aerosol with what can only be sub-monolayer surface coverages of SDS. Due to the polydisperse nature of our aerosol sample, we must take into account a distribution of surface area to volume ratios present for a single population, resulting in a distribution of
SDS surface coverages for a given SDS concentration. If we define \( \theta_i \) to be the fractional surface coverage of the aerosol in the DMA size bin \( i \), we can estimate the fraction of the total available surface area of our poydisperse aerosol population that is occupied by SDS molecules according to:

\[
\Theta = \frac{\sum \theta_i N_i}{\sum N_i},
\]

where \( N_i \) is the number density in bin \( i \). Figure 4b shows the data from Fig. 4a plotted as a function of overall fractional surface coverage, \( \Theta \), calculated for each point according to Eq. (8), and assuming that all surfactant in the aerosol partitions to the surface until a saturation surface coverage is reached. As noted above, based on the behaviour of \( \gamma_{N_2O_5} \) we estimate that monolayer coverage corresponds to \( 2 \times 10^{14} \) molecules cm\(^{-2} \) (3.5 wt\% SDS on a 150 nm radius particle). For a given SDS concentration, the standard deviation of \( \Theta \) due to variation in the particle size distributions between decays was found to be less than 5% of the \( \Theta \) value. We observe roughly a factor of 3 suppression in \( \gamma_{N_2O_5} \) at \( \sim 8\% \) of a monolayer and 15% of a monolayer for NaCl and NSW aerosol, respectively. If adding SDS molecules to the aerosol surface caused a linear decrease in the accessible aerosol surface area, from Eq. (2) we might expect a linear reduction in \( \gamma_{N_2O_5} \) with increasing SDS surface coverage. We would then predict a factor of 3 reduction in \( \gamma_{N_2O_5} \) to correspond to \( \sim 65\% \) of a monolayer, i.e. a factor of \( \sim 5 \) greater coverage than that at which we observe the reduction. In order to produce a linear dependence of \( \gamma_{N_2O_5} \) on \( \Theta \), it is necessary to set the saturation surface coverage to \( 5 \times 10^{15} \) molecules cm\(^{-2} \), a factor of 4 lower than the saturation coverages reported in the literature (Matijevic and Pethica, 1958; Persson et al., 2003). While we can’t rule out a larger average footprint for SDS in the highly ionic aerosol used, it seems unlikely for the footprint to be \( \sim 200 \) \( \AA \)\(^2 \).

Another possible explanation for the observed non-linearity in \( \gamma_{N_2O_5} \) versus \( \Theta \) is that fatty acids with 12–15 carbon atoms form “compressible” or “expanded” films (Schofield and Rideal, 1926). Expanded films can be thought of as a disordered transition phase between sparse surface coverage, in which the molecules behave like a 2-D gas, and a tightly packed “incompressible” inverted micelle configuration. Fatty acids with 16 or more carbon atoms form only incompressible films (Schofield and Rideal, 1926; Rosano and La Mer, 1956). We expect SDS to behave similarly to lauric acid (C\(_{12}\)) and form an expanded film at sub-monolayer surface coverages. Therefore, the degree of packing and thus the SDS “footprint” will change with increasing surface coverage, resulting in a nonlinear relationship between the number of molecules of SDS on the surface and the degree of inhibition of gas-to-aerosol mass transfer. We know of no other studies showing a reduction in gas-aerosol mass transfer reactive uptake to aerosol due to sub-monolayer coverages of a surface active organic, but our observations in this regard are consistent with measurements of water evaporation which show that lauric acid in the expanded state exhibits a resistance to mass transfer (Rideal, 1925).

Thornton and Abbatt (2005) found \( \gamma_{N_2O_5} \) to decrease by a factor of 3–4 in the presence of what they estimated to be monolayer coverage of hexanoic acid. At what we interpret to be full coverage of SDS, we observe greater inhibition of \( N_2O_5 \) uptake by a factor of 3 or more relative to their results. Shorter-chain fatty acids such as hexanoic acid are expected to form less tightly packed films (Schofield and Rideal, 1926; Lawrence et al., 2005; Lawrence et al., 2005). Water evaporation studies have found that the resistance to evaporation of water through a fatty acid film was proportional to the compressibility of the film (Rosano and La Mer, 1956), and generally, resistance increases with increasing carbon chain length. Although we do not have molecular-level information about the surface films in this experiment or those of Thornton and Abbatt (2005), it seems reasonable to conclude that the same resistance behaviour is true for \( N_2O_5 \) transfer to the aerosol surface.

4.3 Atmospheric implications

This work, together with the studies available in the literature, supports the conclusion that the presence of a sufficiently long-chain surfactant coating creates a resistance to the transfer of gas phase species to and from the aerosol surface (Andrews and Larson, 1993; Thornton and Abbatt, 2005; Garland et al., 2005). Our results suggest that even small quantities of C\(_{12}\)-C\(_{15}\) surfactants on the surface of aqueous atmospheric aerosol have the potential to inhibit surface processing via formation of expanded films. We conclude that, under conditions where an effective surfactant barrier on aqueous aerosol exists, NO\(_x\) lifetimes will increase, and in the case of marine aerosol, halogen activation by N\(_2O_5\) will be suppressed. Such films could also affect the rate of other important multiphase processes, such as SO\(_2\) oxidation. The potential for \( \gamma_{N_2O_5} \) to be highly variable and dependent on the speciation and mixing state of aerosol organic matter poses a challenge for accurate parameterization of this process in models. It appears that \( \gamma_{N_2O_5} \) is a useful indicator of organic aerosol phase and mixing states, and together with detailed information on aerosol composition, could be used to infer the prevalence of surfactant barriers in atmospheric aerosol.

Water evaporation studies indicate that long-chain fatty acids could be even more efficient barriers to gas-aerosol mass transfer (Rosano and La Mer, 1956) than the shorter-chained proxy surfactants used in this study and Thornton and Abbatt (2005). However, since long-chain fatty acids (>C\(_{16}\)) form tightly packed, incompressible films even at low surface concentrations, higher surfactant concentrations may be necessary before an effect is observed. The most
realistic scenario for atmospheric aerosol would be a mixed surfactant film containing both long- (>C\textsubscript{16}) and shorter-chained surfactants. Such films have been shown to have a resistance to water evaporation intermediate between those of the single-component films (Rosano and La Mer, 1956). It is also important to note that any surfactants on aerosol surfaces would be exposed to the oxidants in the atmosphere, leading to oxidation and possible shortening of the carbon chains (Bertram et al., 2001; Molina et al., 2004). Shortening of the carbon chains of long chain fatty acids (>C\textsubscript{16}) could increase the effectiveness of the surface layer as a barrier to uptake from the gas phase by inducing the formation of expanded films. Further shortening of the chain length (<C\textsubscript{12}) would likely lead to volatilization of the surfactant layer, or reduced surface-activity, depending on the functional groups of the product and the details of the oxidation pathway. Complete volatilization of saturated alkyl surfactants would be expected to occur on a time scale of a few days, which is on the order of the expected lifetime of submicron aerosol (Molina et al., 2004). Experiments to assess the effect of film oxidation on reactive uptake are currently underway.

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