

This discussion paper is/has been under review for the journal Atmospheric Chemistry and Physics (ACP). Please refer to the corresponding final paper in ACP if available.

Pseudo steady states of HONO measured in the nocturnal marine boundary layer: a conceptual model for HONO formation on aqueous surfaces

P. Wojtal, J. D. Halla, and R. McLaren

Centre for Atmospheric Chemistry, York University, North York, ON, Canada

Received: 7 September 2010 - Accepted: 13 October 2010 - Published: 27 October 2010

Correspondence to: R. McLaren (rmclaren@yorku.ca)

Published by Copernicus Publications on behalf of the European Geosciences Union.

Discussion Paper

Discussion Paper

Discussion Paper

Discussion Paper

ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page

Abstract

Back

Introduction

Conclusions References

Tables Figures

l∢ ≯l

4 -

Full Screen / Esc

Printer-friendly Version



A complete understanding of the formation mechanism of nitrous acid (HONO) in the ambient atmosphere is complicated by a lack of understanding of processes occurring when aqueous water is present. We report nocturnal measurements of HONO, SO₂ and NO₂ by differential optical absorption spectroscopy over the ocean surface in a polluted marine environment. In this aqueous environment, we observed reproducible pseudo steady states (PSS) of HONO every night, that are fully formed shortly after sunset, much faster than seen in urban environments. During the PSS period, HONO is constant with time, independent of air mass source and independent of the concentration of NO₂. The independence of HONO on the concentration of NO₂ implies a 0° order formation process, likely on a saturated surface, with reversible partitioning of HONO to the gas phase, through vaporization and deposition to the surface. We observed median HONO/NO₂ ratios starting at 0.13 at the beginning of the PSS period (with an apparent lower bound of 0.03), rising to median levels of ~ 0.30 at the end of the PSS period (with an upper bound >1.0). The implication of these numbers is that they suggest a common surface mechanism on terrestrial and aqueous surfaces, with an increase in the HONO/NO₂ ratio with the amount of water available at the surface. The levels of HONO during the nocturnal PSS period are positively correlated with temperature, consistent with a partitioning of HONO from the surface to the gas phase with an apparent enthalpy of vaporization of ΔH_{SNI} (HONO)=55.5±5.4 kJ mole⁻¹. The formation mechanism on aqueous surfaces is independent of relative humidity (RH), despite observation of a negative HONO-RH correlation. A conceptual model for HONO formation on ambient aqueous surfaces is presented, with the main elements being the presence of a surface nanolayer (SNL), highly acidic and saturated with N(IV) precursors, production of HNO₃, that diffuses to underlying water layers, and HONO, which partitions reversibly between the SNL and the gas phase. Implications of the conceptual model are discussed.

ACPD

Discussion Paper

Discussion Paper

Discussion Paper

Discussion Paper

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I ◆ ▶I

◆ Back Close

Printer-friendly Version

Full Screen / Esc

Interactive Discussion



25154

The importance of nitrous acid (HONO) in the polluted atmosphere has been known for some time (Harris et al., 1982). Highest concentrations of HONO are typically seen in the boundary layer at night (Kitto and Harrison, 1992) due to significant loss of HONO by photolysis during the day. Photolysis of HONO is often the major source of OH in the morning boundary layer (Platt et al., 1980), and has been estimated to contribute from 17–34% of the total OH production integrated over 24 h in polluted environments (Alicke et al., 2002, 2003). Despite significant research activity and its importance, a complete understanding of the atmospheric sources and sinks of HONO has remained elusive. Some consensus has been achieved though. It is generally believed that in urban areas a direct fuel combustion source of HONO exists with a HONO/NO, ratio typically below 0.01 (Kurtenbach et al., 2001). HONO can also be formed in small amounts through heterogeneous reaction of NO2 on fresh soot aerosol surfaces (Aubin and Abbatt, 2007), an observation that is likely not independent from the fuel combustion source. More importantly, observations in the nocturnal boundary layer indicate a significant surface source of HONO (Harrison and Kitto, 1994), as indicated by negative gradients of HONO and positive gradients of NO₂ with respect to height above ground (Harrison et al., 1996; Stutz et al., 2002; Wang et al., 2006). The ground usually presents a higher surface area in the nocturnal boundary layer compared to the surface area of aerosols for NO₂ conversion (Yu et al., 2009; Stutz et al., 2004); the aerosol surface area gains relative importance with increasing height of the mixed layer (Lammel, 1999). There is also significant laboratory based evidence for a surface source of HONO originating from the heterogeneous reaction of NO2 on a range of laboratory surfaces including glass, quartz, steel and even teflon (Lammel, 1999; Finlayson-Pitts et al., 2003 and references therein). The conversion of NO₂ to HONO on surfaces is generally believed to proceed with the following stoichiometry,

$$NO_2 + NO_2 + H_2O \rightarrow HONO + HNO_3,$$
 (R1)

Discussion Paper

Discussion Paper

Discussion Paper

Discussion Paper

ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I∢ ≯I

■ Back Close

Full Screen / Esc

Printer-friendly Version



ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page **Abstract** Introduction Conclusions References **Figures** Back Full Screen / Esc

Printer-friendly Version

Interactive Discussion



with minor products NO and N₂O observed in laboratory studies, with high NO₂ levels, highly acidic surfaces and high surface to volume (S/V) ratios (Finlayson-Pitts et al., 2003).

Despite the stoichiometry of Reaction (R1), the rate of the surface reaction on these ₅ laboratory surfaces is generally believed to be first order in NO₂ (Jenkin et al., 1988) and was thought originally to be first order in water vapor as well. Recent reviews suggest that the dependence on water vapor is complicated, since the reaction is thought not to proceed in the gas phase. More correctly, relative humidity can be described as a surrogate for the amount of water adsorbed chemically or physically onto the surface where the heterogeneous reaction proceeds (Lammel and Cape, 1996; Finlayson-Pitts et al., 2003; Stutz et al., 2004).

The normalized HONO/NO₂ ratio and/or production rates in ambient air are often used as an indicator of the extent of NO₂ conversion to HONO, with an implicit assumption that the heterogeneous formation mechanism is first order in NO2 on all surface types. Ambient measurements suggest that HONO/NO₂ ratios of less than 0.01 are associated with direct fuel combustion sources (Kurtenbach et al., 2001), ratios up to 0.03 are associated with surface sources in low relative humidity environments while ratios of up to 0.10 are associated with high relative humidity (Stutz et al., 2004). A pseudo steady state (PSS) can often be achieved at night (Harrison and Kitto, 1994) that has been suggested to be a balance between the heterogeneous NO₂ to HONO conversion on the surface and simultaneous deposition of HONO on the surface (Stutz et al., 2004). Several studies mention the unusual temporal profile of constant HONO or a "levelling off" of HONO throughout the night (Lammel and Cape, 1996; Stutz et al., 2004, 2002; Yu et al., 2009), consistent with the existence of a PSS.

A compilation of HONO measurements in urban and rural areas showed that while higher nighttime levels of HONO are generally seen in the urban area, the source strength when normalized to NO₂ is similar in both areas, spanning a HONO/NO₂ range of 1.7-13% determined at the time of maximum HONO (Lammel and Cape, 1996). More recently, HONO/NO₂ ratios up to 30% have been reported (Yu et al.,

2009) in the nocturnal atmosphere of Kathmandu, Nepal where a combination of low inversion layer, high pollution and high relative humidity are invoked to explain the efficient conversion of NO₂ to HONO.

Despite the importance of surface water in the HONO/NO₂ conversion, the role of the 5 air/water interface in bulk water systems as a potential surface source for the heterogeneous reaction remains uncertain (Lammel, 1999). There is evidence that HONO can be formed heterogeneously on the surface of cloud water droplets (Acker et al., 2000). Using a liquid droplet train (Ponche et al., 1993) and liquid jet techniques (Mertes and Wahner, 1995), two studies found much higher rates of uptake of NO₂ on the aqueous surface, faster than could be accounted for by bulk-phase solubility and homogeneous reaction in solution, the latter study also showing partial release of HONO to the gas phase. In contrast, a subsequent study (Cheung et al., 2000) found no evidence for an aqueous surface reaction of NO₂ using droplet train and bubble train flow reactors; instead invoking the bulk-phase solubility and reaction for disappearance of NO₂. In a comprehensive review of all evidence of NO₂-HONO conversion in laboratory systems and outdoor atmospheres, Finlayson-Pitts et al. (2003) proposed a detailed integrated mechanism for HONO production via Reaction (R1) that involves the reaction of NO₂ (or its derivatives) on any surface, where the chemical nature of the surface is not important other than its ability to act as a physical support for thin films of surface water. The suggested mechanism involves the uptake or formation of the symmetric NO₂ dimer on the surface, N₂O₄, isomerization to the asymmetric form ONONO₂, autoionization to NO⁺NO₃, and reaction with water to generate surface adsorbed HNO₃ and gaseous HONO. It was shown therein, that the NO2 to HONO conversion in these systems could not be accounted for by bulk aqueous phase kinetics, although the question of the reaction occurring on the surface of a bulk water system was not addressed directly.

Summaries of HONO observations (Kitto and Harrison, 1992; Lammel and Cape, 1996; Yu et al., 2009) indicate that measurements have been made in a wide range of locations including urban, suburban, rural, remote and arctic atmospheres. How-

ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I4 ≯I

▼Back Close

Full Screen / Esc

Printer-friendly Version



ever, reports of HONO measurements in the marine boundary layer (MBL) are almost nonexistent, with but one brief mention of HONO measured in the open Agean Sea (Večeřa et al., 2008). The absence of HONO measurements in "aqueous" environments (Lammel, 1999), the limited study of the water dependence of HONO formation in the real atmosphere (Stutz et al., 2004), and the levels of HONO that may exist in polluted marine environments (Osthoff et al., 2008) have all been questioned recently. The presence of HONO in the polluted MBL may be important since HONO and chlorine containing photolabile species, CINO₂, CINO, Cl₂, may compete as the main radical sources in these environments in early morning, yielding OH and CI, respectively. There was early evidence that HONO is produced in the reaction of N₂O₅ with wet NaCl aerosols (Junkermann and Ibusuki, 1992). Production of CINO2 from reaction of N₂O₅ with NaCl has been observed in laboratory experiments (Finlayson-Pitts et al., 1989) and the first observations of ambient CINO₂, highly correlated with N₂O₅ were recently reported in the MBL (Osthoff et al., 2008). Still more recently, evidence was presented to show that CINO₂ is an intermediate in the production of Cl₂ from the reaction of N₂O₅ with Cl⁻ in acidic aerosols, and a hypothesized mechanism therein indicates that HONO may be produced in the reaction (Roberts et al., 2008). Thus despite a lack of measurements, HONO formation in the polluted MBL is clearly of strong interest.

In this work, we report measurements of HONO, SO₂, and NO₂ by active differential optical absorption spectroscopy (DOAS) in the MBL south of Vancouver. British Columbia, in a polluted region impacted by surrounding coastal cities and marine vessel emissions. This work continues data analysis of a spectral database from which NO₃ levels in the polluted marine boundary layer were reported (McLaren et al., 2010), establishing a link between nighttime chemistry in the MBL layer and ozone formation in the adjacent populated coastal region. With respect to potential HONO formation mechanisms, the marine boundary layer represents a unique, yet interesting environment due to the apparent simplicity of the surfaces available for HONO formation compared to terrestrial environments. The only surfaces of significance to act as **ACPD**

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page **Abstract** Introduction Conclusions References **Figures**

Full Screen / Esc

Back

source/sink of HONO are the air sea-surface interface and the surface of MBL aerosols, including deliquesced sea salt and anthropogenic aerosols. The highly aqueous surfaces in this environment are expected to be unchanging with time to a first approximation, and independent of relative humidity, compared to terrestrial urban atmospheres. We report analysis of HONO measurements for 12 nights in the MBL. The PSS that develops each night in the MBL has important implications for HONO formation mechanisms.

2 Experimental

2.1 Location and instrumentation

The location of the measurement site is shown in Fig. 1 and has been described in more detail previously (McLaren et al., 2010). Briefly, instrumentation was located at the eastern tip of Saturna Island (East Point) in the Strait of Georgia. The surrounding marine area is polluted by a variety of sources including major populated cities in British Columbia (Vancouver and Victoria) and Washington State (Bellingham and Seattle), as well as emissions from marine vessels arriving and departing from large ports in the major cities. Saturna Island itself has a low population density, especially at its eastern fringes, and has very few anthropogenic emissions that would influence the measurements. The DOAS instrumentation described in detail previously (McLaren et al., 2010) was located on a grassy patch next to the ocean at an elevation of 23 m a.s.l. Meteorological observations were taken at a height of 24.4 m a.s.l. within a few meters of the DOAS telescope (Environment Canada, 2008). The DOAS retro reflector was located on Tumbo Island (Fig. 1) at an elevation of 6 m a.s.l. and distance of 1203.5 m for a return path length of 2.41 km, with 2.16 km of the path over the open ocean at a mean elevation of 15 m a.s.l. The return beam was focused onto a bifurcated quartz fiber optic (Ocean Optics) with two 400 µm fibers, each leading to a different fiber optic spectrometer. One spectrometer was optimized for the UV spectral region (USB2000, Grating

ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page

Conclusions
Tables

Abstract

Figures

Introduction

References

4

Back

Full Screen / Esc

Printer-friendly Version



#10, 295–492 nm, 1800 lines mm $^{-1}$, 2048 element CCD, 25 µm slit, UV2 upgrade, L2 lens, optical resolution of ~ 0.5 nm), and was used for measurement of HONO, NO $_2$ and SO $_2$. The other spectrometer was optimized for measurement of NO $_3$ and will not be discussed here. A small diffuser was placed in the entrance of the fiber to lower atmospheric turbulence noise (Stutz and Platt, 1997). The spectrometers were cooled to -5° C in a portable freezer.

Custom acquisition software was written in LabView to acquire spectra from the spectrometer with typical integration times of 200–300 ms, and 2500 averages for spectra with a time resolution of 8–12 min. Mercury lamp reference spectra were collected periodically for wavelength calibration and for convoluting molecular reference spectra to the slit function of the spectrometer. Xenon lamp, electronic offset and dark noise spectra were collected periodically for use in fitting routines. The spectral measurements were collected during the period, 24 June–9 July 2005. The system was set up to collect data each evening at around sunset and ran overnight. As such, only nocturnal measurements are available here.

2.2 Spectral fitting

Each averaged ambient spectrum was corrected for electronic offset and dark noise. The net spectra were fit using DOASIS software (Kraus, 2006). HONO was fit in the wavelength region 337–358 nm. The fit scenario included an absorption reference spectrum of HONO (Bongartz et al., 1991), with cross sections reduced by 14.5% as recommended (Bongartz et al., 1994) and convoluted to the slit function of our spectrometer using a line from the Hg reference spectrum, NO₂ at 294 K (Vandaele et al., 1998), and the (O₂)₂ collision dimer (Hermans et al., 1999). SO₂ was fit in the wavelength region 303–312 nm. The fit scenario included a convoluted absorption reference spectrum of SO₂ (Vandaele et al., 1994) and NO₂ (Vandaele et al., 1998) at 294 K. NO₂ was fit in the wavelength region 422–437.5 nm. The fit scenario included the convoluted reference spectrum of NO₂ (Vandaele et al., 1998) at 294 K.

ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page

Abstract Introduction

Conclusions References

I∢ ≯I

Figures

■ Back Close

Full Screen / Esc

Printer-friendly Version



All fit scenarios included a Xenon-arc lamp spectrum, a Fraunhofer solar reference (if necessary) and a 3rd order polynomial. Detection limits (3σ) were 0.30 ppb for HONO, 0.45 ppb for SO₂, and 0.20 ppb for NO₂, determined by repetitive determination of a low concentration sample.

3 Results and discussion

An overview of the measurements made during the study is given in Fig. 2. The mixing ratio of HONO typically increased rapidly in the evening (sunset: $20:51\pm11$ min PDT) to a steady state value between 0.5 and 1.5 ppb, where it remained level until its decline in the morning (sunrise: $05:46\pm11$ min PDT) due to photolysis. The mixing ratio of SO_2 was generally about 1 ppb, apart from marine vessel plume incidents during which SO_2 increased to as high as 18 ppb, highly correlated with NO_2 . Evidence points to these events being associated with emissions from major commercial ships that were observed passing around the site throughout the study. The levels of NO_2 were as high as 50 ppb with higher levels after sunset and before sunrise, indicative of a polluted area. Relative humidity varied between 60-90% throughout the study, typical for a marine area. The temperature was at its seasonal maximum, ranging between 25° C and 14° C at night. Continuous clear to mostly clear and sunny conditions were experienced throughout the majority of the study; with a few incidents of morning haze and a 5 min period of rain on the night of 31 July/1 August.

3.1 Case study observations of HONO

Two cases studies are presented to highlight some of general features observed throughout the study. The first was the night of 24/25 July (Fig. 3). The HONO mixing ratio was low shortly before sunset, and rose quickly to just less than 1 ppb. It then stabilized at ca 0.5 ppb for the rest of the evening, before depleting after sunrise. The increase in HONO during sunset may appear to be associated with a large

ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

■ Back Close

Full Screen / Esc

Printer-friendly Version



10, 25153-25204, 2010

HONO formation on aqueous surfaces

ACPD

P. Wojtal et al.

Title Page

Abstract Introduction

Conclusions References

Figures

Back Full Screen / Esc

Printer-friendly Version

Interactive Discussion



increase in NO2 and SO2 during a marine vessel plume event, although we believe this to be coincidental timing. Generally, any changes in HONO during plume events were not statistically significant, and in certain cases, a slight anti-correlation appeared to exist between NO₂ and HONO. Levels of NO₂ were lowest in the latter part of the night on this day. Despite the low NO₂ levels during the latter period, HONO levels remained high apart from the photochemical decrease after sunrise. Two periods of relative chemical and meteorological stability were encountered during the night. The first was between 11:30 p.m. and 02:00 a.m., long after the marine emissions plume was encountered. Chemical levels were constant with NO₂~3.9 ppb, SO₂~1.3 ppb and HONO~0.50 ppb. Winds were from the SW at ~18 km/h. Apart from a few small islands, there is mostly open ocean in the SW direction for a distance of ~40 km. This implies that the air mass had traversed the open ocean for at least 2h prior to arrival at the site and prior to having encountered any major anthropogenic sources. The average HONO/NO₂ ratio during this period was 0.13 (13%). Between 02:30 and 03:30 a.m., an air mass with elevated NO₂ arrived as the wind shifted to the west. The HONO remained largely unchanged despite the change in air mass although the HONO/NO₂ ratio dropped to ~0.04 (4%) during this event. The second stable period was experienced after this from ~03:30–05:30 a.m., just before sunrise with winds from the NW and wind speeds ~5 km/h. During this second period, the pollutant levels were low; average NO₂~0.40 ppb, SO₂~1.1 ppb. It is likely that the air had travelled from the northwest down the Straight of Georgia over the ocean for a period of hours before impacting the site, and without encountering any major NO_x sources. The HONO levels were ~0.65 ppb and the HONO/NO2 ratio was highly variable and uncertain due to the uncertainty in NO2 levels close to the detection limit. The average HONO and NO₂ levels during this second stable period suggest though, that the HONO/NO₂ ratios were greater than 1.0 (100%). Despite the changes in air masses encountered between 11:30 p.m. and 05:30 p.m., HONO levels remained largely unperturbed, typical of a pseudo steady state (PSS) and did not appear to be correlated with the NO₂ concentration or the source of the air mass. The only common feature between the

3 different air masses transported to the site was their passage over the ocean in the MBL for extended periods of time. The $HONO/NO_2$ ratio was inversely correlated with NO_2 during these times, as HONO was clearly independent of NO_2 .

The first case presented here was essentially reproduced on another night, 1/2 August (not shown). On that night, the level of HONO levelled off at ~1 ppb within an hour after sunset and remained relatively constant until 07:30 a.m., despite NO₂ dropping from >15 ppb in the early night to <1 ppb after 01:30 a.m. On that morning, the onset of HONO photolysis was delayed due to hazy conditions and low light levels. For a period of 6 h (01:30–07:30 a.m.) after NO₂ dropped by an order of magnitude, the HONO, NO₂, SO₂ and HONO/NO₂ ratios were relatively constant with average (± standard deviation) levels of 1.10(±0.11) ppb; 0.90(±0.24) ppb; 0.56(±0.24) ppb and 1.83(±0.77) (mol mol⁻¹, average ratio), respectively, with wind speeds of 6–13 km h⁻¹, and a shift in wind from SW to NW at ~03:00 a.m. Thus high HONO/NO₂ ratios, greater than 100% were observed, and HONO remained constant for extended periods, even after a significant drop in NO₂ levels of one order of magnitude. There were several other examples of this type during the study.

The second night to be presented is in Fig. 4. On this night, winds were persistently from the south west, and several ship plume events were detected, starting with one centered at 09:12 p.m. Again HONO levels rose shortly after sunset, reaching a relatively stable level by $\sim\!10$ pm, and decreased following sunrise. The levels of NO₂ decreased progressively throughout the night but did not fall below 2 ppb. The HONO/NO₂ ratios were $\sim\!0.05$ early in the evening and show a gradual steady increase throughout the night to $\sim\!0.40$ at sunrise, coincident with the decrease in NO₂ levels, and again supporting the lack of a dependence of the observed HONO levels on the coincident NO₂ levels. A clear anticorrelation exists between NO₂ and the HONO/NO₂ ratio as various NO₂ plumes were encountered throughout the night, although the HONO levels remained largely unperturbed throughout these events.

All these cases highlight the existence of a HONO PSS that is well established shortly after sunset. After that, HONO levels remain relatively constant regardless of

ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.



the air mass source, and appear not to be correlated with NO₂ to a first approximation, giving rise to an anti-correlation between HONO/NO₂ levels and NO₂. The HONO/NO₂ ratios are surprisingly high, starting out with ratios of 0.05-0.13 early in the evening and rising to levels of >1.0 before sunrise. Some of these points will be explored further in the following sections.

3.2 Study wide observations of HONO

The HONO/NO₂ ratio is frequently used as an index of the efficiency of heterogeneous conversion of NO₂ to HONO, the rationale being that the production of HONO is first order with the concentration of NO2. A tremendous amount of scatter exists in the relationship between HONO and NO₂ (Fig. 5a), using all data from the study. This scatter has been seen in several other studies in which HONO is highly correlated to NO2 close to direct sources of NO_x (urban areas) but significantly less correlation exists between these two pollutants in rural air (Harrison et al., 1996). The data in Fig. 5a appears to be bounded on the lower end by a HONO/NO2 ratio of 0.0084, similar to that observed in a recent tunnel experiment dominated by diesel vehicles (HONO/NO_x=0.008±0.001) (Kurtenbach et al., 2001) and similar to that in the nocturnal boundary layer of Kathmandu, Nepal (Yu et al., 2009). Thus the lower bound in Fig. 5a may represent the HONO/NO_x ratio from diesel sources operating in the region, with NO titrated quickly to NO₂ by O₃ in the MBL. Also similar to the study in Nepal, the zero forced regression slope in Fig. 5 was HONO/NO₂=0.083±0.03 (not shown). What is not similar to that of other studies is that the data in Fig. 5 does not have an upper bound for the HONO/NO₂ ratio, as has been seen in many previous studies in terrestrial regions. For example, it was found that the nocturnal HONO/NO2 ratio rarely exceeded 0.04 in urban Birmingham (Harrison et al., 1996), 0.035 in Milan after correction (Stutz et al., 2002), 0.075 at two rural sites in England (Harrison et al., 1996), 0.09 in Los Angeles (Harris et al., 1982), 0.03-0.10 at various urban locations in the US, increasing with an increase in relative humidity (Stutz et al., 2004), 0.125 in South-East England (Kitto and Harrison, 1992), 0.13 at various locations in Europe (Lammel and Cape, 1996), and most **ACPD**

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page

Abstract Introduction

Conclusions References

Figures

Full Screen / Esc

Back

Printer-friendly Version



recently 0.30 in Nepal (Yu et al., 2009). In the present study, the upper HONO/NO $_2$ ratios were >1.0. During such cases, NO $_2$ is present at very low levels and yet HONO remains unperturbed and maintains a constant level. A case in point was presented in Fig. 3 between 03:30–05:30 a.m., as well as the observations discussed for 2 August, 01:30–07:30 a.m. Note that the interpretation of Fig. 5a is complicated since it includes both light and dark periods with variable lifetimes for HONO with respect to photolysis. For further analysis, we will define the PSS period in this study to be between 10:00 p.m. and 05:00 a.m. PDT, purposely excluding the build up and photolysis periods during sunset and sunrise. The overall statistics for observations in the PSS period are presented in Table 1.

When we plot data from the PSS, Fig. 5b, we now have an apparent lower limit of ~0.033, but still with no upper limit. It is interesting to note that in Milan Italy, Stutz et al. (2002) found that all data (day and night) fell below a HONO/NO₂ line of x=0.035when corrected for direct sources. The interpretation of the upper limit in their study is that it represents the overall conversion efficiency in the reaction, $NO_{2(q)} \rightarrow x HONO_{(q)}$ on the terrestrial surface below. Data falls below the line in their plot due to other losses of HONO such as photolysis. And yet, data we collect during the PSS period fall above such a line. Our best interpretation of Fig. 5b at this point is as follows. First, during the PSS period, we have no losses of HONO from the NBL due to photolysis. Second, we argue that the data falling above the line is an indication of a mixture of HONO formation mechanisms in the aqueous system that are both 0° order and 1° order in NO₂. The significance of the x=0.033 line is not completely understood, but it seems to indicate the HONO to NO₂ conversion on dry surfaces (Stutz et al., 2004). It may represent a fundamental 1° order limit for production of HONO on a surface coated with a single monolayer of water. We will describe the 0° order mechanism in a future section, but for now, we just comment that the 0° order mechanism would result from the heterogeneous formation of HONO on a surface that is saturated with NO2 equivalent precursors, followed by reversible distribution of HONO from the surface to the gas phase (vaporization/deposition), without loss from the MBL system. In Fig. 5a,

ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

© O

the limit for a zero order process is the vertical y-axis.

Study average nocturnal profiles of HONO, NO₂, HONO/NO₂ ratio, relative humidity and temperature presented in Fig. 6, were created by binning all study data into 0.5 h intervals. The average profile for HONO is similar to that discussed for the case stud-5 ies, showing a rapid increase after sunset to a PSS level, which is steady throughout the night, with a small discernable decrease with time until sunrise, when HONO photolysis begins. The increase in HONO to a PSS level is already started at the point of sunset, and is complete within an hour after sunset, much faster than is typically seen in terrestrial locations. In case studies in urban areas with similar values of RH. 60%<RH<100%, the PSS in HONO was not reached until 00:00 a.m. in Nashville, 02:30 a.m. in Houston and ~01:00 a.m. in Phoenix, in each case being several hours after sunset (Stutz et al., 2004). The main difference between the environment in the present study and that in the urban environment is the presence of the ocean surface, and to a lesser extent the presence of marine aerosols. One of these factors is responsible for the fast approach to the steady state. The nocturnal profile of NO₂ (Fig. 6) indicates a general decrease in NO₂ as the night proceeds. The temporal decrease in NO₂ could be due to a number of factors including a reduction in anthropogenic NO₂ sources, a loss of NO₂ to the NO₃/N₂O₅ reservoir with heterogeneous conversion to HNO₃ and photolabile halogen species (Dentener and Crutzen, 1993; McLaren et al., 2010), loss of NO2 due to heterogeneous formation of HONO, dissolution of NO2 in the ocean, and/or a change in transport patterns during the night. An increase in the HONO/NO₂ ratio through the night is also apparent in Fig. 6, with median levels climbing quickly after sunset to 13.0% at the beginning of the PSS period (average 22:00-24:00 PDT), but gradually increasing thereafter, reaching 25-30% by early morning. The 95-th percentiles of HONO/NO₂ ratios at the end of the night are >1.0 (Table 1). The implications of these numbers are that they suggest that the mechanism of HONO formation on terrestrial surfaces and on aqueous water are fundamentally related with a gradual increase in the maximum HONO/NO, ratio from 0.03 on surfaces in dry environments (10% RH) (Stutz et al., 2002, 2004) increasing to maximums of 0.10-0.13 on

ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

14

►I

•

Back



Full Screen / Esc

Printer-friendly Version



surfaces at high relative humidity (Lammel and Cape, 1996; Stutz et al., 2004) and with a continuity on aqueous surfaces observed in this study, with HONO/NO₂ ratios during the PSS observed ranging from 0.13 to >1.0. Thus it appears that not only the presence of water (1 monolayer of H₂O at low RH) but also the amount of water is important in the HONO formation mechanism. The similarity here is that all surfaces are coated with a minimum of 1 monolayer of water. Since H₂O is consumed in Reaction (R1), this could be a limiting reagent.

Several reports of HONO in terrestrial locations exist where authors comment on a PSS for HONO on certain nights (Lammel and Cape, 1996; Stutz et al., 2002, 2004; Yu et al., 2009), and it appears to be associated with humid nights (Stutz et al., 2004; Yu et al., 2009). The constant temporal behaviour of HONO in the PSS, implies a balance between sources and sinks of HONO at these times. Stutz presented a simple model to explain the PSS in more detail (Stutz et al., 2004), in which the source of HONO is the heterogeneous reaction of NO_2 with the surface (ground or aerosol surface) according to Reaction (R1), and the sink is the first order irreversible deposition of HONO to the surface. Assuming that NO_2 is constant with minimal losses, the result is a PSS for both HONO and the HONO/ NO_2 ratio:

$$[HONO] = \frac{\gamma_{NO_2 \to HONO}}{\gamma_{HONO}}[NO_2]$$
 (1)

where $\gamma_{\text{NO}_2 \rightarrow \text{HONO}}$ is the conversion probability (or uptake coefficient) for the reaction of NO₂ on the surface and γ_{HONO} is the uptake coefficient for HONO on the surface. The time constant for the approach to the PSS in the Stutz model (τ = $S/V \times v/4 \times \gamma_{\text{HONO}}$) is essentially the inverse of the first order loss constant for deposition of HONO to the surface, τ_{dep} , and the uptake coefficients were presented as being dependent on relative humidity. Equation (1) predicts that during periods where a PSS exists: i) the concentration of HONO is proportional to the concentration of NO₂, and ii) the [HONO]/[NO₂] ratio is constant. It also suggests that the level of HONO in the PSS should respond to changes in NO₂ with a time constant of τ_{dep} . Observations in our study suggest

ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

14

►I

Back

Close

Full Screen / Esc

Printer-friendly Version



Discussion Paper

Back

Printer-friendly Version

Interactive Discussion



that in this polluted aqueous environment, a PSS for HONO is established consistently and quickly almost every night, with a time constant of <1 h. Further though, the HONO/NO₂ ratio is not constant throughout the night, the steady state HONO concentration is not proportional to NO2, and the HONO PSS level does not respond to changes in NO_2 concentration with a time constant of τ_{dep} . The HONO PSS level we observe is relatively undisturbed with a change in air mass or a change in NO₂ levels by as much as an order of magnitude. These observations lead us to two inescapable conclusions thus far. The first conclusion is that during the PSS period, the source of HONO in the MBL is not proportional to the concentration of NO₂. Despite this, a source of HONO must exist. The second conclusion requires more explanation. The general consensus from field and laboratory investigations is that HONO is lost from the gas phase through deposition to surfaces, which in this environment are coated with water. As the lifetime for deposition of HONO from the PSS is <1 h, and since the HONO PSS can persist for much longer than this (i.e. -7 h), then the source of the HONO must either be: i) a reservoir of nitrogen that is significantly larger than the gas phase reservoir of HONO in the MBL and/or ii) HONO must be recycled into the source reservoir after deposition. Both these conclusions are significant since they modify our current understanding of nocturnal HONO formation. We will return to these points and discuss potential reservoirs of HONO and/or precursors in a future section.

There is still a chance that the phenomena observed here are specific to the MBL. We do not believe this to be true since a careful examination of previous datasets in terrestrial urban/suburban/rural environments can lead to similar conclusions. We provide a compilation of what we believe to be examples of PSSs of HONO in the nocturnal atmosphere in urban terrestrial environments, from existing literature (Table 2). The examples were compiled either through the authors identification of a nocturnal PSS or "levelling off" for an extended period of time, or through our examination of the data for periods of constant HONO during the night which are not perturbed by high local emission sources of NO_x (i.e. high NO) or high surface area of PM₁₀. As discussed by Stutz (2004), high direct sources can increase the amount of HONO beyond the PSS,

ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page **Abstract** Introduction Conclusions References

Figures

Full Screen / Esc

albeit temporarily, before HONO returns to the PSS. At this point we will highlight just one example of a PSS that has properties of our MBL PSSs. In Houston on 5 September 2000 (Stutz et al., 2004), after the establishment of a HONO PSS at 02:30 a.m., NO₂ levels increase from 02:30–05:30 a.m. by more than 60% (from 11.0 to 18.3 ppb, without increase in NO), although HONO remained almost constant (±6%), and the HONO/NO₂ ratio decreased by a factor proportional to the NO₂ increase. The independence of the HONO PSS level on NO₂ concentration is characteristic of what we have observed and central to our arguments above. This is but one example of a PSS in the urban environment during which HONO is independent of NO₂. Arguments of transport can always be invoked to rationalize an individual case, but clearly the case for transport as influencing the PSS (to a first approximation) has been ruled out in the polluted MBL in this study. We suggest that careful examination of PSSs in other urban data sets may result in similar conclusions.

3.3 Relative humidity and temperature dependence of HONO in PSS

To explore the dependence of the HONO mixing ratios in the PSS periods, all data collected during the PSS period (22:00-05:00 PDT) are plotted as a function of relative humidity (RH) in Fig. 7, and temperature in Fig. 8. A negative dependence of HONO with relative humidity is obvious (Fig. 7). A casual observation of the data might suggest lack of a dependence when relative humidity is less than ~75%, although two separate regressions for data in the RH<75% and RH≥75% regions indicate a statistically significant negative correlation over the whole RH range (59-90% RH). It is possible that HONO is coincidentally related to RH through a subtle decrease in HONO throughout the night, and a coincidental temporal change in other parameters as the night proceeds. To rule this out, HONO vs. RH regressions were performed on 1 h binned data from 22:00-05:00, thus removing any temporal relationship. All seven bins indicated a statistically significant negative correlation between HONO and RH. similar in magnitude to that seen in Fig. 7.

ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page

Abstract

Introduction

Conclusions References

Figures

▶I

Full Screen / Esc

Back

Printer-friendly Version



Full Screen / Esc Printer-friendly Version

Interactive Discussion



While a negative correlation between HONO and RH is observed, this is not proof of cause and effect. Relative humidity and temperature are not independent, with relative humidity increasing due to a decrease in temperature for a constant amount of water vapor. Thus the observed correlation between HONO and RH could indicate a positive dependence of HONO with temperature through some other physical or chemical mechanism. With current mechanisms in the literature, the link between HONO concentration and RH are thought to be related to increased water content on surfaces, which enhances the heterogeneous formation rate of HONO via Reaction (R1), and it could also enhance the loss rate of HONO from the gas phase on a surface, or into an aqueous layer formed on a surface. These surfaces in general could be the surface at the base of the air column (i.e. ocean), the surface of aerosols, or any anthropogenic/biogenic surfaces available in the air mass. Since the HONO concentration is a balance between sources and losses during the PSS period, for an increase in water on surfaces due to increase in RH, one would expect that an increase in the source rate of HONO would show a positive correlation of HONO with RH, while any increase in the loss of HONO would show a negative correlation of HONO with RH, all else being equal. Since the available surface is dominated by the ocean surface in the MBL, we expect that changes in RH will not significantly change the amount of water available on the surface of the ocean, but it could increase the amount of water available on marine aerosols. Thus, we conclude that the negative correlation of HONO with RH is either due to increased net loss of HONO on the surface of, or into the aqueous layer of MBL aerosols, or failing this, there exists a positive correlation between HONO and temperature that is somehow related to the HONO PSS mechanism. We next argue that the latter is true.

The loss of HONO on, or into the agueous surface of MBL aerosols arising from an increase in RH would be of significance only if the surface or volume density of H₂O on MBL aerosols is more significant than the surface or volume density of the ocean in the MBL. Despite a lack of supporting aerosol measurements during this study, these parameters can be reasonably estimated. For example, the total surface and volume

ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page **Abstract** Introduction Conclusions References **Figures** Back

Printer-friendly Version

Interactive Discussion



density of H_2O in a typical distribution of deliquesced sea salt aerosols (10 μ g m⁻³) at 96% RH is $4.3 \times 10^{-5} \,\mathrm{m^2 \, m^{-3}}$ ($43 \,\mu\mathrm{m^2 \, cm^{-3}}$) and $1.47 \times 10^{-10} \,\mathrm{m^3 \, m^{-3}}$ ($147 \,\mu\mathrm{m^3 \, cm^{-3}}$), respectively (Chameides and Stelson, 1992). Presuming that the MBL is polluted with a moderate amount of anthropogenic aerosols (10 μ g m⁻³; 50% H₂O, d_p =0.2 μ m), an additional surface and volume density of H_2O can be estimated to be 5.5×10^{-4} m² m⁻³ and $9.1 \times 10^{-12} \,\mathrm{m}^3 \,\mathrm{m}^{-3}$, respectively. The total surface and volume densities of $\mathrm{H}_2\mathrm{O}$ in the MBL aerosols would thus be 5.9×10^{-4} m² m⁻³ and 1.56×10^{-10} m³ m⁻³, respectively. We can compare these to the ocean. The geometric surface area density of the ocean surface in a well mixed MBL column of height H is given by 1/H; a lower estimate since the MBL may not be well mixed at night (Stutz et al., 2002). The volume density of H₂O in the ocean available to the MBL column of air above it can be estimated very conservatively to be equal to $t_{\rm SMI}$ /H (at a minimum), where $t_{\rm SMI}$ is the thickness of the aqueous layer in the sea surface microlayer (SML). Using a typical nocturnal marine boundary layer height, $h_{\text{MBI}} = 100 \, \text{m}$ (Brown et al., 2004) and a typical SML thickness, $t_{\rm SML}$ =50 µm (Zhang et al., 2006), our minimum estimate of the effective ocean surface and volume densities of H₂O in the MBL are 0.01 m² m⁻³ and 5×10⁻³ m³ m⁻³, respectively. These numbers are more than an order of magnitude (×17) and seven orders of magnitude (3×10⁷) greater than those available on MBL aerosols. Thus, we conclude that increased surface or volume density of aerosols coincident with an increase in RH is negligible compared to an unchanging ocean, having a negligible effect on HONO deposition loss. This implies that the negative correlation between HONO and RH is coincidental with a positive correlation between HONO and temperature, that is somehow related to the HONO PSS mechanism.

The positive relationship between HONO and temperature evident in Fig. 8 is statistically significant at the 95% confidence level. The correlation coefficient (R^2 =0.25) accounts for more of the variance in HONO observations than does the HONO vs. RH dependence (R^2 =0.17). Using a linear regression, the corresponding temperature coefficient $((1/P_{HONO}) \times dP_{HONO}/dT)$ of the dependence indicates that HONO increases

ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page **Abstract** Introduction

Conclusions References

Figures

Back

Full Screen / Esc

Back

Interactive Discussion

by $+7.2\pm0.6\%$ K⁻¹ in the temperature region. As we did for RH, HONO vs. T regressions were performed on 1 h binned data to rule out other temporal processes that may occur at night. A statistically significant (>95%) correlation between HONO and temperature was still observed in all seven bins (not shown), similar to that shown in ₅ Fig. 8.

There are many temperature dependent processes that could give rise to the temperature dependence of HONO over the ocean surface. Of interest is a comparison to the temperature dependence of a gas phase-aqueous phase equilibrium process such as that dictated by the temperature dependent Henry's Law coefficient, H(T) for the solubility of HONO in water:

$$H(T) = \frac{[\text{HONO}]_{(\text{aq})}}{P_{\text{HONO}}} = H^{\circ} \exp\left(\frac{\Delta H_{\text{vap}}}{R} \left(\frac{1}{T} - \frac{1}{T^{\circ}}\right)\right)$$
 (2)

where P_{HONO} is the partial pressure of HONO, H° is the Henry's Law constant at reference temperature T° , ΔH_{vap} is the enthalpy of vaporization of HONO_(aq), and R is the gas constant. For this equilibrium process, the temperature coefficient of HONO would be given by:

$$\frac{1}{P_{\text{HONO}}} \frac{dP_{\text{HONO}}}{dT} = \frac{\Delta H_{\text{vap}}}{R} \frac{1}{T^2}$$
 (3)

The value of H° (298 K) and $\Delta H_{\rm vap}/R$ in Eq. (2) are $49\pm1\,{\rm mole\,L^{-1}\,atm^{-1}}$ and 4900±150 K, respectively in water (Park and Lee, 1988; Becker et al., 1996). Using the ocean surface temperature in the Strait of Georgia in July (10-15°C), the temperature coefficient of P_{HONO} predicted by Eq. (3) would be: $+6.0\pm0.2\% \,\mathrm{K}^{-1}$. This is close to that observed in Fig. 8, despite that air in the MBL and the ocean are not in thermal equilibrium, and that the Henry's Law constant in sea water may deviate from that in water. Another way to express the temperature dependence is to determine the value of ΔH_{van} . This was achieved by a least squares regression of the relationship between In P_{HONO} vs. 1/T, the slope of which is equivalent to $-\Delta H_{van}/R$. The regression **ACPD**

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page

Abstract Introduction

Conclusions References

Figures

Printer-friendly Version

yields $\Delta H_{\text{vap}}(\text{HONO}) = 55.5 \pm 5.4 \text{kJ mole}^{-1}$, close to $\Delta H_{\text{vap}} = 40.7 \pm 1.2 \text{kJ mole}^{-1}$ determined from the literature value of H(T) in pure water. While this is not proof of the process, the temperature dependence of HONO PSS observations is at least consistent with an equilibrium process releasing HONO from the surface of the ocean.

3.4 Potential aqueous reservoirs of HONO

As discussed in Sect. 3.2, one of the conclusions from our observations was that there exists either a reservoir of HONO (or precursors) in the polluted MBL that is significantly larger than the gas phase reservoir or, HONO must be recycled into the source reservoir after deposition. We now address three possible aqueous reservoirs in the MBL: i) bulk water in the sea surface microlayer, ii) aerosols in the MBL, iii) the air-sea surface interface.

There is a suggestion that storage of HONO or nitrite on aqueous surfaces can be a source for daytime HONO in the early morning (Acker et al., 2004). This suggests the aqueous phase may be a significant reservoir. In the PSS model presented by Stutz (2004), the ultimate fate of HONO that deposits was not expressed explicitly. While there is evidence that HONO can deposit to laboratory surfaces releasing NO in a heterogeneous reaction (Lammel and Cape, 1996; Finlayson-Pitts et al., 2003; Stutz et al., 2004), it is also likely that HONO at the air-aqueous interface will dissolve if the surface is aqueous, containing more than just a few monolayers of water (Mertes and Wahner, 1995)

$$HONO_{(q)} \rightarrow HONO_{(aq)}$$
 (R2)

In the case of Reaction (R1) being the source of HONO, HNO₃ that is produced will also dissolve resulting in further acidification. The HONO_(aq) would subsequently ionize:

$$HONO_{(aq)} + H_2O(I) \rightarrow H_3O^+_{(aq)} + NO^-_{2(aq)}$$
 (R3)

dictated by the acid dissociation constant of $HONO_{(aq)}$, K_a , and pH. Total nitrite 25173

ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page **Abstract** Introduction Conclusions References **Figures ▶**I Back

Printer-friendly Version

Full Screen / Esc



 $([N(III)]=[HONO]_{(aq)} + [NO_2^-]_{(aq)})$ would build up in the aqueous reservoir to the point where the reservoir becomes a source of HONO to the gas phase.

$$HONO_{(aq)} \rightarrow HONO_{(q)}$$
 (R4)

At the point where the rate of Reacts. (2) and (4) are equal, we have dynamic equilibrium dictated by the Henry's Law constant (Eq. 2). If equilibrium is reached, the total concentration of N(III) in the water reservoir would be given by:

$$[N(III)] = P_{HONO}H\left(1 + \frac{K_{a, HONO}}{[H^+]}\right)$$
(4)

If the reservoir is the ocean, N(III) would be contained in the sea surface microlayer (SML), the top ~50 µm of the ocean that is dynamically isolated from the underlying ocean. The SML is frequently enriched in nutrients, biota, organic matter and anthropogenic pollutants. The SML can have different physiochemical properties than the underlying ocean and can serve as a source/sink for materials in the atmosphere (Hardy, 1982). Enhancement of pollutants in the SML with a sharp gradient in concentration and physiochemical properties at the base of the SML has been observed (Zhang et al., 1998, 2006). The underlying ocean also contains N(III), although it is dynamically separate from the SML through a diffusion barrier at the SML/subsurface layer interface.

The concentration of N(III) contained in the SML reservoir at equilibrium would depend on pH, temperature and SML thickness. At equilibrium, the molar fraction of N(III) stored in the SML reservoir would be given by:

stored in the SML reservoir would be given by:
$$f_{\text{SML}} = \frac{N_{\text{NIII,SML}}}{N_{\text{NIII,SML}} + N_{\text{HONO,MBL}}} = \frac{1}{1 + \frac{h_{\text{MBL}}}{RTHt_{\text{SML}} \times \left(1 + \frac{K_{\text{a}}}{[H^{+}]}\right)}}$$
(5)

where $N_{N(III), SML}$ is the moles of N(III) in the SML of thickness t_{SML} , $N_{HONO, MBL}$ is the moles of HONO in the MBL with effective boundary layer height $h_{\rm MBI}$. Using an unperturbed ocean pH=8.1 in the SML, $t_{SMI} = 50 \mu \text{m}$, $h_{MBI} = 100 \text{ m}$, T = 290 K, $K_a = 5.1 \times 10^{-4}$, 25174

Discussion Paper

Conclusions

Back

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



References

ACPD

10, 25153-25204, 2010

HONO formation on

aqueous surfaces

P. Wojtal et al.

Title Page

Abstract

Figures

Introduction



Printer-friendly Version

Interactive Discussion



 $H(T)=76\,\mathrm{M\,atm}^{-1}$, Eq. (5) predicts that 98% of the N(III) would be partitioned in the ocean SML, independent of the partial pressure of HONO in the gas phase. Thus, the SML, at equilibrium with the ocean, could be a significant reservoir of HONO. However, using Eq. (4) and the median nocturnal HONO mixing ratio of ~1 ppb, the concentration of N(III) in the SML at equilibrium would be 4.9×10^{-3} M. This is significantly higher than reports of concentrations of N(III) measured in the SML (<2 µM) (Zafiriou and True, 1979; Zhang et al., 2001, 2006), or the highest N(III) measured in the SML, 9 µM (Grasshoff, 1975). Only slight enhancements of N(III) and N(V) have been reported in the polluted Bay of Hong Kong (Zhang et al., 2001), and only a slight acidification of the SML compared to the underlying ocean has been recorded (-0.2 pH units) (Zhang et al., 2006). Despite this, it is currently unclear how low the pH of the SML could be in a polluted region such as the Strait of Georgia. If we change the pH to 7.0, we predict from Eq. (5), $f_{SMI} = 0.82$ and $[N(III)] = 4 \times 10^{-4}$ M. In this extreme acidification case, [N(III)] is still two to three orders of magnitude larger than what has been reported in the literature. Values of H_{HONO} in seawater are unavailable, and it is likely smaller than H in pure water used here due to the salt effect, but it is unlikely to be more than a factor of 2 smaller if it behaves similar to most other small molecules. It is unclear what effect the natural organic film (lipids and fatty acid material) on the surface of the SML will have on air/sea exchange and other physiochemical properties. Despite these uncertainties, evidence suggests that bulk seawater contained in the SML is significantly under saturated in N(III) with respect to the atmospheric HONO we measure during PSS periods, and as such, the bulk alkaline ocean would be a net sink for HONO in the gas phase in this region, not a source.

The second potential reservoir to address is aerosols. Sea salt aerosols are likely deliquesced in this region as the RH never fell below the efflorescence relative humidity of sea salt, 43-48% (Tang et al., 1997). We calculate the amount of HONO that could dissolve in marine aerosols and the fraction, f_{aerosol}, of HONO present in this reservoir compared to the total amount in the combined gas/aerosol reservoir. We presume that HONO in the gas phase comes to equilibrium with water in the aerosol phase.

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page **Abstract** Introduction Conclusions References **Figures** Back Full Screen / Esc

Discussion

Interactive Discussion

Using a generously alkaline aerosol pH=8.5 (Chameides and Stelson, 1992) for sea salt aerosols, the same values for H, T, and K_a as previous, and our previous estimate of the volume density of H₂O in the marine aerosols, 1.56×10⁻¹⁰ m³ m⁻³, we calculate $f_{\text{aerosol}} = 0.038$, and the gas phase equivalent of 40 pptV of N(III) that would exist in the aerosol phase. This is a minor reservoir in comparison to gas phase HONO, and would become significantly smaller as pH decreases. Observations of particulate nitrite and gas phase HONO in terrestrial regions also suggest that aerosol is a minor reservoir of N(III) compared to the gas phase (Lammel and Cape, 1996). We conclude that the bulk water present in aerosols could not support the PSS we are observing.

The third reservoir we address is the surface of the aqueous layer itself, which could be the surface of the ocean, lakes, deliquesced aerosols and wet terrestrial surfaces. In the MBL, the ocean surface dominates as discussed previously. Current mechanisms suggest that nocturnal HONO formation occurs at surfaces coated with at least a monolayer of H₂O. Lammel (1999) suggested that the formation begins with a molecule of NO₂ adsorbing to a molecule of water either chemisorbed to the surface (S-OH₂), or adsorbed to another surface molecule of water (S-OH₂*H₂O).

$$S-OH_2+NO_2 \rightarrow SS-OH_2\cdot NO_2 \tag{R5}$$

$$S-OH2·H2O+NO2 \rightarrow SS-OH2·H2ONO2$$
 (R6)

In a similar mechanism, Finlayson-Pitts proposed that the HONO formation begins with adsorption of a dimer of NO₂, N₂O₄, to the surface film, that is more highly stabilized by two water molecules than one. In a model similar to that presented previously (Mertes and Wahner, 1995), we propose that the first step of the mechanisms could also occur on the surface of bulk aqueous water, $(H_2O)_{\infty} \cdot H_2O$

$$(H2O)∞·H2O + NO2 \rightarrow (H2O)∞·H2O·NO2$$
(R7)

$$(H_2O)_{\infty} \cdot 2H_2O + N_2O_4 \rightarrow (H_2O)_{\infty} \cdot 2H_2O \cdot N_2O_4$$
 (R8)

In either case, the maximum density of NO₂ units that could be adsorbed on the surface is equal the maximum density of water molecules at the surface. For a geometrically

ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page **Abstract** Introduction

Conclusions References

Figures

I₫

Back

25176

flat aqueous surface, one can calculate the water molecule radius, $r_{\rm H_2O}$, cross sectional area, $A_{\rm H_2O}$, and surface density of water molecules, $S_{\rm H_2O}=1/A_{\rm H_2O}$, to be: $r_{\rm H_2O}=1.9\times10^{-10}\,\rm m$, $A_{\rm H_2O}=1.2\times10^{-19}\,\rm m^2$ and $S_{\rm H_2O}=8.6\times10^{18}\,\rm molecules\,m^{-2}$. The equivalent NO₂ surface density, $S_{\rm NO_2}^*$, if all the NO₂ in a well mixed MBL were brought to the ocean surface is given by:

$$S_{NO_2}^* \equiv \frac{P}{RT} C_{NO_2} h_{MBL} \tag{6}$$

where C_{NO_2} is the NO_2 mixing ratio, all else previously defined. The amount of nitrogen required to completely saturate the ocean water surface with a monolayer is obtained by equating $S_{NO_2} = S_{H_2O}$; we predict $C_{NO_2} = 3.4$ ppb. Thus the upper limit to the ocean surface reservoir under these conditions would be the equivalent of 3.4 ppb of nitrogen. If those precursors reacted by Reaction (R1) until they reach a PSS, ~1.0 ppb HONO would be produced in the gas phase. Ignoring excess NO₂ in the gas phase, the maximum fraction of HONO precursors existing on the surface in the MBL for this type of surface reservoir system, would be f_{surface} =77% (given our simple parameters). While this potential upper limit to the reservoir is reasonably large, if the HONO that subsequently deposits is lost from the system, the nitrogen in the surface/gas system would be depleted at a rate of [HONO]/ $\tau_{\rm dep}$. The surface-gas system could be replenished with gas phase NO₂ as the loss occurs. But if NO₂ is zero, the combined surface/gas phase nitrogen reservoir would be depleted with a time constant, $\tau_{\text{surface}} = \tau_{\text{dep}}/(1-f)$. Using $\tau_{\rm dep}$ =1 h, we would predict $\tau_{\rm surface}$ =4.4 h, as an upper limit. We almost had such a case on the morning of 2 August (discussed previously), where the PSS remained stable for 6 h, despite the mixing ratio of NO₂ falling from >15 ppb to <1.0 ppb. Although NO₂ was not zero, the absence of any observable decline in the HONO PSS level during this period still leads us to believe that if a surface system is generating HONO in the PSS, the HONO that deposits must be recycled into the surface reservoir.

We conclude this section by saying that the aqueous air-water interface at the surface of the ocean, perhaps confined to the top monolayer of water (0.4 nm thick), but

ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Printer-friendly Version



ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page **Abstract** Introduction Conclusions References **Figures** Back Full Screen / Esc Printer-friendly Version Interactive Discussion

Discussion Paper

thinner than the thickness of the SML, is the most likely reservoir for NO₂ (or its equivalent precursors) to produce HONO via Reaction (R1) in sufficient quantities to support the PSS of HONO that we observe in the MBL. As with any surface based mechanism, this sea surface nanolayer (SNL), would be chemically active through its ability to provide a medium that achieves extremely high surface densities where precursor species can react at high rates. Although we cannot prove through our observations that a monolayer of NO₂ or precursors would exist, the surface tension of water could support a monolayer of precursors (just as the surface tension of water can support the weight of a water spider), if there is a strong adhesive force of attraction between NO₂ (or equivalent) and surface water molecules, but less than the cohesive force of attraction between water bound to itself. Only certain molecules would fulfill this requirement. A more thorough discussion of this topic is beyond the scope of this work.

3.5 A conceptual model for HONO formation on aqueous surfaces

15

We summarize some of our conclusions on the PSS of HONO in the MBL thus far:

- 1) The PSS of HONO in the polluted MBL is characterized by an extended time period during which $d[HONO]/dt \approx 0$. It begins to form just before sunset and is usually fully formed within an hour after sunset or faster, faster than seen in terrestrial locations. The levels of HONO in the PSS are similar to that seen in several terrestrial locations; ~0.5-2.0 ppb; although there is a current lack of temperature and other data to support better analysis.
- 2) During the period of the PSS, HONO is frequently independent of NO₂, signifying a formation process that is 0° order in NO2, suggestive of a saturated surface reaction system. The median HONO/NO₂ ratios increase with time from 0.13 (lower limit 0.03) to 0.30 (upper limit >1.0) from the beginning to the end of the PSS period. The continuity of these numbers from those on terrestrial surfaces (maximum 0.03 at low RH to maximum 0.10-0.13 at high RH) suggest that the

- 3) P_{HONO} in the PSS is largely independent of air mass and transport direction, signifying that the PSS is characteristic of the MBL in the region, where the surface of the ocean is the dominating surface.
- 4) The levels of HONO during the PSS period show a statistically significant negative correlation with RH over the range RH=60–90%, that is only coincidentally related to the HONO PSS levels, through the negative *T*–RH relation for water in air.
- 5) The levels of HONO during the PSS period show a statistically significant positive correlation with temperature. The correlation indicates a temperature coefficient $(1/P_{\rm HONO})(dP_{\rm HONO}/dT)$ =+7.2±0.6% K⁻¹. If all of this temperature dependence is associated with a surface activation process of HONO, the enthalpy of the process that releases HONO from the SNL would be $\Delta H_{\rm SNL}$ =55.5±5.4kJ mole⁻¹, slightly larger than the $\Delta H_{\rm vap}$ =40.7±1.2kJ mole⁻¹ for HONO vaporization from pure water.

15

25

- 6) The bulk alkaline aqueous phase of the SML and sub layers of the ocean are very unlikely to be a reservoir of N(III) that could act as a net source of HONO to the MBL based upon current literature of [N(III)] and pH in the SML. The bulk aqueous phase of marine aerosols are also an unlikely reservoir of N(III) that could act as a net source of HONO to the MBL.
- 7) The top monolayer of water at the air-sea surface interface could represent a significant reservoir of nitrogen precursors for HONO formation, if the monolayer was saturated with NO₂ and/or related nitrogen precursors and products (N₂O₄, ONONO₂, NO⁺NO₃, HONO, HNO₃). This sea surface nanolayer (SNL) would be very acidic through the generation of nitric acid. A single report of NO₃ and NO₂ measurements in the surface microlayer (SML) and sublayers of polluted Daya

ACPD

Discussion Paper

Discussion Paper

Discussion Paper

Discussion Paper

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I∢ ≯I

→

Back Close
Full Screen / Esc

Printer-friendly Version



Bay near Hong Kong (Zhang and et al., 2001) indicate average enhancements of $2.83\,\mu\text{M}$ (NO $_3^-$) and $0.05\,\mu\text{M}$ (NO $_2^-$), perhaps a sign of faster transport of HNO $_3$ from the surface or higher HNO $_3$ /HONO density at the surface of the ocean.

Thus far, we have not discussed a detailed mechanism for HONO formation. Mertes and Wahner (1995) presented a model for NO₂ hydrolysis via Reaction (R1) on experimental aqueous surfaces that used the concept of a surface layer separate from the bulk liquid phase. In an extensive review of all studies to date, Lammel (1999) suggested a surface based formation mechanism and attempted to parameterize the formation of HONO on surfaces according to the following formula:

$$\frac{d[\mathsf{HONO}]}{dt} = a[\mathsf{NO}_2] \left\{ \left[\left(\frac{S}{V} \right)_{\mathsf{dry}} + \left(\frac{S}{V} \right)_{\mathsf{aqueous}} \right] k_{\mathsf{het1}} + \left(\frac{S}{V} \right)_{\mathsf{wet}} k_{\mathsf{het2}} [\mathsf{H}_2 \mathsf{O}] \right\}$$
 (7)

with different parameterizations for dry (dry laboratory and ambient surfaces), wet (humid surfaces), and aqueous (bulk water) surfaces, and a=0.5 from Reaction (R1). It predicts a HONO formation that is 0° order in H₂O for dry and aqueous surfaces (noting that dry surfaces contain a monolayer of H₂O) and 1° order in H₂O for humid surfaces, and a rate that is 1° order in NO₂ for all three surface types. Following from Reaction (R6), the main reaction step to produce HONO proposed by Lammel (1999) is:

$$S-OH2·H2ONO2 + NO2 \rightarrow SS-OH2·HONO2 + HNO2$$
(R9)

Finlayson-Pitts (2003) presented an integrated mechanism for HONO formation on surface films. The detailed mechanism suggested the formation of N_2O_4 in the gas phase, followed by uptake of N_2O_4 onto the surface film (dry or humid). The N_2O_4 isomerizes to asymmetric ONONO $_2$ through reaction with H_2O , which autoionizes to the $NO^+NO_3^-$ complex in the supporting layer of H_2O . The complex then reacts with water to generate surface adsorbed HNO $_3$ and surface HONO. The surface HONO escapes in part to the gas phase, and is presumably reversible. It was also suggested that in highly

ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I∢ ►I

< →

Full Screen / Esc

Back

Printer-friendly Version



acidic surfaces, HNO $_3$ can generate NO $_2^+$, which reacts with HONO to produce NO that escapes from the surface. The rate of formation of HONO in this mechanism was overall 1° order in NO $_2$, +2° due to formation of N $_2$ O $_4$ and -1° due to back reaction of NO $_2$ with ONONO $_2$ to produce N $_2$ O $_4$.

Based upon all the observations and conclusions above, and with considerable aid from previous literature, we can now provide a conceptual model for HONO formation in aqueous systems. The model can explain the PSS state we observe in the MBL that is also frequently seen in terrestrial regions on nights that are humid enough to produce condensed water. The model is presented in Fig. 9, illustrating conceptually the mechanism operating in 1° order and 0° order (in NO₂) generation of HONO. In the model, a sea surface nanolayer (SNL) (or surface nanolayer/surface film in urban areas) develops after exposure of the aqueous surface to NO₂ or its equivalent precursors. The SNL is 1 to n monolayers [$t_{SNI} = n \times 0.38$ nm], where the total thickness is significantly less than the thickness of the SML (~50 μm). The SNL could be a single monolayer of NO₂ (or equivalent) precursors on top of the water that is stabilized by a high adhesive force of attraction between the precursor molecule and water, which is less than the cohesive force of attraction between water molecules. The SNL becomes saturated with nitrogen after exposure to a sufficient amount of NO₂, and as time progresses during the night. If the SNL is completely saturated with N species (NO₂, N₂O₄, ONONO₂, NO⁺NO₃, HONO, HNO₃), the subsequent HONO formation chemistry is independent of the concentration of NO2, i.e. 0° order in NO2. If the surface layer is under saturated, then the system reverts to HONO formation with 1° order kinetics in the extreme, limited by the deposition of NO2 to the surface. The high surface density of the layer would provide the medium for close contact of nitrogen pairs and surface reactions necessary to generate all the surface species present in the mechanism presented by Finlayson-Pitts (2003). We previously calculated the maximum density of N that could be present on the ocean surface, suggesting that at night, exposure of the ocean surface to several ppb of NO2 would be necessary to completely saturate the surface to some equilibrium state. The density of N required to establish an equilibrium system

ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

l∢ ≯l

■ Back Close

Full Screen / Esc

Printer-friendly Version



in the SNL is not certain, although we note that that the SNL would be very acidic due to the production of HNO_3 . While we have no proof of the N density in the SNL, the upper limit is approximately equivalent to the surface density of a monolayer of water, 8.6×10^{18} molec m⁻². Finlayson-Pitts (2003) provided evidence for the existence of N_2O_4 in surface films of laboratory surfaces, and demonstrated that its uptake on the surface is enhanced or stabilized by the presence of HNO_3 . We presume that this important precursor, N_2O_4 , would be present in the SNL as well, possibly formed in the gas phase as suggested, but also potentially formed at high reaction rates in the high N density of the SNL monolayer. The existence of the SNL, saturated with precursors at night and partially saturated at sunset, could explain our fast approach to a PSS at sunset, compared to urban areas. In urban environments, a SNL usually does not exist at sunset, and would form after wetting of surfaces has happened, and after the water has adsorbed sufficient NO_2 .

In Fig. 9, we simplistically represent the detailed chemistry for HONO production by Reaction (R1), with mechanistic details that could be those offered by Finlayson-Pitts (2003) or Lammel (1999) or others. The final reaction by Finlayson Pitts (2003) would be:

$$NO^{+}NO_{3}^{-} + H_{2}O \rightarrow HONO_{SNL} + HNO_{3, SNL}$$
(R10)

with equivalent products to those in Reacts. (R9) and (R1). The HONO that is produced in the SNL would encounter a strongly acidic surface medium, supporting its reversible distribution to the atmosphere above:

$$HONO_{SNI} \rightleftharpoons HONO_{(q)} \qquad \Delta H = \Delta H_{SNI} (HONO)$$
 (R11)

The enthalpy of Reaction (R11) accounts for all, or a significant portion of the temperature dependence we have observed during the PSS period. In Fig. 9, we included the diffusion of products, HONO and HNO $_3$ into the underlying water as a slow process, although turbulence, wave breaking and bubble bursting in the ocean would all be mechanisms for transport of N(III) into sublayers and generation of NO $_2^-$ in a more

ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Printer-friendly Version

Interactive Discussion



alkaline medium (React. R3). We expect that HONO_{SNI} is undissociated, as could be HNO₃, depending on the surface density of nitrogen. The presence of sub layers of water allows diffusion of HNO₃, H₃O⁺ and NO₃, away from the SNL. This transport would prevent the SNL from becoming too acidic, and could be an important distinguishing feature between the aqueous system, and HONO formation on "dry" or "humid" surface films. In the former, NO and N₂O have been observed (Finlayson-Pitts et al., 2003), likely due to the presence of HNO₃ complexes in the concentrated film. The underlying layers of water in urban/suburban/rural aqueous systems would achieve the same goal. Thus the presence of the sublayers of water make Reaction (R11) truly reversible, since deposition of HONO to generate NO in a highly concentrated surface film could result in a loss of N and HONO from the PSS system. It is also worthy to note that there is a significant thermodynamic advantage to Reaction (R1) if the product is HNO_{3 (aq)} $(\Delta H_f = -207.4 \text{ kJ mol}^{-1})$ compared to HNO₃(I) $(\Delta H_f = -174.1 \text{ mol}^{-1})$, the latter existing in extremely concentrated surface films in the limit of zero water.

A distinguishing feature between the two mechanisms in Fig. 9, is that when the mechanism operates under 1° order NO2 kinetics, the reaction is pushed forward (left to right) by the irreversible loss of HONO from the SNL, likely dominated by photolysis of gaseous HONO. The surface layer is depleted at such times creating the conditions necessary for 1° order kinetics. The system will continue to operate under 1° order kinetics when photolysis stops at sunset for a short time, as HONO continues to be rapidly lost from the surface to establish a new steady state level of HONO, the PSS level. When the PSS level of HONO is reached, reversible deposition of HONO to the surface will dominate the loss of gaseous HONO, slowing down the depletion of nitrogen in the surface layer and created a saturated surface system again. Under these conditions, a quasi equilibrium system exists where uptake of NO₂ at the surface will slow considerably, and the PSS of HONO is independent of NO₂ concentration. The whole system would sit in static equilibrium, being driven forward only through the slow losses of HNO₃ and HONO from the SNL (i.e. diffusion from the SNL to the SML). The formation of HONO under these conditions is zero order in NO₂, represented by

ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page **Abstract** Introduction Conclusions References **Figures** Back Full Screen / Esc

the equilibrium established between HONO partitioning between the SNL and the gas

ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page **Abstract** Introduction Conclusions References **Figures ▶**I Back Full Screen / Esc Printer-friendly Version

Interactive Discussion



Discussion and implications for atmospheric chemistry

phase.

The aqueous model presented in Fig. 9 is representative for all aqueous systems, including the ocean, freshwater lakes, the surface of deliquesced aerosols and terrestrial surfaces coated with condensed water or precipitation. In the MBL, the agueous surface is perpetual, day and night. In terrestrial locations and with absence of precipitation, condensed water appears later in the night as the RH increases. Condensed water on grass surfaces for example can be seen at RH~80% (personal observation). The transition from a dry or humid HONO formation mechanism, to an aqueous PSS mechanism would be characterized by the time at which increasing HONO levels out to a period of $d[HONO]/dt \approx 0$, perhaps with overshoot of the PSS level before return to the PSS level (Stutz et al., 2004). The overshoot could be attributed to the presence of a variety of surfaces that form condensed water over a range of relative humidities (Lammel, 1999). If the aqueous system is truly operational with a fully saturated SNL, then the PSS level of HONO, created by a thermodynamic equilibrium, should in principle be independent of NO₂, relative humidity, and S/V ratio, and could be easily predicted with knowledge of the temperature. The zero order dependence in NO₂ during the PSS period makes previous literature difficult to interpret that have used HONO/NO₂ ratios at all times, implicitly implying HONO formation that is 1° order in NO₂. The evaporation of a terrestrial aqueous system after sunrise could also release significant amounts of nitrogen in many forms including HONO, NO, NO₂ and HNO₃, as observed previously for HONO (Acker et al., 2004), that may be difficult to interpret. Thus storage of nitrogen in the SNL of aqueous water could be considerable.

The aqueous system would only operate where a suitable SNL could be established, requiring exposure of surface water to a certain amount of NO₂ and/or HONO. This threshold level is difficult to predict without knowing the nitrogen density that would exist in the SNL, and will likely be related to the S/V ratio of the environment. We have

Conclusions

Back

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



made arguments that the amount of NO₂ to saturate a geometrically flat aqueous layer would have an upper limit of ~3.4 ppb for a 100 m boundary layer when the surface starts clean, as may happen after a precipitation event or rapid condensation on dry surfaces. An additional amount of NO₂ equivalent to the PSS level of HONO would be ₅ also be required, typically 1 ppb. The data in central Birmingham (1–5 February 1993) (Harrison et al., 1996) are very interesting in this regard as they appear to show evidence on a nightly basis of NO₂ depletions of 10-25 ppb, as HONO climbs, generating HONO PSS levels of 1–2 ppb. If an aqueous system were not operating, one would expect NO₂ depletions of only 2×[HONO]_{PSS}, based on Reaction (R1) stoichiometry. The NO₂ depletions are much greater than 2×[HONO]_{PSS}, and could be evidence of the depletion of NO₂ through establishment of an urban SNL where a high S/V ratio exists in the urban environment. Further analysis including transport, recent precipitation, nocturnal boundary layer height analysis would be needed to verify this.

In the remote marine boundary layer, we expect it probable that there is an insufficient amounts of NO2 for the establishment of a saturated SNL. We base this on the only other report of HONO measured in the MBL (Večeřa et al., 2008). In that report, as the ship travelled the Agean Sea, it experienced a range of HONO from below 50 ppt to 2.7 ppb as it encountering air masses with widely different NO₂ levels. The HONO formation in a clean MBL could revert to a formation mechanism that is 2° order in NO₂, as the binary reactions of NO₂ precursors on the surface of the monolayer could become the rate limiting step.

Another important implication of this work is the prediction that a SNL system could be operational both day and night in areas with high NO_v emissions in proximity to oceans and freshwater lakes. As photolysis of HONO during the day would dominate, the SNL system would eventually begin to form HONO under 1° kinetics, representing a significant pump for the production of OH radicals, not just in the early morning, but as long as the system continues to operate, likely limited by the rate of uptake of NO₂ to the surface The following reactions outline a simplified daytime mechanism to illustrate the process, where the production mechanism in the SNL is again represented simply as

10, 25153-25204, 2010

ACPD

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page **Abstract** Introduction

References

Figures

$$\begin{array}{ll} \text{NO}_{2\,(\text{g})} \rightarrow \text{NO}_{2\,(\text{SNL})} & \text{(R12)} \quad \times \, 2 \\ 2 \text{NO}_{2\,(\text{SNL})} + \text{H}_2 \text{O} \rightarrow \text{HONO}_{(\text{SNL})} + \text{HNO}_{3\,(\text{SNL})} & \text{(R13)} \\ \text{HNO}_{3\,(\text{SNL})} \rightarrow \text{HNO}_{3\,(\text{SML})} & \text{(R14)} \end{array}$$

$$HONO_{(SNL)} \rightarrow HONO_{(g)}$$
 (R15)

$$\begin{aligned} &\text{HONO}_{(g)} + h\nu \rightarrow \text{NO}_{(g)} + \text{OH}_{(g)} \\ &\text{NO}_{(g)} + \text{O}_3 \rightarrow \text{NO}_{2(g)} + \text{O}_{2(g)} \end{aligned} \tag{R17} \quad \text{day time}$$

The net daytime reaction is obtained from addition of these reactions (2(R12)+R13-R18), where a sufficient amount of O₃ to titrate the NO is assumed:

$$NO_{2(g)} + O_3 + H_2O + h\nu \rightarrow HNO_{3(SML)} + OH_{(g)} + O_{2(g)}$$
 (R13)

The net reaction predicts loss of odd oxygen (O₃+NO₂) in the boundary layer to generate HNO₃ in the SML and a quantitative amount of OH. It should be possible to detect this mechanism if it exists by measuring a negative flux of odd oxygen from the surface (positive gradient, $d[NO_2 + O_3]/dz > 0$) and a positive flux of OH from the surface (d[OH]/dz<0). This mechanism would break down if odd oxygen is significantly depleted in the MBL.

Conclusions

We report measurements of nocturnal HONO in a polluted marine boundary layer, the first such measurements of their kind, and have performed a detailed analysis of this dataset to gain a better understanding of the formation and destruction mechanism of HONO in the atmosphere. The environment was unique because the surfaces are predominantly aqueous. The temporal behaviour of HONO over aqueous systems is **ACPD**

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page

Abstract Introduction

Conclusions References

Figures

Back

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



Discussion Paper Discussion

Paper

(R17)

day time

25186

Full Screen / Esc Printer-friendly Version

Interactive Discussion



unique, with nightly formation of pseudo steady states of HONO shortly after sunset, a balance between formation and deposition. While nocturnal PSSs of HONO have been observed in terrestrial polluted areas later at night, they were thought to be a balance between a HONO formation process on surfaces that is 1° order in NO₂, and a 1° 5 order deposition loss of HONO to the surface, possibly irreversible. We have found that over the ocean, the levels of HONO during the PSS period are largely independent of NO₂, a clear signal of a 0° order process. This suggests that a reservoir of HONO and/or nitrogen precursors must exist in the MBL and/or a reversible loss of HONO to the surface exists. Our calculations support a conclusion that this reservoir is likely on the surface of the ocean, close to saturation to support a HONO formation process that is 0° order in NO₂. With minor modifications to the mechanisms of others, we present a conceptual model for HONO formation in aqueous systems. The elements of the model include a pool of N(IV) species concentrated on the surface of the ocean in a few monolayers, n, that react to produce HNO₃ (N(V)) and HONO (N(III)) in an acidic environment. We call this layer the sea surface nanolayer (SNL, $t_{SNI} = n \times 0.4$ nm), in keeping with convention in chemical oceanography where research on the sea surface microlayer (SML) proceeds. The HNO₃ is highly soluble and diffuses into the SML representing a slow loss of nitrogen from the SNL and a source to the SML. The HONO produced in the SNL escapes to the gas phase in a reversible process balanced by its deposition to the SNL. This reversible process represents a dynamic equilibrium dictated by Henrys Law. We observe a temperature dependence for this process that is consistent with an enthalpy for release of HONO from the surface with $\Delta H_{\rm SNL} = 55.5 \pm 5.4 \,\rm kJ \, mole^{-1}$, slightly larger than $\Delta H_{\rm vap} = 40.7 \pm 1.2 \,\rm kJ \, mole^{-1}$ for HONO partitioning from pure water. The extra 14.8±5.5 kJ mole⁻¹ could represent an extra stabilization of HONO in the high nitrogen environment of the SNL matrix, or it could indicate an additional enthalpy of a related process occurring in the saturated SNL system.

There is evidence in our data that during the PSS state, the HONO formation mechanism may operate between 0° and 1° states, again indicative of a surface mechanism

ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page Introduction **Abstract** Conclusions References **Figures** Back

Paper



that could become saturated. The evidence for this comes from an increase in the HONO/NO₂ ratio from 0.13 to greater than 1, as the night proceeds and with a lower limit seen at the start of the night of HONO/NO₂=0.03, similar to the maximum seen in other studies in dry environments. These numbers also suggest similarities between the HONO formation mechanisms on dry, humid and aqueous surfaces, with the maximum HONO/NO₂ ratios on dry surfaces of ~0.03, increasing, with RH on humid surfaces to 0.10-0.13, and then increasing further in the case of aqueous surfaces as the night proceeds. This suggests a common surface mechanism to all surfaces, where the top monolayer of water may play an important support role for precursors, and increasing water available to the surface plays a role either because it is a limiting reagent or perhaps due to a thermodynamic advantage of the product state where HNO₃ is dissolved.

Thus, the conceptual aqueous model we present should be representative of all aqueous surfaces including polluted coastal regions, freshwater lakes, deliquesced aerosol surfaces and other surfaces exposed to water. It is possible that the mechanism for HONO production can continue on aqueous surfaces under 1° kinetics with respect to NO₂ during the day, generating high amounts of the hydroxyl radical through HONO photolysis. We have demonstrated that the signal for such a mechanism operating during the day should be the loss of odd oxygen (O₃+NO₂) from the boundary layer, close to the surface with an apparent positive flux of OH from the ocean which would be 1/2 of the total negative flux of odd oxygen from the surface.

We further suggest, that the presence of a thin acid monolayer on top of the ocean on marine aerosols in polluted MBLs could provide the acid necessary to link the mechanism for generation of Cl2 in coastal environments at night, combined with the observations of Roberts (2008) that CINO₂ is a precursor to formation of Cl₂, but only on acidified sea salt surfaces with pH<2.0. The Cl₂ could be formed on the surface of the aerosols or on the ocean surface following deposition of CINO₂ to the surface.

In future, there could be immediate benefit from reanalysis of past HONO data sets where temperature and other supporting meteorological data are available in urban ar**ACPD**

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page **Abstract** Introduction Conclusions References

Figures

Back

Full Screen / Esc

Printer-friendly Version

Discussion Paper

Discussion Paper

Interactive Discussion



eas. Especially missing from the literature is the reporting of supporting temperature and water vapor data. The verification of the existence of PSSs in urban areas would be in order, with a view to understanding negative fluxes of NO₂ necessary to create the SNL, in terms of surface to volume ratios, the temporal extent of HONO production into daylight hours, and the consequences of an evaporation of the SNL at sunrise every day with release of stored nitrogen precursors. The temperature dependence of the HONO PSS should be verified in freshwater systems. On a molecular scale, more research is needed to determine the molecular density and the chemical nature of the SNL formed on aqueous systems. A better understanding of the aqueous mechanism would be obtained in future field studies if gradients and fluxes of HONO, NO2 and O₃ and other relevant species were measured both day and night in aqueous environments. The implications of the heterogeneous mechanism operating in polluted coastal and lakeside cities should be determined, with the help of modelers who would need to develop new algorithms to represent the aqueous mechanism in air quality models.

As a final note, we have not yet explored the implications of the acidic SNL on uptake of other gases on the ocean surface. It is likely that the deposition of anthropogenic NO2 and HNO3 contributes to ocean acidification on a local scale, but at the same time, it could have a negative feedback by slowing the uptake of other acidic gases, such as CO₂, to the bulk water below. This could be the case if the pH at the surface of the ocean deviates significantly from normal alkaline conditions, where the ocean is normally a net sink for atmospheric CO₂. The relationship between these opposing factors should be investigated, especially in light of the recently observed upwelling of acidified ocean currents along the continental shelf of Pacific North America, not far from where our study was conducted (Freely et al., 2008).

Acknowledgement. The authors acknowledge funding from the National Science and Engineering Research Council of Canada. We thank Joy McCourt and Daniel Majonis for assistance in experimental work, and lighthouse keepers Roy and Marie Barrows for their hospitality and generosity during our stay at Saturna Island.

ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page **Abstract** Introduction Conclusions References **Figures** Back Full Screen / Esc Printer-friendly Version

Discussion

Paper

Back

References

- Acker, K., Spindler, G., and Brüggemann, E.: Nitrous and nitric acid measurements during the INTERCOMP2000 campaign in Melpitz, Atmos. Environ., 38, 6497–6505, 2004.
- Acker, K., Wieprecht, W., Auel, R., Kalass, D., and Möller, D.: Evidence for heterogeneous formation of nitrous acid on cloud droplets, J. Aerosol Sci., 31(1), S352-S353, 2000.
- Alicke, B., Geyer, A., Hofzumahaus, A., Holland, F., Konrad, S., Pätz, H. W., Schäfer, J., Stutz, J., Volz-Thomas, A., and Platt, U.: OH formation by HONO photolysis during the BERLIOZ experiment, J. Geophys. Res., 108, 8247, doi:10.1029/2001JD000579, 2003.
- Alicke, B., Platt, U., and Stutz, J.: Impact of nitrous acid photolysis on the total hydroxyl radical budget during the limitation of oxidant production/Pianura Padana Produzione di Ozono study in Milan, J. Geophys. Res., 107(D22), 8196, doi:10.1029/2000JD000075, 2002.
- Aubin, D. G. and Abbatt, J. P. D.: Interaction of NO₂ with hydrocarbon soot: focus on HONO yield, surface modification, and mechanism, J. Phys. Chem. A, 111, 6263-6273, 2007.
- Becker, K. H., Kleffmann, J., Kurtenbach, R., and Wiesen, P.: Solubility of nitrous acid (HONO) in sulfuric acid solutions, J. Phys. Chem., 100, 14984–14990, 1996.
- Bongartz, A., Kames, J., Schurath, U., George, Ch., Mirabel, Ph., and Ponche, J. L.: Experimental determination of HONO mass accommodation coefficients using two different technigues, J. Atmos. Chem., 18, 149-169, 1994.
- Bongartz, A., Kames, J., Welter, F., and Schurath, U.: Near-UV absorption cross sections and trans/cis equilibrium of nitrous acid, J. Phys. Chem., 95, 1076–1082, 1991.
- Brown, S. S., Dibb, J. E., Stark, H., Aldener, M., Vozella, M., Whitlow, S., Williams, E. J., Lerner, B. M., Jakoubek, R., Middlebrook, A. M., DeGouw, J. A., Warneke, C., Goldan, P. D., Kuster, W. C., Angevine, W. M., Sueper, D. T., Quinn, P. K., Bates, T. S., Meagher, J., Fehsenfeld, F. C., and Ravishankara, A. R.: Nighttime removal of NO_v in the summer marine boundary layer, Geophys. Res. Lett., 32, L07108, doi:10.1029/2004GL019412, 2004.
- Chameides, W. L. and Stelson, A. W.: Aqueous-phase chemical processes in deliquescent seasalt aerosols, Ber. Bunsenges. Phys. Chem., 96, 461–470, 1992.
- Cheung, J. L., Li, Y. Q., Boniface, J., Shi, Q., Davidovits, P., Worsnop, D. R., Jayne, J. T., and Kolb, C. E.: Heterogeneous interactions of NO₂ with aqueous surfaces, J. Phys. Chem. A, 104, 2655-2662, 2000.
- Dentener, F. J. and Crutzen, P. J.: Reaction of N₂O₅ on tropospheric aerosols: impact on the global distributions of NO_x , O_3 , and OH, J. Geophys. Res., 98, 7149–7163, 1993.

ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page

Abstract Introduction

Conclusions References

Figures

I₫

Full Screen / Esc

Printer-friendly Version

- **ACPD**
- 10, 25153-25204, 2010
- HONO formation on aqueous surfaces
 - P. Wojtal et al.

Interactive Discussion

. 1

- Environment Canada, National Climate Data and Information Archive, available at: http://www.climate.weatheroffice.ec.gc.ca/Welcome_e.html, 2008.
- Finlayson-Pitts, B. J., Ezell, M. J., and Pitts Jr., J. N.: Formation of chemically active chlorine compounds by reactions of atmospheric NaCl particles with gaseous N₂O₅ and ClONO₂, Nature, 337, 241–244, 1989.
- Finlayson-Pitts, B. J., Wingen, L. M., Sumner, A. L., Syomin, D., and Ramazan, K. A.: The heterogeneous hydrolysis of NO₂ in laboratory systems and in outdoor and indoor atmospheres: an integrated mechanism, Phys. Chem. Chem. Phys., 5, 223–242, 2003.
- Freely, R. A., Sabine, C. L., Hernandez-Ayon, J. M., and Ianson Debby, H. B.: Evidence for upwelling of corrosive "acidified" water onto the continental shelf, Science, 320, 1490–1492, 2008.
- Grasshoff, K.: Hydrochemistry of landlocked basins and fjords, in: Chemical Oceanography, Vol. 2, edited by: Riley, J. P. and Skirrow, G., Academic Press, New York, 456–598, 1975.
- Hardy, J. T.: The sea surface microlayer: biology, chemistry and anthropogenic enrichment, Progress Oceanogr., 11, 307–328, 1982.
- Harris, G. W., Carter, W. P. L., Winer, A. M., Abbas, A. A., Pitts Jr., J. N. Platt, U., and Perner, D.: Observations of nitrous acid in the Los Angeles atmosphere and implications for predictions of ozone-precursor relationships, Environ. Sci. Technol., 16, 414–419, 1982.
- Harrison, R. M. and Kitto, A. M. N.: Evidence for a surface source of atmospheric nitrous acid, Atmos. Environ., 28, 1089–1094, 1994.
- Harrison, R. M., Peak, J. D., and Collins, G.: Tropospheric cycle of nitrous acid, J. Geophys. Res., 101, 14429–14439, 1996.
- Hermans, C., Vandaele, A. C., Carleer, M., Fally, S., Colin, R., Jenouvrier, A., Coquart, B., and Mérienne, M.-F.: Absorption cross-sections of atmospheric constituents: NO₂, O₂, and H₂O, Environ. Sci. Poll. Res., 6, 151–158, 1999.
- Jenkin, M. E., Cox, R. A., and Williams, D. J.: Laboratory studies of the kinetics of formation of nitrous acid from the thermal reaction of nitrogen dioxide and water vapour, Atmos. Environ., 22, 487–498, 1988.
- Junkermann, W. and Ibusuki, T.: FTIR spectroscopic measurements of surface bond products of nitrogen oxides on aerosol surfaces implications for heterogeneous HNO₂ production, Atmos. Environ., 26A, 3099–3103, 1992.
- Kitto, A. M. N. and Harrison, R. M.: Nitrous and nitric acid measurements at sites in South-East England, Atmos. Environ., 26A, 235–241, 1992.

10, 25153-25204, 2010

HONO formation on aqueous surfaces

- Title Page **Abstract** Introduction Conclusions References **Tables Figures** I₫ Back Full Screen / Esc Printer-friendly Version

- Interactive Discussion

- Kraus, S. G.: DOASIS: a framework design for DOAS, Thesis, Universität Mannheim, Germany, 2006.
- Kurtenbach, R., Becker, K. H., Gomes, J. A. G., Kleffmann, J., Lörzer, J. C., Spittler, M., Wiesen, P., Ackermann, R., Geyer, A., and Platt, U.: Investigations of emissions and heterogeneous formation of HONO in a road traffic tunnel, Atmos. Environ., 35, 3385-3394, 2001.
- Lammel, G.: Formation of nitrous acid: parameterisation and comparison with observations. Rep. 286, Max-Planck-Institut für Meteorologie, 1999.
- Lammel, G. and Cape, J. N.: Nitrous acid and nitrite in the atmosphere, Chem. Soc. Rev., 25, 361–369, 1996.
- McLaren, R., Wojtal, P., Majonis, D., McCourt, J., Halla, J. D., and Brook, J.: NO₃ radical measurements in a polluted marine environment: links to ozone formation, Atmos. Chem. Phys., 10, 4187-4206, doi:10.5194/acp-10-4187-2010, 2010.
- Mertes, S. and Wahner, A.: Uptake of nitrogen dioxide and nitrous acid on aqueous surfaces, J. Phys. Chem., 99, 14000-14006, 1995.
- Osthoff, H. D., Roberts, J. M., Ravishankara, A. R., Williams, E. J., Lerner, B. M., Sommariva, R., Bates, T. S., Coffman, D., Quinn, P. K., Dibb, J. E., Stark, H., Burkholder, J. B., Talukdar, R. K., Meagher, J., Fehsenfeld, F. C., and Brown, S. S.: High levels of nitryl chloride in the polluted subtropical marine boundary layer, Nature Geosci., 1, 324-328, doi:10.1038/ngeo177, 2008.
- Park, J.-Y. and Lee, Y.-N.: Solubility and decomposition kinetics of nitrous acid in aqueous solution, J. Phys. Chem., 92, 6294-6302, 1988.
- Platt, U., Perner, D., Harris, G. W., Winer, A. M., and Pitts Jr., J. N.: Observations of nitrous acid in an urban atmosphere by differential optical absorption, Nature, 285, 312-314, 1980.
- Ponche, J. L., George, Ch., and Mirabel, Ph.: Mass transfer at the air/water interface: mass accomodation coefficients of SO₂, HNO₃, NO₂ and NH₃, J. Atmos. Chem., 16, 1–21, 1993.
- Roberts, J. M., Osthoff, H. D., Brown, S. S., and Ravishankara, A. R.: N₂O₅ oxidizes chloride to Cl₂ in acidic atmospheric aerosol, Science, 321, 1059–1100, 2008.
- Stutz, J., Alicke, B., Ackermann, R., Geyer, A., Wang, S., White, A. B., Williams, E. J., Spicer, C. W., and Fast, J. D.: Relative humidity dependence of HONO chemistry in urban areas, J. Geophys. Res., 109, D03307, doi:10.1029/2003JD004135, 2004.
- Stutz, J., Alicke, B., and Neftel, A.: Nitrous acid formation in the urban atmosphere: gradient measurements of NO₂ and HONO over grass in Milan, Italy, J. Geophys. Res., 107, 8192,

- doi:10.1029/2001JD00390, 2002.
- Stutz, J. and Platt, U.: Improving long-path differential optical absorption spectroscopy with a quartz-fiber mode mixer, Appl. Opt., 36, 1105–1115, 1997.
- Tang, I. N., Tridico, A. C., and Fung, K. H.: Thermodynamic and optical properties of sea salt aerosols, J. Geophys. Res., 102, 23269–23275, 1997.
- Vandaele, A. C., Hermans, C., Simon, P. C., Carleer, M., Colin, R., Fally, S., Mérienne, M.-F., Jenouvrier, A., and Coquart, B.: Measurements of the NO₂ absorption cross-section from 42 000 cm⁻¹ to 10 000 cm⁻¹ (238–1000 nm) at 220 K and 294 K, J. Quant. Spectrosc. Ra., 59, 171–184, 1998.
- Vandaele, A. C., Simon, P. C., Guilmot, J. M., Carleer, M., and Colin, R.: SO₂ absorption cross section measurement in the UV using a Fourier transform spectrometer, J. Geophys. Res., 99(D12), 25599–25605, 1994.
 - Večeřa, Z., Mikuska, P., Smolik, J., Eleftheriadis, K., Bryant, C., Colbeck, I., and Lazaridis, M.: Shipboard measurements of nitrogen dioxide, nitrous acid, nitric acid and ozone in the Eastern Mediterranean Sea, Water Air Soil Pollut. Focus, 8, 117–125, 2008.
 - Wang, S., Ackermann, R., and Stutz, J.: Vertical profiles of O_3 and NO_x chemistry in the polluted nocturnal boundary layer in Phoenix, AZ: I. Field observations by long-path DOAS, Atmos. Chem. Phys., 6, 2671–2693, doi:10.5194/acp-6-2671-2006, 2006.
 - Yu, Y., Galle, B., Panday, A., Hodson, E., Prinn, R., and Wang, S.: Observations of high rates of NO₂-HONO conversion in the nocturnal atmospheric boundary layer in Kathmandu, Nepal, Atmos. Chem. Phys., 9, 6401–6415, doi:10.5194/acp-9-6401-2009, 2009.
 - Zafiriou, O. C. and True, M. B.: Nitrite photolysis in seawater by sunlight, Marine Chem., 8, 9–32, 1979.
 - Zhang, Z. B., Pan, M. X., Wang, Z. D., Zhao, W., Huang, H. H., Wei, G. F., Gao, H. L., Peng, Y. H., Zhu, Z. H., and Li, L.: Biological and chemical studies of sea surface microlayer (SML) in Daya Bay, China, Chin. J. Oceanol. Limnol., 19, 272–281, 2001.
 - Zhang, Z. B., Liu, C., and Liu, L.: Physicochemical studies of the sea-surface microlayer, Front. Chem. China, 1, 1–14, doi:10.1007/s11458-005-0003-8, 2006.
 - Zhang, Z., Liu, L., Wu, Z., Li, J., and Ding, H.: Physicochemical studies of the sea surface microlayer: I. Thickness of the sea surface microlayer and its experimental determination, J. Coll. Interf. Sci., 204, 294–299, 1998.

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I ◆ ▶I

◆ ▶ Back Close

Full Screen / Esc



Table 1. Statistical summary of observations during the pseudo steady state period (10:00 p.m.-05:00 a.m.), 24 July-9 August 2005 (432 observations).

	95-th percentile	75-th percentile	Average	Median	25-th percentile	5-th percentile
HONO (ppb)	1.54	1.26	1.08	1.08	0.93	0.52
NO ₂ (ppb)	19.1	10.6	7.8	6.7	3.7	0.90
SO ₂ (ppb)	4.1	1.92	1.52	1.14	0.64	0.19
HONO/NO ₂	1.17	0.30	NA	0.155	0.093	0.048
<i>T</i> (K)	294.6	292.3	291.0	290.8	289.6	288.4
RH (%)	86	83	77.4	79	74	62
WS $(km h^{-1})$	24	19	12.7	13	6	4

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page						
Abstract	Introduction					
Conclusions	References					
Tables	Figures					
I₫	►I					
4	•					
Back	Close					
Full Screen / Esc						
Printer-friendly Version						

Table 2. Evidence of nocturnal HONO pseudo steady states in terrestrial locations.

Reference	Location	Date	Time (local)	HONO (ppb)	NO ₂ (ppb)	₹ (°C)	RH (%)	Comment
Stutz (2004)	Houston, USA	5 Sep 2000	2.5–5.0	0.9–1.1	11–24	_	88–93	
Stutz (2004)	Nashville, USA	13 Jul 1999	0.0–5.0	0.6–0.9	16–22	-	90-100	
Stutz (2004)	Phoenix, USA	26 Jun 2001	0.0–5.7	1.2–1.7	15–33	-	65-75	10–139 m
Acker (2004)	Melpitz, Germany	10 Apr 2000– 12 Apr 2000	1.0–6.0	0.1–0.2	1.1–2.2	-1-1	70–90	3 nights, PSS uncertain
Kleffman (2007)	Wuppertal, Germany	3 Sep 2004	22.5–5.5	1.0–1.2	-	-	-	PSS uncertain
Yu (2009)	Kathmandu, Nepal	3 Feb 2003	0.0–3.0	0.8–1.4	3–15	4–11	70–100	PSS uncertain, NO _v and PM ₁₀
Yu (2009)	Kathmandu, Nepal	4 Feb 2003	1.0–4.0	0.8 –1.2	5–9	3–10	80–100	sources, T and RH are range
Yu (2009)	Kathmandu, Nepal	5 Feb 2003	1.0–4.0	0.9–1.7	8–15	3–10	80–100	for whole study
Yu (2009)	Kathmandu, Nepal	6 Feb 2003	1.0–4.0	1.1–1.7	7–15	3–10	80–100	
This work*	Saturna Is, Canada	24 Jul 2005– 9 Aug 2005	22–5.0	0.5–1.5	4–19	15–21	74–86	* marine PSS

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I ← ►I

Back Close

Full Screen / Esc

Printer-friendly Version





Fig. 1. Study location.

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

■ Back Close
Full Screen / Esc

▶I

14

Printer-friendly Version



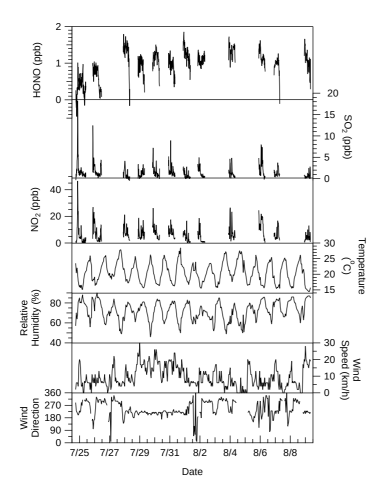


Fig. 2. Chemical and meteorological observations at East Point, Saturna Island.

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

Back Close

Full Screen / Esc

Printer-friendly Version





10, 25153-25204, 2010

HONO formation on aqueous surfaces

ACPD





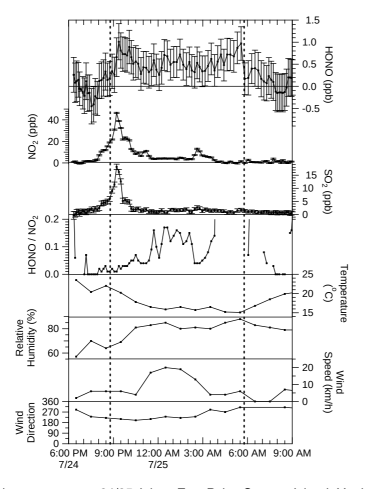


Fig. 3. Nocturnal measurements 24/25 July at East Point, Saturna Island. Vertical dashed lines represent times of sunset and sunrise.



10, 25153-25204, 2010

HONO formation on aqueous surfaces

ACPD





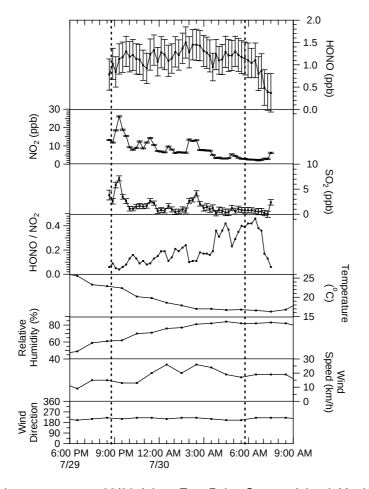


Fig. 4. Nocturnal measurements 29/30 July at East Point, Saturna Island. Vertical dashed lines represent times of sunset and sunrise.



10, 25153–25204, 2010

HONO formation on aqueous surfaces

ACPD





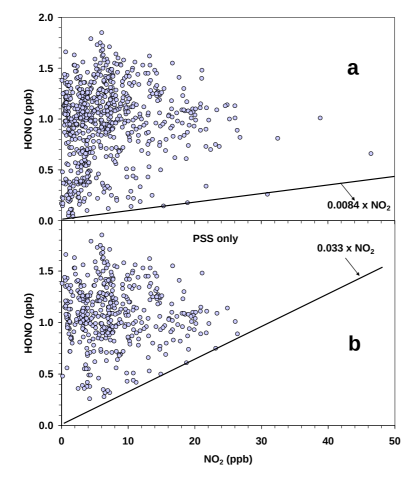


Fig. 5. Relationship between HONO and NO_2 using **(a)** all data. The lower limit line indicates HONO/ NO_2 =0.0084, fuel combustion sources **(b)** only data during the PSS period (10:00 p.m.–05:00 a.m.): the lower limit line indicates HONO/ NO_2 =0.033. See text for details.

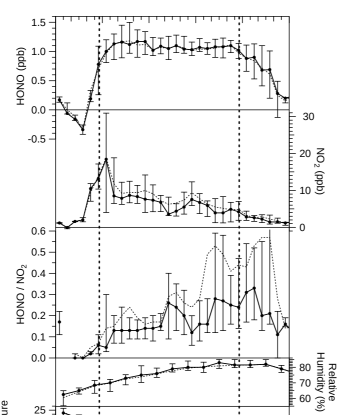


Fig. 6. Study average nocturnal profiles, 24 July-9 August 2005. Shown are median (.), 25th and 75-th percentiles (error bars), and average value (dotted line) within each 30 min bin. Vertical dashed lines represent times of sunset and sunrise.

6:00 PM 9:00 PM 12:00 AM 3:00 AM 6:00 AM 9:00 AM

Temperature

ွ

25

20

ACPD

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Abstract Introduction

Title Page

Conclusions References

Tables Figures

Close Back

Full Screen / Esc

Printer-friendly Version



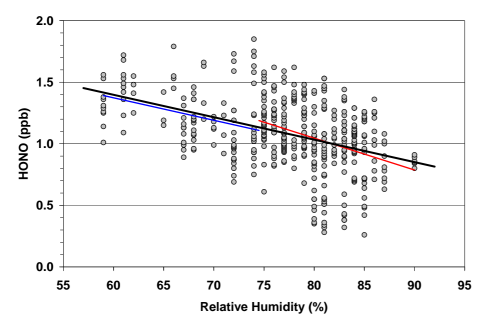


Fig. 7. Relative humidity dependence of HONO measured at Saturna Island during the PSS time period: 22:00–05:00. Black line – linear fit using all data (C_{HONO} =2.47(±0.35) ppb –0.0182(±0.0049)*RH%; R^2 =0.17). Blue and red lines show linear fit for RH<75% and RH \geq 75%, respectively. All negative dependencies are statistically significant at the 95% confidence level.

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page

Abstract Introduction

Conclusions References
Tables Figures

i∢ ⊳i

Back Close
Full Screen / Esc

Printer-friendly Version



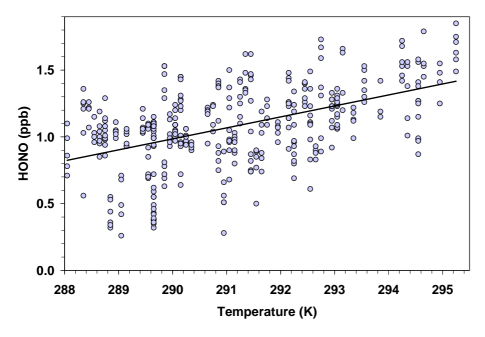


Fig. 8. Temperature dependence of HONO measured at Saturna Island during the PSS time period: 22:00–05:00. The equation of the linear line is $C_{\rm HONO}=0.082(\pm0.007)\times T-22.8(\pm2.1)\,{\rm ppb};~R^2=0.25.$ The positive temperature dependence is statistically significant (95% confidence level). The corresponding temperature coefficient of the dependence in the middle of the temperature range (292 K) is $(1/C_{\rm HONO})dC_{\rm HONO}/dT=+7.2\%(\pm0.6\%)\,{\rm K}^{-1}$.

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

I∢





Back



Full Screen / Esc

Printer-friendly Version



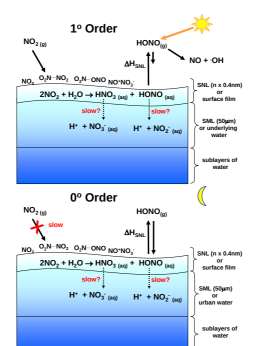


Fig. 9. Conceptual model of HONO formation mechanism on aqueous surfaces. The sea surface nanolayer (SNL) or surface film is from 1 to n monolayers thick, and is highly acidic. Underlying layers include the sea surface microlayer (SML) in the case of the ocean or underlying water (dew, surface water, etc.) in the case of terrestrial surfaces, which would allow a medium for acid transport away from the SNL. **(a)** The formation of HONO is 1° order in NO_2 , when irreversible loss of HONO (i.e. photolysis) from the SNL is faster than deposition of NO_2 to the surface. **(b)** The formation of HONO is 0° order in NO_2 , (i.e. a static system), when the SNL is saturated in NO_2 precursors and irreversible losses of HONO from the SNL are negligible. $HONO_{(g)}$ is dependent on the equilibrium dictated by Henry's Law for distribution of HONO between the SNL and gas phase.

10, 25153-25204, 2010

HONO formation on aqueous surfaces

P. Wojtal et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

I**4**

►I

■
Back

Close

Full Screen / Esc

Printer-friendly Version

