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The influence of gas-particle partitioning and surface-atmosphere exchange on ammonia during BAQS-Met

R. A. Ellis¹, J. G. Murphy¹, M. Z. Markovic¹, T. C. VandenBoer¹, P. A. Makar², J. Brook², and C. Mihele²

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Correspondence to: J. G. Murphy (jmurphy@chem.utoronto.ca)

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¹Department of Chemistry, University of Toronto, 80 St. George St., Toronto, ON, M6P 2S1, Canada

²Environment Canada, Science and Technology Branch, Air Quality Research Division, 4905 Dufferin St., Toronto, ON, M3H5T4, Canada

The Border Air Quality and Meteorology study (BAQS-Met) was an intensive field campaign conducted in Southwestern Ontario during the summer of 2007. The focus of BAQS-Met was determining the causes of the formation of ozone and fine particulate matter (PM_{2.5}), and of the regional significance of trans-boundary transport and lake breeze circulations on that formation. Fast (1 Hz) measurements of ammonia were acquired using a Quantum Cascade Laser Tunable Infrared Differential Absorption Spectrometer (QC-TILDAS) at the Harrow supersite. Measurements of PM_{2.5} ammonium, sulfate and nitrate were made using an Ambient Ion Monitor Ion Chromatograph (AIM-IC) with hourly time resolution. The median mixing ratio of ammonia was 2.5 ppb, with occasional high spikes at night resulting from local emissions. Measurements were used to assess major local emissions of NH₃, diurnal profiles and gas-particle partitioning. The measurements were compared with results from A Unified Regional Airguality Modelling System (AURAMS). While the fraction of total ammonia (NH_v≡NH₂ + NH₄) observed in the gas phase peaks between 0.1 and 0.8, AURAMS tended to predict fractions of either less than 0.05 or greater than 0.8. The model frequently predicted acidic aerosol, in contrast withobservations wherein NH, always exceeded the observed equivalents of sulfate. One explanation for our observations is that the net flux of ammonia from the land surface to the atmosphere increases when aerosol sulfate is present, effectively buffering the mixing ratio of gas phase ammonia, a process not included in the model. We explore the impact of a bi-directional flux parameterization on the predicted gas-particle partitioning of atmospheric ammonia.

1 Introduction

Atmospheric ammonia (NH₃) plays an important role in atmospheric processes and biogeochemical cycles. Ammonia readily neutralizes acidic compounds, reacting preferentially with sulphuric acid (H₂SO₄) to form non-volatile aerosol species ammonium

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bisulfate (NH_4HSO_4), letovicite [(NH_4)₃H(SO_4)₂] or fully neutralized ammonium sulfate [(NH_4)₂SO₄] (Finlayson-Pitts and Pitts, 1999). Any additional NH_3 is then available to take part in gas-particle partitioningwith other atmospheric acids to form semi-volatile salts. In the case of ammonium nitrate (NH_4NO_3), this equilibrium is strongly dependent on temperature and relative humidity, where colder, more humid conditions will shift the equilibrium to the particle phase (Stelson and Seinfeld, 1982). Ammonia is thus an important precursor to fine particulate matter ($PM_{2.5}$), the latter having repercussions with respect to human health (Pope et al., 2002) and to radiative budgets (IPCC, 2007). Fine particulate ammoniumalso has a longer atmospheric lifetime than gaseous NH_3 and can therefore be transported and deposited further downwind (NA) (NA) and NH_4 to the Earth's surface can be beneficial through the fertilization of nitrogen-limited ecosystems, or have detrimental effects such as eutrophication, soil acidification, and biodiversity loss in sensitive ecosystems ... (Fangmeier et al., 1994; Bobbink et al., 1998; Galloway et al., 2003; Krupa, 2003).

Unlike other anthropogenic pollutants, for example sulphur dioxide (SO_2) and nitrogen oxides (NO_x), the dominant emissions for ammonia are area wide-sources (Erisman et al., 2008; Makar et al., 2009; Environment Canada, 2010; U.S. E.P.A., 2009), and are subject to large uncertainty. In Canada, the majority of these emissions are agricultural in origin, such as from the application of fertilizer and volatilization of animal wastes, with lesser impacts from light-duty gasoline vehicles and chemical industries (Environment Canada, 2010). Total Canadian anthropogenic sources, including agriculture, are on the scale of 500 000 tonnes per year whereas natural sources such as forest fires are estimated at 5000 tonnes per year (NPRI, 2010). Because of its contribution to fine aerosol mass loading, modeling efforts have investigated the relative effectiveness of ammonia emission reductions in curtailing $PM_{2.5}$ formation...(Pinder et al., 2007; Tsimpidi et al., 2007; Pinder et al., 2008; Makar et al., 2009). Controls were found to be most effective in the winter when lower ammonia resulted in significant decreases in the mass loadings of ammonium nitrate relative to the base case.

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Emission reductions may be more challenging to implement for NH₃ than for SO₂ and NO_xdue to the nature and distribution of its sources. Additionally, area-wide emissions depend on both past and present land use. There is substantial evidence that over certain land types both emission and deposition of ammonia can occur depending 5 on the ambient mixing ratios (e.g. Farguhar et al., 1980; Sutton et al., 1993; Sutton et al., 1995; Asman et al., 1998; Nemitz et al., 2001). This bi-directional flux implies a compensation point: an atmospheric mixing ratio below which emission is observed and above which deposition is observed. Studies have shown that the compensation point is dependent on surface temperature and on the ratio $\Gamma = [NH_4^+]/[H^+]$ in soil and vegetation. This ratio is dependent on the accumulation of nitrogen from past fertilization (Massad et al., 2010). The deposition rate is also known to be dependent on the surface type and on the co-deposition of acidic species such as SO₂, HNO₃ and HCI (Erisman and Wyers, 1993; Flechard et al., 1999). Recent attempts to parameterize the bi-directional flux of ammonia for inclusion in modelshave suggested improved model-measurement agreement ... (Wu et al., 2009; Kruit et al., 2010; Massad et al., 2010; Zhang et al., 2010). In addition to emission and deposition to the surface, condensation and volatilization from particles plays an important role in determining the gas phase mixing ratio of ammonia. Wolff et al. (2010), showed that the timescale of surface fluxes is similar to the timescale of ammonium nitrate gas-particle partitioning.

Because of the high levels of uncertainty in the processes that govern the abundance of ammonia in the atmosphere, high time resolution measurements of ammonia are needed. Whitehead (2007), Nowak (2006) and Li (2006) studied the NH₃mixing ratios in urban environments and suggested ammonia emissions from traffic as a major source. Other studies have investigated the surface exchange of ammonia over crops and grassland (Walker et al., 2006; Kruit et al., 2007; Sutton et al., 2009; Bash et al., 2010). Relatively few studies have made simultaneous high time resolution measurements of ammonia and particulate ammonium. Nowak et al. (2006) compared measurements of ammonia made by a chemical ionization mass spectrometer to predictions from CMAQ, and found significant underpredictions from the model in

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the afternoon. Using the thermodynamic model ISORROPIA (Nenes et al., 1998) constrained by measurements of the chemical composition of fine particulate matter made by a particle into liquid sampler, it was determined that the thermodynamic partitioning was well-represented. The model underestimate of ammonia at midday led the authors to hypothesize that an additional light- or temperature-dependent source of ammonia was missing from the model.

The Border Air Quality and Meteorology Study (BAQS-Met), which took place in southwestern Ontario in the summer of 2007, offered the opportunity to make observations in a region impacted by regional transport of aerosol and local emissions of ammonia. A suite of instrumentation, including ammonia and aerosol measurements, was deployed at the Harrow supersiteto elucidate the primary andsecondary sources and composition of PM_{2.5} in this region. The field study was a collaborative effort between universities and government scientists with research activities that included measurements from several platforms, and modeling support from Environment Canada's regional air quality model AURAMS (A Unified Regional Air-quality Modelling System). In this paper, we investigate the measurements and modeling associated with gas phase ammonia and particulate ammonium in terms of gas-particle partitioning and the surface-atmosphere exchange of ammonia.

2 Experimental methods

2.1 Site description

Measurements were made at the Harrow ground site (42° 2′ 0″ N, 82° 55′ 0″ W, elevation 173 m), one of three BAQS-Met supersites. The site is situated in an agricultural area in southwesternOntario, Canada, approximately 5 km north of the shoreline of Lake Erie and 30 km south of Windsor, ON. The surrounding area is used to grow soybeans, corn, grapes, apples and tomatoes. The area around Harrow is impacted by long range transport of $PM_{2.5}$ and its precursors from the United States, especially

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areas inOhio, Indiana and Illinois to the south and from Michigan to the northwest. The siteis also impacted by urban and suburban emissions of pollutants from Windsorand Detroitand from localagricultural emissions. Harrow was often subject to lake breezes delivering onshore flow from Lake Erie in the middle of the day, hence our site provided information on the gas and particle chemistry taking place over the lake at night.

2.2 QC-TILDAS

Gas-phase measurements of ammonia were made using a Quantum Cascade Tunable Infrared Laser Differential Absorption Spectrometer (QC-TILDAS), developed by Aerodyne Research Inc. (Billerica, Massachusetts) and described in detail in Ellis et al. (2010). The QC-TILDAS monitors the NH₃ absorption spectrum at 967 cm⁻¹ using a thermoelectrically cooled, pulsed quantum cascade (QC) laser (Alpes Lasers, Neuchatel, Switzerland) which provides higher stability and increased power output over traditional diode lasers (McManus et al., 2002). The laser beam is directed into an astigmatic Herriot type multiple pass absorption cell (0.5 L, 76 m effective pass length) coated with a hydrophobic coating in an effort to reduce the interaction of NH3 with cell walls. A reference cell containing ethylene (C₂H₄), a less surface reactive gas that contains an absorption line near that of NH₃ is used for absorption-line lock. The signal and reference paths are separated temporally by 250 ns and focused on the same thermoelectrically cooled Mercury Cadmium Telluride (HgCdTe) infrared detector (Vigo Systems, Poland).

The laser control, spectral retrieval and mixing ratio calculations are managed by the software package TDLWintel, developed by Aerodyne Research Inc. The laser is scanned across the full NH₃ transition and the resulting spectrum is fit by convolving the laser line shape with a calculated absorption line shape based on the HITRAN (high-resolution transmission) molecular absorption database and the measured pressure, temperature and path length of the optical cell (Herndon et al., 2007). Although TDLWintel calculates absolute mixing ratios, calibration is highly recommended when measuring surface reactive gases such as ammonia. The software allows for automatic

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user defined additions of calibration gas and background gas through the use of two solenoid valves. Further description of the QC-TILDAS design and data acquisition system can be found in Nelson et al. (2004).

The QC-TILDAS uses a 10 cm, heated (to 40+2°C) quartz inlet which is internally coated with a hydrophobic fluorinated silane coating to limit condensation of water and interaction of NH₃ with inlet surfaces. A critical orifice is used to achieve a flow of ~9 L min⁻¹ (measured upstream at 101.3 kPa and 298 K) and maintain the pressure in the sampling lines and optical cell between 5.3–8 kPa in order to reduce absorption line broadening (Whitehead et al., 2008). The flow is split after the critical orifice, with 90% of the flow making a sharp turn and being pulled through the optical cell. The other 10% is pulled directly to the pump, relying on inertia to remove particles and eliminating the use of a particle filter, which may cause interferences from ammonium (Gras, 1984; Cheng and Tsai, 1997; Chow et al., 1998). The inlet also includes two ports for the introduction of calibration gas and background gas, designed so that these flows encounter the inlet in the same way as ambient sampling. Laboratory tests demonstrate the ability to restrict the majority (>90%) of the bi-exponential NH₃ time response to less than 0.4 s when using the quartz inlet in conjunction with a heated line. The ammonia time response is thus sufficient for data acquisition at 1 Hz, with a detection limit of 0.6 ppb.

2.3 AIM-IC

The inorganic composition of $PM_{2.5}$ was measured by Ambient Ion Monitor Ion Chromatography (AIM-IC). The AIM-IC is capable of continuous, hourly online measurements of the inorganic constituents of $PM_{2.5}$ (SO_4^{2-} , NO_3^- , NH_4^+ , CI^- , NO_2^- , Na^+ , K^+ , Ca^{2+} , and Mg^{2+}) and some precursor gases (HNO $_3$, NH_3 , HCI, SO_2 and HONO) with limits of detection as low as 0.1 μ g m $^{-3}$. The instrument consists of an AIM 9000D air sampler (URG Corp., Chapel Hill, NC) outfitted with a 2.5 μ m cyclone for particle size-selection, a constantly-generated wet parallel-plate denuder for the collection of gases, a particle supersaturation chamber and two ICS-2000 ion chromatographs (Dionex

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Corp., Sunnyvale, CA). The anion IC was equipped with a potassium hydroxide eluent generator cartridge (KOH EGC II), TAC-ULP1 (4mm) concentrator column, IonPac AG and AS19 guard and analytical columns, ASRS ULTRA II(4-mm) suppressor and CD25A conductivity detector operated at 30 °C. Anion chromatographic runs were performed isocratically with 20mM KOH at a flow of 1ml/min. The cation IC was equipped with a methanesulphonic acid eluent generator cartridge (MSA EGC II), TCC-ULP1 (4mm) concentrator column, IonPac CG and CS12A guard and analytical columns, CSRS ULTRA II (4-mm) suppressor and CD25A conductivity detector operated at 30°C. Cation runs were performed isocratically with 20 mM MSA at a flow of 1ml/min. Ambient air was sampled at 3L/min for the first 55 min of each hour and hourly averages

Cation runs were performed isocratically with 20 mM MSA at a flow of 1ml/min. Ambient air was sampled at 3L/min for the first 55 min of each hour and hourly averages of all species were reported in $\mu g \, m^{-3}$ via Chromeleon 6.8 and URGAIM software. An off-line calibration of the AIM-IC was performed in-field by directly injecting multiple ion standards into each IC. Following the campaign, it was determined that the sampling inlet (1/2" OD Teflon-coated aluminum tube breaching a 4" diameter PVC pipe) was not appropriate for the quantitative sampling of gases, especially soluble ones such as HNO₃ and NH₃, so only the PM_{2.5} measurements are used in this analysis.

2.4 Air quality modeling

Ground-based measurements were supported in real time by Environment Canada's AURAMS (A Unified Regional Air-quality Modelling System)which consists of three main components: (a) a prognostic meteorological model, GEM: Global Environmental Multiscale model (Cote et al., 1998); (b) an emissions processing system, SMOKE: Sparse Matrix Operator Kernel Emissions (Houyoux et al., 2000; CEP, 2003) and (c) an off-line regional chemical transport model, the AURAMS Chemical Transport Model (cf. Gong et al., 2006; Stroud et al., 2008; Cho et al., 2009; Makar et al., 2009; Smyth et al., 2009)

Following the conclusion of the BAQS-Met measurement campaign, AURAMSv1.4.0 was configured to run with three levels of nesting: a 2.5 km high resolution local domain nested within a 15 km regional domain, nested within a 42 km North American

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domain. Further details on the model setup for these simulations and a more detailed description of the model are given in Makar et al. (2010), this issue. Output of mixing ratios and mass loadings from the 2.5 km resolution domain at the surface for the Harrow grid-square was used to compare to measurements of ammonia and particulate ammonium, sulfate, and nitrate. Aerosol partitioning is calculated in AURAMS as a bulk calculation across all particle sizes using HETV (Makar et al., 2003), a vectorized form of ISORROPIA. AURAMS uses a speciated sectional approach to describe the aerosol size distribution: mass from the first 8 bins, encompassing the 0 to 2.5 µm range, were used to calculate the PM_{2.5} mass loadings of particulate matter for comparison to the size-selection used by the AIM-IC observations.

In order to obtain outputs of mixing ratios and mass loadings of atmospheric species, AURAMS solves a set of complex differential equations that describe the rate of change of each model species. The solution is split into different components (operators) which are solved in sequence, with the solution from each operator becoming the initial concentrations for the next operator in the sequence (Makar et al., 2010). AURAMS can be configured to track the change in mass of model species of interest for each operator for each time step. Using this mass-tracking, it is possible to obtain important model information on gas-particle partitioning, advection, vertical exchange and diffusion of a species of interest.

3 Results and discussion

3.1 Ammonia mixing ratio analysis

Near continuous measurements of ammonia were acquired using the QC-TILDAS at 1 Hz between 20 June to 13 July at the Harrow supersite. The ammonia mixing ratios were averaged to 5 min for viewing simplicity and are displayed in Fig. 1. We observed an average of approximately 2.5 ppb throughout the measurement campaign, with short-lived spikes of up to 30 ppb on the order of minutes and over 50 ppb on the

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order of seconds. The majority of these spikesoccurred at night, which we attribute to local emissions, such as from traffic, into a shallow nocturnal boundary layer. It is known that ammonia can be formed from the incomplete reduction of NO_x in light-duty automobiles equipped with three-way catalytic converters (Kean et al., 2009; Livingston et 5 al., 2009). We investigated the time series of other traffic-related species such as benzene, toluene, nitrogen oxides (NO_v) and carbon monoxide (CO) but could not exactly match ammonia spikes to any of these species. Nights on which we observed high spikes in ammonia coincided with high and variable mixing ratios of these other trafficrelated species. The lack of exact overlap in time may be due to the different location of instrument inlets, different data acquisition times and different types of automobile combustion engines. For example, ammonia is mostly emitted from light-duty vehicles (NPRI, 2010) while NOx is generally associated with heavy-duty diesel vehicles such as trucks. Li et al. (2006)also observed that not all automobiles emit ammonia under on-road driving conditions, and Whitehead et al. (2007) did not see a significant correlation between NO_x and NH₃ in the summer. Meteorological data were examined to further investigate possible sources of the ammonia spikes.

Polar plots of the dependence of ammonia mixing ratios on wind direction are shown in Fig. 2. These plots display 5 minute averaged ammonia data on the radial axis colouredas a function of (a) wind speed, (b) temperature and (c) relative humidity. The polar plots indicate sporadic high mixing ratios of ammonia originate in the east and the southwest of the measurement site, while binning the data into 10-degree intervals (Fig. 2d) show that the largest background source was to the Northwest. In the eastern quadrant, we observe many instances when the mixing ratio was 10 ppb or higher, characterized by low wind speeds, low temperatures (~15 °C) and high relative humidity (>75%), which are consistent with night-time conditions. A local road which was directly east of the measurement site may be the source of these night-time ammonia spikes. We have eyewitness evidence that street racing was occurring on some nights, and Livingston et al. (2009) states that aggressive driving is likely to lead to higher ammonia emissions. A secondary regime with frequent high ammonia events

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exists in the southwest, with higher wind speeds, relative humidity between 50-60% and high temperatures of approximately 25 °C, indicating daytime flow. Southwest of the measurement site was a field managed by Agriculture and AgrifoodCanada, growing mostly soybean and corn crops. On some days, employees were seen turning the 5 soil and working the field. As each point on the polar plot is a 5 min averaged data point, the short-lived but high ammonia mixing ratios events may mask any indication of a dominant source. Figure 2d shows the average ammonia mixing ratiosare approximately 2 ppb in each quadrant except for the northwest. This enhancement in the northwest may be due to a local unidentified agricultural or industrial source or transport from the Windsor/Detroit area, which is approximately 30 km away from Harrow. Although NH3 has a high deposition velocity and is expected to be lost quickly to the surface, transport of NH₃ over such distances has been previously observed (Sakurai et al., 2003).

There is no change in the average ammonia mixing ratio when air is coming from the south in the direction of Lake Erie. One would assume that an air mass passing over a water body would become depleted of ammonia, because the effective Henry's Law constant of ammonia is very high. On the other hand, the deposition velocity of surface reactive gases such as ammonia is lower over water than over land, possibly extending the lifetime of ammonia against deposition (Seinfeld and Pandis, 2006). When southerly air masses, consistent with onshore flow, impacted Harrow, lower ammonia mixing ratios were not observed. We investigated the ammonia time series for each day of the study looking for an increase in relative humidity and shift in wind direction to southerly that would indicate a lake breeze. Because lake breeze passages consistently occurred in the middle of the day, when emissions of ammonia are expected to the highest, it is also possible that the volatilization of ammonia from the agricultural fields in the 5 km between the lakeshore and the measurement site was sufficient to replenish the ammonia mixing ratio in the air that had been over the lake. However, we do not have any evidence that the ammonia mixing ratios at the edge of the lake were lower (or different) than at the measurement site.

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Four significant precipitation events were identified from the meteorological data and air mass flags. During all four precipitation events, the ammonia mixing ratios dropped between 1–2 ppb and remained low for the duration of the rain storm. It is known that ammonia is effectively scavenged by raindrops and can decrease during rainfall (Nowak et al., 2006). We were unable to fit for the scavenging coefficient due to both short duration of the rainfall and the low absolute changes in ammonia mixing ratios during some of the precipitation events. After the rain ceased, significant increases in mixing ratios were observed as demonstrated in Fig. 3. The increase in ammonia mixing ratios after the rain may be from increased microbial processes in soil and vegetation that can release ammonia. Roelle and Aneja (2002) observed soil surface emissions of NH₃ to be enhanced by rainfallin a North Carolina coastal plain.

3.2 Diurnal behaviour

For the remainder of the paper, we use hourly averages of ammonia observations with the short duration spikes removed, to match the acquisition time of the AIM-IC system. We investigated the time of day behaviour for ammonia, which is plotted in Fig. 4 along with particulate ammonium (both in $\mu g m^{-3}$), relative humidity and temperature. For ammonia, there is an average mass loading of 1.5 µg m⁻³ with a sharp morning rise and a slow decrease in concentration in the afternoon. The morning rise between 06:00 and 07:00 could be attributed to one of several reasons, including evaporation of dew, volatilization of particulate ammonium, emission from plant stomata, or mixing down of ammonia from the residual layer. We consider each of these hypotheses in turn. When the sun comes up and heats the surface, the evaporation of dew could release ammonia trapped in liquid water condensed on surfaces. Kruit et al. (2007) also observed a morning peak in concentrations which coincided with a decrease in leaf wetness. The BAQS-Met study did not include measurements of leaf wetness, but the measured relative humidity did exceed 95% on several nights during the measurement campaign. Figure 4illustrates that the relative humidity and temperature did not change significantly during the hours of 06:00-07:00, but increased later in the

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morning. If evaporation of dew is the reason for the NH₃ increase, we would expect it to be coincident witha large increase in temperature, which can heat the surface enough to allow for evaporation. It is possible that the temperature of the soil increases slightly earlier than the temperature of the atmosphere at 2 m (the height of our RH/T probe). However, the observation of a sharp morning rise in ammonia did not specifically correspond to mornings for which the preceding night had been particularly humid and dew was more likely to have formed, so it is unlikely that the evaporation of dew is the main contributor. Because the largest increase in ammonia also precedes the large changes in atmospheric relative humidity and temperature, and because the particulate ammonium is flat during that time, volatilization of ammonium is ruled out as a significant contribution to the morning rise in ammonia. Plant and/or soil processes may be a more likely reason for this morning rise, as observed by Bash et al. (2010) in a corn field. The local sunrise in Harrow at this time of year was around 05:45, and as an immediate response, plant stomata open for photosynthesis. During this time, release of ammonia from the apoplastic fluid can occur through the stomata, which may lead to an increase in concentration at this time. This increase would be more prominent if the emission is occurring into a shallow nocturnal boundary layer. On the other hand, the morning rise may also be due to the breakup of the nocturnal boundary layer. As the surface heats up and vertical mixing begins, the downward mixing of the residual layer, which may contain relatively higher ammonia, could lead to a surface increase. This effect was also observed by Walker et al. (2006) who attributed an early morning increase in NH₃ surface concentrations to a deposition flux. At Harrow, our observations of early morning concentration changes rule out a significant contribution from particle volatilization, but are not sufficient to identify whether the main cause of the morning increase is surface emissions or mixing down from above. In the future, simultaneous flux measurements would be ideal to elucidate the contributing processes.

The diurnal changes in aerosol species and their gaseous precursors are controlled by emission and deposition processes, horizontal and vertical transport and gas-particle partitioning. Between 10:00 and 15:00, we observe a decrease in measured ammonia corresponding in magnitude to an increase in ammonium, which coincides with high mass loadings of sulfate at the site. This decrease in gas phase ammonia is likely the result of uptake onto aerosols to form (NH₄)₂SO₄. In order to investigate this claim further we looked at the total ammonia (NH_x), which is the sum of ammonia and ammonium, shown as a stacked plot in Fig. 5a. The effect of gas-particle partitioning is removed in plots of NH_v, and the diurnal variability is thus only controlled by transport and vertical exchange. We observe higher values of NH, during the daytime hours, likely because the emissions of ammonia are more significant due to higher temperatures or because of the transport to the site of ammonium sulfate particles. Total ammonia is relatively constant during the hours of 10:00 and 15:00, indicating that gas-particle partitioning contributes significantly to the decrease in gas phase ammonia during this time. Figure 5b shows the total ammonia calculated from the AURAMS model output.Between 22:00 and 06:00, both the amount and behaviour of the measured NH, are well-represented by the model. During the hours of 10:00 and 15:00, when measured total ammonia is fairly constant at 5 µg m⁻³, the model results in Fig. 5b predict a large decrease in NH_v. Figure 5c shows the difference in NH_v between the model and the measurements as a function of time of day. We find a consistent low bias for NH_x in the model, which is the most dramatic during the daytime hours of 10:00 and 15:00, when sulfate levels are highest. Despite predicting an appropriate range of midday sulfate levels, the model is biased low for ammonium because it predicts much lower NH_x, which is insufficient to neutralize the aerosol.

Figure 6 displays the distribution of important ammonia mass tracking operators extracted from AURAMS for the lowest 6.89 m above the surface. The heterogeneous operator describes inorganic heterogeneous and aqueous phase chemistry (i.e. gas-

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particle partitioning), the vertical exchange operator includes the net change associated with emissions, deposition and vertical diffusion of ammonia, and the advection term describes mass changes associated with total three-dimensional advective transport. On average, the model predicts heterogeneous chemistry as a sink for ammonia, consistent with the existence of acidic particles in the model. The net vertical exchange term on average is positive, thus vertical exchange on average is a source of ammonia over Harrow, which is most likely to due to emissions. We see a low change in ammonia mass loadings from advection, which points to vertical exchange, local fluxes and associated chemistry as the controlling factors for high concentration of NH_x at our site.

The ability of the model to accurately describe the gas-particle partitioning of the measurements is further investigated by plotting the fraction of total ammonia in the gas phase in Fig. 7 (gas fraction ≡NH₃/NH₄). In the early morning, a higher fraction of ammonia is in the gas phase, while in the late morning to afternoon, the gas phase fraction decreases quickly. This is consistent with an early morning increase in gas phase ammonia concentrations from vertical transport or surface exchange, followed by gas-particle conversion in the afternoon. The model is biased slightly high at night, and low during the day, but appears to represent the overall diurnal cycle well, suggesting that on average, gas-particle partitioning is not the cause of the differences noted in Fig. 5. The low bias during the day is the result of the model also being biased low in NH_x, pointing to issues with the surface-atmosphere exchange. Additionally, when the measured $\mathrm{NH_{x}}$ is repartitioned offline using ISORROPIA and observed $\mathrm{SO_{4}^{2-}}$ and NO₃, the modeled gas-particle partitioning matches the observations very well. With the elimination of inorganic heterogeneous chemistry as a cause for the differences, emission, deposition and vertical diffusion remain as possible causes. As these three terms are described by a single operator in AURAMS, we could not identify the main cause of the discrepancy between the model and measurements.

The variability in the ammonia gas fraction predicted by the model, shown in the error bars, is much larger than observed. To better illustrate the extreme values in the model output, histograms of the observed and modelled gas fractions were calculated

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and displayed in Fig. 8. The observations are clustered in the middle of the histrogram, whilethe model tends to be bi-modal, with all the NH_x predicted in the gas phase (gas fraction >0.8) or in the particle phase (gas fraction < 0.05).In Fig. 9, the gas fraction correlates strongly with the sulfate levels in both measurements (a) and model (b). However, the measured gas fraction rarely drops below 0.1 even when sulfate levelsexceed $10 \, \mu \mathrm{g \, m^{-3}}$, and never rises above 0.8, even when sulfate is less than $1 \, \mu \mathrm{g \, m^{-3}}$. This implies some buffering process that is not represented in the model, which allows additional ammonia to enter the atmosphere in the presence of a large condensation sinkthat exists when there are high mass loadings of SO_4^{2-} present.

3.4 Ammonia bi-directional flux

Like most chemical transport models, AURAMS uses a thermodynamic model based on ISORROPIA (Makar et al., 2003) to solve for the equilibrium concentrations of ammonia and ammonium in each grid cell. When we used our observations as input to the offline version of the same thermodynamic model, ISORROPIA v2.1, the gas-particle partitioning matched the observations very closely, confirming that the assumption of equilibrium partitioning was valid. Thus we turn our attention to other processes in the model that can be responsible for the differing behaviour between model predictions and measurements evident in Fig. 8. The total ammonia predicted in each grid cell in the model is governed by the difference between emissions and deposition, and horizontal advection. According to the model, the contribution of three-dimensional advection was minimal for both NH₃ and NH₄ except at low concentrations. Both the emission and dry deposition of NH₄ are negligible, so the dominant process driving variability of total ammonia in the boundary layer is the surface-atmosphere exchange of gaseous ammonia. For NH₃, AURAMS uses an emission inventory that depends of the time of day and the day of week, and a deposition velocity based on the local meteorology and land surface type. In contrast to the surface-atmosphere exchange parameterizationscurrently used in AURAMS, the net exchange of ammonia between the atmosphere and some surfaces is thought to be governed in part by the equilibrium between the

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atmospheric ammonia and the ammonium in plant apoplastic fluid (Schjoerring et al., 1998). In this pathway, ammonia emissions occur if the stomatalcompensation point is greater than the atmospheric mixing ratios (Farquhar et al., 1980). Conversely, deposition will occur if the compensation point is lower than the atmospheric mixing ratios. Thus atmospheric gas phase ammonia is simultaneously participating in two equilibria, one with atmospheric particulate ammonium, and one with a pool of ammonium at the surface. If the condensation sink increases (e.g. by an increase in aerosol sulfate), the atmosphere is depleted of ammonia, potentially lowering the ambient mixing ratios below the compensation point, and triggering emissions from the plants and soil. This phenomenon was previously implied in a study by Sutton et al. (1994)where high emissions of NH3 were observed at the same time as high SO2 concentrations, from which the authors inferred acidic aerosol loadings.

One interpretation of the difference between the measured and modelled distributions of gas fraction shown in Fig. 8 is that in the real atmosphere, the direction of the surface flux of ammonia may depend on the atmospheric mixing ratio of NH₃, thereby buffering the gas fraction. In the current version of AURAMS, the dry deposition flux of ammonia depends on the ambient ammonia mixing ratios. However, the magnitude of AURAMS' ammonia emissions flux is not coupled to the model's ambient ammonia mixing ratios; the model does not simulate bi-directional fluxes. There have been many recent papers discussing methods to fully couple the bi-directional flux of ammonia into a chemical transport model (van Pul et al., 2009; Kruit et al., 2010; Massad et al., 2010; Zhang et al., 2010), wherein the ammonia emissions flux is to some extent dependant on the mixing ratios of ammonia present in the atmosphere.AURAMS does not currently include a bi-directional flux parameterization for ammonia, so when a large amount of sulfate exists in the atmosphere, all of the ammonia condenses, generating a significant number of points in the model at zero gas fraction. However, bi-directional flux suggests that the depletion of gas phase ammonia by condensation would lead to larger fluxes from the surface to the atmosphere, replenishing the gas phase ammonia and moving the observations more toward the middle of the histogram

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in Fig. 8. Rather than defining the emission and deposition terms in the model as separate processes, these can be represented by a net flux, whose direction depends on how ambient ammonia mixing ratios compare to a compensation point. To accurately represent this process in the model, it would have to be incorporated online so that it was fully coupled with all of the other source and sink processes. As a simple sensitivity test of this concept, starting with the AURAMS output at Harrow, we successively replaced all the model gas phase ammonia values with a single compensation point value, and then repartitioned the total ammonia(NH_v) offline using the ISORROPIA thermodynamic model. For a single compensation point value, we performed several iterationsin ISORROPIA each time replacing the calculated gas phase amounts in the output with the compensation point asthe input of the next run. Eventually, the output converges to the compensation point for gas phase ammonia and enough particulate ammonium for fully neutralized aerosol. Single compensation point values of 0.5, 1.0, 1.5 and 1.7 µg m⁻³ were tested, chosen to reflect the vegetation type and estimated nitrogen content at our measurement site. The results shown in Fig. 10 were achieved using a compensation point of 1 µg m⁻³ and indicate that this approach was successful at drawing down the number of zero gas phase points in the model histogram. The number of high gas fraction cases increased when all of the added NH_x remained as NH₃. This tended to occur in situations where modeled sulfate and nitrate were biased low and could not accommodate additional ammonium. Additionally, we also tested a temperature dependent compensation point as described by Nemitz et al., (2001). Here, we used an average value for the stomatal apoplastic ratio ($\Gamma_{\rm s}$) from Kruit et al. (2010). However, this approach does not accurately reflect our data, as a temperature dependent compensation point will predict the highest NH₃ in the middle of the day, contrary to what was observed (Fig. 4). Possible reasons for this inconsistency could be that there is a time-of-day dependency to the wind direction, and hence upwind Γ_s , or that the full coupling of the gas-particle partitioning and surfaceatmosphere exchange of ammonia is not achieved through the full boundary laver instantaneously. While this simplified approach does not reproduce the observations

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perfectly, it does imply that the inclusion of a bi-directional flux of ammonia may improve model-measurement agreement regarding the gas fraction of ammonia.

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Gas phase mixing ratios of ammonia in an agricultural region in Southwestern Ontarioaveraged 2.5 ppb, with sharp spikes from traffic at night. The diurnal profile of ammonia shows an increase in the early morning thatis attributed to plant/soil emissions or mixing down of stagnant air upon the breakup of the nocturnal boundary layer. Ammonia decreased in the afternoon as a result of condensation to aerosol containing high levels of sulfate. Observations showed the ratio of NH_x/SO_4^{2-} always exceeded 2, however, the regional model AURAMS was consistently biased low in regards to the neutralization of sulfate at this site. These observations in an agricultural region suggest that the presence of large amounts of acidic sulfate aerosol in the atmosphere can induce high net emission rates of ammonia due to bi-directional flux. The timescale of this flux is likely quite short because even airmasses which had recently been over Lake Erie and contained high amounts of sulfate were found to be fully neutralized upon their arrival at Harrow (less than 1 hour from shoreline). This analysis is the first to use simultaneous observations of atmospheric ammonia and ammonium to examine the coupling between gas-particle and surface-atmosphere equilibria governing ammonia mixing ratios. Our observations indicate that in regions where area-wide sources dominate, the accuracy of chemical transport models may be improved by including a fully coupled bi-directional flux parameterization to make accurate predictions of atmospheric ammonia. Because the timescales of the gas-particle and surface-atmosphere equilibrium processes are likely similar, and comparable to the times required for vertical mixing, simultaneous measurements of the concentrations, and ideally fluxes, of gas phase NH₃, and particulate ammonium are needed to provide full observational constraints. Additionally, this work suggests that the emission rates of ammonia in some regions may depend on the atmospheric mixing ratios of not only ammonia, but **ACPD**

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also the mass loadings and composition of aerosol. This will have implications both for efforts to model ammonia emissions and also to implement ammonia emission reductions to control $PM_{2.5}$.

Acknowledgements. The QC-TILDAS and AIM-IC were available through infrastructure support to J.G.M. from the Canada Foundation for Innovation and the Ontario Research Fund. R.A.E. was supported by the Research Affiliate Program at Environment Canada (2008–2010) while carrying out much of this work. The authors acknowledge Agriculture and Agri-food Canada for providing the site at Harrow and Al Melanson, from the Ontario Ministry of Environment, for his efforts in coordinating the site infrastructure and logistics.

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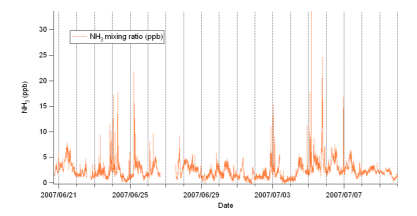


Fig. 1. Five minute averaged ammonia mixing ratios during BAQS-Met fluctuated around 2.5 ppb, with higher, shorter spikes up to 30 ppb mostly occurring during night-time.

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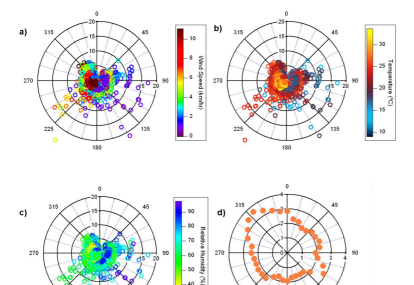


Fig. 2. Wind direction polar plots of ammonia mixing ratios (ppb) as a function of **(a)** wind speed, **(b)** temperature and **(c)** relative humidity. Panel **(d)** shows all data binned into 10 degree bins and indicates a dominant source in the northwest.

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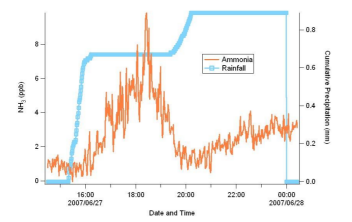


Fig. 3. Increase in ammonia mixing ratios directly after rainfall may be due to increased biological activity.

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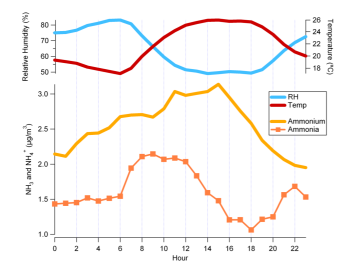


Fig. 4. Average diurnal cycles of relative humidity and temperature with ammonia and ammonium concentrations. Ammonia concentrations display a rise in the early morning hours, followed by decrease in the afternoon.

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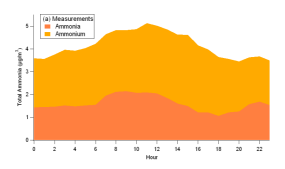
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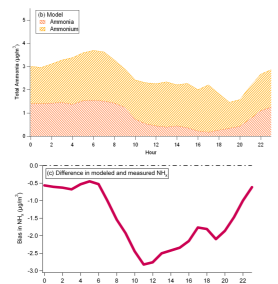


Fig. 5. Stacked plots of the components of total ammonia (top of the trace) for **(a)** measurements and **(b)** model. The difference between the modeled and measured NH_x is show in **(c)** and indicates model is biased low in NH_x .

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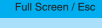


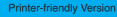
















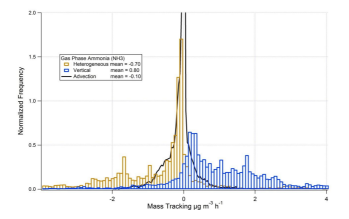


Fig. 6. Distribution of important ammonia mass tracking operators from AURAMS shows heterogeneous chemistry process to be a net sink of ammonia, while vertical exchange processes are a net source.

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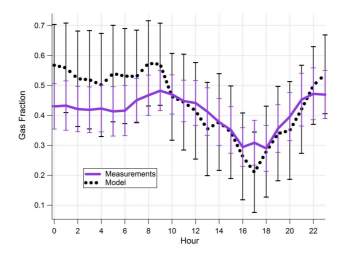


Fig. 7. Diurnal cycles of the measured and modeled gas fraction (NH₃/NH_x) with the error bars representing the 95% confidence interval.

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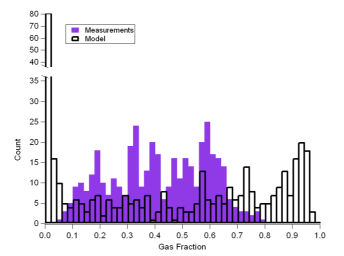


Fig. 8. While the measured gas fraction distribution is clustered in the middle of the histogram, the model is largely bi-modal.

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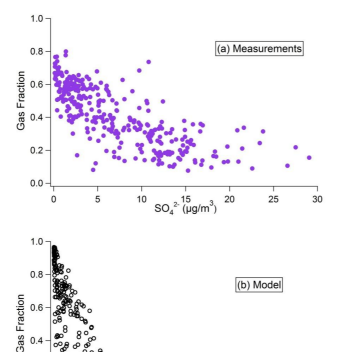


Fig. 9. Gas fraction of (a) measured and (b) modelled ammonia as a function of $PM_{2.5}$ sulfate.

 $SO_4^{2-} (\mu g/m^3)$

25

30

0.2

0.0

5

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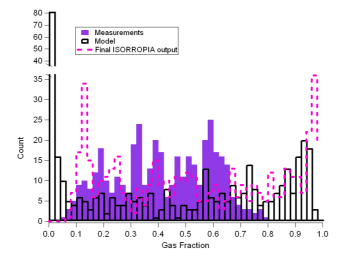


Fig. 10. Histogram of the measured and modelled gas fraction from Fig. 8, now including the result of final offline run of ISORROPIA, iterated until the aerosol was fully neutralized and the gas phase concentration was equal to a pre-determined compensation point.

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