Modelling the impacts of ammonia emissions reductions on North American air quality

P. A. Makar¹, M. D. Moran¹, Q. Zheng¹, S. Cousineau², M. Sassi², A. Duhamel², M. Besner², D. Davignon², L.-P. Crevier², and V. S. Bouchet²

¹Air Quality Research Division, Science and Technology Branch, Environment Canada, Toronto, Ontario, Canada
²Air Quality Model Applications Section, Meteorological Service of Canada, Environment Canada, Montreal, Quebec, Canada

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Correspondence to: P. A. Makar (paul.makar@ec.gc.ca)

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Abstract

A unified regional air-quality modelling system (AURAMS) was used to investigate the effects of reductions in ammonia emissions on regional air quality, with a focus on particulate-matter formation. Three simulations of one-year duration were performed for a North American domain: (1) a base-case simulation using 2002 Canadian and US national emissions inventories augmented by a more detailed Canadian emissions inventory for agricultural ammonia; (2) a 30% North-American-wide reduction in agricultural ammonia emissions; and (3) a 50% reduction in Canadian beef-cattle ammonia emissions. The simulations show that a 30% continent-wide reduction in agricultural ammonia emissions lead to reductions in median hourly PM$_{2.5}$ mass of $<1\mu g m^{-3}$ on an annual basis. The atmospheric response to these emission reductions displays marked seasonal variations, and on even shorter time scales the impacts of the emissions reductions are highly episodic: 95-percentile hourly PM$_{2.5}$ mass decreases can be up to a factor of six larger than the median values.

A key finding of the modelling work is the linkage between gas and aqueous chemistry and transport; reductions in ammonia emissions affect gaseous ammonia concentrations close to the emissions site, but substantial impacts on particulate matter and atmospheric deposition often occur at considerable distances downwind, with particle nitrate being the main vector of ammonia/um transport. Ammonia emissions reductions therefore have trans-boundary and possibly trans-oceanic consequences downwind. Calculations of critical-load exceedances for sensitive ecosystems in Canada suggest that ammonia emission reductions will have a minimal impact on current ecosystem acidification within Canada, but may have a substantial impact on future ecosystem acidification. The 50% Canadian beef-cattle ammonia emissions reduction scenario was used to examine model sensitivity to uncertainties in the new Canadian agricultural ammonia emissions inventory, and the simulation results suggest that further work is needed to improve the emissions inventory for this particular sector.
1 Introduction

The chemistry describing the interactions of atmospheric ammonia (NH$_3$) with other atmospheric constituents has been well established through field and laboratory studies. While very high concentrations of ammonia gas are known to have health impacts (e.g., Stilg, 1994), its role in the creation of airborne particulate matter (PM) at lower concentrations is of interest due to the known effects of fine particulate matter on human health (cf. Schwarze et al., 2006).

The creation of airborne PM from ammonia is dependant on the presence of other precursor gases, primarily directly-emitted sulphur dioxide and nitrogen oxides (SO$_2$ and NO$_x$; Seinfeld and Pandis, 1998). SO$_2$ may oxidize via gas-phase (Stockwell and Calvert, 1983) and/or aqueous-phase (Coste and Courtier, 1936; Junge and Ryan, 1958; Hermann et al., 2000, 2005) reactions, creating sulphuric acid gas or sulphuric acid ions in cloud or rain water, respectively. The vapour pressure of sulphuric acid gas is sufficiently low that almost all of the gas created will partition to the particle phase, either through condensation on existing particles or nucleation of new particles. Sulphuric acid created in cloud and/or rainwater may be transferred to the particle phase upon droplet evaporation.

Ammonia affects aqueous-phase chemistry through the provision of a weak base; the hydrogen ion concentration will be inversely proportional to the ammonia partial pressure. Reductions in the hydrogen ion concentration due to excess ammonia will allow a greater aqueous uptake of SO$_2$ in cloud water in the form of the bisulphite ion (HSO$_3^-$). The latter may be oxidized by hydrogen peroxide, ozone, organic peroxides, or catalytic oxygen reactions to bisulphate and sulphate ions, the ionic equilibrium products of sulphuric acid dissociation (Hermann et al., 2000, 2005). H$_2$O$_2$ is believed to be the dominant aqueous-phase oxidant of HSO$_3^-$, but the strongly pH-dependent oxidation by O$_3$ becomes more important as pH increases or when H$_2$O$_2$ has been depleted (e.g., Fung et al., 1991). The relative contribution of these two oxidants to aqueous-phase sulphate formation is therefore influenced by NH$_3$ levels. An additional process
of importance for ammonia chemistry is the formation of gaseous nitric acid (HNO₃) through well-known “NOₓ termination” reactions (Seinfeld and Pandis, 1998); HNO₃ may in turn participate in aqueous reactions with the ammonium ion, or in particle-phase chemistry.

Laboratory studies and related thermodynamics of high-concentration particle ammonium chemistry are well established (cf. D’Ans, 1913), and observations of ammonium, sulphate, and nitrate in PM have appeared in the literature over the past sixty years (cf. Robbins and Cadle, 1958; Fenn et al., 1963; Spurny and Heard, 1969; Heard and Wiffen, 1969; Gordon and Bryan, 1973; Anlauf et al., 1978; Brosset, 1978; Stelson et al., 1979; Tanner, 1983). The partitioning between different phases, including gases, may be predicted using fundamental thermodynamics theory (cf. Kusik and Meissner, 1978) in box models (e.g., Ansari and Pandis, 1999) or regional models such as AURAMS (Makar et al., 2003; Gong et al., 2006).

The concept of ammonia limitation may be introduced, in order to better understand the aqueous and particulate chemistry of ammonia. Small perturbations in the ammonia concentration will have a significant effect on particle concentrations while the ammonia remains insufficient to charge-balance the available anions. Ammonia emissions changes for this situation may therefore have a significant impact on particle concentrations, and ammonia availability limits the inorganic fine-mode particle mass.

When the ammonia is available in excess, perturbations to the ammonia concentration will have little impact on particulate matter mass; an anion-limited environment. Here, the atmosphere is said to be “ammonia-limited” when there are insufficient moles of total ammonium (ammonia gas + all ammonium as aqueous phase or particulate ions, or crystalline ammonium salts) to charge balance the total number of moles of sulphate. A broader definition of ammonia-limited environments includes a requirement for charge balancing with nitrate and other anions; the use of sulphate restricts the definition to strongly ammonia-limited environments, where changes in ammonia emissions would be expected to result in a significant change in particulate matter mass.

Comprehensive Eulerian regional models are useful tools for studying the poten-
tial impacts of ammonia emissions on atmospheric particle formation and deposition to sensitive ecosystems. The first generation of these models were designed to predict the gas-phase concentrations of acidifying gases, ozone, other reactive gases, and wet and dry deposition of atmospheric acidic species (e.g., Chang et al., 1987; Venkatram and Karamchandani, 1988). Later work extended these models to include size-distributed PM (Binkowski and Shankar, 1995). Further developments within the last decade included the introduction of more detailed inorganic and organic particulate chemistry, and the introduction of size-resolved and speciated PM (e.g., Binkowski and Roselle, 2003; Gong et al., 2006). More recently, these models have begun to be used to investigate the role of ammonia and other nitrogen compounds in atmospheric chemistry and deposition (Mathur and Dennis, 2003; Ying and Kleeman, 2006; Phillips et al., 2006; Luo et al., 2007; Quan and Zhang, 2008; Wang et al., 2008). A recent extension of deposition modelling to assess environmental impacts is the calculation of exceedances of acid-deposition critical loads (defined below) in order to estimate the impact of deposition on sensitive ecosystems (Fowler et al., 1998; Dentener et al., 2006; Spranger et al., 2008; Fenn et al., 2008).

Various emission control strategies have been suggested in recent years to reduce ambient PM levels in North America, including controls on ammonia emissions. In this paper, we describe the application of a comprehensive regional air-quality model to predict the likely effects of reductions in North American emissions of agricultural ammonia on the mass and composition of atmospheric PM and on the amount of acid deposition to sensitive ecosystems. Three one-year simulations, a 2002 base case and two hypothetical NH$_3$ emission scenarios, have been run and analyzed. The next section describes the study methodology. Section 3 summarizes a performance evaluation for the 2002 base case, and Sect. 4 analyzes the results of the two emission scenarios. Finally, conclusions and recommendations for further study are provided in Sect. 5.
2 Methodology

2.1 Modelling system description

AURAMS (A Unified Regional Air-quality Modelling System) consists of three main components: (a) a prognostic meteorological model, GEM (Global Environmental Multiscale model: Côté 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 (CTM: Gong et al., 2006).

The GEM meteorological model is an integrated weather forecasting and data assimilation system that was designed to meet Canada’s operational needs for both short- and medium-range weather forecasts. For the 2002 simulation, GEM version 3.2.0 with physics version 4.2 was run on the variable-resolution global horizontal grid centred on North America.

Files of gridded hourly emission fields (including ammonia) for input by the AURAMS CTM were prepared using version 2.2 of the SMOKE emissions processing system for four major emissions streams: on-road mobile sources; area and offroad-mobile sources, minor point sources; and major point sources. Emitted (i.e., “primary”) PM from these sources is speciated within the AURAMS CTM based on speciation profiles for each emissions stream, and these profiles include some primary particulate ammonium. The mass of ammonium emitted as primary particulate ammonium, however, is much smaller than the mass of gaseous ammonia emissions, and most particulate ammonium (p-NH₄) is created through chemical reactions (i.e., “secondary” sources). A simulation of this partitioning between ammonia gas and p-NH₄ takes place within the chemical modules of the AURAMS CTM.

The multi-pollutant, regional AURAMS CTM was developed as a tool to study the formation of ozone, PM, and acid deposition in a single “unified” framework. The PM size distribution in this study was represented using 12 size bins ranging from 0.01 to 41 µm in Stokes diameter and nine chemical components: sulphate (p-SO₄); nitrate...
(p-NO₃); ammonium (p-NH₄); elemental carbon (EC); primary organic matter (POM); secondary organic matter (SOM); crustal material (CM); sea salt; and particle-bound water. PM is assumed to be internally mixed in each size bin. Process representations in version 1.3.1b of the AURAMS CTM include emissions from surface and from elevated sources, horizontal and vertical advection, vertical diffusion, gas-phase, aqueous-phase, and inorganic heterogeneous chemistry, secondary organic particle formation, dry and wet deposition, and particle nucleation, condensation, coagulation, sedimentation, and activation (Gong et al., 2006). Up to 157 model species (gases and speciated particle size bins) may be selected as model output, although summary measures such as PM₂.₅ bulk mass are compared to observations here.

AURAMS inorganic particle components are reported as the mass of sulphate, nitrate, and ammonium within each particle bin size, but within the model, the inorganic heterogeneous chemistry module (Makar et al., 2003) performs equilibrium calculations to determine the relative amounts of mass of ammonium sulphate ((NH₄)₂SO₄(s)), ammonium bisulphate (NH₄HSO₄(s)), letovicite (((NH₄)₃H(SO₄)₂(s)), ammonium nitrate (NH₄NO₃(s)), and the ammonium (NH₄⁺(aq)), sulphate (SO₄²⁻(aq)), bisulphate (HSO₄⁻(aq)), and nitrate ions (NO₃⁻(aq)). The reported sulphate mass (p-SO₄) is thus the sum of sulphate mass from all particle components containing sulphate, with similar sums for the nitrate and ammonium mass.

The representation of dry deposition of ammonia gas within AURAMS follows Zhang et al. (2002); deposition is parameterized as a weighted combination of the deposition properties of ozone and SO₂. Dry deposition of p-NH₄ is a function of particle size (Zhang et al., 2001). It should be noted that AURAMS does not include the possible “co-deposition” of SO₂(g) and NH₃(g). Some researchers have found evidence of enhanced deposition of both gases when both are present at the same site (e.g., Neirynck et al., 2005; Van Hove et al., 1989; Adema et al., 1986). Others have found no effect (Erisman et al., 1994a, b), and Sutton et al. (1994) found enhanced emissions of ammonia gas from natural surfaces when ambient NH₃ is present. More observational work on co-deposition is needed before parameterizations for this process may
be reliably included in air-quality models.

The time-invariant, vertically-varying chemical lateral boundary conditions used in AURAMS CTM are taken from a variety of sources. Latitudinally-dependent O₃ boundary conditions were taken from a monthly-varying climatology (Logan, 1998). CO boundary conditions were derived from vertical profiles in Wang et al. (1999), with a simple latitudinal dependence of concentration peaking at 45 N, in rough accord with satellite observations. Speciated particulate boundary conditions (including p-NH₄) were based on data collected at an elevated site on Whistler Mountain on the Canadian west coast (MacDonald et al., 2006), with a similar simple latitude dependence assumed as for CO. Seasonally-varying profiles of the concentrations of other reactive gases (including NH₃) were taken taken from a set of “clean” chemical boundary conditions from simulations of the ADOM regional acid-deposition model (Scire et al., 1986; Fung et al., 1991).

2.2 Model domain, grid discretization, and simulation period

The GEM horizontal grid consisted of 353×415 grid points on a rotated latitude-longitude map projection with grid spacing of approximately 24 km (0.22°) on the 270×353 uniform regional “core” grid. In the vertical 28 hybrid-coordinate levels reached from the Earth’s surface to 10 hPa, with layer thickness increasing monotonically with height. A time step of 450 s was used.

The uniform horizontal grid used for the AURAMS CTM was 150×106 in size and spanned the North American continent on a secant polar-stereographic projection true at 60° N, with a horizontal grid spacing of 42 km (see Fig. 1). Twenty-eight terrain-following vertical levels stretched telescopically from the Earth’s surface to 29 km, with the first three levels at 0, 13.9, and 55 m a.g.l. An advective time step of 900 s was used, and AURAMS-predicted fields were output hourly. Both GEM and the AURAMS CTM were run for the 13-month period from 1 December 2001 to 31 December 2002, where the first month was treated as a spin-up period for the AURAMS CTM. GEM was run from analyzed fields for 396 overlapping 30-h segments starting 24 h apart,
where the first six hours of each segment were treated as a “spin-up” period and were discarded. The remaining 24 h of consecutive simulations were then “stitched” together to create a complete set of meteorological fields with a 900 s timestep for input to the AURAMS CTM. The CTM itself was run in three segments, with a one-month spin-up for each segment, allowing an entire year’s simulation to be run in parallel on multiple processors in order to reduce simulation “wall-clock” time.

2.3 Description of emissions scenarios

The hourly gridded anthropogenic emissions files used in this study were generated using SMOKE v2.2 (http://www.smoke-model.org/index.cfm) based on the 2002 Canadian (obtained from Environment Canada), 2002 US (obtained from US EPA), and 1999 Mexican (obtained from US EPA) national criteria-air-contaminant inventories. Biogenic emissions are calculated on-line in the AURAMS CTM using BEIS version 3.09 (Biogenic Emissions Inventory System: CEP, 2003). The following three 2002 annual emissions scenarios were considered.

1. Base Case: Canadian 2002 ammonia emissions from the default national inventory were replaced with those resulting from a more detailed inventory constructed as part of the National Agri-Environmental Standards Initiative (NAESI), a multi-year study which included the collection of Canada-specific emission factors and activity levels (Ayres et al., 2009; Bittman et al., 2009). The scenario using these combined emissions inventories will be referred to hereafter as the “Base Case”. The kilotonne emissions of ammonia gas for the base case on a seasonal basis are shown in Fig. 2.

2. 30% agricultural NH₃ emissions reduction, Canada and US: The base case’s agricultural emissions of NH₃ (including emissions from animal husbandry and from fertilizer application) were reduced by a factor of 30% at all times and locations in both countries.
3. 50% Canadian beef cattle NH$_3$ emissions reduction: a 50% reduction in Canadian emissions from this single sub-sector was implemented in order to test model sensitivity to sub-sector-specific inventory uncertainty estimates of a factor of two.

2.4 Metrics and Diagnostic Fields for Scenario Analysis

The key species of interest in this study are the model-predicted values of gaseous ammonia, SO$_2$, and nitric acid, as well as the total PM$_{2.5}$ mass and PM$_{2.5}$ inorganic composition. Differences between base case and scenario (i.e., base case value – scenario value) for these species show the impacts of the change in emissions of NH$_3$, with positive values indicating decreases in the mass or concentration arising from the reduction in NH$_3$ emissions.

In order to explain the chemistry associated with the base case, and the chemical reasons for the changes associated with the emissions-reduction scenarios, four chemical metrics based on the ambient air concentrations of several species have been employed. The metrics and their interpretation are given in Table 1.

A number of diagnostic deposition fields were also calculated to help quantify the impacts of changes in NH$_3$ emissions on atmospheric chemistry and deposition. The derived deposition fields include the total amount of sulphur deposited per season (sum of wet deposition and dry deposition of all species containing sulphur), the total amount of nitrate deposited, the total amount of ammonium deposited, and the total amount of nitrogen deposited. Subcomponents of these diagnostics will also be occasionally referenced (e.g., the amount of wet-deposited sulphate+SO$_2$ as a fraction of the total sulphur deposition).

Another important set of diagnostic outputs calculated for the AURAMS analysis were exceedances of annual critical loads in Canada. The “critical load” of an ecosystem refers to its ability to buffer acidifying precipitation. The underlying concept is that an ecosystem will have the ability to absorb a certain amount of acidifying sulphur and nitrogen compounds, including NH$_3$ and p-NH$_4$, without damage to the ecosystem itself. If the rate of deposition of these compounds exceeds the rate at which the
ecosystem can naturally absorb the compounds, however, ecosystem damage begins to occur. The maximum amount of acidifying mass that an ecosystem can absorb per unit area in a year is known as its annual critical load, and any additional deposited mass that exceeds that amount is known as an annual critical load exceedance (e.g., Jeffries et al., 1999; Hall et al., 2001; Jeffries and Ouimet, 2004; McNulty et al., 2007). Critical loads depend on local bedrock type, soil type and thickness, and other factors.

Sulphur deposition is essentially entirely acidifying, but nitrogen has a large biological activity, and may be stored in various catchment compartments within an ecosystem (Jeffries and Ouimet, 2004). Eventually, though, an ecosystem may reach a steady state with regard to nitrogen deposition (termed “nitrogen saturation”), after which all further nitrogen deposition is acidifying. Sulphur critical-load exceedance thus describes conditions where immediate ecosystem damage will occur, whereas sulphur + nitrogen critical-load exceedance describes conditions where ecosystem damage will once the ecosystem’s ability to absorb nitrogen is overwhelmed. Critical-load exceedances for sulphur + nitrogen thus describe a worst-case scenario, in which all of the deposited nitrogen is assumed to be acidifying. At the current time, Canadian ecosystems are not nitrogen-saturated (Jeffries and Ouimet, 2004); exceedances of sulphur + nitrogen critical loads thus indicate the potential for future ecosystem damage, as opposed to current ecosystem damage.

3 Model performance evaluation for the base case

The statistics used here for evaluation of the base case for the AURAMS simulations were used previously as part of an extensive AURAMS performance evaluation against measurements for the 2002 calendar year (Moran et al., 2007, 2008). The statistical measures used for the comparison are shown in Table 2. Measurements from 15 different Canadian and US air-chemistry and precipitation-chemistry networks and sub-networks were used to evaluate the base-case fields.

A number of steps were followed in preparing the measurements for comparison...
with model predictions. For example, units were adjusted as required to a common set of units (e.g., concentrations at ambient conditions to concentrations at STP). Data records from individual stations were screened for temporal completeness, and if they passed, then measurements were combined to create seasonal and annual values for the station. In locations where more than one station was located in an AURAMS CTM grid cell, the measurements were averaged. Even so, measurements from multiple air-chemistry networks are quite heterogeneous, since individual networks have different goals and objectives, choose different types of sampling locations, employ different sampling instruments, techniques, and protocols, and measure different species (e.g., Eder and Yu, 2006). For example, individual networks have very different sampling periods, ranging from hourly to weekly, and sampling intervals that vary from hourly, to 1 day in 6, to weekly. Combining measurements from different network does provide the benefits of increased sample size, spatial coverage, and spatial density, but the price paid is greater variability within the combined measurement data set.

The resulting annual statistics for ambient concentrations of the key species related to $p$-$\text{NH}_4$ formation and removal, with the exception of $\text{NH}_3$, for which routine measurements are not available, are shown in Table 3, and those relating to wet deposition are shown in Table 4.

These comparisons to observations show that, on an annual basis, AURAMS underpredicted the base-case PM$_{2.5}$ bulk mass by 31%, the PM$_{2.5}$ inorganic species concentrations by 18% to 19%, the concentration of inorganic ions in precipitation by 11% to 33%, and the wet deposition of inorganic ions by 6% to 24% (negative normalized mean biases). The implication of the comparison is that the model estimates for the base case for the PM$_{2.5}$ components and precipitation species are likely to be lower than the ambient atmosphere, and that model-predicted critical-load exceedances, described below, are likely to be underestimates. The impact of the model bias on the scenarios is harder to quantify. The usual assumption that is made is that the bias will be linear, so that the scenarios will have the same normalized biases as the base case. The impacts of ammonia emission reductions on the real atmosphere are therefore ex-
pected to be similar in relative magnitude and sign as simulated here but may vary in the absolute sense.

4 Scenario analysis

4.1 Analysis of the base case

One means of identifying NH$_3$-limited environments is to calculate the total (ammonia + ammonium) to sulphate mole ratio (cf. Table 1). Seasonal fields of this metric for the 2002 base case are shown in Fig. 3. Yellow to red areas in this figure have a large excess of NH$_3$ and are not strongly NH$_3$-limited: significant reductions in NH$_3$ emissions would be required to reach NH$_3$-limited conditions. These areas tend to correspond to areas of high NH$_3$ emissions (cf. Fig. 2). Green to blue regions, on the other hand, are NH$_3$-limited, with more acidic conditions. Reductions in NH$_3$ emissions in these areas would have an immediate impact on ambient PM$_{2.5}$ levels. Pronounced seasonal and local variations can also be seen; summer and winter have the largest extent of NH$_3$-limited regions.

One implication of the above analysis is that the regions that may benefit from NH$_3$ reductions will not necessarily be the regions that have the highest NH$_3$ emissions. For example, the region of greatest NH$_3$ saturation in Fig. 3 is centered on the US states of Minnesota, South Dakota, Nebraska, and Iowa – this also corresponds to the region of greatest NH$_3$ emissions. Reductions in NH$_3$ in this region are unlikely to significantly impact PM concentrations, due to the locally NH$_3$-saturated chemistry. However, further to the east and downwind from this source region are the NH$_3$-limited regions of the Great Lakes, the Ohio River Valley, and the Appalachian mountains. Depending on the extent to which NH$_3$ is transported, PM reductions are likely to occur downwind. The impact of NH$_3$ reductions will therefore be a combination of local chemistry and transport from higher-emission NH$_3$-saturated regions to lower-emission NH$_3$-limited regions.
4.2 Analysis of continental agricultural NH$_3$ emission reduction scenario

In this scenario, NH$_3$ emissions from agricultural sources were decreased uniformly by 30% in both Canada and the United States. In the following figures, the difference between base-case and scenario concentration fields will be displayed [\((\text{base case}) - (\text{scenario})\)], and the same difference formats will be used for the metrics. Positive values in the difference plots thus indicate decreases in the scenario concentrations relative to the base case; negative values indicate increases in the scenario concentrations relative to the base case. It is also important to note that seasonal average differences are being displayed unless noted otherwise; within each season, shorter duration events will occur with larger (and smaller) impacts than those depicted here.

4.2.1 NH$_3$ concentrations

The greatest decreases in seasonal NH$_3$ concentrations (Fig. 4) are closely matched with the locations of the main NH$_3$ emissions regions (Fig. 2). The effect of reductions in NH$_3$ emissions on NH$_3$ gas concentrations is therefore primarily local; most of the NH$_3$ is removed close to the source (either through deposition or conversion to PM).

The influence of seasonal variations in the NH$_3$ emissions can also be seen in Fig. 4. Agricultural emissions in North America are highest in the spring and lowest in the winter (e.g., Gilliland et al., 2006). As a consequence, the predicted change in concentration of NH$_3$ gas in the winter is lower than in the other seasons. For example, the decrease in ammonia concentrations in the high emissions region of Southern Minnesota in the summer is on the order of 2.5 ppbv, while the wintertime value in the same region is on the order of 0.5 ppbv.

4.2.2 PM$_{2.5}$ concentrations

The impact of the NH$_3$ emissions reductions on average seasonal PM$_{2.5}$ mass is shown in Fig. 5. Seasonally-averaged PM$_{2.5}$ changes resulting from a 30% reduction in NH$_3$
emissions range from an increase of 0.07 to a decrease of 3.99 $\mu g m^{-3}$. The greatest overall reductions in mass occur in spring and summer and occur in specific regions. The largest of these regions is centered over the Ohio River valley and Southern Ontario; most of the Eastern US and Southern Ontario and Quebec experience reductions in PM$_{2.5}$. Another region with reductions greater than 1 $\mu g/m^3$ occurs in the US eastern seaboard corridor. Significantly, these regions do not coincide with the regions of largest NH$_3$ emission reductions (cf. Fig. 4). The San Joaquin valley of central California also shows seasonal average PM$_{2.5}$ reductions greater than 0.5 $\mu g m^{-3}$ and locally as large as 2 $\mu g m^{-3}$. The Vancouver to Seattle region has PM$_{2.5}$ reductions up to 1.5 $\mu g m^{-3}$ in the summer; this effect is highly seasonal however, with a reduction of only 0.25 $\mu g m^{-3}$ in the winter and intermediate reductions in the transition seasons. The Alberta-Saskatchewan area of Western Canada has smaller reductions in PM$_{2.5}$, on the order of 0.25 to 0.5 $\mu g m^3$, with the greatest reductions in the spring and fall.

The episodic nature of the changes in PM$_{2.5}$ mass resulting from ammonia emissions reductions has been examined by constructing annual frequency distributions of hourly mass difference and hourly percent mass difference between the base case and the 30%-reduction scenario. Hourly model values of the changes in PM$_{2.5}$ were first extracted for those model grid cells containing the CAPMoN and CASTNET monitoring stations shown in Fig. 6 and were then used to construct annual frequency distributions of hourly mass change (Fig. 7) and percent mass change (Fig. 8). In both Figs. 7 and 8, the sites are arranged roughly from the west of the continent on the left to the east of the continent on the right, and positive values indicate reductions relative to the base case. Both figures show that median hourly PM$_{2.5}$ reductions are usually less than 1 $\mu g m^{-3}$, or on the order of 5% of the PM$_{2.5}$ mass at any given location, while the reductions during episodes (95 percentiles) may be considerably higher. For the Canadian sites (CAPMoN), the largest median mass reductions (Fig. 7a) are at Abbotsford Airport (0.55 $\mu g m^{-3}$) located in an agricultural area to the east of Vancouver, British Columbia, and Simcoe, Ontario, located in an agricultural area to the east of the De-
troit/Windsor conurbation (0.50 µg m⁻³). The median values at all sites are much lower than the 95-percentile limits (e.g., Abbotsford Airport, 95 percentile value of 3.3 µg m⁻³; Simcoe, 3.2 µg m⁻³). Median percent reductions (Fig. 7b) are more centered in the frequency distributions, although the upper ends of the range are still considerably higher than the medians (e.g., Abbotsford Airport median and 95-percentile values of 3.8% and 17%; Simcoe 8.5% and 23%, respectively). A similar pattern may be observed at US sites (CASTNET), with median and 95 mass percentile values in Indiana and Ohio reaching 0.69 and 3.1 µg m⁻³, and 0.63 and 3.2 µg m⁻³, respectively, and corresponding percent reduction median and 95-percentile values of 8% and 22%, and 7.8% and 22%, respectively. The mass reduction distributions (Figs. 7a, 8a) suggest that the impact of ammonia controls on PM₂.₅ mass will be episodic, with mass reductions during periods of elevated PM₂.₅ levels being as much as 4 to 6 times greater than the median mass reduction. The percent mass reduction distributions (Figs. 7b, 8b) show that median percent mass decreases of 0 to 8% are predicted at the network locations in both countries, with 95-percentile values of up to 22%. However, Figs. 7 and 8 also show that NH₃ emission decreases can also lead to lower-magnitude episodic increases of PM₂.₅ mass.

4.2.3 PM₂.₅ chemistry

The manner in which the reductions in NH₃ emissions create the PM₂.₅ mass reductions described above can be examined by considering the changes in the metrics of Table 1.

The predicted change in particle neutralization ratio resulting from a 30% reduction in NH₃ emissions relative to the base case is shown in Fig. 9. Positive regions indicate areas where the neutralization ratio has decreased (i.e., the particles have become more acidic) compared to the base case, and negative regions indicate areas where the neutralization ratio has increased (i.e., the particles have become less acidic). The main NH₃ emitting regions in the US midwest display relatively little change in particle
charge balance; these regions are NH₃-saturated, so there is little impact on particle charge balance in spite of the predicted concurrent decreases in ambient NH₃ concentrations (cf. Fig. 4). Substantial decreases in the particle neutralization ratio (i.e., increases in particle acidity and changes in particle composition) do occur downwind and on the fringes of the NH₃ emissions regions.

The predicted change in the total ammonium to sulphate ratio resulting from the 30% change in NH₃ emissions is shown in Fig. 10. The total ammonium to sulphate ratio describes the chemical regime, and Fig. 10 shows that the reduction in emitted NH₃ has resulted in a more acidic chemical regime over the NH₃ source regions, with relatively little change outside of those source regions. Figures 9 and 10 suggest that the composition of particles formed over the NH₃ source regions will change. For example, given the minor change in neutralization ratio in Southern Minnesota (Fig. 9), the drop in total ammonium to sulphate in these regions (Fig. 10) suggests that the particles may have more acidic components (ammonium bisulphate, etc.) over the source regions. That is, these two figures taken together suggest that particle nitrate is the dominant means of transport of NH₃ from the source regions to regions downwind; the mass of particle sulphate in the NH₃ source regions is invariant, and the lack of change in the neutralization ratio there shows that reductions in particle ammonium are being accompanied by reductions in particle nitrate. The downwind impact of the NH₃ emissions is therefore due in part to particle ammonium nitrate transport.

A strong seasonal variation in changes to chemical regime can also be seen in Fig. 10, particularly over the Canadian Prairies. Spring and fall have the greatest increase in particle regime acidity, while summer and winter have smaller changes.

The change in the mass fraction of NH₃ (i.e., \([\text{NH}_3(g)] \times [\text{NH}_3(g)] + [\text{PM}_{2.5}-\text{NH}_4]\) on a mass basis) is shown in Fig. 11. The differences are positive over most of the domain and over all seasons, showing that a greater proportion of the remaining (ammonia + ammonium) mass resides in the particle phase instead of the gas phase following a reduction in NH₃ emissions. The effect is strongest in the summer and weakest in the winter. The figure suggests that reductions in NH₃ gas emissions will result in a
nonlinear reduction in NH$_3$ gas concentrations due to chemistry: a shift in chemical regime will reduce the relative amount of NH$_3$ that remains in the gas-phase.

The ratio of the sum of PM$_{2.5}$ nitrate and ammonium mass to total PM$_{2.5}$ mass is shown in Fig. 12. This figure shows that the relative amount of PM$_{2.5}$ composed of these NH$_3$-sensitive species has decreased in the 30% reduction scenario. The figure is also of interest in that it confirms ammonium nitrate as the means of long-range transport of NH$_3$ mass. Comparing the summer (upper left) panels of Figs. 4, 5 and 12, it can be seen that:

1. The largest ammonia source region is located in Southern Minnesota and Northern Iowa (Fig. 4), while the largest particle mass reductions occur further to the east, north of the Ohio River (Fig. 5).

2. The region of greatest particle ammonium and nitrate reduction (Fig. 12) occurs over the state of Illinois i.e., between the ammonia source region and the region of greatest particle mass reduction.

The change in the fraction of directly NH$_3$-sensitive particle mass has a strong seasonal variation, with the greatest impact in the winter (lower left panel, Fig. 12). This is consistent with the strong dependence of particle nitrate formation on temperature, with colder temperatures resulting in a greater proportion of ammonia and nitric acid gas being converted to particle ammonium nitrate, ammonium, and nitrate ions.

4.2.4 Total deposition

AURAMS calculates the wet and dry deposition of various species to the Earth’s surface as moles m$^{-2}$ h$^{-1}$. The hourly wet and dry deposition fields have been added together and summed to seasonal mass totals (kg/ha/season) for the following analysis. As before, scenario values are then subtracted from the base case to determine the impact of the reduced NH$_3$ emissions.
The predicted change in total deposition of all forms of sulphur (SO$_2$+H$_2$SO$_4$+p-SO$_4$ dry deposition and wet deposition) is depicted in Fig. 13. The 30% reduction in the emissions of NH$_3$ has resulted in decreases in sulphur total deposition (red) in many regions in both Canada and the US. Increases in sulphur deposition are also present, in the colder seasons (SE US, Atlantic provinces). It should be noted that these predicted changes to the sulphur deposition associated with ammonia emissions reductions, while significant, are relatively small in magnitude relative to the total sulphur deposited: on the order of 1% of the total sulphur total deposition.

The predicted changes in sulphur total deposition are the result of the following NH$_3$ emissions-reduction-induced changes in the state of atmospheric sulphur:

1. A reduction in NH$_3$ reduces the capacity of cloud water and rain to absorb SO$_2$, via the net equilibrium: \( \text{NH}_3(g) + \text{SO}_2(g) + \text{H}_2\text{O} \rightleftharpoons \text{NH}_4^+(aq) + \text{HSO}_3^-(aq) \).

The concentration of the ammonium ion decreases, hence less SO$_2(g)$ can enter the aqueous phase as HSO$_3^-(aq)$ in the absence of the buffering provided by NH$_3$. The reduction of the sulphur content in cloud water and rain results in less sulphur being removed by wet deposition.

1. A corollary to (1) is that less sulphur is removed in precipitation. The sulphur, which remains in the form of SO$_2(g)$, will therefore be transported longer distances due to decreased rainout/washout. The increases in sulphur deposition that takes place in the colder seasons in Fig. 13 (eastern seaboard of US, Atlantic Ocean) results from the transport and subsequent deposition of SO$_2$ to greater downwind distances.

2. Another corollary to (1) is that reductions in NH$_3$ emissions will reduce the amount of ammonium ion in the cloud droplets and water, and hence will reduce the amount of nitrate taken up in cloud water and rain, and thus the amount of nitrate removed by wet deposition. Total nitrogen deposition therefore decreases due to decreases in both reduced and oxidized nitrogen deposition, as discussed below.
3. Reductions in NH$_3$ emissions may also cause a reduction in the size of ambient particles, since less NH$_3$ leads to less $p$-NH$_4$ and $p$-NO$_3$ in the particle phase, hence smaller particles, which have a smaller deposition velocity.

The change in total deposition of all phases of nitrogen related to ammonia chemistry (sum of \{NH$_4^+(aq)$+$NH_3(g)$+$PM_{2.5}$NH$_4$+$NO_3^-(aq)$+$HNO_3(g)$+$PM_{2.5}$ NO$_3$\}) is shown in Fig. 14. The reduction in NH$_3$ emissions by 30% has resulted in substantial reductions in deposited nitrogen (similar in magnitude to the total deposited nitrogen in many locations). The greatest spatial extent of nitrogen deposition reduction occurs in the spring (lower right panel), when NH$_3$ emissions are highest, and the smallest change occurs in the winter (lower left panel), when NH$_3$ emissions are lowest. The location of the largest reductions in nitrogen deposition occurs over the NH$_3$ emitting areas (compare Figs. 4 and 14). Less than 10% of the total change in deposited nitrogen is associated with the various forms of nitrate, and is instead dominated by the ammonium components (not shown). The change in total ammonia/um deposition is itself dominated by $p$-NH$_4$ wet deposition (approximately 5/6 of the total) and NH$_3$ dry deposition (remaining 1/6), with changes to $p$-NH$_4$ dry deposition being relatively insignificant for the nitrogen budget.

The main results of the deposition analysis for a 30% reduction in agricultural NH$_3$ emissions are thus:

1. Sulphur deposition close to the sources of sulphur decreases slightly, due to a reduction in SO$_2(g)$ uptake in clouds and possibly to a decrease in particle sulphate deposition with decreasing particle size. Sulphur deposition further downwind of the sources may increase as a consequence, depending on the season.

2. Nitrogen deposition decreases significantly, driven largely by decreases in $p$-NH$_4$ wet deposition but also by near-source decreases in $p$-NO$_3$ wet deposition.

3. Hydrogen ion wet deposition increases (not shown). The increase in hydrogen ion deposition is spatially matched with the decreases in nitrogen deposition and is greatest over the regions of NH$_3$ emissions.
4.2.5 Annual critical load exceedances for sensitive ecosystems

Annual critical-load exceedance fields for Canada were calculated for the base case and for the 30% emissions reduction scenario in two ways: for (a) sulphur (S) total deposition and for (b) sulphur+nitrogen (S+N) total deposition.

Annual critical-load exceedances for sulphur were not significantly changed between the base case and the 30% NH3 emissions reduction scenario (annual critical-load exceedances did decrease for the NH3 emissions reduction scenario, but significant reductions only occurred at two model gridpoints: not shown). This indicates that the impact of NH3 emissions reductions on sulphur acidification of ecosystems is expected to be small, in accord with the relatively small changes in total sulphur deposition (Fig. 13).

Figure 15a shows the predicted S+N annual critical-load exceedances for the base case. Figure 15b shows the corresponding reductions in S+N annual critical-load exceedances in many parts of Canada that are predicted to result from a 30% reduction in agricultural NH3 emissions. These substantial decreases in S+N annual critical-load exceedances are in contrast to the small decreases in S annual critical-load exceedances. The implication of this finding is that, if the ecosystem’s ability to absorb nitrogen is compromised at these locations in the future, then the deposition of atmospheric nitrogen resulting from NH3 emissions will contribute to a degradation of these ecosystems.

4.2.6 A conceptual model

The above analysis of AURAMS simulations may be used to provide a simple conceptual model to describe the effect of reductions in NH3 emissions on atmospheric chemistry. The following diagram (Fig. 16) depicts the processes, on a hypothetical transect with NH3 emissions on the left, a source of SO2 and NOx in the centre, and a receptor region downwind on the right. This is similar to the situation on the eastern half of the North American continent, with the US midwest NH3 source on the left, the Ohio Valley and Great Lakes regions in the centre, and the Atlantic provinces and New...
England states on the right. The prevailing wind blows from left to right in this diagram. The upper half of the diagram shows the system in the absence of NH₃ emission controls, the lower half the system including controls in NH₃ emissions. The size of the font for the different chemical variables is intended as a visual guide to the relative magnitude of those species, without (top) and with (bottom) NH₃ emissions controls.

In the absence of emissions controls (Fig. 16, top), excess NH₃ in the source region at left creates particle ammonium nitrate, in addition to particle ammonium sulphate. Winds blowing to the right then transport the particles and NH₃ gas. En route, the NH₃ gas is depleted due to wet and dry deposition, as are the particles by wet deposition. Nevertheless, significant amounts of particle ammonium and nitrate reach the SO₂ and NOₓ emissions source region in the centre of the Figure. The addition of fresh SO₂ and nitric acid to the system cause the particles to locally become more acidic, with some transfer of the transported ammonium from particle nitrate to particle sulphate possible due to inorganic thermodynamics. With subsequent transport further downwind, the particles are deposited; relatively little SO₂ reaches far downwind locations due to NH₃-enhanced aqueous-phase conversion to sulphate closer to the source regions.

With the presence of NH₃ controls (bottom), less ammonium nitrate is created in the Midwest source region, and hence less is available for transport. Smaller amounts of ammonium reach the central SO₂ and NOₓ source region; this reduces the rate of further particle formation and allows more subsequent downwind transport and deposition of SO₂. Downwind nitrogen and sulphur deposition is thus increased due to increased transport of these precursor species.

A neutral charge balance ratio (pink text in Fig. 16) is maintained over the NH₃ source region at the left regardless of the scenario; since this region remains NH₃-saturated, reductions in p-NH₄ here are matched by reductions in p-NO₃ in the denominator. In the SO₂ and NOₓ source region in the centre, however, the upwind reductions in NH₃ emissions result in a net decrease in charge balance and an increase in particle acidity that is then maintained during further downwind transport.
5 Uncertainty analysis of the 50% Canadian beef cattle emissions reduction scenario

The main intent of this scenario was to serve as an uncertainty benchmark for the previous scenario. The beef-cattle NH$_3$ emissions factors used in constructing the updated 2002 Canadian agricultural NH$_3$ emissions inventory were considered to be the most uncertain. Their uncertainty was estimated to be as high as a factor of two; the 50% reduction considered in this scenario thus represents the lower range of the uncertainty envelope.

The model response for PM$_{2.5}$ mass for this emissions scenario, relative to the base case, is shown below in Fig. 17. This figure uses the same colour scale as Fig. 5 to allow a direct comparison: the reduction of beef-cattle emissions by 50% has about the same impact in Alberta (in the area of concentrated beef-cattle operations) as the 30% overall reduction noted above. The only other region where a change is noticeable is in SW Ontario in the fall (upper right panel). There, the effect on PM$_{2.5}$ emissions is much smaller than that of the 30% overall emissions reduction scenario due to the smaller local relative contribution of beef-cattle NH$_3$ emissions to total NH$_3$ emissions.

One important conclusion from this analysis is that for Alberta, the range of uncertainty in model predictions associated with the beef-cattle emissions factors may be as large as the impacts from an across-the-board 30% reduction of NH$_3$ emissions. While the best available information was used to compile the new NH$_3$ emissions inventory, improvements in the beef-cattle subsector of the inventory are recommended for future work.

The other aspect to this sensitivity analysis is to demonstrate the extent to which sector-specific scenario simulations are possible with the updated 2002 NH$_3$ Canadian emissions inventory. An emissions reduction strategy may be “tailored” for the dominant emissions sources in a given region; very specific emissions reduction strategies may be tested in the future.
6 Conclusions and recommendations for future research

A unified regional air-quality modelling system (AURAMS) was used to investigate the effects of reductions in NH₃ emissions on regional air quality, especially PM. Three simulations of one-year duration were performed for a North American domain for different sets of NH₃ emissions. The simulation for a 30% continent-wide reduction in agricultural ammonia emissions predicted decreases in median hourly PM$_{2.5}$ mass of $\leq 1$ µg m$^{-3}$. However, the atmospheric response to these emission reductions has marked seasonal variations, and on even shorter time scales the impacts of the emissions reductions are highly episodic: for example, 95-percentile hourly PM$_{2.5}$ mass decreases may be a factor of six larger than the median values.

A key feature of the above simulations is the manner in which continental-scale long-range transport may play a role in defining the impacts of reductions in NH₃ emissions. The emissions reductions affect local NH₃ gas concentrations, but the largest impacts of these reductions may take place significantly downwind of the NH₃ emissions source in NH₃-limited areas. The interaction between transport and chemistry is complex: NH₃ mass is transported from the source regions as aqueous and particle ammonium, and emissions of other particle precursors play a significant role in the subsequent chemistry. Reductions in NH₃ emissions result in a small but significant decrease in the amount of SO$_2$ gas converted to sulphate. Reductions in aqueous buffering capacity and decreases in particle size decrease local sulphur deposition in favour of SO$_2$ deposition further downwind; a by-product of the NH₃ emissions reduction is to increase the overall transport distance of emitted atmospheric sulphur. Ammonia emissions reductions result in a significant decrease in total ammonia deposition and a smaller decrease in nitrate deposition, in regions of high ammonia emissions.

The frequency-distribution figures of hourly model output for PM$_{2.5}$ mass show that the impact of NH₃ emissions reductions is highly episodic in nature. In both high and low resolution model runs, 95 percentile values of the differences between base case and scenarios are often much larger (up to a factor of 6) than the median differences.
of the distribution. This is in accord with the known chemistry of $p$-$\text{NH}_4^+$ formation, specifically ammonium nitrate. Relatively small changes in local temperature, humidity, and precursor-gas concentrations can give rise to rapid particle formation and/or loss conditions (e.g., Yu et al., 2005). Decreases in NH$_3$ emissions may have a modest or low impact on PM$_{2.5}$ levels in median or average conditions, but a much larger impact when PM$_{2.5}$ levels are high.

Reductions in NH$_3$ emissions may result in decreases in acid deposition and in exceedance of S+N critical loads for sensitive Canadian ecosystems. The predicted small changes to sulphur deposition, on the other hand, have little impact on sulphur critical-load exceedances, implying that NH$_3$ emissions reductions at the current time will not reduce the sulphur acidification of sensitive ecosystems. However, if the ability of these ecosystems to absorb nitrogen becomes saturated, the role of NH$_3$ on the exceedance of critical loads does become substantial. Ammonia emissions reductions hence may eventually be required to reduce acidification of Canadian ecosystems.

The important linkages between transport and chemistry when NH$_3$ reductions are considered suggests that cross-border transport is an important factor with regards to predicting the outcomes of NH$_3$ reduction strategies. Ammonia and PM concentrations may be affected considerably downwind, and sulphur and nitrogen transport distances from sources of sulphur and nitrogen and associated deposition patterns are changed. Future scenario runs should focus on trans-boundary transport.

The beef-cattle emissions scenario shows that the uncertainty associated with the updated 2002 Canadian agricultural NH$_3$ emissions inventories for this source sub-sector is large. Future work on emissions inventories should attempt to reduce this uncertainty. This scenario also serves to show the potential for NH$_3$ emissions reduction scenarios that assess the impacts of agricultural-subsector-specific changes in emitting practices. Scenarios examining the impacts of NH$_3$ management practice changes should therefore be considered in future work.

A potentially significant source of uncertainty in all of the model results is the role of coarse-mode chemistry in the real atmosphere. Coarse-mode particle chemistry may
reduce the impact of ammonia emissions reductions by competing with the fine mode for the available nitric acid, as well as providing sites for condensation of nitric and sulphuric acid during intense dust storm events. Inclusion of coarse-mode chemistry is recommended for future model simulations of ammonia emission scenarios.

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Table 1. Metrics for chemical evaluation of model responses to NH$_3$ emission changes.

<table>
<thead>
<tr>
<th>Metric</th>
<th>Formula</th>
<th>Significance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Particle Neutralization Ratio</td>
<td>$(\rho-\text{NH}_4)/[2(\rho-\text{SO}_4)+(\rho-\text{NO}_3)]$</td>
<td>Ratio of total ammonium charge to the net sulphate and nitrate charge (each particle species variable is the sum over all particle sizes). Values of 1 indicate that the particles are NH$_3$-saturated, so that significant NH$_3$ reductions may be required to reduce particulate mass. Regions with values less than unity are more NH$_3$-limited; smaller reductions in NH$_3$ may result in significant reductions in particulate mass.</td>
</tr>
<tr>
<td>Total ammonia to sulphate mole ratio</td>
<td>$(\text{NH}_3(g)+(\rho-\text{NH}_4))/(\rho-\text{SO}_4)$</td>
<td>Mole ratio of ammonia gas + particle ammonia to particle sulphate. This defines the chemical regime: values less than unity denote acidic conditions (e.g., ammonium bisulphate, sulphuric acid present in the particles); values between 1.0 and 2.0 denote intermediate acidity (ammonium bisulphate, letovicite, ammonium sulphate present), and values greater than 2 indicate less acidic particles (ammonium sulphate, ammonium nitrate may be present in the particles). Note that a decrease in the value of the ratio does not necessarily imply a significant change in the particle composition, if the initial and final values of the ratio are both high.</td>
</tr>
<tr>
<td>Gas-phase fraction ammonia mass</td>
<td>$(\text{NH}<em>3)/(\text{PM}</em>{2.5}-\text{NH}_4+\text{NH}_3)$</td>
<td>Relative mass of NH$_3$ in the gas phase to total ammonia + ammonium mass. Changes in this parameter indicate a change in the mass partitioning of ambient NH$_3$ in response to changes in NH$_3$ emissions.</td>
</tr>
<tr>
<td>PM$<em>{2.5}$ Ammonium + Nitrate to total PM$</em>{2.5}$ mass ratio</td>
<td>$(\text{PM}_{2.5}-\text{NH}<em>4+\text{PM}</em>{2.5}-\text{NO}<em>3)/(\text{PM}</em>{2.5})$</td>
<td>Fraction of fine particle mass that is directly ammonia-sensitive. A diagnostic of the direct impact of emissions reductions.</td>
</tr>
</tbody>
</table>
Table 2. Statistical measures of model performance. \( N \) is the number of paired observed-model values, \( \bar{O} \) is the mean observed value, \( \bar{M} \) is the mean model value.

<table>
<thead>
<tr>
<th>Statistical Measure</th>
<th>Description</th>
<th>Formula</th>
</tr>
</thead>
<tbody>
<tr>
<td>( R )</td>
<td>Pearson Correlation Coefficient</td>
<td>( R = \frac{\frac{N}{\sum_{i=1}^{N}(O_i \times M_i)} - \frac{1}{N} \sum_{i=1}^{N} M_i \sum_{i=1}^{N} O_i}{\sqrt{\frac{1}{N} \sum_{i=1}^{N} (M_i \times M_i) - \frac{1}{N} \sum_{i=1}^{N} M_i \times \frac{1}{N} \sum_{i=1}^{N} O_i}} )</td>
</tr>
<tr>
<td>( b )</td>
<td>Slope of observations vs. model best-fit line</td>
<td>( b = \frac{\sum_{i=1}^{N} [(O_i - \bar{O})(M_i - \bar{M})]}{\sum_{i=1}^{N} [(O_i - \bar{O})^2]} )</td>
</tr>
<tr>
<td>( a )</td>
<td>Intercept of observations vs. model best-fit line</td>
<td>( a = \bar{M} - b \times \bar{O} )</td>
</tr>
<tr>
<td>MB</td>
<td>Mean bias</td>
<td>( MB = \frac{1}{N} \sum_{i=1}^{N} (M_i - O_i) )</td>
</tr>
<tr>
<td>RMSE</td>
<td>Root Mean Square Error</td>
<td>( RMSE = \sqrt{\frac{1}{N} \sum_{i=1}^{N} (M_i - O_i)^2} )</td>
</tr>
<tr>
<td>NMB</td>
<td>Normalized Mean Bias</td>
<td>( NMB = \frac{\sum_{i=1}^{N} (M_i - O_i)}{\sum_{i=1}^{N} O_i} \times 100 )</td>
</tr>
<tr>
<td>NME</td>
<td>Normalized Mean Error</td>
<td>( NME = \frac{\sum_{i=1}^{N}</td>
</tr>
</tbody>
</table>
### Table 3. Annual statistics for selected AURAMS gas- and particle-phase species. Statistical metrics are defined in Table 2.

<table>
<thead>
<tr>
<th>Statistic</th>
<th>SO₂ (ppbv)</th>
<th>HNO₃ (ppbv)</th>
<th>PM₂.₅ (µg m⁻³, STP)</th>
<th>PM₂.₅-SO₄ (µg m⁻³, STP)</th>
<th>PM₂.₅-NO₃ (µg m⁻³, STP)</th>
<th>PM₂.₅-NH₄ (µg m⁻³, STP)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Networks</td>
<td>a,d,e,g</td>
<td>d,e</td>
<td>a,b,c,f,g,h</td>
<td>c,f,h</td>
<td>c,f,h</td>
<td>c,h</td>
</tr>
<tr>
<td>N</td>
<td>451</td>
<td>86</td>
<td>845</td>
<td>265</td>
<td>254</td>
<td>141</td>
</tr>
<tr>
<td>O</td>
<td>3.32</td>
<td>0.53</td>
<td>11.33</td>
<td>2.77</td>
<td>1.28</td>
<td>1.53</td>
</tr>
<tr>
<td>M</td>
<td>3.55</td>
<td>0.66</td>
<td>7.87</td>
<td>2.26</td>
<td>1.05</td>
<td>1.25</td>
</tr>
<tr>
<td>a</td>
<td>1.01</td>
<td>0.08</td>
<td>−0.74</td>
<td>−0.64</td>
<td>0.12</td>
<td>0.23</td>
</tr>
<tr>
<td>b</td>
<td>0.77</td>
<td>1.10</td>
<td>0.76</td>
<td>1.04</td>
<td>0.73</td>
<td>0.66</td>
</tr>
<tr>
<td>R</td>
<td>0.56</td>
<td>0.81</td>
<td>0.65</td>
<td>0.92</td>
<td>0.77</td>
<td>0.76</td>
</tr>
<tr>
<td>MB</td>
<td>0.23</td>
<td>0.13</td>
<td>−3.46</td>
<td>−0.51</td>
<td>−0.23</td>
<td>−0.29</td>
</tr>
<tr>
<td>RMSE</td>
<td>2.88</td>
<td>0.28</td>
<td>5.00</td>
<td>0.95</td>
<td>1.01</td>
<td>0.54</td>
</tr>
<tr>
<td>NMB (%)</td>
<td>7.0</td>
<td>25.2</td>
<td>−30.5</td>
<td>−18.5</td>
<td>−17.9</td>
<td>−18.7</td>
</tr>
<tr>
<td>NME (%)</td>
<td>51.7</td>
<td>38.3</td>
<td>36.8</td>
<td>27.3</td>
<td>43.5</td>
<td>27.2</td>
</tr>
</tbody>
</table>

Networks: a: AQS-continuous; b: AQS-filter; c: AQS-STN; d: CAPMoN; e: CASTNet; f: IMPROVE; g: NAPS-continuous; h: NAPS-filter.
Table 4. Annual statistics for several AURAMS wet deposited species. Measurements were obtained from five Canadian precipitation-chemistry networks (CAPMoN, BCPCSN, NBPMN, PQMPA, REPQ) and one US network (NADP).

<table>
<thead>
<tr>
<th>Statistic</th>
<th>SO$_4^{2-}$ conc. in precip. (mg SO$_4$/L)</th>
<th>NO$_3^-$ conc. in precip. (mg NO$_3$/L)</th>
<th>NH$_4^+$ conc. in precip. (mg NH$_4$/L)</th>
<th>SO$_4^{2-}$ wet dep. (kg SO$_4$/ha/y)</th>
<th>NO$_3^-$ wet dep. (kg NO$_3$/ha/y)</th>
<th>NH$_4^+$ wet dep. (kg NH$_4$/ha/y)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$N$</td>
<td>277</td>
<td>270</td>
<td>271</td>
<td>277</td>
<td>270</td>
<td>271</td>
</tr>
<tr>
<td>$\bar{O}$</td>
<td>1.08</td>
<td>1.11</td>
<td>0.31</td>
<td>10.1</td>
<td>9.39</td>
<td>2.39</td>
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<tr>
<td>$\bar{M}$</td>
<td>0.96</td>
<td>0.94</td>
<td>0.21</td>
<td>9.54</td>
<td>8.30</td>
<td>1.81</td>
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<tr>
<td>$a$</td>
<td>$-0.03$</td>
<td>0.202</td>
<td>0.021</td>
<td>0.449</td>
<td>1.46</td>
<td>0.11</td>
</tr>
<tr>
<td>$b$</td>
<td>0.91</td>
<td>0.67</td>
<td>0.60</td>
<td>0.90</td>
<td>0.73</td>
<td>0.71</td>
</tr>
<tr>
<td>$R$</td>
<td>0.81</td>
<td>0.61</td>
<td>0.76</td>
<td>0.84</td>
<td>0.71</td>
<td>0.78</td>
</tr>
<tr>
<td>MB</td>
<td>$-0.12$</td>
<td>$-0.17$</td>
<td>$-0.10$</td>
<td>$-0.58$</td>
<td>$-1.09$</td>
<td>$-0.58$</td>
</tr>
<tr>
<td>RMSE</td>
<td>0.37</td>
<td>0.50</td>
<td>0.16</td>
<td>3.94</td>
<td>4.24</td>
<td>1.07</td>
</tr>
<tr>
<td>NMB (%)</td>
<td>$-11.4$</td>
<td>$-15.0$</td>
<td>$-33.1$</td>
<td>$-5.8$</td>
<td>$-11.6$</td>
<td>$-24.1$</td>
</tr>
<tr>
<td>NME (%)</td>
<td>24.8</td>
<td>33.4</td>
<td>36.2</td>
<td>28.3</td>
<td>33.7</td>
<td>33.9</td>
</tr>
</tbody>
</table>
Fig. 1. AURAMS CTM North American 150×106 42-km domain.
Fig. 2. Ammonia emissions (kilotonnes) in each season. Upper left: June-July-August; upper right: September-October-November; lower left: December-January-February; lower left: March-April-May.
Fig. 3. Seasonal total NH$_3$ to sulphate mole ratio for base-case simulation: summer – upper left; fall – upper right; winter – lower left; spring – lower right.
Fig. 4. Seasonal average change in NH$_3$ concentration, ppbv, associated with a 30% reduction in agricultural NH$_3$ emissions. Panels arranged as in Fig. 2.
Fig. 5. Seasonal average change in PM$_{2.5}$ mass, $\mu$g m$^{-3}$, associated with a 30% reduction in agricultural NH$_3$ emissions. Panels arranged as in Fig. 2.
Fig. 6. Locations of sites used in analysis of episodic nature of ammonia emissions reductions: (a) Canadian CAPMoN sites; (b) US CASTNet sites.
Fig. 7. AURAMS-predicted annual frequency distributions of the decrease in hourly PM$_{2.5}$ concentrations associated with a 30% decrease in NH$_3$ emissions at CAPMoN stations (numbered sites from Fig. 6a): (a) expressed in mass units; (b) expressed as percentage difference relative to the base case. Median: solid horizontal bar; 5 and 95 percentiles: limits of red vertical bar; 2 and 98 percentiles: thin horizontal bars.
Fig. 8. Same as Fig. 6 but for CASTNet station sites shown in Fig. 5b.
Fig. 9. Change in particle neutralization ratio due to 30% reduction in NH$_3$ emissions. Positive regions indicate increased particle acidity relative to the base case. Panels arranged as in Fig. 2.
Fig. 10. Change in total ammonia to sulphate mole ratio due to 30% reduction in NH$_3$ emissions. Positive regions indicate decreases in the ratio (more acidic chemical regimes) relative to the base case. Panels arranged as in Fig. 2.
Fig. 11. Change in gas-phase NH$_3$ mass fraction due to 30% reduction in NH$_3$ emissions. Positive regions indicate decreases in mass fraction (i.e., proportionately more ammonium in the particle phase) relative to the base case. Panels arranged as in Fig. 2.
Fig. 12. Change in fraction of ammonium + nitrate mass in PM$_{2.5}$ relative to the total PM$_{2.5}$ mass. Positive regions indicate decreases in ammonium and nitrate mass fraction (i.e., particles composed of proportionately less ammonium and nitrate) relative to the base case. Panels arranged as in Fig. 2.
Fig. 13. Change in total – sulphur total deposition (kg S/ha/season) due to 30% reduction in NH₃ emissions. Positive values (green to red colours) indicate decreases in sulphur deposition resulting from decreasing NH₃ emissions; negative values (blue) indicate increases in sulphur deposition resulting from decreasing NH₃ emissions. Panels arranged as in Fig. 2.
Fig. 14. Same as Fig. 13 but for total-nitrogen total deposition (kg N/ha/season).
Fig. 15. AURAMS-predicted Canadian (S+N) critical-load exceedances for 2002: (top) NAESI base case; and (bottom) base case – 30% NH$_3$ emissions reduction scenario. Gray areas in the top panel indicate areas for which critical-load values were not available. Positive values in the bottom panel indicate areas where the critical load exceedance has decreased in response to decreasing NH$_3$ emissions.
Fig. 16. Conceptual model of NH₃ emissions, reaction and transport: (top) without NH₃ emissions controls; (bottom) with NH₃ emissions controls.
Fig. 17. Seasonal average change in PM$_{2.5}$ mass ($\mu$g m$^{-3}$) associated with a 50% reduction in Canadian beef-cattle emissions. Panels arranged as in Fig. 2.