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# Size distribution of alkyl amines in continental particulate matter and their online detection in the gas and particle phase

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be appropriate.

An ion chromatographic method is described for the quantification of the simple alkyl amines: methylamine (MA), dimethylamine (DMA), trimethylamine (TMA), ethylamine (EA), diethylamine (DEA) and triethylamine (TEA), in the ambient atmosphere. Limits of detection  $(3\sigma)$  are in the tens of pmol range for all of these amines, and good resolution is achieved for all compounds except for TMA and DEA. The technique was applied to the analysis of time-integrated samples collected using a micro-orifice uniform deposition impactor (MOUDI) with ten stages for size resolution of particles with aerodynamic diameters between 56 nm and 18 um. In eight samples from urban and rural continental airmasses, the mass loading of amines consistently maximized on the stage corresponding to particles with aerodynamic diameters between 320 and 560 nm. The molar ratio of amines to ammonium (R<sub>3</sub>NH<sup>+</sup>/NH<sub>4</sub>) in fine aerosol ranged between 0.005 and 0.2, and maximized for the smallest particle sizes. The sizedependence of the R<sub>3</sub>NH<sup>+</sup>/NH<sub>4</sub><sup>+</sup> ratio indicates differences in the relative importance of the processes leading to the incorporation of amines and ammonia into secondary particles. The technique was also used to make simultaneous hourly online measurements of amines in the gas phase and in fine particulate matter using an Ambient Ion Monitor Ion Chromatograph (AIM-IC). During a ten day campaign in downtown Toronto,

DMA, TMA+DEA, and TEA were observed to range from below detection limit to 2.7 ppt in the gas phase. In the particle phase, MAH<sup>+</sup> and TMAH<sup>+</sup>+DEAH<sup>+</sup> were observed to range from below detection limit up to 15 ng m<sup>-3</sup>. The presence of detectable levels

of amines in the particle phase corresponded to periods with higher relative humid-

ity and higher mass loadings of nitrate. While the hourly measurements made using

the AIM-IC provide data that can be used the evaluate the application of gas-particle

partitioning models to amines, the strong size-dependence of the R<sub>3</sub>NH<sup>+</sup>/NH<sub>4</sub><sup>+</sup> ratio indicates that using bulk measurements and an assumption of internal mixing may not

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Recent ambient measurements of amines in the gas and particle phase have prompted research into their role in the formation and growth of atmospheric particulate matter. Amine emissions have been detected from animal husbandry operations (Mosier et al., 1973; Schade and Crutzen, 1995), decomposition of organic matter on land and in the oceans (Van Neste et al., 1987; Gibb et al., 1999), and from biomass burning and the incineration of waste (Finlayson-Pitts and Pitts, 2000; Chang et al., 2003). In the future, increased emissions may arise from the proposed use of amines as CO<sub>2</sub> scrubbing agents in industrial applications (Veltman et al., 2010). The amines emitted in the greatest quantity to the atmosphere are thought to be the simple alkyl amines: methylamine (MA), dimethylamine (DMA), trimethylamine (TMA), ethylamine (EA), diethylamine (DEA) and triethylamine (TEA). Collectively, these alkyl amines will be referred to as NR<sub>3</sub> throughout the discussion that follows. The greatest direct source of NR<sub>3</sub> has been suggested to be animal husbandry operations, with global emissions estimated at 0.15 Tg N a<sup>-1</sup>, leading to amine concentrations up to hundreds of ppb near animal housing and waste storage areas (Schade and Crutzen, 1995; Rabaud et al., 2003). Gaseous amine levels in marine and background continental environments are generally on the order of <1-100 ppt (Mopper and Zika, 1987; Grönberg et al., 1992; Gibb et al., 1999; Chang et al., 2003; Akyuz, 2007). Amines are often present concomitantly with NH<sub>3</sub>, typically at mixing ratios 1-3 orders of magnitude smaller (Grönberg et al., 1992; Chang et al., 2003; Huang et al., 2009; Veltman et al., 2010). The primary sinks of the simple alkyl amines are thought to be oxidation by OH and wet deposition via rain scavenging. Lifetimes for NR<sub>3</sub> against oxidation are on the order of a few hours to a day in the presence of atmospherically relevant concentrations of OH, compared to a month for NH<sub>3</sub> (Atkinson et al., 1977a,b, 1997; Pitts et al., 1978). Oxidation by O<sub>3</sub> can also be an appreciable sink for secondary and tertiary amines (lifetimes of hours to days) in a polluted environment (Finlayson-Pitts and Pitts, 2000). Elucidation of the fast loss mechanisms for NR<sub>3</sub> led to a decline in research concerning

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There are several different scenarios in which amines might become associated with atmospheric aerosols. Amines are likely candidates for forming aminium salts with atmospheric acids resulting in the formation of secondary aerosol mass (Pratt et al., 2009). With pKa values ranging from 9.25–10.98 (pKa<sub>NH3</sub>=9.25, Seinfeld and Pandis, 2006) and similar water solubilities to NH<sub>3</sub> (Gibb et al., 1999; Mackay et al., 2004), amines can partition to and neutralize aqueous acidic particles. The partitioning and reactive uptake mechanisms of NH<sub>3</sub> in to particles and aqueous solutions have been well explored (Mozurkewich, 1993; Donaldson, 1999; Dinar et al., 2008) and impacts on aerosol properties continue to be investigated. Amines detected in an aqueous particle or as a salt are likely to be protonated since they are strong bases, and will therefore be denoted with an appended H<sup>+</sup> in the following discussion. The degree to which amines may be scavenged by aqueous aerosols is likely dependent on the effective pH of the solution and the pKa of the amine Reaction (R1). The degree of protonation will have a buffering effect on isolating amine molecules from exchange back to the gas phase (Pankow, 2003):

$$NR_{3(g)} \leftrightarrow NR_{3(aq)} + H_2O_{(l)} \leftrightarrow R_3NH^+_{(aq)} + OH^-_{(aq)}$$
 (R1)

Second, subsequent reaction with acidic solution components could lead to the formation of insoluble salts:

$$R_3NH^+_{(aq)} + A^-_{(aq)} \leftrightarrow R_3NHA_{(s)}$$
(R2)

Where A<sup>-</sup> is a conjugate base in solution and HA is the parent acid. Third, gas phase acid-base reactions to form an insoluble salt:

$$NR_{3(q)} + HA_{(q)} \leftrightarrow R_3NHA_{(s)} \tag{R3}$$

Finally, reactive uptake and displacement of ammonia from pre-existing salts may also occur:

$$NR_{3(g)} + (NH_4)_n(A)_{m(s)} \leftrightarrow NH_{3(g)} + (NH_4)_{n-1}(R_3NH)(A)_{m(s)}$$

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Modeling and laboratory studies indicate that amines can enter the condensed phase by all the routes proposed above in reactions 1–4 (Mmereki et al., 2000; Murphy et al., 2007; Barsanti et al., 2009) and suggest that small chain (C<sub>1</sub>-C<sub>5</sub>) alkyl amines may be important for the initiation of particle formation events and growth of nucleating aerosols. Junninen et al. (2010) used an atmospheric pressure interface time-of-flight mass spectrometer to collect ambient atmospheric ions from the gas phase, including molecular clusters where the formation of inorganic aminium salts might be occurring. Preliminary results showed homologous series of alkyl amines and pyridines in their samples in addition to the typical HNO<sub>3</sub> and H<sub>2</sub>SO<sub>4</sub> acids. These results are indicative of gas phase reaction prior to particle formation between the trace constituents, supporting the hypothesis that amines may promote H<sub>2</sub>SO<sub>4</sub> nucleation through the formation of non-volatile salts. Further support for this from laboratory investigations was made by Berndt et al. (2010) in a series of lab experiments comparing the nucleation rate of H<sub>2</sub>SO<sub>4</sub> as a function of relative humidity (RH) and the presence of NH<sub>3</sub> and a surrogate amine, tert-butylamine. The results showed that both basic species had significant and similar promoting effects on the nucleation rate of H<sub>2</sub>SO<sub>4</sub>. However, the number density of amine was also two orders of magnitude smaller than NH<sub>3</sub>, showing that typical near-source atmospheric concentrations of both gases may result in similar nucleation enhancing effects. Alkyl amines, like ammonia, will form salts (R2, R3) with inorganic and organic acidic species due to their basic nature (Angelino et al., 2001; Murphy et al., 2007; Barsanti et al., 2009). Laboratory studies have shown that in some instances, alkyl amines can displace ammonium from inorganic salts to form aminium salts Reaction (R4) (Murphy et al., 2007; Lloyd et al., 2009; Bzdek et al., 2010). Smoq chamber experiments have also shown that the oxidation of amines can lead to the formation of secondary organic aerosol (SOA) or basic products that can also form salts with atmospheric acids (Murphy et al., 2007). The relative importance of these processes for the incorporation of amines in ambient fine particulate matter is not yet clear.

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Amines have been detected frequently in atmospheric fine and ultrafine particulate matter by a number of online techniques, including: AToF-MS (Angelino et al., 2001; Pastor et al., 2003; Denkenberger et al., 2007), cToF-AMS (Murphy et al., 2007; Sorooshian et al., 2008), APi-TOF-MS (Junninen et al., 2010), TDCIMS (Smith et al., <sub>5</sub> 2008, 2010), and PILS-IC (Murphy et al., 2007; Sorooshian et al., 2008). The R<sub>3</sub>NH<sup>+</sup> contribution of 10-47% of the positive ions in 8-10 nm particles sampled by Smith et al. (2010) indicates that amines may be a significant constituent of newly formed particles. Their frequent presence as detectable fractions of fine mode particles, specifically that which may not be fully neutralized by ammonia, indicates that they may also contribute to condensational growth through reactive uptake (Murphy et al., 2007). A drawback of many mass spectrometric observations is that they cannot provide quantitative mass loadings of specific R<sub>3</sub>NH<sup>+</sup> constituents. For example, AMS response factors for various aminium salts have been reported to range from 5-10 times that of ammonium nitrate depending on the composition of the salt, thereby limiting this method to amine identification only (Silva et al., 2008). Similarly, Murphy et al. (2007) showed that the exchange of ammonium for methylaminium in sulphate particles enhanced the collection efficiency by the cTOF-AMS significantly due to changes in the physical properties of the particles. Simultaneous observations by cTOF-AMS and PILS-IC by Sorooshian et al. (2008) bridged the gap in producing quantitative amine information in addition to detailed knowledge of particle composition. They found instances of EA and DEA in sub-micrometer particles at mass loadings of 50-200 ng m<sup>-3</sup> (~20% of the mass loading of  $NH_4^+$ ), indicating that alkyl amines can contribute significantly to the neutralizing capacity of particles less than 1 µm in diameter. More recently, Huang et al. (2009) described an online derivatization method for the detection of NR<sub>3</sub>/R<sub>3</sub>NH<sup>+</sup>, and reported simultaneous NH<sub>3</sub>/NH<sub>4</sub> and MA/MAH observations near a small animal farm in New York on the order of 10 ppb and 100 ppt, respectively.

The presence of DMAH<sup>+</sup> in nucleation event particles reported by Mäkelä et al. (2001) provided the first indication of a potentially important role that amines might play in particle formation and growth. The presence of DMAH<sup>+</sup> was also reported in

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a small fraction of the total reduced nitrogen in particles, but because many amines are stronger bases than ammonia their presence may significantly impact aerosol composition and properties. An increase in effective basicity of aqueous aerosol has been proposed as a way to reduce the free fraction of protons and therefore significantly alter pH values even in the presence of relatively small R<sub>3</sub>NH<sup>+</sup> concentrations (Angelino et al., 2001; Pratt et al., 2009). Simultaneous measurements of amines and other chemical constituents and physical properties of aerosol can help to understand their impact on the pH of aerosols, which can affect particle chemistry, e.g. SOA formation. To improve our understanding of the mechanisms governing amine exchange between the gas and particle phases, online methods to separately observe and quantify individual amines in both phases with high time resolution are required. The applicability of simple thermodynamic equilibrium algorithms to describe gas-particle partitioning also needs to be evaluated by size-resolved measurements of amine mass loadings in the sub-micrometer range.

the 10 nm fraction for non-event particulate collected, but at much lower mass loadings. Further investigations of the size-distribution of particle phase amines have shown the

presence of DEAH<sup>+</sup> and MAH<sup>+</sup> in the 0.01-1.0 µm range in marine aerosols, their

presence being attributed to biogenic sources metabolizing the labile fraction of watersoluble organic carbon (Facchini et al., 2008; Müller et al., 2009). A number of size-

segregated investigations of the composition of particles impacted by marine sources

have recently reported a ubiquitous contribution of amines to the detectable fraction of the sub-micrometer aerosol water-soluble organic nitrogen (WSON) (Mäkelä et al.,

2001; Facchini et al., 2008; Müller et al., 2009; Violaki and Mihalopoulos, 2010). However, their size-dependent composition in continental air masses has not been reported

In general, online and offline observations of R<sub>3</sub>NH<sup>+</sup> have shown them to be only

to date, despite the fact that most known amine sources are terrestrial.

We present observations of the simple alkyl amines: MA, DMA, TMA, EA, DEA and TEA quantitatively measured using ion chromatography. In Sect. 2.1, we describe the method developed to resolve and detect this suite of alkyl amines at ng levels using ion **ACPD** 

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chromatography. This method was used for the analysis of size-resolved micro-orifice uniform deposit impactor (MOUDI) samples using a protocol described in Sect. 2.2. We also employed the technique for simultaneous hourly gas and particle phase measurements using continuous online sampling with an Ambient Ion Monitor-Ion Chromatography (AIM-IC) system (Sect. 2.3). In Sect. 3 we discuss the observations made using these methods including the size-resolved water-soluble composition of particulate matter in urban and rural locations in Southwestern Ontario (Sect. 3.1) and gas and PM<sub>2.5</sub> observations from Toronto, Canada in July of 2009 (Sect. 3.2).

#### 2 Methods

#### 2.1 Ion chromatography (IC)

Chromatographic resolution of anions and cations was performed using two Dionex ICS-2000 systems operating with reagent-free eluent and suppressed conductivity detection. Anions were separated using a gradient elution method on an AS19 hydroxideselective anion exchange column fitted with a trace anion concentrator column (TAC-ULP1), guard column and an ASRS® 300 conductivity suppressor using KOH as an eluent. A linear gradient from 1-20 mM was employed over the first 12 min, with a linear increase to 85 mM over the next 6 min and then equilibrated back to 1 mM for the final 7 min of the total 25 min run time at a flow rate of 1 mL min<sup>-1</sup>. Two separate methods have been developed for the separation of cations using the Dionex CS17 and CS12A cation exchange columns fitted with a trace cation concentrator column (TCC-ULP1), CG17 or CG12A guard column and a CSRS® 300 suppressor by gradient elution with methanesulfonic acid (MSA). For the CS17, a linear gradient was used at an eluent flow rate of 1.0 mL min<sup>-1</sup> from 2-8 mM over the first 11.55 min, increased to 10 mM over the next 4.45 min and then equilibrated back to 2 mM for 9 min prior to the next run. For the CS12A column, higher eluent strengths were used due to the increased capacity of the column for ion exchange. Over the first 12 min the MSA concentration

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was ramped linearly from 5–10 mM and then from 10–80 mM over the next 10 min and allowed to equilibrate back to 5 mM over the final 3 min of the 25 min run at a flow rate of 1.2 mL min<sup>-1</sup>. Column temperatures, in all cases, were maintained at 30 °C.

All chemicals used were reagent ACS or HPLC grade. Stock solutions of MA (40% w/w), DMA (40% w/w), TMA (45% w/w), EA (71% w/w), DEA (>99.5% w/w), TEA (100%), formic acid (1 mg/mL), acetic acid (1 mg/mL) and oxalic acids (1 mg/mL) were purchased from Sigma-Aldrich (St. Louis, MO). Ethanol (95%) was purchased from Commercial Alcohols (Brampton, ON) and deionised water (18.2 MΩ) was produced in house with a Barnstead Easypure RoDi system (VWR, Mississauga, ON). Premixed analytical standards for inorganic anions (F⁻, Cl⁻, NO₂, Br⁻, NO₃, SO₄⁻, PO₃⁻) and cations (Li⁺, Na⁺, NH₄⁺, K⁺, Mg²⁺, Ca²⁺) and eluent generator cartridges of KOH and MSA were supplied by Dionex Corporation (Sunnyvale, CA).

## 2.1.1 Ion chromatography for resolution of amines and inorganic cations in atmospheric samples

Previous investigations of aliphatic amines have used ion chromatography for the detection and quantification of amines in atmospheric samples (Chang et al., 2003; Murphy et al., 2007; Sorooshian et al., 2008; Erupe et al., 2010), but not for the variety of species investigated in this study. The design of commercial columns has been focused on the separation of inorganic cations, limiting their ability to fully resolve them from protonated organics, such as alkyl amines, especially where their solvated properties are similar. Over the last decade their use in the separation of a number of amines has been demonstrated (Chang et al., 2003; Murphy et al., 2007; Erupe et al., 2010). The amine resolution achieved using the methodology we have developed on the Dionex CS17 column using preconcentration is shown in Fig. 1. This figure includes three overlaid chromatograms: methyl amines (black trace), ethyl amines (blue trace), and inorganic cations (grey dashed line), with evidence of small amounts of inorganic contamination from deionized water in the amine standards. The amines

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are sufficiently separated from one another and from the dominant inorganic species such that valley-to-valley peak integration is possible. The exceptions are TMA and DEA, henceforth TMA+DEA/TMAH++DEAH+, which coelute under all attempted separation conditions, a complication that has been reported previously (Murphy et al., 5 2007) for Dionex columns. The method we have developed for the CS17 column has acceptable resolution performance for use in environmental samples that are not significantly impacted by high loadings of Na<sup>+</sup>, NH<sub>4</sub><sup>+</sup> and K<sup>+</sup>, which might be encountered under conditions with significant mass loadings of sea salt, crustal, or biomass burning particles. An example of the resolution achieved for inorganic cations and amines in a particulate sample from one stage of a MOUDI sampler (aerodynamic particle diameters between 320 and 560 nm) using the CS17 column is shown in Fig. 2. Under conditions of higher concentration, poor separation of Na<sup>+</sup> and NH<sub>4</sub><sup>+</sup> and poor peak shapes become problematic for retrieving accurate peak area integrations, particularly in offline sample analysis.

For analysis of online and offline samples from environments impacted by high loadings of water-soluble cations, a method using a Dionex CS12A column was developed. Improved peak shape due to a higher column exchange capacity was an additional benefit of this alternative method. Figure 3 shows the overlaid chromatograms of a mixed standard of 0.05 µg/mL methyl and ethyl amines (black trace), and inorganic cations (grey dashed line). Increased resolution between Na<sup>+</sup> and NH<sub>4</sub> was achieved, but resolution of TMA and DEA was still not possible. Coelution of MA and EA with NH<sub>4</sub> occurred at relative concentration ratios 5 times lower than in our CS17 methodology, making it more prone to NH<sub>4</sub> interfering in R<sub>3</sub>NH detection. Where required, both methods were used to characterise offline samples to acquire the best possible quantitative data.

Detection of aliphatic amines in any sample using either of these methods is compromised by the presence of high concentrations of NH<sub>4</sub> and/or K<sup>+</sup>. For offline samples, such as the MOUDI samples collected in this study, dilution and reanalysis to improve separation and quantitation of amines and inorganics was utilized to overcome this **ACPD** 

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problem. However, this is not an effective strategy for online sample analysis. Application of deconvolution software could assist in some cases where peaks cannot be resolved due to high loadings. Since NH<sub>3</sub>/NH<sub>4</sub> arise from similar sources as NR<sub>3</sub>/R<sub>3</sub>NH<sup>+</sup> (Van Neste et al., 1987; Schade and Crutzen, 1995), separation of these species is 5 an issue that must be addressed. For online observations, the best approach was the CS17 methodology described above.

Figure 4 illustrates the NH<sub>4</sub> interference in the detection of DMA in a MOUDI sample when using the CS12A column (panel A) and the subsequent improvement in the resolution and detection of DMA and TMA+DEA on the CS17 column (panel B). Some species may not be detected during online analysis with the AIM-IC due to the analytical limitations of separating MA and EA from NH<sub>4</sub><sup>+</sup>. However, when identified by the software as a distinct peak (area >0.001 µS min), a minimum quantity of each alkyl amine can be determined even under conditions of partial coelution. The ability to resolve and quantify amines in an atmospheric sample, therefore, depends not only on the absolute amount of amine present, but on the absolute amount of ammonia and on the relative amounts of amine and ammonia. In an ideal situation, the sampling period will be sufficiently long to collect enough amine to exceed the detection limit, but short enough that NH<sub>4</sub> will not coelute. The occurrence of coelution is most frequent in offline time-integrated sampling and is dependent on local conditions at the sampling site.

#### MOUDI filter preparation and field operation

To ensure particle collection filters were free of residual organic compounds and salts prior to use in particle collection, the following pretreatment methodology was used to ensure their removal. The filters (47 mm, 0.1 mm thickness, 5-6 µm pore size, PTFE, Savillex, MN) were soaked for 24 h in ethanol with intermittent shaking, rinsed thoroughly with 100 mL of deionised water, placed in a drying rack and dried for 12 h. The filters were then transferred to clean petri dishes (50 mm, Pall Canada Ltd, Ville St. Laurent, QC) and stored in a light shielded container for 24 h to equilibrate. The filters

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were mounted on the collection stages of the MOUDI (Model 110, MSP, MN) (Marple et al., 1986) for online sampling just prior to collection.

The MOUDI was used to collect size-resolved aerosol samples in urban and rural environments in Ontario periodically from July 2009-May 2010. Specifically, these locations were: on the roof of the Physical Geography Building of the University of Toronto (43°39'37.60" N, 79°23'47.42" W), on top of a 3 m shed in Dorset (45°14′14.96" N, 78°53′47.33" W) and an open grass field in Lambton Shores (43°9'43.41" N, 81°56'4.33" W), respectively, under a rain shelter that allowed for free ventilation. The sampling interval varied between 20 and 46 h according to the level of particulate matter reported in real time by the Ontario Ministry of the Environment (http://www.airqualityontario.com/history/summary.cfm). The MOUDI was operated at a flow rate of 30 L min<sup>-1</sup> with 10 stages at the following 50% cut-off points for the particle aerodynamic diameters: 10, 5.6, 3.2, 1.8, 1.0, 0.56, 0.32, 0.18, 0.1, and 0.056 µm. No back filters to collect particles less than 0.056 µm were used due to pump limitations.

Filter extracts were obtained by room temperature sonication in 10 mL of deionized water for 1 h, filtration through a 0.45 µm PTFE membrane (Pall Ion Chromatography Acrodisc®, VWR International, Mississauga, ON) and dilution to a final volume of 15 mL with deionised water. Extracts were then analyzed by ion chromatography using one or both of the methods described above. Blanks were collected by transporting filters to and from the sampling site and subjecting them to the same extraction and analysis procedures.

#### Online detection by URG-AIM

The operational principles of the AIM-IC 9000D (URG, Chapel Hill, NC) system have been previously described (Ellis et al., 2010a). Briefly, the AIM-IC system employs two online ion chromatography systems for the simultaneous detection of cations and anions from gas and particle samples concentrated from a continuous flow of ambient air. In our system, the air sample is pulled at 3.0 L min<sup>-1</sup> through a minimum inlet **ACPD** 

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length constructed of Teflon-coated aluminum and fitted with an impactor assembly to select particles with an aerodynamic diameter of less than 2.5 µm. Gases are stripped from the air stream by passing through a liquid parallel plate denuder with continuously replenished solvent flowing across the surface. The particles in the air flow are constrained into a supersaturated steam condensation coil and cyclone assembly and grown hygroscopically for collection as an aqueous solution. The sample flows containing the soluble gas and particle analytes are separated into equal aliquots for anion and cation chromatographic analyses every hour following 60 min of ambient sampling.

The AIM-IC was deployed utilizing the CS17 columns and operated according to the separation methodology described in Sect. 2.1 to monitor ambient air in downtown Toronto, ON at the University of Toronto's St. George Campus from 26 June-3 July 2009. Quantities were determined from offline injections of known aqueous standards of the analytes of interest across the relevant range of observed atmospheric concentrations. Background levels of all compounds detected by the AIM-IC were determined by overflowing the inlet with high purity zero air and subtracting the average peak area acquired over 8 h from the peaks observed while sampling ambient air. If background peaks were not observed for a particular species, the 5 lowest points of the time series were taken as blank values, the average of which was subtracted from the ambient dataset. Simultaneous gas phase NH3 measurements were made using a quantum cascade tunable infrared diode laser absorption spectrometer (QC-TILDAS) system (Aerodyne, USA) (Ellis et al., 2010b).

#### 2.4 Sensitivity and detection limits of aliphatic amines

Previous observations of aliphatic amines have shown their gas phase concentrations to be strongly dependent on the proximity of the amine source due to fast reaction with OH. Near source values can be on the order of several ppb, while concentrations within a day of transport are typically on the order of ppt (Grönberg et al., 1992; Schade and Crutzen, 1995). Therefore, a technique capable of detecting and quantifying concentrations in the ppt-ppb range is ideal. The sensitivity of the conductivity detector to 10, 27435-27477, 2010

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each amine was calculated as the peak area response as a function of the moles of analyte injected. Amine concentrations spanned 3 orders of magnitude. Sensitivities for each compound were determined for each separation methodology and compared to  $NH_4^+$ , the most common nitrogenous base in atmospheric samples. The response is linear for all the amines investigated and their sensitivity is equal to the slope. The response is non-linear for  $NH_4^+$  across its atmospherically relevant range in gas and particulate samples. To calibrate for ammonia, an exponential fit in the form of Eq. (1) has been applied, where k is the indicator of the instrument sensitivity.

$$y = y_0 + Ae^{-kx} \tag{1}$$

The exponential fit applied is linear in two regions: near the origin and as it approaches its asymptote. Since the greatest instrumental response is in the linear region near the origin,  $NH_4^+$  samples are ideally quantified here. This makes the k-value suitable for comparison with the slopes of the amine species that have linear responses across the range of  $8 \times 10^{-12}$  to  $2 \times 10^{-8}$  moles injected. The detection sensitivities of all the alkyl amines explored in this study were found to be an order of magnitude greater than ammonium (Table 1). This is likely due to their stronger basicity. With pKa values up to one unit higher than that of ammonia, a higher fraction of alkyl amines exist in the protonated form in solutions of the same pH. This makes them relatively stronger electrolytes and, therefore, more conductive in solution; hence, the improvement in detection sensitivity when using the MSA eluent. Furthermore, the CS12A method is 1.5–3.5 times more sensitive to the presence of all the alkyl amines studied here except for TMA and DEA, but suffers from being unable to resolve the smaller amines from NH<sub>4</sub> and K<sup>+</sup> when the latter are present at significant concentrations in atmospheric samples. Detection limits of the ion chromatography method were calculated as an absolute number of moles using the slope and three times the standard deviation in the area of the background conductivity observed in sequential blank samples at the elution time of the analyte of interest. The detection limits for each alkyl amine and  $NH_{\perp}^{4}$ are reported in Table 2, detection limits for all other inorganic analytes are <500 pmol.

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By utilizing a continuously regenerated solvent in a parallel plate denuder and a supersaturated condensation coil to collect water soluble gases and particles from atmospheric samples, followed by online preconcentration with Dionex concentrator columns, hourly detection of MA, DMA, TMA, EA, DEA and TEA at the ppt and ng m<sup>-3</sup> levels was achieved with the AIM-IC system. AIM-IC detection limits were then calculated as a function of the sampling rate at standard ambient conditions. The application of ion chromatography with concentrator columns to offline MOUDI samples collected over a minimum of 24 h and at 10 times the flow rate of the AIM-IC reduced the R<sub>3</sub>NH<sup>+</sup> detection limits to sub-ng m<sup>-3</sup> levels for the particulate samples collected, allowing the composition of subsections of the aerosol population to be probed for alkyl amine presence and concentration. Limits of detection were calculated as three times the standard deviation of the background in blank samples run during online or offline analysis and expressed as ambient concentrations based on length of time and volume sampled.

#### Results and discussion

#### Size-resolved particulate amines

Eight samples were collected using the MOUDI under the conditions listed in Table 3. The fine aerosol fraction (PM<sub>1,8</sub>) collected by the MOUDI is determined by summing the six lowest MOUDI size bins. The total water soluble mass was determined from the sum of the species quantified by ion chromatography. The fraction of amines in the total mass of soluble PM<sub>1.8</sub> (%  $m_{sol}$ ) ranges from <1% to 9.1% across our observations. The amine mass fraction in  $PM_{2.5}$  (%  $m_{tot}$ ) was calculated by dividing  $PM_{1.8}$  amines by the total mass loading of PM<sub>2.5</sub> obtained from the nearest Ontario Ministry of Environment air quality monitoring station for the period of observation. The sum of the particulate alkyl amines was never more than 1% of the PM<sub>2.5</sub> aerosol mass loadings. PM<sub>1.8</sub> mass loadings of all the amines, denoted as R<sub>3</sub>NH<sup>+</sup>, and NH<sub>4</sub><sup>+</sup> are shown in Table 3. While the R<sub>3</sub>NH<sup>+</sup> mass loadings only span one order of magnitude among the samples, the

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mass loading of  $NH_4^+$  varies by a factor of 60. The final column includes the molar ratio of the sum of amines to ammonium ( $R_3NH^+/NH_4^+$ ) in  $PM_{1.8}$ , the variability of which depends more on the mass loading of  $NH_4^+$  than that of  $R_3NH^+$ .

The presence of amines was detected throughout the fine mode (0.056–1.8  $\mu$ m) in all cases for DMAH<sup>+</sup> and TMAH<sup>+</sup>+DEAH<sup>+</sup>. The presence of TEAH<sup>+</sup> was found in 5 samples and MAH<sup>+</sup> near the LOD in the sample collected at Dorset, ON. In all cases the mass loadings or moles of TMAH<sup>+</sup>+DEAH<sup>+</sup> were calculated assuming everything was DEAH<sup>+</sup>. DEA has a larger Henry's law constant and pKa value, indicating that it would be more likely to partition to an aqueous particle than TMA under conditions of similar gas phase mixing ratios. Furthermore, the dissociation constants ( $K_p$ ) calculated by Murphy et al. (2007) for the formation of aminium nitrate salts from gas phase precursors (R3) were at least three orders of magnitude lower for DEA than TMA. DEA showed higher analytical sensitivity under the CS17 method described in Sect. 2.1 and has a lower molecular mass than TMA. Therefore, the values we have calculated and report here are the lower estimate for the sum of these species. The true values could be up to 23% greater for mass and moles if an appreciable fraction of the sample is actually TMA.

#### 3.1.1 Aerosol composition of water-soluble ionic species

Figure 5 shows the composition of water-soluble particulate species, as a function of particle size at three different sampling sites. In all cases, the dominant cation at particle diameters less than 1.8  $\mu$ m was NH $_4^+$ , which nearly balanced the major anionic components of SO $_4^{2-}$ , NO $_3^-$ , formic, acetic and oxalic acids. Thus, the sub-micrometer aerosol we sampled was neutralized or nearly neutral, with respect to ion balance, in all cases. In some cases, amines were found to have important contributions to the total mass of the fine mode. Panel A displays a sample collected at Dorset, Ontario, a rural forest site, on 14 August 2009 with the air mass originating over South-Western Ontario. Amines contribute about 5% of the total water-soluble cation fraction in the

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accumulation mode aerosol from 0.056–1.8 μm. This sample was also particularly enhanced in formic, acetic and oxalic acids in addition to the dominant SO<sub>4</sub><sup>2-</sup> anion. The sample composition displayed in panel B was collected from the roof of the Physical Geography Building at University of Toronto, a dense urban location, on 7 August 2009. The aerosol is neutralized in terms of NH<sub>4</sub><sup>4</sup>-NO<sub>3</sub><sup>-</sup>-SO<sub>4</sub><sup>2-</sup> throughout the accumulation mode, with <5% of the water-soluble cation fraction being contributed by amines. Ca<sup>2+</sup> and NO<sub>3</sub><sup>-</sup> dominate the coarse mode, likely as CaCO<sub>3</sub> with some Ca(NO<sub>3</sub>)<sub>2</sub> from processing of lofted particles. The relative composition of this sample is typical for samples collected in downtown Toronto, although the magnitude of mass in the sample can vary considerably, particularly in the accumulation mode. The sample displayed in panel C was collected from Lambton Shores, Ontario, a rural location, on 10 April 2010. The contribution of amines was insignificant to the aerosol neutrality, despite NO<sub>3</sub><sup>-</sup> representing about 30% of the anion water-soluble fraction at the peak of the accumulation mode.

lon chromatography cannot measure  $H^+$  in solution, but it is assumed to balance most of the remaining negative ions in the  $PM_{1.8}$ . No significant unidentified peaks were observed in the cation or anion chromatographs. The dominant cation at particle diameters greater than 1.8 µm was  $Ca^{2+}$ , which is not balanced by a quantifiable anionic species. The most likely constituent that balances the observed  $Ca^{2+}$  is  $HCO_3^-/CO_3^{2-}$ , which cannot be quantitatively detected by the sample handling and ion chromatographic methods used in this study.  $Ca(HCO_3)_2$  and  $CaCO_3$  are known to be major components of concrete and arises from surface erosion in urban environments. This is likely responsible for the large  $Ca^{2+}$  signal seen in panel B of Fig. 5. In panels B and C, coarse mode nitrate was also observed. It has been well-documented that coarse mode nitrate can be formed by acid displacement reactions between  $HNO_3$  and  $Ca(HCO_3)_2$  and  $CaCO_3$  to form  $Ca(NO_3)_2$  (Goodman et al., 2000; Hanisch and Crowley, 2001; Bauer et al., 2004; Vlasenko et al., 2006; Liu et al., 2008), which further supports the presence of unmeasured carbonate in our samples.

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Despite being a small fraction of the total cation composition and the total aerosol mass, the alkyl amines measured may influence the effective pH of aqueous aerosols by several units (Pankow, 2003; Pratt et al., 2009), especially where they represent more than 1% of the water-soluble aerosol mass. Stronger basic properties compared to  $NH_4^+$  make the amines better at limiting the  $H_4^+$  freely available in solution, even when present in molar quantities more than an order of magnitude smaller than  $NH_4^+$ .

#### 3.1.2 Size-dependence of alkyl amine mass

Size distributions of amines by mass in samples collected at Dorset, Toronto and Lambton Shores, ON are shown in the panels of Fig. 6. The size dependence of the alkyl amines shows that their mass maximizes most often in the accumulation mode. The amines detected in our samples maximized in the 320–560 nm size range in 7 of 8 samples. Often, only tens of  $ng\,m^{-3}$  were observed in the size bin with the largest contributions from TMAH<sup>+</sup>+DEAH<sup>+</sup> and DMAH<sup>+</sup>. The maximum mass loading of each amine detected in our samples and the size bin in which it maximized is given in Table 4. TEAH<sup>+</sup>, when present, showed the most variability in the distribution of the mass throughout the aerosol populations we collected. In Sample 6 (collected in Toronto) the abundance of TEAH<sup>+</sup> was 14  $ng\,m^{-3}$ , the highest we encountered in this sample set, and was found to maximize in the 3.2–5.6  $\mu$ m size range.

The variety of amines detected changed based on sampling location and time of year. This could be due to changes in the source region of the air mass sampled, regional temperature and relative humidity, and seasonality as amine emissions are expected to increase with temperature and biological activity (Schade and Crutzen, 1995; Müller et al., 2009; Pratt et al., 2009). Figure 6a illustrates Sample 4 from Dorset, ON, which had the widest array of amines present in any sample collected. The sampled air mass originated over a region of South-Western Ontario with a high density of agricultural activity, including animal husbandry, a likely source region for the primary emission of alkyl amines to the atmosphere. Two samples, one displayed in Fig. 6c were collected in this source region, at Lambton Shores, to investigate par-

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ticulate amines in proximity to presumed sources. During the collection period air originated over Michigan and the northern Great Lakes region of Ontario. Whether the amines we observed in Fig. 6c had partitioned from local emissions or were preexisting components of transported aerosol is not possible to determine from these samples. The collected sample had 19 ng m<sup>-3</sup> of TMAH<sup>+</sup>+DEAH<sup>+</sup> in the 320–560 nm size bin, the highest detected mass of a single amine species in our samples. While the absolute mass loadings of amines seem comparable, or higher, in the downwind Dorset sample compared to the source region sample, it should be kept in mind that these samples were collected in the summer and spring, respectively. The seasonality of amine emissions and partitioning has been related to primary productivity (Müller et al., 2009) and local meteorology (Pratt et al., 2009). Online observations of both gas and PM<sub>2.5</sub> ionic, water-soluble composition could yield more information in identifying source regions and thermodynamic factors that govern the partitioning of amines.

Figure 6b from Toronto, ON shows an atypical amine mass loading compared to our other MOUDI samples collected in Toronto. The sampled air originated over central Ontario and Georgian Bay with similar amine loadings to those collected at the other two locations. Figure 6d shows a more typical size-dependence of the mass loading of individual amines in our urban samples. In general, the observations from Figs. 5 and 6 show that even in instances where very large quantities of NH<sub>4</sub> were present throughout the fine mode, the presence of DMAH<sup>+</sup> and TMAH<sup>+</sup> +DEAH<sup>+</sup> was observed. These observations are consistent with previous quantitative observations of alkyl amines in fine particulate samples (Sorooshian et al., 2008). Our measurements show the sizeresolved distributions throughout the sampled aerosol populations are also consistent with previous observations in marine atmospheres (Mäkelä et al., 2001; Facchini et al., 2008; Müller et al., 2009; Violaki and Mihalopoulos, 2010) with the maximum mass loadings seen in the accumulation mode, specifically from 320-560 nm. Our observations are, to the best of our knowledge, the first to show the size-dependence of particulate alkyl amines in continental air masses.

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The dominance of NH<sub>4</sub><sup>+</sup> throughout the accumulation mode is most likely due to the ambient concentrations of NH<sub>3</sub> being significantly higher than gas phase amines at our sampling sites (Sect. 3.2 provides evidence in Toronto). Typical NH<sub>3</sub> measurements in Southwestern Ontario are on the order of a few ppb (Ellis et al., 2010a). While alkyl amines are not routinely measured, they are expected to range from several ppb close to sources to ppt levels in the background atmosphere due to their short lifetime against OH ( $\tau$ ~1 d). A higher ambient concentration of NH<sub>3</sub> would allow its uptake to dominate the neutralization of the aerosol as growth by condensation occurs and the ratio of surface area to volume becomes more limited for reactive uptake. Theoretical calculations by Kurten et al. (2008) and Loukonen et al. (2010) have shown that small cluster formation of aminium salts with inorganic acids is thermodynamically favoured over ammonium salts. At smaller particle sizes (e.g. clusters – 10 nm), amines may be able to successfully out-compete NH<sub>3</sub> due to thermodynamics of the system as shown experimentally (Murphy et al., 2007; Lloyd et al., 2009; Bzdek et al., 2010). As particles grow by condensation and reactive uptake, competition with NH<sub>3</sub> for transport to the surface becomes dominant and the influence of relative concentrations supersedes the thermodynamic considerations.

To examine the size-dependence of the relative importance of amines versus ammonia, the molar ratio of the sum of amines to ammonium ( $R_3NH^+/NH_4^+$ ) detected in each sample was compared in Fig. 7. The error bars represent the uncertainty calculated based on the propagation of errors for the measured ammonium and only those amines that were detected in the sample. In most cases the observed ratio is well-constrained because the amounts of all the considered analytes are above the detection limits. Small quantities of all the analytes in the coarse mode produced larger uncertainties in those ratios. In all MOUDI samples, the molar ratio  $R_3NH^+/NH_4^+$  showed one maximum below 200 nm and decreased as the particle size increased up to about 1  $\mu$ m; the ratio then increased again in the coarse mode. All samples show this behaviour re-

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gardless of the absolute values of the R<sub>3</sub>NH<sup>+</sup>/NH<sub>4</sub> fraction. In the fine mode, maxima for an individual size bin ranged from ratios of 0.020 to 0.37, with the largest fraction being observed in a sample from Dorset, ON (Fig. 7a). A similar enhancement in R<sub>3</sub>NH<sup>+</sup>/NH<sub>4</sub> in excess of 0.3 was seen in Toronto (Fig. 7b) from an air mass also originating from southwestern Ontario. However, most samples from the Toronto region typically show ratios below 0.04 (Fig. 7d). The samples collected at Lambton Shores (e.g. Fig. 7c), were in closer proximity to agriculturally active regions, but did not have ratios of R<sub>3</sub>NH<sup>+</sup>/NH<sub>4</sub><sup>+</sup> greater than 0.05 despite having the highest absolute amine mass loading of all the samples collected. The average molar ratio for PM<sub>1,8</sub> in each sample is given in the final column on Table 3.

These observations of amine composition in conjunction with other ionic aqueous species can help to inform models of particle formation and thermodynamic partitioning and predictions of aerosol properties, such as pH and hygroscopicity. The observations in Fig. 7 demonstrate that an assumption that the fine aerosol population is internally mixed is not appropriate, especially with reference to the R<sub>3</sub>NH<sup>+</sup>/NH<sub>4</sub> ratio. In general, the amines contribute more importantly in a relative sense to the reduced nitrogen composition of particles with diameters <200 nm. Possible explanations include the higher thermodynamic stability of aminium salt clusters, or the preferential reactive uptake of amines by ultrafine particles with higher surface area-to-volume ratios. Alternately, particulate amines may be more susceptible to condensed phase reactions than ammonium, leading to their preferential degradation as particles age and grow. In the future, a back filter can be included to collect the smallest particles which may provide more information on nucleation mode particles. The maxima in the coarse mode are highly uncertain, but do appear to be significantly higher than the accumulation mode particles. From Figs. 5 and 6, it is evident that the coarse mode represents a population of particles with very different composition than the fine particles, and with lower absolute amounts of amines. The coarse mode amines may result from primary biogenic emissions or reactive uptake on aqueous coarse mode aerosol that has been significantly aged by secondary acidification with HNO<sub>3</sub>. In either case,

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the ammonium signal in these coarse mode samples was near or below our detection limits, enhancing the observed ratio when any amines are above their detection limits.

#### Online detection of amines in the gas phase and PM<sub>2.5</sub>

The utility of the information provided by the MOUDI data is limited by the long integration time necessary for collection and the absence of simultaneous gas phase measurements. To address these limitations, continuous online collection and analysis of gas and particle phase analytes was performed with the AIM-IC from 26 June-6 July 2009. This time period was characterized by cool and humid conditions, with rain events occurring from 09:00-14:00 and at 21:00 on 28 June, from 14:00-16:00 and 20:00-22:00 on 29 June, at 04:00 and 20:00 on 30 June, and at 22:00 on 1 July (all times are Eastern Daylight Savings Time (EDT)). The northwesterly flow at the start and end of the campaign was interrupted by a period of southwesterly flow between 28 June and 1 July, and stagnant air early on 2 July. Amines were detected in both the gas and particle phases throughout the campaign. Figure 8 shows that amines in the gas phase (panel A) were almost always at or near the instrument detection limits. Mixing ratios of DMA, and TMA+DEA ranged from below the detection limit to 2.7 ppt for both sets of compounds. While observations of TEA are reported, the measured values were consistently below the detection limit. Amine emission sources in the Toronto area and their collective emission rates are not well-known, although particulate amines have been previously detected during new particle formation events (Tan et al., 2002). The variability in mixing ratio is not consistent across all species or with NH<sub>3</sub>, indicating different sources may exist for each. If local amine sources are limited, the observed abundances of a few ppt may be due to dilution and/or reaction with OH ( $\tau$ ~1 d) prior to arriving at our sampling location.

Simultaneous observations of NH<sub>3</sub> (Fig. 8a) showed mixing ratios ranging from 1-18 ppb. Therefore, the relative ratios of NR<sub>3</sub>/NH<sub>3</sub> in the gas phase range from 1.6-20×10<sup>-3</sup> over the course of our observations. According to the modelling studies by

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Murphy et al. (2007) and Barsanti et al. (2009), at this range of gaseous ratios, alkyl amines would seldom out-compete ammonia in bulk partitioning to the particle phase. Our particle observations showed only periodic amine mass loadings in PM<sub>2.5</sub> above our online detection limits. Figure 8 shows the sum of TMAH++DEAH+ observed in the particle phase (panel B) at mass loadings in the range of below detection limit to 16 ng m<sup>-3</sup>. The sporadic nature of the MAH<sup>+</sup> observations results from the difficulty in resolving this peak in the presence of much higher amounts of NH<sub>4</sub><sup>+</sup>. On two occasions, the increase in TMAH++DEAH+ coincided with mass loadings of PM<sub>2.5</sub> NO<sub>3</sub> greater than 1 µg m<sup>-3</sup>, suggestive of the formation of aminium nitrate salts. However, early on 30 June TMAH++DEAH+ was not detected when NO<sub>3</sub> was high and early on 1 July, it was observed despite nitrate being lower than  $0.1\,\mu\mathrm{g\,m^{-3}}$ . Therefore, an increased presence of NO<sub>3</sub> in fine particulate matter does not appear to govern the presence and concentration of TMAH++DEAH+. In Sorooshian et al. (2008), high amine mass loadings were measured by PILS-IC at the same time as positive values of excess nitrate, defined as the nitrate left over after sulfate and nitrate have been neutralized by the available ammonium, as measured by C-ToF-AMS. Values of excess nitrate inferred from the AIM-IC were zero within the uncertainty of the calculation, and this uncertainty is much larger than the mass loading of the observed amines. In our observations, it is more likely that the presence of nitrate in the particle phase is indicative of either high relative humidity or high nitric acid, both of which would be conducive to the partitioning of amines to the particle phase.

In most cases, increases in particulate amine loadings coincided with a decrease of amines in the gas phase, but mass balance was not observed (e.g. formation of 7.7 ng m<sup>-3</sup> (2.54 pptv equivalent) of TMAH<sup>+</sup>+DEAH<sup>+</sup> on 1 July and a coincident decrease in 0.85 pptv of TMA+DEA). This may be indicative of a sudden change in the total (gas+particle) amines during the time period or a slow time response to changing amine concentrations in our gas channel because of adsorption in our inlet. Such small amounts of amines partitioning to the particle phase make it challenging to use our online observations to determine if R<sub>3</sub>NH<sup>+</sup> is displacing NH<sub>4</sub><sup>+</sup> from particles because

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changes in the abundance of NH<sub>4</sub><sup>+</sup> smaller than 100 ng m<sup>-3</sup> are below our measurement precision for that species.

While the online sampling campaign did not coincide with the collection dates of the Toronto MOUDI samples, some consistency may be expected between the R<sub>3</sub>NH<sup>+</sup>/NH<sub>4</sub> observed in both sets of samples. Based on the final column in Table 3, the molar ratio for PM<sub>1.8</sub> in MOUDI samples ranged from  $\sim 10^{-3}$  to  $10^{-1}$ . Because the online AIM-IC method has higher detection limits than the time-integrated MOUDI samples, at an  $NH_4^+$  mass loading of  $0.5 \,\mu \mathrm{g} \, \mathrm{m}^{-3}$ , representative of the online sampling campaign, the lowest ratio we could quantitatively measure in hourly samples is  $>10^{-2}$  (see the detection limits in Table 2). In the MOUDI samples with similar PM<sub>1.8</sub> mass loadings of  $NH_4^+$ , the  $R_3NH^+/NH_4^+$  ratio was always less than  $1.5 \times 10^{-2}$ . Based on the observed gas phase NR<sub>3</sub>/NH<sub>3</sub> ratios of 1.6–20×10<sup>-3</sup> (quantifiable because of high NH<sub>3</sub>), a R<sub>3</sub>NH $^+$ /NH $^+_4$  ratio in PM<sub>2.5</sub> of >10 $^{-2}$  would only be expected if amines are able to significantly outcompete ammonia based on thermodynamic considerations. Barsanti et al. (2009) and Murphy et al. (2007) suggested that the formation of stronger aminium salts with organic acids or nitric acid could favour preferential partitioning of amines to the particle phase. Particulate organic acids, such as oxalic acid, were not above the AIM-IC detection limits (0.1 µg m<sup>-3</sup>) during any of the online observation period. Because of the high humidity conditions during much of the campaign, the aerosol were likely aqueous, in which case partitioning according the Reaction (R1) may be most relevant. For example, the effective Henry's law constant for DEA (after accounting for protonation) is an order of magnitude higher than NH<sub>3</sub>, which would result in a stronger preference for partitioning to aqueous aerosol. Observation periods spanning a wider range of atmospheric conditions would be valuable to assess the relative importance of meteorological conditions and chemical composition in dictating the gas-particle partitioning of atmospheric amines.

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These measurements contribute to the small, but expanding, collection of ambient observations of atmospheric amines (Angelino et al., 2001; Mäkelä et al., 2001; Tan et al., 2002; Denkenberger et al., 2007; Facchini et al., 2008; Sorooshian et al., 2008; Müller et al., 2009; Smith et al., 2010; Violaki and Mihalopoulos, 2010). An advantage of the ion chromatography analysis described here is that the calibration protocol provides measurements (mass loadings or mixing ratios) with a high degree of precision for trace quantities (ng m<sup>-3</sup> or ppt). The sensitivity of the IC technique is a factor of 5–10 higher for the individual amines measured compared to ammonia, which contributes to the low detection limits for the amines. Nevertheless, because in atmospheric samples ammonia/ammonium can be present in concentrations several orders of magnitude higher than the amines, it can impede the detection of some amines (e.g. methylamine and ethylamine) by this method. An ongoing issue with IC measurements of amines is the inability to resolve DEA and TMA. Because one, or both, of these amines appear to be among the most significant of the simple alkyl amines in atmospheric samples, and because their  $K_H$  and pKa values are quite different, it is highly desirable to achieve individual quantitative measurements.

Our MOUDI observations in one urban and two rural environments demonstrate consistent trends with respect to size-dependence: an absolute maximum of amine mass loading in 320–560 nm particles, and an increase in importance relative to ammonium for the smallest (56–180 nm) particles measured. The importance relative to ammonium in the coarse mode is less meaningful because ammonium was often below our detection limits, despite the presence of detectable levels of amines. In PM<sub>1.8</sub> measurements, calculated by summing the six lowest MOUDI size bins, amines contributed tens of ng m<sup>-3</sup> and constituted between 0.3 and 10% of the water soluble particulate mass loading. To the best of our knowledge, the AIM-IC data represent the first simultaneous observations of gas and particle phase amine species with accompanying water-soluble ionic speciation in PM<sub>2.5</sub> at hourly time resolution. Through online sam-

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pling in an urban environment, we observed instances of amines increasing in the particle phase at times with elevated PM<sub>2.5</sub> nitrate. The mass loading of amines in PM<sub>2.5</sub> never exceeded 20 ng m<sup>-3</sup> during the online sampling period, a value too low to cause observable changes to the abundance of NH<sub>4</sub> or NO<sub>3</sub>. Observations made using this technique can provide data which is valuable in the evaluation of models designed to predict the contribution of amines to particle nucleation, or to bulk particle composition based on thermodynamic partitioning. However the size dependence of the R<sub>3</sub>NH<sup>+</sup>/NH<sup>+</sup> ratio consistently seen in the MOUDI samples shows that an assumption that the fine aerosol population is internally mixed is not appropriate.

#### References

- Akyüz, M.: Simultaneous determination of aliphatic and aromatic amines in indoor and outdoor air samples by gas chromatography-mass spectrometry, Talanta, 71, 486-492, 2007.
- Angelino, S., Suess, D. T., and Prather, K. A.: Formation of aerosol particles from reactions of secondary and tertiary alkylamines: characterization by aerosol time-of-flight mass spectrometry, Environ. Sci. Technol., 35, 3130-3138, 2001.
- Atkinson, R., Perry, R. A., and Pitts Jr., J. N.: Rate constants for the reaction of the OH radical with CH<sub>3</sub>SH and CH<sub>3</sub>NH<sub>2</sub> over the temperature range 299–426 K, J. Chem. Phys., 66, 1578– 1581, 1977a.
- Atkinson, R., Perry, R. A., and Pitts Jr., J. N.: Rate constants for the reactions of the OH radical with (CH<sub>3</sub>)<sub>2</sub>NH, (CH<sub>3</sub>)<sub>3</sub>N, and C<sub>2</sub>H<sub>5</sub>NH<sub>2</sub> over the temperature range 298–426 K, J. Chem. Phys., 68, 1850-1853, 1977b.
- Atkinson, R., Baulch, D. L., Cox, R. A., Hampson, R. F., Kerr, J. A., Rossi, M. J., and Troe, J.: Evaluated kinetic, photochemical and heterogeneous data for atmospheric chemistry. 5. IU-PAC subcommittee on gas kinetic data evaluation for atmospheric chemistry, J. Phys. Chem. Ref. Data, 26, 521-1011, 1997.
- Barsanti, K. C., McMurry, P. H., and Smith, J. N.: The potential contribution of organic salts to new particle growth, Atmos. Chem. Phys., 9, 2949-2957, doi:10.5194/acp-9-2949-2009, 2009.
- Bauer, S. E., Balkanski, Y., Schulz, M., Hauglustaine, D. A., and Dentener, F.: Global mod-

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eling of heterogeneous chemistry on mineral aerosol surfaces: influence on tropospheric ozone chemistry and comparison to observations, J. Geophys. Res.-Atmos., 109, D02304, doi:10.1029/2003jd003868, 2004.

Berndt, T., Stratmann, F., Sipilä, M., Vanhanen, J., Petäjä, T., Mikkilä, J., Grüner, A., Spindler, G., Lee Mauldin III, R., Curtius, J., Kulmala, M., and Heintzenberg, J.: Laboratory study on new particle formation from the reaction OH + SO<sub>2</sub>: influence of experimental conditions, H<sub>2</sub>O vapour, NH<sub>3</sub> and the amine tert-butylamine on the overall process, Atmos. Chem. Phys., 10, 7101-7116, doi:10.5194/acp-10-7101-2010, 2010.

Bzdek, B. R., Ridge, D. P., and Johnston, M. V.: Amine exchange into ammonium bisulfate and ammonium nitrate nuclei, Atmos. Chem. Phys., 10, 3495-3503, doi:10.5194/acp-10-3495-2010, 2010.

Chang, I. H., Lee, C. G., and Lee, D. S.: Development of an automated method for simultaneous determination of low molecular weight aliphatic amines and ammonia in ambient air by diffusion scrubber coupled to ion chromatography, Anal. Chem., 75, 6141-6146, doi:10.1021/Ac0347314, 2003.

Denkenberger, K. A., Moffet, R. C., Holecek, J. C., Rebotier, T. P., and Prather, K. A.: Real-time, single-particle measurements of oligomers in aged ambient aerosol particles, Environ. Sci. Technol., 41, 5439-5446, 2007.

Dinar, E., Anttila, R., and Rudich, Y.: CCN activity and hygroscopic growth of organic aerosols following reactive uptake of ammonia, Environ. Sci. Technol., 42, 793-799, 2008.

Donaldson, D. J.: Adsorption of atmospheric gases at the air-water interface. I. NH<sub>3</sub>, J. Phys. Chem. A, 103, 62-70, 1999.

Ellis, R. A., Murphy, J. G., Markovic, M. Z., VandenBoer, T. C., Makar, P. A., Brook, J., and Mihele, C.: The influence of gas-particle partitioning and surface-atmosphere exchange on ammonia during BAQS-Met, Atmos. Chem. Phys. Discuss., 10, 21895-21929, doi:10.5194/acpd-10-21895-2010, 2010a.

Ellis, R. A., Murphy, J. G., Pattey, E., van Haarlem, R., O'Brien, J. M., and Herndon, S. C.: Characterizing a Quantum Cascade Tunable Infrared Laser Differential Absorption Spectrometer (QC-TILDAS) for measurements of atmospheric ammonia, Atmos. Meas. Tech., 3, 397–406, doi:10.5194/amt-3-397-2010. 2010b.

Erupe, M. E., Liberman-Martin, A., Silva, P. J., Malloy, Q. G. J., Yonis, N., Cocker, D. R., and Purvis-Roberts, K. L.: Determination of methylamines and trimethylamine-N-oxide in particulate matter by non-suppressed ion chromatography, J. Chromatogr, A, 1217, 2070**ACPD** 

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2073, doi:10.1016/j.chroma.2010.01.066, 2010.

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Facchini, M. C., Decesari, S., Rinaldi, M., Carbone, C., Finessi, E., Mircea, M., Fuzzi, S., Moretti, F., Tagliavini, E., Ceburnis, D., and O'Dowd, C. D.: Important source of marine secondary organic aerosol from biogenic amines, Environ. Sci. Technol., 42, 9116-9121, 2008.

Finlayson-Pitts, B. J. and Pitts Jr., J. N.: Chemistry of the Upper and Lower Atmosphere: Theory, Experiments, and Applications, Academic Press, San Diego, CA, 2000.

Gibb, S. W., Mantoura, R. F. C., and Liss, P. S.: Ocean-atmosphere exchange and atmospheric speciation of ammonia and methylamines in the region of the NW Arabian Sea, Global Biogeochem. Cy., 13, 161-177, 1999.

Goodman, A. L., Underwood, G. M., and Grassian, V. H.: A laboratory study of the heterogeneous reaction of nitric acid on calcium carbonate particles, J. Geophys. Res.-Atmos., 105, 29053-29064, 2000.

Grönberg, L., Lövkvist, P., and Jönsson, J. A.: Measurement of aliphatic amines in ambient air and rainwater. Chemosphere. 24, 1533-1540, 1992.

Hanisch, F. and Crowley, J. N.: The heterogeneous reactivity of gaseous nitric acid on authentic mineral dust samples, and on individual mineral and clay mineral components, Phys. Chem. Chem. Phys., 3, 2474-2482, 2001.

Huang, G., Hou, J., and Zhou, X. L.: A measurement method for atmospheric ammonia and primary amines based on aqueous sampling, OPA Derivatization and HPLC Analysis, Environ. Sci. Technol., 43, 5851-5856, doi:10.1021/Es900988g, 2009.

Junninen, H., Ehn, M., Petäjä, T., Luosujärvi, L., Kotiaho, T., Kostiainen, R., Rohner, U., Gonin, M., Fuhrer, K., Kulmala, M., and Worsnop, D. R.: A high-resolution mass spectrometer to measure atmospheric ion composition, Atmos. Meas. Tech. Discuss., 3, 599-636, doi:10.5194/amtd-3-599-2010, 2010.

Kurtén, T., Loukonen, V., Vehkamäki, H., and Kulmala, M.: Amines are likely to enhance neutral and ion-induced sulfuric acid-water nucleation in the atmosphere more effectively than ammonia, Atmos. Chem. Phys., 8, 4095-4103, doi:10.5194/acp-8-4095-2008, 2008.

Liu, Y., Gibson, E. R., Cain, J. P., Wang, H., Grassian, V. H., and Laskin, A.: Kinetics of heterogeneous reaction of CaCO<sub>3</sub> particles with gaseous HNO<sub>3</sub> over a wide range of humidity, J. Phys. Chem. A, 112, 1561–1571, doi:10.1021/Jp076169h, 2008.

Llovd, J. A., Heaton, K. J., and Johnston, M. V.: Reactive uptake of trimethylamine into ammonium nitrate particles, J. Phys. Chem. A, 113, 4840–4843, doi:10.1021/Jp900634d, 2009.

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- Loukonen, V., Kurtén, T., Ortega, I. K., Vehkamäki, H., Pádua, A. A. H., Sellegri, K., and Kulmala, M.: Enhancing effect of dimethylamine in sulfuric acid nucleation in the presence of water - a computational study, Atmos. Chem. Phys., 10, 4961-4974, doi:10.5194/acp-10-4961-2010, 2010.
- 5 Mackay, D., Shiu, W. Y., Ma, K. C., and Lee, S. C.: Aliphatic Amines, 2nd ed., Handbook of Physical-Chemical Properties and Environmental Fate for Organic Chemicals, Taylor & Francis Group, 3218-3230, 2006.
  - Mäkelä, J. M., Yli-Koivisto, S., Hiltunen, V., Seidl, W., Swietlicki, E., Teinilä, K., Sillanpää, M., Koponen, I. K., Paatero, J., Rosman, K., and Hämeri, K.: Chemical composition of aerosol during particle formation events in boreal forest, Tellus B, 53, 380-393, 2001.
  - Marple, V., Rubow, K., Ananth, G., and Fissan, H. J.: Micro-orifice uniform deposit impactor, J. Aerosol Sci., 17, 489-494, 1986.
  - Mmereki, B. T., Hicks, J. M., and Donaldson, D. J.: Adsorption of atmospheric gases at the air-water interface. 3: Methylamines, J. Phys. Chem. A, 104, 10789-10793, doi:10.1021/Jp0023258, 2000.
  - Mopper, K. and Zika, R. G.: Free amino acids in marine rains evidence for oxidation and potential role in nitrogen cycling, Nature, 325, 246–249, 1987.
  - Mosier, A. R., Andre, C. E., and Viets, F. G.: Identification of aliphatic-amines volatilized from cattle feedyard, Environ. Sci. Technol., 7, 642-644, 1973.
- Mozurkewich, M.: The dissociation-constant of ammonium-nitrate and its dependence on temperature, relative-humidity and particle-size, Atmos. Environ. A-Gen., 27, 261–270, 1993.
  - Müller, C., Iinuma, Y., Karstensen, J., van Pinxteren, D., Lehmann, S., Gnauk, T., and Herrmann, H.: Seasonal variation of aliphatic amines in marine sub-micrometer particles at the Cape Verde islands, Atmos. Chem. Phys., 9, 9587–9597, doi:10.5194/acp-9-9587-2009, 2009.

25

- Murphy, S. M., Sorooshian, A., Kroll, J. H., Ng, N. L., Chhabra, P., Tong, C., Surratt, J. D., Knipping, E., Flagan, R. C., and Seinfeld, J. H.: Secondary aerosol formation from atmospheric reactions of aliphatic amines, Atmos. Chem. Phys., 7, 2313-2337, doi:10.5194/acp-7-2313-2007, 2007.
- Pankow, J. F.: Gas/particle partitioning of neutral and ionizing compounds to single and multiphase aerosol particles. 1. Unified modeling framework, Atmos. Environ., 37, 3323-3333, doi:10.1016/S1352-2310(03)00346-7, 2003.
  - Pastor, S. H., Allen, J. O., Hughes, L. S., Bhave, P., Cass, G. R., and Prather, K. A.: Ambi-

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ent single particle analysis in Riverside, California by aerosol time-of-flight mass spectrometry during the SCOS97-NARSTO, Atmos. Environ., 37, S239-S258, doi:10.1016/S1352-2310(03)00393-5, 2003.

Pitts Jr., J. N., Grosjean, D., Van Cauwenberghe, K., Schmid, J. P., and Fitz, D. R.: Photooxidation of aliphatic amines under simulated atmospheric conditions: formation of nitrosamines, nitramines, amides, and photochemical oxidant, Environ. Sci. Technol., 12, 946–953, 1978.

Pratt, K. A., Hatch, L. E., and Prather, K. A.: Seasonal volatility dependence of ambient particle phase amines, Environ. Sci. Technol., 43, 5276-5281, doi:10.1021/Es803189n, 2009.

Rabaud, N. E., Ebeler, S. E., Ashbaugh, L. L., and Flocchini, R. G.: Characterization and quantification of odorous and non-odorous volatile organic compounds near a commercial dairy in California, Atmos. Environ., 37, 933-940, doi:10.1016/S1352-2310(02)00970-6, 2003.

Schade, G. W. and Crutzen, P. J.: Emission of aliphatic amines from animal husbandry and their reactions-potential source of N<sub>2</sub>O and HCN, J. Atmos. Chem., 22, 319–346, 1995.

Seinfeld, J. H. and Pandis, S. N.: Ammonia-water equilibrium, in: Atmospheric Chemistry and Physics: From Air Pollution to Climate Change, 2nd ed., John Wiley & Sons, Inc., Hoboken, NJ, p. 299, 2006.

Silva, P. J., Erupe, M. E., Price, D., Elias, J., Malloy, Q. G. J., Li, Q., Warren, B., and Cocker, D. R.: Trimethylamine as precursor to secondary organic aerosol formation via nitrate radical reaction in the atmosphere, Environ. Sci. Technol., 42, 4689-4696, doi:10.1021/Es703016v, 2008.

Smith, J. N., Dunn, M. J., VanReken, T. M., Iida, K., Stolzenburg, M. R., McMurry, P. H., and Huey, L. G.: Chemical composition of atmospheric nanoparticles formed from nucleation in Tecamac, Mexico: evidence for an important role for organic species in nanoparticle growth, Geophys. Res. Lett., 35, 5, 2008.

Smith, J. N., Barsanti, K. C., Friedli, H. R., Ehn, M., Kulmala, M., Collins, D. R., Scheckman, J. H., Williams, B. J., and McMurry, P. H.: Observations of aminium salts in atmospheric nanoparticles and possible climatic implications, P. Natl. Acad. Sci. USA, 107, 6634-6639, 2010.

Sorooshian, A., Murphy, S. M., Hersey, S., Gates, H., Padro, L. T., Nenes, A., Brechtel, F. J., Jonsson, H., Flagan, R. C., and Seinfeld, J. H.: Comprehensive airborne characterization of aerosol from a major bovine source, Atmos. Chem. Phys., 8, 5489-5520, doi:10.5194/acp-8-5489-2008, 2008,

Tan, P. V., Evans, G. J., Tsai, J., Owega, S., Fila, M. S., and Malpica, O.: On-line analysis of

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Size distribution of alkyl amines

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- urban particulate matter focusing on elevated wintertime aerosol concentrations, Environ. Sci. Technol., 36, 3512–3518, 2002.
- Van Neste, A., Duce, R. A., and Lee, C.: Methylamines in the marine atmosphere, Geophys. Res. Lett., 14, 711–714, 1987.
- Veltman, K., Singh, B., and Hertwich, E. G.: Human and environmental impact assessment of postcombustion CO<sub>2</sub> capture focusing on emissions from amine-based scrubbing solvents to air, Environ. Sci. Technol., 44, 1496–1502, doi:10.1021/Es902116r, 2010.
  - Violaki, K. and Mihalopoulos, N.: Water-soluble organic nitrogen (WSON) in size-segregated atmospheric particles over the Eastern Mediterranean, Atmos. Environ., 4339–4345, 2010.
- Vlasenko, A., Sjogren, S., Weingartner, E., Stemmler, K., Gäggeler, H. W., and Ammann, M.: Effect of humidity on nitric acid uptake to mineral dust aerosol particles, Atmos. Chem. Phys., 6, 2147–2160, doi:10.5194/acp-6-2147-2006, 2006.

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**Table 1.** Ammonium and alkyl amine detection sensitivities from the injection of known concentrations of the individual compounds in to the ICS-2000 system using the CS17 and CS12A columns. Ammonium sensitivity is reported as the k coefficient in Eq. (1), while amine sensitivities are reported as slopes since they show linear response in the 0.005–0.1  $\mu$ g mL<sup>-1</sup> range typical of samples collected online with the AIM-IC and offline with the MOUDI.

Amine	CS17	CS12A
species	Sensitivity (μS mole <sup>-1</sup> )	Sensitivity (μS mole <sup>-1</sup> )
NH <sub>3</sub>	2.57×10 <sup>7</sup>	1.74×10 <sup>7</sup>
MA	1.72×10 <sup>8</sup>	5.17×10 <sup>8</sup>
DMA	1.55×10 <sup>8</sup>	5.41×10 <sup>8</sup>
TMA	1.12×10 <sup>8</sup>	8.06×10 <sup>7</sup>
EA	1.71×10 <sup>8</sup>	4.16×10 <sup>8</sup>
DEA	1.92×10 <sup>8</sup>	1.12×10 <sup>8</sup>
TEA	1.62×10 <sup>8</sup>	2.40×10 <sup>8</sup>

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**Table 2.** AIM-IC online detection limits and sensitivity for small alkyl amines and ammonia using the CS17 column.

Amine	$3\sigma$ limits of $\sigma$		
species	moles	pptv	ng m <sup>-3</sup>
$NH_3$	$1.32 \times 10^{-09}$	180	130
MA	$1.74 \times 10^{-11}$	2.4	3.1
DMA	$1.93 \times 10^{-11}$	2.6	4.9
TMA	$2.68 \times 10^{-11}$	3.6	8.9
EA	$1.76 \times 10^{-11}$	2.4	4.5
DEA	1.56×10 <sup>-11</sup>	2.1	6.4
TEA	1.85×10 <sup>-11</sup>	2.5	10

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**Table 3.** MOUDI size-resolved particulate matter sample collection and analysis information. Eight samples were collected over the course of a year using either a rotating or fixed set of MOUDI stages. Mass loadings of  $PM_{1.8}$  amines in the particulate MOUDI samples are reported and then calculated as a function of measureable water soluble mass  $(m_{sol})$ , total mass  $(m_{tot})$ , as well as relative to  $PM_{1.8}$   $NH_4^+$ .

Sample	Location	Start date	Sample	Amines	R <sub>3</sub> NH <sup>+</sup>	$m_{sol}$	$m_{\rm tot}$	$NH_4^+$	R <sub>3</sub> NH <sup>+</sup>
#		dd/mm/yyyy	time (h)	detected	$(ng m^{-3})$	(%)	(%)	$(ng m^{-3})$	NH <sub>4</sub>
1 <sup>a</sup>	Toronto, ON	21 Jul 2009	24	DMA, TMA+DEA	16	0.78	0.31	470	0.014
2 <sup>a</sup>	Toronto, ON	7 Aug 2009	22	DMA, TMA+DEA, TEA	55	9.10	0.98	90	0.11
3 <sup>a</sup>	Toronto, ON	7 Aug 2009	22	DMA, TMA+DEA, TEA	35	7.52	0.68	90	0.17
4 <sup>a</sup>	Dorset, ON	14 Aug 2009	46	MMA, DMA, TMA+DEA, TEA	42	7.98	0.71	120	0.14
5 <sup>a</sup>	Toronto, ON	16 Aug 2009	24.25	DMA, TMA+DEA, TEA	81	0.42	0.24	5210	0.0045
6 <sup>a,b</sup>	Toronto, ON	16 Mar 2010	29	DMA, TMA+DEA, TEA	35	0.88		740	0.011
7 <sup>a,b</sup>	Lambton Shores, ON	9 Apr 2010	24.25	DMA, TMA+DEA	19	0.48	0.74	790	0.0065
8 <sup>a,b</sup>	Lambton Shores, ON	10 Apr 2010	24	DMA, TMA+DEA	33	0.58	0.45	1320	0.0067

<sup>&</sup>lt;sup>a</sup> Indicates CS17 separation method.

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b Indicates CS12A separation method.

**Table 4.** The maximum mass loading  $(ng\,m^{-3})$  on a single stage, for each amine species quantified in the MOUDI sample. The median diameter (nm) of the size bin where each species maximizes is given in parentheses when it is not the 320–560 nm size bin. No amines were consistently detected in all size bins from 0.056 to 18  $\mu$ m, so a range was not reported as the lower value would always be below the limit of detection. ND indicates the species was not detected on any stage of the MOUDI.

Sample #	MMA	DMA	TMA+DEA	TEA
1	ND	4	3	ND
2	ND	5	11	1.3
3	ND	3	7	0.3
4	0.05	6	5	0.4
5	ND	10	19	2 (780)
6	ND	2 (100)	10	14 (4400)
7	ND	0.8	5 (780)	ND
8	ND	0.4 (1400)	9	ND

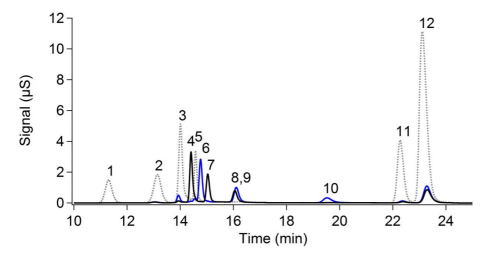
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**Fig. 1.** Overlaid chromatograms of mixed standard of  $0.05 \,\mu\text{g/mL}$  methyl amines (black trace), ethyl amines (blue trace) and inorganic cations (grey dashed line) separated on a Dionex CS17 column with active conductivity suppression with a CSRS-300: 1 – lithium, 2 – sodium, 3 – ammonium, 4 – methylamine, 5 – potassium, 6 – ethylamine 7 – dimethylamine 8 – trimethylamine, 9 – diethylamine, 10 – triethylamine, 11 – magnesium, 12 – calcium.

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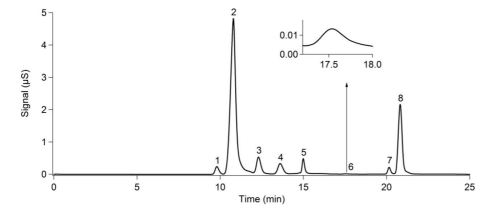
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**Fig. 2.** Sample CS17 chromatogram of particles collected on the 440±120 nm MOUDI stage at Dorset, ON. Peaks are: 1 – sodium, 2 – ammonium, 3 – potassium, 4 – dimethylamine, 5 – trimethlyamine and diethylamine, 6 – triethylamine, 7 – magnesium, 8 – calcium.

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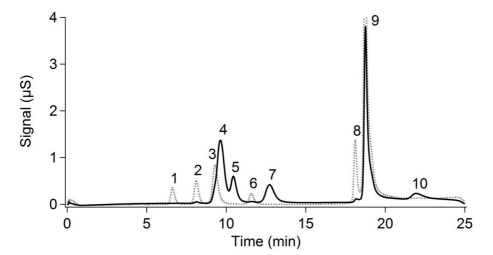
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**Fig. 3.** Overlaid chromatograms of mixed standard of  $0.05\,\mu\text{g/mL}$  methyl and ethyl amines (black trace), and inorganic cations (grey dashed line) separated on a Dionex CS12A column with active conductivity suppression with a CSRS-300: 1 – lithium, 2 – sodium, 3 – ammonium, 4 – methylamine and ethylamine, 5 – dimethylamine, 6 – potassium, 7 – trimethylamine and diethylamine, 8 – magnesium, 9 – calcium, 10 – triethylamine.

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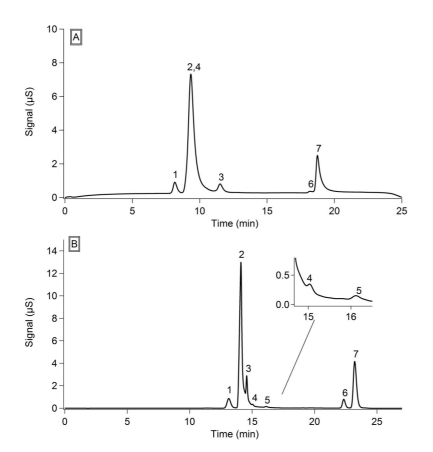


Fig. 4. Chromatograms of particles collected on the 780±200 nm MOUDI stage at Lambton Shores, ON separated on (A) CS12A and (B) CS17 columns. Compounds are 1 - sodium, 2 - ammonium, 3 - potassium, 4 - dimethylamine, 5 - trimethylamine and diethylamine, 6 magnesium, 7 - calcium.



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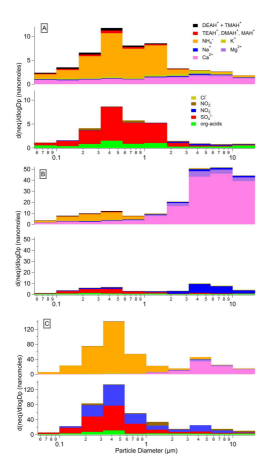
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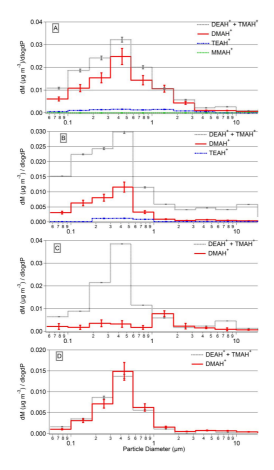
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**Fig. 5.** Mole equivalent distribution of major water-soluble ions in MOUDI samples. **(A)** Sample 4 collected in Dorset, **(B)** Sample 3 collected in Toronto, **(C)** Sample 8 collected in Lambton Shores.



**Fig. 6.** Amine mass distributions as a function of particle diameter for **(A)** sample 4 collected in Dorset, **(B)** sample 3 collected in Toronto, **(C)** sample 8 collected in Lambton Shores, and **(D)** sample 1 collected in Toronto.

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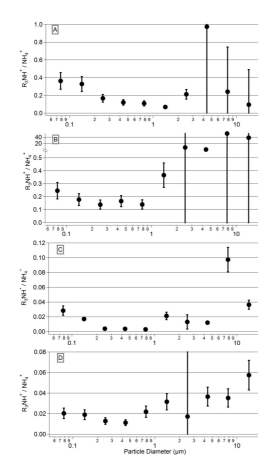
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**Fig. 7.** Ratio of the sum of all particulate amines (R<sub>3</sub>NH<sup>+</sup>) measured to the total ammonium (moles/moles) as a function of particle diameter for **(A)** Sample 4 collected in Dorset, **(B)** Sample 3 collected in Torontom **(C)** Sample 8 collected in Lambton Shores, and **(D)** Sample 1 collected in Toronto.

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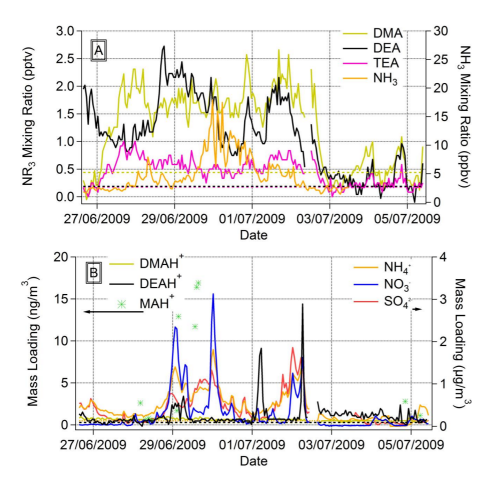
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**Fig. 8.** Online AIM-IC observations of amines and dominant inorganic constituents in **(A)** the gas phase and **(B)**  $PM_{2.5}$ . In each panel, the left axis applies to amines, and the right axis to inorganic consituents. Detection limits for amines are depicted using dashed lines.

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Size distribution of alkyl amines

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