Characterization of the sources and processes of organic and inorganic aerosols in New York city with a high-resolution time-of-flight aerosol mass spectrometer


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Abstract. Submicron aerosol particles (PM1) were measured in-situ using a High-Resolution Time-of-Flight Aerosol Mass Spectrometer during the summer 2009 Field Intensive Study at Queens College in New York, NY. Organic aerosol (OA) and sulfate are the two dominant species, accounting for 54% and 24%, respectively, of the total PM1 mass. The average mass-based size distribution of OA presents a small mode peaking at ∼150 nm (Dv80) and an accumulation mode (∼550 nm) that is internally mixed with sulfate, nitrate, and ammonium. The diurnal cycles of both sulfate and OA peak between 01:00–02:00 p.m. EST due to photochemical production. The average (±σ) oxygen-to-carbon (O/C), hydrogen-to-carbon (H/C), and nitrogen-to-carbon (N/C) ratios of OA in NYC are 0.36 (±0.09), 1.49 (±0.08), and 0.012 (±0.005), respectively, corresponding to an average organic mass-to-carbon (OM/OC) ratio of 1.62 (±0.11). Positive matrix factorization (PMF) of the high resolution mass spectra identified two primary OA (POA) sources, traffic and cooking, and three secondary OA (SOA) components including a highly oxidized, regional low-volatility oxygenated OA (LV-OOA; O/C = 0.63), a less oxidized, semi-volatile SV-OOA (O/C = 0.38) and a unique nitrogen-enriched OA (NOA; N/C = 0.053) characterized with prominent C8H24+2N+ peaks likely from amino compounds. Our results indicate that cooking and traffic are two distinct and mass-equivalent POA sources in NYC, together contributing ∼30% of the total OA mass during this study. The OA composition is dominated by secondary species, especially during high PM events. SV-OOA and LV-OOA on average account for 34% and 30%, respectively, of the total OA mass. The chemical evolution of SOA in NYC appears to progress with a continuous oxidation from SV-OOA to LV-OOA, which is further supported by a gradual increase of O/C ratio and a simultaneous decrease of H/C ratio in total OA. Detailed analysis of NOA (5.8% of OA) presents evidence that organic nitrogen species such as amines might have played an important role in the atmospheric processing of OA in NYC, likely involving both acid-base chemistry and photochemistry. In addition, analysis of air mass trajectories and satellite imagery of aerosol optical depth (AOD) indicates that the high potential source regions of secondary sulfate and aged OA are mainly located in regions to the west and southwest of the city.

1 Introduction

Aerosol particles play significant roles in climate change by altering the radiative balance of the Earth’s atmosphere directly and indirectly (IPCC, 2007). They also constitute a threat to public health by increasing the risk of morbidity and mortality of sensitive groups (Pope et al., 2002, 2009). Fine particulate matter (PM) in densely populated megacity...
environments are of particular concern for their adverse effects on human health and regional air quality (Molina and Molina, 2004). The New York City (NYC) metropolitan area is one of the most populous megacities in the world and among the most polluted cities in the US by fine PM and ozone (American Lung Association’s Report, 2010). It is well known that atmospheric aerosols are influenced by various emission sources and transformation processes. Characterizing the chemical composition and dynamic variations of aerosols in large urban environments such as NYC is important to unravel the complexities of anthropogenic aerosols and supporting health outcome studies (Demerjian and Mohnen, 2008; Wexler and Johnston, 2008).

A number of studies conducted in recent years characterized inorganic species (e.g., sulfate, nitrate, and ammonium), carbonaceous material (e.g., organic and elemental carbon; OC and EC), and trace metal species in aerosols in metropolitan NY (Schwab et al., 2004; Bae et al., 2006; Dutkiewicz et al., 2006; Qin et al., 2006; Venkatachari et al., 2006; Sunder Raman et al., 2008; Rattigan et al., 2010). Sulfate and OC usually dominate fine PM composition in the region. While sulfate is mainly contributed by regional transport, organic aerosol (OA) have contributions from both local and regional accumulation (Dutkiewicz et al., 2004; Lall and Thurston, 2006; Qin et al., 2006). Particulate organic materials can be classified as either primary OA (POA) from direct emissions, e.g., fossil fuel and biomass burning, or secondary OA (SOA) from the oxidation of gas-phase precursors (Kanakidou et al., 2005). SOA has been identified as a major contributor to the fine PM burden in NYC, especially under summertime conditions. Using the EC tracer method, Rattigan et al. (2010) estimated that SOA accounted for 63–73% and 40–50% of total OC during summer and winter, respectively, in South Bronx, NY. In addition, by analyzing ambient aerosol data acquired with an Aerodyne Aerosol Mass Spectrometer (AMS, Canagaratna et al., 2007), Zhang et al. (2007a) reported that oxygenated OA (OOA), a surrogate for SOA, on average accounted for 70% and 50% of total OA mass in Queens, NY during summer and winter, respectively.

Two AMS measurements were conducted in summer 2001 (Drewnick et al., 2004a, b) and winter 2004 (Weimer et al., 2006), respectively, on the campus of Queens College (QC) in NYC as part of the PM$_{2.5}$ Technology Assessment and Characterization Study-NY (PMTACS-NY) – one of the EPA supersite programs (Demerjian and Mohnen, 2008). These real-time, highly time-resolved measurements of the concentrations and size distributions of non-refractory submicron aerosol (NR-PM$_1$) species (i.e., organics, sulfate, nitrate, ammonium, and chloride) offered valuable insights into the chemistry, sources, and evolution processes of fine PM in NYC (Drewnick et al., 2004a, b; Weimer et al., 2006; Demerjian and Mohnen, 2008; Wexler and Johnston, 2008). An interesting observation was a distinct small particle mode (∼70 nm) of organics in summer that was externally mixed with sulfate and nitrate (Drewnick et al., 2004b). This mode appeared to be related to traffic emissions as it intensified during morning rush-hour (Drewnick et al., 2004a). In addition, significant differences in aerosol composition and size distributions were observed in winter at the same site due to different meteorological conditions and photochemical processing (Weimer et al., 2006). The AMS used during PMTACS-NY 2001 and 2004 was equipped with a Quadrupole mass spectrometer(i.e., Q-AMS; Jayne et al., 2000) which generates unit mass resolution (UMR) spectra. In summer 2009 we returned to the same site with a new advanced version of the AMS – a High-Resolution Time-of-Flight AMS (HR-ToF-AMS, termed as HR-AMS hereafter) (DeCarlo et al., 2006). Compared to the Q-AMS, the HR-AMS is significantly improved on chemical resolution and sensitivity. In particular, the high m/z resolution of the HR-AMS allows most ion fragments, especially those at low m/z’s (<100), to be resolved and their nominal elemental compositions (i.e., the ratios among carbon (C), hydrogen (H), oxygen (O), and nitrogen (N)) of OA to be determined (Aiken et al., 2008). Here we report the main findings from this study, including (1) evaluation of HR-AMS based on the comparisons of measurements with collocated instruments; (2) mass concentrations, size distributions, chemical composition, and temporal and diurnal variations of PM$_1$ species; (3) elemental composition of OA; and (4) characteristics and dynamic variations of OA components determined via Positive Matrix Factorization (PMF) of the high-resolution mass spectra (HRMS). Based on these results, the sources and processes of organic and inorganic aerosol components in NYC are also discussed.

2 Methods

2.1 Sampling site and instrumentation

This study took place from 13 July through 3 August 2009 on the campus of QC (40.74° N, 73.82° W, ∼25 m a.s.l.; Fig. S1 in the Supplement). The HR-AMS measurements were conducted inside the state-of-the-art Atmospheric Sciences Research Center Mobile Laboratory (ASRC-ML) (Schwab et al., 2010) along with various fast-response aerosol and gas instruments. These instruments include a DMT single-wavelength Photoacoustic Soot Spectrometer (PASS-1), a TSI Fast Mobility Particle Sizer (FMPS, Model 3091, 5.6–560 nm) spectrometer, a TSI water Condensation Particle Counter (CPC, Model 3781), an Aerodyne Quantum Cascade Laser (QCL) Spectrometer for measurements of formaldehyde and NO$_2$, a Li-COR CO$_2$ analyzer, a BTEX analyzer for benzene, toluene, ethylbenzene, and xylenes, and 2B technologies analyzers for O$_3$, NO, and NO$_2$. The ASRC-ML was parked at Lot 6 (Site B in Fig. S1) during this study except for two mornings on 28 and 30 July, and two evenings on 27 July and 1 August when it was moved to Lot 15 (Site C in Fig. S1) to measure near-road traffic plumes. Lot 6 is located...
approximately 500 m south to the Long Island Expressway (LIE, I-495) and 1.2 km east to the Van Wyck Expressway (I-678), two high-traffic highways in the NYC metropolitan area. Lot 15 is on the immediate south side of I-495 and ∼500 m to the northeast of Lot 6. During part of this study, an Aerodyne Mobile Laboratory was also deployed, in which a novel Soot Particle-AMS (SP-AMS) was operated alongside a number of aerosol and gas instruments.

In addition to measurements from the mobile facilities, aerosol and gas species were also measured inside a one-story building (Site A in Fig. S1) situated at ∼140 m north of Lot 6. This was the same sampling site for the PMTACS-NY 2001 summer and 2004 winter campaigns. Detailed descriptions of this site are given in Drewnick et al. (2004a) and Weimer et al. (2006). A key instrument deployed at Site A is a Particle-into-Liquid Sampler (PILS) coupled with two Metrohm Compact 761 Ion Chromatography (IC) systems (Herisau, Switzerland). The IC systems were equipped with an A Supp 5-250 and a C 4 column, respectively, for in-situ, simultaneous measurements of anions and cations at 30 min time interval. Ambient air was sampled into the PILS, in sequence, after a URG PM$_{2.5}$ cyclone, a VOC denuder, and two URG 2000 annular glass denuders coated with sodium carbonate and citric acid to remove acidic and basic gases, respectively. The IC systems were calibrated every 3–5 days, and the denuders were regenerated every week. In order to account for volatile losses, a correction factor of 1.14 (=0.88) suggested by Sorooshian et al. (2006) was applied for ammonium quantification.

In addition to the PILS-IC, other aerosol instruments deployed in the building include a Sunset Lab OC/EC Analyzer, a long- and a nano- Scanning Mobility Particle Sizer (SMPS), a Thermo Electron 5020 Sulfate Particulate Analyzer (SPA), a Tapered Element Oscillating Microbalance (TEOM), and an Aerodyne Aerosol Chemical Speciation Monitor (ACSM). Note that the Sunset Lab OC/EC Analyzer provides an optical measurement of EC (denoted as optical EC) by laser transmission in addition to the standard thermal measurement of EC (Rattigan et al., 2010). The optical EC shows tight correlation with the thermal EC ($r^2 = 0.86$, slope = 0.93) and also very good correlation with the light absorption coefficient ($B_{abs}$, $r^2 = 0.86$) from DMT PASS-I measurements. Here we use the optical EC due to its high time resolution (1-min) and hence a better characterization of aerosol evolution. All the EC data cited in this study refer to the optical EC unless otherwise noted, while the EC values quoted from other studies refer to the thermal EC.

All the data in this study are reported at ambient temperature and pressure (∼101 kPa) conditions in Eastern Standard Time (EST), which equals Coordinated Universal Time (UTC) minus 5 h or local time (i.e., East Daylight Time – EDT) minus 1 h.

### 2.2 HR-AMS operation

Ambient air was sampled isokinetically into the HR-AMS from a 1.0 inch (outer diameter) stainless steel tube with an inline PM$_{2.5}$ cyclone (URG-2000-30EH). The inlet of sampling line was positioned on the top of ASRC-ML at ∼2 m above the ground. The residence time of air between the inlet and the HR-AMS was estimated at ∼5 s. The HR-AMS was operated under the “V” and “W” ion optical modes alternatively every 5 min. The V-mode is more sensitive while the W-mode has higher mass resolution. Under V-mode operation, the AMS cycled through the mass spectrum (MS) mode and the particle time-of-flight (PTof) mode every 30 s, spending 10 s and 20 s, respectively, in each mode. Size distribution data are reported in units of mass-weighted aerodynamic diameter. No PTof data were sampled in W-mode due to limited signal-to-noise (S/N) ratio. However, the high mass resolution (∼5000–6000) of W-mode allows us to determine the ion-specific mass spectra and thus the elemental compositions of OA (DeCarlo et al., 2006; Aiken et al., 2008). The HR-AMS was calibrated for ionization efficiency (IE) and particle sizing at the beginning and in the middle of this study following the standard protocols (Jayne et al., 2000; Jimenez et al., 2003; Drewnick et al., 2005). The detection limits (DLs) of NR-PM$_1$ species were determined as 3 times the standard deviations (3σ) of the corresponding signals in particle-free ambient air through a HEPA filter (Zhang et al., 2005b). The 5-min DLs of organics, sulfate, nitrate, ammonium, and chloride are 57, 5, 4, 23, and 5 ng m$^{-3}$ respectively for V-mode and 110, 24, 30, 151, and 22 ng m$^{-3}$ respectively for W-mode, which are close to the values reported in previous HR-AMS studies (DeCarlo et al., 2006; Sun et al., 2009).

### 2.3 HR-AMS data analysis

#### 2.3.1 Standard data analysis

The mass concentrations and size distributions of NR-PM$_1$ species were analyzed using the standard AMS data analysis software (SQUIRREL v1.46, Sueper, 2010) written in Igor Pro 6.12A (WaveMetrics, Lake Oswego, OR). A collection efficiency (CE) was introduced to account for the incomplete detection of aerosol species due to particle bounce at the vaporizer ($E_b$) (Matthew et al., 2008) and/or partial transmission of particles by the lens (Huffman et al., 2005). Previous studies have shown that the particle loss is mainly due to $E_b$ which is primarily dependent on particle phase and composition (Matthew et al., 2008). A CE = 0.5 has been widely found to be representative for ambient particles in many AMS studies (Canagaratna et al. 2007 and references therein). High CE values were also observed for the ambient particles with high fraction of nitrate (Crosier et al., 2007; Matthew et al., 2008), high acidity (Quinn et al., 2006; Kleinman et al., 2007), high particle phase water content or
high organic liquid content (Matthew et al., 2008). Given that the aerosol particles in this study are overall neutralized (see Sect. 3.5.4) and that ammonium nitrate accounts for a minor fraction of total NR-PM$_1$ mass for most of time, a CE = 0.5 is used for the AMS quantitative analysis. The value of 0.5 was further validated based on inter-comparisons with collocated measurements (see Sect. 3.1) and is consistent with the CE values observed in previous AMS campaigns at the same site (Drewnick et al., 2003, 2004a; Hogrefe et al., 2004; Weimer et al., 2006). However, using a constant CE value may introduce an uncertainty of ~20–30%. Relative ionization efficiencies (RIEs) of 1.4 for organics, 1.2 for sulfate, 1.1 for nitrate, and 1.3 for chloride were used as previously reported (Allan et al., 2003b; Jimenez et al., 2003). The RIE for ammonium was determined at 4.0 based on the analysis of pure NH$_4$NO$_3$ particles during this study.

2.3.2 High resolution mass spectra analysis

The HRMS of both V- and W-mode were analyzed using the PIKA v1.06 – software toolkit downloaded at http://cires.colorado.edu/jimenez-group/ToFAMSResources/ToFSoftware/index.html. The W-mode spectra were analyzed to determine the elemental compositions of ion fragments, and subsequently the H/C, O/C, N/C, and OM/OC ratios of OA (Aiken et al., 2008). Since the organic H$_2$O$^+$, HO$^+$, O$^+$, and CO$^+$ signals were not measured directly due to large uncertainties, they were scaled according to the intensities of the organic CO$_2^+$ signal using the ratios suggested by Aiken et al. (2008), i.e., CO$^+$/CO$_2^+$. H$_2$O$^+$/CO$_2^+$. HO$^+$/CO$_2^+$. O$^+$/HO$_2^+$. Additionally, elemental analysis was also performed to the V-mode spectra using the same ion fragments identified for the W-mode spectra. Very similar O/C, H/C, and OM/OC ratios as those from W-mode were observed. However, the V-mode N/C ratios are on average a factor of 2 different from the W-mode results, due to larger uncertainties in quantifying C$_n$H$_y$N$_p^+$ ions using the lower resolution V-mode spectra.

2.3.3 PMF analysis of high resolution mass spectra

PMF analysis (Paatero and Tapper, 1994) was performed to the HRMS, i.e., the ion-spectiated W-mode spectra, using the PMF Evaluation Toolkit (PET) v2.02 (Ulbrich et al., 2009). The error matrix for PMF analysis was determined based on propagation of errors of Poisson counting statistics and electronic noise (DeCarlo et al., 2010). The peak fitting errors calculated from the residuals of PIKA fitting were also considered for small ions situated in close proximity to large ions. Ions with S/N ratio <0.2 were removed from the HRMS data and error matrices before PMF analysis. Excluding these noisy ions improves the differentiation of OA components, but has little impact on the mass concentrations since they together account for only ~2% of total OA signal.

The “weak” ions with S/N between 0.2 and 2 were down-weighted by increasing their errors by a factor of 2 (Paatero and Hopke, 2003; Ulbrich et al., 2009). In addition, H$_2$O$^+$, HO$^+$, O$^+$, and CO$_2^+$ were removed from the data and error matrices before the PMF analysis because they were determined according to the relationship with CO$_2^+$. Including these ions may introduce additional weight to CO$_2^+$ (Ulbrich et al., 2009). They were inserted back into the mass spectral matrix after the PMF analysis.

A summary of the PMF results is presented in Fig. S2. After an extensive evaluation of the mass spectral profiles and time series of different number of factors and the rotational forcing parameter, fPeak, the 5-factor solution with fPeak = 0 ($Q/Q_{\text{expected}} = 3.6$) was chosen. The OA components of the 5-factor solution solved under different fPeak values show very similar mass spectral patterns and time series (Fig. S2). The direct comparisons of the mass spectra and time series of 4-factor and 6-factor solution are shown in Fig. S3. The 4-factor solution does not resolve the small, yet distinct, fifth factor (~6% of total OA) (see Sect. 3.5.4 for details). The 6-factor solution splits the semi-volatile OA (SV-OOA) into two components for which we cannot offer a physically meaningful interpretation. We therefore choose the 5-factor solution. A detailed list of reasons for the selection of the 5-factor solution is given in Table S1.

2.4 Air mass trajectories and aerosol optical depth

3-day back trajectories arriving at QC were calculated every 1 h over the time period of this study using the National Oceanic and Atmospheric Administration (NOAA) HYSSPLIT 4.8 model (Draxler and Rolph, 2003) and the meteorological input from Air Resources Laboratory FNL data archive. 490 trajectories in total were obtained. These trajectories were then categorized into four clusters, i.e., cluster 1 from northwest (7.3% of time), cluster 2 from west (37.3% of time), cluster 3 from the Atlantic Ocean (24.5% of time), and cluster 4 from southwest (30.8% of time), using the trajectory cluster analysis of the HYSSPLIT 4.8 model.

Aerosol optical depth (AOD) was retrieved from the observations at wavelength of 550 nm made by National Aeronautics and Space Administration (NASA) Moderate Resolution Imaging Spectroradiometer (MODIS) onboard the Terra satellite. The Collection 5 level 2 data was used in this study (http://ladsweb.nascom.nasa.gov/browse_images/l2_browse.html?form=AADS&browseType=Level+2). The algorithm of retrieval and validation of retrieved AOD is described in detail in Remer et al. (2005).
IC measurements were close to 1:1 for both sulfate and nitrate. In 2001 and 2004 the correlations between Q-AMS and PILS-IC at New York City (Salcedo et al., 2006). Note that during PMTACS-NY 2005, Toronto, Canada (Buset et al., 2006), and Mexico City (Salcedo et al., 2006), Tokyo, Japan (Takegawa et al., 2005), and significant amounts of refractory-chloride are likely present in PM1, and does not respond to refractory components such as black carbon and crustal materials. An intercept of 3.91 µg m⁻³ for the linear regression of AMS total vs. TEOM mass is likely due to the losses of semi-volatile materials at 50 °C inside TEOM (Eatough et al., 2003). The AMS sulfate, nitrate, and ammonium account for ~71–76% of those in PM2.5 from PILS-IC measurements with r² ranging between 0.62–0.90. These comparisons are consistent with previous observations at other urban sites, e.g., Pittsburgh (Zhang et al., 2005b), Tokyo, Japan (Takegawa et al., 2005), Toronto, Canada (Buset et al., 2006), and Mexico City (Salcedo et al., 2006). Note that during PMTACS-NY 2001 and 2004 the correlations between Q-AMS and PILS-IC measurements were close to 1:1 for both sulfate and nitrate (Drewnick et al., 2003; Drewnick et al., 2004a; Weimer et al., 2006). However, a CE value of 0.43 was applied in those studies vs. CE = 0.5 used in this analysis.

While the chloride concentrations measured by the HR-AMS correlate with those by the PILS-IC (r² = 0.47, Fig. 1f'), the HR-AMS on average reports only ~35% of the chloride measured by PILS. In addition, there are times when the AMS-chloride is much lower than the PILS-chloride (e.g., 09:00 a.m.–20:00 p.m., 31 July, Fig. 1f). This is probably because the HR-AMS measures primarily NR-chloride (e.g., in the form of NH₄Cl) and is insensitive to refractory species such as NaCl and KCl at its vaporizer temperature of 600 °C. Indeed, significantly elevated signals of chloride (e.g., ³⁵Cl⁻ and ³⁷Cl⁺) are observed in the “closed” (or “background”) mass spectra acquired during this study, indicating the detection of considerable amounts of refractory-chloride. Refractory species, such as lead, evaporate slowly, yet continuously, on the AMS oven and thus show elevated signals in the “background” spectra (Salcedo et al., 2010). Given the proximity of NYC to the Atlantic Ocean, significant amounts of refractory-chloride are likely present in airborne PM1.

Although the HR-AMS organic concentrations show tight correlation with the OC concentrations measured by a Sunset Lab OC/EC Analyzer (r² = 0.79), the linear regression slope
of 2.59 is higher than the average OM/OC ratio of 1.62 determined via elemental analysis of the HRMS (Sect. 3.3). The typical OM/OC ratios observed at urban sites range ∼1.6–1.8 (Turpin and Lim, 2001; Takegawa et al., 2005; Zhang et al., 2005b; Bae et al., 2006; Aiken et al., 2008). Weimer et al. (2006) also observed relatively high slopes of 2.06–2.72 in winter 2004 at QC comparing Q-AMS vs. Sunset OC measurements. Possible reasons for this discrepancy include: (1) evaporative losses of semi-volatile species during the carbon analysis, which is consistent with the observation that semi-volatile organic species compose a large fraction of OA in NYC (see Sect. 3.5.1), (2) “over-correction” of the OC data using the blank filter values (Bae et al., 2006), and (3) underestimation of the CE value for organics. Note that in a previous AMS study in Pittsburgh, Zhang et al. (2005b) applied a CE value of 0.7 for organics and CE = 0.5 for inorganic species based on the observations that the smaller mode organics appeared to be externally mixed with SO4\(^{2-}\), NO3\(^{-}\) and NH4\(^{+}\) and that the small mode particles appeared to be primarily sooty combustion particles for which the transmission and detection is close to 1 in the AMS according to laboratory studies (Slowik et al., 2004). Although the chemically resolved size distributions of this study also suggested external mixtures of the small mode organics and the inorganics, we did not see clear indication that the CE values should vary significantly according to particle size or composition. For example, the correlations between AMS OM and Sunset Lab OC are independent on the mass fraction of the small mode organics or the fraction of hydrocarbon-like OA (Fig. S4). Given that using CE = 0.5 for inorganic species is supported well by the intercomparisons with other particle measurements (Fig. 1) and that the major fraction of the organics appears to be internally mixed with the inorganics, our decision of using CE = 0.5 for all species seems to be valid and the organic mass concentrations reported in this study are unlikely biased.

### 3.2 Mass concentration, composition, and diurnal variation of submicron aerosol particles

Figure 2 shows the time variations of total PM\(_1\) mass, the mass fractional contributions of individual species, and the meteorological conditions during the entire campaign. The PM\(_1\) concentration and composition vary dynamically with high aerosol loadings generally associated with southerly wind. Low PM\(_1\) loading periods typically occur with northwesterly wind. The total mass concentrations of PM\(_1\) (including EC) vary between 2.08 and 35.8 µg m\(^{-3}\) during this study. Organics frequently comprise the largest fraction of PM\(_1\) with sulfate being the second largest (Fig. 2d). On average, organics and sulfate account for 54.3% (24.5–96.5%) and 24.2% (0.7–54.8%), respectively, of PM\(_1\) mass (Table 1). The dominance of organics and sulfate is similar to previous observations at the same site in summer 2001 (Drewnick et al., 2004a). The average loading of PM\(_1\) is also similar between this study (11.7 µg m\(^{-3}\) and summer 2001 (12.5 µg m\(^{-3}\)) (Drewnick et al., 2004a; Weimer et al., 2006). However, on average sulfate is ∼31% lower and organics ∼9% higher in 2009 compared to 2001. The decrease in sulfate is consistent with the statewide reduction in SO\(_2\) emissions in New York over the years (e.g., by more than a factor of 2 from 2001 to 2009; http://www.dec.ny.gov/chemical/54358.html), for which the introduction of ultra low sulfur diesel fuel starting in 2006 is part of the mitigation measures. However, differences in wind patterns between the 2001 and 2009 study periods might have also played an important role for the concentration differences.

While nitrate overall represents a minor fraction of PM\(_1\) (4.2%), high fraction of nitrate (up to 30–40% of PM\(_1\)) are observed during some time periods (e.g., 02:00–04:00 a.m., 22 July, Fig. 2). The average EC concentration of 0.70 µg m\(^{-3}\) (6% of PM\(_1\)) is close to the values observed in NYC previously, e.g., 0.75 ± 0.20 µg m\(^{-3}\) from 2002–2004 at QC (Bae et al., 2006) and 0.5–1.4 µg m\(^{-3}\) from 2006–2008 at South Bronx (Rattigan et al., 2010). In comparison, the EC concentrations observed in rural NY were typically 2–4 times lower, e.g., 0.18 ± 0.20 µg m\(^{-3}\) at Pinnacle State Park (Bae et al., 2006) and 0.31 and 0.36 µg m\(^{-3}\), respectively, at Potsdam and Stockton (Sunder Raman et al., 2008). A weekday vs. weekend comparison of the concentrations of NR-PM\(_1\) species shows almost no difference (1.5%) for sulfate yet ∼8–28% lower concentrations for the other NR species during weekend. EC shows the largest decrease (on average 35%) during weekend, similar to the observations made in South Bronx (Rattigan et al., 2010). The average diurnal cycles of aerosol species are shown in Fig. 3. Organics show a small early morning peak between 05:00–06:00 a.m. (i.e., 06:00–07:00 a.m. local time) and a distinct early afternoon peak between 12:00–14:00 p.m. The morning peak correlates with that of EC, which is primarily due

### Table 1. Summary of the mass concentrations (µg m\(^{-3}\)) of NR-PM\(_1\) species, OA components, and EC.

<table>
<thead>
<tr>
<th>Species</th>
<th>Average</th>
<th>σ</th>
<th>Median</th>
<th>Minimum</th>
<th>Maximum</th>
</tr>
</thead>
<tbody>
<tr>
<td>Organics</td>
<td>6.34</td>
<td>3.2</td>
<td>5.76</td>
<td>1.03</td>
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<td>1.38</td>
<td>1.72</td>
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<td>0.06</td>
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<td>0.03</td>
<td>0.02</td>
<td>&lt;D. L.</td>
<td>0.20</td>
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<td>0.54</td>
<td>0.57</td>
<td>&lt;D. L.</td>
<td>6.21</td>
</tr>
<tr>
<td>Totalb</td>
<td>11.7</td>
<td>5.43</td>
<td>11.1</td>
<td>2.08</td>
<td>35.8</td>
</tr>
</tbody>
</table>

* Detection Limit.
* Total = organics + sulfate + nitrate + ammonium + chloride + EC.
to local traffic emissions. The early afternoon peak of organics is mostly contributed by oxygenated species as the result of photochemical formation of SOA and by emissions of POA from cooking-related activities as well (Sect. 3.5.3). The high concentration of organics at night is likely due to increased traffic from heavy duty diesel trucks (to avoid commuter traffic; Venkatachari et al., 2006) compounded with less dilution due to shallow boundary layer. Indeed, substantially elevated EC (Fig. 3a) and HOA – a surrogate for traffic related POA (Sect. 3.5.2) – are clearly seen at night.

Sulfate shows an early afternoon peak at \( \sim 13:00–14:00\) p.m., similar to that observed in summer 2001 (Drewnick et al., 2004a). In order to investigate the source of this peak, the diurnal production rate of sulfuric acid was estimated based on the gas phase bi-molecule association reaction of \( \text{SO}_2 + \text{OH} + M \rightarrow \text{HOSO}_2 + M \). The OH concentration data used for the calculation was acquired in the same month during PMTACS-NY 2001 at the same location (Ren et al., 2004). The calculation formula is presented in Appendix A and the diurnal profile of the production rate of sulfuric acid is shown in Fig. 4a, in which gaseous CO\(_2\) is used as an indicator of dilution associated with changes in boundary layer height. The gas phase production rate of sulfuric acid peaks at early afternoon as sulfate does, counteracting the dilution associated with higher boundary layer that clearly leads to a gradual drop in the concentrations of primary emission species (e.g., CO\(_2\), NO\(_x\), EC, and HOA) during daytime. The integrated increase of sulfate due to gas phase production of \( \text{H}_2\text{SO}_4 \), which is estimated at \( \sim 0.35 \text{ µg m}^{-3} \) from 08:00 a.m. to 14:00 p.m., is much lower than the observed \( \sim 1 \text{ µg m}^{-3} \) increase of sulfate (Fig. 4a), indicating that regional transport and/or aqueous processes may have played a significant role controlling sulfate concentration in NYC. Indeed, the relatively flat diurnal pattern and a general trend of multi-day build-up signify the regional influences on sulfate in NYC, consistent with the regional characteristics of sulfate in the northeastern US (Dutkiewicz et al., 2004; Zhang et al., 2005b; Qin et al., 2006).

Both nitrate and chloride show similar diurnal cycles with peak concentrations appearing in early morning (04:00–07:00 a.m., Fig. 3d, f). The diurnal cycle of nitrate is mainly driven by HNO\(_3\) production, gas-to-particle partitioning to form ammonium nitrate, and boundary layer dynamics (Seinfeld and Pandis, 2006). Figure 4b shows the production rate of HNO\(_3\) estimated from the reaction of \( \text{NO}_2 + \text{OH} + M \rightarrow \text{HONO}_2 + M \), which is the main formation mechanism of nitrate during daytime, and the diurnal profile of the equilibrium constant \( K_p \) of \( \text{NH}_3(g) + \text{HNO}_3(g) \leftrightarrow \text{NH}_4\text{NO}_3(s) \) (see Appendix A for calculations). Note that the formation of \( \text{NH}_4\text{NO}_3(s) \) is also controlled by the acidity of particles as acidic particles retain very little nitrate (Zhang et al., 2007b). However, since PM\(_1\) appears to be bulk neutralized for a large fraction of time during this study (Sect. 3.5.4), the overall influence of aerosol acidity on ammonium nitrate production is very small and the gas-to-particle partitioning of \( \text{NH}_4\text{NO}_3 \) is primarily controlled by temperature and RH.

Fig. 2. Time series of (a) relative humidity (RH), temperature (T), and hourly precipitation (Precip), (b) wind direction (WD) colored by wind speed (WS), (c) mass concentration of AMS total + EC, and (d) mass fractions of chemical species in PM\(_1\).
Fig. 3. Diurnal profiles of the mass concentrations (solid black circles) and the size distributions of EC, organics, sulfate, nitrate, ammonium, and chloride.

Fig. 4. Diurnal cycles of (a) sulfate, estimated gas phase production rate of H₂SO₄, and CO₂ and (b) nitrate, estimated gas phase production rate of HNO₃, equilibrium constant of K_p for ammonium nitrate, and CO₂.

Despite intense daytime production of HNO₃ (at peak rate of 5.6 µg m⁻³ h⁻¹, Fig. 4b), the diurnal cycle of nitrate shows overall similar variation pattern as that of K_p, which decreases during daytime and reaches minimum at ~02:00 p.m. EST. However, the higher concentrations of nitrate at night (Fig. 4b), as well as the occurrence of several nighttime elevated nitrate events (Fig. 2d), may also be due to nighttime heterogeneous hydrolysis of N₂O₅ – a reaction product of NO₂ and nitrate radical (Brown et al., 2003). The hydrolysis processes is favored by the high RH at night. These results suggest that gas-to-particle partitioning of HNO₃ and nocturnal processing control the diurnal variation of nitrate in NYC. Similar diurnal pattern of nitrate was observed in Pittsburgh (Zhang et al., 2005b), Ontario, Canada (Rupakheti et al., 2005) and at the same site in summer 2001 (Drewnick et al., 2004a). However, significantly different patterns were observed in both Mexico City (Salcedo et al., 2006; Aiken et al., 2009) and Beijing in 2006 (Sun et al., 2010), where daytime photochemical production of HNO₃ is much faster likely due to considerably higher NOₓ emissions. The diurnal cycle of ammonium is very similar to the sum of sulfate and nitrate as it presents mainly in the form of ammonium sulfate and ammonium nitrate.

3.3 Bulk composition and elemental ratios of OA

Figure 5 shows the average HRMS of OA based on the contributions of elements and ion categories. The m/z 44 peak, mainly contributed by CO₂⁺ (92%, Fig. 5c), is the base peak in the spectrum. m/z 57 (~2% of OA), which has been suggested as an AMS spectral tracer for HOA (Zhang et al., 2005a), is also a significant peak. However, on average only 70% of signal at m/z 57 is the hydrocarbon ion – C₄H₉⁺ and the rest 30% is associated with C₃H₃O⁺ (Fig. 5c). The bulk mass-based composition of OA in NYC is dominated by carbon (62.1%) and oxygen (29.3%) with a minor contribution from hydrogen (7.7%) and nitrogen (0.8%). All of the elements peak at ~01:00 p.m. due to photochemical production and enhanced cooking emissions despite deeper boundary layer (Fig. S5). A small morning peak of C, H, and N occurs between ~05:00–06:00 a.m. due to local traffic emissions and shallow boundary layer, consistent with that of EC. In contrast, the average mass fractions of elements show relatively flat diurnal patterns except for slightly higher contributions of C in the morning and of O during the afternoon (Fig. 5d). On a daily basis, the O/C ratio of PM₁ varies between 0.31–0.40 and OM/OC between 1.56–1.67 (Fig. 5e).

The average (±1σ) OM/OC ratio for the entire study is 1.62 (±0.11), which is consistent with the value of 1.6 (±0.2) suggested for urban aerosols (Turpin and Lim, 2001). The OM/OC ratios determined from a multi-year study at QC by comparing the reconstructed PM₂.₅ mass and OC vary between 1.27 and 2.01 (Bae et al., 2006). The O/C ratio (0.36 ± 0.09) of this study is also consistent with AMS observations at other urban sites, e.g., Mexico City (Aiken et al., 2008), Tokyo (Takegawa et al., 2005), and London (Allan et al., 2010). The OM/OC and O/C ratios both increase gradually from morning to late afternoon due to enhanced photochemical SOA formation (Fig. 5e). The H/C ratio (1.49 ± 0.08) shows an opposite diurnal pattern with a
Fig. 5. Average OA spectrum colored by the contributions of (a) elements (C, O, H, and N) and (b) six ion categories. (c) The ion compositions of m/z 44 and 57; (d) diurnal profiles of the mass fractions of elements; and (e) diurnal profiles of organics, O/C and OM/OC ratios. The inset pie charts in (a) and (b) show the average mass fractions of elements and ion categories, respectively. The average elemental and OM/OC ratios of OA are also shown in the legend of (a).

significant morning peak due to local traffic influence. While the average N/C ratio of 0.012 (±0.004) is similar to the values observed from previous HR-AMS studies (DeCarlo et al., 2008; Aiken et al., 2009; Sun et al., 2009), periods with much higher N/C ratio (∼0.03–0.04) are also observed, likely due to the formation of N-containing organic compounds (Sect. 3.5.4).

3.4 Chemically-resolved size distributions of submicron aerosol particles

The average size distributions of aerosol species and the size-resolved aerosol composition for the entire campaign are shown in Fig. 6. We derived the size distribution of EC based on m/z 57 after removing the contribution of C₃H₅O⁺ (Fig. 7), assuming that the distribution pattern of EC mirrors that of C₄H₇⁺ – a dominant hydrocarbon ion at m/z 57 (Fig. 5c). The rationales behind this assumption are: (1) C₃H₅O⁺ (or m/z 57 of the unit resolution AMS data) is an AMS spectral tracer for HOA (Canagaratna et al., 2004; Zhang et al., 2005a; Aiken et al., 2008) and (2) HOA is a surrogate for combustion-related POA in urban areas (Zhang et al., 2005c, 2007a; Jimenez et al., 2009; Ulbrich et al., 2009; Allan et al., 2010; Ng et al., 2010). Indeed, C₃H₅O⁺ correlates well with EC (r² = 0.42) and NO₂ (r² = 0.61). The assumption is also supported by the similar size distributions between m/z 57 and BC from the SP-AMS measurements of exhaust plumes of heavy duty trucks and Metropolitan Transportation Authority (MTA) standard buses during this campaign (Massoli et al., 2010). However, since C₃H₅O⁺
Fig. 7. Diurnal evolution of the size distributions of (a) m/z 44, (b) m/z 57, and (c) C₄H₉⁺. (d) The average size distributions of m/z 57, m/z 44, C₄H₉⁺, and sulfate for entire study.

contributes ~30% of the m/z 57 signal during this study, the size distributions of m/z 57 may be influenced by oxygenated species that are secondary in origin. The poor correlation between C₃H₅O⁺ and C₄H₉⁺ (r² = 0.14) indicates their differences in sources. Similarly, Liggio et al. (2010) observed at a rural site of Ontario, Canada that the ratios of C₃H₅O⁺ to C₄H₉⁺ in aerosols increased with the photochemical age, suggesting the association of C₃H₅O⁺ with SOA or oxidized POA.

We estimated the size distribution of C₃H₅O⁺ according to that of m/z 44, which is an AMS spectral tracer for OOA (Zhang et al., 2005a). C₃H₅O⁺ correlates well with m/z 44 (r² = 0.73) and moderately with sulfate (r² = 0.29), a well-known secondary inorganic aerosol species. The size distributions of m/z 44 and sulfate also closely resemble each other (Figs. 7 and 3c). Note that upon these treatments, the accumulation mode of EC may nevertheless be overestimated because of the contribution of OOA to C₄H₉⁺ (∼10% of the total C₄H₉⁺ signals on average, Sect. 3.5.2), especially in late afternoon when SOA production is intense. A support for this possibility is the better correlation of C₄H₉⁺ with EC (r² = 0.60) and NOₓ (r² = 0.76) during morning rush hours (04:00–08:00 a.m., EST or 05:00–09:00 a.m local time) than other times of day.

All aerosol species show an overlapping accumulation mode peaking at ~550 nm in vacuum aerodynamic diameter (Dva) (Fig. 6a). This mode dominates the size distributions of secondary aerosol species, i.e., sulfate, nitrate and ammonium, and m/z 44 as well. In addition, it usually persists throughout the day with relatively little variations in intensity (Fig. 3). However, a mode peaking at 100–150 nm shows up in the size distributions of organics, EC (Fig. 6), and m/z 57 (Fig. 7b), suggesting that primary carbonaceous particles are externally mixed with secondary species. Similar phenomena were observed at various other urban sites (Allan et al., 2003a; Alfarra et al., 2004; Drewnick et al., 2004b; McFiggans et al., 2005; Zhang et al., 2005b; Takegawa et al., 2006; Weimer et al., 2006; Huang et al., 2010; Sun et al., 2010) and in a chamber study on mixing and partitioning of semi-volatile SOA and POA related to vehicle emission (Asa-Awuku et al., 2009).

During this study, ultrafine particles (<100 nm) are almost completely composed of carbonaceous material with EC contributing 24% of total mass and OA contributing an average 73%. EC is only a minor mass fraction (<5%) of particles larger than 200 nm. The accumulation mode aerosols are mainly composed of organics and sulfate, which together account for ~80% of the total mass.

3.5 Determination of OA components and characterization of their sources and processes

PMF analysis of the HRMS of OA identified five OA components, each of which demonstrates distinct temporary variation and mass spectral patterns (Fig. 8) that are indicative of their associations with unique sources and processes. In brief, the OA components include: (1) a highly oxidized and low-volatility OOA (LV-OOA; O/C = 0.63), (2) a less oxidized and semi-volatile OOA (SV-OOA; O/C = 0.38), (3) a nitrogen-enriched OA (NOA) with a much higher N/C ratio (0.0523) than other OA components (~0.004–0.011), (4) a cooking-emission related OA (COA) which shows spectral features similar to those of POA from cooking emissions and a unique diurnal pattern peaking during lunch and dinner times, and (5) a hydrocarbon-like OA component (HOA) that represents POA from incomplete fossil fuel combustion given its low O/C ratio (0.06) and good correlation with combustion emission tracers such as NOₓ and EC. Detailed discussions on each component, including the mass spectral profiles, correlations with the tracer species, temporal variation patterns, and associations with different sources and processes, are given in the following subsections.
3.5.1 Semi-volatile and low-volatility OOA (SV-OOA & LV-OOA)

The mass spectrum of LV-OOA (Fig. 8a) is characterized by a dominant peak at m/z 44 (CO$_2^+$), similar to the more oxidized OOA-1 component determined at other urban sites (Lanz et al., 2007; Ulbrich et al., 2009; Sun et al., 2010). Its O/C and OM/OC ratios (0.63 and 1.96, respectively) are in the range of the values of LV-OOA and total OOA (when LV- and SV-OOA are not separated) observed worldwide (Jimenez et al., 2009; Ng et al., 2010). The mass spectrum of SV-OOA (Fig. 8b), which has lower O/C ratio (0.38) and higher fraction of m/z 43 ($f_{43} = 9.3\%$, mainly C$_2$H$_3$O$^+$) compared to LV-OOA, tracks the spectral pattern of previously reported SV-OOA (Morgan et al., 2010; Ng et al., 2010). In addition, the spectrum of SV-OOA resembles those of biogenic SOA observed in ambient atmosphere (Chen et al., 2009; Slowik et al., 2010) and chamber experiments (Kiendler-Scharr et al., 2009; Shilling et al., 2009; Chhabra et al., 2010).

A survey of the correlations between the five OA components and individual ions in the HRMS is shown in Fig. 9. LV-OOA shows most significant correlation with oxygenated ions containing two O atoms (C$_4$H$_7$O$_2^+$) followed by C$_6$H$_7$O$_4^+$, and very weak correlation with C$_4$H$_3$N$^+$. On average LV-OOA accounts for 51% and 38% of C$_4$H$_7$O$_2^+$ and C$_6$H$_7$O$_4^+$, respectively, for the entire study (Fig. 10). Apportionment of the ions of six major m/z’s (Fig. 11) further reveals that that LV-OOA accounts for 53% of CO$_2^+$ (m/z 44), 61% of C$_2$H$_2$O$_2^+$ (m/z 58), and 46% of C$_2$H$_4$O$_2^+$ (m/z 60). Compared to LV-OOA, SV-OOA shows even tighter correlations with C$_6$H$_7$O$_4^+$ and slightly less, yet still high, correlations with C$_4$H$_7$O$_2^+$. It also correlates much better with C$_4$H$_3$N$^+$ ions than LV-OOA does. On average, SV-OOA contributes 42% of the C$_x$H$_y$O$_z^+$ signal and 31% of the C$_x$H$_y$O$_z^+$ (Fig. 10). These results clearly indicate that the chemical compositions of the two OOA types are significantly different: SV-OOA comprises less oxygenated, possibly freshly oxidized species, while LV-OOA comprises highly oxidized species including di- and poly-carboxylic acids.

LV-OOA correlates well with sulfate ($r^2 = 0.66$, Fig. 8a and Table 2), consistent with previous studies at various sites showing that these two species are secondary in nature and mostly driven by regional production (Zhang et al., 2005c; Lanz et al., 2007; Ulbrich et al., 2009; Sun et al., 2010). SV-OOA in general correlates with chloride and nitrate (Fig. 8b), indicating its semi-volatile characteristics (Lanz et al., 2007; Ulbrich et al., 2009; Sun et al., 2010).
correlation coefficient between SV-OOA and nitrate appears low ($r^2 = 0.05$), mainly due to the occurrence of several nighttime nitrate plumes (e.g., 22 July) likely from nocturnal processing of N$_2$O$_5$. For instance, upon excluding the nitrate plume on July 22 (5% of total data points), the correlation between SV-OOA and nitrate improves to $r^2 = 0.17$. In comparison to the tight correlation between LV-OOA and sulfate, the relatively weaker correlation between SV-OOA and chloride/nitrate might also be due to the contribution of biogenic OA to SV-OOA in summer. Total, OOA (=LV-OOA + SV-OOA) correlates well with total secondary inorganic species (=sulfate + nitrate + chloride, $r^2 = 0.50$, Fig. 12a), marking the link between OOA and SOA. The ratio of total OOA to sulfate in NYC is 1.29 (Table 2), much higher than 0.38 observed in Pittsburgh (Zhang et al., 2005c), highlighting the importance of SOA contribution to fine PM loading in NYC. Total OOA does not show much correlation with O$_x$ (=O$_3$ + NO$_2$, $r^2 = 0.08$), similar to the observations made in Pittsburgh (Zhang et al., 2005c).

**Fig. 9.** Correlations of each OA component with the HRMS ions colored by four ion categories, i.e., $C_xH_y^+$, $C_xH_yO_1^+$, $C_xH_yO_2^+$, and $C_xH_yN_p^+$. The formulas of the ions that show significant correlations with individual OA components are marked.

**Table 2.** Correlations between OA components and tracers from collocated measurements.

<table>
<thead>
<tr>
<th></th>
<th>$r^2$</th>
<th>Slope</th>
<th></th>
<th>$r^2$</th>
<th>Slope</th>
</tr>
</thead>
<tbody>
<tr>
<td>HOA vs. EC</td>
<td>0.44</td>
<td>1.29$^a$</td>
<td>SV-OOA vs. Chloride</td>
<td>0.30</td>
<td>48.9$^a$</td>
</tr>
<tr>
<td></td>
<td>0.72</td>
<td>1.41$^b$</td>
<td>Total OOA vs. Sulfate</td>
<td>0.51</td>
<td>1.29$^a$</td>
</tr>
<tr>
<td></td>
<td>0.65</td>
<td>1.25$^c$</td>
<td></td>
<td>0.74</td>
<td>0.38$^b$</td>
</tr>
<tr>
<td>HOA vs. NO$_x$</td>
<td>0.59</td>
<td>0.045$^a$</td>
<td>Total OOA vs. O$_x$</td>
<td>0.08</td>
<td>0.11$^a$</td>
</tr>
<tr>
<td></td>
<td>0.82</td>
<td>0.042$^b$</td>
<td></td>
<td>0.55</td>
<td>0.12$^d$</td>
</tr>
<tr>
<td>LV-OOA vs. Sulfate</td>
<td>0.66</td>
<td>0.65$^a$</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

$^a$ NYC, This study.
$^b$ Pittsburgh Zhang et al. (2005c).
$^c$ BC, Mexico City Aiken et al. (2009).
$^d$ Mexico City Aiken et al. (2009).
LV-OOA accounts for 30.3% of the OA mass on average, with the average contribution up to 46% during late afternoon and down to 22% at night (Fig. 13). SV-OOA accounts for an average 34% of the OA mass, showing a major contribution (~50%) in early morning. LV-OOA and SV-OOA together account for 64% of OA on average, similar to the estimated SOA contribution (63–73%) in NYC using the EC tracer method (Rattigan et al., 2010). In addition, OOA in total accounts for 81% of the O mass, 58% of the C mass, 52% of the H mass, and 49% of the N mass in OA (Fig. S6).

LV-OOA displays a pronounced diurnal cycle that is characterized by a gradual increase in concentration from ~09:00 a.m. to ~17:00 p.m. (Fig. 13a). SV-OOA also starts to increase at ~09:00 a.m. but peaks at ~13:00 p.m. The decrease of SV-OOA in the afternoon is likely due to higher ambient temperature that evaporates more semi-volatile species and the rising boundary layer, counteracting the photochemical production of SOA. Since the decrease of SV-OOA is accompanied with a gradual increase of LV-OOA, it is also possible that further oxidation of less oxidized, more-volatile species to highly oxidized, less-volatile species has occurred. Indeed, the O/C ratio of total OAOA gradually increases from 0.49 to 0.54 between 09:00 a.m. and 17:00 p.m. The continuous oxidation of SV-OOA to LV-OOA was observed at a remote coastal site in Europe as well (Hildebrandt et al., 2010).

### 3.5.2 Hydrocarbon-like OA (HOA)

The mass spectrum of HOA (Fig. 8e), which is characterized with the prominent C$\text{C}_x\text{H}_y^+$ and C$_{x\text{H}}^{2x+1}$ ion series, is very similar to previously reported reference spectra of POA emitted from gasoline and diesel combustion sources (Canagaratna et al., 2004; Schneider et al., 2006) and the spectra of HOA components determined at other sites (Zhang et al., 2005a; Aiken et al., 2009). The O/C (0.06) and OM/OC (1.24) ratios of HOA are also close to the values (0.03–0.04 and 1.22–1.24) determined for POA in motor vehicle exhaust (Mohr et al., 2009). As shown in Fig. 9e, HOA correlates best with hydrocarbon ions C$_{x\text{H}}^{2x+1}$ and C$_{x\text{H}}^{2x-1}$. It contributes 25% of the C$_{x\text{H}}^{2+}$ signal and only 3% of the C$_{x\text{H}}^{+}$ and the C$_{x\text{H}}^{2+}$ signals (Fig. 10). In addition, HOA correlates well with combustion tracers such as EC ($r^2 = 0.44$) and NO$_x$ ($r^2 = 0.59$) (Fig. 8e and Table 2) and presents a pronounced morning peak associated with traffic emissions (Fig. 13a). These facts together indicate that HOA is likely a surrogate for combustion POA, a conclusion achieved in a number of other studies (Zhang et al., 2005a, c; Aiken et al., 2009; Ulbrich et al., 2009). HOA on average accounts for 14.2% of the total OA mass for the whole campaign, showing the highest contribution (~28%) in the morning and the lowest (~8%) in the afternoon (Fig. 13b). As summarized in Table 2, the average ratios of HOA/EC (1.29) and HOA/NO$_x$ (0.043 µg m$^{-3}$ ppbv$^{-1}$) observed in this study are similar to the values reported for Pittsburgh (Zhang et al., 2005c) and Mexico City (Aiken et al., 2009).

### 3.5.3 Cooking OA (COA)

Although emissions related to cooking activities have been observed to contribute a significant fraction (~5–20%) of OA at various urban sites (Rogge et al., 1991; Zheng et al., 2002; Chow et al., 2007), reports of COA from AMS measurements are rare (Allan et al., 2010; Huang et al., 2010). In this study, a COA component was identified. The HRMS spectrum of COA is significantly different from those of other OA components but very similar to the reference spectra of POA from cooking emissions (Mohr et al., 2009) as well as the spectra of COA components observed in London (Allan et al., 2010) and Beijing (Huang et al., 2010).

The COA spectrum is characterized by a high ratio of m/z 55/57 (≈2.9) and a high fraction of m/z 55 (f$_{55} = 8\%$), which could be used as a diagnostic indicator for the presence of COA at urban locations. As shown in Fig. 11, both m/z 55 and 57 have significant contributions from COA, especially their oxygenated ions. In specific, COA contributes 38% and 29%, respectively, of C$_3\text{H}_5\text{O}^+$ and C$_4\text{H}_7^+$ at m/z 55 and 25% and 20%, respectively, of C$_3\text{H}_5\text{O}^+$ and C$_4\text{H}_7^+$ at m/z 57. In addition, COA shows the most significant correlation with a few C$_x\text{H}_y\text{O}_z^+$ ions (e.g., C$_5\text{H}_8\text{O}^+$, C$_6\text{H}_{10}\text{O}^+$, and C$_7\text{H}_{12}\text{O}^+$ in Fig. 9d), all of which are prominent peaks in the sources spectra of cooking emissions, thus could be used as spectral markers for COA.

The HRMS of COA reveals that it is on average more oxidized than HOA (O/C=0.18 vs. 0.06), containing much higher contribution of oxygenated ions C$_x\text{H}_y\text{O}_4^+$ and C$_x\text{H}_y\text{O}_3^+$ (30% vs. ~9%), thus higher O content mass
Fig. 11. Average contribution of each OA component to the ions at m/z (a) 43, (b) 44, (c) 55, (d) 57, (e) 58, and (f) 60. Note that only the major ions at each m/z are shown.

Fig. 12. Correlations of (a) total OOA (=LV-OOA + SV-OOA) vs. total secondary inorganic anions (=Sulfate + Nitrate + Chloride). The data fitting was performed using the orthogonal distance regression (ODR), and (b) total OOA vs. POA (=HOA + COA), colored by the mass fraction of OA in NR-PM$_1$. The marker size is proportional to the concentration of total NR-PM$_1$.

Fig. 13. Diurnal cycles of (a) mass concentrations and (b) mass fractions of five OA components. The dash line in (b) shows the mass fraction of POA (HOA + COA).
(10% vs. 3%), consistent with the detection of more oxygenated organic species in meat cooking emissions (Mohr et al., 2009). The O/C and H/C ratios of COA are 0.18 and 1.58, respectively (Fig. 8d). The diurnal cycle of COA shows distinctive peaks corresponding to lunch and dinner times. Similar diurnal patterns of COA were observed in London (Allan et al., 2010) and Beijing (Huang et al., 2010). On average, COA accounts for 16% of the total OA mass in NYC. Similar fractions of cooking aerosol were determined using the molecular tracer approach in Los Angeles (Rogge et al., 1991), Fresno, CA (Chow et al., 2007), and some urban sites in the southeastern US (Zheng et al., 2002).

During this study, POA (=HOA + COA) accounts for an average 30% of the OA mass and up to ~40% during morning rush hour and at night (Fig. 13b). It is interesting to note that despite the fact that our sampling site is close to two major highways (<1 mile), the average mass concentration of COA (1.02 µg m\(^{-3}\)) is higher than that of HOA (0.91 µg m\(^{-3}\)). This is an indication that cooking activities are a significant source of primary particles in NYC and that efforts to reduce particle pollution in this densely populated metropolitan area should focus on controlling both cooking and traffic emissions. POA and total OOA show almost no correlation (\(r^2 = 0.01\), Fig. 12b), confirming their different source origins. However, there is a general trend that low particle loading periods are characterized with elevated POA that accounts for a high mass fraction of OA (Fig. 12b), due to stronger influences from local primary emissions. High PM events, on the other hand, are mainly contributed by secondary inorganic species and oxygenated organics species formed regionally.

3.5.4 Nitrogen-enriched OA (NOA)

A unique N-enriched OA (NOA) component was determined for this study. Compared to the other four OA components, it shows a much higher N/C ratio (0.053; Fig. 8c) and correlates particularly tight (\(r^2 = 0.91\)) with \(\text{C}_2\text{H}_4\text{N}^+\), \(\text{C}_3\text{H}_6\text{N}^+\), and \(\text{C}_4\text{H}_{10}\text{N}^+\), which were likely generated from amino compounds (McLafferty and Turecek, 1993). It, however, shows almost no correlation with \(\text{C}_3\text{H}_7\) and \(\text{C}_6\text{H}_5\text{O}^-\) (Fig. 9c). NOA on average accounts for 5.8% of the total OA mass, yet 39% of the \(\text{C}_3\text{H}_7\text{N}^+_\text{p}\) signal (Fig. 10) and 33% of the nitrogen mass (Fig. S6). Note that the NOA of this study is clearly different from the “local N-containing reduced OA” (LOA) detected in Mexico City (Aiken et al., 2009), despite similar N/C ratios (0.06 for LOA). Specifically, the mass spectrum of NOA resembles more to that of OOA, while LOA in Mexico City is more similar to HOA and biomass burning OA (BBOA). The O/C and OM/OC ratios of NOA are 0.37 and 1.69, respectively, close to those of SV-OOA but much higher than 0.13 and 1.40, respectively, of LOA in Mexico City.

The diurnal pattern of NOA is marked with a pronounced noon-time peak (Figs. 13a and 8c) in association with southerly wind (Fig. 14a). To investigate the sources and processes of NOA, periods of highest NOA concentrations (~17% of time; denoted as Hi-NOA) are compared to the rest of the time (denoted as Lo-NOA). The average concentration of NOA during Hi-NOA (1.29 µg m\(^{-3}\)) is ~6 times higher than that during Lo-NOA (0.22 µg m\(^{-3}\)). Note that the average NR-PM\(_1\) composition is very similar between the two types of periods except for a substantially higher fraction of NOA during Hi-NOA. Based on comparing measured NH\(_4\)\(^+\) vs. predicted NH\(_4\)\(^+\) (=\(\text{SO}_4^2-/\text{NO}_3^-/\text{Cl}^-/\text{NH}_3^-/35.5\); Zhang et al., 2007b), particles appear to be slightly anion-rich during Hi-NOA while fully neutralized during Lo-NOA (Fig. 14b), suggesting that NOA might be carrying positive charges. In addition, Fig. 14c illustrates a broader size distribution of \(m/\ell\) 58 – the \(m/\ell\) contributed the most by NOA (Fig. 11e) – during Hi-NOA than during Lo-NOA while the size distributions of both \(m/\ell\) 44 and 57 show essentially no difference. Other \(m/\ell\)'s with relatively high contribution of NOA such as \(m/\ell\) 42 (\(\text{C}_3\text{H}_7\text{N}^+\)) and 30 (\(\text{CH}_3\text{N}^+\)) also show broader size distributions during Hi-NOA. These results together suggest a stronger influence of gas-particle partitioning on the size distribution of NOA, consistent with the hypothesis that NOA is mainly produced via acid-base reactions of amino compounds with acidic gases (Ge et al., 2010b) followed by the condensation of low volatility products onto pre-existing particles. A recent study in Cape Verde observed non-negligible contribution of aliphatic amines, e.g., dimethylamine (\(\text{C}_2\text{H}_7\text{N}\)) and diethylamine (\(\text{C}_4\text{H}_{11}\text{N}\)), to OC during algal blooms (Müller et al., 2009). Given the proximity of NYC to the Atlantic Ocean, it is possible that the NOA spikes are associated with marine emissions of aliphatic amines. In addition, given that amines are emitted from various industrial operations (Ge et al., 2010a), the large industrial region to the southwest of NYC could also be an important source of NOA.

3.6 Potential source regions of PM\(_1\) in NYC

Figure 15a and Table S2 present the average composition of PM\(_1\) and OA corresponding to four principal clusters of air masses, i.e., northwest (NW), west (W), southwest (SW), and Atlantic Ocean determined via cluster analysis of the HYPLIT back-trajectories (BT). The air flow patterns in NYC during this study were predominantly westerly and southwesterly (at frequencies of 37% and 25%, respectively) with infrequent flows from NW (7.3%). Aerosol loadings are on average the lowest (6.6 µg m\(^{-3}\)) for the NW cluster and the highest (14 µg m\(^{-3}\)) for the SW cluster, consistent with the spatial distribution of emission sources of air pollutants in the northeastern US (Fig. 15a). The correlation of aerosol loadings with different BT clusters is also consistent with the average spatial distribution of AOD determined from satellite

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Fig. 14. (a) Statistical distributions of NOA concentrations corresponding to different wind sectors, (b) correlations between measured NH$_4^+$ and predicted NH$_4^+$ ($=\text{SO}_2^-/96 \times 36 + \text{NO}_3^-/62 \times 18 + \text{Cl}^-/35.5 \times 18$; (Zhang et al., 2007b)), and (c) average size distributions of $m/z$ 44, 57, and 58 during high NOA periods vs. the rest of the times, respectively. The size distributions in (c) are all scaled to their corresponding maxima for better comparisons of peak shapes.

Fig. 15. (a) Average composition of PM$_1$ (pie charts) and OA (bar charts) for each cluster. The markers on the trajectories indicate 12 h interval. The area of the pie chart is proportional to the total mass of PM$_1$. The solid grey circles show the distribution of SO$_2$ emission sources from the EPA’s National Emission Inventory database (http://www.epa.gov/air/data/netemis.html). Bigger size indicates larger emissions. (b) Time series of total mass of PM$_1$ and AOD retrieved from Terra MODIS at 550 nm and (c) distribution of average AOD for the entire study.

retrieval. As shown in Fig. 15b, the daily averaged AOD values observed above NYC agree well with the average PM$_1$ concentrations measured during this study, indicating that the satellite AOD data (Fig. 15c) may be a useful indicator for the spatial distribution of PM$_1$ during this study.

The chemical compositions of PM$_1$ are also significantly different among the four clusters, likely reflecting the characteristics of different source regions of air pollutants. For example, the contributions of EC and HOA to PM$_1$ are the highest in the NW cluster, suggesting that emissions of primary...
particles from local sources are a key controller of PM$_1$ concentration and composition in NYC during relatively clean periods. In comparison, secondary species, e.g., OOA and sulfate, dominate the PM$_1$ composition in air masses originated from SW. The westerly air masses also correlate with high aerosol pollution with large fraction of secondary species. The observed correlations between aerosol loading and air mass trajectories are consistent with the high AOD and the large source emissions in the regions W and SW to NYC, e.g., large coal-fired plants and industrial sources in the Ohio River Valley, West Virginia, and Pennsylvania (Polissar et al., 2001; Qin et al., 2006). In addition, the large contribution of SV-OOA (∼70%) to total OOA (=SV-OOA+LV-OOA) in the NW cluster (Fig. 15a) might reflect large biogenic emissions in regions to the NW of NYC (Slowik et al., 2010). In addition, for ∼25% of the time during this study, air masses were traced back to the Atlantic Ocean with very low flow speed. Relatively high loadings of PM$_1$ composed of higher fraction of sulfate (26.6% of PM$_1$) and LV-OOA (40% of OA) appear to correlate with these air masses, which likely reflects the recirculation air pollutions in the US continental outflow. Indeed, the high AOD along the coastal line and near shore (Fig. 15c) seems to support this possibility.

4 Conclusions

The mass concentration, chemical composition, and size distributions of submicron aerosol species in NYC were measured in-situ using an HR-AMS in summer 2009. Aerosol composition varies dynamically with OA and sulfate on average comprising 54% and 24% of total PM$_1$ mass. Analysis of aerosol ion balance indicates that submicron particles in NYC are overall bulk neutralized. The size distributions of NR-PM$_1$ species, all of which vary dynamically during this study, on average present a similar large accumulation mode peaking at ∼550 nm with an additional small mode (∼150 nm) for OA. In addition to boundary layer dynamics, local photochemical production plays an important role in controlling the variation of sulfate and OA during daytime. Gas-to-particle partitioning and nighttime hydrolysis of N$_2$O$_5$ drive the diurnal cycle of nitrate. The average O/C and OM/OC ratios of OA are 0.36 (±0.08) and 1.62 (±0.11), respectively.

PMF analysis of the HRMS of OA identifies five components, i.e., a traffic-related HOA, a highly aged, regional LV-OOA, a less oxidized, more volatile SV-OOA, a cooking-related COA, and a NOA enriched of organic nitrogen compounds. SV-OOA and LV-OOA constitute the largest fraction of OA, accounting for 34% and 30%, respectively, on average. HOA and COA, which have similar mass concentrations, together account for 30% of the OA mass. HOA correlates well with the primary traffic emission tracers, with HOA/EC and HOA/NO$_x$ ratios of 1.29 and 0.045 (µg m$^{-3}$ ppbv$^{-1}$), respectively. LV-OOA correlates tightly with sulfate and C$_x$H$_{2y}$O$_z^+$ ions while SV-OOA correlates better with chloride and the C$_x$H$_y$O$_z^+$ ions. The diurnal cycle of HOA shows a pronounced morning traffic peak, so do EC and NO$_x$. The diurnal cycles of SV-OOA and LV-OOA demonstrate a trend of chemical evolution of SOA in NYC that likely involves a continuum of oxidation from SV-OOA to LV-OOA. The SV-OOA to LV-OOA evolution is consistent with a gradual increase of O/C ratio and a corresponding decrease of H/C ratio in total OA. NOA, which accounts for 5.8% of the OA mass, yet 33% of the organic N mass, shows a pronounced peak around noon time and correlates well with amines related peaks, e.g., C$_{x}$H$_{2x+2}$N$.^+$ Analysis of the sources and ion balance of NOA suggests that N-containing organic species such as amines might have played an important role in the atmospheric processing of OA, likely via acid-base chemistry coupled with photochemical production of acidic organic species. Air mass trajectory analysis, together with satellite AOD observations, indicates that emissions of primary particles from local sources likely play an important role in controlling PM$_1$ concentration and loading in NYC during relatively clean periods associated with northerly flows. High PM pollution events, however, are mainly contributed by secondary inorganic and organic particles typically associated with air masses originated from regions to the west and southwest of the city.

Appendix A

A1 Estimation of production rates of HNO$_3$ and H$_2$SO$_4$

The production rate of HNO$_3$ and H$_2$SO$_4$ can be estimated from bi-molecule association reactions (Seinfeld and Pandis, 2006). The effective second-order rate constant for a given temperature and pressure (altitude) can be calculated using Eq. (A1).

\[
k([M], T) = \left( \frac{k_o(T)[M]}{1 + \frac{k_o(T)[M]}{k_\infty(T)}} \right) ^{-1} 0.6 \left[1 + \log \left( \frac{k_o(T)[M]}{k_\infty(T)} \right) \right]^{-1}
\]

(cm$^6$ molecule$^{-2}$ s$^{-1}$)

\[
k_o(T) = k_o 300 \left( \frac{T}{300} \right)^{-n} (cm^6$ molecule$^{-2}$ s$^{-1}$) \quad (A2)
\]

\[
k_\infty(T) = k_\infty 300 \left( \frac{T}{300} \right)^{-m} (cm^3$ molecule$^{-1}$ s$^{-1}$) \quad (A3)
\]

NO$_2$ + OH $\xrightarrow{M} $ HONO$_2$ \quad (R1)

SO$_2$ + OH $\xrightarrow{M} $ HOSO$_2$ \quad (R2)

Where $k_o(T)$ (Eq. A2) is low-pressure-limiting rate constant and $k_\infty(T)$ is the high-pressure-limiting rate constant. The four parameters $k_o 300$, $n$, $k_\infty 300$, and $m$ for Reactions (R1 and R2) are listed in Table A1.
Table A1. Kinetic parameters used to determine the production rates of HNO$_3$ and H$_2$SO$_4$ (JPL, 2009).

<table>
<thead>
<tr>
<th>Reaction</th>
<th>$k^{300}$</th>
<th>$n$</th>
<th>$k^{300}$</th>
<th>$m$</th>
</tr>
</thead>
<tbody>
<tr>
<td>NO$_2$ + OH $\rightarrow$ HONO$_2$</td>
<td>$2.0 \times 10^{-30}$</td>
<td>3.0</td>
<td>$2.5 \times 10^{-11}$</td>
<td>0</td>
</tr>
<tr>
<td>SO$_2$ + OH $\rightarrow$ HOSO$_2$</td>
<td>$3.0 \times 10^{-31}$</td>
<td>3.3</td>
<td>$1.5 \times 10^{-12}$</td>
<td>0</td>
</tr>
</tbody>
</table>

A2 Calculation the equilibrium constant of gas-to-particle partitioning

Formation of ammonium nitrate involves an equilibrium reaction between the gas phase NH$_3$ and HNO$_3$, and particle phase NH$_4$NO$_3$ (Reaction R3). The gas-to-particle partitioning is strongly temperature dependent, and the equilibrium constant of Reaction (R3) can be calculated as Eq. (A4) (Seinfeld and Pandis, 2006)

$$\text{NH}_3(g) + \text{HNO}_3(g) \leftrightarrow \text{NH}_4\text{NO}_3(s)$$  \quad (R3)

$$k(T) = k(298)\exp\left\{a \left(\frac{298}{T} - 1\right) + b\left[1 + \ln\left(\frac{298}{T}\right) - \frac{298}{T}\right]\right\}$$  \quad (A4)

where $T$ is the ambient temperature in Kelvin. $k(298) = 3.35 \times 10^{16}$ (atm$^{-2}$), $a = 75.11$, and $b = -13.5$.

Supplementary material related to this article is available online at: http://www.atmos-chem-phys.net/11/1581/2011/acp-11-1581-2011-supplement.pdf.

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