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Sources and Secondary Production of Organic Aerosols in the Northeastern United States during WINTER

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Key Points:

- Wintertime organic aerosol (OA) was almost as oxidized as in summer studies, with wood burning OA being an important contributor to primary OA
- SOA formation rate and amount is similar to summer (accounting for lower OH). A box model reproduces SOA, but fast chemistry may be missing
- The default GEOS-Chem formulation with nonvolatile POA shows large errors. The SIMPLE and semivolatile parameterizations perform better

[Supporting Information:](http://dx.doi.org/10.1029/2018JD028475)

[•](http://dx.doi.org/10.1029/2018JD028475) [Supporting Information S1](http://dx.doi.org/10.1029/2018JD028475)

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Sources and Secondary Production of Organic Aerosols in the Northeastern United States during WINTER

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Abstract Most intensive field studies investigating aerosols have been conducted in summer, and thus, wintertime aerosol sources and chemistry are comparatively poorly understood. An aerosol mass spectrometer was flown on the National Science Foundation/National Center for Atmospheric Research C-130 during the Wintertime INvestigation of Transport, Emissions, and Reactivity (WINTER) 2015 campaign in the northeast United States. The fraction of boundary layer submicron aerosol that was organic aerosol (OA) was about a factor of 2 smaller than during a 2011 summertime study in a similar region. However, the OA measured in WINTER was almost as oxidized as OA measured in several other studies in warmer months of the year. Fifty-eight percent of the OA was oxygenated (secondary), and 42% was primary (POA). Biomass burning OA (likely from residential heating) was ubiquitous and accounted for 33% of the OA mass. Using nonvolatile POA, one of two default secondary OA (SOA) formulations in GEOS-Chem (v10-01) shows very large underpredictions of SOA and O/C (5×) and overprediction of POA (2×). We strongly recommend against using that formulation in future studies. Semivolatile POA, an alternative default in GEOS-Chem, or a simplified parameterization (SIMPLE) were closer to the observations, although still with substantial differences. A case study of urban outflow from metropolitan New York City showed a consistent amount and normalized rate of added OA mass (due to SOA formation) compared to summer studies, although proceeding more slowly due to lower OH concentrations. A box model and SIMPLE perform similarly for WINTER as for Los Angeles, with an underprediction at ages <6 hr, suggesting that fast chemistry might be missing from the models.

1. Introduction

Atmospheric aerosols are known to affect human health (Pope III et al., 2002), visibility (Boers et al., 2015), and climate (Myhre et al., 2013). In particular, considerable uncertainty remains in the radiative forcing by aerosols. Therefore, efforts to improve our understanding of aerosol sources, transport, processing, and forcing continue to be important.

Field studies provide our primary source of information on aerosol concentrations, sources, and evolution in the real atmosphere. However, the large majority of intensive field studies at northern midlatitudes have been conducted in warmer and more photochemically active times of the year, especially those targeting detailed organic aerosol (OA) composition, sources, and evolution. Several studies that have looked at seasonal variations of aerosol composition around the world have shown distinct differences between summer and winter (Allan, Alfarra, et al., 2003; Hu et al., 2016; Huang et al., 2013; Poulain et al., 2011; Takegawa et al., 2005; Weimer et al., 2006). Midlatitude winter has significantly lower temperatures, reduced solar radiation, and often more precipitation, all of which can significantly alter the processes that govern the production, processing, and lifetimes of aerosols. In addition, dry deposition velocities of semivolatile species have been shown to be significantly different for some species in winter (Jaeglé et al., 2017). Although in the last

few decades there have been a number of wintertime studies of aerosols in Asia (e.g., Cao et al., 2003; Kim et al., 2017; Rengarajan et al., 2007, 2011; Takegawa et al., 2005; Xu et al., 2016; Zhang et al., 2015) and Europe (e.g., Allan, Alfarra, et al., 2003; Bardouki et al., 2003; Castro et al., 1999; Crippa et al., 2013; Lanz et al., 2008; Louvaris et al., 2017; Viana et al., 2006), very few have been conducted in North America (e.g., Budisulistiorini et al., 2016; Öztürk et al., 2013; Song et al., 2001; Strader et al., 1999; Tan et al., 2002; Weimer et al., 2006; Xu et al., 2015). However, most of the North American studies were conducted at least a decade ago, and none were conducted in the northeastern United States (NE-US). Moreover, of these wintertime aerosol studies, only a few have investigated the source contributions of OA and none have investigated the rates of secondary OA (SOA) formation from wintertime pollution emissions. In general, wintertime studies have shown lower total concentrations of submicron aerosols than summer. Similar to summer studies, OA is a major (and often the largest) fraction of submicron aerosol in the winter, typically with increases in primary OA (POA), mostly driven by increases in biomass burning OA (BBOA) from residential wood burning (Bressi et al., 2016; Budisulistiorini et al., 2016; Crippa et al., 2013; Lanz et al., 2008; Xu et al., 2015, 2016). The amount of oxidized OA (OOA) observed in warmer months of the year has been documented extensively in the literature, throughout much of the world (Jimenez et al., 2009, and references within). The amount of OOA reported in urban environments varies greatly depending on the geographical location, but in general, accounts for approximately 50–80% of the total OA mass during the summer. In the winter, fewer studies have reported on measurements of urban OOA (Crippa et al., 2013; Kim et al., 2017; Öztürk et al., 2013; Pirjola et al., 2017; Zhang et al., 2015), showing fractions of OA mass between 20% and 70%.

OOA is often used as a proxy for SOA, which forms from gas-to-particle conversion via gas-phase or aqueousphase mechanisms. During the summer, the solar radiation, temperature, and vegetation leaf coverage are highest, resulting in biogenic emissions that contribute significantly to local and regional chemistry (Ding et al., 2016; Ehn et al., 2014; Henze et al., 2008; Marais et al., 2016; Palm et al., 2016), even in polluted regions. In winter, however, biogenic emissions are often much smaller or negligible (Ding et al., 2016, 2017; Guenther et al., 2012) due to reduced sunlight, temperatures, and foliage. Several studies have investigated the production of urban SOA in polluted areas under warm weather (Cevik et al., 2016; DeCarlo et al., 2010; Freney et al., 2014; Hayes et al., 2013, 2015; Kleinman et al., 2008; Ma et al., 2017; McMeeking et al., 2012; Weber et al., 2007). However, no study (to the best of our knowledge) has presented a detailed evaluation on the regional production of SOA during winter using aircraft data.

Chemical transport models often underpredict OA concentrations (Tsigaridis et al., 2014), particularly near and downwind of anthropogenic sources. Recently, significant efforts have gone into improving the capability of global models to better predict OA evolution by implementing a variety of SOA formation and aging schemes, with differing levels of complexity (Hodzic et al., 2016; Hodzic & Jimenez, 2011; Jo et al., 2013; Kim et al., 2015; Pye & Seinfeld, 2010; Tsigaridis et al., 2014; Tsimpidi et al., 2016). As part of the AeroCom II model intercomparison, Tsigaridis et al. (2014) compared OA from 31 different global chemical transport and general circulation models. That study found that in general, global models can reasonably predict the dominantly secondary nature of OA, yet the absolute amount of OA is still significantly underpredicted, with a mean normalized bias of -0.62 for all surface urban data compared, -0.15 in remote regions, and -0.30 in marine locations. Although the spatial resolution of most 3-D models is too coarse to accurately capture OA mass concentrations in smaller urban areas, megacities like Los Angeles, Beijing, or New York City are of more similar size to model resolutions thus reducing error in the comparisons. In addition, when examining seasonality, the models are unable to predict observed wintertime trends in several urban locations, leading the authors to hypothesize that capturing the relative importance of SOA versus anthropogenic POA is a critical factor for the models to more accurately simulate seasonal trends. Similarly, Tsimpidi et al. (2016) used the general circulation model ECHAM combined with the Modular Earth Submodel System (MESSy) MESSy to evaluate modeled OA combustion sources compared to 84 different observational data sets. From that study, the authors concluded that a major weakness in the model is the poor ability to simulate wintertime POA and SOA concentrations, which they hypothesized is due to the model missing an important wintertime OA source or SOA formation pathway.

Wintertime measurements are needed to better understand the changes in aerosol sources, atmospheric processing, and lifetimes, and to better constrain regional and global models to more accurately predict wintertime concentrations and seasonal differences. Here we present wintertime submicron aerosol composition measurements from the NE-US from an airborne platform. We compare wintertime concentrations to

previously reported summertime studies, apply source apportionment analysis to explore the sources of OA and contrast them to observations from warmer periods of the year, investigate the evolution of OA (amount and composition), and compare it to predictions from the GEOS-Chem model. We report the first aircraft measurements of SOA production efficiency in winter from a case study of urban outflow from New York City, and we compare it to a box model and a simple parameterization of SOA formation.

2. Methods

The Wintertime INvestigation of Transport, Emissions, and Reactivity (WINTER) aircraft campaign (WINTER, 2018) took place out of the NASA Langley Research Center (Hampton, VA) from 1 February to 15 March 2015 aboard the National Center for Atmospheric Research (NCAR) C-130 aircraft. Aircraft studies provide insight into changes in chemical properties and processes spatially, both vertically and horizontally, that stationary ground campaigns cannot. The campaign objectives for WINTER were to characterize both gas-phase and particle-phase wintertime emissions and chemical transformations. Thirteen research flights (RF) were carried out (see supporting information Figure S1), with 102 hr of sampling time. Aircraft parameters and ambient meteorological measurements were provided by the NCAR Research Aviation Facility. More details about WINTER, the payload, and logistics can be found online, and all data are public and available at (WINTER-Data, 2018). Data reported here are at standard temperature (273 K) and pressure (1,013 mbar) (STP), and all linear fits within this manuscript use the orthogonal distance regression method.

In general the 2015 winter season over the NE-US was colder than average. The NCEP/NCAR reanalysis data indicate that surface temperatures in the NE-US for February 2015 were about 5 °C colder than the February 1981–2010 average.

2.1. Instrumentation

2.1.1. PM₁ Nonrefractory Composition (AMS)

A highly customized high-resolution time-of-flight aerosol mass spectrometer (Aerodyne Research Inc., hereafter referred to as "AMS") measured nonrefractory composition of PM₁ (NR-PM₁; DeCarlo et al., 2006; Dunlea et al., 2009). The theory and operation of the AMS, including the key adaptations for aircraft sampling, have previously been described in detail (Canagaratna et al., 2007; DeCarlo et al., 2006; Dunlea et al., 2009; Jayne et al., 2000; Kimmel et al., 2011) and is only briefly summarized here. Ambient particles were sampled through an NCAR High-Performance Instrumented Airborne Platform for Environmental Research Modular Inlet (HIMIL, 2018; Moharreri et al., 2014) at a flow rate of 10 L/min, into a pressure-controlled inlet operated at 325 Torr (Bahreini et al., 2008). The particles were then focused into the high-vacuum region of the mass spectrometer through an aerodynamic lens and transmitted into a detection chamber, where particles impacted on a porous tungsten standard vaporizer (600 °C). Nonrefractory species in the particles were flash vaporized and then ionized with 70-eV electron impact ionization. The generated ions were then extracted and analyzed by time-of-flight mass spectrometry. The air residence time from the outside tip of the HIMIL inlet to the vaporizer was 0.5–0.6 s, depending on altitude. Transmission of particles through the AMS inlet and lens was similar to previous publications for well-performing instruments (Figure S2; Hu et al., 2017).

During WINTER, the AMS was operated in the Fast Mass Spectrum mode (Kimmel et al., 2011) in order to obtain high time resolution measurements (1 Hz). For more information on Fast Mass Spectrum operation see supporting information section S1.1 and Figure S3. For all flights the AMS sampled in V mode with a resolution of 2,300 at m/z 44 and 2,900 at m/z 184. Collection efficiency (CE) was estimated with the algorithm of Middlebrook et al. (2012; supporting information section S1.1.2). Relative ionization efficiencies (Canagaratna et al., 2007) for sulfate, ammonium, and chloride were determined from repeated infield calibrations (Figure S5). Typical detection limits during WINTER were 137 (57) ng/sm³ for sulfate (SO₄), 12 (5) ng/sm³ for ammonium (NH₄), 69 (30) ng/sm³ for nitrate (NO₃), 77 (32) ng/sm³ for chloride (ChI), and 474 (194) ng/sm³ for OA for a 1-s (1-min) sampling interval (sm³ refers to cubic meters at standard temperature and pressure). Note that the charge symbols are not used (i.e., we write NO₃ and not NO₃ $\,$), as there can be contributions from organonitrates and organosulfates to AMS $NO₃$ and $SO₄$, respectively (e.g., Docherty et al., 2011; Fry et al., 2013; Liao et al., 2015). Accuracy for aircraft AMS measurements of inorganic (organic) species is estimated at ±35% (±38%; Bahreini et al., 2009). Semirefractory species, such as NaNO₃ (sodium nitrate) and Na₂SO₄ (sodium sulfate), are inefficiently detected by the AMS (Hayes et al., 2013). AMS data have sometimes been used to estimate the fractions of ammonium nitrate and

organic nitrates using the measured NO_2^+ /NO⁺ ion ratio (Farmer et al., 2010; Fry et al., 2013). However, in this study, the presence of particulate nitrite (Guo et al., 2016) and the likely partial detection of NaNO₃ made that method indeterminate. For that reason, in addition to total measured nitrate, estimates were made of total inorganic nitrate concentrations based on comparisons of the AMS data to the ion chromatography-based instruments. For a complete discussion on the measured nitrate during WINTER see supporting information section S1.1.4.2 and Figures S8–S10. Several corrections were made to the AMS data to account for recently discovered, small vaporizer artifacts that affect OA and chloride quantification, also discussed in supporting information section S1.1.3 and Figure S6. The AMS was fully operational for all flights during WINTER except RF05 leading to an overall 96% data coverage.

2.1.2. Collocated WINTER Measurements Used in This Study

PM₁ size-resolved volume concentration was measured by two optical particle counters mounted on the wings of the C-130, an Ultra High Sensitivity Aerosol Spectrometer (Droplet Measurement Technology; Brock et al., 2016; Cai et al., 2008) and a Passive Cavity Aerosol Spectrometer Probe (PCASP-100X, Droplet Measurement Technologies; Liu et al., 1992). The Ultra High Sensitivity Aerosol Spectrometer, as used in this study, had a geometric size range of 0.075–1 μm and the PCASP measured from 0.1–3 μm (geometric). Only data collected within bins that correspond to a vacuum aerodynamic diameter of ≤ 1 µm were used in our analyses, and both particle counters were corrected for differences in inlet transmission to achieve more accurate comparisons with the AMS (Hu et al., 2017; Knote et al., 2011; see supporting information S1.1.4.1 for more details). Both probes substantially dry particles on the way to the laser, due to their diffusing intakes up front (which slow the air, leading to heating), sample flow passage though warm anti-iced components, internal components near room temperature, and recirculating sheath flow at that internal temperature. Thus, the aerosol volume measured by these probes approximately corresponds to the dry particle volume and does not include liquid water that was present on those particles under ambient conditions (Strapp et al., 1992). Both instruments were serviced and calibrated using polystyrene latex spheres (refractive index $[n] = 1.595$) by the manufacturer just prior to deployment. To more accurately estimate the total volume concentrations of ambient particles, both instruments were size corrected (1.12×diameter) similar to Kupc et al. (2017) and Liu et al. (2017) to estimate the measured diameters with a more atmospherically relevant refractive index (dry ammonium sulfate, $n = 1.53$ [Hand & Kreidenweis, 2002]).

PM₁ water-soluble mass concentration was measured by two particle-into-liquid samplers (PILS). The first PILS-IC, described in Guo et al. (2016), was coupled to an online ion chromatograph ("Online-PILS-IC," Metrohm 761 Compact ICs; Hennigan et al., 2006; Orsini et al., 2003; Peltier et al., 2007) and reported the concentrations of sulfate, nitrate, and chloride at a time resolution of 3 min. The second PILS-IC was coupled to a Bretchel fraction collector system (Sorooshian et al., 2006) for offline analysis ("Offline-PILS-IC") of levoglucosan and the inorganic ions sodium, ammonium, potassium, chloride, nitrate, and nitrite at a time resolution of 2 min (Sullivan et al., 2017). The inorganic ions were measured by anion/cationexchange chromatography (Guo et al., 2016) and levoglucosan by high-performance anion-exchange chromatography-pulsed amperometric detection (Sullivan, Frank, Kenski, et al., 2011; Sullivan, Frank, Onstad, et al., 2011; Sullivan et al., 2014). PM₅ bulk aerosol composition was measured offline from filter samples ("SAGA Filters"; Dibb, 2003; Dibb et al., 2002), at a time resolution of ~7 min. Volatile organic compounds (VOCs) were measured for 35 s every 2 min with the Trace Organic Gas Analyzer (Apel et al., 2010). For more details on specific WINTER configurations for SAGA-Filters and Trace Organic Gas Analyzer see supporting information sections S1.2 and S1.3, respectively. Carbon monoxide (CO) was quantified by vacuum ultraviolet resonance fluorescence with a customized commercial instrument (Aero-Laser AL-5002), at a rate of 0.5 Hz. NO, NO₂, and NO_v were measured by a six-channel cavity ring-down instrument. $NO₂$ was measured directly by laser absorption at 405 nm. NO was converted and measured as $NO₂$ after an addition of excess O_3 (Wagner et al., 2011). NO_y was thermally dissociated in a heated quartz inlet (650 °C) and similarly converted and measured as $NO₂$ with an addition of excess $O₃$ (Wild et al., 2014; Womack et al., 2017).

2.2. Modeling

2.2.1. FLEXPART

European Reanalysis-Interim meteorological fields with a horizontal resolution of $0.5^{\circ} \times 0.5^{\circ}$, a temporal resolution of 3 hr, and 60 vertical hybrid pressure-terrain following coordinate levels were used to drive

the FLEXPART Lagrangian particle dispersion model (Seibert & Frank, 2004; Stohl et al., 1998, 2005; Stohl & Thomson, 1999). The FLEXPART output has a resolution of 0.1° \times 0.1° over the domain of 100.0°W to 60.0°W and 25.0°N to 50.0°N and includes residence time in the surface layer (<100, <250, and <500 m). Thirty thousand back trajectory particles were released every 20 s, or for every altitude change of 100 m, along the aircraft flight tracks. The back trajectories were calculated for 48 hr with outputs at 15-min intervals.

2.2.2. GEOS-Chem

GEOS-Chem is a 3-D chemical transport model driven by assimilated meteorological fields from NASA Global Modeling and Assimilation Office (GMAO) GEOS-5 FP system (Rienecker et al., 2008). GEOS-Chem simulates the tropospheric chemistry of CO, HO_x, NO_x, VOC, O₃, BrO_x, SO₂, and aerosols (Bey et al., 2001; Park, 2003; Park et al., 2004; Parrella et al., 2012). We use GEOS-Chem v10-01 for this study. The model is run in a nested-grid configuration with a resolution of 0.5° latitude × 0.625° longitude over North America (10°N to 60°N, 130°W to 60°W), and a 4° latitude \times 5° longitude resolution for the rest of the globe. Anthropogenic emissions over the United States are from U.S. Environmental Protection Agency's (EPA) 2011 Version 6 Emissions Modeling Platform based on the National Emissions Inventory (NEI) 2011 v1 (NEI, 2018) and from the Emissions Database for Global Atmospheric Research (EDGAR) v4.2 (EDGAR, 2018) for the rest of the globe.

We perform three simulations using different approaches for the simulation of OA. The first approach follows the OA simulation of Pye and Seinfeld (2010). This approach assumes nonvolatile POA and is referred to as the "Complex SOA scheme without semivolatile POA" in the GEOS-Chem online manual (GEOS-Chem, 2017) but referred to as "nvPOA" here. This approach includes SOA formation from oxidation of aromatics (benzene, toluene, and xylene), monoterpenes, sesquiterpenes, and isoprene. Aerosol yields are parameterized using a volatility basis set with saturation concentrations (C*) at 298 K of 0.1, 1, 10, and 100 μg/m³ and using an enthalpy of vaporization (ΔH_{vap}) of 42 kJ/mol to determine the temperaturedependent $\check{\mathsf{C}}^*$. SOA formation from semivolatile volatility organic carbon (SVOC) and intermediate volatility organic carbon (IVOC), as well as heterogeneous oxidation of POA are not included in the nvPOA GEOS-Chem simulation. The second approach is similar to nvPOA with the exception of replacing nonvolatile POA with semivolatile POA, which includes partitioning of SVOC precursors to the gas phase, as well as the addition of primary IVOC (P-IVOC) precursors, with oxidation of both types of precursors to form SOA (Pye et al., 2010; Pye & Seinfeld, 2010). This approach is referred to as the "Complex SOA scheme with semivolatile POA" in the GEOS-Chem online manual (GEOS-Chem, 2017) but referred to as "svPOA" here. We assume that all SVOC emissions are captured by the NEI-2011 emissions. P-IVOC emissions are estimated by scaling naphthalene emissions, which are based on benzene emissions following Pye and Seinfeld (2010). Thus, our assumptions of IVOC emissions and chemistry are the same as those in (Pye & Seinfeld, 2010). Both Pye simulations are currently the default formulations implemented in GEOS-Chem v10-01 and can be selected by the user.

The third approach implements the SIMPLE ("SIMPLifiEd parameterization of combustion SOA") parameterization of Hodzic and Jimenez (2011) and Hayes et al. (2015) for anthropogenic and biomass burning SOA, as described by Kim et al. (2015). This approach is referred to as the "Simple SOA Scheme" in the GEOS-Chem online manual (GEOS-Chem, 2017) and is also referred to as "SIMPLE" here. This approach is planned to be the default formulation starting with v11-02 of GEOS-Chem. A single lumped SOA precursor VOC, which represents the SOA formation potential from all anthropogenic VOCs, P-IVOCs, and primary SVOCs (P-SVOCs), is emitted proportionally to anthropogenic CO, with a ratio of 0.069 g VOC (g CO) $^{-1}$. Residential burning emissions are calculated separately based on the literature, and the ratio of the precursor VOC to CO is 0.013 g VOC (g CO) $^{-1}$. The lumped VOC makes nonvolatile SOA upon OH oxidation with a 100% mass yield. Multigenerational aging is not treated explicitly in SIMPLE but is included implicitly since the parameterization is based on fits to ambient observations. The precursor VOCs are oxidized by OH with a rate coefficient of 1.25 \times 10⁻¹¹ cm³ per molecule per second to form SOA, which is assumed to be nonvolatile. No SOA is formed from heterogeneous oxidation of POA. The limited net SOA formation (with OA increasing only slightly above the initial POA emissions) from biomass burning emissions implemented in SIMPLE is consistent with the overwhelming evidence from field studies of this source, including 17 aircraft studies (Cubison et al., 2011; Jolleys et al., 2012; Liu et al., 2016; Shrivastava et al., 2017). In SIMPLE, POA is modeled similarly as is in the nvPOA scenarios, and biogenic SOA is assumed to form with yields of 3% and 5% from isoprene and monoterpene oxidation, respectively, at the point of emissions.

In GEOS-Chem, OA species are represented on the basis of OC mass, and the OA mass is calculated from specified OA/OC values listed in supporting information Table S7. For comparison with the aircraft-based observations, GEOS-Chem is sampled along the flight track at 1-min intervals.

2.2.3. SOA Formation Box Model

SOA formation and evolution was also simulated with a 0-D box model, in which the oxidation of SOA precursors follows the ROB + ZHAO + MA parameterization (Ma et al., 2017). This parameterization performed reasonably well to simulate SOA formation in the Los Angeles area during CalNex-2010, with some underestimation at short photochemical ages, and agreement within the uncertainties for ages >1 day (Ma et al., 2017). The parameterization is described previously in Ma et al. (2017) and Hayes et al. (2015) and is documented in more detail in supporting information section S1.4. Briefly, oxidation products from VOCs and P-IVOCs are described by a volatility basis set where the precursor species are oxidized into five volatility bins $(C^* = 0.1, 1, 10, 100, 1,000)$ according to experimental chamber measurements of OA yields for speciated VOCs and P-IVOCs (Ma et al., 2017; Presto et al., 2010). The VOC oxidation yields are corrected for vapor wall losses in the experimental chambers as described in Ma et al. (2017). Oxidation and multigeneration aging of S/IVOCs follow a bin-hopping approach where oxidation with hydroxyl radicals lowers species volatility by 1 order of magnitude for each generation (Robinson et al., 2007). Semivolatile gases are treated by equilibrium partitioning to the particle phase, using the reformulation of Pankow theory as described by Donahue et al. (2006). Initial concentrations are specified based on available WINTER measurements whenever possible or using scaled ratios to CO determined for Los Angeles when WINTER measurements are not available (see supporting information section S1.4). The SIMPLE parameterization of urban SOA formation, with the parameters of Hayes et al. (2015) derived for Los Angeles during CalNex-2010 (also used in Kim et al., 2015), is also compared to the results of the box model to evaluate the utility of the derived summertime parameters for use in wintertime conditions. We note that biogenic precursors made a very small contribution to SOA formation from urban precursors (which is what SIMPLE is parameterizing) in the CalNex study, independently of some biogenic background SOA being present over larger regional scales (Hayes et al., 2015). Biomass burning OA in field studies is observed to oxidize but without significant net gain of mass (Cubison et al., 2011; Jolleys et al., 2012; Liu et al., 2016; Shrivastava et al., 2017), and for this reason it is treated as inert in the box model.

3. Results and Discussion

3.1. Total Submicron Nonrefractory Aerosol Composition During WINTER

The average fractional composition of the nonrefractory submicron aerosol during WINTER (\leq 0.5 km), is shown in Figure 1, along with paired summer/winter studies at several urban and rural sites around the world. The WINTER boundary layer height (BLH) was estimated to be 0.5 km (supporting information section S2.2.2 and Figure S16). The average NR-PM₁ (NH₄ + NO₃ + SO₄ + Chl + OA) concentration during WINTER in the BL was 5 μ g/sm³ with 13% NH₄, 22% NO₃, 26% SO₄, 1% Chl, and 37% OA. In contrast, the total concentration calculated from data measured in a similar region during the summer of 2011, which offers the most direct comparison with the WINTER data set, was 2.5 times greater. Four of six comparison data sets also show higher concentrations in summer than winter. On average across all data sets, winter concentrations are about 20% (by mass) lower in winter than summer, consistent with Hand et al. (2012). However, since wintertime BLH are significantly lower than summer, these results suggest that the total BL column amounts are substantially lower in winter. Other notable trends that are apparent from Figure 1 are (1) the fraction of nitrate tends to be larger ($2 \times$ on average) in winter than summer. (2) Sulfate fraction is typically lower (0.7x) in winter. (3) The fraction of OA also tends to be lower (0.8×) during winter, presumably due to the importance of biogenic SOA sources in summer and faster SOA formation from anthropogenic emissions.

To investigate the components of OA present during WINTER, positive matrix factorization (PMF) analysis was performed (Ulbrich et al., 2009; Zhang et al., 2011), and four components were identified: hydrocarbon-like OA (HOA); BBOA; less-oxidized, oxidized OA (LO-OOA); and more-oxidized, oxidized OA (MO-OOA). For a discussion on the details of the PMF analysis, see supporting information section S2.1 and Figure S12. The average fraction of each OA component is shown in Figure 2a: 9% is apportioned to HOA, 33% to BBOA, 22% to LO-OOA, and 36% to MO-OOA. These data yield a total POA of 43% and 57% of the OA mass as OOA. BBOA is approximately 79% of the POA mass, which is not surprising, since residential wood burning as a source of household heating is ubiquitous in this region during

Figure 1. Overview of main NR-PM₁ species fractions and concentrations near the surface during WINTER (\leq BLH, 0.5 km) and during ground-based studies at several other Northern Hemisphere locations. Averages during WINTER are summarized under "Northeast, US, 2015, Winter." Fractional compositions are also measured with an AMS, from several urban sites: New York City (NYC; Drewnick et al., 2004; Weimer et al., 2006), Manchester, UK (Allan, Alfarra, et al., 2003; Allan, Jimenez, et al., 2003), Shanghai, China (Huang et al., 2013), and Tokyo, Japan (Takegawa et al., 2005); from an urban downwind site 80 km east of NYC (Zhou et al., 2016); and one rural/remote site in Melpitz, Germany (Poulain et al., 2011), are also shown. The averages of all sites are shown in the gray-shaded box. These sites were chosen because data from both summer and winter were available. All data used in Figure 1, with the exception of data from this study, and data for NYC Downwind US, 2011 (Zhou et al., 2016), were taken from the summary Table S1 compiled by Zhang et al. (2007; and references within).

Figure 2. (a) Average fractional contribution of the identified positive matrix factorization factors to the average OA mass measured during WINTER. The thick outline surrounding HOA and BBOA represents the total fraction of primary organic aerosol (POA), while the remaining segment represents the total fraction of oxidized OA (OOA). Binned vertical concentration profiles (0–3 km) of (b) all four positive matrix factorization factors, (c) POA and OOA, and (d) atomic ratios H/C and O/C, during the WINTER campaign are also shown. The dashed lines in b–d are averages for each 153-m (500-ft) bin, and the shaded regions represent the standard error. BBOA = biomass burning OA; HOA = hydrocarbon-like OA; LO-OOA = less-oxidized, oxidized OA; MO-OOA = more-oxidized, oxidized OA.

the winter (Bond et al., 2007; Zhang et al., 2010). The contribution of BBOA to total OA is within the range (~20–40%) reported in several other studies that have reported BBOA during the winter in the United States, Asia, and Europe (Gilardoni et al., 2016; Lanz et al., 2008; Louvaris et al., 2017; Pirjola et al., 2017; Zhang et al., 2015). Some BBOA may age into OOA over regional timescales (Cubison et al., 2011). In 2015 the U.S. EPA implemented a new national policy (EPA, 2015) that increased the regulations and standards for new residential wood heating appliances, which may over time lead to a decrease of the impact of wood burning BBOA in the United States during winter. This recent change in emission standards highlights the need for continuing wintertime studies in this region to track the impact of this regulation. The fraction of OOA is similar to that observed in other winter studies, such as in urban sites in China (Zhang et al., 2015), Finland (Pirjola et al., 2017), and Switzerland (Lanz et al., 2008).

3.1.1. Spatial Patterns of Submicron Aerosol During WINTER

Figure 3 shows the spatial distributions of the submicron aerosol mass of NH_4 , SO_4 , NO_3 , Chl, OA, and O/C measured during WINTER. In general, sulfate (Figure 3b) tends to have higher concentrations (>2 μ g/sm 3) off the coast of the northeastern seaboard with intermediate concentrations (0.75 $<$ SO₄ $<$ 2 μ g/sm³) in the Ohio River Valley region, where SO_2 sources are expected to be larger. This observation is consistent with slower conversion of SO₂ to SO, due to longer lifetime of SO₂ in winter (Lee et al., 2011), which reduces the concentration of SO_4 over the source regions and increases it downwind. The distribution of nitrate (Figure 3c) shows similar regions of higher concentrations (>1.5 μg/sm³) as sulfate, yet perhaps with higher frequency near urban regions, consistent with shorter lifetimes of NO_x in winter (Fibiger et al., 2018). Chloride (Figure 3e) was typically very low ($<$ 0.03 μ g/sm 3) throughout the entire sampling region, with the exception of one flight (RF03), which was off the coast of the New York City Metropolitan Area ("NYC" hereinafter; >0.1 μg/sm³). OA (Figure 3d) was fairly widespread during WINTER with concentrations along the eastern seaboard ranging from about 1 to 4 μ g/sm³ and elevated regions downwind of Atlanta and also in the Ohio River Valley. The extent of oxidation (O/C) of the OA was relatively consistent at around 0.8 (Figure 3f), which is surprisingly high for wintertime conditions due to expected reduced oxidant levels and photochemical activity.

Figure S13 shows the spatial distribution of the fraction of each species in NR-PM₁. Sulfate tends to dominate (≥50% of aerosol mass) in coal-fired power plant-rich regions (when over land) or when sampling far off the eastern seaboard, where the total submicron concentrations are low ($<$ 1 μ g/sm 3). Nitrate is consistently above 25% when sampling in, or downwind, of major urban centers (Atlanta, Cincinnati, and Pittsburgh) and reaches >50% in the NYC Metro area. OA was ubiquitous, with fractions persistently above 25–30% and sometimes >50% when near urban centers (Atlanta and NYC). Higher fractions of OA are also often seen during high-altitude (>3 km) transits. Chloride, on the other hand, was typically $<1\%$ of the total mass throughout the entire sampling region, with the exception of one flight (RF03), which was off the coast of NYC, and reached a maximum of about 2–3%.

The vertical distributions of submicron aerosols are presented in supporting information section 2.2.1 and Figure S14a for the entire campaign, where all species show a trend of decreasing concentration with altitude. Vertical profiles were also investigated for data filtered for day, night, over land, and over the ocean sampling (Figures S14b–S14g). However, no major differences were observed for these sampling conditions.

3.1.2. Spatial Patterns of OA Composition During WINTER

To investigate the vertical distribution of OA components and the extent of oxidation of OA in the vertical, the identified PMF factors (supporting information section s2.1) as well as H/C and O/C ratios are plotted as a function of altitude in Figure 2. The OA components tend to rapidly decrease in magnitude with an increase in altitude, until an enhanced layer is observed at around 1.1 to 1.8 km. These layers are predominantly driven by a few short segments, during only a couple of flights (RF03 and RF06), and therefore, do not represent a ubiquitous campaign-wide stratification (supporting information Figure S15). Near the surface, the OA is found to be a 50/50 mixture of POA and OOA. At these altitudes the POA is dominated by BBOA and OOA is ~60% MO-OOA. The H/C and O/C vertical profiles (Figure 2d) show that OA near the surface is substantially oxidized, with an O/C~0.65. As altitude increases, OA tends to be more oxidized. This increase in oxidation with altitude may be explained by (a) fresh emissions being oxidized (leading to SOA formation and heterogeneous oxidation of POA) as they loft to higher altitudes; (b) as fresh emissions loft, they are mixed into more aged air; or (c) more likely some combination of both.

Summarizing the OA oxidation over the entire sampling region during WINTER, the normalized frequency distribution of O/C is presented in Figure 4b. The WINTER O/C distribution is compared to typical ranges for OA components from previous studies and to distributions from two previous U.S. aircraft studies: DC3 over the Central and southeastern United States (spring; Barth et al., 2015) and SEAC⁴RS over much of the continental United States (summer; Toon et al., 2016). Only data within the boundary layer (Figure S16) is included for each campaign. Figure S17 shows that the distributions are insensitive to the specific BLH chosen, to within a few kilometers. Surprisingly, OA is almost as oxidized in WINTER as in the previous campaigns. These results indicate a substantial contribution of SOA to OA during WINTER (see section 3.1.3.), in agreement with the interpretation of PMF results, despite the slower photochemistry. DC3 and SEAC⁴RS also had significant impacts from biomass burning POA (~1/4 of the OA; Liu et al., 2016; Saide et al., 2015; Yang et al., 2015), which is more oxidized than HOA (e.g., Aiken et al., 2008; Canagaratna et al., 2015). Therefore, the presence of BBOA during WINTER could not explain these data if SOA was not a major component during WINTER.

Figure 4. (a) Average and one standard deviation of O/C for hydrocarbon-like organic aerosol (HOA), biomass burning OA (BBOA), and oxidized OA (OOA) compiled in the supporting information of Canagaratna et al. (2015) are shown for reference. (b) Normalized O/C distributions for the Wintertime INvestigation of Transport, Emissions, and Reactivity (WINTER) campaign and three different simulations from GEOS-Chem using the NEI-2011 inventories with the nonvolatile primary OA (nvPOA), semivolatile POA (svPOA), and SIMPLifiEd parameterization of combustion (SIMPLE) secondary OA (SOA) formulations (only data from RF01-07 are included for these distributions). Distributions measured during the summertime SEAC⁴RS and the springtime DC3 campaigns are also shown for comparison. Only data within the boundary layer (see Figures S16 and S17) and when OA is larger than its detection limit are included in the elemental composition distributions for all campaigns (to remove the effect of noise at very low concentrations). The boundary layer height for WINTER was determined to be 0.5 km, and for both SEAC⁴RS and DC3 a value of 2 km was determined. The concentration of measured and modeled POA and SOA components are shown in c and as relative contributions in d (RF01-RF07 only).

3.1.3. Evolution of Bulk OA Composition During WINTER

The mass weighted average O/C for WINTER was 0.75. The O/C from WINTER is more oxidized than in several prior winter studies (Figure 5a). For example, an O/C of 0.35 was measured at the surface in Fresno, CA, during the month of January (Ge et al., 2012), and in Southern China an O/C was measured at 0.43 during December (Huang et al., 2013). The WINTER O/C data indicates that the OA is even more oxidized than found in a few summertime studies in a similar region to WINTER. For example, Sun et al. (2011) measured OA at Queens College, NYC, during July and found an O/C of 0.46 and Zhang found an O/C in Upton, NY, of 0.61 (Chen et al., 2015). At the summer ground-based study of Sun et al. (2011), the fraction of chemically reduced POA components (HOA and cooking OA) was 30%, compared to 9% in our study. BBOA, a more oxidized POA, was not reported by Sun et al., but accounts for 32% of the OA in our study, which contributes to the observed differences.

The atomic ratios of H/C and O/C can be plotted against each other (Figure 5a) in a Van Krevelen (VK) diagram (Van Krevelen, 1950), which is useful to visualize the evolution of OA (Hayes et al., 2013; Heald et al., 2010; Ng et al., 2010, 2011). Different slopes on the VK diagram may be due to different oxidation mechanisms or result from sampling a variable mixture of POA and OOA. A linear fit to the WINTER VK diagram yields a slope of -0.85 , which is approximately consistent with the formation of carboxylic acids or hydroxycarbonyls. Chen et al. (2015) recently synthesized numerous campaigns into VK space for a range of different locations (aircraft, remote/rural, urban, and downwind sites). Air masses sampled during WINTER were predominantly urban and downwind targets. The average slopes calculated, from data summarize by Chen et al. (2015), for urban sites was -0.87 and for downwind sites was -0.86, very similar to the WINTER slope. Average data for different components and several other winter studies are also shown on Figure 5a for reference.

Ng et al. (2010, 2011) utilized the fractions of integrated signal in the AMS at m /z 44 (mostly CO $_2^{\text{+}}$) versus m /z 43 (mostly C₃H₇⁺ from POA and C₂H₃O⁺ from OOA) to the total OA signal as a simpler way to characterize the evolution of OA to OOA, which can also be performed with the now ubiquitous Aerosol Chemical Speciation Monitors (Ng et al., 2011). The WINTER data (Figure 5b) are consistent with Ng et al. (2010, and references within), in that theyfall within a region well defined by a triangle and that most of the data lie within the defined range of mixed POA/OOA, extending into the MO-OOA area. The fraction of data, in WINTER, within the HOA region is very small, which is consistent with the low average fraction of HOA calculated with PMF (~9%).

Figure 5. (a) Van Krevelen (VK) diagram and (b) f_{44} versus f_{43} plot for 1-min data colored by altitude (above ground level [AGL]) for the entire Wintertime INvestigation of Transport, Emissions, and Reactivity (WINTER) study. Shown for reference on a are average hydrocarbon-like organic aerosol (HOA), biomass burning OA (BBOA), less-oxidized, oxidized OA (LO-OOA), and more-oxidized, oxidized OA (MO-OOA) from WINTER along with the same species plus cooking OA (COA) and total OOA summarized in Canagaratna et al. (2015) Table S1 (and references within). Also shown for comparison are data from several other wintertime studies (W in legend); Davis, CA (Chen et al., 2015), Fresno, CA (Ge et al., 2012), Melpitz, Germany (Poulain et al., 2011), Barcelona, Spain (Mohr et al., 2012), China (Huang et al., 2013), and two summertime studies (S in legend) that took place in Queens, NY (Sun et al., 2011), and Upton, NY (Chen et al., 2015). All the older studies have been corrected using the updated H/C and O/C calibrations of Canagaratna et al. (2015). Linear fits for the WINTER data (slope $= -0.85$), WINTER LO-OOA and MO-OOA (slope $= -0.49$), and the average urban (slope $= -0.87$) and downwind (slope = 0.86) from Chen et al. (2015) are also shown on a. In b, regions of OA oxidation, based on the observations by Ng et al. (2010), are shown in the shaded boxes, and the average f_{44} and f_{43} data from each OA component for RF01-07, as well as for each transect during RF03.

Cubison et al. (2011) proposed a plot of f_{44} versus f_{60} as useful to investigate the presence and chemical evolution of BBOA. f_{60} is the fraction of integrated signal in the AMS at m/z 60 to the total OA signal and is typically dominated by the $\mathsf{C_2H_4O_2}^+$ ion (under biomass burning-dominated conditions), which originates from levoglucosan and related species (cellulose pyrolysis products). In WINTER, there is some evidence (Figure S18) that other ions not typically associated with biomass burning are present at m/z 60 in appreciable amounts. Therefore, to more accurately investigate the evolution of BBOA during WINTER, we plot the dominant ions f_{CO2+} and $f_{C2H4O2+}$ in Figure S19a. This shows that there is a large portion of data that is greater than a background line of non-BBOA sources of ~0.3% (Cubison et al., 2011), consistent with the conclusion that there is a ubiquitous regional source of BBOA mixed with other sources. Results from the OA source apportionment analysis (Figure S12), indeed confirms this conclusion, as the time series data of the fraction of BBOA is consistently >10% and on average is 33% of the OA mass. Some BBOA may age into OOA (Cubison et al., 2011), although the contribution of this evolution for our case study (section 3.2) appears to be small.

In addition to the above characteristic plots, Hu et al. (2015) examined the relationship of f_{CO2+} versus $f_{\text{CH6O+}}$ to explore the evolution of SOA production from $low-NO_x$ isoprene oxidation. Given that this study took place during the winter, when deciduous trees have lost their leaves and the solar irradiance is reduced, the mass contribution to OA from biogenic sources is expected to be negligible (Budisulistiorini et al., 2016). A plot of WINTER data of f_{CO2+} as a function of f_{CBH6O+} (Figure S19b) clearly shows that this is the case; as the majority of the data tends to scatter around the 0.0175% background level determined for urban + biomass burning influenced air (Hu et al., 2015).

3.1.4. Comparison of OA Components and Oxidation to GEOS-Chem

The O/C distribution is also compared in Figure 4b to model results from three simulations of GEOS-Chem using the nvPOA, svPOA, and SIMPLE SOA formulations (see supporting information section S2.4 for details on this analysis). Using the nonvolatile default SOA formulation in GEOS-Chem v10-01 (nvPOA) and the 2011 NEI emissions result in an O/C mode of ~0.15, which is an underprediction of the measured peak O/C by about a factor of 5. The nvPOA results in a POA concentration that is more than $2\times$ greater than the measured HOA + BBOA (since BBOA is not tracked separately from POA in nvPOA and svPOA) and accounts for 97% of the total OA in this simulation (Figures 4c and 4d and supporting information Table S7). The unrealistically low production of SOA (18× too low) and high POA by this model version explains the very large underprediction of O/C.

In contrast, the svPOA simulation yields an O/C mode of 0.65, which is much closer to the observations. In this case, the concentration of POA is about 1/3 of the measurements (HOA + BBOA). The relative contribution of SOA (86%) is about 1.4× greater than measured (58%), despite being ~10% lower in concentration. This large fraction of SOA leads to better agreement with the observed O/C, albeit partially due to cancellation of errors. The SIMPLE formulation results in an O/C mode of 0.45, which falls between the nvPOA and svPOA O/C modes, and underpredicts the observations by a factor of 1.6. Here the concentration of POA (which is simulated similarly to the nvPOA case) is greater than the observations by ~2×, resulting in a lower relative SOA contribution than measured.

Summarizing these results indicates that (1) using the default nvPOA formulation is clearly inadequate at capturing the production of SOA in polluted air and use of this formulation should be avoided in future studies (either winter or summer); (2) the default svPOA formulation more closely predicts the extent of OA oxidation, reasonably predicts SOA, but significantly underestimates POA; and (3) the less computationally expensive SIMPLE formulation performs reasonably well at reproducing the mass concentration of SOA, although it overpredicts POA (presumably due to too high POA emissions in NEI2011). We therefore recommend using the SIMPLE or svPOA formulations in GEOS-Chem in the future. In addition, a reduction in O/C for primary OA and an increase in O/C for secondary OA, using the SIMPLE formulation, better predicts the extent of oxidation in this formulation (Figure S21) and should be considered when evaluating wintertime OA oxidation in other regions.

3.2. Wintertime Evolution of OA: A Case Study Downwind of the NYC Metropolitan Area

Several studies have investigated the production and downwind evolution of urban SOA (de Gouw et al., 2008; DeCarlo et al., 2010; Hayes et al., 2013; Kleinman et al., 2007, 2008), but most of these studies were conducted during warmer seasons (e.g., spring or summer). To investigate these processes during winter, the NYC metropolitan area sampled during RF03 (7 February 2015) was selected as a case study. The portion of the flight path, for this case study, followed a pseudo-Lagrangian pattern heading east of NYC. This flight was selected because the flight path was predominantly off the coast of the northeastern seaboard, and as anthropogenic emissions essentially stop at the coastline, the plume sampled is dominated by advection, dilution, and aging of urban emissions from metro NYC, thus providing a unique case to investigate the formation and evolution of SOA during winter. The relevant portion of the flight path for this analysis is shown in Figure 6a. The average wind direction during this time was southwesterly at 225° and with an average speed of 7 m/s, and the flight path crossed the metro NYC plume several times. Based on the flight path, wind direction, and time series concentration profiles (Figure 6c), five plume transects were identified as T1–T5. Although the plane crossed the plume two more times downwind of T5, these transects showed strong evidence of mixing from unrelated air masses and therefore were excluded in this analysis. All gas-phase species mixing ratios increased as the aircraft entered a plume transect and decreased to background levels upon exiting the transect. This, along with the average wind direction, strongly suggests that we were sampling NYC urban emissions injected into a regional background. To confirm this, 48-hr FLEXPART back trajectories (section 2.2.1) are shown in Figures 6a and 6b and supporting information Figure S22. All back trajectory altitude profiles show slight uplift approximately 12 hr prior to the air masses descending to the point of the aircraft transect. The back trajectories for T1 and T2 pass directly through NYC, whereas T3–T5 show the air coming from the northern metropolitan area. The average hourly surface (<140 m) CO, simulated in GEOS-Chem, is also overlaid on the map (Figure 6a) and represents the areas impacted by urban emissions in the area. All back trajectories distinctly cross a region of high CO emissions, further indicating that all transects sampled were directly influenced by metro NYC urban emissions. In addition, based on transport times determined from the back trajectories (see section 3.2.1), the amount of transport during daylight ranged from 3 to 10 hr for T1 and T5, respectively (supporting information Table S8).

OA/ΔCO can be used as a metric for evaluating the evolution of OA in an urban plume, where ΔCO is a conserved tracer used to account for dilution of primary emissions. The method allows quantifying the increase

Figure 6. (a) Map of the flight track for RF03 colored by day (yellow) and night (black), according to sunrise/sunset, and median 48-hr FLEXPART back trajectories colored by transect number. Overlaid on a is the GEOS-Chem (NEI-2011) CO, which is the modeled average hourly surface CO (<140 m) concentrations, from 10:00 to 23:00 UTC (i.e., from the estimated emission time to the end of sampling of T5). Also shown on the map are cities and populations (in millions) for reference. (b) Altitude profiles of the median FLEXPART back trajectories (colored by transect number), along with the estimated time of emission (determined from linearly interpolating across the T0 line shown in a for each transect). (c) Time series data of organic aerosol (OA), CO, toluene, NO_x , and flight altitude for the relevant period of RF03. Transects T1-T5 are colored uniquely and identified in the legend in b. The 25th and 75th percentiles for all back trajectories and altitude profiles are not shown here for clarity (see supporting information Figure S22). AGL = above ground level.

in OA/ΔCO (i.e., ΔOA/ΔCO) with photochemical age due to SOA formation (e.g., DeCarlo et al., 2010; Docherty et al., 2008; Hayes et al., 2013; Kleinman et al., 2007, 2008; Ortega et al., 2016; Takegawa et al., 2006). As mentioned above, Figure 6c suggests that there is a relatively constant OA baseline of \sim 1.6 μ g/sm³, , indicating that the NYC plume is emitted into a preexisting regional background. Therefore, each of the plume transects in this case study were analyzed for the ratio of $\Delta X_{\text{species}}/\Delta CO$, where $\Delta X_{\text{species}} = \text{total}$ X_{Species} – regional background X_{Species}. Those ratios represent the impact of metro NYC emissions.

Figure 7a shows the evolution of ΔOA/ΔCO, which shows an increasing trend from T1 to T5. To evaluate the causes of the increases, the decay of the measured aromatic VOCs were similarly analyzed (Figure 7d and supporting information Figure S23), since aromatic VOCs are thought to be significant contributors to urban SOA

Figure 7. (a) Evolution of ΔOA/ΔCO, (b, c) the fraction of organic aerosol (OA) components from positive matrix factorization (PMF) analysis, and (d) decay of an example volatile organic compound (see Figure S23 for all volatile organic compound ratios) used for estimating photochemical age. The standard errors for the y variables in a and d are smaller than the data points shown. BBOA = biomass burning OA; HOA = hydrocarbon-like OA; LO-OOA = less-oxidized, oxidized OA; MO-OOA = more-oxidized, oxidized OA (MO-OOA); POA = primary organic aerosol.

formation (Hayes et al., 2015; Odum et al., 1997). As the evolution of the plume progressed from T1 to T5, all aromatic ΔVOC/ΔCOs (with the exception of the least reactive benzene) decreased as ΔOA/ΔCO increased, which is consistent with photochemical aging and thus should lead to SOA formation.

To investigate the causes of OA evolution, PMF results are shown in Figures 7b and 7c. The fraction of OOA increased from T1 (33%) to T5 (56%). Even at T5, this fraction is lower than what was reported by de Gouw et al. (2005) and Kleinman et al. (2007; 57–75%) in the Northeastern Air Quality Study (NEAQS) 2002 study, which also took place in the NE-US but during the summer. During the same study de Gouw et al. (2005) reported that only 11% of the OA mass was from anthropogenic primary sources, which is significantly lower than the POA at T1 (67%) or even T5 (44%). The larger fraction of POA measured in this study is driven by the large fraction of BBOA, attributed to residential wood burning, which was absent from the 2002 summertime study, as well as the absence of biogenic OA during WINTER (which was very likely an important contributor to the summer study). The progression of OA components from T1 to T5 shows that the fraction of primary components decreased over time and the OOA components increased as the air was photochemically aged, consistent with many previous studies (de Gouw et al., 2008; DeCarlo et al., 2010; Hayes et al., 2013; Kleinman et al., 2007, 2008). The increase in SOA fraction is predominately a result of gas-to-particle conversion since the POA remained relatively constant throughout these transects (see Figure 8b). This suggests that aging of BBOA into OOA (Cubison et al., 2011) played a minor role during this case study.

This study took place during the winter, where clouds can be prevalent. The evidence for SOA formation in clouds is mixed (Ervens, 2015; Gilardoni et al., 2016; Wagner et al., 2015) but cannot be discounted. Vertical profiles of ambient temperature and relative humidity (RH) were constructed from data collected at two nearby sounding sites, OKX (40.87°N, 72.87°W, near T1/T2) and CHH (41.67°N, 69.97°W, near T5) at 0 UTC and 12 UTC (Figure S24). These data indicate that at the average aircraft altitude (0.3 km, pressure Altitude) during RF03 (and up to 2 and 3 km), RH was persistently below 100%. The time series of ambient RH and air temperature (Figure S25) measured from the aircraft, during the relevant portion of RF03, as well as forward facing camera images from the C-130 taken at roughly the middle of each transect from T1 to T5 (Figure S26) confirm that we were not flying directly in clouds and that the RH was <100% during all plume transects. Three of four nearby airports (John F. Kennedy, NY; Islip/Macarthur, NY; Montauk, NY; and Nantucket Memorial, MA) reported trace amounts of precipitation (Local Climatological Data, 2018) at some point during the sampling period. Furthermore, during these transects, there was no evidence of a significant increase in sulfate (Figure S27) as one would expect if aqueous phase chemistry was a substantial factor (Ervens, 2015, and references within). These data combined suggest that wet deposition and cloud processing were most likely insignificant contributors (but cannot be completely excluded) to the production or loss of SOA in this case study.

3.2.1. Air Mass Age

To enable a quantitative analysis of the rate of SOA formation, the age of the air (since injection of urban emissions into the NYC metropolitan area) for each transect was estimated based on both the transport time and

Figure 8. (a) RF03 plume transect ΔOA/ΔCO versus photochemical age using an standardized OH (bottom x axes) of 1.5 \times 10⁶ molecules per cubic centimeter (for ease of comparison to literature studies) and the estimated OH during our study of 6.7 \times 10⁵ molecules per cubic centimeter (top x axes). Shown for comparison are ΔOA/ΔCO values measured in the northeastern United States (NE-US) during summer (NEAQS-2002; de Gouw et al., 2005; DeCarlo et al., 2010; Kleinman et al., 2007), Los Angeles in summer (CalNex; Hayes et al., 2013; Ortega et al., 2016), and Mexico in spring (MILAGRO-2006) (DeCarlo et al., 2010; Kleinman et al., 2008). For a complete discussion on the comparison studies see supporting information section S2.5.1. ΔFactor/ΔCO for (b) primary organic aerosol (POA) components and for (c) secondary organic aerosol (SOA) components are also shown with linear fits, where "m" and "b," in the legend, denote the slope and intercept, respectively, using the bottom x axes. Note, the standard errors for the y variables for all WINTER data in a–c are smaller than the data points shown. A small apparent increase in biomass burning organic aerosol (BBOA) with age may be due to noise or to small variations in BBOA emissions with time of day. HOA = hydrocarbon-like OA; LO-OOA = less-oxidized, oxidized OA; MO-OOA = more-oxidized, oxidized OA.

photochemical processing. The transport age is described as the average amount of time the air mass took to travel to each point along the transect's flight path, from a defined time zero (T0), which was chosen as the longitude of NYC Center, and is shown on Figure 6a. For comparison, the transport age calculated based on the average wind speed, measured from onboard the aircraft at each transect, and distance from NYC was within 8% of the transport age determined using FLEXPART. To evaluate the amount of chemical processing of the air mass intercepted in each transect, photochemical ages were estimated using three different methods: (1) VOC ratio clock, (2) the NO_x clock, and (3) a novel approach to the VOC ratio clock, termed the VOC/CO clock. We note that CO formation from VOCs is a very small contributor to its concentrations in urban plumes (Griffin et al., 2007). The first two methods have been described at length in previous literature (de Gouw et al., 2005; Kleinman et al., 2003, 2007, 2008; Olszyna et al., 1994; Parrish et al., 2007) and are only briefly described here. All three methods are based on the first-order decay of a species under OH oxidation (equation (1)).

$$
Y_{t=t} = Y_{t=0} \exp^{-kt[OH]} \tag{1}
$$

where Y_i is the concentration of any OH reactive species *i*, *t* is time, *k* is the rate constant for a given species with OH, and the subscript $t = 0$ ($t = t$) denotes Y_i at the time of emission (aircraft measurement).

The VOC ratio clock is based on the concept that if two VOCs are coemitted from a given source with a characteristic initial emission ratio, the more reactive VOC will be removed faster from the atmosphere, through oxidation, as the air moves downwind. As a result, any ratio of VOCs (VOC_a/VOC_b), where the more reactive VOC is in the numerator, will decrease with time and can be substituted in equation (1) for Y. The rate constant for a ratio of VOCs is represented by the difference, or $k_{(VOCa-VOCb)}$. The VOC/CO clock is identical to the VOC ratio clock except CO is used in the denominator. On timescales relative to VOCs, CO is considered inert ($k_{OH, VOCi}$ > > $k_{OH, CO}$) thus, $k_{OH, CO} \approx 0$, and only $k_{OH, VOCi}$ is needed for this method.

Table 1

Estimated OH and Initial Emission Ratio (eER) Determined for Each Pair of Species (Figure 7d and supporting information S23)

Note. The average OH (\pm 1σ) and the OH rate constant (k_{OH}) used are also shown. k_{OH} for toluene and NO_x were calculated at the average temperature of the plume transects (269 K), and all other k_{OH} are temperature independent (Calvert et al., 2015).

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aUnits of pptv/ppmv.

The NO_x clock is defined such that at the time of emission (i.e., $t = 0$) NO_y = NO_x. Therefore, as air is transported away from the source, the ratio (NO_x/NO_y) will decrease and can also be substituted in equation (1) for Y. Since, in this method, the main loss process of NO_x is assumed to be photochemically driven from the reaction of OH with NO₂, k is simply that of NO₂ with OH. The impact of nighttime N₂O₅ chemistry is more important at T5 and may reduce the accuracy of the NO_x clock for this transect.

For all three methods described, equation (1) can be used to determine the photochemical age (t) or the concentration of OH. In this analysis, 13 ratios (Δ1,2,3-TMB/Δtoluene; Δ1,2,4-TMB/Δtoluene; Δ1,2,3-TMB/Δ o-xylene; Δ1,2,4-TMB/Δo-xylene; Δ (m,p-xylene+ethylbenzene)/Δtoluene; Δo-xylene/Δtoluene; Δtoluene/ ΔCO; Δo-xylene/ΔCO; Δ (m,p-xylene+ethylbenzene)/ΔCO; Δ1,2,3-TMB/ΔCO; Δ1,2,4-TMB/ΔCO; and ΔNOx/ ΔNOy) were evaluated by fitting versus transport time to estimate OH (Figures 7d and S23), since no direct measurement of OH was available during WINTER. Here OH is assumed to be the dominant daytime oxidant available for reaction with the precursor VOCs and NO_y during atmospheric transport. In NEAQS-2002 Warneke et al. (2004) investigated VOC oxidation, by O_{3} , NO₃, and OH, and found that anthropogenic and oxygenated VOCs were primarily oxidized during the daytime by OH. During winter, OH is expected to be lower than in the summer due to (a) reduced water vapor decreasing the fraction of O(1 D), which converts to OH, and (b) decreased photolysis of OH precursors such as O₃ as a result of reduced solar radiation (Gligorovski et al., 2015; Heard et al., 2004; Ren et al., 2006). However, the potential importance of the radical precursors CINO₂, HONO, and CH₂O during WINTER may lead to larger OH concentrations than expected otherwise (Heard et al., 2004). Based on our analysis, the average daytime OH was estimated to be 6.7 \pm 2.6 \times 10⁵ molecules per cubic centimeter (Table 1), which is indeed lower than typical summer average daytime OH (1.2 \times 10⁶–3 \times 10⁶ molecules per cubic centimeter; de Gouw et al., 2005; Hayes et al., 2013; Ren et al., 2006; Volkamer et al., 2006) and within a range of what has been observed in previous winter studies (1.1 \times 10⁵–7 \times 10⁵ molecules per cubic centimeter; Berresheim et al., 2013; Emmerson et al., 2005; Heard et al., 2004; Ren et al., 2006). Our estimated OH is also consistent with a mean daytime OH of 8 \times 10⁵ molecules per cubic centimeter for the GEOS-Chem simulations of WINTER.

To enable a direct comparison to results of prior publications that mostly used a standardized OH concentration of 1.5 \times 10⁶ molecules per cubic centimeter (Hayes et al., 2013; Hu et al., 2016; Mao et al., 2009; Ortega et al., 2016; Palm et al., 2016), we used this value to determine equivalent (eq.) ages using equation (2) as follows:

$$
eq.Age = transport time \times \frac{[OH]_{estimated}}{[OH]_{standard}}
$$
 (2)

Figure 9. Comparison of measured to predicted secondary organic aerosol (SOA) formation from the box model, with unscaled primary intermediate volatility organic carbons. Model simulations were initialized at (a) time zero (T0) and (b) Transect 1 (T1). SOA formed from T0 to T1 is the amount of measured LO-OOA + MO-OOA at T1. The SIMPLE parameterization of urban SOA (Hayes et al., 2015) is also shown for comparison. In the figure legend I-SOA is SOA from primary intermediate volatility organic carbons, V-SOA is SOA from volatile organic compounds, and S-SOA is SOA from primary semivolatile volatility organic carbons. The y-error bars in a represent the uncertainty of the OA and CO measurements, and the x-error bars show the uncertainty in equivalent age as a result of the uncertainty in estimated OH concentration. Only two error bars are shown for clarity. BBOA = biomass burning OA; HOA = hydrocarbon-like OA; LO-OOA = less-oxidized, oxidized OA; MO-OOA = more-oxidized, oxidized OA.

3.2.2. Wintertime Evolution of SOA

The average ΔOA/ΔCO versus equivalent photochemical age for the case study are shown in Figure 8a. As in previous field studies (e.g., DeCarlo et al., 2010; Hayes et al., 2013), ΔOA/ΔCO increases steeply with age. For comparison, Figure 8 also shows results from summertime measurements in Los Angeles during CalNex (Hayes et al., 2013; Ortega et al., 2016), summer NE-US from the NEAQS-2002 study (de Gouw et al., 2005; DeCarlo et al., 2010; Kleinman et al., 2007), and springtime in Mexico City (MILAGRO; DeCarlo et al., 2010; Kleinman et al., 2008). The ΔOA/ΔCO in WINTER reaches ~32 μ g·sm $^{-3}$ ·ppmv $^{-1}$ in about 1 eq. hr of aging at T1 and steadily increases to ~54 μ g·sm $^{-3}$ ·ppmv $^{-1}$ after about 5 eq. hr of aging by T5, indicating an SOA formation rate of 4 μ g·sm $^{-3}$ ·ppmv $^{-1}$ ·eq. hr $^{-1}$. For comparison, the average rate of formation in the NE-US summer was observed to be ~3 μg·sm $^{-3}$ ·ppmv $^{-1}$ ·eq. hr $^{-1}$, ~6 μg·sm $^{-3}$ ·ppmv $^{-1}$ ·eq. hr $^{-1}$ in Los Angeles summer, and ~5 μ g·sm $^{-3}$ ·ppmv $^{-1}$ ·eq. hr $^{-1}$ in Mexico City spring. The formation rate has been observed to be steeper at short ages and eventually plateaus at ages of ~1 day (Hodzic & Jimenez, 2011; Ortega et al., 2016). Overall, the WINTER results show that urban SOA formation is proceeding with similar intensity and rate as during summer studies, once oxidant and emission/dispersion differences are accounted for.

The primary OA components (HOA, BBOA, and total POA) and oxidized components (LO-OOA, MO-OOA, and total OOA) are analyzed identically to OA in Figures 8b and 8c. HOA and BBOA show nearly constant levels, as does total POA, which is consistent with the identification as primary components, the good pseudo-Lagrangian approximation of the case study, and their slow aging due to heterogeneous oxidation (e.g., DeCarlo et al., 2008; Ortega et al., 2016). OOA (Figure 8c) clearly captures nearly all the increase in OA with photochemical age, which is consistent with SOA production. Total OOA increases at a rate of 3.8 μ g·sm $^{-3}$ ·ppmv $^{-1}$ ·CO eq. hr $^{-1}$. These results further corroborate the efficient wintertime production of SOA observed in the NYC metro area.

3.2.3. SOA Production in Pseudo-Lagrangian Study: Measurements Versus Box Model

A box model (Hayes et al., 2015; Ma et al., 2017) was used to simulate the formation of SOA in the NYC metro case study (Figure 9). The box model was simulated with two different initializations: (1) from T0 (Figure 9a, referred to as I_{T0}) by using the estimated initial emission ratios from the fits shown in Figure S23 and (2) from T1 (Figure 9b, referred to as I_{T1}), in which all SOA formed before T1 is accounted for by the measured average OOA at T1. Additionally, I_{T_0} and I_{T_1} were run with two cases for P-IVOC inputs: (a) constrained using the CalNex measurements of Zhao et al. (2014) and (b) scaled down according to the ratio of measured VOCs between WINTER and CalNex (see supporting information section S1.4).

The unscaled P-IVOC results (Figure 9) more closely reproduce the measured results, when compared to the scaled P-IVOCs (Figure S29), suggesting that the VOCs and P-IVOCs have dominant contributions from different sources. This is consistent with Zhao et al. (2014), who attributed P-IVOCs to non-on-road sources, and McDonald et al. (2018), who showed that volatile chemical products (e.g., personal care and cleaning products, coatings, adhesives, or pesticides) are major sources of P-IVOCs. Results from both initializations (with unscaled P-IVOCs, Figure 9) are consistent and capture \sim 2/3 of the observed SOA formation (I_{T0} is within 63%, and I_{T1} is within 60% of the observed SOA formation, on average over the sampling period). This model was also lower by a very similar amount (~53%, see supporting information Figure S30) than the observations for the same range of short photochemical ages in the Los Angeles case (Ma et al., 2017). This persistent result suggests that some fast-reacting precursors might be missing or that the SOA formation chemistry of SVOCs and P-IVOCs are faster than implemented in these parameterizations, for example, if fast autoxidation reactions played a role (Crounse et al., 2013), although this is less likely under colder winter temperatures.

Similar to what has been observed in CalNex, the fraction of the total SOA formed from VOCs (according to the model) is ~25–30%. However, during WINTER, the fractional contribution to SOA from SVOCs is ~3× less, whereas SOA from P-IVOCs is about 2x greater than what was observed in summertime LA, after ~6 hr of photochemical aging. Lower SOA from SVOCs is consistent with lower temperatures driving the SVOCs to the particle phase where they remain as POA. Similarly, P-IVOCs more rapidly condense to form SOA at these temperatures, thus increasing their relative contribution at younger ages (together with their higher relative input concentrations for the default model case).

The box model was also used to simulate O/C during this case study for each of the four initializations (see supporting information section S1.4 and Figure S32). All four simulations capture the general trend of increasing O/C with age observed but are consistently on the low end of the uncertainty range of the measured O/C.

In addition to the box model, the SIMPLE parameterization (Hayes et al., 2015; Hodzic & Jimenez, 2011; Kim et al., 2015), with the parameters derived by Hayes et al. (2015) for Los Angeles (Figure 9), is also consistent with the observed SOA formation, albeit slightly lower (~80% of observed amount at T5). This result suggests that the parameters derived from Los Angeles region in summertime can also be used to reasonably predict the SOA formation from NYC in winter. Again, the ratio of the SIMPLE parameterization to the observations is very similar to the Los Angeles results for the same range of short photochemical ages.

Unfortunately, the range of photochemical ages reached in this case study was limited. Extending the I_{T0} simulation to 3 days (supporting information Figure S31) shows that both the box model and the SIMPLE parameterization plateau after ~1–2 days, due to the depletion of the precursor VOC. However, they reach significantly different final values. It is not possible to constrain the SIMPLE parameterization for longer ages in this domain with the information available to this manuscript. However, some of the air masses sampled contain air that has been oxidized for longer periods than in this case study. Thus, we recommend optimizing the SIMPLE parameters by performing multiple simulations in GEOS-Chem (or other 3-D models) and finding the ones that results in optimal agreement with the regional observations, similar to what was done by Murphy et al. (2017), who used a different 3-D model.

4. Conclusions

Wintertime ambient aerosol measurements were made aboard the NCAR C-130 in the NE-US from February to March 2015, as part of the WINTER campaign. These measurements offered a unique opportunity to investigate aerosol sources and processes during wintertime, which are understudied in the scientific literature. This manuscript focuses on (1) general trends observed in measured submicron aerosol throughout the NE-US; (2) sources and processing of submicron OA; and (3) wintertime production of SOA, through a pseudo-Lagrangian case study.

On average, the submicron aerosol concentration was 5 μ g/sm 3 , with 13% being NH₄, 22% NO₃, 26% SO₄, 1% Chl, and 37% OA. Comparing to summer measurements in a similar region, wintertime values are lower by about a factor of 2.5. However, accounting for differences in BLH between winter and summer, these results suggest that wintertime total column concentrations are significantly lower. This observation is consistent with several other studies in that wintertime concentrations tend to be approximately 20% lower, on average, than in summer, but with substantial variability (2–63%). Based on PMF analysis, primary HOA (a surrogate for

vehicle exhaust and similar sources) was a minor contributor to the overall OA mass (<10%), while BBOA was fairly ubiquitous, accounting for 33% of the OA. BBOA in this study is thought to be dominated by residential wood burning, consistent with wintertime studies in other regions. OA was also fairly oxidized, with ~60% of the OA mass apportioned to OOA, and was almost as oxidized as in recent spring and summer U.S. aircraft campaigns. The atomic O/C mode in WINTER was 0.75 (<0.5-km altitude).

SOA and O/C are significantly underestimated for the nvPOA default formulation within GEOS-Chem v10-01, which produces an O/C mode 5 times lower than the measurements, and with 97% of the OA mass as primary. The nvPOA GEOS-Chem simulation only produced a small fraction of SOA (3%), far lower than observed (~60%). This result, clearly indicates that the GEOS-Chem v10-01 formulation with nvPOA and anthropogenic SOA from benzene, toluene, and xylene (based on Pye et al., 2010) is inadequate to capture SOA formation from urban emissions, and we recommend completely avoiding its use in future studies (for either summer or winter studies). When GEOS-Chem was used with the alternative default parameterization, which includes SVOC and P-IVOC (svPOA) or the SIMPLE SOA parameterization (Hayes et al., 2015; Hodzic & Jimenez, 2011) as implemented in GEOS-Chem by Kim et al. (2015), model SOA concentrations were closer to the observations. These two parameterizations differed by about an order of magnitude for POA, with svPOA underpredicting it by a factor of 5, while SIMPLE overpredicted POA by a factor of ~2. O/C was closest to the observations for the svPOA parameterization, however, partially due to a cancellation of errors (too low POA together with too high O/C for SOA). Considering the reasonable performance of the SIMPLE SOA parameterization during WINTER and its computational efficiency, we recommend this parameterization for future GEOS-Chem studies. The svPOA parameterization should be considered for studies where additional details for SOA formation are useful.

A case study from a research flight that flew a pseudo-Lagrangian flight pattern off the coast of the NYC metropolitan area was used to investigate the production of SOA. Given that it was winter, this case study provided a unique opportunity to examine the impact of urban emissions, isolated from biogenic influences, on SOA, something that cannot be done easily during the summertime. Evolution of the ratios of different species are consistent with the pseudo-Lagragian nature of this case study, and through comparison with the transport age, allowed estimation of the concentration of OH (6.7 \times 10⁵ molecules per cubic centimeter, as expected lower than in summer), and therefore photochemical age. SOA production from urban emissions in the NE-US during winter was as efficient and rapid (when accounting for the differences in OH) as what has been observed in summer studies. This supports previous literature evidence of the importance of SOA formation from urban emissions and the need to represent this source in atmospheric models. A box model that approximately reproduced the Los Angeles observations, as well as the SIMPLE parameterization, performed well for this case study, slightly underpredicting SOA at short photochemical ages, and with consistent results with GEOS-Chem for the comparable simulations. This persistent difference may indicate that some fastreacting precursors are missing from the model, and/or that some fast reaction pathways may be missing from the models.

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