Trends in emissions and concentrations of air pollutants in the lower troposphere in the Baltimore/Washington airshed from 1997 to 2011

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Abstract. Trends in the composition of the lower atmosphere (0–1500 m altitude) and surface air quality over the Baltimore/Washington area and surrounding states were investigated for the period from 1997 to 2011. We examined emissions of ozone precursors from monitors and inventories as well as ambient ground-level and aircraft measurements to characterize trends in air pollution. The US EPA Continuous Emissions Monitoring System (CEMS) program reported substantial decreases in emission of summertime nitrogen oxides (NOx) from power plants, up to ~80% in the mid-Atlantic States. These large reductions in emission of NOx are reflected in a sharp decrease of ground-level concentrations of NOx starting around 2003. The decreasing trend of tropospheric column CO observed by aircraft is ~0.8 Dobson unit (DU) per year, corresponding to ~35 ppbv yr−1 in the lower troposphere (the surface to 1500 m above ground level). Satellite observations of long-term, near-surface CO show a ~40% decrease over western Maryland between 2000 and 2011; the same magnitude is indicated by aircraft measurements above these regions upwind of the Baltimore/Washington airshed. With decreasing emissions of ozone precursors, the ground-level ozone in the Baltimore/Washington area shows a 0.6 ppbv yr−1 decrease in the past 15 yr. Since photochemical production of ozone is substantially influenced by ambient temperature, we introduce the climate penalty factor (CPF) into the trend analysis of long-term aircraft measurements. After compensating for inter-annual variations in temperature, historical aircraft measurements indicate that the daily net production of tropospheric ozone over the Baltimore/Washington area decreased from ~20 ppbv day−1 in the late 1990s to ~7 ppbv day−1 in the early 2010s during ozone season. A decrease in the long-term column ozone is observed as ~0.2 DU yr−1 in the lowest 1500 m, corresponding to an improvement of ~1.3 ppbv yr−1. Our aircraft measurements were conducted on days when severe ozone pollution was forecasted, and these results represent the decreasing trend in high ozone events over the past 15 yr. Back trajectory cluster analysis demonstrates that emissions of air pollutants from Ohio and Pennsylvania through Maryland influence the column abundances of downwind ozone in the lower atmosphere. The trends in air pollutants reveal the success of regulations implemented over the past decades and the importance of region-wide emission controls in the eastern United States.

1 Introduction

Ozone controls much of the chemistry in the lower atmosphere, such as hydroxyl radical (OH) production and the lifetimes of atmospheric species including methane (CH4), carbon monoxide (CO), and volatile organic compounds (VOCs) (Levy, 1971; Logan et al., 1981; Seinfeld and Pan-
Tropospheric ozone is a good absorber of thermal radiation, acting as the third most important anthropogenic contributor to radiative forcing of climate (Fishman et al., 1979a; Ramanathan and Dickinson, 1979; IPCC, 2007). High concentrations of ground-level ozone also threaten human health (WHO, 2003; Anderson, 2009; Jerrett et al., 2009) and cause damage to ecosystems (Adams et al., 1989; Chameides et al., 1999; Ashmore, 2005). Thus, ozone is one of the six criteria pollutants regulated by the US Environmental Protection Agency (EPA) through National Ambient Air Quality Standards (NAAQS).

As a secondary air pollutant, the majority of ozone in the lower troposphere is produced by photochemical reactions involving CO and VOCs in the presence of nitrogen oxides (NOx) (Crutzen, 1974; Fishman et al., 1979b; EPA, 2006), while troposphere-stratosphere exchange of air contributes substantial amounts of ozone to the upper troposphere (Levy et al., 1985; Holton et al., 1995; Wild et al., 2003; Stevenson et al., 2006). In the US, ozone pollution drew public attention starting with the “Los Angeles smog” events in the 1950s (Haagensmit, 1952; Haagensmit and Fox, 1956). Emissions of ozone precursors, mainly NOx and VOCs from anthropogenic sources, have decreased significantly under regulation, resulting in lower ground-level ozone (Cooper et al., 2012). Long-term records of tropospheric ozone in the US have been investigated through ground-level observations (Oltmans et al., 2006; Jaffe and Ray, 2007; Lefohn et al., 2008, 2010; Oltmans et al., 2013), however trends of ozone are not monotonic (Cooper et al., 2012; Oltmans et al., 2013). Analysis of these long-term ozone measurements can also shed light on the change of “baseline” ozone in order to estimate the flux of air pollutants entering and exiting North America, relevant to regulation of ambient air quality (Oltmans et al., 2008; Parrish et al., 2009; Chan and Vet, 2010; Cooper et al., 2010; Lin et al., 2012).

In the mid-Atlantic States, ozone is a major air pollutant during summer (EPA, 2012a). The Baltimore/Washington area has failed to meet mandated attainment deadlines (EPA, 2012b); despite tremendous improvements in local air quality over the past several decades, a significant number of exceedances of the current 75 parts per billion by volume (ppbv), 8 h surface ozone standard, were experienced. The design value of daily maximum 8h ozone for Edgewood, MD (generally downwind of Baltimore), was 92 ppbv in 2011 (http://www.epa.gov/airtrends/values.html).

The Regional Atmospheric Measurement Modeling and Prediction Program (RAMMPP, http://www.atmos.umd.edu/~RAMMPP) led by the University of Maryland was created to conduct state-of-the-art scientific research pertaining to surface ozone, through in situ measurements, air quality forecasting, mesoscale dynamics modeling, and chemical transport modeling. As a key component of RAMMPP, aircraft measurements of air pollutants have been conducted during the ozone season (May to September) over the past 15 yr (Taubman et al., 2006; Hains et al., 2008). At present, the RAMMPP aircraft has the capability to measure CO, O3, sulfur dioxide (SO2), nitrogen dioxide (NO2), aerosol absorption, aerosol scattering, particle counts, aerosol size distribution, as well as meteorological variables.

The RAMMPP aircraft program started with measurements of tropospheric O3 and CO in 1997, providing a 15 yr record of summertime tropospheric ozone for investigating summertime air quality trends in the Baltimore/Washington airshed. Many air quality studies focus on the trends in ground-level ozone (Fiore et al., 1998; Jaffe and Ray, 2007; Lefohn et al., 2008, 2010; Cooper et al., 2012). In the free troposphere (FT), ozone and its precursors aloft have longer lifetimes and can be transported farther downwind than constituents near the surface (Luke et al., 1992; Jaffe, 2011; Newman et al., 2012). The Baltimore/Washington nonattainment area is downwind of the Ohio River valley, where a large number of power plants are located, and upwind of other nonattainment areas such as Philadelphia, New Jersey and New York City. Previous studies using ozonesonde measurements have revealed the importance of regional transport and Chesapeake Bay meteorological effects on ozone pollution in the Baltimore/Washington area (Yorks et al., 2009; Martins et al., 2013; Stauffer et al., 2013). Distinguishing local ozone production from regional transport is critical for effective policy and control measures.

We quantify trends in summertime air pollution over the past 15 yr using measurements of tropospheric ozone and its precursors (CO and NOx) in the mid-Atlantic region and upwind states, as well as national and regional emissions data. Section 2 presents the data sets and methods. In Sect. 3, we discuss the trends in emissions near the Baltimore/Washington airshed, and the trends in tropospheric O3 and CO observed by the RAMMPP aircraft and EPA Air Quality System (AQS) network. A case study on regional transport is conducted using a back trajectory clustering technique to reveal the relationship between upwind anthropogenic emissions and downwind air quality. Finally, we discuss the importance of emission regulation for improving ozone pollution in the nonattainment Baltimore/Washington airshed, as well as implications for future control measures.

2 Data and methods

For our study of ozone pollution in the Baltimore/Washington airshed, two emissions data sets were used: the EPA National Emissions Inventory (NEI) and the CEMS data for states of Maryland (MD), Virginia (VA), West Virginia (WV), Pennsylvania (PA) and Ohio (OH) (the mid-Atlantic and one upwind state, hereafter named the research domain). The NEI emissions inventory, a comprehensive estimate of emissions for both criteria and hazardous air pollutants from all sources on the county scale, is prepared every three years, based on emission survey and specialized emission modeling tools (e.g.,
http://www.epa.gov/ttnchie1/net/2008inventory.html). In the NEI, emissions are grouped into five categories: point, nonpoint, onroad, nonroad, and event (details given in Table S1 of the Supplement). Here, detailed NEI data (2002, 2005, and 2008) and summaries of annual national data (available at: http://www.epa.gov/ttn/chief/trends/index.html) were utilized to study the trends of ozone precursor emissions from point and mobile sources. The NEI only provides annual emission estimates. To investigate daily changes of emissions, we examined CEMS data from continuous monitoring of point sources; i.e., major power plants and industries, covered by the EPA Clean Air Markets Division (data available at: http://ampd.epa.gov/ampd/). The CEMS data include NOx, SO2 and carbon dioxide (CO2) emissions, with temporal resolution as fine as hourly from individual point source. Previous studies indicate that NOx emissions from point sources substantially influence surface ozone (Frost et al., 2006; Kim et al., 2006; Hains et al., 2008). Here, daily CEMS data for NOx emissions from sources within the research domain from 1997 to 2011 were used.

Measurements of NOx, CO and O3 from ground stations throughout the Baltimore/Washington area archived on the EPA AQS website (http://www.epa.gov/ttn/airs/airsaqs/detaildata/downloadaqsdata.htm) were used to quantify trends in surface air pollution. NOx measurements provided by the EPA are achieved through conversion of NO2 to NO using a hot molybdenum catalyst, followed by the chemiluminescence reaction of NO with O3. Interferences from other oxidized reactive nitrogen species such as peroxyacyl nitrates (PAN) and alkylnitrates are almost certain (Fehsenfeld et al., 1987; Dunlea et al., 2007). Correcting the EPA measurements of NOx for potential interferences is beyond the scope of this study. Any interfering compounds originate from chemical reactions involving NOx, so the EPA NOx record is consistent over the course of this study and almost certainly provides a surrogate for trends in actual ambient NOx.

Since CO is emitted mainly by automobiles and incomplete combustion such as biomass burning, anthropogenic emissions of CO are not monitored as are NOx and SO2 emissions from point sources in the CEMS program. The EPA estimates national emissions of CO using models such as the Motor Vehicle Emissions Simulator (MOVES). MO-PITT (Measurements of Pollution In the Troposphere) on Terra (Deeter et al., 2003), launched in 1999, has together with measurements from numerous other satellite instruments proven useful for tracking the long-term trend of tropospheric CO on regional scales (Worden et al., 2013). A ∼15 % decrease was reported for column of CO over the eastern US between 2000 and 2011. Here we used the MOPITT version 5 level 3 monthly products that exploit both near and thermal infrared radiances (MOP03JM.005, https://eosweb.larc.nasa.gov/project/mopitt/mopitt_table). We selected the CO values from the surface to 900 hPa (Deeter et al., 2012) to compare with RAMMPP aircraft measurements of CO in the lower troposphere. We note that although MOPITT multispectral retrievals have highly variable sensitivity to near-surface CO over different surface types, retrievals over the mid-Atlantic region considered in this study showed consistently good sensitivity to near-surface CO (Worden et al., 2010). For MOPITT data used in this analysis, the sensitivity to CO typically peaks in the surface layer from 1000 to 900 hPa and falls off exponentially with a scale height of ∼1 km, making them suitable for comparison with our aircraft measurements.

RAMMPP aircraft observations of tropospheric O3 and CO date back to 1997, providing a 15 yr record of summertime air quality. The sampling platform for the early 2000s was discussed extensively in previous studies (Taubman et al., 2006; Hains et al., 2008); details of the current sampling platform are listed in Table S2 of the Supplement. Here, a brief summary of O3 and CO measurements is provided. Ozone is measured using a commercially available analyzer (Model 49/49C, Thermo Environmental Instruments, TEI, Franklin Massachusetts), based on the absorption of ultraviolet radiation. The analyzer is routinely serviced and calibrated with an in-house primary ozone calibrator (TEI Model 49PS) using zero-grade air. The instrument has been compared to the National Institute of Standard and Technology (NIST) standard, and the precision can reach 1 ppbv for 10 s average data (Taubman et al., 2006). Observations of ambient CO are conducted using a modified commercially available nondispersive infrared gas filter correlation analyzer (TEI Model 48) with enhanced precision of about 30 ppbv for 1 min moving average of 10 s data (Dickerson and Delany, 1988). The detector is regularly calibrated with a NIST traceable CO gas standard (Scott Marrin Inc., Riverside, CA). Processed airborne measurements are archived at http://www.atmos.umd.edu/~RAMMPP/archives/ArchiveFlightData.html (access to this website is available upon request).

RAMMPP flight plans are based on air quality forecasts, issued by the Maryland Department of the Environment (MDE), usually for days with predicted poor air quality. Most of the research flights were performed on hot summer days with weak surface winds and/or stagnation. In the past 15 yr, around 1000 research profiles (defined as a spiral over a fixed location to measure the vertical distribution of air pollutants, hereafter named research spirals) were carried out over more than 100 airports from Georgia to Vermont. Here we examine ∼500 research spirals over more than 20 airports in the Baltimore/Washington airshed. Figure 1 shows the locations of the selected airports and a typical flight route. On average, ∼40 research spirals were conducted each year (except 2006), providing robust statistics for long-term trend analysis (the number of research spirals per year is given in Fig. S1 of the Supplement).

To quantify the effects of regional transport and photochemical production of tropospheric ozone in the Baltimore/Washington airshed, two flights are conducted during
a typical day: one flight over the upwind area in the morning and one over the downwind area in the afternoon. On a typical westerly transport day (Fig. 1), the morning flight samples air over Luray, VA (38.67° N, 78.50° W, “Lu”, 65 spirals), Winchester, VA (39.14° N, 78.14° W, “Wi”, 57 spirals), Cumberland, MD (39.62° N, 78.76° W, “Cu”, 71 spirals), Harford County, MD (39.57° N, 76.20° W, “Ha”, 64 spirals), and Easton, MD (38.80° N, 76.07° W, “Ea”, 72 spirals).

Fig. 1. Locations of the selected RAMMPP research spirals in the Baltimore/Washington airshed. Cyan and Green lines show the route of morning and afternoon flights, respectively, during a typical westerly transport flight pattern. Blue and Red dots show the locations of morning and afternoon spirals, respectively. Five airports (the spiral locations) extensively covered by this flight pattern are, from the lower left, clockwise: Luray, VA (38.67° N, 78.50° W, “Lu”, 65 spirals), Winchester, VA (39.14° N, 78.14° W, “Wi”, 57 spirals), Cumberland, MD (39.62° N, 78.76° W, “Cu”, 71 spirals), Harford County, MD (39.57° N, 76.20° W, “Ha”, 64 spirals), and Easton, MD (38.80° N, 76.07° W, “Ea”, 72 spirals).

3 Trends of emissions and RAMMPP aircraft FV measurements

3.1 Trends of NEI and CEMS emissions

NO$_x$ and CO are important anthropogenic precursors of tropospheric ozone. Onroad and nonroad internal combustion sources release the majority of anthropogenic CO and around half of the anthropogenic NO$_x$, with the other half emitted from point sources such as electricity generating units, construction equipment, and other industrial sources. In this study, CO and NO$_x$ emissions estimates from onroad and nonroad sources were obtained from the NEI emission inventory; daily NO$_x$ emissions from point sources were acquired from CEMS measurements.

The EPA NEI program provides two emission estimates: (1) a comprehensive and detailed estimate of air pollution emissions from different sectors at county level every three years; and (2) annual estimates of national emissions, used to define national trends. The national emissions trends in mobile sources between 1997 and 2011 (CO and NO$_x$) were scaled from total emissions inside the research domain based on the detailed inventory years of 2002, 2005, and 2008, in order to estimate the trends in mobile source emissions near the Baltimore/Washington airsheded during the past 15 yr. We found that the research domain accounts for $\sim 4$ and $\sim 5\%$ of the national NO$_x$ and CO emissions, respectively (Fig. S2 in the Supplement). Between 1997 and 2011, large decreases of NO$_x$ and CO emissions, $\sim 50$ and $\sim 60\%$, respectively (using 1997 emissions as the reference values), were observed in the US (Fig. S2 in the Supplement). Here, we assume that onroad and nonroad emissions have the same trends in the research domain as for the entire US; consequently, the large reductions of precursor emissions should affect tropospheric ozone in the Baltimore/Washington airshed.
Long-term trends in monthly CEMS data (Fig. 2a) show that NO\textsubscript{x} emissions from point sources decreased by \(\sim 80\%\) from 1997 to 2011 for each state, except for PA, where they decreased by only \(\sim 50\%\) (using 1997 emissions as a baseline). Several characteristics are observed: (1) between 1997 and 2002, a gradual decrease of NO\textsubscript{x} emissions for all states; (2) a large reduction after 2003, due to the EPA NO\textsubscript{x} State Implementation Plan (SIP) Call; (3) a square wave pattern in NO\textsubscript{x} emissions from OH, PA, and WV between 2003 and 2009 when NO\textsubscript{x} in these states was only controlled during the ozone season; and (4) through 2003–2008 further slow decreases for all states, followed by substantial reductions in MD, OH, and WV, and slight increases in PA and VA after 2009. During the same period, CO\textsubscript{2} emissions from these point sources, representing the total amount of fossil fuel burned, stayed flat (Fig. S3 in the Supplement). Emissions of NO\textsubscript{x} from point sources are regulated by the EPA NO\textsubscript{x} Budget Trading Program (NBP), later superseded by the Clean Air Interstate Rule (CAIR) (EPA, 2009; Butler et al., 2011), which covers 27 eastern states and the District of Columbia (http://www.epa.gov/cair/). These EPA programs regulated power plant emissions in the past 15 yr, and the NO\textsubscript{x} SIP call substantially reduced the NO\textsubscript{x} emissions from point sources in the research domain in 2003.

Figure 2b presents locations of the major point sources of NO\textsubscript{x} in the Ohio River valley and western PA. These air pollutants released upwind of the Baltimore/Washington airshed exert a significant influence on downwind air quality under the prevailing westerly winds during the ozone season (Marufu et al., 2004; Frost et al., 2006; Kim et al., 2006). We also computed the ratio of NO\textsubscript{x} and CO\textsubscript{2} emissions for each source. Figure 2c shows that the point sources with high NO\textsubscript{x}/CO\textsubscript{2} ratios exist mainly in the Ohio River valley, and these “hot spots” (details given in Table S3 in the Supplement) are suitable targets for future NO\textsubscript{x} emission reductions.

3.2 Trends of ground-level observations

To study ozone photochemistry, we analyzed the EPA CO, NO\textsubscript{x}, and O\textsubscript{3} observations in the afternoon (12:00–18:00 LT (local time)) during the ozone season. O\textsubscript{3} monitors are located throughout this area, while CO and NO\textsubscript{x} observations are mainly along the Interstate 95 corridor between Baltimore and Washington, DC (Fig. S4 in the Supplement). Figure 3 shows the long-term trends of annual mean ground-level concentrations of air pollution during the summer. The long-term ozone measurements show large interannual variations, because ozone production is not only determined by emissions but also weather, especially temperature (EPA, 2006). A discernible decrease (\(\sim 0.6\) ppbv yr\(^{-1}\), Fig. 4) is observed, suggesting a general decreasing trend of ambient ozone in the Baltimore/Washington area, similar to decreasing trends reported across the eastern US (Lefohn et al., 2010; Cooper et al., 2012).

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Long-term ground-level CO observations show a non-monotonic but steadily decreasing trend from \( \sim 600 \) to \( \sim 350 \) ppbv or \( \sim 18 \) ppbv yr\(^{-1} \) decrease (linear regression coefficient \( r = -0.90 \)). If the regional baseline CO mixing ratio is assumed to be \( \sim 150 \) ppbv, then the reduction in the added regional CO mixing ratio is from \( \sim 450 \) to \( \sim 200 \) ppbv, corresponding to a \( \sim 55 \% \) decrease since 1997. During the same period, the NEI annual CO emissions decreased from \( \sim 13.5 \times 10^6 \) to \( \sim 5.5 \times 10^6 \) tons (Fig. S2 in the Supplement), a \( \sim 59 \% \) decrease since 1997. Therefore, after correcting for baseline CO, the trends of ground-level CO observations and the NEI CO emissions are similar.

Long-term ambient NO\(_x\) measurements demonstrated a sharp decrease (from \( \sim 20 \) to \( \sim 8 \) ppbv) after 2003, while annual NEI emissions (CEMS data plus automobile emissions) declined gradually. This sharp decrease coincides with the large reduction of point source emissions of NO\(_x\) during the EPA NO\(_x\) SIP call (Fig. 2a). Here we estimate daily mean NO\(_x\) emissions during the ozone season by incorporating daily CEMS emissions (point sources) and scaled mobile emission estimates from national NEI annual data (4 % of the total emissions in the US). Figure 5 indicates that total emissions from both point sources and mobile sources decreased significantly, \( \sim 60 \% \) and \( \sim 50 \% \), respectively. Point source emissions exhibited a sharp decrease around 2003 due to the EPA NO\(_x\) SIP call, while mobile emissions decreased gradually. The contribution of NO\(_x\) from point sources in this region to the total emissions decreases from \( \sim 70 \% \) in the late 1990s to \( \sim 50 \% \) in the early 2010s, which confirms the efficacy of the EPA NO\(_x\) SIP call.

Long-term aircraft measurements show that tropospheric ozone has large day-to-day variations; to illustrate this, all aircraft measurements of tropospheric ozone obtained in the summer of 2001 are shown in Fig. 7. In the lower atmosphere, ozone is reasonably well mixed with a local maximum near 1000 m AGL, around the top of planetary boundary layer (PBL). In the morning upwind flights (Fig. 7b), ozone mixing ratios near the surface are relatively low compared with the values in the FT (higher than 1000 m), suggesting that ozone is lost due to dry deposition and reactions with NO\(_x\) during the night. With low photochemical pro-

**Fig. 3.** Long-term trends of ground-level O\(_3\), CO, and NO\(_x\) observed in the Baltimore/Washington area in EPA AQS sites (sites shown in Fig. S4 of the Supplement). CO mixing ratios have been scaled by 10 for clarity. To investigate the ozone chemistry, measurements in the afternoon (12:00–18:00 LT) during the ozone seasons were used to calculate annual means.

**Fig. 4.** Linear regression analysis of long-term EPA AQS observations of surface ozone in the Baltimore/Washington area. Ozone data are the same as in Fig. 3; and the linear regression of ozone is with respect to year.

We compared the CEMS, scaled NEI, and total NO\(_x\) emissions with EPA AQS observations using linear regression analysis (Fig. 6), revealing good correlation between these emission estimates and ground-level NO\(_x\) observations. The slope of observations versus emissions is 0.0042 ppbv ton\(^{-1}\) for CEMS and 0.0024 ppbv/ton for NEI mobile emissions alone. Therefore, if we assume emissions from point sources and mobile sources have an equivalent effect on surface concentrations of NO\(_x\), point sources contribute \( \sim 60 \% \) of the ambient ground-level NO\(_x\) concentrations. The daily average CEMS measurements provide a unique opportunity to investigate the effects of emissions on surface NO\(_x\) observations. Considering that ground-level NO\(_x\) concentrations are controlled by other parameters such as advection, temperature, turbulence, and concentration of VOCs, the CEMS data correlate surprisingly well with observed ground-level NO\(_x\) concentrations.

**3.3 Trends of aircraft measurements**

**3.3.1 Vertical distribution of tropospheric O\(_3\) and CO**

Long-term aircraft measurements show that tropospheric ozone has large day-to-day variations; to illustrate this, all aircraft measurements of tropospheric ozone obtained in the summer of 2001 are shown in Fig. 7. In the lower atmosphere, ozone is reasonably well mixed with a local maximum near 1000 m AGL, around the top of planetary boundary layer (PBL). In the morning upwind flights (Fig. 7b), ozone mixing ratios near the surface are relatively low compared with the values in the FT (higher than 1000 m), suggesting that ozone is lost due to dry deposition and reactions with NO\(_x\) during the night. With low photochemical pro-
production rates and weak mixing in the morning, ozone mixing ratios stay low until close to noon. Relatively high concentrations of ozone observed in the FT are in the residual layer, reflecting ozone production on previous days (Yorks et al., 2009). With the rise of the PBL during daytime, air in the residual layer is entrained and mixed, transporting the residual-layer pollutants downward. In the afternoon, tropospheric ozone is produced through photochemical reactions throughout the PBL and lower FT; the mean ozone altitude profile shows a quasi-uniform concentration (Fig. 7c). The afternoon mean ozone profile also has a local maximum at \( \sim 1000 \text{ m} \), with ozone \( \sim 18 \text{ ppbv} \) higher than the ground-level value. Pollutants aloft observed in both morning and afternoon do not immediately influence local air quality, but transport of these pollutants exerts significant effects on air pollution in downwind areas.

We conducted a similar analysis on all the CO profiles obtained in 2001 (Fig. 8). A relatively uniform mixing ratio of \( \sim 250 \text{ ppbv} \) CO was observed in the FT, and maximum CO concentrations were observed near the surface. The CO mixing ratios observed in the FT during summer 2001 are consistent with previous observations at high elevation in western VA (Hallock-Waters et al., 1999). These measurements reflect atmospheric CO sources on the Earth’s surface and on average an exponential decrease within the PBL. With a lifetime of about one month, CO can be transported upward and well mixed in the FT, so CO is a good chemical tracer to investigate advection in the lower atmosphere (Castellanos et al., 2011; Loughner et al., 2011). The mean CO profiles from upwind and downwind research spirals exhibit similar vertical distributions (Fig. S5 in the Supplement), indicating that CO pollution in the Baltimore/Washington airshed is a regional problem.

### 3.3.2 Estimate of photochemical production of tropospheric ozone

Ozone production is closely related to temperature (EPA, 2006; Camalier et al., 2007; Bloomer et al., 2009; Jacob and Winner, 2009), so we must take the inter-annual variations of ambient temperature into account to study the long-term trend of tropospheric ozone. Here, we use the climate penalty factor (CPF), defined as \( \frac{\partial \text{O}_3}{\partial T} \). One previous study reported CPF as \( \sim 3.2 \text{ ppbv} \text{ °C}^{-1} \) prior to 2002 and \( \sim 2.2 \text{ ppbv} \text{ °C}^{-1} \) after 2002 in the eastern US (Bloomer et al., 2009). Here, we first calculated a “standard” temperature vs. altitude profile by averaging all the temperature profiles. We then normalized ozone mixing ratio profiles by applying the CPF (3.2 ppbv °C\(^{-1}\) before 2002 and 2.2 ppbv °C\(^{-1}\) after 2002) to the differences from the “standard” temperature profile and calculated column ozone.

Figure 9 shows the long-term trend of lower tropospheric ozone column abundances after adjustment for variations in temperature using the CPF. The upwind ozone column abundances are taken as the baseline ozone profile entering the Baltimore/Washington area, with ozone mainly in the residual layer; the downwind column abundances represent the total of baseline ozone plus ozone generated by local photochemical reactions. The afternoon flights usually finish around 16:00 LT, when ozone reaches its maximum, so the difference between morning and afternoon flights represents the daily production of tropospheric ozone.
in the Baltimore/Washington area. Negative values are observed for 1999 and 2008, when a relatively small number of research flights were conducted. The mean net production in the past 15 yr is estimated as $2.3 \pm 2.0 \, \text{DU day}^{-1}$ (1 DU $= 2.69 \times 10^{16} \, \text{mol cm}^{-2}$), where 2.0 DU represents the ±1 σ variability. When assuming ozone is well-mixed in the lower 1500 m, this value equals $\sim 15 \pm 13 \, \text{ppbv}$ of ozone production.

It is notable that the net daily production of ozone decreased from $\sim 3.0 \, \text{DU day}^{-1}$ in the late 1990s to $\sim 1.0 \, \text{DU day}^{-1}$ in the early 2010s, corresponding to $\sim 20$ to $\sim 7 \, \text{ppbv day}^{-1}$, respectively. This shows that the daily ozone production within the Baltimore/Washington area has decreased significantly, $\sim 60 \%$ in total or $\sim 1.0 \, \text{ppbv yr}^{-1}$. This magnitude of decrease is higher than the decreasing trends observed in ground-level measurements (Fig. 4). The ground-level measurements were collected during the whole ozone season, while the RAMMPP research flights were only conducted on air pollution action days when the transport is forecasted to play an important role in the episodic ozone pollution, either under westerly transport or southerly transport patterns. These results suggest that ozone transported into the Baltimore/Washington airshed plays an important role in determining downwind ozone pollution; likewise, emissions in Baltimore can impact air quality over downwind areas such as Philadelphia. Since most of the RAMMPP research flights were conducted on poor air quality days, our estimates can be treated as the upper limit of net daily production of ozone in the Baltimore/Washington airshed.

### 3.3.3 Long-term trends of tropospheric O$_3$ and CO

Here we focus on O$_3$ and CO measurements on afternoon flights, when maximum ground-level ozone is usually observed. The annual mean O$_3$ and CO column abundances are plotted, showing their long-term trends in Fig. 10. We used the Savitzky–Golay Smoothing technique (Savitzky and Golay, 1964; Bromba and Ziegler, 1981; Gorry, 1990), which preserves the temporal features of the data while reducing the influences of noise and missing data. Figure 10a shows a $\sim 0.2 \, \text{DU yr}^{-1}$ decrease of ozone column abundances. Assuming ozone in the lower 1500 m is well mixed, this decrease of column ozone is equivalent to $\sim 1.3 \, \text{ppbv yr}^{-1}$ throughout this part of the atmosphere. This value is much higher than a 0.4 to $\sim 0.6 \, \text{ppbv yr}^{-1}$ decrease of ground-level ozone reported in the eastern US (Bloomer et al., 2010; Cooper et al., 2012). The explanation of this difference could be that RAMMPP research flights were usually conducted on air quality action days, suggesting that for these conditions (i.e., meteorological conditions conducive to poor air quality) ozone pollution has improved more than under others. Cooper et al. (2012) observed a $\sim 15 \, \text{ppbv}$ decrease in the 95th percentile of eastern US summertime ozone from the early 1990s to the late 2000s. These results are consistent with the observed decrease in ozone production.
Fig. 8. Vertical distributions of tropospheric CO over the Baltimore/Washington region in 2001 (all spirals). The 25th percentile, mean, and 75th percentile are shown as in Fig. 7.

tent with the results from our long-term RAMMPP aircraft measurements.

To test the effects of variations of ambient temperature, we also calculated the trends in long-term ozone column abundances without applying the CPF (Fig. 10b). The difference between the ozone column abundances with and without CPF adjustment is discernible, especially for years 2009 to 2011. Figure 10b shows a monotonic increase of column ozone from 2009 to 2011, suggesting an increasing trend. However, the statistics of daily mean temperature show that the summer of 2009 has far less hot days, especially when compared with the hot summers of 2010 and 2011 (Fig. S6 in the Supplement). After compensating for the effects of ambient temperature with the CPF, the ozone column abundances in 2010 and 2011 decreased, and the increasing trend in ozone over those years disappeared.

The long-term CO column abundances show a $\sim 0.8$ DU yr$^{-1}$ decrease (Fig. 10c). As shown in Fig. 8, CO concentrations decay exponentially above the surface within the PBL to a near constant value aloft. So the $0.8$ DU yr$^{-1}$ decrease in column CO can be interpreted as a $\sim 35$ ppbv yr$^{-1}$ decrease at the ground level, less than the nationwide decrease (data available at http://www.epa.gov/airtrends/). One possible explanation is that the national trend is calculated based on the 2nd maximum of annual 8 h average, i.e., measurements in CO episodes, which usually arise on cold winter days with a temperature inversion. RAMMPP research flights are normally carried out during the ozone season, when CO pollution is not as severe and strong convection can transport CO upward. Figure 10 also shows that O$_3$ and CO column abundances are correlated, in particular after the EPA NO$_x$ SIP call. For instance, in 2009, both high CO and high O$_3$ column abundances were observed. These large inter-annual variations in column CO cannot be explained by NEI emissions trends (Fig. S2) and have not been observed by ground-level observations (Schwab et al., 2009). CO is usually emitted alongside with NO$_x$ from automobile emissions or other incomplete combustion such as biomass burning. Therefore, a high concentration of CO might be an indicator of high NO$_x$, which would also boost the production of ozone in the largely NO$_x$-limited eastern US. Airborne NO$_2$ measurements were not available from RAMMPP aircraft before 2011. Thus, this hypothesis of intercepting automobile exhaust or wildfire plumes cannot be verified. However, when we used the 7 yr Savitzky–Golay method to smooth the data, the $\sim 0.8$ DU yr$^{-1}$ decrease, $\sim 15\%$ relatively, can be treated as the long-term trend of column CO in the lower atmosphere, subject to the uncertainties associated with limited airborne sampling.

To investigate the long-term CO trends from space, we used the monthly MOPITT near-surface CO retrievals from 2000 to 2011 (Deeter et al., 2003, 2012). Since the MOPITT Equator crossing time is around 10:30 LT, we used the morning RAMMPP CO column abundances observed in western MD. Only data in summer (JJA) were selected to reduce the effect of CO seasonal variations. Figure 11 shows the observed annual mean. The oxidation of VOCs is an important source of tropospheric CO; in the eastern US, biogenic VOC emissions such as isoprene dominate (Kesselmeier and Staudt, 1999). We assume that the amount of CO oxidized from VOCs has not changed in the past 15 yr, so the CO trends are only discussed with reference to anthropogenic CO emissions in this study. Linear regression analyses of long-term EPA AQS observations, RAMMPP aircraft measurements, MOPITT observations, and NEI emissions were conducted, with high correlation of concentrations ($Y$) as a
function of time ($X$): for AQS ($r = -0.89$), MOPITT observations ($r = -0.86$) and NEI emissions ($r = -0.99$). However, the correlation is low for RAMMPP aircraft measurements in the morning, which could be caused by the sampling uncertainty mentioned above. The overall reduction (relative to year of 2000) is estimated as $\sim 40\%$ ($\sim 3.5\%$ yr$^{-1}$) for AQS observations, RAMMPP measurements, and MOPITT observation, compared to $\sim 60\%$ ($\sim 4.5\%$ yr$^{-1}$) for the NEI emissions. The decreasing trend of MOPITT CO observed over western MD, $\sim 3.5\%$ yr$^{-1}$, is higher than the value ($\sim 1.5\%$ yr$^{-1}$) reported in the eastern US (Worden et al., 2013). However, in Worden et al. (2013), MOPITT total column CO data in all seasons were analyzed, while this study only focused on summertime observations of near-surface CO. We would expect to see a larger trend in near-surface concentrations compared to total column CO because the near-surface CO is substantially influenced by surface-based pollution sources (Fig. 8).

4 Clustering analysis on the regional transport of CEMS emissions

The CEMS program monitors emissions from point sources equipped with tall smokestacks, such as power plants and industrial boilers. A study conducted by the US Government Accountability Office (US GAO) tracked 284 tall smokestacks at 172 coal-fired power plants in 34 states, of which 205 are 62.5 to 213 m tall, 63 are 213 to 305 m tall, and 12 are higher than 305 m (GAO, 2011). About one-third of these tall stacks are located in PA, WV and OH along the Ohio River valley. Such high smokestacks can disperse air pollutants over great distances. Since the Baltimore/Washington region is often downwind of the Ohio River valley, emissions from point sources are expected to exert substantial influence on downwind air quality. We aggregated the CEMS emission data by state, because point sources are regulated by each state.

We studied regional transport with a hierarchical clustering technique using measurements from 63 RAMMPP research spirals over Harford County airport from 1998 to 2011. We utilized the National Oceanic and Atmospheric Administration (NOAA) Air Resource Laboratory (ARL) Hybrid Single-Particle Lagrangian Integrated Trajectory (HYSPLIT) model (version 4, http://www.arl.noaa.gov/readyyysplit.html) to calculate 48 h back trajectories. The time of each aircraft spiral was used to initialize each back trajectory with release heights of 500, 1000, and 1500 m AGL, respectively. Details on the model and meteorological fields are given in Table S4 of the Supplement. The accuracy of these trajectories is adequate to distinguish various transport pathways of air pollutants that reach the Baltimore/Washington airshed (Stohl et al., 1995; Stohl, 1998).

Location and altitude information of each HYSPLIT back trajectory was archived and back trajectories ending at 1000 m over Harford County airport were classified based on air mass origin, as shown in Fig. 12. Three major clusters of air mass origin, denoted OH/PA, PA, and WV/VA, arose from the analysis. Two minor clusters, recirculation and stagnation (containing less than 10 members each), are embedded underneath these spaghetti-like clusters (Table 1). Here we establish a “conceptual” model to evaluate the relationship between the upward power plant emissions of NO$_x$ and downwind ozone pollution. Previous studies suggest that single back trajectories might not be sufficient to relate the upward sources to downwind pollution (Stohl et al., 2002, 2003). For this analysis, we focus on transport from upward sources over a 24–48 h period of time, much shorter than the period of time considered by Stohl et al. (2002, 2003). Our comparison of single back trajectories and 27 member back trajectory ensembles from HYSPLIT demonstrates that single back trajectories can accurately represent the ensemble behavior over the period of time important for our analysis (Fig. S7 in the Supplement). Precise determination of the relationship between upward sources and downwind pollution is complicated by the release altitude of NO$_x$ from power plants mostly at 200–300 m AGL (GAO, 2011), followed by a small-scale, rapid-plume rise that is difficult to quantify. Nonetheless, we have used HYSPLIT to show that the 24 h backward concentration dispersion of air pollutants observed by the RAMMPP aircraft occurs for emissions over a region comparable to the area of a state (Fig. S8 in the Supplement).

**Table 1.** Characteristics and emission estimates of each cluster. Correlation coefficients are for lower tropospheric column ozone over Maryland, regressed against emissions.

<table>
<thead>
<tr>
<th>Cluster</th>
<th>Upwind states</th>
<th>Emissions estimate</th>
<th>$r^b$</th>
<th>$n^c$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>OH and PA</td>
<td>$MD_l + 0.5PA_{l-1} + 0.25OH_{l-1}$</td>
<td>0.86</td>
<td>18</td>
</tr>
<tr>
<td>2</td>
<td>PA</td>
<td>$MD_l + 0.5PA_{l-1}$</td>
<td>0.16</td>
<td>22</td>
</tr>
<tr>
<td>3</td>
<td>WV or VA</td>
<td>$MD_l + 0.5VA_{l-1}$ or $MD_l + 0.5WV_{l-1}$</td>
<td>-0.08</td>
<td>12</td>
</tr>
<tr>
<td>4</td>
<td>Recirculation</td>
<td>$MD_l + 0.5 VA_{l-1} + 0.5 PA_{l-1}$</td>
<td>0.53</td>
<td>4</td>
</tr>
<tr>
<td>5</td>
<td>Stagnation$^a$</td>
<td>$MD_l$</td>
<td>0.04</td>
<td>7</td>
</tr>
</tbody>
</table>

$^a$ The background flight (conducted on 3 September 2009) was incorporated into the stagnation cluster because it shares the same estimate of upwind emissions; $^b$ linear regression coefficient $r$ is calculated for each group in Fig. 13a; $^c$ number of research spirals.
Fig. 10. Long-term trends of annual mean afternoon O\textsubscript{3} and CO column abundances (from the surface to 1500 m AGL) over the region downwind of the Baltimore/Washington area: (a) column O\textsubscript{3} with CPF adjustment; (b) column O\textsubscript{3} without CPF adjustment; (c) column CO. Error bars represent one standard deviation. The red lines are with the 7 yr Savitzky–Golay smoothing.

The “conceptual” model approach described below demonstrates the general, quantitatively consistent nature of the relationship between upwind sources and downwind pollution. The more precise determination of this relationship would benefit from a Lagrangian particle dispersion model and cluster analysis treatment, such as described by Stohl et al. (2002, 2003); this type of analysis is beyond the scope of our present study. The “conceptual” model that relates upwind sources at a state level to downwind pollution is of great use to air quality managers.

The OH/PA cluster corresponds to the prevailing westerly winds during the ozone season. Emissions of NO\textsubscript{x} from OH and PA are ~4 and ~6 times higher than emissions from MD (Fig. 2a). A simplified approach was developed to evaluate the influence of upwind emissions: if the back trajectory passes over OH and PA, total emissions are estimated as

\[
\text{Emission}_{\text{total}} = \text{Emission}_{\text{MD},i} + 0.5\text{Emission}_{\text{PA},i-1} + 0.25\text{Emission}_{\text{OH},i-1},
\]

(1)

where the subscript \(i\) represents the flight date, and \(i-1\) represents upwind emissions from the day before (Table 1). The coefficients of 0.5 and 0.25 are used under a simple assumption based on the distance between the upwind sources and air pollutants over eastern MD, i.e., PA is next to MD so half of the CEMS emissions are assumed to be transported into MD while OH is farther away, so the coefficient is assumed to be 0.25. Values of the coefficients used for other clusters are given in Table 1. Our analysis is insensitive to the exact value of these coefficients, as discussed below.

Figure 13 shows a scatter plot of CEMS emissions found using coefficients given in Table 1 versus measured column ozone between the surface and 1500 m AGL. Linear regression analyses were conducted for each cluster (\(r\) values are listed in Table 1), and only the OH/PA cluster shows a strong correlation between upwind emissions and downwind air pollution. We also correlated ozone column abundances with emission estimates assuming double (high estimate) and half (low estimate) of these coefficients in Table 1 (see Fig. S9 in the Supplement). Ozone over the Harford County airport shows a strong positive correlation with emissions from the OH/PA cluster with little sensitivity to the exact value of these coefficients. However, the correlation between column ozone and MD emissions is low (Fig. S10 in the Supplement), suggesting that emissions from MD alone are a poor predictor of ozone pollution within this state. In summary, emissions of NO\textsubscript{x} in OH and PA substantially influence the ground-level air quality in the Baltimore/Washington area during the ozone season. Interstate/regional transport should be taken into account in plans to improve future air quality in the Baltimore/Washington area.

5 Conclusions and discussion

We have investigated the long-term trends of air pollution in the lower troposphere in the Baltimore/Washington airshed through a comprehensive study using emissions, aircraft, ground-based, and satellite measurements, as well as a clustering back trajectory analysis.

The EPA CEMS program continuously monitors emissions of NO\textsubscript{x} from major point sources including power plants and industrial boilers. Emissions data and atmospheric observations were used to investigate the regional transport...
of air pollutants. The long-term trends of CEMS emissions demonstrate a significant decrease in the emission of NO\(_x\) over the past 15 yr. The EPA NO\(_x\) SIP call (2003–2004) has been effective in reducing the emission of NO\(_x\) from point sources, which improved local and regional air quality, especially ground-level ozone. The EPA AQS NO\(_x\) observations declined by an amount similar to that of the CEMS emissions, with both time series exhibiting a sharp decrease after the EPA NO\(_x\) SIP call.

The aircraft measurements of O\(_3\) and its precursor CO were integrated to obtain column abundances, and categorized into upwind and downwind groups. The difference of tropospheric column ozone between the downwind and upwind group was used to estimate the daily net photochemical production of ambient ozone: ∼3.0 DU day\(^{-1}\) in the late 1990s and ∼1.0 DU day\(^{-1}\) in the early 2010s, equivalent to ∼20 ppbv day\(^{-1}\) and ∼7 ppbv day\(^{-1}\) averaged over the lowest 1500 m of the atmosphere. The decreasing trends of CO and O\(_3\) were found to be ∼0.8 and ∼0.2 DU yr\(^{-1}\), respectively, equivalent to decreases of ∼35 and ∼1.3 ppbv yr\(^{-1}\) in the lowest 1500 m. Because RAMMPP research flights are mostly conducted on air quality action days, the measurements indicate ozone pollution has significantly improved on days when the local meteorological conditions have historically been associated with poor air quality. Ground-based observations, RAMMPP aircraft measurements, and MOPITT spaced-based observations of CO show a ∼40 % (∼3.5 % yr\(^{-1}\)) decrease over western MD between 2000 and 2011, while the NEI emissions inventory suggests a ∼60 % (∼4.5 % yr\(^{-1}\)) reduction over this same period of time.

To investigate the effects of long-range transport of air pollutants, in particular from power plants with high smokestacks, back trajectories were calculated using the NOAA HYSPLIT model. These trajectories fell into several clusters, and total NO\(_x\) emissions were estimated for the geographic origins corresponding to each cluster. We performed a linear regression analysis for each cluster, and found that transport from OH and PA through MD has greatest effect on column abundances of ozone in the Baltimore/Washington airshed. Ozone over eastern MD correlates strongly (\(r = 0.86\)) with emissions of NO\(_x\) from upwind states such as OH and PA, when the trajectory analysis indicated westerly transport. The next highest correlation is with a recirculation pattern involving MD, VA, and PA (\(r = 0.53\)).
According to the NEI inventory, $\sim 90\%$ of CO and $\sim 50\%$ NO$_x$, important precursors for ozone pollution, are emitted by mobile sources (onroad and nonroad vehicles). In this study, annual mobile emissions from the EPA NEI inventory were used to track the long-term trend in air quality, because seasonal and diurnal variations of mobile emissions are not directly available. However, the seasonal and diurnal variations of mobile emissions could be important, so future work on quantifying these variations is essential to refine our understanding of the effects of regional and local emissions of ozone precursors on air quality.

Overall, this study reveals that recent regulation of emissions of ozone precursors substantially improved the air quality of the Baltimore/Washington airshed. We also find that the regional transport of NO$_x$ from point sources in upwind PA and OH still exert considerable influence on the downwind ozone pollution of the Baltimore/Washington area. These results identify the effects of emission controls on local air quality under complex regional transport patterns, which can inform future air quality planning and emission regulations in the eastern US.

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