Higher surface ozone concentrations over the Chesapeake Bay than over the adjacent land: Observations and models from the DISCOVER-AQ and CBODAQ campaigns

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HIGHLIGHTS

• Observations of ozone are higher over the Chesapeake Bay than areas upwind on land.
• Dry deposition rates, boundary layer depth, and photolysis play an integral role.
• Model resolution plays a role in determining accurate surface ozone concentrations.
• Observations of total reactive nitrogen are much lower than model simulations.

ABSTRACT

Air quality models, such as the Community Multiscale Air Quality (CMAQ) model, indicate decidedly higher ozone near the surface of large interior water bodies, such as the Great Lakes and Chesapeake Bay. In order to test the validity of the model output, we performed surface measurements of ozone (O_3) and total reactive nitrogen (NO_y) on the 26-m Delaware II NOAA Small Research Vessel experimental (SRVx), deployed in the Chesapeake Bay for 10 daytime cruises in July 2011 as part of NASA’s GEO-CAPE CBODAQ oceanographic field campaign in conjunction with NASA’s DISCOVER-AQ air quality field campaign. During this 10-day period, the EPA O_3 regulatory standard of 75 ppbv averaged over an 8-h period was exceeded four times over water while ground stations in the area only exceeded the standard at most twice. This suggests that on days when the Baltimore/Washington region is in compliance with the EPA standard, air quality over the Chesapeake Bay might exceed the EPA standard. Ozone observations over the bay during the afternoon were consistently 10–20% higher than the closest upwind ground sites during the 10-day campaign; this pattern persisted during good and poor air quality days. A lower boundary layer, reduced cloud cover, slower dry deposition rates, and other lesser mechanisms, contribute to the local maximum of ozone over the Chesapeake Bay. Observations from this campaign were compared to a CMAQ simulation at 1.33 km resolution. The model is able to predict the regional maximum of ozone over the Chesapeake Bay accurately, but NO_y concentrations are significantly overestimated. Explanations for the overestimation of NO_y in the model simulations are also explored.

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with a strong summer anticyclone aid in the rapid growth of ozone near the surface during the late morning and early afternoon.

Air quality models such as the Community Multiscale Air Quality (CMAQ) model indicate decidedly higher ozone near the surface of large interior waters bodies such as the Great Lakes and Chesapeake Bay (e.g., Godowitch et al., 2008). In order to test the validity of the model output, we performed surface measurements of ozone (O\textsubscript{3}) and total reactive nitrogen (NO\textsubscript{x}) on 26-m Delaware II NOAA Small Research Vessel experimental (SRVx), deployed in the Chesapeake Bay for ten daytime cruises in July 2011. The objectives of this paper are to:

- Compare ozone observations over the Bay to nearby land areas
- Determine if ozone concentrations are indeed higher over the Bay
- Determine if known meteorological and chemical processes can explain the observed differences
- Investigate whether model grid resolution plays a role in determining the simulated surface ozone concentrations over the Bay
- Investigate NO\textsubscript{x} observations to determine if this group of precursors is accurately predicted by the model simulations

### 1. Current EPA regulations and designations

The U.S. EPA has designated a “moderate” non-attainment region in the counties surrounding Baltimore, MD and a “marginal” non-attainment region throughout the New York City-Philadelphia-Washington, D.C. metropolitan corridor (U.S. EPA, 2008). A “moderate” attainment region is an area that has a ground station that exceeds an 8-h maximum concentration of 85 ppbv O\textsubscript{3} in its 3rd highest day, while a marginal attainment region has a threshold of 75 ppbv.

The Baltimore/Washington metropolitan region exceeds the threshold due to favorable meteorological conditions for ozone production during the summer months and substantial ozone precursor emissions, generated locally as well as advected to the region during strong westerly transport conditions (Ryan et al., 1998; He et al., 2013). Peaks in surface ozone are highest just downwind of major metropolitan areas due to the enhanced emissions from the metropolitan city centers (Kleinman et al., 2000). This has been shown in many air quality model simulations (Yegorova et al., 2011; Castellanos et al., 2011) and has been verified by ground monitoring stations (U.S. EPA, 2006; Castellanos et al., 2009). In the Baltimore–Washington region there are complex interactions that arise with the influence of the Chesapeake Bay breeze (Loughner et al., 2011; Stauffer et al., 2012), which have not been fully investigated.

### 1.2. Previous field campaigns over interior water bodies

Measurements of ozone over the southern Great Lakes during a 2007 summer field campaign show higher concentrations of ozone over the lakes than over the adjacent land with the biggest difference detected at night (Levy et al., 2010). A similar study was conducted over Lake Michigan in the summers of 1990 and 1991, where O\textsubscript{3} and NO\textsubscript{x} were monitored from aircraft (Luria et al., 1992). High levels of ozone are shown only at the lowest levels of the boundary layer, which they attribute to a lack of vertical mixing over the lake (Dye et al., 1995). An experiment in 2003 measured ozone at the Chesapeake Bay Lighthouse, located on an island 15 miles to the east of the entrance to the Chesapeake Bay, as a means to test ozone monitoring on ocean buoys and towers (Hintsa et al., 2004). This field campaign found ozone at the surface consistently exceeding 80 ppbv during an air quality episode from June 24 to 28, 2003.

### 1.3. DISCOVER-AQ And GEO-CAPE CBODAQ field campaigns

During the month of July 2011, the National Aeronautics and Space Administration (NASA) conducted a comprehensive air quality field study, DISCOVER-AQ (Crawford and Pickering, 2011), in the Washington, DC–Baltimore, MD metropolitan area and over the Chesapeake Bay to investigate air quality with the primary goal of providing data to better interpret observations from current and future satellites for air quality applications. In conjunction with DISCOVER-AQ, NASA conducted the oceanographic field campaign GEO-CAPE CBODAQ (Geostationary Coastal and Air Pollution Events-Chesapeake Bay Oceanographic Campaign with DISCOVER-AQ), to address questions related to both estuarine biogeochemical processes as well as atmospheric pollution over the Chesapeake Bay urban estuarine environment (Tzortziou et al., 2013). There were seven instrument platforms during the field campaign. A detailed description of the modes, locations, types, and days of observations is provided in Table 1. This paper focuses on observations from the 26 m Delaware II NOAA Small Research Vessel experimental (SRVs) deployed in the Chesapeake Bay as part of the CBODAQ campaign from July 11 to 20, 2011.

### 2. Materials and methods

#### 2.1. Measurements description

The SRVx was equipped with a Thermo Environmental Model 49 UV photometric ozone (O\textsubscript{3}) analyzer and a modified Thermo Environmental Model 42C chemiluminescence nitric oxide (NO) analyzer retrofitted with an external molybdenum catalyst to also measure total reactive nitrogen (NO\textsubscript{x}) (Delany et al., 1982). The NO\textsubscript{x} analyzer was zeroed for 10 min each hour during the campaign and

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<th>Mode of measurement</th>
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<th>Location</th>
<th>Days in which Active during</th>
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<td>July 11, July 14, July 16, July 20</td>
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<td>UC-12 king air</td>
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<td>July 11, July 14, July 20</td>
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<td>Cessna 402B</td>
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<td>Ozone sondes</td>
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<td>O\textsubscript{3}, NO, total NO\textsubscript{x}</td>
<td>See Fig. 1</td>
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measurements were adjusted based on the drift of the instrument. The NO\textsubscript{y} analyzer was calibrated in-situ on July 19, 2011 using a NO\textsubscript{2} standard reference material (SRM) from the National Institute of Standards and Technology (NIST).

### 2.2. Model description

In this study, we use U.S. EPA’s Community Multiscale Air Quality (CMAQ) (Byun and Schere, 2006) model Version 5.0, driven off-line by output from the Weather Research and Forecasting (WRF) (Skamarock et al., 2008) model Version 3.3 to simulate the state of the atmosphere covering the entire months of June and July 2011. Passing the meteorology into an air quality model at a high temporal resolution or running the chemistry online within a meteorological model is preferable, but requires significantly more computational resources (Grell et al., 2004).

The WRF and CMAQ model simulations are at 36, 12, 4 & 1.33 km resolution in the area of interest with 34 verticals from the surface to 100 mbar and 16 levels within the lowest 2 km in order to accurately simulate boundary layer processes. The 1.33 km model domain covers the Baltimore–Washington metropolitan region and nearby Chesapeake Bay. The North American Regional Reanalysis (NARR) is used for the model initial and outermost lateral boundary conditions in WRF. The Multi-scale Ultra-High Resolution (MUR) dataset was used to set the sea surface temperatures. The WRF model was re-initialized every 3 days and run in 3.5 day increments. The first 12 h of each simulation was thrown out (i.e., not passed to CMAQ). WRF model output is input into the Meteorology–Chemistry Interface Processor (MCIP; Otte and Pleim, 2010) to create meteorological input fields for CMAQ. Chemical initial and boundary conditions come from a MOZART-4 simulation (Emmons et al., 2010). The Carbon-Bond-05 (CB05) gas-phase chemical mechanism (Yarwood et al., 2005) was used in CMAQ. The CMAQ and WRF simulations began May 24, 2011, which allows ample spin-up time for our comparison in mid-July.

Anthropogenic emissions input files for CMAQ are created with the Sparse Matrix Operator Kernel Emissions (SMOKE) modeling system (Houyoux and Vukovich, 1999). We use a projected 2012 emissions inventory because a 2011 emissions inventory is not yet available. Annual projected point and countywide area emissions are temporally distributed based on the time of day, day of the week, and season based on temporal surrogates from the EPA. Mobile emissions estimates from cars, trucks, and motorcycles are computed with the Motor Vehicle Emission Simulator (MOVES; Kota et al., 2012). Point sources are vertically distributed based on the meteorology, stack height, and the temperature and velocity of the emissions exiting the stack. Biogenic emissions are calculated using Biogenic Emissions Inventory System (BEIS) and lightning NO\textsubscript{y} emissions are calculated in-line within the CMAQ model. Further model details on the parameterization options are described in the Supplementary Material.

### 3. Results

#### 3.1. Observational comparisons: ozone

The SRVx was deployed in the Chesapeake Bay for 10 daytime cruises during the DISCOVER-AQ Maryland campaign extending from July 11, 2011 through July 20, 2011. This overlapped with four flights of the NASA P3-B (a four-engine turboprop capable of long duration flights of 8–12 h) three flight days (2 flights per day) of the UC–12B King Air (a twin-engine turboprop capable of 6 h flights) and three flight days (2 flights per day) of the University of Maryland (UMD) Cessna 402B (a twin-piston engine, unpressurized aircraft) (Table 1). The SRVx docked each night in Annapolis, MD and had different cruise route each day (Fig. 1). The instruments were running while the SRVx was in port overnight in Annapolis, MD, but the data are subject to frequent local emissions.

A time series of O\textsubscript{3} for the 10-day period can be seen in Fig. 2. On four days ozone exceeded the 8-h maximum 75 ppbv NAAQS threshold on the moving vessel in the Chesapeake Bay: July 12, 13, 19 & 20. During this same time period, ground stations in the DISCOVER-AQ field campaign region (stations denoted in Table 2) exceeded the 75 ppbv threshold an average of 0.71 times per ground station. This alone is an indicator that the ozone may be higher near the surface of the Chesapeake Bay than nearby ground stations.

Comparing the hourly ozone at the SRVx’s location and closest upwind ground station reinforces the idea that higher ozone concentrations exist over the Bay. The closest upwind ground station was determined by using the backward trajectories at 10 m, 500 m, 1500 m heights ending at 1800 UTC (2 PM local time) using Global Data Assimilation System (GDAS) meteorological data in the NOAA HYSPLIT trajectory model (Draxler and Rolph, 2003). Eight-hour maximum ozone from all relevant ground sites and the SRVx can be seen in Table 2. The closest upwind ground station was often only 20–30 km away and was chosen to ensure that comparisons were made following the same parcel of air. Fig. 3a and b shows that during an exceedance day (July 13) and non-exceedance day (July 14) on the boat, the ozone near the surface of the Chesapeake Bay is uniformly higher. During the afternoon of July 13 the ozone measurement on the SRVx was 10–40 ppbv greater than at the Calvert County MDE site. Ozone was consistently 10–20 ppbv greater over the Bay than at the Essex MDE site throughout the day on July 14.

The 8-h maximum ozone concentration over the Bay during each day of the 10-day cruise averaged 12.7 ± 6.1 ppbv higher than the closest upwind ground site. The systematic high anomaly over the Chesapeake Bay can be seen in Fig. 4. The closest upwind ground site never experienced higher 8-h maximum ozone and...
only during three days did any ground station in the region have an 8-h maximum ozone concentration that was 10 ppbv higher than the SRVx’s location. This was especially pronounced on July 13 when the SRVx saw an 8-h maximum of 85 ppbv and none of the ground stations in the region exceeded the 75 ppbv NAAQS standard. When compared to 8-h maximum ozone at the ground stations in the Baltimore “moderate” non-attainment area, the 8-h ozone at the SRVx’s location was 4.6 ± 14.3 ppbv higher suggesting that the Chesapeake Bay has just as poor if not worse air quality than the surrounding “moderate” non-attainment area.

The ozone concentration remained higher over the Chesapeake Bay later into the afternoon than over the ground stations, suggesting that there must be a mechanism to maintain high O₃ concentrations later into the day. A plot of the median hourly ozone concentrations at the SRVx’s location and closest upwind ground station (Fig. 5) illustrates the late afternoon high anomaly. Ozone concentrations over the Bay are greater and exist for longer durations than over the upwind land area due to several potential causes:

1. A difference in ozone deposition rates over land and water;
2. A shallower PBL depth over the Chesapeake Bay than the nearby land causing emissions from shipping to be trapped near the surface;
3. Fewer fair-weather cumulus clouds over the Chesapeake Bay allowing for increased photolysis; and
4. Decreased boundary layer venting caused by a meso-high pressure that develops over the Bay as part of the bay-breeze circulation trapping pollutants.

Furthermore, when meteorological conditions are conducive, a low-level jet can form overnight transporting polluted air over the Chesapeake Bay from the Norfolk/Virginia Beach, VA metropolitan region bypassing ground stations allowing for increased ozone production over the Bay. This phenomenon, however, was not observed during this field campaign.

3.2. Model comparisons: ozone

Our CMAQ model simulation results typically reproduce the systematically higher ozone concentrations over the Chesapeake Bay than in the Baltimore-Washington region. As shown in Fig. 6, median ozone concentrations for the 10-day period output by both the 1.33 km and 4 km resolution CMAQ model simulations closely match the observations from the SRVx throughout the day. Model mean bias of ozone at the boat’s location was 0.78 ppbv, but a root-mean square error (RMSE) of 10.14 ppbv Table 3 shows the mean model bias, normalized mean bias, root-mean square error, and normalized mean error. At a grid cell size of 12 km, the surface ozone output by the model begins to lose correlation and at a grid cell size of 36 km, there was very little correlation throughout the

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Table 2

Maximum 8 h ozone at various sites in the Maryland & Delaware region.
day; both the 12 km and 36 km model runs show a high model bias in the late morning and afternoon. Model resolution seems to play an integral role in predicting ozone concentrations over the Bay.

Furthermore, we conducted a validation of the 2-m temperature in the 1.33 km WRF model simulation to ensure that the meteorology is indeed representative of the actual conditions during the 10-day period. Model mean bias of 2-m temperature at the boat’s location over the 10-day period was $0.52 \pm 0.14^\circ C$ and a RMSE of $1.59 \pm 0.14^\circ C$, which was a lower error than the nearby BWI airport. Table 4 shows the hourly mean model bias and RMSE at the boat’s location and the BWI airport. WRF at 1.33 and 4 km resolution was able to reasonably capture wind speed and direction. For example, on July 20 the boat showed light winds that veered dramatically from the NE to SW at 1 PM local time. The WRF runs at both resolutions indicated a similar wind shift but closer to 4 PM local time. This will shift the ozone maximum by only a few hours.

Although the 1.33 km resolution CMAQ model simulation closely matched the median for the 10-day period, on certain days the model was unable to predict ozone accurately, with a high bias shown in Fig. 7a and a low bias shown in Fig. 7b. During an exceedance day (July 12), the model had a consistent 10–15 ppbv high bias and on an earlier exceedance day (July 13), the model had a 10–15 ppbv low bias. The high bias of the model can likely be attributed to the boundary layer depths calculated by WRF and input into CMAQ, while the low bias of the model may be related to a lower temperature at the surface or perhaps a more stratified PBL inhibiting downward mixing.

Measurements of the aerosol-based boundary layer height were determined by a High Spectral Resolution Lidar (HSRL) instrument onboard the UC-12B aircraft on July 20. The HSRL dataset includes aerosol extinction at 532 nm, aerosol backscatter at 532 nm and 1064 nm and depolarization at 532 nm and 1064 nm (Hair et al., 2008) and profiles of aerosol backscatter are used to derive the mixed layer height (Scarino et al., 2013). Observations from the HSRL were compared to the modeled boundary layer from WRF on July 20. Observations were used only for July 20 because this was the only day the SRVx was in the north part of the Chesapeake Bay and the UC-12B aircraft simultaneously conducted a flight. On July 20, the modeled boundary layer in the morning agreed to within 100 m, but in the afternoon the modeled boundary layer was 300–500 m lower over the Chesapeake Bay than the observed aerosol-based boundary layer (Fig. 8).

3.3. Observational comparisons: total reactive nitrogen

Observations of reactive nitrogen species are critical since the eastern United States lies in the NO$_x$-limited regime of ozone

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**Fig. 4.** 8-h maximum ozone concentrations (ppbv) at the SRVx’s location and the closest upwind ground monitoring station from July 11, 2011 through July 20, 2011.

**Fig. 5.** Median hourly ozone concentrations (ppbv) at the SRVx’s location and the closest upwind ground monitoring station from July 11, 2011 through July 20, 2011 as a function of time.

**Fig. 6.** Median ozone concentrations (ppbv) at the SRVx’s location and at the closest CMAQ (1.33 km) grid point for each hour from July 11, 2011 through July 20, 2011 as a function of time.

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**Table 4.** Mean hourly ozone concentrations (ppbv) at the SRVx’s location and the closest upwind ground monitoring station from July 11, 2011 through July 20, 2011 as a function of time.
production (Chameides et al., 1992; Trainer et al., 1993; Frost et al., 2006) due to the excess of largely biogenic isoprene. Accurate model output of NO\textsubscript{y} species is especially important due to reactive nitrogen’s critical role in ozone formation in the NO\textsubscript{y}-limited regime found in eastern United States during the summer.

Observations from the SRVx were compared to the UC-Berkeley thermal dissociation laser-induced fluorescence (TD-LIF) instrument (Day et al., 2002) used on the P3-B when it flew spirals over the Chesapeake Bay. The TD-LIF does not measure NO\textsubscript{y}, so all comparisons are NO\textsubscript{y} – NO. The observations of NO\textsubscript{y} – NO from the SRVx using a chemiluminescence instrument with external molybdenum converter are higher than the data from the TD-LIF. This is an expected outcome since NO\textsubscript{y} concentrations decrease exponentially with height (Brent et al., 2013) due to emissions that come from the surface and relatively short lifetimes compared to other trace gases. There were no other suitable ground observations of NO\textsubscript{y} during this campaign.

### 3.4. Model comparisons: total reactive nitrogen

Observations of NO\textsubscript{y} from the SRVx were compared to 1.33 km CMAQ results over the Bay. On each day of the 10 daytime cruises, with the exception of July 19 when the instrument was taken offline for calibration, NO\textsubscript{y} observations were consistently lower than the output from the nearest grid point in CMAQ. The model regularly overestimated NO\textsubscript{y} and on July 12, it was overestimated by 100% in the mid-afternoon as shown in Fig. 9. The data from the TD-LIF instrument (Day et al., 2002) on the P3-B aircraft during a spiral on July 20 also indicate a significant overestimation of NO\textsubscript{y} species by CMAQ as shown in Fig. 10. While the vertical profiles of NO\textsubscript{2} and HNO\textsubscript{3} match well, alkyl nitrates (ANs) and peroxy nitrates (PNs) are overestimated by factors of 2 and 4 respectively. This overestimation of reactive nitrogen species has also been seen in other modeling studies (Brioude et al., 2013; Yu et al., 2012).

To understand whether the overestimate is an emissions issue, chemistry issue, or both, we examined the partitioning of the NO\textsubscript{y} species. If partitioning is correct, then the issue is likely due to high emissions or low dispersion rates. To gain insight on this issue, we took the ratio of NO/NO\textsubscript{y} during the morning hours when the two species are positively correlated and the NO measurement is above the detection limit. As seen in Fig. 11, the NO/NO\textsubscript{y} ratios between the model simulation and observations often lie below the 1-to-1 line. The mean of the data shows NO concentrations are 10.0% of total NO\textsubscript{y} in the observations, while NO concentrations are 7.6% of total NO\textsubscript{y} in the CMAQ simulation. This indicates that CB05, as employed, partitions more NO\textsubscript{y} species as higher oxides (i.e., ANs, PNs, HNO\textsubscript{3}) than is observed. This suggests that gas-phase chemistry scheme (CB05) overestimates the lifetimes of higher order NO\textsubscript{y} species such as ANs and PNs, deposition rates are too slow, or conversion rates of NO\textsubscript{y} to NO\textsubscript{2} are slower than observed.

To minimize computing time, the CB05 chemical mechanism simplifies the alkyl nitrates by grouping all alkyl nitrates in a single chemical species (NTR). The lifetime of NTR calculated during a simulation of CMAQ using 2007 summer conditions, yields a lifetime of 10 days. It has been shown that isopropyl nitrate has a lifetime of 10 days (Luke et al., 1989), but higher-order alkyl nitrates have a much shorter lifetime (1–2 days) (Horowitz et al., 2007; Perring et al., 2009) due to a lack of electronegativity holding the gas phase species together. The shorter lifetimes of the high-order alkyl nitrates species are not accounted for in the CB05 gas-phase chemistry scheme. After decomposition, the alkyl nitrates split into an alkyl chain and NO\textsubscript{2}. If the lifetime of NTR in CB05 were to be shorter, then this would yield lower concentrations of alkyl nitrates, which would be more consistent with observations.

To represent peroxy nitrates in the model, the CB05 mechanism simplifies the species into peroxyacyl nitrates (PAN), all higher order peroxyacyl nitrates (PANX) and peroxyacitic acid (PNA), with the latter being a very small fraction of the first two at high temperatures. The sum of the concentrations of peroxy nitrate species (PNs) in the model is higher than observed. The primary destruction of peroxyacyl nitrates is via thermal dissociation. At higher temperatures, PAN and PANX disassociate more rapidly into acetylperoxy radicals (CH\textsubscript{3}C(O)O\textsubscript{2}) and higher order acylperoxy radicals (C\textsubscript{2}H\textsubscript{5}C(O)O\textsubscript{2}) respectively. The concentration of PAN and PANX is therefore governed by the kinetic equilibrium rate constant, which is a function of temperature, and the concentrations of the products, CH\textsubscript{3}C(O)O\textsubscript{2}, C\textsubscript{2}H\textsubscript{5}C(O)O\textsubscript{2}, and NO\textsubscript{2}. There are stark differences in the kinetic equilibrium rate constant (K\textsubscript{eq}) between IUPAC (2010) and JPL (2011), with the latter being 24% less than the former (3.03 × 10\textsuperscript{-8} vs. 2.3 × 10\textsuperscript{-8}) at 298 K. The CB05 mechanism uses the higher IUPAC (2010) kinetic equilibrium rate constant, which favors a higher production rate of PAN. Furthermore, some studies (Turnipseed et al., 2006; Wu et al., 2012) have suggested that the dry deposition rates of PAN in the air quality models are too slow. Updating the rate constants of PAN formation as well as changing the dry deposition velocities, may better align the model output with observations.

### 4. Discussion

The observations from the SRVx show, with a 95% confidence level certainty, that ozone concentrations are elevated over the Bay when compared to upwind ground sites. The extended period of high ozone causes a larger number of days to exceed the U.S. EPA 8-h 75 ppbv NAAQS threshold over the Bay than over nearby land areas. Here we discuss potential reasons for this phenomenon and attempt to apportion a relative importance for each mechanism.

During a day that lacks precipitation, which is the case for most ozone exceedance days in the Baltimore–Washington metropolitan region, ozone is primarily destroyed by the mechanisms listed in Table 5 (Seinfeld and Pandis, 2006). Dry deposition (mechanism 1)
is the primary mode of ozone destruction near the surface. Titration
due to NO (mechanism 2) also occurs near the surface, but this
serves as a reservoir for O$_3$ as NO$_2$ is re-generated. Mechanisms 3–5
are most prominent in the upper troposphere and isolated ocean
regions where dry deposition rates are minimal.
Deposition is the primary mode of destruction in the boundary
layer and occurs fastest in heavily forested areas (Fowler et al.,
2001; Nowak et al., 2006). Differences in ozone dry deposition
rates have been widely studied. A list of 24-h-averaged dry
deposition velocities from the literature is given in Table 6 (Wesely
and Hicks, 2000; Chang et al., 2004; Nowak et al., 2006).
For a mix of 50% deciduous forest, 25% grass, and 25% pavement,
the 24-h averaged dry deposition velocity for ozone is 0.50 cm s$^{-1}$.
However, estimates for dry deposition of ozone in coastal envi-
ronments are 0.15 cm s$^{-1}$. The slower deposition velocity is due to a
lack of vegetation and surface roughness in coastal areas (Gallagher
et al., 2001). To calculate the difference in ozone deposition over an
hour, we can use the formula described in Table 7. For a boundary
layer depth of 800 m, which is typical over the Chesapeake Bay
during the mid-afternoon, ozone concentration would be approx-
imately 1.6% higher after an hour than an air parcel of similar
concentration over land due to slower deposition velocities over
water, assuming all other environmental conditions are the same.
If winds are from the southwest, maximizing residence time
over the Bay, an air parcel that entered the southwest portion of
Chesapeake Bay may have been over the Bay for approximately 5 h.
By the time an air parcel leaves the Bay, its ozone concentration
theoretically could be 8% higher than transport over land.
Boundary layer height also plays a major role in determining
concentrations of ozone near the surface (Rao et al., 2003). Pol-
lutants are primarily confined within the boundary layer due to a
strong subsidence inversion during anticyclonic events. The only
mechanism by which pollutants can be vented out of the boundary
layer during strong anticyclonic setups is through fair-weather
cumulus clouds (Dacre et al., 2007). However, cumulus clouds are
largely non-existent over the Chesapeake Bay during strong sub-
sidence events (Loughner et al., 2011).
The boundary layer over land tends to be deeper because the
surface temperature is higher over land during clear-sky conditions
in the mid-afternoon. As the boundary layer depth decreases,
emissions of ozone precursors, such as NO$_x$ compounds,
accumulate in a smaller volume of the atmosphere leading to higher concentrations. On July 20 between 20 and 21 UTC or 4 PM and 5 PM local time, the HSRL aboard the UC-12B aircraft measured the aerosol-based boundary layer depth to be 1000–1200 m over land and 400–600 m over the Chesapeake Bay within 10 min as seen in Fig. 12. If there were no boundary layer venting and environmental conditions and emissions were identical, the concentrations of NO2 could be up to a factor of 2 higher over the Bay than over land leading to a substantial increase in O3, since the mid-Atlantic region is in the NOx-limited regime. However, there is likely some vertical mixing and emissions are likely lower over the Bay. Although there were no direct measurements of NO2 at the surface of the bay during this particular campaign, data on the P-3-B shows that at 0.3 km, the lowest altitude of the flight spirals, NO2 is higher by as much as 0.5 ppbv over water than land. Using ozone efficiency rates from the DISCOVER-AQ campaign, for every 1 ppbv increase in NOx, ozone production will increase by an average of 8.26 ppbv with a 90% confidence interval of 4.93–19.4 ppbv (He et al., 2013); this is slightly higher than an urban study in Houston, which showed an average ozone production efficiency of 5.9 (Neuman et al., 2009).

A bay-breeze circulation often develops over the Chesapeake Bay during the late spring and early summer (Ryan et al., 1998; Staufer et al., 2012) impacting the coastal temperature structure and associated meteorological conditions. The bay-breeze yields a meso-high pressure directly over the Chesapeake Bay, and a meso-low pressure just inland from the Bay. This creates stagnation and clear skies directly over the Bay. Fewer cumulus clouds develop over the Chesapeake Bay than over land because of the lower surface temperature, shallower boundary layer depth and relative lack of thermals over the water. Decreased cloud cover increases photolysis rates by allowing more UV radiation to reach the lowest levels of the atmosphere creating an environment more favorable for ozone production. On July 20, visible satellite imagery, seen in Fig. 13, shows an expanse of low level fair cumulus clouds over the Baltimore—Washington region, with no clouds over the Bay. Cloud coverage is estimated to be 10–30% over land and 0% over the Bay leading to a higher j(NO2) value over the Bay.

During the DISCOVER-AQ campaign, the P-3-B aircraft measured j(NO2) throughout its flights. In the mid-afternoon, 3:30 PM local time, on July 20, 2011 when the P-3-B flew at an altitude of 390 m over land in an absence of clouds, the j(NO2) rate constant was 0.0082 s⁻¹, while 30 s later underneath a fair-weather cumulus cloud, which was confirmed by looking at the forward camera on the P-3-B, the j(NO2) rate constant dropped to 0.0043 s⁻¹. If we assume the sky over land is filled with 20% cumulus clouds and the sky over the Bay has no clouds, the average j(NO2) would be 0.0074 s⁻¹ over land and 0.0082 s⁻¹ over the Bay. Therefore, dissociation of NO2 into NO and odd oxygen may be up to 10.5% faster during the mid-afternoon of a summer day.

It is estimated that NOx emissions from barges that travel the Chesapeake Bay account for 10% of all mobile emission sources (U.S.

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**Table 5**

Loss mechanisms for ozone in the lower troposphere.

<table>
<thead>
<tr>
<th>Mechanism</th>
<th>Dry deposition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mechanism 2</td>
<td>O3 + NO → O2 + NO2</td>
</tr>
<tr>
<td>Mechanism 3</td>
<td>O3 + OH → O2 + HO2</td>
</tr>
<tr>
<td>Mechanism 4</td>
<td>O3 + HO2 → OH + O2</td>
</tr>
<tr>
<td>Mechanism 5</td>
<td>O3 + hν → O2 + O(1D) at hν &lt; 320 nm</td>
</tr>
</tbody>
</table>

**Table 6**

Ozone deposition velocities for various surface types.

<table>
<thead>
<tr>
<th>Surface Type</th>
<th>Forest</th>
<th>Coastal</th>
<th>Ocean</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nowak et al. (2006)</td>
<td>0.5 cm s⁻¹</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Wesely and Hicks (2000)</td>
<td>0.8 cm s⁻¹</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Chang et al. (2004)</td>
<td>–</td>
<td>–</td>
<td>0.05 cm s⁻¹</td>
</tr>
<tr>
<td>Gallagher et al. (2001)</td>
<td>–</td>
<td>0.148 cm s⁻¹</td>
<td>–</td>
</tr>
</tbody>
</table>
In March 2010, the U.S. EPA adopted a regulation requiring large barges to burn cleaner fuel that emits less NO\textsubscript{x} when they are within 200 nautical miles of the North American coastline (U.S. EPA, 2010). However, this regulation was not enforceable by the U.S. EPA until August 2012, which is after the Maryland DISCOVER-AQ field study. Many large transport tankers burn bottom-of-the-barrel bunker fuel, which releases a higher proportion of NO\textsubscript{x} than diesel fuel (Eyring et al., 2005). To date, there has been little quantification of barge emissions (Mason et al., 2008). Using the 8.26 ppbv O\textsubscript{3} per ppbv NO\textsubscript{x} ozone production efficiency calculated during the DISCOVER-AQ campaign (He et al., 2013), we estimate that a 0.1 ppbv increase in NO\textsubscript{x} concentrations over the Chesapeake Bay could yield a 0.8 ppbv increase in ozone, since the mid-Atlantic region is characterized by the NO\textsubscript{x}-limited regime of ozone production.

Halogen chemistry may play a role in ozone formation over the Chesapeake Bay. Recent modeling studies suggest that Cl\textsubscript{2} photochemistry may result in an increase of 5–8 ppbv in daily maximum ozone levels (Finley and Saltzman, 2006). To see if more chlorine is available over the Bay, we looked at the 5-year average (between 2007 and 2011) of Cl\textsuperscript{-} dry and wet deposition at two Clean Air Status and Trends Network (CASTNET) sites in Maryland. The Blackwater National Refuge site is located on Maryland’s eastern shore and is generally downwind of the Bay, while the Beltsville site is located upwind of the Bay. Dry deposition rates of Cl\textsuperscript{-} are 2.14 times higher over a 5-year average at the Blackwater site and wet deposition rates of Cl\textsuperscript{-} are 3.62 times higher.

One factor inhibiting ozone production over the Bay is the lower tropospheric temperature profile. Coastal areas in extratropical...
latitudes heat up more slowly than nearby inland locations during the summer due to the influence of the cooler waters. During the 10-day campaign, temperatures on the SRVx at 2 PM local time were on average 3.4 °C cooler than the Baltimore-Washington International (BWI) airport which is located 30 km inland from the Chesapeake Bay.

The dissociation of PAN into NO₂ has a strong temperature dependence (Seinfeld and Pandis, 2006). A calculation of the rate constant using IUPAC (2010) shows that PAN dissociates 1.66 times quicker at 304.3 K than 300.9 K. The quicker dissociation of PAN at higher temperatures over land shifts the equilibrium reaction toward NO₂, the primary precursor to ozone in the NOₓ-limited regime over the Mid-Atlantic. However, the dissociation of PAN is slower over the Bay, keeping more NO₂ tied up as PAN, and thereby decreasing O₃ production.

5. Conclusions

Observations from the NOAA SRVx vessel during the DISCOVER-AQ and GEO-CAPE CBODAQ campaigns show with a certainty exceeding the 2-sigma level, that daytime ozone concentrations are elevated over the Bay when compared to the closest upwind ground station. We posit that this high anomaly is influenced by a number of mechanisms, in approximate descending order:

- Shallower boundary layers trapping shipping emissions near the surface
- Higher photolysis rates due to clear skies over the bay
- Decreased boundary layer venting due to a lack of fair-weather cumulus clouds
- Slower deposition velocity over the Bay

The ozone concentrations exhibit a high anomaly over the Bay even though temperatures are cooler and allow precursors to ozone such as PAN to remain more stable. The observed high anomaly over the Chesapeake Bay is of primary importance since many citizens spend their leisure time on or near the Chesapeake Bay during the summertime, and are exposed to the unhealthy air quality conditions. Onshore winds can bring these pollutants to local coastal and inland communities. Expanded monitoring of ozone directly over the Chesapeake Bay is needed to precisely quantify the extent of this high anomaly.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at http://dx.doi.org/10.1016/j.atmosenv.2013.11.008.

References
