

Ozone distributions over southern Lake Michigan: Comparisons between ferry-based observations, shoreline- based DOAS observations and air quality forecast models

P.A. Cleary¹, N. Fuhrman¹, L. Schulz^{2*}, J. Schafer³, J. Fillingham³, H. Bootsma³,
J. McQueen⁴, Y. Tang^{4†}, T. Langel⁵, S. McKeen⁶, E.J. Williams^{5,6}, S.S. Brown⁵

[1]{University of Wisconsin-Eau Claire, Department of Chemistry, 105 Garfield Ave, Eau
Claire, WI 54702}

[2]{University of Wisconsin-Parkside, 900 Wood Road, Kenosha, WI 53144}

[3]{University of Wisconsin-Milwaukee, School of Freshwater Science, 600 E Greenfield Ave,
Milwaukee, WI 53204}

[4]{National Centers for Environmental Prediction/Environmental Modeling Center, 5830
University Research Court, College Park, MD 20740}

[5]{Chemical Sciences Division, NOAA Earth System Research Laboratory, Boulder, CO
80305}

[6]{Cooperative Institute for Research in Environmental Sciences, University of Colorado,
Boulder, CO 809309}

*now at University of Wisconsin-Madison College of Agricultural Life and Sciences, 1450
Linden Drive, Madison, WI 53706

† Now at Air Resources Laboratory, NOAA, 5830 University Research Court, College Park,
MD 20740

Correspondence to: P.A. Cleary (clearypa@uwec.edu)

Abstract

Air quality forecast models typically predict large summertime ozone abundances over water relative to land in the Great Lakes region. While each state bordering Lake Michigan has dedicated monitoring systems, offshore measurements have been sparse, mainly executed through specific short-term campaigns. This study examines ozone abundances over Lake Michigan as measured on the Lake Express ferry, by shoreline Differential Optical Absorption Spectroscopy (DOAS) observations in southeastern Wisconsin, and as predicted by the Community Multiscale Air Quality (CMAQ) model. From 2008-2009 measurements of O₃,

1 SO₂, NO₂ and formaldehyde were made in the summertime by DOAS at a shoreline site in
2 Kenosha, WI. From 2008-2010 measurements of ambient ozone conducted on the Lake
3 Express, a high-speed ferry that travels between Milwaukee, WI and Muskegon, MI up to 6
4 times daily from spring to fall. Ferry ozone observations over Lake Michigan were an average
5 of 3.8 ppb higher than those measured at shoreline in Kenosha with little dependence on
6 position of the ferry or temperature but with greatest differences during evening and night.
7 Concurrent 1-48h forecasts from the CMAQ model in the upper Midwestern region surrounding
8 Lake Michigan were compared to ferry ozone measurements, shoreline DOAS measurements
9 and EPA station measurements. The bias of the model O₃ forecast was computed and evaluated
10 with respect to ferry-based measurements. Trends in the bias with respect to location and time
11 of day were explored showing non-uniformity in model bias over the lake. Model ozone bias
12 was consistently high over the lake in comparison to land-based measurements with highest
13 biases for 25-48h after initialization.

14 **1 Introduction**

15 Air quality near Lake Michigan has been under study for more than 30 years (Lyons and Cole,
16 1976; Keen and Lyons, 1978; Dye et al., 1995). The shoreline air quality has gone from a highly
17 impacted environment for surface ozone in the 1970's-80's to persistent non-attainment status
18 in the 2008 ground-level ozone standards for counties near to Lake Michigan in Wisconsin
19 (Sheboygan and Kenosha), Illinois (Cook, Lake, Grundy, Kane, Kendall, McHenry, Will) and
20 Indiana (Lake, Porter). The number of critical ozone events in the Chicago metro area region
21 has been reduced in the past 20 years (EPA, 2014), but stricter measures for particulates have
22 maintained a steady pattern of particulate matter exceedances for this region (Katzman et al
23 2010, Stanier 2012). Non-attainment of federal ozone standards are still of concern. Kenosha
24 remains in marginal non-attainment of federal ozone standards (as of 2012) and Sheboygan
25 County, north of Milwaukee, remains in non-attainment. The proposed rule as of Nov. 26th,
26 2014 is to reduce the 8-hour primary standard to between 65 and 70 ppb ozone, which has the
27 possibility of maintaining the non-attainment status for these counties in the future (EPA, 2014).
28 These Wisconsin counties in non-attainment are unique in that they are both suburban, Lake

1 Michigan shoreline counties as opposed to urban or rural counties. Studies have been addressing
2 the role of lake breeze in air quality near the Great Lakes of North America (Levy et al., 2010;
3 Sills et al., 2011; Makar et al., 2010), with a whole campaign, BAQS-MET, dedicated to the
4 evaluation of lake breezes. Complexities in the reduction of precursors and continued increases
5 in ozone are of current concern in the Toronto area (Pugliese et al., 2014). Here, we evaluate
6 the Lake Michigan ozone mixing ratios off-shore with those on-shore, including agreement
7 with ozone forecast models overwater and at the shoreline.

8 Ozone is generated in the troposphere by the reaction of precursors (nitrogen oxides
9 (NO_x) and volatile organic compounds (VOCs)) in a photochemical cycle that is typically most
10 active during high pressure events in summer. The Milwaukee-Chicago-Gary urban corridor
11 constitutes a large emissions source for ozone precursors and is home to significant populations
12 impacted by poor air quality. The understanding of ozone production and distribution around
13 Lake Michigan requires monitoring of land-based sites year-round, but no regular observations
14 of offshore air quality exist. Some land-based monitors are situated farther from Lake Michigan
15 than others, but no specific quantification of the difference between surface level offshore air
16 quality and onshore air quality exists on a routine basis. Forecast models typically produce large
17 ozone mixing ratio maxima over Lake Michigan (Lennartson and Schwartz, 1999, 2002). The
18 nature of the distribution of ozone precursor emissions near to the Lake Michigan shoreline
19 from an urban corridor is in stark contrast to the reduced anthropogenic and biogenic emissions
20 over the lake. This, combined with the unique meteorological effects from this large body of
21 water, like the lake breeze, which can trap, stratify and recirculate air offshore, highlights the
22 need for ozone measurements at a near shore site and across the lake.

23 The study of high ozone events in the region has centered around mesoscale
24 meteorological effects that contribute to the formation of ozone and the movement of air masses

1 over land (Lennartson and Schwartz, 2002; Lyons and Cole, 1976). Lyons and Cole (1976)
2 outlined the influence of the land-breeze effect on shoreline air quality. Lennartson and
3 Schwartz (2002) indicated a pattern of high pressure anticyclonic events as coincident with
4 higher ozone abundances at land-based sites. Recently, Levy et al. (2010) investigated the
5 impact of local-scale flows in Great Lakes air quality in the region of Lake Erie. Levy et al.
6 determined that local-scale emissions play a significant role in ozone production, and the
7 meteorological constraints on air movement aid in isolating and stratifying air pockets from
8 which ozone is generated on a next-day basis.

9 A few studies have investigated offshore air quality in regional-scale monitoring of
10 ozone around Lake Michigan. The two most notable studies are the Lake Michigan Air Quality
11 Study in 1991, which used aircraft for monitoring (Dye et al., 1995) and the LADCO Aircraft
12 Project (Foley et al., 2011). Dye et al. (1995) determined that stratification over Lake Michigan
13 lead to limited vertical and horizontal mixing beyond the lake area during the summer, allowing
14 for the confinement of ozone precursors. The LADCO Aircraft Project (LAP) was a 9-year
15 aircraft-based study to evaluate air quality in the region, where flights were conducted on days
16 of suspected high ozone in non-attainment of hourly federal standards (Foley et al., 2011). The
17 work from LAP is consistent with the interpretation presented by Dye et al. in that inversions
18 over the lake created stable layers of urban plumes, and that air sampled at greater distance
19 from the Chicago - Milwaukee shoreline tended to be more processed. Foley, et al. (2011)
20 determined in the late 1990's and early 2000's that in lower altitude air (< 200 m above ground
21 level (AGL)) ozone formation switched between VOC-limited conditions in the morning to
22 NO_x-limited in the afternoon, and that above 200 m AGL, ozone formation was always NO_x
23 limited. The observations from LAP showed a progression of the "photochemical clock" during
24 northward aircraft transects over the lake where more aged plumes were found farther north of
25 Chicago. Fast and Heilman (2003, 2005) developed a regional coupled meteorological and

1 chemical model to describe ozone formation on or near the Great Lakes. For offshore
2 measurements they used ozone observations from the SS *Badger*, which operates between
3 Luddington, Michigan and Manitowoc, Wisconsin. The comparison between the model and
4 measurements was restricted to specific times of the day due to the ferry movement where the
5 agreement of model to measurement was poorest for the eastern side of Lake Michigan in 1999
6 (Fast and Heilman, 2003). Their model results from 1999 and 2001 showed distinct features in
7 the ozone spatial distribution over Lake Michigan but did not reproduce eastern Wisconsin
8 shoreline observations when ozone mixing ratios were high (>60 ppb) (Fast and Heilman,
9 2005).

10 The Lake Michigan land/lake breeze is a well-documented phenomenon that influences
11 local scale air flow due to differential heating of air masses over land and water on a daily basis
12 (Lyons and Cole, 1976; Foley et al., 2011; Hanna and Chang, 1995; Lennartson and Schwartz,
13 2002). Offshore flow (the land breeze) is dominant during the nighttime during summer when
14 surface waters are higher in temperature than land surface temperatures. For counties along the
15 western side of Lake Michigan, this westerly pattern follows typical westerly synoptic flow for
16 the region. Onshore flow (the lake breeze) is more common in the summer daytime when land
17 temperatures exceed water surface temperatures. The lake breeze has been seen to coincide with
18 higher ozone and the transport of aerosol in Chicago (Harris and Kotamarthi, 2005; Lyons and
19 Olsson, 1973) and larger-scale high pressure anticyclonic flows have been implicated in the
20 higher Lake Michigan shoreline ozone observations (Lennartson and Schwartz, 1999), which
21 enhance the flow of photochemically aged air from the Chicago urban plume northward along
22 the Lake Michigan shoreline to southeastern Wisconsin.

23 In this study, the deployment of both a long path Differential Optical Absorption
24 Spectrometer (DOAS) at the shoreline and an ozone monitor on a ferry has several benefits: the

1 long path length for the DOAS instrument creates an averaged signal that is unaffected by small
2 spatial scale point-source emissions, and allows for simultaneous observations of several
3 compounds (NO_2 , SO_2 , O_3 , formaldehyde). This combination of species provides relevant
4 information about air masses, where O_3 is the pollutant of interest to compare with offshore
5 observations, NO_2 is a proxy for NO_x and a precursor to O_3 production, formaldehyde is a proxy
6 for total VOC which are other necessary ozone precursors, and SO_2 is used as a tracer for
7 industrial emissions and electric power generation. The use of a DOAS instrument for
8 monitoring atmospheric species at a shoreline has proven effective in other environments, such
9 as the observatory on the west coast of Ireland, (Carpenter et al., 1999; Seitz et al., 2010), Crete
10 (Vrekoussis et al., 2004), Galapagos Islands (Martin et al., 2013), Okinawa Island (Takashima
11 et al., 2011), Houston (Rivera et al., 2010), Helgoland (Martinez et al., 2000) and Appledore
12 Island, NH (White et al., 2008), to name a few. In the study described here, the four constituents
13 measured by DOAS are used to show the change in chemical composition of air masses from
14 offshore and onshore and to evaluate the spatial distribution of the species at the Lake Michigan
15 shoreline. The routine monitoring of ozone over Lake Michigan on the ferry platform allows
16 for an evaluation of the spatial distribution of ozone over the lake, comparison of over-water
17 ozone to shoreline ozone, and comparison to forecast models of surface-level ozone. This
18 investigation is the first to present high resolution, regular observations of ozone at the surface
19 over Lake Michigan in comparison to air quality model output. Results have been analyzed to
20 show the difference between shoreline and over-water ozone as a function of time of year, time
21 of day, location over the lake and meteorology.

22 **2 Methods**

23 Kenosha, Wisconsin is located along the shoreline of Lake Michigan in the southeast corner of
24 the state, bordering Illinois (Figure 1). The commercial DOAS instrument was mounted to two
25 municipal buildings at the Kenosha Harbor along Lake Michigan spanning the harbor with a

1 one-way single-beam path length of 596 m. The light source was mounted to the roof of the
2 Kenosha Municipal Building at 625 52nd St and the detector was housed at the Kenosha Water
3 Utility Water Production Plant located at 100 51st Place on Simmons Island. The beam passed
4 over land and water at 10-14 meters above ground level. At this location, the shoreline of Lake
5 Michigan is oriented North-South, with a small residential area directly south of the
6 measurement site (see inset of Figure 1). The measurement site is located in downtown
7 Kenosha, a city of 100,000 located 35 miles south of Milwaukee (metropolitan area population
8 2 million) and 50 miles north of Chicago (metropolitan area population 9.5 million). The DOAS
9 unit was calibrated with known standards in Sept. of 2008 ($\pm 4\%$ yearly drift). In-beam standards
10 were used to test the calibration Nov 7, 2008 and Aug 8, 2009. The instrument was operated
11 from Sept. 19 to Nov. 24, 2008 and April 28, to Nov. 10, 2009. Meteorological data
12 (temperature, relative humidity, wind speed and direction) were obtained in 2009 by the
13 addition of a meteorological station at the Kenosha Harbor site of the DOAS detector. The
14 meteorological sensors were mounted to a pole extending 3 meters above the rooftop where the
15 DOAS detector was mounted. Data were collected as 1-minute averages for each compound
16 (NO_2 , SO_2 , O_3 and formaldehyde) sequentially, which resulted in single data points every 5
17 minutes (1% precision). Data was filtered for low light levels when the instrument required
18 realignment. No post-processing filters, (e.g. omitting data with low wind speeds) were placed
19 on meteorological measurements.

20 The *Lake Express* ferry runs from May to October from Milwaukee, WI to Muskegon,
21 MI (Figure 1) at 06:00 (eastbound), 09:15 (westbound), 12:30 (eastbound), 15:45 (westbound)
22 CDT and in late July/August also at 19:00 (eastbound) and 22:00 (westbound) CDT. Time
23 zones for Wisconsin and Michigan differ, but all times given here are in Central Daylight Time.
24 The ferry stays in port overnight in Milwaukee and the average trip duration of the ferry for this
25 study was 2.25 hours. The inlet for air monitoring was installed at the bow above the

1 wheelhouse (3 m starboard of center and 10 m above water line) and approximately 15 meters
2 of ¼" PTFE tubing was routed through the interior conduit into a utility closet where a
3 commercial CO₂ (Li-Cor) and O₃ (Thermo Scientific Model 49) monitor were housed. The
4 sample line had a teflon cartridge filter (changed approx. weekly) and tee fitting to the two
5 instruments (each with independent pumps) with a sampling time lag of approximately 10 s.
6 The inlet was positioned to the stern so as to minimize water spray entering the sample lines,
7 with intake tubing surrounded by a larger tubing as a rain/spray cover. The O₃ instrument was
8 installed on the ferry from July 9-Sept 21, 2008, May 12 to Oct. 28, 2009 and June 23-Nov. 1,
9 2010. GPS coordinates and gas measurements were recorded every 30 seconds resulting in a
10 frequency/spatial resolution of ~1 min/km, with an average speed of ferry at 30 knots. Zeros on
11 the ozone monitor were conducted during powerdown of the ferry (typically twice per day when
12 ferry was docked in port). Ozone data was excluded from data set when the ferry was in port
13 because measurements were also influenced by engine emissions of NO. On occasion, due to
14 inclement weather or mechanical problems, the ferry did not follow its posted schedule. The
15 ozone instrument had a manufacturer stated accuracy of ± 2 ppbv. The ozone instrument was
16 calibrated at NOAA before and after deployment each year by comparison of the instrument
17 deployed on the ferry to a standard ozone monitor (Thermo Scientific Model 49i-PS)
18 maintained in the laboratory for comparison purposes. Comparisons were always within 2%.

19 **3 Results**

20 **3.1 Shoreline DOAS Observations as a function of wind direction**

21 Observations from the Kenosha Harbor DOAS instrument were evaluated with respect to
22 offshore versus onshore air mass origin by sorting the data with respect to observed wind
23 direction in 2009. For 2009, all 30-minute averaged data were binned to median mixing ratio
24 per 30 degree increment of wind direction. Figure 2 shows the distribution of gases O₃, NO₂,

1 SO₂ and formaldehyde median mixing ratios with respect to wind direction. The highest median
2 ozone and SO₂ mixing ratios observed at the Kenosha Harbor location arise from air masses
3 flowing from the lake (0-180° are from offshore), whereas the highest NO₂ and formaldehyde
4 observations arise from air masses originating on land. So few formaldehyde measurements in
5 the onshore flow were above the detection limit that average data from those wind directions
6 were omitted from Figure 2d. The observation of NO₂ from land-based air masses is consistent
7 with localized fossil-fuel combustion sources of short-lived NO_x (=NO+NO₂) coming from
8 land-based mobile and point sources as NO_x oxidizes rapidly to other nitrogen species during
9 the daytime. Formaldehyde can serve as a proxy for VOCs, with anthropogenic and biogenic
10 emissions arising from sources on land, and can also be produced *in situ* as an oxidation product
11 of VOCs. Formaldehyde can be lost to reaction with OH and photolysis during the day. The
12 longer-lived atmospheric species of O₃ and SO₂ were observed in higher abundance from
13 offshore. The O₃ and SO₂ mixing ratios were otherwise not correlated in individual days, which
14 is typical as the chemistry and emissions driving the evolution of each were quite different. O₃
15 is produced by catalytic photochemical cycles which require the presence of NO_x and VOCs
16 and can be titrated by fresh emissions of NO. Sulfur dioxide is most commonly emitted by
17 fossil fuel combustion at coal-fired power plants, many of which lie at the Lake Michigan
18 shoreline in the Gary-Chicago-Milwaukee urban corridor from Indiana to Wisconsin. The
19 diurnal wind patterns (Figure 3) at the Kenosha Harbor site also contribute to the apparent
20 higher mixing ratios of ozone and SO₂ over the lake because the lake breeze wind pattern drives
21 winds from land offshore at night (when NO₂ and formaldehyde losses by photolysis and
22 reaction with OH were minimized) and from the lake onshore during the day (when ozone
23 mixing ratios were at a maximum). This maximum for NO₂ arriving from off-shore air masses
24 we interpret as an artifact of NO₂ minima mid-day coinciding with on-shore air masses, even
25 though sources (both mobile and stationary) of NO₂ are predominantly located on land. The

1 night-time NO₂ maxima is likely from lower night time losses of NO₂ (not to be mistaken for
2 nighttime NO_x losses which can still be significant (Brown et al., 2004)) instead of indicating
3 some high NO_x emissions source from off-shore at night.

4 These DOAS observations align with past studies of Lake Michigan air quality in that
5 they implicate higher O₃ mixing ratios over Lake Michigan (Dye et al., 1995; Foley et al., 2011;
6 Lennartson and Schwartz, 1999, 2002). The higher SO₂ mixing ratios may show the influence
7 of power plant emissions mixing over longer distances and timescales over the lake. The nearest
8 power plants to the DOAS site are located to the southwest (Pleant Prairie), north (Oak Creek)
9 and south (Waukegan) and yet SO₂ observations are highest from the southeastern quadrant,
10 including from the south and east. The lifetime of SO₂ is long enough (approx. 1 week) that
11 sources from other powerplants neighboring Lake Michigan (see Fig. 1) may contribute to these
12 observations. Foley et al (2011) described sampling high NO_x plumes over Lake Michigan that
13 appeared to remain aloft. They suggested that these plumes originated from power plants in
14 the region, which would also be a source of SO₂. The shoreline observations presented here do
15 not constrain the extent to which ozone was higher over the lake, nor the distribution of ozone
16 across the lake, but only show that air with enhanced ozone was observed during afternoon
17 hours when the air moved inland during the lake breeze. At the intersection between the
18 offshore environment and the onshore environment, titration of O₃ occurs via emissions from
19 local NO_x sources, and therefore the additional offshore processing cannot be distinguished
20 from chemistry at the shoreline with this DOAS measurement alone.

21 **3.2 Comparison between shoreline DOAS and ferry observations**

22 Kenosha shoreline DOAS observations of O₃ were compared with the *Lake Express* ferry O₃
23 observations in order to understand the regional distribution of ozone. The two measurements
24 were compared by averaging the ferry measurements to 30 minute intervals at the timescale of

1 the Kenosha harbor DOAS measurements. The two instruments were never intercompared at
2 the same location so we estimate an uncertainty in their intercomparison at 5% (which is higher
3 than the stated drift of either instrument as evaluated independently). The differences in 30-
4 minute averaged data from 2009, as measured as O_3 (Lake Express Ferry) - O_3 (Kenosha Harbor), fluctuated
5 from as high as 45 ppb to -37 ppb, with a median difference of 2.8 ppb, mean of 3.8 ppb and
6 standard deviation of 9.1 ppb. The daily maximum data (30-minute average) had a range of 39
7 ppb to -9 ppb, a median of 4.2 ppb, mean of 5.0 ppb, standard deviation 7.6 ppb. The time of
8 peak ozone for ferry measurements was approximately 14-17h CDT for the whole campaign
9 and for the DOAS measurements was from 14-16h CDT, which are not considerably different.
10 Day-to-day variations in the time of peak ozone off-shore versus onshore can occur from
11 changes in wind direction and local NO_x sources at the shoreline Kenosha site, and therefore
12 cannot be used to indicate differences in chemical processing over the day. There is a
13 statistically significant difference in the O_3 distribution over land vs. lake from summer (June,
14 July, August) to fall (September, October) with median difference of 3.3 ppb for summer and
15 1.6 ppb for fall (Kruskal-Wallis $p=0.05$).

16 In order to demonstrate the agreement between ozone measurements of both platforms,
17 Figure 4 shows the wind direction, O_3 measurements, the difference in ozone measurements,
18 temperature, NO_2 , SO_2 and formaldehyde for Aug. 12 to Aug. 18, 2009. This week was chosen
19 because of the range of ozone maxima depicted (with daily maxima ranging from 40-70ppb)
20 and the example of a wind shift event that correlated to temperature and atmospheric
21 composition changes at the shoreline on August 14th. In the example of Aug. 12, 2009, the
22 ozone mixing ratios for both instruments appear quite similar. Note that the discontinuities in
23 ferry data represent times when the ferry was in port, and each of the segments between the
24 data gaps represents an entire transect of Lake Michigan. In some cases, such as Aug. 12, there
25 was very little variation in the difference between ferry and shoreline O_3 with respect to the

1 location of the ferry. For Aug. 13, the maximum ozone as measured at the shoreline (~50 ppb)
2 was observed by the ferry upon return to the western side of Lake Michigan and again when it
3 left with roughly a 15 ppb difference between the eastern and western sides of Lake Michigan
4 in the afternoon hours. NO₂ measurements in Figure 4d peaked at night as high as 30 ppb and
5 were at a minimum during the day, particularly after noon. The mixing ratios of NO₂ for this
6 period do not correlate with SO₂ mixing ratios and so can be considered to be from different
7 emissions sources, such as urban non-point source NO_x and power-plant or industrial sources
8 of SO₂.

9 Evidence of lake breeze shifts in the data was most clearly shown on Aug 14th (indicated
10 by dotted lines in Fig. 4). The wind direction shifted abruptly from southwest (offshore flow)
11 until about 10:00 CDT, when it shifted to southeast (onshore flow). The temperature change
12 between these two air masses is evident in Figure 4c, where the ambient temperature dropped
13 3 °C as the wind direction shifted. The NO₂ mixing ratio increased to 30 ppb after the wind
14 shift, which may be evidence of recent land-based NO₂ emissions from the northern Chicago
15 area flowing offshore during rush-hour and then returning onto land after the wind shift.
16 Following the rapid NO₂ decrease, O₃ increased as measured at the shoreline and also as
17 measured on the ferry. By 18:00 CDT, the wind shifted back to arriving at the Kenosha Harbor
18 site from the southwest, the shoreline ozone decreased precipitously but the ferry observations
19 of ozone remained high. The shoreline NO₂ mixing ratios also rebounded to 12 ppb. In this
20 case, the maximum SO₂ observations arrived at the Kenosha harbor site from offshore later in
21 the afternoon before the wind shifted. A Hysplit back trajectory model was calculated for the
22 morning of Aug 14th for synoptic winds at 250 m AGL and indicated an air mass arriving from
23 the northeastern suburbs of Chicago, Illinois which would intercept the rush-hour traffic
24 emissions. Thus, the low O₃ mid-morning was a result of near-source and early-day NO_x
25 titration. On Aug. 13, 14 and 15, NO₂ increased following the wind shift between south-

1 westerly and south-easterly wind flows. Hysplit back trajectories were generated for each of
2 these days, which showed air masses from Chicago transported northward along the shoreline at
3 the same time of day. Emissions were likely brought back on land from lake breezes which
4 could not be resolved from back trajectories.

5 Differences between ferry O_3 and shoreline DOAS O_3 mixing ratios were evaluated with
6 respect to temperature (Figure 5), location of the ferry (Figure 6) and wind direction (Figure 7).
7 Each figure shows the data for all times of the day, and for distinct time windows (06:00-12:00
8 CDT, 12:00-18:00 CDT, 18:00-02:00 CDT) in box plots which represent mean (line), median
9 (\square), 25-75% (box), and 10-90% (whiskers) for the 30-minute average difference between O_3
10 (Lake Express) and O_3 (Kenosha Harbor). Differences between ozone observations from the ferry and
11 shoreline with respect to temperature were investigated (Figure 5). There was no observed trend
12 in difference in ozone versus temperature for all data (Figure 5a). A minor trend in median
13 difference in ozone versus temperature is observed for morning times (06:00-12:00 CDT,
14 Figure 5b) where the difference changed from a positive difference to a more negative
15 difference with increasing temperature above 15.5 °C, and an opposite trend toward higher
16 ozone over the lake in the afternoon (12:00-18:00 CDT) and for temperatures above 26 °C.
17 Ozone differences after 18:00 CDT show consistently higher ozone mixing ratios over the lake
18 for all temperatures (Figure 5d), with a trend for larger differences above 21.1°C.

19 For each of the graphs in Figure 5, we conducted a Kruskal-Wallis (K-W) non-
20 parametric statistical test whether the distributions depicted in each box plot in the figure could
21 be considered different from each other. For Figure 5a, the K-W test indicated that no difference
22 in the distributions (as depicted by box plots across different temperatures) could be determined
23 within 95% confidence. For Figures 5b and 5c the distributions (as depicted as box plots across
24 different temperatures) could be considered different with 75% confidence. In Figure 5d, the

1 K-W test indicated that the distributions were significantly different with 95% confidence. The
2 K-W test does not indicate trend, just whether one or more distributions with the comparison
3 are different from each other. K-W tests applied across different times of day for a given
4 temperature range (a vertical comparison in the stacked plots) consistently showed a significant
5 difference (95% confidence) in distributions.

6 While the chemistry can drive more ozone production at higher temperatures, the fact
7 that the largest differences were observed in the evening and at night can arise from the isolation
8 of air masses at this time from the lake/land breeze effects. If the airmasses observed at the
9 shoreline arrived from inland in the late evening, they could have been chemically different
10 from those found far offshore. The only time when shoreline DOAS ozone observations tended
11 to be higher than those from the ferry was at 06:00-12:00 CDT for temperatures above 26.7 °C.
12 This may be due to days when temperatures were high in the morning, thus stagnating the air
13 and limiting the influence of lake/land breeze on horizontal movement of airmasses.
14 Differences in offshore and shoreline observations of ozone with respect to temperature were
15 largest later in the day and at higher temperatures when ozone was typically at a maximum. The
16 range in temperatures observed from different wind directions was higher in wind arriving from
17 land (180°-360°) in comparison to over water (0°-180°), such that the median temperature of all
18 masses arriving at the site from the east was 12.8°C and from the west was 9.3°C. The highest
19 differences depicted in Figure 7 are showing the highest ozone differences between shoreline
20 and offshore measurements from a wind direction where temperatures are not as extreme.

21 Investigations into the ozone differences between shoreline and ferry observations with
22 respect to ferry location were conducted as a test of the east-west gradient over Lake Michigan.
23 Figure 6 depicts the difference of O_3 (Lake Express) – O_3 (Kenosha Harbor) with respect to ferry distance
24 from Milwaukee. For all data the mean and median difference was positive (i.e., greater as

1 measured over water from the ferry). The median differences were not significantly positive or
2 negative for the morning, slightly positive for the early afternoon time window, and consistently
3 positive for the late afternoon/evening. In the case of the late evening time window, the mean,
4 median and extremes (25%-75%) of the data all lie above 0, which is a strong suggestion that
5 at these times the ozone mixing ratios over the lake are consistently higher than at the shoreline.
6 However, there does not appear to be a significant variation with respect to longitude, meaning
7 that evaluated as a whole, the land-lake differences in ozone did not depend on the ferry's
8 distance from the shoreline. All K-W tests for each plot in Figure 6 show no difference (95%
9 confidence) in distributions across different locations, corroborating this assertion. The only K-
10 W tests that showed a difference in distributions in Figure 6 was comparisons with respect to
11 time of day, which is similar to the time of day tests for Figure 5. This demonstrates a widely
12 regional distribution of ozone over the lake.

13 In order to distinguish between meteorological effects at the shoreline, the differences
14 in ozone observations from the ferry and shoreline DOAS ozone mixing ratios with respect to
15 wind direction at Kenosha Harbor were evaluated. All data (Figure 7a) show a trend in which
16 the differences between offshore and onshore observations of ozone are positive (i.e., greater
17 ozone over water as measured from the ferry) when wind arrives at the Kenosha Harbor site
18 from 180-360 degrees (inland) where the median and mean lie above 0. However, a K-W test
19 for all data does not show a significant difference in the distributions with 95% confidence.
20 When broken up into time windows of morning, afternoon and evening/night, the largest
21 differences were observed after 18:00 CDT if winds were arriving from 180-360°. The K-W
22 tests only showed a difference in distributions across wind direction for Figure 7d with 75%
23 confidence. This picture is consistent with land breezes developing in the evening and
24 producing surface winds which draw from land and move over the lake. The sampled air masses
25 at the shoreline, thus, were of different origin (or sampled air masses over the lake were isolated

1 from land-based air masses). The number of data points ($n < 15$) were acquired when the wind
2 blew from $30\text{--}160^\circ$ from 18:00-02:00 CDT were insufficient for analysis. For the morning and
3 early afternoon times, the trend with respect to wind direction was not large.

4 The two key differences between ferry and shoreline ozone observations in these
5 comparisons were those after 18:00 CDT and into the night, as shown in Figures 5, 6, and 7 (all
6 of which were significantly different from other times of day based on K-W tests with 95%
7 confidence) and the trend with the wind direction for all times of the day with the mean
8 difference for wind directions from $0\text{--}180^\circ$ at 0.2 ppb and for wind directions from $180\text{--}360^\circ$ at
9 6.3 ppb. This trend in the dependence of the observed ozone difference with respect to wind
10 direction is magnified after noon. One possible key driver of differences between observed
11 offshore and shoreline ozone could be the differences in NO_x emissions from each wind
12 direction. The trends with respect to temperature are small in comparison to the trends with
13 respect to wind direction and may be a subtle indicator of the strength of lake breeze effects.
14 Trends with temperature may demonstrate some differences in photochemistry, where some
15 aspects of photochemical ozone production are enhanced with temperature (water vapor
16 content, VOC emissions). Trends with location could be influenced by the distance from
17 emissions sources at the western Lake Michigan shoreline, or lower losses of O_3 to water
18 surfaces compared to terrestrial surfaces (Levy et al., 2010). One complicating factor is that
19 the ferry intercepted air near the surface, whereas urban plumes might reside aloft over an
20 inversion above the lake (Foley et al., 2011; Dye et al., 1995). However, the subtleties of these
21 effects appear to be outweighed by the magnitude of air-mass isolation effects due to local
22 meteorology, as indicated by the large ozone mixing ratio trends with wind and time of day.
23 More complex yet similar observations near Lake Erie were made in summer 2007 during
24 BAQS-Met by Levy et al. (2010) where oscillations in inland ozone were observed at times
25 associated with lake-breeze front movement. The extent to which inversion occurs over the lake

1 at night and ozone precursors and ozone mixing ratios remain high aloft, as suggested by Dye
2 et al. and Foley et al. (Foley et al., 2011; Dye et al., 1995) cannot be evaluated by our
3 measurements at the surface.

4 **3.3 Comparison of ferry ozone with CMAQ experimental model forecasts**

5 The National Air Quality Forecast Model (NAQFM) was developed with the
6 collaboration of the National Oceanic and Atmospheric Administration (NOAA) and the
7 Environmental Protection Agency (EPA) (Eder 2009). The NAQFM is made up of two
8 components: the National Center for Environmental Prediction's (NCEP) North American
9 Mesoscale (NAM) meteorological model and the Environmental Protection Agency's (EPA)
10 Community Multiscale Air Quality (CMAQ) modeling system (Janjic, 2003, Eder 2009, Byun
11 and Schere 2006). The NAM is used to input meteorological conditions into the CMAQ to
12 generate 48h forecasts. Initialization steps to the forecasts are conducted every 12 hours at 06:00
13 and 12:00 UTC (Eder 2009, Chai 2010). The NAQFM provides real-time predictions for
14 ground-level ozone mixing ratios over the contiguous US (Eder 2009) with a 12 km grid size.
15 The NAQFM CMAQ runs in 3 modes: operational, experimental and developmental, with the
16 operational product displayed publicly on the NAQFM web-site (Figure 8, for illustration
17 purposes only, shows an example of the operational product for June 24, 2009, along with the
18 *Lake Express* ferry measurements on that day). Here we compare observations with the
19 developmental model product which used the Carbon Bond Mechanism 5 (CB05) gas-phase
20 chemical mechanism. The emissions inventory used in model forecasts is adopted from from
21 the EPA's 2005 National Emissions Inventory (NEI) (Pan 2014).

22 Hourly output from the developmental CMAQ forecasts were saved for the monitoring
23 season of 2009 from June 18-Sept. 15 2009. The CMAQ output ozone mixing ratios were
24 reported to 1 ppb precision. Figure 9 depicts O₃ forecast levels consistently higher than ferry

1 measurements with 57 days of overlapping data. These forecasts produce a distinct ozone
 2 maximum over the water surfaces of the Great Lakes and, in particular, southern Lake Michigan
 3 (e.g. Figure 8). Statistical comparisons with the Lake Express observations use model grid and
 4 time values determined from ship tracks through the model domain and with no spatial or
 5 temporal interpolation. Figure 10 depicts the sample numbers within distinct model grid cells
 6 for the 3 month time period according to model longitude and central daylight time for the ferry
 7 transects. The extreme western and eastern points are within ports and the Milwaukee model
 8 grid are over land. The model comparison may not be reliable for the shoreline grids due to
 9 local sources and contamination by ferry exhaust. Figure 11 shows the median ozone values for
 10 the forecast 1-24 hours after model initialization (11a), 25-48 hours after initialization (11b),
 11 and *Lake Express* monitor (11c). Figure 11 depicts distinct higher model median O₃ forecasts
 12 in comparison to observations. The maxima in the model forecast O₃ are mid-lake from 15:00-
 13 18:00 CDT. The forecast O₃ mixing ratios are highest after 25-48 hours after initialization,
 14 especially between 14:00 and 21:00 h CDT. The location of the daily maximum ozone from the
 15 ferry is similar to the distribution given by the CMAQ for 1-24h after initialization (Figures
 16 11a,c). The CMAQ predicts the highest median daily maximum O₃ just offshore on the eastern
 17 side of Lake Michigan for 1-24h after initialization (Figure 11a) and a larger area for 25-48h
 18 after initialization (Figure 11b). The correlation coefficients between model and measurement
 19 are high (R=0.85 to 0.95) from 14:00 - 17:00 h CDT for the 1-24h forecast (Figure 12a). The
 20 correlations were reduced for the 25-48h forecast (Figure 12b).

21 The comparison between the ozone forecast and the ferry observations were computed
 22 as bias:

$$23 \quad \text{bias} = p_i - o_i \quad (1)$$

1 where p_i is the model-predicted O_3 concentration and o_i is the observed O_3 concentration on the
2 ferry. Bias was determined for each sample location and time referenced in Figure 10. Model
3 bias is shown in Figure 13. The forecast from 1-24h after initialization in Figure 13a shows an
4 11-16 ppb median O_3 bias for offshore locations, which is highest between 12:00 and 17:00 h
5 CDT. The 24-48h forecast (Figure 13b) has higher biases extending to time periods later in the
6 day. Components of the model were investigated to evaluate differences that may lead to the
7 higher model bias to the eastern side of Lake Michigan. Winds tend to start the day with a north-
8 to-south median wind component, with a switch to south-to-north wind component in the region
9 of 11:00-15:00h CDT for the 1-24h forecast, and an earlier at 8:00h CDT for the 25-48h
10 forecast. This difference in modeling Chicago's northward travelling plume in the 25-48h
11 forecast may lead to the higher O_3 biases for that forecast.

12 CMAQ developmental model biases were also determined at the Kenosha site for ozone,
13 NO_2 , SO_2 and formaldehyde (Figure 14). Ozone was overpredicted in the model for this
14 shoreline measurement for daylight times, with correlations lower than those obtained over
15 water ($R^2 = 0.67$ 1-24h, $R^2 = 0.58$ 25-48h). NO_2 is underpredicted during daylight hours, but not
16 of the same magnitude as the overprediction of ozone ($R^2 = 0.38$ 1-24h, $R^2 = 0.30$ 25-48h).
17 Formaldehyde is consistently underpredicted when it is measured, with effectively no
18 correlation ($R^2 = 0.03$ for both 1-24h and 25-48h forecasts). Gaps in formaldehyde bias are from
19 gaps in formaldehyde data at the Kenosha site. Bias in SO_2 show little trend with respect to time
20 of day and little-to-no correlation ($R^2 = 0.16$ 1-24h, $R^2 = 0.18$ 25-48h).

21 The mid-afternoon O_3 (20:00 UTC) was also determined for all EPA station monitors
22 in the region (Figure 15). The *Lake Express* ferry data were also used to obtain the bias at a
23 similar time (12:30-15:00 h CDT transect), shown in squares in Figure 15. Note that there is an
24 upwind bias in central in western Wisconsin of ~7-8ppb and high biases are observed at some

1 locations near Chicago and the northern Indiana region. The high biases in the Chicago area,
2 and possibly northern Indiana, are likely due to high bias at low O_3 , where the effect of O_3
3 titration by NO_x is not properly captured. The ferry biases are the only ones that are very high
4 in a downwind region with a much smaller effect from local ozone titration, implicating other
5 causes such as the depth of the lake inversion, or too much photochemistry in the model rather
6 than too little titration. The high biases seen over Lake Michigan don't appear to extend strongly
7 inland on either side of the lake.

8 Others have also found the CMAQ to predict ozone mixing ratios that were biased high
9 (Eder et al., 2009; Tang et al., 2009; Zhang et al., 2012a, b; Wilczak et al., 2006). Simon et al.
10 (2012) completed an exhaustive comparison of photochemical performance statistics reported
11 from 2006-2012, whereby national median in mean bias for hourly ozone was approximately 4
12 ppb, for 1-hour maximum ozone was approximately 8 ppb (Simon et al., 2012). In comparison,
13 the bias determined in this study would be higher than 75th percentile of studies of hourly ozone
14 mean bias for 40 studies compiled by Simon et al. (2012). The work presented here represents
15 the first study of CMAQ model bias over the water of Lake Michigan and show a higher bias
16 than over the surrounding land.

17 **4 Conclusions**

18 Observations of shoreline O_3 and ferry O_3 in comparison to forecast O_3 by the developmental
19 NAQFM show more agreement between shoreline and the ferry measurements than between
20 ozone forecasts over the lake and ferry measurements. Shoreline Lake Michigan measurements
21 of O_3 , NO_2 , SO_2 and formaldehyde demonstrated the differences between onshore and offshore
22 air masses. The comparison between ferry-based O_3 observations and shoreline DOAS O_3
23 observations indicated that diurnal changes in ozone mixing ratio were larger than spatial
24 gradients across Lake Michigan, and ozone tended to be higher over Lake Michigan,

1 particularly in the evening. Mesoscale meteorologic processes involving differential heating
2 between the lake and land surfaces produced diurnal cycles of air mass flow between shoreline
3 environments and offshore, which complicated the understanding of offshore ozone dynamics.
4 Model forecast O₃ is highly correlated with ferry monitor observations, but with afternoon
5 median biases ranging from 11-16 ppb, compared to 6-9 ppb biases for land-based monitors
6 just west of Lake Michigan. The model O₃ overpredictions over water are similar to those
7 determined for the Kenosha site, though formaldehyde and NO₂ are underpredicted. The
8 developmental NAQFM showed a trend of increasing O₃ bias to the eastern side of Lake
9 Michigan, and a larger bias for the second day forecast compared to the first 24 hours. Further
10 analyses are required to determine whether NAQFM predictions might be improved by
11 adjusting model parameters related to emission sources, localized shoreline meteorology, or
12 atmospheric chemistry.

13 **Acknowledgements**

14 The authors would like to thank Kaya Sims, Lindsey Kuettner and Renee Hanson for their
15 assistance in this experiment, the *Lake Express* ferry, University of Wisconsin-Eau Claire
16 Office of Sponsored Programs Faculty and Student Collaboration Grant, Great Lakes Water
17 Institute, Kenosha Water Utility and the Great Lakes Observing System for their cooperation
18 and support of this project. The authors would like to thank Bruce E. Brown for assistance with
19 collection and calibration of ozone data from the Lake Express ferry, and Kenneth Aikin for
20 archiving of NAQFM images. Thomas Langel acknowledges the NOAA Hollings Scholar
21 Program for fellowship support during 2010. SSB acknowledges support from NOAA's
22 Atmospheric Chemistry, Carbon Cycle and Climate Program.

23

24

References

- Brown, S. S., Dibb, J. E., Stark, H., Aldener, M., Vozella, M., Whitlow, S., Williams, E. J., Lerner, B. M., Jakoubek, R., Middlebrook, A. M., DeGouw, J. A., Warneke, C., Goldan, P. D., Kuster, W. C., Angevine, W. M., Sueper, D. T., Quinn, P. K., Bates, T. S., Meagher, J. F., Fehsenfeld, F. C., and Ravishankara, A. R.: Nighttime removal of NO_x in the summer marine boundary layer, *Geophysical Research Letters*, 31, 10.1029/2004gl019412, 2004.
- Carpenter, L. J., Sturges, W. T., Penkett, S. A., Liss, P. S., Alicke, B., Hebestreit, K., and Platt, U.: Short-lived alkyl iodides and bromides at Mace Head, Ireland: Links to biogenic sources and halogen oxide production, *Journal of Geophysical Research-Atmospheres*, 104, 1679-1689, 10.1029/98jd02746, 1999.
- Dye, T. S., Roberts, P. T., and Korc, M. E.: Observations of transport processes for ozone and ozone precursors during the 1991 Lake Michigan Ozone Study, *Journal of Applied Meteorology*, 34, 1877-1889, 10.1175/1520-0450(1995)034<1877:ootpfo>2.0.co;2, 1995.
- Eder, B., Kang, D. W., Mathur, R., Pleim, J., Yu, S. C., Otte, T., and Pouliot, G.: A performance evaluation of the National Air Quality Forecast Capability for the summer of 2007, *Atmospheric Environment*, 43, 2312-2320, 10.1016/j.atmosenv.2009.01.033, 2009.
- EPA: National Air Quality Standards for Ozone; Proposed Rule, in: 40 CFR Parts 50, 51, 52, et al., 242, *Federal Register*, 75234-754111, 2014.
- Fast, J. D., and Heilman, W. E.: The effect of lake temperatures and emissions on ozone exposure in the western Great Lakes region, *Journal of Applied Meteorology*, 42, 1197-1217, 10.1175/1520-0450(2003)042<1197:teolta>2.0.co;2, 2003.
- Fast, J. D., and Heilman, W. E.: Simulated sensitivity of seasonal ozone exposure in the Great Lakes region to changes in anthropogenic emissions in the presence of interannual variability, *Atmospheric Environment*, 39, 5291-5306, 10.1016/j.atmosenv.2005.05.032, 2005.
- Foley, T., Betterton, E. A., Jacko, P. E. R., and Hillery, J.: Lake Michigan air quality: The 1994-2003 LADCO Aircraft Project (LAP), *Atmospheric Environment*, 45, 3192-3202, 10.1016/j.atmosenv.2011.02.033, 2011.
- Hanna, S. R., and Chang, J. C.: Relations between meteorology and ozone in the Lake Michigan region, *Journal of Applied Meteorology*, 34, 670-678, 10.1175/1520-0450(1995)034<0670:rbmaoi>2.0.co;2, 1995.
- Harris, L., and Kotamarthi, V. R.: The characteristics of the Chicago Lake breeze and its effects on trace particle transport: Results from an episodic event simulation, *Journal of Applied Meteorology*, 44, 1637-1654, 10.1175/jam2301.1, 2005.
- Keen, C. S., and Lyons, W. A.: Lake/Land Breeze circulations on the western shore of Lake Michigan, *Journal of Applied Meteorology*, 17, 1843-1855, 10.1175/1520-0450(1978)017<1843:lbcotw>2.0.co;2, 1978.

- 1 Lennartson, G. J., and Schwartz, M. D.: A synoptic climatology of surface-level ozone in
2 Eastern Wisconsin, USA, *Climate Research*, 13, 207-220, 10.3354/cr013207, 1999.
- 3
- 4 Lennartson, G. J., and Schwartz, M. D.: The lake breeze-ground-level ozone connection in
5 eastern Wisconsin: A climatological perspective, *International Journal of Climatology*, 22,
6 1347-1364, 10.1002/joc.802, 2002.
- 7
- 8 Levy, I., Makar, P. A., Sills, D., Zhang, J., Hayden, K. L., Mihele, C., Narayan, J., Moran, M.
9 D., Sjostedt, S., and Brook, J.: Unraveling the complex local-scale flows influencing ozone
10 patterns in the southern Great Lakes of North America, *Atmospheric Chemistry and Physics*,
11 10, 10895-10915, 10.5194/acp-10-10895-2010, 2010.
- 12
- 13 Lyons, W. A., and Olsson, L. E.: Detailed mesometeorological studies of air pollution
14 dispersion in Chicago lake breeze, *Monthly Weather Review*, 101, 387-403, 10.1175/1520-
15 0493(1973)101<0387:dmsop>2.3.co;2, 1973.
- 16
- 17 Lyons, W. A., and Cole, H. S.: Photochemical oxidant transport - Mesoscale lake breeze and
18 synoptic-scale aspects, *Journal of Applied Meteorology*, 15, 733-743, 10.1175/1520-
19 0450(1976)015<0733:potmlb>2.0.co;2, 1976.
- 20
- 21 Makar, P. A., Gong, W., Mooney, C., Zhang, J., Davignon, D., Samaali, M., Moran, M. D.,
22 He, H., Tarasick, D. W., Sills, D., and Chen, J.: Dynamic adjustment of climatological ozone
23 boundary conditions for air-quality forecasts, *Atmospheric Chemistry and Physics*, 10, 8997-
24 9015, 10.5194/acp-10-8997-2010, 2010.
- 25
- 26 Martin, J. C. G., Mahajan, A. S., Hay, T. D., Prados-Roman, C., Ordonez, C., MacDonald, S.
27 M., Plane, J. M. C., Sorribas, M., Gil, M., Mora, J. F. P., Reyes, M. V. A., Oram, D. E.,
28 Leedham, E., and Saiz-Lopez, A.: Iodine chemistry in the eastern Pacific marine boundary
29 layer, *Journal of Geophysical Research-Atmospheres*, 118, 887-904, 10.1002/jgrd.50132,
30 2013.
- 31
- 32 Martinez, M., Perner, D., Hackenthal, E. M., Kulzer, S., and Schutz, L.: NO₃ at Helgoland
33 during the NORDEX campaign in October 1996, *Journal of Geophysical Research-*
34 *Atmospheres*, 105, 22685-22695, 10.1029/2000jd900255, 2000.
- 35
- 36 Pugliese, S. C., Murphy, J. G., Geddes, J. A., and Wang, J. M.: The impacts of precursor
37 reduction and meteorology on ground-level ozone in the Greater Toronto Area, *Atmospheric*
38 *Chemistry and Physics*, 14, 8197-8207, 10.5194/acp-14-8197-2014, 2014.
- 39
- 40 Rivera, C., Mellqvist, J., Samuelsson, J., Lefer, B., Alvarez, S., and Patel, M. R.:
41 Quantification of NO₂ and SO₂ emissions from the Houston Ship Channel and Texas City
42 industrial areas during the 2006 Texas Air Quality Study, *Journal of Geophysical Research-*
43 *Atmospheres*, 115, 10, 10.1029/2009jd012675, 2010.
- 44
- 45 Seitz, K., Buxmann, J., Pohler, D., Sommer, T., Tschritter, J., Neary, T., O'Dowd, C., and
46 Platt, U.: The spatial distribution of the reactive iodine species IO from simultaneous active
47 and passive DOAS observations, *Atmospheric Chemistry and Physics*, 10, 2117-2128, 2010.
- 48

- 1 Sills, D. M. L., Brook, J. R., Levy, I., Makar, P. A., Zhang, J., and Taylor, P. A.: Lake breezes
2 in the southern Great Lakes region and their influence during BAQS-Met 2007, *Atmospheric*
3 *Chemistry and Physics*, 11, 7955-7973, 10.5194/acp-11-7955-2011, 2011.
- 4
- 5 Simon, H., Baker, K. R., and Phillips, S.: Compilation and interpretation of photochemical
6 model performance statistics published between 2006 and 2012, *Atmospheric Environment*,
7 61, 124-139, 10.1016/j.atmosenv.2012.07.012, 2012.
- 8
- 9 Takashima, H., Irie, H., Kanaya, Y., and Akimoto, H.: Enhanced NO₂ at Okinawa Island,
10 Japan caused by rapid air-mass transport from China as observed by MAX-DOAS,
11 *Atmospheric Environment*, 45, 2593-2597, 10.1016/j.atmosenv.2010.10.055, 2011.
- 12
- 13 Tang, Y. H., Lee, P., Tsidulko, M., Huang, H. C., McQueen, J. T., DiMego, G. J., Emmons,
14 L. K., Pierce, R. B., Thompson, A. M., Lin, H. M., Kang, D. W., Tong, D., Yu, S. C., Mathur,
15 R., Pleim, J. E., Otte, T. L., Pouliot, G., Young, J. O., Schere, K. L., Davidson, P. M., and
16 Stajner, I.: The impact of chemical lateral boundary conditions on CMAQ predictions of
17 tropospheric ozone over the continental United States, *Environmental Fluid Mechanics*, 9, 43-
18 58, 10.1007/s10652-008-9092-5, 2009.
- 19
- 20 Vrekoussis, M., Kanakidou, M., Mihalopoulos, N., Crutzen, P. J., Lelieveld, J., Perner, D.,
21 Berresheim, H., and Baboukas, E.: Role of the NO₃ radicals in oxidation processes in the
22 eastern Mediterranean troposphere during the MINOS campaign, *Atmospheric Chemistry and*
23 *Physics*, 4, 169-182, 2004.
- 24
- 25 White, M. L., Russo, R. S., Zhou, Y., Mao, H., Varner, R. K., Ambrose, J., Veres, P.,
26 Wingenter, O. W., Haase, K., Stutz, J., Talbot, R., and Sive, B. C.: Volatile organic
27 compounds in northern New England marine and continental environments during the
28 ICARTT 2004 campaign, *Journal of Geophysical Research-Atmospheres*, 113, 16,
29 10.1029/2007jd009161, 2008.
- 30
- 31 Wilczak, J., McKeen, S., Djalalova, I., Grell, G., Peckham, S., Gong, W., Bouchet, V.,
32 Moffet, R., McHenry, J., McQueen, J., Lee, P., Tang, Y., and Carmichael, G. R.: Bias-
33 corrected ensemble and probabilistic forecasts of surface ozone over eastern North America
34 during the summer of 2004, *Journal of Geophysical Research-Atmospheres*, 111,
35 10.1029/2006jd007598, 2006.
- 36
- 37 Zhang, Y., Bocquet, M., Mallet, V., Seigneur, C., and Baklanov, A.: Real-time air quality
38 forecasting, part I: History, techniques, and current status, *Atmospheric Environment*, 60,
39 632-655, 10.1016/j.atmosenv.2012.06.031, 2012a.
- 40
- 41 Zhang, Y., Bocquet, M., Mallet, V., Seigneur, C., and Baklanov, A.: Real-time air quality
42 forecasting, part II: State of the science, current research needs, and future prospects,
43 *Atmospheric Environment*, 60, 656-676, 10.1016/j.atmosenv.2012.02.041, 2012b.
- 44
- 45
- 46

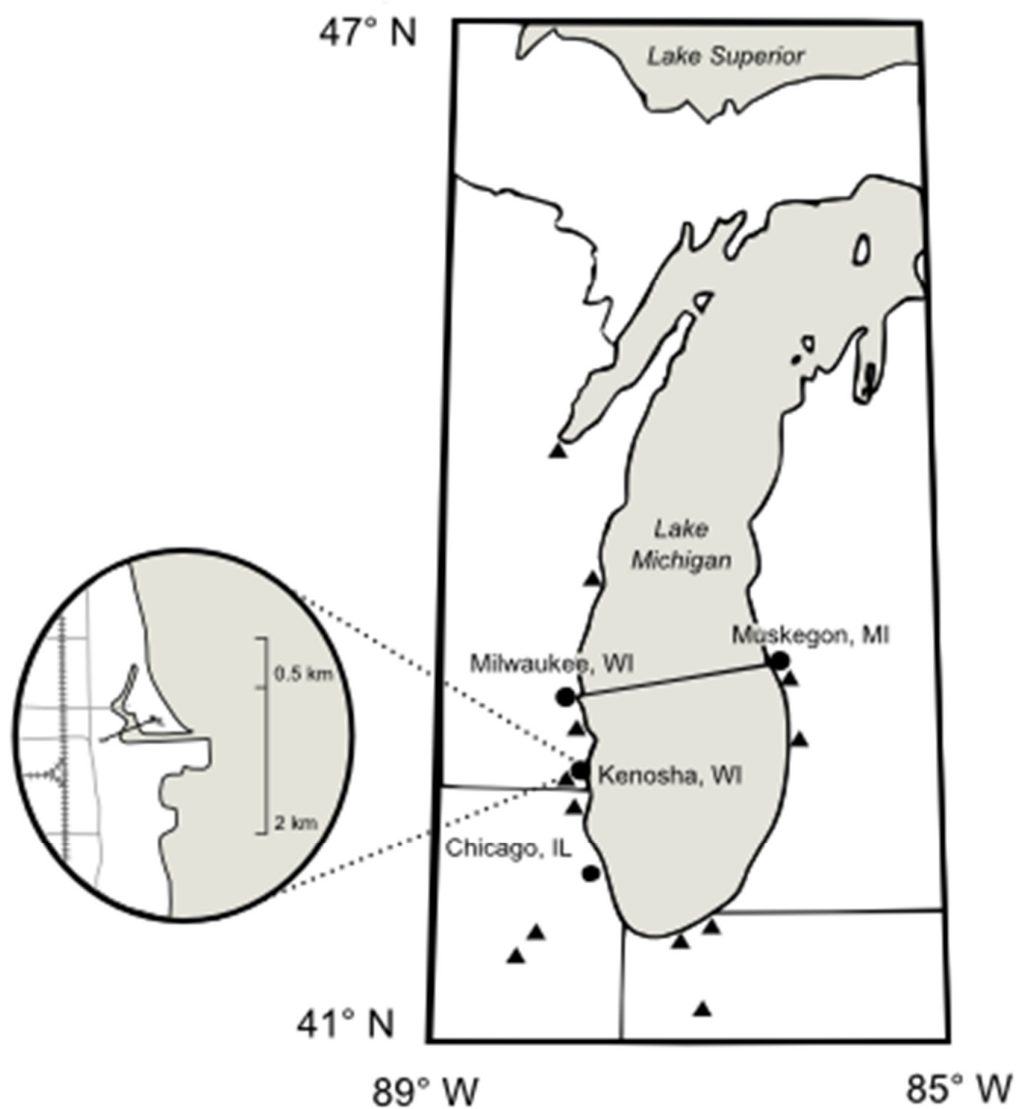


Figure 1: Map of experiment. Path of ferry from Milwaukee, Wisconsin to Muskegon, Michigan is shown with black line across the lake in the map. The DOAS instrument was placed at the Kenosha, Wisconsin harbor with the beam path shown (inset) as the dark line across the harbor. Coal fired power plants with power capacity greater than 400 MW are shown as black triangles.

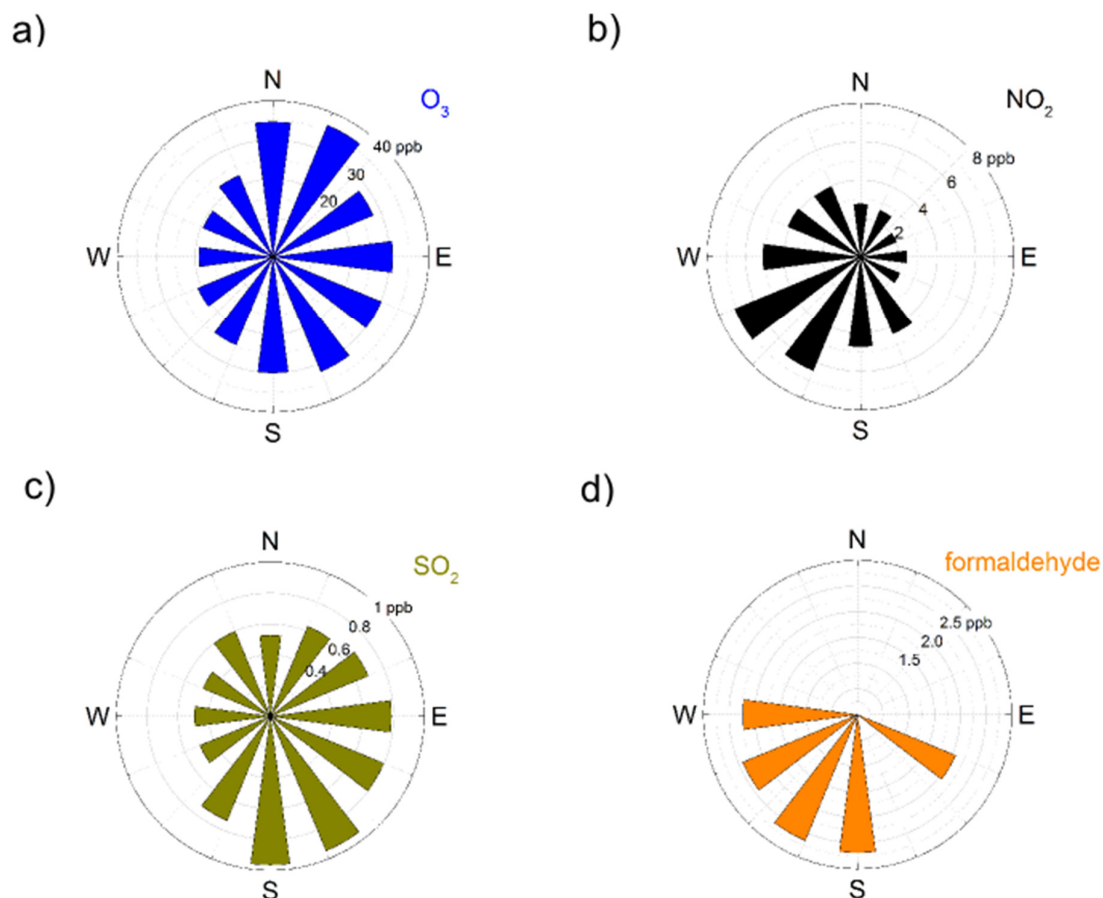


Figure 2. Wind rose depictions of median mixing ratio of a) O_3 b) NO_2 c) SO_2 and d) formaldehyde with respect to wind direction as measured by DOAS at Kenosha harbor from April-November of 2009. Medians are not reported for wind directions where few measurements ($n < 75$ for 30 minute averaged data points) were above the detection limit (d.l. = 1.5 ppb for formaldehyde).

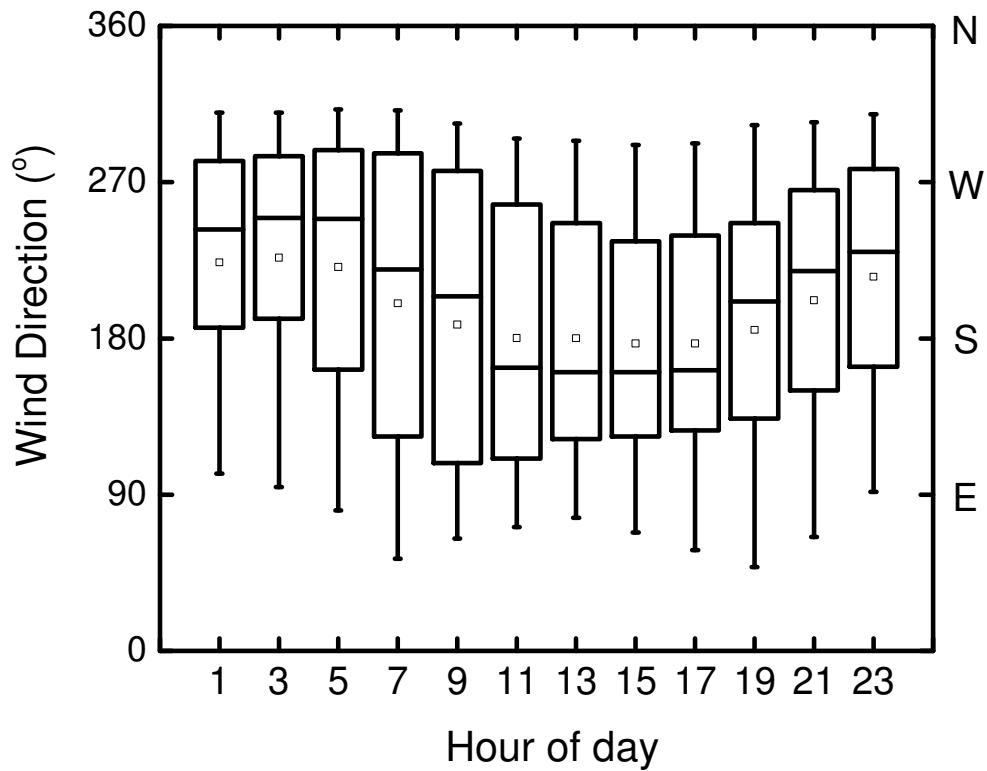


Figure 3: Wind direction as a function of time of day as measured at Kenosha harbor from April-November of 2009. Box plots show mean (\square), median (centerline), 25%-75% (box) and 10-90% (whiskers).

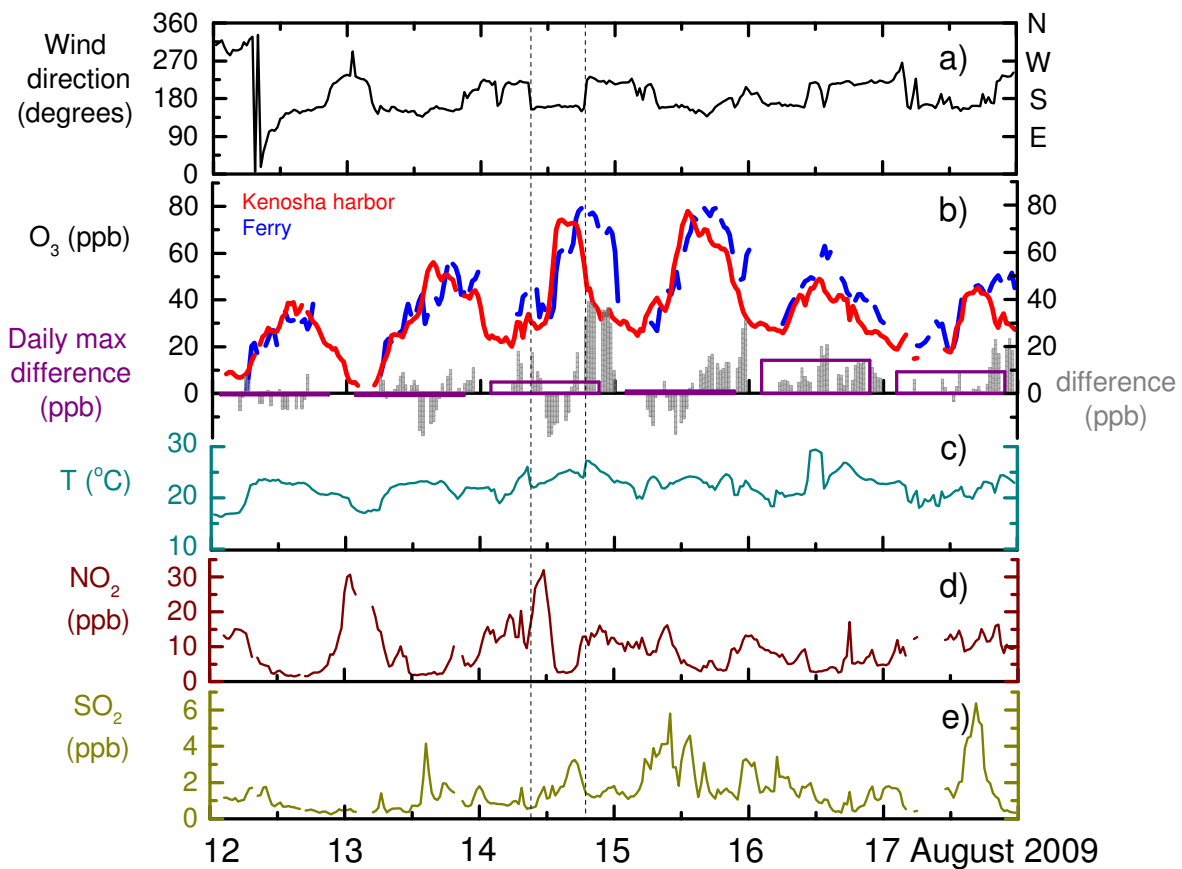


Figure 4: Example period of observations from Aug. 12, 2009 to Aug. 18, 2009 a) wind direction at Kenosha Harbor site, b) concurrent O₃ observations from Kenosha Harbor and *Lake Express* in transit, their 30 minute average O₃ (Ferry) - O₃ (Kenosha Harbor) difference and daily max difference c) temperature at Kenosha Harbor in Celsius d) NO₂ observations from Kenosha Harbor and e) SO₂ observations from Kenosha Harbor.

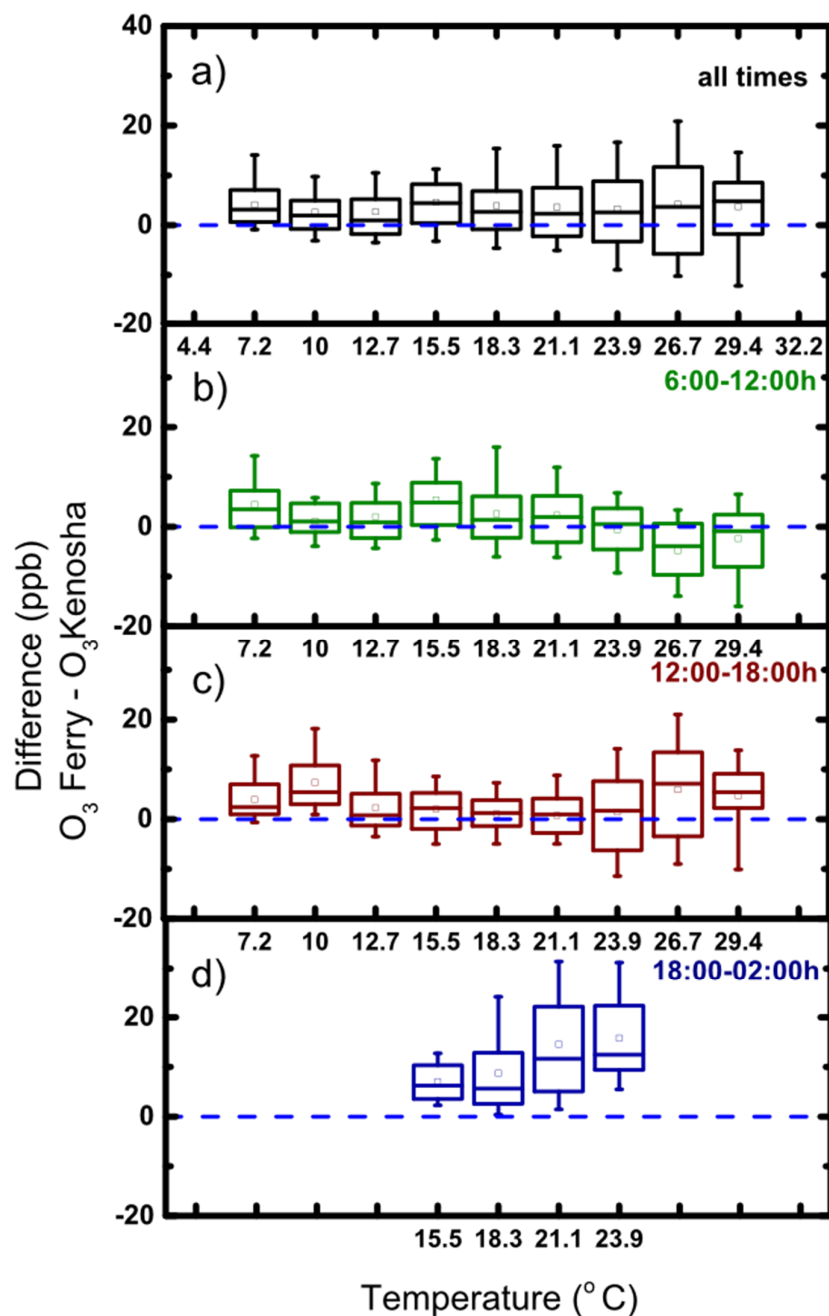


Figure 5: Difference in O₃ observations between platforms with respect to temperature (°C) measured at the shoreline for a) all times, b) morning (06:00-12:00h CDT), c) early afternoon (12:00-16:00h CDT) and d) late afternoon/evening (16:00-02:00h). Box plots show mean (□), median (centerline), 25%-75% (box) and 10-90% (whiskers). Each box represents a minimum of 15 points.

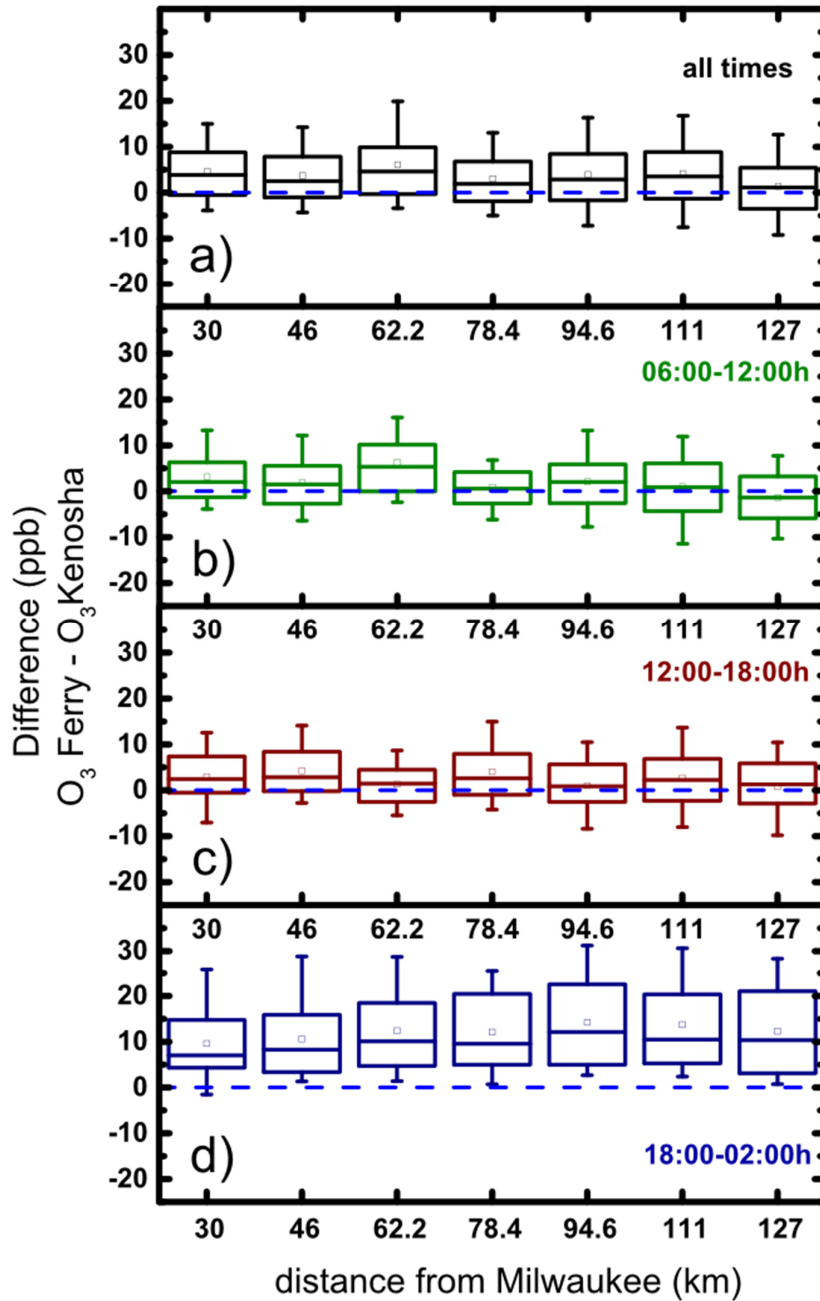


Figure 6: Difference in O₃ observations between platforms with respect to position of the ferry as indicated by km from Milwaukee along ferry path at: a) all times, b) morning (06:00-12:00h CDT), c) early afternoon (12:00-16:00h CDT) and d) late afternoon/evening (16:00-02:00h). Box plots show mean (\square), median (centerline), 25%-75% (box) and 10-90% (whiskers). Each box plot represents a minimum of 12 points.

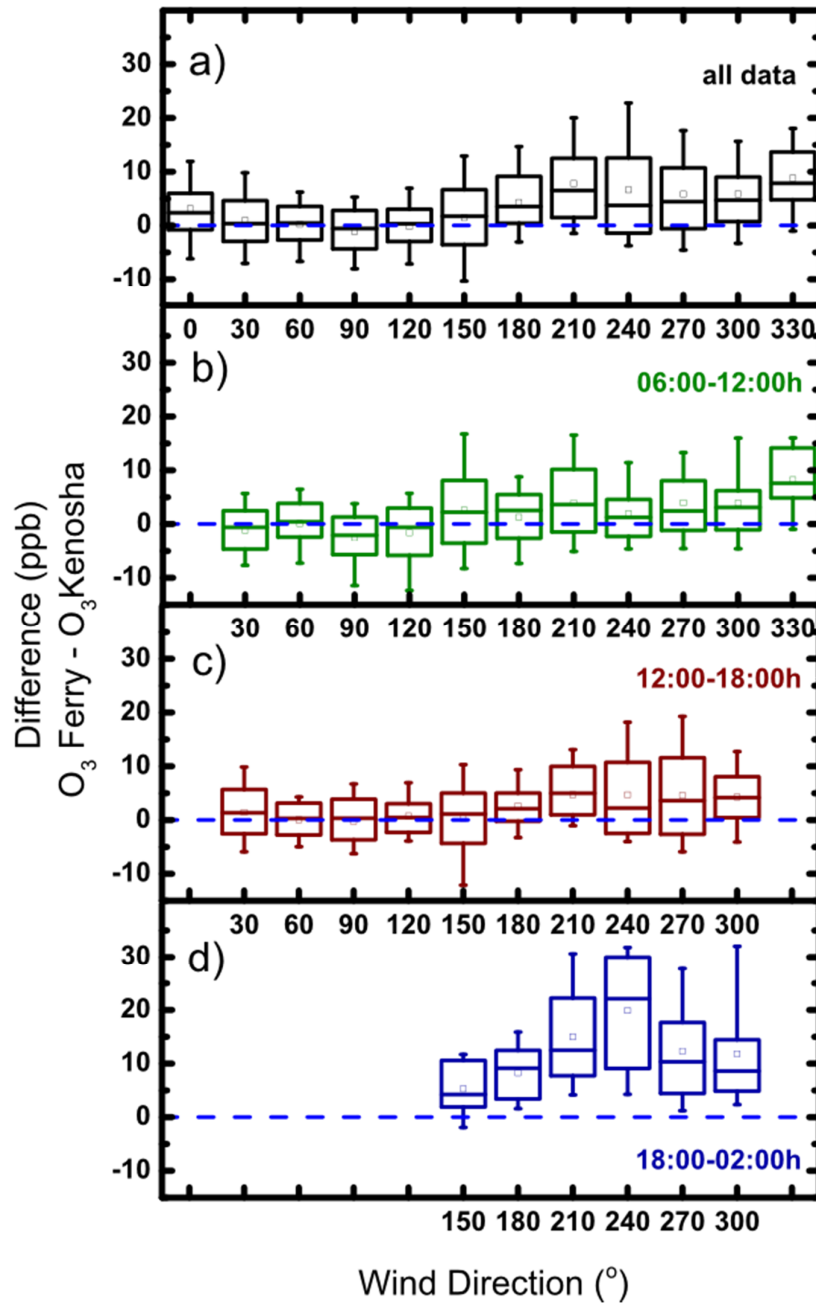


Figure 7 : Difference in O₃ observations between platforms with respect to wind direction measured at Kenosha harbor for a) all times, b) morning (06:00-12:00h CDT), c) early afternoon (12:00-16:00h CDT) and d) late afternoon/evening (16:00-02:00h). Box plots show mean (\square), median (centerline), 25%-75% (box) and 10-90% (whiskers). Each box represents a minimum of 15 points.

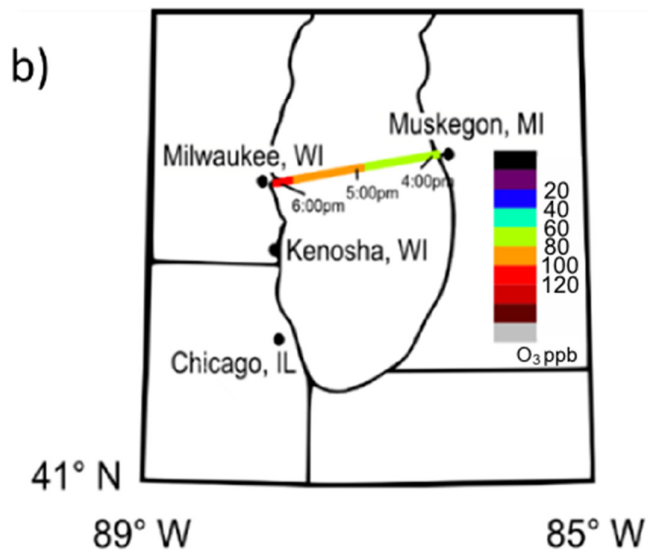
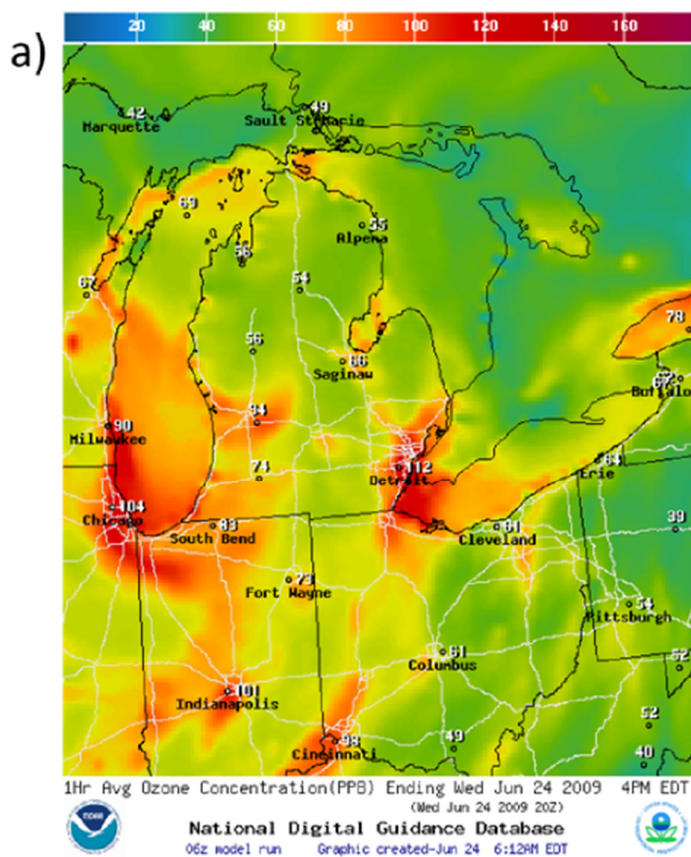


Figure 8: a) Sample image of National Air Quality Forecast Model (NAQFM) during the campaign period, b) O₃ measurements for one ferry trip on June 24, 2009 where the ferry was in transit from 3:50 pm (CDT) to 6:15 pm (CDT).

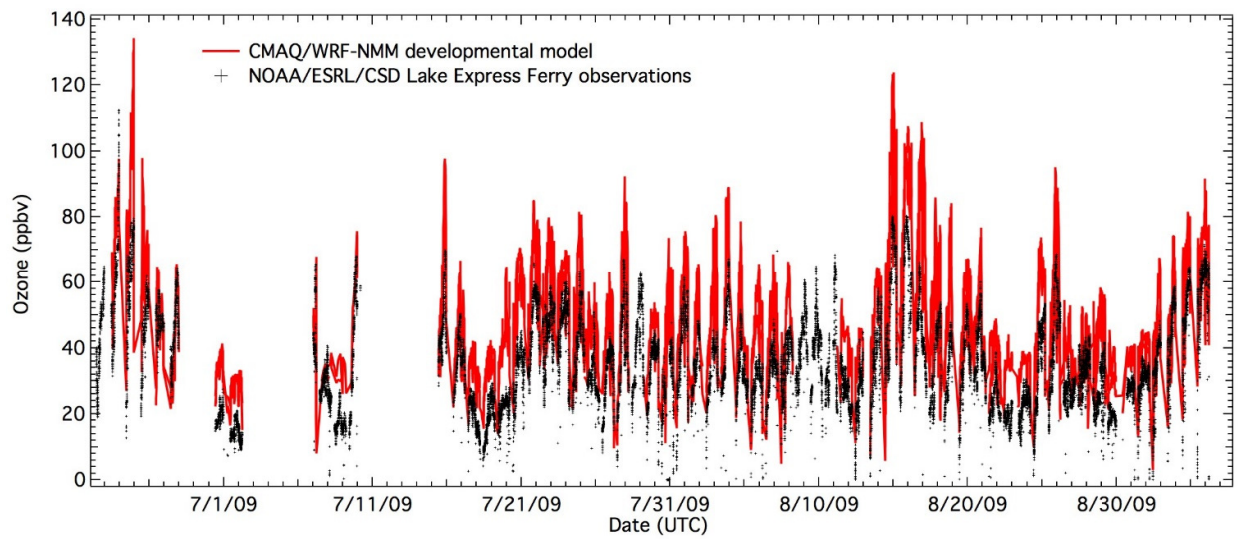


Figure 9: Graph of all CMAQ model forecast ozone mixing ratios in red with *Lake Express* Ferry observations in black from 2009.

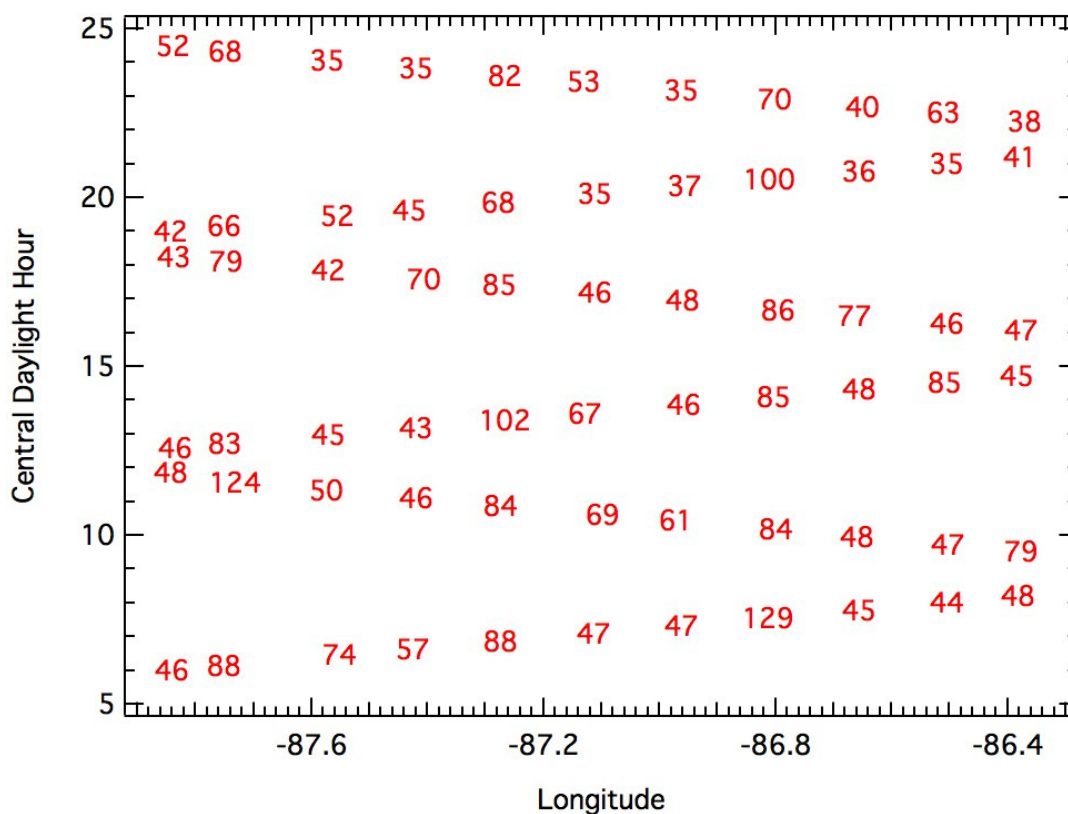
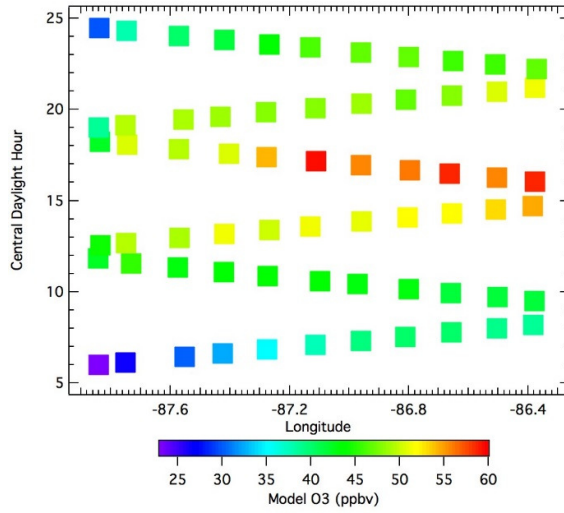
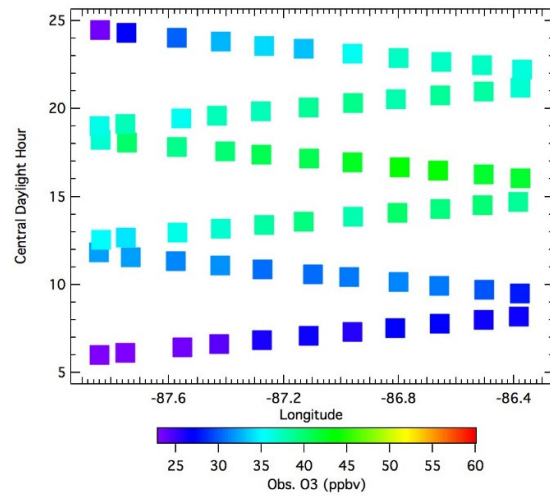


Figure 10: Statistical data for CMAQ model and ferry measurement comparison. Each model grid value and observation averages were binned according to model west-east grid number and CDT time of the ferry transect. The 1-min O₃ observations were averaged over model grid and hourly output. The numbers here are the number of hourly comparisons between model grid values and hourly averaged O₃ observations via ferry.

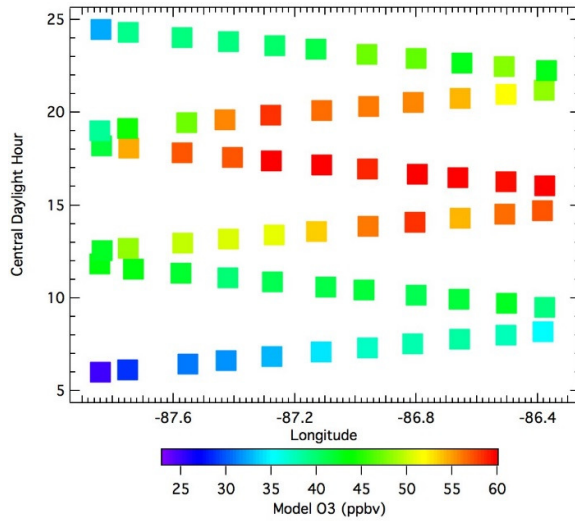
1 a)



c)



2 b)

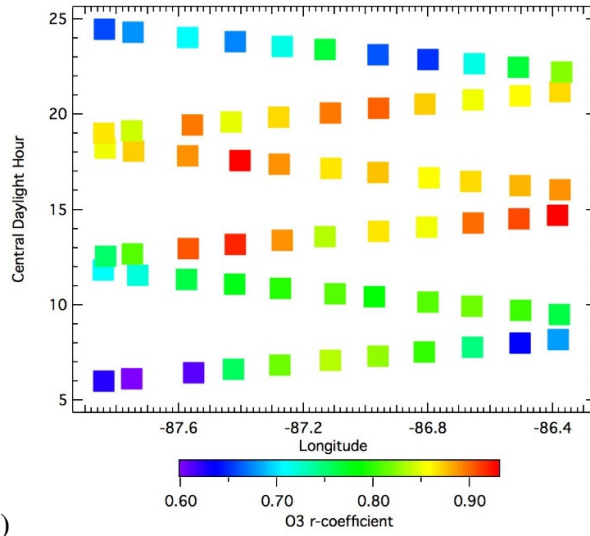


11

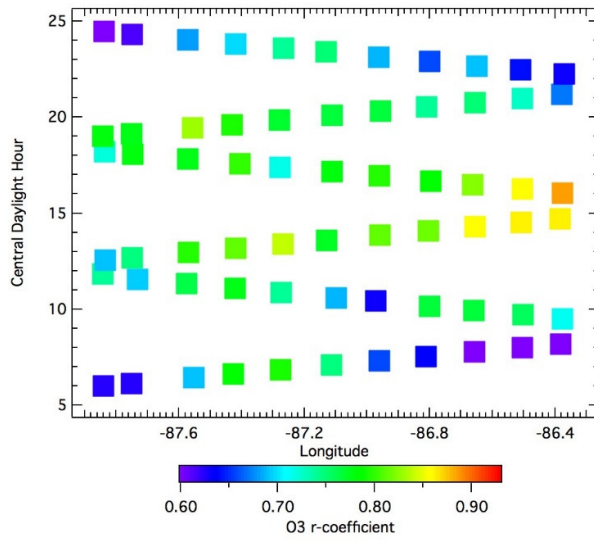
12 **Figure 11:** Median O₃ from a) 1-24hr CMAQ forecasts b) 25-48 CMAQ forecasts and c) ferry
13 observations.

14

1 a)



2 b)



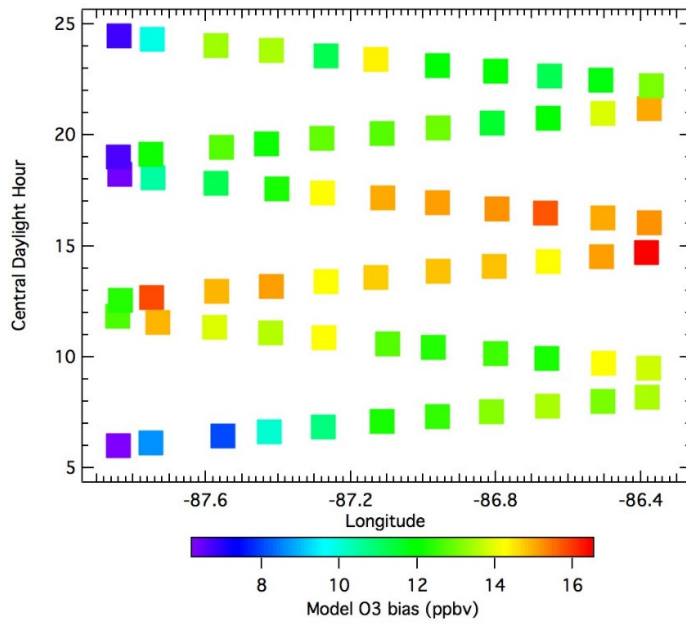
3

4 **Figure 12:** Correlation coefficients for model-measurement comparison for each bin a) 1-24h
5 forecast b) 25-48h forecast

6

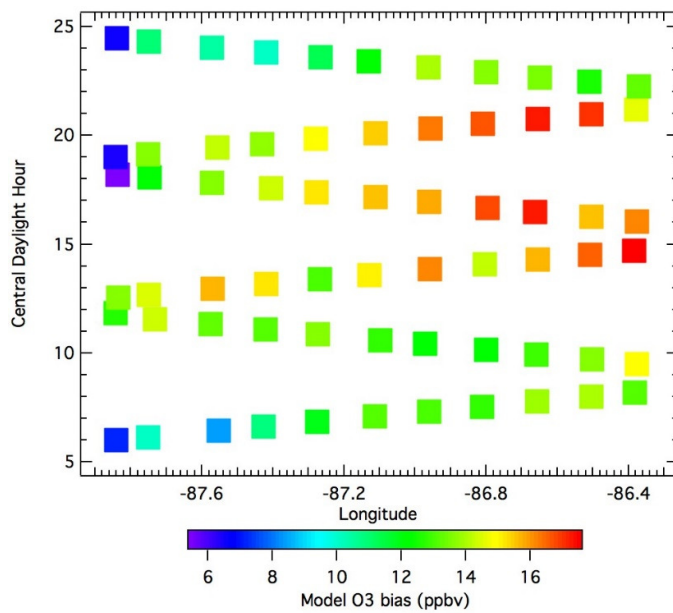
7

1 a)



2

3 b)



4

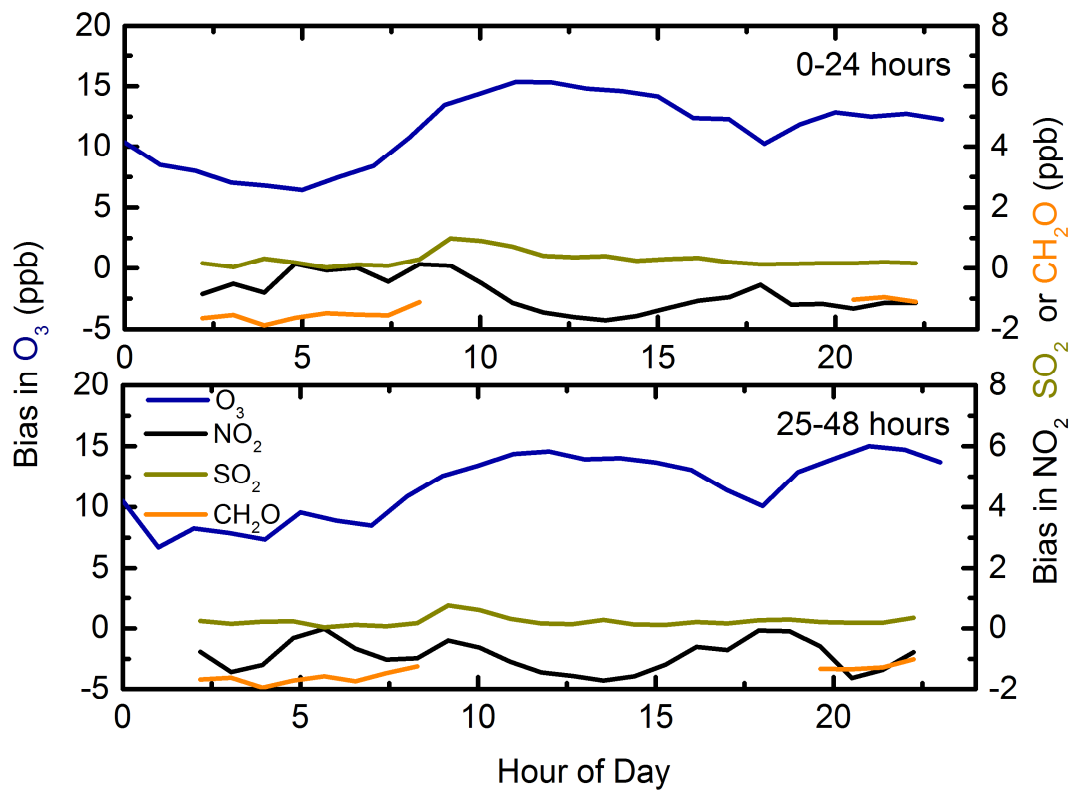
5 **Figure 13:** CMAQ model bias from a) 1-24h forecast and b) 25-48h forecast

6

7

8

1

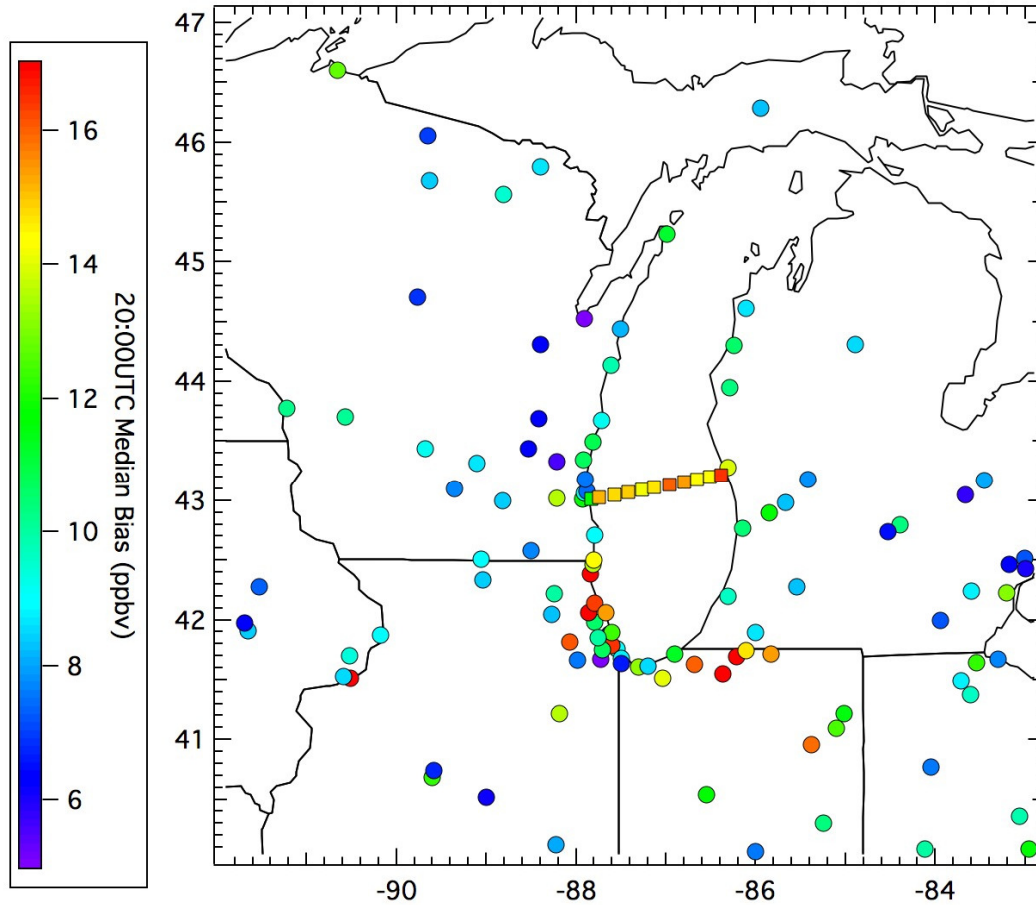


2

3 **Figure 14:** CMAQ model bias at Kenosha for O₃ (in blue, left axis), NO₂ (black), SO₂ (brown),
 4 or formaldehyde (orange) (right axis) for a) 1-24h forecast and b) 25-48 h forecast.

5

6



2 **Figure 15:** CMAQ model O₃ bias for air quality EPA station monitors (circles) and *Lake*
 3 *Express* ferry (boxes). EPA monitor biases are calculated at 20:00 UTC (3:00pm CDT), and
 4 the data has been windowed for only those days when *Lake Express* ferry
 5 data is available. For the *Lake Express* ferry data are from the 12:30 to 3:00 pm (CDT)
 6 transect statistics.