Response to Comments by Anonymous Referee #1

Comment: In this article, the authors describe the hemispheric version of the Community Multiscale Air Quality (CMAQ) modeling system and present a variety of applications of the system for evaluation. In general, the paper is of good quality and should be published with minor revisions as detailed below, although it is a description paper of a new model version and does not contain any truly new science.

Response: We thank the reviewer for the overall positive assessment of our manuscript and for the constructive suggestions for improvements. Detailed below is our response to the specific reviewer comments and the changes incorporated in the revised manuscript. We believe the incorporation of the reviewer suggestions has helped improve the quality of the revised manuscript.

The extension of CMAQ to hemispheric scales required enhancements to both the model’s structural and process attributes, to adequately represent the expanded space and time-scales. We do feel that the incorporation, synthesis, and systematic evaluation of these enhancements conveyed in this paper, does represent new modeling science.

Comment: I think the authors should make it clearer in the Introduction that Figure 1 is simply a characterization of the regional-scale CMAQ model and does not necessarily represent the what actually happens in the real atmosphere. CMAQ is known to be rather diffusive (Emery et al. 2011; Garcia-Menendez et al. 2010; Mathur 2008) and probably does not represent this transport very faithfully. Some readers might be fooled into believing that the fractions presented in Figure 1 are realistic, when they are probably not.

Response: Figure 1 illustrates the impact of lateral boundary conditions on the simulated free tropospheric and surface concentration variability. The discussion explicitly states that Figure 1 illustrates the influence of “LBC specification on simulated surface-level concentrations across a typical regional modeling domain covering the contiguous U.S.”. The subsequent sentences then describe the model tracer species calculations that were used in constructing the Figure. The figure caption also states that these are model simulated concentrations from CMAQ.

The reviewer’s somewhat philosophical question on “what happens in the real atmosphere?” speculates on whether this specific model characterization agrees with those from other models and measurements. The results illustrated in Figure 1 agree with the well-established conceptual understanding (built off both measurements and modeling analysis) of long-range transport where in source regions, pollutants within the boundary layer are convectively lofted to the free troposphere where they are intercontinentally transported over long distances by efficient winds and can subsequently be entrained to the surface (subsidence, cloud and boundary layer mixing) in receptor regions thereby regulating “background” concentrations of the receptor region. This conceptual view also implicitly suggests that surface background concentrations are likely influenced by free tropospheric values since that’s where transport is more efficient, and that’s what the model tracer concentrations suggest. Of course the extrapolation of the illustrated fractions to a specific atmospheric species would also require consideration of specific chemical and physical sinks for that species, and in our discussion of results we have attempted to be deliberate of that distinction. Nevertheless, the relative importance of specification of free-
tropospheric LBCs (representative of long-range pollutant) transport in any model characterization of “background” pollution cannot be diminished, and that’s what the figure is intended to illustrate.

We reread the three papers the reviewer brought to our attention. Mathur et al. (2008) analyze the impact of long-range transport of pollution from the Alaskan fires through complementary analysis of CMAQ simulations, in-situ surface and aloft measurements, and satellite retrievals and show impact on surface concentrations at distant sites in the continental U.S. We are not aware of any suggestions of impacts of excessive diffusive transport characteristics in that analysis. Garcia-Menendez et al. (2010) describe the implementation of an adaptive grid methodology in CMAQ to better resolve horizontal concentration gradients in plumes as they undergo lateral transport, but relative to a fixed resolution coarser grid. Their results are applicable to any model that utilizes a uniform discretization, but do not necessarily point to a systematic issue in the CMAQ modeling system. Emery et al. (2011) analyzed vertical transport in the CAMx model, found excessive vertical transport in mountainous regions due to a combination of using a low-order advection scheme, coarse vertical resolution, and likely artifacts of translation of dynamical information between different vertical grid structures employed by the meteorological model and their CTM. Though they did not present any analysis of CMAQ based calculations, the discussion in their manuscript speculates that models such as CMAQ may exhibit similar characteristics. The extrapolation of the Emery et al (2011) analysis with a different modeling system to the current CMAQ tracer results presented here is not straightforward given the differences in model formulations employed in these tests (advection in Emery vs advection, turbulent and cloud transport, dry deposition and wet scavenging here), set-up (different grid resolutions and coupling with meteorological model both in terms of layer configuration and process representation) and since significant changes have occurred in the CMAQ modeling system over the past 6 years. We thus respectfully disagree with the reviewer’s assertion that “CMAQ is known to be rather diffusive”, but acknowledge that model inferences of impacts of vertical transport from advection, turbulent mixing, and cloud transport are influenced by the choice of horizontal and vertical grid structures, accuracy of numerical methods employed, and consistency in coupling of these aspects with the driving dynamical model (as also discussed in Section 2.1 of our manuscript).

Comment: The results of Figure 3 are fascinating. Increasing the vertical resolution from 35 layers to 44 layers substantially reduced ozone profiles in the lower atmosphere. This immediately begs the question as to what would happen if the number of layers was increased to 60 or 70! A pet peeve of this reviewer is that air quality and atmospheric chemistry models are not rigorously evaluated as numerical models. In any basic numerical modeling class, one is taught to increase grid resolutions until the solution converges to a consistent result. This is _never_ done in 3-D atmospheric chemistry modeling! How much different would the results be if this simple numerical procedure was carried out? My guess is quite different.

Response: The reviewer raises a pertinent issue on the optimal vertical grid structure that should be employed in atmospheric chemistry models. As indicated by Figure 3, the associated discussion in Section 2.1, and our response to the previous comment, more attention needs to be devoted to the vertical grid structure employed by the models. The 44-layer structure depicted in Figure 2 was judiciously designed to provide higher resolution above the boundary layer (nominally at altitudes
and near the tropopause to resolve the sharp gradients in O₃ mixing ratios. Clearly, employing 60 or 70 layers will improve the resolution further, but greater impacts will be seen if the additional layers are deployed near the tropopause. We whole heartedly agree with the reviewer that rigorous modeling protocols need to be developed to define appropriate layer configurations and vertical and horizontal grid resolutions for specific model applications, and numerical convergence tests that establish the order of accuracy of the models be conducted on a more routine basis. Pragmatically, this is a challenging endeavor as it is not only application-dependent (for instance, an optimal layer configuration designed specifically for resolving stratosphere-troposphere exchange cases may be different from one for resolving boundary layer venting) and also because some parameters (e.g., mixing height, definition of cloud base and top, representation of wind-shear) depend on the discrete layer structure. We echo the reviewer’s concerns in the Summary and Concluding Remarks section by re-emphasizing the need for additional model sensitivity simulations to assess the impact of different “model vertical extent and vertical grid resolution” (see page 21).

Comment: In the description of the model, the authors in several cases describe what is in the version being presented in this paper, but also describe improvements that are or have been worked on. Examples of this include ... (i) seven NTR species rather than 1; (ii) the marine environment chemistry and deposition; (iii) windblown dust parameterization; and, (iv) the ozone-PV parameterization. In these discussions, it’s not always clear what’s included in this model version used in this paper and what’s just an "advertisement" of the improvements to come in the future.

Response: Since the manuscript describes the extension of the CMAQ modeling system from the traditional regional scale to the expanded hemispheric scale, we have outlined the process representations in CMAQ and how they were enhanced for the extended space and time scales for hemispheric applications. This manuscript is an overview of a modeling system that has evolved over several years, so results are presented from model applications that spanned the development period and contributed to the extension of the CMAQ system. For instance, the initial constant PV-scaling approach helped in the conduct of the multi-decadal simulations. The analysis and availability of the fields from these simulations then enabled the development of a more robust space and time varying PV-scaling parameterization. Similarly, the analysis of NTR species and the initial simple approach to modulate its atmospheric lifetime prompted the investigation of the expanded NTR scheme as well as examination of the RACM2 mechanism.

We agree with the reviewer that it is important to clearly convey what process state is being examined and how the model configurations across the different applications may differ. To address the reviewer’s concern we reviewed the descriptions of the model configuration in each section to ensure that the model/process state used in the specific runs is stated clearly.

Comment: For Figure 6c, the authors make the dubious statement ... "The comparisons in Figure 6c further show that CMAQ captures the SO₄ enhancements in the free troposphere associated
with this episodic event." In looking at this figure, I find it very hard to not laugh out loud when reading this sentence! The observed and modeled SO4 values are of the same general magnitude, but don’t seem to be correlated at ALL. I think the authors should be more truthful in their comments about this figure.

Response: The discussion was intended to convey the elevated SO4 concentrations at 4-6 km altitudes measured by the C-130 aircraft and previously attributed to long-range transport from Asia. The modeled transport patterns in Figure 6b also suggest the influence of long-range transport. We agree that the model does not do a good job at capturing the space-time variability indicated in the measurements; the coarse (108 km) grid resolution is a likely contributor to such discrepancies. To address the reviewer’s concern, in the revised manuscript (Page 13; lines 9-12) we have reworded the discussion as follows: “As illustrated in Figure 6c, SO4 levels >1 μg m\(^{-3}\) were often measured in the free troposphere. Both the observations and model show these enhanced SO4 levels at altitudes of 4-6 km, which in conjunction with the large scale simulated SO4 distributions in Figure 6b suggest that CMAQ captures the SO4 enhancements in the free troposphere associated with this episodic event. Some discrepancies in space-time matched model and observed concentrations are also apparent in Figure 6c, which likely result from the relatively coarse (108 km) horizontal grid resolution employed in the model calculations”.

Comment: It is admirable that the authors have implemented the RACM2 chemical mechanism into the hemispheric version of CMAQ, which is likely more suitable than CB05 for larger domain applications. However, the authors should give thought to taking the next step and implementing a mechanism that is even more applicable for domains containing regions remote from major sources. Both CB05 and RACM2 were designed for regional-scale applications where NOx concentrations are relatively large compared to values found in the remote troposphere.

Response: We agree with the reviewer that representation of chemistry of the remote troposphere is an important area of future research. Further evaluation of the current mechanisms options in CMAQ for the remote troposphere is an area of current and future research. In the manuscript discussions we acknowledge the need to represent the chemistry of longer-lived species (e.g., acetone) that are important for chemistry of the upper troposphere (page 21; lines 21-25) and also additional analyses of NOx partitioning and HOx predictions from the current and any additional chemical mechanism (Page 16; lines 3-4).
Response to Comments by Anonymous Referee #2

General Comments: This paper provides an overview of the development and some initial applications of a major extension to the off-line Community Multiscale Air Quality (CMAQ) regional chemistry-transport model. Some limitations of regional air quality models are first described and the case is made for the use of hemispheric (or global) air quality models to better address some important research and policy questions. The paper then describes a number of model enhancements that were required to extend CMAQ from its traditional regional-scale configuration to the hemispheric scale, followed by a survey of a number of different evaluations and applications using this new model version.

This is a well-written paper that describes hemispheric CMAQ, an important enhancement of a widely used regional air quality model to enable it to be applied for larger spatial scales and longer time scales. The process enhancements that were required to achieve this design goal should be of general interest. A diverse set of six different applications of the new hemispheric CMAQ are then presented. Several of these applications have been presented elsewhere while others are presented for the first time. In the former case, however, additional perspective and discussion are provided.

I recommend acceptance of this manuscript with minor revisions. I have made a number of specific comments and suggestions below related to clarity and completeness that I would ask the authors to consider. I have also included a number of editorial comments and corrections that I hope will improve the final version.

Response: We thank the reviewer for the overall positive assessment of our manuscript. We thank the reviewer for the thorough review and suggestions, the incorporation of which has led to an improved manuscript. Detailed below is our response to the specific reviewer comments and the changes incorporated in the revised manuscript.

Comment: In Section 2 there were a few places where I asked myself "but what about ...?". In order to provide a more complete description of hemispheric CMAQ, I would suggest that some text could be added to address the following points.

Response: Detailed below is our response to the specific points raised by the reviewer and the changes incorporated in the revised manuscript.

Comment: (Section 2.1) There are many regional chemistry-transport models and there are many global chemistry-transport models, but I am not aware of any other hemispheric chemistry-transport models. Could you add some text to explain the rationale for choosing a hemispheric rather than a global extension, and are there any other models that you are aware of that have also taken your hemispheric approach?

Response: CMAQ’s governing equations are cast in a generalized coordinate form which allows the system to accommodate commonly used horizontal map projections (i.e., Lambert conformal, Mercator, and polar stereographic). The polar stereographic projection allows for convenient
representation of a single hemisphere, and thus as already discussed on lines 1-6 of Section 2.1, this formulation and flexibility enables CMAQ to be used on a horizontal domain covering the Northern Hemisphere set on a polar stereographic projection without altering CMAQ or its input/output file structure. This flexibility was the primary motivation to extend CMAQ to hemispheric versus the global scale. However, it should be noted that the hemispheric expansion is only the first step in creating a multi-scale modeling system that spans urban to global scales which is an area of active research and development in our group. To our knowledge, there are a couple of other active efforts on developing and applying hemispheric scale air quality models: (1) the Danish Eulerian Hemispheric Model (DEHM) (Brandt et al., 2012; https://doi.org/10.1016/j.atmosenv.2012.01.011) and (2) a hemispheric version of the CHIMERE model (https://doi.org/10.5194/gmd-10-2397-2017).

To address the reviewer’s point, we have modified the discussion in Section 2.1 as follows: “CMAQ’s governing three-dimensional equations for species mass conservation and moment dynamics (number, surface area, and volume) describing modes of particulate size distribution are cast in generalized coordinates (cf., Mathur et al., 2005; Byun and Schere, 2006). This formulation allows CMAQ to accommodate horizontal map projections and vertical coordinates from various meteorological models. This flexibility enables CMAQ to be used on a horizontal domain covering the Northern Hemisphere set on a polar stereographic projection (Figure 2a) without altering CMAQ or its input/output file structure. Polar stereographic projections are also used in the Danish Eulerian Hemispheric Model (Brandt et al., 2012) and a hemispheric version of the CHIMERE model (Mailler et al., 2017).”

Comment: (Section 2.1) Limited-area models require lateral boundary conditions. Although you have greatly expanded your model domain by choosing a hemispheric domain, hemispheric CMAQ is still a limited-area model. However, there is no discussion in the text of the LBC that you have used for hemispheric CMAQ. Also, Figure 1 showed CMAQ sensitivity to ozone lateral boundary conditions for a regional configuration over the continental U.S.: has a similar sensitivity test been performed for hemispheric CMAQ to show its sensitivity to LBC specification?

Response: The reviewer is correct that lateral boundary conditions also need to be specified for the discrete lateral boundaries of the hemispheric domain. To address the reviewer’s concern, in the revised manuscript, we have included the following discussion at the end of Section 2.2 (page 6, line 1-9):

“Similar to regional applications, chemical boundary conditions also need to be specified along the discrete lateral boundaries of the hemispheric domain. In current applications, these are set to same values as in the clean IC case discussed above. Note that the boundaries of the hemispheric domain (shown in Figure 2) are in the area encircling the Earth near the equator. Because of the presence of the intertropical convergence zone (ITCZ) in this region, the mixing of air masses originating in the Northern and Southern Hemispheres occurs relatively slowly, with exchange times of typically about 1 year (e.g., Jacob et al., 1987). Since the atmospheric lifetimes of most modelled species are significantly shorter, any impacts of chemical lateral boundary condition
specification are typically confined to the lower latitudes and do not propagate into the domain. Additional model sensitivity tests should however be conducted in the future to quantify any likely seasonal influence of LBC specification on model predictions in lower latitude regions of the Northern Hemisphere”.

**Comment:** (Section 2.1) *What is the vertical coordinate used by WRF and hemispheric CMAQ?*

**Response:** Both WRF and CMAQ use the $\sigma$-P vertical coordinate. To clarify this point, we have modified the description in section 2.1 to now read “Current WRF and CMAQ hemispheric applications have utilized a horizontal discretization of a 187x187 grid configuration with a grid spacing of 108km and a $\sigma$-P vertical coordinate system”

**Comment:** (Section 2.2) *You recommend here that hemispheric CMAQ should be started from clean tropospheric conditions, and you mention initializing O$_3$ at 30 ppb throughout the model column for the clean IC case. Do you similarly recommend that other CMAQ model species should be set to a uniform clean value in both the horizontal and vertical? Also, can you mention at least some of the clean IC values used for other CMAQ species (e.g., NO, NO$_2$, CO, NH$_3$, SO$_2$, ...)? And given its complex suite of sources, have you examined how CO responds to a 9- or 12-month spin-up from clean conditions?*

**Response:** As suggested in the reviewer’s comments, the ideal initialization period is species dependent. In our experiments the initial conditions for other chemical species were based on the profiles presented in Byun and Ching (1999), which nominally represent clean tropospheric conditions. We focused our initialization experiments on O$_3$ because of the extensive available measurements on its vertical variations in the troposphere which was used to guide the analysis. To address the reviewer’s suggestion on mentioning the initial conditions for other species we point the reader to the profiles in Byun and Ching (1999) by including the following additional sentence on Page 5 (line 20-21): “Initial conditions for all other chemical species were based on clean tropospheric conditions prescribed in Byun and Ching (1999)”.

**Comment:** (Section 2.3.1) *Is the ARCTAS emissions inventory associated with a nominal base year?*

**Response:** As stated in the manuscript, the ARCTAS inventory was compiled to support pre-mission planning for the ARCTAS study. As stated on the website for the data, the inventory was compiled using the most recent data available to represent emissions at that time ([http://bio.cger.uiowa.edu/arctas/emission.html](http://bio.cger.uiowa.edu/arctas/emission.html)). Thus the data are not associated with a specific base year.

**Comment:** (Section 2.3.1) *Are the GEIA biogenic VOC and lightning NOx emissions climatological or year-specific?*

**Response:** The GEIA biogenic and lightning NO$_x$ emissions represent climatological averages.
The temporalization of the monthly mean biogenic emissions and the annual lightning NOx emissions to the hourly scale used by CMAQ is described in Xing et al. (2015a) as already mentioned in the manuscript discussion. To address the reviewer’s comment, we modified the discussion in Section 2.3.1 as follows: “In applications to date, biogenic VOC (Guenther et al., 1995) and lightning NOx (Price et al., 1997) emissions were obtained from GEIA (Global Emission Inventory Activity; http://www.geiacenter.org). The monthly biogenic VOC emissions were further temporalized to hourly resolution for each simulation day. Monthly lightning NOx emissions were distributed evenly to each hour of each simulation day. Xing et al. (2015a) further describe the processing of global emission inventories for CMAQ, including temporalization of the annual estimates to hourly model inputs, vertical distributions of anthropogenic and lightning emissions, and speciation of PM2.5 and NMVOC emissions to model primary aerosol constituents and gas-phase species”

Comment: (Section 2.3.1) Natural emissions take on increased importance for a global or hemispheric chemistry-transport model. Were sea-salt emissions, biomass-burning emissions, soil NO emissions, or volcanic SO2 emissions considered by hemispheric CMAQ?

Response: We agree with the reviewer that representation of the impact of natural emissions on tropospheric composition is important not only for the hemispheric and global scales but also for regions with rapidly declining anthropogenic emissions. Year-specific large-scale biomass burning emissions are included in the EDGAR inventory and have been used in our simulations – this is now explicitly mentioned in Section 2.3.1. We also include parameterizations for natural wind-blown dust emissions (detailed in Section 2.3.2) as well as sea-salt emissions based on Kelly et al. (2010) (already mentioned in Section 2.3.3). The current model simulations however did not include soil NO emissions or volcanic SO2 emissions – this is now clarified in Section 2.3.1 by including the following sentence: “Emissions of NO from soil or SO2 from volcanos are not considered in the applications presented here”.

Comment: (Section 2.3.3) What about marine DMS emissions?

Response: DMS emissions and chemistry are not yet considered in the hemispheric CMAQ model. We however are currently testing implementations of: (1) a DMS emission parameterization scheme, and (2) modifications to the chemical mechanisms to include DMS oxidation pathways to represent their effects on atmospheric SO42- distributions.

Comment: (Section 2.4.3) It could be mentioned here that RACM2 does not include all (any?) of the chlorine, bromine, and iodine species discussed in Section 2.4.2 – this is relevant to Section 3.4.

Response: Yes, the base RACM2 mechanism (Goliff et al. 2013) does not include halogen chemistry. We however have included a version of the mechanism augmented with halogen chemistry and discuss those results in section 3.4. We feel that the sentence (Page 16 lines 7-8) already conveys this distinction: “Also shown are predictions with a model configuration in which
the RACM2 mechanism was augmented with the halogen chemistry described in section 2.4.2”

Comment: (Section 2.5) In essence this section deals with chemical upper boundary conditions. How are other CMAQ model species treated at the top of the model?

Response: Section 2.5 only discusses the impact of stratosphere-troposphere exchange on O3 distributions. We do not modify other species concentrations in the upper layers with this parameterization. The concentrations of all other species are dictated by the modeled transport and chemistry processes. To be specific to the discussion in this section and to avoid any confusion we have modified the title of section 2.5 to “Representing Impacts of Stratosphere-Troposphere Exchange on O3 Distributions”.

Comment: It is not made clear in Section 3.1 (p. 15, l. 12) exactly which forms of the two gas-phase chemistry mechanisms were used. Although Sarwar et al. (2013) is referenced (l. 11), I am not sure that the exact mechanism versions used for that paper were also used in this study. For example, I think but I am not sure that a modified version of CB05TU was used in this study that included the modifications described in Sections 2.4.1 and 2.4.2. I am not sure whether or not the version of RACM2 used in this study included the modifications described in Section 2.4.1. The analysis presented in this section does state that two different versions of RACM2 were used, one with the modifications described in Section 2.4.2 and one without those changes, but some additional clarification would be very helpful.

Response: To address the reviewer’s comment, in Section 3.1 we have now clarified the mechanisms used by explicitly stating the following: “Note that the simulation “PV, NoHalogen” employed the CB05TU mechanism, while the other two simulations employed a version of the CB05TU mechanism augmented with the halogen chemistry discussed in Section 2.4.2”.

We believe the reviewer’s query on the RACM2 mechanism is referring to the discussion related to Figure 12 (b-d) in Section 3.4. These figures show results from 3 simulations denoted CB05TU, RACM2, and RACM2_HAL. As discussed in the text and noted in the figure caption RACM2_HAL represents the RACM2 mechanism augmented with halogen chemistry. We see the reviewer’s point that it may not be readily apparent whether the simulation denoted CB05TU employed halogen chemistry or not. To address this concern we now explicitly point this out in Section 3.4 as: “Note that the simulations denoted CB05TU and RACM2 did not include representation of halogen chemistry”.

Comment: In preparing figure panels 13d, 13e, and 13f, was the SCIAMACHY averaging kernel applied to the CMAQ NO2 fields?

Response: We did not use the averaging kernel in comparisons with satellite column data because it was not available for the entire 2000-2010 period. Thus the model NO2 column was estimated without any averaging kernel and was just a straight integration of model NO2 through the model column from surface to ~50mb. To address the reviewer’s comment, we clarify this in the
discussion in Section 3.5 as: “Note that the model NO₂ column is estimated by integrating the predicted NO₂ fields through the model column from the surface to ~50hPa and did not utilize an averaging kernel”.

**Comment:** In Section 3.5, how were the station trends that are shown in Figure 15 calculated?

**Response:** The trends are estimated as a linear regression of the June-July-August (JJA or summer) average values of a concentration metric (for O₃ JJA average of the daily maximum 8-hr average value and for SO₄²⁻, the JJA average of weekly average data; as also indicated in the Figure caption) at each site for the 21 years – 1990-2010. We modified the discussion in the last paragraph of section 3.5 as: “Figure 15 presents comparisons of model and observed trends in summer average daily maximum 8-hour average O₃ and summer average weekly-average SO₄²⁻ at each CASTNET monitor site. Trends are estimated as the slope of the linear regression of these concentration metrics for the 21-year period”.

**Comment:** In Section 3.6 how were the CMAQ SWR fields calculated in conjunction with predicted cloud fields?

**Response:** Clear and all-sky shortwave radiation fields can be obtained from the RRTMG radiation scheme used in the WRF-CMAQ modeling system. The aerosol direct feedbacks modify these fields in the 2-way coupled WRF-CMAQ configuration and the difference between these is being analyzed.

**Comment:** In Section 3.6 I am not sure that the discussion of Figure 16 is completely correct, in particular the following sentence: “The change in the SWR and AOD for each summer month in the 2001-2010 period was estimated relative to the corresponding year-2000 value, and the relationship between these changes is examined in Figure 16 for both model simulations with and without direct aerosol feedback effects.” If this were true, then for East China, which has been experiencing dimming over the past decade, wouldn’t most of the observed SWR changes be negative, whereas for Europe and the eastern U.S., which have been experiencing brightening over the past decade, wouldn’t most of the observed SWR changes be positive? Instead, the difference patterns in Figure 16 seem to be more consistent with the subtraction of the 11-year monthly means; that is, they are centered.

**Response:** We are grateful to the reviewer for noting this discrepancy between text and what is shown in the Figure. We re-examined our calculations and verified that for this figure we indeed did not estimate the change relative to 2000, as erroneously noted in the original manuscript discussion. The reviewer is correct that the characteristics of the data shown in Figure 16 suggest that it is an anomaly (i.e., the mean value subtracted out). We have updated Figure 16 to indicate that the relationship examined is between the AOD anomaly and SWR anomaly and also clarified this in Section 3.6 (on page 19) as: “The relationship between these deseasonalized values (or anomaly) of SWR and AOD for each summer month in the 2001-2010 period is examined in Figure 16 for both model simulations with and without direct aerosol feedback effects. Also shown
in Figure 16 is the corresponding observed relationship between similarly estimated AOD anomaly and SWR anomaly derived from retrievals from the MODIS and the Clouds and the Earth's Radiant Energy System (CERES; Wielicki et al., 1998) instruments, respectively.”

The caption of Figure 16 has also been updated to: “Relationship between regional and monthly average (only summer months) changes in aerosol optical depth and changes in surface clear-sky shortwave radiation during the 2001-2010 period for (a) East China, (b) Europe, and (c) East U.S. Observed values are shown in grey, CMAQ calculations with direct aerosol radiative feedbacks in red, and CMAQ calculations without aerosol radiative feedbacks in blue. Also, indicated are the slope and correlation coefficient (R) for the individual linear regressions. For each data set (model or observed) there are 33 values, corresponding to each summer month over the 11-year (2001-2010) period. The anomaly is estimated by subtracting the corresponding 11-year average for that month.”

Technical corrections: The manuscript reads very well but it would still benefit from a careful copyediting to add commas, hyphens, and definite articles in some places but remove them in other places (e.g., change “space and time varying” to “space-and time-varying”).

Response: We thank the reviewer for the thorough review and the editorial suggestions, majority of which have been incorporated as detailed below.

Some acronyms are used but never defined: ECMWF, GFS, ADP, PBL, NOAA, AQS
Response: The acronyms have now been defined.

p. 1, l. 13 WRF is also included as a keyword so it could be defined here on the same page.
We have defined WRF in the abstract as suggested.

p. 2, l. 4 Perhaps “… implementation of the U.S. National Ambient Air Quality Standards …”
Limited area models have and are helping with the design and implementation of air quality standards across the world. We have thus left the sentence unaltered.

p. 2, l. 11 Perhaps “… postulated that in limited-area chemistry-transport models, …”
We have modified the sentence following the reviewer’s suggestion.

p. 2, l. 14 Perhaps “… derived from the global Integrated Forecasting System of…”
We have modified the sentence following the reviewer’s suggestion.

p. 2, l. 16 hPa is the equivalent SI unit to mb.
We have replaced mb with hPa throughout the manuscript.

p. 2, l. 26 “higher values in the high elevation regions”: concentration values or variability values?
We have modified the sentence to clarify that they are higher normalized concentration values.

p. 2, l. 26 Perhaps “Additionally, higher contributions from background levels are estimated”
Since these boundary tracers represent “model background” values, they may not necessarily imply higher fractional contributions to the net concentrations. We thus have left the sentence as is.

p. 2, l. 30  “Expectedly” makes me think of “bigly” – perhaps “As expected” would be a better choice.
We use both “As expected” and “expectedly” interchangeably through the manuscript and thus have left the sentence as is.

p. 2, l. 34  Perhaps “... that other pollutants with atmospheric lifetimes greater than a few days”
As written the sentence implies “any pollutant” with atmospheric lifetimes greater than a few days, which we feel is more appropriate. Thus we have left the sentence as is.

p. 3, l. 17  Perhaps “Section 2 provides an overview of the ...”
We feel the sentence as written is ok, and have left it as is.

p. 3, l. 21  Perhaps “Lastly, Section 4 summarizes the current model status ...”
We have added “Lastly” to the sentence but feel “model state” is more appropriate than “model status”.

p. 4, l. 28  Could give a reference for the NCEP/NCAR Reanalysis data set?
We now include the following reference for the NCEP/NCAR Reanalysis data set:

p. 4, l. 31  Check weblink (extraneous blank after “V3.1/”?)
The “extraneous blank” was put in just top space the sentence wording within the formatting – it should be correct in the final typeset.
p. 5, l. 8  Perhaps “... closer attention to model chemical initialization ...”
We have modified the sentence following the reviewer’s suggestion.

p. 5, l. 13  Perhaps “... based on the model emissions, physics, and chemistry ...”
We have modified the sentence following the reviewer’s suggestion.

p. 5, l. 24  Perhaps “... Clean IC and Profile IC cases by August, nine months after the start of the simulation, suggests the diminishing impact of initialization ...”
We have modified the sentence following the reviewer’s suggestion.

p. 8, l. 33  Should it be Xie et al. (2013)?
Thank you for catching the typo – we have corrected the year of the publication.

p. 9, l. 23  The Goliff et al. (2013) reference is missing.
Thank you for pointing that out. We have added the Goliff et al. (2013) reference.

p. 10, l. 5  “... in the modelled upper troposphere/lower stratosphere”
We have modified the sentence following the reviewer’s suggestion.

p. 10, l. 29  Perhaps “These hemispheric O3 fields can then be used ...”.
We have modified the sentence following the reviewer’s suggestion.

p. 11, l. 20  There are a number of references to the "Pacific": perhaps some could refer to the "Pacific Ocean" instead.
We have modified some of the instances as suggested.

p. 11, l. 24  “DC-8”
We have modified “DC8” to “DC-8” throughout the manuscript following the reviewer’s suggestion.

p. 11, l. 25  “... were based ... and sampled ...”
We feel the sentence as written is fine and thus have not changed it.
p. 11, l. 27 There are some unexpected capitalizations, such as Pollution, Spring, Winter, and Continental. We have corrected the capitalization of “pollution” in this sentence.

p. 12, l. 33 “Particle Into Liquid Sampler” is the more common usage. We have modified the sentence following the reviewer’s suggestion.

p. 13, l. 23 Perhaps “… air pollutants, dating back almost a century” We have modified the sentence following the reviewer’s suggestion.

p. 15, l. 15 Perhaps “… are illustrated in Figure 12a for August” We have modified the sentence following the reviewer’s suggestion.

p. 17, l. 1 Perhaps “… in the Savanna region of Africa both in …” We have modified the sentence following the reviewer’s suggestion.

p. 17, l. 2 “SCHIAMACHY” (spelling) Thank you for pointing out the typo – we have corrected it.

p. 17, l. 21-22 The Figure 15 caption and labels state that the analysis shown is for the summer months only and not all year as stated in this sentence. Thank you for noting this discrepancy. The figure caption is correct but the text erroneously suggested annual average – we have rewritten the sentence to correct this inconsistency.

p. 17, l. 23 Perhaps “…are results from an additional 21-year simulation with CMAQ” We have modified the sentence following the reviewer’s suggestion.

p. 17, l. 28 The “Conclusions” section states a range for the underestimation (p. 20, l. 12) that should be mentioned here. We now mention the quantitative range of underestimation in the discussion, as suggested by the reviewer.

p. 18, l. 12-21 As a lead in to the next paragraph, it could be mentioned here that the aerosol
optics calculations in WRF-CMAQ include the calculation of AOD.

We feel it should be apparent to the ACP readers that since aerosol optical properties are calculated in WRF-CMAQ, the AOD can be easily estimated and available. We have thus left the discussion unaltered.

p. 18, l. 29  Perhaps “... using regional monthly averages ...” and “Eastern U.S.”

We have modified the sentence following the reviewer’s suggestion.

p. 18, l. 33  Perhaps “... but AOD at noon (local-time) for model values to be consistent”

We have modified the sentence following the reviewer’s suggestion.

p. 20, l. 11  Perhaps “... at U.S. CASTNET monitors, ...”

We have modified the sentence following the reviewer’s suggestion.

p. 21, l. 33  IONS and WOUDC are not networks.

We have deleted “networks” following the reviewer’s suggestion.

p. 29  Figure 1 caption: Perhaps “Impact of ozone lateral boundary conditions (LBC)”

Since our calculations did not include any chemical decay mimicking that of O₃ we do not think we should suggest that the calculation necessarily represents impacts of O₃ LBCs on simulated O₃ concentrations.

p. 31  Figure 3: Would it be useful to give associated states or longitudes for these four measurement sites?

We have modified the caption of Figure 3 to also provide the state names for the four sites.

p. 32  Figure 4 caption: Would it be useful to indicate the elapsed simulation time for these two panels; that is, four months and nine months after the start of the simulation?

Following the reviewer’s suggestion, we now indicate the elapsed simulation time for the two panels.

p. 33  Figure 5 caption: Does not mention the aircraft altitude time series in the first four panels.

The figure caption now also indicates that the aircraft altitude is shown in (a-d).
p. 43 Figure 14 caption: Perhaps “... changes in regional- and monthly-average modeled ...”

We have modified the caption following the reviewer’s suggestion.
Response to Short Comment by Tim Canty

Comment: This is a very interesting paper and an admirable endeavor to extend CMAQ to hemispheric scales. I was especially interested in the handling of alkyl nitrate lifetime. Our group has done similar analysis of alkyl nitrates with comparisons to satellite and aircraft observations in using the CMAQ with CB05 chemistry and CAMx with CB6r2 chemistry (Canty et al., 2015, Goldberg et al., 2016).

What is the lifetime of NTR in this new model framework? Our modification of NTR so that its lifetime is much shorter (~1 day), expected if hydroxynitrates are the most abundant species, has led to a better representation of NTR in CB05 when compared to aircraft observations taken during DISCOVER-AQ. Across our model domain, tropospheric column NO2 from CMAQ was in better agreement with satellite observations when the NTR lifetime was decreased. The faster decomposition of NTR also led to an increase in modeled surface ozone. Based on this, the decrease in ozone reported in Fig 4 of this manuscript was a surprise. Is this decrease due to transport or to increased deposition processes? The improved speciation of NTR in the CB6r2 chemical mechanism led to a shorter lifetime of NTR, in the model, without any needed changes to the NTR chemistry. Great job with this analysis!

Response: Thank you for the positive comments on the manuscript and the analysis. We are aware of the analysis reported in Canty et al. (2015) and read with interest the work reported in Goldberg et al. (2016). As discussed in our manuscript and previously (Schwede and Luecken, 2014; Canty et al., 2015, Appel et al., 2017), a thorough description of the NTR family of gases is needed to accurately represent their atmospheric lifetimes and rate of NOx recycling that eventually will influence O3 distributions on local to hemispheric scales. As indicated in the discussion, both physical and chemical sinks of NTR influence its atmospheric lifetimes, but an important distinction is that chemical sinks recycle back NOx (on varying time scales) while the physical sinks (wet and dry deposition) do not. Clearly, the relative roles of these two NTR removal processes depend on the form of the organic nitrate species. In our initial implementation, we followed Xie et al. (2013) and modified the rate constant for the NTR+OH reaction to that for isoprene nitrates, since on the hemispheric scales organic nitrates formed from isoprene are the largest contributor to the simulated tropospheric NTR burden. More importantly, the dry deposition velocity for NTR was mapped to that for HNO3 and the Henry’s law constant for NTR was also mapped to that of HNO3, thereby enhancing wet scavenging of NTR. Thus, for the calculations based on these assumptions, at least at the surface one can expect the lifetime of NTR to be comparable to HNO3 and on the same order or shorter than what was invoked by increasing its photolysis rate by a factor of 10 in your analysis. However, unlike your approach which recycles the NOx back more rapidly locally, an important distinction is that the enhancement of the physical sinks removes the reactive nitrogen from the atmosphere. Thus on the hemispheric scales, less NOx is recycled (due to lower amounts of NTR) and so there is lower ozone shown in Figure 4.
Recently, in CMAQ we have replaced the single alkyl nitrate species (NTR) in CB05TU with seven species to better capture the range of chemical reactivity and Henry’s law constants (and thus the physical sinks) – see Appel et al. (2017, GMD). We believe such an approach helps better regulate the organic nitrate budget (also evaluated through comparisons with wet deposition measurements), increases the amount of NOx recycled locally (resulting in local increases in O3), but reduces the NTR burden on the larger hemispheric scale and better modulates hemispheric background O3. The impacts of the rate of NOx recycling from organic nitrates is also seen in the RACM2 results presented in Figure 12, which show higher O3 in polluted regions, but lower values are seen in the remote areas relative to the CB05TU mechanism; these differences in part arise from higher rates of NOx recycling from organic nitrates in RACM2 relative to CB05TU. The recently released version of CMAQ (v5.2) includes the CB6 mechanism with updated organic nitrate chemistry; application and evaluation of the CB6 mechanism over hemispheric scales is currently underway.
Extending the Community Multiscale Air Quality (CMAQ) Modeling System to Hemispheric Scales: Overview of Process Considerations and Initial Applications

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Abstract. The Community Multiscale Air Quality (CMAQ) modeling system is extended to simulate ozone, particulate matter, and related precursor distributions throughout the Northern Hemisphere. Modeled processes were examined and enhanced to suitably represent the extended space and time scales for such applications. Hemispheric scale simulations with CMAQ and the Weather Research and Forecasting (WRF) model are performed for multiple years. Model capabilities for a range of applications including episodic long-range pollutant transport, long-term trends in air pollution across the Northern Hemisphere, and air pollution-climate interactions are evaluated through detailed comparison with available surface, aloft, and remotely sensed observations. The expansion of CMAQ to simulate the hemispheric scales provides a framework to examine interactions between atmospheric processes occurring at various spatial and temporal scales with physical, chemical, and dynamical consistency.

Keywords: Hemispheric CMAQ, WRF, long-range transport, background pollution

1 Introduction

Comprehensive atmospheric chemistry-transport models must constantly evolve to address the increasing complexity arising from emerging applications that treat multi-pollutant interactions at urban to hemispheric spatial scales and hourly to annual temporal scales. To assist with the design of emission control strategies that yield compliance with more stringent air quality standards, such models must accurately simulate ambient pollutant levels across the entire spectrum ranging from background to extreme concentrations. The adverse impacts of airborne pollutants are not confined to a region or even a continent (NRC, 2009). Both observational (e.g., Andrea et al., 1988; Fishman et al., 1991; Jaffe et al., 1999; Zhang et al., 2008; Uno et al., 2009) and modeling studies (e.g., Jacob et al., 1999; Fiore et al., 2009; HTAP, 2010) have demonstrated that pollutants near the Earth’s surface can be convectively lofted to higher altitudes where strong winds can efficiently transport them from one continent to another, thereby impacting air quality on intercontinental to global scales.
As air quality standards are tightened, the need to quantify the contributions of long-range transport on local pollution becomes increasingly important. Limited-area models such as the Community Multiscale Air Quality (CMAQ) (Byun and Schere, 2006; Foley et al., 2010; Appel et al., 2017) have played a central role in guiding the development and implementation of the National Ambient Air Quality Standards (NAAQS). These models are now being routinely applied to examine variability in surface-level air pollutants across the continental U.S. over annual cycles. Since transport is efficient in the free-troposphere and since simulations over continental scales and annual cycles provide sufficient opportunity for “atmospheric turn-over”, i.e., exchange between the free-troposphere and the boundary-layer, it can be argued that accurate simulation of the variability in free-tropospheric pollutant concentrations is important for the model’s ability to capture the variability in surface-level concentrations, especially at moderate-to-low concentrations. Based on typical advective time scales, it can further be postulated that in limited-area chemistry-transport models, this free-tropospheric variability in simulated concentrations is largely dictated by the specification of lateral boundary conditions (LBC). This is exemplified in Figure 1 which illustrates the influence of LBC specification on simulated surface-level concentrations across a typical regional modeling domain covering the contiguous U.S.; the space and time varying LBCs themselves were derived from the global Integrated Forecasting System of European Center for Medium-Range Weather Forecasts (ECMWF) (Flemming et al., 2015). In these calculations, three tracer species were added to CMAQ to track the ozone (O3) LBC specified for three vertical zones: (i) surface to 750 hPa, nominally representing the atmospheric boundary layer (BL), (ii) 750-250 hPa representing the free-troposphere (FT), and (iii) 250-50 hPa (the model top) representing possible stratospheric influences (cf. Mathur et al., 2008). These tracer species were subject to transport processes associated with 3-D advection, turbulent mixing in the vertical and horizontal diffusion, and cloud transport on resolved and sub-grid scales. The tracer species were deposited at the surface using space- and time-varying deposition velocity estimates for O3, and they were also subjected to wet scavenging and rainout processes that mimic modelled O3. If the tracer background is defined as the amount of the tracer imported into the regional domain, then the sum of the three simulated tracers can be viewed as the modelled background for this species. Since we did not include chemical sinks for the tracers and since the intent here is to assess the relative influence of LBC, we examine the distribution of the normalized concentrations (normalized by the domain maximum value across all seasons) in Figures 1a-d. Significant spatial and seasonal variability is seen in the estimated tracer background levels, with higher normalized concentration values in the high elevation regions of the inter-mountain west. Additionally, higher background levels are estimated in the warmer seasons, with highest levels during spring. More importantly, the free tropospheric concentrations dominate the surface background levels (Figures 1e-h). During summer, across most of the continental U.S. more than half (and up to 90%) of the tracer background originated in the FT. Though significant seasonal variability is noted in the FT fractional contributions to surface-level background concentrations, the contributions are still substantial during other seasons and expectedly the highest contributions are seen in the high elevation regions of the Western U.S. across all seasons. These results clearly illustrate the importance of accurately characterizing the long-range transport that occurs in the free troposphere and its influence on surface background pollution levels via subsidence and entrainment into the boundary layer. It can be expected that pollutants with atmospheric lifetimes
greater than a few days would exhibit similar characteristics, thereby highlighting the need to accurately characterize long-range transport influences on regional model simulations spanning seasonal to annual time-scales.

One approach to capture the effects of long-range transport in regional models is through deriving space- and time-varying LBC from global chemistry transport models. However, efforts linking regional and global scale models have met with mixed success because biases in the global model can propagate and influence regional calculations and often confound interpretation of model results (e.g., Tang et al., 2008; Schere et al., 2012). Additionally, inconsistencies in process representations, species mapping, and grid structures could also introduce errors in the model linkage if not examined and handled carefully. A modeling framework is thus needed wherein interactions between processes occurring at various spatial and temporal scales can be consistently examined. Expanding comprehensive regional models to the hemispheric scale enables a consistent representation of atmospheric processes across spatial and temporal scales. Motivated by this need, the applicability of the CMAQ modeling system has been extended to hemispheric scales through systematic investigation of key model processes and attributes influencing simulated distributions of O₃, fine particulate matter (PM₂.₅), and precursor species. The hemispheric modeling system also facilitates examination of linked air pollution-climate across a region in context of the changing global atmosphere.

Section 2 overviews the key CMAQ model structural attributes and process representations that were refined to fully simulate the Northern Hemisphere. Section 3 summarizes a variety of applications with the hemispheric CMAQ configuration, highlights the model performance relative to a variety of observational data sets, and identifies aspects that would benefit from further model development. A variety of surface, aloft, and remotely sensed observations used to guide and evaluate the model changes are presented in sections 2 and 3. Lastly, section 4 summarizes the current model state and discusses future development and applications of the hemispheric CMAQ.

2 Model Setup and Process Enhancements

Atmospheric chemistry and transport of trace species occur across the continuum of spatial and temporal scales. For instance, transport across intercontinental to hemispheric scales occurs over timescales ranging from days to months, which influences the distribution of trace species with lifetimes within this range. Transport on these scales can also influence shorter lived radical budgets through chemical reactions involving intermediate-lived species, especially reservoir species such as organic nitrates. Thus, the expansion of CMAQ to hemispheric scales required re-examination of process representations and grid structures so that interactions amongst various processes occurring over the disparate scales is adequately captured. The key changes to CMAQ that were considered in this effort are summarized below.
2.1 Domain and Grid Configuration

CMAQ’s governing three-dimensional equations for species mass conservation and moment dynamics (number, surface area, and volume) describing modes of particulate size distribution are cast in generalized coordinates (cf., Mathur et al., 2005; Byun and Schere, 2006). This formulation allows CMAQ to accommodate horizontal map projections and vertical coordinates from various meteorological models. This flexibility enables CMAQ to be used on a horizontal domain covering the Northern Hemisphere set on a polar stereographic projection (Figure 2a) without altering CMAQ or its input/output file structure. Polar stereographic projections are also used in the Danish Eulerian Hemispheric Model (Brandt et al., 2012) and a hemispheric version of the CHIMERE model (Mailler et al., 2017). Current WRF and CMAQ hemispheric applications have utilized a horizontal discretization of a 187x187 grid configuration with a grid spacing of 108 km and a σ-P vertical coordinate system.

Current regional modeling applications with CMAQ typically utilize 35 layers of variable thickness to resolve the model vertical extent between the surface and 50 hPa. Longer-term calculations over the Northern Hemisphere must be able to capture potential impacts of stratosphere-troposphere exchange (STE) as well as that between the free troposphere and the BL. At altitudes above 10 km (Figure 2b), the 35-layer configuration has relatively coarse resolution with layer thickness >1.5 km and the top-most layer is nearly 4 km deep. To improve the representation of three-dimensional transport processes on modelled vertical profiles, the vertical resolution employed in hemispheric CMAQ calculations is increased. The revised layer structure uses 44 layers, with significantly finer resolution above the boundary layer (Figure 2b) to better represent long-range transport in the free troposphere, STE processes, and influences from cloud mixing both at the sub-grid- and resolved scale. The impacts of using these alternate layer configurations are illustrated in Figure 3 for a case where emissions only across the U.S. were zeroed out to isolate the impacts of model vertical resolution on representing the downward transport of pollutants in the region. Both the 35-layer and 44-layer model simulations were initialized with the same conditions in mid-February 2006, utilized a constant potential vorticity scaling to specify O₃ in the model top-layer (discussed further in Section 2.5), and were driven by meteorological information from the Weather Research and Forecasting (WRF) model simulations using the respective layer configurations (discussed in Section 2.2). Figure 3 shows systematically higher simulated O₃ below 10 km in the 35-layer configuration compared to the 44-layer configuration, indicating that the coarser vertical resolution will likely overestimate the downward transport of both long-range transport effects as well as stratospheric influences.

2.2 Coupling of WRF and CMAQ and Initialization

To minimize interpolation error and to avoid introducing mass imbalances, hemispheric simulations with CMAQ inherit the projection and grid structure from the WRF model, which provides the driving meteorological fields. In applications presented in Section 3, meteorological inputs for grid nudging used in WRF over the Northern Hemisphere domain were derived from the NCEP/NCAR Reanalysis data (Kalnay et al., 1996) which has 2.5 degree spatial and 6-hour temporal resolution; other reanalysis products such as Global Forecast System (GFS) can also be used instead. Surface reanalysis based on a fusion of the NCEP/NCAR Reanalysis and NCEP Automated Data Processing (ADP) Operational Global Surface Observations on the
WRF grid using the NCAR distributed objective analysis tool Obsgrid ([http://www2.mmm.ucar.edu/wrf/users/docs/user_guide_V3.1/users_guide_chap7.htm#techniques](http://www2.mmm.ucar.edu/wrf/users/docs/user_guide_V3.1/users_guide_chap7.htm#techniques)) is used for the indirect soil moisture and temperature nudging in the Pleim-Xiu land surface model (Pleim and Gilliam, 2009). The WRF configuration over the Northern Hemisphere also used MODIS land-use classification with 20 categories, RRTMg shortwave and longwave radiation scheme (Iacono et al., 2008), and the ACM2 planetary boundary layer model (Pleim 2007). WRF’s simulation of hourly surface temperature, relative humidity, wind speed and direction was evaluated by Xing et al. (2015a) through comparison with observations from NOAA’s National Center for Environmental Information (NCEI) Integrated Surface Data and no significant bias in the meteorological fields were detected. WRF and CMAQ can be run either in the traditional off-line sequential manner or in the coupled mode with or without aerosol feedback effects (Mathur et al., 2010; Wong et al., 2012).

Expectedly, the application over expanded space and time scales necessitates closer attention to model chemical initialization, especially in the free troposphere wherein typical residence time for most atmospheric pollutants of concern are long enough so that initial conditions can persist. If the FT is poorly represented, model predictions within the boundary layer will be adversely affected. Thus, unlike regional simulations with CMAQ which are initialized with a prescribed vertical profiles for different species or with concentration fields derived from global chemistry-transport models, for hemispheric applications it is recommended that CMAQ be initialized to “clean” tropospheric background values and allowed to build up based on the model emissions, physics and chemistry. The impact of these different initializations are illustrated through comparisons of the cases denoted Profile IC and Clean IC in Figure 4 which compares monthly mean model and observed O3 profiles at Trinidad Head, CA. In the Profile IC case, a vertical O3 profile that monotonically increased from 35 ppb at the surface to 100ppb at model top was used for initialization. The clean IC case initialized O3 at 30 ppb through the model column. In both cases, the hemispheric model simulations were initiated on 1 November 2015. Initial conditions for all other chemical species were based on clean tropospheric conditions prescribed in Byun and Ching (1999). Large overestimations in O3 through the troposphere are noted for the Profile IC simulation and arise from the profile used to initialize O3 in the mid-troposphere. In contrast in the Clean IC case wherein the model was initialized to “clean” tropospheric background values and allowed to build up based on the model physics and chemistry, resulted in much better agreement with the measured profile during spring (Figure 4a); however, by summer, overestimations developed (discussed further in section 2.4.1). Note that the clean IC case also utilized a fixed potential vorticity scaling for O3 at the model top, described further in Section 2.5. Also the similarity in simulated O3 profiles for the Clean IC and Profile IC cases by August, nine months after the start of the simulation, suggests the diminishing impact of initialization. Based on these results, the inherent seasonality in atmospheric transport and chemistry, and practices employed in previous global chemistry-transport model applications (e.g., Fiore et al., 2009), a spin-up of 12 months from clean tropospheric conditions is recommended for new CMAQ applications over the Northern Hemisphere. Additional future studies would be helpful to further constrain this spin-up period recommendation.
Similar to regional applications, chemical boundary conditions also need to be specified along the discrete lateral boundaries of the hemispheric domain. In current applications, these are set to same values as in the clean IC case discussed above. Note that the boundaries of the hemispheric domain (shown in Figure 2) are in the area encircling the Earth near the equator. Because of the presence of the intertropical convergence zone (ITCZ) in this region, the mixing of air masses originating in the Northern and Southern Hemispheres occurs relatively slowly, with exchange times of typically about 1 year (e.g., Jacob et al., 1987). Since the atmospheric lifetimes of most modelled species are significantly shorter, any impacts of chemical lateral boundary condition specification are typically confined to the lower latitudes and do not propagate into the domain. Additional model sensitivity tests should however be conducted in the future to quantify any likely seasonal influence of LBC specification on model predictions in lower latitude regions of the Northern Hemisphere.

### 2.3 Emissions

Specifying emissions across the Northern Hemisphere is challenging because the distributions and compositions of emissions across the globe are rapidly changing and because emissions are poorly quantified in many regions. In addition, simulations of CMAQ at broader spatial scales are influenced by emissions from marine environments (which are less prominent in regional/continental applications), and intercontinental transport of other sources (e.g., wind-blown dust). Changes to emissions used by CMAQ for the Northern Hemisphere application are described below.

#### 2.3.1 Global Emission Inventories

Two primary sources of global emission estimates have been used in hemispheric CMAQ applications to date. The first is based on global emission inventory compiled by Argonne National Laboratory in support of the ARCTAS pre-mission planning and includes estimates for anthropogenic, international shipping, and biomass burning ([http://bio.cgrer.uiowa.edu/arctas/emission.html](http://bio.cgrer.uiowa.edu/arctas/emission.html)). This inventory was used in early testing of the hemispheric CMAQ model (e.g., Mathur et al., 2014) and will be referred to as the ARCTAS inventory in subsequent discussions. More recent applications have relied on year specific estimates from the EDGAR (Emission Database for Global Atmospheric Research, version 4.2; European Commission, 2011) database which reports emissions for 17 anthropogenic sectors and large-scale biomass burning on a 0.1° × 0.1° resolution grid. Since EDGARv4.2 provides only PM$_{10}$ emissions, PM$_{2.5}$ emissions were estimated by deriving the ratio of PM$_{2.5}$ to PM$_{10}$ from the 2000-2005 EDGAR HTAP (Hemispheric Transport of Air Pollution, version 1) inventory (Janssens-Maenhout et al, 2012) and then applying this ratio to partition EDGARv4.2 PM$_{10}$ emissions into PM$_{2.5}$ and PM$_{2.5-10}$ (Xing et al, 2015a). In applications to date, biogenic VOC (Guenther et al., 1995) and lightning NOx (Price et al, 1997) emissions were obtained from GEIA (Global Emission Inventory Activity; http://www.geiacenter.org). The monthly biogenic VOC emissions were further temporalized to hourly resolution for each simulation day. Monthly lightning NO$_x$ emissions were distributed evenly to each hour of each simulation day. Xing et al. (2015a) further describe the processing of global emission inventories for CMAQ, including temporalization of the annual estimates to hourly model inputs, vertical distributions of anthropogenic and lightning emissions, and speciation of PM$_{2.5}$ and NMVOC emissions to model primary aerosol constituents.
and gas-phase species. Emissions of NO from soil or SO2 from volcanos are not considered in the applications presented here. It should also be noted that several efforts are underway to harmonize regional emission estimates and incorporate them into global emission inventories with improved spatial and temporal resolution (e.g., Janssens-Maenhout et al., 2015). Furthermore, the SMOKE modeling system typically used to prepare emissions for regional CMAQ applications has recently been updated to support hemispheric CMAQ applications to allow for a more streamlined implementation of the various emission processing steps described above (Eyth et al., 2016).

### 2.3.2 Wind-blown Dust

The wind-blown dust emission parameterization employed in CMAQ (Tong et al., 2008) was adapted for hemispheric applications by making two primary modifications. First, the mapping for land-use categories representing potentially erodible dust sources was updated to map to the categories of the MODIS land-use types used in the hemispheric WRF-CMAQ configuration. Second, the threshold friction velocity (above which dust emissions occur due to wind action) for desert regions was reduced to mobilize sufficient episodic dust emissions over the Sahara. The original value for threshold friction velocity, derived from the work of Gillette et al. (1980), was based on data from the Mojave Desert. However, Li et al. (2007) suggest a much lower (about half) threshold friction velocity based on dust samples from the northern China desert. Fu et al. (2014) found that the default threshold friction velocity for loose, fine grained soil with low surface roughness was too high for Asian dust sources and that reducing it to the Li et al. (2007) values yielded much better agreement of simulated airborne dust relative to observations. We found similar underestimations in PM2.5 concentrations and AOD over the Sahara with the default values, and have thus followed an approach similar to Fu et al. (2014) in the hemispheric CMAQ applications presented here. Concurrent with the development of this paper, a newer physics-based windblown dust emission parameterization was developed and implemented in CMAQ, and that parameterization includes a dynamic relation for the surface roughness length relevant to small-scale dust generation processes (Foroutan et al., 2017). The new dust emission parameterization is currently being tested for hemispheric applications and will be available in future public releases of CMAQ.

### 2.3.3 Emissions in Marine Environments

Natural emissions of particulate matter and gas-phase species from the oceans can impact air quality in coastal regions, influence global burdens of atmospheric trace species and radiative budgets, and modulate lifetimes of tropospheric O3 thereby influencing its long-range transport. A detailed representation of sea-spray particle emissions and chemistry is already available in CMAQ (Kelly et al., 2010), and it can be used for hemispheric scale applications without any modifications.

Reactive halogen emissions can play an important role in dictating lifetimes of O3 in marine environments. Parameterizations to estimate marine emissions of bromine and iodine containing compounds for the three categories (halocarbons, inorganic bromine, and inorganic iodine) were developed for inclusion in the hemispheric-CMAQ. The halocarbons include five bromocarbons (CHBr3, CH2Br2, CH2BrCl, CHBrCl2, CHBr2Cl) and four iodocarbons (CH3I, CH2ICl, CH2IBr, CH2I2). The
halocarbon emissions are estimated using monthly average climatological chl-α concentrations derived from the Moderate Resolution Imaging Spectroradiometer (MODIS). Sarwar et al. (2015) provide details on estimating halogen emission and comparisons with other existing estimates.

2.4 Enhancements to Gas-Phase Chemistry

The 2005 Carbon Bond Mechanism with updated toluene chemistry (CB05TU; Sarwar et al., 2011), commonly used in regional CMAQ applications, was also used for initial hemispheric-scale applications. Important enhancements to CB05TU were implemented to improve (1) its ability to represent multi-day chemistry associated with cycling of NOx through reservoir organic nitrate species in the mechanism, and (2) representing chemical sinks for tropospheric O3 due to halogen mediated chemistry in marine environments. Additionally, the more detailed RACM2 mechanism has also been implemented (Sarwar et al., 2013) to facilitate its use in follow-on hemispheric applications.

2.4.1 Organic Nitrate Lifetime

Organic nitrates form during the atmospheric photo-degradation of hydrocarbons in the presence of nitrogen oxides (NOx) through reactions of peroxy alkyl radicals (RO2) with NO as well as through reactions with NO3, and they act as a reservoir for oxides of nitrogen. In the CB05TU mechanism, the species NTR is used to represent organic nitrates. Depending on its modelled lifetime, NTR can potentially redistribute NOx from source regions to NOx-sensitive remote areas where additional ozone may be produced. Representing inert and reservoir organic nitrate species in condensed mechanisms used in chemistry-transport models is challenging (e.g., Kasibhatla et al., 1997) since they can dramatically influence simulated O3 and NOy distributions. In the CB05TU implementation in CMAQ, the chemical sinks for NTR include photolysis (producing NO2) and reaction with OH (producing HNO3). Additionally, defining a Henry’s law constant for a single lumped species representing several alkyl nitrates such as NTR is challenging. In previous CMAQ versions, the Henry’s law constant for PAN was also used for NTR, resulting in its very slow removal either through scavenging by clouds or through dry deposition at the Earth’s surface. However, the Henry’s law constants for several alkyl nitrates and hydroxyalkyl nitrates have been suggested to be much higher (some comparable to HNO3), especially those that are of biogenic origin (Shepson et al., 1996; Treves and Rudich, 2003). On the hemispheric scale, organic nitrates formed from isoprene are the largest contributor to the simulated tropospheric NTR burden and can consequently modulate the simulated tropospheric O3 burden. Based on recent work by Xie et al. (2013), we updated the rate constant for the NTR+OH reaction to that for isoprene nitrates. The Henry’s law constant for NTR was also mapped to that of HNO3, thereby enhancing wet scavenging of NTR. Additionally, the dry deposition velocity for NTR was mapped to that for HNO3. Collectively, these changes result not only in faster NOx recycling from NTR but also faster removal of NTR through the enhancement of its dry deposition and wet scavenging physical sinks.

The impacts of these changes to representing NTR in CMAQ on simulated O3 distributions are illustrated in Figure 4, which presents a comparison of monthly mean profiles of simulated O3 mixing ratios for various cases with ozonesonde
measurements at Trinidad Head, California, a site nominally representing inflow conditions to North America. The comparisons shown in Figure 4 illustrate the relatively large effects of modulating the resultant NTR burden on the simulated O$_3$ distribution through much of the lower to mid-troposphere, especially during summer when isoprene emissions are high. In limited-area calculations with the CB05TU mechanism, it is likely that the NTR produced is transported out of the regional domains before it can significantly alter O$_3$ production. However, over the spatial and temporal scales of northern hemispheric calculations, NO$_x$ recycled from NTR can modulate the simulated background O$_3$; consequently, accurate characterization of its sources and sinks becomes critical. Thus the hemispheric calculations provide a framework for examining the role of various physical and chemical processes on atmospheric chemical budgets in a consistent manner.

Additional improvements to representing NTR in the CB05TU mechanism in CMAQ are underway. In particular, replacing the single alkyl nitrate species (NTR) in CB05TU with seven species to better capture the range of chemical reactivity and Henry’s law constants (and thus the physical sinks) is being investigated (Schwede et al., 2014; Appel et al., 2017). Early comparisons of this expanded treatment with the simpler changes discussed above suggest that the approximations invoked through mapping the OH reactivity to isoprene nitrates (from Xie et al., 2013) and mimicking NTR’s wet and dry removal rates to HNO$_3$, yield similar simulated O$_3$ distributions to the ones obtained from the expanded treatment.

2.4.2 Representation of Marine Environments

More than half of the Northern Hemisphere is covered by oceans. To accurately represent the inter-continental transport of pollutants, it is important to accurately represent how continental air masses evolve as they traverse the vast oceanic regions. The fate of O$_3$ in marine environments directly affects inflow to continental regions and background O$_3$ concentrations. Though O$_3$ photolysis in the presence of high water vapor results in chemical O$_3$ loss and is well quantified, additional loss of O$_3$ in these environments through deposition as well as chemical reactions with halogens emitted from the ocean is expected (Vogt et al., 1999; Read et al., 2008), but still uncertain and not represented in most tropospheric models. In expanding CMAQ to hemispheric scales, particular attention was devoted to the role of deposition and halogen chemistry in marine environments, which can serve as sinks for O$_3$ exported from continental outflow and in transit to other regions via long-range transport. An enhanced O$_3$ deposition treatment that accounts for the interaction of iodide in seawater with O$_3$ was implemented (Sarwar et al., 2015) and found to increase deposition velocities in marine environments by an order of magnitude. In addition, the gas-phase chemical mechanisms were expanded to include 25 chemical reactions involving 7 chlorine species (Sarwar et al., 2012), 39 chemical reactions involving 14 bromine species, and 53 chemical reactions involving 17 iodine species (Sarwar et al., 2015).

2.4.3 Alternate Gas-Phase Mechanism

As discussed in Section 2.4.1, characterizing multi-day chemistry and long-lived reservoir species is important for representing the long-range transport of pollutants and their distributions on hemispheric scales. To enable practical model applications
over extended spatial and temporal scales, the chemical mechanisms must be sufficiently condensed to run efficiently while faithfully representing the chemistry over the space and time scales modelled. However, the impacts on model predictions of using different condensation rules and assumptions on species lumping and intermediate compounds are largely unquantified. To enable such investigation in the future over the hemispheric scale, an alternate chemical mechanism, RACM2 has also been implemented and tested in the hemispheric CMAQ (Sarwar et al., 2013). The RACM2 mechanism is designed to simulate remote to polluted conditions from the Earth’s surface through the upper troposphere (Goliff et al., 2013). It consists of 363 chemical reactions including 33 photolytic reactions among 120 chemical species.

2.5 Representing Impacts of Stratosphere-Troposphere Exchange on O₃ Distributions

Though the role of cross-tropopause transport of O₃ is acknowledged as a significant contributor to the tropospheric O₃ budget, the distribution of O₃ in the troposphere that originates from the stratosphere is still uncertain. Tightening O₃ NAAQS and decreasing amounts of photo-chemically derived O₃ due to continuously declining anthropogenic precursor emissions across large parts of North America and Europe, now put greater emphasis on accurately characterizing the fraction of O₃ in the troposphere, especially at the surface, that is of stratospheric origin. For instance, Roelofs and Lelieveld (1997) using a global chemistry-transport model estimated that stratospheric contributions to surface O₃ varied between 10-60% depending on season and location. Clearly this fraction varies spatially and seasonally in response to the tropopause height, and perhaps even more episodically, from deep intrusion events associated with weather patterns and frontal movement. Potential vorticity (PV) has been shown to be a robust indicator of air mass exchange between the stratosphere and the troposphere with strong positive correlation with O₃ and other trace species transported from the stratosphere to the upper troposphere (Danielsen, 1968). Numerous modeling studies have used this correlation to develop scaling factors that specify O₃ in the modelled upper troposphere/lower stratosphere (UTLS) based on estimated PV. The reported O₃/PV ratios (e.g., Ebel et al, 1991; Carmichael et al, 1998; McCaffery et al, 2004) however exhibit a wide range: 20-100 ppb/PVu (1 PV unit = 10⁻⁶ m² K kg⁻¹ s⁻¹), and vary as a function of location, altitude and season. Based on extensive ozonesonde measurements available during the summer of 2006 from the IONS network (http://croc.gsfc.nasa.gov/intexb/ions06.html) and PV fields from the WRF model matched to the location and time of the ozone-sonde launch, we examined the UTLS O₃-PV correlation at sites across North America. At 12 sites with sufficient number (11 or greater) of launches during August 2006, strong linear correlations (r²>0.7) were noted, with slopes of the linear regression varying between 20 and 39 ppb/PVu (Mathur et al., 2008). Based on this analysis, in the initial implementation of STE impacts on tropospheric O₃ in hemispheric CMAQ, we scale the space and time varying model estimated PV in top-most model layer with a scaling factor of 20 ppb/PVu to specify O₃ at the model top. This initial conservative choice for the O₃/PV ratio was in part dictated by lack of additional information on seasonality and also by the relatively coarse model resolution in the UTLS. As indicated in Figure 3, layer configuration influences the representation of STE and subsequent simulation of 3-D O₃ distributions. Thus the initial conservative choice of 20ppb/PVu was motivated by the desire to reduce any possible effects of excessive and artificial downward entrainment associated with inadequate vertical model resolution.
To overcome some of these challenges and to develop a more robust representation of STE impacts, we have recently developed a dynamic O$_3$-PV function based on 21-year ozonesonde records from World Ozone and Ultraviolet Radiation Data Centre (WOUDC) with corresponding PV values from WRF-CMAQ simulation across the Northern Hemisphere from 1990 to 2010. Analyses of PV and ozonesonde data suggests strong spatial and seasonal variations of O$_3$/PV ratios which exhibits large values in the upper layers and in high latitude regions, with highest values in spring and the lowest values in autumn over an annual cycle. The new generalized parameterization, detailed in Xing et al. (2016a), can dynamically represent O$_3$ in the UTLS across the Northern Hemisphere. The implementation of the new function significantly improves CMAQ's simulation of UTLS O$_3$ in both magnitude and seasonality compared to observations, which results in a more accurate simulation of the vertical distribution of O$_3$ across the Northern Hemisphere (Xing et al., 2016a). These hemispheric O$_3$ fields can then be used to derive more realistic vertically and temporally varying LBCs for regional nested model calculations.

3 Hemispheric-Scale Applications, Analysis, and Evaluation

The hemispheric CMAQ model is now being used for a variety of process-based air pollution studies across the Northern Hemisphere over seasonal (e.g., Mathur et al., 2014; Sarwar et al., 2014) to decadal (Xing et al., 2015a) time scales. In this section we present an overview of these diverse evolving applications with the hemispheric CMAQ model. In some instances, the applications have been detailed before, but the analysis summarized here builds upon that previous work and those distinctions are stated in the individual application discussion. Detailed comparisons of model predictions of pollutant concentrations (and radiative properties) with corresponding observations are conducted to establish credibility in the model’s use in these applications that range from representing episodic long-range pollutant transport to quantifying long-term trends in air pollution across the Northern Hemisphere, to emerging applications examining air pollution-climate interactions. Model applications are performed over the hemispheric domain and 44-layer structure illustrated in Figure 2. The CMAQ configuration is based on version 5.0 (CMAQv5.0) with the process updates detailed in Section 2. Two sets of applications are analyzed and evaluated in the subsequent discussion: (1) a 21-year simulation over 1990-2010 (in section 3.5 and 3.6), and (2) process sensitivity studies for the calendar year 2006, which were each initialized in September 2005 using fields from the prior 21-year simulation set (in sections 3.1-3.4).

3.1 Comparing model predictions with measurements from the 2006 INTEX-B campaign

The Intercontinental Chemical Transport Experiment-B (INTEX-B) was a NASA-led, multi-partner atmospheric field experiment conducted in the spring of 2006. A major objective of the second phase of the campaign during 17 April–15 May 2006 was characterizing the long-range transport and evolution of Asian pollution and implications for air quality across western North America (Singh et al., 2009). Airborne measurements of a variety of trace species were made over the remote Pacific as well along the inflow region to western North America from extensively-instrumented aircrafts and provide a unique
data set to test and evaluate the ability of hemispheric CMAQ to represent the 3-D structure of air pollutants as they are transported long distances across the Pacific Ocean to eventually impact U.S. background pollution levels.

The NASA DC-8 flights during 17 April–1 May 2006 were based out of Honolulu, Hawaii, and sampled the sub-tropical Pacific, while the flights during 4–15 May 2006 based out of Anchorage, Alaska, sampled the troposphere over the sub-Arctic Pacific region. Figures 5a-d present comparisons of modelled and observed O3 mixing ratios along selected flight paths of the DC-8 aircraft; several of these flights were designed to sample aged and fresh Asian pollution over the Pacific (Table 5a in Singh et al., 2009). Modelled mixing ratios were extracted by “flying” the aircraft through the 3-D modeling domain; the spatial locations of the aircraft were mapped to the model grid, whereas hourly model output was temporally interpolated to the time of the measurement. Figure 5 shows results from three different CMAQ configurations aimed at isolating the impacts of STE and marine halogen chemistry on simulated 3-D O3 distributions. Differences between the simulations denoted “PV+Halogen” and “Halogen, NoPV” are used to estimate the O3 sources due to modelled STE processes, while differences between the simulations “PV+Halogen” and “PV, NoHalogen” help quantify the model O3 sinks due to halogen chemistry in marine environments. Note that the simulation “PV, NoHalogen” employed the CB05TU mechanism, while the other two simulations employed a version of the CB05TU mechanism augmented with the halogen chemistry discussed in Section 2.4.2. Comparisons of model predictions from the “PV+Halogen” simulation with observations along the various flight paths suggest that the model exhibits skill in capturing the vertical variations in O3 observed in the region. The simulation that did not employ any PV scaling (green trace in Figure 5) systematically underestimates O3. The improved comparisons with measurements along the different flight paths for the PV+Halogen simulation suggest that O3 in the lower to mid troposphere in this region during this period is often influenced by sizeable contributions from the stratosphere, and these enhancements are generally captured by the simulation employing the PV scaling.

The model's ability to simulate the regionally averaged vertical profiles sampled by the aircraft over the subtropical and sub-Arctic Pacific is illustrated in Figures 5e and 5f, respectively. In constructing these composite average vertical O3 profiles, the observed and the modelled data were averaged within each 500 m vertical bin and over all the flights in that region; the figure also shows standard deviation for the observations. These vertical profiles represent the mean conditions that occurred over subtropical and sub-Arctic Pacific during the study period. The model tracks the composite average gradients within the lower and upper troposphere in these regions and accurately simulates that there is higher O3 in the sub-Arctic Pacific upper troposphere relative to the subtropical Pacific. Also apparent in these comparisons is the systematic and large underestimation of O3 throughout the troposphere in the simulation that did not account for any contributions due to STE processes. The much closer agreement of the observed composite profile with that derived from the simulations with the PV scaling further suggest that on average ~10 ppb (or greater) of the O3 in the troposphere over the Pacific during Spring could be of stratospheric origin. Thus O3 in air masses entering western North America is comprised of both anthropogenic contributions due to long-range transport of aged pollution from Asia and central America as well as a natural stratospheric component. The composite O3
vertical profile during this period derived from the hemispheric CMAQ is within the range of those predicted by other global atmospheric chemistry models illustrated in Singh et al. (2009).

Anthropogenic emissions from Asia are often lifted into the free troposphere and transported across the Pacific to North America in 5-10 days (e.g., Jaffe et al., 1999). Enhancements to free tropospheric SO$_4^{2-}$ measurements over northwest North America have been attributed to Asian sources (e.g., Andreae et al., 1988; Barrie et al., 1994). Increasing SO$_2$ emissions in Asia could potentially increase the amount of SO$_4^{2-}$ imported to North America and impact local efforts to reduce regional haze and improve visibility in national parks. Consequently, developing tools that accurately characterize the long-range transport from source regions and the amount of aerosols (both natural and anthropogenic) in air masses imported into a region is needed. To assess the ability of the hemispheric CMAQ model to represent airborne SO$_4^{2-}$ levels and gradients off the Pacific coast of North America, we compared model predictions of SO$_4^{2-}$ (total of SO$_4^{2-}$ in the Aitken and accumulation modes) distributions with measurements taken during the INTEX-B study. In addition to SO$_4^{2-}$ measurements from bulk aerosol filters on the DC-8, measurements from the Particle into Liquid Sampler (PILS) on board the C-130 aircraft (11 flights) were also analyzed to evaluate the simulated SO$_4^{2-}$ distributions within both the boundary layer and free-troposphere over the eastern Pacific and western North America. Analysis of the evolution of the MODIS aerosol optical depth (AOD) retrievals during mid-April 2006 (van Donkelaar et al., 2008) documents the development and transport of an Asian plume to western north America, and transects of C-130 flight on 21 April 2006 during 2000-2300 UTC sampled this plume in the free troposphere off the coast of western U.S. Figure 6 illustrates the simulation of this episodic Asian plume transport event. The flight path (color coded by UTC time) and sampling region are shown in Figure 6a. Simulated transport features of the Asian SO$_4^{2-}$ plume in the model layer at approximately 4 km altitude at 2100 UTC on 21 April are illustrated in Figure 6b while Figure 6c presents space and time matched comparisons of the model results with measurements along the C-130 flight path. Both the MODIS retrievals (in van Donkelaar et al., 2008) and model simulations in Figure 6b show the export of SO$_4^{2-}$ from East Asia and its eastward transport across the Pacific Ocean to the western coast of North America. As illustrated in Figure 6c, SO$_4^{2-}$ levels $>$1 μg m$^{-3}$ were often measured in the free troposphere. Both the observations and model show these enhanced SO$_4^{2-}$ levels at altitudes of 4-6 km, which in conjunction with the large scale simulated SO$_4^{2-}$ distributions in Figure 6b suggest that CMAQ captures the SO$_4^{2-}$ enhancements in the free troposphere associated with this episodic event. Some discrepancies in space-time matched model and observed concentrations are also apparent in Figure 6c, which likely result from the relatively coarse (108 km) horizontal grid resolution employed in the model calculations.

Comparisons of campaign average composite vertical profiles for SO$_4^{2-}$ for all the DC-8 and C-130 flights are shown in Figures 7a and 7b, respectively. Relative to the observations, CMAQ tends to overestimate mean SO$_4^{2-}$ levels especially in the lower troposphere as seen in the comparisons with the bulk filter measurements from the DC-8. It should be noted that the C-130 PILS measurements represent SO$_4^{2-}$ mass only for particles size $<$1μm, while the model values which are total mass in the Aitken and accumulation modes, nominally represents particle sizes $<$2.5 μm. This discrepancy in particle size cut-offs between
the measured and modelled SO$_4^{2-}$ in part contributed to the systematic overestimation relative to the C-130 PILS measurements. In their comparisons with model results, van Donkelaar et al. (2008) for instance used a factor of 1.4 to scale the PILS SO$_4^{2-}$ during INTEX-B to account for particle size restrictions. Using a similar scaling here would result in a much closer comparison of the composite measured and modelled SO$_4^{2-}$ profiles in Figure 7.

3.2 Episodic inter-continental transport of Saharan dust and impact on U.S. air quality

Some of the earliest recognition of long-range transport of air pollutants, dating back almost a century, was through observations of inter-continental transport of dust (Husar, 2004). North Africa is one of the largest sources of wind-blown dust, and the frequent transport of Saharan dust across the North Atlantic Ocean to the Caribbean has long been studied (e.g., Prospero and Carlson, 1972). Trans-Atlantic transport of major Saharan dust outbreaks can episodically influence tropospheric particulate matter loading in the southeastern U.S. The ability of the hemispheric CMAQ to simulate such long-range transport events is investigated through analysis of a Saharan dust transport event in late July-early August, 2006. The simulated development and trans-Atlantic transport of a Saharan dust plume during this period is illustrated in Figure 8, which presents daily average enhancements in PM$_{2.5}$ concentrations attributable to wind-blown dust. Large amounts of dust lofted into the atmosphere were transported west across the Atlantic, eventually impacting Gulf coast region of the U.S. Surface-level PM$_{2.5}$ measured in the U.S. Gulf states showed enhanced values as seen in the average concentrations across monitoring sites in Florida (Figure 9b).

A demonstration of CMAQ’s ability to simulate episodic long-range Saharan dust transport is shown using comparisons with surface-level PM$_{2.5}$ measurements at the Air Quality System (AQS) monitors in the Gulf states (Figure 9). The average change in bias in modelled PM$_{2.5}$ between simulations with and without dust emissions is shown in Figure 9a, which indicates a reduction in bias in the simulation incorporating the impact of Saharan dust emissions and transport. Collectively, the analysis here and in section 3.1 demonstrate that the hemispheric CMAQ modeling system can represent with reasonable skill the impacts of episodic trans-Atlantic (Figures 8 and 9) and trans-Pacific (Figures 6 and 7) transport events on air pollution over North America.

3.3 Assessing the influence of stratosphere-troposphere exchange on surface O$_3$

The analysis of 3D O$_3$ distributions from model sensitivity simulations relative to aircraft measurements, discussed in section 3.1, indicated the influence of STE processes on tropospheric O$_3$ distributions over the Pacific during the INTEX-B study period. To further analyze impacts of STE on tropospheric and surface-level O$_3$ over different seasons and regions, two simulations for the calendar year 2006 were conducted with the hemispheric CMAQ: with and without the dynamic PV-scaling approach discussed in section 2.5. Figure 10 presents the simulated seasonal average influence of STE processes on daily maximum 8-hour average (DM8) O$_3$, estimated as the difference between the simulations with and without the dynamic PV-scaling parameterization. As can be seen, the amount of O$_3$ at the surface that is of stratospheric origin varies substantially
both spatially as well as seasonally. As expected, high-latitude regions typically show greater influence of STE at the surface. Also the contributions to surface O3 from STE are much higher during Spring and Winter when height of the tropopause is lower (e.g., Elbern et al., 1998) and the stratospheric influence can penetrate far down to the lower troposphere (e.g., Wang et al., 2002).

Figure 11 presents an evaluation of the PV-scaling parameterization for representing the seasonal impacts of STE processes on surface DM8O3 relative to measurements from the CASTNET monitoring network in the U.S. A third simulation of 2006 was conducted using a constant O3-PV scaling factor of 20 ppb/PV unit rather than the dynamic scaling approach. The model-estimated stratospheric contribution to surface DM8O3 at the CASTNET locations can be estimated as the difference between the DM8O3 from the simulations with and without the dynamic PV scaling. The bias in the DM8O3 predictions was computed at each location for the simulations with constant-PV and dynamic-PV parameterization. A reduction in bias between these two simulations is a relative measure of the improvements in surface O3 predictions from using the dynamic-PV parameterization. Figure 11 correlates the seasonal average of this bias change with the estimated stratospheric contribution. The calculated seasonal means at each location, were restricted to days with observed DM8O3 >40 ppb. This helps screen out days where low O3 may not be captured due to model grid resolution or other process limitations, and limited the analysis to periods where STE influences were likely greater. A strong linear relationship is noted in Figure 11 between the bias change and estimated stratospheric contribution. Across all seasons and at most locations, the dynamic-PV parameterization reduced the bias in predicted surface DM8O3 relative to the constant-PV scaling. More importantly, when the estimated stratospheric contribution to surface O3 is high, greater reductions in model DM8O3 error are realized through the use of the dynamic-PV scaling parameterization, demonstrating the ability of the PV-based parameterization in representing the effects of STE on surface O3 levels and its seasonal and spatial variability. Additionally, the improvements in model predictions (i.e., reduction in model bias) of DM8O3 are also greater during Spring and Winter when the stratospheric contributions are higher (Figure 10). These evaluation results help build further confidence in the use of the dynamic-PV scaling parameterization in the hemispheric CMAQ model and for representing the impact of STE processes on surface O3 levels.

3.4 Comparison of O3 predictions using the RACM and CB05 mechanisms

As mentioned in section 2.4.3, the RACM2 is also available as an alternate and more detailed representation of gas-phase atmospheric chemistry for hemispheric-scale CMAQ applications. A detailed comparison of the CB05TU and RACM2 predictions for regional scale applications over the continental U.S. was described in Sarwar et al. (2013). A brief summary of comparisons of tropospheric O3 predictions using the two mechanisms in hemispheric CMAQ is presented in Figure 12. The simulations were conducted for May-August, 2006 and were initialized using chemical fields from an existing longer-term simulation. Differences in predictions of surface-level monthly mean O3 mixing ratios across the Northern Hemisphere using the RACM2 and CB05TU mechanisms are illustrated in Figure 12a for August 2006. Note that the simulations denoted
CB05TU and RACM2 did not include representation of halogen chemistry. In the simulation using the RACM2, higher O$_3$ is noted in polluted regions (regions with higher NO$_x$ in Figure 13e), but lower values are seen in the remote areas. These differences arise primarily due to higher rates of NO$_x$ recycling from organic nitrates and more active organic chemistry in RACM2.

To further assess the impacts of using different chemical mechanisms on 3-D O$_3$ predictions, modelled O$_3$ distributions were compared with ozonesonde measurements at Sable Island, Nova Scotia; Trinidad Head, California; and Hilo, Hawaii. Comparisons of monthly mean O$_3$ vertical profiles simulated using different chemical mechanisms with corresponding observed profiles are shown in Figures 12b-d. Also shown are predictions with a model configuration in which the RACM2 mechanism was augmented with the halogen chemistry described in section 2.4.2. At Sable Island, which often receives outflow from the U.S. northeast corridor, RACM2 overpredicts O$_3$ at lower altitudes. The higher O$_3$ relative to CB05TU in the North American outflow is likely due to the faster NO$_x$ recycling in RACM2 as discussed earlier. At Trinidad Head, both RACM2 and CB05TU overestimate O$_3$ near the surface, though RACM2 is closer to the observations at altitudes of 1000-3000 m. In contrast at Hilo, CB05TU overestimates O$_3$ and RACM2 is much closer to the observed profile. In general, the addition of halogen chemistry in RACM2 helps reduce the overestimation at lower altitudes. At altitudes > 1km, the RACM2 O$_3$ predictions are generally in closer agreement with the observations at all three sites. These mixed performance results indicate that neither mechanism is necessarily better suited over the other for hemispheric scale calculations. Nevertheless, analysis with both the CB05TU and RACM2 demonstrate the importance of NO$_x$-recycling from isoprene nitrates and halogen chemistry on simulated O$_3$ distributions. Additional analyses of NO$_y$ partitioning and HO$_x$ predictions is needed to gain further insights on the reasons for the differences between the behaviors of the two mechanisms.

3.5 Simulating long-term trends in tropospheric composition

Over the past two decades significant and contrasting changes have occurred in anthropogenic air pollutant emissions across the globe. Emissions control measures implemented in North America and western Europe have led to improvements in air quality in these regions. In contrast, due to increasing energy demand associated with rapidly growing economies and population, many regions in Asia and Africa are experiencing a dramatic increase in emissions of pollutants. These spatially heterogeneous emission trends across the globe have not only resulted in contrasting changes in human exposure levels to air pollution (e.g., Wang et al., 2017), but are likely impacting long-range transport patterns and influencing background air pollution levels in receptor regions. Accurate characterization of these changes in the chemical state of the troposphere (and potential influences on atmospheric dynamics) is needed to guide future control measures aimed at protecting human and environmental health. To assess these contrasting changes in air pollution levels, the hemispheric CMAQ was used to simulate trends in air quality across the Northern Hemisphere over a 21-year period (1990-2010). Year specific emission inputs were
derived from the Emission Database for Global Atmospheric Research (EDGAR, version 4.2) database (European Commission, 2011) as discussed earlier in Section 2.3.1 and detailed in Xing et al., (2015a).

Satellite-based tropospheric NO2 measurements are now providing valuable observable information on the changing emission patterns and air quality across the globe (e.g., Richter et al., 2005; van der A et al., 2008). To determine if the model can capture the impact of these changing emissions on tropospheric composition, trends in model tropospheric vertical column distributions (VCD) of NO2 were compared with those derived from radiances measured by satellite instruments GOME (Global Ozone Monitoring Experiment) and SCIAMACHY (Scanning Imaging Absorption spectrometer for Atmospheric Cartography). Note that the model NO2 column is estimated by integrating the predicted NO2 fields through the model column from the surface to ~50 hPa and did not utilize an averaging kernel. GOME NO2 observations are available during 1995-2003, while SCIAMACHY NO2 retrievals are available since 2002. Figures 13a, 13b, 13d, and 13e compare annual mean tropospheric vertical column NO2 for the calendar years 2003 and 2010 derived from SCIAMACHY retrievals and hemispheric CMAQ. Spatial distributions of the estimated 2003-2010 trends in NO2 VCD from SCIAMACHY and the model are presented in Figures 13c and 13f, respectively.

Figure 13 shows that the spatial distributions of NO2 VCD across the Northern Hemisphere are generally well correlated between CMAQ and SCIAMACHY, with higher NO2 in the industrial and urban areas of North America, Europe, and Asia. Some discrepancies are noted in central Africa where CMAQ simulates higher tropospheric NO2 in the Central African Republic and its northern border with Chad. Trends in NOx emissions derived from the EDGAR inventory show a similar increasing trend in this region (see Figure 2b in Xing et al., 2015a) and indicates that this discrepancy is associated with the underlying emission data set used in these simulations. In contrast, SCIAMACHY distributions in the region show a signal associated with biomass burning in the Savanna region of Africa both in 2003 and 2010 – the spatial extent of which is not captured by the model. Comparison of 2003-2010 trends in tropospheric NO2 between the CMAQ simulations and SCIAMACHY also indicate that the model captures increases in East China, and many cities in India and Middle East as well as the decreases across the eastern U.S., southern California, and western Europe. Figure 14 presents comparisons of time-series of regional-average monthly mean variations in tropospheric NO2 simulated by the model with corresponding values based on the GOME and SCIAMACHY retrievals for three regions: East China, United States, and Europe (see Figure 3 of Xing et al., 2015a for sub-domain extents). The domain-mean seasonal variability in tropospheric NO2 (as represented by the GOME and SCIAMACHY retrievals) is captured reasonably well by the model with a cool season maximum and warm season minimum. The model accurately simulates the amplitude of this variability for the U.S. as well as its inter-annual variability. For East Asia the model underestimates the peak values. In contrast for Europe, relative to both GOME and SCIAMACHY, the model consistently overestimates the summertime minimum values. Note that these simulations did not account for aerosol radiative feedback effects which, due to scattering and absorption of incoming solar radiation, reduce the amount of radiation impinging the Earth’s surface. The resulting stabilization can reduce boundary layer ventilation and increase surface-level
concentrations. As shown in Xing et al. (2015b, 2015c) these effects are especially important in polluted environment such as East Asia. Consequently, some of the underestimation in tropospheric NO2 over East Asia during the cooler months (when particulate matter pollution is the highest) could also arise from ignoring the aerosol direct radiative effects in simulated concentrations.

In addition to NOx, anthropogenic emissions of SO2 and volatile organic compound (VOCs) have also been reduced significantly in the U.S. To assess the impact of these precursor emission changes on trends in concentrations of secondarily produced species, we compared model simulated trends in ambient O3 and aerosol SO4^{2-} with those inferred from measurements at the CASTNET monitors. Figure 15 presents comparisons of model and observed trends in summer average daily maximum 8-hour average O3 and summer average weekly-average SO4^{2-} at each CASTNET monitor site. Trends are estimated as the slope of the linear regression of these concentration metrics for the 21-year period. Also shown in Figure 15 are results from an additional 21-year simulation with CMAQ that used a 36-km regional domain focused on the contiguous U.S. (see Figure 2). This regional simulation used an updated long-term emission inventory for the U.S. (Xing et al., 2013) and was driven by space and time varying lateral boundary conditions from the hemispheric CMAQ simulations for 1990-2010 (Gan et al., 2015a). Figure 15 shows that both the hemispheric-scale and the nested regional model capture the decreasing trends in both DM8O3 and SO4^{2-} as well as the spatial variability in the magnitude of the trends across the CASTNET sites, though the hemispheric model tends to underestimate the magnitude of the trends by 25-47%. However, the finer resolution of the nested simulation in conjunction with the updated emission inventory better captures the observed trends in surface level DM8O3 and SO4^{2-} as indicated by slopes closer to unity. This suggests the need to further explore finer resolution model calculations with the hemispheric CMAQ. As computing resources increase in the future it may be possible to conduct hemispheric scale simulations utilizing grid spacing finer than the 108 km utilized here.

3.6 Assessment of representation of aerosol direct radiative effects

Both aerosol scattering and absorption reduce the shortwave radiation impinging on the Earth’s surface. The variability in surface solar radiation plays a prominent role in the Earth’s climate system as it contributes to the modulation of the surface temperature, intensity of the hydrological cycle, and potentially the net ecosystem productivity (Wild, 2009). Observed trends in solar radiation reaching the Earth’s surface are very likely associated with changes in aerosol and aerosol-precursor emissions governed by economic development and air pollution regulations, which have modulated the trends in regional and global tropospheric aerosol burden over the past two decades (cf. Wild, 2009; Streets et al., 2009). Surface solar radiation “dimming” and “brightening” effects respectively dampen and enhance the warming trends induced by greenhouse gasses, so it is essential to accurately characterize these trends and quantify the role of regional variability in tropospheric aerosol burdens on these trends.
Simulations over the Northern Hemisphere can also be conducted using a two-way coupled WRF-CMAQ configuration (Wong et al., 2012), where CMAQ-simulated aerosol composition and size distribution are used to estimate their optical properties which are then fed back to the WRF radiation module to influence the simulated radiation by WRF. Thus the effects of aerosol scattering and absorption of incoming radiation can further impact the simulated atmospheric dynamics (boundary layer heights, temperature, simulated resolved and sub-grid scale clouds), which then impact emission rates, transport and dispersion, deposition, and temperature and actinic flux dependent chemical rate constants. The aerosol optical properties in the two-way coupled WRF-CMAQ are calculated using the BHCOAT coated-sphere module approach (Bohren and Huffman, 1983), i.e., particles in the Aitken and accumulation model are assumed to have a core composed of elemental carbon with a shell coating of other species. The aerosol optics calculations in the WRF-CMAQ have been evaluated against field measurements as detailed in Gan et al. (2015b).

In addition to the long-term (1990-2010) simulations discussed in Section 3.5, additional feedback simulations were conducted over the Northern Hemisphere with the two-way coupled WRF-CMAQ configuration for the summer months (June, July and August) of this 21-year period. Comparison of these two sets of simulations (with and without aerosol feedbacks) provides an indication of the impact of the aerosol direct radiative effects (ADRE) and an assessment of its trends associated with the changing tropospheric aerosol burden over the multi-decadal period. Figure 16 examines the modelled and observed relationships between changes in aerosol optical depth (AOD; representing changes in the tropospheric aerosol burden) and changes in clear-sky surface shortwave radiation (SWR) using regional monthly averages for East China, Europe, and East U.S. Figure 16 examines 2000-2010, when satellite-based data were available. Monthly regional averages of SWR and AOD were calculated for each of the summer months (June, July, and August). To minimize the influence of month-to-month variability, monthly averaged SWR and AOD were deseasonalized by subtracting the average of 11-year data for the corresponding month. Additionally, we used 24-hour-averaged SWR but AOD at noon (local-time) for model values to be consistent with the observation-derived data from satellite-retrievals (Xing et al., 2015b). The relationship between these deseasonalized values (or anomaly) of SWR and AOD for each summer month in the 2001-2010 period is examined in Figure 16 for both model simulations with and without direct aerosol feedback effects. Also shown in Figure 16 is the corresponding observed relationship between similarly estimated AOD anomaly and SWR anomaly derived from retrievals from the MODIS and the Clouds and the Earth’s Radiant Energy System (CERES; Wielicki et al., 1998) instruments, respectively. Note that the CERES mission estimates clear-sky surface SWR through radiative transfer calculations using satellite-retrieved surface, cloud, and aerosol properties as input (Kato et al., 2013), which have also been shown to agree with surface observations (Wild et al., 2013).

All three regions show a strong relationship between observed changes in AOD and clear-sky surface SWR, with reductions in SWR associated with increases in AOD and increases in SWR with reductions in AOD. This observational comparison clearly suggests that as tropospheric aerosol burden increases, scattering and absorption reduces the amount of surface SWR.
The magnitude of these changes is comparatively larger for East China than for Europe and East U.S., due to the higher levels of tropospheric particulate matter in East China. In all three regions, the model simulation without direct aerosol feedback fails to capture the changes in SWR and its association with AOD. In contrast the model simulation incorporating the aerosol direct radiative effects replicates the relationship between the observed AOD and SWR changes during the 2000-2010 period in all three regions as reflected by the higher R² and slopes of the linear regression closer to those inferred from the observed data. These results suggest that the hemispheric two-way coupled WRF-CMAQ configuration can represent the differing regional changes in surface SWR with contrasting changes in regional aerosol burden. Accordingly, this tool could also be used to examine chemistry climate interactions on hemispheric to regional scales.

4 Summary and Concluding Remarks

The applicability of the Community Multiscale Air Quality (CMAQ) modeling system was extended to the entire Northern Hemisphere to enable consistent examination of interactions between atmospheric processes occurring at various spatial and temporal scales. Improvements were made in model process representation (stratospheric O₃ influences, representation of NOₓ recycling through organic nitrates, halogen chemistry in marine environments, deposition over water), structure (model vertical extent and layer resolution), and input data sets (allocation of global emission estimates). These improvements to CMAQ were investigated and evaluated through comparison of model predictions with measurements from surface, aloft, remote sensing and specialized field campaign platforms. Comparisons with measurements from the INTEX-B field campaign indicate that hemispheric CMAQ captures the mean variability in O₃ and SO₄²⁻ distributions observed over the tropical and sub-Arctic Pacific regions, and episodic transport of Asian pollution across the Pacific as indicated by comparisons of model and observed SO₄²⁻ along specific flight tracks. The ability to capture the development and evolution of inter-continental transport (i.e., the lofting of pollutants in the source region, multi-day transport in the free troposphere and subsequent subsidence and mixing down to the surface in receptor regions) is also demonstrated by evaluating a trans-Atlantic Saharan dust transport event and its contributions to elevated surface PM₂.₅ in the U.S. Gulf region. These results suggest that regional CMAQ applications can now be driven by space and time varying lateral boundary conditions derived from consistent hemispheric applications, enabling examination of air quality across the U.S. in context of the changing global atmosphere.

Hemispheric CMAQ model can reproduce historical trends in tropospheric air pollution, as shown by comparing simulated results with surface and remote-sensing observation-derived records during 1990-2010. Trends in modelled tropospheric NO₂ vertical column distributions agree with those inferred from GOME and SCIAMACHY retrievals and indicate the contrasting and heterogeneous changes in emissions across the Northern Hemisphere, with increases in rapidly developing regions of Asia and decreases in Europe and North America resulting from implementation of control measures. Additionally, comparisons with observed trends at the U.S. CASTNET monitors, indicate that though the model captures the resultant decreasing trends in surface-level air pollution (for O₃ and SO₄²⁻) in the U.S., the current configuration underestimated (by 25-47%) the
magnitude of the trend at some monitoring locations. The underestimation in the magnitude of the trend is however significantly reduced in a nested regional simulation utilizing finer horizontal grid resolution and updated historical regional emissions. The changing emission patterns across the Northern Hemisphere will likely influence future long-range pollutant transport patterns and potentially impact background pollution levels in receptor regions. The hemispheric CMAQ provides a framework to explore such changing impacts on air pollution exposure. For instance, Wang et al. (2017) estimated trends in PM$_{2.5}$ premature mortality during 1990-2010 using hemispheric CMAQ predictions, and show that correlations between population and PM$_{2.5}$ have become weaker in Europe and North America due to air pollution controls but stronger in East Asia due to deteriorating air quality.

Analysis of aerosol optical and radiative effects inferred from the two-way coupled WRF-CMAQ applications over the Northern Hemisphere also indicate the association between changing tropospheric aerosol burden and clear-sky surface shortwave radiation. In rapidly developing regions such as East Asia, the increasing tropospheric aerosol burden results in greater scattering and absorption by aerosols, and that acts to reduce the amount of clear-sky surface shortwave radiation. In contrast, increases in clear-sky surface shortwave radiation are noted in regions with declining tropospheric aerosol burden, where emissions controls have been more active during that period. The two-way coupled WRF-CMAQ configuration that incorporates direct aerosol radiative effects captures these contrasting observed changes in clear sky shortwave radiation across the Northern Hemisphere during 2000-2010, with “brightening” in U.S. and western Europe and “dimming” in East China. Using these modeling results, Xing et al. (2016b) show that because of reduced atmospheric mixing resulting from direct aerosol feedbacks, air pollutants become more concentrated locally, especially in highly polluted and populated regions. Thus modulation of air pollution due to aerosol direct effects also translates into an additional human health dividend in regions (e.g., U.S. and Europe) with air pollution control measures but a penalty for regions (e.g., East Asia) witnessing rapid deterioration in air quality.

Analysis of three-dimensional O$_3$ distributions across the Northern Hemisphere from model sensitivity simulations and comparisons with surface and aloft measurements also highlight the need to better quantify the contribution of STE processes on surface O$_3$. A non-trivial contribution of up to ~10 ppb from the stratosphere to seasonal mean surface-level O$_3$ mixing ratios is inferred from the current applications. An accurate characterization of this contribution is essential for source attribution of background O$_3$. Since measurements of 3-D O$_3$ distributions alone are insufficient to directly quantify this contribution, model estimates need to be better constrained. To that end additional CMAQ simulations that explore the sensitivity of STE process representation to model vertical extent and vertical grid resolution are warranted. Model calculations presented here also indicate the possible influence of horizontal grid resolution on model evaluation results. Hemispheric CMAQ simulations to date have employed a horizontal grid resolution of 108 km, which is insufficient to resolve local gradients. Emerging environmental problems will likely require the simultaneous characterization of air pollution from local-to-global scales. Variable resolution nonuniform grids can simultaneously and accurately resolve local gradients and
large-scale features in air pollutant distributions (e.g., Odman and Russell, 1991; Mathur et al., 1992; Srivastava et al., 2000). The emergence of variable resolution atmospheric dynamical models (e.g., Skamarock et al., 2012) provides opportunities to develop comprehensive atmospheric modeling systems that seamlessly represent the scale interactions from urban to global scales. The use of such approaches could improve the representation of scale interactions in air pollution modeling.

Several efforts are underway to harmonize regional emission estimates and incorporate them into global emission inventories with improved spatial and temporal resolution (e.g., Janssens-Maenhout et al., 2015). It can be expected that future improvements in performance of hemispheric CMAQ will also be realized through improvements in these underlying global emission inventories used to drive model calculations. Additional improvements in sector specific emissions and additional details on chemical speciation of the emissions will also lend themselves to the use of more detailed chemical mechanisms such as RACM2 that explicitly treat the chemistry of longer-lived species (e.g., acetone) that are important for chemistry of the upper troposphere, and help further assess the relative benefits of the use of different chemical mechanisms at hemispheric scales. Predictions of a variety of atmospheric pollutant species from hemispheric CMAQ are also being compared to those from other modeling systems (Hogrefe et al., 2015) through activities of Air Quality Model Evaluation International Initiative (AQMEII). The adequacies and inadequacies of the lateral boundary conditions derived from hemispheric CMAQ to drive regional CMAQ simulations are further being analyzed through comparisons with those from other large scale models and observations (Hogrefe et al., 2017) and will also guide the future evolution of the hemispheric CMAQ.

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Figure 1: Impact of lateral boundary conditions (LBC) on simulated seasonal surface-level concentrations. (a-d): Spatial variation in seasonal-mean surface concentrations normalized by the maximum value within the model domain across all seasons. (e-h): Fractional contribution of free-tropospheric (FT) LBCs (specified between 750-250 hPa) to the total LBC-derived concentrations at the surface. Seasons are defined as: Winter (December-February), Spring (March-May), Summer (June-August), Fall (September-November).
Figure 2: Left: The Northern Hemisphere modeling domain discretized using a 108 km resolution grid. The shaded region shows the extent of the typical nested Continental U.S. nested domain discretized using a 12 km resolution horizontal grid. Right: Comparison of two layer configurations used to discretize the vertical extent ranging from the surface to 50 hPa.
Figure 3: Impact of layer configuration on simulated mean O$_3$ vertical profiles for August 2006 at selected locations for a case involving zero-out of emissions across the U.S.: (a) Trinidad Head, CA, (b) Boulder, CO, (c) Huntsville, AL, and (d) Narragansett, RI.
Figure 4: Comparisons of simulated average vertical profiles of O₃ with ozonesonde measurements at Trinidad Head, California, USA: (a) March 2006 (four months after start of simulation), and (b) August 2006 (nine months after start of simulation). Also shown is ±1 standard deviation of the observed mixing ratios. “Profile IC” uses the default profile for initialization as in regional CMAQ applications, “Clean IC” is the case where the model is spun up from clean conditions, and “Revised NTR” is the simulation with “Clean IC” with updates to the physical and chemical sinks for the species NTR representing organic nitrates.
Figure 5: Comparison of simulated 3-D distributions of O₃ mixing ratios with observations from the DC-8 aircraft during the INTEX-B field campaign: (a-d) comparison of observed O₃ and simulated values from various model configurations along flight tracks on specific days; (e) comparison of model and observed campaign average O₃ vertical profile for flights over the sub-tropical Pacific during 17 April-1 May 2006; (f) comparison of model and observed campaign average O₃ vertical profile for flights over the sub-Arctic Pacific during 1-15 May 2006. Also shown in (e) and (f) is ±1 standard deviation for the observed values. Also shown in (a-d) is the aircraft altitude along the flight path.
Figure 6: Simulation of impact of trans-Pacific transport on 3-D distributions of SO$_4^{2-}$ aerosol on 21 April 2006. (a) Flight path of C-130 aircraft color-coded by hour (UTC). (b) Simulated SO$_4^{2-}$ distribution at 4 km altitude on 21 April 2006 at 2100 UTC. (c) Comparison of modelled and observed SO$_4^{2-}$ aerosol concentrations along C-130 flight path.
Figure 7: Comparison of modelled and observed campaign average $\text{SO}_4^{2-}$ vertical profiles: (a) against measurements from the DC-8 aircraft, and (b) against measurements from the C-130 aircraft. Also shown is $\pm 1$ standard deviation for both observed and modelled values.
Figure 8: Trans-Atlantic transport of a Saharan dust plume, 29 July – 3 August 2006, as simulated by hemispheric CMAQ. Shown in the panels is the difference in daily-average PM$_{2.5}$ concentrations (μg m$^{-3}$) between CMAQ simulations with and without considering dust emissions.
Figure 9: The impact of simulated trans-Atlantic transport of Saharan dust on daily-average surface PM$_{2.5}$ in the Gulf region. (a) Average change in bias in daily average surface PM$_{2.5}$ (in μg m$^{-3}$) at AQS monitor locations between CMAQ simulations with and without considering dust emissions, 29 July-2 August 2006. Negative changes in bias denote improvement in model performance by including Saharan dust emissions and representing its trans-Atlantic transport. (b) Comparison of modelled and observed daily-average surface PM$_{2.5}$ averaged over all AQS monitor locations in Florida.
Figure 10: Impact of dynamic PV scaling on surface-level seasonal mean maximum daily 8-hour average O$_3$ (MD8O$_3$) mixing ratios (in ppb) estimated as the difference between the simulation with and without the dynamic PV formulation.
Figure 11: Impact of dynamic PV scaling parameterization on model performance for surface-level seasonal mean maximum daily 8-hour average O$_3$ (MD8O$_3$) at CASTNET sites in the U.S. The model estimated stratospheric contribution is estimated as the difference between the simulation with the dynamic PV scaling and one without. The bias change is estimated as the difference between the absolute bias in the simulation with a constant-PV (20 PV) scaling and the absolute bias in the dynamic-PV scaling simulations. Positive changes in bias represent reduction in bias due to dynamic-PV, while negative changes represent an increase in bias in simulated surface MD8O$_3$. Seasonal means are computed based on model-observed pairs when the observed MD8O$_3$ >40 ppb.
Figure 12: Differences in simulated O₃ distributions using the RACM2 and CB05TU gas-phase chemical mechanisms in the hemispheric CMAQ model. (a) Mean difference (RACM2 minus CB05TU) in monthly mean surface-level O₃ (in ppb) for August 2006. Comparisons of mean (for August 2006) O₃ mixing ratio vertical profiles simulated with the CB05TU and RACM2 mechanisms with ozonesonde measurements at (b) Sable Island, Nova Scotia, (c) Trinidad Head, California, and (d) Hilo, Hawaii. RACM2_HAL is from a simulation in which the RACM2 mechanism was augmented with halogen chemistry (described in section 2.4.2).
(a) SCIAMACHY; 2003

(b) SCIAMACHY; 2010

(c) SCIAMACHY 2003-2010 Trend

(d) CMAQ; 2003

(e) CMAQ; 2010

(f) CMAQ 2003-2010 Trend
Figure 13: Comparison of observed (left) and model (right) changes in NO$_2$ vertical column density (VCD) across the northern hemisphere. (a) 2003 SCIAMACHY NO$_2$ VCD; (b) 2010 SCIAMACHY NO$_2$ VCD; (c) SCIAMACHY NO$_2$ VCD trend; (d) 2003 CMAQ NO$_2$ VCD; (e) 2010 CMAQ NO$_2$ VCD; (f) CMAQ NO$_2$ VCD trend. VCD is units of $10^{15}$ molecules cm$^2$, and VCD trend is in units of $10^{15}$ molecules cm$^2$ year$^{-1}$.
Figure 14: Comparison of changes in regional- and monthly-average modelled NO$_2$ vertical column density with satellite retrievals from SCIAMACHY and GOME for (a) East Asia, (b) United States, and (c) Europe.
Figure 15: Comparison model and observed summertime (JJA) 1990-2010 trends at CASTNET monitoring sites in the U.S. for (a) JJA average daily maximum 8-hour average O$_3$, and (b) JJA average SO$_4^{2-}$. Model results from the 108 km resolution hemispheric CMAQ simulation are shown in blue, while results from a 36-km resolution nested model calculation over the contiguous U.S. are shown in red.
**Figure 16**: Relationship between regional and monthly average (only summer months) changes in aerosol optical depth and changes in surface clear-sky shortwave radiation during the 2001-2010 period for (a) East China, (b) Europe, and (c) East U.S. Observed values are shown in grey, CMAQ calculations with direct aerosol radiative feedbacks in red, and CMAQ calculations without aerosol radiative feedbacks in blue. Also, indicated are the slope and correlation coefficient (R) for the individual linear regressions. For each data set (model or observed) there are 33 values, corresponding to each summer month over the 11-year (2001-2010) period. The anomaly is estimated by subtracting the corresponding 11-year average for that month.