Comparison of Ozone Formation Attribution Techniques in the Northeast United States

Qian Shu¹, Sergey L. Napelenok¹, William T. Hutzell¹, Kirk R. Baker¹, Benjamin Murphy¹, Christian Hogrefe¹, Barron H. Henderson¹

⁵ ¹U.S. Environmental Protection Agency, Research Triangle Park, NC, 27711, USA.

Correspondence to: Sergey L Napelenok (sergey.napelenok@epa.gov)

Abstract.

The Integrated Source Apportionment Method (ISAM) has been revised in the Community Multiscale Air Quality (CMAQ) model. This work updates ISAM to maximize its flexibility, particularly for ozone (O₃) modeling, by providing multiple attribution options, including products inheriting attribution fully from nitrogen oxide reactants, fully from volatile organic compound (VOC) reactants, equally to all reactants, or dynamically to NO_x or VOC reactants based on the indicator gross production ratio of hydrogen peroxide (H₂O₂) to nitric acid (HNO₃). The updated ISAM has been incorporated into the most recent publicly accessible versions of CMAQ (v5.3.2 and beyond). This study's primary objective is to document these ISAM updates and demonstrate their impacts on source apportionment results for O₃ and its precursors. Additionally, the ISAM results are compared with the Ozone Source Apportionment Technology (OSAT) in the Comprehensive Air-quality Model with Extensions (CAMx) and the brute force method (BF). All comparisons are performed for a 4km horizontal grid resolution application over the northeast U.S. for a selected two-day summer case study (August 9th and 10th, 2018). General similarities among ISAM, OSAT, and BF results add credibility to the new ISAM algorithms. However, some discrepancies in magnitude or relative proportions among tracked sources illustrate the distinct features of each approach while others may be related to differences in model formulation of chemical and physical processes. Despite these differences, OSAT and ISAM still provide useful apportionment data by identifying the geographical and temporal contributions of O₃ and its precursors. Both OSAT and ISAM attribute the majority of O₃ and NO_x contributions to boundary,

mobile, and biogenic sources, whereas the top three contributors to VOCs are found to be biogenic, boundary, and area sources.

1 Introduction

40

Tropospheric O₃ is a critical air pollutant that endangers human health (WHO, 2013) and sensitive vegetation (Booker et al., 2009), and contributes to climate change (Jacob and Winner, 2009). It is produced through non-linear photochemical reactions of carbon monoxide (CO), volatile organic compounds (VOC), and nitrogen oxides (NO_x = NO + NO₂) with sunlight (Atkinson, 2000). In the United States, the national average ambient O₃ concentration has decreased by 22% since 1990, owing to regulations such as the Clean Air Act (CAA) on NO_x and VOC emissions (Simon et al., 2015). Long-term space observations have also confirmed the improvement in air quality (Duncan et al., 2013; Lamsal et al., 2015). However, many major metropolitan areas continue to exceed the O₃ national ambient air quality standards (NAAQS) set by the US Environmental Protection Agency (US EPA). To continue to reduce O₃ levels, it is critical to develop effective emission control strategies as has been done for other pollutants (Lefohn et al., 1998; Reitze, 2004; Cooper et al., 2015). The effectiveness of any O₃ control strategy hinges on accurately quantifying the contributions of various precursor emissions to O₃ formation.

Numerous techniques have been used to characterize and quantify the relationship between emission sources and O₃ concentrations, including statistical methods, model sensitivity simulations, and model source apportionment approaches, each with its own set of advantages and disadvantages (Cohan and Napelenok, 2011). While some traditional receptor-based methods based on chemical mass balance (CMB, Hidy and Friedlander, 1971), such as Effective Variance solution (EV, Watson et al., 1984) and Positive Matrix Factorization (PMF, Paatero and Tapper, 1994), produce insightful results when measurements are taken at a specific receptor, they are typically applied to speciated VOC and particulate matter (PM) and are also constrained by the relative sparsity of observations in space and time, rendering them unsuitable for regional and national O₃ precursor emission control strategies. Alternatively, three-dimensional air quality models (AQM) allow for the quantification of O₃ source contributions at regular intervals over longer periods and wider spatial distributions. The most basic

source apportionment (SA) technique in the context of an AQM is to conduct source sensitivity simulations using the brute force (BF) method, in which several simulations are conducted, each with one source eliminated or reduced. The differences in the output fields compared to the baseline simulation are then attributed to the eliminated or reduced source (e.g., Marmur et al., 2005). BF has some limitations when used to determine total source culpability of O₃ due to the pollutants' nonlinear dependence on both relative and absolute VOC and NO_x concentrations. For example, removing NO_x may lead to an increase of O₃ concentrations in the vicinity of large NO emissions (e.g., power plants), as the result of net conversion of O₃ to NO₂ (Gillani et al., 1996) or at night-time when NO_x titration cannot be balanced by the photolysis of NO₂. In some cases, where a source contributes a substantial portion of total NO_x or VOC emissions, complete source removal for the purposes of source apportionment calculation may also substantially alter the underlying chemical regime for formation of secondary pollutants such as O₃. Further, to separate the contributions and interactions of "n" sources, Stein and Alpert (1993) showed that BF would require two to the power of the number of sources (2ⁿ). This is quickly impractical leading to a subset of BF simulations with unknown interactions. As a result, summarizing the O₃ change in response to multiple brute force emission source simulations can make it difficult to interpret the cumulative effect of those emissions on O₃ (Kwok et al., 2015).

Reactive tracer or tagged species SA methods for O₃ have also been incorporated in AQMs.

These tracers are usually additional species added to the AQM to track the contributions of pollutants from specific source categories. They undergo the same atmospheric processes as the bulk chemical species within the model (Kwok et al, 2015). As one example, OSAT within CAMx quantifies the contributions of various emission sectors, source regions, as well as initial and lateral boundary conditions, to simulated O₃ concentrations (Ramboll Environ, 2015). OSAT allocates instantaneous O₃ formation to either NO_x or VOCs based on the ratio of hydrogen peroxide (H₂O₂) to nitric acid (HNO₃) production (Dunker et al., 2002). O₃ formation is classified as being NO_x-limited or VOC-limited based on the gross production of H₂O₂ (PH₂O₂) and HNO₃ (PHNO₃). When the ratio (PH₂O₂/PHNO₃) is above 0.35, the formation is classified as NO_x-limited and VOC-limited otherwise (Sillman, 1995). If the photochemical formation of O₃ (PO₃) occurs in a NO_x-limited regime, the NO_x tracers are used to attribute PO₃ proportionally to the emissions sources that contributed to the NO_x concentrations.

Otherwise, VOC tracers are used to attribute PO₃ to the sources that contributed to the VOC concentrations (Dunker et al., 2002; Kwok et al., 2015). The OSAT formulation was recently changed (OSAT3) to track all forms of NO_x to account for NO_x recycling, which occurs when NO_x is converted to another form of NO_x (e.g., peroxyacetyl nitrate (PAN) or HNO₃) and then converted back to NO_x. OSAT has been used to support policy assessments (e.g., U.S. EPA, state government agencies, etc., Ramboll Environ, 2015, 2022) as well as for scientific research purposes (Li et al., 2012; Zhang et al., 2017; Shu et al., 2020).

Additionally, the Integrated Source Apportionment Method (ISAM) within CMAQ has shown promising results for O₃ tagging (Kwok et al., 2015). Recent ISAM experiments have quantified the contribution of O₃ sources to air pollution in several major cities throughout the United States and Europe (Kwok et al., 2015; Valverde et al., 2016; Karamchandani et al., 2017; Butler et al., 2018; Pay et al., 2019). The attribution of O₃ and precursors from specific sources estimated by ISAM implemented in version 5.0 of CMAQ compared well with source-specific aircraft transect measurements (Baker and Woody, 2016). The ISAM algorithms have also been updated several times following the original implementation in CMAQv5.0.2.

95

100

105

ISAM updates presented in this study substantially increase the flexibility to the user of the CMAQ source apportionment model. These updates were intended to provide long term flexibility within the model to accommodate newer chemical mechanisms and changed the attribution approach as detailed in the methods section. These flexibilities allow for apportionment of more species and allow for more methods of apportionment. Further in the manuscript we apply the changes to CMAQ-ISAM for a Northeastern U.S. O₃ air quality episode and compare the results to CMAQ-BF and CAMx-OSAT. The manuscript is organized as follows: Section 2 documents the ISAM updates in detail; Section 3 describes the methodology for this study, which includes the base modeling configurations, simulation designs for source apportionment, tracked species classes, evaluation methods, and case study development; Section 4 presents the findings, including model evaluation results and comparisons of source apportionment for several species; Section 5 documents the running speed comparisons between CMAQ-ISAM, CAMx-OSAT and CMAQ-BF; and finally, the findings and their implications for future research are discussed in Section 6.

2 Source apportionment methods

110 2.1 Updates in ISAM

115

120

125

130

135

The ISAM implementation in the version 5.0 release of CMAQ was based on Kwok et al. (2013 and 2015). That approach was then updated starting from CMAQ version 5.3 to an attribution based on integrated reaction rates and product yields (US EPA, 2019). The later versions (v5.3.2 and beyond) of CMAQ-ISAM (US EPA, 2022a) employ an apportionment scheme that assigns products of each chemical reaction to sources based on reactant stoichiometry. For example, the isoprene peroxy radical (ISO₂) reacts with nitric oxide (NO) to produce several different stable and radical species as represented in the CB6R3 chemical mechanism by the following reaction R1.

 $ISO_2 + NO = 0.1*INTR + 0.9*NO_2 + 0.673*FORM + 0.9*ISPD + 0.818*HO_2 + 0.082*XO_2H + 0.082*RO_2 (R1)$

In addition to nitrogen dioxide (NO₂), the products include isoprene nitrate (INTR), formaldehyde (FORM), hydroperoxy radicals (HO₂), alkoxy radicals (XO₂H), peroxy radicals (RO₂), and other isoprene reaction products (ISPD). ISO₂ is a product of the oxidation of isoprene, which originates from overwhelmingly biogenic sources. NO is typically emitted from anthropogenic combustion processes, with a much smaller natural component originating from lightning strikes and microbial soil processes on the global scale (Jacquemin et al., 1990; Yienger et al., 1995). Thus, the reactants are approximately half from biogenic and half from anthropogenic sources, so the reaction's products have the same attribution distribution. However, source attribution approaches, both receptor-based (such as PMF) and source-based (such as ISAM), are often used to understand how originally emitted NO_x and VOC from particular sources ultimately contribute to model-predicted O₃ production.

The loss of source identity through processes such as the NO_x cycle and the role of organic peroxy radicals from sources not controlling O₃ production make it difficult to determine the culpability of emission sources. In the preceding example, the NO₂ produced by R1 is assigned a source that is approximately 50% biogenic and 50% anthropogenic. These source assignments propagate quickly when catalytic processes cause NO₂ to cycle back to NO through photooxidation and radical oxidation

Because NO_x cycling is fast in regional air pollution models, anthropogenically emitted nitrogen species can be assigned to biogenic (or other nearby) sources downwind, so the original source identity was not

retained. R1 is just one example that illustrates the complex relationship between precursors and subsequent source identities of secondary pollutants. Many such reactions exist in modern chemical mechanisms. Some source apportionment applications, such as O₃ source attribution assessments, focus on how sources induce O₃ production above background levels. Nitrogen molecules should then retain their original source signatures. This approach is used by other apportionment models such as OSAT, earlier ISAM implementations (Kwok et al., 2015), and other tagging methods (Butler et al., 2018; Grewe et al., 2010).

Because attribution objectives may vary based on scale (e.g., global compared to urban) or

purpose (e.g., policy or tracing chemical reactions), ISAM has been enhanced to provide additional
configuration options for the user to define how secondarily formed gaseous species are assigned to
sources of parent reactants (Table 1) (US EPA 2022b). The existing scheme based on stoichiometrically
proportional product attribution introduced in CMAQ version 5.3.2 has been retained as ISAM option 1
(ISAM-OP1). Four new options have been added so the user can configure their simulation based on the
application's goal. Each option allows for greater retention of source identity based on subsets of
species in the chemical mechanism. ISAM-OP2 apportions products according to the source identity of
reactive nitrogen species, including NO, NO2, nitrate radical (NO3), nitrous acid (HONO), HNO3,
dinitrogen pentoxide (N2O5), and aerosol nitrate (ANO3). For example, CB6R3 contains the following
reaction between the methyl peroxy radical (MEO2) and NO:

155
$$MEO_2 + NO = FORM + HO_2 + NO_2 (R2)$$

140

160

In the original ISAM-OP1 configuration, the products of R2, FORM, HO₂, and NO₂ inherit source identities proportional to the source identities of the reactants (MEO₂ and NO). However, ISAM-OP2 apportions the product to be from the source identity of NO (presumed predominantly anthropogenic), because NO is a weighted nitrogen-containing species. When a reaction's reactants do not include any of the weighted species, products are apportioned to source identities using the same methodology used in OP1.

ISAM-OP3 expands OP2's list of weighted species to include VOC species identified as important to O₃ production. In CB6R3, this includes aldehydes (ALD₂ and ALDX), FORM, acetone (ACET), lumped ketones (KET), peroxy operators (XO₂ and XO₂H), ISO₂, acetyl peroxy radicals (C₂O₃

and CXO₃). Therefore, products of reactions containing these VOCs in addition to the nitrogen species of OP2 as reactants would inherit these species' source identities. For example, ALD₂ reacts with the NO₃ as follows in CB6R3.

$$ALD_2 + NO_3 = C_2O_3 + HNO_3$$
 (R3)

The reaction's products, C₂O₃ and HNO₃, inherit identities equally divided between the sources of the reactants because ALD₂ and NO₃ are on the list of OP3 species. Reactions without any of these species in the reactants list, like OP2, have their products apportioned to source using OP1's methodology when the reactants are not among the weighted ones.

ISAM-OP4 lists only VOC species and daughter products instrumental in O₃ chemistry as defined in OP3. In the R1 example, the products are apportioned to the source identity of ISO₂, because the other reactant, NO, is not on the list of weight species. Similarly, the products of R3 are attributed to the source identity of ALD₂. As in options 2 and 3, reactions (such as R2) without any listed species are attributed as in OP1's method.

Finally, ISAM-OP5 was added to account for the instantaneously calculated O₃ formation regime or limiting case. The regime is determined using the ratio of PH₂O₂/PHNO₃. The transition point between regimes has a default value equal to 0.35 (Sillman, 1995). For the NO_x-limited regime (PH₂O₂/PHNO₃>0.35), source identity is passed from the nitrogen species of OP2, while for the VOC-limited regime (PH₂O₂/PHNO₃≤0.35), source identity is passed from the organics of OP4. These CMAQ-ISAM options, including the regime threshold value (or transition point), are accessible at runtime through the standard model run script.

185 Table 1. Expanded CMAQ-ISAM options.

175

CMAQ ISAM option	Reaction product source identity assignment	Representative CB6R3* Species
ISAM-OP1	Proportional to stoichiometry of all reactants.	All tracked model species
ISAM-OP2	Proportional to stoichiometry of nitrogen containing reactants, otherwise same as ISAM-OP1.	NO, NO ₂ , NO ₃ , HONO, HNO ₃ , N ₂ O ₅ , ANO ₃

ISAM-OP3	Proportional to stoichiometry of key O ₃ chemistry	NO, NO ₂ , NO ₃ , HONO, HNO ₃ ,
	reactants (reactive VOCs, radicals, nitrogen species),	N ₂ O ₅ , ANO ₃ , ALD ₂ , ALDX, FORM,
	otherwise same as ISAM-OP1.	ACET, KET, XO ₂ , XO ₂ H, ISO ₂ ,
		C_2O_3 , CXO_3
ISAM-OP4	Proportional to stoichiometry of VOC and radical	ALD ₂ , ALDX, FORM, ACET, KET,
	containing reactants, otherwise same as ISAM-OP1.	XO ₂ , XO ₂ H, ISO ₂ , C ₂ O ₃ , CXO ₃
ISAM-OP5	According to the ratio of PH ₂ O ₂ to PHNO ₃ if O ₃	NO _x -limited: NO, NO ₂ , NO ₃ ,
	chemistry reactants present, otherwise same as ISAM-	HONO, HNO ₃ , N ₂ O ₅ , ANO ₃
	OP1.	VOC-limited: ALD ₂ , ALDX,
		FORM, ACET, KET, XO ₂ , XO ₂ H,
		ISO ₂ , C ₂ O ₃ , CXO ₃

^{*}Species are based on CB6R3 and may vary based on different chemical mechanisms implemented in CMAQ. Details can be found in SA DEFN.F in the CMAQ source code.

2.2 OSAT description

190

195

200

The source apportionment approach implemented in CAMx is briefly recapped here. Detailed updates of all OSAT versions can be found in the CAMx official user guide (https://camx.com/Files/CAMxUsersGuide_v7.10.pdf). All available versions of OSAT (including OSAT3) in CAMx separately solve for production and destruction of O3 with production being attributed to either NOx or VOC emissions, depending on which is estimated to be limiting O3 production. When the ratio of PH2O2/PHNO3 exceeds 0.35, the produced O3 is attributed to NOx emissions, and VOC emissions below that threshold. The CAMx source apportionment implementation includes an option (OSAT-APCA) that allows for a redirection of attribution to anthropogenic emissions in situations where the limiting precursor is biogenic. In CAMx-OSAT, O3 attributed to NOx and VOCs is tracked as separate tracer groups. O3 tracers are first adjusted to account for O3 destruction processes and subsequently for net O3 production, which is defined as the difference between O3 production and O3 destruction based on a subset of photochemical reactions that result in O3 destruction. In situations where the net O3 production is negative (destruction reactions dominate), all the O3 tracers are proportionally decreased. When net O3 production is positive, production is assigned proportionally to the sources of those emissions (NOx and VOC precursor tracers) at the time and place

where O₃ was made. OSAT includes a group of tracers that track odd-oxygen that is consumed when O₃ reacts with NO to form NO₂ that can quickly photolyze and reform O₃ through a reaction with oxygen. In this situation, the O₃ removed from the O₃ tracers due to the NO + O₃ reaction is moved to the odd-oxygen tracers (which have separate NO_x and VOC tracer groups). When NO₂ is photolyzed and O₃ formed a proportional amount of O₃ is taken from the odd-oxygen tracers and moved to the O₃ tracers.

210 **3 Method**

3.1 Base model configurations

Two models, CMAQ version 5.3.2 with modified ISAM and CAMx version 7.10 with OSAT3, are used to simulate a one-month period during the summer of 2018 (July 29th to August 30st). The summary of the two model configurations is presented in Table 2. Both models are applied to the same 215 horizontal modeling domain with 4 km x 4 km resolution encompassing the northeastern United States. This domain is nested within a larger 12 km domain that encompasses the entire contiguous United States which is used for providing simulation boundary and initial conditions (BC and IC) for the 4 km domain. BCs were generated for the 12 km simulations using a hemispheric application of the GEOS-Chem model (Henderson et al., 2014) that was run for 2018. Anthropogenic emissions were based on 220 version 1 of the 2016 National Emission Inventory (NEI, US EPA, 2021). Electrical Generating Unit emissions were based on continuous emissions monitoring data from 2018 where available. Onroad emissions were projected to 2018 to reflect decreases in emissions due to vehicle fleet turnover and the implementation of emission control technology in 2017. The Biogenic Emission Inventory System (Bash et al., 2016) was used to generate biogenic volatile organic compound emissions, and offline 225 meteorology was created using the Weather Research and Forecasting (WRF, Skamarock et al., 2008) model version 3.8. CMAQ was configured using Carbon Bond 6 version 3 (CB6R3, Emery et al., 2015) for chemistry. Similarly, all base meteorological and emissions inputs for CAMx were identical to those for CMAQ but were processed using CAMx appropriate data pre-processors (https://www.camx.com). The CAMx model was configured with Carbon Bond 6 version 4 (CB6R4, Emery et al., 2016a) 230 chemical mechanism. It is noteworthy that the major updates for CB6R4 from CB6R3 are to (1) replace

full marine halogen chemistry with a condensed iodine mechanism called "I-16," which could reduce O_3 depletion over marine areas, and (2) add dimethyl sulfide (DMS) chemistry. Emery et al. (2016b) demonstrated that the difference in O_3 decrements between full halogen chemistry and I-16 is small and can be neglected over land.

Table 2. CMAQ and CAMx model configurations

Model option	CMAQ	CAMx
Model version	Version 5.3.2	Version 7.10
Horizontal resolution	4 km	4 km
Vertical layers	35	35
Meteorology	WRF3.8	WRF3.8
Anthropogenic Emissions	2016 NEI version 1 ^a	2016 NEI version 1 ^b
Biogenic Emissions	BEIS ^c	BEIS ^c
BC/IC	12km US CONUS	12km US CONUS
Gas phase chemistry	CB6R3	CB6R4
Source apportionment	ISAM	OSAT3

^aEGU were based on continuous emissions monitoring data from 2018 where available. Onroad emissions were projected to 2018.

240 3.2 Source apportionment simulation designs

As discussed in Section 2, ISAM has been updated to include a user option with five possible configurations for source apportionment approach. Here, we conduct CMAQ source apportionment simulations for all these options: ISAM-OP1, ISAM-OP2, ISAM-OP3, ISAM-OP4 and ISAM-OP5, hereafter referred to as OP1, OP2, OP3, OP4 and OP5. The OSAT3 approach was also used in the CAMx v7.10 base model for comparison with the five ISAM simulations. Hereafter OSAT3 is referred to as OSAT. A brute force method (zeroing out the entire emission stream for tracked sources in CMAQ, hereafter referred to as CMAQ-BF) was also used to compare with the ISAM options and OSAT. Eleven different emission source categories were tracked using each apportionment technique. The source categories comprise four point-source categories including electricity generating units (EGU), non-electricity generating units (NONEGU), fires (FIRE), and commercial marine vessels (CMV), and six area-source categories including on-road mobile (ONROAD), non-road mobile

^bCAMx EGU and Onroad were identically processed as CMAQ.

^cBELD v4.1 vegetation data for biogenic emissions, BEIS version is 3.61.

(NONROAD), biogenic (BIO), railway (RAIL), airports (AIRP), and other (AREA). Additionally, OILGAS was tracked as a mixed category (both point and area) of emissions from the oil and natural gas industry in the domain. Total emissions from the above sectors have been displayed in Table 3. 255 Finally, three predefined tracers for lateral boundary conditions (BCON), initial conditions (ICON), and other sources (OTHR) were also tracked for O₃ and its precursors. OTHR is used for all remaining untagged emission categories. In this study, all emissions sectors were tracked as previously mentioned above for OSAT and ISAM. For CMAQ-BF, a unique CMAQ simulation for each emission source category listed above was performed by fully removing the category's entire emission stream. CMAQ-BF apportionment was then calculated by subtracting the resulting pollutant fields from a base model 260 simulation. However, for ICON and BCON, each was reduced by 50%, and the output field difference with the base model was scaled up by a factor of 2 to avoid numerical issues associated with very low model ICON and BCON values. As for OTHR, there is no suitable way to retain an appropriate chemical state of the troposphere after subtracting necessary emission categories, initial and boundary 265 conditions from an original CMAQ simulation. Thus, OTHR is not being compared among CMAQ-BF, ISAM and OSAT in this study.

Table 3. Total emissions from each sector for 4km Northeast U.S. domain (month of August 2018)

	Tons/month		Percent of Total (%)	
Sector	NOx	VOC	NOx	VOC
AIRP	2,536	1,198	1.6	0.1
AREA	10,617	95,434	6.8	8.7
BIO	8,721	895,829	5.5	81.6
CMV	6,262	684	4.0	0.1
EGU	22,458	791	14.3	0.1
FIRE	400	5,007	0.3	0.5
NONEGU	15,020	11,323	9.6	1.0
NONROAD	23,958	33,561	15.2	3.1
OILGAS	11,053	22,526	7.0	2.1
ONROAD	49,361	30,578	31.4	2.8
RAIL	6,847	318	4.4	0.0
Total	157,233	1,097,247	100	100

270 **3.3 Tracked species classes**

275

280

285

O₃, NO_x and VOC species were tracked by each method. As mentioned above, ISAM tracks individual oxidized nitrogen and VOC species based on selected chemical mechanism in CMAQ, whereas OSAT tracks tracer families for each. To facilitate the comparison between the two models, the ISAM species were aggregated in the same fashion as OSAT (Table 4). However, some differences still exist since species representations between the two models are not completely the same. The nitrogen groupings NO_y and RNO_x (Table 4) were added to better elucidate the behavior of each model under different O₃ producing chemical regimes.

Table 4. Tracked species classes between ISAM and OSAT.

OSAT	ISAM
O ₃	O ₃
$RGN=NO_2+NO_3+2*N_2O_5+INO_3$	¹ RGN=NO ₂ +NO ₃ +2*N ₂ O ₅
NIT=NO+HONO	NIT=NO+HONO
TPN=PAN+PNA+PANX+OPAN+INTR	² TPN=PAN+PNA+PANX+INTR
NTR=NTR ₁ +NTR ₂ +CRON	³ NTR=NTR ₁ +NTR ₂
HNO ₃	HNO ₃
RNOx=RGN+NIT	RNOx=RGN+NIT
NOy=RGN+NIT+TPN+NTR+HNO ₃	NOy=RGN+NIT+TPN+NTR+HNO ₃
⁴ VOC=1.0*PAR+1.0*MEOH+1.0*FORM+1.0*KET+2.0*ETHA	VOC=1.0*PAR+1.0*MEOH+1.0*FORM+1.0*KET+2.0*ETH
$+2.0*ETOH + 2.0*ETH + 2.0*OLE + 2.0*ALD{\scriptstyle 2} + 2.0*ALDX + 2.0*E$	A+2.0*ETOH+2.0*ETH+2.0*OLE+2.0*ALD ₂ +2.0*ALDX+2.
THY+3.0*PRPA+3.0*ACET+4.0*IOLE+5.0*ISOP+6.0*BENZ+	0*ETHY+3.0*PRPA+3.0*ACET+4.0*IOLE+5.0*ISOP+6.0*B
7.0*TOL+8.0*XYL+10.0*TERP	ENZ+7.0*TOL+8.0*XYL+10.0*TERP

¹ISAM does not track INO₃

3.4 Evaluation method and case study development

Although identical emissions and meteorological inputs are used for CAMx and CMAQ (Table 2), potential differences still exist in multiple scales and processes. Shu et al. (2017, 2022) have reported that deposition is one of the largest uncertainties between the two models when other processes are constrained. For inter-comparing ISAM and OSAT, it is not feasible to constrain all process uncertainties. Thus, we established criteria to choose representative days for ISAM and OSAT

²ISAM does not track OPAN

³ISAM does not track CRON

⁴OSAT VOC has been pre-calculated as equation in Table 4

comparisons based on the performance of their parent models rather than comparing them throughout the entire simulation period to reduce the difference that may be brought on from their parent models. We initially set the correlation relationship (R²) criteria of maximum daily 8-hour averaged (MDA8) O₃ between CMAQ and CAMx to be above 0.7 to ensure that the performance of the two parent models is comparable. Next, MDA8 O₃ was also used as the indicator for case study selection since ISAM and OSAT normally are used as regulatory application with this metric. We assess the mean bias (MB) of MDA8 O₃ for every day to choose the days on which both models have the lowest MB for predicted MDA8 O₃. Therefore, CMAQ and CAMx simulated ambient concentrations were paired in space and time with observed data from the Air Quality System (AQS, https://www.epa.gov/aqs) monitoring network. Hourly concentrations of total O₃, NO and NO₂ were also compared to the AQS observations, and their bias statistical metrics were calculated as well.

4 Results

290

295

300

305

310

4.1 Model performance evaluation and case study selection

Figure 1 shows observed site averaged MDA8 O₃ and its corresponding biases predicted by CMAQ and CAMx over paired AQS sites for the entire episode. Observed site averaged MDA8 O₃ ranges from 30 to 50 ppbv. The performance of two models for predicting MDA8 O₃ varies by paired day and monitor site with the range of biases from -23 to 35 ppbv, approximately. Table S1 summarizes R² and MB of MDA8 O₃ for each day for both models. Based on our criteria introduced in Section 3.4, there are 13 days on which the two models show very good correlation relationships. Among these days, two models both show good performance on predicting MDA8 O₃ with closest MB on Aug 09th (CMAQ/CAMx = 3.09/2.99 ppbv) and 10th (CMAQ/CAMx: 2.42/2.61 ppbv). For other days, either two models both have higher MB (> 10 ppbv), or their predictions do not agree well with each other, with a difference of MBs up to 8 ppbv. Therefore, Aug 09th and 10th were selected as a two-day case study for source apportionment comparisons. Additional evaluations of hourly O₃, NO and NO₂ is available in Fig. S1 of the supplemental information (SI). From Fig. 2, MDA8 O₃ is relatively higher over east coastal urban areas with generally over 50 ppbv but reduces to 35 ppbv at other rural areas of northeast

315 U.S. domain. The two models predicted MDA8 O₃ show very good agreement spatially, underestimating MDA8 O₃ at sites where observed MDA8 O₃ is high but overestimating MDA8 O₃ at sites where O₃ is low. Similar spatial plots of hourly paired O₃, NO and NO₂ can be found in SI (Fig S2). Table 5 and 6 respectively summarize statistical metrics for MDA8 O₃, hourly O₃, NO and NO₂ at all paired monitoring sites for the monthly O₃ episode and the selected two-day case study episode.

The metrics in Table 5 and 6 both show consistent results with Fig. 1 as discussed above. The changes of NO and NO₂ metrics are marginal from the monthly episode to the two-day case. As in Fig S1, NO and NO₂ concentrations are less variable than O₃ across days in the monthly episode, as a result, the comparison of NO and NO₂ are less dependent on which day is selected. Unlike NO and NO₂, CAMx and CMAQ performance is statistically better in the two-day case study with lower MB for hourly O₃ (CMAQ/CAMx = 4.67/7.02 ppbv) and MDA8 O₃ (CMAQ/CAMx = 2.75/2.80) than the monthly episode (hourly O₃: CMAQ/CAMx = 6.49/7.99 ppbv; MDA8 O₃: CMAQ/CAMx = 5.30/4.18). The differences of MB, NMB and R² between the two models also diminish for MDA8 O₃ but increase for hourly O₃ from the monthly episode to the two-day episode. The statistical metrics of hourly O₃ and MDA8 O₃ demonstrate that the selected two-day case is suitable for a source apportionment comparison in which CAMx and CMAQ not only both have the least-biased predictions compared to observations but also show a good agreement with each other.

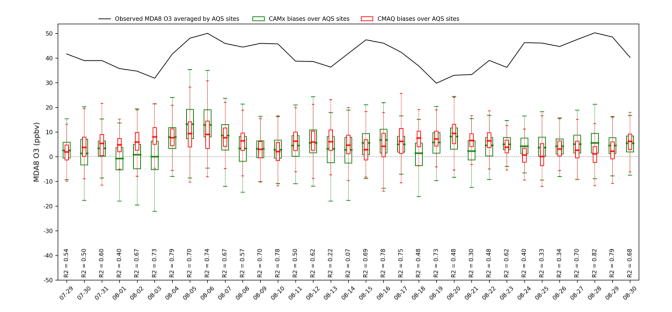


Fig.1 observed site averaged MDA8 O₃ and its corresponding biases predicted by CMAQ and CAMx over paired AQS sites for the entire episode. R² shows correlation relationship between CMAQ and CAMx.

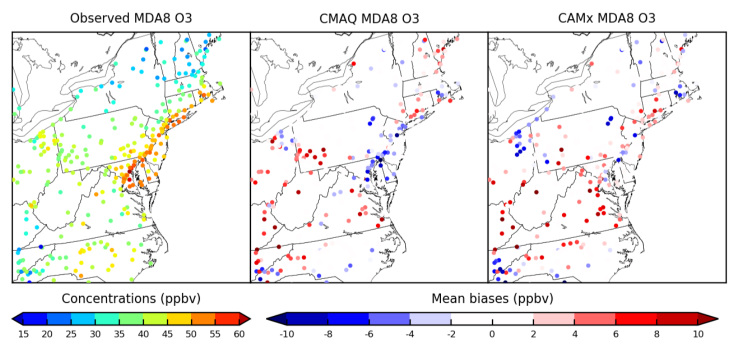


Fig.2 Two-day averaged observed MDA8 O₃ over paired sites for northeast US domain and its corresponding mean biases predicted by CMAQ and CAMx for selected case study

Table 5. Model performance summary at paired AQS surface monitoring sites. (Monthly episode)

Species	Model	Number of Observations	MB ^a	NMB ^b	RMSE ^c	dR ²
Hourly	CMAQ	72987	-1.05	-44.50	6.24	0.07
NO	CAMx	72987	-1.23	-52.25	6.39	0.05
Hourly	CMAQ	61987	0.64	10.21	6.39	0.32
NO_2	CAMx	61987	1.86	29.78	7.57	0.28
Hourly	CMAQ	232768	6.49	23.11	11.73	0.59
O_3	CAMx	232768	7.99	28.47	14.46	0.42
MDA8	CMAQ	9409	5.30	12.80	8.23	0.64
O_3	CAMx	9409	4.18	10.09	9.26	0.58

Table 6. Model performance summary at paired AQS surface monitoring sites. (Two-day case study episode)

Species	Model	Number of Observations	MB	NMB	RMSE	\mathbb{R}^2
Hourly	CMAQ	4264	-1.15	-48.30	6.44	0.05
NO	CAMx	4264	-1.38	-58.14	6.57	0.04
Hourly	CMAQ	3612	0.15	2.20	6.83	0.28
NO_2	CAMx	3612	0.83	11.88	7.51	0.25
Hourly	CMAQ	13486	4.67	15.06	10.88	0.61
O_3	CAMx	13486	7.02	22.65	13.26	0.49
MDA8	CMAQ	567	2.75	6.00	6.28	0.62
O_3	CAMx	567	2.80	6.10	6.95	0.63

^a Mean bias: $MB = \frac{1}{N} \sum M_i - O_i$, MB ranges from negative infinity to positive infinity with 0 indicating unbiased data, unit here is ppbv.

c Root mean square error: RMSE = $\sqrt{\frac{1}{n}} \sum_{i=1}^{n} (M_i - O_i)^2$, is the standard deviation of the prediction errors. d $R^2 = \left\{ \frac{\sum (O_i - \bar{O})(M_i - \bar{M})}{\sqrt{\sum (O_i - \bar{O})^2 \sum (M_i - \bar{M})^2}} \right\}^2$, R^2 ranges from 0 to 1 with 1 indicating perfect correlation and 0 indicating an uncorrelated relationship.

4.2 Comparison of model source apportionment

4.2.1 Temporal variations of sector contributions

355 To better understand how the ISAM model apportionment approach simulated source contributions at each time step, time-series comparisons for each source were examined for O₃ and its

^b Normalized mean bias: $NMB = \frac{1}{N} \sum_{i=0}^{M_i - O_i}$, ranges from negative 1 to positive infinity with 0 indicating unbiased data. The values shown in the table were multiplied by 100.

precursors, RNO_x and VOC for the two-day case study. Figure 3 shows hourly variations of domain averaged predicted total O₃ (bulk) concentrations and sector contributions for seven source apportionment simulations (OSAT, BF, ISAM OP1 to OP5). In Fig. 3, CMAO and CAMx predict similar O₃ concentrations during the day, but differences appear at night, with a maximum difference of 5 ppb. This disparity was discussed in Section 4.1 and can be mitigated by employing the MDA8 O₃ metric. The seven source apportionment simulations yield similar diurnal trends via the trajectory of the total concentrations, but they apportion concentrations to each sector somewhat differently. Comparisons of five ISAM options reveals significant variability. OP1, which apportions uniformly according to stoichiometry, shows similar trends of apportionments for each sector as OP4, an option that always allocates products to sources with reactive VOCs and their radicals. They both apportion more BCON and BIO O₃ but fewer contributions from all other sectors than the other three ISAM options (OP2, OP3 and OP5). Results of OP1 and OP4 would likely overestimate sensitivity to emissions to these reactants because VOCs are often available in excess. OP2 always allocates products to sources with nitrogen reactants, which prevents the attribution of NO_x to non-nitrogen reactants. Typically, these non-nitrogen reactants are common in transported (e.g., BCON) or natural sources (e.g., isoprene in BIO). As a result, OP2 decreases BCON and BIO contributions while increasing contributions from other sectors relative to OP1 and OP4.

360

365

370

OP5 assigns products to either reactive VOCs or NO_x based on the ratio of PH₂O₂/PHNO₃,

375 placing O₃ contribution results for all sectors between the previous four ISAM options. OSAT, which
utilizes a similar methodology as OP5, shows consistent diurnal patterns of domain averaged total O₃
and sector contributions as the ISAM options, but with varying magnitudes. OSAT has the largest
BCON O₃ but the least contributions from AREA, BIO and FIRE. The rest of the OSAT sector
contributions are between the ISAM options. Consistent with earlier findings, CMAQ-BF estimates

380 systematically smaller O₃ contributions for all sectors besides EGU and BCON (Kwok et al., 2015).
While ISAM and OSAT appear to retain bulk mass as intended, CMAQ-BF shifts the chemical system
into a different nonlinear O₃ response to source change.

In Fig. 4, CAMx and CMAQ predict comparable total RNO_x except for the first 12 hours of the two-day example, when OSAT values deviate from those of the other six simulations. As the total

concentrations of the two models converge, OSAT exhibits similar patterns to OP2 and OP3. OP1, OP4 and OP5 show comparable results, with increased BCON and BIO RNO_x but decreased contributions from other sectors. CMAQ-BF show comparable results with OSAT, OP2 and OP3 except for BCON and BIO, which are negative for CMAQ-BF, suggesting that removing these source sectors results in a slight rise in RNO_x. In previous source sensitivity and allocation investigations, it has been shown that
BF may have limits when the model response contains an indirect effect coming from the influence of substances other than the direct precursors (Kwok et al., 2015; Burr and Zhang, 2011; Koo et al., 2009; Jimenez and Baldasano, 2004; Zhang et al., 2009). This would be particularly true in situations where emissions are a large percentage of total NO_x or VOC in a particular area. The nonlinear impacts on gas phase chemistry realized in a source sensitivity model simulation would not be a relevant representation of culpability from that same source group.

Figure 5 illustrates the hourly variability of domain-averaged VOC concentrations and sector contributions. CAMx only gives pre-lumped VOC (Table 4) for OSAT outputs. For consistency, VOC for CMAQ ISAM and BF has also been carbon-weighted by summing all individual VOC species in CMAQ outputs using the same method as OSAT (Table 4). In Fig. 5, CAMx consistently simulates higher attribution to total VOC concentrations than CMAQ, with a maximum difference of 30 ppb. These larger CAMx VOC concentrations are also reflected in apportioned OSAT sectors, particularly those with substantial contributions, such as BCON and BIO. Given that the difference is present in the total concentration, this is unlikely caused by different source apportionment formulation between CMAQ and CAMx. As CAMx only gives pre-lumped VOC, it is challenging to compare individual VOC species between CMAO and CAMx to explain this difference at current stage. Another possible reasons to cause it could be that models have different internal treatments for advection and diffusion, which can impact surface-level concentrations and indirectly impact chemical reactions. The five ISAM options have comparable diurnal patterns for most sectors, with the exception of CMV, EGU, and RAIL, however the magnitudes for these three sectors are relatively minor, which is consistent with earlier findings (Kwok et al., 2015). CMAQ-BF estimates notably lower sector contributions for VOCs, which is similar to O_3 results (Fig. 4), with negative contributions for small sectors (e.g., CMV, EGU,

400

405

and RAIL). Additional figures of other grouped nitrogen species tracked in Table 4 (e.g., RGN, HNO₃ and NO_y) can be found in SI.

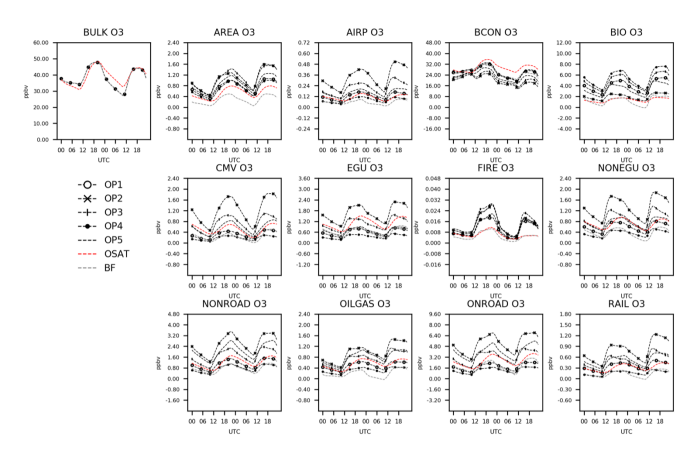


Fig.3 Total and attributed O₃ concentrations to various sectors as a function of hour of day and apportionment technique.

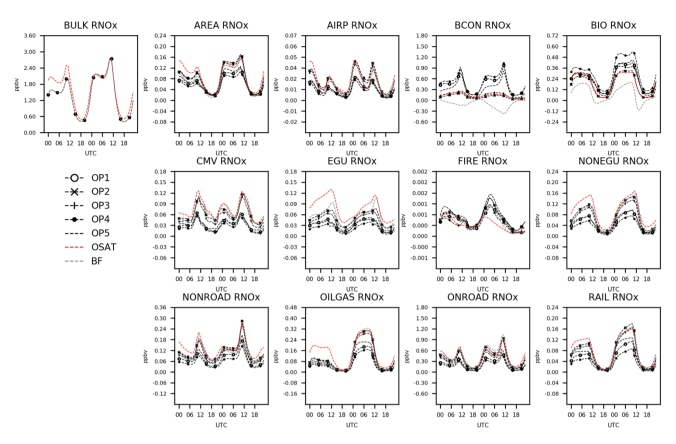


Fig. 4 Total and attributed RNO_x concentrations to various sectors as a function of hour of day and apportionment technique.

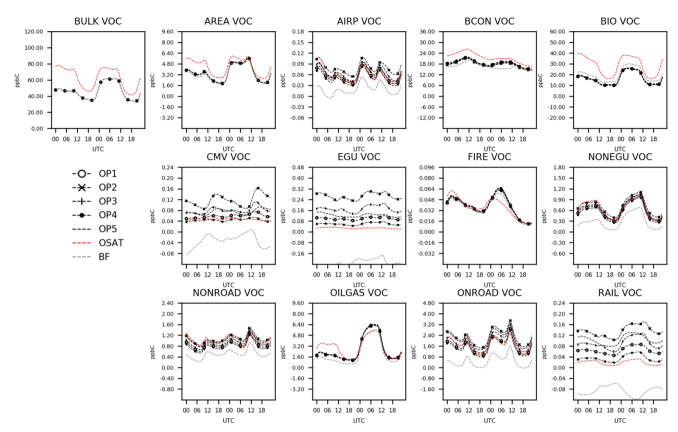


Fig. 5 Total and attributed VOC concentrations to various sectors as a function of hour of day and apportionment technique.

4.2.2 Spatial distribution of source apportionment simulations

420

425

430

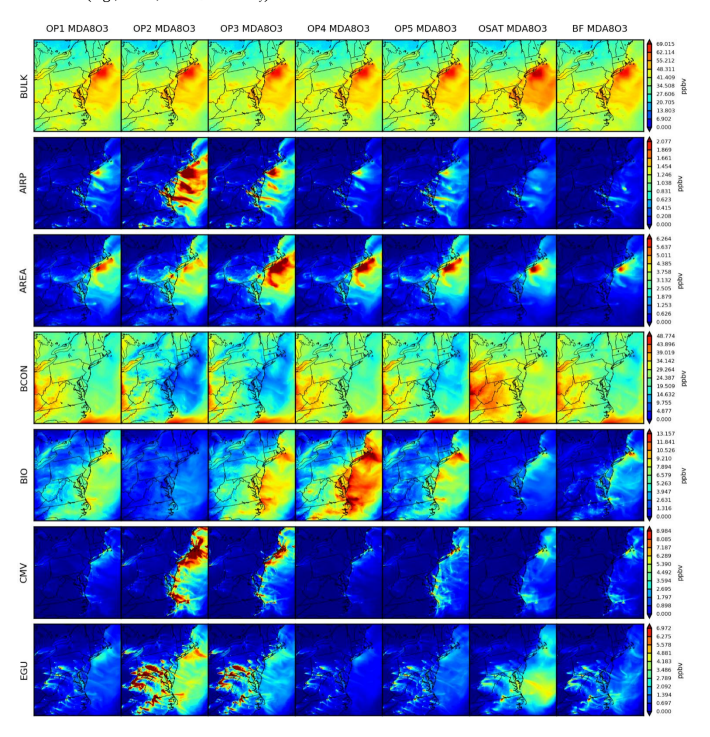
Spatial patterns of total and sector contributions of MDA8 O₃ (Fig. 6), RNO_x (Fig. 7) and VOC (Fig. 8) have been examined for the seven simulations. In Fig. 6, OSAT exhibits the same spatial distribution of MDA8 O₃ total concentrations as other CMAQ-based simulations (OP1, OP2, OP3, OP4, OP5, and CMAQ-BF), with the exception of OSAT's relatively high marine and offshore total concentrations (> 5 ppbv), which could be explained by the difference in planetary boundary layer dynamics or different marine chemistry configuration between the two parent models. CMAQ CB6R3 uses a rough parameterization for full marine halogen chemistry to destroy O₃, depending only on landuse category and sunlight (Sarwar et al., 2015, 2019), whereas CAMx CB6R4 handles O₃ depletion in the marine boundary more efficiently by including the 16 most important reactions of inorganic iodine (I-16b, Emery et al., 2016b). According to a sensitivity test conducted by Emery et al. (2016b), I-16b could reduce O₃ depletions by 2-5 ppbv in comparison to full halogen chemistry. Regarding sector

concentrations, the spatial distributions of seven simulations are comparable. They can all capture geographic contribution hot spots from each sector, although their magnitudes vary. OP2 stands out with fewer contributions from BIO than the other four ISAM options, and subsequently assigns larger concentrations to other sectors, particularly over east coastal regions, as shown in Fig. 3 and 6. Since OP2 assigns all products to sources with nitrogen reactants, the influence of reactants from biogenic sources is diminished, as intended.

440 Figure 7 depicts the associated outcomes of RNO_x. Except for BCON, the seven simulations produce geographically and quantitatively consistent findings. From the spatial distributions, we can conclude that local sources govern RNO_x more than long-transported sources compared to O₃. Anthropogenic RNO_x is either more concentrated in the urban areas (e.g., AREA, NONEGU, NONROAD), gasoline industry (OILGAS) and electric facilities (EGU) or along with transportation (e.g., AIRP, ONROAD, CMV and RAIL). Biogenic RNO_x is more prevalent in rural locations with 445 vegetation. It should be noted that OP1, OP4 and OP5 show more BCON RNO_x across the entire domain because of the way to assign products in nitrogen related reactions (Section 2). OP1, OP4 and OP5 show local hotspots of RNO_x attributed to BCON. Since there is no physical reason to suspect hotspots over urban areas, we conclude that these contributions represent RNO_x attributed based on 450 VOC or oxidants transported from the boundary. Figure 8 depicts the outcomes associated with VOC. Higher VOC concentrations from CAMx already shown in Fig. 5 are primarily from Virginia and North Carolina (OSAT bulk). As CMAQ and CAMx both use the same BEIS inventory data, the difference in total VOC concentrations may result from other differences between two models, like chemistry or deposition, accordingly, leading to higher biogenic sources in CAMx (BIO). For the rest of sectors, 455 OSAT and ISAM options are fairly consistent except that the OP2 predicts more contributions from EGU, CMV and RAIL. CMAQ-BF predicts consistently lower source contributions for MDA8 O₃, RNO_x, and VOC, as shown in Section 4.2.1. This yet again illustrates that brute force represents an integrated sensitivity while the OSAT and ISAM represent attribution at a point in the nonlinear chemical systems. Monthly averaged spatial maps for MDA8 O₃, RNO_x, and VOC are also included in 460 Fig. S4(a-c) and show consistent results as two-day averaged maps. This demonstrates that our case

study is appropriate, efficiently selecting representative days as well as minimizing the uncertainties

from parent models (CMAQ and CMAQ). Additional figures of other grouped nitrogen species tracked in Table 4 (e.g., RGN, HNO₃ and NO_y) can also be found in SI.



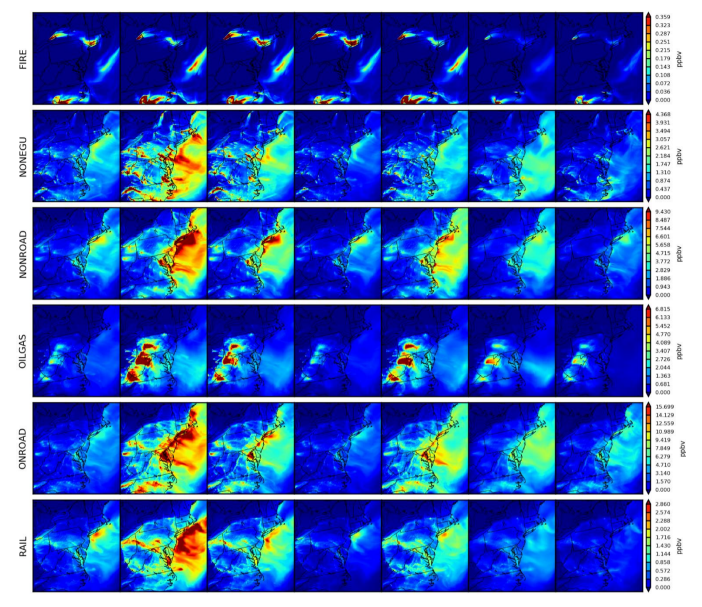
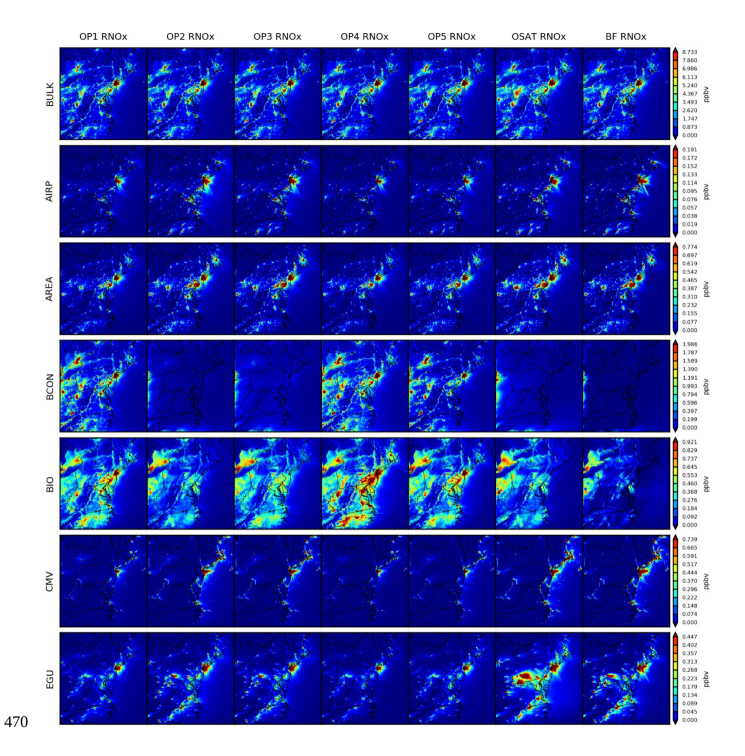


Fig. 6 Spatial comparisons of seven simulations for two-day averaged O_3 (08/09 and 08/10).



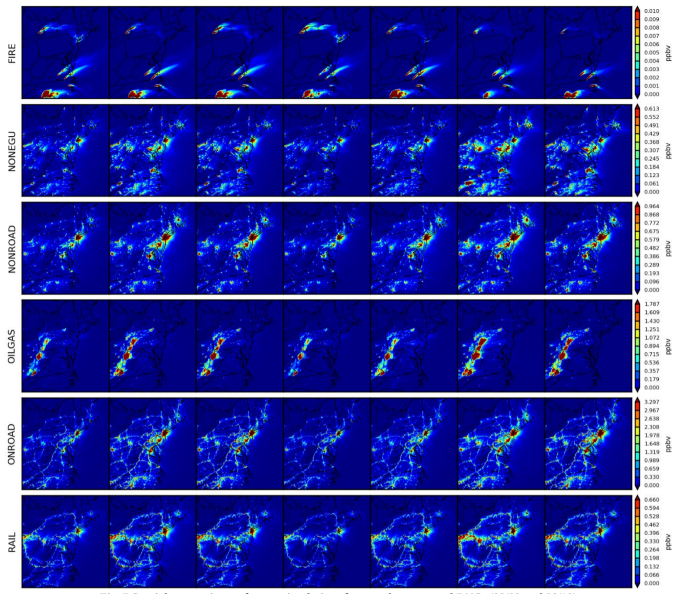
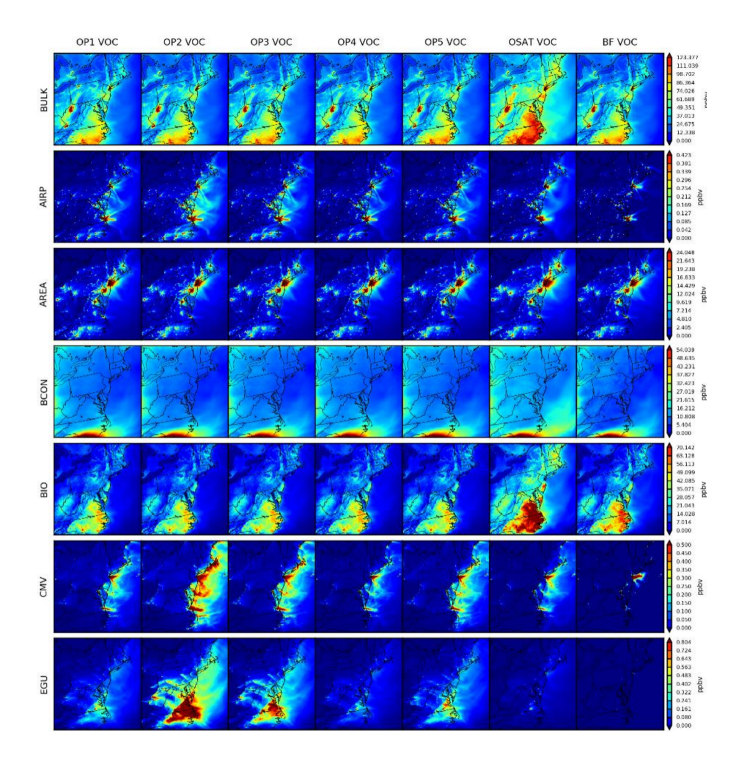


Fig. 7 Spatial comparisons of seven simulations for two-day averaged RNO $_{x}$ (08/09 and 08/10).



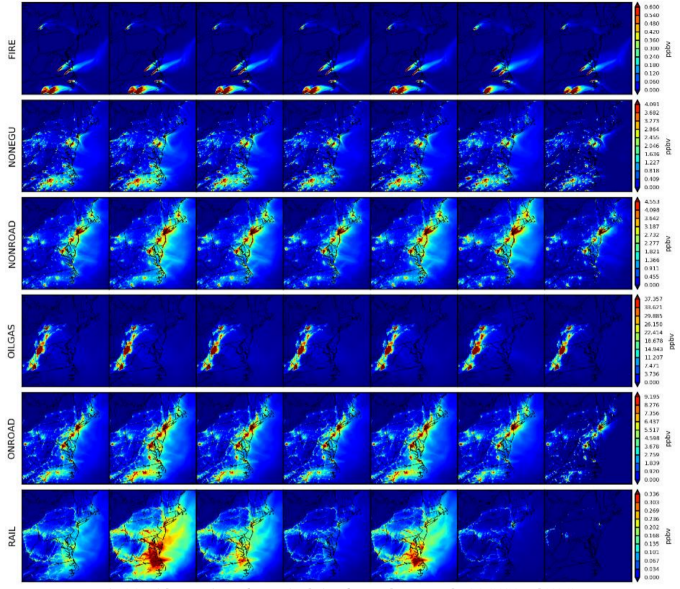


Fig. 8 Spatial comparisons of seven simulations for two-day averaged VOC (08/09 and 08/10).

5 Model Simulation Time

475

480

The CPU time required to complete a source apportionment simulation in a 3D AQM is an important consideration for usability. For a 4 km x 4 km simulation domain encompassing the northeast U.S., the model run times for OSAT and ISAM are similar. Using 128 processors, base CMAQ (without ISAM) and CMAQ-ISAM simulations (11 source categories) are tested. Base CMAQ requires around

60 minutes per simulation day (24 hours), whereas CMAQ-ISAM requires approximately 120 minutes. If the number of processors is increased to 256, the simulation time for CMAO-ISAM can be reduced to 30 minutes showing good scalability. It is worth noting that our CMAO-ISAM simulations simultaneously track all additional species classes, such as sulfate, nitrate, ammonium, elemental carbon, organic carbon, and chloride. It would shorten simulation times if related species were only tracked for O₃ Base CAMx (without OSAT) and CAMx OSAT are also tested with 128 processors, taking 37 and 67 minutes, respectively. CAMx also provides an optional tool for particles that can be simultaneously applied similarly to ISAM (PSAT, Yarwood et al., 2007). When additional pollutants are selected for tracking (e.g., sulfate, primary PM_{2.5} species, etc.) total simulation time will increase for 490 both ISAM and OSAT/PSAT. CMAQ-BF speed is based on CMAQ base simulation (60 mins/day x (1 base + 11 sources + 1 boundary condition + 1 initial condition + 1 other) = 900 mins/day.

6 Discussions and Conclusions

485

495

Source attribution approaches are generally intended to determine culpability of precursor emissions sources to ambient pollutant concentrations. Source-based apportionment approaches such as ISAM and OSAT provide similar types of information, specifically an estimate of which sources or groups of sectors (e.g., a sector) contributed to the air quality measured or estimated at a particular location. The assumptions in each technique have implications for interpretation in the context of air quality management.

Source attribution of secondarily formed pollutants cannot be explicitly measured, which makes evaluation of source apportionment approaches challenging. Here, the ISAM approach was evaluated 500 by 1) a comparison with a source apportionment approach implemented in a different photochemical modeling system and 2) a comparison with a simple source sensitivity (brute-force difference) approach in the same modeling system that is most comparable to source apportionment in more linear systems and less useful when formation and transport is nonlinear. Further, this section notes qualitative 505 consistency between the spatial nature of sector emission and the attribution of precursors and O₃ as another method to generate confidence in these approaches.

In this study, multiple apportionment approach comparisons show common features but still reveal wide variations in predicted sector contribution and species dependency. The attribution to sources emitting NO_x and VOC is consistent with the spatial nature of these sources, which provides confidence in the approach. However, nitrogen species (e.g., NO_x), for instance, are more sensitive to the choice of ISAM options than VOC. For example, although the attribution of NO_x to EGUs matches the location of these sources (e.g., New York urban area) for all ISAM options, OP1, OP4 and OP5 predict more BCON NO_x. This is because the fast NO_x cycling process assigns anthropogenically emitted nitrogen species to other sources, as the original emitted source identity is not retained through these complex reactions. Further, sources entirely located offshore, such as commercial marine vessels, do not have culpability assigned to distant inland regions of the model domain. Most of the time, the amount of attribution to a certain sector depends on the number of emissions from that sector, how far away those emissions are, and whether the prevailing winds carried emissions from those places to the monitor or grid cell where air quality was predicted.

The designed five ISAM options maximize its flexibility, particularly for modeling source apportionment of O₃ and its precursors, but the choice of option depends on target species. Among all ISAM options, the OP5 option, after making the assignment decision based on the ratio of PH₂O₂ to PHNO₃, is expected to predict generally similar spatial and temporal patterns for O₃ to the OSAT source apportionment approach implemented in CAMx. However, it still shows disparity for some sectors (e.g., biogenic sectors for O₃). This result may be because of the OSAT formulation which differs from the ISAM options presented here. The OP5 option was also similar to brute-force sensitivity estimates predicted in CMAQ with the exception of source groups that dominate regional emissions or O₃, such as biogenic VOC and O₃ introduced into the model through boundary inflow. In those situations, it is not reasonable to expect a source sensitivity approach to provide a useful comparison for source attribution given the highly nonlinear change in atmospheric chemistry. After assigning products to sources emitting nitrogen reactants, the OP2 option can predict results of RNO_x attributions that are more comparable to OSAT and BF. It demonstrated that the OP2 works better for RNO_x because it makes it easier to find the original source and lessens the effect of other sources when these species are cycling quickly through an integrated chemical reaction system. Unlike O₃ and RNO_x, the VOC contribution for

the majority of source categories depends very little on the ISAM option. We expect that the user will use OP5 for O₃ and OP2 for RNO_x, but this is not a firm suggestion. In turn, we give the user this flexibility so that ISAM can be used for a wide range of purposes.

By comparing the multiple approaches in the Northeast U.S., we found that both OSAT and ISAM attribute the majority of O₃ and NO_x contributions to boundary, mobile, and biogenic sources, whereas the top three VOC contributions are attributed to biogenic, boundary, and area sources. However, comparisons of OSAT and ISAM have some limits, especially when they are under the two different parent models, CAMx and CMAQ. Although we have put efforts into diminishing the differences between the two models by making most configuration options as similar as possible, some inevitable uncertainties cannot be eliminated at the current stage of this study (e.g., an imperfect match of chemical mechanisms, different internal treatments for advection, diffusion, and deposition processes). Further, it is also worthwhile to note that our results in this study are based on limited duration and specific regions, and they may not comprehensively reflect all situations. Given that the source attribution of secondary pollutants cannot be explicitly measured, these inter-comparisons between ISAM and OSAT are still useful for reference. We continue to need further efforts that combine field experiment studies and model evaluations for longer terms and multiple regions to better understand source attribution given the highly nonlinear change in nature of O₃-NO_x chemistry.

Code availability

540

545

550

555

The updated ISAM code used in this study has been permanently archived at https://doi.org/10.5281/zenodo.6266674 and has also been implemented in the newer version of CMAQ (v5.4). The CMAQ model documentation is available at https://github.com/USEPA/CMAQ and www.cmaq-model.org. Model post-processing scripts, are available upon request.

Data availability

560

565

The raw observation data used are available from the sources identified in Sect. 3, while the post-processed observation data are available upon request. The CMAQ model data utilized are available upon request as well. Please contact the corresponding author to request any data related to this work.

Author contributions

QS, SN, KB designed this study and experiments. QS led the development of this manuscript and was responsible for most of the model evaluation components in this study. SN, WH and BM developed the ISAM code. KB provided all the input data for the CMAQ simulations. QS carried out the CMAQ preprocessing, simulations, and post-processing, produced the figures, and prepared the initial paper draft. SN contributed directly to the writing of Sect. 2 of this paper. KB contributed directly to the writing of Sect. 6 of this paper. WH, BM, CH and BH discussed all results throughout the ISAM development and contributed to the final writing of this paper.

Competing interests

570 The authors declare that they have no conflict of interests.

Disclaimer

The views expressed in this article are those of the authors and do not necessarily represent the views or policies of the U.S. Environmental Protection Agency.

Acknowledgments

This project was supported in part by an appointment to the Research Participation Program at the Office of Research and Development, US Environmental Protection Agency, administered by the Oak Ridge Institute for Science and Education through an interagency agreement between the US Department of Energy and the EPA.

References

- 580 Atkinson, R.: Atmospheric chemistry of VOCs and NOx, Atmospheric Environment, 34, 2063–2101, https://doi.org/10.1016/S1352-2310(99)00460-4, 2000.
 - Bash JO, Baker KR, Beaver MR, 2016. Evaluation of improved land use and canopy representation in BEIS v3. 61 with biogenic VOC measurements in California. Geoscientific Model Development 9, 2191.
- Baker, K. R., M. C. Woody, G. S. Tonnesen, W. Hutzell, H. O. T. Pye, M. R. Beaver, G. Pouliot, and T. Pierce. "Contribution of regional-scale fire events to ozone and PM2. 5 air quality estimated by photochemical modeling approaches." Atmospheric Environment 140 (2016): 539-554.
- Booker, F., Muntifering, R., McGrath, M., Burkey, K., Decoteau, D., Fiscus, E., Manning, W., Krupa, S., Chappelka, A., and Grantz, D.: The Ozone Component of Global Change: Potential Effects on Agricultural and Horticultural Plant Yield, Product Quality and Interactions with Invasive Species, Journal of Integrative Plant Biology, 51, 337–351, https://doi.org/10.1111/j.1744-7909.2008.00805.x, 2009.
 - Burr, Michael J., and Yang Zhang. "Source apportionment of fine particulate matter over the Eastern US Part I: source sensitivity simulations using CMAQ with the Brute Force method." Atmospheric Pollution Research 2, no. 3 (2011): 300-317.
 - Butler, T., Lupascu, A., Coates, J., and Zhu, S.: TOAST 1.0: Tropospheric Ozone Attribution of Sources with Tagging for CESM 1.2.2, Geosci. Model Dev., 11, 2825–2840, https://doi.org/10.5194/gmd-11-2825-2018, 2018.
- Cohan, D. S. and Napelenok, S. L.: Air Quality Response Modeling for Decision Support, Atmosphere, 2, 407–425, https://doi.org/10.3390/atmos2030407, 2011.
 - Cooper, O. R., Langford, A. O., Parrish, D. D., and Fahey, D. W.: Challenges of a lowered U.S. ozone standard, Science, 348, 1096–1097, https://doi.org/10.1126/science.aaa5748, 2015.
- Duncan, B. N., Yoshida, Y., de Foy, B., Lamsal, L. N., Streets, D. G., Lu, Z., Pickering, K. E., and Krotkov, N. A.: The observed response of Ozone Monitoring Instrument (OMI) NO2 columns to NOx emission controls on power plants in the United States: 2005–2011, Atmospheric Environment, 81, 102–111, https://doi.org/10.1016/j.atmosenv.2013.08.068, 2013.
 - Dunker, A. M., Yarwood, G., Ortmann, J. P., and Wilson, G. M.: Comparison of Source Apportionment and Source Sensitivity of Ozone in a Three-Dimensional Air Quality Model, Environ. Sci. Technol., 36, 2953–2964, https://doi.org/10.1021/es011418f, 2002.
- Emery, C., Jung J, Koo, B., Yarwood, G. 2015. Improvements to CAMx Snow Cover Treatments and Carbon Bond Chemical Mechanism for Winter Ozone. Final report for Utah Department of Environmental Quality, Division of Air Quality, Salt Lake City, UT, August 2015, available at http://www.camx.com/files/udaq_snowchem_final_6aug15.pdf (last accessed 13 December 2019).
- Emery, C., Koo, B., Hsieh, W. C., Wetland, A., Wilson, G., Yarwood, G. 2016(a) Technical Memorandum for Updated Carbon Bond Chemical Mechanism. EPA Contract EPD12044, available at https://www.camx.com/files/emaq4-07_task7_techmemo_r1_laug16.pdf (last accessed 1 February 2023).

- Emery, C., Liu, Z., Koo, B., Yarwood, G. 2016(b). Improved Halogen Chemistry for CAMx Modeling. Final report for Texas Commission on Environmental Quality WO 582-16-61842-13, May 2016, available at https://www.tceq.texas.gov/airquality/airmod/project/pj_report_pm.html (last accessed 13 December 2019).
 - Gillani, N. V., & Pleim, J. E. (1996). Sub-grid-scale features of anthropogenic emissions of NOx and VOC in the context of regional Eulerian models. *Atmospheric Environment*, *30*(12), 2043-2059.
- Grewe, V., Tsati, E., and Hoor, P.: On the attribution of contributions of atmospheric trace gases to emissions in atmospheric model applications, Geosci. Model Dev., 3, 487–499, https://doi.org/10.5194/gmd-3-487-2010, 2010.
 - Henderson, B. H., F. Akhtar, H. O. T. Pye, S. L. Napelenok, and W. T. Hutzell. "A database and tool for boundary conditions for regional air quality modeling: description and evaluation." Geoscientific Model Development 7, no. 1 (2014): 339-360.
 - Hidy, G. M., Friedlander, S. K., "The Nature of the Los Angeles Aerosol," p 391 in "Proceedings of the Second International Clean Air Congress" Academic Press, London, 1971.
 - Jacob, D. J. and Winner, D. A.: Effect of climate change on air quality, Atmospheric Environment, 43, 51–63, https://doi.org/10.1016/j.atmosenv.2008.09.051, 2009.

- Jacquemin, B. and Noilhan, J.: Sensitivity study and validation of a land surface parameterization using the HAPEX-MOBILHY data set, Boundary-Layer Meteorol, 52, 93–134, https://doi.org/10.1007/BF00123180, 1990.
- Jiménez, Pedro, and José M. Baldasano. "Ozone response to precursor controls in very complex terrains: Use of photochemical indicators to assess O3-NOx-VOC sensitivity in the northeastern Iberian Peninsula." Journal of Geophysical Research: Atmospheres 109, no. D20 (2004). Karamchandani, P.,
- Long, Y., Pirovano, G., Balzarini, A., and Yarwood, G.: Source-sector contributions to European ozone and fine PM in 2010 using AQMEII modeling data, Atmos. Chem. Phys., 17, 5643–5664, https://doi.org/10.5194/acp-17-5643-2017, 2017.
- Koo, B., Wilson, G. M., Morris, R. E., Dunker, A. M., and Yarwood, G.: Comparison of Source Apportionment and Sensitivity Analysis in a Particulate Matter Air Quality Model, Environ. Sci. Technol., 43, 6669–6675, https://doi.org/10.1021/es9008129, 2009.
 - Kwok, R.H.F., Baker, K.R., Napelenok, S.L. and Tonnesen, G.S., 2015. Photochemical grid model implementation and application of VOC, NO x, and O 3 source apportionment. Geoscientific Model Development, 8(1), pp.99-114.
- Kwok, R. H. F., Napelenok, S. L., and Baker, K. R.: Implementation and evaluation of PM2.5 source contribution analysis in a photochemical model, Atmospheric Environment, 80, 398–407, https://doi.org/10.1016/j.atmosenv.2013.08.017, 2013.
 - Lamsal, L. N., Duncan, B. N., Yoshida, Y., Krotkov, N. A., Pickering, K. E., Streets, D. G., and Lu, Z.: U.S. NO2 trends (2005–2013): EPA Air Quality System (AQS) data versus improved
- observations from the Ozone Monitoring Instrument (OMI), Atmospheric Environment, 110, 130–143, https://doi.org/10.1016/j.atmosenv.2015.03.055, 2015.
 - Lefohn A. S., Shadwick D. S. and Ziman S. D., 1998. The Difficult Challenge of Attaining EPA's New Ozone Standard. Environmental Science & Technology. 32(11):276A-282A.

- Li, Ying, AK-H. Lau, JC-H. Fung, J. Y. Zheng, L. J. Zhong, and Peter Kwok Keung Louie. "Ozone source apportionment (OSAT) to differentiate local regional and super-regional source contributions in the Pearl River Delta region, China." Journal of Geophysical Research: Atmospheres 117, no. D15 (2012).
- Marmur, A., Unal, A., Mulholland, J. A., and Russell, A. G.: Optimization-Based Source
 Apportionment of PM _{2.5} Incorporating Gas-to-Particle Ratios, Environ. Sci. Technol., 39, 3245–3254, https://doi.org/10.1021/es0490121, 2005.
 - Paatero, Pentti, and Unto Tapper. "Positive matrix factorization: A non-negative factor model with optimal utilization of error estimates of data values." Environmetrics 5.2 (1994): 111-126.
- Pay, M. T., Gangoiti, G., Guevara, M., Napelenok, S., Querol, X., Jorba, O., and Pérez García-670 Pando, C.: Ozone source apportionment during peak summer events over southwestern Europe, Atmos. Chem. Phys., 19, 5467–5494, https://doi.org/10.5194/acp-19-5467-2019, 2019.
 - Ramboll Environ. Final Report "Improved OSAT, APCA and PSAT Algorithms for CAMx". Contract. 2015 Aug; 582:15-50417.
- Ramboll Environ. Final Report "Implementation of the Piecewise Parabolic Method for Vertical Advection in Comprehensive Air Quality Model with Extensions (CAMx)". Contract. 2022 June; 582-19-90500.
 - Reitze Jr AW. Air Quality Protection Using State Implementation Plans-Thirty-Seven Years of Increasing Complexity. Vill. Envtl. LJ. 2004; 15:209.
- Sarwar, G., Gantt, B., Foley, K., Fahey, K., Spero, T. L., Kang, D., Mathur, R., Foroutan, H., Kang, J., Sherwen, T., and Saiz-Lopez, A.: Influence of bromine and iodine chemistry on annual,
- seasonal, diurnal, and background ozone: CMAQ simulations over the Northern Hemisphere, Atmospheric Environment, 213, 395–404, https://doi.org/10.1016/j.atmosenv.2019.06.020, 2019.

- Sarwar, G., Gantt, B., Schwede, D., Foley, K., Mathur, R., and Saiz-Lopez, A.: Impact of Enhanced Ozone Deposition and Halogen Chemistry on Tropospheric Ozone over the Northern Hemisphere, Environ. Sci. Technol., 49, 9203–9211, https://doi.org/10.1021/acs.est.5b01657, 2015.
- Shu, L., Wang, T., Han, H., Xie, M., Chen, P., Li, M., and Wu, H.: Summertime ozone pollution in the Yangtze River Delta of eastern China during 2013–2017: Synoptic impacts and source apportionment, Environmental Pollution, 257, 113631, https://doi.org/10.1016/j.envpol.2019.113631, 2020.
- Shu, Q., Koo, B., Yarwood, G., and Henderson, B. H.: Strong influence of deposition and vertical mixing on secondary organic aerosol concentrations in CMAQ and CAMx, Atmospheric Environment, 171, 317–329, https://doi.org/10.1016/j.atmosenv.2017.10.035, 2017.
- Shu, Q., Murphy, B., Schwede, D., Henderson, B. H., Pye, H. O., Appel, K. W., ... & Perlinger, J. A. (2022). Improving the particle dry deposition scheme in the CMAQ photochemical modeling system. Atmospheric Environment, 289, 119343.
 - Sillman, Sanford. "The use of NO y, H2O2, and HNO3 as indicators for ozone-NO x-hydrocarbon sensitivity in urban locations." Journal of Geophysical Research: Atmospheres 100, no. D7 (1995): 14175-14188.
- Simon, H., Reff, A., Wells, B., Xing, J., and Frank, N.: Ozone Trends Across the United States over a Period of Decreasing NOx and VOC Emissions, Environ. Sci. Technol., 49, 186–195, https://doi.org/10.1021/es504514z, 2015.

- Skamarock, W., Klemp, J., Dudhia, J., Gill, D., Barker, D., Wang, W., Huang, X.-Y., and Duda, M.: A Description of the Advanced Research WRF Version 3, UCAR/NCAR, https://doi.org/10.5065/D68S4MVH, 2008.
- Smith, W.P., Nicholls, M.E. and Pielke Sr, R.A., 2020. The role of radiation in accelerating tropical cyclogenesis in idealized simulations. Journal of the Atmospheric Sciences, 77(4), pp.1261-1277.
 - T. Pierce and L. Bender, Examining the Temporal Variability of Ammonia and Nitric Oxide Emissions from Agricultural Processes Proceedings of the Air and Waste Management Association/U.S. Environmental Protection Agency Emission Inventory Conference, Paleigh October 26, 28, 1999.
- 710 Environmental Protection Agency Emission Inventory Conference, Raleigh October 26-28, 1999, Raleigh NC.
 - U.S. EPA (2019). CMAQ Zenodo. https://doi.org/10.5281/zenodo.3585898.
 - U.S. EPA (2021). https://www.epa.gov/air-emissions-modeling/2016-version-1-technical-support-document.
 - U.S. EPA (2022a). CMAO Zenodo. https://doi.org/10.5281/zenodo.7218076.

- U.S. EPA (2022b). CMAQ Zenodo. https://doi.org/10.5281/zenodo.6266674.
- Valverde, V., Pay, M. T., and Baldasano, J. M.: Ozone attributed to Madrid and Barcelona onroad transport emissions: Characterization of plume dynamics over the Iberian Peninsula, Science of The Total Environment, 543, 670–682, https://doi.org/10.1016/j.scitotenv.2015.11.070, 2016.
- World Health Organization. Global tuberculosis report 2013. World Health Organization; 2013. Watson, John G., John A. Cooper, and James J. Huntzicker. "The effective variance weighting for least squares calculations applied to the mass balance receptor model." Atmospheric Environment (1967) 18.7 (1984): 1347-1355.
- Yarwood G, Morris RE, Wilson GM. Particulate matter source apportionment technology (PSAT) in the CAMx photochemical grid model. In Air Pollution Modeling and Its Application XVII 2007 (pp. 478-492). Springer, Boston, MA.
 - Yienger, J. J. and Levy, H.: Empirical model of global soil-biogenic NO $_\chi$ emissions, J. Geophys. Res., 100, 11447, https://doi.org/10.1029/95JD00370, 1995.
- Zhang, L., Jacob, D. J., Kopacz, M., Henze, D. K., Singh, K., and Jaffe, D. A.: Intercontinental source attribution of ozone pollution at western U.S. sites using an adjoint method, Geophys. Res. Lett., 36, L11810, https://doi.org/10.1029/2009GL037950, 2009.
 - Zhang, R., Cohan, A., Pour Biazar, A., and Cohan, D. S.: Source apportionment of biogenic contributions to ozone formation over the United States, Atmospheric Environment, 164, 8–19, https://doi.org/10.1016/j.atmosenv.2017.05.044, 2017.