



TOAST 1.0: Tropospheric Ozone Attribution of Sources with Tagging for CESM 1.2.2

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Abstract. A system for source attribution of tropospheric ozone produced from both NO_x and VOC precursors is described, along with its implementation in the Community Earth System Model (CESM) version 1.2.2 using CAM4. The user can specify an arbitrary number of tag identities for each NO_x or VOC species in the model, and the tagging system rewrites the model chemical mechanism and source code to incorporate tagged tracers and reactions representing these tagged species, as well as ozone produced in the stratosphere. If the user supplies emission files for the corresponding tagged tracers, the model will produce tagged ozone tracers which represent the contribution of each of the tag identities to the modelled total tropospheric ozone. Our tagged tracers preserve O_x. The size of the tagged chemical mechanism scales linearly with the number of specified tag identities. Separate simulations are required for NO_x and VOC tagging, which avoids the sharing of tag identities between NO_x and VOC species. Results are presented and evaluated for both NO_x and VOC source attribution. We show that northern hemispheric surface ozone is dominated year-round by anthropogenic emissions of NO_x, but that the mix of corresponding VOC precursors changes over the course of the year; anthropogenic VOC emissions contribute significantly to surface ozone in winter-spring, while biogenic VOC are more important in summer. The system described here can provide important diagnostic information about modelled ozone production, and could be used to construct source-receptor relationships for tropospheric ozone.

1 Introduction

Tropospheric ozone is an important air pollutant, as well as an contributor to anthropogenic radiative forcing of the climate (Monks et al., 2015). Major sources of ozone in the troposphere are transport from the stratosphere, and photochemical production involving reactions of oxides of nitrogen (NO and NO₂, collectively NO_x) and Volatile Organic Compounds (VOC), including methane. Almost all of this photochemical production is related to the conversion of NO to NO₂ by reaction with a peroxy radical produced during the oxidation of VOC (Atkinson, 2000). Due to its long lifetime in the troposphere (several weeks), ozone can be transported over intercontinental distances. Concentrations of ozone observed at any given location can be due to both transported ozone from elsewhere, and ozone produced from precursors emitted nearby.

Global chemistry-climate models are important tools for understanding the complex processes of chemistry and transport which affect tropospheric ozone, simulating its evolution and distribution under future climate change, and projecting how



this may change in response to precursor emission controls. Based on a suite of model simulations from the ACCMIP model intercomparison project (Lamarque et al., 2013), Young et al. (2013) found that while the ensemble average of modelled ozone mixing ratios generally agreed with the present-day distribution of tropospheric ozone well, the individual models showed large differences from each other. Furthermore, the models generally agreed on the sign of the difference between present-day and both pre-industrial and future (late 2100s) conditions, but they tended to disagree strongly on the magnitude of these changes. The current state-of-the-art models show differing sensitivities in tropospheric ozone to changes in both climate and precursor emissions. In a more detailed comparison of the ACCMIP models with observed datasets, Parrish et al. (2014) showed that the models are not able to simulate the observed long term changes in tropospheric ozone, and concluded that more work is needed to improve the representation of chemistry and transport processes in models, as well as our understanding of historical emission changes, before the models could be reliably used to simulate future changes in tropospheric ozone. Young et al. (2013) identified the need for improved diagnostic information about modelled ozone budgets in order to understand the differences between models.

Intercontinental source-receptor relationships for tropospheric ozone have been modelled in the HTAP (Hemispheric Transport of Air Pollution) project using a “perturbation” methodology in which emissions of ozone precursors in source regions are reduced by some fraction (eg. 20%), and the resulting modelled ozone concentrations in receptor regions are compared with a base simulation in which the emissions were not perturbed (Fiore et al., 2009). An alternative approach for determining source-receptor relationships in model runs is a technique known as “tagging”, in which ozone molecules are labelled with the identity of their source, allowing direct attribution of ozone concentrations to these sources in receptor regions (eg. Wang et al., 1998; Dunker et al., 2002; Sudo and Akimoto, 2007; Grewe et al., 2010; Emmons et al., 2012; Derwent et al., 2015; Kwok et al., 2015; Grewe et al., 2017; Guo et al., 2017). Tagging (source apportionment) methodologies are complementary to perturbation (sensitivity) methodologies (Emmons et al., 2012; Grewe et al., 2017; Clappier et al., 2017). Since tagging methods can deliver detailed information about the provenance of modelled ozone concentrations, they could potentially be a useful tool for understanding the differences between models.

In this manuscript we describe and characterise a novel method for tagged source attribution of tropospheric ozone, and contrast our approach with previous work. We present a review of prior tagging approaches in Section 2, then describe the implementation of our method in CESM 1.2.2 with CAM4 (Tilmes et al., 2015; Lamarque et al., 2012) in Section 3. The design of our model evaluation experiments is described in Section 4, and we evaluate and compare the results for both NO_x- and VOC-tagging in Section 5. Conclusions and outlook are presented in Section 6.

2 Review of tagging methods

There are many different examples of several different approaches to ozone tagging in both regional and global models. In this study, we focus on the attribution of ozone production to emitted precursors. Studies such as Wang et al. (1998), Sudo and Akimoto (2007) and Derwent et al. (2015) each tag ozone molecules based on the geographical model domains in which the



ozone molecules are formed, so do not directly attribute chemical ozone production to emissions of particular precursors, and will not be discussed further here.

Attribution of ozone production to emissions in models of atmospheric chemistry involves several design decisions and associated trade-offs:

- 5 – Is ozone production attributed to emissions of NO_x , VOC, or both? And if both, how is the chemical regime (NO_x - or VOC-limited) accounted for?
- Is ozone production attributed explicitly for each chemical reaction producing ozone, or is the total instantaneous ozone production in each grid cell attributed according to the proportion of each precursor present?
- Are tagged precursor species simulated explicitly, or are they grouped into chemical “families”?
- 10 – How does the tagging system treat the O_3 - NO_x null chemical cycle?

Since both NO_x and VOC are involved in the chemical production of ozone, most tagging schemes attempt to attribute ozone production to both of these types of precursors. Two approaches for simultaneous attribution of ozone to both NO_x and VOC have been used: determination of the chemical regime with attribution to the limiting precursor (either NO_x or VOC); and equal attribution to both NO_x and VOC precursors. In each case, additional tracers are added to the model, which track the emissions of NO_x and VOC species, which are typically labelled with the identities of their source sectors (eg. transport, industry, etc. . .) or source regions (eg. East Asia, North America, etc. . .).

Determination of the chemical regime is typically made according to the indicator ratio $\text{PH}_2\text{O}_2/\text{PHNO}_3$ (the ratio between the production rates of hydrogen peroxide and nitric acid). According to (Sillman, 1995), the chemical regime is NO_x - or VOC-limited if the ratio is above or below 0.35 respectively. This approach somewhat simplifies the highly complex chemistry of ozone production, in which there is a transition regime of sensitivity to both NO_x and VOC emissions. This approach is also typically used in regional modelling studies, where model grid cells are relatively small (compared with global models). VOC-limited chemical regimes are typically found in regions of very high NO_x emissions, such as urban areas, which are not well-resolved by global models. Dunker et al. (2002) and Kwok et al. (2015) describe the use of this technique in the regional models CAMx and CMAQ respectively. In both cases, the tagging scheme determines whether ozone production in each model grid cell is in a NO_x -limited or a VOC-limited chemical regime, and attributes all instantaneous ozone production to the limiting precursor, with tagged ozone tracers added in proportion to the relative concentrations of the tagged precursor tracers present in that grid cell. Tagged ozone tracers are chemically destroyed according to the modelled instantaneous ozone chemical loss rate. Such tagging schemes account for the rapid null cycles involving O_x species by not considering their cycling reactions as part of the instantaneous ozone production or loss rates.

30 We are not aware of any global modelling study which has attempted to attribute ozone production to NO_x or VOC precursors based on the chemical regime in each grid cell. Instead, ozone tagging at the global scale has been done either by focusing on only NO_x precursors (eg. Emmons et al., 2012), or by giving equal weight to both NO_x and VOC precursors (eg. Grewe et al., 2010, 2017; Guo et al., 2017). In each case, the production rate of tagged ozone tracers is determined explicitly with respect



to the rates of the underlying chemical reactions producing ozone, rather than the bulk instantaneous ozone production rate. Grewe et al. (2010, 2017) use rate constants from the base chemical mechanism as well as the full set of concentrations of the tagged species to explicitly calculate the production rate of each tagged species considering all of the possible combinations between differently tagged precursor reactants. In their scheme, tagged ozone is produced from reactions between tagged
5 NO and tagged peroxy radicals, and the ultimately produced ozone molecules inherit their tag identities from both types of precursors. Emmons et al. (2012) and Guo et al. (2017) take a different approach, and add extra reactions to the base chemical mechanism representing the transformations of the tagged precursors and the production of tagged ozone, relying instead on the chemical solver of their model to calculate the production and loss rates of tagged species.

Similarly to Grewe et al. (2010, 2017), Guo et al. (2017) also takes a combinatorial approach to the simultaneous attribution
10 of tagged ozone to both NO_x - and VOC-tagged precursors. They avoid the chemical mechanism becoming too large by only considering two tag identities (“East Asia” (EA) and “everywhere else” (EE)). Each reaction between a peroxy radical and NO then requires four corresponding tagged reactions: EA+EA, EA+EE, EE+EA, and EE+EE. The size of their tagged mechanism thus increases quadratically with the number of tag identities. In the case of the cross-reactions (EA+EE and EE+EA), the NO_2 produced from the reaction between NO and a peroxy radical is split into equal parts NO_2 from EA and NO_2 from EE, despite
15 the fact that the NO reactant in any given reaction can only have come from one of these regions. By using such a combinatorial approach, Grewe et al. (2010, 2017) and Guo et al. (2017) allow the transfer of tag identities between NO_x and VOC species, which can produce tagged tracer concentrations which have no physical meaning. For example, Figure 5(b) of Grewe et al. (2017) attributes approximately 10 Tg of CO production per year to lightning, despite the fact that lightning is only a source of NO_x in their model. Such an unphysical result could be obtained in their tagging scheme after decomposition of a molecule
20 of PAN (peroxy acetyl nitrate, an organic nitrate) which had been tagged as coming from NO_x due to lightning. A similarly unphysical transfer of tag identity would be obtained in the approach used by Guo et al. (2017) if lightning were chosen as one of their tag identities.

The treatment of the NO_x - O_3 chemical cycle is another area in which ozone tagging schemes can produce unphysical results. As pointed out by Kwok et al. (2015), the approach of Emmons et al. (2012) treats the reaction between NO and O_3 (forming
25 NO_2) as chemical destruction of O_3 . The subsequent rapid re-formation of O_3 from NO_2 photolysis is treated as new ozone production due to an emitted NO_x precursor, effectively “overwriting” the identity of tagged ozone from remote sources with the identity of tagged NO_x emissions from more nearby sources. The work of Grewe et al. (2017) does not suffer from this problem, because ozone is included in a chemical family (O_x = odd oxygen = O_3 + O + NO_2 + others) which is preserved during fast chemical exchanges. Guo et al. (2017) do not give enough information to determine whether their approach also
30 suffers from this tag-overwriting problem.

While the use of the O_x chemical family is essential to preserve the correct identity of tagged ozone species, the use of other chemical families for ozone precursors can introduce additional problems with tagging schemes. For example, Grewe et al. (2017) do not explicitly follow the propagation of tags through the full set of VOC oxidation intermediates, but instead only tag
35 a single “NMHC” chemical family, which includes all VOC oxidation intermediate species, including the oxidation products of methane, but excludes PAN. Their use of this NMHC family leads to the unphysical result from their Figure 5(d), in which



formation of PAN has been partially attributed to methane. There is no known chemical pathway in the atmosphere capable of transforming methane into PAN. This is not an inherent weakness of their tagging approach, but rather results from their choice of one chemical family to represent all VOC ozone precursors. In order to avoid such unphysical results, the choice of chemical families must be made carefully. Ideally, each individual VOC oxidation intermediate should be explicitly tagged.

5 Butler et al. (2011) introduced a method for recursively tagging all reactions involving VOC species in a chemical box model. They followed and tagged the oxidation pathways of all VOC intermediate products until they were fully oxidised, and thus no longer included in the chemical mechanism. Butler et al. (2011) used this method to determine the time-dependent ozone production potential of all VOC species in the MCM (Master Chemical Mechanism Saunders et al., 2003) by tagging each of the “primary“ (emitted) VOC with its own identity, and were thus able to attribute ozone production from intermediate VOC species back to the emissions of each primary VOC species, thus avoiding the use of a generic VOC chemical family. Butler et al. (2011) showed that the chemistry of VOC intermediate products can contribute significantly to the total ozone production from VOC over the timescales of several days after emission. Using this approach, it was feasible to tag each primary VOC and all of its intermediate oxidation products in the MCM with a unique tag, due to the way in which the interactions between different organic peroxy radicals are treated in the MCM. The peroxy-peroxy chemistry of each individual peroxy radical in the MCM is represented as a unimolecular decay reaction with a rate constant proportional to the total concentration of all other peroxy radicals. As also noted by Ying and Krishnan (2010), if these peroxy-peroxy reactions are treated explicitly in a tagged chemical mechanism, the size of the tagged mechanism would scale quadratically with the number of tags, which would rapidly become too large for practical use. The technique of Butler et al. (2011) was subsequently applied for comparison of several VOC oxidation mechanisms by Coates and Butler (2015). In order to avoid the quadratic scaling problem, the chemistry of the organic peroxy radicals in each chemical mechanism was rewritten in the MCM style, allowing the size of the tagged chemical mechanism to scale linearly with the number of tag identities.

In this manuscript we describe an extension to the ozone tagging system first described fully by Emmons et al. (2012). This extended tagging system improves upon the earlier work of Emmons et al. (2012), avoiding the various problems with previous tagging schemes described above.

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- Our tagging scheme allows an arbitrary number of user-defined tag names in a single model run, with the size of the chemical mechanism increasing linearly with the number of tag identities.
 - Our tagging scheme introduces new tagged tracers for members of the O_x chemical family (which avoids the problem that ozone tags are destroyed by the null cycle involving NO_x).
 - Our tagging scheme incorporates the recursive VOC tagging system of Butler et al. (2011), explicitly tagging each intermediate VOC and avoiding the use of precursor families.
 - Our tagging scheme avoids the possibility of VOC species being tagged with identities of NO_x species (and vice-versa) by requiring that two separate model runs be performed, one with NO_x tagging, and another with VOC tagging. The tagged O_x produced during the conversion of NO to NO_2 can only be assigned to the tagged identity of the NO precursor, or the
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tagged identity of the peroxy radical involved in each such transformation, depending on whether NO_x- or VOC-tagging is being used.

The extended tagging system allows a completely closed source attribution of tropospheric ozone to all precursors to be performed in two model runs, one with NO_x tagging, and another with VOC tagging.

5 3 Implementation of NO_x and VOC tagging

The tagging system is implemented as software which takes as input an arbitrary list of chemical species to be tagged (typically precursor emissions), and for each of these species, and arbitrary list of tags to be applied. The full suite of tagging tools, input files, and machine-readable tagged mechanism files are included in the online supplement to this manuscript. Due to the different requirements of NO_x and VOC tagging, the user must explicitly choose whether NO_x or VOC tagging is to be performed. The NO_x tagging approach is described in Section 3.1, and the VOC tagging approach is described in Section 3.2. The resulting complete lists of both NO_x-tagged reactions and VOC-tagged reactions are included in both machine- and human-readable form in the supplementary material to this manuscript.

The tagging system rewrites the model chemical mechanism and CAM4 source files to include a new set of tracers and reactions corresponding to these user-specified tagged species and their associated chemical reactions. For example, if the user specifies that the tags “anthropogenic” and “biogenic” are to be applied to the species NO and NO₂, the chemical mechanism file will be modified to include all necessary species and reactions such that the model will be able to simulate ozone due to NO_x emitted by anthropogenic and biogenic sources. The user must supply appropriate emission files containing the names of each of the tagged species in order for the additional tagged reactions and tracers to have any effect. The size of the modified chemical mechanism scales linearly with the number of tag identities requested by the user. The tagging system also modifies all model source files which contain code in which the tagged species are modified by other modelled processes such as deposition (dry and wet) and input or removal due to boundary conditions. The source code modification, including a full list of the source files which are modified, is described in more detail in Section 3.3.

Due to the potentially large number of additional reactions and species introduced into the chemical mechanism, it was necessary to modify the chemical mechanism preprocessor shipped with CESM1.2.2 to raise some hard-coded limits and ensure that the addition of the tagged reactions containing untagged species from the base mechanism does not alter the treatment of the untagged species in the chemical solver. The modified source code of the chemical preprocessor is included in the online supplement to this manuscript.

3.1 NO_x-tagged mechanism

The base chemical mechanism used here is taken from Emmons et al. (2012). The same base mechanism is used for both NO_x and VOC tagging. Following Emmons et al. (2012), the reactions of tagged species are implemented as additional reactions in the model chemical mechanism file involving both tagged and untagged reactants. Untagged reactants appear in stoichio-



metrically identical amounts in the reactants and products of each tagged reaction, so that tagged reactions do not alter the concentrations of untagged species.

3.1.1 Separation of NO_y and O_x tagged species

In order to allow an arbitrary number of tags in a single model run, and to avoid the tag overwriting problem described in Section 2, the chemical families NO_y (which includes NO_x and all NO_x reservoir species) and O_x are tagged separately. The following species from the base chemical mechanism belong to the NO_y family: NO; NO_2 ; NO_3 ; N_2O_5 ; HNO_3 ; HO_2NO_2 ; ISOPNO₃; ONIT; ONITR; PAN; MPAN. The following species from the base chemical mechanism belong to the O_x family: O_3 ; $\text{O}(^1\text{D})$; O; NO_2 ; NO_3 ; N_2O_5 ; HNO_3 ; HO_2NO_2 ; ISOPNO₃; ONIT; ONITR; PAN; MPAN. When performing VOC tagging, HO_2 is added to the O_x family (see Section 3.2.1 for more details).

Following Butler et al. (2011) we regard the reaction of NO with any peroxy radical (HO_2 and all organic peroxy radicals) and subsequent production of NO_2 as the process which effectively generates tropospheric ozone.



Reaction R1 from the base chemical mechanism is represented in our tagging system as follows:



Since NO_2 is in both the NO_y and O_x chemical families, two different tagged versions of NO_2 are produced in Reaction R2, which represent the distinct roles of NO_2 in each of these chemical families: NO_2_TAG is NO_y -tagged NO_2 ; while $\text{NO}_2_X_TAG$ is O_x -tagged NO_2 . The suffix “_TAG” is a placeholder which can be replaced by the tagging system with an arbitrary number of user-chosen tag identities, each of which is represented by a unique reaction added to the tagged chemical mechanism. The suffix “_X_TAG” represents members of the O_x chemical family produced from emitted NO_x species tagged with the identity “TAG”. Additional production pathways of O_x species are discussed below.

In the base chemical mechanism, ozone is produced from NO_2 via photolysis:



The NO produced from Reaction R3 is then available for additional reaction with a peroxy radical, while the atomic O goes on to produce O_3 . In the tagged chemical mechanism, the fate of O_x -tagged NO_2 is different from that of NO_y -tagged NO_2 :



In the tagged versions of Reactions R3 - R4, tagged ozone is produced from tagged O_x in Reaction R6, while the tagged O_x precursor NO remains available for further subsequent conversion of NO to NO_2 after its regeneration in Reaction R5.

The tag overwriting problem (Section 2) emerges from the reaction between ozone and NO:



Because Emmons et al. (2012) did not clearly distinguish the NO_y and O_x chemical families in their tagging system, their tagged NO_2 effectively inherited its tag from NO , leading to the replacement of tagged ozone identities by the O_3 - NO_x null chemical cycle. This has the effect that tag identities from nearby sources of NO_x are over-represented in the tagged O_3 in the study of Emmons et al. (2012).

In our tagging system, we avoid this problem by handling Reaction R8 as follows:



The tagged identity of the emitted NO_x precursor is preserved in Reaction R9, while the tagged identity of O_x is preserved in Reaction R10.

3.1.2 Preparation of the chemical mechanism for NO_x tagging

Before the tagging system can automatically generate a tagged chemical mechanism file including the user-specified tag identities, a set of placeholder reactions must be added by hand to the base chemical mechanism. In future versions of our tagging system, it may be possible to identify these reactions automatically. These reactions can be classified into a number of different categories based on their chemical characteristics:

1. Reactions of emitted NO_x and corresponding NO_y reservoir species. This category includes all reactions between NO and peroxy radicals which generate O_x -tagged NO_2 ($\text{NO}_2_X_TAG$).
2. Reactions of O_x species, including transformations between O_x family members, and sinks of O_x . This category changes slightly depending on whether NO_x or VOC are being tagged; for NO_x tagging, reactions of OH radicals with atomic O and molecular O_3 are sinks of O_x , while for VOC tagging these reactions preserve O_x (see Section 3.2.1 for more details).
3. Reactions which endogenously generate NO_y or O_x species. Stratospheric O_3 is produced in this category of reactions, through the photolysis of O_2 and N_2O which ultimately produce the specially tagged species $\text{O}_3_X_STR$. A small amount of atomic O is produced from the self-reaction of OH radicals, producing the specially tagged species “ O_X_XTR ” (“extra” sources). This category also changes slightly depending on whether NO_x or VOC are being tagged. When NO_x are being tagged, photolysis of N_2O in the stratosphere produces NO_STR , and the reactions of HO_2 with certain organic peroxy radicals produce $\text{O}_3_X_XTR$.

Similarly to Emmons et al. (2012), the species N_2O_5 , which is formed by reaction between NO_2 and NO_3 , is duplicated to account for the possibility that its tag is inherited from either NO_2 or NO_3 . The species NO_3 is also subject to a tag inheritance problem when being tagged as a member of the O_x chemical family in the following reaction:





In this case, the O_x tag identity could be inherited from either NO_2 or O_3 . Following Emmons et al. (2012) we simply let the O_x tag be inherited from NO_2 in this case.

A full list of NO_x -tagged reactions is given in the supplementary material to this manuscript, including reactions producing species specially tagged as “STR” and “XTR”.

5 3.2 VOC-tagged mechanism

Butler et al. (2011) introduced a methodology to recursively follow the chemistry of VOC species, starting from the emitted VOC, following all intermediate species, and ending when only unreactive products remain. For each intermediate species, additional reactions and tracers are added to the chemical mechanism which are tagged with the same identity as the originally emitted species. The added species include tagged organic peroxy radicals (generically represented here as RO_2 , but which are explicitly tagged by our tagging system). These RO_2 produce O_x by converting NO to NO_2 :



The “tagged products” of Reaction R12 include tagged versions of all of the intermediate VOC associated with the corresponding reaction from the base chemical mechanism. Many such reactions also include HO_2 as a product, which may go on to produce O_x by converting NO to NO_2 (Reaction R1). In order to attribute this O_x production to the appropriate tag identity, HO_2 is included in the O_x family when doing VOC tagging, and the HO_2 produced in tagged organic reactions is given the identity of the organic reactant responsible for its production. The $HO_2_X_TAG$ thus-produced gives its O_x tag to NO_2 when reacting with NO :



The tagging software automatically identifies reactions involving the user-specified “primary” (or emitted) VOC species in the base chemical mechanism, and automatically generates tagged reactions of these species and their intermediates, including $NO_2_X_TAG$ and $HO_2_X_TAG$ in the products where appropriate in order to attribute production of O_x to these emitted VOC species.

3.2.1 Preparation of the chemical mechanism for VOC tagging

In the case of VOC tagging, a number of reactions must be identified and categorised by hand, similarly to the case of NO_x tagging described in Section 3.1.2.

1. Reactions involving HO_2 . These include Reaction R13, reactions of the HO_2 reservoir species HO_2NO_2 , and sinks of HO_2 which do not pass the tag identity onto their products (typically reactions of HO_2 with RO_2 species).
2. Reactions of O_x species, including transformations between O_x family members and sinks of O_x . This category has substantial overlap with reactions involved in NO_x tagging, but with one small difference: since HO_2 is considered a member of the O_x chemical family when tagging VOC, the production of HO_2 from reactions of OH radicals with



atomic O and molecular O₃ is not treated as a sink for O_x as it is for NO_x tagging (Section 3.1.2). Instead, the tagged identity is preserved as HO₂_X_TAG.

3. Reactions which endogenously generate NO_y or O_x species. This category also has substantial overlap with NO_x tagging, including the production of stratospheric O₃ from photolysis O₂. An additional reaction which is considered during VOC tagging is the production of the specially-tagged species HO₂_X_XTR from the reaction between OH and H₂O₂.

Following Coates and Butler (2015), the chemistry of the organic peroxy radicals in the base chemical mechanism is modified here to use the permutation approach employed by the MCM, in which the cross reactions of individual RO₂ are represented as unimolecular decay reactions with rates proportional to the total concentration of all RO₂ species. Further details are given in Coates and Butler (2015).

- 10 A full list of VOC-tagged reactions is given in the supplementary material to this manuscript, including reactions producing species specially tagged as “STR” and “XTR”.

3.3 Automatic source code rewriting

Several of the CAM source code files must be modified in order to correctly handle the processes involving the tagged tracers. Source files are first modified by hand in such a way that they can be automatically rewritten by the tagging software to accommodate the tagged tracers, and will also compile and run correctly when the CAM is run without tagging enabled. This is accomplished by enclosing sections of relevant code between FORTRAN comments. For example, model variables which index the concentration array for tagged species are declared as follows:

```
! START TAGGING CODE
integer :: no_tag_ndx, no2_tag_ndx, no2_x_tag_ndx
! END TAGGING CODE
```

The tagging logic itself is similarly enclosed between comments. The tagging software scans each source file for these comment lines, and expands the code where appropriate, adding code for each tagged tracer which has been added to the chemical mechanism.

The modified files are listed here, along with short summaries of the changes made in each case. The hand-modified source files themselves, along with the tagging software and all other necessary input files are available in the supplementary material to this manuscript.

- cam_history.F90 Code is modified to account for the larger number of tracers which could potentially be written to history files.
- mo_aerosols.F90 Code for gas/aerosol partitioning of tag identities between ammonium nitrate and nitric acid is added.
- mo_airplane.F90 Code is added to tag emissions from aircraft with the hard-coded identity “AIR”.



- mo_drydep.F90 Dry deposition fluxes are calculated for tagged species using deposition velocities of the corresponding untagged species.
- mo_flbc.F90 Species added at the lower model boundary are appropriately tagged if tags are defined for these species.
- mo_fstrat.F90 Tagged tracers are adjusted at the upper model boundary based on the adjustments made to the corresponding non-tagged species. Any O_x or NO_y added to the model is tagged as being of stratospheric origin. Other species are added or removed in proportion to their share of the corresponding untagged species.
- mo_gas_phase_chemdr.F90 Indices into the model concentration array for tagged species are determined during initialisation.
- mo_imp_sol.F90 Relative error parameters for tagged species in the implicit solver are set to the same values as for the corresponding untagged species.
- mo_lightning.F90 Code is added to tag NO production from lightning with the hard-coded identity “LGT”.
- mo_neu_wetdep.F90 Wet deposition fluxes are calculated for tagged species using removal rates of the corresponding untagged species.
- mo_photo.F90 Photolysis rates for the tagged reactions are set equal to the corresponding untagged reactions.
- mo_setext.F90 Code is added to facilitate the tagging of lightning NO and aircraft emissions.
- mo_sethet.F90 Loss rates due to heterogeneous chemistry are calculated for tagged species using removal rates of the corresponding untagged species.
- mo_srf_emissions.F90 Emissions of isoprene and monoterpenes are tagged appropriately if tags have been specified for these species.
- mo_usrrxt.F90 Rate constants of several of the tagged reactions are set equal to the rate constants of the corresponding untagged reactions.

4 Experiment Design

We use CESM version 1.2.2 (Tilmes et al., 2015; Lamarque et al., 2012) with the component set “FSDCHM” at a horizontal resolution of 1.9×2.5 degrees, with 56 vertical levels. This component set includes the tropospheric chemistry version of CAM4-chem forced with specified dynamics from year 2010 of the MERRA reanalysis (Rienecker et al., 2011). NO_x , O_3 , HNO_3 , N_2O_5 , N_2O , CO, and CH_4 are relaxed towards climatological values in the stratosphere. For this study, we replace the default chemical mechanism with the base mechanism from Emmons et al. (2012), modified as described in Section 3. Emissions of anthropogenic species are taken from the HTAP_v2.2 emission inventory (Janssens-Maenhout et al., 2015).



Biomass burning emissions are from GFEDv3 (van der Werf et al., 2010). Emissions of NO_x from lightning are calculated online within the model according to Price et al. (1997). Biogenic emissions of NO_x (from soils) and VOC (from vegetation) are prescribed as in Tilmes et al. (2015). Mixing ratios of CH_4 and N_2O are fixed at the surface as in Tilmes et al. (2015).

Model runs are done using both NO_x and VOC tagging, with the base chemical mechanism and model source code modified in each case as described in Section 3. We specify separate tag identities for emissions from anthropogenic (ANT), biogenic (BIO), biomass burning (BMB), and aircraft (AIR) sources. For NO_x tagging runs we specify an additional tag for NO_x from lightning (LGT), and for VOC tagging runs we specify an additional tag for methane (CH_4). In both cases (NO_x and VOC tagging) we include tags representing chemical production in the stratosphere (STR), “extra” chemical production (XTR, as described in Section 3), and a special tag representing the initial conditions (INI), allowing us to monitor the progress of the model spinup. This choice of tag identities allows us to compare our source attribution with that of Emmons et al. (2012), who used a similar set of tag identities, and on which our new tagging scheme is based.

Initial conditions for O_x species were tagged with STR in the stratosphere, and INI in the troposphere. Following Emmons et al. (2012), we used a chemical tropopause definition of 150 ppb of ozone. Initial methane in the VOC-tagging run was tagged with CH_4 . The concentration of INI-tagged and STR-tagged species was set equal to the mixing ratio of the corresponding species in the initial conditions, and all other tagged tracers were set to zero at the beginning of the model run. The model was run with annually repeating meteorology from 2010 until the maximum contribution of surface ozone attributable to the initial conditions was less than 1% of the total surface ozone, and the maximum difference between the stratospheric contribution to surface ozone in December from the stratospheric contribution to surface ozone in the previous December was also less than 1%. For VOC tagging we imposed the additional constraint that the difference between the contribution of methane to surface ozone in December and the contribution in the previous December was less than 1%. This was achieved after 2 years of simulation for NO_x -tagged runs, and 3 years of simulation for VOC-tagged runs. For the final year of simulation in each case, we verified that the method was working as expected by comparing the sum of the tagged ozone tracers with the actual ozone simulated by the model. At the lowest model level, the maximum monthly average difference was of the order of 1×10^{-5} ppb, while in the free troposphere the maximum monthly average difference was of the order of 1 ppb. The final year of simulation for both NO_x - and VOC-tagged runs is presented and discussed in Section 5.

5 Results

January average surface ozone mixing ratio, along with the mixing ratios of major contributing sources are shown from the NO_x -tagging run in Figure 1 and from the VOC-tagging run in Figure 2. Similarly, July average surface ozone mixing ratio is shown for NO_x - and VOC-tagging in Figures 3 and 4.

In the northern mid-latitudes, the land-sea gradient of modelled surface ozone reverses sign between January and July. Over the mid-latitude continental regions, modelled surface ozone has its maximum in summer, and its minimum in winter. Over the remote ocean regions, the opposite is the case; modelled surface ozone concentrations are higher in winter than they are in summer. Low modelled surface ozone mixing ratios over the northern mid-latitudes in winter are consistent with high

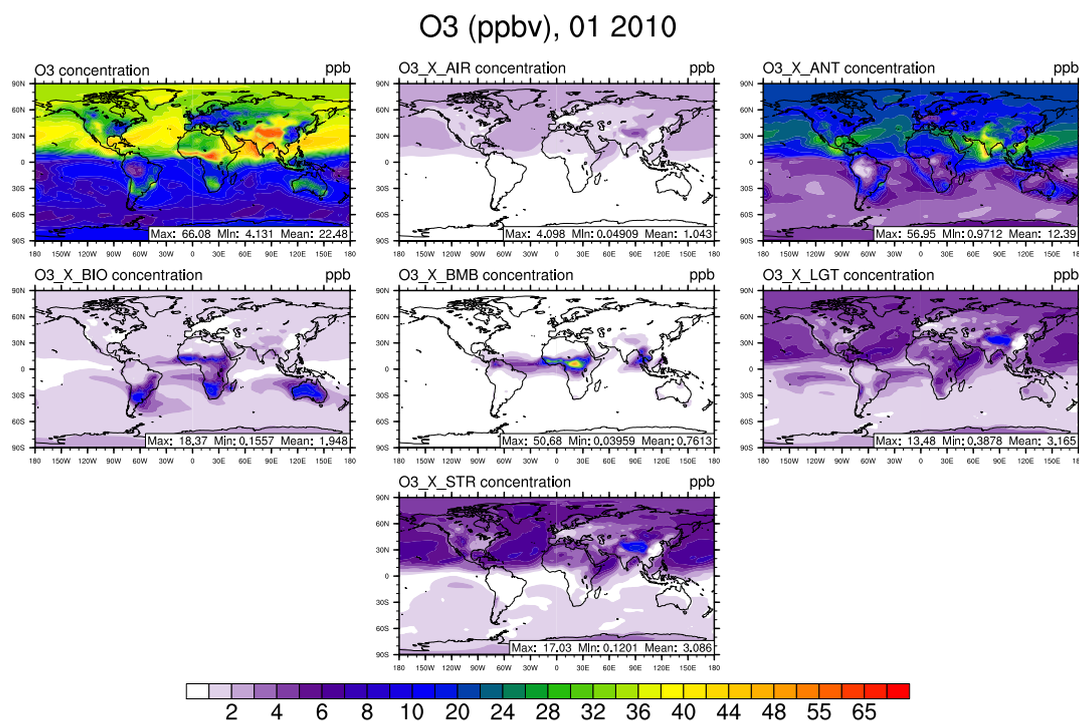


Figure 1. Surface ozone in January from the NO_x-tagging run. Total surface ozone is shown in the top-left panel. Other panels show the contribution to surface ozone due to NO_x precursors emitted by aircraft (AIR), anthropogenic sources (ANT), biogenic sources (BIO), biomass burning (BMB), lightning (LGT), and transport from the stratosphere (STR).

local emissions of NO_x, and ozone removal by Reaction R8. High modelled surface ozone mixing ratios over the northern mid-latitudes in summer are primarily attributable to a combination of anthropogenic NO_x emissions and biogenic NMVOC emissions, combined with more active photochemistry due to higher insolation. Anthropogenic NMVOC contribute relatively little to modelled high surface ozone mixing ratios in the boreal summer. This difference is consistent with the relatively high reactivity of biogenic NMVOC, especially isoprene, as well as the strong seasonal cycle in biogenic NMVOC emissions in mid-latitude regions, being emitted almost exclusively during the growing season.

Low modelled surface ozone mixing ratios over the remote northern hemispheric ocean regions in summer are consistent with a stronger chemical sink due to photolysis of ozone with subsequent production of OH radicals from water vapor (Johnson et al., 1999). The strength of this sink decreases during the winter, allowing modelled ozone to build up over large regions of the remote northern hemisphere. This hemispheric background ozone reaches a maximum in March/April (not shown) before the

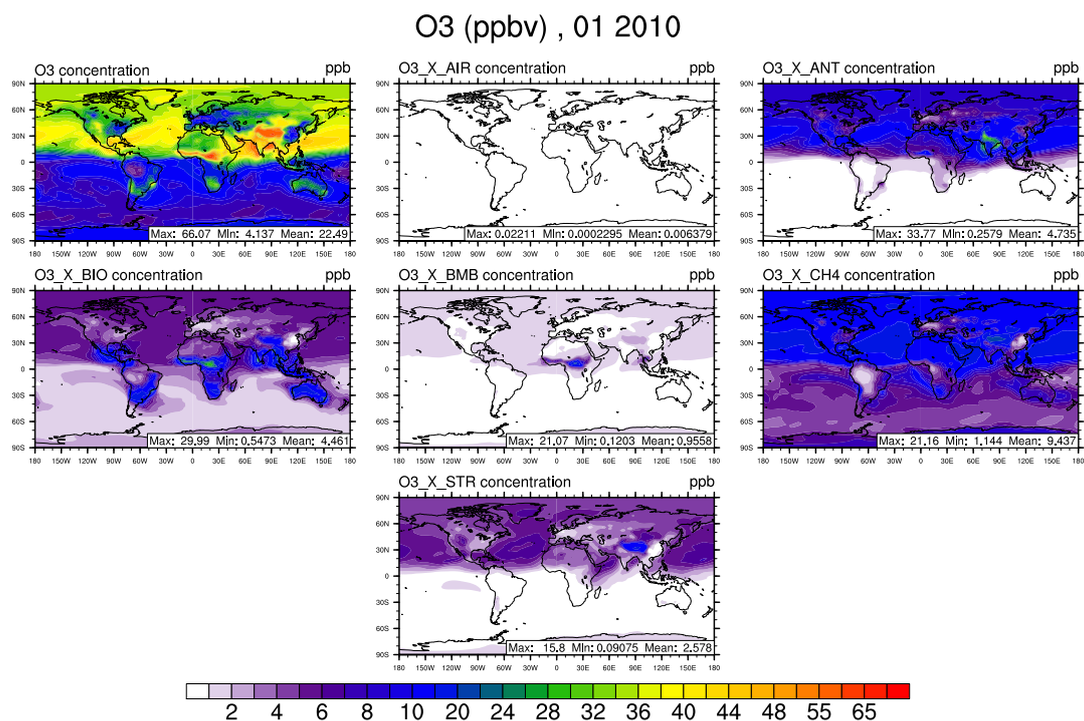


Figure 2. Surface ozone in January from the VOC-tagging run. Total surface ozone is shown in the top-left panel. Other panels show the contribution to surface ozone due to organic precursors emitted by aircraft (AIR), anthropogenic sources (ANT), biogenic sources (BIO), biomass burning (BMB), methane (CH₄), and transport from the stratosphere (STR).

chemical sink increases again. Examination of the tagged ozone tracers shows that this winter-spring remote maritime buildup of ozone is primarily attributable to both anthropogenic NO_x and NMVOC emissions. This is in contrast to the summer maximum in surface ozone modelled over continental regions, for which the primary responsible NMVOC precursor is of biogenic origin. A strong sensitivity of tropospheric ozone to anthropogenic NO_x and biogenic VOC emissions has been noted in previous studies (eg. Young et al., 2013; Stevenson et al., 2013), but we are not aware of any previous work in the peer reviewed literature showing that anthropogenic non-methane VOC contribute disproportionately to springtime ozone over remote regions of the northern hemisphere.

Another noteworthy feature of Figures 2 and 4 is the strong contribution of methane to the modelled mixing ratio of ozone at the surface, in both January and July. A strong sensitivity of modelled tropospheric ozone to the mixing ratio of methane has been noted in previous work (eg. Fiore et al., 2008; Young et al., 2013). Here, we show that the contribution of surface ozone

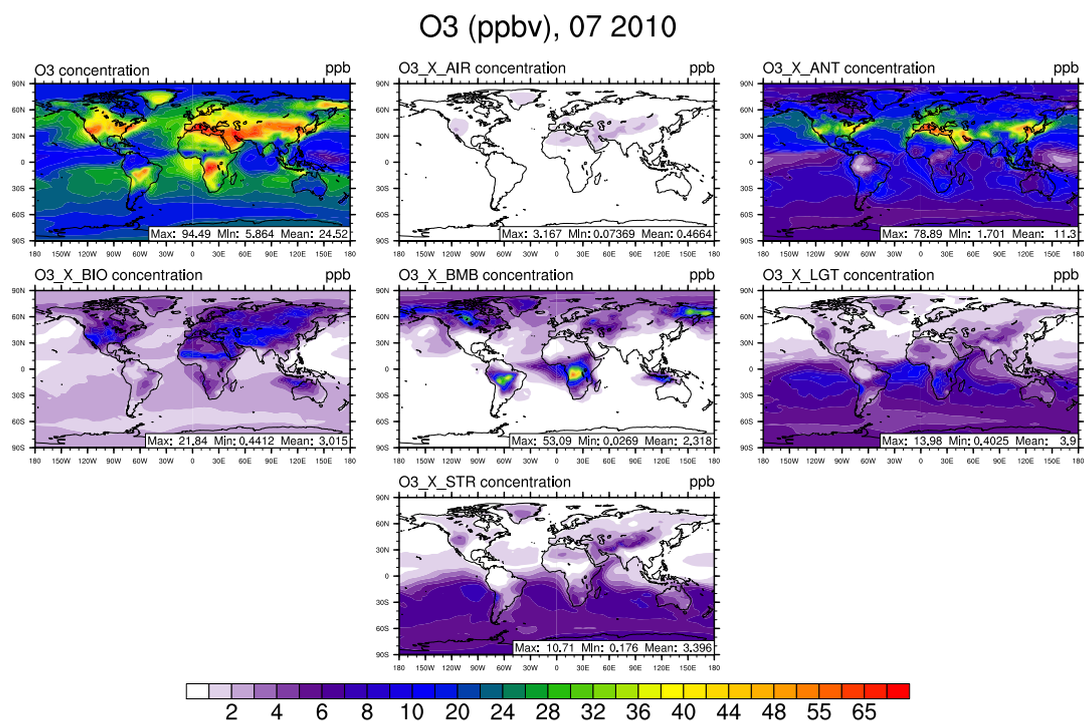


Figure 3. Surface ozone in July from the NO_x-tagging run. Total surface ozone is shown in the top-left panel. Other panels show the contribution to surface ozone due to NO_x precursors emitted by aircraft (AIR), anthropogenic sources (ANT), biogenic sources (BIO), biomass burning (BMB), lightning (LGT), and transport from the stratosphere (STR).

attributable to methane as an organic precursor remains remarkably constant at about 15 ppbv over large regions of the northern hemisphere year-round (at least in our model).

The influence of the stratosphere on the modelled ozone mixing ratio at the surface is stronger in winter than in summer. The stratospheric influence on northern hemisphere surface ozone is smallest in July and August, and reaches a maximum in March (not shown), when the contributions from the stratosphere, and the organic precursors methane and anthropogenic VOCs to the northern hemispheric background ozone are approximately equal. The late-winter early-spring maximum in the stratospheric contribution to surface ozone is consistent with both an increased lifetime of tropospheric ozone during this period, as well as the increasing flux of ozone from the stratosphere, which is consistent with the earlier work of Roelofs and Lelieveld (1997), who also used a stratospheric ozone tracer to determine the contribution of the stratosphere to surface ozone.

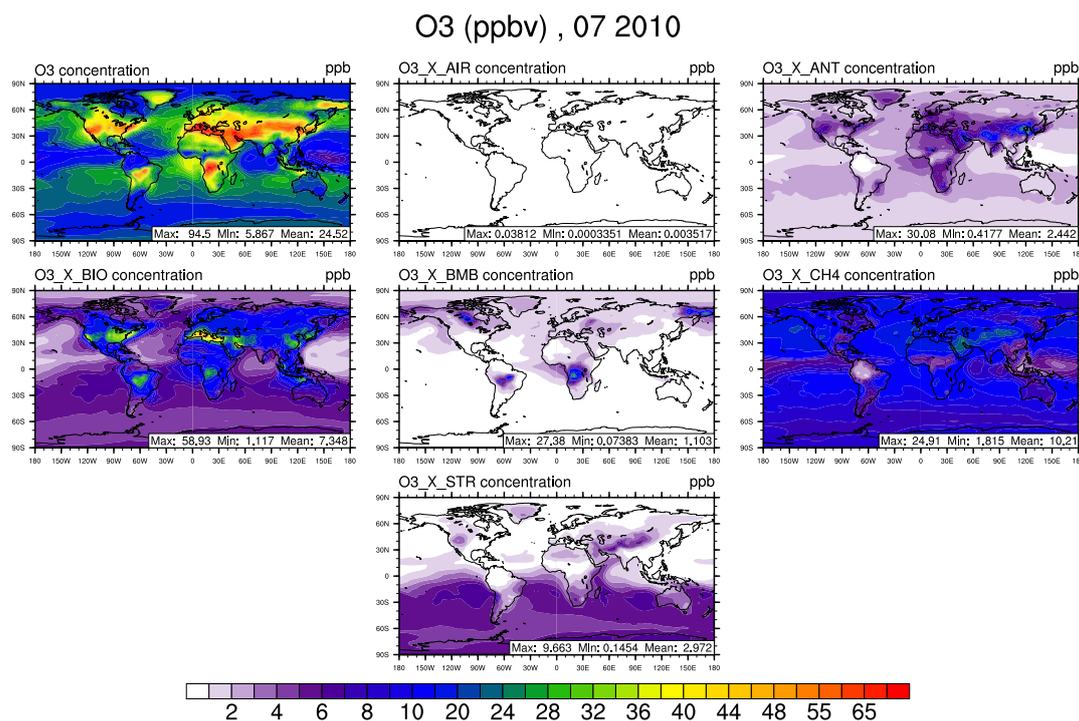


Figure 4. Surface ozone in July from the VOC-tagging run. Total surface ozone is shown in the top-left panel. Other panels show the contribution to surface ozone due to organic precursors emitted by aircraft (AIR), anthropogenic sources (ANT), biogenic sources (BIO), biomass burning (BMB), methane (CH₄), and transport from the stratosphere (STR).

Emmons et al. (2012) determined the contribution of stratospheric ozone to the modelled mixing ratio of ozone at the surface using their tagging approach. Since they did not explicitly tag the ozone originating in the stratosphere, they calculated the stratospheric contribution to tropospheric ozone as the residual after subtracting all of the ozone which had been produced from tagged tropospheric sources. They found that their residual stratospheric contribution to surface ozone was less than half of the contribution determined using a stratospheric tracer. Emmons et al. (2012) pointed out that such a stratospheric ozone tracer is likely to give an upper bound on the stratospheric contribution to surface ozone due to the fact that the tagged stratospheric ozone is set equal to the total ozone mixing ratio in the stratosphere, which effectively overwrites any tropospheric ozone which may have been imported into the stratosphere. We also regard the residual estimate of Emmons et al. (2012) as a lower bound on the contribution of stratospheric ozone to surface ozone, due to the “overwriting” problem mentioned above, in which their ozone tag identities are overwritten with the identity of nearby sources.

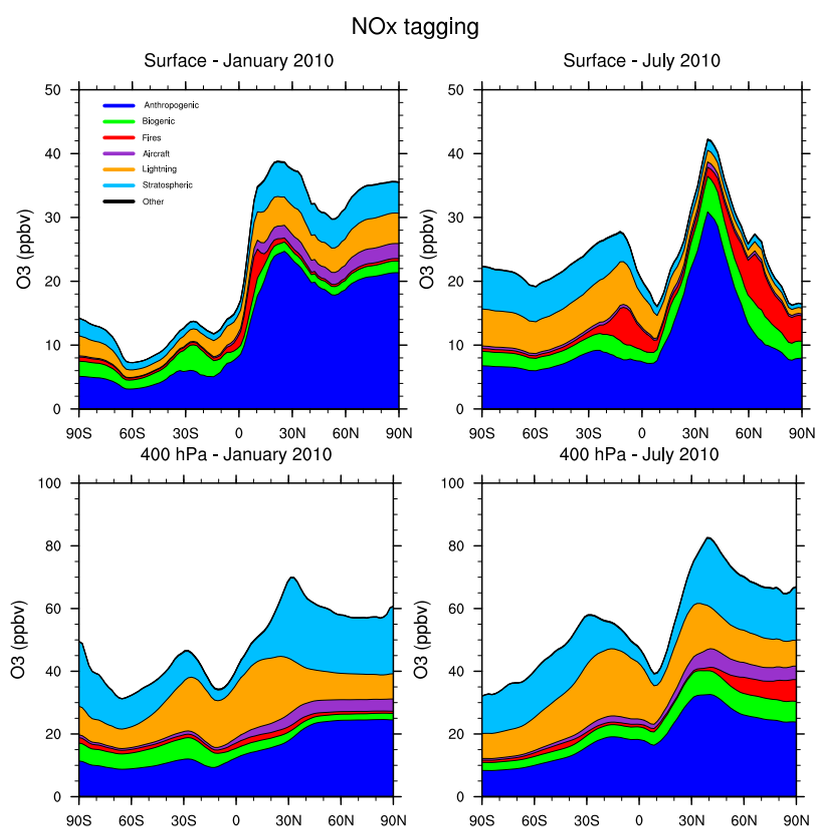


Figure 5. Zonal average of tagged ozone source contributions at the surface (top panels) and at 400 hPa (bottom panels) for January (left panels) and July (right panels) from the NO_x-tagging run

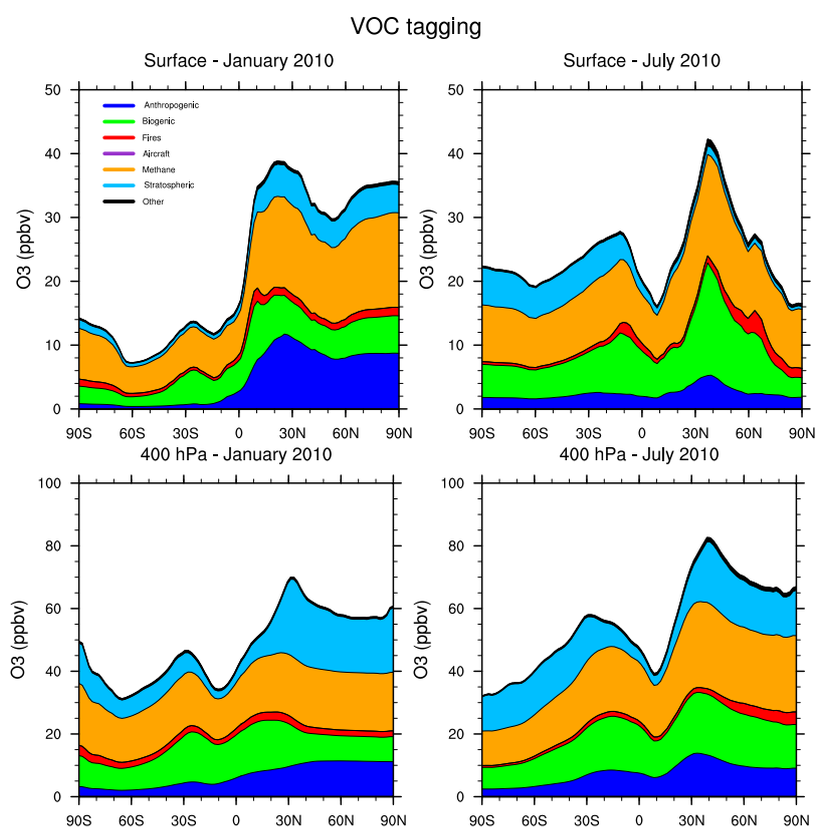


Figure 6. Zonal average of tagged ozone source contributions at the surface (top panels) and at 400 hPa (bottom panels) for January (left panels) and July (right panels) from the VOC-tagging run



Figure 5 shows the contribution of each of our tag identities to the zonally averaged ozone at the surface and at 400 hPa from our NO_x tagging run. This figure is designed to be directly comparable with Figure 6 of Emmons et al. (2012). Our simulated zonal average total ozone mixing ratio is broadly similar with that of Emmons et al. (2012) in both January and July, but there are some noteworthy differences in contributions of the tagged tracers; the stratospheric contribution to surface ozone shows particularly large differences. We model a zonally averaged stratospheric contribution to surface ozone of approximately 8 ppb in each winter hemisphere (NH in January and SH in July). These results are similar to those of Emmons et al. (2012) in the southern hemisphere, but approximately double those in the northern hemisphere winter, where Emmons et al. (2012) attribute only about 4 ppb of surface ozone to stratospheric origin. The lower stratospheric contribution to northern hemisphere surface ozone from Emmons et al. (2012) is consistent with their bias towards nearby sources due to the tag overwriting problem, as noted above. Similarly, Emmons et al. (2012) estimate a higher (by approximately 5 ppb) contribution of anthropogenic emissions to zonal average surface ozone than we see in our Figure 5, and show effectively no stratospheric contribution in July, while our run shows a small contribution of about 3 ppb stratospheric ozone to surface ozone in July. These results illustrate the importance of explicitly separating tagged species which are members of both the NO_y and O_x families to preserve the tagged identities of ozone transported over long distances.

The contribution of the tagged VOC precursors to zonal average surface ozone is shown in Figure 6. The widespread, year-round contribution of methane to ozone production is clearly visible, as is the increased importance of anthropogenic non-methane VOC as an ozone precursor during winter, noted earlier. Figure 6 also includes a contribution from the stratospherically tagged ozone in our VOC tagging run. It is immediately apparent from comparison with Figure 5 that the stratospheric contribution to tropospheric ozone is lower in the VOC tagging run than in the NO_x tagging run. Since the direct production of stratosphere-tagged ozone is identical in both runs, this difference must be due to ozone production involving stratosphere-tagged NO, produced from photolysis of N₂O as described in Section 3.1.2. By comparison with Figure 5, we see that this NO_x from the stratosphere contributes approximately an additional 2 ppb to the surface ozone ultimately attributable to the stratosphere (or approximately one quarter of the total stratospheric contribution). We are not aware of any previous work quantifying the contribution of the photolysis of N₂O in the stratosphere to the photochemical production of ozone in the troposphere. We note that our model does not include a comprehensive treatment of stratospheric chemistry and associated stratosphere-troposphere exchange. While our model does explicitly represent the photolysis of O₂ and N₂O in the stratosphere, the mixing ratios of O_x and NO_y species are also relaxed towards climatological values in the stratosphere. Future work examining the contribution of stratospheric NO_x to tropospheric ozone production should implement our tagging methodology in a fully coupled stratosphere-troposphere model.

6 Conclusions

We have introduced and described a technique for attribution of tropospheric ozone to emitted precursors of both NO_x and VOC, as well as transport from the stratosphere. The results obtained using this technique are consistent with understanding of tropospheric ozone chemistry based on previous work. Our work shares features with many earlier methodologies for



attribution of tropospheric ozone, but combines these features in unique ways which allow a unique and deeper understanding of the processes influencing tropospheric ozone in our model, and avoid many of the problems associated with previous work such as over-attribution of ozone to locally emitted precursors, and the unphysical transfer of tag identities between NO_x and VOC species.

- 5 By performing simultaneous but separate attribution of ozone to both its NO_x and VOC precursors, we have quantified, for example, the changing contributions of anthropogenic and biogenic sources to modelled seasonal cycles of surface ozone over the populated and remote regions of the northern hemisphere. In particular, we have identified the combination of anthropogenic NO_x and anthropogenic VOC as a significant contributor to the widespread buildup of ozone over the northern hemisphere during winter-spring in our model, in contrast with a relatively insignificant role for anthropogenic VOC in summer ozone production, for which biogenic VOC play a more important role. Further experiments using this tagging technique should examine the winter-spring contribution of anthropogenic VOC in more detail. Such experiments could instead tag anthropogenic VOC emissions according to their source sector, geographical region, time of emission, or even according to the particular kinds of VOC molecules emitted, in order to understand more about the ultimate sources of this springtime ozone in different receptor regions.
- 15 Given the problems of the current generation of global chemistry-climate models in simulating amounts, trends, and seasonal cycles of tropospheric ozone, the deeper understanding provided by our tagging methodology may yield information about deficiencies in these models and point the way towards improvements. If implemented in additional chemistry-climate models, our methodology could be a useful tool in understanding the differing responses of different models to changes in precursor emissions. Given the large number of alternative methodologies for attribution of tropospheric ozone, including the several
- 20 different ways of implementing tagging which have been reviewed here, we also believe that the community would benefit from a systematic intercomparison of the different techniques for constructing source-receptor relationships of tropospheric ozone.

Code availability. The full suite of tagging tools, input files, and machine-readable tagged mechanism files are included in the supplementary material to this manuscript.

- 25 *Author contributions.* TB conceived and designed the study. TB implemented the automatic mechanism rewriting and code generation tools. ZS and AL adapted the CESM source code. AL performed the model runs and subsequent analysis. AL and JC both contributed tools for analysing the model runs. TB wrote the paper.

Competing interests. The authors declare no competing interests.



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References

- Atkinson, R.: Atmospheric chemistry of VOCs and NO_x, *Atmos. Environ.*, 34, 2063–2101, 2000.
- Butler, T., Lawrence, M., Taraborrelli, D., and Lelieveld, J.: Multi-day ozone production potential of volatile organic compounds calculated with a tagging approach, *Atmos. Environ.*, 45, 4082 – 4090, <https://doi.org/10.1016/j.atmosenv.2011.03.040>, <http://www.sciencedirect.com/science/article/pii/S1352231011003001>, 2011.
- 5 Clappier, A., Belis, C. A., Pernigotti, D., and Thunis, P.: Source apportionment and sensitivity analysis: two methodologies with two different purposes, *Geoscientific Model Development*, 10, 4245–4256, <https://doi.org/10.5194/gmd-10-4245-2017>, 2017.
- Coates, J. and Butler, T. M.: A comparison of chemical mechanisms using tagged ozone production potential (TOPP) analysis, *Atmospheric Chemistry and Physics*, 15, 8795–8808, <https://doi.org/10.5194/acp-15-8795-2015>, 2015.
- 10 Derwent, R. G., Utembe, S. R., Jenkin, M. E., and Shallcross, D. E.: Tropospheric ozone production regions and the intercontinental origins of surface ozone over Europe, *Atmospheric Environment*, 112, 216–224, <https://doi.org/10.1016/j.atmosenv.2015.04.049>, 2015.
- Dunker, A., Yarwood, G., Ortmann, J., and Wilson, G.: Comparison of source apportionment and source sensitivity of ozone in a three-dimensional air quality model, *Environmental Science and Technology*, 36, 2953–2964, <https://doi.org/10.1021/es011418f>, 2002.
- Emmons, L. K., Hess, P. G., Lamarque, J.-F., and Pfister, G. G.: Tagged ozone mechanism for MOZART-4, CAM-chem and other chemical transport models, *Geosci. Model Dev.*, 5, 1531–1542, <https://doi.org/10.5194/gmd-5-1531-2012>, <http://www.geosci-model-dev.net/5/1531/2012/>, 2012.
- 15 Fiore, A. M., West, J. J., Horowitz, L. W., Naik, V., and Schwarzkopf, M. D.: Characterizing the tropospheric ozone response to methane emission controls and the benefits to climate and air quality, *J. Geophys. Res.*, 113, <https://doi.org/10.1029/2007JD009162>, 2008.
- Fiore, A. M., Dentener, F. J., Wild, O., Cuvelier, C., Schultz, M. G., Hess, P., Textor, C., Schulz, M., Doherty, R. M., Horowitz, L. W., MacKenzie, I. A., Sanderson, M. G., Shindell, D. T., Stevenson, D. S., Szopa, S., Van Dingenen, R., Zeng, G., Atherton, C., Bergmann, D., Bey, I., Carmichael, G., Collins, W. J., Duncan, B. N., Faluvegi, G., Folberth, G., Gauss, M., Gong, S., Hauglustaine, D., Holloway, T., Isak-
sen, I. S. A., Jacob, D. J., Jonson, J. E., Kaminski, J. W., Keating, T. J., Lupu, A., Marmer, E., Montanaro, V., Park, R. J., Pitari, G., Pringle, K. J., Pyle, J. A., Schroeder, S., Vivanco, M. G., Wind, P., Wojcik, G., Wu, S., and Zuber, A.: Multimodel estimates of intercontinental source-receptor relationships for ozone pollution, *J. Geophys. Res.*, 114, article number D04301, <https://doi.org/10.1029/2008JD010816>,
25 2009.
- Grewe, V., Tsati, E., and Hoor, P.: On the attribution of contributions of atmospheric trace gases to emissions in atmospheric model applications, *Geoscientific Model Development*, 3, 487–499, <https://doi.org/10.5194/gmd-3-487-2010>, 2010.
- Grewe, V., Tsati, E., Mertens, M., Frömming, C., and Jöckel, P.: Contribution of emissions to concentrations: the TAGGING 1.0 submodel based on the Modular Earth Submodel System (MESSy 2.52), *Geoscientific Model Development*, 10, 2615–2633, <https://doi.org/10.5194/gmd-10-2615-2017>, <https://www.geosci-model-dev.net/10/2615/2017/>, 2017.
- 30 Guo, Y., Liu, J., Mauzerall, D. L., Li, X., Horowitz, L. W., Tao, W., and Tao, S.: Long-Lived Species Enhance Summertime Attribution of North American Ozone to Upwind Sources, *Environmental Science & Technology*, 51, 5017–5025, <https://doi.org/10.1021/acs.est.6b05664>, <http://dx.doi.org/10.1021/acs.est.6b05664>, 2017.
- 35 Janssens-Maenhout, G., Crippa, M., Guizzardi, D., Dentener, F., Muntean, M., Pouliot, G., Keating, T., Zhang, Q., Kurokawa, J., Wankmüller, R., Denier van der Gon, H., Kuenen, J. J. P., Klimont, Z., Frost, G., Darras, S., Koffi, B., and Li, M.: HTAP_v2.2: a mosaic of regional



- and global emission grid maps for 2008 and 2010 to study hemispheric transport of air pollution, *Atmospheric Chemistry and Physics*, 15, 11 411–11 432, <https://doi.org/10.5194/acp-15-11411-2015>, <https://www.atmos-chem-phys.net/15/11411/2015/>, 2015.
- Johnson, C., Collins, W., Stevenson, D., and Derwent, R.: Relative roles of climate and emissions changes on future tropospheric oxidant concentrations, *Journal of Geophysical Research Atmospheres*, 104, 18 631–18 645, 1999.
- 5 Kwok, R. H. F., Baker, K. R., Napelenok, S. L., and Tonnesen, G. S.: Photochemical grid model implementation and application of VOC, NO_x, and O₃ source apportionment, *Geoscientific Model Development*, 8, 99–114, <https://doi.org/10.5194/gmd-8-99-2015>, 2015.
- Lamarque, J.-F., Emmons, L. K., Hess, P. G., Kinnison, D. E., Tilmes, S., Vitt, F., Heald, C. L., Holland, E. A., Lauritzen, P. H., Neu, J., Orlando, J. J., Rasch, P. J., and Tyndall, G. K.: CAM-chem: description and evaluation of interactive atmospheric chemistry in the Community Earth System Model, *Geoscientific Model Development*, 5, 369–411, <https://doi.org/10.5194/gmd-5-369-2012>, [https://www.](https://www.geosci-model-dev.net/5/369/2012/)
- 10 [geosci-model-dev.net/5/369/2012/](https://www.geosci-model-dev.net/5/369/2012/), 2012.
- Lamarque, J.-F., Shindell, D., Josse, B., Young, P., Cionni, I., Eyring, V., Bergmann, D., Cameron-Smith, P., Collins, W., Doherty, R., Dalsoren, S., Faluvegi, G., Folberth, G., Ghan, S., Horowitz, L., Lee, Y., MacKenzie, I., Nagashima, T., Naik, V., Plummer, D., Righi, M., Rumbold, S., Schulz, M., Skeie, R., Stevenson, D., Strode, S., Sudo, K., Szopa, S., Voulgarakis, A., and Zeng, G.: The atmospheric chemistry and climate model intercomparison Project (ACCMIP): Overview and description of models, simulations and climate diagnostics,
- 15 *Geosci. Model Dev.*, 6, 179–206, 2013.
- Monks, P., Archibald, A., Colette, A., Cooper, O., Coyle, M., Derwent, R., Fowler, D., Granier, C., Law, K., Mills, G., Stevenson, D., Tarasova, O., Thouret, V., Von Schneidmesser, E., Sommariva, R., Wild, O., and Williams, M.: Tropospheric ozone and its precursors from the urban to the global scale from air quality to short-lived climate forcer, *Atmospheric Chemistry and Physics*, 15, 8889–8973, <https://doi.org/10.5194/acp-15-8889-2015>, 2015.
- 20 Parrish, D., Lamarque, J.-F., Naik, V., Horowitz, L., Shindell, D., Staehelin, J., Derwent, R., Cooper, O., Tanimoto, H., Volz-Thomas, A., Gilge, S., Scheel, H.-E., Steinbacher, M., and Fröhlich, M.: Long-term changes in lower tropospheric baseline ozone concentrations: Comparing chemistry-climate models and observations at northern midlatitudes, *Journal of Geophysical Research*, 119, 5719–5736, <https://doi.org/10.1002/2013JD021435>, 2014.
- Price, C., Penner, J., and Prather, M.: NO_x from lightning 1. Global distribution based on lightning physics, *Journal of Geophysical Research Atmospheres*, 102, 5929–5941, 1997.
- 25 Rienecker, M. M., Suarez, M. J., Gelaro, R., Todling, R., Bacmeister, J., Liu, E., Bosilovich, M. G., Schubert, S. D., Takacs, L., Kim, G.-K., Bloom, S., Chen, J., Collins, D., Conaty, A., da Silva, A., Gu, W., Joiner, J., Koster, R. D., Lucchesi, R., Molod, A., Owens, T., Pawson, S., Pegion, P., Redder, C. R., Reichle, R., Robertson, F. R., Ruddick, A. G., Sienkiewicz, M., and Woollen, J.: MERRA: NASA's Modern-Era Retrospective Analysis for Research and Applications, *Journal of Climate*, 24, 3624–3648, <https://doi.org/10.1175/jcli-d-11-00015.1>,
- 30 2011.
- Roelofs, G.-J. and Lelieveld, J.: Model study of the influence of cross-tropopause O₃ transports on tropospheric O₃ levels, *Tellus, Series B: Chemical and Physical Meteorology*, 49, 38–55, 1997.
- Saunders, S. M., Jenkin, M. E., Derwent, R. G., and Pilling, M. J.: Protocol for the development of the Master Chemical Mechanism, MCM v3 (Part A): tropospheric degradation of non-aromatic volatile organic compounds, *Atmos. Chem. Phys.*, 3, 161–180, 2003.
- 35 Sillman, S.: The use of NO_y, H₂O₂, and HNO₃ as indicators for ozone-NO_x-hydrocarbon sensitivity in urban locations, *J. Geophys. Res.*, 100, 14 175–14 188, 1995.
- Stevenson, D., Young, P., Naik, V., Lamarque, J.-F., Shindell, D., Voulgarakis, A., Skeie, R., Dalsoren, S., Myhre, G., Berntsen, T., Folberth, G., Rumbold, S., Collins, W., MacKenzie, I., Doherty, R., Zeng, G., Van Noije, T., Strunk, A., Bergmann, D., Cameron-Smith, P., Plummer,



- D., Strode, S., Horowitz, L., Lee, Y., Szopa, S., Sudo, K., Nagashima, T., Josse, B., Cionni, I., Righi, M., Eyring, V., Conley, A., Bowman, K., Wild, O., and Archibald, A.: Tropospheric ozone changes, radiative forcing and attribution to emissions in the Atmospheric Chemistry and Climate Model Intercomparison Project (ACCMIP), *Atmos. Chem. Phys.*, 13, 3063–3085, 2013.
- Sudo, K. and Akimoto, H.: Global source attribution of tropospheric ozone: Long-range transport from various source regions, *Journal of Geophysical Research*, 112, <https://doi.org/10.1029/2006jd007992>, 2007.
- 5 Tilmes, S., Lamarque, J.-F., Emmons, L. K., Kinnison, D. E., Ma, P.-L., Liu, X., Ghan, S., Bardeen, C., Arnold, S., Deeter, M., Vitt, F., Rye-son, T., Elkins, J. W., Moore, F., Spackman, J. R., and Martin, M. V.: Description and evaluation of tropospheric chemistry and aerosols in the Community Earth System Model (CESM1.2), *Geoscientific Model Development*, 8, 1395–1426, <https://doi.org/10.5194/gmd-8-1395-2015>, 2015.
- 10 van der Werf, G. R., Randerson, J. T., Giglio, L., Collatz, G. J., Mu, M., Kasibhatla, P. S., Morton, D. C., DeFries, R. S., Jin, Y., and van Leeuwen, T. T.: Global fire emissions and the contribution of deforestation, savanna, forest, agricultural, and peat fires (1997–2009), *Atmospheric Chemistry and Physics*, 10, 11 707–11 735, <https://doi.org/10.5194/acp-10-11707-2010>, 2010.
- Wang, Y., Jacob, D. J., and Logan, J. A.: Global simulation of tropospheric O₃-NO_x-hydrocarbon chemistry: 3. Origin of tro-
15 pospheric ozone and effects of nonmethane hydrocarbons, *Journal of Geophysical Research: Atmospheres*, 103, 10 757–10 767, <https://doi.org/10.1029/98jd00156>, 1998.
- Ying, Q. and Krishnan, A.: Source contributions of volatile organic compounds to ozone formation in southeast Texas, *Journal of Geophysical Research Atmospheres*, 115, <https://doi.org/10.1029/2010JD013931>, 2010.
- Young, P., Archibald, A., Bowman, K., Lamarque, J.-F., Naik, V., Stevenson, D., Tilmes, S., Voulgarakis, A., Wild, O., Bergmann, D., Cameron-Smith, P., Cionni, I., Collins, W., Dalsoren, S., Doherty, R., Eyring, V., Faluvegi, G., Horowitz, L., Josse, B., Lee, Y., MacKenzie,
20 I., Nagashima, T., Plummer, D., Righi, M., Rumbold, S., Skeie, R., Shindell, D., Strode, S., Sudo, K., Szopa, S., and Zeng, G.: Pre-industrial to end 21st century projections of tropospheric ozone from the atmospheric chemistry and climate model intercomparison project (ACCMIP), *Atmos. Chem. Phys.*, 13, 5401–5402, 2013.