A Comparison of Chemical Mechanisms using Tagged Ozone Production Potential (TOPP) Analysis:

Supplementary Material

J. Coates¹ and T. Butler¹

¹Institute for Advanced Sustainability Studies, Potsdam, Germany

January 15, 2015

S1 Introduction

2

This is the supplementary material to the research paper "A Comparison of Chemical Mechanisms using Tagged Ozone Production Potential (TOPP) Analysis" and provides further information about the methodology as well as additional analysis.

S2 Mechanism Setup

All chemical mechanisms were adapted from their original format into the modularised KPP (Damian et al., 2002) format for use in the MECCA boxmodel (Sander et al., 2005) as modified by (Butler et al., 2011).

The MCM v3.2 (Jenkin et al., 1997, 2003; Saunders et al., 2003; Bloss et al., 2005;
Rickard et al., 2014) is the reference mechanism and its approach to dry deposition,
photolysis and peroxy radical-peroxy radical reactions were applied to all mechanisms.

8 S2.1 Photolysis

Photolysis was parameterised as a function of the solar zenith angle following the MCM approach (Saunders et al., 2003). Species from reduced mechanisms with a direct

counterpart in the MCM v3.2 were assigned the corresponding MCM v3.2 photolysis rate parameter. Otherwise, the recommended rate parameter in the mechanism determined the appropriate MCM v3.2 photolysis rate parameter. In some cases, the MCM v3.2 photolysis rate parameter closest in magnitude to that specified by the mechanism.

For example, the organic nitrate species ONIT in RACM2 has a photolysis rate parameter of 1.96×10^{-6} s⁻¹ that was compared to the MCM v3.2 organic nitrate photolysis rate parameters (J₅₁ – J₅₇). The rate parameter J₅₄ is the most similar in magnitude and was assigned as the ONIT photolysis rate parameter in RACM2.

Photolysis reactions of a species in reduced mechanisms were sometimes represented
by more than one MCM v3.2 photolysis reaction. The product yields of the original
mechanism reactions were preserved using combinations of the MCM v3.2 rate parameters.

2 For example, glyoxal photolysis described by (R1) and (R2) in RADM2.

$$GLY + h\nu \longrightarrow 0.13 \text{ HCHO} + 1.87 \text{ CO} + 0.87 \text{ H}_2$$
 (R1)

$$\mathrm{GLY} + \mathrm{h}\nu \ \longrightarrow \ 0.45 \ \mathrm{HCHO} + 1.55 \ \mathrm{CO} + 0.8 \ \mathrm{HO}_2 + 0.15 \ \mathrm{H}_2 \quad \mathrm{(R2)}$$

Whereas in the MCM v3.2, (R3), (R4) and (R5) are prescribed for glyoxal photolysis with the rates J_{31} , J_{32} and J_{33} respectively.

$$GLYOX + h\nu \longrightarrow 2 CO + 2 H_2$$
 (R3)

$$\mathrm{GLYOX} + \mathrm{h}\nu \ \longrightarrow \ 2 \ \mathrm{CO} \, + \, 2 \ \mathrm{HO}_2 \quad \mathrm{(R4)}$$

$$GLYOX + h\nu \longrightarrow HCHO + CO$$
 (R5)

The product yields in (R1) were retained using a photolysis rate parameter of $0.87 J_{31} + 0.13 J_{32}$, whilst for (R2) the rate $0.15 J_{31} + 0.45 J_{32} + 0.4 J_{33}$ was used. Table S1 illustrates the product yield calculations.

40 S2.2 Organic Peroxy Radical Self and Cross Reactions

36

Reactions of organic peroxy radicals (RO₂) with other organic peroxy radicals are divided into self (RO₂ + RO₂) and cross (RO₂ + R'O₂) reactions. These reactions are typically represented in chemical mechanisms as bimolecular reactions which would cause ambiguities when implementing the tagging scheme. Namely, which tag to be used for the products of reactions between RO₂ reactants having different tags. The MCM v3.2 approach to self and cross RO₂ reactions (each RO₂ species reacts with the pool of all

| | Rate Parameter | MCM v3.2 Products and Yields |
|-------|---|--|
| | $0.87 \mathrm{J}_{31}$ | $1.74~{\rm CO}+0.87~{\rm H}_2$ |
| (R1) | $0.13 \mathrm{J}_{32}$ | $0.13~{ m CO}+0.13~{ m HCHO}$ |
| | $0.87~{ m J}_{31}+0.13~{ m J}_{32}$ | $1.87~{\rm CO}+0.13~{\rm HCHO}+0.87~{\rm H}_2$ |
| | $0.15 \mathrm{J}_{31}$ | $0.30~{\rm CO}+0.15~{\rm H}_2$ |
| (R2) | $0.45 \mathrm{J}_{32}$ | $0.45~\mathrm{CO}+0.45~\mathrm{HCHO}$ |
| (112) | $0.4 \mathrm{J}_{33}$ | $0.80~{\rm CO}+0.80~{\rm HO}_2$ |
| | $ \overline{ 0.15 \; \mathrm{J}_{31} + 0.45 \; \mathrm{J}_{32} + 0.4 \; \mathrm{J}_{33} } $ | $1.55~{\rm CO}+0.45~{\rm HCHO}+0.80~{\rm HO}_2+0.15~{\rm H}_2$ |

Table S1: Calculation of glyoxal MCM v3.2 photolysis rate parameters retaining RADM2 glyoxal photolysis product yields.

| Mechanism | Reaction | Rate Constant |
|----------------------|---|---|
| | C2H5O2 = C2H5O | $k*RO2*0.6 s^{-1}$ |
| MCM v3.2 | C2H5O2 = C2H5OH | $k*RO2*0.2 \text{ s}^{-1}$ |
| | C2H5O2 = CH3CHO | $k*RO2*0.2 \text{ s}^{-1}$ |
| MOZART-4 | $	ext{C2H5O2} + 	ext{CH3O2} = 0.7 	ext{ CH2O} + 0.8 	ext{ CH3CHO} + 	ext{HO2} \\ + 0.3 	ext{ CH3OH} + 0.2 	ext{ C2H5OH} 	ext{}$ | $2 \times 10^{-13} \text{ cm}^3$ $\text{molecules}^{-1} \text{ s}^{-1}$ |
| | $\begin{array}{l} {\rm C2H5O2+C2H5O2} = 1.6{\rm CH3CHO}+1.2{\rm HO2} \\ +0.4{\rm C2H5OH} \end{array}$ | $6.8 \times 10^{-14} \text{ cm}^3$ molecules ⁻¹ s ⁻¹ |
| MOZART-4 modified | $	ext{C2H5O2} = 0.8 \; 	ext{CH3CHO} + 0.6 \; 	ext{HO2} + 0.2 \; 	ext{C2H5OH}$ | $2 \times 10^{-13} * RO2 s^{-1}$ |

Table S2: Ethyl peroxy radical $(C_2H_5O_2)$ self and cross organic peroxy reactions in MCM v3.2 and MOZART-4 including rate constants. $k=2(6.6\times 10^{-27}\exp(365/T))^{\frac{1}{2}}$ molecules⁻¹ s⁻¹ and RO2 is the sum of all organic peroxy radical mixing ratios.

| Reactants | Products | Rate Constant |
|------------|--|--|
| MO2 + MO2 | $0.74~{ m HO2}+1.37~{ m HCHO}+0.63~{ m MOH}$ | $9.4 \times 10^{-14} \exp{(390/T)}$ |
| MOZ MOZ | 0.71 H 02 1.97 H 0H 0 0.00 H 0H | ${ m cm^3~molecules^{-1}~s^{-1}}$ |
| MO2 | $0.37~{ m HO2}+0.685~{ m HCHO}+0.315~{ m MOH}$ | $9.4 \times 10^{-14} \exp{(390/T)} * \text{RO2}$ |
| MOZ | 0.51 1102 0.005 110110 0.515 WOII | s^{-1} |
| ETHP + MO2 | $ m HO2+0.75\;HCHO+0.75\;ACD$ | $1.18 \times 10^{-13} \exp{(158/T)}$ |
| | $+\ 0.25\ {\rm MOH}\ +\ 0.25\ {\rm EOH}$ | ${ m cm^3~molecules^{-1}~s^{-1}}$ |
| ETHP | $0.63~{\rm HO2}+0.065~{\rm HCHO}+0.75~{\rm ACD}$ | $1.18 \times 10^{-13} \exp{(158/T)} * RO2$ |
| | + 0.25 EOH | s^{-1} |

Table S3: Dermination of ETHP pseudo-unimolecular reaction and rate constant in RACM2 including rate constants. RO2 is the sum of all organic peroxy radical mixing ratios.

other RO₂ at a single uniform rate) is used to avoid such ambiguities. The MCM v3.2
approach represents RO₂-RO₂ reactions as a pseudo-unimolecular reaction whose rate
constant includes a factor 'RO2' which is the sum of the mixing ratios of all organic peroxy
radicals (Saunders et al., 2003).

The pseudo-unimolecular reaction products and their yields were determined by one of two methods. Firstly, by using the $RO_2 + RO_2$ reaction and halving the product yields, demonstrated for the MOZART-4 treatment of the ethyl peroxy radical in Table S2.

- Alternatively, the RO₂+CH₃O₂ reaction was used to determine the products due to CH₃O₂ and these products are then removed.
- Table S3 demonstrates the steps determining the ETHP pseudo-unimolecular reaction in RACM2. First the products due to MO2 (CH₃O₂ in RACM2) are determined as outlined previously using the MO2 + MO2 reaction. The MO2 product yields are subtracted from the ETHP + MO2 reaction. Any products having a negative yield are not included in the final pseudo-unimolecular reaction.

The methyl acyl peroxy radical $(CH_3C(O)O_2)$ was the exception to the above approach. Although most mechanisms include a $CH_3C(O)O_2 + CH_3C(O)O_2$ reaction, the $CH_3C(O)O_2$ pseudo-unimolecular reaction was derived by subtracting the CH_3O_2 product yields from the $CH_3C(O)O_2 + CH_3O_2$ reaction. This approach was used as the $CH_3C(O)O_2 + CH_3O_2$ reaction is the most significant reaction for $CH_3C(O)O_2$.

- The rate constant for each pseudo-unimolecular reaction was taken as that of the $RO_2 + CH_3O_2$ reaction multiplied by an 'RO2' factor, which is the sum of the mixing ratios of all organic peroxy radicals. The $RO_2 + CH_3O_2$ rate constant was chosen as this is the most likely reaction to occur.
- Model runs using the original and modified approach to the RO₂-RO₂ reactions for each mechanism were performed. The resulting O₃ concentration time series were compared and are shown in Figure S1.

S2.3 Dry Deposition

Dry deposition velocities are taken from the MCM v3.2. The MCM v3.2 dry deposition velocities of the same chemical functional group were used for mechanism species without direct MCM v3.2 analogues. For example, the dry deposition velocity of PAN-like species

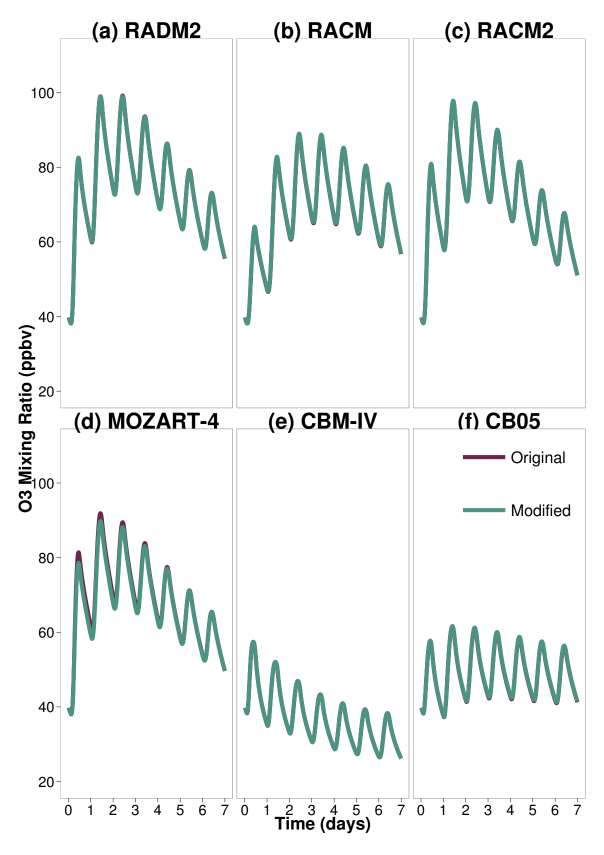


Figure S1: The time series of the O_3 for the reduced mechanisms using the original and modified approach to RO_2 – RO_2 reactions

in all mechanisms was equivalent to that of the PAN species in the MCM v3.2.

78 S2.4 Negative Product Yield Treatment

Some mechanisms include reactions where products have a negative yield. These reactions
were re-written including an operator species with a positive yield as the analysis tools
used in this study do not allow negative product yields. The operator species acts as a
sink for the original product by immediately reacting with the original product generating
a 'NULL' product.

For example, in RADM2 the OH + CSL (cresol) reaction has negative OH yield in (R6) (Stockwell et al., 1990).

$$CSL + OH \longrightarrow 0.1 HO_2 + 0.9 XO_2 + 0.9 TCO_3 - 0.9 OH (R6)$$

The negative OH yield was adapted to a positive operator (OHOP) yield in (R7). OHOP then reacts immediately with OH giving a 'NULL' product with a rate constant of $8.0 \times 10^{-11} \text{ cm}^3 \text{ s}^{-1}$ (R8). Thus preserving the OH yields from (R6) in RADM2.

S3 Mapping Emitted NMVOC to Mechanism Species

The emitted NMVOC are typical of Los Angeles as described in Baker et al. (2008). The MCM v3.2, v3.1 (Jenkin et al., 1997; Saunders et al., 2003; Jenkin et al., 2003) and CRI v2

(Jenkin et al., 2008) explicitly represent all of these NMVOC.

NMVOC representation in all other mechanisms required mapping to specific mechanism species. This mapping followed the recommendations on the literature of the mechanism. Table S4 describes the mechanism species used for mapping the initial NMVOC. Table 2 of the main article shows the final mapping of each NMVOC to each mechanism species.

o References

86

Angela K. Baker, Andreas J. Beyersdorf, Lambert A. Doezema, Aaron Katzenstein, Simone
Meinardi, Isobel J. Simpson, Donald R. Blake, and F. Sherwood Rowland. Measurements

| Mechanism | Species | Description | Mechanism | Species | Description |
|--------------------------|---------|---|----------------------|---------|---|
| | C2H6 | Ethane | | ETH | Ethane |
| | C3H8 | Propane | | HC3 | OH rate constant (298 K, 1 atm) less than 3.4×10^{-12} $\rm cm^3~s^{-1}$ |
| MOZART-4 (Emmons | BIGALK | Lumped alkanes C $>$ 3 | | HC5 | OH rate constant (298 K, 1 atm) between 3.4×10^{-12} and 6.8×10^{-12} cm 3 s $^{-1}$ |
| et al., 2010) | C2H4 | Ethene | m RACM2 | HC8 | OH rate constant (298 K, 1 atm) greater than $6.8\times10^{-12}~{\rm cm}^3~{\rm s}^{-1}$ |
| | C3H6 | Propene | (Goliff et al., | ETE | Ethene |
| | BIGENE | Lumped alkenes C >3 | 2019) | OLT | Terminal alkenes |
| | ISOP | Isoprene | | OLI | Internal alkenes |
| | TOLUENE | Lumped aromatics | | OSI | Isoprene |
| | ETH | Ethane | | BEN | Benzene |
| | HC3 | OH rate constant (298, 1 atm) between 2.7×10^{-13} and 3.4×10^{-12} | | TOL | Toluene and less reactive aromatics |
| m RADM2 | HC5 | OH rate constant (298, 1 atm) between 3.4×10^{-12} and 6.8×10^{-12} | | XYM | m-Xylene |
| (Stockwell | HC8 | OH rate constant (298, 1 atm) greater than 6.8×10^{-12} | | XYO | o-Xylene |
| et al., 1990) | OL2 | Ethene | | XYP | p-Xylene |
| | OLT | Terminal Alkenes | | PAR | Paraffin carbon bond C—C |
| | OLI | Internal Alkenes | | ETH | Ethene |
| | OSI | Isoprene | 1 | OLE | Olefinic carbon bond C=C |
| | TOL | Toluene and less reactive aromatics | CBM-IV | ALD2 | High molecular weight aldehydes |
| | XXL | Xylene and more reactive aromatics | (Get) et a, 1989) | ISOP | Isoprene |
| | ETH | Ethane | ` | TOL | Toluene |
| | HC3 | OH rate constant (298 K, 1 atm) less than 3.4×10^{-12} $\rm cm^3~s^{-1}$ | | XXL | Xylene |
| RACM | HC5 | OH rate constant (298 K, 1 atm) between 3.4×10^{-12} and $6.8\times10^{-12}~\rm cm^3~s^{-1}$ | | FORM | Formaldehyde |
| (Stockwell et al., 1997) | HC8 | OH rate constant (298 K, 1 atm) greater than 6.8×10^{-12} | | ЕТНА | Ethane |
| | ETE | Ethene | | PAR | Paraffin carbon bond C-C |
| | OLT | Terminal alkenes | CB05 | OLE | Terminal olefin carbon bond R-C=C |
| | OLI | Internal alkenes | (Yarwood | FORM | Formaldehyde |
| | OSI | Isoprene | Ct at.; 2000) | ISOP | Isoprene |
| | TOL | Toluene and less reactive aromatics | | TOL | Toluene and other monoalkyl aromatics |
| | XYL | Xylene and more reactive aromatics | | XYL | Xylene and other polyalkyl aromatics |

Table S4: Description of primary mechanism species used for mapping emitted NMVOCs.

- of nonmethane hydrocarbons in 28 United States cities. *Atmospheric Environment*, 42: 170–182, 2008.
- C. Bloss, V. Wagner, M. E. Jenkin, R. Vollamer, W. J. Bloss, J. D. Lee, D. E. Heard,
 K. Wirtz, M. Martin-Reviejo, G. Rea, J. C. Wenger, and M. J. Pilling. Development of a detailed chemical mechanism (MCMv3.1) for the atmospheric oxidation of aromatic
 hydrocarbons. Atmospheric Chemistry and Physics, 5:641-664, 2005.
- T. M. Butler, M. G. Lawrence, D. Taraborrelli, and J. Lelieveld. Multi-day ozone production potential of volatile organic compounds calculated with a tagging approach.

 Atmospheric Environment, 45(24):4082–4090, 2011.
- V. Damian, A. Sandu, M. Damian, F. Potra, and G.R. Carmichael. The kinetic preprocessor KPP A software environment for solving chemical kinetics. Computers and Chemical Engineering, 26(11):1567–1579, 2002.
- L. K. Emmons, S. Walters, P. G. Hess, J.-F. Lamarque, G. G. Pfister, D. Fillmore,
 C. Granier, A. Guenther, D. Kinnison, T. Laepple, J. Orlando, X. Tie, G. Tyndall,
 C. Wiedinmyer, S. L. Baughcum, and S. Kloster. Description and evaluation of the Model
 for Ozone and Related chemical Tracers, version 4 (MOZART-4). Geoscientific Model
 Development, 3:43-67, 2010.
- Michael W. Gery, Gary Z. Whitten, James P. Killus, and Marcia C. Dodge. A photochemical kinetics mechanism for urban and regional scale computer modeling.
 Journal of Geophysical Research, 94(D10):12,925-12,956, 1989.
- Wendy S. Goliff, William R. Stockwell, and Charlene V. Lawson. The regional atmospheric chemistry mechanism, version 2. *Atmospheric Environment*, 68:174–185, 2013.
- M. E. Jenkin, S. M. Saunders, V. Wagner, and M. J. Pilling. Protocol for the development of
 the Master Chemical Mechanism, MCM v3 (Part B): Tropospheric degradation of aromatic
 volatile organic compounds. Atmospheric Chemistry and Physics, 3(1):181–193, 2003.
- M. E. Jenkin, L. A. Watson, S. R. Utembe, and D. E. Shallcross. A Common Representative Intermediates (CRI) mechanism for VOC degradation. Part 1: Gas phase mechanism
 development. Atmospheric Environment, 42:7185–7195, 2008.

- Michael E. Jenkin, Sandra M. Saunders, and Michael J. Pilling. The tropospheric degradation of volatile organic compounds: A protocol for mechanism development.

 Atmospheric Environment, 31(1):81–104, 1997.
- Andrew Rickard, Jenny Young, and Stephen Pascoe. The Master Chemical Mechanism:

 Major updates in MCM v3.2. http://mcm.leeds.ac.uk/MCM/project.htt#New_3.2,

 2014. [Online; accessed 15-January-2015].
- R. Sander, A. Kerkweg, P. Jöckel, and J. Lelieveld. Technical Note: The new comprehensive
 atmospheric chemistry module MECCA. Atmospheric Chemistry and Physics, 5:445–450,
 2005.
- S. M. Saunders, M. E. Jenkin, R. G. Derwent, and M. J. Pilling. Protocol for the development of the Master Chemical Mechanism, MCM v3 (Part A): Tropospheric
 degradation of non-aromatic volatile organic compounds. Atmospheric Chemistry and Physics, 3(1):161–180, 2003.
- William R. Stockwell, Paulete Middleton, Julius S. Chang, and Xiaoyan Tang. The Second Generation Regional Acid Deposition Model Chemical Mechanism for Regional Air Quality
 Modeling. Journal of Geophysical Research, 95(D10):16,343–16,367, 1990.
- William R. Stockwell, Frank Kirchner, Michael Kuhn, and Stephan Seefeld. A new mechanism for regional atmospheric chemistry modeling. *Journal of Geophysical Research D: Atmospheres*, 102(22):25,847–25,879, 1997.
- Greg Yarwood, Sunja Rao, Mark Yocke, and Gary Z. Whitten. Updates to the Carbon Bond Chemical Mechanism: CB05. Technical report, U. S Environmental Protection
 Agency, 2005.