

Derivation of Atmospheric Reaction Mechanisms for Volatile Organic Compounds by the SAPRC Mechanism Generation System (MechGen)

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ABSTRACT

This paper describes the methods that are used in the SAPRC mechanism generation system, MechGen, to estimate rate constants and derive mechanisms for gas-phase reactions of volatile organic compounds (VOCs) in the lower atmosphere. Versions of this system have been used for over 20 years in the development of the SAPRC mechanisms for air quality models, but this is the first complete documentation of the scientific basis for the chemical mechanisms it derives. MechGen can be used to derive explicit gas-phase mechanisms for most compounds with C, H, O, or N atoms. Included are reactions of organic compounds with hydroxy (OH) and nitrate (NO₃) radicals, O₃, O³P, or by photolysis or unimolecular reactions, and the reactions of the radicals they form in the presence of O₂ and oxides of nitrogen (NO_x) at lower atmospheric temperatures and pressures. Measured or theoretically calculated rate constants and branching ratios are used when data are available, but in most cases rate constants and branching ratios are estimated using various structure-reactivity or other estimation methods. Types of reactions include initial reactions of organics with atmospheric oxidants or by photolysis, and unimolecular and bimolecular reactions of carbon-centered, alkoxy, and peroxy radicals and Criegee and other intermediates that are formed. This paper documents the methods, assignments, and estimates currently used to derive these reactions, and provides examples of MechGen predictions. Many of the estimation methods discussed here have not been published previously, and others have not been used previously in developing comprehensive mechanisms. Our knowledge of the of atmospheric reactions of organic compounds rapidly and continuously evolves, and therefore mechanism generation systems such as MechGen also need to evolve to continue to represent the current state of the science. This paper points out areas where MechGen may need to be modified when the system is next updated. This paper concludes with a summary of the major areas of uncertainty where further experimental, theoretical, or mechanism development research are most needed to improve predictions of atmospheric reaction mechanisms of volatile organic compounds.

25 1. Introduction

26 1.1. Background

27 When most volatile organic compounds (VOCs) are introduced into the atmosphere in the
28 presence of sunlight, they can react to form various radicals that then further react to form oxidized
29 organic products, including gas-phase toxics and secondary organic aerosol (SOA). In the presence of
30 oxides of nitrogen (NO_x) from combustion sources, these radicals also promote the formation of ozone
31 (O_3) and oxidized nitrogen compounds that further affect air quality. Many hundreds of types of organic
32 compounds are emitted, from both anthropogenic and biogenic sources. The atmospheric reaction
33 mechanisms for these compounds are complex, and for larger molecules can involve an extremely large
34 number of reactive intermediates and result in a large number of stable oxidized organic products. Most
35 of these organic products also react in the atmosphere, forming even more intermediates and products. In
36 most cases these mechanisms involve reactions whose rate constants, and in some cases mechanisms, are
37 unknown and thus have to be estimated. Because of the complexity, it is necessary either to greatly
38 simplify the mechanisms for most VOCs, use extensive lumping or condensations in representations of
39 VOCs, or use an automated chemical mechanism generation system to derive the mechanisms.

40 In the case of tropospheric reaction mechanisms of VOCs, existing automated chemical
41 mechanism generation systems include the Generator of Explicit Chemistry and Kinetics of Organics in
42 the Atmosphere (GECKO-A) developed by Aumont and co-workers (Aumont et al., 2005) and the
43 SAPRC mechanism generation system, MechGen, (Carter, 2024) that is the subject of this paper.
44 GECKO-A has been used in a number of studies of varying chemical complexity (e.g., Aumont et al.,
45 2005; Camredon et al., 2007; Lee-Taylor et al., 2011; Aumont et al., 2012; Lannuque et al., 2018; Afreh
46 et al., 2021) and is designed primarily to derive and carry out model simulations using multi-generation
47 mechanisms of selected compounds and all of their oxidation products. The SAPRC MechGen system
48 was developed primarily to support the development of versions of the SAPRC atmospheric gas-phase
49 chemical mechanisms (Carter, 2000, 2010a, b, 2016, 2020), and has been focused on single-generation
50 reactions of individual compounds, with mechanisms for representative organic oxidation products being
51 derived separately. Therefore, the objectives and operations of the two systems are somewhat different. In
52 addition, although the two systems employ similar structure-activity relationships (SARs) and can give
53 very similar predictions, there are some differences in the mechanisms they derive, reflecting both
54 different treatments of uncertainties and also different areas where updates are needed to incorporate
55 recent results. For example, GECKO-A does not yet predict autoxidation reactions of peroxy radicals that
56 are believed to occur (and are predicted by MechGen), while GECKO-A employs more detailed and
57 updated estimates for bimolecular reactions of peroxy radicals and photolysis reactions of larger
58 molecules.

59 Perhaps the most important difference between GECKO-A and MechGen is that GECKO-A is
60 described in the peer-reviewed literature (e.g., Aumont et al., 2005), while the description of MechGen is
61 largely incomplete in the literature, with only an abbreviated description in applied studies (Jiang et al.,
62 2020; Li et al., 2022). Different versions of MechGen, incorporating our changing and evolving
63 understanding and ability to estimate the underlying chemistry, have been used in the development of
64 SAPRC-99 (Carter, 2000), SAPRC-07 (Carter, 2010a, b), SAPRC-11 (Carter and Heo, 2013), SAPRC-16
65 (Carter, 2016; Venecek et al., 2018) and SAPRC-18 (Carter, 2020). Additional updates to the system have
66 been made since the release of SAPRC-18. The lack of a stable and peer-reviewed version has inhibited
67 the use of MechGen for atmospheric chemistry research beyond the development of SAPRC mechanisms,
68 despite its potential utility as an alternative to or for comparison with GECKO-A or the semi-explicit
69 Master Chemical Mechanism (Jenkin et al., 1997, 2003; Bloss et al., 2005; MCM, 2023). The purpose of
70 this paper is to document the chemical basis of the estimates and assignments as it currently exists, so it

71 can be appropriately cited and more widely used. The version of MechGen described here has been used
72 in the preparation of the recently-completed SAPRC-22 mechanisms (Carter, 2023a). This paper can also
73 be used to as a starting point for future updates and to illustrate areas of uncertainty where experimental
74 or theoretical studies are needed.

75 **1.2. Scope**

76 MechGen is capable of generating fully explicit mechanisms for the atmospheric reactions of
77 most types of organic compounds emitted into the atmosphere and the intermediate radicals they form. It
78 is designed to generate mechanisms for lower tropospheric modeling only, so its mechanisms are
79 applicable primarily for temperatures at or near 300 K and pressures at or near 1 atmosphere. Although
80 temperature-dependent rate constants are assigned or estimated for many reactions, for others the rate
81 constants or branching ratios are only applicable for 300 K and 1 atmosphere. In particular, MechGen is
82 not currently designed for estimating mechanisms for combustion modeling or for low temperature or low
83 pressure systems.

84 Table 1 lists the types of stable compounds and the categories of initial atmospheric reactions that
85 can be generated for those compounds. Table 2 lists the types of reactions within each category that can
86 be generated, including reactions of intermediate radicals and reactions of stable compounds. These
87 include H-atom abstractions from stable compounds by OH, NO₃, and Cl radicals; additions to double
88 bonds by these radicals and by O₃ and O³P; and photolyses at various groups. The types of radicals that
89 can be generated include carbon-centered radicals that in most cases react primarily with O₂; peroxy
90 radicals that in most cases react with NO, NO₂, NO₃, HO₂, or other peroxy radicals and in some cases also
91 have unimolecular reactions; alkoxy radicals that can react with O₂ or by various types of unimolecular
92 reactions; and excited and stabilized Criegee intermediates (CI). More information about their reactions
93 and how they are generated are described below, with additional detail given in the Supplementary
94 Information (SI) as referenced below.

95 Although the MechGen system can represent chemical structures containing halogen atoms and
96 estimate some types of reactions of compounds with Cl atoms, not all types of reactions of halogen-
97 containing compounds are currently supported. Therefore, a discussion of reactions of halogen-containing
98 compounds is not included here.

99 The operations and capabilities of MechGen as a software system are described in a separate
100 manuscript and user's manual (Carter, In Prep.), while this paper focuses on the chemical basis for the
101 derived mechanisms.

102 **1.3. Designation of Structures**

103 Information on how to input and designate structures for MechGen is given in the software
104 description and users' manual (Carter, In Prep.). Briefly, the structure of an organic reactant or radical is
105 specified by giving the "groups" in the molecule or radical and indicating the groups each are bonded to,
106 the type of bond, and in some cases (e.g., *cis* or *trans* isomers) the orientation of groups around the bonds.
107 Groups are parts of molecules that are treated as units in the system and are used to determine which
108 types of reactions can occur and what products are formed and used by the SARs or other methods to
109 estimate rate constants. These groups contain no more than one carbon or nitrogen atom, and also contain
110 zero to three hydrogen atoms and zero to three oxygen atoms. Structure designations are summarized in
111 Section S1.1 of the SI.

112 Table 1. Types of stable compounds whose reactions are supported by MechGen.

Type of Compound	Reactions	Exceptions not Supported
Alkanes	OH, NO ₃	Some estimates for polycyclic compounds are not reliable
Alkenes (including multiple double bonds)	OH, O ₃ , NO ₃ , O ³ P	See above; not all reactions of Cl-addition radicals can be generated
Alkynes	OH, NO ₃	See above
Aromatic hydrocarbons	OH	PAHs other than naphthalenes
Aldehydes, ketones, hydroperoxides, organic nitrates	OH, NO ₃ , hv	Photolysis reactions of larger compounds are over-simplified
Alcohols, ethers, esters, other oxygenates	OH, NO ₃	
Unsaturated aldehydes, ketones, hydroperoxides, and nitrates	OH, O ₃ , NO ₃ , O ³ P, hv	Ketenes
Phenols	OH, NO ₃	
Furans	OH, O ₃ , NO ₃ , O ³ P	
Other unsaturated oxygenates	OH, O ₃ , NO ₃ , O ³ P	
Amines	OH, NO ₃	Amines with no α or β hydrogens; imines
Peroxy nitrates	Unimolecular, OH, NO ₃ , hv	Formation of alkyl peroxy nitrates from RO ₂ + NO ₂ is not generated because of assumed rapid decomposition back to reactants
Bi- and polyfunctional compounds	OH, O ₃ , NO ₃ , O ³ P, hv, as applicable	Estimates are generally much more uncertain and less reliable

113

114

115 The generated reactions and estimated rate constants depend not only on the group(s) where the
 116 reaction occurs but also on the immediate neighbor group and in some cases groups some distance away
 117 in the molecule. In many cases, different groups or combination of groups are estimated to react similarly
 118 or have similar effects on reactions at neighboring groups, so this documentation uses designations that
 119 refer to such combinations. Examples include -CX₂H- to refer to carbon-centered groups with at least one
 120 hydrogen (where "X" designates either H or any neighboring group bonded to the carbon with a single
 121 bond), -CH_x-OH to any carbon-centered group bonded to an OH group. G refers to any group and -G-G-
 122 to chains of groups, etc. Some group designations used in the text will be somewhat different than those
 123 used by MechGen in order to be more familiar to chemists. Generally the group designations are noted in
 124 cases where they may not be obvious.

125

126 Table 2. Summary of types of reactions supported by MechGen. "VOC" indicates any stable
 127 compound supported by the system (see Table 1).

Reactant(s)	Type of Reactions
VOC + OH	H-atom abstraction Addition to double and triple bonds Addition to aromatic rings (OH only)
VOC + O ₃	Addition to double bonds followed by Criegee intermediate formation Excited adduct addition to amines, followed by decomposition forming OH
VOC + NO ₃	H-atom abstraction Addition to double bonds
VOC + O ³ P	Addition to double bonds
VOC + hv	Breaking the weakest bond in aldehydes, ketones, hydroperoxides, α-dicarbonyls, PAN compounds, and organic nitrates More rapid photolysis of dinitrates and carbonyl nitrates and carbonyl hydroperoxides
VOC Uni.	Unimolecular decompositions of peroxy nitrates
Carbon-centered Radicals	Unimolecular decompositions of radicals with α-nitro, α-nitrato, or α-peroxy groups Reaction with O ₂ with H-abstraction from α-OH groups Addition of O ₂ to radicals with allylic resonance Reactions of O ₂ with aromatic -OH adducts Addition of O ₂ to other alkyl radicals Unimolecular reactions of excited OH + alkene and aromatic adducts
Peroxy or Acyl Peroxy Radicals	Reactions with NO forming the corresponding alkoxy radical or organic nitrate Cyclization of aromatic OH-O ₂ adducts Unimolecular H-shift reactions forming hydroperoxides Reactions with NO ₂ forming the corresponding peroxy nitrate or PAN Reaction with HO ₂ forming the corresponding hydroperoxide or other products Reaction with NO ₃ forming NO ₂ and the corresponding alkoxy radical Reaction with the total of other peroxy or acyl peroxy radicals forming the corresponding alkoxy radical, carbonyl compound, or alcohol
Alkoxy Radicals	α-H abstraction by O ₂ forming the corresponding carbonyl compound β-scission decompositions H-shift isomerizations Ester rearrangement
Excited Criegee Intermediates	Decompositions, stabilization, or rearrangements of saturated intermediates Internal addition to the double bond of unsaturated intermediates, followed by O-O scission and epoxide formation

128 2. Initial Reactions of Organic Compounds

129 2.1. Bimolecular reactions with Atmospheric Oxidants

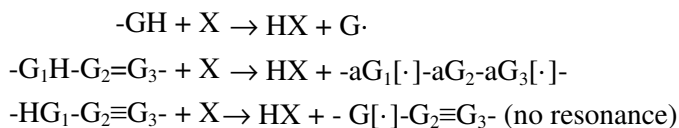
130 The methods MechGen uses to estimate rate constants for reactions of organic compounds with
131 OH, NO₃, O₃, O³P, and Cl have been documented by Carter (2021), with updates as discussed below and
132 in the SI. In most cases the rate constant estimates are made separately for each reaction route, with the
133 total rate constant being the sum of the estimated rate constants for each route. The estimates are made for
134 the various types of reactions using SARs based on the groups in the molecule, the neighboring groups
135 bonded to them (in some cases the 2nd neighbor as well), and the structure of the molecule around the
136 group(s) where the reaction occurs (e.g., groups in a ring). The equations and parameters used for
137 estimating the rate constants are summarized in Table 3 and the parameter values are given in various
138 tables in Section S1.2 of the SI. Most adjustable parameters needed for mechanism generation were
139 derived by optimizations to fit the measured OH, NO₃, O₃, and Cl rate constants tabulated by McGillen et
140 al. (2020), with the rate constants for most O³P reactions taken from Calvert et al. (2000). There were
141 insufficient data to derive all parameters using this approach, so some parameters were estimated based
142 on assumed relationships with the adjustable parameters.

143 The derivation or estimation of most of the parameters and the ability of the estimates to predict
144 the experimental data were given by Carter (2021). However, parameters for predicting rate constants for
145 reactions of phenols with OH and NO₃ were recently updated to improve the ability to model
146 environmental chamber experiments with generated mechanisms, as discussed in the SI. Rate constants
147 for most hydrocarbons and monofunctional compounds can be estimated to within ±30%, though
148 predictions are not as good for multifunctional compounds and predictions for ~15% of the rate constants
149 are off by more than a factor of 2. Estimates are more uncertain in the case of NO₃ and O₃ reactions
150 compared to OH, Cl, and O³P. This reflects the current state of the science in estimating these rate
151 constants with empirical structure-reactivity methods. Note that our estimates perform comparably to
152 those developed for the GECKO-A system by Jenkin et al. (2018a, b) for OH, Kerdouci et al. (2014) for
153 O₃, which were developed independently using similar methods (see Carter (2021)).

154 The following sections describe the mechanisms generated for the various types of reactions and
155 how branching ratios, if applicable, are derived. Note that some compounds may have more than one
156 reaction for a given oxidant, so branching ratios are estimated using ratios of estimated rate constants,
157 unless the branching ratios have been assigned explicitly for the compound and oxidant, as indicated in
158 the tables in Section S1.2 of the SI. If only the total rate constant is assigned, the branching ratios are
159 unaffected because they are derived from ratios of estimated rate constants. Note that additional
160 branching ratio estimates are needed for additions of O₃ or O³P to double or triple bond systems, as
161 discussed below.

162 2.1.1. H-Atom Abstractions by OH or NO₃

163 OH and NO₃ radicals can react with groups containing H atoms as follows, with rate constants
164 estimated as indicated in Table 3a and the parameters as indicated in Table S-3.



165

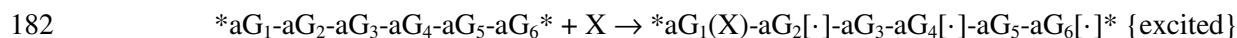
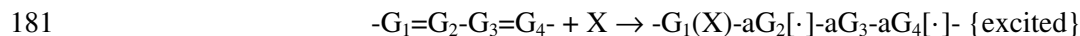
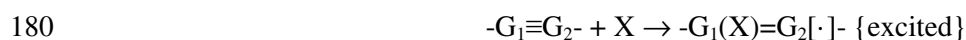
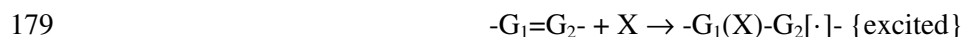
165 Table 3. Structure-reactivity equations and parameters used to estimate rate constants for reactions
 166 of VOCs with OH, NO₃, Cl, or O³P. Parameter values are given in the SI. Rate constants
 167 are per group unless otherwise indicated. The table numbers in the SI giving the assigned
 168 parameter values are also indicated.

a) <u>H-Atom abstractions from a group by OH or NO₃</u>
$kH(\text{group, site, nbrs}) = kH^{\text{base}}(\text{group}) \times FH^{\text{ring}}(\text{site}) \times FH^{\text{nbrs}}(\text{nbrs, group})$ $FH^{\text{nbrs}}(\text{nbrs, group}) =$ <p style="margin-left: 40px;">If no more than one non-alkyl substituent $FH^{\text{nbr}}(\text{nbr, group})$</p> <p style="margin-left: 40px;">If $\prod_{\text{nbrs}} FH^{\text{nbr}}(\text{nbr, group}) \leq 1$ $\prod_{\text{HCnbrs}} FH^{\text{nbr}}(\text{nbr, group}) \times \prod_{\text{nHCnbrs}} FH^{\text{nbr}}(\text{nbr, group})$</p> <p style="margin-left: 40px;">If $\prod_{\text{nbrs}} FH^{\text{nbr}}(\text{nbr, group}) > 1$ $\prod_{\text{HCnbrs}} FH^{\text{nbr}}(\text{nbr, group}) \times \text{Avg}_{\text{nHCnbrs}} [FH^{\text{nbr}}(\text{nbr, group})]$</p> <p style="margin-left: 40px;">Where HCnbrs and nHCnbrs refer to sets of alkyl and non-alkyl substituents, respectively</p> <p>$kH^{\text{base}}(\text{group})$: Base rate constant for abstraction from group (Table S-37 in the SI) $FH^{\text{ring}}(\text{site})$: Ring strain correction based on the smallest ring containing the group (Table S-42) $FH^{\text{nbrs}}(\text{nbrs, group})$: Correction factors for all substituents groups (defined above) $FH^{\text{nbr}}(\text{nbr, group})$: Correction factors for single substitution on groups (S-38, S-39, S-40)</p>
b) <u>Addition to a group on isolated or conjugated double or triple bonds by OH or NO₃</u>
<p>(Addition to group G₁ on a bond defined by G₁=G₂ or G₁≡G₂; nC = number of carbons)</p> $kA(G_1, G_2, \text{site, nbrs}) = kA^{\text{base}}(G_1, G_2) \times FA^{\text{struct}}(\text{site}) \times \prod_{\text{nbrs1}} FA_1^{\text{nbr}}(\text{nbr}) \times \prod_{\text{nbrs2}} FA_2^{\text{nbr}}(\text{nbr})$ $FA^{\text{struct}}(\text{nC}) = \{1 - \exp(-\min(\text{nC}, 12) \times FS^{\text{radical}}) / \{1 - \exp(-4 \times FS^{\text{radical}})\}\}; \text{ where } FS^{\text{NO}_3} = 0.101$ <p style="margin-left: 40px;">and $FS^{\text{OH}} = FS^{\text{Cl}} = 0$ (FA^{struct} correction for NO₃ only)</p> <p>$kA^{\text{base}}(G_1, G_2)$: Base rate constant for addition to this group on the double or triple bond (S-37) $FA^{\text{struct}}(\text{site})$: Structural correction factor for addition to this site (NO₃ only, see above) (S-39) $FA_1^{\text{nbr}}(\text{nbr})$: Correction factor for each substituent on G₁ (other than G₂) (S-38, S-39, S-40) $FA_2^{\text{nbr}}(\text{nbr})$: Correction factor for each substituent on G₂ (other than G₁) (S-39, S-39, S-40)</p>
c) <u>Additions to groups on aromatic rings by OH or NO₃</u>
$kAro(\text{aGrp, aSubs}) = kAro^{\text{base}}(\text{aGrp}) \times FA^{\text{ipso}}(\text{aSub}_1) \times FA^{\text{ortho}}(\text{aSub}_2) \times FA^{\text{meta}}(\text{aSub}_3) \times$ $FA^{\text{para}}(\text{aSub}_4) \times FA^{\text{meta}}(\text{aSub}_5) \times FA^{\text{ortho}}(\text{aSub}_6)$ <p>$kAro^{\text{base}}(\text{aGrp})$: Base rate constant for addition to this aromatic group (S-37) $FA^{\text{ipso}}(\text{aSub}_1)$: Correction for substituents on the same group where the addition occurs (S-41) $FA^{\text{ortho}}(\text{aSub}_2)$, etc: Correction factors for ortho, meta, or para substituents on aromatics (S-41)</p>
d) <u>Additions to non-aromatic double or triple bonds by O₃ or O³P (rate constant per bond)</u>
$kA'(\text{bond, site, nbrs}) = kA'^{\text{base}}(\text{bond}) \times FA'^{\text{ring}}(\text{site}) \times FA'^{\text{struct}}(\text{site}) \times \prod_{\text{nbrs}} FA'^{\text{nbr}}(\text{nbr})$ $FA'^{\text{struct}}(\text{n}\beta) = 1 - (F\beta^{\text{O}_3} \times \{(\min[4, \max(1, \text{n}\beta)] - 1)\}), \text{ where } F\beta^{\text{O}_3} = 0.268, F\beta^{\text{O}_3\text{P}} = 0, \text{ and } \text{n}\beta \text{ is}$ <p style="margin-left: 40px;">the number of substituents on non-H groups bonded to groups around the unsaturated bond</p> <p>$kA'^{\text{base}}(\text{bond})$: Base rate constant for addition to this type of bond (O₃ only) (S-43) $FA'^{\text{ring}}(\text{site})$: Ring strain correction factor for addition to this site (O₃ only) (S-44) $FA'^{\text{struct}}(\text{site})$: Correction factors for branched structures (see above) or furans (O₃ only) (S-44) $FA'^{\text{nbr}}(\text{nbr})$: Correction factor for each substituent on groups at ends of the bond (S-44)</p>
e) <u>Additions to the center group with cumulated double bonds by OH or NO₃, and Additions to the N in amino groups by OH, NO₃, O₃, or O³P</u>
$kA''(\text{group, nbrs}) = kA''^{\text{base}}(\text{group, nbrs}) \text{ (substituent effects incorporated in } kA''^{\text{base}} \text{) (S-37)}$

169 Here "G" is any group where H is removed, "X" is OH or NO₃, and "aG" refers to groups in a resonance
 170 structure (bonded to alternating double and single bonds). Separate estimates are made for each H-
 171 containing group in the molecule, so the branching ratio can be derived from the ratio of estimated rate
 172 constants to the sum of all the estimated rate constants for the VOC, including addition reactions (see
 173 below) where applicable. The subsequent reactions of the carbon-centered radicals are discussed in
 174 Section 3.

175 2.1.2. Additions of OH or NO₃ to Groups with Unsaturated Bonds

176 The OH and NO₃ radicals can also add to groups with double, triple, or aromatic bonds as
 177 follows, with rate constants estimated as indicated in Table 3b for alkenes and alkynes and Table 3c for
 178 aromatics. The parameters are given in the SI as indicated in Table S-3.



183 Here "G" refers to any group with an unsaturated bond, "X" refers to the reacting radical, "{excited}"
 184 indicates vibrational excitation, and "*" indicates ring closure. Separate estimates are made for addition to
 185 each group so these can also be used to make the branching ratio estimates. Most of the excited carbon-
 186 centered radicals are stabilized and react with O₂ as discussed in Section 3.1, but some have fast
 187 unimolecular reactions, as discussed in Section 3.3 for radical + alkene and alkyne adducts and Section
 188 3.4 for radical and aromatic adducts.

189 2.1.3. Additions of OH or NO₃ to Amines

190 Both OH and NO₃ radicals are known to react rapidly with amines (McGillen et al., 2020; Carter,
 191 2021), but the fact that the reaction with tertiary amines is about as rapid as reaction with secondary or
 192 primary amines indicates that the reaction is due to initial addition of the radical to the amino group,
 193 rather than direct H-abstraction from the amino group [see discussion in Section S1.9 of Carter (2021)].
 194 Based on this, MechGen assumes that the rate-determining step is addition of the radical to the amino
 195 group forming an addition complex that rapidly decomposes. An example is shown below.



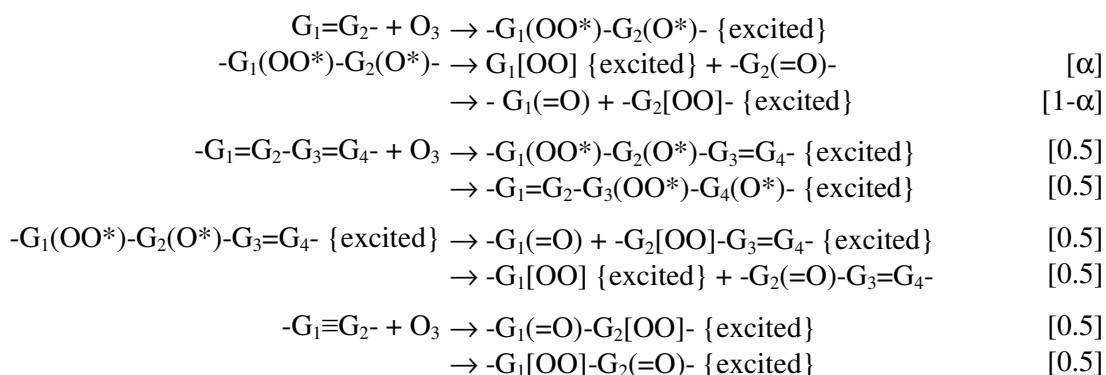
198 In the case of tertiary amines, where reactions analogous to (1) are not possible, the adduct is assumed to
 199 decompose by abstraction of an α hydrogen if one is present, forming the same products that would be
 200 formed if the radical abstracted from the α hydrogen directly. Both decomposition modes are possible for
 201 primary or secondary amines with α hydrogens, but the formation of the α -amino carbon-centered radical
 (analogous to 2) is estimated to be more exothermic and is assumed to dominate. If there is no α
 hydrogen, then only the reaction forming the amino radical (analogous to 1) is generated.

202 Neither of these decomposition modes are possible for tertiary amines with no α hydrogens (e.g.,
 203 tri-*t*-butyl amine), so MechGen assumes that the adduct decomposes back to reactants and there is no net
 204 reaction at the amino group. We are aware of no measurements or calculations of rate constants for these
 205 reactions, but they would be interesting to study.

206

2.1.4. Additions of O₃ to Unsaturated Bonds

207 O₃ is assumed to react only by additions to double or triple bond systems as follows, with rate
 208 constants estimated as indicated in Table 3d and with parameters as indicated in Table S-3. Reactions of
 209 O₃ with aromatics or cumulated double bonds are assumed to be negligible based on low measured rate
 210 constants (McGillen et al., 2020). Note that only total rate constants for additions to unsaturated bond
 211 systems are estimated, so it is necessary to assume branching ratios for reactions of the initially formed
 212 adducts or, in the case of additions to conjugated bond systems, for the initial reaction. The branching
 213 ratios used are shown in brackets where applicable.



214 Here α and $(1-\alpha)$ are branching ratios assigned for the additions to separated double bonds, $G(=O)$ refers
 215 to a carbonyl ($-\text{CO}-$ or $-\text{CHO}$) group, and $G[OO] \{ \text{excited} \}$ refers to an excited Criegee intermediate (CI)
 216 whose subsequent reactions are discussed in Section 7, and "*" indicates rings. The additions to double
 217 bonds are assumed to proceed via initial formation of excited primary ozonides, which decompose to the
 218 products as shown above; while additions to alkynes are assumed to form the excited CIs directly. The
 219 assumption that the Criegee intermediates are formed entirely in the excited state is an approximation that
 220 is made to simplify the estimations of their subsequent reactions, as discussed in Section 7. In most cases
 221 it is assumed that formation of all the possible Criegee and carbonyl combinations are equally likely, as
 222 indicated above. However, following the recommendations of Jenkin et al. (2020), we estimate that $\alpha =$
 223 0.1 if G_3 is $-\text{HCO}$, $-\text{CO}-$, $-\text{O}-$, or $-\text{OH}$ and no such group is bonded to G_1 , or 0.5 otherwise (or if G_3 is
 224 absent). In other words, formation of the carbonyl bonded to the oxygenated group is favored.

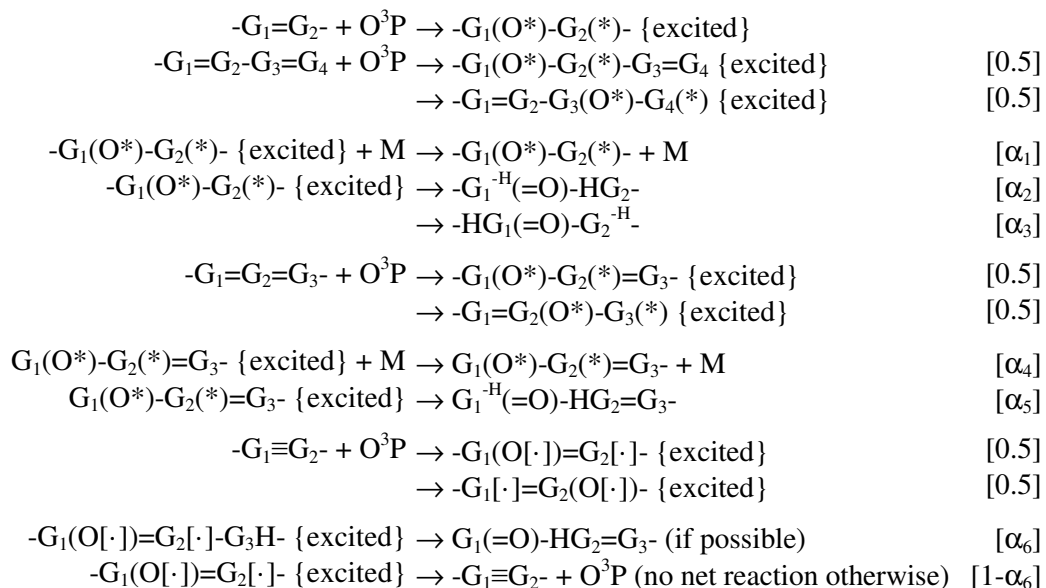
225 In the case of alkynes, the primary ozone adduct is assumed to directly form excited CIs with α -
 226 carbonyl substituents, with the two possible CIs formed in equal yields. However, it is assumed that the
 227 intermediates can rapidly interconvert by O shifting from the Criegee to the carbonyl group, so reactions
 228 of both of the isomers in this equilibrium need to be considered. The Criegee reactions are discussed in
 229 Section 7.

230

2.1.5. Additions of O³P to Unsaturated Bonds

231 O³P is assumed to react only by additions to double bonds, with rate constants estimated as
 232 indicated in Table 3d for isolated or conjugated double bonds and in Table 3e for additions to cumulated
 233 double bonds. Addition reactions involving O³P are expected to initially form a highly excited oxirane
 234 compound or biradical, which can stabilize, rearrange, or decompose to radicals. Based on model
 235 simulation results of high NO_x environmental chamber experiments performed when developing the
 236 SAPRC mechanisms for alkenes (e.g., Carter, 2000, 2010b), it is assumed that radical formation from
 237 reactions of O³P with C₃₊ organics are negligible, so only stabilization or rearrangements forming stable
 238 compounds are assumed to occur (as shown below). This is assumed for conjugated alkenes and alkynes
 239 as well, though this assumption is more uncertain. Although it is assumed that additions occur to both
 240 sides of the double bond at the same time, it is necessary to assign branching ratios in cases where there

241 are different possible rearrangements of the excited adducts. The general mechanisms are assumed as
 242 follows, with branching ratios assumed shown to the right in brackets.

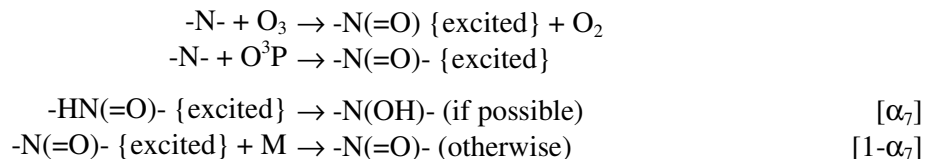


243 The only type of rearrangement considered for the initially formed adducts are H-shifts from one
 244 carbon to another, forming a carbonyl compound. In the case of additions to double bonds, it is assumed
 245 that stabilization occurs 50% of the time if at least one rearrangement is possible (i.e., $\alpha_1=\alpha_4=0.5$,
 246 $\alpha_2+\alpha_3=\alpha_5=0.5$) and all of the time if it is not (i.e., $\alpha_1=\alpha_4=1$ and $\alpha_2=\alpha_3=\alpha_5=0$). If more than one
 247 rearrangement is possible, they are assumed to be equally likely ($\alpha_2=\alpha_3=0.25$). In the case of additions to
 248 triple bonds, it is assumed that stabilization (not shown) is unlikely, and that there is no net reaction if the
 249 rearrangement shown is not possible ($\alpha_6=0$) and no back decomposition of the adduct otherwise ($\alpha_6=1$).

250 These assumptions are uncertain, but they have relatively little effect on model simulations of
 251 most atmospheric systems because reactions of O^3P are generally not important unless NO_2 is very high,
 252 such as in high NO_x environmental chamber simulations or in combustion/biomass burning plumes.

253 2.1.6. Reactions of O_3 and O^3P with Amines

254 The reactions of amines with O_3 or O^3P are assumed to form excited N-oxides, with rate constants
 255 estimated as shown in Table 3e, and parameters for other addition reactions as indicated in Table S-3.



256 In the case of primary or secondary amines, these excited N-oxides are assumed to rapidly rearrange and
 257 then stabilize to form -N-OH compounds ($\alpha_7=1$). In the case of tertiary amines, it is assumed that the N-
 258 oxide is stabilized ($\alpha_7=0$). The current system does not handle reactions of stabilized N-oxides so it treats
 259 them as unreactive. Their reactions and mechanisms are unknown.

260 2.2. Photolysis Reactions

261 Compounds with the groups -CHO, -CO-, -ONO₂, or the group pairs -O-OH, -O-O- are expected
262 to undergo photolysis at non-negligible rates under lower tropospheric conditions. These groups or group
263 pairs are referred to as "photoreactive groups" hereafter. Compounds not containing these photoreactive
264 groups may photolyze with UV wavelengths less than the atmospheric cutoff of ~290 nm, but photolyses
265 at this low wavelength regime are not currently considered in MechGen. Photolysis of radicals and CIs
266 are also ignored because it is assumed that loss of these intermediates by other reactions will dominate
267 over photolysis under atmospheric conditions.

268 Rate constants for photolysis reactions depend on the compound, reaction, and lighting
269 environment, and can be calculated by:

$$270 \quad J(\text{reaction, environment}) = J(\text{photolysis set, environment}) \times \phi_{\text{overall}}(\text{reaction})$$
$$271 \quad J(\text{photolysis set, environment}) = \sum_{\lambda} \sigma_{\lambda}(\text{photolysis set}) \times \phi_{\lambda}(\text{photolysis set}) \times F_{\lambda}(\text{environment})$$

272 where J is the first order rate constant for a photolysis reaction in a particular lighting environment,
273 "photolysis set" is a set of wavelength-dependent absorption cross-sections and quantum yields assigned
274 to various types of reactions, ϕ_{overall} is an overall wavelength-dependent quantum yield used for the
275 reaction of interest; if needed, λ is the wavelength, σ_{λ} and ϕ_{λ} are the wavelength-dependent absorption
276 cross-sections and quantum yields at wavelength λ in the photolysis set, and F_{λ} is the intensity of the light
277 source at wavelength λ , which is a function only of the environment, not the reaction.

278 MechGen makes no assumptions about the light source (other than having no significant intensity
279 at wavelengths below ~290 nm) and therefore does not output estimated or assigned photolysis rate
280 constants when generating photolysis reactions. Instead, it outputs the name of the photolysis set and the
281 value of overall quantum yield (ϕ_{overall}), if applicable, assigned to the reaction. The photolysis sets
282 assigned by MechGen are a subset of those incorporated in the SAPRC-16, -18 and -22 mechanisms
283 (Carter, 2016, 2020, 2023a), and are summarized in the SI on Table S-5. That table also indicates how
284 they were derived and gives half lives (with $\phi_{\text{overall}}=1$) at direct overhead sun using the actinic fluxes in the
285 reactivity scenarios of Carter (1994), and orders them by increasing half life. The files containing the data
286 for these photolysis sets are available online at the SAPRC website (Carter, 2023a, b). Photolysis rate
287 constants can be calculated using these data and are required for model application of mechanisms
288 generated using MechGen. Photolysis rate constants should be calculated for the light conditions of the
289 experimental or atmospheric conditions of interest. The mechanisms, photolysis sets, and (where
290 applicable) overall quantum yields for the various types of photolysis reactions are summarized below.
291 The photolyses of compounds with only a single photolysis group are described first, followed by a
292 discussion of treatments of photolyses of compounds with more than one such group. Note that if there is
293 more than one reaction, the branching ratios shown are implemented as overall wavelength-independent
294 quantum yields and are assumed not to depend on the spectrum of the light source.

295 In most cases the estimated mechanisms, absorption cross-sections, and quantum yields for
296 compounds with photoreactive groups are based on those derived for the smallest C₃₊ or C₄₊ compound
297 with the photoreactive group(s), with all the other groups being alkyl (-CH_x-) groups. This is clearly an
298 oversimplification, especially since the structure and size of the molecule, and the presence of non-alkyl
299 substituents, can affect the absorption cross-sections or quantum yields for at least some of the different
300 types of photoreactive compounds (Calvert et al., 2011). Although MechGen currently does not take these
301 differences into account, this needs to be considered in future versions.

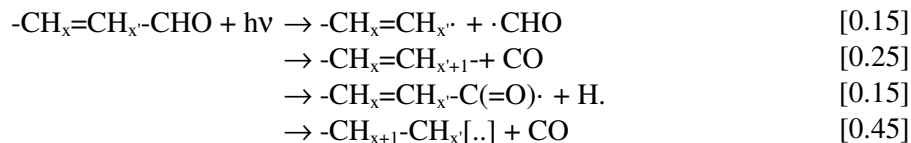
302 **2.2.1. Compounds with a Single Aldehyde Group**

303 Aldehydes with a single -CHO group that are not directly bonded to any unsaturated or
 304 photoreactive group, other than acetaldehyde and glycolaldehyde for which photolysis assignments are
 305 given explicitly (see SI), are assumed to photolyze as follows:



307 where "R" is any carbon-centered group other than methyl that does not contain a double bond on the
 308 group next to the -CHO. The absorption cross sections used are those recommended by IUPAC (2023) for
 309 propionaldehyde, and are given in photolysis set "C2CHOabs". IUPAC makes no specific
 310 recommendations concerning quantum yields or branching ratios, but it appears likely that the quantum
 311 yield $\cdot\text{CHO}$ formation for $\lambda > \sim 295$ nm is near unity, based on the data of Chen and Zhu (2001). MechGen
 312 assumes that this is applicable for higher aldehydes of this type as well.

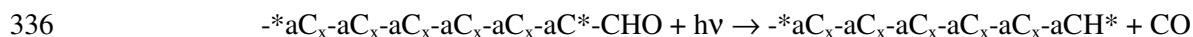
313 Aldehydes where the -CHO group is bonded to a C=C double bond group are assumed to
 314 photolyze with a mechanism based on that derived for acrolein, which is taken as representative for all α -
 315 unsaturated aldehydes, including those with conjugated or cumulated double bonds, and with non-alkyl
 316 (but non-photoreactive) substituents. The photolysis set giving the absorption cross sections and quantum
 317 yields used for acrolein is ACROL-16, while MACR-15 (based on methacrolein) is used for the others,
 318 though the acrolein mechanism is used in all cases to predict the subsequent reactions that occur. This
 319 gives half lives of ~ 50 and ~ 60 hours, respectively, for loss by photolysis for direct overhead sunlight,
 320 which means that photolysis is relatively slow but not necessarily negligible. The assumed mechanism is
 321 as follows:



322 where $x=0-2$, $x'=0-1$, and $[\cdot]$ indicates a carbene radical group, whose subsequent reactions are discussed
 323 in Section 6.3. These branching ratios are based on the data of Gardner et al. (1987) for acrolein, as
 324 shown on Figure IX-C-7 of Calvert et al. (2011) for air at 1 atm. These are assumed to apply to photolyses
 325 of other unsaturated aldehydes as well.

326 Aldehydes where the -CHO is bonded to aromatic groups are assumed to photolyze with the same
 327 rate constant and a similar mechanism as estimated for benzaldehyde, regardless of substituents. The
 328 photolysis set used is "BALD-11", where absorption cross-sections are those recommended for
 329 benzaldehyde by Calvert et al. (2002). In addition, $\phi_{\text{overall}} = 0.06$ is used to approximately fit consumption
 330 rates of benzaldehyde in chamber experiments (Carter, 2000)).

331 Benzaldehyde acts as an efficient radical terminator when added to environmental chamber
 332 experiments (Carter et al., 1982; Carter, 2000), suggesting that photolyses of aromatic aldehydes do not
 333 form radicals. (The reaction with OH radicals is also predicted to be radical terminating, but that would
 334 not be sufficient to compensate for radical initiation due to photolysis if the quantum yield for radical
 335 production were non-negligible.) For mechanism generation purposes, these are represented as follows:



337 where $\text{-aC}_x\text{-}$ refers to either -aC- or -aCH- and "*" indicates ring closure. The actual photolysis reaction is
 338 highly uncertain, and it is likely that other products are actually formed, since we know of no evidence
 339 that aromatic hydrocarbons are formed in high yields from the photolyses of aromatic aldehydes.

340 Aldehydes where the -CHO is bonded to an alkyne group, e.g., -C≡C-CHO, are assumed not to
 341 photolyze. We could not find data concerning absorption cross-sections or photolyses of these
 342 compounds, and assume that they either do not photolyze, or do so relatively slowly. Data are needed to
 343 test this assumption.

344 2.2.2. Compounds with a Single Ketone Group

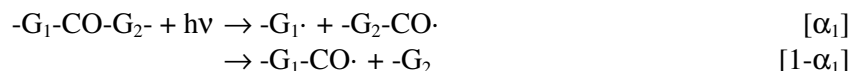
345 C₄₊ ketones with a single -CO- group that is not directly bonded to any unsaturated or
 346 photoreactive group are assigned photolysis sets giving absorption cross-sections and overall quantum
 347 yields as indicated in Table 4. In the case of acetone, the wavelength-dependent quantum yields
 348 recommended by IUPAC (2023) for atmospheric temperature and pressure are used. In the case of the
 349 higher ketones, the photolysis set only has absorption cross-sections, and overall quantum yield factors
 350 are derived so that the SAPRC mechanism simulates environmental chamber data for ketones (Carter,
 351 2010b). These data suggest that the photolysis rate constants, and therefore overall quantum yield factors,
 352 decrease with carbon number for C₄₊ ketones. The structure of the ketone and the presence of non-alkyl
 353 but non-photoreactive substituents, other than -O- groups next to the carbonyl (i.e., esters), are assumed
 354 not to affect the absorption cross-sections or quantum yields.

355
 356 Table 4. Assignments used to estimate photolysis rate constants for compounds with single ketone
 357 groups based on the numbers of carbons in the ketone.

C's	Photolysis Set	ϕ_{overall}	$t_{1/2}$ [a]	Based on Data for Compound
3	ACET-06 (σ_{λ} and ϕ_{λ} for acetone)	1	~180 hours	Acetone
4	MEK-06 (σ_{λ} 's for MEK)	0.175	~70 hours	methyl ethyl ketone (MEK)
5	Same as above	0.1	~120 hours	2-pentanone
6	Same as above	0.1	~120 hours	methyl isobutyl ketone
7+	Same as above	0.02	~600 hours	2-heptanone

358 [a] Half life relative to the photolysis rate constant for ketone photolysis calculated for direct overhead
 359 sunlight (z=0) using the actinic fluxes used in the reactivity scenarios of Carter (1994).

360
 361
 362 The photolysis of these ketones are assumed to proceed only by breaking one of the two C-CO
 363 bonds, forming an acyl and an alkyl radical.

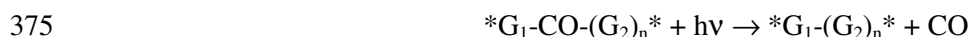


364 Here G is any non-photoreactive group, and α_1 is the branching ratio for the formation of radical G₁·. The
 365 relative branching ratios for breaking CO-alkyl bonds are assumed to depend on whether the radical
 366 formed is a methyl, secondary, primary, or tertiary, with equal branching ratios if both radicals are of the
 367 same type. Based on IUPAC (2023) data for methyl ethyl ketone, we assume methyl vs. ethyl branching
 368 ratios of 15% and 85%, respectively, for that compound. Extrapolating this to other higher compounds,
 369 we use the following branching ratios, where $\alpha=0.5$ is used if both are of the same type:

\underline{G}_1	\underline{G}_2	α_1	\underline{G}_1	\underline{G}_2	α_1	\underline{G}_1	\underline{G}_2	α_1
-CH ₂ -	-CH ₃	0.85	-CH()-	-CH ₃	1	-C()()-	-CH ₃	1
-CH()-	-CH ₂ -	0.85	-C()()-	-CH ₂ -	1	-C()()-	-CH()-	0.85

370 Note that this is somewhat arbitrary and uncertain (except for methyl ethyl ketone), and is assumed not to
 371 be affected by the presence of non-alkyl substituents, which is even more uncertain.

372 If the ketone group is in a ring, the reaction would form excited biradicals that should re-form the
373 starting ketone if stabilized. In this case, it is assumed that the excited biradical primarily decomposes by
374 CO elimination, i.e.:

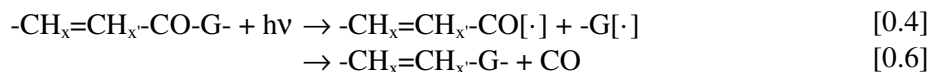


376 where "*" indicates closure for any size ring.

377 Esters, carbonates, and acids, i.e., compounds with -CO-O- or -CO-OH group combinations, do
378 not appear to have significant absorption in the atmospheric wavelength region, and are assumed not to
379 photolyze.

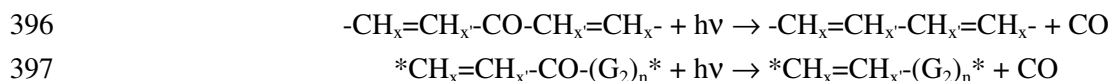
380 Estimations of photolyses of ketones with an adjacent double bond group are based on data for
381 methyl vinyl ketone (MVK). These reactions are assigned the photolysis set "MVK-16", which contains
382 IUPAC (2023) recommendations for absorption cross-sections and quantum yields of methyl vinyl ketone
383 at 1 atm. This gives an estimated half life of ~36 hours for photodecompositions of these unsaturated
384 ketones with direct overhead sunlight, and gives reasonably good fits to NO oxidation rates in MVK -
385 NOx chamber experiments.

386 The photolysis reactions for ketones with only a single adjacent double bond are assumed to be as
387 follows:



388 Here, x=0-2, x'=0-1, and G is any non-photoreactive group with only saturated bonds. This is based on
389 IUPAC (2023) recommendations for MVK but is assumed to apply to all compounds of this type.

390 Ketones with double bond groups on both sides of the carbonyl group are estimated to photolyze
391 with the same absorption cross-sections and quantum yields as those bonded to only one double bond
392 group, but it is assumed that formation of vinylic radicals is not as favorable based on thermochemical
393 estimates. These reactions are assumed to occur only via CO formation, as shown in the 2nd reaction
394 above. This is also assumed for unsaturated ketones where the -CO- group is in a ring, where the first
395 reaction is not expected to occur for the reasons discussed above.



398 Estimations of photolyses of ketones with an adjacent aromatic or triple bond group are not
399 supported by MechGen, so no photolysis reactions are generated for these compounds. The need to
400 represent photolyses of these compounds has not been investigated, but it is expected that
401 photodecompositions of aromatic ketones will be relatively slow, and ketones with triple bonds are not
402 expected to be important in atmospheric systems.

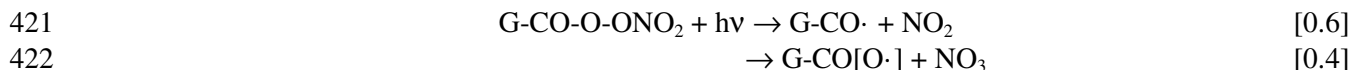
403 **2.2.3. Organic Nitrates, Carbonyl Nitrates and Carbonyl Peroxynitrates**

404 Compounds with a single organic nitrate group and no other photoreactive groups are assumed to
405 photolyze entirely by scission of the weak O-NO₂ bond, forming NO₂ and an alkoxy or acyloxy radical
406 whose reactions are discussed in Section 5.



408 These reactions are assigned the "IC3ONNO2" photolysis set with quantum yields of 1 at all wavelengths,
409 which gives a half life of ~50 hours for loss due to photolysis with direct overhead sunlight. This is based
410 on the IUPAC (2023) recommendation for isopropyl nitrate, but is assumed to apply to all organic nitrates
411 without other photoreactive groups, regardless of structure, the presence of non-alkyl substituents, and the
412 presence of unsaturated bonds. This is an approximation because isopropyl nitrate has stronger
413 absorptions than for n-propyl and smaller nitrates, but is taken as representative. Although the accuracy of
414 this approximation is uncertain in some cases, loss by photolysis is relatively unimportant compared to
415 other reactions.

416 In the case of acyl peroxy nitrates (e.g., PANs), the photolyses are assigned "PAN-11" or
417 "PPN-11" photolysis sets, with the former used for PAN itself and the latter used for all other PAN
418 analogues based on peroxypropionyl nitrate (PPN). These contain only absorption cross-sections, since
419 unit quantum yields are assumed. They give approximate photodecomposition half lives of ~200 and
420 ~100 hours, respectively, for direct overhead sunlight. The assumed mechanism is as follows:



423 This is based on Sander et al. (2006) recommendations for PAN, but is applied to all PAN analogues
424 regardless of substituents or the presence of multiple bonds.

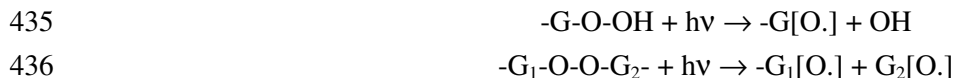
425 In the case of non-acyl (e.g., alkyl) peroxy nitrates, photolysis is assumed to proceed primarily by
426 NO_2 formation,



428 with the same rate constant assignments as used for simple organic nitrates. However photolysis is not
429 important for these compounds because they are assumed to undergo rapid thermal unimolecular
430 decomposition, forming the same products, as discussed in Section 2.3.

431 **2.2.4. Compounds with a Single set of Peroxide Groups**

432 Compounds with peroxy or hydroperoxy groups and no other photoreactive group are assumed to
433 photolyze entirely by scission of the weak O-O bond, forming alkoxy radicals and (for hydroperoxides)
434 OH.



437 The reactions are assumed to have unit quantum yields at all wavelengths, and are assigned the photolysis
438 set "COOH", which contains the absorption cross-section of methyl hydroperoxide (IUPAC, 2023), and
439 gives a half life of ~30 hours for loss by photolysis with direct overhead sun. Note that it is assumed that
440 organic peroxides (which tend to be relatively less important than hydroperoxides in atmospheric
441 systems) photolyze with approximately the same rate constant as hydroperoxides, and that the size,
442 structure, and presence of non-photoreactive substituents or unsaturated bonds do not have a significant
443 effect. The most questionable assumption in this case is the assumption that the presence of adjacent
444 unsaturated bonds has no effect on hydroperoxide or peroxide photolysis.

445 2.2.5. Compounds with More than One Photoreactive Group

446 The photolysis sets, quantum yields, and photolysis reactions used for compounds with various
447 combinations of photoreactive groups are summarized in Table 5. If the ϕ_{overall} column is blank, then no
448 overall quantum yield correction is applied and the photolysis rate constant is the same as that calculated
449 for the photolysis set. The half lives relative to photodecomposition for direct overhead sun, which
450 incorporate the overall quantum yield corrections if applicable, are also shown in the table. No corrections
451 are made for non-alkyl substituents or unsaturated bonds on the molecule except as indicated in the table.
452 The considerations used when making these assignments are discussed below.

453 The assignments for some (but not all) types of reactants with adjacent photoreactive groups are
454 made based on available data for the simplest or representative molecule of the types, i.e., methyl glyoxal,
455 biacetyl, peroxy acetic acid, PANs, carbonyl nitrates, dinitrates, and various unsaturated 1,4-dicarbonyls.
456 The applicability to higher molecular-weight compounds, compounds with non-alkyl substituents, or
457 compounds with multiple bonds is uncertain, but not as uncertain as the assignments for the reactants in
458 the table where data for representative compounds are either not available or were not considered.

459 It is assumed that -CO-O- groups are not photoreactive, so the presence of this group combination
460 is not considered when making photolysis assignments. The exception is compounds with -CO-O-ONO₂
461 group combinations (PAN analogues), where assignments are made based on data for PAN and
462 peroxypropynyl nitrate (PPN) as discussed above.

463 The unsaturated 1,4-dicarbonyls with at least one aldehyde group are highly photoreactive
464 aromatic ring opening products (Calvert et al., 2011; see also Section 3.4, below) that account for the
465 relatively high reactivity of aromatic hydrocarbons observed in environmental chamber experiments. The
466 AFGS photolysis file is used to give absorption cross sections for all these compounds, with an overall
467 wavelength-independent quantum yield adjusted to give satisfactory fit to results of aromatic - NO_x
468 chamber experiments, which are highly sensitive to this photolysis (e.g., Carter, 2010b; Carter and Heo,
469 2013). The SAPRC-22 mechanism (Carter, 2023a) that was derived using this version of MechGen best
470 fit the data using effective quantum yields close to 25%. However, the quantum yields in the AFGS file
471 do not take into account absorption cross sections reported for 4-oxo-2-pentenal reported by Xiang et al.
472 (2007), which are the basis of the current IUPAC (2023) recommendations for this compound. This is
473 because these quantum yields, which are much lower than the previous data for these compounds, could
474 not be used as a basis for a predictive mechanism for aromatics because fitting the chamber data required
475 using an effective quantum yield of greater than 3. Therefore, either there are additional sources of
476 radicals in aromatic photooxidation systems that are not taken into account in the current estimated
477 mechanisms for aromatics, the absorption cross sections of these 1,4-dicarbonyls are highly dependent on
478 structure, or the data of Xiang et al. (2007) have systemic issues. Until this situation is resolved and
479 MechGen can be updated to derive a more predictive mechanism, we will continue to use the present
480 cross sections in AFGS to derive aromatics mechanisms for airshed models.

481 It is assumed that unsaturated 1,4-diketones do not photolyze sufficiently rapidly in ambient
482 sunlight that it is necessary to include their photolysis in atmospheric models. This is consistent with the
483 discussion of photolyses studies of 3-hexene, 2,5-dione given by Calvert et al. (2011), which indicate that
484 radical formation is relatively minor compared to isomerization. This is also consistent with the fact that
485 para-substituted dialkyl benzenes, whose reactions can form these compounds, tend to be less reactive
486 than other isomers, all else being equal (Carter and Heo, 2013).

487 Table 5. Summary of photolysis sets, overall quantum yields, approximate photolysis half lives,
 488 and products formed for compounds with more than one photoreactive group.

Reactant [a]	Photolysis Set [b]	ϕ_{overall}	$t_{1/2}$ [c]	Products Formed [d]	
<u>Adjacent photoreactive groups</u>					
HCO-CO-G-	MGLY-13 (methyl glyoxal)		40 min.	HCO[·] + G-CO[·]	
-G ₁ -CO-CO-G ₂ -	BACL-11 (biacetyl)		25 min.	G ₁ -CO[·] + G ₂ -CO[·]	
-G-CO-ONO ₂	IC3ONO2 (isopropyl nitrate σ 's)		50 hrs.	G-CO[O.] + NO ₂	
-G-CO-O-OH	PAA (peroxy acetic acid σ 's)		200 hrs.	G-CO[O.] + OH	
-G ₁ -CO-O-O-G ₂ -	COOH (methyl hydroperoxide σ 's)		30 hrs.	G ₁ -CO[O.] + -G ₂ O[·]	
-G-CO-O-ONO ₂	PPN-11 (PPN σ 's) [e]		200 hrs.	See Section 2.2.3	
<u>Separated photoreactive groups (x>0, no upper limit)</u>					
HCO-G ₁ =G ₂ -CHO	AFGS (various 1,4-unsaturated dicarbonyl σ 's, see text)	0.45	4 min.	Gs ₁ -CO[·] + H., & Gs ₂ -CO[·] + H. (equal)	
HCO-(G) _x -CHO [f]	C2CHO (propionaldehyde)		8 hrs.	Same as -Gs-CHO	
HCO-(G) _x -CO-	C2CHO (propionaldehyde)		8 hrs.	Same as -Gs-CHO	
HCO-G ₁ =G ₂ -CO-G ₃ .	AFGS (see above)	0.45	4 min.	Gs ₂ -CO[·] + G ₃ [·]	
HCO-(G) _x -ONO ₂	C2CHOabs (propionaldehyde σ 's)		3 hrs.	Same as -Gs-ONO ₂	
xCO-G ₁ =G ₂ -(G) _x -O-OX	HPALDS (acroleins σ 's, see text)	0.1	3 hrs	Same at -Gs-O-OX	
HCO-(G) _x -O-OX	C2CHOa (propionaldehyde σ 's)		3 hrs.	Same at -Gs-O-OX	
HCO-(G) _x -O-ONO ₂	C2CHOa (propionaldehyde σ 's)		3 hrs.	Same as -Gs-O-ONO ₂	
G ₁ -CO-G ₂ =G ₃ -CO-G ₄ -	Does not photolyze (see text)				
-G ₁ -CO-(G) _x -CO-G ₂ - [f]	MEK-06 (methyl ethyl ketone [MEK] σ 's)	6 C's 7+ C's	0.1 0.02	70 hrs. 120 hrs.	Same as G ₁ -CO-Gs-, & G ₂ -CO-Gs- (equal)
-G ₁ -CO-(G) _x -ONO ₂	CRBNIT (carbonyl nitrate σ 's)		4 hrs.	Same as -Gs-ONO ₂	
-G-CO-(G) _x -O-OX	MEK-06 (MEK σ 's)		12 hrs.	Same at -Gs-O-OX	
-G-CO-(G) _x -CO-ONO ₂	CRBNIT (carbonyl nitrate σ 's)		4 hrs.	Same as Gs-CO-ONO ₂	
-G-CO-(G) _x -CO-O-ONO ₂	CRBNIT (carbonyl nitrate σ 's)		4 hrs.	Same as Gs-CO-O-ONO ₂	
X ₁ O-O-(G) _x -O-OX ₂	COOH (methyl hydroperoxide σ 's)		30 hrs.	Same as Gs-O-OX ₁ + Gs-O-OX ₂ , each equal	

489 [a] Group combinations that give unique compounds for which specific assignments are used (e.g.,
 490 glyoxal and HNO₃) are not shown. "-OX" is -O- or -OH. "xCO-" is HCO- or -CO-.

491 [b] The derivations of the photolysis sets are given in Table S-5 in the SI. " σ 's" means that the photolysis
 492 set contains only absorption cross-sections. "(equal)" means that both possible reactions occur with
 493 equal probability.

494 [c] Approximate half lives relative to loss of the compound by photolysis for direct overhead sunlight,
 495 including the contribution by the assigned wavelength-independent quantum yield, if applicable.

496 [d] "Gs" refers to a combination of other groups on the molecule (other than G, G₁ or G₂), including the
 497 other photoreactive group, which is assumed not to be transformed by the reaction.

498 [e] The photolysis set "PAN-11", giving an approximate photolysis half live of 100 hours for direct
 499 overhead sun, is used for PAN itself (G = CH₃).

500 [f] Not applicable if -(G)_x- is -CH_x=CH_x-, as shown for previous reactant.

501 If a reactant has more than one non-adjacent photoreactive group of the same type it is assumed
502 that the photolysis reaction at each group occurs with equal probability, with the mechanism the same as
503 if there were no other photoreactive group. However, the total photolysis rate constant for reactions at
504 both groups is estimated to be the same as if the compound had only one group since the same photolysis
505 set is used. This is uncertain.

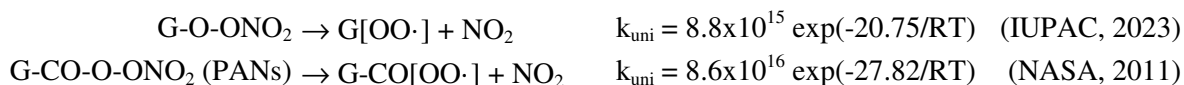
506 If a reactant has more than one non-adjacent photoreactive group of different types it is assumed
507 that the absorption cross-sections can be approximated by those of compounds with the single group that
508 has the highest absorption cross-section at atmospheric wavelengths. On the other hand, the mechanisms
509 (e.g., photolysis products) are assumed to be those resulting from reaction at the group with the weakest
510 bond. This is based on the assumption that once the photon is absorbed the energy is rapidly distributed
511 around the molecule. This is consistent with the data of Wolfe et al. (2012), which suggest that α -
512 unsaturated carbonyls with hydroperoxide groups photolyze at rates consistent with those calculated using
513 absorption cross-sections of α -unsaturated carbonyls, but with unit quantum yields and with the reaction
514 breaking the peroxy bond forming OH. This gives a photolysis rate that is about 100 times faster than
515 simple hydroperoxides. However isoprene and 1,3-butadiene NO_x experiments are not well simulated
516 with this high of a photolysis rate for α -unsaturated hydroperoxy carbonyls (Carter et al, unpublished
517 results), so we arbitrarily cut the rate by a factor of ~ 10 by using an effective quantum yield of 0.1 for
518 these compounds. The current system assigns unit quantum yields for saturated hydroperoxy or nitrate-
519 substituted carbonyls, though this is highly uncertain and the estimates are probably upper limits.

520 Despite the differences in assigned overall quantum yields, the photolysis rate constants
521 calculated for saturated carbonyls with other photoreactive groups are about the same as for those that are
522 unsaturated (see Table 5). This is due to the higher absorption cross-sections assigned to unsaturated
523 aldehydes.

524 2.3. Unimolecular Reactions

525 2.3.1. Peroxynitrates

526 Peroxynitrates are formed from the reactions of peroxy or acyl peroxy radicals with NO_2 , but the
527 O- ONO_x bond is weak enough that these can thermally decompose back to reactants at significant rates at
528 atmospheric temperatures.



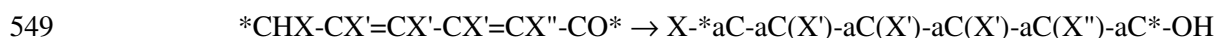
529 Here, "G" is any saturated or unsaturated group and the rate constants are in s^{-1} , T is the temperature in K,
530 and $R = 1.99 \times 10^{-3} \text{ K mole kcal}^{-1}$. The presence of non-alkyl substituents, unsaturated bonds or other
531 structural factors are assumed not to affect these rate constant estimates or decomposition mechanisms.
532 The OO- NO_2 bonds are currently believed to be the only bonds in compounds formed in atmospheric
533 systems to be weak enough to undergo simple scission at non-negligible rates at atmospheric
534 temperatures. (The weak O-O bonds in peroxy compounds are at least 10 kcal/mole stronger.)

535 The decompositions of the non-acyl peroxynitrates are estimated to have a half life of $\sim 5 \text{ sec}^{-1}$ at
536 298K, which is sufficiently fast that the formation of these compounds is rapidly reversed. Therefore, the
537 default is for MechGen to ignore the formation and therefore the decomposition of these compounds
538 when multi-step mechanisms are generated for atmospheric conditions. However, they do show up when
539 single step mechanisms are generated (Carter, In Prep.). On the other hand, the acyl peroxynitrates
540 (PANs) decompose much slower and are observed as photooxidation products in atmospheric systems, so
541 their formation cannot be ignored. It cannot be ruled out that some α substituents other than -CO- may

542 slow down the decomposition sufficiently that peroxy nitrates formation for some non-acyl peroxy nitrates
543 may be non-negligible. However, this is not currently considered in MechGen.

544 **2.3.2. Aromatic formation from Cyclohexadienones**

545 The other type of non-radical unimolecular reaction that is currently considered involves
546 formation of aromatic rings from 2,4-cyclohexadienone compounds containing a -CHX-CO- group pair
547 in the ring, where an H-shift from the -CHX to the carbonyl group is expected to be highly favorable
548 energetically.



550 MechGen predicts that cyclohexadienones are formed in high yields in the reactions of phenolic
551 compounds if this reaction does not occur (see Section 3.4), but they are not observed as products. On the
552 other hand, catechols (the products of this type of isomerization reaction) are observed in high yields in
553 the reactions of OH with phenols (e.g., Olariu et al., 2002; Berndt and Böge, 2003), which would not be
554 expected if this reaction did not occur (Xu and Wang, 2013). The nature of the "X" group should not have
555 an obvious effect on how fast this reaction occurs, so if this is fast then it is reasonable to expect
556 compounds with other "X" substituents, including X=alkyl or H, are also fast. MechGen assumes that this
557 is the case. The rate constant is unknown but it is assumed to be high in order to account for the observed
558 formations of catechols from phenols. This assumption is implemented by treating such compounds like
559 rapidly reacting radicals when the reactions are generated.

560 It is possible that this reaction may require the presence of surfaces and/or water to occur at
561 sufficient rates. The H-shift involves a strained 4-member ring transition state, which may be catalyzed by
562 surfaces, while the involvement of water would result in the same H-shift but with a 6-member ring
563 transition state. Given surface- or water-free environments rarely occur in the ambient atmosphere it is
564 probably better that these reactions be included for modeling applications than being neglected. For
565 example, the predicted formation of catechols from phenols due to this reaction has implications in
566 predictions of SOA from aromatics.

567 **2.3.3. Other Compounds**

568 Decompositions of peroxy nitrates and rearrangements of cyclohexadienones are currently the
569 only types of non-radical unimolecular reaction considered by MechGen. In particular, MechGen does not
570 consider other types of water- or surface-catalyzed conversion of non-radical products that could affect
571 the environmental fate of multi-functional products. For example, geminal diols, compounds with HO-
572 CX₂-OH groups, are predicted to be formed in some cases and would be expected to be in equilibrium
573 with H₂O and XC(=O)X in the environment. It is uncertain whether this or similar reactions would occur
574 at significant rates in the gas phase, and MechGen currently assumes that this is not the case. However,
575 ignoring these potentially heterogeneous or H₂O-catalyzed reactions may also have implications in SOA
576 predictions.

577 **3. Reactions of Carbon-Centered Radicals**

578 Carbon-centered radicals are formed in the initial reactions of most VOCs with atmospheric
579 oxidants, from H-shift isomerization reactions of peroxy and alkoxy radicals, and from most alkoxy
580 radical decompositions. Under lower atmospheric conditions, their major fate is expected to be reaction
581 with O₂, which is estimated to occur with a pseudo-unimolecular rate constant of $\sim 3.8 \times 10^7 \text{ sec}^{-1}$ (see
582 Table S-6 for measured radical + O₂ rate constants and associated text in the SI). However, rapid
583 unimolecular reactions are estimated to dominate for some types of carbon-centered radicals formed in

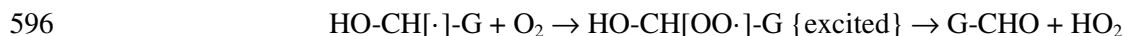
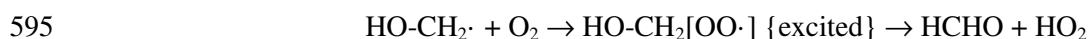
584 generated mechanisms, as discussed below. Note that there are currently no cases where MechGen
 585 assumes that both O₂ and unimolecular reactions of carbon-centered radicals occur at competitive rates --
 586 one or the other is assumed to dominate. This means that estimates of carbon-centered radical + O₂ rate
 587 constants are not currently needed or used in the generated mechanism, though estimates of these rate
 588 constants are used when considering when unimolecular reactions may dominate.

589 3.1. Reaction with O₂

590 Most carbon-centered radicals are assumed to primarily add O₂ and form the corresponding
 591 peroxy radical, whose reactions are discussed in Section 4.

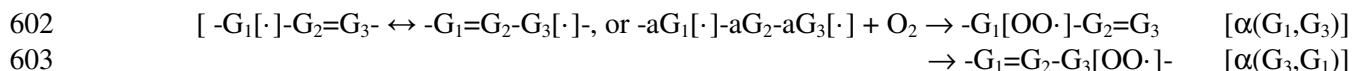


593 However, the peroxy radicals formed from α -OH substituted radicals are assumed to rapidly decompose
 594 to form HO₂ and the corresponding carbonyl.

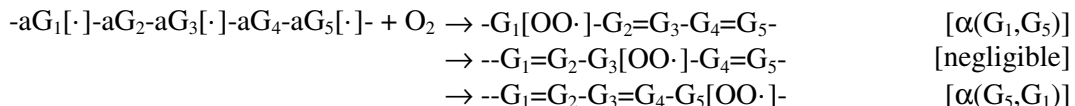


598 MechGen treats these as concerted processes when reactions of α -OH substituted carbon-centered
 599 radicals are generated, since stabilization and other reactions are expected to be unimportant.

600 If the carbon-centered radical is adjacent to one or more double bond groups, it is assumed to be a
 601 resonance structure where O₂ can add to any of the radical positions, e.g.:



604 where $\alpha(G_a, G_b)$ is the fraction of adding to the G_a position when G_b is the other position on the radical.
 605 Additions to radicals with more than one double bond are assumed to form only those adducts with the
 606 most conjugated structures, e.g.:



607 Because of lack of data indicating otherwise, we assume that the branching ratios, α , depend only on the
 608 nature of the radical groups where the addition occurs, regardless of the number of double bond groups in
 609 the radical. Although this has not been examined comprehensively, we believe that all allylic radical
 610 structures that may arise in generated mechanisms have only two radical centers where O₂ addition forms
 611 the most conjugated structures. This is supported by the fact that we have not found examples of where
 612 this is not the case when generating many mechanisms.

613 Available information and estimates concerning branching for allylic additions is discussed in
 614 Section S1.4.2. There are very limited data, but what is known does not support the assumption that
 615 additions at each position are equally likely. Product data for the reactions of 1,3-butadiene and isoprene
 616 with OH indicate that in radicals with both primary (-aCH₂[\cdot]) and secondary (-aCH[\cdot]-) radical positions,
 617 the addition occurs ~65% of the time at the secondary position. This is consistent with the fact that
 618 thermochemical analysis indicates that addition to the secondary radical is ~2 kcal/mole more favorable
 619 (see SI). In addition, theoretical calculations of Xu and Wang (2013) on phenol mechanisms and of Yuan
 620 et al. (2017) on furan mechanisms both indicate that O₂ preferentially adds to radical centers with -OH or

621 -OR substituents, instead of those with unsubstituted secondary radical centers. However, if this is
 622 assumed then the mechanisms significantly underpredict reactivities observed in environmental chamber
 623 experiments with phenolic compounds, and better fits to the data are obtained if it is assumed that
 624 addition to non-OH-substituted secondary radical centers occur approximately 15% of the time (Carter et
 625 al, unpublished results). This is highly uncertain, but it provides an additional data point for deriving
 626 general estimation method incorporating -OH substituted radicals.

627 As discussed in Section S1.4.2, we assume that the branching ratios can be estimated based on the
 628 differences in heats of addition of O₂ to the different radical positions, with a correction included if one of
 629 the radical centers is -OH or -OR substituted. The estimate used is as follows:

630
$$\alpha(G_1, G_2) = \text{WtF}(G_1) / [\text{WtF}(G_1) + \text{WtF}(G_2)]$$

631 where
$$\text{WtF}(G) = \text{Weighting factor for } G = e^{-\beta \times [\Delta\Delta\text{Hr}(G) - O_{\text{corr}}(G)]}$$

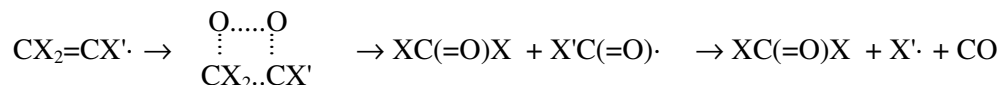
632 and $\Delta\Delta\text{Hr}(G)$ is the difference of heats of reaction for O₂ addition to G compared to a primary radical
 633 center in a radical that has both, β is a parameter assigned a value of 0.33 to be consistent with the
 634 product data for isoprene and 1,3-butadiene, and O_{corr} is set at 5.78 kcal/mole if G is -OH or -OR
 635 substituted to be consistent with modeling results for phenols, or zero if there are no such substituents.
 636 The weighting factors so derived are as follows (see Section S1.4.2 for derivations):

<u>G·</u>	<u>WtF(G)</u>	<u>G·</u>	<u>WtF(G)</u>	<u>G·</u>	<u>WtF(G)</u>
-aCH ₂ [·]	1	-aCH[·]-CO-X	0.35	-aCH[·]-OX	21.9 *
-aCH[·]-R	1.9	-aC[·](R)-CO-X	0.28	-aC[·](R)-OX	13.6 *
-aC[·](R)-R	1.13	(O _{corr} not used)		* O _{corr} used	

637 Note that use of this methodology to estimate branching ratios in general is highly uncertain because only
 638 two data points are available to derive the two parameters used (β and O_{corr}), with no data available to test
 639 this estimation method. However, this is the best that can be done at the present time.

640 The above discussion is not applicable to radicals that are estimated to have rapid unimolecular
 641 reactions as discussed in Section 3.2. If such a rapid reaction is possible for one of the radical centers in
 642 an allylic radical, it is assumed that all the reaction goes by this route unless a rapid reaction is possible at
 643 the other radical center, in which case it is assumed that they occur with equally probability. The above
 644 discussion is also only applicable to stabilized radicals, and not necessarily to excited adducts formed
 645 when OH or NO₃ adds to double bonds. In most cases excited adducts formed following additions to
 646 double bonds are assumed to be primarily stabilized and then react as discussed above, but that is not
 647 always the case, as discussed in Section 3.3.

648 A different mechanism is assumed for the addition of O₂ to vinylic radicals where the radical
 649 center is on an unsaturated carbon. In this case, the reaction with O₂ is expected to involve the O₂ adding
 650 to each side of the bond, with the bonds rearranging to form a carbonyl product and a carbonyl radical,
 651 i.e.,



652 This is based on data of Slagle et al. (1984) for vinyl and Slagle et al. (1988) for methylvinyl and i-C₄H₉·.
 653 More recently, Matsugi and Miyoshi (2014) directly measured the yield of HCO radicals from vinyl, and
 654 observed that the yields were only about 0.2, independent of pressure, with the remainder of the reaction
 655 being prompt dissociation to H + CO. This is assumed to be applicable to additions to radicals of any size,
 656 though it is possible that rapid energy distribution following reaction of larger vinyl species with O₂ could

657 lead to stabilized vinylperoxy radicals. However, insufficient data are currently available to quantitatively
 658 assess the degree to which this occurs, so MechGen assumes that the stabilization is not important. The
 659 issue of whether the HCO promptly decomposes is not relevant to atmospheric modeling, since the same
 660 products (HO₂ and CO) are formed under atmospheric conditions in either case. Therefore, MechGen
 661 represents the reactions of CX₂=CH· as forming H· + CO alone, since this is the major overall process.

662 However, the issue of prompt dissociation is relevant in the case of RC(=O)·, since the stabilized
 663 radical is expected to primarily add O₂ to form an acyl peroxy radical, which can give different products
 664 than if RC(=O) decomposes to R· + CO. Decomposition is expected to be more favorable because the
 665 lower estimated heat of reaction, though the rate of stabilization of the RC(=O)· should also be greater
 666 than for HC(=O)·. MechGen assumes that the energetics is the larger factor in the case of additions of O₂
 667 to vinylic radicals, assumes that the prompt dissociation dominates, and also represents the overall
 668 reaction as forming the carbonyl, CO, and R· as a single process.

669 3.2. Unimolecular Reactions of Stabilized Radicals

670 Several types of carbon-centered radicals are expected to undergo unimolecular reactions
 671 sufficiently rapidly that unimolecular reaction is assumed to dominate over reaction with O₂, even for
 672 thermalized radicals. Note that if these reactions are assumed to be fast for thermalized radicals, they are
 673 necessarily assumed to be fast when the radicals are excited.

674 3.2.1. Rapid α-Scission Decompositions

675 Carbon-centered radicals with adjacent hydroperoxy, peroxide, nitrate, or dicarbonyl groups are
 676 expected to rapidly undergo decompositions by simple scission of an adjacent bond, forming products
 677 that are more energetically favorable.



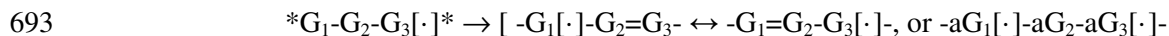
682 We expect that there should not be large entropy or energy barriers to these reactions (e.g. Green et al.,
 683 1990; Vereecken et al., 2004, Vereecken, 2008), so MechGen assumes they all dominate over O₂ addition
 684 whenever radicals of these types are formed. Note that if the peroxy, nitrate, or carbonyl substituent is
 685 next to an allylic radical center, the bond scission reaction is still expected to dominate over O₂ addition,



687 so there is no O₂ addition to the other radical centers. Reactions of allylic radicals with peroxy, nitrate, or
 688 dicarbonyl groups are treated similarly.

689 3.2.2. Cyclopropyl Ring Opening Decompositions

690 Carbon-centered radicals with the radical center in a cyclopropyl ring can decompose by breaking
 691 the bond opposite the radical center, relieving the ring strain and forming an energetically favorable
 692 allylic stabilized radical.



694 This reaction is not expected to have large or any barriers, so is assumed to dominate over O₂ addition.
 695 On the other hand, the analogous reactions when the radical center is in a cyclobutane ring is estimated to

696 be much less energetically favorable and is only expected to occur in chemically activated radicals, as
697 discussed in Section 3.3. The reactions for larger rings are estimated to be endothermic and are assumed
698 to be unimportant relative to stabilization and/or O₂ addition even for excited radicals.

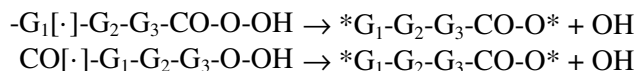
699 3.2.3. Cyclizations forming 5-Member Ring Cyclic Esters

700 As discussed in Section S1.4.3, Curran et al. (1998) provided estimates for the Arrhenius
701 parameters for decompositions of various QOOH radicals of the type,

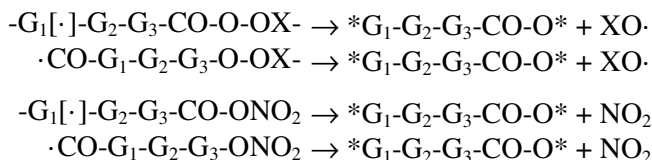


703 as a function of cyclic ether ring size. These can be used as a basis for estimating the approximate
704 importance of these and similar reactions in atmospheric systems. In general, both the estimated A factors
705 and activation energies decreased with ring size, with the net result of these opposing factors being that
706 formation of 5-member cyclic ethers is generally estimated to be most favorable for atmospheric
707 conditions. Nevertheless, these cyclization reactions are estimated to be too slow for such radicals with
708 only alkyl or hydroperoxide groups to compete with reaction with O₂ at atmospheric temperatures.

709 However, as discussed in Section S1.4.4 in the SI, extrapolating the relation between the rate
710 constant and heats of reaction derived from the results of Curran et al. (1998) results in estimates that
711 cyclizations of carbonyl-containing radicals forming 5-member ring esters are ~10 times faster than O₂
712 addition for atmospheric conditions.



713 In addition, analogous cyclizations forming cyclic esters can also occur with carbon-centered radicals
714 with peroxy, nitrate, or peroxy nitrate substitution in the δ position, are also estimated to be faster than O₂
715 addition, i.e.,



716 where "X" = G or NO₂, and "XO." is G-O· or NO₃.

717 However, as also discussed in A1.4.3, Miyoshi (2011) reports results of theoretical calculations of
718 hydroperoxyalkyl rate constants that extrapolate to much lower rate constants at atmospheric
719 temperatures, suggesting that the cyclizations forming 5-member ring esters may not as fast as estimated
720 using the results of Curran et al. (1998). On the other hand, Vereecken and Nozière (2020) indicated that
721 some rate coefficients given by Miyoshi (2011) may not be accurate, and other theoretical work may yield
722 higher rate constants when extrapolated to room temperature (e.g., Ali et al., 2023). These reactions may
723 still be important for excited radicals. This will need to be re-examined in a subsequent version of
724 MechGen, but presently this reaction is still assumed to be fast even for thermalized radicals.

725 These reactions are assumed to be fast for mechanism generation purposes if (1) there is a
726 carbonyl group either at the radical center or next to the peroxy oxygen that becomes bonded to the
727 radical center, (2) the cyclic ester, carbonate, or anhydride has a 5-member ring, and (3) the radical is not
728 stabilized by resonance. O₂ addition is assumed to dominate if not all of these conditions are met. For
729 simplicity and to avoid the necessity of making highly uncertain quantitative rate constant estimates, we
730 assume that either decomposition or O₂ addition dominates, with the other being negligible. This

731 assumption can be revisited in the future if the ability to estimate these decomposition rate constants for
732 atmospheric conditions improves.

733 Although it is not likely in reactions of most VOCs and their oxidation products, in some cases
734 carbon-centered radicals may have more than one peroxy group in positions where decompositions are
735 estimated to be rapid. In these cases, it is assumed that the reactions occur at equal rates.

736 **3.3. Unimolecular Reactions of Chemically Activated Radicals**

737 Carbon-centered radicals formed by the addition of OH to double bonds are initially formed with
738 internal excitation, and in some cases might undergo unimolecular reactions that may not be important for
739 stabilized radicals. Reactions of the chemically activated radicals formed following additions to aromatic
740 rings are discussed separately in Section 3.4. Reactions that are assumed to be fast even for thermalized
741 radicals were discussed in the previous section, and they are also assumed to dominate for excited radicals
742 as well. Reactions that are assumed to dominate for radicals excited by additions of OH radicals to double
743 bonds, but not be important for stabilized radicals, are listed in Table 6 and are discussed in more detail in
744 Sections A1.4.3 and A1.4.4 in the SI.

745 Note that additions of NO₃ to double bonds also result in the formation of excited radicals, but the
746 excitation energy is estimated to be ~10 kcal/mole less than OH addition. Although this is uncertain, we
747 currently assume that stabilization is correspondingly more important for NO₃ adducts, and the
748 unimolecular reactions shown in Table 6 are assumed not to occur when mechanisms for reactions of NO₃
749 with alkenes are generated.

750 **3.4. Reactions of Radicals Formed by Additions to Aromatic Rings**

751 OH radicals can react by adding to various positions around the aromatic ring, with the branching
752 ratios derived using SARs for OH reactions with aromatics, as discussed in Section 2.1.2 and by Carter
753 (2021). These excited OH-aromatic adducts can either decompose back to the aromatic + OH or be
754 stabilized and then react with O₂. The effect of the back decomposition is taken into account in the SARs
755 for OH addition, which are based on total rate constants, so only reaction with O₂ is considered. O₂
756 reaction can occur either by abstracting the H-atom on the carbon where the OH added, forming HO₂ and
757 a phenol, or addition to the ring forming an OH-aromatic-O₂ adduct. As discussed below, one of the
758 possible reactions of this adduct is decomposition to form the same products as the abstraction reaction,
759 so for simplicity we assume that this is the major pathway for formation of phenolic products, so only the
760 addition reaction is considered.

761 The OH-aromatic adduct is an allylic carbon-centered radical with three possible radical centers
762 where O₂ can add. Although these adducts have some excitation, we assume that the branching ratios for
763 O₂ addition to the different positions are the same as estimated for stabilized allylic radicals, as discussed
764 in Section 3.1 and Section S1.4.2. This means that all the addition is assumed to occur at the *ortho*
765 positions since this forms the adducts with the most conjugation, that addition to an alkyl-substituted
766 *ortho* position is ~60% less likely than adding to the other *ortho* position, and addition to an -OH
767 substituted *ortho* position (as occurs in reactions with phenols) is ~7 times more likely, assuming that the
768 other *ortho* position is unsubstituted.

769 The aromatic-OH-O₂ adducts are expected to be formed with some excitation, and are assumed to
770 undergo various types of unimolecular reactions as discussed below. Although peroxy radical reactions
771 are discussed separately in Section 4, MechGen incorporates the rapid aromatic-OH-O₂ isomerization
772 reactions into the overall reactions generated for excited OH-aromatic adducts, so these reactions are
773 discussed in this section.

774 Table 6. Summary of types of unimolecular reactions that are assumed to be non-negligible for
 775 chemically activated carbon-centered radicals formed following additions of OH to
 776 double bonds.

Type [a]	Reactant [b]	Reaction	Ratio [c]
<i>Exo</i> ring opening (A1.4.3)	$-G_1[\cdot]-*G_2-G_3-G_4^*- \{excited\}$	$\rightarrow -G_1=G_2-G_3-G_4[\cdot]-$	
	$-G_1[\cdot]-*G_2-G_3-G_4-G_5^*- \{excited\}$	$\rightarrow -G_1=G_2-G_3-G_4-G_5[\cdot]-$ $+ M \rightarrow -G_1[\cdot]-*G_2-G_3-G_4-G_5^*- + M$	70% 30%
<i>Endo</i> ring opening (A1.4.3)	$-*G_1-G_2[\cdot]-G_3^*- \{excited\}$	$\rightarrow -G_1=G_2-G_3[\cdot]-$	
	$-*G_1-G_2[\cdot]-G_3-G_4^*- \{excited\}$	$\rightarrow -G_1=G_2-G_3-G_4[\cdot]-$ $+ M \rightarrow -*G_1-G_2[\cdot]-G_3-G_4^*- + M$	70% 30%
Peroxy Cyclizations (A1.4.4)	$-G_1[\cdot]-G_2-O-OX \{OHadd\}$	$-*G_1-G_2-O^* + [\cdot]OX$	
	$-G_1[\cdot]-G_2-ONO_2 \{OHadd\}$	$-*G_1-G_2-O^* + NO_2$	
	$-G_1(O'H)-G_2[\cdot]-CO-O-OH \{OHadd\}$	$\rightarrow G_1(O'H)-*G_2-CO-O^* + OH$ $\rightarrow *O'-G_1-G_2^*-CO-OH + OH$ $+ M \rightarrow G_1(O'H)-G_2[\cdot]-CO-O-OH + M$	14% 5% 82%
	$-G_1(O'H)-G_2[\cdot]-CO-O-ONO_2 \{OHadd\}$	$\rightarrow G_1(O'H)-*G_2-CO-O^* + NO_3$ $\rightarrow *O'-G_1-G_2^*-CO-OH + NO_3$ $+ M \rightarrow G_1(O'H)-G_2[\cdot]-CO-O-ONO_2 + M$	60% 20% 20%
	$-G_1(O'H)-G_2[\cdot]-CO-O-O-G_3- \{OHadd\}$	$\rightarrow G_1(O'H)-*G_2-CO-O^* + -G_3[O\cdot]$ $\rightarrow *O'-G_1-G_2^*-CO-OH + -G_3[O\cdot]$ $+ M \rightarrow G_1(O'H)-G_2[\cdot]-CO-O-O-G_3- + M$	45% 15% 40%

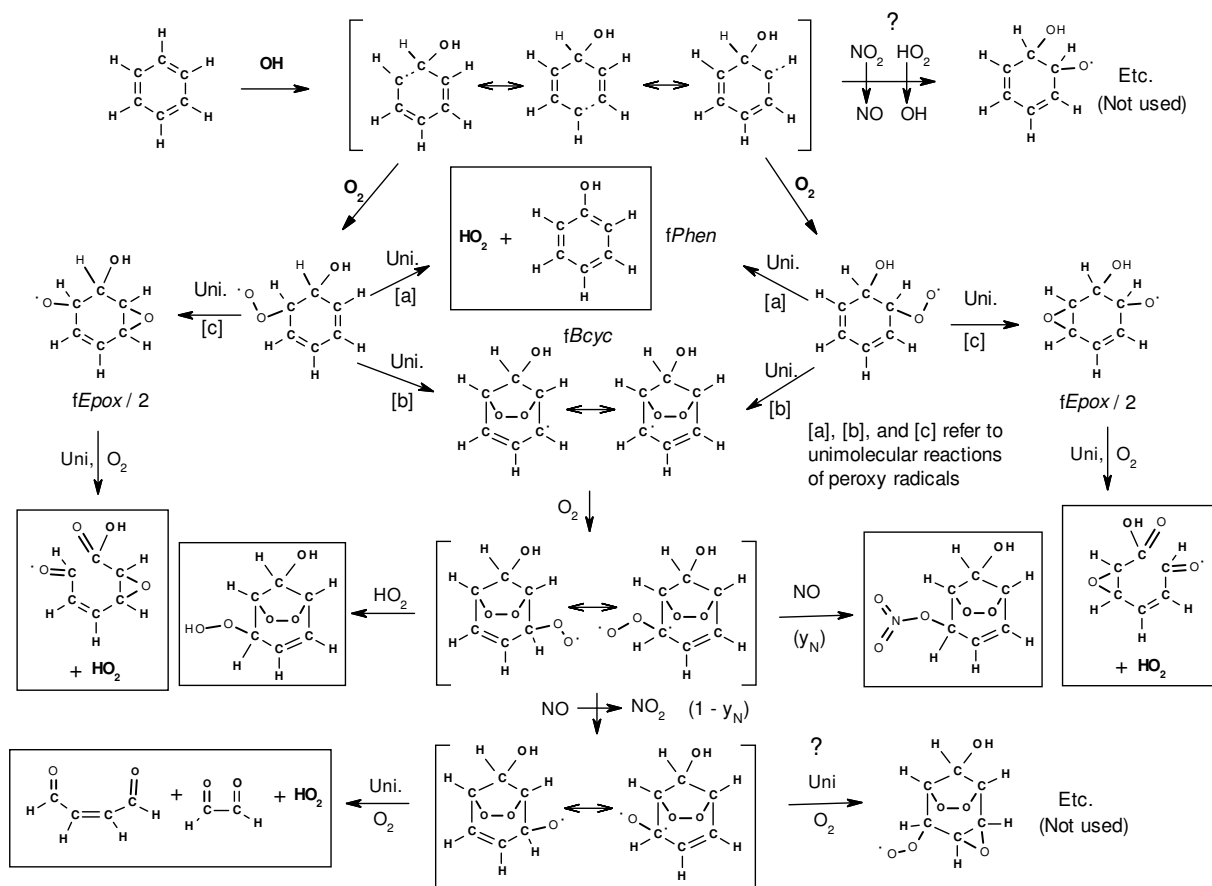
777 [a] The section in the SI where the reaction and the estimated branching ratios are discussed is shown in
 778 parentheses.

779 [b] "*" indicates ring closure, "{excited}" indicates excitation, {OHadd} indicates excitation by OH
 780 addition only, "+ M" indicates stabilization, and "X" is H or any group and $\cdot OX$ is OH, or G-O.

781 [c] Branching ratio assumed for 1 atmosphere pressure (the default) if stabilization is assumed to be non-
 782 negligible. Note that MechGen allows for changes in atmospheric pressure, in which case this ratio
 783 would change assuming that the stabilization rate constant is proportional to pressure.

784
 785

786 The subsequent reactions of the excited adducts are assumed to be as shown in Figure 1, which
 787 uses benzene as an example. This is based on the scheme recommended by Jenkin et al. (2018b), and
 788 incorporated in the MCM (Bloss et al., 2005; MCM 2023). It shows the initial formation of the OH-
 789 aromatic adducts, their subsequent reactions with O₂, and the fast unimolecular reactions assumed for the
 790 excited OH-aromatic- O₂ adducts. The codes for the branching ratios, which depend on the substituents
 791 about the aromatic ring are also shown. Three types of overall reactions are assumed. These include
 792 formation of a phenol + HO₂ via a 6-member ring transition state where the H transfers from the -OH to
 793 the peroxy group, followed by elimination of HO₂, with branching ratio designated *fPhen*; cyclization to
 794 form a bicyclic allylic-stabilized carbon-centered radical with branching ratio *fBcyc*; and the third
 795 involving a different type of cyclization where a peroxy O transfers to the opposite double bond, forming
 796 an epoxide and an alkoxy radical, with branching ratio *fEpoxy*. The alkoxy radical can form two different
 797 products depending on where the O₂ added and the substituents about the aromatic ring, which are



798

799 Figure 1. Initial reactions following the addition of OH to aromatic rings, using benzene as an
 800 example, showing the formation of the OH-aromatic adduct, followed by unimolecular
 801 reactions assumed for the OH-aromatic-O₂ adduct. Major products assumed to be formed
 802 are indicated, and additional pathways that may occur but are not currently used are also
 803 shown.

804

805 assumed to be equally likely. Therefore, up to four overall reactions can be generated following addition
 806 of OH to aromatic rings.

807 The *Phen* and *Bcyc* pathways in aromatic mechanisms (routes [a] and [b] on Figure 1) are well
 808 supported by existing product data (Calvert et al., 2002) and theoretical calculations (e.g., Calvert et al.,
 809 2002; Vereecken, 2018, 2019). However, the existence of the *Epoxy* pathway (routes [c]) is speculative
 810 and much more uncertain. It is included in the mechanism to account for carbon balance and is consistent
 811 with reactivities of aromatics observed in environmental chamber experiments (Carter 2010b, Carter and
 812 Heo, 2013). This pathway is also assumed in the aromatics mechanisms recommended by Jenkin et al.
 813 (2018b) and is incorporated in the MCM (Bloss et al., 2005; MCM 2023). However, theoretical
 814 calculations discussed by Vereecken (2018) indicate that the direct epoxide formation from aromatic-OH-
 815 O₂ adducts (route [c]) is unlikely to be important. Vereecken (2018) suggests alternative pathways that are
 816 shown on the top and bottom right side of Figure 1 as alternatives that appear to be more consistent with
 817 theoretical calculations and recent experimental data (Xu et al., 2020). These alternatives will be
 818 considered when MechGen is updated. The possible alkoxy cyclization reaction shown on the bottom
 819 right of Figure 1 is discussed in Section 5.6.

820 The mechanisms for the subsequent reactions of the carbon-centered and alkoxy radicals formed
 821 are estimated using the same procedures discussed elsewhere in this paper. The major stable products
 822 predicted to be formed are also indicated by the boxes in Figure 1. Note that as discussed in Section 2.2.5,
 823 the unsaturated 1,4-dicarbonyl aldehydes or aldehyde-ketones are predicted to be highly photoreactive, so
 824 the yields of these products significantly affect predictions of reactivity in aromatic hydrocarbon - NO_x
 825 environmental chamber experiments (e.g., Carter and Heo, 2013).

826 The branching ratios were derived based on reported yields of phenolic products and α-
 827 dicarbonyls from the reactions of OH with benzene and various alkylbenzenes, as well as simulations of
 828 environmental chamber experiments. Values of *fPhen* were derived based on phenolic yields, *fBcyc*
 829 values are based on 1,2 dicarbonyl yields and modeling environmental chamber experiments, and *fEpo*
 830 are obtained as 1 - *fPhen* - *fBcyc*. It was sufficient to assume that the branching ratios depend only on the
 831 presence of substituents in the *ipso* and *ortho* positions, with substituents in the *meta* or *para* positions
 832 assumed not to have an effect on these branching ratios. The branching ratios used for the various
 833 substituent cases are summarized in Table 7. Note that *fPhen* is necessarily zero if there is an *ipso*
 834 substituent, and the *fBcyc* / *fEpo* ratios for such compounds are estimated to be the same as those for
 835 compounds without *ipso* substituents, but with the magnitudes increased so they sum up to 1. The ratios
 836 in Table 7 are used regardless of substituents, though estimates for compounds with substituents other
 837 than alkyl or -OH, and therefore generated mechanisms for such compounds, should be considered to be
 838 unreliable and probably should not be used for mechanism development.

839
 840 Table 7. Branching ratios derived for the reactions of aromatic + OH and aromatic + OH + O₂
 841 adducts, based on the presence or absence of substituents in the *ipso* and *ortho* positions.

Branching Ratio	No <i>ipso</i> substituents			1 <i>ipso</i> substituent [a]		
	No <i>ortho</i>	1 <i>ortho</i>	2 <i>ortho</i>	No <i>ortho</i>	1 <i>ortho</i>	2 <i>ortho</i>
<i>fPhen</i>	57%	16%	5%	-	-	-
<i>fBcyc</i>	31%	57%	83%	73%	68%	88%
<i>fEpo</i>	12%	27%	12%	27%	32%	12%

842 [a] Derived from *fBcyc* and *fEpo* assigned for adducts with no *ipso* substituents.
 843

844 The above discussion is applicable only to reactions following additions of OH radicals to
 845 aromatic rings, not to additions by NO₃ radicals. As discussed by Carter (2021), the rate constants for
 846 these additions are estimated to be slow, based on the relatively low total rate constants reactions for
 847 aromatic + NO₃ reactions. MechGen currently assumes that all aromatic-NO₃ adducts rapidly decompose
 848 back to reactants, resulting in no net reaction. However, this cannot strictly speaking be true, since the
 849 experimental rate constants give SARs that predict nonzero, though low, rate constants for these reactions
 850 (Carter, 2021; see also Section 2.1.2). However, the predicted rate constants are too low to be important in
 851 atmospheric systems, so this oversimplification has no practical consequences for atmospheric
 852 mechanisms.

853 The reactions of radicals formed by the additions of OH radicals to the aromatic rings in phenols
 854 are expected to be analogous to those discussed above, but the presence of OH substitution affects
 855 branching ratios for the addition to the allylic OH + phenol adduct as discussed in Section 3.1, and some
 856 of the OH + phenolic + O₂ adducts form cyclohexadienones, which undergo are assumed to undergo rapid
 857 unimolecular reactions to form catechols as discussed in Section 2.3.2. The mechanisms assumed for OH
 858 + phenolic adducts are discussed in Section S1.8 of the SI.

859

4. Reactions of Peroxy Radicals

860 Peroxy radicals are formed primarily by the reactions of alkyl radicals with O₂ and are critical
 861 intermediates in atmospheric oxidation mechanisms. Peroxy radicals can be classified alkyl peroxy,
 862 -G[OO.] or RO₂, or acyl peroxy, -G-CO[OO.] or RCO₃ (where "G" is any group that does not have an α
 863 carbonyl substituent). In the following discussion, we will use the term "acyl" peroxy radical to refer to
 864 any peroxy radical bonded to a carbonyl group and "alkyl" peroxy radicals otherwise, including peroxy
 865 radicals with non-alkyl groups or substituents.

866 Peroxy radicals can react with NO, NO₂, NO₃, HO₂, and other peroxy radicals, and some can also
 867 undergo unimolecular reactions at significant rates under atmospheric conditions. The rate constants
 868 assigned when generating bimolecular rate constants are summarized in Table 8, and their derivations are
 869 summarized in the following subsections where the mechanisms for the different types of reactions are
 870 discussed. More details concerning the derivations of the bimolecular rate are given in Section S1.5.1 of
 871 the SI. The unimolecular reactions are discussed separately below.

872

873 Table 8. Summary of bimolecular rate constants used when generating reactions of peroxy
 874 radicals under atmospheric conditions.

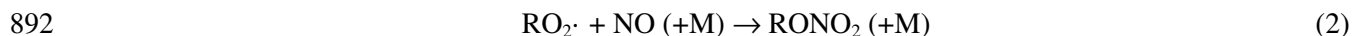
Reactant	Rate constant around 300K (cm ³ molec ⁻¹ s ⁻¹) [a]						RCO ₃
	Non OH-Substituted RO ₂			OH-Substituted RO ₂			
	Primary	Secondary	Tertiary	Primary	Secondary	Tertiary	
NO			9.13x10 ⁻¹²				2.10x10 ⁻¹¹
NO ₂			8.8x10 ⁻¹²				7.7x10 ⁻¹²
NO ₃			2.3x10 ⁻¹²				4.0x10 ⁻¹²
HO ₂	10 ⁻¹² x min [20.3, 2.48 + (2.48 x nC)]			10 ⁻¹² x min [21.1, 9.37 + (1.68 x nC)]			2.1x10 ⁻¹¹
Generic RO ₂	2.9x10 ⁻¹⁴	1.6x10 ⁻¹⁴	3.7x10 ⁻¹⁶	9.1x10 ⁻¹⁴	1.7x10 ⁻¹³	1.3x10 ⁻¹⁴	1.6x10 ⁻¹¹
Generic RCO ₃			1.6x10 ⁻¹¹				1.4x10 ⁻¹¹

875 [a] See Table S-45 in the SI for the sources of the rate constants used. Note that the same rate constant is
 876 used for reactions of alkyl peroxy radicals with NO_x species and with acyl peroxy radicals, while the
 877 rate constants depend on the carbon number for reactions with HO₂, and on the structure of the radical
 878 for reactions with other alkyl peroxy radicals (See Sections 4.4 and 4.5, respectively).
 879

880 4.1. Reaction with NO

881 The major atmospheric loss process of most peroxy radicals when formed in the presence of NO_x
 882 is reaction with NO. Available data on the rate constants for peroxy + NO reactions are given in Table S-
 883 45 in the SI, and the rate constants measured for ethyl peroxy and propionyl peroxy are estimated to apply
 884 to all peroxy radicals for mechanism generation. There is some variability in the measured rate constants
 885 for the alkyl peroxy radicals, but the data are insufficient to derive estimates for dependences of the rate
 886 constant on the structure or size of the radicals. As indicated in Table 8, although different rate constants
 887 are used for reactions of acyl peroxy vs peroxy radicals with NO, because of insufficient information we
 888 assume that other structural effects have no effects on the rate constant.

889 The reactions of peroxy radicals with NO can occur via two possible routes, either forming NO₂
 890 and the corresponding alkoxy radical, or forming a stable organic nitrate in the presence of a bath gas.



893 Both reactions are known to be important in the case of alkyl peroxy radicals, and the nitrate yield,
 894 $y_N = k_2 / (k_1 + k_2)$, is an important parameter affecting model predictions in atmospheric systems since the
 895 conversion of NO to NO₂ in Reaction (1) contributes to O₃ formation and the formation of alkoxy radicals
 896 results in radical propagation, while nitrate formation in Reaction (2) causes radical termination and is
 897 also a NO_x sink, at least for the intermediate term.

898 Available data and estimates concerning nitrate yields from various peroxy radicals are discussed
 899 in Section S1.5.2 of the SI. There are extensive measurements of nitrate yields from various unsubstituted
 900 peroxy radicals formed from the reactions of alkanes, including data at various temperatures and
 901 pressures, allowing a parameterization to be derived that fits these data as a function of temperature,
 902 pressure, and carbon number. The parameterization used is similar to that employed previously (e.g.,
 903 Atkinson et al., 1983), updated based on the mechanism considerations discussed by Zhang et al. (2004)
 904 and re-optimized to fit available data as discussed in the SI. The predicted nitrate yields at around 298K
 905 and 1 atmosphere pressure are given in Table 9 and are plotted against carbon number in Figure 2, where
 906 they are compared to available data as discussed below.

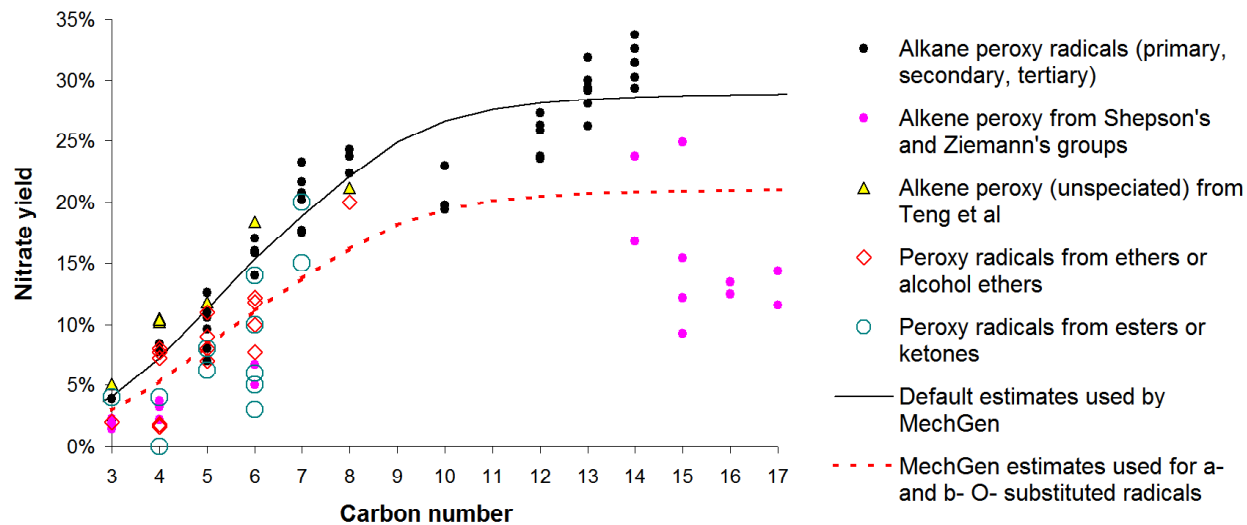
907
 908 Table 9. Estimated nitrate yields as a function of carbon number calculated for ~298K and 1
 909 atmosphere pressure.

Carbon Number	2	3	4	5	6	7	8	9	10	11	12	13	14+
Default nitrate yield	2%	4%	7%	11%	15%	19%	22%	25%	27%	28%	28%	28%	29%
Nitrate yield if α- or β -O substituted	2%	3%	5%	8%	11%	14%	16%	18%	19%	20%	20%	21%	21%

910
 911 Figure 2 shows that the parameterization derived to fit measured nitrate yields for unsubstituted
 912 peroxy radicals from alkanes (black line) fit those data reasonably well, but its applicability for
 913 substituted radicals from other VOCs is much more uncertain. The limited available measurements of
 914 nitrate yields from β OH-substituted radicals are inconsistent. Data obtained from Shepson's (Muthuramu
 915 et al., 1993; Shepson et al., 1985; O'Brien et al., 1998) and Ziemann's (Matsunaga and Ziemann, 2009,
 916 2010) groups gave yields that are about a factor of two lower than from unsubstituted radicals, while more
 917 recent data from Teng et al. (2015) gave yields that agree with those from the unsubstituted radicals, at
 918 least at carbon numbers <10. The estimated nitrate yields from the OH-substituted radicals significantly
 919 affect results of model simulations of alkene - NO_x chamber experiments (Carter, 2000, 2010b), and we
 920 found that some data are better fit using the higher yields estimated for radicals from alkanes, while some
 921 are better fit using the yields that are a factor of ~2 lower (see Section S1.5.2 in the SI).

922 Figure 2 also shows scatter in the available data concerning nitrate yields from peroxy radicals
 923 formed from ethers, esters, and ketones, though they tend to be lower than observations for unsubstituted
 924 peroxy radicals (Jenkin et al., 2019 and references therein). As discussed in the SI, these yields are subject
 925 to greater uncertainties because they are derived either from unspiciated total nitrate measurements in
 926 complex chemical systems, or by fits to environmental chamber data, where uncertainties are even
 927 greater.

928



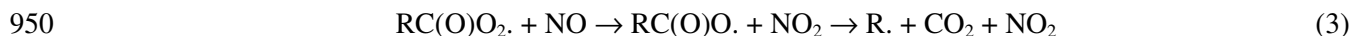
929
930

931 Figure 2. Measured and estimated nitrate yields from reactions of various types of peroxy radicals
932 with NO against numbers of carbon atoms in the radicals.

933 Until more information is available to resolve these discrepancies and provide a better basis for
934 making nitrate yield estimates for non-alkyl-substituted peroxy radicals, MechGen uses a
935 parameterization (red dotted line in Figure 2) that predicts yields halfway between those derived for
936 radicals from alkanes and the lower yields for the β OH-substituted radicals, when estimating nitrate
937 yields for peroxy radicals bonded to oxygen groups in the α or β positions, e.g., -OH, ether, ester,
938 peroxides, or nitrates. This is similar to the approach used in the nitrate yield estimation method
939 developed by Jenkin et al. (2019) for alcohols, ethers, and esters. Based on the limited data, Jenkin et al.
940 (2019) also assumed carbonyl groups in the α or β positions caused even lower nitrate yields, but the
941 current version of MechGen does not yet incorporate corrections for these cases.

942 The nitrate yields predicted using this parameterization, derived as discussed in Section S1.5.2 of
943 the SI, are included in Table 9. Note that we assume that substituents farther away than the β position are
944 too far away from the reaction site to affect the nitrate yield. However, the estimates for nitrate yields
945 from substituted peroxy radicals are uncertain by at least a factor of two, and are more likely to be biased
946 high rather than low.

947 There is no evidence for nitrate formation in the case of acyl peroxy radicals, so the formation of
948 NO_2 and the corresponding acyl oxy radical is assumed to be the only pathway of significance (e.g., see
949 IUPAC, 2023).



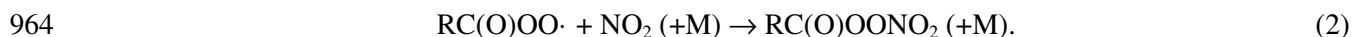
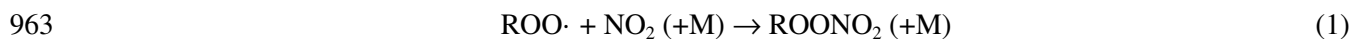
951 As discussed above, the acyl oxy radicals are expected to rapidly decompose to form CO_2 and the
952 corresponding alkyl radical, making the overall process for acyl + NO reactions as indicated above.

953 4.2. Reaction with NO_2

954 Alkyl and acyl peroxy radicals also react with NO_2 under atmospheric conditions, and the rate
955 constants assigned to them are included in Table 8. The assigned rate constants are based on measured
956 high pressure rate constants for ethyl peroxy and ethyl acyl peroxy radicals, since we could not find data
957 for larger peroxy radicals (see Table S-45 in the SI). Although the reactions of ethyl and ethyl acyl peroxy

958 radicals are not in the high pressure limit under atmospheric conditions, most of the radicals are larger and
959 their rate constants should be closer or at the high pressure limit, so the high pressure rate constant is used
960 for general estimates.

961 The reactions for both alkyl and acyl peroxy radicals with NO₂ are assumed to result entirely in
962 the formation of the corresponding alkyl or acyl peroxy nitrate.



965 Reactions forming NO₃ + alkoxy or acyloxy radicals are estimated to be endothermic, and are assumed
966 not to be important. The peroxy nitrate compounds formed are thermally unstable at atmospheric
967 temperatures, decomposing back to NO₂ and the peroxy radical, as discussed in Section 2.3. It is assumed
968 that the presence of non-alkyl substituents on the "R" does not affect either the formation or
969 decomposition rate constants, but this is uncertain.

970 **4.3. Reaction with NO₃**

971 Reactions with NO₃ can be a potentially important sink for peroxy radicals at nighttime and
972 potentially other conditions when NO concentrations are low. The rate constants used for mechanism
973 generation (Table 8) are those measured for ethyl peroxy and acyl peroxy radicals because we could not
974 find data for larger radicals (see Table S-45).

975 Peroxy radicals react with NO₃ to form the corresponding alkoxy radical, NO₂ and O₂, with no
976 evidence of other reaction routes or products being reported (e.g., IUPAC, 2023).



979 Note that the same organic products are formed in this reaction as in the reaction of peroxy radicals with
980 NO, other than the lack of the nitrate formation route in the NO₃ reaction. Although nitrate formation in
981 this reaction is energetically possible, there is no evidence that it occurs under atmospheric conditions.
982 The presence of non-alkyl substituents on "R" is assumed not to affect the rate constants or mechanisms.

983 **4.4. Reaction with HO₂**

984 Reactions with HO₂ can be the major sink for peroxy radicals under conditions when NO_x is low,
985 being generally much more important than reactions with other peroxy radicals. Information concerning
986 measurements and estimates for mechanisms and rate constants for the reactions of peroxy radicals with
987 HO₂ are discussed in Section S1.5.3 of the SI. In the case of alkyl peroxy radicals, the data suggest that
988 the rate constants depend on the size of the radical and the presence of OH substitution on the radical,
989 with the following equations being derived for estimation purposes:

990
$$k(\text{RO}_2 + \text{HO}_2) = 10^{-12} \times \min [20.3, 2.48 + (2.48 \times n\text{C})] \text{ cm}^3 \text{ molec}^{-1} \text{ s}^{-1}$$

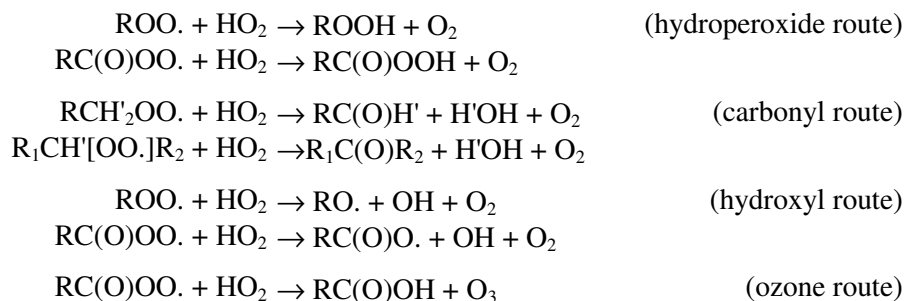
991
$$k(\text{HO-RO}_2 + \text{HO}_2) = 10^{-12} \times \min [21.1, 9.37 + (1.68 \times n\text{C})] \text{ cm}^3 \text{ molec}^{-1} \text{ s}^{-1}$$

992 where "HO-RO₂" refers to alkyl peroxy with OH substituents anywhere on the radical, and "RO₂" refers
993 to those without. Note that there is some inconsistency in the measured rate constants for OH substituted
994 radicals (see SI), so there is some uncertainty in these estimates.

995 The only data we found for reactions of acyl peroxy radicals with HO₂ concerned acetyl peroxy
996 radicals, so the rate constant measured for that reaction (on Table S-45) is used for estimation purposes.

997 This is uncertain, especially in view of the variability of the measured rate constants for the alkyl peroxy
 998 radicals.

999 Available information concerning the products of the reactions of various types of peroxy radicals
 1000 with HO₂ have been evaluated by IUPAC (2023), and their recommendations of branching ratios for
 1001 methyl and acetyl peroxy radicals, and for methyl peroxy substituted with -CH₃, -OH, -OCH₃, and
 1002 -C(O)CH₃ are given in S-12 of the SI. Their recommendations include the following four routes:



1003 with the "ozone" route only considered for acyl peroxy radicals. The "carbonyl" route occurs via an H
 1004 shift from an α hydrogen to the HO₂ via a 6-member ring transition state, and is not possible for tertiary
 1005 alkyl or acyl peroxy radicals. Although there are data only for the C₁ or C₂ peroxy radicals, they are
 1006 assumed to be applicable to all alkyl peroxy radicals with the same types of α substituents for mechanism
 1007 generation purposes, and are the basis for the branching ratios that are summarized in Table 10. However,
 1008 the applications of these recommendations to the many types of larger peroxy radicals predicted in
 1009 generated mechanisms is highly uncertain.

1010
 1011 Table 10. Branching ratios assigned for the four possible routes for the reactions of HO₂ with
 1012 various types of peroxy radicals.

Route	CH ₃ O ₂	RO ₂ [a]	HO-RO ₂ [b,c]		RO-RO ₂ [b,c]		RCO-RO ₂ [b]	RCO ₃
			P, S	T	P, S	T		
Hydroperoxide	0.9	1.0	0.5	0.8	0.6	1.0	0.85	0.37
Carbonyl	0.1	-	0.3	-	0.4	-	-	-
Hydroxyl	-	-	0.2	0.2	-	-	0.15	0.5
Ozone	-	-	-	-	-	-	-	0.13

1013 [a] "RO₂" refers to peroxy radicals with no -OH, -OR, or carbonyl substituents in the α position. Note
 1014 that they can have these and other non-alkane substituents or groups elsewhere on the molecule.

1015 [b] "The "HO-", "RO-" or "RCO-" refer to the substituent in the α position only. "RCO-" refers to either
 1016 -CHO or -CO-, though data are only available for -CO- substitution.

1017 [c] "P", "S", and "T" refer to primary, secondary, and tertiary radicals, respectively. Since the carbonyl
 1018 route is not possible for tertiary radicals, the branching ratio estimate for this route for primary or
 1019 secondary radicals is added to the estimate for the hydroperoxy route for such radicals.
 1020

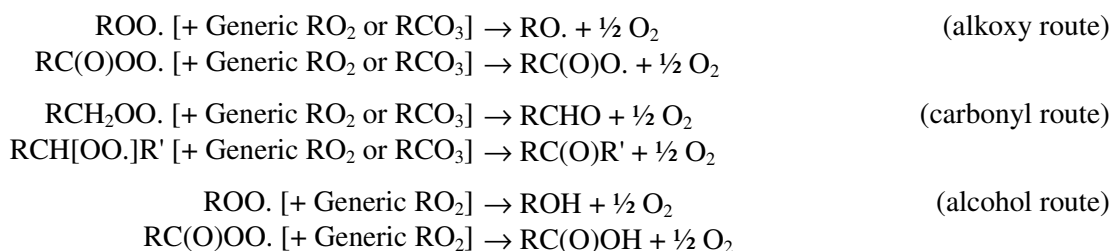
1021 4.5. Reactions with Other Peroxy Radicals

1022 Organic peroxy radicals can also be consumed under atmospheric conditions by reactions with
 1023 other organic peroxy radicals. Although this is generally a less important loss process under atmospheric
 1024 conditions than reactions with NO_x or HO₂, such reactions can be non-negligible under relatively high

1025 VOC and low NO_x atmospheric conditions, and also in some laboratory experiments. Available
1026 information concerning measured rate constants for these reactions is given in Table S-45 of the SI. The
1027 rate constants vary by orders of magnitude, with the most important factor being whether the alkyl peroxy
1028 radicals are primary, secondary, or tertiary. However, the presence of α-OH substitution also appears to
1029 affect the rate constant, and this is also taken into account. Table S-45 in the SI includes a summary of the
1030 averages or estimated rate constants for various types of radicals that can be derived from the
1031 experimental data, and the rate constants used by MechGen when generating mechanisms for peroxy
1032 radicals are included in Table 8.

1033 Because it is not practical to represent all of the peroxy + peroxy combinations in large
1034 mechanisms, MechGen does not assign rate constants or generate mechanisms for all combinations of the
1035 different types of radicals, but instead represents the radicals as reacting with a generic alkyl (RO₂) and a
1036 generic acyl (RCO₃) peroxy radical. The rate constants and also mechanism assignments for the generic
1037 RO₂ are based on those estimated for secondary peroxy radicals because they are intermediate in
1038 reactivity and are also expected to be most frequently formed. The generic acyl peroxy radical is based on
1039 acetyl or propionyl peroxy radicals, the only type for which data are available. Although this
1040 representation can be inaccurate because it lumps reactions whose rate constants vary by orders of
1041 magnitude, it is considered to be relatively unbiased, and is better than the alternative of ignoring these
1042 reactions altogether. Fortunately this is a relatively unimportant loss process for peroxy radicals under
1043 most conditions, so this is not considered to be a significant problem compared to other uncertainties.

1044 Available information concerning the mechanisms of peroxy + peroxy reactions is discussed in
1045 Section S1.5.4 of the SI, and recommended and assigned branching are given in Table S-14. Three types
1046 of generic reactions are considered.



1047 The products formed from the generic radicals are not shown because they are not generated because they
1048 are represented separately by the reactions of the individual radicals. (This is an approximation because it
1049 undercounts the contributions of self-reactions in the kinetic differential equations.) Note that the
1050 "carbonyl" and "alcohol" routes are two instances of the disproportionation reaction, where an α H
1051 migrates from the carbon next to the peroxide group on one of the radicals to a peroxy oxygen on the
1052 other radical. Formation of carbonyl products is not possible for acyl radicals and tertiary alkyl radicals,
1053 and the alcohol route is not possible for generic acyl radicals because they lack the necessary α hydrogen.

1054 The branching ratios used when generating reactions of the various types of peroxy radicals are
1055 summarized in Table 11. These are based roughly on the branching ratios for individual reactions given in
1056 Table S-14 in the SI, and extrapolating them to other types of radicals where necessary as discussed in
1057 Section S1.5.4. These assignments are uncertain when applied to larger radicals and radicals with non-
1058 alkyl substituents or groups, but the effects of this uncertainty are probably less than the uncertainty and
1059 necessary inaccuracies when estimating the rate constants.

1060

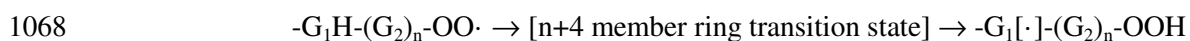
1061 Table 11. Branching ratios used when generating mechanisms of peroxy + peroxy reactions.

Peroxy radical type	Reaction with Generic RO ₂			Reaction with Generic RCO ₃	
	Alkoxy	Carbonyl	Alcohol	Alkoxy	Alcohol
Primary or secondary RO ₂	0.5	0.25	0.25	0.8	0.2
Tertiary RO ₂	0.5	-	0.5	1.0	-
Any RCO ₃	0.8	-	0.2	1.0	-

1062
1063

1064 4.6. H-shift Isomerizations

1065 Organic peroxy radicals can undergo H-shift isomerizations where the radical center abstracts a
1066 hydrogen from elsewhere in the molecule, via a cyclic transition state, forming a hydroperoxide and a
1067 carbon-centered radical.



1069 This reaction has been known to occur at higher temperatures in combustion systems, where the α -
1070 hydroperoxy carbon-centered radicals formed are designated "QOOH", though they had not been
1071 expected to be important at atmospheric temperatures until recently. Recent experimental data indicate
1072 that these reactions occur at measurable rates and may be non-negligible under atmospheric conditions for
1073 certain radicals (e.g., Crouse et al., 2012; Teng et al., 2015; Praske et al., 2018, 2019; Nozière and
1074 Vereecken, 2019). This has been supported by quantum theoretical calculations of rate constants for a
1075 wide variety of such reactions (e.g., Davis and Francisco, 2010; Møller et al., 2019; Vereecken and
1076 Nozière, 2020 and references therein), whose results are reasonably consistent with the available
1077 experimental data and indicate that in some cases these isomerizations would be the dominant fate of such
1078 radicals.

1079 Vereecken and Nozière (2020) reviewed the available theoretical and experimental rate constant
1080 data for peroxy H-shift isomerizations for a wide variety of peroxy radicals, and provided recommended
1081 SARs for predicting their rate constants for automated mechanism generation applications. These consist
1082 of lookup tables giving the rate constant for various pairs of peroxy radical types and H-groups (the
1083 groups with the H that is transferred) involved in the H-shift, giving the rate constants as a function of
1084 transition state ring size, with correction factors for a few types of substituents around the H-group. The
1085 effects of -OH, -OOH, -OR, carbonyl groups, and double bonds at or near the H-group were incorporated
1086 into the lookup tables; separate rate constant recommendations were given for reactions of -CH₂OO·, -
1087 >CH(OO·)- and -C(OO·)< radicals, and rate constants were given for ring sizes up to 8 or 9. The
1088 recommended correction factors covered the effects of α -ONO₂, β -OH, β -endo oxo and β -exo oxo
1089 substituents on the H-group. These recommendations were adopted as the starting point for use in this
1090 work. However, the recommended lookup tables did not cover all the types of peroxy radicals that might
1091 be formed in atmospheric photooxidation systems, including reactions of acyl peroxy radicals, of radicals
1092 with both β double bonds and α -OH groups, abstractions from an HCO group with a β double bond, H-
1093 shifts with transition state rings larger than 8 or 9, and several other cases. Therefore, it was necessary in
1094 this work to extrapolate or extend the work of Vereecken and Nozière (2020) to allow predictions for
1095 these other types of radicals.

1096 The adaptation of the peroxy H-shift SARs for use in this work is described in detail in Section
1097 S1.5.5 of the SI. In order to provide a basis for estimating rate constants, the various factors affecting the

1098 rate constants are split up into components that are somewhat more straightforward to estimate when
1099 expanding the lookup tables to cover the other cases. The expression used for estimation purposes was:

$$1100 \quad k(T) \approx nH \times A(n,ts) \times e^{-\{Ea^0(H,R) + Ea^{strain}(H,R,n) + Ea^{corr}(subs,n)\} / RT} \quad (I)$$

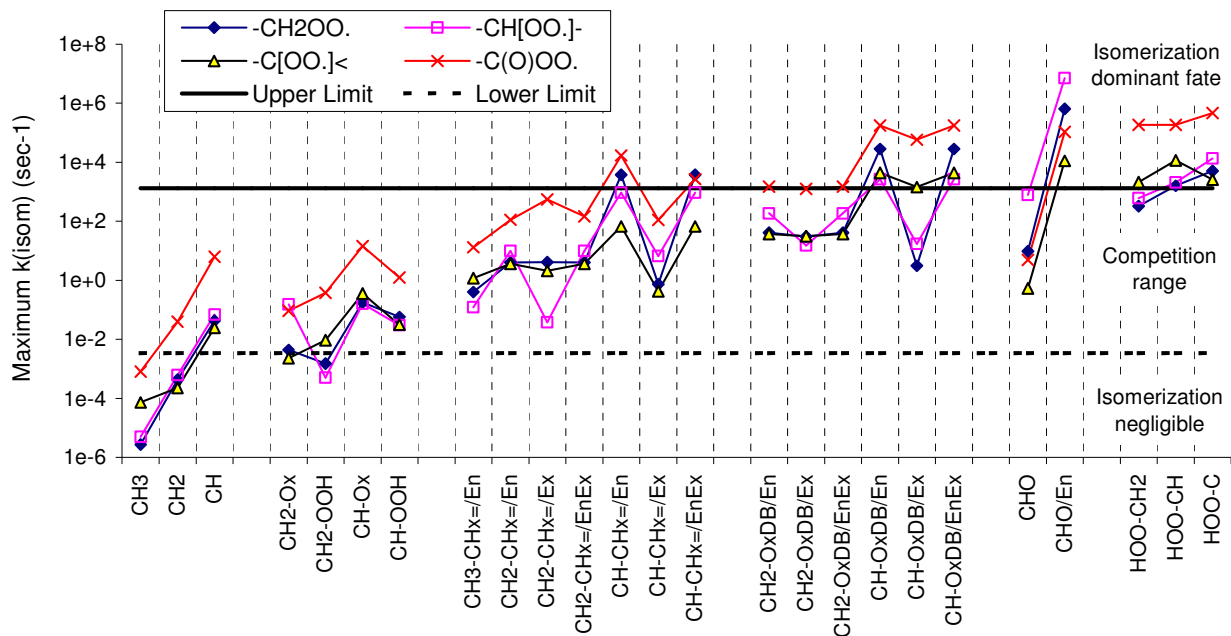
1101 where nH is the number of equivalent abstractable hydrogens; $A(n,ts)$ is an Arrhenius A factor assumed to
1102 be dependent only on the size of the transition state ring, n , and the type of transition state, ts ; $Ea^0(H,R)$ is
1103 the activation energy in cases where there is no strain or substituent corrections, and is assumed to be
1104 dependent only on the type of H-group, H , and the type of radical, R ; $Ea^{strain}(H,R,n)$ is the ring strain that
1105 depends on the ring size as well as the types of groups involved; and $Ea^{corr}(subs,n)$ are corrections for
1106 substituents on the H-group, some of which depend on ring size. The temperature-dependence
1107 recommendations given by Vereecken and Nozière (2020) were used to derive approximate A factors
1108 which were then used, in conjunction with the recommended rate constants in the lookup tables, to create
1109 lookup tables for Ea^0 and for Ea^{strain} for the various ring sizes. These were then used to estimate Ea^0 and
1110 Ea^{strain} for cases not covered by the original SARs. The recommended substituent correction factors were
1111 used to derive Ea^{corr} to use in Equation (I), where applicable. No new types of substituent correction
1112 factors were added in this work.

1113 The range of magnitudes of estimated rate constants for the peroxy H-shift isomerization for the
1114 various types of radicals is presented in Figure 3, which shows the rate constants for the transition state
1115 ring size (generally 6-8) that gives the highest rate constant. These maximum rate constants for the
1116 different types of reaction range in magnitude from $\sim 3 \times 10^6$ to $\sim 7 \times 10^6 \text{ sec}^{-1}$, and the lookup tables cover
1117 25 types of H-groups and transition states, as indicated on the x-axis of the figure. The figure also has
1118 lines giving the upper and lower limits of these rate constants used by MechGen when generating peroxy
1119 radical reactions -- reactions faster than the upper limit are assumed to be the only fate of the peroxy
1120 radical, while those that are below the lower limit are treated as negligible and not generated. For those
1121 with rate constants between these limits there is a competition between unimolecular and bimolecular
1122 reactions of these radicals, so the estimated rate constant would have an impact in predicted mechanisms.

1123 The H-shift reaction from unsubstituted $-CH_3$ and $-CH_2-$ groups in alkyl peroxy radicals was
1124 generally negligible, while abstractions from hydroperoxy groups and aldehyde groups and abstractions
1125 of allylic hydrogens by acyl peroxy groups were generally fast. Most of the other types were in the
1126 intermediate range where there is a competition between unimolecular and bimolecular reactions of these
1127 peroxy radicals. This indicates that this is an important process for many radicals that must be considered
1128 when implementing mechanisms into models.

1129 The rate constant estimates that were derived directly from the SARs of Vereecken and Nozière
1130 (2020) are based on results of quantum theory calculations and may be good to within a factor of ~ 10 , and
1131 can predict the experimental 298 K rate constants to within a factor of 3, as shown in Figure S-12 in the
1132 SI. Those that are based on our extrapolations or estimates are much more uncertain, though many of
1133 these are estimated to be high enough that the actual fate of the peroxy radical may not be sensitive to this
1134 estimate. However, experimental and theoretical data are needed to verify or improve our estimated
1135 extensions of the Vereecken and Nozière (2020) SARs as discussed in Section 1.5.5 of the SI, as well as
1136 more experimental data to validate the extensive set of theoretical calculations that form the primary basis
1137 for the SARs that are currently employed.

1138 No attempt was made to make separate estimates of rate constants for H-shift isomerizations of
1139 cyclic peroxy radicals by either Vereecken and Nozière (2020) or in this work. However, Vereecken et al.
1140 (2021) does include calculations for cyclic peroxy radicals that could be used when MechGen is updated,
1141 but gives insufficient data to derive SARs at this time. For the purpose of making general estimates, we
1142 assume that the isomerization reaction does not occur if three or more of the atoms in the transition state
1143 ring are in another ring, since they would usually (but not necessarily always) result in a highly strained



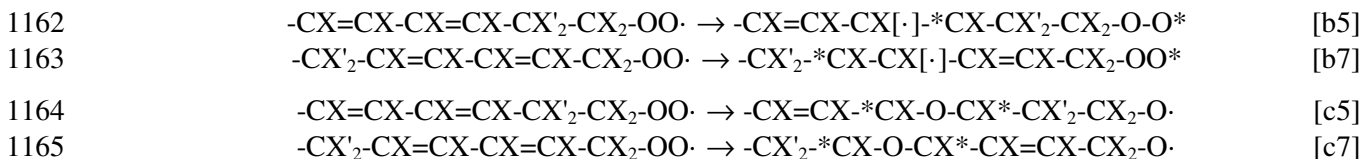
Nomenclature: H-group with substituents (if applicable), where -Ox = α -OH or -OR substituents; -CH_x=/En: β double bond by the H-group, with at least one double bond group in the transition state ring; -CH_x/Ex: β double bond not in the transition state; -OxDB = both α -Ox and a double bond substituent; /EnEx = two β double bonds by the H-group.

1144
 1145 Figure 3. Estimated peroxy H-shift isomerization rate constants for the most favorable ring size for
 1146 various H- and R-group pairs without β -substitution corrections. Also shown are upper
 1147 and lower rate constant limits used during mechanism generation.

1148
 1149 structure. Although these are not unreasonable assumptions, some isomerizations will be overestimated
 1150 and others overlooked. The practical effects of these uncertainties have not been assessed, but such an
 1151 assessment may give an indication for priorities for future theoretical or experimental studies of
 1152 unimolecular reactions of peroxy radicals formed from terpenes and other cycloalkenes.

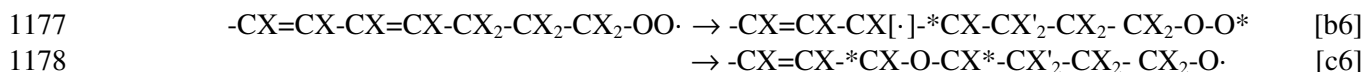
1153 4.7. Ring Closure Reactions of Unsaturated Peroxy Radicals.

1154 The current estimated mechanisms for aromatics, discussed above in Section 3.4, assume two
 1155 additional types of unimolecular reactions of OH-aromatic-O₂ adducts to account for observed aromatic
 1156 product formation and reactivities, both involving ring formation. These are shown as reactions labeled
 1157 "[b]" and "[c]" in Figure 1. These reactions are accounted for in the process of generating reactions of
 1158 OH-aromatic adducts as discussed in Section 3.4, and thus their rate constants do not need to be estimated
 1159 to generate mechanisms for aromatics. However, these reactions may also be non-negligible in non-
 1160 aromatic systems, since neither the reactants, products, nor the transition states have aromatic rings. The
 1161 reactions shown in Figure 1 can be generalized as follows:



1166 where X is H or any group, and * indicates ring closure. The reactions labeled [b] involve the peroxy
 1167 group adding to a double bond forming a cyclic ether and an allylic carbon-centered radical, while those
 1168 labeled "[c]" have a similar cyclic transition state but the O-O bond breaks, with one O adding to the
 1169 double bond forming an epoxide and the other becoming an alkoxy radical center. Note that the reactions
 1170 labeled [b5] and [c5] involve 5-member ring products or transition states, while those labeled [b7] and
 1171 [c7] involve a 7-member ring. Since these reactions apparently dominate over bimolecular reactions when
 1172 formation or intermediacy of both 5 and 7 member rings are involved, one would reasonably expect them
 1173 to be even faster in acyclic systems where only one new ring is involved.

1174 Since 6- or 7-member rings tend to have less ring strain than 5 member rings, (see discussion of
 1175 peroxy H-shift reactions in Section 1.5.5 of the SI) one would expect the analogue involving formation or
 1176 intermediacy of 6 member rings also to be fast, e.g.,



1179 Therefore, to be consistent with the aromatics mechanisms, the above six types of reactions are generated
 1180 for peroxy radicals with conjugated double bonds in the α -, β -, or γ - positions, relative to the carbon
 1181 bonded to the peroxy group, and are assumed to dominate over bimolecular reactions and H-shift
 1182 isomerizations. Because of this, it is not necessary to estimate their total rate constants, only the branching
 1183 ratios for the [b] and [c] reaction routes, or the "fEpoxfBcyc" ratio using the terminology of Section 3.4
 1184 in the context of the aromatic system. The fractions reacting via the [b] or Bcyc route used when
 1185 generating aromatics mechanisms ranged from 68% to 88%, depending on the number of *ortho*
 1186 substituents in the OH-aromatic adduct. It is highly uncertain whether this branching ratio is an
 1187 appropriate basis for estimations for acyclic peroxy radicals with conjugated double bonds, but no
 1188 information is available to justify other assumptions. Therefore, when generating mechanisms for peroxy
 1189 radicals with conjugated double bonds in the α -, β -, or γ - positions, we assume that:

$$1190 \quad \text{Cyclic peroxide Fraction} = k_{b5}/(k_{b5} + k_{c5}) = k_{b8}/(k_{b6} + k_{c6}) = k_{b7}/(k_{b7} + k_{c7}) \approx 0.75$$

$$1191 \quad \text{Epoxy alkoxy Fraction} = 1 - \text{Cyclic peroxide Fraction} \approx 0.25$$

1192 where the fractions are approximately the averages of those given in Table 7. This is highly uncertain, but
 1193 at least it is reasonably consistent with our treatment of aromatic ring addition reactions. However,
 1194 formation of these types of peroxy radicals is probably relatively rare in atmospheric systems, so the
 1195 practical effect of this uncertainty on atmospheric mechanisms may be relatively small.

1196 Note that, as discussed in Section 3.4, the epoxide forming cyclization reaction may not be as
 1197 important in aromatic systems as currently estimated, and if this is the case then their analogues (reactions
 1198 labeled [c] above), may also not occur in acyclic systems. This will need to be examined when MechGen
 1199 is updated.

1200 It is also possible that ring closure involving only single double bonds may be non-negligible.
 1201 However, if such reactions were to be significant, they should be important in the isoprene + OH reaction
 1202 system, resulting in formation of different products than what are observed (Calvert et al., 2011, and
 1203 references therein). Therefore, we currently assume that these cyclizations are only important in radicals
 1204 with conjugated double bonds.

1205 **5. Reactions of Alkoxy Radicals**

1206 Alkoxy radicals are formed in atmospheric oxidation mechanisms primarily from the reactions of
 1207 peroxy radicals with NO and NO₃, and to a lesser extent by their reactions with other peroxy radicals, as

1208 discussed above. These can undergo a variety of reactions in atmospheric systems, and this leads to much
1209 of the complexity in atmospheric reactions of organic compounds. The types of alkoxy radical reactions
1210 that MechGen considers, and how their rate constants are estimated, are summarized in Table 12. These
1211 reactions are discussed further below.

1212 Note that alkoxy radicals formed in the reactions of peroxy radicals with NO could be formed
1213 with up to ~12 kcal/mole of excitation energy, which would affect estimates for unimolecular reactions
1214 that are not highly thermochemically favorable (Orlando et al., 2003). This is not taken into account in
1215 current estimates of unimolecular reactions of these radicals, but will need to be considered for future
1216 updates of the system.

1217 **5.1. Bimolecular Reactions**

1218 Primary and secondary alkoxy radicals can react with O₂ via abstraction of an α hydrogen to form
1219 HO₂ and the corresponding carbonyl compound, as shown for the first three reactions in Table 12.
1220 Available data concerning rate constants for this reaction are discussed in Section S1.7.1 of the SI. These
1221 data suggest that the rate constant may only depend on whether the radical is primary or secondary, and
1222 the following rate constants are assigned for estimation purposes:

$$1223 \quad k(-\text{CH}_2\text{O}\cdot + \text{O}_2) = 2.38 \times 10^{-14} \exp(-0.38/RT) = 1.25 \times 10^{-14} \text{ cm}^3 \text{ molec}^{-1} \text{ s}^{-1} \quad (k^{\text{uni}} = 6.4 \times 10^4 \text{ sec}^{-1})$$

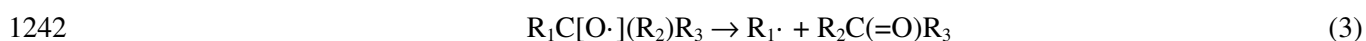
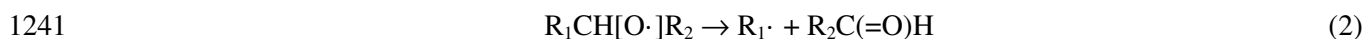
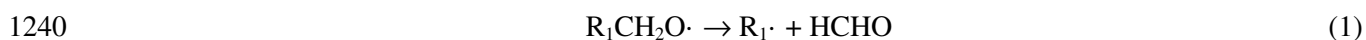
$$1224 \quad k(-\text{CH}[\text{O}\cdot] + \text{O}_2) = 1.95 \times 10^{-14} \exp(-0.59/RT) = 7.27 \times 10^{-14} \text{ cm}^3 \text{ molec}^{-1} \text{ s}^{-1} \quad (k^{\text{uni}} = 3.8 \times 10^4 \text{ sec}^{-1})$$

1225 where the rate constant is given for 298 K and k^{uni} is the pseudo-first order rate constant for 1 atmosphere
1226 of air. Note that separate assignments are given for rate constants for reactions of O₂ with methoxy or
1227 ethoxy radicals (see Table S-56 in the SI), but there is no indication of effects of radical size for larger
1228 radicals, though this cannot be ruled out for C₆₊ radicals. There are also no data on effects of non-alkyl
1229 substituents or groups on this rate constant, so these estimates are highly uncertain for non-alkyl radicals.

1230 Although alkoxy radicals can also react with NO_x and radical species with relatively high rate
1231 constants, the atmospheric levels of NO_x are many orders of magnitude less than those of O₂, and the rate
1232 constants are not sufficiently high that these reactions are competitive. Therefore, reaction with O₂ is the
1233 only type of bimolecular reaction generated for alkoxy radicals. The only exceptions for the types of
1234 radicals handled by MechGen are tertiary alkoxy radicals, where β-scission decompositions are estimated
1235 to be favorable, or phenoxy radicals, which are discussed separately in Section 6.1.

1236 **5.2. β-Scission Decompositions**

1237 Most alkoxy radicals can undergo β-scission reactions, where a bond on the carbon bonded to the
1238 oxy radical center breaks, forming a carbonyl group and a fragment radical. These types of reactions can
1239 be represented as:



1244 Table 12. Summary and examples of reactions of alkoxy radicals as derived using MechGen.

Types	Reactions [a]	Rate Constant Estimation
Reaction with O ₂	-CH ₂ O· + O ₂ → HO ₂ + -CHO -CH(O·) + O ₂ → HO ₂ + -C(O)-	See Sections 5.1 and A1.7.1
	HC(O)O· + O ₂ → HO ₂ + CO ₂	Assumed to be fast [b]
	<u>X</u> ₂ C=C <u>X</u> O· + O ₂ → <u>X</u> ₂ C(OO·)C(O) <u>X</u>	Assumed to be fast [b, c]
β-Scission decompositions	XCH ₂ O· → X· + HCHO XCHX'O· → X· + X'CHO XCX' ₂ O· → X· + X'C(O)X'	See Sections 5.2 and A1.7.2.
	XC(O)O· → X· + CO ₂	Assumed to be fast [b]
H-Elimination decompositions	XCH ₂ O· → XCHO + H· XCH(O·)X' → XC(O)X' + H·	See Sections 5.3 and A1.7.3.
H-Shift isomerizations	HC <u>X</u> ₂ -(Y) _n -C <u>X</u> ₂ O· → ·C <u>X</u> ₂ -(Y) _n -C <u>X</u> ₂ OH	See Sections 5.4 and A1.6.
α Ester and α Nitrate Rearrangement	<u>X</u> CH(O·)OC(O) <u>X</u> ' → <u>X</u> C(O)· + <u>X</u> 'C(O)OH <u>X</u> CH(O·)ONO ₂ → <u>X</u> C(O)· + HONO ₂	See Sections 5.5 and A1.7.4.

1245 [a] The following codes used for unspecified groups: X = Any non-radical group except -H; X = -H or
1246 any non-radical group; Y = Any stable compound group that is bonded to at least two other groups,
1247 except =C=, ≡C-, and -aC_x; Y = As above, but also excluding =CH- and =C<

1248 [b] This is assumed to dominate over competing reactions of this radical, so the rate constant does not
1249 need to be estimated.

1250 [c] Vinyloxy radicals are expected to exist in resonance with a carbon-centered radical form, e.g.,
1251 X₂C=CXO· ↔ X₂C[·]C(O)X, which rapidly adds O₂ under atmospheric conditions.

1252

1253

1254 where the R's are any groups next to the alkoxy radical group except -H. Decompositions of acyl oxy
1255 radicals (Reaction 4) are expected to be fast and thus are the only fate of these radicals considered in
1256 MechGen. However, this is not the case for alkoxy radicals in general (Reactions 1-3) where, depending
1257 on the structure of the radical, these reactions can be very fast and dominate over other reactions, or they
1258 can be negligible compared to reaction with O₂ or other types of unimolecular reactions such as H-shifts.
1259 Note that the relative importance of the decomposition reactions affects the extent to which the reactions
1260 of a VOC leads to fragmentation to lower molecular-weight products, as opposed to becoming more
1261 oxidized and forming lower volatility products that may contribute to SOA formation.

1262 Available information concerning rate constants for β-scission reactions, and methods for their
1263 estimation, are discussed in Section S1.7.2 of the SI. These include experimentally measured rate
1264 constants given by Orlando et al. (2003), theoretically calculated rate constant parameters from
1265 Vereecken and Peeters (2009), and rate constants derived from results of experimental product studies.
1266 Based on the data and estimates in these resources, MechGen uses the following equations to estimate
1267 rate constants for β-scission reactions of alkoxy radicals:

1268 $k(T) (\text{sec}^{-1}) = n \times 1.0 \times 10^{14} \times \exp(-E_a/RT)$
 1269 $E_a(\text{kcal/mole}) = E_aR(\text{Rad}) + E_aP(\text{ProdType}) + E_aRc(\text{R.Subst}) + E_aPc(\text{P.Subst}) + R.\text{Corr}(\text{Ring})$

1270 Here k is the unimolecular rate constant at temperature T in sec⁻¹; n is the number of equivalent R₁ groups
 1271 on the radical; 1.0 x 10¹⁴ sec⁻¹ is the estimated A factor for all β-scission reactions; E_a is the calculated
 1272 activation energy that depends on the reaction; and E_aR, E_aP, E_aRc, E_aPc, and R.Corr are estimated or
 1273 adjustable parameters that depend on the reaction. In addition, "Rad" refers to the radical R₁· formed in
 1274 the reaction; "ProdType" refers to whether the carbonyl product formed is formaldehyde, a higher
 1275 aldehyde, or a ketone (reactions 1-3, respectively); "R.Subst" refers to substituents on R₁; "P.Subst" refers
 1276 to substituents R₂ or R₃ if applicable; and "Ring" refers to the presence of rings on the alkoxy radical at or
 1277 adjacent to the radical center. The derived or estimated values of these parameters are given in various
 1278 tables in Section S1.7.2 of the SI. Values of representative parameters are given in Table 13.

1279
 1280 Table 13. Representative values of parameters used to estimate activation energy of β-scission
 1281 reactions.

Radical Formed [a]		Product Formed		Substituent Corrections			Ring Corrections		
Type	E _a R	Type	E _a P	Substituent	E _a Rc	E _a Rp	Type	Size	R.Corr
Methyl	18.5	HCHO	0	-Alkyl	0	0	Ring is	3	-7.1
Primary	15.5	C ₂₊ Aldehyde	-2.2	-Allyl	-5.0	-2.0	broken	6	0.0
Secondary	12.8	Ketone	-3.9	-C(=O)X	-1.3	0.8			
Tertiary	10.7			-OX	-3 [b]	-4 [b]	Ring on	3	2.0
HC(O).	11.4			-ONO2	0 [b]	-1.1	radical	6	1.8
RC(O).	9.3			-CH _x -O-	0	1.1			
Alkoxy	18 [b]			-CH _x -OH	0	-0 [b]	Ring on	3	7.2
							product	6	-0.3

1282 [a] Parameters are all in units of kcal/mole. "X" refers to -H or any other substituent.

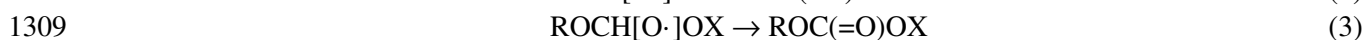
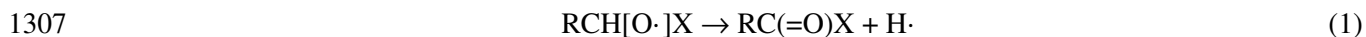
1283 [b] The value of this parameter varies somewhat depending on the specific groups involved, and the
 1284 average value is given. See the tables in Section S1.7.2 of the SI for the specific values.
 1285

1286 From Table 13 it can be seen that the estimated activation energies range from near zero to ~20
 1287 kcal/mole or higher. Note that an activation energy of ~12 kcal/mole would give a rate constant that is
 1288 comparable to the pseudo-unimolecular rate constant for the reactions of primary or secondary alkoxy
 1289 radicals with O₂, so reactions with higher activation energies would generally be negligible, while those
 1290 with lower E_a values could potentially be the dominant process unless another type of unimolecular
 1291 reaction (e.g., a β-scission involving a different group or an H-shift) is also estimated to be fast.
 1292 Decompositions are usually important for tertiary alkoxy radicals not only because reactions with O₂ are
 1293 not possible, but because their β-scission decompositions are generally estimated to be relatively fast.
 1294 Decompositions can also be important for primary or secondary radicals.

1295 As discussed in the SI, there are sufficient data on rate constants for decompositions of alkoxy
 1296 radicals formed from alkane systems that estimates in these cases are not considered to be highly
 1297 uncertain. However, if non-alkyl groups are present, data to derive estimates are highly limited or lacking
 1298 and thus needed. One uncertainty is the lack of measured or theoretically calculated rate constants for
 1299 decompositions forming alkoxy radicals, which might occur in atmospheric photooxidation mechanisms
 1300 of ethers and esters and compete with other reactions discussed below.

1301 **5.3. H-Elimination Decompositions**

1302 Primary and secondary alkoxy radicals can also undergo β -scission reactions by H-elimination,
1303 forming a carbonyl product and a hydrogen atom. Available information and estimates for the rate
1304 constants for these reactions are discussed in Section S1.7.3 of the SI. The rate constants for these
1305 reactions appear to be enhanced by the presence of α -oxygen substituents, so three types of these
1306 reactions are considered.



1310 Here X is -H or any group, and R is any group except -OX. The data are limited, but sufficient to derive
1311 the following expression to estimate the rate constant:

1312
$$k(\text{T}) (\text{sec}^{-1}) = n \times 3.3 \times 10^{13} \times \exp(-E_a/\text{RT})$$

1313
$$E_{a1} = E_{a2} (\text{kcal/mole}) = 13.5 + 0.52 \Delta H_r$$

1314
$$E_{a3} (\text{kcal/mole}) = 11.5 + 0.52 \Delta H_r$$

1315 where E_{a1} , E_{a2} , E_{a3} refer to the activation energies for reactions of types 1, 2, and 3, respectively. The
1316 estimate for E_{a3} is uncertain and is strictly speaking an upper limit -- i.e., rate constants for H-elimination
1317 reactions of alkoxy radicals with two -OX may be higher than estimated.

1318 Note that under atmospheric conditions, the H-elimination reactions yield exactly the same
1319 products as the reactions the alkoxy radicals with O_2 , because the eliminated $\text{H}\cdot$ rapidly forms HO_2 . These
1320 estimates indicate that the H-elimination reaction is negligible compared to O_2 reaction if there are no α -
1321 O substituents, non-negligible but slower than O_2 reaction if there is only one, and faster than the O_2
1322 reaction if there are two such substituents. Therefore, this reaction is only important in the third case, so
1323 only estimates for E_{a3} have a practical effect on generated mechanisms.

1324 **5.4. H-Shift isomerizations**

1325 Alkoxy radicals with sufficiently long chains ($\approx \text{C4}$ or larger) can also undergo H-shift
1326 isomerization reactions, where the alkoxy group abstracts an H from elsewhere in the molecule via a
1327 cyclic transition state, forming a carbon-centered radical with an -OH substituent. These are similar to the
1328 H-shift isomerizations that were previously discussed for peroxy radicals (Section 4.6), but in this case
1329 the reactions are much more exothermic, and their importance in atmospheric systems has been
1330 recognized for some time. Methods to estimate their rate constant have been developed for previous
1331 versions of the SAPRC mechanism, with the version used for developing SAPRC-99 being the most
1332 completely documented (Carter, 2000). The method used for SAPRC-99 has been subsequently updated
1333 for SAPRC-18 to incorporate evaluated experimental data (Atkinson, 2007; IUPAC, 2023), results of
1334 theoretical quantum chemistry calculations of Vereecken and Peeters (2010), and upper limit rate
1335 constants derived from various product studies, as given in Table S-50 of the SI. This updated method is
1336 documented in the SI in Section S1.6.3.

1337 The rate constants for alkoxy H-shift reactions are estimated in the same way as for peroxy
1338 radicals, using Equation (I) in Section 4.6, which is based on the types of groups from which the H is
1339 abstracted, their substituents, and also substituents elsewhere in the transition state ring. Because of more
1340 limited amount of theoretically calculated data available at the time this method was last updated in the
1341 mid-2010's, only a single set of ring-size-dependent A factors and E_a^{strain} values were derived, and E_a^0
1342 values were derived using only four types of H-groups (- CH_3 , - CH_2 -, - CH -, and - OOH), with no
1343 distinction being made between primary, secondary, or tertiary alkoxy radicals. To compensate for this, a

1344 larger number of substituent correction factors were employed. The A factors and strain energies used are
 1345 given in Table S-25, the E_a^0 values are given in Table S-26, and the E_a^{corr} values are given in Table S-16
 1346 and Tables S-27 and S-28 in the SI. Footnotes to these tables indicate how these were derived or
 1347 estimated.

1348 Estimated 298 K rate constants for H-shift isomerizations of various unsubstituted alkoxy radicals
 1349 are listed in Table 14, which also shows the ratios of these rate constants to the pseudo-unimolecular rate
 1350 constant for the reaction of secondary alkoxy radicals with O_2 in the atmosphere. It can be seen that 1,4-H
 1351 shift isomerizations, with 6-member ring transition states, are ~10-350 times faster than O_2 reaction, while
 1352 those involving 1,5-H shifts are ~4-130 times faster, while 1,2, 1,3, and 1,8+ H shifts are negligible to
 1353 minor.

1354 Note that the current estimation methods do not consider effects of functional groups that are
 1355 between the radical center and the H atom being transferred. This may be considered in the future as more
 1356 experimental or theoretical data become available.

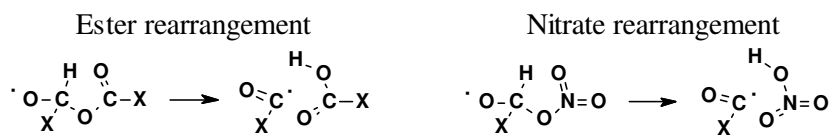
1357 Although there are uncertainties in these estimates, particularly if non-alkyl groups or substituents
 1358 are present, their effects on generated mechanisms would be minor if these reactions are estimated to be
 1359 either very slow or very fast compared to competing processes. Uncertainties will affect generated
 1360 mechanisms primarily when there are competing β -scission decompositions with comparable estimated
 1361 rate constants.

1362
 1363
 1364 Table 14. Representative 298 K rate constants for H-shift isomerizations of unsubstituted alkoxy
 1365 radicals.

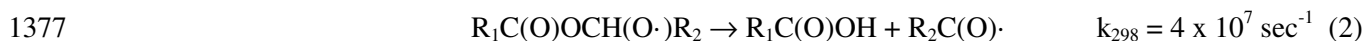
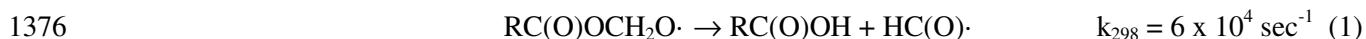
Ring size	Shift from $-CH_3$		Shift from $-CH_2-$		Shift from $-CH<$		Shift from $-CHO$	
	k(sec-1)	k/k[O_2]	k(sec-1)	k/k[O_2]	k(sec-1)	k/k[O_2]	k(sec-1)	k/k[O_2]
5	8.9e-3	2e-7	8.5e-2	2e-6	2.8e-1	8e-6	3.3e-1	9e-6
6	3.5e+5	9.30	3.3e+6	89	1.1e+7	295	1.3e+7	346
7	1.3e+5	3.48	1.2e+6	33	4.1e+6	110	4.9e+6	130
8	4.3e+2	0.01	4.1e+3	0.1	1.4e+4	0.4	1.6e+4	0.4
9	7.2e+1	2e-3	6.8e+2	0.02	2.3e+3	0.06	2.7e+3	0.07

1366
 1367
 1368 **5.5. Ester and Nitrate Rearrangements**

1369 Alkoxy radicals with an α hydrogen and either an ester or nitrate group in the α position can
 1370 undergo a second type of H-shift isomerization, where the α hydrogen transfers to the carbonyl or nitrate
 1371 oxygen and the bonds rearrange to form a carbonyl radical and a carboxylic or nitric acid, via a 5-member
 1372 ring transition state, e.g.,:



1373
 1374 Evidence for these reactions and their measured or estimated rate constants are discussed in Section
 1375 S1.7.4 of the SI. Based on limited data, the following rate constants are estimated:



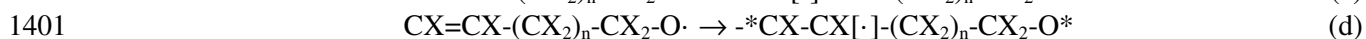
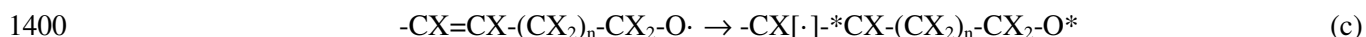
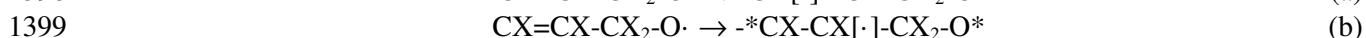
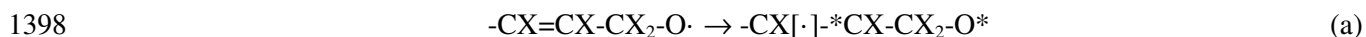
1379 This predicts that the ester rearrangement of primary alkoxy radicals with ester substituents occurs at a
 1380 similar rate as reaction with O₂ under atmospheric conditions, while it is the dominant fate for most
 1381 secondary alkoxy radicals with ester substituents, being predicted to be faster than possible competing H-
 1382 shift or β-scission decompositions, as well as the reaction with O₂. On the other hand, the nitrate
 1383 rearrangement is predicted to be slower than the O₂ reaction, so it is not expected to be important in
 1384 atmospheric systems.

1385 Formation of α-nitrato alkoxy radicals that might undergo the nitrate rearrangement is not
 1386 expected to be as important in atmospheric oxidation systems, because α-nitrato carbon-centered radicals
 1387 formed in reactions of organic nitrates that could be precursors to these radicals rapidly decompose by
 1388 eliminating NO₂, as discussed above in Section 3.2.1, rather than reacting to form the alkoxy radical.
 1389 Therefore, any uncertainties in estimating rate constants for this reaction are not expected to be important.

1390 Radicals that can undergo the ester rearrangement are predicted to be formed in the atmospheric
 1391 oxidation of esters, so this reaction is a factor in predicting their atmospheric mechanisms. The
 1392 uncertainties in the estimated rate constant have a significant effect on predictions for esters forming
 1393 primary alkoxy radicals because it is competitive with the O₂ reaction, but is not as important for those
 1394 forming secondary radicals because the ester rearrangement is predicted to dominate.

1395 5.6. Cyclization Reactions

1396 Unsaturated alkoxy radicals could possibly undergo cyclization reactions where the alkoxy
 1397 radical adds to the double bond, e.g.



1402 These reactions are not considered in the current version of MechGen because they are not consistent with
 1403 available product data on the reactions of OH with isoprene (Calvert et al., 2000), where the observation
 1404 of methacrolein and methyl vinyl ketone as major products indicates that these reactions do not compete
 1405 with the β-scission reaction forming α-hydroxy alkyl radicals. However, theoretical calculations and
 1406 experimental data suggest that the epoxide formation reaction of β-unsaturated alkoxy radicals [process
 1407 (a), above] is important following the addition of NO₃ radicals with isoprene; Carlsson et al. (2023) and
 1408 Vereecken (2018) suggested that this reaction may also account for the missing mass balance in aromatic
 1409 systems (see Section 3.4). This epoxide forming reaction being non-negligible in aromatic systems but
 1410 not in the case of isoprene oxidation could be attributed to beta-scission decompositions forming a-nitrato
 1411 or a-peroxy-substituted alkyl radicals in the aromatic systems being much faster than decompositions
 1412 forming α-hydroxy radicals formed in the isoprene systems. However, our current estimates discussed in
 1413 Section 5.2 and the SI are based on the isoprene system. Note, however, that the competing cyclization
 1414 reaction (b) is estimated to be somewhat more thermochemically favored, and cyclizations forming larger
 1415 rings [e.g. (c) and (d), above] are even more thermochemically favored, though they have not been
 1416 suggested in published mechanisms to our knowledge.

1417 None of these alkoxy cyclization reactions are currently considered by MechGen. However,
 1418 based on recent results discussed above, they will need to be considered when MechGen is updated.

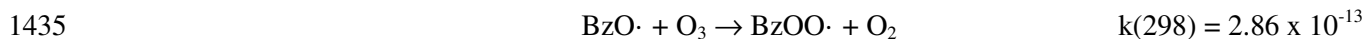
1419 6. Reactions of Other Types of Radicals

1420 The previous sections focused on the reactions of unsubstituted and substituted alkyl radicals, and
 1421 the peroxy and alkoxy radicals formed from them in the presence of O₂ and NO_x, which are the major
 1422 types of radicals formed in atmospheric oxidation systems. Other types of radicals are also formed in
 1423 some circumstances and the treatments of these are discussed in the subsections below. Note that CIs are
 1424 discussed separately in Section 7.

1425 6.1. Phenoxy Radicals

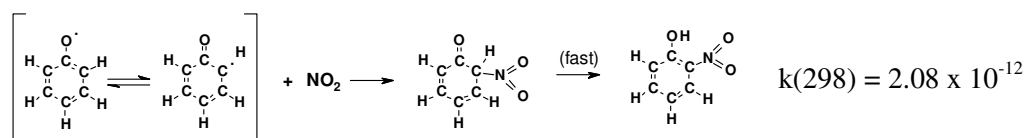
1426 Phenoxy radicals are formed in the reactions of phenols, aromatic aldehydes, and potentially
 1427 other aromatic compounds. They cannot react with O₂ or undergo any of the unimolecular alkoxy radical
 1428 reactions discussed in Section 5. Therefore, their only atmospheric fate is expected to be reactions with
 1429 NO_x species or other radicals. MechGen assumes that their major sinks in atmospheric systems are either
 1430 reaction with O₃, NO₂ or HO₂. Reactions with NO are ignored because they are expected to form nitroso
 1431 species or nitrites that would rapidly photolyze to re-form the reactants, and because the concentrations of
 1432 NO are usually lower than NO₂ in the atmosphere.

1433 MechGen assumes that the mechanisms for the reactions of phenoxy with O₃ and HO₂ are as
 1434 expected for general alkoxy systems and involve simple O or H-transfers, as follows:



1437 The rate constants are in cm³ molec⁻¹ s⁻¹ and that for O₃ is from Tao et al. (1999) and that for HO₂ is
 1438 estimated to be the same as for methoxy + HO₂ (Assif et al., 2018). The O₃ reaction is not a net sink for
 1439 phenoxy radicals because they can be regenerated by reactions of phenyl peroxy radicals with NO. This
 1440 reaction will also convert NO to NO₂, which can also regenerate O₃ under atmospheric conditions. On the
 1441 other hand, the reaction with HO₂ is a net sink for both HO₂ and phenoxy, though it will regenerate the
 1442 phenol whose reactions are the source of the phenoxy radicals.

1443 The reactions of phenoxy radicals with NO₂ are expected to be the main sink for these radicals in
 1444 the presence of NO_x. Alkoxy radicals are expected to react with NO₂ forming the corresponding nitrate,
 1445 but for most alkoxy radicals this is minor compared to reaction with O₂ or unimolecular reactions, so
 1446 these reactions can be ignored. In the case of phenoxy, the analogous formation of benzyl nitrate is not
 1447 expected to be an overall sink for the phenoxy radical because the BzO-NO₂ bond is so weak (Batiha et
 1448 al., 2012) that the phenyl nitrate would be expected to rapidly decompose to reactants resulting in no net
 1449 reaction, analogous to the formation of alkyl peroxy nitrates discussed in Section 2.3. Instead, the reaction
 1450 is assumed to occur as follows:



1451 where the rate constant is from Platz et al. (1998). If the initially formed product has an H atom on the
 1452 same carbon where the NO₂ added it will undergo a fast unimolecular rearrangement to a nitrophenol as
 1453 shown above and discussed in Section 2.3.2. This is consistent with theoretical studies (e.g., Batiha et al.,

1454 2012) and products formed in reactions of phenols with NO₃ (e.g., Calvert et al., 2002, 2011). However,
1455 this fast reaction is not possible if NO₂ adds to a carbon with another substituent, in which case a 2-nitro-
1456 3,5-cyclohexadienone with a substituent in the 2-position is predicted to be the final product.

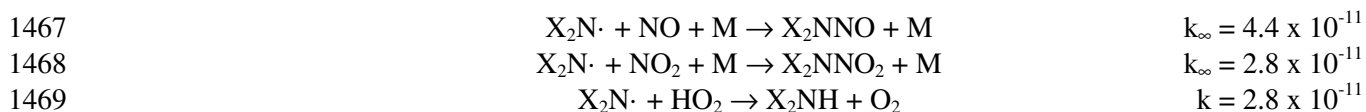
1457 The presence of substituents on the aromatic rings are assumed not to affect the rate constants or
1458 mechanisms of the bimolecular reactions of phenoxy radicals. Note that the *ortho* NO₂ addition can occur
1459 in two different positions for unsymmetrical molecules, and currently MechGen assumes that the reaction
1460 at each position is equally likely, regardless of substituents.

1461 **6.2. Nitrogen-Centered Radicals**

1462 Nitrogen centered radicals can be formed in the atmospheric reactions of some amines, as
1463 discussed in Section 2.1.3 above. Those with α hydrogens are assumed to react primarily with O₂,
1464 forming HO₂ and the corresponding amide,



1466 while those lacking α hydrogens are assumed to undergo only bimolecular reactions, as follows:

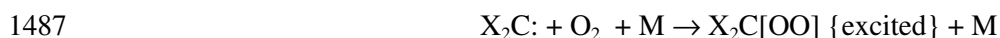


1470 where the rate constants are in units of cm³ molec⁻¹ s⁻¹ and are at the high pressure limit where applicable.
1471 The high pressure rate constants for the reactions with NO and NO₂ are estimated to be the same as the
1472 IUPAC (2023) recommendation for their reactions with ethoxy radicals, while the rate constant for the
1473 reaction with HO₂ is estimated to be approximately the same as given in the NIST kinetics database for
1474 methoxy + HO₂ (Sander et al., 2009). MechGen does not generate reactions of amino radicals without α
1475 hydrogens, but they would only be formed from reactions of amines with more than one tertiary
1476 substituent, which do not appear to be important in current atmospheric emissions.

1477 Note that this mechanism predicts that formation of N-centered radicals in the reactions of
1478 amines without α hydrogens is a radical terminating process, which means that such amines tend to be
1479 radical inhibitors when they react in the atmosphere. This is consistent with results of environmental
1480 chamber experiments with t-butyl amine and 2-amino-2-methyl-1-propanol (Carter, 2008). This is not
1481 expected to be the case with other amines, where results of chamber experiments do not indicate
1482 significant radical inhibition in their reactions.

1483 **6.3. Carbenes**

1484 Carbenes are predicted to form in some photolysis reactions, so their reactions also need to be
1485 considered. It is assumed that their major fate is the addition of O₂ to form highly excited Criegee
1486 intermediates, e.g.,

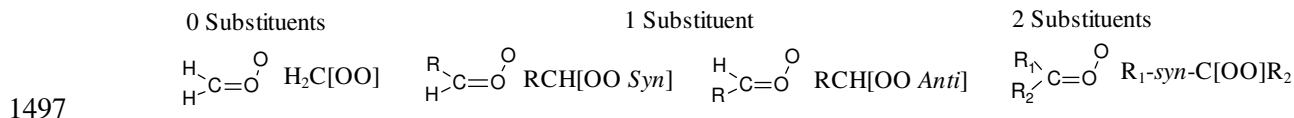


1488 where “{excited}” indicates excitation. Reactions of excited and stabilized CIs are discussed in the
1489 following section. Note that CIs formed from reactions of O₂ with carbenes are expected to have much
1490 higher levels of excitation than those formed from the reactions of O₃ with alkenes, which affects
1491 estimates of stabilization in some cases.

1492

7. Reactions of Criegee Intermediates

1493 Criegee Intermediates (CIs) are highly reactive intermediates with general structure X_2COO that
 1494 are formed primarily in the reactions of O_3 with alkenes. These can have either zero, one, or two
 1495 substituents and those with one or two different substituents can exist in either the *syn* or *anti* form
 1496 depending on the configuration of the substituent(s) relative to the two oxygens:



1498 They are also formed to a lesser extent from the reactions of O_3 with alkynes, and in the reactions of O_2
 1499 with carbenes formed in some photolysis reactions. In all cases they are expected to be formed in a
 1500 vibrationally excited state, with the amount of excitation depending on the formation reaction, though the
 1501 amount of excitation is expected to be variable because, except for reactions of O_3 with cyclic
 1502 compounds, the available energy is distributed among two fragments. In some cases the excitation may be
 1503 sufficient to allow interconversions between *syn* and *anti*, though Vereecken et al. (2017, 2022, and
 1504 references therein) calculated that the isomerization barriers are high, and interconversions of excited CIs
 1505 may be slow in many cases. In any case, the interconversion between *syn* and *anti* is assumed to be slow
 1506 once the intermediate is stabilized. The excited intermediates can undergo unimolecular decompositions
 1507 or be collisionally stabilized, and the stabilized intermediates can also undergo unimolecular reactions or
 1508 react with water, though reactions of stabilized intermediates involving interactions between the
 1509 substituent and the Criegee oxygens requires that the substituent be in the *syn* configuration.

1510 For estimation purposes, we make the assumptions that CIs are initially formed in excited states
 1511 and that the unimolecular reactions of the excited CIs do not depend on *syn* or *anti* configurations. Their
 1512 configurations are determined only after they are stabilized. This is an oversimplification because the
 1513 excited CIs may actually have different configurations and may not always interconvert rapidly compared
 1514 to decompositions. However, available experimental and theoretical data are insufficient to derive the
 1515 additional parameters required to make separate configuration-dependent estimates for excited as well as
 1516 stabilized CIs. In many cases the estimates are constrained by experimental or theoretical results, and this
 1517 informs the parameters used in the estimates. Thus while the parameters may have been different if
 1518 configuration were taken into account for excited CI reactions, the predictions would be similar.
 1519 However, when applied to systems for which no data are available, the parameters and the predictions
 1520 may be different, and are therefore more uncertain.

1521 Information and estimates about the reactions of excited and stabilized CIs are discussed in
 1522 Section S1.9 of the SI. Briefly, the mechanisms adopted for MechGen are based largely on the IUPAC
 1523 (2023) reviews of the reactions of O_3 with ethene and alkenes with methyl substituents, and on the
 1524 theoretical calculations of Vereecken et al. (2017) on the possible unimolecular reactions of the
 1525 intermediates. Note that Vereecken et al. (2022) provided a more recent update on the reactions of
 1526 oxygenated and unsaturated CIs, but these have not yet been taken into account in the current estimates.
 1527 Table 15 lists the types of intermediates and the reactions and branching ratios considered for them. The
 1528 types of reactions considered are summarized below.

1529 Excited intermediates with an alkyl group with α hydrogens, and also stabilized intermediates
 1530 with such a group in the *syn* position, are assumed to undergo the vinyl hydroperoxide (VHP)
 1531 rearrangement, where the α hydrogen transfers to the OO group forming an excited VHP, which then
 1532 decomposes to OH and an α -carbonyl radical (e.g., reactions 8, 24, 25-27, 36, and 38 in Table 15).
 1533 Vereecken et al. (2017) calculated these were relatively fast even for stabilized intermediates.

1534 Table 15. Summary of types of Criegee intermediates and the reactions generated for them

Criegee Type [a]	Reactions [b]	Branching Ratios, notes [c]	
<u>Excited CH₂[OO]</u>	1 Stabilization	0.42 1	
	2 → HCO· + OH	0.17	
	3 → CO ₂ + H ₂	0.18	
	4 → CO + H ₂ O	0.18	
	5 → CO ₂ + 2 H	0.05	
<u>Excited RCH[OO]</u> , where R = X ₂ CH-	6 Stab. to X ₂ CHCH[OO <i>syn</i>]	0.33 x SF ₁ 2, 3, 4	
	7 Stab. to X ₂ CHCH[OO <i>anti</i>]	0.67 x SF ₁	
	8 → ·CX ₂ CHO + OH	0.49 x (1 - SF ₁) 2	
	9 → R· + ·C(=O)OH	0.10 x (1 - SF ₁)	
	10 → RH + CO ₂	0.26 x (1 - SF ₁)	
	11 → ROH + CO	0.14 x (1 - SF ₁)	
	R = OH	12 → HC(=O)O· + OH	Reaction fast 5
	R = CHX ₂ -CX'=CX''-	13 → ·CX ₂ CX'=CX''CHO + OH	Reaction fast 5
	R = CX ₂ =CX-	14 Stab. to CX ₂ =CXCH[OO <i>syn</i>]	0.33 x SF ₁ 3, 4
		15 Stab. to CX ₂ =CXCH[OO <i>anti</i>]	0.67 x SF ₁
		16 → ·CX ₂ C(=O)X + ·C(=O)H	1 - SF ₁ 6
R = none of the above	17 Stab. to RCH[OO <i>syn</i>]	0.33 x SF ₁ 3, 4	
	18 Stab. to RCH[OO <i>anti</i>]	0.67 x SF ₁	
	19 → R· + ·C(=O)OH	0.10 x (1 - SF ₁) 7	
	20 → RH + CO ₂	0.26 x (1 - SF ₁)	
	21 → ROH + CO	0.14 x (1 - SF ₁)	
SF ₁ (nC) = min (1.0 , 0.21 + 0.065 x nC); SF ₁ (2) = 0.35; SF ₁ (12) = 1.0		4	
<u>Excited R₁C[OO]R₂</u> , where R ₁ = X ₂ CH- and R ₂ = X ₂ CH-	22 Stab. to R ₁ -syn-C[OO]R ₂	0.5 x SF ₂ 4	
	23 Stab. to R ₁ C[OO]-syn-R ₂	0.5 x SF ₂	
	24 → ·CX ₂ C(=O)R ₂ + OH	0.5 x (1 - SF ₂) 8	
	25 → R ₁ C(=O)CX' ₂ · + OH	0.5 x (1 - SF ₂)	
	R ₁ = OH, any R ₂	26 → R ₂ C(=O)O· + OH	Reaction fast 8
R ₁ = CHX ₂ -CX'=CX''-, R ₂ not OH	27 → ·CX ₂ CX'=CX''C(=O)R ₂ + OH	Reaction fast 8	
R ₁ = CX ₂ =CX-; R ₂ = not OH or CHX ₂ -CX=CX-	28 Stab. to CX ₂ =CX-syn-C[OO]-R ₂	0.5 x SF ₂ 4	
	29 Stab. to CX ₂ =CX-C[OO]-syn-R ₂	0.5 x SF ₂	
	30 → ·CX ₂ C(=O)X + ·C(=O)R ₂	1 - SF ₂ 6	
R ₁ = X ₂ CH- and R ₂ none of above	31 Stab. to R ₁ -syn-C[OO]R ₂	0.5 x SF ₂ 4	
	32 Stab. to R ₁ C[OO]-syn-R ₂	0.5 x SF ₂ 8	
	33 → ·CX ₂ C(=O)R ₂ + OH	1 - SF ₂	
R ₁ , R ₂ = none of the above	34 Stab. to R ₁ -syn-C[OO]R ₂	0.5 9	
	35 Stab. to R ₁ C[OO]-syn-R ₂	0.5 9	
SF ₂ (nC) = min (1.0 , 0.13 + 0.072 x nC); SF ₂ (3) = 0.35; SF ₂ (12) = 1.0			
<u>Stabilized X₂CHCH[OO <i>syn</i>]</u>	36 → ·CX ₂ CHO + OH	Reaction fast 5	
<u>Stabilized X₂C=CXCH[OO <i>syn</i>]</u>	37 → ·CX ₂ C(=O)X + HC(=O)·	Reaction fast 6	
<u>Stabilized X₂CH-syn-C[OO]R₂</u>	38 → ·CX ₂ C(=O)R ₂ + OH	Reaction fast 6	

Table 15 (continued)

Criegee Type [a]	Reactions [b]	Branching Ratios, notes [c]	
Stabilized $X_2C=CX\text{-syn-C[OO]R}$	39 $\rightarrow \cdot CX_2C(=O)X + \cdot C(=O)R$	Reaction fast	6
Other stabilized $XC[OO]X'$	40 $+ H_2O \rightarrow X_2C(OH)OOH$	If H_2O option	10
	41 $\rightarrow *X_2COO*$ (dioxirane)	If not H_2O option	10

1535 [a] X = H or any group; R = any group other than -H or as indicated below; "*syn*" and "*anti*" refer to the
 1536 orientation of the group with respect to the C-OO bond; "*" refers to ring closure; SF is stabilization
 1537 fraction; and nC is the number of carbons in the intermediate.

1538 [b] The first column gives the reaction numbers, which are referenced in the discussion in the text.

1539 [c] Notes for the derivations of the branching ratios are as follows.

1540 1 The reactions and branching ratios of excited $CH_2[OO]$ are based on the recommendations of
 1541 IUPAC (2023) in their review of ethene + O_3 reactions.

1542 2 The reactions and branching ratios of excited $CH_3CH[OO]$ are based primarily on the
 1543 recommendations of IUPAC (2023) in their review of the reactions of O_3 with propene and the 2-
 1544 butenes, with some adjustments as discussed in the SI.

1545 3 The assumption that the *anti* isomer is favored is necessary to improve simulations of OH yields
 1546 from some alkenes. The assumption that the *syn* isomer is formed $\sim 1/3$ of the time gives
 1547 predictions that are reasonably consistent with the data.

1548 4 The stabilization fractions are assumed to increase linearly with the size of the intermediate from
 1549 35% for methyl-substituted intermediates to 100% for intermediates with 16 carbons. This
 1550 improves model predictions but is somewhat arbitrary and other methods for estimating size
 1551 dependence may perform as well or better. It is also assumed that the stabilization fraction is not
 1552 affected by how the CI are formed, except that SF is assumed to be zero for the highly excited
 1553 intermediates formed from O_2 + carbene reactions.

1554 5 These reactions are assumed to be fast based on the calculations of Vereecken et al. (2017) and
 1555 are assumed to dominate over stabilization or competing reactions of the excited intermediates
 1556 when they are possible.

1557 6 These reactions are assumed to be fast for stabilized as well as excited intermediates based on the
 1558 calculations of Vereecken et al. (2017), though not so fast that they dominate over stabilization of
 1559 excited radicals where these are possible. Vereecken et al. (2017) also calculated that they are
 1560 more rapid than the VHP rearrangement, so they dominate in excited disubstituted radicals where
 1561 both are possible.

1562 7 The branching ratios for the "hot acid" decompositions are derived from those used for methyl-
 1563 substituted intermediates, with the yield of the vinyl hydroperoxide (VHP) reaction set to zero
 1564 and the yields of the others adjusted upward to take this into account.

1565 8 The VHP reaction is assumed to be the only fate of this excited intermediate other than
 1566 stabilization. If two different reactions are possible they are assumed to have equal probability;
 1567 otherwise the only possible VHP reaction dominates.

1568 9 Stabilization is assumed to dominate if the VHP, "hot acid" or the fast decomposition reactions
 1569 are not possible.

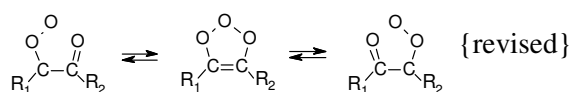
1570 10 Rate constants for these reactions are not estimated for this version, so the " H_2O " option
 1571 determines the major fate assumed for unreactive stabilized intermediates. Note that this ignores
 1572 loss by photolysis, which may be non-negligible.

1573 Vereecken et al. (2017) calculated that intermediates with -OH substituents, rare in the
 1574 atmosphere, will very rapidly rearrange by the O from the -OH group transferring to the [OO] group and
 1575 then decomposing to OH and an acyl oxy radical (which then decomposes to form CO₂ and a radical)
 1576 (reactions 12 and 26 in Table 15). These reactions are assumed to be so rapid that they dominate over
 1577 stabilization.

1578 Excited intermediates with a double bond group in the α position, and also stabilized
 1579 intermediates with such a group in the *syn* position, can react in one of two ways depending on whether
 1580 there is an abstractable hydrogen in the γ position. If there is such a hydrogen, it is assumed that the
 1581 Criegee group abstracts the γ hydrogen to form a highly excited allylic hydroperoxide, which then rapidly
 1582 decomposes to form OH and an allylic carbonyl; shown as a single process in Table 15, reactions 13 and
 1583 27. Vereecken et al. (2017) calculated this to be extremely fast, and we assume that this is sufficiently fast
 1584 that stabilization does not have a chance to occur. If there are no γ hydrogens, it is assumed that the
 1585 Criegee group adds to the double bond forming a highly excited unsaturated peroxy ether, which then
 1586 rearranges and decomposes to ultimately form the ring opening products shown as reactions 16, 30, 37
 1587 and 39 in Table 15. Vereecken et al. (2017) calculates that internal cyclization reactions are more rapid
 1588 than the VHP rearrangement, so it is assumed that cyclization is the major process in excited disubstituted
 1589 radicals where both are possible. Note, however, that Vereecken et al. (2017) only calculated rate
 1590 constants for the formation of the excited unsaturated peroxy ether, but we assume that the subsequent
 1591 rearrangement and decomposition is also fast (see Section S1.9.2 in the SI).

1592 Excited intermediates with zero or one substituent can undergo "hot acid" rearrangements where
 1593 an H bonded to the Criegee group is involved in a rearrangement forming a highly excited acid, which
 1594 then decomposes in several ways, forming either radicals or stable compounds (e.g., reactions 1-5, 9-11,
 1595 and 19-21 on Table 15). These reactions are assumed only to occur when the intermediate is excited.
 1596 These estimates are based on data for simple alkenes and may not correctly predict products formed from
 1597 more complex CIs with a single substituent. Other "hot acids" may not decompose in this way or may be
 1598 mostly stabilized. These pathways will need to be re-examined when MechGen is updated.

1599 MechGen assumes that excited CIs with α-carbonyl groups are in rapid equilibrium with the
 1600 primary ozonide and the isomer where the O is transferred to the other group, forming a different
 1601 carbonyl-substituted intermediate if the groups bonded to them are different, i.e.:



1602
 1603 It further assumes the most reactive isomer determines the subsequent reactions, or that reaction of either
 1604 form is equally likely if they undergo the same types of reactions. Although Vereecken et al. (2017)
 1605 discussed a possible reaction forming primary ozonides, they did not include discussion of this particular
 1606 O-atom rearrangement reaction. However, calculations by Cremer et al. (2011) on the O₃ + acetylene
 1607 system suggests a relatively high barrier to this rearrangement, so this ring closure reaction may be much
 1608 slower than MechGen assumes. This will need to be re-examined when MechGen is updated.

1609 How MechGen processes stabilized intermediates that cannot undergo the above reactions
 1610 depends on whether the "H₂O" option is specified in MechGen. If it is, they are assumed to react with
 1611 H₂O, forming an α-hydroxy hydroperoxide (reaction 40 in Table 15). If not, they are assumed to
 1612 isomerize to the corresponding dioxirane (reaction 41). MechGen treats both of these as non-radical
 1613 products when generating mechanisms. MechGen does not consider effects of substituents in these cases.

1614 MechGen currently does not consider photolysis of stabilized CIs, but loss by photolysis may
1615 well be non-negligible for the less reactive intermediates. IUPAC (2023) evaluated data concerning
1616 photolyses of the simpler Criegee intermediates and gave recommended cross sections for unsubstituted
1617 and methyl substituted intermediates. They also recommended assuming unit quantum yields for the
1618 photolysis forming O³P and the corresponding carbonyl. The recommended absorption cross sections
1619 correspond to rate constants for loss by photolysis of ~0.15 sec⁻¹ with direct overhead sun, which is faster
1620 than the estimated rate constants calculated by Vereecken et al. (2017) for the dioxirane-forming reaction,
1621 but may not be faster than loss by reaction with H₂O, depending on the intermediate and the environment.
1622 Therefore, photolysis of the less reactive stabilized CIs may need to be considered in future versions of
1623 MechGen.

1624 The stabilization factors and branching ratios are given in Table 15 for the various types of CIs,
1625 and footnotes indicate how they were derived. More details are given in Section S3.7 of the SI. The
1626 branching ratios used for unsubstituted and methyl-substituted intermediates are based primarily on
1627 IUPAC (2023) recommendations and include VHP and "hot acid" reactions of the excited intermediate,
1628 with the stabilization fraction being 42% for H₂C[OO] and 35% for the two methyl-substituted
1629 intermediates. It is assumed that the VHP reactions also occur with stabilized intermediates if the
1630 substituent with the group with the α hydrogen is in the *syn* orientation.

1631 Estimates for larger alkyl-substituted intermediates are more uncertain, but the reactions are
1632 assumed to be analogous to those of the methyl-substituted intermediates as long as the VHP reaction is
1633 possible. If the VHP reaction is not possible, then only the "hot acid" reactions are assumed for
1634 monosubstituted intermediates, and only stabilization is assumed to occur for disubstituted intermediates,
1635 where the hot acid reactions are not possible.

1636 The largest uncertainties concern stabilization fractions for the larger CIs, or intermediates
1637 formed in reactions where there is not a separate carbonyl co-product, such as reactions of O₃ with double
1638 bonds in a ring or with triple bonds. As discussed in the SI, best fits to the observed OH yields in O₃ +
1639 alkene reactions (Calvert et al., 2000) and in simulations of NO oxidation and O₃ formation rates in some
1640 chamber experiments with alkenes (e.g., see Carter, 2000, 2010b) are obtained if it is assumed that (1)
1641 stabilization increases with the size of the molecule, (2) stabilization is the same even if all of the energy
1642 is in the intermediate formed, and (3) formation of the *anti* isomer is favored when monosubstituted
1643 Criegees are stabilized, with the enhancement (somewhat arbitrarily) being assumed to be a factor of two.
1644 The assumption that stabilization energies do not affect stabilization of intermediates formed with higher
1645 excitation energies is questionable but is necessary to avoid significantly overpredicting OH yields from
1646 cycloalkenes.

1647 On the other hand, we assume that CIs formed from reactions of carbenes with O₂ are so highly
1648 excited that they react before they are stabilized. In the case of unsubstituted carbenes (reactions 1-5 in
1649 Table 15), we use a branching ratio of 0 for reaction 1 and increase the others accordingly. In the case of
1650 substituted intermediates, the branching ratios are as shown in S-33, calculated using SF₁ or SF₂=0. This
1651 is uncertain, but carbene formation is generally not predicted except in some photolysis reactions (see
1652 Section 2.2.1).

1653 MechGen does not consider all the many possible reactions of CIs or complications considered by
1654 Vereecken et al. (2017) or other more recent theoretical or experimental results (e.g., Vereecken et al.,
1655 2022), somewhat arbitrary assumptions had to be made in some cases, and treatments of excited or
1656 unreactive stabilized intermediates are likely to be overly simplistic. MechGen also does not consider
1657 bimolecular reactions with species other than H₂O, though such reactions may be non-negligible in some
1658 circumstances, such as some laboratory experiments. Therefore, treatments of reactions of CIs must be
1659 considered to be uncertain and are an appropriate area to update in future versions of MechGen.

1660

8. Examples of Results

1661 Although application and analysis of mechanisms for individual compounds are beyond the scope
1662 of the present paper, examples of generated mechanisms, including numbers of reactions and numbers
1663 and types of products, are given for illustrative purposes. We generated mechanisms and product yields at
1664 representative atmospheric NO_x and radical levels (see Section S3 of the SI) for 38 representative
1665 compounds. These include propane, the C₄, C₈, C₁₂ and C₁₆ n-alkanes, 2-methyl alkanes, 1-alkenes, 2-
1666 alkanes, other representative C₈ branched alkanes, alkenes, aromatics, and oxygenates, and representative
1667 terpenes. Note that the generated mechanisms considered only the reactions of the starting compounds
1668 and the radicals they form, not the subsequent reaction of the non-radical products formed. Therefore,
1669 these are only single generation mechanisms; multi-generation mechanisms, such as those derived using
1670 GECKO-A (Aumont et al., 2005) will predict many more products. The specific compounds and selected
1671 results for these compounds are given in Table S-36.

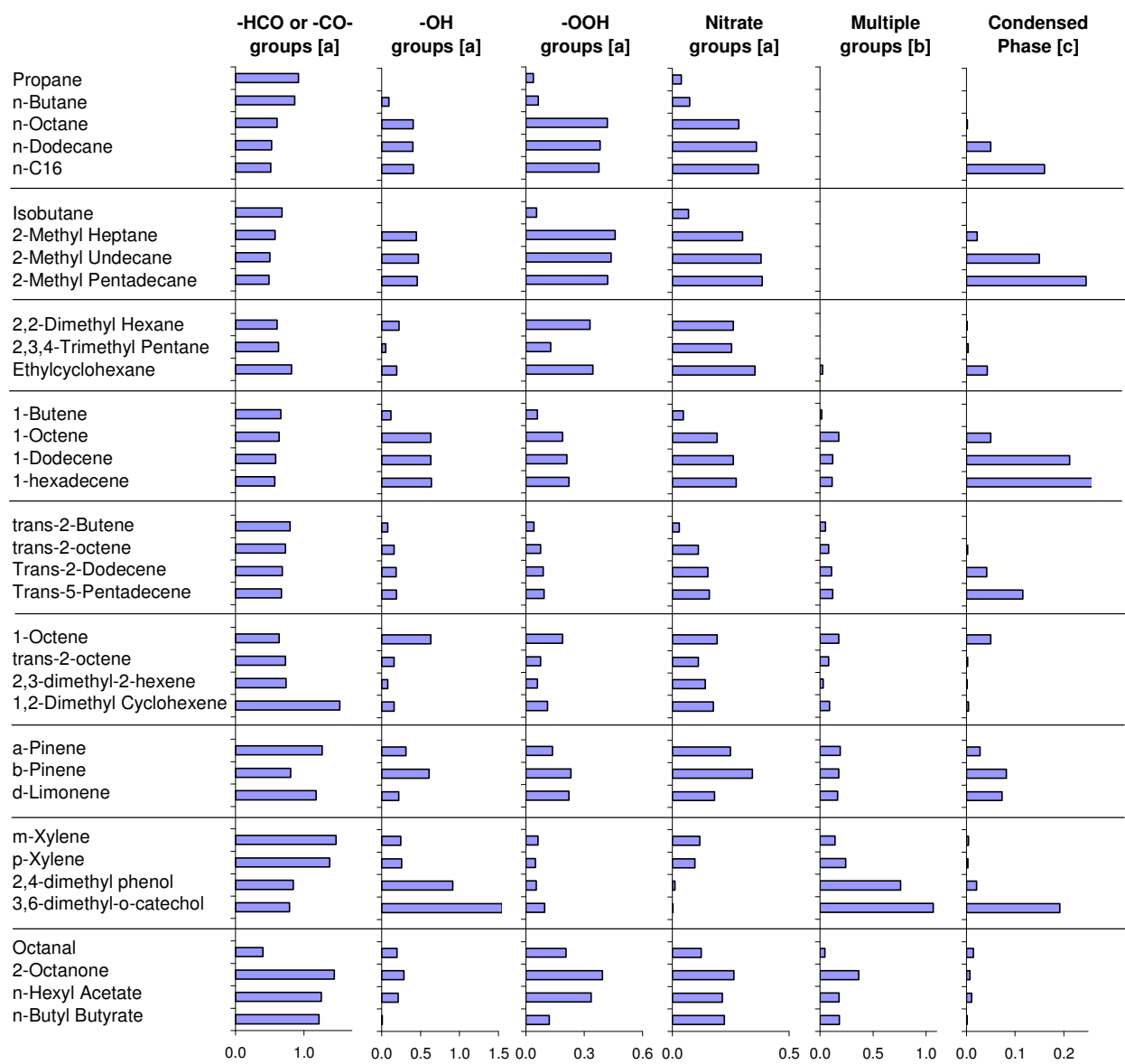
1672 Selected results of the example mechanism generations by MechGen are given in Table S-36 in
1673 the SI and in Figure 4. As discussed in the corresponding MechGen system manuscript (in preparation),
1674 the exact numbers of reactions generated and products predicted depend on the mechanism generation
1675 parameters designed to minimize the numbers of very minor reactions and species predicted, with the
1676 values given in Table S-36 being representative of results when default options are used. The distribution
1677 of products also depends on the environments where the compounds are reacted because they are affected
1678 by competitions involving bimolecular reactions with atmospheric species such as OH, O₃, NO_x species
1679 and peroxy radicals. The yield data shown here were calculated based on reactions under urban conditions
1680 where O₃ formation is equally sensitive to changes in VOC and NO_x, which roughly represent conditions
1681 of the single-day box model scenarios used to derive EBIR (equal benefit incremental reactivity) ozone
1682 reactivity scales (Carter, 1994). More information about how the yields were derived is given in Section
1683 S3.1 in the SI.

1684 Figure 4 shows total yields of compounds containing the types of structural groups formed in the
1685 highest yields. Many or most of the products contain more than one such group on the molecule, and
1686 some compounds can contribute to total yields of more than one group, or (if they have more than one of
1687 the same type of group), may contribute more than one mole to their group total. The category labeled
1688 "multiple groups" gives the total yields of compounds with more than one non-alkyl group bonded to a
1689 carbon, which are types of compounds whose mechanisms and estimated rate constants are particularly
1690 uncertain. The yields of such multifunctional compounds are the highest for some of the terpenes, but are
1691 non-negligible for most types of compounds except alkanes.

1692 MechGen has the capability to estimate vapor pressures of generated compounds, using either the
1693 SIMPOL (Pankow and Asher, 2008) or EVAPORATION (Compernelle et al., 2011) group contribution
1694 methods. The SIMPOL method was used here for illustrative purposes because it provides estimates for a
1695 wider range of compounds. Estimated vapor pressures were used to estimate the equilibrium fractions of
1696 the first-generation products in the condensed phase at an assumed total organic aerosol level of 50
1697 μg/m³. The estimated total fractions of the products in the condensed phase (sum of product yields × the
1698 fraction of each in the condensed phase) are given in Table S-36 in the SI and included in Figure 4. More
1699 information about the method used is given in Section S2.2 of the SI. Figure 5 shows the fraction of
1700 products in the condensed phase as a function of carbon number for all 38 compounds. As expected, the
1701 yields of condensed-phase products increase with carbon number, with the yields becoming significant for
1702 most C₁₂₊ compounds, though highly variable with type of compound. These volatility estimates are based
1703 on uncertain vapor pressure estimates in many cases (Bilde et al., 2015), and, most importantly, ignore the
1704 contributions from products formed in subsequent generation reactions, which tend to form lower
1705 volatility products than first generation reactions (e.g., see Aumont et al., 2012, 2013). A comprehensive

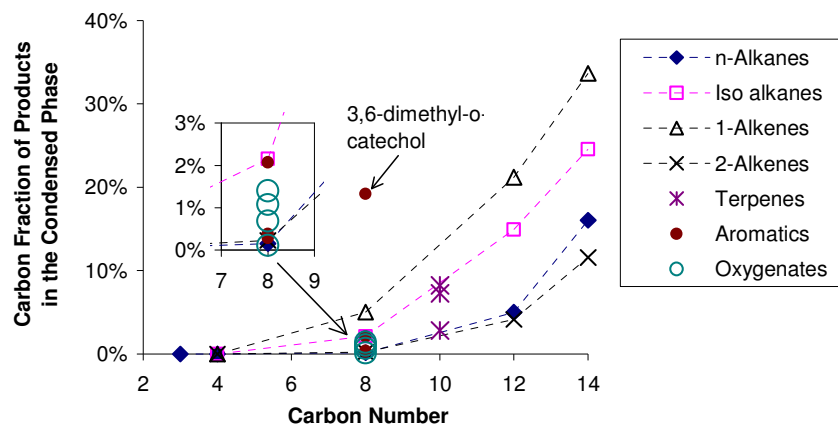
1706 investigation of the use of MechGen to predict low volatility products is beyond the scope of this paper,
 1707 but this is a potentially useful area of future study and comparison with GECKO-A predictions.

1708
 1709



1710 Notes: [a] Sum of carbons in products with the indicated groups; [b] Sum of carbons in multifunctional products with
 1711 two or more different non-alkyl groups on the same carbon; [c] Sum of carbons in products x the fraction of the
 1712 product in equilibrium with the condensed phase for atmospheric organic aerosol levels of 50 $\mu\text{g}/\text{m}^3$.
 1713
 1714

1715 Figure 4. Comparisons of total mole carbon yields of various types of products for generated
 1716 mechanisms for reactions of representative C₈ compounds and terpenes.
 1717



1718
 1719 Figure 5. Estimated fraction of products in the condensed phase at 298 K and with $50 \mu\text{g}/\text{m}^3$ of
 1720 background organic aerosol as a function of carbon number for representative
 1721 compounds.
 1722

1723 9. Discussion and Conclusions

1724 The MechGen system provides a means to derive nearly fully explicit mechanisms for the
 1725 reactions of most organic compounds under lower atmospheric conditions. These can be used as a basis
 1726 for deriving more condensed mechanisms for use in urban, regional, and global models, and serve as
 1727 standards for developing and evaluating mechanism reduction approaches. The level of chemical detail
 1728 that generated mechanisms potentially provide may not be necessary for modeling urban O_3 formation,
 1729 but may well be necessary for developing or evaluating chemically-based models for formation of SOA
 1730 in the atmosphere. MechGen, like GECKO-A (Aumont et al., 2005) also provides a potentially useful
 1731 alternative to the widely-used MCM (MCM, 2023) for evaluating chemical systems for laboratory and
 1732 other studies. Like MCM, it is available to the general user online; but unlike MCM, it can be used to
 1733 derive mechanisms for compounds whose reactions have not been previously estimated.

1734 Explicit chemical detail in itself does not guarantee chemical accuracy or predictive capability.
 1735 Detailed mechanisms that incorporate incorrect rate constants and branching ratios, incorporate reactions
 1736 that do not occur, omit important reactions, or incorporate inappropriate assumptions, cannot be expected
 1737 to give reliable predictions and may provide an illusion of accuracy and understanding that does not exist.
 1738 On the other hand, if the mechanism generation process appropriately incorporates our current knowledge
 1739 and estimates based on best available expert judgment, then the generated mechanism provides our best
 1740 estimate for the reactions of the compound(s) of interest, and, if our knowledge is sufficient, has a
 1741 reasonable chance of predictive capability. However, our current knowledge of atmospheric reactions of
 1742 organics is not complete, so our best judgment as to the most likely detailed mechanism will evolve over
 1743 time, and any mechanism generation system will need to evolve as well so its predictions continue to
 1744 represent the state of the science.

1745 **9.1. Summary of Areas of Mechanism Uncertainty**

1746 Most of the types of chemical reactions discussed in this paper have at least some areas of significant
1747 uncertainty or areas where simplifying assumptions have to be made to avoid excessive complexity that is
1748 either impractical to currently implement, or that is not justified by available knowledge. The impacts of
1749 these uncertainties are expected to vary widely depending on the compounds whose mechanisms are
1750 being generated, the environment where the mechanism will be applied, and the objectives of the model
1751 application. Table 16 lists what are considered to be the major areas of uncertainty in the current
1752 mechanism generation system and gives very brief indications of the associated problems and impacts.
1753 Sections of this paper and SI where the issues are discussed in more detail are also given. This can serve
1754 both as a guideline for areas of research that could be most beneficial for improving detailed mechanisms
1755 and also areas of MechGen that need to be the focus of future updates. These are certainly not the only
1756 areas of uncertainty in developing comprehensive mechanisms but appear based on this work to be
1757 priority areas of research. However, the greatest uncertainty may be reactions or reaction mechanisms that
1758 are not currently considered and are therefore omitted entirely in atmospheric mechanisms or models.

1759 **9.2. Recommendations**

1760 Mechanism generation systems such as MechGen provide an important link between basic kinetic
1761 and mechanistic data and theories, and applications such as air quality modeling for research and
1762 regulatory applications and analysis of laboratory and field data. With regard to regulatory modeling,
1763 Kaduwela et al. (2015) proposed an approach for developing atmospheric chemical mechanisms for the
1764 future, in which the development and applications of detailed chemical mechanism generation systems
1765 played an essential role. Although generated mechanisms are much too large for full three-dimensional
1766 atmospheric modeling, they can serve as a basis for deriving more condensed mechanisms for modeling
1767 application that still have a direct link to the underlying basic data and theories, and where the effects of
1768 the simplifications and condensations can be systematically characterized. Without these links, research
1769 and regulatory modeling may not benefit from the emerging science in atmospheric chemistry.

1770 Collection and evaluation of relevant laboratory and theoretical data are essential to detailed
1771 mechanism development, both for direct incorporation into the mechanisms and to support the
1772 development of the many types of SARs needed for comprehensive mechanism development. The
1773 ongoing IUPAC (2023) and NASA (Sander et al., 2006, 2009) evaluations provide an essential role in
1774 this effort, as have the books by Calvert et al. (2000, 2002, 2008, 2011, 2015) on atmospheric chemistry,
1775 and efforts like these need to continue. Recently McGillen et al. (2020) made available a comprehensive
1776 collection of rate constants for reactions of organics with OH, O₃, NO₃, and Cl atoms, and this data
1777 collection effort is continuing, with the latest update being available at [https://data.eurochamp.org/data-
1778 access/kin/](https://data.eurochamp.org/data-access/kin/). Similar collections are needed for other types of reactions, particularly data to support SAR
1779 development for the many types of radical reactions. The NIST kinetics database (NIST Chemical
1780 Kinetics Database, 2023) is a useful resource, but recommendations are not provided and it is limited to
1781 reactions of simpler molecules and radicals. Quantum theory has become an increasingly important tool
1782 in developing SARs for radical reactions, since experimental data are insufficient for this purpose.
1783 Compilations and evaluations of available theoretical data are needed to take full advantage of this
1784 ongoing work.

1785 Table 16. Summary of major areas of uncertainty in the current mechanism generation system

<u>Uncertainty</u>	<u>Discussion</u> (Section discussed)
Organic + oxidant rate constants	Affects VOC lifetimes and product formation. Not a large uncertainty for the major VOCs or mechanisms for mixtures, but potentially significant for multifunctional products. (Secs. 2.1, A1.2)
Organic photolysis reactions	Affects radical initiation rates as well as VOC lifetimes and fates. Major oversimplifications for larger and multifunctional compounds. Priority area for system update. Photolytic radical from aromatic ring fragmentation products are uncertain. (Secs. 2.2, A1.3)
Unimolecular reactions of non-radical compounds	Affects product formation. MechGen includes one type of reaction that may actually be heterogeneous or H ₂ O catalyzed but ignores others that may be equally important in real environments. (Sec. 2.3)
O ₂ additions to carbon-centered allylic radicals	Affects product formation and predictions of effects of phenolic compounds on radical levels. Insufficient data are available to reliably estimate branching ratios for additions to the different allylic radical centers. (Sec. 3.1)
Cyclization of carbon-centered radicals forming cyclic esters	Affects product formation. Experimental data needed. Existing theoretical studies are inconsistent. Reaction may not be as fast as assumed. (Secs. 3.2.3, A1.4.3)
Other unimolecular reactions of carbon-centered radicals	Affects product formation. Current assumptions are considered reasonable but some need experimental or theoretical verification. (Secs. 3.2, 3.3, A1.4)
Aromatic ring opening reactions	Affects product formation and radical initiation. Current estimates are likely oversimplifications and not all are consistent with all theoretical and product studies. Arbitrary assumptions had to be used in some cases. (Sec. 3.4)
Nitrate yields from NO + substituted peroxy radicals	Affects radical and NO _x sink and recycling for almost all compounds. Experimental data for OH-substituted radicals are inconsistent and there are no direct data for effects of other substituents. (Secs. 4.1, A1.5.2)
H-shift isomerization reactions of peroxy radicals	Affects product formation, particularly hydroperoxides and low volatility compounds. Estimates are uncertain for multifunctional radicals and are especially uncertain for radicals formed from reactions of cyclic compounds such as terpenes. (Secs. 4.6, A1.5.5)
Ring closure reactions of unsaturated peroxy radicals	Affects product formation. Current estimates are based on analysis of uncertain aromatic ring-opening reactions. Estimates need experimental or theoretical verification. (Sec. 0)
β-scission decompositions of alkoxy radicals	Affects amounts of fragmentation to lower molecular weight products. Data insufficient to derive reliable estimates for decompositions forming alkoxy radicals or effects of some substituents. Current estimates do not take into account chemical activation effects. (Secs. 5.2, A1.7.2)
Cyclization reactions of unsaturated alkoxy radicals	These reactions are not currently considered by MechGen, but recent theoretical and experimental data suggest that they may be non-negligible and need to be considered when MechGen is updated (Sec. 5.6). They may also affect aromatic ring opening mechanisms (Sec. 3.4).

Table 16 (continued)

<u>Uncertainty</u>	<u>Discussion</u> (Section discussed)
Other alkoxy radical reactions	Affects product formation. Uncertainty is variable, but even small estimation errors can affect product predictions. Chemical activation effects uncertain. (Secs. 5, A1.7.3, A1.7.4)
Reactions of Criegee intermediates	Affects radical initiation and product formation from alkenes. Stabilization and reactions of larger and substituted intermediates are particularly uncertain. Simplifications and arbitrary assumptions had to be used, and some of the current estimates are not consistent with recent data. (Secs. 7, A1.9)
Thermochemical estimates	Affects assessments of what reactions need to be considered and some estimates of radical reactions. Certain thermochemical group values used are extremely uncertain and possibly inappropriate. A complete review and update of the thermochemical estimates for mechanism development is required. (Sec. A2).
Appropriate cutoff rules for mechanism generation efficiency	Affects the size of the mechanisms that are generated and the number of products that have to be reacted in multi-generation mechanisms. Too high a cutoff will affect mechanism accuracy, and too low will affect mechanism efficiency and simulation times. (Secs 8; Carter, 2023b)

1786

1787 Vereecken et al. (2018) reviewed the current status and research needs for development of SARs
 1788 for detailed chemical mechanisms. In addition to discussing data needs and the importance of data
 1789 collection, they also discussed the need to evaluate and improve existing SARs, particularly with regard
 1790 to multifunctional compounds. Many of the SARs used in this work may not incorporate all the relevant
 1791 data and need to be updated or possibly replaced by newer versions. Work on SAR development and
 1792 improvement needs to be ongoing for chemical mechanisms to continue to represent the state of the
 1793 science and retain their link to laboratory data and theories.

1794 The predictive capabilities of atmospheric chemical mechanisms, whether detailed or condensed,
 1795 need to be evaluated for comparing their predictions against laboratory data. In some cases, MechGen had
 1796 to be modified so that developed mechanisms gave predictions of NO oxidation and O₃ formation
 1797 consistent with results of environmental chamber experiments. Consistency with O₃ formation in well-
 1798 characterized environmental chamber data needs to be a priority for mechanisms used for regulatory
 1799 modeling, though this is not a sufficient evaluation for detailed mechanisms. Comparison of experimental
 1800 vs. predicted product yields is necessary for a comprehensive evaluation, but this is a major effort with
 1801 many components and experimental difficulties. This would include comparisons with ambient data as
 1802 well as results of laboratory and environmental chamber studies. Doing this comprehensively is a multi-
 1803 year effort but it needs to be carried out.

1804 Many if not most of the areas of uncertainty in generated mechanisms may not be reduced
 1805 significantly in the near term. These uncertainties have variable effects on model predictions of interest,
 1806 with some highly uncertain or questionable estimates having almost no effects on predictions, and in
 1807 some cases only moderate uncertainties in rate constants having large effects. A systematic study of the
 1808 effects of the various types of uncertainties on predictions of interest is needed not only to prioritize areas
 1809 where basic research is needed, but also to prioritize which SARs and estimates used in systems like
 1810 MechGen should be given the highest priorities for future work.

1811 The research needs do not end once a detailed mechanism is developed and its predictive
 1812 capabilities have been evaluated. Mechanisms output by systems such as MechGen or GECKO-A are far
 1813 too large and complex for most modeling applications and need to be reduced to be useful. The reduction

1814 method will depend on the intended application, with, for example, detailed product predictions being a
1815 relatively low priority in O₃ models but a high priority in models for SOA and toxics. Further discussion
1816 of reduction issues is beyond the scope of this work, though it should be noted that MechGen as
1817 documented in this paper has been used to develop updated versions of the SAPRC mechanisms for use in
1818 airshed models for predicting O₃ and toxics, designated SAPRC-22 (Carter, 2023a). It can also be used as
1819 a basis for developing future versions of the mechanism for predicting SOA.

1820 **Author Contributions**

1821 This manuscript largely represents the body of work on chemical mechanism development led by
1822 W. P. L. Carter, and he is the main contributor to this publication. J. J. Orlando provided consultation on
1823 the methods, including rate constant and branching ratio estimates, and the resulting mechanism
1824 predictions. J. J. Orlando and K. C. Barsanti contributed to writing and editing, and J. Jiang contributed to
1825 editing and accuracy checking of the IUPAC references.

1826 **Competing Interests**

1827 Authors J. J. Orlando and K. C. Barsanti are members of the editorial board of Atmospheric
1828 Chemistry and Physics. The peer-review process was guided by an independent editor, and the authors
1829 have also no other competing interests to declare.

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1840 The opinions and conclusions in this paper are entirely those of the authors.

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