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2
3 **THEThe SAPRC Atmospheric Chemical Mechanism**
4 **GENERATION SYSTEM (MECHGEN)Generation System**
5 **(MechGen)**
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20 **ABSTRACT**

21 MechGen is a software system designed to derive gas-phase reaction mechanisms for reactive
22 organic compounds under atmospherically relevant conditions for use in chemical models and for data
23 analysis and interpretation. It has been used to derive versions of the SAPRC mechanisms used in airshed
24 models, with SAPRC-22 being the most recent. MechGen derives fully explicit mechanisms for many
25 types of organic compounds and their oxidation products when they react in the atmosphere in the
26 presence of oxides of nitrogen and other pollutants, and then uses these explicit mechanisms to derive
27 reduced or lumped mechanisms more suitable for use in airshed models. This paper gives an overview of
28 the system, describes the procedures it uses to generate explicit and reduced mechanisms, and presents
29 several types of applications. The assignments and estimates used to derive individual chemical reactions
30 and assign rate constants are discussed in a separate companion paper. The system is publicly accessible
31 for generating explicit mechanisms for single compounds and viewing associated documentation using a
32 web-based interface. A separate terminal login is available for deriving mechanisms for multiple
33 compounds, multi-generation mechanisms, and portions of lumped mechanisms for airshed models, as
34 well as for system programming and management. MechGen is designed to accommodate updates to the
35 chemical estimates and assignments that it uses to reflect our evolving knowledge of and ability to
36 estimate atmospheric reactions of organic compounds.

1. INTRODUCTION

1.1. Background

Many hundreds of ~~types of~~ volatile organic compounds (VOCs) are emitted into the lower atmosphere, from both anthropogenic and biogenic sources. Once emitted, they can undergo reactions to form oxidized organic products, including gas-phase toxics, criteria pollutants, and secondary organic aerosol (SOA). For example, in the presence of nitrogen oxides (NO_x), these reactions can generate radicals that react to form ozone (O_3) and oxidized nitrogen compounds that affect air quality. The atmospheric reaction mechanisms for most of these compounds are complex, particularly for larger molecules that involve a large number of reactive intermediates and form a large number of oxidized organic products that can continue to react in the atmosphere. In addition, in most cases these mechanisms involve reactions with rate constants that have not been measured and must be estimated. Because of the complexity, for practical reasons it is necessary either to greatly simplify the mechanisms for most VOCs, use extensive lumping or ~~condensations~~reduction in VOC representations, or use an automated chemical mechanism generation system to generate estimated mechanisms.

An overview of automated generation of reaction mechanisms was presented by Green (2019), which included references to systems that have been developed. A notable example is the RMG system of Green and co-workers (RMG, 2025), originally developed to model combustion systems (Gao et al., 2016) but later extended to apply to other areas such as liquid-phase systems and heterogeneous catalysis (Liu et al., 2021). RMG has an extensive set of tools, links to kinetics and other databases, and has extensive documentation (RMG, 2025). Kirchner (2005) discussed an atmospheric chemical mechanism generator program called CHEMATA, but that work focused on its application to condensed mechanisms, and literature searches revealed no subsequent work with that system. The most comprehensive systems specifically designed to predict detailed mechanisms for lower tropospheric air pollution models, are the Generator for Explicit Chemistry and Kinetics of Organics in the Atmosphere (GECKO-A) system (Aumont et al., 2005) and the SAPRC mechanism generation system (MechGen) that is the subject of this paper.

MechGen was developed for deriving the portions of the SAPRC-90 through SAPRC-16 atmospheric chemical mechanisms (Carter, 1990, 2000, 2010a, b, 2016; Carter and Heo, 2012, 2013; Venecek et al., 2018) that concerned reactions of C_{2+} organic compounds. The first version, used for SAPRC-90 (Carter, 1990), generated mechanisms only for alkanes, using the procedures and estimates documented by Carter and Atkinson (1985). When SAPRC-99 was being developed, the mechanism generation system was re-written and extended to cover a much wider range of acyclic and monocyclic compounds, including monoalkenes, alcohols, ethers, esters, aldehydes, ketones and organic nitrates in addition to alkanes (Carter, 2000). That version was used to determine the net effects of these compounds in the presence of NO_x that was incorporated into SAPRC-99, though reactions in the absence of NO_x were not generated. A number of updates to the system were made when SAPRC-99 was updated to SAPRC-07 (Carter, 2010a, b), including the ability to generate mechanisms for a wider variety of compounds; however, it remained limited to generation of mechanisms in the presence of NO_x , and aromatics were still not supported. The system was further updated for use in the development of the SAPRC-16 (Carter, 2016; Venecek et al., 2018) mechanism and unpublished updated versions, incorporating capabilities to generate reactions in the absence of NO_x , reactions of aromatics, autoxidation reactions of peroxy radicals, and other enhancements. Further updates were made for the current version, which has been used in the latest of the SAPRC mechanisms, SAPRC-22 (Carter, 2023).

The chemical basis of MechGen and the methods it uses to assign or estimate rate constants or mechanisms are documented by Carter et al. (2025a). The current, and first publicly documented, version of MechGen, v1.0, v1.1, is available online, along with a comprehensive User Manual and a quick start guide. These are available at the MechGen website (Carter, 2025) and GitHub page (<https://github.com/SAPRC/MechGen>), and archived versions can also be downloaded from these websites. This paper describes the software system and the procedures it uses for mechanism generation and processing.

1.2. Chemical Systems Represented

MechGen is capable of generating fully explicit mechanisms for the atmospheric reactions of most types of organic compounds emitted into or formed in the lower atmosphere, and of the intermediate radicals they form. While temperature-dependent rate constants are provided for many reactions, for others the rate constants or branching ratios are only applicable for conditions representing the troposphere, i.e., around 298 K and 1 atm. MechGen is not currently designed for estimating mechanisms for combustion modeling or for the low temperature and pressures characteristic of the upper troposphere.

Table 1 lists the types of stable compounds whose reactions can be generated, and shows the types of initial atmospheric reactions that can be generated for them. Table 2 lists the types of reactions that are generated, including reactions of intermediate radicals as well as reactions of stable compounds. The reactions of stable compounds include H-atom abstractions by OH, NO₃, and Cl radicals, additions to double bonds by these radicals and by O₃ and O³P, and reactions by photolysis. The types of radicals generated include carbon-centered radicals that in most cases react primarily with O₂; peroxy radicals that in most cases react with NO, NO₂, NO₃, HO₂, or other peroxy radicals and in many cases also have unimolecular reactions; alkoxy radicals that can react with O₂ or by various types of unimolecular reactions; and excited and stabilized Criegee intermediates formed in reactions of alkenes with O₃. Measured rate constants or branching ratios derived from measured product yields are used when data are available. In the absence of such data, which is typically the case, rate constants or branching ratios are estimated using various structure-activity relationships (SARs) or other estimation methods. The details of the types of reactions and how rate constants or branching ratios are assigned or estimated are given by Carter et al. (2025a).

1.3. System Software and Access

The current version of MechGen is incorporated into an online Multi-User Object Oriented (MOO) system that was originally developed as a programmable text-based virtual reality system (Curtis, 1997; Fox, 2004; Wikipedia, 2025). Features of the MOO object-oriented programming language, which is similar to Python, made it well suited for mechanism generation applications compared to the original Fortran version used for SAPRC-90 (Carter and Atkinson, 1985; Carter, 1990). The MOO system also has online access capabilities that make it relatively straightforward to permit multiple users to access it online simultaneously. ~~While MOO is no longer widely used or maintained, MechGen could be migrated to Python or another actively supported software platform, allowing ongoing maintenance and collaborative development in the future.~~

Since MechGen is written in an object-oriented language, the system employs various software "objects" that include properties defining their characteristics and programs or subroutines controlling their operations. The MechGen software objects referenced in this paper include reactants objects that represent molecules or radicals that can be reacted; group objects that represent portions of the molecule that determine how they react; reactor objects that are assigned to each user login that provides a user interface and gives mechanism generation options; environment objects that contain concentrations of

atmospheric species that can be used to determine product yields from mechanisms; multi-generation mechanism objects that control multi-generation mechanism derivations for a given reactant; and lumping objects that control the full mechanism generation process and whether and how lumped mechanisms are also produced. The MechGen system employs other types of objects, but a discussion of these details of the software system is beyond the scope of this paper.

MechGen is accessible online via a web interface available at the MechGen website (Carter, 2025) or through a terminal interface such as Telnet (Postel and Reynolds, 1983), with instructions available at the MechGen and MechGen GitHub sites. Terminal access provides the highest degree of system capability, while web access is more user-friendly for basic operations and is better suited for providing information about the system and the reactions being generated. Due to the high computational demand, resource-intensive operations such as deriving multi-generation mechanisms, or generating mechanisms for large or for large numbers of molecules, are not currently available using the online system. To access the full capabilities of MechGen, users need to install the system locally by following the instructions provided in the User Manual.

1.4. Overview of Documentation

The methods used by the current version of MechGen to derive or estimate rate constants and mechanisms for gas-phase atmospheric reactions of organic are described by Carter et al. (2025a). While that paper describes the chemical basis for the mechanisms MechGen generates, it does not explain the system itself or its functionality. This paper fills that gap by discussing the following specific capabilities of MechGen:

- Deriving unimolecular or bimolecular reactions of individual compounds and radicals under atmospheric conditions, using assigned or estimated rate constants and products formed (Section 2);
- Deriving explicit single-generation mechanisms of a selected compound by reacting the compound and all the rapidly reacting intermediates formed, but not reacting the stable, non-radical products (Section 3.1);
- Deriving minimally reduced mechanisms that give the same predictions as the explicit mechanism at a selected temperature, but with about 1/3 the numbers of reactions and 1/2 the numbers of species (Section 3.2);
- Estimating yields of products formed when reacting a compound under selected environmental conditions (Section 3.3);
- Deriving explicit multi-generation mechanisms of a selected compound by fully reacting the compound and all stable reactive products formed in non-negligible yields (Section 4); and
- Optionally driving lumped mechanisms by using that use a limited number of lumped model species to represent chemically similar compounds, and by using various approaches to reduce the numbers of reactive intermediates in the mechanisms (Section 5).

Additional details about the methods and algorithms are also given in the Supplement. Details concerning how to work with MechGen and a description of features not covered here are given in the User Manual that is available at the MechGen website (Carter, 2025). The MechGen GitHub site (<https://github.com/SAPRC/MechGen>) also contains the User Manual and downloadable software and files used by MechGen (see Section 7.3). The examples shown here and the Supplement are based on the current chemistry assignments and methods discussed by Carter et al. (2025a). The system is expected to operate similarly when the chemistry assignments and estimates are updated in the future.

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

Type of Compound	Reactions [a]
Alkanes	OH, NO ₃
Alkenes (including multiple double bonds)	OH, O₃, NO₃, O³P
<u>Alkenes (including those with multiple double bonds)</u>	<u>OH, O₃, NO₃, O³P</u>
Alkynes	OH, NO ₃
Aromatic hydrocarbons	OH
Aldehydes, ketones, hydro-peroxides, organic nitrates	OH, NO₃, hv
<u>Aldehydes, ketones, hydroperoxides, organic nitrates</u>	<u>OH, NO₃, hv</u>
Alcohols, ethers, esters, other oxygenates	OH, NO ₃
Unsaturated aldehydes, ketones, hydroperoxides, and nitrates	OH, O ₃ , NO ₃ , O ³ P, hv
Phenols	OH, NO ₃
Furans and other unsaturated oxygenates	OH, O ₃ , NO ₃ , O ³ P
Amines	OH, NO ₃
Peroxynitrates	Unimolecular, OH, NO ₃ , hv
Bi- and polyfunctional compounds	OH, O ₃ , NO ₃ , O ³ P, hv, as applicable

168 [a] See Carter et al. (2025^a) for more detail and types of compounds not supported.
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172 Table 2. Summary of types of reactions supported by MechGen.

Reactant(s) [a]	Type of reactions [b]
VOC + OH	H-atom abstraction. Addition to double, aromatic, and triple bonds
VOC + O₃	Addition to double bonds and N atoms in amines
<u>VOC + O₃</u>	<u>Addition to double bonds and to N atoms in amines</u>
VOC + NO ₃	H-atom abstraction. Addition to double bonds
VOC + O ³ P	Addition to double bonds
VOC + hv	Usually breaking the weakest bond, but depends on the compound
VOC Uni.	Unimolecular decompositions of peroxy nitrates
Carbon-centered Radicals	O ₂ addition. H-abstraction from α-OH groups, unimolecular reactions of excited OH adducts, unimolecular decompositions
Peroxy or Acyl peroxy Radicals	Reactions with NO forming the corresponding alkoxy radical and organic nitrate; reactions with NO ₂ , NO ₃ , HO ₂ or other peroxy radicals; and unimolecular reactions
Alkoxy Radicals	α-H abstraction by O ₂ forming the corresponding carbonyl compound. β-scission and H-elimination decompositions. H-shift and ester rearrangement isomerizations
Criegee biradicals	Decompositions, stabilization, or rearrangements.
<u>Criegee intermediates</u>	<u>Decompositions, stabilization, or rearrangements.</u>

173 [a] "VOC" indicates any stable volatile organic compound supported by the system.

174 [b] See Carter et al. (2025^a) for more detail concerning types of reactions.

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2. REACTANT SPECIFICATION AND SINGLE-STEP REACTIONS

2.1. Creation and Specification of Reactants

Reactants are created as objects within MechGen by specifying their structures. The structure of an organic reactant or radical is specified by giving the "groups" in the molecule or radical, and indicating the groups each are bonded to, and the type of bond. As listed in Table 3, groups are parts of molecules that are treated as units in the system, and contain no more than one carbon, nitrogen, or halogen atom, but can also contain one or several hydrogen or oxygen atoms. The leading or trailing "-", "=", "#", or "-a" indicate groups that can bond to neighboring groups by single, double, triple bonds, or aromatic/allylic bonds. Any groups with such designations can bond to other groups with such designations to form chains. Mechanisms are generated and rate constants are estimated based on the groups present, the groups they are bonded to, and in some cases on groups elsewhere in the molecule. Groups can be radical or non-radical, with reactants containing a radical group referred to as radicals or intermediates. Examples of reactant designations are given in Table 4, and additional discussion on reactant designation and creation by specifying structures is available in the User Manual.

MechGen also includes objects representing elementary species that can react with reactants (e.g., OH or O₃), react with radicals (e.g., O₂ or NO), or be formed in reactions (e.g., H₂O or CH₄). These are listed in Table 5.

2.2. Single Step Reactions

Single step mode consists of generating only the initial reaction(s) of a specified compound or intermediate, without reacting the resulting products or intermediates. When reacting stable compounds, it is necessary to specify the type of reaction (e.g., whether unimolecular or a bimolecular reaction with a specified oxidant), whereas all possible reactions are automatically generated for radicals. Generating reactions in single step mode is useful to obtain information on how the system estimates a compound's reactions, since the results show not only the products formed and the rate constants or branching ratios derived, but also documentation on the estimates or assignments used during reaction generation.

The algorithm for generating single step reactions is shown in Figure 1, and examples of output are shown in Figure S1 in the Supplement. Note that Figure S1 shows not only the generated reactions but also text documenting how the reactions were derived. Reaction types and derivation methods are discussed by Carter et al. (2025a), and the specific procedures shown in Figure 1 depend on whether the reactant is a stable compound or an intermediate, and also on the type of intermediate. Assigned reactions and rate constants are used for reactants for which assignments are made, but for most reactants, reactions and rate constants ~~are~~ have to be estimated using methods discussed by Carter et al. (2025a). In some cases, assignments are made for rate constants but not for mechanisms or branching ratios; in those cases, the reactions are generated using the general estimation methods, but the rate constants output are the assigned rather than the estimated values.

Some radicals are estimated to undergo unimolecular or O₂ reactions so rapidly that it is not necessary to estimate their total rate constants to predict their atmospheric fates. For example, most alkyl radicals are consumed only by reaction with O₂, and many "explicit" mechanisms simply replace them with the peroxy radicals they form, resulting in these alkyl radicals not appearing in the generated mechanism. MechGen does include these radicals and their reactions, though it flags them as being "fast" and only outputs their branching ratios, when applicable.

For photolysis reactions, the system outputs a "photolysis set" name that gives absorption cross sections and optionally quantum yields as a function of wavelength, and also a wavelength-independent quantum yield if the quantum yields are not given in the photolysis set. The photolysis sets referenced by MechGen are those used for the SAPRC-22 mechanism as listed and documented by Carter et al. (2025a), and these files can be downloaded with other files needed to implement that mechanism (Carter 2024). Single step reaction output also includes the calculated photolysis frequency for a representative solar light intensity and spectrum, but this should not be considered part of the mechanism because it depends on the light environment.

In the case of peroxy radicals, the system first derives unimolecular reactions if they are estimated to be non-negligible for this radical, then derives the bimolecular reactions with NO, NO₂, NO₃, HO₂, generic alkyl peroxy radicals (RO₂), and generic acyl (RCO₃) peroxy radicals in that order. Deriving all the possible organic peroxy + peroxy reactions is impractical due to the large number of peroxy radicals formed in realistic photooxidation systems and the fact that the other peroxy radicals present are unknown at the time of mechanism generation. Instead, MechGen treats peroxy radicals as reacting with the totals of all alkyl and all acyl peroxy radicals. The products from each reaction type include those formed from the subject radical, plus counter species used to represent the type of co-products formed from the sum of peroxy radicals it reacts with. Those counter species can be either generic alkoxy radicals, designated "RO." or "RCO2" [representing RC(O)O $\cdot$]; or carbonyl or alcohol H-transfer products, designated "RO-alpha-H" and "ROH" or "RCO-OH" (Carter et al., 2025a). They are included as products of these reactions during reaction generation for tracking if desired, but are deleted as products when the mechanisms are reduced or lumped.

The user-modifiable parameters that affect the single step reaction generation process are shown in Table 6. The temperature affects the thermal rate constants. The pressure affects rate constants assigned to falloff reactions and also estimated nitrate yields in the reactions of peroxy radicals with NO, and in some cases, fractions of excited radicals that are stabilized. The presence of water affects estimated reactions of low reactivity stabilized Criegee intermediates, as indicated in Figure 1 (Carter et al., 2025a). When users select single step operations, all estimated reactions are displayed, allowing users to see which reactions are considered and how their rate constants are estimated. However, negligible reactions are removed when the outputs of single step reaction operations are processed when called during full mechanism generation, as discussed in ~~the next section~~ [Section 3](#). In addition, reactions of non-acyl peroxy radicals with NO₂ forming alkyl peroxy nitrates (shown in Figure S1c) are not included during full mechanism generation by default because these peroxy nitrates are assumed to rapidly decompose at temperatures of interest (Carter et al., 2025a). These reactions can be included by either increasing the kFastUni parameter (see Table 8), or decreasing the reactor temperature such that the estimated rate constant is less than kFastUni.

2.3. User Modifiable Mechanism Assignments

As indicated above, MechGen uses assigned rate constants and branching ratios when deriving single-step mechanisms whenever possible, and otherwise uses estimates and SARs when no assignments are available. While updates will be made as new data become available, the assignments will not always be up to date. In addition, some users may wish to employ MechGen to derive mechanisms using recently measured or newly derived rate constants and branching ratios, or to explore how varying uncertain estimates impact mechanisms. To address this, MechGen allows users to add custom (or "user") mechanism assignments, which can supplement or replace the default settings. The steps involved in making or managing user assignments are described in the User Manual.

Table 3. List of groups and group designations used to specify C₂₊ organic reactants whose reactions can be generated using MechGen.

Type	Group designations [a]			
	<u>Non-Radical Groups</u>			
Alkane	-CH3	-CH2-	-CH()-	-C()-
Alkene	=CH2	=CH-	=C()-	=C=
Alkyne	#CH	#C-		
Aromatic or Allylic [b]	-aCH-	-aC()-		
Oxygenate	-CHO	-CO-	-OH	-O-
Nitrate, nitro, nitroso	-ONO2	-NO2	-NO	
Amino and amine-oxy	-NH2	-NH-	-N()-	-N[O]()-
Imine	=NH	=N-		
Criegee Intermediates	-CHOO[syn]	-CHOO[anti]	-syn-C[OO]-	-C[OO]-syn-
<u>Criegee intermediates</u>	<u>CH2OO-</u>			
	<u>-C[OO]-syn-</u>	<u>-CHOO[syn]-</u>	<u>-CHOO[anti]</u>	<u>-syn-C[OO]--</u>
Halogen	-F	-Cl	-Br	-I
	<u>Radical or Reactive Intermediate Groups</u>			
Carbon Centered radicals	-CH2.	-CH[.]	-C[.]()	=CH.
<u>Carbon centered radicals</u>	<u>-CH2.</u>	<u>-CH[.]</u>	<u>-C[.]()</u>	<u>=CH.</u>
	=C[.]	-pC[.]	-CO.	
Allylic radicals [b]	-aCH2.	-aCH[.]	-aC[.]()	
Peroxy radicals	-CH2OO.	-CH[OO.]	-C[OO.]()	
	-CO[OO.]	=CHOO.	=C[OO.]	-pC[OO.]
Alkoxy radicals	-CH2O.	-CH[O.]	-C[O.]()	=CH[O.]
	=C[O.]	-CO2.	-pC[O.]	
Criegee Intermediates [c]	-CHOO {excited}	-C[OO] {excited}		
<u>Criegee intermediates [c]</u>	<u>-CHOO {excited}</u>	<u>-C[OO] {excited}</u>		
Radicals from amines	-NH.	-N[.]	-NH2[O]	-NH[O]
Carbenes	-CH[..]	-C[..]		

[a] Bond types are indicated by "-" (single or aromatic), "=" (double), or "#" (triple). The number of bonds shown indicate if it is bonded to one or two or more groups. The "()" notation is not part of the group designation itself, but is used to indicate that the group is bonded to a 3rd or 4th group. An "a" or "p" prefix in the group name indicates that it has alternating single and double bonds. The "p" prefix is used for phenyl, phenoxy or phenyl peroxy radicals. Some groups can have different bond designations depending on their location in the molecule, e.g., "CH3-" or "-C[.]=" rather than "-CH3" and "=C[.]". See examples in Table 4.

[b] Allylic groups are used in reactants with two or more resonance structures involving adjacent double bonds and radical centers. Allylic radical groups are used for the portions where that the radical center is located in at least some of the resonance structures. Aromatic groups are used for the portions where the radical center is not located in any resonance structures.

275 [c] The "{excited}" designation is given at the end of the structure designation and indicates how the
276 excited intermediate was formed. This designation is also used following full reactant structure
277 designations to indicate excited adducts formed when radicals add to double bonds, but in this case
278 the excitation is associated with the entire molecule and not a single group.
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282 Table 4. Examples of designations of selected representative compounds or intermediates.

Compound	MechGen Structures [a]
<u>Stable Compounds</u>	
propene	CH ₂ =CH-CH ₃ ; CH ₃ -CH=CH ₂
methyl acetylene	CH#C-CH ₃ ; CH ₃ -C#CH
methyl acetylene	CH#C-CH₃; CH₃-C#CH
2-methyl-2-ethyl butane	CH₃-CH₂-C(CH₃)(CH₃)-CH₂-CH₃; CH₃-C(CH₃)(CH₂-CH₃)-CH₂-CH₃
3,3-dimethyl pentane	CH₃-CH₂-C(CH₃)(CH₃)-CH₂-CH₃; CH₃-C(CH₃)(CH₂-CH₃)-CH₂-CH₃
trans-3-methyl-2-pentene	CH₃-^CH=C(vCH₃)-CH₂-CH₃; CH₃-CH₂-C(^CH₃)-CH-^CH₃
trans-3-methyl-2-pentene	CH₃-^CH=C(vCH₃)-CH₂-CH₃; CH₃-CH₂-C(^CH₃)=CH-^CH₃ [b]
cyclopropane	CH ₂ *-CH ₂ -CH ₂ *; *CH ₂ -CH ₂ -*CH ₂
<u>cyclopropane</u>	<u>CH₂*-CH₂-CH₂*; *CH₂-CH₂-*CH₂ [c]</u>
bicyclo [1.1.1] heptane	CH ₂ *1-CH*2-CH ₂ -CH*1-CH ₂ *2; CH*12-CH ₂ -CH(CH ₂ *1)-CH ₂ *2
toluene	CH₃-aC*-aCH-aCH-aCH-aCH-aCH*; CH₃-C*=CH-CH=CH-CH=CH*
<u>bicyclo [1.1.1] heptane</u>	<u>CH₂*1-CH*2-CH₂-CH*1-CH₂*2; CH*12-CH₂-CH(CH₂*1)-CH₂*2 [c]</u>
<u>toluene</u>	<u>CH₃-aC*-aCH-aCH-aCH-aCH-aCH* [d]; CH₃-C*=CH-CH=CH-CH=CH*</u>
naphthalene	aC*12-aCH-aCH-aCH-aCH-aC*1-aCH-aCH-aCH-aCH*2
2-propyl nitrate	CH ₃ -CH(CH ₃)-ONO ₂ ; CH ₃ -CH(ONO ₂)-CH ₃
carbitol acetate	CH ₃ -CH ₂ -O-CH ₂ -CH ₂ -O-CH ₂ -CH ₂ -O-CO-CH ₃
<u>Intermediates</u>	
2-propyl radicals	CH₃-CH[.] -CH₃
<u>2-propyl radical</u>	<u>CH₃-CH[.] -CH₃</u>
2-propyl peroxy radicals	CH₃-CH[OO.] -CH₃
<u>2-propyl peroxy radical</u>	<u>CH₃-CH[OO.] -CH₃</u>
methyl allyl radicals	CH₃-aCH[.] -aCH-aCH₂; CH₃-CH[.] -CH=CH₂; CH₃-CH=CH-CH₂.
<u>methyl allyl radical</u>	<u>CH₃-aCH[.] -aCH-aCH₂ [d]; CH₃-CH[.] -CH=CH₂; CH₃-CH=CH-CH₂.</u>
OH + benzene adduct	HO-CH*-aCH[.] -aCH-aCH[.] -aCH-aCH[.]*
Excited MEK oxide from O₃ + alkene	CH₃-CH₂-C[OO]-CH₃ {*O₃Ole}; CH₃-C[OO]-CH₂-CH₃ {*O₃Ole}
<u>Excited Criegee int. from O₃ + alkene</u>	<u>CH₃-CH₂-C[OO]-CH₃ {*O₃Ole}; CH₃-C[OO]-CH₂-CH₃ {*O₃Ole}</u>
Stabilized MEK oxide with syn-CH₃	CH₃-syn-C[OO]-CH₂-CH₃; CH₃-CH₂-C[OO]-syn-CH₃

Stabilized Criegee int. with *syn* -CH₃ CH3-*syn*-C[OO]-CH2-CH3; CH3-CH2-C[OO]-*syn*-CH3

[a] The first MechGen structure codes given are those generated by the system. If subsequent structures are given, they are alternatives that can be used to create the same compound.

[b] The symbols "v" and "^" are used to indicate "cis" or "trans" configurations, analogous to "\" and "/" in Smiles notation.

[c] The symbol "*" is used to indicate ring closure. "*1", "*2", etc used for multiple ring structures, analogous to "1", "2", etc. in Smiles notation.

[d] The symbol "aC" used to indicate carbon centers where with single and double bonds are in resonance, as in aromatic species or allylic radicals.

Table 5. List of elementary species used by MechGen.

Type	Group designation				
Non-reacting products [a]	H2 HCHO	H2O NH3	HONO HCl	HNO3	CH4
Species that react with VOCs	OH Cl.	O3 Uni [b]	NO3	O3P	hv [b]
Species that react with radicals	O2	NO	NO2	RO2. [c]	RCO3. [c]
Counter species [c]	RO.	ROH	RCO2.	RCO-OH	RO-alpha-H
<u>Counter species [c]</u>	<u>RO.</u>	<u>ROH</u>	<u>RCO2.</u>	<u>RCO-OH</u>	<u>RO-alpha-H [d]</u>

[a] These species are treated as unreactive by MechGen.

[b] Pseudo-species used to indicate photolysis or unimolecular reactions of stable compounds.

[c] Generic species used as reactants (RO2., RCO3.) or products (RO., etc.) in peroxy + peroxy [reactions](#).

~~reactions. Note that~~[\[d\]](#) "RO-alpha-H" refers to the carbonyl formed when an α -hydrogen on an alkoxy radical is abstracted.

Table 6. List of parameters that affect single-step mechanism generation.

Parameter (code)	Affects	Default	Note
Temperature (T)	Thermal rate constants and O ₂ concentration	298K	
Pressure (P)	Pressure-dependent rate constants and O ₂ level	1 atm	
O ₂ fraction (O2)	Percentage of O ₂ in reacting environment	20.95%	[a]
Light Source (light)	Actinic flux values used to calculate rate constants for photolysis reactions in output displays	As used by Carter (1994)	[b]
Water present (H2O)	Parameter indicating whether water is present.	H ₂ O is present	[c]

[a] Affects output displays and estimates of product yields, but not reactions in explicit generated mechanisms. However, this affects the full mechanism generation process discussed in Section 3.

[b] The actinic fluxes are used to calculate rates of photolysis reactions given the absorption cross sections and quantum yields assigned to the photolysis reactions. The photolysis frequencies are not part of the mechanism but are included as outputs of single step mechanism generation and are also used to derive overall product yields for various environments, as discussed in Section 3.3 and to derive multi-generation mechanisms as discussed in Section 4.

[c] Currently applicable only to low reactivity Criegee intermediates (Carter et al., 2025[a](#)).

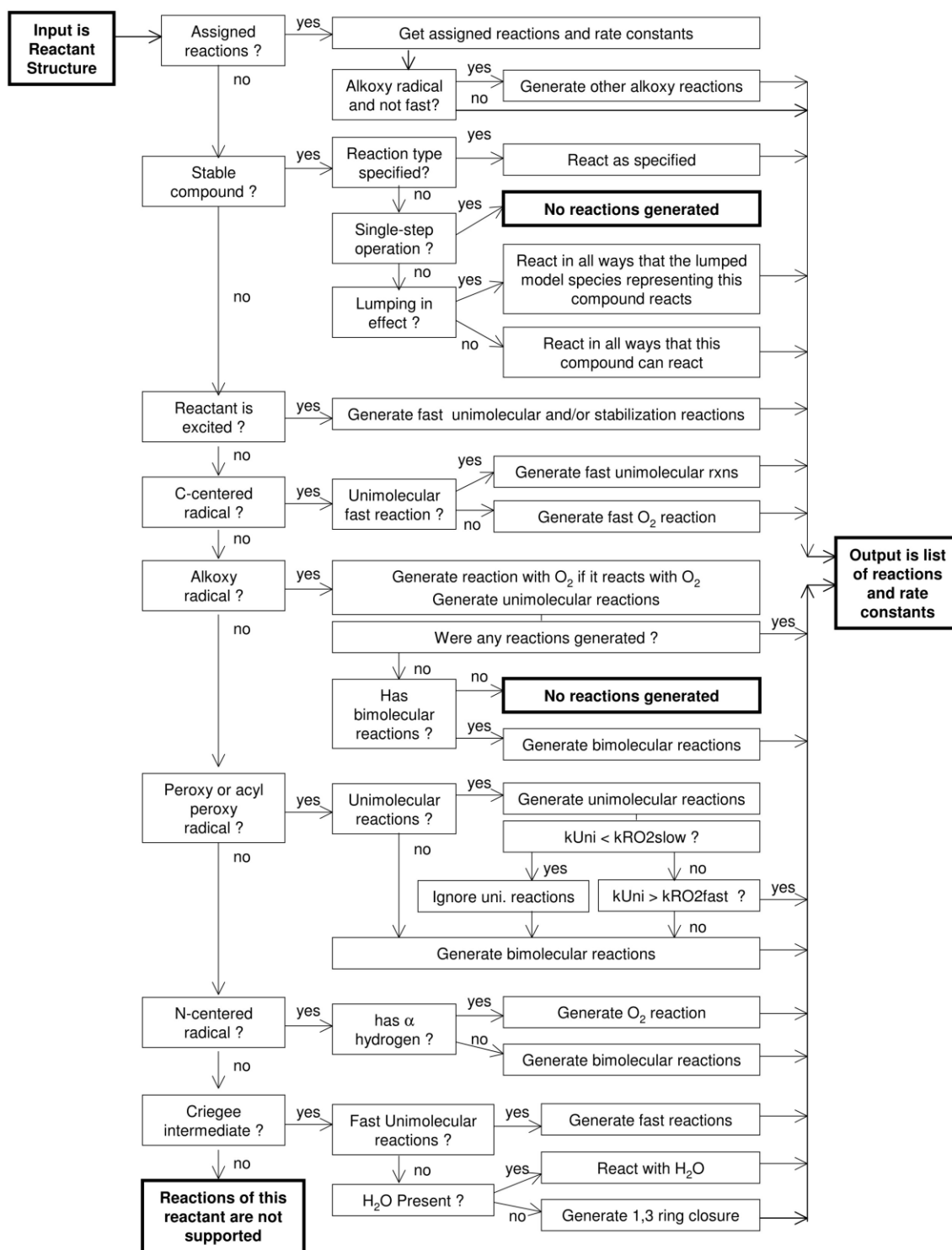


Figure 1. Diagram of algorithm used to generate single-step reactions.

3. FULL MECHANISM DERIVATION

3.1. Description of Process

Full mechanism derivation for a compound involves first reacting it through all, or a designated subset, of its possible initial atmospheric reactions; and then reacting any reactive intermediates that are formed. Stable products formed are not reacted, so the mechanism output reflects only a single generation of reactions, as opposed to multi-generation mechanism derivations discussed in Section 4. Reactive intermediates are defined as products that have radical or intermediate groups (see Table 3) or non-radical compounds that undergo unimolecular reactions with rate constants greater than ~~0.0167 sec⁻¹ (0.7 min half life)~~a specified value, which is 0.0167 s⁻¹ (1 min⁻¹) by default. The result is a sequence of reactions that were generated and their rate parameters, lists of intermediates involved and products formed, and information about the products and intermediates that may be useful.

In order to provide the capability to generate explicit mechanisms of appropriate sizes for specific modeling applications, and to estimate relative yields of products formed under various conditions, MechGen provides standard environments that specify concentrations of atmospheric species that can be used for these purposes. The currently available standard environments are summarized in Table 7, with additional information given in Section S2 of the Supplement.

The currently available standard environments listed in Table 7 all represent urban conditions, but users can define new environments as described in the User Manual to include a wider range of conditions. The default use of these environments is indicated in Table 7, with three environments representing the range of urban conditions with regard to NO_x availability, and an additional environment representing nighttime conditions. By default, the "Standard" or mid-NO_x environment is not used for mechanism generation, as reactions important under those conditions would also be important under either high or low NO_x conditions, if not both. The mid NO_x environment is useful, however, for determination of relative product yields under urban conditions that are equally sensitive to both VOC and NO_x. It is also the default environment used when deriving multi-generation mechanisms (see Section 4).

The user-modifiable options controlling full mechanism derivation are summarized in Table 8, and the algorithm employed is shown in Scheme S1 of the Supplement, which outlines the overall process, and Scheme S2, which details the portion that determines which reactions are deleted. A key option affecting the sizes of generated mechanisms is the "MinYld" parameter, which determines which competing reactions in the single-step reactions can be neglected. A related option concerns whether and which environments are used during mechanism generation to estimate upper limit yields for competing bimolecular reactions. The "MinYld" test is implemented by assigning an "upper limit yield" for the initial reactant as unity, and estimating upper limit yields of intermediates whose reactions are generated as the product of the assigned or estimated upper limit yield of the reactant forming it, multiplied by the upper limit relative yield of the intermediate in the reactions of the reactant forming it. Upper limit yields are also derived for stable products formed in the generated mechanisms, for the purpose of estimating their approximate importance in the mechanism once it is generated. Reactions forming products with estimated upper yields less than MinYld are neglected, unless there is only one reaction of the reactant. When reactions are neglected, the rate constants and yields of the competing reactions are adjusted to ensure no mass or moles are lost (see Scheme S2). Upper limit yields are also derived for non-reacting products, but this is used for information purposes only.

The relative yields of products from reactions of the initial reactant or reacting intermediate can depend on the environment. MechGen assumes constant temperature, pressure, and O₂ levels when doing full mechanism generations. As a result, relative yields of products from reactants that undergo only unimolecular reaction or with O₂ are treated as environment independent. However, environments significantly affect yields of products from reactants that undergo other bimolecular reactions. In such cases, either a single environment must be specified to calculate yields for the MinYld test, or upper limit relative yields must be estimated. If no environment is specified, upper limit yields are derived by assuming that each type of bimolecular reaction is equally important. If more than one environment is used for mechanism generation, relative yields are determined for each environment, and the highest yield among these environments is then used to derive upper limit yields for the MinYld test. Note that the total of upper limit yields for competing reactions can exceed 100% unless only one environment is used.

As shown in Scheme S2, the MinYld test is not used for reactions of peroxy radicals with NO and for photolysis reactions of initial reactants. The only reactions with NO that may be generated are reactions of peroxy radicals forming alkoxy radicals or organic nitrates. Since the formation of alkoxy radicals is always relatively more important, neglecting the nitrate-forming reaction using MinYld test will result in underestimation of radical and NO_x sink processes without significantly reducing mechanism size. Similarly, the MinYld test is not used for photolysis reactions because MechGen does not generate many competing such reactions and deleting any of them may affect predictions of radical sources without significantly reducing the size of the mechanism. Generally, it is unimolecular reactions that tend to have many competing processes where some may be negligible according to the MinYld test.

For a given MinYld value, assuming no environments gives the largest mechanisms that are applicable to a full range of conditions, while assuming a single environment gives the smallest mechanisms that are suitable for that environment, but they may not be as accurate when applied to other conditions. Using an appropriate set of environments produces mechanisms of intermediate size that may be optimum if the environments used represent the range of conditions where the mechanism may be used. As indicated in Table 8, the default is to use three environments that represent the range of urban conditions. Use of additional environments may be appropriate if the mechanisms are to be used for more remote scenarios.

An example of a mechanism derived using a full mechanism generation process is shown in Figure S2 of the [S](#)upplement, with a representative subset displayed in Figure [3-2](#). These figures show reactions generated for 1,3-butadiene under default conditions. Note that designations such as "{*OHadd}" or "{*O3ole}" indicate excited intermediates. This output includes the rate constants at the default temperature of 298 K ("k"), the branching ratio for various competing reactions ("Fac"), the estimated upper limit weighting factor to determine which reactions can be neglected ("Weight"), and the reactants and products involved of the reactions. If a rate constant is not listed, the reaction is assumed to be fast and the reactant to be in steady state, making the fate of the reactant independent of the rate constant. In such cases, the branching ratios for competing fast routes are given in the "Fac" column.

The counter species such as "RCO2." and "RCO-OH", shown ~~in in Figure 3, and others in~~ Figures S1 and S2, are used for reactions of peroxy radicals with other peroxy radicals, as indicated in Table 5. The output shown in ~~Figure 3 and Figure~~ S2 ~~does~~ not include the assigned or estimated Arrhenius parameters for calculating rate constants at different temperatures for some reactions, but this information can be obtained by selecting other output formats for the reactions as discussed in the User Manual.

The numbers of reacting intermediates, stable products, and reactions for mechanisms derived for representative compounds using various mechanism generation options are discussed in Section 3.4 and ratios of these are shown in Figure S7 in the Supplement. In most cases, the ratios of the numbers of

reacting intermediates in the explicit mechanisms are about 30-50% the numbers of reactions, tending to be smaller when no environments or lower MinYld values are used, with no clear dependence on mechanism sizes. The ratios appear to be higher for the representative aromatics than for other types of compounds, at least for these examples. The ratios of numbers of stable products to numbers of reactions tend to be in the 30-40% range and are less variable than the ratios for intermediates.

3.2. Derivation of Minimally Reduced Mechanisms

The numbers of reactions and intermediates in generated mechanisms can be significantly reduced by combining parallel reactions of the same reactants and by eliminating intermediates that always rapidly form the same product(s) regardless of the environment. Combining parallel reactions gives product yields derived from the ratios of the rate constants for individual reactions to the total rate constant, and reduces the numbers of reactions, but does not affect the numbers of species in the mechanism. The numbers of predicted stable products cannot be reduced without further lumping, but many reacting intermediates whose fates do not depend on environmental conditions can be removed by assuming steady state so that they can be replaced by the products they form. This results in reducing the total number of intermediates by factors of 3 or higher (see below).

The intermediates removed by this process include species that undergo only unimolecular reactions or reactions with O₂, H₂O, or by stabilization, meaning their products do not depend on the environment if O₂, temperature, [water content](#), and pressure are assumed constant. These include all carbon-centered radicals, all excited intermediates, almost all alkoxy and nitrogen-centered radicals, and many Criegee intermediates. These all react fast enough on the time scale involved with atmospheric modeling that they can be assumed not to build up in concentration, so replacing them by their products should not significantly affect model predictions.

However, this process does not eliminate peroxy or acyl peroxy radicals that do not undergo fast unimolecular reactions, since they are consumed primarily or at least significantly by bimolecular reactions involving NO_x and other peroxy species, whose relative concentrations depend on the reacting environment. Nevertheless, as discussed below, this results in the removal of most reactive intermediates in the explicit generated mechanisms. -

Vereecken and Nozière (2020) calculated that most hydroperoxy-substituted peroxy radicals should interconvert to other such radicals with relatively high rate constants, and these are incorporated into the estimated mechanisms produced by MechGen (Carter et al., 2025a). These interconversion reactions are not considered when estimating whether competing reactions may be negligible because they are not net sink processes for such radicals. However, including these rapid interconversion reactions in the processed or lumped mechanisms used in models can potentially cause numerical "stiffness" problems in model simulations and also needs to be taken into account when product yields are estimated, as discussed in Section 3.3.

Therefore, as part of this initial processing, rapidly interconverting hydroperoxy-peroxy radicals are identified and represented in the processed mechanism by a lumped radical species representing the sets of interconverting radicals that are assumed to be in equilibrium. Reactions forming these radicals are represented as forming this lumped radical species, and reactions of these radicals are replaced by reactions of this lumped species, forming products of the interconverting radicals with yields multiplied by the equilibrium fraction of the radical derived from the rate constants involved. This process is discussed in Section S3.1 of the Supplement. This removes these rapid interconversion reactions from the mechanism, while retaining their effects on overall product yields.

Mechanisms processed this way are referred as "minimally reduced" or "processed" mechanisms. They should give essentially the same predictions in models as the explicit generated mechanisms as long as the temperature and O₂ levels are constant at the defaults set for the reactor when the mechanism was generated (see Table 6), and as long as the steady state approximation is appropriate for the reacting intermediates removed during the minimal reduction process. This has been verified in test calculations using representative chemical systems.

MechGen automatically derives a minimally processed mechanism after any successful full mechanism generation process. These are used as the input when deriving product yields, deriving multi-generation mechanisms, or when deriving lumped mechanisms using various approaches, as discussed in the following sections. The algorithm employed to derive minimally reduced mechanisms from the explicit generated mechanisms is outlined in Section S3.2 of the Supplement.

The minimal reduction process tends to reduce the numbers of reactions by factors of 2 - 2.5, and the numbers of reacting intermediates by factors of 3 - 5. This is shown in Figure S8 in the Supplement, which shows plots of these ratios against a measure of the mechanism sizes for the mechanisms of the 27 representative compounds and mechanism generation options discussed in Section 3.4. The extent of reduction in numbers of reactions is variable and has no clear dependence on the mechanism size, but the reduction in numbers of intermediates is less variable and clearly ~~increases~~decreases with mechanism size. Again, the two representative aromatics appear to be outliers in this regard, having much greater reductions in numbers of intermediates. This may be related to the fact that the aromatics tend to have the larger numbers of intermediates than comparably sized mechanisms for other compounds (see Figure S7).

Representative examples of MechGen output for a minimally processed mechanism are shown in Figure 23 and Figure S3, which use the explicit mechanism from Figure 32 or S2 as the starting point. Figure S3 shows the complete output for the products of the reactions of 1,3-butadiene in the default standard environment, while Figure 23 shows the first parts of the species and reaction listings. The format aligns with that used by the SAPRC modeling software (Carter, 2024), which is similar to formats used by other modeling software systems. The top part of the figures shows the names used for reactants and products in the mechanisms and their corresponding structures. Intermediates are named using the name of the reactant followed by "F-" and a sequence number, while stable species use various naming methods. Compounds that are important in emissions or the atmosphere are generally given 2-8 character SAPRC names based on abbreviating their actual names; compounds that may not be as important but where permanent names are or have been needed when developing SAPRC mechanisms are given permanent "ORG-nnnn" names; and others are given temporary "VOC-nnnn" names that are applicable only to this processed mechanism. Additionally, processed mechanisms using actual structures strings to name the reactants and products can also be output, as discussed in the User Manual.

3.3. Estimation of Product Yields

The relative yields of products formed in the generated mechanisms can be determined if the conditions of the environment where the compound reacts are specified and assumed to be constant. Derivations of product yields for specified environments are useful to provide a means for assessing which types of products are important and how their yields vary with environmental conditions. It is also useful for improving the efficiency and optimizing the size of multi-generation mechanisms, as discussed below in Section 4.

The algorithm MechGen uses to derive relative product yields from a processed mechanism is described in Section S3.3 of the Supplement. It uses in part the fact that most radical intermediates can be ordered such that intermediates are formed only by reactions of intermediates above it on the list. This

assumption is generally true for mechanisms derived using MechGen, with the exceptions of interconversion of phenoxy and phenyl peroxy radicals due to their reactions with O₃ and NO, respectively, and to interconversions of hydroperoxy-substituted peroxy radicals due to rapid H-shift reactions, as ~~mentioned above~~[discussed by Carter et al \(2025a\) and Vereecken and Nozière \(2020\)](#). The most rapid of the hydroperoxy-peroxy interconversions are removed by employing an equilibrium approximation, but some such interconversions are too slow for this approximation to be appropriate, as discussed in Section S3.1. For such cases, a matrix inversion procedure derived from the steady-state equations for the pseudo-first-order reactions of the interconverting radicals is employed, as discussed in Section S3.3.

Figure S4 in the Supplement shows an example of product yield output corresponding to the processed mechanism shown in Figure 23 and S3 and the explicit mechanism in Figure S2. Selected portions of this output are shown in Figure 4. This product yield output consists of three parts:

- The first part summarizes the environments used and, for the web system, gives links to obtain their oxidant concentrations.
- The second part shows the fractions of the compound that react with the various oxidants, and is not shown if compound reacts with only one reactant. For this example, most of the reaction is with OH, except in the nighttime environment where reaction with NO₃ dominates.
- The third part shows the product yields for the various environments, sorted in descending order by average yield in the environments, with organic products hyperlinked in the web output so they can be readily created and reacted if desired, and with inorganic products such as NO₂ (e.g., from peroxy + NO reactions), HO₂, O₂, and H₂O [included](#). The entry for "NO-loss" shows how many moles of NO are consumed by reaction with peroxy radicals, which is directly responsible for ~~ozone~~[O₃](#) formation when organics react in the presence of NO_x.

If the reactant forms non-volatile or semi-volatile products, the product yield output will also include mass percent yields of various volatility bins. If a representative atmospheric organic PM level is specified as a reactor option (50 µg/m³ is the default), the product yield output will also include total SOA yields corresponding to that organic PM level. This is discussed in Section S4 of the Supplement. The total mass yield of non-volatile products is also output. Note that these mass bin yields will sum up to more than 100% because of the oxygen and nitrate groups added to the molecules during the oxidation process.

3.4. Effects of Varying Full Mechanism Derivation Options

The mechanisms derived using the full mechanism operation depend on the values of the MinYld parameter and the use of environments, both of which affect which reactions and intermediates are neglected due to low relative yields. The effects of different choices in this regard are discussed in this section. This was examined by varying MinYld and environment options when deriving mechanisms for the representative compound α -pinene, and by comparing results for a more limited set of choices for 27 representative compounds of various types (listed in Table S3 in the Supplement). These include the series of n-alkanes from propane through hexadecane, various C₈ alkanes, alkenes, oxygenates, and aromatics, and two terpenes.

Figure 5 shows the effects of varying the MinYld parameter and environment options on the numbers of explicit reactions, stable products, and peroxy intermediates in mechanisms generated for α -pinene for various environment options. As expected, the number of reactions increases as the MinYld is

decreased, with no tendency to level off at either the high or low MinYld range. As also expected, using no environments resulted in the largest mechanisms, while using the three default environments (shown in Table 7) resulted in about three times fewer reactions in the case of α -pinene, and using only one environment reduces the number of reactions by almost an additional factor of two. This is because estimated upper limit yields are highest if no environments are used and increase with the numbers of environments if more than one is used. Note that the numbers of peroxy radicals are important because they determine the ~~minimize~~minimum size of the processed mechanisms, even if product lumping is employed.

Although removing reactions through the MinYld test does not affect mass balance because they are replaced by competing reactions determined to be more important, these removals will change predicted product yields, reducing the potential accuracy of the mechanism. This can be assessed by assuming that a mechanism derived with very small MinYld values and no environments can approximate the mechanism derived without deletions. Comparing product yields predicted by a mechanism generated with a given set of options with those predicted by this reference mechanism gives an indication of the effects of these deletions on predictions. The total changes in product yields are quantified as:

$$\text{Total yield change (Set, Env)} = \sum_{\text{Products}} | \text{Yield(Product, Set, Env)} - \text{Yield(Product, Ref)} |$$

where "Set" refers to a mechanism derived for a compound using the mechanism generation options being considered, "Env" is a representative environment, "Ref" refers to reference mechanism for the compound derived with the lowest MinYld and no environments, ~~"Env" is a representative environment~~, and "Yields" ~~are-is the~~ molar yield of products per mole VOC reacted under the conditions of that environment. The summation includes all organic products with yields greater than 0.0005 moles per mole VOC reacted and also amounts of NO consumed, and NO₂, HO₂ and OH radicals formed. ~~Only products with yields greater than 0.0005 moles per mole VOC reacted are included, because y~~ Yields for other products less than 0.0005 moles per mole VOC reacted are insignificant individually, and thus excluded in the summation, though not necessarily insignificant collectively. The values for the total yield change can range from zero to a maximum of the total moles of products formed per mole reactant, which is usually more than one.

Figure 6a shows the total yield changes predicted for the mid-NO_x standard urban environment for all the α -pinene mechanisms shown in Figure 5, as a function of the MinYld parameter. The yields predicted by the reference mechanism derived using a very low MinYld value of 0.01% and no environments (with ~26,000 explicit reactions) are used as the standard. As expected, yield changes increase as the value of MinYld increases, though the dependence on environment options is relatively small except at higher MinYld values. The total changes exceed 0.05 moles product per mole α -pinene reacted when MinYld > 1%, but are around 0.01 moles or less when the default MinYld value of 0.5% is used. Decreasing MinYld to 0.1% resulted in changes being decreased to 0.003 moles or less.

Figure 6b shows the effects of MinYld on moles of NO consumed per mole of α -pinene reacted, and Figure ~~6b~~6c shows effects on the yields of organic products in the condensed phase, assuming equilibrium and atmospheric aerosol levels of 50 $\mu\text{g}/\text{m}^3$ (as discussed in Section S4 of the ~~supplement~~Supplement). The effects on NO consumption reflect effects on O₃ formation because reactions of intermediates with NO are the processes responsible for ~~O₃ formation~~ ozone formation in the lower atmosphere (e.g., see Finlayson-Pitts and Pitts Jr., 1999). The amount of organic products predicted to be in the condensed phase assuming equilibrium provides an indication of the effect of the VOC's reaction on SOA formation, though this should be considered to be highly approximate because effects on SOA depend on environmental conditions, estimates of vapor pressure are uncertain, and the equilibrium

approximation is an oversimplification. Figure 6b and c indicate that, for α -pinene at least, the apparent effects on both O₃ and SOA are relatively small, being only 0.3% or less when the default MinYld parameter is used.

Figure 6 shows effects of MinYld on product yield changes for α -pinene under mid-NO_x urban conditions. The effects on yield changes for other standard environments incorporated in MechGen are shown in Figure S6 in the Supplement. The total yield changes for the daytime environments are similar, though the effect on NO consumption is lower under lower NO_x conditions because there is less NO consumed. The total yield changes are much less under nighttime conditions, presumably because fewer reactions tend to be important. Note that no NO is present in the nighttime scenario, and MechGen predicts that α -pinene forms much lower yields of low-volatility products at night compared to daytime.

The effects of varying MinYld and environment options for other compounds are shown in Figure 7. This shows effects on (a) numbers of explicit reactions, and (b) total yield changes, for the mid-NO_x environment for 27 representative compounds of various types (listed in Table S3 in the Supplement). Plots showing ratios of numbers of reacting intermediates or stable products to numbers of reactions for these compounds with different mechanism generation options are shown on Figure S7. As expected, the mechanism size increases with the size of the molecules, but the type of molecule is also important, as indicated by the large variability for the representative C₈ compounds. Highly branched compounds and aromatics tend to have the smallest generated mechanisms, whereas cyclic alkenes, such as terpenes, tend to have the largest. The effects on product yield tend to correlate with mechanism size, though the results are variable and the correlation is relatively weak (~50%) except at the lowest MinYld level.

595 Table 7. Standard environments available for determinations of which reactions are negligible
 596 during mechanism generation and for estimation of product yields.

Scenario	High NO _x	Standard (mid NO _x)	Low NO _x	Nighttime
Description [a]	Maximum VOC reactivity (MIR)	Equal sensitivity of O ₃ to VOC and NO _x (EBIR)	NO _x levels 1/10 those giving maximum O ₃ (MOIR NO _x / 10)	EBIR scenario at nighttime
Default use [b]	Gen, PY	PY, MGen	Gen, PY	Gen, PY
		Component concentrations (molec/cm ³)		
OH	7.51e+6	1.11e+7	3.69e+6	2.86e+5
O ₃	1.78e+12	2.26e+12	9.20e+11	2.90e+12
O ³ P	5.50e+4	2.84e+4	8.85e+3	~ 0
NO	9.58e+10	1.59e+10	4.12e+9	7.43e+5
NO ₂	3.15e+11	8.63e+10	1.32e+10	4.88e+10
NO ₃	8.54e+6	9.92e+6	1.25e+6	2.25e+9
HO ₂	~ 0	6.00e+8	7.17e+8	1.27e+8
Total RO ₂ 's	~ 0	4.59e+8	6.69e+8	2.35e+8
Total RCO ₃ 's	~ 0	7.30e+7	7.27e+7	1.04e+8

597 [a] See Section S2 in the Supplement for more information on these scenarios. All these represent urban
 598 conditions similar to those used to derive the (Carter, 1994) reactivity scales except that the total
 599 VOC levels were reduced to more closely represent modern ambient conditions.

600 [b] Gen = used to determine negligible reaction during full mechanism generation; PY = used for
 601 determination of product yields; MGen = used for multi-generation mechanism derivations discussed
 602 in Section 4.

|

|603
|604

605 Table 8. List of parameters that affect full mechanism generation.

Parameters	Affects	Default
<u>Parameters that affect all full mechanism generation operations</u>		
Minimum yield (MinYld)	Reactions with estimated maximum yields below this value are excluded during full mechanism generation. See Scheme 2 in the Supplement. [a]	0.5%
Environmental conditions	Sets of environmental conditions to determine upper limit yields involving bimolecular reactions and for displaying product yields. Users can select which environments, if any, are used for mechanism generation.	High NOx Low NOx Nighttime
Lumping	Set of lumping options that control generation and processing for lumped mechanisms. See Section 5.	Explicit Lumping
Lumping	Set of lumping options that control generation and processing for lumped mechanisms. See Section 5.	No Lumping
kFastUni	Minimum unimolecular rate constant for a non-radical product to be treated as an intermediate and reacted.	$1.67 \times 10^{-2} \text{ sec}^{-1}$
kMinUni	Minimum rate constant for unimolecular reactions of peroxy radicals not to be neglected.	$2.25 \times 10^{-3} \text{ sec}^{-1}$
kFastUni	Minimum unimolecular rate constant for a non-radical product to be treated as an intermediate and reacted.	$1.67 \times 10^{-2} \text{ s}^{-1}$
kMaxUni	Total unimolecular rate constant for peroxy radicals above which bimolecular reactions are ignored. [a]	1000 sec^{-1}
kRO2slow	Minimum rate constant for unimolecular reactions of peroxy radicals not to be neglected.	$1.0 \times 10^{-3} \text{ s}^{-1}$
kEqConv	Peroxy radical interconversion rate constant above which the interconverting radicals can be treated as in equilibrium when the mechanisms are processed.	30 s^{-1}
kRO2fast	Total unimolecular rate constant for peroxy radicals above which bimolecular reactions are ignored. [a]	50 s^{-1}
Explicit radical list	Organic radicals that are treated as final products and not reacted during "fully react" operations.	None (list empty)
<u>Parameters that affect only multi-generation operations. See Section 4</u>		
Reacting environment	Environment used to estimate product yields to determine if the organic product should be reacted.	Standard (mid NO _x)
MGminYld	Minimum estimated yield for a product formed in a multi-generation operation to react.	0.01%
MinVP	Minimum estimated vapor pressure for a product formed in a multi-generation operation to react.	$1.0 \times 10^{-13} \text{ atm.}$

RxnHours

Reaction time used to estimate amounts of products reacted in the environment for the purpose of determining if its products may be non-negligible.

6 hours

Parameter affecting estimates of SOA formation. See Section S4 in the Supplement.

PM

Concentration of organic particle matter in $\mu\text{g}/\text{m}^3$

50 $\mu\text{g}/\text{m}^3$

[a] These tests are not used when hydroperoxy-substituted peroxy radicals rapidly isomerize to form an another such radical that can isomerize back to the initial radical, resulting in no net loss of the radical, making competing reactions potentially non-negligible.

```
.ACT
13-BUTDE ! CH2=CH-CH=CH2
ACROLEIN ! CH2=CH-CHO
VOC-0016 ! CH2=CH-CH*-O-O*
ORG-4682 ! CH2=CH-CH(ONO2)-CH2-OH
ORG-1004 ! CH2=CH-CH(CH2-OH)-O-OH
ORG-1048 ! CH2=CH-CO-CH2-OH
...

.STS
13-BUTDer1 ! CH2=CH-CH[OO.]-CH2-OH
...

.RXN
R) 1.12e-11 -1.053 0.00 ;13-BUTDE + OH = #.62 13-BUTDer1 + #.33 13-BUTDer2 + #.05 13-BUTDer3
R) 1.34e-14 4.537 0.00 ;13-BUTDE + O3 = #.5 ACROLEIN + #.14 VOC-0016 + #.36 13-BUTDer9 + #.09 H2O + #.08 OH + #.5 HO2 +
#.54 CO + #.12 CO2 + #.5 HCHO + #.21 HCHO2 + #.09 H2
R) 1.10e-13 ;13-BUTDE + NO3 = #.32 13-BUTDer7 + #.18 13-BUTDer8 + #.5 13-BUTDer6
R) 9.13e-12 ;13-BUTDer1 + NO = #.95 ACROLEIN + #.05 ORG-4682 + #.95 HO2 + #.95 NO2 + #.95 HCHO
R) 2.30e-12 ;13-BUTDer1 + NO3 = ACROLEIN + O2 + HO2 + NO2 + HCHO
R) 1.61e-11 ;13-BUTDer1 + HO2 = ORG-1004 + O2
R) 1.60e-11 ;13-BUTDer1 + RC03 = #.8 ACROLEIN + #.2 ORG-1048 + O2 + #.8 HO2 + #.8 HCHO
...
```

Figure 2. Portions of MechGen output showing the minimally reduced processed mechanism for the reactions of 1,3-butadiene under default conditions. The complete output is shown in Figure S3 in the Supplement.

Rxn	k	Fac	Weight	Reaction
1	6.30e-11	95%	95.0%	CH2=CH-CH=CH2 + OH -> .aCH2-aCH-aCH[.]-CH2-OH {*OHadd}
2	3.31e-12	5%	5.0%	CH2=CH-CH=CH2 + OH -> CH2=CH-CH(CH2.)-OH {*OHadd}
3	3.15e-18	50%	50.0%	CH2=CH-CH=CH2 + O3 -> CH2=CH-CHO + CH2OO {*O3ole}
4	3.15e-18	50%	50.0%	CH2=CH-CH=CH2 + O3 -> HCHO + CH2=CH-CHOO {*O3ole}
...				
7		100%	95.0%	.aCH2-aCH-aCH[.]-CH2-OH {*OHadd} -> .aCH2-aCH-aCH[.]-CH2-OH
8		65%	61.7%	.aCH2-aCH-aCH[.]-CH2-OH + O2 -> CH2=CH-CH[OO.]-CH2-OH
9		35%	33.3%	.aCH2-aCH-aCH[.]-CH2-OH + O2 -> .OOCH2-CH=CH-CH2-OH
10	8.64e-12	95%	58.5%	CH2=CH-CH[OO.]-CH2-OH + NO -> CH2=CH-CH[O.]-CH2-OH + NO2
11	4.80e-13	5%	3.2%	CH2=CH-CH[OO.]-CH2-OH + NO -> CH2=CH-CH(ONO2)-CH2-OH
12	2.30e-12	100%	61.7%	CH2=CH-CH[OO.]-CH2-OH + NO3 -> CH2=CH-CH[O.]-CH2-OH + NO2 + O2
13	1.61e-11	100%	61.7%	CH2=CH-CH[OO.]-CH2-OH + HO2 -> CH2=CH-CH(CH2-OH)-O-OH + O2
...				
16	2.45e+8	100%	169.6%	CH2=CH-CH[O.]-CH2-OH -> CH2=CH-CHO + .CH2-OH
17	1.16e+0	100%	33.3%	.OOCH2-CH=CH-CH2-OH -> HO-aCH[.]-aCH-aCH[.]-CH2-O-OH
18	8.47e-12	93%	30.8%	.OOCH2-CH=CH-CH2-OH + NO -> .OCH2-CH=CH-CH2-OH + NO2
...				
91		14%	6.8%	CH2=CH-CHOO {*O3ole} -> CH2=CH-CHOO[syn]
92		28%	13.8%	CH2=CH-CHOO {*O3ole} -> CH2=CH-CHOO[anti]
93		59%	29.5%	CH2=CH-CHOO {*O3ole} -> .CH2-CHO + HCO.
...				

Figure 3. Portions of MechGen output showing representative results of a full mechanism generation operation for the reactions of 1,3-butadiene with default options. The full output is shown in Figure S2 in the Supplement.

```
.ACT
13-BUTDE      ! CH2=CH-CH=CH2
ACROLEIN      ! CH2=CH-CHO
VOC-0014      ! CH2=CH-CH*-O-O*
ORG-4682      ! CH2=CH-CH(ONO2)-CH2-OH
ORG-1004      ! CH2=CH-CH(CH2-OH)-O-OH
ORG-1048      ! CH2=CH-CO-CH2-OH
...

.STS
13-BUTDE-1    ! CH2=CH-CH[OO.]-CH2-OH
...

.RXN
R) 1.12e-11 -1.053 0.00 ;13-BUTDE + OH = #.62 13-BUTDE-1 + #.33 13-BUTDE-2 +
                               #.05 13-BUTDE-3
R) 1.34e-14 4.537 0.00 ;13-BUTDE + O3 = #.5 ACROLEIN + #.14 VOC-0014 +
                               #.36 13-BUTDE-8 + #.09 H2O + #.08 OH + #.5 HO2 +
                               #.54 CO + #.12 CO2 + #.5 HCHO + #.21 HCHO2 +
                               #.09 H2
R) 1.10e-13 ;13-BUTDE + NO3 = #.32 13-BUTDE-6 + #.18 13-BUTDE-7 +
                               #.5 13-BUTDE-5
R) 9.13e-12 ;13-BUTDE-1 + NO = #.95 ACROLEIN + #.05 ORG-4682 +
                               #.95 HO2 + #.95 NO2 + #.95 HCHO
R) 2.30e-12 ;13-BUTDE-1 + NO3 = ACROLEIN + O2 + HO2 + NO2 + HCHO
R) 1.61e-11 ;13-BUTDE-1 + HO2 = ORG-1004 + O2
R) 1.60e-11 ;13-BUTDE-1 + RCO3 = #.8 ACROLEIN + #.2 ORG-1048 +
                               O2 + #.8 HO2 + #.8 HCHO
...
```

Figure 3. Portions of MechGen output showing the minimally reduced processed mechanism for the reactions of 1,3-butadiene under with default conditions. The complete output is shown in Figure S3 in the Supplement.

Mid NOx	High NOx	Low NOx	Night	Explicit product or reacted
98.0%	97.6%	97.6%	6.7%	OH reacted
1.9%	2.2%	2.3%	6.4%	O3 reacted
0.1%	0.2%	0.1%	86.9%	NO3 reacted
63.97%	78.14%	49.30%	117.28%	NO2
89.31%	93.43%	78.80%	13.06%	HO2.
67.42%	82.55%	51.98%	0.07%	NO-loss
5.00%	-	17.48%	82.13%	O2
57.16%	61.01%	47.01%	59.51%	HCHO
54.74%	58.34%	45.00%	58.91%	CH2=CH-CHO
26.48%	17.03%	28.71%	13.59%	HCO-CH=CH-CH2-O-OH
3.74%	-	13.83%	0.95%	CH2=CH-CH(CH2-OH)-O-OH
3.10%	11.98%	0.87%	2.61%	HCO-CH=CH-CH2-OH
...				
0.06%	-	0.16%	-	SOA mass yield from products for atm PM = 50 ug/m3

(lower yield products not shown)

Figure 4. Portions of MechGen output showing estimated product yields the reactions of 1,3-butadiene with OH, O₃ and NO₃ radicals for various environments, derived using default mechanism generation options and environments.

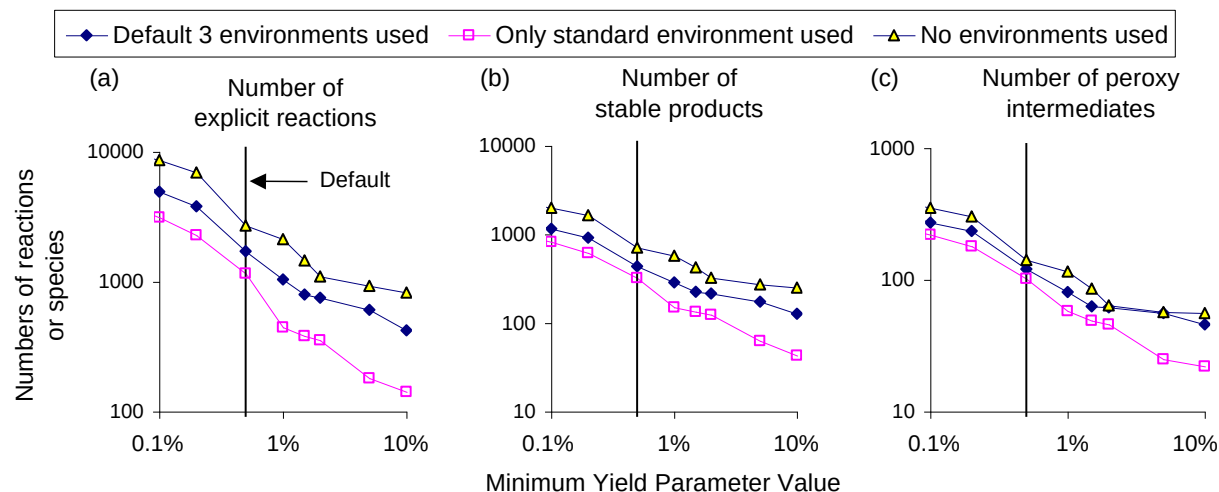


Figure 5. Effects of varying the MinYld parameter and environment options on the numbers of explicit reactions in full mechanisms generated for α -pinene.

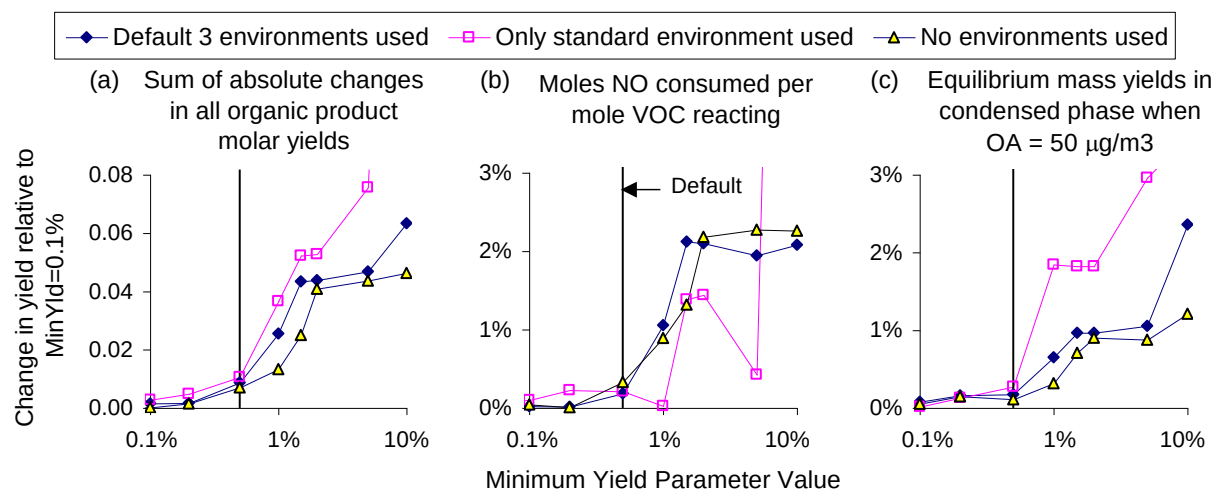


Figure 6. Effects of varying the MinYld parameter and environment options on relative changes in yields of selected groups of products from α -pinene when reacted under mid- NO_x urban environmental conditions.

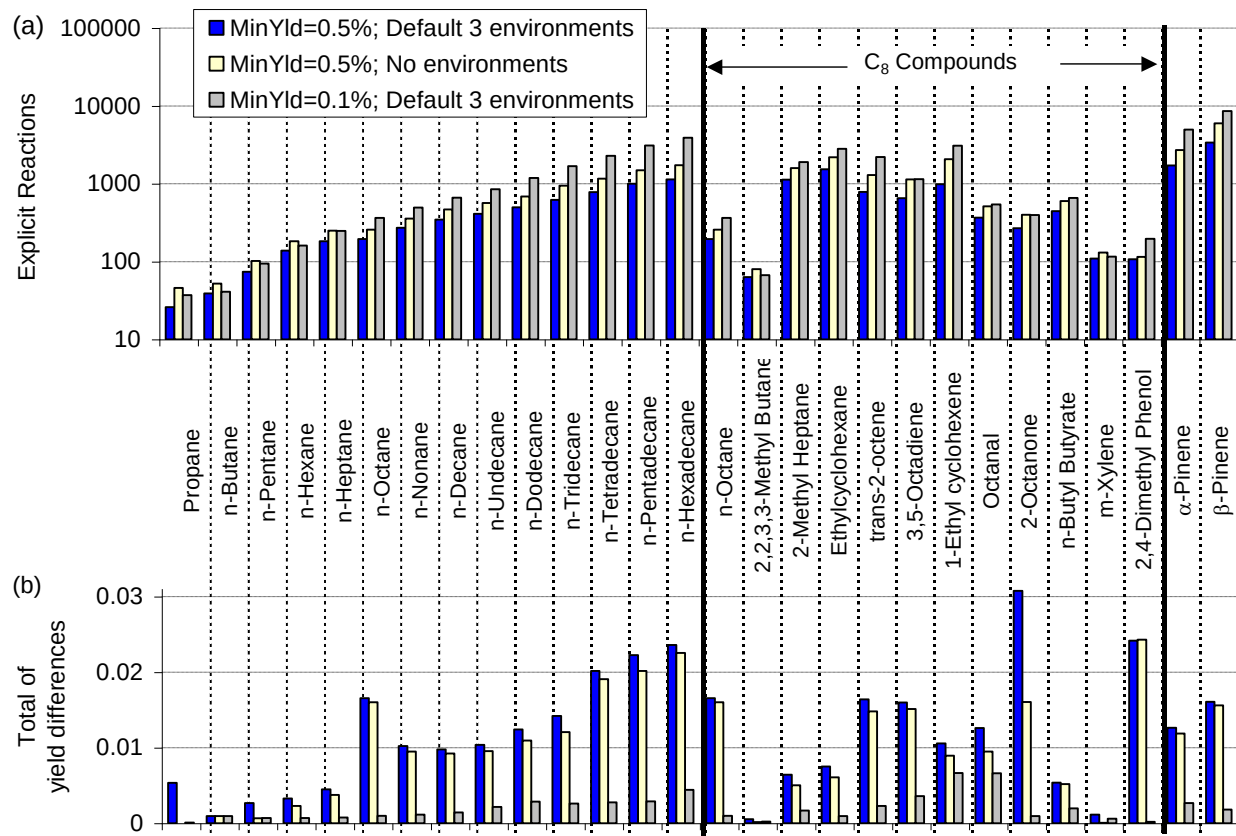


Figure 7. Numbers of explicit reactions and sum of differences in product yields relative to the largest generated mechanisms for the various mechanism generation options.

4. DERIVATION OF MULTI-GENERATION MECHANISMS

4.1. Description of Process and Outputs

The full mechanism generation process discussed above produces only a single generation mechanism because it does not react the stable products formed. MechGen can also derive multi-generation mechanisms where all reactive products formed in non-negligible yields, including those formed after reacting multiple generations, are reacted. It involves carrying out full mechanism generation for a selected starting compound, then determining the stable products formed, then generating mechanisms for products in non-negligible yields, then repeating the procedure until no reactive, non-negligible products remain. This would be relatively straightforward, except that reacting all products regardless of their yields would result in unmanageably large mechanisms, dominated by reactions and species of negligible importance under conditions of interest. Therefore, much of the multi-generation derivation by MechGen ~~focused~~ focuses on determining which products are in fact non-negligible and which don't have to be reacted.

The procedures used for deriving multi-generation mechanisms are described in the User Manual. They involve accessing the system using the terminal interface, creating a software object to control the process for a specified compound and to store the results, and then using it to carry out the operations and transmit the results. These objects have several parameters that control which products are formed in sufficiently high yields to react and which can be treated as reactive. One of these is the specification of an environment used to estimate product yields and another is the "MGminYld" parameter that determines which yields are treated as negligible. In addition, to determine which product can be treated as reactive, a "MinVP" parameter gives the minimum vapor pressure for product compound to react in the gas phase, and a "RxnHours" parameter determines the time for compounds to react and form products. These options can be changed by the users, but by default, the environment used for the multi-step mechanism generation is the first one listed to derive product yields as discussed Section 3.3. The default values of MGminYld, MinVP, and RxnHours are 0.01%, 10^{-13} atm, and 6 hours, respectively (Table 8). These need to be specified to minimize the numbers of totally negligible reactions and processes in the multi-generation mechanisms that are derived.

The algorithm MechGen uses to derive multi-generation mechanisms is summarized below, and a flowchart showing the process is given in Scheme S6 in the Supplement. The process starts by generating full single-generation mechanisms for the initial reactant and for all products formed in non-negligible yields. Each mechanism generation is followed by deriving minimally reduced mechanisms, estimating product yields for the selected environment, and determining what products are formed in sufficiently high yield to react. The estimated upper limit yields are derived based on:

$$\frac{\text{Yield of initial reactant}}{\text{Yield of initial reactant}} = \frac{\text{Reactivity correction for initial reactant}}{\text{Reactivity correction for initial reactant}} = 100\%$$

$$kP_{\text{Uni}} = k_{\text{Uni}} + k_{\text{hv}} + \sum_{\text{OX}} k_{\text{OX}} \times [\text{OX}]^{\text{environment}}$$

$$\frac{\text{Reactivity correction for reacting product}}{\text{Reactivity correction for reacting product}} = 1 - e^{-kP_{\text{Uni}} \times \text{RxnTime}}$$

$$\text{Est. yield of product} = \frac{\text{Est. yield of reactant forming it}}{\text{Est. yield of reactant forming it}} \times \frac{\text{Reactivity correction for reactant forming the product}}{\text{Reactivity correction for reactant forming the product}} \times \text{Yield of product from reactant}$$

where "kPUni" is the pseudo-unimolecular rate constant for the reactions of a reactant in the environment, "kUni" and "khv" are the unimolecular and photolysis rate constants for the reactant (if applicable), "OX" and "k_{OX}" refer to all oxidants with which the compound reacts and their rate constants, "RxnTime" is derived from "RxnHours" to be consistent with the units used for kPUni, and "Est." refer to upper limit estimates. The yields of products from reactants in the environment are derived as discussed in Sections 3.3, and the kPUni values are derived by multiplying the rate constants for the initial reactions multiplied by the concentration of the oxidant with which it reacts in the environment, if applicable. The reactivity correction factors are calculated as fractions in the environment and are used to account for the fact that slower reacting compounds form lower amounts of products during the reaction time in the environment. The formula for calculating fractions reacted, derived from integrating the kinetic differential equation, is

$$\text{Factor Reacted (Reactant, Environment)} = 1 - e^{-k\text{PUni(Reactant, Environment)} \times \text{RxnTime}}$$

This gives near 100% fractions reacted for rapidly reacting compounds, and fractions that are approximately proportional to the kPUni for slowly reacting compounds. Note that the estimated upper limit yields tend to decrease as the number of generations of reactions increases.

The standard environments used (or not used) when carrying out the full, single-generation mechanism derivations for individual compounds, as discussed in Section 3.1, do not necessarily need to be the same as the one required to carry out the multi-generation process. If standard environments are used during the single-generation mechanism derivation process for individual compounds, they should also represent the environment used for the multi-generation process. This is assured if the environment used for the multi-generation process is one of those used for the single-generation process, or if no environments are used for the single-generation process. By default, the three standard environments used for single-generation processes incorporate the conditions of the default standard environment for multi-generation derivations, as indicated in Table 7.

The results of this process consist of lists of reacting species, their reactions and rate constants, and lists of low reactivity, low volatility, or low yield products that are not reacted. As discussed in the User Manual, these results can be transmitted to users in several types of files. These include: (1) files containing lists of reactants, non-negligible products, and counter species representing atom numbers and total masses of low yield products and their approximate yields; (2) files giving information about the reactants and products in terms of groups in the molecules; (3) files with listings of the explicit and processed reactions of the reacting compounds; and (4) files that can be used as input to prepare the mechanisms for model simulations using SAPRC box modeling software (Carter, 2024). The model preparation input files come in two forms: one containing the minimally reduced processed mechanisms for all the reactants, and the other giving pseudo-unimolecular reactions of the reactants that are applicable only to the conditions of the environment used to generate the mechanism. The specific types of information that can be obtained are listed in Table S4 of the Supplement.

4.2. Examples of Results

To illustrate the results of multi-generation mechanism derivation operations and their dependence on selected options, mechanisms were derived for several example compounds ranging from propane to α -pinene using a standard set of options, and additional multi-generation mechanisms were derived for α -pinene using differing sets of options. The compounds and mechanism derivation options are summarized in Table 9, which gives the sets of options, the numbers of explicit reactions, information on lost or low yield carbon, and estimated SOA formation for the mechanisms. In ~~all cases,~~the cases where the compound or MGminYld parameter was varied, the single-generation mechanisms for individual reactants were derived using the three default environments listed in Table 7 and the default

mechanism generation options given in Table 8. In the cases where the environments were varied, the single-generation mechanisms were derived using the stated environment only, with defaults used for the other parameters. Table 9 shows that, as expected, the results depend significantly on the MGMinYld parameter and the multi-generation environment used.

The pseudo-unimolecular reaction output was used to prepare input for the SAPRC box modeling software (Carter, 2024) to carry out model simulations for the evolution of product concentrations and negligible atom counter species over time, under conditions of the environment used to derive the multi-generation mechanism. The use of this output is equivalent to assuming that the atmospheric oxidant levels are constant at the levels associated with the environment, as given in Table 7, and at constant light intensity with photolysis rates calculated for the light associated with the reactor. The model simulations were carried out for 6 hours of simulated time, with the starting reactant being initially present at 1 ppm concentration, mixing ratio, with no dilution or variation in reaction conditions or oxidant species. The calculated concentrations were used in conjunction with separate MechGen output giving structural information about the compounds (Carter et al., 2025a) for the purpose of computing total yields of compounds with different structures (structural counter species), and total carbons in compounds that were not reacted because of estimated low yields or reactivities.

Figure 8 presents concentration-time results for the highest yield structural counter species or C₁ products for α -pinene, and these results for all representative compounds listed in Table 9 are given in Figure S10 of the Supplement. These mechanisms were all generated using a MGminYld value of 0.001% and the mid NO_x urban standard environment, which are the defaults. These figures also show the O/C ratios for the reactant and products as a function of time. The structural counter species are listed and briefly described in Table S4 in the Supplement. Concentrations relative to the amount of starting material reacted at the times are shown at 30-minute intervals up to 2 hours of simulation time, then at 2-hour intervals after that. The evolution of the ratio of total carbon in low yield unreacted compounds (NegC) to total reacted carbon is also shown in Figure 8 or S10, except for cases when they are near zero.

Figure 8 and S10 show that the O/C ratios increase with time, as expected due to ongoing oxidation. The yields of the C₁ products also increase significantly and make relatively high contributions to the total products formed by the end of the simulations. Relative concentrations of other types of products tend to go down with time after the first half hour, particularly more reactive products such as aldehydes. As expected, the distribution of products significantly depends on the type of compound. Note that these simulations treat formaldehyde as a non-reacting product, while in fact a significant fraction of it would react to form CO and CO₂ by the end of the simulations.

Examples of effects of using different environments on multi-generation mechanisms derived for α -pinene are shown in Table 9 and Figure 9. In those examples, the single-generation mechanisms were derived using only the subject environment rather than the three default environments, which is the reason that the number of reactions derived for α -pinene in these calculations is smaller than in those where the compound or MGminYld parameter was varied (see Figure 5), though the overall product yields are essentially the same. As discussed earlier, the standard environment represents intermediate NO_x conditions, and thus includes both high and low NO_x reactions. Therefore, as shown in Table 9, the mechanism derived for it has more reactions and reactants than those derived separately for high and low NO_x conditions. On the other hand, fewer types of reactions are predicted to be important in the nighttime environment, so significantly smaller mechanisms are derived. As shown in Figure 9, the concentrations of compounds with hydroperoxide groups are highest under low NO_x conditions and the yields of N-containing compounds are lower, though the yields of nitrates or peroxyacylnitrates (PANs) are not as strongly affected as one might expect based on the relative NO_x levels in the environments. The yields of compounds with ester, alcohol, or peroxy acid groups are significantly lower under nighttime conditions,

though the predicted levels of the other compounds are generally within the ranges predicted for the daytime environments. Further discussion of the mechanistic implications of these results is beyond the scope of this paper, but this is worthy of additional study.

The effects of varying the MGminYld parameter on products or counter species in the α -pinene mechanisms are shown in Figure S11 in the Supplement. Increasing MGminYld raises yields of neglected compounds (reflected by NegC), resulting in a slight decrease in yields of most products, with the yields decreasing by ~10% when MinYld is increased from 0.25% to 1%. This means that yields in multi-generation mechanisms are necessarily lower limits unless very low values of MGminYld are used to minimize the amount of untreated carbon (NegC).

Table 9. Summary of multi-generation mechanism derivation examples.

Compound	Options [a]	Mech Size [c]	Neg /Rct'd	SOA
Compound	MGminYld Environ [b]	Rxns Ret'd	C [d]	[e]
	MGminYld Environ [b]	Reactions Reacted	C [c]	[d]
Propane	0.01% (default)	Mid NO _x urban (default)	339 37	0.03% ~0
Propane			299 37	0.03% ~0
n-Butane			1457 110	0.2% 0.01%
n-Butane			1203 109	0.2% 0.01%
1-Butene			1582 125	0.3% 0.02%
1-Butene			1381 126	0.3% 0.02%
n-Octane	0.01% (default)	Mid NO _x urban (default)	82,076 1433	6% 7%
n-Octane			50235 1466	6% 7%
Trans-2-octene			32,698 934	4% 5%
Trans-2-octene			24570 945	4% 5%
α -Pinene			113,379 1698	12% 52%
α -Pinene			82375 1695	12% 52%
α -Pinene	0.01%	High NO _x	76,652 1278	8% 35%
		High NO _x	20819 1252	8% 36%
		Mid NO _x	113,379 1698	12% 52%
		Mid NO _x	56446 1683	12% 52%
α -Pinene	0.01%	Low NO _x	92,495 1340	9% 50%
		Night	8493 95	0.3% 7%
		Low NO _x	44933 1337	9% 51%
		Night	5961 95	0.3% 7%
α -Pinene	0.02%	Mid NO _x	78,098 1080	16% 50%
	0.01%		113,379 1698	12% 52%
	0.02%		56952 1088	16% 49%
α -Pinene	0.01%	Mid NO _x	82375 1695	12% 52%
	0.005%		151,139 2569	9% 55%
	0.005%		116426 2587	9% 54%

[a] All other options were held at defaults shown in Table 8.

[b] Environment(s) used when deriving the multi-generation mechanism.

~~[c] Rxns = numbers of reactions in the minimally reduced processed mechanisms derived for all the reacting reactants. Ret'd = number of reacting reactants.~~

~~[d][c]~~ Ratio of moles carbon in compounds that are not reacted due to low yields divided by moles carbon in the reacting starting compound after reacting for RxnTime = 6 hours.

~~[e][d]~~ SOA = total moles of all low-volatility products multiplied by their estimated fractions in the particle phase when total atmospheric SOA levels were set at $50 \mu\text{g}/\text{m}^3$ (see Section S4).

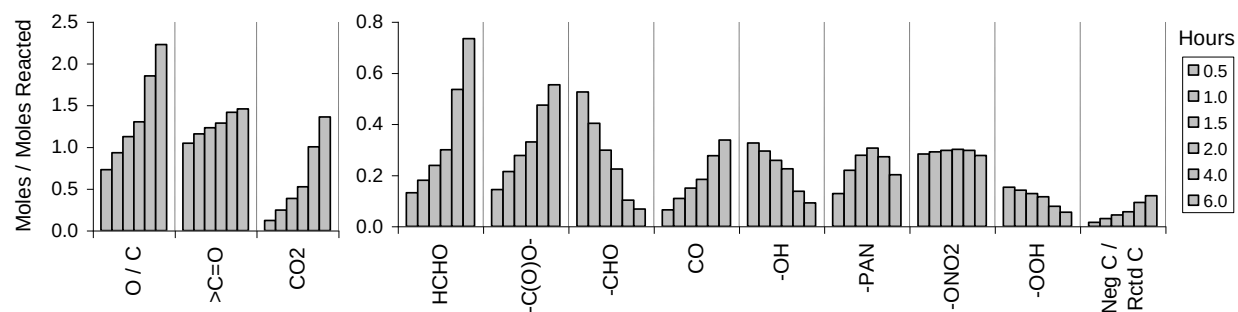


Figure 8. Moles of various types of products or functional groups formed as a function of time for 6-hour simulations using multi-generation mechanisms for α-pinene. Products are ordered from high to low maximum yields.

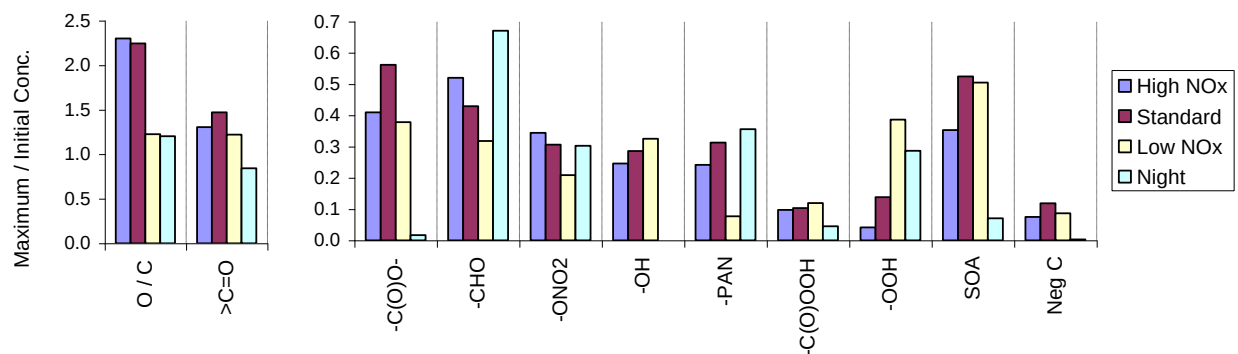


Figure 9. Effects of varying the environment on maximum molar concentrations of various types of products or functional groups for the simulations using multi-generation mechanisms for α-pinene. ~~The corrected yields assuming that the distribution of products from unreacted compounds to those that were reacted are also shown.~~

5. DERIVATION OF LUMPED MECHANISMS

Though smaller than the explicit mechanisms, the minimally reduced mechanisms are still too large for use in practical modeling applications involving complex atmospheric mixtures. Further reduction is necessary, both by using a limited number of lumped model species to represent chemically similar compounds, and by using various approaches to reduce the numbers of reactive intermediates in the mechanisms. MechGen can optionally be used to derive lumped mechanisms for organics that are consistent with the lumping approaches used in recent SAPRC mechanisms (Carter, 2010a, b, 2023; Carter and Heo, 2013). ~~Lumping involves representing the many predicted organic products by lumped model species representing groups of products assumed to react similarly, and by using various approaches to reduce the numbers of intermediates in the mechanism.~~ As indicated in Table 8, the lumping approach must be specified as one of the options used for full mechanism generation. The default lumping option is "explicit" or no lumping, which involves generating all reactions expected to be non-negligible and then preparing only minimally reduced processed mechanisms from the results, with no additional processing. This was used for all the examples discussed in this paper. Other lumping options currently available in MechGen are briefly summarized below, with additional details given in the User Manual, as well as relevant SAPRC mechanisms papers and reports. A more complete discussion of lumping algorithms is beyond the scope of the present paper and will be presented in more detail in a subsequent paper ~~on the SAPRC-22 mechanism that is in preparation~~ (see also Carter, 2023).

SAPRC-11 lumping is the approach used in the SAPRC-11 mechanism (Carter and Heo, 2013), and is essentially the same as that used for SAPRC-07 (Carter, 2010a, b) except as discussed by Carter and Heo (2013). All organic products are represented by one of ~50 lumped or explicit model species, and all peroxy intermediates are represented by ~30 chemical operators that represent products formed and how their yields may change with NO_x conditions. (An "operator" in this discussion refers to a model species that does not correspond directly to a chemical species but is added to represent overall effects of various reactions [e.g., Carter, 2020a,b]). In addition, all acyl peroxy, phenoxy, and phenyl peroxy intermediates are represented using lumped radical model species in all current SAPRC mechanisms. Note that the peroxy operator method as originally implemented for SAPRC-07 and SAPRC-11 ~~did not consider~~ does not provide for peroxy radical isomerization reactions, so strictly speaking they are not compatible with mechanisms currently generated using MechGen, which includes such reactions when predicted. This issue is addressed by assigning an "effective NO" concentration to take competitions between unimolecular and bimolecular reactions into account when determining yields of operators representing the products formed, as discussed by Carter (2023). The default effective NO for SAPRC11 lumping is 0.5 ppb, reflecting intermediate NO_x urban conditions.

SAPRC-22 lumping is the approach used in the "standard" version of SAPRC-22 mechanism (Carter, 2023). This approach represents organic products using ~110 lumped or explicit model species, with the numbers of lumped species increased in part to better represent NO_x recycling and SOA formation potentials. Peroxy radicals are represented using a modified version of the operator approach used for SAPRC-07 and -11, in which operator yields are calculated at both low and high effective NO levels, and an extra operator species is added for each reaction where the yields are different at different NO levels, so the lumped mechanism is applicable for a wide range NO conditions. This lumping approach was used to derive the SAPRC-22 mechanism (Carter, 2023), using procedures discussed in the User Manual.

If SAPRC-22 lumping is selected, users have the option to generate SAPRC-22-compatible lumped mechanisms that represent selected specific organic compounds explicitly in models, rather than having them lumped as in the standard SAPRC-22 mechanism. The products formed in the reactions of

these explicitly represented compounds will then be lumped to be compatible with SAPRC-22 and these lumped reactions can be output along with the rest of the SAPRC-22 mechanism to be used as input to the SAPRC model preparation programs (Carter, 2024). These expanded versions of SAPRC-22 can then be used in airshed models to evaluate the effects of the selected compounds on O₃ or other measures of air quality, or to allow comparisons of their predicted concentrations with observations.

If any type of SAPRC lumping is selected, the mechanism generation process will exclude the reactions of acyl peroxy, phenoxy, or phenyl peroxy radicals, as these are all represented by lumped model species, so their explicit reactions are not used when deriving the lumped mechanism. In addition, if a non-explicit lumping method is employed, the types of initial reactions the organic compounds undergo are determined by the types of initial reactions of the lumped model species used to represent them, which is part of the input used to define the lumping approach, as discussed in the User Manual. In some cases, this is a subset of the types of initial reactions generated when the default "explicit" ~~(no)~~ lumping(no lumping) option is in effect.

6. DISCUSSION

Because of the complexity of the atmospheric reactions of most organic compounds, automated mechanism generation systems such as MechGen provide an important link from chemical kinetic and mechanistic data and theories to model predictions of chemical transformations in the atmosphere. As discussed by Kaduwela et al. (2015) and Ervens et al. (2024), the stages of mechanism development for models that provide this link consist of (1) compiling and evaluating available basic mechanistic data; (2) developing SARs and other estimation methods needed to develop complete mechanisms; (3) deriving detailed mechanisms that incorporate these data and estimation methods; (4) reducing the detailed mechanisms so they can be used in various modeling applications; and (5) evaluating the predictions of the mechanisms against observations. MechGen is a tool that can provide the link between mechanistic data and theory and detailed mechanisms, and can also be used for mechanism reduction. The development of the more recent SAPRC mechanisms for airshed models (Carter, 2010a, b, 2023; Carter and Heo, 2013) was undertaken with this approach in mind, using MechGen to derive the SAPRC lumped mechanisms from the explicit mechanisms for the many compounds that are represented.

Detailed mechanisms derived by MechGen can also be used in analyses of laboratory (e.g., Li et al., 2022) or ambient data, providing an alternative or supplement to the Master Chemical Mechanism (MCM) (Bloss et al., 2005; Jenkin et al., 1997, 2003; Saunders et al., 2003), which has been widely used for this purpose. MechGen offers the flexibility of being applicable to any compound within the scope of its predictive capabilities, whereas MCM, although comprehensive, is a manually-developed mechanism that covers a finite number of compounds. Another alternative without this limitation is the GECKO-A mechanism generation system (Aumont et al., 2005), which also has some online functionality (<https://geckoa.lisa.u-pec.fr/>, <https://www2.acom.ucar.edu/modeling/gecko>).

~~The main reason that MechGen has not been as widely used as MCM or GECKO-A for applications other than SAPRC mechanism development is that it has not been described or documented in the peer-reviewed literature until now, including the continuous revisions that it has undergone. Most researchers have either not been aware of MechGen and have not had a stable version to cite if they do use it. This paper, along with the recently published paper of Carter et al. (2025), addresses its lack of adequate documentation. Future updates to MechGen will be documented and made available as separate versions, so work using versions prior to the updates can be duplicated.~~

One advantage of using MechGen is its relatively user-friendly online interface that allows new users to easily examine predictions for single-step reactions or complete mechanisms for selected compounds without extensive training. It also offers terminal-based access that can be obtained using accounts created using the online system to use features of the system not available to web users, and users can download and install their own copies of this software to use full features, including resource-intensive operations like multi-generation mechanism derivation. This is discussed in the User Manual that is available at the MechGen web site (Carter, 2025). MechGen output can serve as input to the SAPRC box modeling software (Carter, 2024) to carry out model simulations with either minimally reduced detailed mechanisms or extended versions of the SAPRC-22 mechanism (Carter, 2023) with selected compounds represented explicitly. It can also be used for comparing its predictions with those of GECKO-A or incorporated in MCM to assess differences in predictions that indicate areas where additional experimental, theoretical, or SAR development work are needed.

One of the reasons that MechGen has not been as widely used as MCM or GECKO-A for applications other than SAPRC mechanism development is that it has not been described or documented in the peer-reviewed literature until now, including the continuous revisions that it has undergone. Most

researchers have either not been aware of MechGen or have not had a stable version to cite if they do use it. An additional factor that may inhibit its use as an alternative to MCM is that MechGen outputs mechanisms in the format used by the SAPRC model simulation software, which is not as widely used as other modeling software systems. This paper, along with the recently published paper of Carter et al. (2025a), addresses its lack of adequate documentation. Future updates to MechGen will be documented and made available as separate versions to ensure reproducibility. We are developing software converting MechGen mechanism output to other formats and have already released a converter for the FOAM modeling system (Wolfe et al, 2016), which is available at from the MechGen GitHub repository.

MechGen, along with GECKO-A and MCM, represent the current state of the science in atmospheric reactions of organics in the lower troposphere, though they all require updates in some respects to be consistent with recent advances in the continuously evolving science in this area. They all incorporate different estimates or assumptions regarding some uncertain processes, though they also have many assumptions and estimates in common that may not necessarily be correct. The assignments and estimates incorporated in MechGen that need to be updated are discussed by Carter et al. (2025a). Although GECKO-A is also continually being updated, it does not yet incorporate H-shift isomerization (including autooxidation) reactions of peroxy radicals, which are important for many compounds such as monoterpenes (Li et al., 2022). On the other hand, GECKO-A has a more detailed representation of peroxy + peroxy and photolysis reactions. Comparing MechGen predictions with those of GECKO-A and MCM can help identify areas where further experimental, theoretical, or SAR development work is most needed, though this does not rule out the possibility that they all share assumptions or estimates for these highly complex and uncertain mechanisms that may in fact be incorrect.

The atmospheric chemistry of organics and development of SARs or estimation methods continues to be an active area of research, and MechGen and other mechanism generation systems need to be periodically updated to continue to represent the state of the science and thus continue to be appropriate tools for developing mechanisms for models. Updating the estimates for the many types of chemical systems that need to be represented, testing the associated software changes, and evaluating the predictions against available data, is a time-consuming process, making it essentially impossible for any system to be completely up to date at any point of time. This is one reason why comparisons of predictions of different mechanism generation systems or mechanisms are important. Carter et al. (2025a) discuss the many types of estimates and points out the areas where updates are needed and planned.

MechGen is designed to accommodate updates to the underlying chemistry assignments and SARs. Modifying chemical assignments or parameters used in existing SARs is straightforward and discussed in the User's Manual, but adding or deleting types of reactions or adding or changing the structures of SARs will require programming changes. Since implementing such updates would result in new versions of the system that will give different predictions, it is important that older versions be archived and made available so that previous work that used these versions can be reproduced or evaluated. This was not the case for previous versions of MechGen, but will be for newer versions going forward.

The development and maintenance of MechGen has largely been the work of the primary author (Carter), who is nearing full retirement. Unique features and strengths of MechGen (including its integration with other SAPRC tools and models) have led to a growing base of MechGen users, and as this user base has grown, so has the need for a collaborative team to carry it into the future. Steps to ensure the sustainability of this resource thus far have included the publication of the chemical basis of the MechGen system (Carter et al., 2025a), this paper describing the software system, the development of a GitHub page (<https://github.com/SAPRC/MechGen>), publications describing the use of MechGen beyond the derivation of SARPC mechanisms (Jiang et al., 2020; Li et al., 2022), and dissemination of

knowledge through the growing user community. In addition, the MechGen is written in a programming language (MOO) that is no longer widely used or maintained, so conversion to a more actively supported platform would enhance ongoing maintenance and collaborative development in the future. The development of collaborative teams, including those working on the chemical mechanism estimates and databases, ~~and on~~ as well as the software, will ensure that MechGen remains a valuable, adaptable, and robust mechanism generation system well into the future.

SUPPLEMENTARY INFORMATION

The ~~supplement related to this article is available online at: [give URL when published].~~ ~~This~~Supplementary information published with this manuscript contains additional information ~~or~~and details on procedures discussed this paper. These include the following: additional output examples (Section S1); derivation of the standard environments currently available in MechGen (Section S2); additional details concerning procedures used (Section S3); the method used to estimate fractions of semi-volatile products in the ~~condensed~~condensed phase (Section S4); additional information regarding multi-generation mechanisms (Section S5); and additional information on the results with the representative compounds (Section S6).

USER MANUAL

A complete User Manual describing the operation of the MechGen system using both the web and terminal interfaces is available with the MechGen source code (see below) and at the MechGen web site (Carter, 2025) andSAPRC MechGen GitHub page (<https://github.com/SAPRC/MechGen/tree/master/docs>). It discusses how to obtain, install, and configure the system on user's own computer. Updated versions of the user manual will be made available on the MechGen and GitHub sites when the source code or documentation has been updated. A quick start web users guide for new users is also available at the MechGen web site and the SAPRC MechGen GitHub page, where it will also be updated when appropriate.

CODE AND DATA AVAILABILITY

The MechGen source code (~~v1.0,~~v1.1, last updated ~~February 21,~~July, 25, 2025) and its documentation, including the full User Manual and Quick Start Guide, are available on Zenodo (<https://zenodo.org/records/14907949>)-via Carter et al. (2025b) at <https://zenodo.org/records/16622705>. Detailed system information, including online access and further development, is provided on the MechGen website (<https://mechgen.cert.ucr.edu/>), while the GitHub repository at <https://github.com/SAPRC/MechGen> offers the latest software releases and installation resources, with updates archived on Zenodo-~~(<https://zenodo.org/records/14907948>)~~.

All the source code and data used by MechGen are contained within a single MOO code database file as described in available MOO documentation (Curtis, 1997; Fox, 2004; Wikipedia, 2025). The program source code and the data can be viewed and modified by downloading and installing the MechGen database and the MOO server software as described in the User Manual. Once the system is installed and configured, the user can access the system using Telnet and log in as a systems programmer (see User Manual) and use MOO commands to view or modify the source code (Curtis, 1997; Fox, 2024)-2004).

The data used by MechGen to implement the chemical assignments and SARs are described by Carter et al., (2025)-al. (2025a). The detailed parameters, mechanism assignments, and rate constants used are given in tables in the paper or its Supplementary Information.

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