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Modelling the evolution of organic carbon during its gas-phase tropospheric oxidation: development of an explicit model based on a self generating approach

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Abstract. Organic compounds emitted in the atmosphere are oxidized in complex reaction sequences that produce a myriad of intermediates. Although the cumulative importance of these organic intermediates is widely acknowledged, there is still a critical lack of information concerning the detailed composition of the highly functionalized secondary organics in the gas and condensed phases. The evaluation of their impacts on pollution episodes, climate, and the tropospheric oxidizing capacity requires modelling tools that track the identity and reactivity of organic carbon in the various phases down to the ultimate oxidation products, CO and $CO₂$. However, a fully detailed representation of the atmospheric transformations of organic compounds involves a very large number of intermediate species, far in excess of the number that can be reasonably written manually. This paper describes (1) the development of a data processing tool to generate the explicit gas-phase oxidation schemes of acyclic hydrocarbons and their oxidation products under tropospheric conditions and (2) the protocol used to select the reaction products and the rate constants. Results are presented using the fully explicit oxidation schemes generated for two test species: n-heptane and isoprene. Comparisons with wellestablished mechanisms were performed to evaluate these generated schemes. Some preliminary results describing the gradual change of organic carbon during the oxidation of a given parent compound are presented.

1 Introduction

Volatile organic compounds (VOCs) emitted in the atmosphere are oxidized in complex reaction sequences that produce a myriad of intermediates. The intermediates are typically organic radicals involved in radical oxidation chains (peroxy, oxy and Criegee radicals) and oxygenated secondary (i.e. produced in situ) species such as alcohols, aldehydes, ketones, nitrates, peroxides, carboxylic acids and peracids (e.g. Atkinson, 2000). These intermediates play a key role in the chemistry of the troposphere. For example, these organic intermediates are directly involved in the production of tropospheric ozone and in the HO_x ($OH + HO_2$) and $NO_x (NO + NO₂)$ budgets on regional and global scales (e.g. Seinfeld and Pandis, 1998; Finlayson-Pitts and Pitts, 2000, and references therein). Furthermore, many secondary organic species are multifunctional with low vapour pressure. These semi-volatile compounds lead to the production of secondary organic aerosol (SOA) by nucleation and/or condensation on pre-existing aerosols (e.g. Pankow et al., 2001; Hoffmann et al., 1997; Odum et al., 1997; Bowman et al., 1997) and/or polymerisation (Kalberer et al., 2004; Tolocka et al., 2004). The SOA components can contribute to the absorption and scattering of solar radiation (Jacobson, 1999), as well as affect the hygroscopic behaviour of particles (Saxena et al., 1995) and therefore their ability to act as cloud condensation nuclei by changes in surface tension (Facchini et al., 1999, 2000) or growth rates of cloud droplets (Lohmann et al., 2004; Shantz et al., 2003). Most secondary VOCs are water soluble, i.e. associated with high Henry's law coefficients. A significant fraction of secondary organics thus dissolves in the aqueous droplets during cloud

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Fig. 1. Simplified representation of the reaction sequences involved in the tropospheric VOC oxidation.

events. This dissolved organic matter directly or indirectly interacts with the aqueous chemistry of radical, non metal and metal ion and oxidant pools (e.g. Faust, 1994; Ervens et al., 2003). Organic chemistry inside clouds may therefore modify the chemical budget of both aqueous and gas phases. For example, some organics can have an inhibiting effect on aqueous Fe-catalyzed SIV oxidation (Wolf et al., 2000), and aqueous VOC oxidation can have a significant impact on the tropospheric ozone budget (Monod et al., 1999). Organic oxidation inside clouds may also be an additional source of SOA, after the evaporation of the cloud droplets (Blando and Turpin, 2000). Clearly, these and other processes involving organic intermediates may have significant effects on atmospheric reactivity, composition, and radiative budgets on wide-ranging geographic scales. However, the magnitude of these effects remains poorly understood, due largely to a critical lack of information concerning the detailed composition of the highly functionalized secondary organics in the gas and condensed phases.

Three-dimensional chemistry-transport models (e.g. those used for studying pollution episodes, regional and global oxidants, and climate) typically represent organic chemistry with highly simplified mechanisms that include the initial reactions of a few hydrocarbons and only a very limited subset of surrogate or lumped species to represent the intermediates (e.g. Gery et al., 1989; Stockwell et al., 1990, 1997; Bey et al., 2001; Brasseur et al., 1998; Poisson et al., 2000). These simplifications are frequently adequate to represent the nearfield gas phase reactivity (e.g. ozone formation in the vicinity of emission sources), but fall short of providing the chemical speciation necessary for a detailed understanding of the gas phase composition of aging air parcels, and of the contributions that organic intermediates make to the composition and properties of aerosols and clouds. In the past decade, major advances have been made towards the development of more explicit hydrocarbon oxidation mechanisms, such as the NCAR Master Mechanism (Madronich and Calvert, 1990; Aumont et al., 2000) with over 5000 reactions to represent the oxidation of about 20 emitted non-methane hydrocarbon, the SAPRC-99 mechanisms which describes the oxidation of about 400 emitted VOC (Carter, 2000), and the University of Leeds' Master Chemical Mechanism (MCM) with over 13000 reactions to describe the oxidation of about 100 emitted VOC (Jenkin et al., 1997, 2003; Saunders et al., 1997, 2003b). But even these large mechanisms neglect many reaction pathways by assuming that they individually make only a minor contribution, though it remains unclear whether their cumulative contributions could be important.

The fully explicit representation of hydrocarbon oxidation, from the initial compounds to the final product $CO₂$, requires a very large number of chemical reactions and intermediate species, far in excess of the number that can be reasonably written manually. Nevertheless, several potential benefits from the availability of such full mechanisms can be identified. Knowledge of the precise chemical identity of intermediates enables the calculation of their atmospheric lifetimes and thus of their contribution to the gas phase reactivity of aging air masses (e.g. HO_x and NO_x budgets). Their vapour pressure and solubility can also be calculated and used to determine their partitioning to condensed phases, ultimately providing better estimates of the optical and microphysical properties of aerosols and clouds. Moreover, a fully explicit mechanism can serve as a benchmark for the systematic development and testing of simplified chemical mechanism for use in three-dimensional models where computer resources are limited. Finally, an explicit modelling approach offers increased opportunities for comparison with atmospheric measurements, particularly in view of recent advances in techniques for detecting many specific organic compounds.

Here, we describe a "self generating approach" to develop fully explicit gas-phase oxidation schemes of organic compounds under general tropospheric conditions. This approach was codified in a computer program, hereafter referred as the "generator" of the chemical schemes. The generator and the protocol used to select the reactions and the rate constants are similar to the approaches described by Jenkin et al. (1997) and Saunders et al. (2003a). The generator and the protocol are presented in Sects. 2 and 3, respectively. Section 4 presents some tests performed to evaluate the performance of the generated scheme by comparison with existing mechanisms. Finally, some preliminary results showing the major chemical characteristics of organic carbon during the oxidation of a given parent compound are presented (Sect. 5).

2 The generator

The construction of the chemical schemes generator is based on identifying all the reactions for a given primary (i.e.

Fig. 2. Flow diagram of the generator.

emitted) species as well as for every intermediate up to CO or CO² production. Atmospheric oxidation of organic compounds can be represented by a limited number of reaction types, repeated many times up to full oxidation of the given parent compound. With a few exceptions, these steps can be summarized as follow (see Fig. 1):

- **–** The initiation of the radical organic chains by reaction with OH, $NO₃$, $O₃$ or via bond breaking after the absorption of a photon. This first step generally leads to the formation of an organic peroxy radical $(RO₂)$.
- **–** Peroxy radical reactions with NO, NO2, NO3, HO² or with others $RO₂$ radicals. These reactions either lead to a stable (i.e. a non radical) secondary organic species or to an organic alkoxy radical (RO).

 $-$ Alkoxy radical reaction with O_2 , unimolecular decomposition (i.e. C-C bond breaking) or isomerization. These reactions either lead to a stable secondary and/or to a new peroxy radical.

The redundancy in these reaction sequences is the basis of the generator.

The general philosophy followed to develop the generator is similar to the methods recently described by Saunders et al. (2003a). The generator can be viewed as a computer program that mimics the steps by which chemists might develop chemical schemes (i.e. writing the list of the reactions involved in the oxidation of a given species and their associated rate constants). These steps are illustrated Fig. 2 and can be summarized as follow. First the generator analyses the chemical structure of the molecule to identify the reactive sites and determine all reaction pathways. For each identified

Fig. 3. Description of molecular structures in the generator: the methacrolein example.

reaction, the generator then "searches" into a database of laboratory measurements of reaction rate constants and/or products to see if they may already be available. If not (as it is usually the case for the oxidation intermediates), an estimation of the rate constant and reaction products is made based on structure/activity relationships (see Sect. 3 below). The full reaction (reactants, products, stoichiometric coefficients, rate constant, and its temperature dependence) is then written in a file and the products, if new, are added to a "stack" for further reactions. These steps are repeated as long as species are present in the stack. The process stops when the stack is empty, which occurs when the full oxidation scheme has been written.

The molecular structure of the species needs to be known by the generator to select the appropriate set of reactions. This molecular structure is described using a matrix and a vector (see Fig. 3). The bond matrix provides the carbon skeleton of the molecule (each carbon is identified by an index; a bond between the i th and j th carbon is marked with non-zero values in the (i, j) cell of the bond matrix). The group vector holds the information about the functional groups borne by each carbon. To avoid duplications, care is taken that each molecule (and its corresponding bond matrix and group vector) is written in a unique way based on the longest carbon chain and a pre-defined priority order for the functional groups.

The chemical schemes written by the generator are not conceptually different than those that could be written manually given enough time. However, several important advantages of the generator code should be noted:

- **–** Its speed: explicit chemical schemes usually contain thousands of reactions and handwriting such schemes requires months of work. In fact, fully explicit chemical schemes are extremely large (see examples below) and the self-generating approach is probably the only realistic approach to develop such fully explicit schemes.
- **–** Accuracy: it is almost impossible to write large schemes without introducing "errors" (for example in setting the rate constants or simply typographical errors in naming the species).

– Maintenance: updating chemical scheme can be a very tedious task. This is particularly true when the scheme must be updated as a result of a change in a structure/activity relationship. Updating the generator code is relatively straightforward and the new mechanism is easily re-generated.

The generator can therefore be viewed as an expert system, assimilating physico-chemical data from laboratory experiments (e.g. rate constants, vapor pressure, products yields) and estimating all the missing information. It is a very natural link between fundamental laboratory studies and the chemical models.

The current version of the generator treats the chemistry of every species (including C, H, O, N atoms) except cyclic molecules. This includes alkanes (linear or branched), alkenes (conjugated or not), ethers, esters, alcohols, carbonyls, nitrates, peroxyacylnitrates, carboxylic acids, peracid, hydroperoxides, alkyl radicals, peroxy radicals and alkoxy radicals. The chemistry of cyclic molecules is currently read in a file (up to ring-opening) and corresponding mechanisms are added to the generator output file (see Sect. 3.10 below). The generator's code has been fully written in Fortran 77.

Despite the fairly limited number of reaction types implemented in the generator, the total number of species generated to describe the full oxidation of a given parent compound can be extremely large. Figure 4 shows the total number of species generated (hereafter denoted N_g) for the n-alkanes series as a function of the number of carbons in the parent compound. N_g grows from a few hundreds for propane to about 2.5 million species for octane. The increase follows an exponential law, with a growth factor of about 5.6 for each carbon added.

The chemistry codified in the generator can only lead to a limited number of functionalities. For the n-alkane series, possible groups for each carbon are: non-substituted alkyl, ketone (or aldehyde), alcohol, hydroperoxide, nitrate, peroxy radical and alkoxy radical. Four additional groups must be considered for the carbons at a terminal position viz.: carboxylic acid, peracid, peroxy acyl nitrate and peroxy acyl.

Fig. 4. Number of species created by the generator for the n-alkane series without a threshold for selecting reactions (cross) and with a 5% branching threshold (circle). The line shows the maximum number of distinct species that can be produced for the n-alkane series by permuting all the possible functional groups codified in the generator.

Furthermore, we did not identify chemical pathways leading to a carbon bearing more than one functionality. Thus for the n-alkane series, 11 and 7 distinct group substitutions can be identified for the terminal and internal carbon positions, respectively. Figure 4 shows the total number of "distinct" species that can be produced by permuting all the possible groups. For fairly large molecules (i.e. C>4), the total number of distinct species (denoted Ω below) can be very well approximated by:

$$
\Omega \approx \sum_{i=2}^{n} \frac{1}{2} (11)^2 (7)^{i-2},\tag{1}
$$

where n is the total number of carbons in the parent compound, the $\frac{1}{2}$ factor corrects approximately for symmetries, and the sum over i takes into account the possible scissions of C-C bonds. The exponential growth of N_g as a function of the chain length can thus be explained by the exponential increase of Ω . The exponential growth factor of 5.6 found for the n-alkane series is however somewhat lower than the growth factor of \approx 7 predicted by Eq. (1). Some group distributions can indeed not be produced, due for example to C−C bond scission before reaching them. Note however that for n-octane the total number of species produced by the generator still represents about 30% of all the possible functional group permutations (see Fig. 4). Similar results were found for other series (e.g. i-alkanes, 1-alkenes): the key factor determining the number of species and reactions generated is always the total number of carbon atoms in the parent molecule (see Fig. 5). Additional tests were performed to evaluate the sensitivity of N_g to a given constrain for the activation of a particular reaction. Figure 4 shows that N_g remains extremely large even if reactions contributing for less

Fig. 5. Number of species (open symbols) and reactions (grey symbols) created by the generator for various series (using a 5% threshold for selecting reactions).

Fig. 6. Number of species generated as a function of the generation depth (circle, left scale) and species in the generator's stack (cross, right scale) for n-octane.

than 5% in the removal of a given species are ignored. We finally note that the number of the generated reactions is typically 10 times the number of species, i.e. reaching 25 million for the full oxidation of octane. Various tests were performed to reduce the number of species and reactions in the chemical scheme for 3-D applications. These tests are described in a companion paper (Szopa et al., 2005).

The number of generations (i.e. of successive non radical intermediates, see Fig. 1) required to fully oxidize a given parent compound was found to be slightly larger than the number of carbons in the parent compound. The main factor controlling this behaviour is that, on average, at least one generation is required per carbon to oxidize it from $-CH₂$ − to −CO−. Figure 6 shows the number of species generated for a given generation depth as well as the species remaining in the stack (i.e. species for which removal reactions still have to be written) for the n-octane case. The S-shaped and Gaussian profiles shown Fig. 6 were also found for the other species. These profiles can qualitatively be explained as follow. Each species in the generation n leads to the production of a given number of species in the generation $n + 1$ after reactions. For the first generations, these reaction products are mainly new species for which chemistry must be written. The total number of species as well as the number of species in the stack thus grows exponentially. However, a given reaction product can usually be produced by several distinct reaction pathways and must obviously be treated only once. Hence, after few generations, most reaction products are already treated in the scheme (i.e. other pathways already led to the same products) and the species is not added to the stack. The number of species in the stack starts to decrease, as well as the growth rate of the total number of generated species. As stated above, the chemical pathways involved in the full oxidation of every generated species are described when the stack is empty.

3 Protocol description for the development of the organic schemes

Explicit chemical schemes would obviously be a faithful representation of oxidation mechanism occurring in the atmosphere if all reactions were known with a high accuracy. From the above section, it is clear that the number of species involved in such schemes far exceeds that for which direct measurements are available. Reaction pathways and rate constants must therefore be estimated for most species on the basis of their molecular structures. Numerous experimental data are currently available in the literature concerning the VOC oxidation. Compilations, data evaluations and recommendations for use in models are regularly provided by the NASA and IUPAC panels (e.g. DeMore et al., 1997; Atkinson et al., 1999) or in review papers (e.g. Atkinson, 1994; Atkinson and Arey, 2003; Orlando et al., 2003; Mellouki et al., 2003; Calvert et al., 2000). Reasonable estimates can be made by analogy to these known reactions, with the aid of several structure/activity relationships (SARs). The main feature of the generator described above is to codify the various estimation methods to produce consistent and comprehensive oxidation schemes on a systematic basis. We emphasize that the accuracy of the generated explicit schemes is linked directly to the accuracy of SAR predictions. Most rate constants are typically estimated with at least a factor 2 of uncertainty and some reaction pathways are highly hypothetical due to the lack of experimental data. Given these uncertainties, explicit organic schemes must rather be viewed as useful exploratory tools reflecting the current understanding of VOC oxidation than a new more precise representation of it. Note, however, that whenever the products and kinetics of a specific reaction are known directly from laboratory measurements, the generator gives priority to these experimental data rather than attempting to recalculate them from SARs.

Table 1. List of the organic radicals at steady state and their associated reaction products.

Reaction
$O2$ addition
$CH_3(\cdot) + O_2 \rightarrow CH_3(OO \cdot)$
$-CH_2(\cdot) + O_2 \rightarrow -CH_2(OO \cdot)$
$>CH(\cdot) + O_2 \rightarrow CH(OO \cdot)$
$>C(\cdot) - +O_2 \rightarrow C(OO)$
$-C(·)(0) + O_2 \rightarrow -C(0)(00)$
$>C=C(\cdot)-+O_2 \rightarrow BC=C(OO \cdot)$
Decomposition
$HC(·)O (+O2) \rightarrow CO + HO2$
$CH_2(\cdot)(OH)$ $(+O_2) \rightarrow HO_2 + HCHO$
$CH_2(\cdot)$ (OOH) \rightarrow OH + HCHO
$CH_2(\cdot)(ONO_2) \rightarrow NO_2 + HCHO$
$RCH(\cdot)(OOH) \rightarrow RCHO + OH$
$RCH(\cdot)(OH)$ $(+O_2) \rightarrow RCHO + HO_2$
$RCH(\cdot)(ONO_2) \rightarrow RCHO + NO_2$
$RR'C(\cdot)(OOH) \rightarrow RCOR' + OH$
$RR'C(\cdot)(OH)$ $(+O_2) \rightarrow RCOR' + HO_2$
$RR'C(\cdot)(ONO_2) \rightarrow RCOR' + NO_2$
$C(\cdot)(O)(OH) \rightarrow OH + CO$
$C(\cdot)(O)(OOH) \rightarrow HO_2 + CO$
$C(\cdot)(O)(ONO_2) \rightarrow NO_3 + CO$
$C(\cdot)(O)(OONO_2) \rightarrow NO_2 + CO + O_2$
$(O·)CO(OH) \rightarrow CO2 + OH$
$(O·)C(O)(OOH) \rightarrow HO2 + CO2$
$(O·)C(O)(ONO2) \rightarrow NO3 + CO2$
$(O \cdot) C(O)(O O NO2) \rightarrow NO2 + CO2 + O2$
$(O·)C(O)H (+O2) \rightarrow HO2 + CO2$
$>C=C(OO)$ - \rightarrow $>C(O)$ - CO -
$>C=C(O)$ - \rightarrow $-C(\cdot)$ -CO-
$R-CO(O.) \rightarrow CO_2 + R.$
$HC(O)(O \cdot) (+O_2) \rightarrow CO_2 + HO_2$
$R-O-C(O)(OO \cdot)$ $(+O2) \rightarrow R(OO \cdot) + CO2 + O2$
$R-CO-C(·)(O) \rightarrow R-C(·)O + CO$
$-C(\cdot)(OH)(OO\cdot)$ $(+O_2) \rightarrow HO_2 + -C(O)(OO\cdot)$
$-C(\cdot)(OOH)(OO\cdot) \rightarrow OH + -C(O)(OO\cdot)$
$-C(\cdot)(ONO_2)(OO\cdot) \rightarrow NO_2 + -CO(OO\cdot)$

The protocol used in this study is very similar to the approaches already described by Jenkin et al. (1997), Saunders et al. (2003b) and Carter (2000) for the purpose of systematic development of organic schemes. Here we present only a brief overview of the protocol as implemented in the generator. A more detailed discussion concerning the rules used to set the chemical pathways as well as the SAR selected to estimate their associated rate constants can be found in Laval-Szopa (2003), based on data published before 2003. Inorganic and C_1 organic chemistry (i.e. methane oxidation) is out of the frame of the protocol presented here. Direct measurements are in fact available for these species and estimates are not necessary. Rate constants for the inorganic

Structures	k_{add} molecule ^{-1} cm ³ s ^{-1}	Branching ratio	Reference compound	Notes			
alkenes							
$CH2=CH-$	1.4×10^{-14}	1.0	1-butene	a, b			
$CH2=CH2$	3.3×10^{-13}	1.0	<i>i</i> -butene	a, b			
$-CH=CH-$	3.7×10^{-13}	0.5	2-butene	a, b, c			
$-CH=C<$	9.4×10^{-12}	1.0	2-methyl-2-butene	a, b			
$>C=C<$	5.7×10^{-11}	0.5	2,3-dimethyl-2-butene	a, b			
dienes							
$CH2=CH-CH=CH-$	1.5×10^{-12}	1.0	pentadiene	a, f, g			
$-CH=CH-CH=CH-$	1.6×10^{-11}	0.5	2,4-hexadiene	a, e, f, g			
unsaturated carbonyls							
$CH2=C(R)CO-$	1.4×10^{-15}	1.0	methacrolein	b, h			
$-$ CH=CHCO $-$	8.5×10^{-15}	0.5	crotonaldehyde	b, h			
$>CH=CHCO-$	2.1×10^{-13}	1.0		b, i			
$-CH=C(R)CO-$	1.2×10^{-13}	1.0		b, j			
$>CH=C(R)CO-$	7.7×10^{-13}	0.5		b, k			

Table 2. Generic rate constants and branching ratios for alkene $+NO₃$ reactions.

^a rate constants from Atkinson (1997); ^b branching ratio for NO₃ addition at the less substituted carbon; ^c average value of cis- and trans-2butene; e also used for >C=CH−CH=CH− and >C=CH−CH=CH< structures; f if the diene is alkyl substitued or disubstitued in position 2 or 3, the rate constant is multiplied by a factor 6.8 and 21, respectively, based on the isoprene/butadiene and 2,3-dimethyl-butadiene/butaniene rate constant ratios; ^g branching ratio for NO₃ addition at the less substituted terminal carbon (i.e. position 1 or 4). In the resulting allylic structure, the radical center is located on the most substituted carbon; h see text for references of the rate constants; $ⁱ$ rate constant for</sup></sup> crotonaldehyde multiplied by the −CH=C< /-CH=CH− rate constant ratio; j average rate for crotonaldehyde and methacrolein multiplied by the $-CH=C<$ /CH₂=CH< and >C=C−/−C=C− rate constant ratios, respectively; k average rate for crotonaldehyde and methacrolein multiplied by the $-CH=C<$ /CH₂=CH< and >C=C−/-C=C− rate constant ratio, respectively.

reactions are based on NASA and IUPAC recommendations (Sander et al., 2000; Atkinson et al., 1997) as well as on the recent recommendations of Troe (2001) for the $NO₂ + OH \rightarrow HNO₃$ reaction. Methane oxidation is based on the IUPAC recommendations (Atkinson et al., 1999) and the recommendations of Tyndall et al. (2001) for the methyl peroxy reactions. In the following subsections, we first present the management of radical species within the generator, then reactions associated with non-radical organic species (i.e. reactions with OH, $NO₃$, $O₃$, thermal decomposition and photolysis) and finally reactions associated with the organic radicals (i.e. the peroxy, alkoxy and Criegee radicals). The particular case of cyclic molecules is presented last.

3.1 Radical check in the generator

To decrease the number of species as well as the stiffness of the set of ordinary differential equations that must ultimately be solved, the steady state approximation is used for highly reactive species having a unique significant reaction pathway. Such species are not considered in the chemical scheme but are directly replaced by their reaction products. For example, the OH reaction with a paraffin carbon (RH) is written as $RH + OH(+O₂) \rightarrow RO₂ + H₂O$, in which the alkyl radical $(R \cdot)$ first produced by the reaction with OH is directly replaced by the peroxy radical $(RO₂)$, its only significant reaction product under tropospheric conditions. Organic radical moieties for which the steady state approximation has been systematically applied are listed in the Table 1. The only exception to the list presented in the Table 1 is the HCO−CO(·) radical which was experimentally found to react both by decomposition and O_2 reaction. We used the yields provided by Orlando and Tyndall (2001) for this radical. Note that the reactions listed in the Table 1 for peroxy and alkoxy radicals supersede the rules described in the Sects. 3.7 and 3.8 for these types of radicals.

3.2 VOC + OH reactions

Rate constants for H atom abstraction are estimated using the SAR of Kwok and Atkinson (1995), in which we added a group contribution for the hydroperoxide moiety (−OOH). Based on the IUPAC recommendations for CH3−OOH (Atkinson et al., 1999), an activation factor of 14 at 298 K was set for H abstraction to a carbon bearing the −OOH functionality and the rate constant for H abstraction of the −OOH moiety was set to 1.9×10^{-12} exp(190/T) molecule⁻¹ cm³ s⁻¹. OH additions

Table 3. Generic rate constants and branching ratios for alkene+ O_3 reactions.

Structures	Rate constant molecule ^{-1} cm ³ s ^{-1}	Branching ratio	Notes					
alkenes								
$CH2=CH-$	1.0×10^{-17}	0.5/0.5	a, b					
$CH2=C5$	1.2×10^{-17}	0.65/0.35	a, b					
$-CH=CH-$	1.1×10^{-16}	0.5/0.5	a, b					
$-CH=C<$	3.5×10^{-16}	0.7/0.3	a, b					
$>C=C<$	1.1×10^{-15}	0.5/0.5	c, b					
dienes								
$-CH=CH-CH=CH2$	3.5×10^{-17}	0.25 for each carbon	d, e, f					
$>C=CH-CH=CH2$	3.9×10^{-17}	0.25 for each carbon	d, e, f, g					
$-C=CH-CH=CH-$; $>C=CH-CH=CH-$	3.4×10^{-16}	0.25 for each carbon	h, e, f					
$>C=CH-CH=C<$	3.1×10^{-15}	0.25 for each carbon	i, e, f					
unsaturated carbonyls								
$C=C-CHO$	2.9×10^{-19}	0.9/01	j, l					
$C=C-CO-$	5.4×10^{-18}	0.95/0.05	k, l					
$HCO-C=C-CHO$	1.6×10^{-18}	0.5/0.5	m, e					
$HCO-C=C-CO-R$	4.8×10^{-18}	0.5/0.5	m, e					
R –CO–C=C–CO–R	3.6×10^{-18}	0.5/0.5	m, e					

 a Rate constants from Carter (2000), average values at 300 K for various species having the appropriate structure; b branching ratio from Carter (2000). If not equal, the most substituted criegee radical is formed with the highest yield; c rate constant for 2,3-dimethyl-2-butene, Atkinson (1997); d average rate constant for (E) and (Z) 1,3 pentadiene, Atkinson (1997); e no experimental data available for branching ratio. An equal probability is set to each pathway; f rate constant is enhanced by a factor 2 for each alkyl substitution at position 2 or 3 of the diene structure. Enhancement factor based on the isoprene/butadiene and 2,3-dimethyl-butadiene/isoprene rate constant ratio. g rate constant for 1,3 pentadiene enhanced by the rate constant ratio isobutene/propene; h rate constant for 2,4 hexadiene, Atkinson (1997); i rate constant for 2,5-dimethyl-2,4-hexadiene, Lewin et al. (2001); j rate constant for acrolein, see text; k rate constant for methyl vinyl ketone, see text; ^l rate constant enhanced by a factor of 3.9 and 6.0 for each H substitution by an alkyl group in α or β of the carbonyl group, see text; m based on the rate constants provided by Liu et al. (1999).</sup>

to >C=C< bonds are estimated using the SAR of Peeters et al. (1997), except for alkenes having conjugated carbonyl groups. For these unsaturated carbonyls, the SAR of Kwok and Atkinson (1995) was used. We used a branching ratio of 0.2 and 0.8 for OH addition to the carbon in α and β position relative to the carbonyl group, respectively, based on the yields provided by Tuazon and Atkinson (1989, 1990) for methacrolein and methyl-vinyl-ketone. For 1,4 and 1,6 unsaturated dicarbonyls, branching ratio for OH addition to each olefin carbon was set with an equal probability.

3.3 VOC + $NO₃$ reactions

Two VOC $+$ NO₃ reaction types are considered: H abstraction from aldehydes and $NO₃$ addition to alkenes.

Experimental data, including rate constants, are available for aldehydes of low molecular weight. The rate for the butanal+ $NO₃$ reaction was used as generic rate for unknown $RCHO + NO₃$ reactions. The rate constant was set to 1.22×10^{-14} molecule⁻¹ cm³ s⁻¹, based on the data provided by D'Anna et al. (2001). The rate constants for H- abstraction on conjugated dicarbonyls are estimated using the method of Carter (2000), leading to a rate of 4.8×10^{-16} and 2.4×10^{-15} molecule⁻¹ cm³ s⁻¹ for CHO–CHO and RCO−CHO, respectively. According to Atkinson (1991), $NO₃$ reaction with acrolein (CH₂=CH–CHO) mainly occurs by H-abstraction at the aldehyde group. The rate constant for H-abstraction in >C=C−CHO structures was set to 2.5×10^{-15} molecule⁻¹ cm³ s⁻¹, based on the rate constant provided by Cabanas et al. (2001) for the acrolein+NO₃ reaction.

Reaction rates for $NO₃$ addition to non-conjugated alkenes are evaluated using the method of Jenkin et al. (1997) and Carter (2000). The generic constants as well as the branching ratio used in this study are summarized in Table 2. This method was extended to conjugated alkenes (see Table 2). For conjugated unsaturated carbonyls, reaction rates are available at room temperature for (1) methyl vinyl ketone: $k=1.2\times10^{-16}$ (Rudich et al., 1996), (2) methacrolein: $k=3.9\times10^{-15}$ (average value of the rate constants provided by Chew et al., 1998; Kwok et al., 1996), and (3) crotonaldehyde: $k=1.1\times10^{-14}$ molecule⁻¹ cm³ s⁻¹ (average value of

the rate constants provided by Cabanas et al., 2001; Atkinson, 1994). Using a typical nocturnal $NO₃$ concentration of 10^9 molecule cm⁻³, the lifetime of methyl vinyl ketone with respect to $NO₃$ reaction is estimated to be of about 100 days, i.e. well above the corresponding lifetime due to OH removal $(\approx 10 h)$. We therefore did not considered the NO₃ addition to CH₂=CH–CO– structures. Note that this very slow reaction rate is in agreement with the expected minor contribution of $NO₃$ addition to the C=C bond pointed above for acrolein. Furthermore, assuming that H atom abstraction from the −CHO group contributes 2.5×10^{-15} to the total rate constant (see above), the rate for $NO₃$ addition to C=C bond can be estimated to 1.4×10^{-15} and 8.5×10^{-15} molecule⁻¹ cm³ s⁻¹ for methacrolein and crotonaldehyde, respectively. These rate constants were (1) directly used as generic constants for $CH₂=C(R)-CO-$ and R−CH=CH−CO− structures, and (2) combined with the generic constants for $NO₃$ addition to simple alkenes to evaluate NO₃ addition rates for more substituted structures. Rules used for this last evaluation are given in Table 2. Finally, no data are available to our knowledge concerning the reaction rates of $NO₃$ reaction with 1,4 unsaturated dicarbonyls. The data listed Table 2 suggest however that the substitution of an alkyl group with a carbonyl group typically decreases the rate constant by 2 orders of magnitude. For 1,4 unsaturated dicarbonyl, we therefore scaled by a factor of 10−² the generic constant provided in Table 2 for unsaturated carbonyls. Although crude, the evaluation suggests very low rate constants for the $NO₃$ reaction with 1,4 unsaturated dicarbonyl and these reactions should not be a significant loss process under tropospheric conditions.

3.4 VOC + O_3 reactions

Rate constant estimations for the reaction of non-conjugated alkenes with O_3 are based on the method of Jenkin et al. (1997) and Carter (2000). Generic constants as well as the branching ratios for carbonyl and Criegee radical formation at both side of the $>C=C<$ bond are given in the Table 3. This method was extended to conjugated dienes (see Table 3). Evaluations for unsaturated carbonyl were based on the following rate constants at room temperature $(in molecule⁻¹ cm³ s⁻¹ units):$

- **–** acrolein: k=2.9×10−¹⁹ (Atkinson, 1994);
- **–** methyl vinyl ketone: k=5.4×10−¹⁸ (Neeb et al., 1998)
- **–** methacrolein: k=1.1×10−¹⁸ (Atkinson, 1994)
- **–** crotonaldehyde: k=1.7×10−¹⁸ (Grosjean and Grosjean, 1998)

Rates for acrolein and methyl vinyl ketone were used as generic constants for unsaturated aldehydes and ketones, respectively. Contribution of alkyl groups in α and β positions hancement factor of 3.9 and 6.0, respectively, based on the rate constant ratio of methacrolein and crotonaldehyde with respect to acrolein. Rate constants provided by Liu et al. (1999) were used as generic values for 1,4 unsaturated dicarbonyls (see Table 3). O_3 reactions with 1,6 di-unsaturated dicarbonyls are of the order of 10^{-19} molecule⁻¹ cm³ s⁻¹ Klotz et al. (1995). Using a typical O_3 concentration of 10¹² molecule cm−³ , the lifetime for 1,6 diunsaturated dicarbonyls with respect to O_3 reaction is estimated to be about $10²$ days, i.e. well above the corresponding lifetime due to OH removal (\approx 10 h). The reaction of O₃ with 1,6 diunsaturated dicarbonyls was therefore not considered.

relative to the carbonyl were taken into account using an en-

3.5 VOC photolysis

Photolytic reactions are considered for 3 types of chromophores: carbonyl, hydroperoxide, and nitrate. The cross sections and quantum yields of 54 species are tabulated in the generator (see list in Laval-Szopa, 2003). Some of these photolysis parameters are used to directly assign the photolytic frequencies to a given chemical structure. The reference compounds used to assign rate constants to structures are given in the Table 4. For multifunctional species, each chromophore leading to photolysis is treated independently, except conjugated carbonyls.

3.6 Thermal decomposition

Thermal decomposition is only considered for PAN like species: $RC(O) O O NO₂ \rightarrow RC(O) (OO.) + NO₂.$ Rate constant recommended by Tyndall et al. (2001) for $CH_3C(O)OONO_2$ is used for all C_2 compounds. For longer carbon chain $(C_{>2})$, the rate constant is set to the high pressure limit recommended by Atkinson et al. (1999) for $C_2H_5C(0)OONO_2$: $k=2.0\times10^{15}$ exp(-12 800/T) molecule⁻¹ cm³ s⁻¹.

3.7 Peroxy radical reactions

The following reactions of organic peroxy radicals with NO, $NO₃$, $HO₂$ and other $RO₂$ are considered in the generator:

$$
R(OO \cdot) + NO \rightarrow R(O \cdot) + NO_2 \tag{R1}
$$

$$
\rightarrow R(ONO_2) \tag{R2}
$$

$$
RC(O)(OO·) + NO2 \rightarrow RC(O)(OODO2)
$$
 (R3)

$$
R(OO \cdot) + NO_3 \rightarrow R(O \cdot) + NO_2 + O_2 \tag{R4}
$$

$$
R(OO \cdot) + HO_2 \rightarrow R(OOH) + O_2 \tag{R5}
$$

$$
R(OO \cdot) + HO_2 \rightarrow R(OH) + O_3 \tag{R6}
$$

- $R(OO \cdot) + R'(OO \cdot) \rightarrow R(O \cdot) + R'(O \cdot) + O_2$ (R7)
	- $\rightarrow R(=O) + R'(OH) + O_2$ (R8)
	- $\rightarrow R(OH) + R' (=O) + O_2$ (R9)

Note that the Reaction (R2) is only considered for alkyl peroxy radicals while Reactions (R3) and (R6) are considered

Table 4. Reference compounds used to assign photolysis parameters to various chemical structures.

for acyl peroxy radicals only. We first briefly present below the rate constant selected for alkyl peroxy radicals, then for acyl peroxy radicals.

Rate constants for alkyl peroxy reactions with NO were set to $2.7 \times 10^{-12} \exp(360/T)$ molecule⁻¹ cm³ s⁻¹, based on Atkinson (1997) recommendations. Yields for organic nitrate and alkoxy radical formation are estimated using the SAR of Carter (2000), and depend on the peroxy class, the presence of polar functional groups (e.g. hydroxyl, carbonyl, nitrate) and the length of the carbon chain. The reaction rate recommended by Atkinson et al. (1999) for the $C_2H_5O_2 + NO_3$ reaction (i.e. 2.5×10^{-12} molecule⁻¹ cm³ s⁻¹) was used as a generic constant for the $RO₂ + NO₃$ reactions. Rate constants for the

^a quantum yields are from Desai et al. (1986); b this structure was also used for di substituted aldehydes on the α carbon; c quantum yields of the radical channel are from Zhu et al. (1999), quantum yield for the Norrish type II reaction are from Moortgat et al. (1999) for n-pentanal; d enols (>C=C(OH)−) produced by the Norrish type II reaction pathways are assumed to rearrange to the carbonyl form (>CH−CO−); e the more labile H is used to perform the Norrish II reaction if distinct H are available on the γ carbon; f dissociation occurs at the weakest CO–C bond. Bond energies are estimated using the Benson method (Benson, 1976); ^g total quantum yield based on 2-butanone values from Raber and Moortgat (1995); h this reaction is also used for $-CH_2-CO-C\equiv$ structures; i average value of 3-pentanone and 2,4-dimethyl-3-pentanone; j this reaction is also used for >CH−CO−C≡ and ≡C−CO−C≡ structures; k branching ratios for the molecular and radical channel set to 0.7 and 0.3, respectively. These ratios are based on an absolute quantum yield of 0.1 estimated by Carter (2000) for methyl isopropyl ketone; l quantum yields from Koch and Moortgat (1998) and Raber and Moortgat (1995); m Quantum yield from Carter (2000); n quantum yield from Magneron et al. (1999); o the carbene radical is expected to add O_2 to form a criegee like radical; p quantum yield from Carter (1996); q both channels are set with an equal probability.

 $RO₂ + HO₂$ reactions are evaluated using the SAR of Boyd et al. (2003) and depend on the chain length. For these reactions, the temperature dependencies of the rate constants show that Arrhenius activation energy factors are in the range of 1000–1500 K (Lesclaux, 1997). We therefore used an average activation energy factor of 1250 K and modified the SAR of Boyd et al. (2003) accordingly. The counter species method of Madronich and Calvert (1990) was used to parameterize the $RO₂+RO₂$ reactions. However, while Madronich and Calvert (1990) treated the counter species analogously to all other species (i.e. solved by the integration of differential equations), here we simply compute counter concentrations at each time step by summing the concentrations of the various peroxy radicals. Ten reactivity classes of $RO₂$ are considered in the generator, based on the review of Lesclaux (1997). These classes are given in the Table 5 as well as their corresponding kinetic parameters for the self-reactions (i.e. the reaction between 2 peroxy belonging to the same class). Rate constant for cross-reaction (i.e. the reaction between 2 peroxy belonging to distinct classes) are evaluated as by Madronich and Calvert (1990). The only exception is for the cross reactions of acyl and alkyl peroxy radicals, for which the rate constant was set to 1.0×10^{-11} molecule⁻¹ cm³ s⁻¹, independently of the alkyl peroxy class (Villenave et al., 1998).

Rates for $CH_3C(O)(OO)$ reaction with NO, HO_2 , NO₃ and $RO₂$ were used as generic constants for the corresponding acyl peroxy radical reactions. Rate constants were set to (1) 8.1×10^{-12} exp(270/T) for the reaction with NO (Tyndall et al., 2001), (2) 5.0×10^{-12} for the reaction with $NO₃$ (Canosa-Mas et al., 1996) and (3) 1.4×10^{-13} exp(925/T) molecule⁻¹ cm³ s⁻¹ for the reaction with $HO₂$ (Tomas et al., 2001). A branching ratio of 0.8 and 0.2 was set for the Reactions (R5) and (R6), respectively, based on Tomas et al. (2001). The $CH_3C(0)(OO_1) + NO_2$ reaction is in the fall off regime. The rate constants recommended by Tyndall et al. (2001) for this reaction was directly used for the corresponding reactions of C_2 species and the high pressure rate constant was assigned for $C_{>2}$ species, according to the recommendations of Atkinson (1994).

3.8 Alkoxy radical reactions

Alkoxy radicals undergo three reaction types under tropospheric conditions, namely reaction with O_2 , beta scission decomposition and 1,5-H-shift isomerization:

$$
\gt{CH(O\cdot)} + O_2 \to \gt{C(=O)} + HO_2 \tag{R10}
$$

$$
>C(O\cdot)-R \rightarrow C(=O)+R(\cdot) \qquad (R11)
$$

$$
>CH-(C)_2-C(O\cdot) < \ \to >C(\cdot)-(C)_2-C(OH)< \qquad (R12)
$$

Peroxy class	$k(298 \text{ K})$	\overline{A} molecule ^{-1} cm ³ s ^{-1}	E/R	α	notes			
	molecule ^{-1} cm ³ s ^{-1}		K					
CH ₃ O ₂	3.5×10^{-13}	9.5×10^{-14}	-390	0.37	b			
peroxy acyl								
RC(O)(OO·)	1.4×10^{-11}	2.5×10^{-12}	-500	1.0	\boldsymbol{b}			
primary alkyl peroxy								
linear chain	3.0×10^{-13}	5.6×10^{-14}	-500	0.6	\mathfrak{a}			
branched chain on β carbon	1.2×10^{-12}	7.8×10^{-15}	-1500	0.5	$\mathfrak a$			
O or N substitution on α or β carbon	4.0×10^{-12}	7.1×10^{-14}	-1200	0.6	\mathfrak{a}			
secondary alkyl peroxy								
CH ₃ CH(OO·)CH ₃	1.1×10^{-15}	1.7×10^{-12}	2200	0.6	\mathfrak{a}			
$C_{>3}$ peroxy	6.3×10^{-14}	1.0×10^{-10}	2200	0.6	\mathcal{C}			
O or N substitution on α or β carbon	6.6×10^{-13}	8.4×10^{-15}	-1300	0.3	a, d			
tertiary alkyl peroxy								
unsubstitued peroxy	3.1×10^{-17}	4.1×10^{-11}	4200	1.0	a, e			
O or N substitution on α or β carbon	5.0×10^{-15}	3.0×10^{-13}	1220	1.0	a, f			

Table 5. Rate constants at 298 K, Arrhenius parameters and radical channel branching ratios α for peroxy radical self reactions.

^a Kinetic and mechanistic parameters from Lesclaux (1997); b kinetic and mechanistic parameters from Tyndall et al. (2001); c rate constant</sup></sup> based on Boyd et al. (1999) for the reaction of $C_5H_{11}O_2$ and $C_{10}H_{21}O_2$ and using the activation energy and the branching ratio for ipropyl peroxy radical.; ^d based on kinetic and mechanistic parameters for CH₃CH(OH)CH(OO·)CH₃; ^e based on kinetic and mechanistic parameters for $(CH_3)_3C(OO·)$; f based on kinetic and mechanistic parameters for $CH_3C(CH_3)(OH)C(OO·)(CH_3)CH_3$.

Table 6. Arrhenius parameters for alkoxy radical reactions (adapted from Carter, 2000).

a A parameters are given in the [molecule – cm – s] unit system; $\frac{b}{2}$ Ea is set to a minimum of 0.4 kcal mol⁻¹ if the expression leads to lower values; ^c the heats of reaction are evaluated in the generator using the group contribution of Benson (1976) for formation heats estimation, with additional groups taken in Carter (2000) and Laval-Szopa (2003); d F_I factors are the neighbouring group activation parameters used for H abstraction by OH at 298 K in the SAR of Kwok and Atkinson (1995); ^e Activation energy for isomerization involving −O−, −CO− or $-O$ –CO– groups in the cyclic transition state are increased by an additional 3.5 kcal mol⁻¹; f the Eaa parameter is a function of the leaving group. Numerical values for the various groups can be found in Carter (2000).

Absolute rate constants for alkoxy radical reactions are very sparse and most data available concern branching ratios only. Note that alkoxy radicals are very reactive species (lifetime

lesser than \approx 10⁻⁵ s) and their concentrations do not directly contribute to the organic carbon or radical budgets to any significant extent. The branching ratio is therefore the critical **Table 7.** Branching ratios for the reactions of excited Criegee radicals (adapted from Carter (2000).

 a External stabilized biradicals are assumed to be converted into the corresponding carboxylic acid; \bar{b} branching ratio for the OH channel decreases with the chain length. Yields are set to 0.46, 0.36, 0.24, 0.16, 0.08 and 0.0 for C_2 , C_3 , C_4 , C_5 , C_6 and $C_{>7}$ biradical, respectively, based on Paulson et al. (1999); c if the 2 substituents on the biradical have abstractable H, branching ratio is set to the rate constant ratio of the corresponding H abstraction with OH; d if no abstractable H is available, then branching ratio is set to 0.9/0.1 for stabilization by collision and decomposition to $2 R(\cdot) + CO_2$, respectively.

parameters that must accurately be evaluated for the development of atmospheric chemical schemes. Structure/activity relationships for alkoxy reactions were recently developed by Carter (2000) for the purpose of chemical scheme generations and provide a consistent set of rules for branching ratio evaluations. The expressions recommended by Carter (2000) were implemented in the generator and systematically used to evaluate the rate constants associated to the alkoxy reactions. These expressions are summarized in the Table 6.

3.9 Criegee radical reactions

Energy rich Criegee biradicals (i.e. $>C(\cdot)(OO \cdot)$ structures) are produced by the O_3+ alkene reactions. These excited biradicals can either be stabilized by collision or undergo various unimolecular decompositions, leading to the production of radicals (in particular OH) or "stable" species. Branching ratios depend (1) on the identity of the substituents at**Table 8.** Representative C_7 secondary organics contributing to the organic budget during heptane oxidation.

tached to the biradicals (e.g. Paulson et al., 1999; Calvert et al., 2000), (2) on the chemical origins of the biradical that may lead to different ranges of excitation energies or ratio in the syn or anti configuration (e.g. Fenske et al., 2000b; Kroll et al., 2002). Numerous studies have been devoted these last 10 years to quantify the evolution of Criegee radicals, especially with respect to OH radical production (e.g. Orzechowska and Paulson, 2002; Siese et al., 2001; Lewin et al., 2001; Fenske et al., 2000a; Rickard et al., 1999; Paulson et al., 1999; Calvert et al., 2000, and references therein). However, large uncertainties still remain concerning the branching ratios and, to our knowledge, no parameterization is available concerning the formation and evolution of Criegee radicals that takes into account the detailed chemical identity of the groups bonded to the biradicals or the syn/anti ratios. Fixed branching ratios are assigned in the generator for external (i.e. $-CH(\cdot)(OO\cdot)$ structures) and internal (i.e. $>C(\cdot)(OO(\cdot))$ structures) biradicals, based on the SAR of Carter (2000). These branching ratios are summarized in the Table 7.

3.10 Cyclic molecules

As mentioned above, the chemistry of cyclic molecules is not currently implemented in the generator. The chemical schemes of several cyclic compounds (e.g. terpenes and aromatic compounds) were written manually up to the formation of ring opening products. Full oxidation of these noncyclic reaction products is then handled by the generator. The chemical scheme for cyclic compounds is based on the SAPRC-99 mechanism (Carter, 2000).

Fig. 7. Mechanism comparison for n-heptane oxidation. Solid line: generator scheme; dashed lines: MCM-v3 scheme; dotted lines: SAPRC99 scheme. The shaded area corresponds to $\pm 20\%$ deviation around the mixing ratio simulated with the generator scheme. The SAPRC99 mechanism does not include methyl glyoxal for heptane.

4 Evaluation of the chemical schemes

To evaluate the schemes written by the generator, some comparisons were made with well-established chemical mechanisms: (1) the SAPRC99 scheme developed by Carter (2000) which has been developed and validated using experimental smog chamber data and (2) the master chemical mechanism (MCM-v3.1) developed at the University of Leeds (Saunders et al., 2003b; Jenkin et al., 2003) which is to our knowledge the most detailed scheme available in the literature. Both mechanisms treat the chemistry induced by a given VOC specifically and are therefore especially suitable to test the chemical schemes provided by the generator. All the schemes were implemented in the same box model and the simulations were carried out with identical conditions. Photolysis frequencies were computed for mid-latitude equinox

conditions using the TUV model (Madronich and Flocke, 1998). These frequencies were assigned to the corresponding reactions according to the recommended rules for each mechanism. Furthermore, the same set of inorganic reactions was implemented in each scheme, so that differences could only be ascribed to the various organic schemes. Time integration was performed using the two-step solver developed by Verwer (1994); Verwer et al. (1996). This solver is based on the implicit second order two-step backward differentiation formula (BDF) which is solved approximately using Gauss-Seidel iteration. Initial mixing ratios are respectively set to 10, 20 and 40 ppb for NO_x , the VOC under consideration and ozone. With these conditions, the chemistry is characterized by a NO_x saturated regime in the first day but a NO_x limited regime on the last two days.

Fig. 8. Mechanism comparison for isoprene oxidation. Solid line: generator scheme; dashed lines: MCM-v3 scheme; dotted lines: SAPRC99 scheme. The shaded area corresponds to $\pm 20\%$ deviation around the mixing ratio simulated with the generator scheme.

Figures 7 and 8 show the concentration profiles simulated for n-heptane and isoprene, respectively. The comparisons of the profiles simulated with SAPRC99, MCM-v3.1 and the generator scheme show a fairly good agreement. For most of the tested species, mixing ratio simulated with the generator scheme match within 20% those obtained with the SAPRC99 and MCM-v3.1 schemes. In particular, an excellent agreement between the generator scheme and the SAPRC99 scheme was obtained in the case of n-heptane oxidation for the $O_3/NO_x/HO_x$ species (see Fig. 7). On the other hand, the generator scheme was found to best match the MCM-v3.1 for the simulation using isoprene as a precursor. In that case, both mechanisms agree within few percents for the $O_3/NO_x/HO_x$ species (see Fig. 8). As expected, the largest discrepancies were found for secondary organic species produced after many successive oxidation steps. This point is highlighted in the Figs. 7 and 8 for methyl glyoxal: the agreement with MCM-v3.1 is worse than a factor 2; no source of methyl glyoxal is considered for n-heptane oxidation in SAPRC99. Nevertheless, some secondary species produced with large yields like formaldehyde show a surprisingly good agreement between the 3 mechanisms, at least for the conditions tested here.

Additional tests were performed to evaluate the chemical schemes provided by the generator. These tests are presented in a companion paper (Szopa et al., 2005). All the tests performed so far did not highlight any problem or major error on the generator's code. Furthermore, no significant numerical bias was found by the time integration of the very large number of species and reactions generated (e.g. for heptane, we solved the chemistry of about 4.6×10^5 species and 4.5×10^6 reactions).

Fig. 9. Evolution of carbonaceous species during the oxidation of n-heptane. Panel **(a)**: distribution of the carbon. Panel **(b)**: distribution of secondary organic carbon for various chain lengths. Panel (c): distribution of secondary C_7 organics as a function of the number of functional groups borne by the molecules.

5 Example of organic budgets during gas phase oxidation

Some very preliminary simulations were performed to examine the oxidation pathways of a given VOC. We present here some results obtained for the case of n-heptane with a particular focus on the gradual change of organic compounds during oxidation. To our knowledge, such simulations are the first using a fully detailed oxidation scheme. Simulation conditions are identical to those described in the previous section, except that simulation is run over 5 days.

Speciation of the primary carbon (i.e. heptane), secondary organic carbon, and inorganic carbon (i.e. $CO + CO₂$) is shown as a function of time in Fig. 9a. For the conditions used here, more than 90% of heptane is predicted to be removed after 5 days. However, only 20–30% of the carbon available was found to be oxidized into CO or $CO₂$ at that time. Most of the carbon lies in the secondary organic fraction, which is the dominant fraction after 1.5 simulated days. Figure 9b shows the distribution of secondary organic species as a function of the chain length. The secondary organic fraction is shown to be dominated by species holding 7 carbon atoms, i.e. of the same size as the parent compound. The organic carbon included in C_1 to C_6 species grow with time, but this fraction remains lower than the C_7 fraction even after 5 days of simulated oxidation (see Fig. 9b). Figure 9c gives additional insight into the distribution of secondary C_7 and shows how these species are distributed as a function of the number of functional groups borne by the molecules. Organics appear to have typically 2 or more functional groups. Henry's law coefficients for many organic compounds were recently summarized by Sander (1999). Available data shows that the Henry's law coefficient for most difunctional species typically lies in $10^3 - 10^6$ M atm⁻¹ range (or higher when the number of functions is greater than 2). For a typical liquid water content of $L=10^{-6}$ g m⁻³ in clouds, species in that range can be classified as moderately to very soluble (Seinfeld and Pandis, 1998), i.e. dissolving from significantly to almost totally in cloud droplets at thermodynamical equilibrium. These results thus strongly suggest that a significant fraction of the secondary organic carbon would dissolve into water droplets during cloud events. Furthermore, a nonnegligible fraction of C_7 organics is found to bear at least 4 functional groups (see Fig. 9c). Vapor pressure estimates using the Myrdal and Yalkowsky (1997) group contribution method lead to extremely low vapor pressure for these kinds of species, typically in the range of 10^{-10} to 10^{-7} atm. These compounds may thus be candidate compounds for the formation of Secondary Organic Aerosols (SOA).

The distribution of the different organic moieties is shown as a function of time in the Fig. 10. The results are expressed as a functional group per organic carbon ratio $(R_{OF/C})$. About 30% of the organic carbon is found to be substituted after 5 days (see Fig. 10). The dominant functionality is the ketone group $(R_{-CO-}/C=5-15%)$, with a significant contribution of the alcohol and nitrate groups $(R_{\text{ROH/C}} \approx R_{\text{RONO2}/\text{C}} \approx 2-4\%)$. As previously stated, oxidation during the first day occurs in a NO_x saturated regime (i.e. the $RO₂ + NO$ is the dominant reaction for peroxy radicals). The hydroperoxide, peracid and carboxylic acid moieties were found to be negligible during the first day, in agreement with the chemistry expected for a NO_x saturated regime. These moieties appear to be significant only after the second day, when the NO_x limited regime is reached. Finally, the carboxylic acid moiety was found to be one of the major groups after 5 days. This is a particularly interesting

Fig. 10. Evolution of the organic functionalities to carbon ratio $(R_{OF/C})$ during the oxidation of n-heptane.

outcome since the carboxylic acid functional group is highly water-soluble and decreases to a large extent the vapor pressure of the species. Carboxylic acids were measured to be a major functional group in organic aerosols (e.g. Sempere and Kawamura, 1994) and have been suggested as key compounds for organic aerosol nucleation, especially in the diacidic form (Hoffmann et al., 1998). Some representative C⁷ secondary organics predicted during heptane oxidation are reported in the Table 8. The contribution of the species to each group listed in the Table 8 was predicted to be above a few percents.

The nitrogen budget during the n-heptane oxidation is shown in the Fig. 11. About 30% of the initial NO_x load was found to be held as organic nitrate during the first day (i.e. in the NO_x saturated regime). As expected, secondary organics are shown to be potentially key contributors to the long range transport of nitrogen oxides (if not removed by dry or wet deposition when included in multifunctional species).

Although very preliminary, these results are very promising and show the benefit of explicit schemes for modeling the evolution of organic compound during oxidation. These results also highlight the potential of the self generating approach for the development of multiphase organic chemical schemes.

6 Conclusions

The development of explicit chemical schemes is a necessary step to test our current understanding of the evolution

Fig. 11. Evolution of the major nitrogen compounds during the oxidation of n-heptane.

time (day)

and impact of organic carbon during its atmospheric oxidation. However, the myriad of intermediates involved preclude any attempt to describe the numerous physical and chemical transformations manually. Data processing tools are required to (1) assimilate the various experimental data provided by laboratory studies (2) codify the various estimation methods, and (3) generate consistent, comprehensive oxidation schemes on a systematic basis.

We showed in this study that a self generating approach can be used to generate explicit gas-phase oxidation schemes.

A very promising application of the generator is the modelling of secondary organic aerosol formation. Current models describing SOA formation are developed on an empirical approach, based on yields measured in smog chambers (e.g. Odum et al., 1997; Hoffmann et al., 1997; Schell et al., 2001) and justify the development of exploratory models describing in detail the various processes leading to SOA formation (Griffin et al., 2002, 2003; Pun et al., 2002; Jenkin, 2004). The main difficulties lie in describing the sources and sinks of the myriad of VOC that may contribute to SOA and their thermodynamic equilibrium with the condensed phases. The generator is clearly a powerful tool to study SOA formation by its ability to explicitly describe the chemical dynamics of the various precursors.

Furthermore, organic chemistry inside cloud droplets raises many questions, e.g. its role in the O_x/HO_x budget, the formation of organic acids and organic aerosols after the evaporation of the cloud droplets, and the atmospheric removal of organic matter. Answering these kind of questions by a modelling approach clearly requires the development of chemical schemes describing the organic oxidation in the aqueous phase. Currently, laboratory studies dealing with the oxidation in atmospheric aqueous phases are receiving increasing interest and the available kinetic database for these processes is becoming substantial. Extension of the self generating approach to model multiphase system would provide a valuable exploratory tool to assess the role of clouds in the evolution of organic compounds.

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