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Organic particulate matter formation at varying relative humidity using surrogate secondary and primary organic compounds with activity corrections in the condensed phase obtained using a method based on the Wilson equation

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Abstract

Secondary organic aerosol (SOA) formation in the atmosphere is currently often modeled using a multiple lumped "two-product" (N·2p) approach. The N·2p approach neglects: 1) variation of activity coefficient (ζ_i) values and mean molecular weight MW ⁵ in the particulate matter (PM) phase; 2) water uptake into the PM; and 3) the possibility of phase separation in the PM. This study considers these effects by adopting an $(N\cdot 2p)^{\zeta, MW, \theta}$ approach (θ is a phase index). Specific chemical structures are assigned to 25 lumped SOA compounds and to 15 representative primary organic aerosol (POA) compounds to allow calculation of ζ_i and MW values. The SOA structure assignments are based on chamber-derived 2p gas/particle partition coefficient values coupled with 10 known effects of structure on vapor pressure $p_{L,i}^{\circ}$ (atm). To facilitate adoption of the $(N\cdot 2p)^{\zeta,MW,\theta}$ approach in large-scale models, this study also develops CP-Wilson.1, a group-contribution ζ_i -prediction method that is more computationally economical than the UNIFAC model of Fredenslund et al. (1975). Group parameter values required by CP-Wilson.1 are obtained by fitting ζ_i values to predictions from UNIFAC. The $(N\cdot 2p)^{\zeta,MW,\theta}$ approach is applied (using CP-Wilson.1) to several real α -pinene/O₃ chamber cases for high reacted hydrocarbon levels ($\Delta HC \approx 400$ to $1000 \,\mu g \,m^{-3}$) when relative humidity (RH) ≈50%. Good agreement between the chamber and predicted results is obtained using both the $(N \cdot 2p)^{\zeta, MW, \theta}$ and $N \cdot 2p$ approaches, indicating relatively small water effects under these conditions. However, for a hypothetical α -pinene/O₃ 20 case at $\Delta HC = 30 \,\mu g \,m^{-3}$ and RH = 50%, the $(N \cdot 2p)^{\zeta, \overline{MW}, \theta}$ approach predicts that water uptake will lead to an organic PM level that is more double that predicted by the N.2papproach. Adoption of the $(N \cdot 2p)^{\zeta, \overline{MW}, \theta}$ approach using reasonable lumped structures for SOA and POA compounds is recommended for ambient PM modeling.

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1 Introduction

A significant fraction of the fine particulate matter (PM) in the atmosphere can be organic in nature, and so that fraction is of interest for visibility, health effect, and climate effect reasons (Mazurek et al., 1997; Pope, 2000; Bates et al., 2006). Organic PM

⁵ (OPM) is always a complex mixture, and usually contains compounds loosely categorized as primary organic aerosol (POA) compounds and secondary organic aerosol (SOA) compounds. OPM can also contain compounds that have been formed by a variety of accretion reactions in which reactive SOA and POA compounds combine to yield products of appreciable molecular weight and low vapor pressure (Kalberer et al., 2004; Barsanti and Pankow, 2004, 2005, 2006).

Absorptive gas/particle (G/P) partitioning may be parameterized according to the model of Pankow (1994a). In the case of one absorbing phase within the PM, for compound *i* the equilibrium partitioning constant $K_{p,i}$ (m³ μ g⁻¹) is given by

$$K_{p,i} = \frac{c_{p,i}}{c_{g,i}} = \frac{RT f}{10^6 \overline{MW} \zeta_i \rho_{L,i}^\circ}$$
(1)

¹⁵ where: c_{p,i} (ng μg⁻¹)=P-phase concentration; c_{g,i} (ng m⁻³)=G-phase concentration; *R* is the gas constant (=8.2×10⁻⁵ m³ atm mol⁻¹ K⁻¹); *T*(K)=temperature; *f* is the weight fraction of the PM that is the absorbing phase (often taken to be unity for OPM calculations); MW(g mol⁻¹)=mean molecular weight of the absorbing phase; p[°]_{L,i} (atm)=vapor pressure of *i*; and ζ_i=mole-fraction-based activity coefficient of *i*. K_{p,i} values generally
²⁰ depend strongly on *T* because p[°]_{L,i} is usually a strong function of *T*. Significant temporal and spatial variations in K_{p,i} values can also be caused by variations in ζ_i and MW due to changes in the types and levels of the compounds (including water) in the P phase mixture.

Many of the applications of the Pankow (1994a, b) model for predicting secondary OPM formation in the atmosphere have been based on the "two-product" simplification

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of Odum et al. (1996). This implementation acknowledges that a parent hydrocarbon (HC, e.g. toluene, α -pinene, etc.) will be oxidized to a range of secondary products, but assumes that the mix of products can be represented using up to two hypothetical "lumped" surrogate compounds. For each parent HC, yield and compound character-

- ⁵ istics for two lumped compounds are obtained by fitting chamber yield data to fourparameters: two stoichiometric formation α_i factors and two $K_{p,i}$ values. (With one lumped product, one α_i and one $K_{p,i}$ value are invoked.) With Δ HC (μ g/m³) giving the amount of reacted HC, for each of the lumped products, the total (G+P) amount formed is assumed to be given by $\mathcal{T}_i = \alpha_i \Delta$ HC (μ g/m³).
- ¹⁰ Many two-product parameters obtained have been acquired in a chamber at a single temperature and under "dry" conditions (i.e., very low relative humidity (RH)). Extrapolations of $K_{p,i}$ values for a given OPM composition to another temperature under dry conditions have proceeded using the Clausius-Clapeyron equation with an estimate of the enthalpy of vaporization ($\Delta H_{vap,i}$) for each hypothetical lumped product. Utilizing ¹⁵ chamber data in a theoretical consideration of the effects of RH is more difficult, and is a topic of this work.

In a chamber study of the oxidation of a mix of parent HCs, Odum et al. (1997) sought to predict the amount of OPM formed by using the collection of two-product α_i and $K_{p,i}$ values measured for oxidation of the individual parent HCs. This approach implicitly assumes similarity in both the MW and the polarity characteristics of all the various two-product compounds so that in the OPM formed from all mixes of parent HCs, \overline{MW} remains approximately constant and all $\zeta_i \approx 1$. Pankow and Barsanti (2008)¹ have designated this the "*N*·2p approach"; its range of applicability in the atmosphere remains uncertain, but nevertheless the *N*·2p approach has been widely utilized in the prediction of secondary OPM levels in the ambient atmosphere (e.g., Hoffman et al., 1997; Kanakidou et al., 2000; Pun et al., 2001;Tsigaridis and Kanakidou, 2003).

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¹Pankow, J. F. and Barsanti, K. C.: Framework for Considering the Complexity of the Compounds Present in Organic Particulate Matter in the Atmosphere, in preparation, 2008.

The computational advantage of the N·2p approach in 3-D air quality models (e.g., as in MADRID 1 as described by Pun et al., 2001) may be understood as follows. Any multi-component G/P model requires an iterative solution to determine the PM composition and level at each point in space and time. The N·2p approach assumes

- ⁵ a limited number of secondary products, and provides a fixed $K_{p,i}$ value for each *i* for each iteration cycle at the *T* of interest. In contrast, if the $K_{p,i}$ values were allowed to vary because of dependence of the ζ_i and $\overline{\text{MW}}$ on PM composition, then each solution within the series of solutions performed during each iteration cycle would require added computation time to estimate the ζ_i and $\overline{\text{MW}}$ for the PM phase. Bowman and Melton
- ¹⁰ (2004) compare the computational requirements of a number of ζ_i prediction methods, including the UNIFAC method of Fredenslund et al. (1975).

Parent HCs considered in the MADRID 1 model are known to produce oxidation products with a range of polarities. For example, α -pinene quickly leads to products like hydroxyacids and diacids that contain moderate polarity, while humulene initially leads

- ¹⁵ to products of considerably lower polarity. The current assumption within MADRID 1 that all ζ_i =1 for the OPM from all mixes of parent HCs is thus problematic. Moreover, the assumption that the organic portion of the PM formed is essentially free of water will certainly be in error whenever a significant portion of the PM is comprised of relatively high polarity compounds, and the RH is not low. In such circumstances, RH-driven
- ²⁰ water uptake into the PM phase can occur, further affecting ζ_i values (especially of the lower polarity products), and the value of \overline{MW} . (A consideration of the potential magnitude of the effects of changing RH on PM levels at high Δ HC values (244 to $501 \,\mu g \,m^{-3}$) is provided by Seinfeld et al. (2001) for the ozone oxidation of several different biogenic HCs as well as cyclohexene.) Also, increasing RH levels will increase
- ²⁵ the likelihood of phase separation in the PM, especially whenever the OPM contains significant mass fractions of both SOA and POA compounds: the generally significant polarities of the former contrast with the generally low polarities of the latter. When phase separation does occur, a phase index θ is needed, with $\theta = \alpha$ referring to a relatively more polar, hydrophilic phase (α mnemonically suggesting "aqueous"), and



 $\theta = \beta$ referring to the relatively less polar, less hydrophilic phase (Erdakos and Pankow, 2004). When the possibility of variation in the ζ_i and $\overline{\text{MW}}$ and the possibility of phase separation are added to the *N*·2p approach, the result is referred to as the " $(N \cdot 2p)^{\zeta, \overline{\text{MW}}, \theta}$ approach" (Pankow and Barsanti, 2008¹).

With a superscript * used to denote a value determined under particular chamber conditions with a specific parent HC, Bowman and Karamalegos (2002) employ Eq. (1) to extrapolate a K^{*}_{p,i} to different conditions. In the 2p view, the OPM formed from a given parent HC could be composed of significant amounts of both lumped compounds. Thus, given the level of approximation already allowed in that view, it may be reasonable to assume that ζ^{*}_i≈1 for both products in that OPM. For partitioning to a significantly different type of OPM, however, it may be that ζ^{*}_i≠1. Thus, with MW

and *T* also subject to variation, a $K_{p,i}^*$ value may be extrapolated using the ratios: a) $\zeta_i^*/\zeta_i \approx 1/\zeta_i$; b) $\overline{MW}^*/\overline{MW}$; c) T/T^* ; and d) $p_{L,i}^\circ(T^*)/p_{L,i}^\circ(T)$. Assuming that $\Delta H_{vap,i}$ is constant over the temperature interval of interest (i.e., from T^* to *T*), correction for the effect of *T* on $p_{L,i}^\circ$ occurs according to the integrated Clausius-Clapeyron equation which gives $p_{L,i}^\circ(T^*)/p_{L,i}^\circ(T) = \exp[\Delta H_{vap,i}(1/T - 1/T^*)]$. Assuming that f = 1, the overall result is (Bowman and Karamalegos, 2002)

$$\mathcal{K}_{\mathrm{p},i}(T, x_i, x_2, x_3, \dots, x_n) = \mathcal{K}_{\mathrm{p},i}^* \left(\frac{\overline{\mathrm{MW}}^*}{\overline{\mathrm{MW}}}\right) \left(\frac{1}{\zeta_i}\right) \left(\frac{T}{T^*}\right) \exp\left[\frac{\Delta H_{\mathrm{vap},i}}{R} \left(\frac{1}{T} - \frac{1}{T^*}\right)\right]$$
(2)

where $K_{p,i}(T, x_i, x_2, x_3, ..., x_n)$ here denotes that $K_{p,i}$ depends on *T* and on the PM composition, the latter being characterized by the set of mole fraction values x_j . Equation (2) has been applied in global modeling of SOA in the troposphere by Tsigaridis and Kanakidou (2003), with the needed ζ_i values estimated using the Wilson (1964) equation. However, while Tsigaridis and Kanakidou (2003) demonstrate the computational practicality of using Eq. (2) in a large-scale 3-D model with the Wilson equation used for the ζ_i corrections, the Wilson equation parameters were assigned without regard to probable compound structure and functionality.

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This work has four goals: 1) assign reasonable, specific surrogate structures to 25 lumped secondary compounds pertaining to a range of parent HCs of interest, and to 15 surrogate primary OPM compounds; 2) develop and implement a Wilson-equationbased group contribution method for prediction of ζ_i values for use with the 40 surrogate compounds that is computationally more economical than UNIFAC; 3) relax four key assumptions of the *N*·2p approach (all $\zeta_i=1$; no RH effects; $\overline{MW}=$ constant; and a single OPM phase) thereby permitting use of the $(N\cdot2p)^{\zeta,\overline{MW},\theta}$ approach; then 4) use the $(N\cdot2p)^{\zeta,\overline{MW},\theta}$ approach to calculate OPM formation in selected cases using: a) experimentally determined or estimated values of the $K_{p,i}^*$; b) Eq. (2); c) the assigned chemical structures; and d) the ζ_i prediction method developed here. For comparison, calculations were also made using the *N*·2p approach.

2 Methods

2.1 Partitioning SOA compounds

A total of 25 lumped secondary products were considered to arise from a total of 14 HC oxidation processes. Based on Odum et al. (1996), Griffin et al. (1999), Pun et al. (2003), Henze et al. (2006), and Griffin (personal communication, 2007), 11 of the processes assume two lumped products, and three of the processes (β -pinene reacting with NO₃ radical, humulene reacting with OH radical, and an *n*-alkane (C₁₆) reacting with OH radical) assume one lumped product each. Reaction (13) (a 2-ring polycyclic aromatic hydrocarbon (PAH) with OH) and Reaction (14) (C₁₆*n*-alkane with OH) were included as representative secondary reactions involving intermediate volatility parent HC compounds. Table 1 summarizes the information on the final set of the 40 surrogate compounds considered (25 lumped secondary products and 15 primary compounds). As noted above, if ζ_i values are to be estimated in a mixture of interest, specific

²⁵ structural information is required for the compounds in the mixture. Griffin (personal



communication, 2007) and this study considered known gas phase reaction mechanisms and kinetics (Griffin et al., 1999, 2002a, b, 2003; Surratt et al., 2006) to obtain the assignments for each lumped secondary product used in Table 1 for: 1) number of carbon atoms $v_{C,i}$; 2) whether cyclic or acyclic; (3) whether aromatic; and 4) retention

⁵ (or not) of a double bond found in the parent HC. As summarized in Eq. (5) below, a corresponding initial estimate of MW_i was then assigned herein for each lumped product compound. Then, as summarized in Eq. (6) below, an initial estimate of MW * was computed as the mean of the MW_i estimates for the lumped products (two or one) from a given HC oxidation reaction (for two lumped products, this is equivalent to assuming that the OPM can be approximated as a 1:1 molar mixture of the two products). An initial estimate of each log₁₀ p[°]_{L,i}(T*)^{chamber} was then obtained as summarized in Eq.(7)

below, i.e., as based on Eq. (1) and $K_{p,i}^*(T^*)$ at T^* using the \overline{MW}^* estimate and assuming $\zeta_i^* \approx 1$ for each lumped product (see discussion preceding Eq. 2).

With the task denoted in Eq. (7) below completed, an approximation of the spe-¹⁵ cific functionality was needed for each of the 25 surrogate secondary compounds (ultimately, for use as input to the ζ_i prediction method). The approach taken was to utilize known relationships between $p_{L,i}^{\circ}$ values and structure. For organic compounds, $p_{L,i}^{\circ}$ values decrease as $v_{C,i}$ increases, and as compound polarity increases. In the SIM-POL.1 group-contribution model for prediction of $p_{L,i}^{\circ}$ (atm) values, Pankow and Asher (2007) write

$$\log_{10} p^{\circ}_{\mathrm{L},i}(T) = \sum_{k} v_{k,i} b_{k}(T) \qquad k = 0, 1, 2, 3, \text{ etc.}$$
(3)

where: $v_{k,i}$ is the number of groups of type *k* in compound *i*; $b_k(T)$ is the *T*-dependent contribution to $\log_{10} p_{L,i}^{\circ}(T)$ from one group of type *k*. Equation (3) provides a means to assemble the aggregate effects of structure on $p_{L,i}^{\circ}(T)$. SIM-POL.1 utilizes a zeroeth group (*k*=0), with $v_{0,i}=1$ for all *i* and $b_0(293.15)=1.99$. The *k*=1 group pertains to molecular carbon; e.g., for hexane, $v_{1,i}\equiv v_{C,i}=6$. Because $b_1(293.15)\equiv b_C(293.15)=-0.47$, within any given compound class, $p_{L,i}^{\circ}$ values decrease by ~ $\frac{1}{2}$ order of magnitude for every unit increase in carbon number. And, $p_{L,i}^{\circ}$

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can also be decreased for a given carbon skeleton by adding polar functional groups. Eq. (3) may thus be re-written as

$$\log_{10} p_{L,i}^{\circ}(T) = v_{0,i} b_0(T) + v_{C,i} b_C(T) + \omega_i(T)$$

wherein all the structural aspects beyond carbon number are placed in the term $\omega_i(T)$. With $v_{0,i}=1$ and using the assumed value of $v_{C,i}$, then $p_{L,i}^{\circ}(T^*)^{\text{chamber}}$ allows an estimate of $\omega_i(T^*)^{\text{chamber}}$ (see also Eq. 8 below). Four groups are of special interest: hydroxyl ($b_7(293.15) \equiv b_{OH}(293.15) = -2.29$); aldehyde ($b_8(293.15) \equiv b_{CHO}(293.15) = -1.06$); ketone ($b_9(293.15) \equiv b_{CO}(293.15) = -0.99$); and carboxylic acid ($b_{10}(293.15) \equiv b_{COOH}(293.15) = -3.59$). Thus, for example, the conversion of cyclohexene to adipic acid is accompanied by about a seven order of magnitude drop in $\log_{10} p_{1}^{\circ}$ (293.15).

Compound-to-compound differences in polarity are the primary drivers of differences among the ζ_i values in a mixture. Considering the groups that contribute significantly to $\omega_i(T)$ by adding polarity to a molecule (e.g., the four groups noted above, nitrate (ONO₂), nitrite (NO₂)), as well as other structural groups assumed to be retained from the parent HC (e.g., rings), the goal was to manually vary the $v_{k,i}$ to obtain an estimate of $\omega_i(T^*)^{\text{fitted}}$ that would match $\omega_i(T^*)^{\text{chamber}}$ (see Eq. 9). The goal was to thereby derive a reasonable approximation of the overall polarity for each surrogate lumped secondary product for subsequent use in ζ_i prediction. Each resulting new $v_{k,i}$ set (including $v_{C,i}$)

- ²⁰ implied a new MW_{*i*} for the lumped product. When executed in concert with the other lumped product (if it exists) from each particular parent HC, as summarized in Eq. (10) below the process led to updated estimates of the MW_{*i*} and $\overline{\text{MW}}^*$. Consequently, the approach used led naturally to an iterative process producing a possible best-fit $v_{k,i}$ set for each lumped product.
- The overall scheme by which the values of each $v_{k,i}$ set were assigned is summarized below. (The character " \rightarrow " should be read as "gives".)

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(4)

(5)

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$$MW_i \rightarrow \overline{MW}^*$$
 estimated as the simple mean of the MW_i values from Eq. (5)

$$\overline{\text{MW}}^*$$
 with $\overline{\mathcal{K}_{p,i}^*(\mathcal{T}^*)} \rightarrow \text{estimate of } p_{L,i}^\circ(\mathcal{T}^*)^{\text{chamber}}$ by Eq. (1), with $\zeta_i^* = 1$

$$\rho_{\mathrm{L},i}^{\circ}(T^*)^{\mathrm{chamber}}$$
 with $v_{0,i}$ and $v_{\mathrm{C},i} \rightarrow \omega_i(T^*)^{\mathrm{chamber}}$ by Eq. (4) (8)

$$\omega_i(T^*)^{\text{chamber}} \to \text{ fitted } \nu_{k,i} \text{ set } \to \omega_i(T^*)^{\text{fitted}} \text{ by inverse application of Eq. (4)}$$
 (9)

5 fitted
$$v_{k,i}$$
 set (including $V_{C,i}$) $\rightarrow MW_i$

Boxed values represent quantities that were held fixed during the iteration. A set of preliminary structures for the 25 SOA surrogate compounds obtained prior to full convergence of the process is given in the Supplementary Materials (http://www. atmos-chem-phys-discuss.net/8/995/2008/acpd-8-995-2008-supplement.pdf). At the

end of the process, for every surrogate SOA product, the iteration yielded two converged values of ω_i(*T*), namely ω_i(*T**)^{chamber} and ω_i(*T**)^{fitted}: the agreement was within a few percent in every case. Even though binary mixtures of different compounds will not in general reflect exact ideality, the *N*·2p assumption of ζ_i=1 was maintained throughout the iteration (see Eq. 7). However, because a ζ_i prediction method such as CP-Wilson.1 will not in general yield ζ_i=1 in any OPM mixture, after the fitting, re-

sults obtained for one parent HC using the $(N \cdot 2p)^{\zeta, MW, \theta}$ approach will not collapse at RH=0% to the corresponding 2p results if that parent HC is considered to lead to two products.

The final assumed structures for the surrogate SOA products are given in Fig. 1; ²⁰ corresponding inferred molecular parameters are given in Table 1. While there is some arbitrariness in the selection of each final $v_{k,i}$ set and the corresponding structure (including the insertion of ether linkages to fine tune the $\omega_i(T)$ fit), this is not considered problematic given the considerable approximations that are already built into the 2p model: simply finding a structure that matches the value of $\omega_i(T)$ derived using Eq. (4)

provides meaningful insight regarding aggregate compound polarity that can be used to predict ζ_i effects in OPM systems.

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(6)

(7)

(10)

2.2 Partitioning POA compounds

15 POA surrogate compounds (P1–P15) were selected to cover a broad range of source types; all 15 compounds were considered subject to G/P partitioning (Fig. 1). Structures for P1–P8 were obtained from Griffin et al. (2003) for mobile (P1–4, P7), mo⁵ bile/natural (P5), and general cooking (P6) sources. P9, P10, and P15 were selected based on Nolte et al. (1999) as being relevant for meat cooking sources. Levoglucosan (P11) was selected as relevant for biomass burning (Simoneit et al., 2000; Fraser et al. 2002; Zhao et al., 2007). P8, P12, and P13 were selected as representatives of unresolved complex mixture (UCM) materials found in primary mobile emissions. P8
¹⁰ was considered by Griffin et al. (2003). P12 and P13 were included here to expand the volatility range of UCM related materials considered.

2.3 Chang-Pankow-Wilson (CP-Wilson) activity coefficient method

Equations. The approach developed here to calculate ζ_i values is based on the equation of Wilson (1964), with modifications introduced for use in a group-contribution ¹⁵ manner and for consideration of T effects by application of a 1/T factor as suggested by the Scatchard-Hildebrand equation (Flory, 1953). For each neutral compound *i* in a mixture of other such compounds, each group is therefore assumed here to contribute additively to ζ_i according to

$$\ln \zeta_{i} = \frac{\sum_{k} (-n_{k,i} \ln \Gamma_{k}) - C_{i}}{T/300}$$
(11)

²⁰ where: $n_{k,i}$ is the number of groups of type k in i; Γ_k is the activity coefficient of group k; and C_i is a compound specific constant that functions as a reference-state correction term. While k is again used as the group index for the summation, the set of groups used for the CP-Wilson method with coefficients $n_{k,i}$ is not synonymous with the set used by Asher and Pankow (2007) with coefficients $v_{k,i}$.

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The equation for Γ_k is assumed here to take the same form as that originally proposed by Wilson (1964) so that

$$\ln \Gamma_k = -\ln(\sum_j x_j \Lambda_{kj}) + 1 - \sum_j \frac{x_j \Lambda_{jk}}{\sum_j x_j \Lambda_{jj}}$$
(12)

The summations occur over all groups in the mixture wherein: *j* and *l* providing in-⁵ dexing through the groups; x_j is the group mole fraction; and Λ_{kj} is the interaction parameter between groups *k* and *j*. For each compound *i*, the constant C_j is evaluated according to

$$C_i = \sum_k n_{k,i} \ln \Gamma_k^{(i)} \tag{13}$$

where $\Gamma_k^{(i)}$ is the activity coefficient of group *k* in pure *i* and is evaluated using Eq. (12). Equations (11–13) compose the CP-Wilson method. Values of C_i for the compound structures considered here are given Table 1.

In its original form, the Wilson equation is less general than the CP-Wilson method because it is not a group-contribution method. Rather, it handles each compound in the mixture as a full chemical entity, and so its implementation requires specific chem-

- ¹⁵ ical property information for all the compounds in the mixture. This poses an obvious problem for atmospheric applications: even if the composition of a given atmospheric OPM sample could be accurately characterized, the property information needed for use with the Wilson equation would not be available, not even for some appropriate list of lumped/surrogate compounds. The group contribution approach utilized in the
- ²⁰ CP-Wilson method overcomes this problem: this method only requires parameter information for the constituent groups, and not property data for all compounds of interest. In general, then, the CP-Wilson method will be significantly more computationally economical than UNIFAC because it requires fewer logarithm and double summation operations, and because the empirical treatment for the *T* dependence in Eq. (11) allows the *C* to be computed encourse and thereafter acquired from a lock up table.
- the C_i to be computed once, and thereafter acquired from a look-up table.

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Parameter Fitting for CP-Wilson.1. The parameter values needed for a group contribution method are generally obtained by a fitting that minimizes some measure of the difference, for the parameter of interest, between: a) the group-contribution predicted values; vs. b) corresponding experimental values. The particular fitting parameters obtained here combined with the governing equations compose version CP-Wilson.1. Ideally, the fit carried out here would utilize experimental ζ_i values obtained for mixtures involving compounds similar to those of interest. Since such experimental data are not currently available, UNIFAC-generated values (i.e., ζ_i^U values) were used as the best, readily available substitute. The ζ_i^U values were obtained for mixtures involving compounds with the mix of functionalities and structures 10 of interest, plus water. For SOA compounds, the preliminary structures given in the Supplementary Materials (http://www.atmos-chem-phys-discuss.net/8/995/2008/ acpd-8-995-2008-supplement.pdf) were used; for the POA compounds, the structures in Fig. 1 were used. While the nitrate (ONO_2) group is an SOA functionality of interest here, it is not currently a UNIFAC group, and needed experimental thermodynamic data 15 do not exist. The UNIFAC group CHNO₂ was therefore substituted for nitrate during the fitting. Some degree of caution should therefore be exercised with considering predictions made using CP-Wilson.1 for nitrate-containing compounds. Overall, fitting to ζ_i^U values was considered adequate given the high general merits of the UNIFAC method, and because use of the 2p model and the assumed structures for the SOA compounds 20

- (Fig. 1) already represents a significant degree of approximation. At some future point, the CP-Wilson method could be re-fit using the extensive experimental ζ_i data set used by Fredenslund et al. (1975) to fit the UNIFAC method, and using new data for organic nitrate compounds (see above), the result perhaps designated as version CP-Wilson.2.
- ²⁵ A total of 13 338 ζ_i^U values were generated for various binary mixtures of the 41 compounds (40 organic compounds and water) over the mole fraction range 0.2 to 0.8 within the temperature range –10 to 50°C. Although some of these mixtures are not stable (i.e., would separate into two phases), that did not affect the inherent utility of the associated ζ_i^U values. Parameter optimization was performed on the total of 441 Λ_{ki}

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Phase separation considerations. Some of the surrogate compounds considered possess substantial polarity (e.g., the SOA products of isoprene with OH radical), and some are completely non-polar (e.g., the POA compound *n*-nonacosane). A liquid PM mixture containing significant proportions of both types of compounds will be unstable

⁵ ing 4446 ζ_i^U values for mixtures at 20°C was extracted from the entire pool to perform a preliminary optimization. Seven different sets of initial values for the Λ_k were involved in the fitting: all Λ_{ki} =250, all Λ_{ki} =500; all Λ_{ki} =1000; all Λ_{ki} =3000; all Λ_{ki} =5000; all Λ_{kj} =7000; and all Λ_{kj} =10000. During the fitting runs, the Λ_{kj} were restricted within $0 < \Lambda_{kj} < 10000$. (Due to the presence of the natural logarithm term in Eq. 12, it is required that each Λ_{ki} >0.) The mean and standard deviation of the seven χ^2 were 228 10 and 205. The best fit yielded χ^2 =24 and 0< Λ_{kj} <6000. The set of Λ_{kj} yielding χ^2 =24 was further refined by performing 10 additional optimizations in which the initial Λ_{ki} were varied randomly within $\pm 30\%$, but still so that $0 < \Lambda_{ki} < 6000$. The resulting best fit yielded χ^2 =22.8. Consideration of other initial Λ_{ki} sets outside the ±30% range did not improve χ^2 . In the second fitting stage, an optimization involving the entire set 15 of 13338 ζ_i^{U} values was performed five times using the best preliminary Λ_{ki} fit, but randomly varying the values within $\pm 30\%$ (but still so that $0 < \Lambda_{ki} < 6000$). The resulting χ^2 range was 124 to 127. The Λ_{kj} set giving $\chi^2 = 124$ was then used as input for a final optimization during which the step size was reduced three times. The resulting Λ_{kj} set gave $\chi^2 = 120$; further optimization did not reduce χ^2 . 20

2.4 $(N \cdot 2p)^{\zeta, \overline{MW}, \theta}$ Approach implementation

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parameters describing interactions among the 21 constituent groups. The fitting (optimization) occurred by use of the Levenberg-Marquardt algorithm (Levenberg, 1944) to minimize the function $\chi^2 = \sum_{i=1}^{n} \left(1 - \zeta_i^{\text{CPW.1}} / \zeta_i^{\text{U}}\right)^2$ wherein *n*=13338.

The optimization was performed in three stages. In the first stage, a subset contain-

relative to phase separation (Erdakos and Pankow, 2004). At constant P and T in a one-phase liquid system, phase separation will tend to occur when the liquid can find a lower Gibbs free energy by separating into two phases. Similarly, in a gas+liquid system (also at constant P and T) that is initially at equilibrium between the gas and a single-phase liquid phase separation in the liquid will tend to occur when the overall

- ⁵ a single-phase liquid, phase separation in the liquid will tend to occur when the overall system can find a lower Gibbs free energy by transformation into a three-phase system (a gas phase and two liquid phases); the transformation is likely to be accompanied by some net exchange with the gas phase of the partitioning compounds. Following Erdakos and Pankow (2004), when two liquid phases exist, then $\theta = \alpha$ and β (see above).
- ¹⁰ $K_{p,i}^{\theta}$ values were calculated by means of Eq. (2) using $K_{p,i}^{*}$ values. For most of the SOA surrogate compounds, actual chamber-derived $K_{p,i}^{*}$ and α_{i} values were used. For three SOA surrogate compounds (S23, S24, and S25), because the experimental data were not available, the Caltech Atmospheric Chemistry Mechanism (Griffin et al., 2002) and the Model to Predict the Multi-phase Partitioning of Organics (MPMPO) (Griffin et al.,
- ¹⁵ 2003) were used to predict a yield versus organic PM mass concentration (M_o) curve so that $K_{p,i}^*$ and α_i values could be predicted. For the POA surrogate compounds, chamber-derived $K_{p,i}^*$ do not exist, so $K_{p,i}^*$ values were calculated directly by means of Eq. (1) assuming $\zeta_i = 1$ and $\overline{MW} = MW_i$ and using the SIMPOL.1 method of Pankow and Asher (2007) to estimate $p_{L,i}^\circ(T^* = 293 \text{ K})$ based on structure.
- ²⁰ *PM mass calculations.* All 40 lumped surrogate compounds and water were assumed subject to G/P partitioning. \mathcal{F}_i (μ g m⁻³) represents the PM-associated level of *i*. (The related parameter F_i (ng m⁻³) has been used in prior work from this group.) If two PM phases are present, then

$$\mathcal{F}_i = \mathcal{F}_i^{\alpha} + \mathcal{F}_i^{\beta} \tag{14}$$

²⁵ $\mathcal{T}_i (=\mathcal{A}_i + \mathcal{F}_i)$ is the sum of the G- and total P-phase concentrations. At equilibrium the G-phase concentration $\mathcal{A}_i (\mu \text{gm}^{-3})$ can be calculated based on the value of $K_{\text{p},i}$ and

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 \mathcal{F}_i . In the case of phase separation,

$$\mathcal{T}_{i} = \frac{\mathcal{F}_{i}^{\theta}}{M_{\mathsf{TPM}}^{\theta} K_{\mathsf{p},i}^{\theta}} + \mathcal{F}_{i}$$

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wherein the first term on the RHS represents A_i , as based on equilibrium with either liquid phase ($\theta = \alpha$ or β), whichever is more convenient. $M_{\text{TPM}}^{\theta}(\mu \text{gm}^{-3})$ represents the total mass concentration of the θ phase. In this work, a constant RH was assumed in each case considered. At equilibrium, the statement of equality of water activity between the gas and particle phases is

 $\mathsf{RH}/100 = \zeta_{\mathsf{w}}^{\theta} x_{\mathsf{w}}^{\theta} \tag{16}$

which is thermodynamically equivalent to Eq. (1). As with Eq. (15), $\theta = \alpha$ or β , and in the absence of phase separation, the θ is dropped.

Iterational solutions of the overall G/P distribution problem represented by Eqs. (14– 16) were obtained by applying a liquid-liquid-equilibrium (LLE) flash calculation in each iteration as described by Chang and Pankow (2006). Using the index θ as needed, relationships used in consideration of the results are:

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$$M_{\rm o}^{\theta} = \sum_{\rm organici} \mathcal{F}_i^{\theta}$$
 (17)

$$M_{\rm o} = M_{\rm o}^{\alpha} + M_{\rm o}^{\beta} \tag{18}$$

$$M_{\rm w} = M_{\rm w}^{\alpha} + M_{\rm w}^{\beta} \tag{19}$$

$$M_{\rm TPM} = M_{\rm o} + M_{\rm w} = M_{\rm TPM}^{\alpha} + M_{\rm TPM}^{\beta}$$
(20)

where: $M_o^{\theta}(\mu \text{gm}^{-3})$ is total organic mass concentration associated with phase θ ; $M_o(\mu \text{gm}^{-3})$ is the total organic mass concentration over all PM phases; M_w^{α} and M_w^{β}

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(15)



 $(\mu g m^{-3})$ are the water mass concentrations associated with the α and β phases; M_w $(\mu g m^{-3})$ is the total water mass concentration over all PM phases; and M_{TPM} $(\mu g m^{-3})$ is the total PM mass concentration. As noted in Eq. (20), for the systems considered here, M_{TPM} $(\mu g m^{-3})$ is considered to be comprised of organic compounds and water (and no salt), and a maximum of two phases.

2.5 Cases

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Computational Efficiency (CE) Test Case (Liquid Phase Only). For a given group-contribution ζ_i prediction method, PM properties that affect the computation time are the number of constituent groups and the number of compounds. Thus, a one-phase liquid mixture at T = 300 K was invoked containing water and 40 organic compounds (the 25 preliminary surrogate SOA compounds in the Supplementary Materials, http://www.atmos-chem-phys-discuss.net/8/995/2008/ acpd-8-995-2008-supplement.pdf) and the 15 surrogate POA compounds in Fig. 1), all at $x_i = 1/41$. The fact that such a system would not remain a single phase at equilibrium at 200 K was appeared of the compounds of the compounds.

- ¹⁵ librium at 300 K was not a problem because the only issue was the speed of the ζ_i calculations. The speed of the calculations was compared for CP-Wilson.1 vs. five existing ζ_i estimation methods, namely UNIFAC, NRTL (Renon and Prausnitz, 1968), TK-Wilson (Tsuboka and Katayama, 1975), UNIQUAC (Abram and Prausnitz, 1975), and the unmodified Wilson equation. For each method, the CPU time required to compute the ζ_i values for 41 components in the mixture was obtained 100,000 different
- times, the large number allowing an averaging of the fluctuations in the CPU operation due to temporal variations in the system resource availability.

Performance Evaluation (PE) Case for ζ_i Prediction. A performance evaluation (PE) aerosol case at *T*=298 K for consideration of CP-Wilson.1 and UNIFAC in PM calculations with the (*N*·2p)^{ζ,MW,θ} approach was selected to involve all 40 final surrogate compounds in Table 1 (and Fig. 1) each at T_i =0.3 µg m⁻³, plus water at RH=50%. *Chamber Based (CB) Cases with RH=41 to 58%*, Δ*HC*=386 to 986µg m⁻³). Cases

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CB.1 to CB.3 involve α -pinene/O₃ at RH values in the range 41 to 58% (Table 2), and were studied experimentally in the chamber study of Cocker et al. (2001). With α -pinene as the only parent HC, only two surrogate product compounds from Table 1 (and Fig. 1) were considered, namely S3 and S4. The goal here was to allow a com-

⁵ parison of observed chamber PM levels with predictions based on: a) the $(N\cdot 2p)^{\zeta,\overline{\text{MW}},\theta}$ approach using the structures in Fig. 1 with CP-Wilson.1 for the ζ_i calculations; and b) the conventional $N\cdot 2p$ approach.

Hypothetical Varying Humidity Cases. To investigate RH effects at a lower Δ HC than in the CB cases, an α -pinene/O₃ series was considered assuming Δ HC=30 μ g m⁻³ with RH=20 to 80% at T=301 K. Also, a hypothetical mixed SOA+POA series was

developed with T = 301 K and two values of RH (5 and 80%); all compounds were considered subject to G/P partitioning. The individual \mathcal{T}_i values for the SOA+POA series are given in Table 3 with $\sum_{SOA} \mathcal{T}_i = 10 \,\mu \text{g m}^{-3}$ and $\sum_{POA} \mathcal{T}_i = 10 \,\mu \text{g m}^{-3}$.

3 Results

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15 3.1 Fit quality for CP-Wilson.1 relative to UNIFAC

Table 4 gives the best-fit values for the 441 Λ_{kj} parameters for CP-Wilson.1. The averaged unsigned percentage error for ζ_i relative to UNIFAC was calculated based on the 13 338 pairs of predicted $\zeta_i^{\text{CPW.1}}$ and ζ_i^{U} values according to:

$$\sigma_{\mathsf{FIT}}(\%) = \frac{\frac{13\,338}{\sum} \left| \frac{\zeta_i^{\mathsf{CPW.1}} - \zeta_i^{\mathsf{U}}}{\zeta_i^{\mathsf{U}}} \right| \times 100\%}{13\,338}$$
(21)

²⁰ The overall fit quality was very good (σ_{FIT} =6%). Figure 2 provides a plot of the 13 338 points for $\zeta_i^{CPW.1}$ vs. ζ_i^U . When ζ_i^U <1000, where >99% of the points are located, the values are in good agreement. (The good quality of the fit is masked to a considerable



extent by the fact that many of the 13 338 points are near the 1:1 line, and plot essentially on top of one another.) For $\zeta_i^U > 1000$, the agreement is still within a factor of ~2. Moreover, contributions to prediction errors for the mass totals given by Eqs. (17–20) are not likely to be caused simply by incorrectly estimating a large ζ_i value. Indeed, when there is one liquid phase, such an error can only be significant if the corresponding x_i is also of a significant magnitude. However, in that case, the phase would very likely be unstable relative to phase separation, in which case the *i*-related prediction error for the mass total would become small because most of the *i* would retreat into the new, second phase in which ζ_i would be relatively close to 1 and thus reliably estimated.

3.2 CP-Wilson.1 vs. other methods for the Computational Efficiency (CE) case

Table 5 compares the CPU requirements of CP-Wilson.1 with five other ζ_i methods for the CE case. CP-Wilson.1 gave the best result. The economy of this method is achieved by the combination of its group contribution nature (21 groups for the CE case instead of 41 compounds), and its relatively small need for logarithm and double summation operations. While not implemented here, the computational efficiency of a CP-Wilson.1 code can be assisted by utilizing a lookup table to evaluate the logarithm term in Eq. (12).

3.3 CP-Wilson.1 vs. UNIFAC for Performance Evaluation (PE) case

²⁰ When CP-Wilson.1 is used in the $(N \cdot 2p)^{\zeta, \overline{MW}, \theta}$ approach to predict ζ_i in the liquid PM formed in the PE case (all $\mathcal{T}_i = 0.3 \,\mu \text{g m}^{-3}$ for the organic components, and RH=50%), two phases are revealed as being present in the PM at equilibrium; use of UNIFAC leads to the same result. For each phase θ , relative to UNIFAC, the unsigned prediction



difference (%) for x_i^{θ} is defined

$$\delta_{x,i}^{\theta} = \frac{\left| x_{i}^{\theta, \text{CPW.1}} - x_{i}^{\theta, \text{U}} \right|}{x_{i}^{\theta, \text{U}}} \times 100\%$$

where the superscripts on x_i denote the phase and ζ_i method. For the α phase, a plot of $\delta_{x_i}^{\alpha}$ vs. $\log_{10} x_i^{\alpha,0}$ is given in Fig. 3a for the 40 compounds and water. A corresponding plot for the β phase is given in Fig. 3b. Consistent with the results in Fig. 2, δ_{v}^{α} , is small when $x_i^{\alpha,U}$ > 0.01, and the corresponding $\delta_{x_i}^{\beta}$ are small when $x_i^{\beta,U}$ > 0.01.

Table 6 provides observed and predicted results for M_{o} , M_{w} , and M_{TPM} for the PE case. While UNIFAC does not provide perfect estimates of ζ_i , the similarity of the predicted results as obtained using CP-Wilson.1 vs. UNIFAC supports the view discussed above that errors associated with large ζ_i^{θ} are not likely to have significant effects on 10 the quality of predictions for gross parameters such as M_o , M_w , and M_{TPM} (= $M_o + M_w$). The extent to which an error in a given x_i^{θ} value translates into an error in M_{TPM} depends on the magnitude of x_i^{θ} and on the size of the θ phase. For UNIFAC-based predictions, the fraction (%) of the total PM phase identified with i in the θ -phase equals ¹⁵ $\mathcal{F}_{i}^{\theta,U} \times 100\%/M_{TPM}^{U}$. While UNIFAC is certainly also subject to increasing prediction error as any given ζ_i increases, it is again perhaps the best available benchmark for evaluating the results obtained using CP-Wilson.1. Thus, as an means to evaluate the implications of incorrectly predicting \mathcal{F}_{i}^{θ} we define

$$\varepsilon_i^{\theta} = \left| \frac{(\mathcal{F}_i^{\theta, \text{CPW.1}} - \mathcal{F}_i^{\theta, \text{U}}) \times 100\%}{(M_{\text{TPM}})^{\text{U}}} \right|$$
(23)

²⁰ If ε_i^{θ} is small, either $\mathcal{F}_i^{\theta, CPW, 1} \approx \mathcal{F}_i^{\theta, U}$, or both are small relative to M_{TPM} . Figures 4a and b provide plots of ε_i^{θ} vs. $\log_{10} x_i^{\theta, U}$ for the PE case. For both $\theta = \alpha$, and $\theta = \beta$, ε_i^{θ} becomes small as $\log_{10} x_i^{\theta, U}$ increases. This is a consequence of the fact that when 8, 995-1039, 2008

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phase separation occurs, if ζ_i^{α} is large (and thus subject to some error regardless of the prediction method used), then ζ_i^{β} is relatively close to 1 (and vice versa), the overall result being a thermodynamically driven minimization of the mass amount of *i* in the less-hospitable phase where ζ_i is more difficult to predict (see discussion above). This type of hyperbolic behavior in $\log_{10} \zeta_i^{\alpha}$ vs. $\log_{10} \zeta_i^{\beta}$ is clearly evident in Fig. 5. For compounds that fall in the middle of the hydrophobicity \leftrightarrow hydrophilicity scale, both ζ_i^{α} and ζ_i^{β} are neither near 1 nor very large. E.g., for P6, both ζ_i^{α} and ζ_i^{β} are of order 10 because P6 is not particularly "comfortable" in either phase.

3.4 Predictions for Chamber Based (CB) and hypothetical cases

CB Cases for α -Pinene/O₃ with RH=41 to 58%. Table 2 provides M_{TPM} values mea-10 sured by Cocker et al. (2001) along with the predicted values of M_o , M_w , and M_{TPM} using the $(N \cdot 2p)^{\zeta, \overline{\text{MW}}, \theta}$ approach with CP-Wilson.1 and the assumed structures for products S3 and S4. For all three CB cases, use of CP-Wilson.1 indicates a single PM phase. (In this and all other respects for these cases, UNIFAC gives similar results.) At these moderate RH values, water uptake is low and ζ_{S3} and ζ_{S4} values are ~1.3, i.e., relatively close to unity. The prediction errors for M_{TPM} as compared to the chamber experiments ranged from -17 to 3% for the three cases. Table 2 also provides M_0 as predicted using the N·2p approach for which, as has been noted, all $\zeta_i \equiv 1$, $M_0 \equiv M_{\text{TPM}}$, and MW = constant. The prediction errors for the N·2p approach ranged from -22 to -5%. The two approaches give nearly the same results because the as-20 sumed structures for S3 and S4 have similar polarities and MW, values, and the water uptake is relatively low. Also, the good agreement under these conditions between the experimental results of Cocker et al. (2001) and the predictions based on the $N \cdot 2p$ approach indicate good consistency between the Cocker et al. (2001) yield results and the $K_{p,i}$ and α_i values given in Table 1 for α -pinene/O₃. 25

Hypothetical α *-pinene/O*₃ *Series*. Results based on the *N*·2p and (*N*·2p)^{ζ ,MW,\theta} ap-



proaches for the α -pinene/O₃ series are given in Fig. 6. As with the CB cases, use of CP-Wilson.1 in the $(N\cdot2p)^{\zeta,\overline{\text{MW}},\theta}$ approach indicates a single PM phase for the entire RH range (20 to 80%). However, the difference in the M_o predictions obtained using the *N*·2p approach with no water uptake and $(N\cdot2p)^{\zeta,\overline{\text{MW}},\theta}$ with water uptake is much greater in the Fig. 6 series than in the CB cases: M_o by the $(N\cdot2p)^{\zeta,\overline{\text{MW}},\theta}$ approach in Fig. 6 increases by more than $3 \times as$ RH increases from 20 to 80%. The increase in M_o by the $(N\cdot2p)^{\zeta,\overline{\text{MW}},\theta}$ approach is driven mostly by the decreasing $\overline{\text{MW}}$ brought about by water absorption ($\overline{\text{MW}}$ =151 g mol⁻¹ at RH=20%, and 64 g mol⁻¹ at RH=80%), but is also compounded somewhat by the behavior of ζ_{S3} , which changes from 1.6 at RH=20%, to 1.1 at RH=80% (ζ_{S4} remains essentially unchanged at 1.3).

For mono-phasic PM, based on Liang and Pankow (1996), the fraction of i in the particle phase is given by

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$$f_{p,i} = \frac{K_{p,i}M_{\text{TPM}}}{1} + K_{p,i}M_{\text{TPM}}$$
(24)

(Donahue et al., 2006 denote $f_{p,i}$ as ξ_i .) When $K_{p,i}M_{TPM}$ is large relative to 1, $f_{p,i} \approx 1$, and the contribution that *i* makes to M_{TPM} is relatively insensitive to changes in $K_{p,i}$ that may be caused by changes in RH, T, and other factors. However, as $f_{p,i}$ decreases away from 1, that sensitivity increases. Thus, as compared to the CB cases and as compared to the monoterpene cases considered by Seinfeld et al. (2001), M_{TPM} in the series in Fig. 6 is much lower, and M_0 is thus much more sensitive to RH. Pankow and Chang (2008)² provide additional perspective on why the sensitivity of M_0 and M_{TPM} predictions will tend to increase as the levels of condensable compounds decrease.

SOA+POA System. Table 7 provides predicted results assuming the Table 3 levels of SOA+POA compounds at T=301 K for RH=5% and 80%. In each of these two

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²Pankow, J. F. and Chang, E. I.: Variation in the sensitivity of predicted levels of atmospheric organic particulate matter (OPM), submitted, 2008.

cases, the $(N \cdot 2p)^{\zeta, MW, \theta}$ approach using CP-Wilson.1 predicts two liquid phases in the PM, with the α phase containing mostly SOA compounds and water, and the β phase containing mostly POA compounds and little water. At RH=5%, for the major components in each PM phase, $\zeta_i^{\theta} \approx 1$. Because of the considerable mutual exclusion of the SOA and POA compounds, the effective size of the PM compartment at RH=5% is significantly lower than is predicted using the N·2p approach: M_w and M_o based on the $(N\cdot 2p)^{\zeta,\overline{\text{MW}},\theta}$ approach are 0.001 and 6.5 μ g m⁻³. By the N·2p approach, the corresponding values are 0 and 9.2 μ g m⁻³. For the RH=80% case, significant mutual exclusion of the SOA and POA compounds still operates. However, significant water uptake into the α phase is now predicted. This tends to increase the $f_{p,i}$ values of the 10 more polar compounds because M^{α}_{TPM} is relatively larger and $\overline{\text{MW}}^{\alpha}$ relatively smaller than at RH=5%. Moreover, because of the resulting increased M_w^{α} value, several of the rather polar compounds take on significantly reduced ζ_i^{α} values (for S8, 10, 12, 14, 20 and P11, the range for ζ_i^{α} is 0.2 to 0.6). The overall result is that at RH = 80%, $M_{\rm w}$ and $M_{\rm o}$ based on the $(N \cdot 2p)^{\zeta, \overline{\rm MW}, \theta}$ approach are 1.3 and 10.0 μ g m⁻³ vs. 0 and 9.2 μ g m⁻³ 15 by the $N \cdot 2p$ approach.

4 Conclusions

The approximations for the particulate matter (PM) phase incorporated in the multiple lumped "two-product" *N*·2p approach for SOA PM (*i.e.*, all ζ_i =1, $\overline{\text{MW}}$ is constant, and no water uptake occurs at ambient RH levels) will become increasingly problematic as M_0 levels decrease. Also, the approximation of a single-absorbing phase as utilized in the *N*·2p approach can become invalid when RH levels are high, and/or when significant levels of both SOA and POA compounds are present. The structures proposed here for partitioning SOA and POA compounds will allow first stage usage of the $(N\cdot2p)^{\zeta,\overline{\text{MW}},\theta}$ approach for ambient PM modeling; the CP-Wilson.1 ζ_i -prediction



method developed here allows consideration of computationally intensive space-time domains.

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Table 1. Assumed properties of 25 SOA and 15 POA surrogate compounds, and water.

SC	DA Compo	ounds		chamb	er-derived pa	rameters				other parar	neters	
compound	reaction	parent HC	oxidant	<i>Т</i> * (К)	$\mathcal{K}^*_{p,i}(\mathcal{T}^*)$	α,	ref.	MW ^{* a} (g mol ⁻¹)	MW ^b (g mol ⁻¹)	CP- Wilson parame- ter C ^c _i	Δ <i>H</i> _{vap} (303) ^d (kJ mol ⁻¹)	p _L °(<i>T</i> *) ^d (atm)
S1 S2	1	<i>a</i> -pinene	ОН	310	0.171 0.004	0.038 0.326	А	189 189	188 190	70.90 59.07	99.1 85.7	7.38 E-10 3.82 E-08
S3 S4	2	α -pinene	O ₃	310	0.088 0.0788	0.125 0.102	А	194 194	214 174	41.76 69.75	74.3 89.7	1.34 E-09 1.52 E-09
S5 S6	3	β -pinene	ОН	310	0.044 0.0049	0.13 0.041	А	179 179	186 172	56.44 52.63	100.6 78.6	3.63 E-09 2.75 E-08
S7 S8	4	β -pinene	O ₃	310	0.195 0.003	0.026 0.485	А	188 188	202 174	56.92 64.41	103.9 77.1	7.32 E-10 4.18 E-08
S9	5	β -pinene	NO ₃	310	0.0163	1.000	А	245	245	69.29	80.3	7.61 E-09
S10 S11	6	isoprene	ОН	295	0.0086 1.62	0.232 0.029	В	177 177	136 218	52.68 76.27	90.3 87.5	1.36 E-08 8.68 E-11
S12 S13	7	limonene	ОН	310	0.055 0.0053	0.239 0.363	А	195 195	188 202	51.90 43.32	79.8 90.0	2.33 E-09 2.57 E-08
S14 S15	8	ocimene	ОН	310	0.174 0.0041	0.045 0.149	A	152 152	146 158	32.46 37.15	105.5 90.2	1.06 E-09 3.36 E-08
S16 S17	9	terpinene	ОН	310	0.081 0.0046	0.091 0.367	A	174 174	202 146	62.45 24.88	111.3 79.7	1.46 E-09 3.52 E-08
S18 S19	10	toluene	ОН	310	0.053 0.0019	0.071 0.138	C,D	173 173	148 197	38.63 48.12	95.6 81.4	2.54 E-09 8.21 E-08
S20 S21	11	xylene	ОН	310	0.042 0.0014	0.038 0.167	C,D	187 187	176 197	31.03 48.12	87.5 81.4	2.64 E-09 8.21 E-08
S22	12	humulene	ОН	310	0.0501	1.000	А	270	270	72.94	73.9	1.80 E-09
S23 S24	13	2-ring PAH	ОН	298	0.015 0.002	1.000 1.000	Е	175 175	186 164	68.19 47.75	94.5 81.5	8.14 E-09 7.05 E-08
S25	14	C ₁₆ n-alkane	ОН	298	0.0229	1.000	Е	301	301	94.58	100.9	3.28 E-09

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POA Compo	unds					
Compound		K [*] _{p,i} (293) ^c	MW _i (g mol ⁻¹)	CP-Wilson parameter C_i^d	$\Delta H_{vap}(303)^e$ (kJ mol ⁻¹)	p _L °(293) ^e
P1	2,6-naphthalene diacid	101.7	216	69.62	118.9	1.09 E-12
P2	benzo[ghi]perylene	43.83	276	119.40	112.7	1.99 E-12
P3	butanedioic acid	0.0025	118	26.59	84.0	8.31 E-08
P4	$17(\alpha)$ H-21(β)H-hopane	72.83	412	172.24	123.1	8.01 E-13
P5	<i>n</i> -nonacosane	33.62	409	166.96	149.2	1.75 E-12
P6	octadecanoic	1.142	284	107.03	123.4	7.41 E-11
P7	phthalic acid	0.4801	166	47.60	101.4	3.01 E-10
P8	UCM2 (unresolved complex mixture 2)	10.45	390	162.46	132.4	5.90 E-12
P9	monoglyceride	434.0	330	123.00	138.8	1.68 E-13
P10	triglyceride	1.72 E+17	860	299.04	280.0	1.63 E-28
P11	levoglucosan	0.1670	162	57.14	94.4	8.88 E-10
P12	UCM1 (unresolved complex mixture 1) ^f	1.42 E-05	210	87.12	79.2	8.07 E-06
P13	UCM3 (unresolved complex mixture 3)	1.64 E+05	487	202.35	158.2	3.01 E-15
P14	hexadecanoic acid	0.1427	256	95.49	114.3	6.58 E-10
P15	glycerol	0.0005	92	39.10	78.0	5.02 E-07
Water	H ₂ O		18	7.15	g	g

Table 1. Continued.

Footnotes:

^a Calculated as the mean of the inferred MW_i values

^b Inferred based on chamber data using iterative process outlined in Eqs. (5–10).

^c Although these $K_{p,i}$ values were not determined in chamber experiments, the asterisk is maintained for clarity regarding usage in Eq. (2).

^d Calculated based on Eq. (13).

^e Calculated based on parameters given in Pankow and Asher (2007).

^f Included for the sake of completeness, even though its $K_{p,i}^*$ (293) value is so low that it will not contribute significantly to OPM levels.

^g Not required since all water calculations were made based on specified RH values.

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Table 2. Results for chamber-based (CB) cases with α -pinene/O₃ as measured by Cocker et al. (2001) and as predicted using the *N*·2p approach and using the (*N*·2p)^{ζ , \overline{MW} , θ} approach with the CP-W.1 method for prediction of ζ_i values.

				Measured		N⋅2p Pre	edictions		(N·2p) ^{ζ, ₩₩,6}	Predictions	3
case	7 (K)	ΔHC	RH	M _{TPM}	Mo	M _w	M _{TPM}	error ^a in	Mo	M _w	M _{TPM}	error ^a in
		$(\mu g m^{-3})$	(%)	(µg m ^{−3})	$(\mu g m^{-3})$	$(\mu g m^{-3})$	$(\mu g m^{-3})$	M _{TPM}	$(\mu g m^{-3})$	$(\mu g m^{-3})$	$(\mu g m^{-3})$	M _{TPM}
CB.1	301.8	386.3	41.0	86	82	0 ^b	82	-5%	83	6	89	+3%
CB.2	301.8	675.0	57.7	177	148	0 ^b	148	-16%	150	18	168	-5%
CB.3	302.7	986.5	37.3	281	218	0 ^b	218	-22%	219	13	232	-17%

Footnotes:

^a Error based on measured value of Cocker et al. (2001).

^b by definition

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SOA	Compounds				8, 995–	1039,
Rxn.	Parent	Oxidant	Compound	$\mathcal{T}_i ~(\mu { m g~m}^{-3})$		
1	a-pinene	ОН	S1 S2	0.104 0.896	OPM fo	ormat
2	a-pinene	O ₃	S3 S4	0.551 0449	varyin hu	ig rel midi
3	β -pinene	ОН	S5 S6	0.380 0.120	E. I. C	Chang
4	β -pinene	0,	S7	0.025	J. F.	Pank
5	β -pinene	NO3	S9	0.500		
6	isoprene	ОН	S10 S11	0.111 0.014	Tit	le Pag
7	limonene	ОН	S12 S13	0.397 0.603	Abstract	In
8	ocimene	ОН	S14 S15	0.029	Conclusions	s R
9	terpinene	ОН	S16	0.025	Tables	
10	toluene	ОН	S18	0.849	14	
		0	S19	1.650	•	
11	xylene	OH	S20	1.430	Back	
12	humulene	ОН	S22	0.125	Full S	creen /
13	2-ring PAH	ОН	S23	0.125		
			S24	0.125	Printer-fr	iendly \

Table 3. Total mass concentration \mathcal{T}_i values for the hypothetical SOA + POA cases.

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Table 3. Continued.

POA Compounds		
2,6-naphthalene diacid	P1	0.083
benzo-ghi-perylene	P2	0.083
butanedioic acid	P3	0.083
$17(\alpha)$ H-21(β)H-hopane	P4	0.083
n-nonacosane	P5	0.083
octadecanoic acid	P6	0.083
phthalic acid	P7	0.083
UCM2	P8	3.000
monoglyceride	P9	0.083
triglyceride	P10	0.083
levoglucosan	P11	0.083
UCM1	P12	3.000
UCM3	P13	3.000
hexadecanoic acid	P14	0.083
glycerol	P15	0.083

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Table 4. Fitted interaction parameters (Λ_{ki}) for 21 groups.

									-	
	CH3	CH ₂	CH	С	CH=CH	(Aromatic C)-H	Aromatic C	(Aromatic C)-CH ₃	(Aromatic C)-CH ₂	OH
CH ₃	3.88E+02	3.75E+01	9.26E+01	1.96E+01	6.73E+02	1.31E+02	1.75E+01	5.44E+02	1.21E+02	2.34E+03
CH ₂	8.74E+02	3.19E+02	3.85E+02	3.40E+02	3.94E+01	6.59E+01	1.56E+02	5.48E+02	3.66E+02	1.89E+03
CH	6.39E+02	2.42E+02	3.02E+02	2.71E-02	3.33E-07	2.15E+01	1.50E+02	5.05E+02	4.22E+02	1.52E+03
С	9.70E+02	2.05E+02	3.65E+02	1.20E-02	6.33E+02	4.89E+02	3.92E+02	4.95E+02	5.19E+02	8.80E+02
CH=CH	3.52E+00	1.23E+02	9.92E+01	3.37E+01	6.52E-03	2.01E+02	1.37E+01	6.78E-05	2.95E-02	7.22E+02
(Aromatic C)-H	5.44E+02	4.49E+02	4.27E+02	7.76E-02	4.83E+01	2.55E+02	4.10E+02	3.59E+02	3.06E+02	7.54E+02
Aromatic C	7.70E+02	4.95E+02	4.06E+02	6.70E-05	3.19E+02	2.33E+01	2.98E+02	3.97E+02	3.28E+02	1.72E+03
(Aromatic C)-CH ₃	3.27E+01	3.12E+01	2.57E+01	1.98E+01	2.04E+00	8.87E+00	6.52E+01	1.63E+02	1.23E+02	5.67E+02
(Aromatic C)-CH ₂	8.08E+01	7.63E+01	1.70E+01	4.39E+01	2.83E-01	2.06E+01	7.85E+01	1.42E-01	1.70E+02	8.87E+02
OH	9.39E+01	7.06E+01	5.13E+01	1.79E+02	4.30E+02	1.92E+02	2.15E-04	2.90E-03	2.22E-02	8.20E+02
H ₂ O	5.88E+01	7.68E+01	4.15E+01	1.19E+02	1.02E-01	2.49E-01	1.38E+02	5.90E-03	3.62E+02	2.31E+02
(Aromatic C)-OH	8.85E+01	3.89E+01	1.34E-02	1.77E-05	4.47E+03	1.32E+02	2.33E+02	8.82E-04	3.22E-02	6.12E+02
CH ₃ CO	6.08E+01	7.34E+01	4.81E+02	4.40E+02	7.32E+02	1.82E+02	2.17E+01	9.17E+02	1.36E-12	8.79E+02
CH ₂ CO	1.75E+00	3.94E+01	7.04E+01	6.78E-02	1.94E+02	5.62E+01	9.94E-04	1.36E+00	1.17E-12	3.53E+02
CHO	9.86E+02	3.67E+02	1.53E-02	2.13E+02	5.00E-02	3.57E+02	1.08E-01	2.97E-12	3.24E-12	9.65E+02
CH2COO	8.59E+02	3.81E+02	1.53E+00	5.27E-07	4.40E-01	7.33E+02	4.83E-01	4.46E+02	2.64E-02	7.65E+02
CH ₂ O	4.55E+00	1.43E+02	1.24E+00	8.39E+01	5.47E+02	8.17E+01	1.01E-12	5.64E+02	1.22E-12	6.98E+02
CH-O	1.95E+03	9.69E+02	1.54E+02	1.00E+00	1.50E+03	4.01E+02	6.40E+01	7.16E+02	9.19E+02	2.23E+02
COOH	4.30E+02	1.23E+02	4.79E-05	6.64E+02	1.60E+03	3.78E+02	1.38E-05	5.49E-01	2.80E-03	1.14E+03
NO ₃	3.44E+00	8.56E+01	1.47E+02	9.11E-02	1.02E-12	5.41E-04	8.61E+01	2.08E+02	6.45E+02	4.26E+02
(Aromatic C)-NO ₂	4.18E+02	2.63E+02	1.46E+02	2.45E-01	1.96E-03	2.33E+02	7.32E-02	1.79E-02	4.76E+02	1.19E+03
_										

H₂O Aromatic OH CH₃CO CH2CO CHO CH₂COO CH₂O CH-O COOH NO₃ (Aromatic C)- NO₂ 2.71E+03 3.50E+03 1.94E+03 1.94E+03 1.25E+03 7.66E+02 6.04E+02 1.23E+03 1.69E+03 2.08E+03 4.51E+02 1.94E+03 2.47E+03 1.79E+03 1.18E+03 1.03E+03 9.28E+02 5.16E+02 4.19E+02 1.63E+03 1.12E+03 3.23E+02 1.50E+03 2.03E+03 5.53E+02 4.96E+02 9.81E+02 1.24E+03 4.19E+02 3.01E+02 1.25E+03 4.09E+02 7.65E-08 1.25E+03 1.34E+03 7.47E+02 7.64E+02 4.86E+02 2.51E+02 1.69E-02 1.19E+03 3.48E+02 7.63E+02 6.05E+02 1.11E+03 2.77E-01 7.88E-04 2.29E-01 4.32E+02 7.14E+02 8.36E-02 3.09E-12 2.66E+02 4.90E+02 5.08E+02 1.12E+03 1.01E+03 7.18E+02 6.13E+02 3.54E+02 2.13E+02 4.15E+02 5.35E+02 3.84E+02 3.65E+02 2.44E-01 2.15E+03 1.69E+03 1.26E+03 9.01E+02 1.04E+03 4.19E+02 4.18E+02 1.09E+03 1.42E+03 1.24E+03 5.70E+02 7.21E+02 9.31E+02 1.01E+02 4.45E+02 3.71E+02 1.73E+02 1.59E-12 1.27E-12 4.75E+02 2.12E+02 1.07E-01 4.25E+02 2.96E+02 3.76E-12 3.67E+02 2.87E+02 1.13E+03 2.04E+03 3.83E+02 1.10E-02 1.07E+03 4.02E-03 1.52E+03 1.45E-12 2.00E+03 1.21E+03 1.35E+03 1.04E+03 4.19E+01 6.16E+02 9.18E+02 7.23E+02 2.41E+03 1.28E+03 1.09E-12 3.58E+02 2.62E+02 1.68E+02 6.76E+01 1.07E-12 1.10E-12 2.39E+02 1.45E+03 8.47E-02 4.98E+02 3.96E+03 1.00E+03 2.01E-04 1.36E-02 4.14E-01 3.96E+02 4.78E+02 1.52E+03 2.00E-12 4.27E-04 2.25E+03 1.55E+03 1.42E+03 5.74E+02 1.40E+03 1.37E+03 9.32E+02 8.14E+02 1.03E+03 1.40E+02 5.79E-02 6.15E+02 5.82E+02 6.60E+02 2.98E+02 3.14E+02 3.23E+02 6.32E+02 4.43E+02 3.44E+02 8.54E-02 4.30E-03 2.27E+03 1.82E+03 9.71E+02 4.01E+02 7.51E+02 5.14E+02 7.41E+02 9.91E+02 5.55E+02 5.31E+03 2.79E+03 2.18E+03 9.61E-12 2.49E-01 3.71E+02 9.29E+02 1.13E+03 3.94E+00 1.05E-11 2.42E-03 1.10E-12 2.15E+00 3.08E+02 1.03E-12 1.94E+03 4.98E-02 8.19E+02 9.52E-12 4.40E+02 4.40E+02 8.98E+02 1.22E+03 8.06E-07 5.01E+02 2.00E-04 4.58E+02 5.88E+02 1.47E+00 4.10E+03 1.78E+03 1.78E+03 3.46E+02 3.03E+03 3.39E+03 1.98E+03 3.44E+03 7.47E+02 1.19E+02 1.15E+03 7.10E+02 1.03E-02 1.01E-12 1.09E+03 3.67E+03 2.98E+03 1.32E+03 1.13E+03 8.20E+02 2.21E+02 3.86E-04 1.22E+00 1.25E+00 1.34E+03 4.59E+02 1.29E+03 2.11E-01 2.39E+03 1.05E+03 3.51E+02 1.08E-12 2.62E-12 6.03E+02 3.60E+02 4.78E+02 8.69E+02 3.43E-02 1.51E+0.3

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Table 5. Comparison of relative computer processing time required for six ζ_i prediction methods.

ζ_i Method	Method type	Number of groups or compounds	Relative computer processing time
CP-Wilson.1	group	21	0.1
Wilson	compound	41	0.6
UNIQUAC	compound	41	0.6
TK-Wilson	compound	41	0.7
NRTL	compound	41	0.8
UNIFAC	compound	21	1.0

Table 6. Comparison of predictions for the performance evaluation (PE) case by the $(N \cdot 2p)^{\zeta, \overline{\text{MW}}, \theta}$ approach using the CP-Wilson.1 method and the UNIFAC method (*T*=298 K, RH=50%).

	<i>ζ</i> , Μ	ethod
Result	CP-Wilson.1	UNIFAC
Number of PM phases	2	2
$M_{ m o}^{lpha}$, $M_{ m o}^{eta}$ (μ g/m ³)	3.63, 1.79	3.62, 1.79
$M_{\rm o} = M_{\rm o}^{\alpha} + M_{\rm o}^{\beta} \ (\mu {\rm g/m}^3)$	5.42	5.41
$M_{\rm w}^{\alpha}$, $M_{\rm w}^{\beta}$ (μ g/m ³)	0.29, 0.001	0.29, 0.0005
$M_{\rm w} = M_{\rm w}^{\alpha} + M_{\rm w}^{\beta} \ (\mu {\rm g/m}^3)$	0.29	0.29
$M^{lpha}_{ m TPM}$ (μ g/m ³)	3.92	3.92
$M_{\rm TPM}^{\beta}$ (μ g/m ³)	1.79	1.79
M _{TPM} (μg/m ³)	5.71	5.71



	<i>N</i> ·2p Prediction				$(N\cdot 2p)^{\zeta, \overline{MW}, \theta}$ Prediction				
RH	Mo	M _w	M _{TPM}	PM	Mo	<i>M</i> _w	M _{TPM}	PM	phase mass
(%)	$(\mu g m^{-3})$	$(\mu g m^{-3})$	$(\mu g m^{-3})$	phases	(µg m ^{−3})	$(\mu g m^{-3})$	$(\mu g m^{-3})$	phases	distribution
									$M^{lpha}_{ m TPM}/M^{eta}_{ m TPM}$
5	9.23	0 ^a	9.23	1 ^a	6.48	0.001	6.48	2	0.19/6.29
80	9.23	0 ^a	9.23	1 ^a	10.00	1.28	11.28	2	5.02/6.26

Table 7. Results for the hypothetical SOA + POA cases at 300 K as predicted using the *N*·2p approach and using the $(N \cdot 2p)^{\zeta, \overline{\text{MW}}, \theta}$ approach with the CP-Wilson.1 method for prediction of ζ_i values. (See Table 3 for all T_i values.).

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Footnotes: ^a by definition



Fig. 1a. Molecular structures of lumped SOA products and surrogate POA compounds. ${}^{t}R_{1}=C_{15}$ alkyl chain; $R_{2}=C_{17}$ alkyl chain; $R_{3}=C_{17}$ alkyl chain with one double bond. For some structures, ether linkages have been included to fine tune the estimated polarity, even when such linkages may be unlikely consequences of the relevant oxidation reactions.

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Fig. 1c. Continued.

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Fig. 1d. Continued.

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Fig. 2. ζ_i calculated by CP-Wilson.1 vs. ζ_i calculated by UNIFAC for 13338 points used in the fitting of CP-Wilson.1 to UNIFAC.

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Fig. 3. Unsigned percentage difference between the values of x_i^{θ} as predicted using CP-W.1 and UNIFAC plotted vs. log $x_i^{\theta,U}$ (U=UNIFAC) for $\theta = \alpha$ and for $\theta = \beta$ in the performance evaluation (PE) case.

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