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Holger Tost
Editor, Atmospheric Chemistry and Physics

Re: Response to Reviews

Manuscript Number:

Manuscript Title: Metrics to quantify the importance of mixing state for CCN activity

Manuscript Authors: Joseph Ching, Jerome Fast, Matthew West, and Nicole Riemer

Dear Sir,

Please find attached our responses to your comments and the reviewer's comments regarding the above paper. We have updated the manuscript accordingly and marked the changes in blue.

The main changes consist of the addition of a new section in the paper (Section 6, including new Figures 10 and 11). This section compares the mixing state metric χ to other measures of heterogeneity in hygroscopicity, as suggested by Reviewer 2.

Yours sincerely,

Joseph Ching

1 Response to Reviewer #1's comments

We greatly appreciate the reviewer's comments. We revised the manuscript accordingly with changes marked in blue. Our responses are as follows:

(1.1) Figure 1 and Table 1 and 2 are 100% identical to figures and tables in Riemer et. al. 2013. Even with the authors and the Journal being the same I would consider this a questionable practice. The least the authors can do is to add a note to the captions saying "Taken from Riemer et. al. 2013".

We updated the captions to Figure 1 and Tables 1 and 2 accordingly.

We also added text to emphasize that we summarize the important key points from Riemer and West (2013) for the convenience of the reader:

- page 3, line 4: "The salient points are summarized as follows."
- page 3, lines 7–8: "From this quantity, all other mass-related quantities can be defined, as detailed in Riemer and West (2013) and here listed in Table 1, and the diversity metrics can be constructed as shown in Table 2."

(1.2) Most presented findings are statistical observations. I am convinced it would add great value to the manuscript if the authors discussed potential physical origins of their findings. E.g. why do different populations experience different amounts of error cancellations (page 12, line 9)? What physical characteristic might explain the dependence of the relative error on supersaturation threshold (page 12&13)?

We have made a number of changes and clarifications to the paper in an attempt to explain the physical origins of our findings. We renamed Section 5 to "Relationship of error in CCN concentration and mixing state index χ " to highlight that this is where we will discuss this relationship. In this section we discuss Figure 5, which clarifies that "Error is caused only by the *difference* in the number of particles that activate in the reference case, but do not activate after composition averaging [...], and the cases that activate after composition averaging, but did not activate in the reference case."

In this section we also explain the physical basis of zero error for internally mixed populations as the fact that "For populations with $\chi = 100\%$, the particles' composition is the same across the entire population, and hence [...] the number concentrations in quadrants B and D are zero." Correspondingly, we explain the physical basis of the error for partially-internally mixed populations as depending on "both the extent of distortion of the critical supersaturation distribution and the supersaturation threshold."

To explain the dependence of the relative error on the supersaturation threshold, we clarified that "for a given supersaturation threshold and a given χ value, different populations experience different amounts of the error cancellations" described above. A detailed investigation of this is provided by Figure 8, which together with Figure 5 provides physical understanding of the cancellation phenomenon. We explained in the paper that "the change in particle number concentration per supersaturation interval is smaller for s_c of 0.1% compared to $0.3 < s_c < 1.5\%$ " due to the shape of the population \mathbb{P} in Figure 5.

(1.3) Figure 7: the x-axis looks like a ratio rather than %. Also, the caption mentions insets which do not seem to be there.

Thanks for pointing this out. We fixed both of these issues.

2 Response to Reviewer #2's comments

We thank the reviewer for their comments and suggestions. We revised the manuscript accordingly with changes marked in blue. Our responses are as follows:

(2.1) Importance of the mixing state metrics: My major concern is how the new metrics (χ) will help to quantify the mixing state effect? In this study, the determination of mixing state effect was done by comparing CCN predictions of cases with and without composition averaging. If I understand correct, it means that the mixing state effect is determined without the metric χ . So, why would we need such a parameter if it is not even used?

The reviewer is correct—since we have all the per-particle information we can determine the error, and if this was all we wanted to do then the metric χ (or any other mixing state metric) is not needed.

However, our goal is to relate the error in CCN concentration due to the internal mixture assumption to a quantitative measure of mixing state. The paragraph in the introduction, p. 2, line 13 states this goal, and we added text to clarify this further, p. 2, lines 18–20: “The central question that we address is: For aerosol populations of a given mixing state, what magnitude of errors can we expect for estimating CCN concentrations when assuming that the population is internally mixed?”

(2.2) Performance of the mixing state metrics: One of my questions during my reading is that if a single χ corresponds to a unique error in CCN predictions and if it can be used in the CCN prediction or even better than existing parameters. The authors answered my first question, and showed that the relationship of χ and the error in CCN predictions is not unique. According to the size-resolved hygroscopicity distribution in Fig. 4, there are two kappa modes and my feeling is that the fraction of the low hygroscopic mode (F_{LH}) is a critical parameter for the errors when neglecting the mixing state information. Could you make similar plots as in Fig. 6 and Fig. 7 but using F_{LH} instead of χ ? If the error shows more converged dependence on F_{LH} , χ may not be a better parameter for the CCN prediction. Besides, χ is hard to determine in practice by available measurement techniques.

Thanks for this suggestion, which together with reviewer's point 2.3 inspired us to add another section to the paper. We think that including F_{LH} and the geometric standard deviation of the κ -distribution in the discussion will answer the questions that many readers might have.

The new section (Section 6) is titled “Relationship of χ and CCN error to other metrics of hygroscopic mixing state”, and we added two figures.

Figure 10 shows the relationship of (a) size-restricted mixing state parameter χ_{res} and mixing state parameter, χ , (b) size-restricted mixing state parameter χ_{res} and number fraction of particles with low hygroscopicity, F_{LH} , and (c) size-restricted mixing state parameter χ_{res} and geometric standard deviation of the κ -distribution, σ_{κ} , for all 384 aerosol populations in \mathbb{P} .

Figure 11 is analogous to Figure 6 and shows (a) Relative error $\epsilon(s_{\text{env}}, \chi_{\text{res}})$, (b) relative error $\epsilon(s_{\text{env}}, F_{\text{LH}})$, and (c) Relative error $\epsilon(s_{\text{env}}, \sigma_{\kappa})$ for each individual aerosol population in \mathbb{P} , evaluated for 20 supersaturation values between 0.05% and 1%.

We also added text in the conclusions, p. 18, lines 19–23: “We also explored the relationship of CCN error other measures of mixing state, specifically a size-restricted χ , the fraction of particles with low hygroscopicity, and the geometric standard deviation of the κ -distribution. These other measures also capture aspects of the heterogeneity of the particle population and the dependence of CCN error on these quantities are qualitatively similar to the one when using χ . However, χ has advantages as a mixing state metric due to its defined range (0 to 100%) and well-defined extremes (0% is fully internally mixed and 100% is full externally mixed).”

(2.3) Comparison of χ to existing parameters: χ is a single parameter containing more intensive information. The authors have nicely presented its general concept by a nice illustration of Fig. 1. But it is still hard to fully understand it. Can you plot the series of χ and compare it to other well-established parameters, e.g., F_{LH} , or the (geometric) standard deviation of kappa distribution, etc.? Does a higher χ correspond to a larger F_{LH} or a smaller standard deviation? The potential link to other mixing state parameter may help people to accept the new parameter.

This was addressed in conjunction with the response to comment (2.2), and forms the content of the new Section 6.

(2.4) Design of experiments and discussion: In this study, the performance of χ is evaluated by comparing the error with kinds of averaged diversity value over the whole size range. I suggest the authors to reconsider this. The errors in CCN prediction are controlled by multiple parameters, i.e., the evaluated supersaturation, the size distribution and the kappa distribution. We know that the particle size has a dominant effect on the CCN activation. But if we want to quantify the effects of particle size on the CCN prediction, can we plot the error against the averaged particle size as what was done for χ ? It is not clear what's the better solution but maybe if the authors could try to use size-resolved χ and check how to use it in CCN prediction or parameterization, e.g., maybe there is a compact empirical relation between χ and the averaged activation fraction at each size.

We did explore what the error distribution would look like if size-resolved composition information was retained. This is described on p. 14, lines 17–26. We did not include a figure for these results because they are qualitatively similar to Figure 6.

In the new Section 6, we now also added some material to show what happens if χ is calculated based on a size restricted population (Figure 10a and 11a, see response to comment (2.2) for details). Again, the error distribution looks qualitatively very similar to Figure 6.

(2.5) Abstract “However, it has been difficult to rigorously investigate this assumption because appropriate metrics for mixing state were lacking”

I think the kappa distribution and the corresponding parameters (mean kappa, mode kappa, and standard deviation) in Su et al. (2010) may be as good as χ in representing the CCN-relevant mixing state.

We agree with the reviewer and removed this sentence.

(2.6) Page 8 ln 10, Can the authors specify which kappa values were used for the two surrogate groups and how to calculate kappa for internally mixed particle?

We added the specifics about this on p. 8, lines 11/12– p. 9, lines 1–4:

“Since we track the composition evolution of each individual particle throughout the simulation, we can calculate the critical supersaturation s_c for each particle as described in Riemer et al. (2010), using the concept of the dimensionless hygroscopicity parameter κ (Petters and Kreidenweis, 2007). The overall κ for a particle is the volume-weighted average of the κ values of the constituent species. Based on Petters and Kreidenweis (2007) we assume $\kappa = 0.65$ for all salts formed from the $\text{NH}_4^+ - \text{SO}_4^{2-} - \text{NO}_3^-$ system. For all MOSAIC model species that represent SOA we assume $\kappa = 0.1$, and for POA and BC we assume $\kappa = 0.001$ and $\kappa = 0$, respectively.”

Note that we do not assign kappa values for the surrogate species as such, but calculate the overall κ for a particle as the volume-weighted average of the κ values of the constituent species, and assign κ values of the constituent species as specified above.

References

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N. Riemer and M. West. Quantifying aerosol mixing state with entropy and diversity measures. *Atmos. Chem. Phys.*, 13:11423–11439, 2013.

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Metrics to quantify the importance of mixing state for CCN activity

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Abstract. It is commonly assumed that models are more prone to errors in predicted CCN concentrations when the aerosol populations are externally mixed. However, it has been difficult to rigorously investigate this assumption because appropriate metrics for mixing state were lacking and metrics needed to quantify the error in CCN concentrations due to mixing state effects were unavailable. In this work we investigate this assumption by using the mixing state index (χ) proposed by Riemer and

5 West (ACP, 13, 11423–11439, 2013) to quantify the degree of external and internal mixing of aerosol populations. We combine this metric with particle-resolved model simulations to quantify error in CCN predictions when mixing state information is neglected, exploring a range of scenarios that cover different conditions of aerosol aging. We show that mixing state information does indeed become unimportant for more internally-mixed populations, more precisely for populations with χ larger than 60%. For more externally-mixed populations (χ below 20%) the relationship of χ and the error in CCN predictions is not unique, 10 and ranges from lower than –40% to about 150%, depending on the underlying aerosol population and the environmental supersaturation. We explain the reasons for this behavior with detailed process analyses.

1 Introduction

The mixing state of an aerosol population depends on the distribution of chemical compounds across the population (Riemer and West, 2013). Field observations reveal that ambient aerosol mixing states can be complex (e.g. Healy et al., 2013; Moffet 15 and Prather, 2009). Even freshly emitted particles can contain multiple chemical species depending on the source characteristics (Ault et al., 2010; Toner et al., 2006), and the initial particle composition is further modified in the atmosphere as a result of aging processes such as coagulation, condensation of secondary aerosol species, and heterogeneous reactions (Weingartner et al., 1997). This profoundly impacts the aerosol optical properties (Jacobson, 2001), their CCN activity (Wang et al., 2010) and the particle lifetime (Koch et al., 2009).

20 In this study we focus on CCN activity, and indeed, many CCN closure studies show that the quality of the closure depends crucially on the assumptions about aerosol mixing state (McFiggans et al., 2006; Wang et al., 2010; Bhattu and Tripathi, 2015; Ervens et al., 2010). Based on these observational findings, it is commonly assumed that the “internal mixture assumption” works well for regions that are not directly influenced by fresh emission sources. By “internal mixture assumption” we mean

the assumption that the composition of individual aerosol particles equals the composition of the bulk, at least within a certain size range. In contrast, closure studies show that, in areas close to emission sources, a certain degree of external mixing needs to be assumed to obtain good closure. This is confirmed by modeling studies showing that, without introducing fresh emissions, aging processes transform the mixing state of aerosol populations so that the CCN properties can be deduced from the bulk 5 aerosol composition, and mixing state details become negligible (Zaveri et al., 2010; Ching et al., 2012; Fierce et al., 2013). However, the timescale for this transformation depends on the local conditions, such as total number concentration of existing CCN and the amount of condensable aerosol material (Fierce et al., 2015). Introducing fresh emissions additionally complicates the picture.

10 The importance of aerosol mixing state to CCN concentration is also demonstrated by studies that experimentally determine the hygroscopicity of organic species. For example, Mei et al. (2013) showed for the CalNex field campaign that the uncertainty in the derived organic hygroscopicity depended on the uncertainties in the derived CCN hygroscopicity and the volume fraction of chemical species contained in the CCN, which are related to the aerosol mixing state.

15 Our aim for this study is to quantitatively explain how mixing state and error in CCN concentrations are related. Previous work has approached this question from a Lagrangian point of view, considering the aging history of the particle population (Zaveri et al., 2010; Ching et al., 2012, 2016a) as a plume of aerosol particles is evolving. While this approach yields valuable process-level understanding, it is difficult to apply to field observations, because following a particle population in the real atmosphere is inherently challenging. In this study, we therefore will not focus on the temporal evolution of the particle population, but instead individually sample populations from a set of simulations. **The central question that we address is: For 20 aerosol populations of a given mixing state, what magnitude of errors can we expect for estimating CCN concentrations when assuming that the population is internally mixed?**

Our study combines a recently developed metric for aerosol mixing state with particle-resolved modeling and a strategy for error quantification. We use the mixing state index (χ) proposed by Riemer and West (2013) to rigorously quantify the degree of external/internal mixing of aerosol populations. This mixing state index is a scalar quantity, and varies between 0% (for completely external mixtures) and 100% (for completely internal mixtures) for any given aerosol population. The metric 25 has been applied to field observations in Paris during the MEGAPOLI campaign (Healy et al., 2014), in Northern California during CARES (O'Brien et al., 2015), and in central London (Giorio et al., 2015). It has proven useful to gain insight into the processes that govern diurnal changes in mixing state and mixing state changes related to air mass origin. For example, Healy et al. (2014) were able to determine a characteristic diurnal cycle of χ for Paris with low values during the day when daytime 30 primary emissions dominated, compared to the night when formation of ammonium nitrate moved the population towards a more internal mixture, indicated by increasing χ values.

Sections 2 and 3 give a brief background on the definition of the mixing state index and on the particle-resolved model PartMC-MOSAIC. Section 4 describes the model simulations that are the basis of this work and our framework for error quantification, followed by the results in Section 4 and 4. Section 7 summarizes our findings.

2 Mixing state metrics

Riemer and West (2013) put forward a framework to quantify aerosol mixing state, which was inspired by diversity metrics used in other disciplines such as ecology (Whittaker, 1972), economics (Drucker, 2013), neuroscience (Strong et al., 1998), and genetics (Falush et al., 2007). [The salient points are summarized as follows](#). Given a population of N aerosol particles, 5 each consisting of some amounts of A distinct aerosol species, this concept is based on the knowledge of mass of species a in particle i , denoted μ_i^a , for $i = 1, \dots, N$, and $a = 1, \dots, A$. From this quantity, all other mass-related quantities can be defined, [as detailed in Riemer and West \(2013\) and here listed in Table 1](#), and the diversity metrics can be constructed as shown in [Table 2](#).

The particle diversity D_i represents the number of “effective species” of particle i . For a particle i that consists of A species, 10 the highest possible value for D_i is A , and this occurs when all A species are present in equal mass fractions. Knowing the D_i values for all particles, the population-level quantities D_α and D_γ can be calculated, with D_α being the average effective number of species in each particle, and D_γ being the effective number of species in the bulk. Finally, the mixing state index χ is defined as

$$\chi = \frac{D_\alpha - 1}{D_\gamma - 1}. \quad (1)$$

15 The mixing state index χ varies from 0% (a fully externally mixed population) to 100% (a fully internally mixed population). To fully quantify mixing state, two of the three metrics (D_α, D_γ, χ) are needed, and the third can be derived. As shown in Riemer and West (2013) and Healy et al. (2014), it is instructive to map the mixing state metrics of aerosol populations into a mixing state diagram (D_α, D_γ), as shown in Figure 1. Particle populations with single-species particles, i.e., “externally mixed” populations, have $D_\alpha = 1$ and D_γ between 1 and A , and are therefore mapped onto the vertical axis ($\chi = 0\%$). Populations 20 consisting of particles with identical mass fractions map onto the diagonal $\chi = 100\%$. Since χ has the intuitive interpretation of the “degree of internal mixing”, we use it here as a metric for error quantification as shown in Section 4.

Note that the definition of “species” for calculating the mass fractions depends on the application. It can refer to chemical species, as in Riemer and West (2013), Healy et al. (2014), O’Brien et al. (2015), and Giorio et al. (2015), or it can refer to species groups, as in Dickau et al. (2016) who quantified mixing state with respect to volatile and non-volatile components. 25 Since we are concerned with CCN properties in this paper, we will group the chemical model species according to hygroscopicity, defining two surrogate species. Black carbon (BC) and primary organic aerosol (POA) are combined into one surrogate species, since their hygroscopicity is very low. All other model species (inorganic and secondary organic aerosol species) are combined into a second surrogate species and χ is calculated from these two surrogate species.

3 Particle-resolved modeling with PartMC-MOSAIC

30 A detailed description of the numerical methods used in PartMC-MOSAIC is given in Riemer et al. (2009). In brief, PartMC (Particle-resolved Monte Carlo) is a 0-D, or box, model which explicitly resolves the composition of many individual particles within a well-mixed computational volume representing a much larger air parcel. During the evolution of the air parcel moving

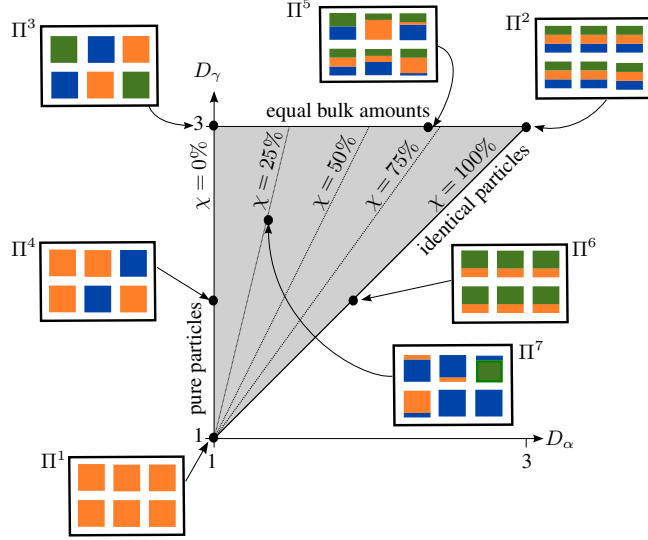


Figure 1. Mixing state diagram to illustrate the relationship between average particle diversity D_α , bulk population diversity D_γ , and mixing state index χ for seven example populations (Π_1 to Π_7). Each population consists of six particles, and the colors represent different chemical species ($A = 3$). [This figure is taken from Riemer and West \(2013\).](#)

Quantity	Meaning
μ_i^a	mass of species a in particle i
$\mu_i = \sum_{a=1}^A \mu_i^a$	total mass of particle i
$\mu^a = \sum_{i=1}^N \mu_i^a$	total mass of species a in population
$\mu = \sum_{i=1}^N \mu_i$	total mass of population
$p_i^a = \frac{\mu_i^a}{\mu_i}$	mass fraction of species a in particle i
$p_i = \frac{\mu_i}{\mu}$	mass fraction of particle i in population
$p^a = \frac{\mu^a}{\mu}$	mass fraction of species a in population

Table 1. Aerosol mass and mass fraction definitions and notation. The number of particles in the population is N , and the number of species is A . [This table is taken from Riemer and West \(2013\).](#)

Quantity	Name	Units	Range	Meaning
$H_i = \sum_{a=1}^A -p_i^a \ln p_i^a$	mixing entropy of particle i	—	0 to $\ln A$	Shannon entropy of species distribution within particle i
$H_\alpha = \sum_{i=1}^N p_i H_i$	average particle mixing entropy	—	0 to $\ln A$	average Shannon entropy per particle
$H_\gamma = \sum_{a=1}^A -p^a \ln p^a$	population bulk mixing entropy	—	0 to $\ln A$	Shannon entropy of species distribution within population
$D_i = e^{H_i} = \prod_{a=1}^A (p_i^a)^{-p_i^a}$	particle diversity of particle i	effective species	1 to A	effective number of species in particle i
$D_\alpha = e^{H_\alpha} = \prod_{i=1}^N (D_i)^{p_i}$	average particle (alpha) species diversity	effective species	1 to A	average effective number of species in each particle
$D_\gamma = e^{H_\gamma} = \prod_{a=1}^A (p^a)^{-p^a}$	bulk population (gamma) species diversity	effective species	1 to A	effective number of species in the bulk
$D_\beta = \frac{D_\gamma}{D_\alpha}$	inter-particle (beta) diversity	—	1 to A	amount of population species diversity due to inter-particle diversity
$\chi = \frac{D_\alpha - 1}{D_\gamma - 1}$	mixing state index	—	0% to 100%	degree to which population is externally mixed ($\chi = 0\%$) versus internally mixed ($\chi = 100\%$)

Table 2. Definitions of aerosol mixing entropies, particle diversities, and mixing state index. In these definitions we take $0 \ln 0 = 0$ and $0^0 = 1$. [This table is taken from Riemer and West \(2013\)](#).

along a specific trajectory, the mass of each constituent species within each particle is tracked. Emission, dilution, nucleation and Brownian coagulation are simulated with a stochastic Monte Carlo approach. The relative positions of particles within the computational volume are not tracked. To improve efficiency of the method we use weighted particles in the sense of DeVille et al. (2011) and efficient stochastic sampling methods (Michelotti et al., 2013).

5 PartMC is coupled with the aerosol chemistry model MOSAIC (Model for Simulating Aerosol Interactions and Chemistry) (Zaveri et al., 2008) which includes the gas phase photochemical mechanism CBM-Z (Zaveri and Peters, 1999), the Multicomponent Taylor Expansion Method (MTEM) for estimating activity coefficients of electrolytes and ions in aqueous solutions (Zaveri et al., 2005a), the multi-component equilibrium solver for aerosols (MESA) for intraparticle solid-liquid partitioning (Zaveri et al., 2005b) and the adaptive step time-split Euler method (ASTEM) for dynamic gas-particle partitioning over size-10 and composition-resolved aerosol (Zaveri et al., 2008), as well as a treatment for SOA (secondary organic aerosol) based on the SORGAM scheme (Schell et al., 2001). The CBM-Z gas phase mechanism treats a total of 77 gas species. MOSAIC treats key aerosol species including sulfate (SO_4), nitrate (NO_3), ammonium (NH_4), chloride (Cl), carbonate (CO_3), methanesulfonic acid (MSA), sodium (Na), calcium (Ca), other inorganic mass (OIN), BC, POA, and SOA. The model species OIN represents species such as SiO_2 , metal oxides, and other unmeasured or unknown inorganic species present in aerosols. SOA includes 15 reaction products of aromatic precursors, higher alkenes, α -pinene and limonene.

PartMC-MOSAIC has been used in the past for process studies of mixing state impacts on aerosol properties in various environments. For example, Ching et al. (2012) quantified the impact of aerosol mixing state on cloud droplet formation, and Fierce et al. (2013) investigated the sensitivity of CCN activity to mixing state characteristics at emission. The model was also used to explain the observed diurnal variations of aerosol hygroscopicity in the North China Plain (Liu et al., 2011) and to 20 characterize the evolution of aerosol mixing state in a ship plume (Tian et al., 2014).

4 Framework for error quantification

The basis for the error quantification framework is the library of eight urban plume scenarios described in Ching et al. (2016a), Sections 3.2.1 and 4.1. These scenarios were designed to simulate the aging process of black carbon-containing particles from combustion sources. With “aging” we refer here to the transition from CCN-inactive to CCN-active at a given supersaturation 25 threshold due to coagulation with other aerosol particles or due to condensation of secondary aerosol material (Riemer et al., 2010). We consider the formation of ammonium nitrate and ammonium sulfate as well as secondary organic aerosol from biogenic or anthropogenic gaseous precursors as represented in the SORGAM module.

The eight scenarios are derived from the base case, which is the scenario presented in Ching et al. (2012). To construct the scenario library, the BC emission rate was set to 100% (E100), 25% (E25), and 2.5% (E2.5) of the base-case, and the number 30 concentration of background aerosol particles was set to 100% (B100), and 10% (B10) of the base-case. Combining the emission cases and the background aerosol concentration cases results in six scenarios, and the gas phase emissions for these six scenarios were the same as in Ching et al. (2012). For the base case (E100-B100) we performed two additional simulations by reducing the emission rate of all gaseous components by 50% (G50) and 25% (G25), respectively, of the case presented

Initial/Background	N_a / cm^{-3}	$D_g / \mu\text{m}$	σ_g	Composition by mass
Aitken mode	1800	0.02	1.45	49.64% $(\text{NH}_4)_2\text{SO}_4$ + 49.64% SOA + 0.72% BC
Accumulation mode	1500	0.116	1.65	49.64% $(\text{NH}_4)_2\text{SO}_4$ + 49.64% SOA + 0.72% BC
<hr/>				
Emission	$E_a / \text{m}^{-2} \text{s}^{-1}$	$D_g / \mu\text{m}$	σ_g	Composition by mass
Meat cooking	9×10^6	0.086	1.91	100% POA
Diesel Vehicles	1.6×10^8	0.05	1.74	30% POA + 70% BC
Gasoline Vehicles	5×10^7	0.05	1.74	80% POA + 20% BC

Table 3. Number concentration, N_a , of the initial/background aerosol population, and area source strength, E_a , of the three types of emission. All aerosol size distributions are assumed to be lognormal and defined by the geometric mean diameter, D_g , and the geometric standard deviation, σ_g .

in Ching et al. (2012). The BC mass concentrations of all the plume scenarios ranged from $0.05 \mu\text{g m}^{-3}$ to $3.6 \mu\text{g m}^{-3}$. The formation of secondary aerosol material was similar in all G100 cases, since the emissions of gaseous precursors were the same. On the first simulation day, the total mass concentration was dominated by ammonium nitrate formation, which evaporated after ammonia emissions ceased (Ching et al., 2012). Secondary production of sulfate and organic mass occurred 5 on the first and second days of the simulation. The total mass concentration varied by a factor of about two between plume scenarios G100 and G50 due to differences in secondary aerosol mass formation.

As a result of these changes in input parameters, the aging of BC particles proceeded at different rates, and different mixing states arose over the course of the simulation, which are further discussed below. The simulation time for each scenario was 48 hours, and during the first 12 hours gas and aerosol emissions entered the air parcel. Background aerosol particles were 10 introduced over the entire simulation time owing to dilution with background air. Table 3 specifies the details of the initial and background aerosol distribution as well as the aerosol emissions for the base case run.

Here we are not focused on the process analysis of the temporal evolution of BC-containing particles, but rather on the aerosol state; we use the hourly aerosol state from the scenarios as a basis for our analysis. We will refer to the set of these populations as set \mathbb{P} , which comprises $N_{\text{pop}} = 48 \times 8 = 384$ elements. The populations in \mathbb{P} cover a wide range of mixing 15 states. Figure 2 illustrates how these aerosol populations map to the mixing state diagram. Each symbol represents an aerosol population from \mathbb{P} , colored by the scenario the population is sampled from.

As explained in Section 2, the maximal possible range for both D_α and D_γ is from 1 to 2, however this range is not entirely accessed with the populations investigated here. The effective number of species in the population (D_γ) is always larger than 1, i.e., there are no populations that consist solely of hygroscopic or hydrophobic species. It reaches values of almost 2 for some 20 populations, meaning that for these populations the bulk mass fractions of hydrophobic and hygroscopic material are about the same. On a per-particle level, the average number of effective species (D_α) is close to 1 for some populations, indicating that populations exist for which the particles consist of purely hygroscopic or hydrophilic material. The maximum value of D_α is about 1.7. Figure 2 also shows lines of constant mixing state index χ . The populations cover values of χ from 7.5%

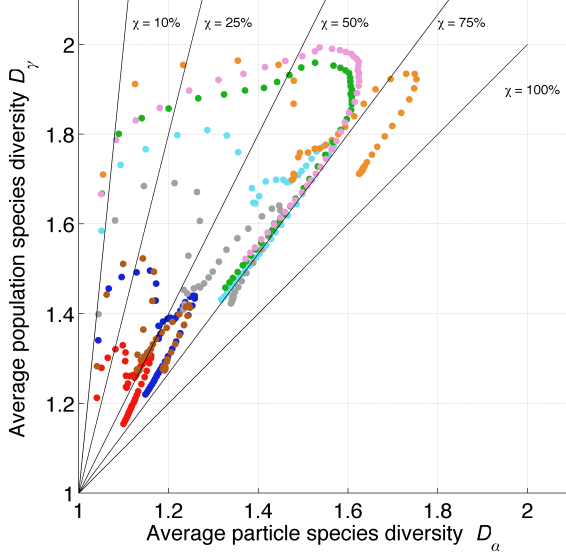


Figure 2. Mixing state diagram for the 384 populations in \mathbb{P} . The colors indicate the Ching et al. (2016a) scenario from which the populations were sampled: B100-E100-G100 (light blue), B10-E100-G100 (orange), B100-E25-G100 (blue), B10-E25-G100 (grey), B100-E2.5-G100 (red), B10-E2.5-G100 (brown), B100-E100-G50 (green) and B100-E100-G25 (pink). The abbreviations of the scenarios follow Ching et al. (2016a), “B” indicates the level of background particle concentration (100% and 10% of the base case scenario), “E” indicates the level of the black carbon emission rate (100%, 25%, and 2.5% of the base case scenario) and “G” indicates the level of gas emission rate (100%, 50%, and 25% of the base case scenario).

to 88%. The lack of populations with χ between 88% and 100% does not impact the generality of the conclusions. As we will see later, the error in CCN concentration when neglecting mixing state information vanishes for $\chi > 60\%$. The pattern that each scenario forms in this mixing state diagram can be explained by the temporal evolution of D_γ and D_α , which is determined by coagulation, condensation of secondary aerosol material, dilution, and emission (Riemer and West, 2013). All 5 scenarios start with low average particle species diversity, D_α , because at the beginning of the simulation the particles consist mainly of one of the surrogate species, either hydrophobic BC or POA, or hygroscopic background species. As the simulation proceeds, secondary aerosol species condense and coagulation occurs, moving D_α towards larger values. This process does not necessarily proceed monotonically, i.e., D_α can also decrease with time. The reason could be either that secondary aerosol material is evaporating or that so much secondary aerosol material is condensing that it dominates in terms of composition. A 10 similar argument can be made for the temporal evolution of D_γ , but applied to the composition of the bulk population.

For each of the 384 aerosol populations we determine CCN concentrations as follows: Since we track the composition evolution of each individual particle throughout the simulation, we can calculate the critical supersaturation s_c for each particle as

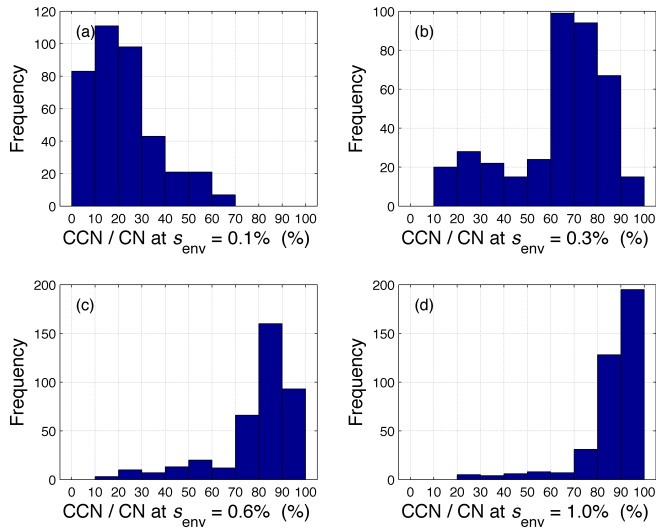


Figure 3. Frequency distribution of CCN/CN fractions for all 384 aerosol populations in \mathbb{P} at four selected environmental supersaturations s_{env} . The 384 populations were derived from eight scenarios, each with 48 hourly snapshots.

described in Riemer et al. (2010), using the concept of the dimensionless hygroscopicity parameter κ (Petters and Kreidenweis, 2007). The overall κ for a particle is the volume-weighted average of the κ values of the constituent species. Based on Petters and Kreidenweis (2007) we assume $\kappa = 0.65$ for all salts formed from the $\text{NH}_4^+ - \text{SO}_4^{2-} - \text{NO}_3^-$ system. For all MOSAIC model species that represent SOA we assume $\kappa = 0.1$, and for POA and BC we assume $\kappa = 0.001$ and $\kappa = 0$, respectively.

5 We use s_c to define a cumulative number distribution $N_{\text{CCN}}(s_c)$, which represents the number of particles per volume with critical supersaturation less than s_c . The CCN concentration at a chosen environmental supersaturation threshold s_{env} for a population with mixing state index of χ_0 is then denoted as $N_{\text{CCN}}(s_{\text{env}}, \chi_0)$.

Figure 3 provides an overview of the CCN activity for the populations. It shows the frequency distribution of the CCN/CN fraction, evaluated at four different supersaturation thresholds (0.1%, 0.3%, 0.6%, and 1%). These four supersaturation thresholds bound the CCN activity of our populations well. As expected, for the lowest supersaturation threshold of 0.1%, most populations only have small CCN/CN fractions with the median at $\text{CCN}/\text{CN} = 20\%$. For the highest supersaturation threshold of 1% the median is at $\text{CCN}/\text{CN} = 90\%$.

We determine the error that is introduced by neglecting mixing state information as shown schematically in Figure 4. The starting point is an aerosol population from the particle-resolved simulations as shown in Figure 4(a). This particular population had a mixing state index of $\chi_0 = 13\%$. Figure 4 shows the distribution density function $\partial^2 N(D, \kappa) / (\partial \log_{10} D \partial \kappa)$ based on the two-dimensional cumulative number distribution $N(D, \kappa)$ in terms of particle diameter D and hygroscopicity parameter κ (Su et al., 2010; Fierce et al., 2013). We observe that at a particular particle size a distribution of hygroscopicity parameter

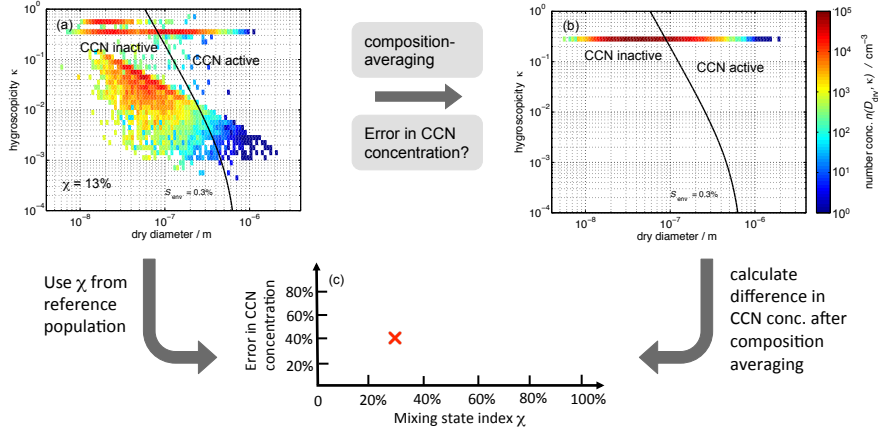


Figure 4. Conceptual framework for error quantification, see Section 4 for details.

values exists. The solid black line indicates the chosen supersaturation threshold (here 0.3% as an example) that separates CCN-active from CCN-inactive particles.

We then perform composition averaging on this particle population as described by Ching et al. (2016a). Composition averaging preserves the bulk aerosol mass concentrations, the number concentration and the particle diameters, but modifies 5 the per-particle composition, so that each particle is assigned a composition equal to the average composition of the population. Figure 4(b) shows the population after composition averaging. Because all particles were assigned the same composition (equal to the average composition), the spread in κ vanishes, and the CCN number concentration was altered as a result.

After composition averaging we re-calculate the CCN concentration, $\bar{N}_{\text{CCN}}(s_{\text{env}})$, and define the relative error

$$\epsilon(s_{\text{env}}, \chi_0) = \frac{\bar{N}_{\text{CCN}}(s_{\text{env}}) - N_{\text{CCN}}(s_{\text{env}}, \chi_0)}{N_{\text{CCN}}(s_{\text{env}}, \chi_0)}. \quad (2)$$

10 Finally, we graph the error $\epsilon(s_{\text{env}}, \chi)$ as a function of the mixing state parameter χ for a given value of s_{env} of the reference population as shown in Figure 4(c).

5 Relationship of error in CCN concentration and mixing state index χ

To aid the interpretation of the resulting error distributions as defined in Equation 2, we first show the effect of composition 15 averaging on CCN concentrations in Figure 5 for one example population in \mathbb{P} . This normalized 2D histogram relates the critical supersaturation of each particle before composition averaging to its critical supersaturation after composition averaging. Points above the 1:1 line represent particles for which the critical supersaturation increased after composition averaging, and points below the 1:1 line represent particles for which the critical supersaturation decreased after composition averaging. Figure 5 also shows a vertical and horizontal line that marks an assumed environmental supersaturation threshold of 0.3%. The particles to

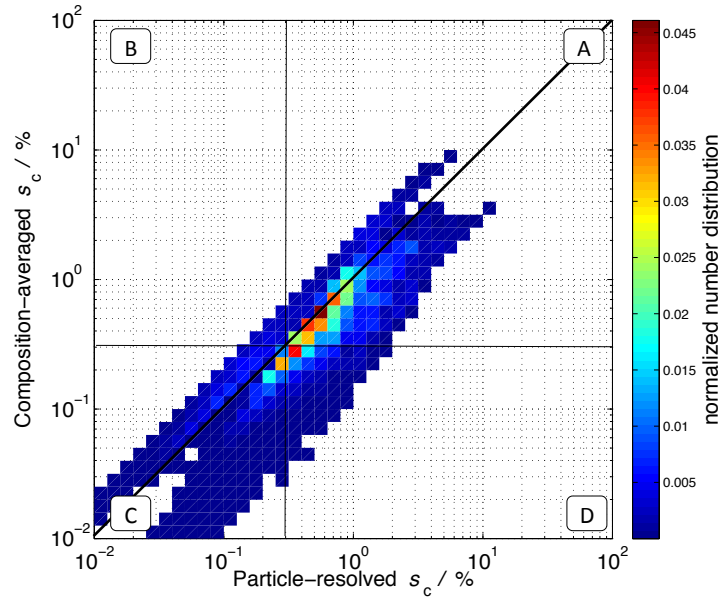


Figure 5. 2D histogram of particles' critical supersaturations before and after composition averaging for one aerosol population from set \mathbb{P} . Particles in quadrant A are not CCN active before or after composition averaging. Particles in quadrant C are CCN active before and after composition averaging. Particles in quadrant B activate in the reference case, but not after composition averaging. Particles in quadrant D activate after composition averaging, but not in the reference case.

the left of the vertical line activate in the reference case, and the particles below the horizontal line activate in the composition-averaged case.

Based on this we can define four quadrants. Particles in quadrant C activate both in the reference case and in the composition-averaged case, while in quadrant A they do not activate in either case. Particles in these two quadrants do not incur any error in 5 CCN concentration even though composition averaging may have changed their critical supersaturation value. Error is caused only by the *difference* in the number of particles that activate in the reference case, but do not activate after composition averaging (quadrant B), and the cases that activate after composition averaging, but did not activate in the reference case (quadrant D). If the number concentration of particles in quadrants B and D was equal, the overall error in CCN concentration would be zero even though a large number of particles might have been mis-classified.

10 The number concentrations in quadrants B and D are determined by the supersaturation threshold and the extent to which composition averaging simplifies mixing state, i.e., the distribution $n(D, \kappa)$ is distorted (compare Figure 4). This determines the error in the critical supersaturation distribution for the composition-averaged population compared to its particle-resolved counterpart. For populations with $\chi = 100\%$, the particles' composition is the same across the entire population, and hence all 15 the particles have the same hygroscopicity values. Composition averaging therefore does not distort $n(D, \kappa)$ and the number concentrations in quadrants B and D are zero. In contrast, for populations with $\chi < 100\%$, composition averaging distorts

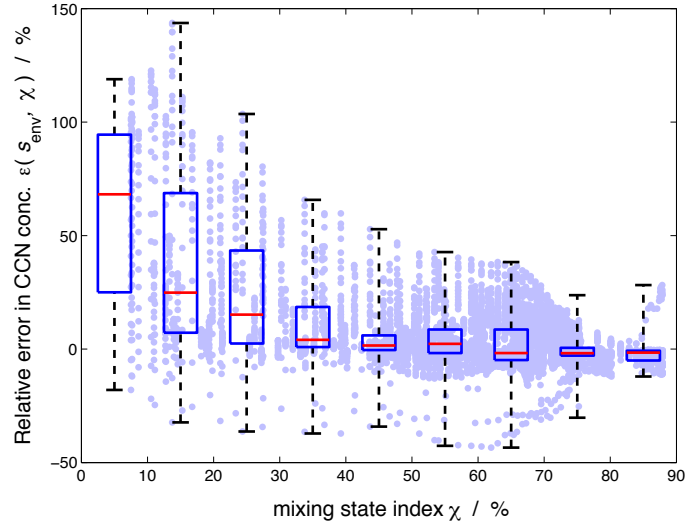


Figure 6. Relative error $\epsilon(s_{\text{env}}, \chi)$ for each individual aerosol population in \mathbb{P} (light purple dots), evaluated for 20 supersaturation values between 0.05% and 1%, see Equation 2 for definition. The boxes show the 25th and 75th percentiles of the distribution of relative errors in each χ -bin. The red line is the median of that distribution, and the dashed line extends to the minimum and maximum error.

$n(D, \kappa)$ depending on the distribution of $n(D, \kappa)$. The 2D histogram in Figure 5 therefore varies from population to population, and the number concentrations in quadrants B and D depend on both the extent of distortion of the critical supersaturation distribution and the supersaturation threshold (the black horizontal and vertical lines) as shown in Figure 5.

In summary, the dependence of the relative error on supersaturation threshold and mixing state index (χ) is determined by 5 the number concentrations in quadrants B and D. Whenever the number concentrations in quadrants B and D are of similar magnitude, the error is small. As we will show in Figures 7 and 8 this can occur for all χ values. The situation of number concentrations in quadrants B and D having different magnitudes tends to occur only for small χ values.

Figure 6 shows the relative errors $\epsilon(s_{\text{env}}, \chi)$ for all populations, evaluated for 20 supersaturation values ranging between 0.05% and 1% in steps of 0.05%. The boxes show the 25th and 75th percentiles of the distribution of relative errors in each 10 χ -bin ($\Delta\chi = 10\%$). The red line is the median of that distribution, and the dashed line extends to the minimum and maximum. This figure confirms that the error tends to decrease as χ increases, and that for a given χ value a range of error values exists, which narrows as χ increases as presented by the boxes in Figure 6. This range is caused by the assumed supersaturation threshold at which the CCN concentration is evaluated, but also reflects that, even for a given supersaturation threshold and a given χ value, different populations experience different amounts of the error cancellations illustrated in Figure 5.

15 Further, the relative error $\epsilon(s_{\text{env}}, \chi)$ is positive for most populations, meaning that composition averaging causes the CCN concentration to be overestimated. However, for some populations the relative error is negative, i.e., composition averaging results in an underestimation of the CCN concentration. Cases with negative relative error mostly occur at low environmental supersaturation ($s_{\text{env}} \leq 0.1\%$) and are most prevalent for χ values between 60% and 80%. For most of these cases the error

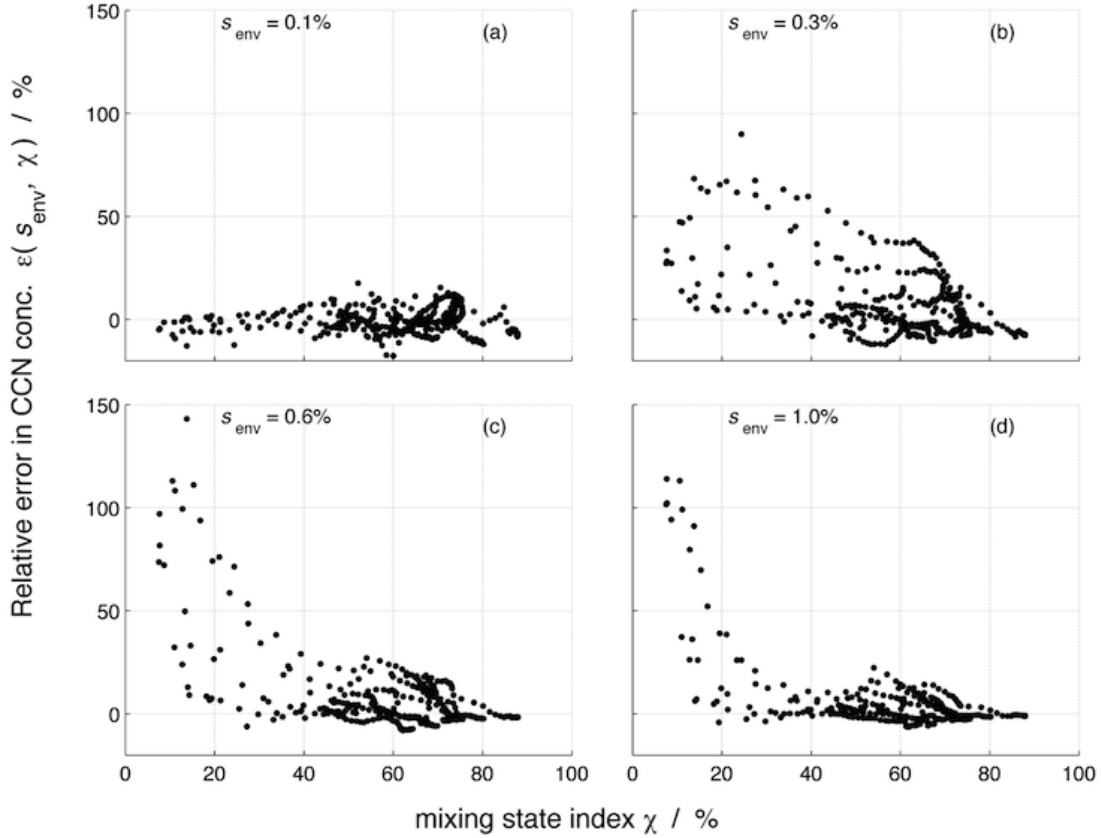


Figure 7. Relative error in CCN concentration $\epsilon(s_{\text{env}}, \chi)$ for different environmental supersaturations, see Equation 2 for definition. The insets show the range for χ between 40% and 80% in more detail.

magnitude is less than 10% (i.e., $-10\% \leq \epsilon \leq 0\%$), however some points stand out with negative relative errors of up to -43% . These originate from the cases with $s_{\text{env}} = 0.05\%$.

To further investigate the dependence of the relative error $\epsilon(s_{\text{env}}, \chi)$ on supersaturation threshold, Figure 7 shows $\epsilon(s_{\text{env}}, \chi)$ for four selected environmental supersaturation thresholds. For low supersaturation thresholds ($s_{\text{env}} = 0.1\%$, Figure 7(a)), the error in CCN concentration is independent of the mixing state index and the magnitude is within $\pm 18\%$. We can explain this fact by consulting Figure 8, which maps the number concentrations in quadrant D and quadrant B. If the concentrations are the same, the points line up on the 1:1 line and the overall error is small because of the cancellation effect explained above. Figure 8(a) shows that this is indeed generally the case for $s_{\text{env}} = 0.1\%$.

At higher supersaturation thresholds (Figure 7(b), (c), and (d)), the highest errors are observed for the populations with the lowest χ values, however at a given χ value a range of errors is possible. The exact outcome depends on to what extent the mis-classified particles in quadrants B and D cancel out. Errors in the intermediate regime of χ between 40% and 80% are up

to about 50%. The largest errors occur below $\chi = 20\%$ (up to about 150%). We see in Figure 8 that this behavior can again be explained with the way the quadrants B and D are populated. The points furthest away from the 1:1 line tend to have the lowest χ values. This is consistent with our expectation that accounting for mixing state is important for more “externally mixed” populations. However, some points are close to the 1:1 line even though their χ value is low, which explains why small errors 5 can still be found for low- χ populations.

These results illustrate that the dependence of CCN concentration error on χ depends on the chosen supersaturation threshold. The reason for this can be understood from Figure 5, which shows that the change in particle number concentration per supersaturation interval is smaller for s_c of 0.1% compared to $0.3 < s_c < 1.5\%$. Therefore, for the case of $s_{\text{env}} = 0.1\%$, the number concentrations in B and D for this population are less sensitive to composition averaging. While this figure shows 10 one specific population as an example, this is generally the case for our aerosol populations and explains why at low supersaturation threshold, the CCN error appears to be independent of χ . As the supersaturation threshold increases, the number concentrations in B and D vary more significantly as demonstrated in Figure 5, and such variations depend on the distortion of $n(D, \kappa)$ by composition averaging. This results in a higher sensitivity of CCN error to χ at higher supersaturation thresholds. It is important to keep in mind that while this behavior applies to the aerosol populations in this study, it cannot be expected 15 that it necessarily applies to any arbitrary aerosol population.

It should be noted here that this calculation represents an upper bound for the error, because composition averaging was performed for the entire population. If size-resolved composition information is retained, the magnitude of error decreases. The error decrease is more pronounced for higher environmental supersaturations. For example, the range of error for $s_{\text{env}} = 0.1\%$ reduces from $(-18\%, +18\%)$ to $(-12\%, +16\%)$ when the particles are sorted into five size bins per decade, and composition 20 averaging is performed within each size bin. For $s_{\text{env}} = 0.3\%$ the range reduces from $(-12\%, +90\%)$ to $(-2\%, +43\%)$, and for $s_{\text{env}} = 1\%$ the range reduces from $(-6\%, +114\%)$ to $(-2\%, +49\%)$. This finding is consistent with observations by Che et al. (2016). They analyzed CCN concentration measurements from the Yangtse River Delta area of China and found that at low supersaturations ($\leq 0.1\%$) using only aerosol bulk composition information was sufficient to predict CCN, whereas at higher supersaturations ($\geq 0.2\%$) using size-resolved chemical composition information improved the prediction of CCN 25 concentrations.

To produce a summary of the above information, we calculated a normalized root mean square deviation metric as a function of mixing state index χ , integrating over all supersaturations as follows:

$$\epsilon_{\text{NRMSD},j} = \frac{1}{\frac{1}{n_{\text{ss}} m_j} \sum_{m=1}^{m_j} \sum_{i=1}^{n_{\text{ss}}} N_{\text{CCN},m}(s_i)} \sqrt{\frac{\sum_{m=1}^{m_j} \sum_{i=1}^{n_{\text{ss}}} (\bar{N}_{\text{CCN},m}(s_i) - N_{\text{CCN},m}(s_i))^2}{n_{\text{ss}} m_j}}, \quad (3)$$

where $n_{\text{ss}} = 20$ is the number of supersaturation values at which we evaluated CCN concentrations, and m_j is the number 30 of aerosol populations in a given χ -bin ($j = 1, \dots, 10$). The green line in Figure 9 shows the relationship of ϵ_{NRMSD} and χ . Figure 9 emphasizes that composition averaging causes larger errors as χ decreases. The average error decreases from about 90% to 30% when χ increases from 0% to 30%. For χ between 40% and 90%, the average error is less than 20%. The average value ϵ_{NRMSD} across the whole range of χ is 17.7%. This is comparable to the error of 14.6% found in the size-resolved simulation by Ching et al. (2016b).

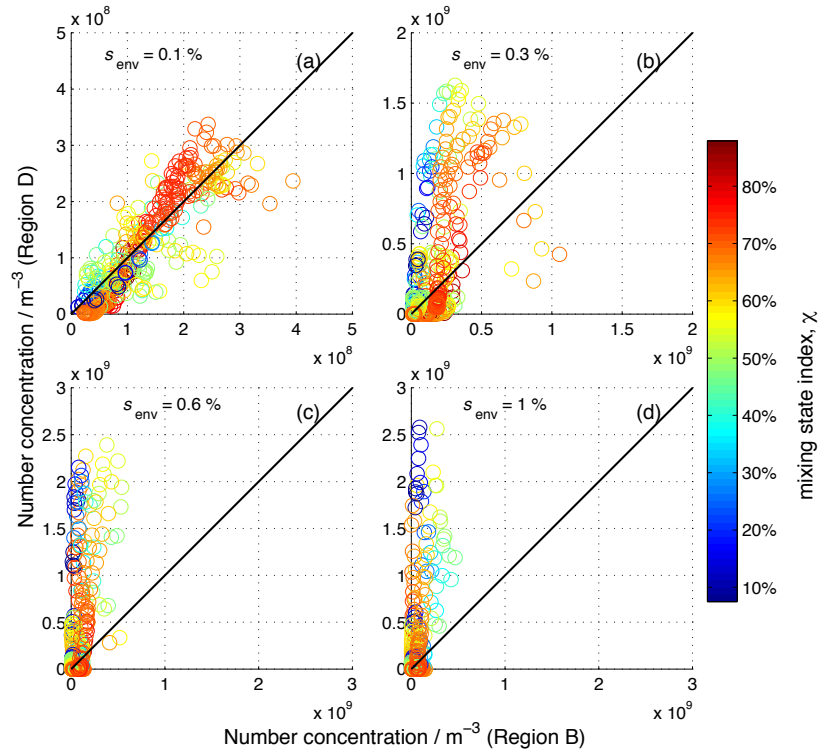


Figure 8. Relationship of number concentration in quadrants B and D (compare Figure 5) for different environmental supersaturations. The data points are colored corresponding to the mixing state index χ .

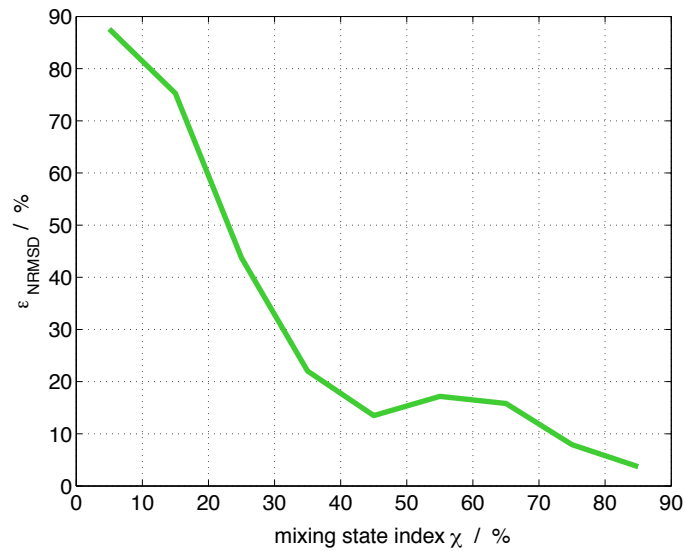


Figure 9. Normalized root mean square deviation in CCN concentration due to composition averaging (ϵ_{NRMSD}) as a function of mixing state index χ , see Equation 3 for definition.

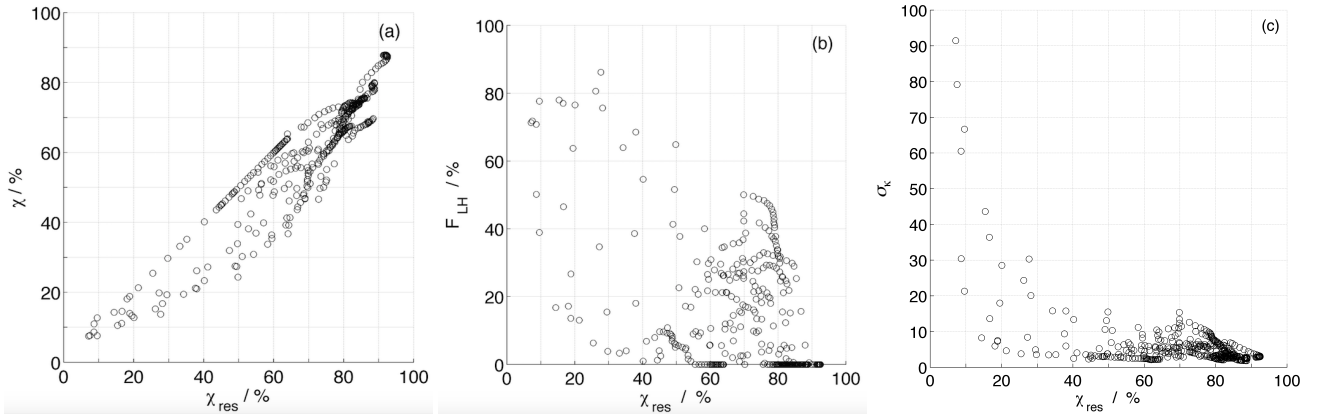


Figure 10. Relationship of (a) size-restricted mixing state parameter χ_{res} and mixing state parameter, χ , (b) size-restricted mixing state parameter χ_{res} and number fraction of particles with low hygroscopicity, F_{LH} , and (c) size-restricted mixing state parameter χ_{res} and geometric standard deviation of the κ -distribution, σ_{κ} , for all 384 aerosol populations in \mathbb{P} .

6 Relationship of χ to other metrics of hygroscopic mixing state

The mixing state index χ can be interpreted as a measure of the heterogeneity of a particle distribution with respect to its hygroscopic properties. In Section 4 we showed how χ relates to the error in CCN concentration that is introduced by assuming that the particle population is internally mixed. To illustrate the concept further, in this section we will show how χ relates to other possible metrics of hygroscopicity heterogeneity, and how each of them relates to the error in CCN due to neglecting mixing state information.

We focus here on three other metrics: (1) A size-restricted version of χ , χ_{res} , only considers particles in the diameter size range of 30 nm to 150 nm when calculating the mixing state index. The rationale for this is that this constitutes the size range for which composition matters most because particles smaller than 30 nm are unlikely to activate, and particles larger than 150 nm are very likely to activate, regardless of their composition. (2) The number fraction of particles with low hygroscopicity, F_{LH} . Here we considered particles with $\kappa < 0.1$ as having low hygroscopicity. (3) The geometric standard deviation of the aerosol distribution with respect to the hygroscopicity parameter κ , σ_{κ} . The choice of this parameter was motivated by Su et al. (2010) who used the concept of hygroscopicity distribution to characterize aerosol particle mixing state with regard to CCN properties.

For consistency, σ_{κ} and F_{LH} were also evaluated using only particles with diameters between 30 and 150 nm.

Figure 10a shows that the mixing state index χ for the entire population and the size-restricted mixing state index χ_{res} are strongly correlated. The sub-population with particle diameters between 30 and 150 nm is in many cases more internally mixed than the whole population. From Figure 10b we conclude that, at least for our study, populations with a high fraction

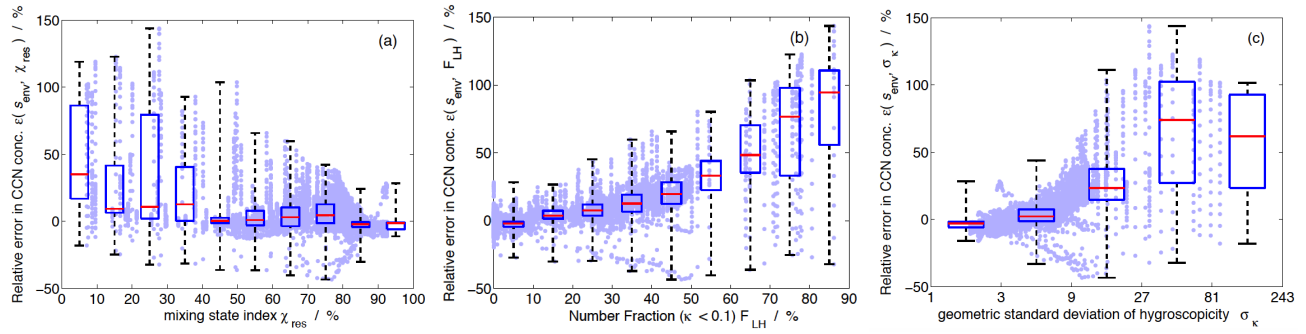


Figure 11. Relative error of (a) $\epsilon(s_{\text{env}}, \chi_{\text{res}})$, (b) $\epsilon(s_{\text{env}}, F_{\text{LH}})$, and (c) $\epsilon(s_{\text{env}}, \sigma_{\kappa})$ for each individual aerosol population in \mathbb{P} (light purple dots), evaluated for 20 supersaturation values between 0.05% and 1%, analogous to Figure 6.

of low-hygroscopic particles tend to be associated with low values of χ_{res} , and vice versa. However, the scatter is large, and populations can be found that do not contain particles in the low-hygroscopicity category, but exhibit a wide range of mixing state index values. It is also plausible that populations exist that are internally mixed, i.e. have high mixing state index, but contain mainly non-hygroscopic species, i.e. have a high value of F_{LH} . The fact that we do not show such populations in 5 our study is due to our set up, where a hygroscopic background population is always present into which we emit fresh, less-hygroscopic particles. Lastly, Figure 10c shows that more externally mixed populations (small χ_{res}) are associated with large values of σ_{κ} .

Figure 11 relates the relative error in CCN concentration to the mixing state metrics χ_{res} , F_{LH} , and σ_{κ} , analogous to Figure 6. These figures looks qualitatively similar, which is expected given that all these measures are measures of heterogeneity of the 10 population with respect to hygroscopic properties. However, fundamentally, the mixing state index χ is appealing because it has a defined range (from 0 to 100%), and the property that when χ is 100%, the population is perfectly internally mixed with a CCN error of zero. It can also be easily generalized to alternate mixing state definition, e.g. based on optical properties, by defining the surrogate species appropriately.

The measure F_{LH} also has a defined range (from 0 to 100%), and a value of 0 implies that the population is fairly (although 15 not necessarily totally) homogeneous in the sense that all particles have a hygroscopicity larger than 0.1. However, the reverse implication does not hold. If the population is homogeneous (perfectly internally mixed), F_{LH} could be 0 or 100%. The geometric standard deviation σ_{κ} is 1 for all perfectly internally mixed populations, but does not have a defined upper limit.

7 Conclusions

Our analysis used particle-resolved modeling and a metric for aerosol mixing state to develop a framework for quantifying error 20 in CCN activity due to simplifying assumptions about mixing state. Mixing state information does indeed become unimportant for more internally-mixed populations. The NRMSD error decreased to less than 10% when χ was larger than 75%. For more externally-mixed populations (χ below 60%), a wide range of error in CCN concentration existed, with errors as large as 150%

for χ lower than 20%. Because the composition averaging was performed for the whole population, without retaining any size-resolved composition information, the maximum CCN error as a function of χ gives an upper bound of CCN error. It should be noted that even for low χ values, results close to the true result may be obtained because of error cancellations. In summary, the CCN error depends on both χ and s_{env} . As the population becomes more internally-mixed (χ approaching 100%), the CCN 5 error is small regardless of the s_{env} threshold. For the aerosol populations in this study, we found higher sensitivity to χ at high s_{env} threshold and lower sensitivity to χ at low s_{env} threshold.

We also explored the relationship of CCN error to other measures of mixing state, specifically a size-restricted χ , the fraction of particles with low hygroscopicity, and the geometric standard deviation of the κ -distribution. These other measures also capture aspects of the heterogeneity of the particle population and the dependence of CCN error on these quantities are 10 qualitatively similar to the one when using χ . However, χ has advantages as a mixing state metric due to its defined range (0 to 100%) and well-defined extremes (0% is fully internally mixed and 100% is full externally mixed).

Several avenues exist for applying and extending our approach in future work. To confirm the validity of this approach, it would be useful to perform CCN closure studies on field campaign data with different mixing state assumptions, and to obtain concurrent aerosol diversity measurements, using for example single-particle mass spectrometry or off-line single-particle 15 analytical techniques. The same framework, with appropriately defined surrogate species, could also be applied to quantify how error for aerosol optical properties depends on aerosol mixing state.

Aerosol-cloud interactions continue to be one of the major sources of future climate prediction uncertainties (IPCC, 2013). One of the reasons for this is the challenge to represent the many microscale aerosol processes in large-scale global climate 20 models (Seinfeld et al., 2016). Using a high-fidelity aerosol model, our study provides quantitative support that mixing state is important for determining the aerosol impact on clouds. As a consequence we conclude that the aerosol representation in global models should account for mixing state. The aerosol microphysics schemes in several current global climate models have been moving into this direction (Ghan and Schwartz, 2007; Bauer et al., 2008; Jacobson, 2002; Aquila et al., 2011; Kaiser et al., 2014; Mann et al., 2010; Liu et al., 2016; Stier et al., 2005) by separating the particles into interacting subpopulations, e.g. freshly emitted BC and aged BC. The central question is whether this degree of resolution of mixing state is sufficient 25 to accurately predict CCN concentration and aerosol optical properties, and whether this framework allows for an accurate prediction of the mixing state *evolution*. In situ measurements of size-resolved aerosol mixing state, as suggested by Seinfeld et al. (2016), are needed to constrain our models and the estimates of aerosol-cloud forcing in climate models.

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