Response to the comments of Anonymous Referee #1

Referee General Comment:

This manuscript explores the equilibrium timescale and mixing timescale of IVOC and LVOC with particles considering different phase states. The work combines the authors’ previous KM-GAP model (Shiraiwa et al., 2012) with the authors’ recent glass transition model (Shiraiwa et al., 2017; DeRieux et al. 2018) to understand the interplay among equilibrium timescale, temperature, relative humidity, and the glass transition temperature of the aerosols.

Besides the numerical results obtained from the model shown in Figure 1-5, the manuscript provides two more useful results: (1) when there is no diffusion limitation within the particle, the gases that have higher saturation mass concentrations will reach gas-particle equilibrium faster; (2) when there is strong diffusion limitation within the particle, gases that have lower volatility may reach gas-particle equilibrium (locally) faster than VOCs with higher volatility. One of the implications is that at a lower temperature (upper troposphere) or when dealing with highly viscous particles, the particle growth process may need to be treated kinetically.

The authors used a numerical model to obtain result #1 above, and it agrees with the findings in Liu et al. 2012 in which an analytical model was used. It is nice to see two different approaches come with the same results and can validate each other. On the other hand, result #2 is more interesting because it shows that local equilibrium could be reached relatively faster for low volatility species when the particle phase is highly viscous. The manuscript also illustrates some concepts that can be commonly misused by folks, such as the difference between equilibrium timescale and the mixing timescale. Because some of the results have been previously discussed in or can be easily inferred from other publications (Liu et al. 2012, Shiraiwa et al. 2011&2012), the novelty of the manuscript needs to be improved. I suggest that the author focus on result #2, which is novel, and use it to make further predictions regarding the physical and chemical processes of aerosols.

Response: We thank Anonymous Referee #1 for the review and the positive evaluation of our manuscript. Following your constructive suggestions, in the revised manuscript, we add (1) a contour plot of $\tau_{eq}$ as a function of bulk diffusivity and volatility to illustrate under what conditions the fast local equilibrium may be achieved to highlight the result #2 you are interested; and (2) simulations for open systems and the results are compared with $\tau_{eq}$ in a closed system. We find partitioning of LVOC is very different in open and closed systems and the corresponding implications in SOA evolution in ambient air and chemical transport models are further broadened. As Referee #2 pointed, this publication essentially closes the loop between predicting phase state and calculating the gas/particle equilibration time. We believe after addition of above two aspects, the novelty of the revised manuscript is improved. Please see the detailed response below.

Referee Major Comment:

(1) For instance, what is the relationship between particle sizes and condensation/evaporation kinetics of gases with different volatility interacting with particles with various viscosity?
Response: Thanks for this helpful comment. In our ACPD manuscript, Figure 5 (Figure 6b and d in the revised manuscript) has shown the relationship between $\tau_{eq}$ and particle size for LVOC condensing on less viscous as well as highly viscous particles. In the revised manuscript, we add comparable calculation for SVOC (Fig. 6a, c). This issue has also been discussed in previous studies, e.g., Liu et al. (2013) and Mai et al. (2015). For example, Mai et al. (2015) presented $\tau_{eq}$ as a function of particle diameter and volatility, showing that $\tau_{eq}$ increases as the particle diameter increases or the volatility of the condensing species decreases when particles are liquid with partitioning limited by interfacial transport. When particles are highly viscous with bulk diffusion-limited partitioning, the time to reach full equilibrium depends on mixing timescale. The following discussions have been added in the revised manuscript:

Lines 259-263: “Previous studies have shown that $\tau_{eq}$ depends on particle size (Liu et al., 2013; Zaveri et al., 2014; Mai et al., 2015) and particle mass loadings (Shiraiwa and Seinfeld, 2012; Saleh et al., 2013). For further examination of these effects at different $T$, Figure 6 shows the dependence of $\tau_{eq}$ of SVOC ($C^0 = 10 \, \mu g \, m^{-3}$) and LVOC ($C^0 = 0.1 \, \mu g \, m^{-3}$) on the mass concentration and the diameter of pre-existing particles”.  

Lines 271-274: “When particles are less viscous at 298 K ($D_b = 10^{-11} \, cm^2 \, s^{-1}$) $\tau_{eq}$ of SVOC is shorter than that of LVOC for the same particle size and mass loadings. When partitioning into highly viscous particles at 250 K ($D_b = 10^{-18} \, cm^2 \, s^{-1}$), SVOC takes longer time than LVOC to reach equilibrium”.  

Lines 275-285: “Typical ambient organic mass concentrations in Beijing, Centreville in southeastern US, Amazon Basin, and Hyytiälä, Finland are indicated in Fig. 6. The particle phase state was observed to be mostly liquid in highly polluted episodes in Beijing (Liu et al., 2017), under typical atmospheric conditions in the southeastern US (Pajunoja et al., 2016), and under background conditions in Amazonia (Bateman et al., 2017). At these conditions $\tau_{eq}$ should be mostly less than 30 minutes (Fig. 6a, b). Particles were measured to be semi-solid or amorphous solid in clear days in Beijing (Liu et al., 2017), in Amazonia when influenced by anthropogenic emissions (Bateman et al., 2017), and the boreal forest in Finland (Virtanen et al., 2010). Under these conditions and also when particles are transported to the free troposphere, $\tau_{eq}$ can be longer than 1 hour especially in remote areas with low mass loadings (Fig. 6c, d)".
Figure 6. Equilibration timescale ($\tau_{eq}$) for (a, c) SVOC ($C^0 = 10 \mu g m^{-3}$) and (b, d) LVOC ($C^0 = 0.1 \mu g m^{-3}$) as a function of particle diameter (nm) and mass concentration ($\mu g m^{-3}$) of pre-existing particles at 60% RH and $T$ of (a-b) 298 K and (c-d) 250 K in the closed system. The glass transition temperature of pre-existing particles under dry conditions ($T_{g,org}$) is set to be 270 K, which leads to $D_b$ of (a-b) $10^{-13}$ cm$^2$ s$^{-1}$ and (c-d) $10^{-18}$ cm$^2$ s$^{-1}$. Ambient organic mass concentrations are indicated with arrows.

(2) I would also be curious to know what is the boundary criteria for result #2 to happen, i.e., how viscous would the particles have to be or how low the volatility of the VOC have to be in order to reach relatively fast local equilibrium? The author can also explore the effects of equilibrium partitioning when the gases can both partition in and react with the particle phase.

Response: This is a very interesting point. To address your question, we conducted additional simulations for $\tau_{eq}$ as a function of bulk diffusivity and volatility for both open and closed systems. The results of such simulations are shown in new Figure 4. The effect of particle-phase reactions on SOA partitioning is an important question, which is beyond the scope of this study. We plan to follow up on this issue in our future study. The following discussions have been added in the revised manuscript:

Lines 194-201: “We further computed $\tau_{eq}$ as a function of $D_b$ and $C^0$ in the closed system. As shown in Fig. 4a, when $D_b$ is higher than $\sim 10^{-13}$ cm$^2$ s$^{-1}$, $\tau_{eq}$ is insensitive to bulk diffusivity but sensitive to volatility: decreasing volatility increases $\tau_{eq}$ in this regime. In the regime with $D_b$ lower than $\sim 10^{-13}$ cm$^2$ s$^{-1}$ and $C^0$ higher than $\sim 10 \mu g m^{-3}$, $\tau_{eq}$ is controlled by bulk diffusivity: $\tau_{eq}$ increases from 30 s to longer than 1 year as $D_b$ decreases from $10^{-13}$ cm$^2$ s$^{-1}$ to $10^{-20}$ cm$^2$ s$^{-1}$. In the regime with $D_b < \sim 10^{-13}$ cm$^2$ s$^{-1}$
and $C^0 < \sim 10 \, \mu g \, m^{-3}$, $\tau_{eq}$ depends on both diffusivity and volatility. Decreasing volatility would lead to shorter $\tau_{eq}$ due to an establishment of local equilibrium of LVOC”.

Lines 221-225: “Figure 4b shows simulated evaporation timescales as a function of $D_b$ and $C^0$ in an open system, which agrees very well with Fig. 3 in Liu et al. (2016). It shows that for less viscous particles $\tau_{eq}$ is limited by volatility, while for highly viscous particles $\tau_{eq}$ is insensitive to volatility and controlled by bulk diffusivity”.

Lines 127-128: “Particle-phase reactions and their potential impacts on particle viscosity are also not considered in this study”.

Lines 361-369: “Incorporation of the particle-phase formation of oligomers and other multifunctional high molar mass compounds can lead to a reduced bulk diffusivity (Pfrang et al., 2011; Hosny et al., 2016), which may prolong the equilibration timescales. Decomposition of highly oxidized molecules (e.g., organic hydroperoxides) in water may also affect gas-particle partitioning (Tong et al., 2016).

Current simulations are focused on trace amount of SVOC or LVOC condensing on mono-dispersed particles with negligible particle growth. Potential phase transition in the course of particle growth/evaporation should also be incorporated in future simulations”.

Figure 4. Contour plot of equilibration timescale ($\tau_{eq}$) as a function of bulk diffusivity ($D_b$) and saturation mass concentration ($C^0$) for (a) condensation in the closed system and (b) evaporation in the open system. The initial mass concentration of pre-existing particles is set to be $20 \, \mu g \, m^{-3}$ with the number concentrations of $3 \times 10^4 \, cm^{-3}$ and the initial particle diameter of 100 nm. Viscosity is calculated from the Stokes-Einstein equation assuming the effective molecular radius of $10^{-8} \, cm$ at $T$ of 298 K.

(3) My other question is that most of the modeling results shown the manuscript assumed that the gas-particle is in a closed system. How realistic is the closed system in ambient environment? Would the ambient environment often be an open system for
evaporation kinetics?
Response: Thanks for this helpful comment. To address this question, we add simulations for an open system (Fig. 4b, S5, and S7) in the revised manuscript. The following discussions have been added in the revised manuscript.

Lines 202-208: “In an open system with fixed vapor concentration (Fig. S5), $\tau_{eq}$ of SVOC is slightly longer but on the same order of magnitude as $\tau_{eq}$ in the closed system, as relatively small amounts of SVOC need to condense to reach equilibrium. In contrast, $\tau_{eq}$ of LVOC in the open system become dramatically longer as LVOC continue to condense into the particle phase because of low volatility (Pankow, 1994). For further simulations we focus mainly on the closed system and the corresponding simulations for the open system are provided in the supplement”.

Lines 221-225: “Figure 4b shows simulated evaporation timescales as a function of $D_b$ and $C^0$ in an open system, which agrees very well with Fig. 3 in Liu et al. (2016). It shows that for less viscous particles $\tau_{eq}$ is limited by volatility, while for highly viscous particles $\tau_{eq}$ is insensitive to volatility and controlled by bulk diffusivity”.

Lines 242-244: “The corresponding simulations of SVOC partitioning in the open system (Fig. S7) show a similar pattern as $\tau_{eq}$ in the closed system”.

Lines 306-312: “The timescale of gas-particle partitioning can be different in closed or open systems especially for LVOC (Fig. 4, S7). The closed system simulations represent SOA partitioning in chamber experiments and in closed atmospheric air mass, which could be justified well within seconds-to-minutes timescales and possibly up to hours depending on meteorological conditions. The real atmosphere may be approximated better as an open system due to dilution and chemical production and loss especially at longer timescales”.

**Figure S5.** Temporal evolution of mass concentrations of the condensing compound Z in the gas phase ($C_g$), just above the particle surface ($C_s$), and in the particle phase ($C_p$) in the open system. $\tau_{eq}$ is marked with the red circle. RH = 60% and $T$ is (a–b) 298 K and (c–d) 250 K. The $C^0$ of Z is (a, c) 10 µg m$^{-3}$ and (b, d) 0.1 µg m$^{-3}$. The
glass transition temperature of pre-existing particles under dry conditions ($T_{g,\text{org}}$) is set to be 270 K, which leads to $D_b$ of (a–b) $10^{11}$ cm$^2$ s$^{-1}$ and (c–d) $10^{18}$ cm$^2$ s$^{-1}$. The initial mass concentration of pre-existing particles is set to be 20 µg m$^{-3}$ with the number concentrations of $3 \times 10^4$ cm$^{-3}$ and the initial particle diameter of 100 nm.

**Figure S7.** Equilibration timescale ($\tau_{eq}$) as a function of temperature and relative humidity in the open system. The glass transition temperatures of pre-existing particles at dry conditions ($T_{g,\text{org}}$) are (a) 240 K, (b) 270 K, and (c) 300 K, respectively. The saturation mass concentration ($C^0$) of the condensing compound is 10 µg m$^{-3}$ (SVOC). The mass concentration of pre-existing particles is set to be 20 µg m$^{-3}$ with the number concentrations of $3 \times 10^4$ cm$^{-3}$ and the initial particle diameter of 100 nm.

**Referee Minor Comments:**
The author should also include Price et al. 2015 in the reference list in line 115.

Reference: Shiraiwa et al. 2011: doi.org/10.1073/pnas.1103045108
Shiraiwa et al. 2012: doi.org/10.1029/2012GL054008
Price et al. 2015: doi.org/10.1039/C5SC00685F
Response: Price et al. (2015) has been included on Line 123 in the revised manuscript. Besides this, Liu et al. (2013) has been included on Lines 181, 259 and 357.

Response to the comments of Anonymous Referee #2

Referee General Comment:
In this manuscript, the authors use simulations to calculate the gas/particle equilibration time for secondary organic aerosol as a function of temperature, relative humidity, and SOA microphysical properties. The topic of SOA partitioning, equilibration, and phase state are highly relevant in atmospheric chemistry, the topic is timely, and will be of interest to the readers of ACP. This work is a logical extension of the previous work done by the PI. It essentially combines work in predicting the phase state of particles as a function of atmospheric conditions with work on calculating gas/particle equilibration times (Shiraiwa et al., 2017; Shiraiwa and Seinfeld, 2012). Based on past results from the PI on these topics, the present results are not particularly surprising, but I think there is enough new material here to warrant publication. This publication essentially closes the loop between predicting phase state and calculating the gas/particle equilibration time. With that said, there are a few areas that could be improved before publication. The authors could do a better job of calling out, in the manuscript, what is new unique about this manuscript relative to the previous publications by the PI. A few of the conclusions reached in this work also seem to contradict the PI’s previous publications and the authors should clear this up. The authors could help readers put this work into context if they explain why they make certain assumptions in their model (i.e., a closed system), offer some insights into how realistic these assumptions are relative to the ambient atmosphere, and explain how their conclusions would be different if/when the assumptions are not atmospherically representative. While the manuscript is generally well written and clear, there was one section that was somewhat confusing and should be clarified before publication. With that said, there are no major shortcomings with the manuscript and, providing the authors make some revisions, I have no reservations about recommending this manuscript for publication in ACP.

Responses: We thank Anonymous Referee #2 for the review and the positive evaluation of our manuscript. As you pointed out, this is the first study to directly relate equilibration timescale of SOA partitioning to ambient temperature and relative humidity, which has important implications in treatment of SOA evolution in chemical transport models. The novelty of the revised manuscript is further strengthened by two additional new results. Firstly, we add a contour plot of $\tau_{eq}$ as a function of bulk diffusivity and volatility to define the regimes of diffusivity-limited and volatility-limited partitioning. Secondly, we add simulations for open systems and the results are compared with $\tau_{eq}$ in closed systems. The implications of $\tau_{eq}$ in closed and open systems are further broadened for SOA evolution in ambient atmosphere and chemical transport models. Following your suggestion, we clarify that apparently contradicting conclusions regarding $\tau_{eq}$ of LVOC actually are consistent with PI’s previous publication (e.g., Shiraiwa & Seinfeld, 2012). We also revise the last figure and associated section for better presentation of our results. Please see the detailed response below.

Referee Major Comment:
In the present manuscript the authors conclude that the equilibration timescale for low volatile compounds is shorter than for semi-volatile compounds when other conditions are equal. However, Shiraiwa and Seinfeld 2012 report the opposite (see for example Figure 2), with ELVOC’s having longer equilibration times than SVOCs (Shiraiwa and Seinfeld, 2012). The authors should comment in the text on why these studies reach opposite conclusions.

Response: The results in this study are actually consistent with Shiraiwa and Seinfeld (2012) even though our previous statements on Lines 23-25 were somewhat misleading. Figure 2 in Shiraiwa and Seinfeld (2012) was presented for liquid particles showing that $\tau_{eq}$ of LVOC is longer, which agreed with the simulations in our current study showing that for less viscous particles LVOC takes longer time than SVOC to reach equilibrium (Fig. 2a-b). Shiraiwa and Seinfeld (2012) did not compare $\tau_{eq}$ of LVOC and SVOC condensing on highly viscous particles, which has been simulated in current study showing that $\tau_{eq}$ of LVOC is shorter (Fig. 2c-d). We clarified this point on Line 23 in the revised manuscript. In addition, we add Fig. 4 in the revised manuscript to systematically evaluate the dependence of $\tau_{eq}$ on both volatility and bulk diffusivity. Please also refer to our response to Comment (2) of Referee #1.

(2) Lines 115-123. I understand that the authors need to make some assumptions or approximations in their model/calculations. I am trying to understand how atmospherically realistic these assumptions are. The two main assumptions in the present model are of a closed system and that the condensation of molecule Z does not alter the composition and microphysical properties of the pre-existing particles. It is clear that the real atmosphere isn’t a closed system. The argument could be made that on seconds-to-minutes timescales, it may approximate a closed system, but the processes that authors are modeling are sometimes occurring on timescales of hours or even days. In addition, one of the author’s major conclusions is that low volatility material reaches equilibrium more slowly than higher volatility material. I understand why this is the case in a closed system, but would this conclusion also hold in an open system like in the atmosphere with a constant dilution and/or loss of gas-phase molecules? A plume transported from the surface to the upper troposphere would experience an evaporative driving force where this model seems focused on cases where the driving force is toward the particle phase (condensation). With respect to a single compound (Z) changing the composition and microphysical properties, it may be true that a single molecule or even a few molecules rarely make up the bulk of the SOA mass. However, in the real atmosphere, particles obviously grow and their composition and microphysical properties change as SOA condenses. The PI of this manuscript has previously used kinetic modeling to reproduce particle growth, so I’m not really clear on why these assumptions needed to be made (Shiraiwa et al., 2013). In the both cases, I think it is important that the authors explain: why they chose to make these assumptions, how likely it is that these assumptions are representative of the atmosphere, and how their conclusions would likely be different if the assumptions are not correct.

Response: Thanks for this helpful comment. Based on your suggestions we add
simulations for an open system (Fig. 4b, S5 and S7) in the revised manuscript. Figure S5 and S7 show that for condensation of SVOC, $\tau_{eq}$ is slightly longer but on the same order of magnitude as $\tau_{eq}$ in the closed system. In contrast, $\tau_{eq}$ of LVOC condensation in the open system become dramatically longer as LVOC keep condensing into the particle phase because of low volatility. For evaporation in an open system with continuous removal/dilution of gas-phase LVOC, $\tau_{eq}$ of LVOC is also much longer than that in a closed system due to continuous evaporation (Fig. 4b). For the details please refer to the response to Comment (3) of Referee #1. The implications of $\tau_{eq}$ in closed versus open systems in SOA evolution are broadened. In the revised manuscript we state that:

Lines 306-312: “The timescale of gas-particle partitioning can be different in closed or open systems especially for LVOC (Fig. 4, S7). The closed system simulations represent SOA partitioning in chamber experiments and in closed atmospheric air mass, which could be justified well within seconds-to-minutes timescales and possibly up to hours depending on meteorological conditions. The real atmosphere may be approximated better as an open system due to dilution and chemical production and loss especially at longer timescales”.

We agree that condensation of substantial amounts of materials may lead to changes in particle microphysical properties including phase state and viscosity, which is beyond the scope of current study, even though this is indeed an important aspect. In this study we let only trace amounts to condense so that physical properties including size and phase state would remain unaffected. As KM-GAP can indeed treat evolution of particle properties upon particle growth/evaporation, we plan to explore this aspect systematically by varying particle-phase reaction rates and resulting impacts on phase state in future studies. Following your suggestion, in the revised manuscript we broaden the discussion as below:

Lines 361-369: “Incorporation of the particle-phase formation of oligomers and other multifunctional high molar mass compounds can lead to a reduced bulk diffusivity (Pfrang et al., 2011; Hosny et al., 2016), which may prolong the equilibration timescales. Decomposition of highly oxidized molecules (e.g., organic hydroperoxides) in water may also affect gas-particle partitioning (Tong et al., 2016). Current simulations are focused on trace amount of SVOC or LVOC condensing on mono-dispersed particles with negligible particle growth. Potential phase transition in the course of particle growth/evaporation should also be incorporated in future simulations”.

(3) Lines 276-292 and Figure 5. This figure and associated text was confusing. The figure was confusing because particle diameter and total mass loading are typically not independent of one another in a model or in the atmosphere. I eventually understood the point the authors were trying to make. Perhaps providing some context by pointing out where different atmospheric regimes lie (i.e., remote, typical continental, polluted) in the figure would help. The associated text is also confusing; it wasn’t clear what point the authors were trying to make here. They seem to postulate several different processes which could determine particle equilibration
timescales (e.g., bulk diffusion, gas-diffusion). Can’t the KM-GAP model be used to clear this up? Overall, I’m not sure what message the authors are trying to convey here.

Response: Following your suggestion, we indicate typical ambient organic mass concentrations in Beijing (Liu et al., 2017), southeastern US (Pajunoja et al., 2016), Amazon Basin (Bateman et al., 2017), and Hyytiälä, Finland (Virtanen et al., 2010) in Fig. 6, where ambient phase state measurements are available. Figure 6 indeed has implications on how different ambient conditions have effect on SOA partitioning. For clarification, the following discussions have been added in the revised manuscript.

Lines 259-261: “Previous studies have shown that $\tau_{eq}$ depends on particle size (Liu et al., 2013; Zaveri et al., 2014; Mai et al., 2015) and particle mass loadings (Shiraiwa and Seinfeld, 2012; Saleh et al., 2013). For further examination of these effects at different $T$…”.

Lines 275-285: “Typical ambient organic mass concentrations in Beijing, Centreville in southeastern US, Amazon Basin, and Hyytiälä, Finland are indicated in Fig. 6. The particle phase state was observed to be mostly liquid in highly polluted episodes in Beijing (Liu et al., 2017), under typical atmospheric conditions in the southeastern US (Pajunoja et al., 2016), and under background conditions in Amazonia (Bateman et al., 2017). At these conditions $\tau_{eq}$ should be mostly less than 30 minutes (Fig. 6a, b). Particles were measured to be semi-solid or amorphous solid in clear days in Beijing (Liu et al., 2017), in Amazonia when influenced by anthropogenic emissions (Bateman et al., 2017), and the boreal forest in Finland (Virtanen et al., 2010). Under these conditions and also when particles are transported to the free troposphere, $\tau_{eq}$ can be longer than 1 hour especially in remote areas with low mass loadings (Fig. 6c, d)”.


Figure 6. Equilibration timescale ($\tau_{eq}$) for (a, c) SVOC ($C^0 = 10$ µg m$^{-3}$) and (b, d) LVOC ($C^0 = 0.1$ µg m$^{-3}$) as a function of particle diameter (nm) and mass concentration (µg m$^{-3}$) of pre-existing particles at 60% RH and $T$ of (a-b) 298 K and (c-d) 250 K in the closed system. The glass transition temperature of pre-existing particles under dry conditions ($T_{g,org}$) is set to be 270 K, which leads to $D_b$ of (a-b) $10^{-11}$ cm$^2$ s$^{-1}$ and (c-d) $10^{-18}$ cm$^2$ s$^{-1}$. Ambient organic mass concentrations are indicated with arrows.

Referee Minor Comments and Technical Corrections:

(1) Line 89. It is a little confusing here about what temperature was used in the calculations. I wasn’t clear whether 273.15 K was used (the most common definition of standard temperature), or if the temperature was variable as a function of pressure altitude. Adding to the confusion, Table S2 lists standard temperature as 288.15 K. Please clarify.

Response: In our simulations the temperature is varied from 220 K to 310 K (Fig. 5) while atmospheric pressure is calculated as a function of $T$ based on the International Standard Atmosphere (ISA): $P/P_{standard} = (T/T_{standard})^{P_{LR}}$, where $P_{standard}$ and $T_{standard}$ are standard sea level $P$ and $T$ in ISA, and $L$ is the lapse rate of 6.5 K/km in the troposphere. This has been clarified on Lines 94-95 and Table S2 in the revised manuscript.

(2) Lines 108-110. An assumption here is that the organic and aqueous phases are not phase separated. The authors point out later in the manuscript that phase separations may occur, but I suggest also briefly mentioning that phase separation has been observed for laboratory generated SOA (You et al., 2012) here, since it is very relevant to their modeling results.
Response: The following sentence has been added on Lines 116-119 in the revised manuscript:
“For simplicity we assume particles are ideally-mixed, even though phase-separated particles are observed for ambient and laboratory generated SOA particles under certain conditions (You et al., 2012; Renbaum-Wolff et al., 2016”).

(3) Lines 144 and elsewhere. I found the use of $C_0$ as an abbreviation more confusing than necessary. It seems $C_0$ is identical to the much more commonly used $C^*$. Why not use the commonly accepted $C^*$? The authors also use $C_{p,0}$ and $C_{g,0}$, which have a different meaning and cause some confusion with $C_0$. Whatever symbol the author use for the saturation vapor pressure please define it the first time it is used.
Response: Instead of $C_0$, $C^0$, which is commonly used for the pure compound saturation mass concentration, is used throughout the revised manuscript. The effective saturation mass concentration $C^*$ is not used as it includes the effect of non-ideal thermodynamic mixing which is not considered in this study. Lines 154-157 have been re-written in the revised manuscript as:
“Figure 2a presents simulations for a semi-volatile organic compound (SVOC) with the pure compound saturation mass concentration ($C^0$) of 10 µg m$^{-3}$ condensing on particles with $D_b$ of 10$^{-11}$ cm$^2$ s$^{-1}$ at RH = 60% and $T = 298$ K (Fig. S2)”.

(4) Figures 4, A1, A2. The labels on different contours were illegible on the printed document.
Response: The resolution of the figures has been improved in the revised manuscript.
Timescales of Secondary Organic Aerosols to Reach Equilibrium at Various Temperatures and Relative Humidities

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Abstract:

Secondary organic aerosols (SOA) account for a substantial fraction of air particulate matter and SOA formation is often modeled assuming rapid establishment of gas-particle equilibrium. Here, we estimate the characteristic timescale for SOA to achieve gas–particle equilibrium under a wide range of temperatures and relative humidities using a state-of-the-art kinetic flux model. Equilibration timescales were calculated by varying particle phase state, size, mass loadings, and volatility of organic compounds in open and closed systems. Model simulations suggest that the equilibration timescale for semi-volatile compounds is on the order of seconds or minutes for most conditions in the planetary boundary layer, but it can be longer than one hour if particles adopt glassy or amorphous solid states with high glass transition temperature at low relative humidity. In the free troposphere with lower temperatures it can be longer than hours or days even at moderate or relatively high relative humidity due to kinetic limitations of bulk diffusion in highly viscous particles. The timescale of partitioning of low-volatile compounds into highly viscous particles is shorter compared to semi-volatile compounds in the closed system, as it is largely determined by condensation sink due to very slow re-evaporation with relatively quick establishment of local equilibrium between the gas phase and the near-surface bulk. The dependence of equilibration timescales on both volatility and bulk diffusivity provides critical insights into thermodynamic or kinetic treatments of SOA partitioning for accurate predictions of gas- and particle-phase concentrations of semi-volatile compounds in regional and global chemical transport models.
1. Introduction

Secondary organic aerosols (SOA) play a central role in climate, air quality and public health. Accurate descriptions of formation and evolution of SOA remain a grand challenge in climate and air quality models (Kanakidou et al., 2005; Shrivastava et al., 2017a). Current chemical transport models usually employ instantaneous equilibrium partitioning of semi-volatile oxidation products into the particle phase (Pankow, 1994), assuming that SOA partitioning is rapid compared to the timescales of other major atmospheric processes associated with SOA formation. The timescale of SOA to reach equilibrium with their surrounding condensable vapors needs to be evaluated under different ambient conditions to validate this assumption.

SOA particles can adopt liquid (dynamic viscosity $\eta < 10^2$ Pa s), semi-solid ($10^2 \leq \eta \leq 10^{12}$ Pa s), or glassy or amorphous solid states ($\eta > 10^{12}$ Pa s), depending on chemical composition, temperature ($T$) and relative humidity (RH) (Virtanen et al., 2010; Koop et al., 2011; Zhang et al., 2015; Reid et al., 2018). The occurrence of glassy or amorphous solid states may lead to kinetic limitations and prolonged equilibration timescale in SOA partitioning (Shiraiwa and Seinfeld, 2012; Booth et al., 2014; Zaveri et al., 2014; Mai et al., 2015), affecting evolution of particle size distribution upon SOA growth (Maria et al., 2004; Shiraiwa et al., 2013a; Zaveri et al., 2018). A number of experimental studies have indeed observed kinetic limitations of the bulk diffusion of organic molecules (Vaden et al., 2011; Perraud et al., 2012; Ye et al., 2016a; Zhang et al., 2018), while chamber experiments probing the
intraparticle mixing did not find kinetic limitations at moderate and high RH and room temperature (Ye et al., 2016b; Gorkowski et al., 2017; Ye et al., 2018).

Recently, global simulations predicted that SOA particles are expected to be mostly in a glassy solid phase state in the middle and upper troposphere and also in dry lands in the boundary layer (Shiraiwa et al., 2017), which can lead to prolonged characteristic bulk diffusion timescales of organic molecules within SOA particles (Shiraiwa et al., 2011; Maclean et al., 2017). Slow bulk diffusion associated with a glassy phase state can prevent atmospheric oxidants to react with organic compounds such as polycyclic aromatic hydrocarbons (Shrivastava et al., 2017b; Mu et al., 2018), contributing to long-range transport of organic compounds. Recent ambient observations have shown that the condensation of highly oxygenated molecules (HOMs), which play an important role in new particle formation, would be governed by kinetic partitioning in the free troposphere (Bianchi et al., 2016). Diffusivity measurements of volatile organics in levitated viscous particles have shown strong temperature dependence of bulk diffusivity and evaporation timescale (Bastelberger et al., 2017). Slow bulk diffusion may impact multiphase processes such as browning of organic particles (Liu et al., 2018), cloud droplet activation (Slade et al., 2017), and ice nucleation pathways (Knopf et al., 2018).

Given these observations and strong implications of SOA phase states, it is important to evaluate common assumption of gas-particle partitioning equilibrium at different ambient conditions. In this study we provide theoretical analysis of partitioning kinetics of organic compounds using the kinetic multi-layer model of
gas-particle interactions in aerosols and clouds (KM-GAP) (Shiraiwa et al., 2012), which accounts for mass transport in both gas and particle phases. The equilibration timescale ($\tau_{\text{eq}}$) of organic compounds partitioning into mono-dispersed particles is evaluated systematically under a wide range of temperatures and RH, considering the effects of particle phase state, particle size, mass loadings, and volatility of organic compounds in a closed system with finite amount of vapor. For comparison we also present simulations in an open system with vapor concentration maintained as constant. This is the first study to directly relate equilibration timescale of SOA partitioning to ambient temperature and relative humidity, which has important implications in treatment of SOA evolution in chemical transport models.

2. Methods

We evaluate the timescale to achieve gas-particle equilibrium by simulating condensation of a compound Z into pre-existing non-volatile mono-dispersed particles using the KM-GAP model. KM-GAP consists of multiple model compartments and layers, respectively: gas phase, near-surface gas phase, sorption layer, surface layer, and a number of bulk layers (Shiraiwa et al., 2012). The following processes are treated as temperature-dependent in KM-GAP: gas phase diffusion, adsorption/desorption, surface-bulk exchange, and bulk diffusion (Fig. S1). The physical and kinetic parameters are summarized in Table S1. The gas-phase diffusion coefficient depends on temperature ($T$) and ambient pressure ($P$). $P$ is calculated as a function of $T$ based on the International Standard Atmosphere.
The adsorption rate coefficient is related to the mean thermal velocity as a function of $T$ and the surface accommodation coefficient, which is assumed to be 1 (Julin et al., 2014). The $T$-dependence of desorption rate coefficient is described by an Arrhenius equation with an assumed typical adsorption enthalpy of 40 kJ mol$^{-1}$.

Phase state and viscosity can be characterized by the glass transition temperature ($T_g$), at which phase transition between amorphous solid and semi-solid states occurs (Koop et al., 2011). When $T_g$ of organic particles under dry conditions ($T_{g,org}$) is known, $T_g$ of organic-water mixtures at given RH can be estimated considering hygroscopic growth combined with the Gordon-Taylor equation. In this work, we assumed the effective hygroscopicity parameter as 0.1 (Petters and Kreidenweis, 2007; Gunthe et al., 2009) and the Gordon-Taylor constant as 2.5 (Koop et al., 2011). Then, the $T$-dependence of viscosity is calculated using the Vogel-Tammann-Fulcher equation (Angell, 1991; Rothfuss and Petters, 2017; DeRieux et al., 2018; Li and Shiraiwa, 2018).

Figure 1 shows the $T$- and RH-dependent viscosity of SOA particles with $T_{g,org}$ of (a) 240 K, (b) 270 K, and (c) 300 K. We chose these three $T_{g,org}$ values to represent different phase states of liquid, semi-solid, and glassy states, respectively, at $T$ of 298 K under dry conditions and these values are within the range recently reported for monoterpene-derived SOA (Petters et al., 2019). The decrease of $T$ leads to increase of viscosity, while the increase of RH leads to decrease of viscosity due to the plasticizing effect of water (Koop et al., 2011). For simplicity we assume particles are
ideally-mixed, even though phase-separated particles are observed for ambient and laboratory generated SOA particles under certain conditions (You et al., 2012; Renbaum-Wolff et al., 2016). The bulk diffusion coefficient $D_b$ (Fig. S2) is calculated by the Stokes–Einstein equation, which has been shown to work very well for organic molecules diffusing through materials with viscosity below $\sim 10^3$ Pa s (Chenyakin et al., 2017). Note that the Stokes–Einstein equation may underpredict $D_b$ in highly viscous SOA thus it gives lower limits of $D_b$ (Price et al., 2015; Marshall et al., 2016; Bastelberger et al., 2017; Reid et al., 2018). $D_b$ is fixed at any given depth in the particle bulk in each simulation, assuming that condensation of Z would not alter particle viscosity and diffusivity as only trace amounts of Z condense to pre-existing particles in our simulations. Particle-phase reactions and their potential impacts on particle viscosity are also not considered in this study.

We mainly consider a closed system, in which condensation of Z would lead to a decrease of its gas-phase mass concentration ($C_g$) and an increase of its particle-phase mass concentration ($C_p$). The particle diameter stays practically constant throughout each simulation, as the amount of condesing Z is set to be much smaller than the non-volatile pre-existing particle mass ($C_{OA}$). The gas-phase mass concentration of Z right above the surface ($C_s$) is also calculated based on the Raoult’s law and partitioning theory (Pankow, 1994) in equilibrium with the near-surface bulk, which is resolved by KM-GAP (Shiraiwa and Seinfeld, 2012). We also calculate the mass fraction of Z in the near-surface bulk ($f_s$) and the average mass fraction of Z in
the entire bulk ($f_b$) to infer the radial concentration profile (Fig. S3). The equilibration timescale ($\tau_{eq}$) is calculated as the e-folding time $t$ when the following criterion is met,

$$\frac{|c_p(t) - c_{p,eq}|}{|c_{p,0} - c_{p,eq}|} < \frac{1}{e}$$

(1)

where $c_{p,0}$ and $c_{p,eq}$ are the initial and equilibrium mass concentration of $Z$ in the particle phase, respectively. Note that practically the same values can also be obtained by using initial and equilibrium gas-phase concentrations in Eq. (1), as the mass change of $Z$ in the gas and particle phases are the same in these simulations.

3 Results

3.1. Impacts of volatility and diffusivity on equilibration timescales

Figure 2 shows exemplary simulations of temporal evolution of $c_g$ (blue line) and $c_p$ (red line) of the compound $Z$ in the closed system along with $\tau_{eq}$ marked with red circles. The initial mass concentration of pre-existing non-volatile mono-dispersed particles ($C_{OA}$) is assumed to be 20 µg m$^{-3}$ with the number concentrations of $3 \times 10^4$ cm$^{-3}$ and the initial particle diameter of 100 nm. Initial mass concentrations of $Z$ in the gas ($C_{g,0}$) and particle ($C_{p,0}$) phases are set to be 0.3 µg m$^{-3}$ and 0 µg m$^{-3}$, respectively. $T_{g,org}$ is assumed to be 270 K. Figure 2a presents simulations for a semi-volatile organic compound (SVOC) with the pure compound saturation mass concentration ($C^0$) of 10 µg m$^{-3}$ condensing on particles with $D_b$ of $10^{-11}$ cm$^2$ s$^{-1}$ at RH = 60% and $T$ = 298 K (Fig. S2). Upon condensation $c_g$ decreases, while $c_s$ and $c_p$ increase, and the gas-particle equilibrium is reached within about 20 s as indicated by $\tau_{eq}$. For low-volatile organic compounds (LVOC) with $C^0 = 0.1$ µg m$^{-3}$, it takes longer time to
reach the equilibrium with $\tau_{eq}$ of $\sim 30$ s (Fig. 2b), as the partial pressure gradient between the gas phase and the particle surface (represented by the difference between $C_g$ and $C_s$) is larger for lower $C^0$. For both cases SOA growth is governed by gas-phase diffusion as indicated by $C_s < C_g$. The mass fraction of Z in the near-surface bulk is identical to the average mass fraction in the entire bulk (Fig. S3 a–b), indicating that Z are homogeneously well-mixed in the particle without kinetic limitations of bulk diffusion in low viscous particles (Fig. 3a).

At lower $T$ of 250 K, the phase state of pre-existing particles occurs as highly viscous with $D_b$ of $\sim 10^{-18}$ cm$^2$ s$^{-1}$ (Fig. S2), resulting in much longer equilibration timescales ($\sim 10^5$ s) for SVOC with $C^0 = 10$ µg m$^{-3}$ (Fig. 2c). After $C_g$ and $C_s$ converge, they continue to decrease simultaneously while $C_p$ increases slowly, showing that the particle undergoes quasi-equilibrium growth (Shiraiwa and Seinfeld, 2012; Zhang et al., 2012). For LVOC ($C^0 = 0.1$ µg m$^{-3}$) condensation, $\tau_{eq}$ is short ($\sim 140$ s) because of a local thermodynamic equilibrium between the gas phase and the near-surface bulk established relatively quickly (as mostly controlled by the condensation sink: Riipinen et al., 2011; Tröstl et al., 2016) due to very slow re-evaporation of LVOC.

The characteristic timescale of mass transport and mixing by molecular diffusion $\tau_{mix}$ can be calculated by $\tau_{mix} = r_p^2 / (\pi^2 D_b)$, where $r_p$ is the particle radius (Seinfeld and Pandis, 2006). Figure 3 shows dimensionless radial concentration profiles of Z ($C^0 = 0.1$ µg m$^{-3}$) in the particle at (a) $D_b = 10^{-11}$ cm$^2$ s$^{-1}$ and (b) $10^{-18}$ cm$^2$ s$^{-1}$, respectively. For low viscous particles, $\tau_{mix}$ is very short and particles are
homogeneously well-mixed at $\tau_{eq}$, which is consistent with previous analytical calculations (Liu et al., 2013; Mai et al., 2015). In contrast, there exists a large concentration gradient between the particle surface and the inner bulk (Fig. 3b, S3d) at $\tau_{eq}$ in highly viscous particles due to strong kinetic limitations of bulk diffusion (as indicated by very long $\tau_{mix}$), which prevents the entire particle bulk to reach complete equilibrium. Thus, for LVOC condensation on highly viscous particles (Fig. 2d), $\tau_{mix}$ represents the timescale to establish full equilibrium with homogeneous mixing in the entire particle bulk. These results are consistent with Mai et al. (2015) and Liu et al. (2016), which showed that an establishment of full equilibrium is limited by bulk diffusion in highly viscous particles, even though the local equilibrium of LVOC may be achieved faster. Note that $\tau_{mix}$ is solely a function of particle size and bulk diffusivity, while $\tau_{eq}$ is also affected by volatility and mass loadings. At lower particle concentrations, the total accommodation of molecules to the particle surface decreases, resulting in longer equilibration timescales (Fig. S4).

We further computed $\tau_{eq}$ as a function of $D_b$ and $C^0$ in the closed system. As shown in Fig. 4a, when $D_b$ is higher than $\sim10^{-13}$ cm$^2$ s$^{-1}$, $\tau_{eq}$ is insensitive to bulk diffusivity but sensitive to volatility: decreasing volatility increases $\tau_{eq}$ in this regime. In the regime with $D_b$ lower than $\sim10^{-13}$ cm$^2$ s$^{-1}$ and $C^0$ higher than $\sim10$ µg m$^{-3}$, $\tau_{eq}$ is controlled by bulk diffusivity: $\tau_{eq}$ increases from 30 s to longer than 1 year as $D_b$ decreases from $10^{-13}$ cm$^2$ s$^{-1}$ to $10^{-20}$ cm$^2$ s$^{-1}$. In the regime with $D_b < \sim10^{-13}$ cm$^2$ s$^{-1}$ and $C^0 < \sim10$ µg m$^{-3}$, $\tau_{eq}$ depends on both diffusivity and volatility. Decreasing
volatility would lead to shorter $\tau_{eq}$ due to an establishment of local equilibrium of LVOC.

In an open system with fixed vapor concentration (Fig. S5), $\tau_{eq}$ of SVOC is slightly longer but on the same order of magnitude as $\tau_{eq}$ in the closed system, as relatively small amounts of SVOC need to condense to reach equilibrium. In contrast, $\tau_{eq}$ of LVOC in the open system become dramatically longer as LVOC continue to condense into the particle phase because of low volatility (Pankow, 1994). For further simulations we focus mainly on the closed system and the corresponding simulations for the open system are provided in the supplement.

We also simulated evaporation in the closed system with same parameters as the condensation simulations (Table S2). Initially $C_g = 0 \ \mu g \ m^{-3}$ and trace amounts of semi-volatile or low-volatile species were assumed to be homogeneously well-mixed in pre-existing particles. Figure S6 shows that for the evaporation of SVOC species with $C^0 = 10 \ \mu g \ m^{-3}$, decreasing $D_b$ from $10^{-11} \ \text{cm}^2 \ \text{s}^{-1}$ to $10^{-18} \ \text{cm}^2 \ \text{s}^{-1}$ would increase $\tau_{eq}$ from $\sim 20 \ \text{s}$ to $\sim 10^5 \ \text{s}$. These evaporation timescales are close to those derived from condensation (Fig. 2a,c) and consistent with previous kinetic simulations (Liu et al., 2016). In the closed system, the evaporation of a very small amount of LVOC species from the particle surface is already sufficient to reach the particle-phase equilibrium concentration, resulting in a short $\tau_{eq}$ (Fig. S6b,d). For an open system with continuous removal of gas-phase compounds, which is often employed in evaporation experiments, the equilibrium timescale in the evaporation of the LVOC species from highly viscous particles can be longer than hours or days (Vaden et al.,
2011; Liu et al., 2016). Figure 4b shows simulated evaporation timescales as a function of $D_b$ and $C^0$ in an open system, which agrees very well with Fig. 3 in Liu et al. (2016). It shows that for less viscous particles $\tau_{eq}$ is limited by volatility, while for highly viscous particles $\tau_{eq}$ is insensitive to volatility and controlled by bulk diffusivity.

3.2. Equilibration timescales at different RH and $T$

We conducted further simulations to estimate $\tau_{eq}$ with a wide range of atmospherically-relevant temperatures (220 - 310 K) and relative humidities (0 - 100%). Figure 5 shows the temperature and humidity-dependent diagrams of $\tau_{eq}$ for SVOC ($C^0 = 10 \mu g m^{-3}$) condensation on particles with $T_{g,org}$ of 240 K, 270 K, and 300 K, respectively, in the closed system. For particles with $T_{g,org}$ of 240 K (panel a), $\tau_{eq}$ is on the order of seconds under boundary layer conditions ($T > 270$ K). In these conditions particles are liquid with high bulk diffusivity (Fig. 1a and S2a), thus gas-particle partitioning is controlled by gas-phase diffusion and interfacial transport (Shiraiwa and Seinfeld, 2012; Mai et al., 2015). At low $T$ (< 260 K) with low or moderate RH (< 70%), $\tau_{eq}$ can increase from minutes to one year with decreasing $T$ and RH mainly due to strong kinetic limitations of bulk diffusion with low $D_b$ (Fig. S2a). With $T_{g,org}$ of 270 K (panel b) or 300 K (panel c), $\tau_{eq}$ is still on the order of minutes in most of boundary layer conditions. At low RH $\tau_{eq}$ can be extended to hours when particles may occur as amorphous (semi-)solid. When $T < 270$ K, $\tau_{eq}$ can be longer than months even at moderate RH, while $\tau_{eq}$ may stay very short at very high
The corresponding simulations of SVOC partitioning in the open system (Fig. S7) show a similar pattern as $\tau_{eq}$ in the closed system.

$\tau_{eq}$ for $C^0 = 10^3$ and 0.1 µg m$^{-3}$ in the closed system are presented in Fig. A1. In general, $\tau_{eq}$ would be shorter at higher $C^0$ when particles are liquid, as the partial pressure gradient between the gas phase and the particle surface is smaller for higher $C^0$ (Shiraiwa and Seinfeld, 2012; Liu et al., 2016). For example, the increase of $C^0$ from 10 µg m$^{-3}$ to $10^3$ µg m$^{-3}$ leads to $\tau_{eq}$ decrease from 30 s to 1 s with $T_{g,org}$ of 240 K at boundary layer conditions (Fig. 5a, A1a). At low $T$ and RH (e.g., $T < 250$ K and RH < 50 %) where particles are highly viscous, $\tau_{eq}$ is on the same order of magnitude for the condensation of IVOC and SVOC, as gas-particle partitioning is limited by bulk diffusion. Figure A2 shows bulk diffusion and mixing timescales ($\tau_{mix}$) as a function of RH and $T$. It is interesting to note that $\tau_{mix}$ is very similar to $\tau_{eq}$ of IVOC (Fig. A1(a-c)) as gas diffusion and interfacial transport of IVOC are fast. For LVOC $\tau_{eq}$ is generally shorter than $\tau_{mix}$ as its mass transfer to the particle surface is governed by condensation sink with negligible re-evaporation, while $\tau_{mix}$ is still long to achieve homogeneous mixing in the particle phase if particles are viscous.

Previous studies have shown that $\tau_{eq}$ depends on particle size (Liu et al., 2013; Zaveri et al., 2014; Mai et al., 2015) and particle mass loadings (Shiraiwa and Seinfeld, 2012; Saleh et al., 2013). For further examination of these effects at different $T$, Figure 6 shows the dependence of $\tau_{eq}$ of SVOC ($C^0 = 10$ µg m$^{-3}$) and LVOC ($C^0 = 0.1$ µg m$^{-3}$) on the mass concentration and the diameter of pre-existing particles, over the range of 0.1 – 100 µg m$^{-3}$ and 30 – 1000 nm, respectively, with
particle phase state to be less viscous with $D_b = 10^{-11} \text{ cm}^2 \text{ s}^{-1}$ at 298 K and highly
viscous with $D_b = 10^{-18} \text{ cm}^2 \text{ s}^{-1}$ at 250 K. In this comparison, when ambient particle
mass concentration is held constant, increasing particle size will translate to a
decrease of the number and surface area concentration of particles, and a decrease of
total accommodation of molecules to the particle surface, thereby leading to an
increase of $\tau_{eq}$. When particle diameter is held constant, an increase of particle
concentration leads to an increase of surface area concentration, resulting in shorter
$\tau_{eq}$. When particles are less viscous at 298 K ($D_b = 10^{-11} \text{ cm}^2 \text{ s}^{-1}$) $\tau_{eq}$ of SVOC is
shorter than that of LVOC for the same particle size and mass loadings. For
partitioning into highly viscous particles at 250 K ($D_b = 10^{-18} \text{ cm}^2 \text{ s}^{-1}$), SVOC takes
longer time than LVOC to reach equilibrium.

Typical ambient organic mass concentrations in Beijing, Centreville in
southeastern US, Amazon Basin, and Hyytiälä, Finland are indicated in Fig. 6. The
particle phase state was observed to be mostly liquid in highly polluted episodes in
Beijing (Liu et al., 2017), under typical atmospheric conditions in the southeastern US
(Pajunoja et al., 2016), and under background conditions in Amazonia (Bateman et
al., 2017). At these conditions $\tau_{eq}$ should be mostly less than 30 minutes (Fig. 6a, b).
Particles were measured to be semi-solid or amorphous solid in clear days in Beijing
(Liu et al., 2017), in Amazonia when influenced by anthropogenic emissions
(Bateman et al., 2017), and the boreal forest in Finland (Virtanen et a., 2010). Under
these conditions and also when particles are transported to the free troposphere, $\tau_{eq}$
can be longer than 1 hour especially in remote areas with low mass loadings (Fig. 6c,
Particles in nucleation mode (diameter < 30 nm) are not considered in this study, as the particle size may affect the phase transition of these nanoparticles (Cheng et al., 2015). The role and impact of phase transition on nucleation and growth of ultrafine particles are beyond the scope of current simulations and need further investigations in future studies.

4 Discussion

The timescale to reach equilibrium for SOA partitioning has been investigated in several laboratory experiments at room temperatures (Vaden et al., 2011; Saleh et al., 2013; Liu et al., 2016; Ye et al., 2016a; Gong et al., 2018; Ye et al., 2018). These experiments monitored particle mass or composition, finding that equilibration timescales are longer at low RH, consistent with our model simulations. Note that, for condensation on highly viscous particles, even though particle mass or particle-phase concentrations appear to reach equilibrium, complete equilibrium with homogeneous mixing in the particle may not have been reached driven by strong kinetic limitations and concentration gradients in the particle bulk (Fig. 2d and 3b). This is also supported by evaporation experiments showing that the local thermodynamic equilibrium established between the vapor and the near-surface bulk should be differentiated from the global equilibrium between the vapor and the entire bulk (Liu et al., 2016). Note that SOA evaporation is also influenced by volatility and oligomer decomposition (Roldin et al., 2014; Yli-Juuti et al., 2017). The timescale of gas-particle partitioning can be different in closed or open systems especially for
LVOC (Fig. 4, S7). The closed system simulations represent SOA partitioning in chamber experiments and in closed atmospheric air mass, which could be justified well within seconds-to-minutes timescales and possibly up to hours depending on meteorological conditions. The real atmosphere may be approximated better as an open system due to dilution and chemical production and loss especially at longer timescales. Thus, particular care needs to be taken in comparing modeling results with different field conditions or experiments on probing equilibration timescale (i.e., evaporation vs. condensation, open vs. closed system, local vs. full equilibrium).

The simulated equilibration timescales of atmospheric SOA are mostly on the order of minutes to hours under conditions of atmospheric boundary layer (Fig. 5, A1). This agrees with previous experimental results that the gas-particle interactions can be regulated by both thermodynamic and kinetic partitioning (Booth et al., 2014; Liu et al., 2016; Saha and Grieshop, 2016; Ye et al., 2016a; Gong et al., 2018), depending on several factors including particle phase state, size, mass loadings, and volatility. Organic particles containing high molar-mass compounds tend to have high glass transition temperatures (Koop et al., 2011) and the occurrence of kinetic limitation will increase with higher $T_{g,org}$ (Fig. 5). This is consistent with the results of intraparticle mixing experiments showing that as the carbon number of precursor (e.g. terpene) increased (that would lead to higher $T_{g,org}$), it took longer time for SVOCs (evaporated from another type of SOA, e.g. toluene SOA) to partition into the terpene SOA, leading to slower molecular exchange among different types of SOA (Ye et al., 2018).
At low temperatures, the particles can occur as highly viscous at relatively high RH (Fig. 1), and $\tau_{eq}$ of SVOC partitioning can be longer than hours or days (Fig. 5, S7). Equilibration timescales of LVOC condensation at low particle mass loadings (Fig. 6) may represent the clean conditions where new particle formation and growth often occur (Wang et al., 2016). It has been reported that highly oxygenated molecules play an important role in the initial growth of atmospheric particles in the free troposphere (Bianchi et al., 2016). Bulk diffusion would likely to be a limiting step in the condensation of semi-volatile and low volatility compounds at low temperatures, where particles may occur as highly viscous (Shiraiwa et al., 2017). In this case, particle growth would need to be treated kinetically, rather than thermodynamic equilibrium partitioning, as it would affect SOA growth kinetics and size distribution dynamics, with significant implications for the growth of ultrafine particles to climatically relevant sizes (Riipinen et al., 2011; Riipinen et al., 2012; Shiraiwa et al., 2013a; Zaveri et al., 2018). Chemical transport models usually have time steps on the order of minutes, within which the partitioning equilibrium may not be reached, for most SVOC species ($C^0 > 1 \mu g \text{ m}^{-3}$) when $D_b$ is less than $10^{-15} \text{ cm}^2 \text{ s}^{-1}$ (Fig. 4). Note that condensation of extremely low volatility organic compounds (ELVOC; Tröstl et al., 2016) into highly viscous particles may be governed by gas-phase diffusion and timescales to reach local equilibrium could be shorter as determined by the condensation sink (Riipinen et al., 2011) (see also Fig. S4b), which may be more relevant for the practical application in chemical transport models.
In this study we assume that the bulk diffusivity within organic particles is independent of particle mixing state and morphology. Chamber experiments have demonstrated that evaporation of organic aerosol may be hindered if it is coated with organic aerosol from a different precursor (Loza et al., 2013; Boyd et al., 2017). Moreover, the phase separation has been observed in organic particles mixed with inorganic salts (You et al., 2014) and even without inorganic salts (Pöhlker et al., 2012; Riedel et al., 2016). Future simulations on equilibration timescale should consider the effects of the immiscibility (Barsanti et al., 2017; Liu et al., 2013) and the phase separation (Shiraiwa et al., 2013b; Pye et al., 2017; Fowler et al., 2018) as well as composition-dependent bulk diffusivity (O’Meara et al., 2016) and the evolution of the particle phase due to reactive uptake and condensed-phase chemistry (Hosny et al., 2016). Incorporation of the particle-phase formation of oligomers and other multifunctional high molar mass compounds can lead to a reduced bulk diffusivity (Pfrang et al., 2011; Hosny et al., 2016), which may prolong the equilibration timescales. Decomposition of highly oxidized molecules (e.g., organic hydroperoxides) in water may also affect gas-particle partitioning (Tong et al., 2016). Current simulations are focused on trace amount of SVOC or LVOC condensing on mono-dispersed particles with negligible particle growth. Potential phase transition in the course of particle growth/evaporation should also be incorporated in future simulations. The shift in particle phase state and gas-particle partitioning in response to temperature and RH may need to be considered in chemical transport models and laboratory experiments to better understand the fate of organic compounds.
Author contribution.

YL and MS designed and conducted modeling and wrote the manuscript.

Acknowledgments.

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References


Figure 1. Viscosity of pre-existing particles as a function of temperature and relative humidity. The glass transition temperatures under dry conditions ($T_{g,org}$) are (a) 240 K, (b) 270 K, and (c) 300 K, respectively.

Figure 2. Temporal evolution of mass concentrations of the condensing compound Z in the gas phase ($C_g$), just above the particle surface ($C_s$), and in the particle phase ($C_p$) in the closed system. $\tau_{eq}$ is marked with the red circle. RH = 60% and $T$ is (a–b) 298 K and (c–d) 250 K. The $C_0$ of Z is (a, c) 10 µg m$^{-3}$ and (b, d) 0.1 µg m$^{-3}$. The glass transition temperature of pre-existing particles under dry conditions ($T_{g,org}$) is set to be 270 K, which leads to $D_b$ of (a–b) 10$^{-11}$ cm$^2$ s$^{-1}$ and (c–d) 10$^{-18}$ cm$^2$ s$^{-1}$. The initial mass concentration of pre-existing particles is set to be 20 µg m$^{-3}$ with the number concentrations of $3 \times 10^4$ cm$^{-3}$ and the initial particle diameter of 100 nm.
Figure 3. Dimensionless radial concentration profiles in the particle for the condensation of the LVOC species ($C^0 = 0.1 \, \mu g \, m^{-3}$) at RH = 60% and (a) $T = 298$ K with $D_b = 10^{-11} \, cm^2 \, s^{-1}$ and (b) $T = 250$ K with $D_b = 10^{-18} \, cm^2 \, s^{-1}$. The x-axis indicates the radial distance from the particle center ($r$) normalized by the particle radius ($r_p$), ranging from the particle core ($r / r_p \approx 0$) to the surface ($r / r_p =1$). The y-axis indicates the bulk concentration of the condensing compound at a given position in the bulk ($r$) normalized by the bulk concentration at particle surface ($r_p$).
**Figure 4.** Contour plot of equilibration timescale ($\tau_{eq}$) as a function of bulk diffusivity ($D_b$) and saturation mass concentration ($C_0$) for (a) condensation in the closed system and (b) evaporation in the open system. The initial mass concentration of pre-existing particles is set to be 20 µg m$^{-3}$ with the number concentrations of $3 \times 10^4$ cm$^{-3}$ and the initial particle diameter of 100 nm. Viscosity is calculated from the Stokes-Einstein equation assuming the effective molecular radius of 10$^{-8}$ cm at $T$ of 298 K.

**Figure 5.** Equilibration timescale ($\tau_{eq}$) as a function of temperature and relative humidity in the closed system. The glass transition temperatures of pre-existing particles at dry conditions ($T_{g,org}$) are (a) 240 K, (b) 270 K, and (c) 300 K, respectively. The saturation mass concentration ($C_0$) of the condensing compound is 10 µg m$^{-3}$ (SVOC). The mass concentration of pre-existing particles is set to be 20 µg m$^{-3}$ with the number concentrations of $3 \times 10^4$ cm$^{-3}$ and the initial particle diameter of 100 nm.
Figure 6. Equilibration timescale \( \tau_{eq} \) for (a, c) SVOC \( (C^0 = 10 \, \mu g \, m^{-3}) \) and (b, d) LVOC \( (C^0 = 0.1 \, \mu g \, m^{-3}) \) as a function of particle diameter (nm) and mass concentration (\( \mu g \, m^{-3} \)) of pre-existing particles at 60% RH and \( T \) of (a-b) 298 K and (c-d) 250 K in the closed system. The glass transition temperature of pre-existing particles under dry conditions \( T_{g,org} \) is set to be 270 K, which leads to \( D_b \) of (a-b) \( 10^{-11} \, \text{cm}^2 \, \text{s}^{-1} \) and (c-d) \( 10^{-18} \, \text{cm}^2 \, \text{s}^{-1} \). Ambient organic mass concentrations are indicated with arrows.
Appendix:

Figure A1. Equilibration timescale ($\tau_{eq}$) as a function of temperature and relative humidity in the closed system. The glass transition temperatures of pre-existing particles at dry conditions ($T_{g,org}$) are set to be (a, d) 240 K, (b, e) 270 K, and (c, f) 300 K. The mass concentration of pre-existing particles is 20 $\mu$g m$^{-3}$. The saturation mass concentration ($C_0$) of the condensing compound is (a, b, c) $10^3$ $\mu$g m$^{-3}$ and (d, e, f) 0.1 $\mu$g m$^{-3}$.

Figure A2. Characteristic timescale of bulk diffusion or mixing timescale ($\tau_{mix}$) as a function of temperature and relative humidity. The particle diameter is assumed to be 100 nm with the glass transition temperatures of pre-existing particles at dry conditions ($T_{g,org}$) of (a) 240 K, (b), 270 K and (c) 300 K.
Supplement of

Timescales of Secondary Organic Aerosols to Reach Equilibrium at Various Temperatures and Relative Humidities

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Table S1. Temperature-dependent kinetic parameters used in the simulations.

<table>
<thead>
<tr>
<th>Parameter (Unit)</th>
<th>Description</th>
<th>Equation(^a)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(\omega) (cm s(^{-1}))</td>
<td>mean thermal velocity</td>
<td>(\omega(T) = (8RT/(\pi M))^{1/2})</td>
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<tr>
<td>(D_g) (cm(^2) s(^{-1}))</td>
<td>gas-phase diffusion coefficient</td>
<td>(D_g(T,P) = (T/T_{\text{standard}})^{1.75} (P_{\text{standard}}/P)^b D_g(T_{\text{standard}}, P_{\text{standard}})^b)</td>
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<td>(D_b) (cm(^2) s(^{-1}))</td>
<td>bulk diffusion coefficient</td>
<td>(D_b(T,RH) = kT/((6\pi \alpha \eta(T,RH)))</td>
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<td>first-order adsorption rate coefficient</td>
<td>(k_a(T) = \alpha \omega(T)/4)</td>
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<td>(k_d) (s(^{-1}))</td>
<td>first-order desorption rate coefficient</td>
<td>(k_d(T) = Ae^{-E_{des}/(RT)})</td>
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<tr>
<td>(k_{ss,s}) (cm s(^{-1}))</td>
<td>first-order rate coefficient for quasi-static-to-sorption layer transport</td>
<td>(k_{ss,s}(T,RH) = 2D_b(T,RH)/(\delta_{ss} + \delta Z))</td>
</tr>
<tr>
<td>(k_{ss,s}) (s(^{-1}))</td>
<td>first-order rate coefficient for sorption-to-quasi-static surface transport</td>
<td>(k_{ss,s}(T,RH) = k_{ss,s}(T,RH)k_d(T)[Z]<em>{ss,eq}/(k_a(T)[Z]</em>{g,eq}))</td>
</tr>
<tr>
<td>(k_{b1,ss}) (cm s(^{-1}))</td>
<td>rate coefficient of bulk layer 1-to-quasi-static surface transport</td>
<td>(k_{b1,ss}(T,RH) = 2D_b(T,RH)/(\delta_{ss} + \delta(1)))</td>
</tr>
<tr>
<td>(k_{ss,b1}) (cm s(^{-1}))</td>
<td>rate coefficient of surface-to-bulk layer 1 transport</td>
<td>(k_{ss,b1}(T,RH) = k_{b1,ss}(T,RH))</td>
</tr>
<tr>
<td>(k_{b,b}) (cm s(^{-1}))</td>
<td>rate coefficient of transport between bulk layers</td>
<td>(k_{b,b}(T,RH) = 2D_b(T,RH)/(\delta(k) + \delta(k + 1)))</td>
</tr>
<tr>
<td>(\tau_d) (s)</td>
<td>desorption lifetime</td>
<td>(\tau_d = k_d^{-1})</td>
</tr>
</tbody>
</table>

\(^a\) Description and the values of the symbols shown in the equations are summarized in Table S2.

\(^b\) \(D_g(T_{\text{standard}}, P_{\text{standard}})\) is calculated by the EPA on-line tools: (https://www3.epa.gov/ceampubl/learn2model/part-two/onsite/estdiffusion-ext.html).
Table S2. Description and the values of the symbols in the equations of Table S1.

<table>
<thead>
<tr>
<th>Variable (Unit)</th>
<th>Description</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$R$ (J K⁻¹ mol⁻¹)</td>
<td>gas constant</td>
<td>8.314</td>
</tr>
<tr>
<td>$T$ (K)</td>
<td>temperature</td>
<td>varied from 220 to 310</td>
</tr>
<tr>
<td>RH (%)</td>
<td>relative humidity</td>
<td>varied from 0 to 100</td>
</tr>
<tr>
<td>$M$ (g mol⁻¹)</td>
<td>molar mass of compound Z</td>
<td>200</td>
</tr>
<tr>
<td>$T_0$ (K)</td>
<td>room temperature</td>
<td>298</td>
</tr>
<tr>
<td>$\rho$ (g cm⁻³)</td>
<td>density of organic particles</td>
<td>1.4</td>
</tr>
<tr>
<td>$P$ (Pa)</td>
<td>atmospheric pressure</td>
<td>$P = P_{\text{standard}} \times (T/T_{\text{standard}})^{g/LR}$</td>
</tr>
<tr>
<td>$g$ (m s⁻²)</td>
<td>gravitational acceleration</td>
<td>9.8</td>
</tr>
<tr>
<td>$R$ (m² s⁻² K⁻¹)</td>
<td>gas constant of air</td>
<td>287</td>
</tr>
<tr>
<td>$L$ (K m⁻¹)</td>
<td>lapse rate</td>
<td>0.0065</td>
</tr>
<tr>
<td>$T_{\text{standard}}$ (K)</td>
<td>sea level standard temperature in the International Standard Atmosphere</td>
<td>288.15</td>
</tr>
<tr>
<td>$P_{\text{standard}}$ (Pa)</td>
<td>sea level standard atmospheric pressure in the International Standard Atmosphere</td>
<td>101325</td>
</tr>
<tr>
<td>$K$ (J K⁻¹)</td>
<td>Boltzmann constant</td>
<td>$1.38 \times 10^{-23}$</td>
</tr>
<tr>
<td>$a$ (cm)</td>
<td>effective molecular radius</td>
<td>$10^{-8}$</td>
</tr>
<tr>
<td>$\delta_Z$ (cm)</td>
<td>effective molecular diameter</td>
<td>$2 \times 10^{-8}$</td>
</tr>
<tr>
<td>$\alpha_{s,0}$</td>
<td>surface accommodation coefficient on free-substrate</td>
<td>1</td>
</tr>
<tr>
<td>$A$ (s⁻¹)</td>
<td>pre-exponential factor</td>
<td>$10^{12}$</td>
</tr>
<tr>
<td>$E_{\text{des}}$ (kJ mol⁻¹)</td>
<td>desorption energy</td>
<td>40</td>
</tr>
<tr>
<td>$[Z]_{g,eq}$ (cm⁻³)</td>
<td>equilibrium (saturation) number concentrations of Z in the gas phase</td>
<td>variable</td>
</tr>
<tr>
<td>$[Z]_{ss,eq}$ (cm⁻²)</td>
<td>equilibrium (saturation) number concentrations of Z in the quasi-static surface layer</td>
<td>variable</td>
</tr>
<tr>
<td>$\delta_{ss}$ (cm)</td>
<td>thickness of the quasi-static surface layer</td>
<td>variable</td>
</tr>
<tr>
<td>$\delta(k)$ (cm)</td>
<td>thickness of the bulk layer $k$</td>
<td>variable</td>
</tr>
</tbody>
</table>
Figure S1. Temperature-dependent kinetic processes simulated in the KM-GAP model. Parameters in blue are treated as a function of temperature (Table S1). [Z] are concentrations of species Z in the gas (g) and near-surface gas phases (gs), at the sorption layer (s) and in the surface (ss) and in the bulk (b) layers. J are the transport fluxes between each layer, including the gas-phase diffusion flux (J_{diff}), the adsorption (J_{ads}) and desorption (J_{des}) fluxes, surface–bulk exchange fluxes (J_{ss,ss}, J_{ss,s}, J_{ss,b1}, J_{b1,ss}), and bulk diffusion fluxes (J_{b,b}).

Figure S2. Bulk diffusion coefficient ($D_b$) in pre-existing particles as a function of temperature and relative humidity. The glass transition temperatures under dry conditions ($T_{g,org}$) are set to be (a) 240 K, (b) 270 K and (c) 300 K, respectively.
Figure S3. Temporal evolution of the mass fraction of Z in the near-surface bulk ($f_s$), and the average fraction of Z in the entire bulk ($f_b$). RH = 60% and $T$ is (a, b) 298 K and (c, d) 250 K in the closed system. The $C^0$ of Z is (a, c) 10 µg m$^{-3}$ and (b, d) 0.1 µg m$^{-3}$. The glass transition temperature of pre-existing particles under dry conditions ($T_{g,org}$) is set to be 270 K, which leads to $D_b$ of (a, b) $10^{-11}$ cm$^2$ s$^{-1}$ and (c, d) $10^{-18}$ cm$^2$ s$^{-1}$. The initial mass concentration of pre-existing particles is assumed to be 20 µg m$^{-3}$ with the number concentrations of $3 \times 10^4$ cm$^{-3}$ and the initial particle diameter of 100 nm.

Figure S4. Temporal evolution of mass concentrations of the condensing compound Z in the gas phase ($C_g$), just above the particle surface ($C_s$), and in the particle phase ($C_p$) in the closed system. The mass fraction of Z in the near-surface bulk ($f_s$), and the average fraction of Z in the entire particle bulk ($f_b$) are also shown. $D_b$ is $10^{-18}$ cm$^2$ s$^{-1}$. The $C^0$ of Z is (a) 0.1 µg m$^{-3}$ and (b) $10^{-9}$ µg m$^{-3}$. The initial mass concentration of pre-existing particles is set to be (a) 0.7 µg m$^{-3}$ and (b) 20
µg m$^{-3}$. $\tau_{eq}$ is marked with the red circle. $\tau_{eq}$ (~28 s) in (b) is consistent with the inverse of the condensation sink (29 s).

Figure S5. Temporal evolution of mass concentrations of the condensing compound Z in the gas phase ($C_g$), just above the particle surface ($C_s$), and in the particle phase ($C_p$) in the open system. $\tau_{eq}$ is marked with the red circle. RH = 60% and $T$ is (a–b) 298 K and (c–d) 250 K. The $C^0$ of Z is (a, c) 10 µg m$^{-3}$ and (b, d) 0.1 µg m$^{-3}$. The glass transition temperature of pre-existing particles under dry conditions ($T_{g,org}$) is set to be 270 K, which leads to $D_b$ of (a–b) $10^{-11}$ cm$^2$ s$^{-1}$ and (c–d) $10^{-18}$ cm$^2$ s$^{-1}$. The initial mass concentration of pre-existing particles is set to be 20 µg m$^{-3}$ with the number concentrations of $3 \times 10^4$ cm$^{-3}$ and the initial particle diameter of 100 nm.
Figure S6. Temporal evolution of mass concentrations of the evaporation compound Z in the gas phase \(C_g\), just above particle surface \(C_s\), in the particle phase \(C_p\), the mass fraction of Z in the near-surface bulk \(f_s\), and the average fraction of Z in the entire particle bulk \(f_b\) in the closed system. \(\tau_{eq}\) are marked with red circles. RH = 60% and \(T\) is (a–b) 298 K and (c–d) 250 K. The \(C^0\) of the evaporation compound is (a, c) 10 µg m\(^{-3}\) and (b, d) 0.1 µg m\(^{-3}\). The glass transition temperature of pre-existing particles under dry conditions \(T_{g,org}\) is set to be 270 K, which leads to \(D_b\) of (a, b) \(10^{-11}\) cm\(^2\) s\(^{-1}\) and (c, d) \(10^{-18}\) cm\(^2\) s\(^{-1}\). The initial mass concentration of pre-existing non-volatile particles \(C_{OA}\) is assumed to be 20 µg m\(^{-3}\) with the number concentrations of \(3 \times 10^4\) cm\(^{-3}\) and the initial particle diameter of 100 nm.
Figure S7. Equilibration timescale ($\tau_{eq}$) as a function of temperature and relative humidity in the open system. The glass transition temperatures of pre-existing particles at dry conditions ($T_{g,\text{org}}$) are (a) 240 K, (b) 270 K, and (c) 300 K, respectively. The saturation mass concentration ($C_0^\text{SVOC}$) of the condensing compound is 10 µg m$^{-3}$ (SVOC). The mass concentration of pre-existing particles is set to be 20 µg m$^{-3}$ with the number concentrations of $3 \times 10^4$ cm$^{-3}$ and the initial particle diameter of 100 nm.