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. 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 RH due to kinetic limitations of bulk diffusion in highly viscous particles. The timescale of partitioning of low-volatile compounds is shorter compared to semi-volatile compounds, as it is largely determined by condensation sink due to very slow re-evaporation. These results provide 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; 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; 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., 2016; 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 ($\tau_{eq}$) 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.

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 both $T$ and ambient pressure, which is chosen to match $T$ in 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 ($T_{g,\text{org}}$) 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 $\kappa$ 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 $T_{g,\text{org}}$ of (a) 240 K, (b) 270 K, and (c) 300 K. We chose these three $T_{g,\text{org}}$ 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). 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 $D_b$ (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.

We 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 condensing 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,

\[
\left| \frac{C_p(t) - C_{p,eq}}{C_p,0 - C_{p,eq}} \right| < \frac{1}{e}
\]

(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
Figure 2 shows exemplary simulations of temporal evolution of $C_g$ (blue line) and $C_p$ (red line) of the compound Z 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 $\mu$g 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 $\mu$g m$^{-3}$ and 0 $\mu$g m$^{-3}$, respectively. $T_{g,org}$ is assumed to be 270 K. Figure 2a presents simulations for a semi-volatile compound (SVOC, $C_0 = 10$ $\mu$g m$^{-3}$) at RH = 60% and $T = 298$ K with $D_{b}$ of $10^{-11}$ cm$^2$ s$^{-1}$ (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 $C_0 = 0.1$ $\mu$g m$^{-3}$, it takes longer time to reach the equilibrium with $\tau_{eq}$ of ~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 $C_0 = 10$ $\mu$g 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 \((C_0 = 0.1 \, \mu \text{g m}^{-3})\) condensation, \(\tau_{eq}\) is short (~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 \((C_0 = 0.1 \, \mu \text{g m}^{-3})\) in the particle at (a) \(D_b = 10^{-11} \, \text{cm}^2 \, \text{s}^{-1}\) and (b) \(10^{-18} \, \text{cm}^2 \, \text{s}^{-1}\), respectively. For low viscous particles, \(\tau_{mix}\) is very short and particles are homogeneously well-mixed at \(\tau_{eq}\). 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 also simulated evaporation in the closed system with same parameters as the condensation simulations (Table S2). Initially \( C_g = 0 \) and trace amounts of semi-volatile or low-volatile species were assumed to be homogeneously well-mixed in pre-existing particles. Figure S5 shows that for the evaporation of SVOC species with \( C_0 = 10 \mu g \text{ m}^{-3} \), 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. S5b,d). For an open system with continuous removal of gas-phase compounds, which has been 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).

We conducted further simulations to estimate equilibration timescales (\( \tau_{eq} \)) with a wide range of atmospherically-relevant temperatures (220 - 310 K) and relative humidities (0 - 100%). Figure 4 shows the temperature and humidity-dependent diagrams of \( \tau_{eq} \) for SVOC (\( C_0 = 10 \mu g \text{ m}^{-3} \)) condensation on particles with \( T_{g,org} \) of 240 K, 270 K, and 300 K, respectively. For particles with \( T_{g,org} \) of 240 K (panel a), \( \tau_{eq} \) is on the order of seconds under boundary layer conditions (\( T > 270 \text{ K} \)). 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 RH.

$\tau_{eq}$ for $C_0 = 10^3$ and 0.1 µg m$^{-3}$ are presented in Fig. A1. In general, $\tau_{eq}$ would be shorter at higher $C_0$, 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. 4a, 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 shorter than $\tau_{mix}$ as its mass transfer to the particle surface is governed by condensation sink with negligible
re-evaporation, while $\tau_{\text{mix}}$ is still long to achieve homogeneous mixing in the particle phase if particles are viscous.

Figure 5 shows the dependence of $\tau_{\text{eq}}$ of LVOC ($C_0 = 0.1 \mu g m^{-3}$) on the mass concentration and the diameter of pre-existing particles, over the range of $0.1 - 100 \mu g m^{-3}$ and $30 - 1000$ nm, respectively, with particle phase state to be less viscous with $D_b = 10^{-11} cm^2 s^{-1}$ at 298 K and highly viscous with $D_b = 10^{-18} cm^2 s^{-1}$ 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 $\tau_{\text{eq}}$. When particle diameter is held constant, an increase of particle concentration leads to an increase of surface area concentration, resulting in shorter $\tau_{\text{eq}}$. Transition of the particle phase state from less viscous at 298 K ($D_b = 10^{-11} cm^2 s^{-1}$) to highly viscous at 250 K ($D_b = 10^{-18} cm^2 s^{-1}$) leads to longer $\tau_{\text{eq}}$. 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; 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). Thus, particular care needs to be taken in comparing modeling results with different 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. 4, 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; 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. 4). 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 $t_{eq}$ can be longer than hours or days (Fig. 4, A1). Equilibration timescales of LVOC condensation at low particle mass loadings (Fig. 5) 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). Note that condensation of extremely low volatility organic
Compounds (ELVOC; Tröstl et al., 2016) 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) and the phase separation (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). Particle size dependent phase transition (Cheng et al., 2015) and phase change in the course of particle growth (Shiraiwa et al., 2013b) may also need to be considered. 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.

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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 ($T_{g,\text{org}}$) 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 ($C_p$). $\tau_{eq}$ 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 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 \(D_b = 10^{-19} cm^2 s^{-1}\). 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. Equilibration timescale \((\tau_{eq})\) as a function of temperature and relative humidity. 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.
Figure 5. Equilibration timescale (τ_{eq}) for 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) 298 K and (b) 250 K. 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) 10^{-11} \, cm^2 \, s^{-1} and (b) 10^{-18} \, cm^2 \, s^{-1}.
Appendix:

Figure A1. Equilibration timescale ($\tau_{eq}$) as a function of temperature and relative humidity. The glass transition temperatures of pre-existing particles at dry conditions ($T_{g,\text{org}}$) 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 µg m$^{-3}$. The saturation mass concentration ($C_0$) of the condensing compound is (a, b, c) $10^3$ µg m$^{-3}$ and (d, e, f) 0.1 µg m$^{-3}$.

Figure A2. Mixing timescale ($\tau_{\text{mix}}$) 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 ($T_{g,\text{org}}$) of (a) 240 K, (b) 270 K and (c) 300 K.