Monte Carlo simulations of homogeneous nucleation and particle growth in the presence of background particles

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ABSTRACT

The application of the Monte Carlo (MC) simulation technique for the modelling of nucleation processes with an existing background particle concentration is presented in this paper. Next to the nucleation of novel particles, the coagulation of an existing particle population as well as the condensational growth and evaporation of unstable particles (whose diameter is smaller than the critical Kelvin diameter) are included into the simulation. The usage of statistically weighted MC particles allows the description of particle size distribution (PSD), whose concentrations differ in several orders of magnitude. It is shown, that this approach allows to model the complex interplay between freshly nucleated particles and an existing background particle population. In this work, the nucleation of novel particles is modelled by three different nucleation theories discussed by Girshick, S. L. and C.-P. Chiu (1990), The Journal of Chemical Physics 93, which comprise of (1) the classical nucleation theory, (2) a mathematical correction to (1) and (3) a self-consistency correction of (2). For the chosen simulation conditions, the resulting PSDs are independent of the used nucleation theories discussed by Girshick, S. L. and C.-P. Chiu (1990), The Journal of Chemical Physics 93, which comprise of (1) the classical nucleation theory, (2) a mathematical correction to (1) and (3) a self-consistency correction of (2). For the chosen simulation conditions, the resulting PSDs are independent of the used nucleation theories discussed by Girshick, S. L. and C.-P. Chiu (1990), The Journal of Chemical Physics 93, which comprise of (1) the classical nucleation theory, (2) a mathematical correction to (1) and (3) a self-consistency correction of (2). For the chosen simulation conditions, the resulting PSDs are independent of the used nucleation theories discussed by Girshick, S. L. and C.-P. 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computational resources and times), the Monte Carlo (MC) method can be applied; hence, the increment of the necessary computational resources depends only linearly on the number of used components.

It has already been shown that MC methods are well suited to describe multicomponent coagulation of aerosols (Efendiev and Zachariah, 2003; Zhao and Kruis, 2014). The inclusion of the homogeneous nucleation into a MC simulation is much more challenging, due to the fact, that continuous nucleation leads to the increase of the number of MC simulation particles. It was already shown, that the application of weighted MC particles makes this problem accessible. The designation of different statistical weights to each simulation particle makes the development of the concept of ‘particle merging’ possible (Kotalczyk and Kruis, 2017). We have also already shown, that this method can be applied to nucleation and growth in aerosol reactors (Kotalczyk et al., 2016, 2017). In atmospheric systems, however, a population of earlier formed background particles has to be taken into account (e.g. Olenius and Riipinen, 2017). The simultaneous description of (1) the background particle population and (2) the freshly nucleating particles poses a numerical challenge; hence, the number-concentrations of those two populations can differ several orders of magnitude. If the number-concentration of the background particles is $10^{10}$ m$^{-3}$ and the freshly nucleated particle are described by a large number-concentration, like $10^{14}$ m$^{-3}$, the application of non-weighted MC particles becomes nearly impossible: a huge number of 1,000,000 simulation particles would be needed, if the background particle population is to be modelled by a modest number of 100 particles. We show in the following, that such systems can be simulated by means of weighted MC simulation particles, but we refrain the simulated system to one component, only.

The theoretical description of the nucleation poses a difficult task, as described by the reviews (Oxtoby, 1992; Ford, 2004; Bennett and Barrett, 2012; Zhang et al., 2012). The great difficulty in the investigation of the nucleation phenomena lies in the fact, that several reasonable theories can be formulated, leading to nucleation rates, which differ in several orders of magnitude. Most theories resort to cluster–cluster interactions of unstable molecules which grow to a critical size and become stable, assumptions about interaction potentials (in case of molecular dynamics simulations, e.g. Tanaka et al., 2014) or about the kinetic collision rates (in case of kinetic formulations, e.g. Oxtoby, 1992; Laaksonen et al., 1995) have to be made. These assumptions cannot be verified by experiments (yet), because an experimental technique, which measures the concentration of the smallest (unstable) clusters would be necessary for this purpose (Wyslouzil and Wölk, 2016).

In this work, we focus on one component nucleation as a result of rapidly changing atmospheric conditions, e.g. a rapid temperature drop leading to a strong increase of the supersaturation. This is numerically a more challenging system than the otherwise applied supersaturation profiles used in atmospheric modelling, which describe a slow increase of the supersaturation at lower supersaturation levels. This system is tested by comparing three well-known expressions for the nucleation rate with each other. The decrease of the supersaturation, the total number concentration and the particle size distributions (PSD) are compared for this purpose. We show, that early stages of the simulation lead to strongly different results, while for longer simulation times the results of different nucleation models become undistinguishable from each other.

2. Methods

In the first subsection, a general parallel simulation algorithm for the solution of the PBE by means of weighted simulation particles is sketched. In the following, second section, the simulated system, consisting of the initial conditions and the used nucleation theories, is discussed in detail.

2.1. MC simulation algorithm

The use of statistically weighted MC simulation particles (Zhao et al., 2009; Lee et al., 2015) allows new modelling approaches, hence no limitations have to be set on the statistical weight of the newly included simulation particles (Menz et al., 2012; Hao et al., 2013). We combine a stochastic coagulation process of weighted MC particles with a deterministically described growth and evaporation process, which can be combined with the nucleation of novel particles, too.

The discrete coagulation events are separated from the continuous growth processes in a simple operator splitting approach (also used to describe the chemical decomposition of a precursor concentration during particle synthesis) (Celnik et al., 2007), so that the simulation of a time step consists of two steps: the MC-driven coagulation step and a growth (resp. evaporation) step which is solved by the solution of the corresponding ODEs. During the nucleation time step many particles could be inserted into the simulation as described further below. Treating the condensation as a continuous process instead of formulating single events for events describing monomer attachments or detachments leads to remarkable speed-ups of the simulations (Patterson et al., 2006). These approximations are valid for the description of larger particle sizes (i.e. >1 nm), discretization errors
might be found for lower particle sizes and a different approach might become necessary, if the here presented approach is coupled to a kinetic MC simulation describing the nucleation (like e.g. in Filipponi and Giammatteo, 2016 or Davari and Mukherjee, 2018).

2.1.1. MC coagulation step. First, the coagulation time step $\tau_{mc}$ is estimated from the given PSD, which is defined by the reciprocal of the sum of all possible event rates: $\tau_{mc} = 1/\sum_{i<j} (\beta_{ij}^{SR})$, where $\beta_{ij}^{SR} = \max(W_i, W_j) \cdot \beta(i, j)$, is the coagulation rate within the concept of ‘stochastic resolution’, according to (Kotalczyk and Kruis, 2017), where $\beta(i, j)$ is the coagulation rate of particles $i$ and $j$. The coagulation rate $\beta$ describes the rate of coagulation between two real particles.

The particles are selected for coagulation by selecting the particle indices $i$ and $j$ by random and selecting this random pair for coagulation, if an additional random number $r$ is smaller than $\beta_{ij}^{SR}/\beta_{ij}^{max}$. The maximal coagulation rate $\beta_{ij}^{max}$ can thereby be quickly approximated by a fast parallel algorithm introduced by (Wei and Kruis, 2013). The parallel implementation of this algorithm can be found in (Wei and Kruis, 2013) or (Kotalczyk and Kruis, 2017). We point out, that the applied coagulation time step $\tau_{mc} = 1/\sum_{i<j} \max(W_i, W_j) \cdot \beta(i, j)$ is also dependent on the number of used simulation particles, and that higher numbers of simulation particles lead to lower values for the applied time step (the values of $\max(W_i, W_j)$ decrease linearly with the increase of the simulation particles, while the number of summands increases quadratically). So that the tuning of shorter separation times of the coagulation and the other processes becomes possible.

2.1.2. Continuous growth step. Second, in order to describe the condensational growth (or evaporation) of the particles, the corresponding growth equation is solved for each particle:

$$\frac{dv_j}{dt} = G(v_j, nG)$$  \hspace{1cm} (1)

The monomer concentration in the gaseous phase $nG$ determines thereby the Kelvin diameter $d^*$. It is thus determined whether the particle grows ($G>0$, if $d>d^*$) or evaporates ($G<0$, if $d<d^*$) and the exact rate of this process. The depletion (resp. increase) of the monomers is taken into account by the following mass balance (i.e. the first term on the r.h.s. of Equation (2)):

$$\frac{dnG}{dt} = - \sum_i W_i \cdot G(v_i, nG)/v_M - N_R \cdot \bar{v}$$  \hspace{1cm} (2)

The second term on the r.h.s. of Equation (2) describes the monomer depletion of the monomer concentration due to the nucleation of novel particles; each nucleated particle consists thereby of $i^*$ monomers (where $i^* = \pi \cdot d^3/(6 \cdot v_M)$).

The coupled set of ordinary differential equations, consisting of Equation (2) and as many equations of the form of Equation (1), as there are MC particles (in the here presented simulations, 10,000) is solved by the Runge-Kutta technique for the time step $\tau_{mc}$, which is set by the coagulation process discussed above.

The total mass of the nucleated material is stored as a special buffer variable $M_B$, so that the set above is extended by one simple equation:

$$\frac{dM_B}{dt} = +N_R \cdot \bar{v} \cdot v_M$$  \hspace{1cm} (3)

If the value $M_B$ surpasses a pre-set limit $M_{TR}$ (in the simulated scenario presented in this paper: $M_{TR} = 10^{-20}$kgm$^{-3}$), a novel MC particle with the statistical weight of $W_{nuc} = M_B/(v_M \cdot \bar{v})$ is inserted into the simulation by the procedure described in the following subsection. It should be noted, that many nucleation events are possible during one coagulation step, dependent on the set threshold $M_{TR}$ and on the number of performed Runge Kutta (RK) time steps. These steps are set by comparison of the RK result of the 4th order technique to the 5th order technique. The difference between both techniques is compared to a predefined acceptable error in order to adjust the next time step or repeat the actual RK step for a much smaller interval of time (like described in Press et al., 2007).

2.1.3. Inclusion of newly nucleated particles: Low weight merging. Hence, the continuous inclusion of novel particles would lead to large numbers of simulation particles and thus higher computational times, the ‘low weight merging’ of particles is applied. This technique selects two ‘nearly similar’ MC particles $(i, j)$ and it stores their properties as one particle in the memory space of the former particle $j$, so that the newly nucleated particle can be stored in the memory space of the former particle $i$. The statistical weight $W_{new}$ of the novel particle and its new volume $v_{new}$ are calculated by:

$$W_{new} = W_i + W_j \quad \text{and} \quad v_{new} = (W_i \cdot v_i + W_j \cdot v_j)/W_{new}$$  \hspace{1cm} (4)

This applied scheme preserves the total mass of the involved particles exactly; hence, $v_{new} \cdot W_{new} = (W_i \cdot v_i + W_j \cdot v_j)$. As a measure for the magnitude of the introduced error, the value $E_{mer}$ is introduced:

$$E_{mer} = (v_i - v_j)^2/\min(v_i, v_j)$$  \hspace{1cm} (5)

An algorithm is applied, which merges particles with low statistical weights (in order to introduce the lowest alternations to the PSD). The particles with low statistical
weights are compared to all other particles, and the pair with the lowest merging error, $E_{\text{mer}}$ is merged. The exact computational procedure for the selection of two ‘nearly similar’ particles is presented in (Kotalczyk and Kruis, 2017) in more detail.

### 2.2. Simulated system

The system consists of a background particle population and a condensable monomer concentration $n_G$ (describing thus the supersaturation $S$ of the system). It is assumed that the background particles and the gaseous monomers are one compound, with the properties summarized in Table 1. These values could designate a quasi-urban atmospheric aerosol consisting of sulphuric acid and organic compounds in a scenario describing sulphuric acid induced nucleation of the organic molecules (Olenius and RiiPinen, 2017). The system is simulated for isothermal conditions (280 K) and 24 hr.

The background particle population is modelled as a lognormal distribution with a geometric mean of 50 nm and a geometric standard deviation of 1.64, which corresponds to the geometric standard deviation of the self-preserving PSD for the coagulation process (Vemury et al., 1994). These initial particles encompass a total number concentration of $N_0 = 10^{10}$ m$^{-3}$, this might describe a polluted environment.

The initial supersaturation is set to the value of $S_0 = 10^5$ at the beginning, which also corresponds to a polluted environment—but is also tested for lower values ($S_0 = 10^4, 10^3$ and $10^2$). This modelling does not take into account a slowly increase and decrease of the monomer concentration, which is sometimes modelled in a sinusoidal way reflecting the diurnal cycle (Olenius and RiiPinen, 2017). It is assumed that these conditions are reached (due to drastic cooling or convection) at the beginning of the simulation and that the condensational growth or evaporation of the existing particles as well as the nucleation are the only reasons for changes of the supersaturation. This assumption is a rather artificial modelling of the complex interplay of an existing background particle population with condensable material in the form monomers in the gaseous phase, whose saturation is slowly increased by cooling or monomer convection. It poses, however, a much more challenging system for the here discussed MC methodology, hence a slow increase would lead to less nucleation and thus less new particles which have to be included into the simulation.

The kernel for coagulation in the free molecule regime is presented in (Kotalczyk and Kruis, 2017), and in this work, or as $N_0^{(\text{cou})}$ in Equation (10). A correction to this expression in the framework of a kinetic formulation has been proposed by (Courtney, 1961) and its mathematical necessity is stressed by many works (e.g. Girshick and Chiu, 1990; Oxtoby, 1992), it will be marked as $N_0^{(\text{cou})}$ in Equation (9). A self-consistent correction to this kinetic nucleation theory (called ‘Courtney’ in the following) has been proposed by (Girshick and Chiu, 1990) and is denoted as $N_0^{(\text{gir})}$ in Equation (10).

The used symbols are summarized in Table 1 and $k_B$ is the Boltzmann’s constant.

The condensational growth is described by the formula for the free-molecule regime as well and depends on the volume $v_i$ (resp. the diameter $d_i$, with $v_i = \pi d_i^3 / 6$) of the particle and the monomer concentration $n_G$ in the gaseous phase:

$$ G(v_i, n_G) = \frac{d v_i}{d t} = \frac{v_M \cdot \pi \cdot d_i^2}{\sqrt{2 \pi \cdot m_1 \cdot k_B \cdot T}} \cdot \exp \left\{ 4 \cdot \frac{\gamma \cdot v_M}{(k_B \cdot T \cdot d_i)} \right\} $$ (7)

The used symbols and its values are described in Table 1. It is assumed, that the nucleated particles and the background particle population consist of the same material and that all simulated particles grow (resp. evaporate) like described by Equation (7). The exponential term describes thereby the Kelvin correction, this leads to an evaporation of all particles whose diameter $d_i$ is smaller than the Kelvin diameter $d^*$, with:

$$ d^* = 4 \cdot \frac{\gamma \cdot v_M}{(k_B \cdot T \cdot \ln(S))} $$ (8)

This diameter is also the diameter of the inserted particles due to nucleation.

Three different expressions are investigated for the nucleation rate $N_R$. First the classic nucleation theory by (Becker and Döring, 1935) has been investigated, denoting it as ‘classic’ in this work, or as $N_0^{(\text{cou})}$ in Equation (10). A correction to this expression in the framework of a kinetic formulation has been proposed by (Courtney, 1961) and its mathematical necessity is stressed by many works (e.g. Girshick and Chiu, 1990; Oxtoby, 1992), it will be marked as $N_0^{(\text{cou})}$ in Equation (9). A self-consistent correction to this kinetic nucleation theory (called ‘Courtney’ in the following) has been proposed by (Girshick and Chiu, 1990) and is denoted as $N_0^{(\text{gir})}$ in Equation (10).

$$ N_R^{(\text{gir})} (n_G) = N_G \cdot \sqrt{\frac{2 \gamma}{\pi \cdot m_1 \cdot k_B \cdot T}} \cdot \frac{P_s}{v_i} \cdot \exp \left\{ \frac{16 \cdot \pi \cdot \gamma^3 \cdot v_M^2}{3 \cdot k_B^2 \cdot T^3 \cdot \ln(S)} \right\} \quad \text{with} \quad S = \frac{n_0 \cdot k_B \cdot T}{P_s} $$ (9)

$$ N_R^{(\text{cou})} (n_G) = N_R^{(\text{gir})} \cdot \exp \left\{ \frac{36 \cdot \pi \cdot \gamma^3 \cdot v_M^2}{k_B^2 \cdot T} \right\}; \quad N_R^{(\text{chi})} (n_G) = N_R^{(\text{cou})} \cdot S $$ (10)

### 3. Results

The compared nucleation theories influence directly the characteristics of the simulated particle population. Due to different nucleation rates, a different depletion of the
monomer concentration takes place, as can be seen in Fig. 1. These variations of the saturation in time affect directly the nucleation rates, shown in Fig. 2. One can coarsely distinct the simulation time into three regions: (1) an initial time span (simulation times smaller than 1 min), for which nearly constant nucleation rates can be observed, which correspond to constant monomer concentration values—meaning that the depletion of the monomer concentration due to condensational growth and nucleation can be neglected during this time-span; (2) a transition time span (simulation times between 1 min and 1 hr), in which a depletion of the monomer concentration can be seen (caused by the condensational growth on the existing particle population and the nucleation of novel particles); (3) a steady state region (simulation times larger than 1 hr), which is described by the continuous growth/evaporation and coagulation and marked by a slow decrease of the surplus supersaturation. In the here presented simulation conditions, the contributions of the growth/evaporation term are nearly negligible for longer simulation times and the supersaturation seems to reach a steady state. Therefore, one can consider the coagulation to be the only one driving force for particulate growth—in contrast to the system describing industrial Fe production conditions (Kotalezyk et al., 2016), for which the complex interplay of coagulation and growth/evaporation has to be taken into account for the correct description of the PSD (Kotalezyk et al., 2017).

The different nucleation rates have, of course a direct impact on the overall number concentration of the particles, as can be seen in Fig. 3. A steep increase in the particle concentration is found in the initial stage of the simulation for the classic and Girshick nucleation theory. The Courtney theory is not describing such high nucleation rates as the other two theories (as can be seen in Fig. 2), which leads to no drastic increase of the particle concentration. The transition time span is marked by a decline of the particle concentration, which can be explained by the onset of the coagulation of the particles.

These findings can also be drawn from the PSDs shown in Figs. 4–6 describing the particle populations after a total simulation time of 1 min, 1 hr and 24 hr. The 10,000 particles are sorted into 80 logarithmically spaced bins and mean values for 100 simulations are shown. Figure 4 shows that after 1 min the differences of the particle concentrations presented in Fig. 3 can be attributed
to the nucleated particles, which represent the smaller sizes of the particle size spectrum. The PSD based on the Girshick nucleation theory exhibits not only higher concentrations of the nucleated particles as the other nucleation theories, but the sizes of the particles are larger as well. This indicates that the newly nucleated particles have coagulated with each other. This explains the number concentration plateau for the initial stage of the simulation shown in Fig. 3 for the Girshick nucleation theory: the increase of the particle number concentration is limited by the onset of the coagulation, which decreases the particle number concentration.

The part of the PSD representing mainly the background particle population changes its shape slightly. It can be seen that the parts representing smaller concentrations of the PSD are not rendered in the results based on the Girshick and the Classic theory. This can be attributed to the ‘low weight merging’, necessary for the novel inclusion of a nucleated particle. The higher the nucleation rate and the more new particles have to be included into the simulation, the less accurate becomes the rendering of the fringes of the background particle population. It should be noted, that the regions, in which these inaccuracies are observed constitute a small part of the PSD (note the logarithmic plot). This small part (especially the larger particles), could describe the part of the PSD, which is responsible for the light-scattering behaviour of the aerosol or act as cloud condensation nuclei (CCN).

The ‘removal’ of these particles due to the merging technique might therefore introduce serious errors in the context of the simulation of an atmospheric system. These errors could be prevented by the following 2 approaches: (1) the particles which are large enough to act as CCNs could be locked against merging, simply by not comparing and merging them with other particles; (2) the light-scattering cross-section (resp. power) $\sigma_i$ of each particle $i$
could be considered for the calculation of the merging error, in this approach, the expression in Equation (5) could be replaced by:
\[ E_{\text{mer}} = \left( \frac{v_i}{C_0} - \frac{v_j}{C_0} \right)^2 / \min(v_i, v_j) + (\sigma_i - \sigma_j)^2 / \sigma_{\text{gauche}} \]
where \( \sigma_{\text{gauche}} \) is some pre-set light-scattering cross-section. This second approach would penalize the merging of particles with different light-scattering cross-sections, if one of them is higher than \( \sigma_{\text{gauche}} \).

Although the fringes of the PSDs change due to the here applied merging techniques, the total volume \( V_{\text{tot}} \) is conserved in the course of the simulation, where:
\[
V_{\text{tot}} = \sum_i W_i \cdot v_i + n_G \cdot v_m \tag{11}
\]

The mean value for \( V_{\text{tot}} \) (for all 100 simulations) is compared with the initial value \( V_0 \) at the beginning of the simulation in Fig. 7. It can be seen that it changes only slightly. The relative deviation of \( 10^{-7} \) corresponds to the magnitude of the floating point precision applied for the calculations. This value accumulates to levels of \( 10^{-6} \) for simulations of high nucleation scenarios. These levels are still acceptable.

We also observed, that the application of a higher number of simulation particles leads to less deviations within the background PSDs, so that the number of simulation particles can be applied as a control parameter for these deviations. The same values for the moments of the PSDs (such as mean geometric diameter or total number-concentration) could be reproduced, independent on the number of the applied simulation particles.

The PSD based on the Courtney theory expresses the background condition with the same accuracy as the initial condition, hence little to none particles have to be inserted into the simulation in this scenario. In turn, much higher differences between the concentrations of the nucleated particles and the concentrations of background particles are described by the PSD based on the Girshick theory as the one based on the Courtney theory.

In the time span, between 1 min and 1 hr (the transition time span), the nucleation of novel particles ceases completely (see Fig. 2). The evolution of the PSD in this time span is mainly described by the coagulation of the particles, the condensational growth of the particle population might be considered to be very small, hence the corresponding driving force \( S - S_\infty \) decreases several orders of magnitude, as can be seen in Fig. 1. The result is a PSD, which consists of two peaks, which can be attributed to the nucleated population and the initial background particle population. Figure 5 shows that the growth of the background particle population due to condensation and due to coagulation with the nucleated particles is relatively small. The part of the PSD, which represents the background particles (around 50 nm), is nearly of the same shape as the initial condition (disregarding the shape changes due to the merging on the edges of this distribution).

The part of the PSD, which can be attributed to the nucleated particles (covering the whole range between 1 and 20 nm) exhibits the form of the self-preserving form for coagulation. The differences between the PSDs are clearly visible, making a distinction between each other.
possible. The attribution of the corresponding nucleation theory based on the size of the PSD in this size spectrum would therefore be possible, while the form of the background particles (all particles with the size 20 nm or higher) would not allow to draw conclusions on the nucleation mechanisms.

These distinctions vanish, however, in the further course of the simulation and the resulting PSDs attain a self-preserving form for the coagulation, making no more distinctions between each other possible, as can be seen in Fig. 6. The depicted results describe the PSDs after 24 hr. A clear difference between the PSDs for 24 hr and the initial PSD can be seen, which can be attributed to the coagulation within this long ‘steady state time span’.

Each PSD is rendered with the same accuracy. This indicates that the differences of the rendering of the background particle population in Figs. 4 and 5, which originate from the ‘low weight merging’ technique did not propagate into the simulation and were in fact not critical for the correct description of the general behaviour of the PSD. However, the negligence of the larger particles might lead to a wrong description of the light scattering behaviour of the aerosol—or its ability to act as CCNs—both points could be addressed by modifying the merging technique as discussed above.

Similar findings can be made for different values for the initial concentration $S_0$, exemplary plots of the supersaturation in Fig. 8 and the number concentration in Fig. 9 show, that the nucleation takes place mostly during the first minute of the simulation and that the depletion of the supersaturation takes place between the first minute and hour of the simulation. This time frame may also be identified as the one, which allows to pose assumptions on the nucleation mechanisms from the shapes of the PSDs.

Higher nucleation rates can be—obviously—observed for higher initial supersaturations $S_0$ as can be seen in Fig. 8, for the lowest supersaturation, $S_0 = 100$, no nucleation can be observed at all and the total monomer concentration is depleted due to condensational growth of the background particles. This means, that high supersaturations are necessary for the nucleation of novel particles in the presence of a background particle population.

The decrease of the supersaturation is much faster for $S_0 = 10^5$ than for $S_0 = 10^4$ or $S_0 = 10^3$. This can be attributed to the much higher concentration of nucleated particles of this system, which in turn act as a source for monomer depletion through condensational growth—an effect, which would not become apparent, if the growth (resp. evaporation) of the particle population would not be included into the simulation.

4. Conclusions

We investigate the homogeneous nucleation of an aerosol under isothermal conditions and in the presence of a background particle population. Three different nucleation theories are simulated. We show that each of these
Theories results in different rates of nucleation and thus different particle concentrations at the beginning of the simulation. For longer simulation times (>1 hr), the resulting PSD become undistinguishable from each other, as can be expected from different systems with the same total particulate mass, which are driven by the coagulation only. In this way, the identification of a time frame relevant for measurements can be established.

The applied MC simulation algorithm makes use of weighted simulation particles. It is shown, that this simulation technique is able to describe PSDs, which are characterized by huge differences in the number-concentrations of the freshly nucleated particles and the already existing background particles. This shows that the presented MC technique could also be used for the description of a multicomponent atmospheric aerosol system. An investigation of the influence of more complex, kinetic MC simulation based nucleation theories (like Davari and Mukherjee, 2018) could also be simulated in a future work.

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