

Soot aging time scales in polluted regions during day and night

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Abstract. The aging of soot is one of the key uncertainties in the estimation of both the direct and indirect climate effect. While freshly emitted soot is initially hydrophobic and externally mixed, it can be transferred into an internal mixture by coagulation, condensation or photochemical processes. These aging processes affect the hygroscopic qualities and hence the growth behaviour, the optical properties and eventually the lifetime of the soot particles. However, due to computational limits the aging of soot in global climate models is often only parameterised by an estimated turnover rate resulting in a lifetime of soot of several days. Hence, the aging process of soot is one of the key uncertainties governing the burden and effect of black carbon. In this study, we discuss the time scale on which diesel soot is transferred from an external to an internal mixture based on the results of our simulations with a comprehensive mesoscale model. For daytime conditions during summer condensation of sulphuric acid is dominant and the aging process occurs on a time scale of $\tau = 8$ h close to the sources and $\tau = 2$ h above the source region. During winter comparable time scales are found but ammonium nitrate becomes more important. During night time condensation is not effective. Then coagulation is the most important aging process and our results show time scales between 10 h and 40 h.

1 Introduction

Soot particles are an important constituent of the atmospheric aerosol, since they participate in tropospheric chemistry (Saathoff et al., 2001), affect human pulmonary health (Pope and Dockery, 1996) and scatter and absorb light (Horvath, 1993). The size distribution of soot particles peaks in the accumulation range, therefore dry deposition veloc-

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ities are small and soot particles can attain long lifetimes and be transported over long distances. The source of soot particles is the incomplete combustion of carbon containing material, which means that except for natural biomass burning all sources of soot are anthropogenic. In cities of the northern hemisphere the combustion of fossil fuel dominates the sources. Typical concentrations of soot range between several hundreds of nanograms per m³ in unpolluted areas to several micrograms per m³ in urban regions (Heintzenberg, 1988; Brémond, 1989; Baltensperger et al., 2002). While freshly emitted soot particles are hydrophobic and present in an external mixture their hygroscopic qualities can change due to the coagulation with soluble aerosols, condensation, and photochemical processes (Weingartner et al., 1997). Hereby the particle growth in response to ambient relative humidity, the optical properties, and the ability of being activated as cloud condensation nuclei are determined. Measurements show that both the external and the internal mixing state exist in the atmosphere (Okada, 2001) and that the hydrophobic portion of the aerosols decreases significantly as the distance from the sources increases.

It is well recognized that soot particles contribute to both the direct and indirect climate effect. While Lesins et al. (2002) and Jacobson (2000) recently studied the direct effect; Nenes et al. (2002) highlighted the indirect effect. A number of studies in the past have been devoted to the representation of soot in global scale models (Liousse, 1996; Cooke et al., 1996, 1999; Lohmann et al., 2000) and the impact of soot aerosol on global climate (Jacobson, 2002). Due to computational limits the aging of soot in global scale models is often not represented explicitly but parameterised. For example, Liousse (1996) assume that soot is hydrophilic as soon as it is emitted. Myhre et al. (1998), in contrast, hypothesize soot to be hydrophobic. Tsigaridis and Kanakidou (2003) assume that the aging process occurs due to photochemical reactions. Cooke et al. (1996) parameterise the aging process by an estimated turnover rate of 1.25% h⁻¹,

which translates to an exponential lifetime of 80 h. Lohmann et al. (2000) assume an exponential lifetime of 40 h emphasizing however, that the model results are highly sensitive to the turnover rate. Koch (2001) comes to a similar conclusion comparing three different methods for the parameterisation of the aging process.

Recently, Wilson et al. (2001) have provided a more detailed representation of the aging process in a global model. In their approach they explicitly calculate the aging of soot depending on the abundance of sulphuric acid. Nevertheless, parameterisations of certain processes are still necessary when such global models are run in the climate mode, that is when several hundreds of years are simulated.

Since the aging process is one of the key uncertainties governing the burden and effect of black carbon, we find it worthwhile to carry out 3D simulations with the coupled mesoscale- γ model KAMM/DRAIS (Vogel et al., 1995) that provides a highly resolved boundary layer and allows for an explicit treatment of the aging process of soot by coagulation and condensation to investigate the variability of the aging time scale of soot in more detail. The particle phase is treated with the aerosol model MADEsoot (Riemer et al., 2003b) and calculates both the composition and the size distribution of the aerosol particles. Based on the results of our simulations, we discuss the time scale on which soot is transferred from an external to an internal mixture, and calculate the turnover rates for a domain size that compares to the grid size of a global climate model. In doing so, we focus on continental conditions in an industrialised environment and investigate two different meteorological scenarios, that is a summer and a winter episode.

2 Model description

The comprehensive model system KAMM/DRAIS couples the non-hydrostatic meteorological driver KAMM and the submodule DRAIS that calculates the transport and diffusion of the reactive trace gases and the aerosol particles. Since the model system KAMM/DRAIS has been described in detail in previous papers (Vogel et al., 1995; Hammer et al., 2002) and has been extensively validated against observations in the past (Vogel et al., 1995; Nester et al., 1995; Fiedler et al., 2000; Corsmeier et al., 2002; Hammer et al., 2002) we only give a short summary with the focus on the aerosol model MADEsoot (Riemer et al., 2003b).

In MADEsoot, several overlapping modes represent the aerosol population, which are approximated by log-normal functions. Currently, we use five modes for the sub-micron particles. Two modes $(i_f \text{ and } j_f)$ represent secondary inorganic particles consisting of sulphate, ammonium, nitrate, secondary organic compounds, and water, one mode (s) represents pure soot and two more modes $(i_c \text{ and } j_c)$ represent particles consisting of sulphate, ammonium, nitrate, organic compounds, water and soot. The modes i_f , j_f , i_c and j_c are

assumed to be internally mixed. Thus, the modes i_f and j_f represent soot-free particles whereas the modes i_c and j_c represent soot containing particles or, in other words, the aged soot particles. All modes are subject to condensation and coagulation. The growth rate of the particles due to condensation is calculated following Binkowski and Shankar (1995) depending on the available mass of the condensable species and the size distribution of particles. With coagulation, the assignment to the individual modes follows Whitby et al. (1991): (1) Particles formed by intramodal coagulation stay in their original modes. (2) Particles formed by intermodal coagulation are assigned to the mode with the larger median diameter. Furthermore, a thermodynamic equilibrium of gas phase and aerosol phase is applied to calculate the concentrations of sulphate, ammonium, nitrate and water (Kim et al. 1993). The calculation of the thermodynamic equilibrium follows a bulk approach. This means that the aerosol concentrations summed over all modes enter the calculation. After the equilibrium concentrations are obtained the concentrations of ammonium, nitrate and water are distributed over the modes depending on the mass fraction of sulphate. The source of the secondary inorganic particles in modes i_f and j_f is the binary nucleation of sulphuric acid and water. The secondary organic compounds are treated according to Schell et al. (2001). The soot particles in mode s are directly emitted into the atmosphere. The source of the particles in modes i_c and j_c is due to the aging process described below. Additionally, sedimentation, advection and turbulent diffusion can modify the aerosol distributions.

In this framework, two processes can impact the transfer of soot from the external into the internal mixture, namely coagulation and condensation. Coagulation of soot particles in mode s with particles in modes i_f , j_f , i_c or j_c transfers the mass of mode s into the modes i_c or j_c . As a second process, condensation of sulphuric acid on the surface of the soot particles and the subsequent formation of ammonium nitrate can transfer soot into an internal mixture as well. To retain the soot mode for pure soot particles, a criteria must be chosen that determines when the aged soot together with the soluble mass is moved to the modes i_c and j_c . Weingartner et al. (1997) show, that atmospheric particles can be divided into "more hygroscopic" and "less hygroscopic" depending on their growth behaviour when they are exposed to a relative humidity above 90%. Based on their findings we define that all material of mode s is moved to modes i_c and j_c if the soluble mass fraction of mode s rises above the threshold value $\varepsilon = 5\%$.

We do not consider the aging process due to photochemical reactions in this study. This is justified since it was shown by Saathoff et al. (2003) that these reactions do not contribute significantly to this process.

We apply the model to an area in south-western Germany. It covers main parts of Baden – Württemberg and the adjacent regions $(248 \times 248 \text{ km}^2)$. The horizontal grid size is $4 \times 4 \text{ km}^2$. The biogenic VOC emissions are calculated online



Fig. 1. Horizontal distribution of the externally mixed soot at 20 m above the surface, 12:00 CET, day 2 (left: summer, right: winter).



Fig. 2. Same as Fig. 1, but for internally mixed soot.

depending on the land use, the modelled temperatures and the modelled radiative fluxes (McKeen et al., 1991, Lamb et al., 1987, Vogel et al., 1995). We parameterise the NO emissions from the soil according to Ludwig et al. (2001). The anthropogenic emissions of SO₂, CO, NO_x, NH₃, 32 classes of VOC and diesel soot are pre-calculated with the spatial resolution of $4 \times 4 \text{ km}^2$ and a temporal resolution of one hour (Obermeier et al., 1995; Wickert et al., 1999; Pregger et al., 1999; Seier et al., 2000). The diesel soot emissions represent the sources by traffic. In addition to the source strength, we must prescribe the median diameter and the standard deviation of the emitted soot particle distribution. The median diameter is fixed to 60 nm and the standard deviation to 1.8 according to measurements by Vogt et al. (2000). We investigate two scenarios. First, we simulate a typical summer situation with a geostrophic wind of 4.5 m s^{-1} blowing from East. Second, we consider a winter day with a westerly flow with a geostrophic wind of 4 m s^{-1} . Three consecutive days were simulated in each case where the first day is used as spin-up and we only consider the results of the second and the third day. Considering further details about the initialisation and boundary conditions we refer to Riemer et al. (2003b).



Fig. 3. Vertical cross section of the externally mixed soot concentration for the summer case at y=80 km, 12:00 CET, day 2.

3 Calculation of the turnover rate

In the following we will explain the approach we use to determine the turnover rate of the externally mixed soot and how a turnover rate for a domain comparable to the grid size of a global climate model can be calculated. In MADEsoot the temporal development of the mass density \overline{m}_s of the externally mixed soot is determined by the following equation:

$$\frac{\partial m_s(\mathbf{x},t)}{\partial t} = \frac{\overline{A}(\mathbf{x},t) + \overline{S}(\mathbf{x},t) + \overline{D}(\mathbf{x},t) + \overline{E}(\mathbf{x},t) - \overline{T}(\mathbf{x},t)}{(1)}$$

 \overline{A} , \overline{S} , \overline{D} , and \overline{E} describe the rate of change due to advection, sedimentation, turbulent diffusion, and the emission, respectively. The bar indicates that the operators A, S, D and E are applied to the Reynolds-averaged quantities representing a typical spatial and temporal scale. \overline{T} is the turnover rate of the externally mixed soot and the sum of two processes:

$$\overline{T}(\mathbf{x},t) = \overline{Ca}(\mathbf{x},t) + \overline{Cn}(\mathbf{x},t)$$
(2)

Ca describes the transfer due to the coagulation of the externally mixed soot particles with soot containing internally mixed particles or soot free internally mixed particles. \overline{Cn} describes the transfer due to condensation of sulphuric acid and organic compounds. The externally mixed soot particles are transferred into internally mixed soot if the soluble mass fraction of mode *s* rises above the threshold value ε =5%. In our mesoscale model system KAMM/DRAIS both processes are explicitly determined as described in Riemer et al. (2003b).

Typical applications of the model simulations of KAMM/DRAIS cover areas of about $250 \times 250 \text{ km}^2$. This equals more or less the typical grid size of a global climate model. In the ideal case averaging a variable over the



Fig. 4. Same as Fig. 3, but for internally mixed soot.

mesoscale domain A should give the variable of the global climate model (assigned by a tilde).

$$\tilde{a}(z,t) = \frac{1}{A} \int \int \overline{a}(x, y, z, t) dA'.$$
(3)

In the global climate model the equivalent equation to Eq. (1) would read as:

$$\frac{\partial \tilde{m}_{s}(\mathbf{x},t)}{\partial t} =$$

$$\tilde{A}(\mathbf{x},t) + \tilde{S}(\mathbf{x},t) + \tilde{D}(\mathbf{x},t) + \tilde{E}(\mathbf{x},t) - \tilde{k}(z,t) \cdot \tilde{m}_{s}(\mathbf{x},t).$$
(4)

Here the transfer from the external to the internal mode is already parameterised as a first order reaction since in global climate models the aerosol processes usually cannot be treated as detailed as it is done in regional scale model. The problem is now reduced to the parameterisation of the rate coefficient \tilde{k} . Following the ideas explained so far \tilde{k} can be derived from results of the mesoscale simulations. This means:

$$\tilde{k}(z,t) = \frac{\int \int \int \int \overline{T}(x, y, z, t') dt' dA'}{\int \int \int \int \overline{T}(x, y, z, t') dt' dA'}.$$
(5)

with $\Delta t = 1$ h. This time interval was chosen as it is comparable to the time steps that are used in global climate models. In this way we determine time-dependent vertical profiles of \tilde{k} . The time scale that represents the exponential time scale for the decay of externally mixed soot is given by

$$\tau(z,t) = \frac{1}{\tilde{k}(z,t)}.$$
(6)

4 Results

Figures 1 and 2 show the horizontal distributions of the soot concentration in external and internal mixtures at 20 m above



Fig. 5. Composition of the total aerosol concentration averaged over the model domain at 12:00 CET at 25 m above the surface.

the surface at 12:00 CET (day 2) for the summer and the winter episode. The distributions of the soot in external mixture reflect the distribution of the sources, i.e. the main motorways and urban centres in the model domain. In the plumes of the urban areas and at some distance from the sources, soot is internally mixed. Comparing the winter and summer results we find that the total soot concentrations reach significantly higher values in winter due to the shallower boundary layer and the reduced vertical mixing in winter.

Figures 3 and 4 show the vertical cross sections of the soot concentration at y=80 km for the summer case. In the elevated layers soot exists mainly internally mixed. The values for the total soot concentrations range between 0.1 and $2 \,\mu g \, m^{-3}$ near the surface at 12:00 CET (day 2). However, higher concentrations (up to $10 \,\mu g \, m^{-3}$) are reached in the morning hours. Figure 5 displays an overview of the composition of the aerosol phase for the summer and the winter case. Shown is the aerosol concentration summed over all modes and averaged over the whole model domain for 12:00 CET (day 2) and 25 m above the surface.

During summer, the averaged sulphate concentration reaches a value of $6 \,\mu g \,\mathrm{m}^{-3}$. Due to temperatures around 27°C the nitrate concentration is low during the afternoon reaching only $0.3 \,\mu g \,\mathrm{m}^{-3}$.

During winter the sulphate concentration is much lower and reaches only $0.7 \,\mu g \,m^{-3}$ since the production via homogeneous oxidation of SO₂ in the gas phase through reaction with OH is less efficient. We do not consider cloud processes; hence the effect of the heterogeneous reaction of SO₂ in aqueous solution with ozone and/or peroxides is not included. Nitrate is much more abundant during the winter and reaches a value of $11.5 \,\mu g \,m^{-3}$. The order of magnitude of our model results for the aerosol composition are in agreement with measurements for continental Europe as the



Fig. 6. Vertical profiles for the aging time scale for summer (top) and winter (bottom) at day 2.

comparison with measurements shows (e.g. Neusüss et al., 2002; Zappoli et al., 1999; Mehlmann and Warneck, 1995). On the basis of the simulation results for both episodes we derive the aging time scale as described in Sect. 3. As explained there, τ may depend on height and time. Figure 6 shows time height sections of τ for the summer and winter episode, respectively. In both cases τ is below 2 h during the day and above 250 m above the surface. During the same time interval and below this height τ varies between 1 hour during the morning and up to about 20 h in the afternoon. During night time τ is much larger than during the day at all heights and there is a strong increase of τ from midnight of day 2 to midnight of day 3 in the summer and the winter case. While τ is almost constant with height during the night in the summer case it decreases with height in the winter case. Considering the temporal and spatial variations of τ in Fig. 6, it becomes obvious that the parameterisation of the aging process of soot using a fixed turnover rate represents an oversimplification.





Fig. 7. Daily cycles of the aging time scale (day 2).

Figure 7 (top) displays the daily cycle of τ in three different heights within the boundary layer for the summer episode (day 2) in more detail. At the heights of 250 m and 1100 m, we can clearly distinguish between the daytime and the night time regime. During daytime at these heights, τ is very short $(\sim 1 h)$. From detailed process studies based on our model results we find that condensation of sulphuric acid is very efficient in transferring externally mixed soot into the internal mixture when sulphuric acid is abundant. During the night, the production of sulphuric acid ceases. Once the available sulphuric acid has been depleted by condensation on the existing particles, condensation of sulphuric acid therefore stops to occur. Under these conditions coagulation is the only process for the transfer into internal mixture, operating very slowly. This leads to much higher values for τ during the night compared to the day.



Fig. 8. Composition of the internally mixed soot particles averaged over the model domain at 12:00 CET at 25 m above the surface.

At the height of 25 m, which is identical with the source height used for our simulations, the distinction between day and night is not as apparent and τ remains at a relatively high level of about 20 h in the afternoon. The reason is that at this height, fresh soot is permanently emitted and the concentration of externally mixed soot reaches its maximum. Given a certain amount of condensable species, it takes a longer time until the shell mass has grown to the certain threshold that marks the transition between externally and internally mixed soot which eventually results in the high values for τ .

At all heights a characteristic trend of the time scale is obvious. This trend can be attributed to the fact that the coagulation rate depends on the particles number densities. The particle number densities of the mode s are largely dominated by the emissions. However, the number densities of the modes i_f , j_f , i_c and j_c are decreasing with time in our simulation, because nucleation on the third day is weaker than on the second day due to the meteorological conditions. Consequently, since these are the particles that provide the coagulation partner for the externally mixed soot for the transfer into the internal mixture, the time scale increases. Similar arguments apply to the winter episode as displayed in Fig. 7 (bottom). Compared to the summer conditions, however, important differences concern the following issues: Obviously, the daytime is shorter, solar radiation less intense and temperatures are significantly lower. This causes a shallower boundary layer and decelerated photochemistry, in particular a lower production rate of H₂SO₄. However, the lower temperatures favour the formation of ammonium nitrate. It turns out that ammonium nitrate contributes significantly to the aging of soot in winter during daytime and hence compensates the lack of H₂SO₄ leading to time scales during the day that are comparable to the summer case. To underline these findings Fig. 8 shows the average mass densities of



Fig. 9. Time height section of the mass transfer rate of pure soot from the external to the internal mixture by condensation only averaged over the whole model domain and integrated for a time interval of one hour.

the individual species for the internally mixed soot aerosol (modes i_c and j_c) at 12:00 CET (day 2) at 25 m above the surface. While the soluble mass fraction of these aerosol particles is clearly dominated by sulphate for the summer case nitrate represents the main contribution for the winter case. Of course the formation of ammonium nitrate also occurs during the night when the gas phase precursor HNO₃ is formed due to the heterogeneous hydrolysis of N₂O₅ (Riemer et al., 2003a). However, since the formation and hence the condensation of H₂SO₄ ceases during the night due to the lack of sunlight, the soot particles that are emitted during the night cannot be coated with sulphate. Since in our current model formulation the total amount of nitrate is distributed over the modes according to the sulphate content the available nitrate ends up in the modes i_f , j_f , i_c and j_c and not in mode s. This means that the aging of soot due to condensation is not effective during the night, both in winter and in summer. Figure 9 which displays a time height section of the soot transfer due to condensation (winter case only) shows this fact.

We are aware of the fact that this treatment of aerosol chemistry represents a limitation of our model. With this assumption it is not possible to form ammonium nitrate on the soot particles without sulphate being involved. In reality this process might occur which means that we might overestimate the time scale for the situations where ammonium nitrate is present. This issue will be addressed in future work.

The pronounced minimum for the time scale in 25 m (Fig. 7, bottom) can be attributed to the interaction of two processes that act in the opposite direction: First, the condensation of sulphate and the formation of ammonium nitrate sets in, which leads to a decrease in τ . Concurrently, the surface area of the particles in the modes i_f , j_f , i_c and j_c increases compared to the surface of the particles in mode s. Therefore the low value of τ cannot be sustained and increases again.



Fig. 10. Correlation of τ and the number density of the internally mixed particles. The solid line shows the fit according to Eq. (7). The blue dots indicate the results for the winter day, the red ones those for the summer day.

Summarizing our findings we can state the following:

Our model results suggest to distinguish between two regimes, a daytime and a night time regime. During daytime, condensation of sulphate and nitrate is the dominant process, and an additional separation according to the height above the source region appears to be appropriate. Above 250 m τ is approximately 2 h, regardless of the season. In order to derive the time scale during daytime and below 250 m we apply the following procedure. The time scales shown in Fig. 6 are averaged below 250 m and for the time interval where the global radiation is above zero. We find an average time scale of 9.38 h for the summer case and 6.89 h for the winter case.

For the night time coagulation has been found to be the governing process. Compared to the day time results the time scales during the night show a much higher variability with height and especially with time, with a clear increase from night to night. Apart from other parameters such as the mean diameters, the coagulation coefficients, the temperature coagulation depends on the number densities of the internally mixed particles. Therefore Fig. 10 shows the simulated values of τ versus the number density of the internally mixed particles for the summer days and the winter days. All grid points are shown for heights below 1100 m above the surface during night time. As expected a decrease of τ with increasing number density is obvious. For these data we calculate the following fit, the result of which is depicted in Fig. 10:

$$\tau = \frac{a}{N} + \frac{b}{N^2} \tag{7}$$

with $a=6\times 10^{4}$ h cm⁻³ and $b=3\times 10^{8}$ h cm⁻⁶

N is the number density in the modes i_f , j_f , i_c and j_c in cm⁻³ and the dimension of τ is h.

The limitations inherent to our case studies should be stated clearly:

We only include soot emissions from traffic. While they contribute a large fraction of the emissions in densely populated areas, more sources contribute to the total soot than diesel cars alone, especially in winter. However, we did not have emission data for these emission categories.

Cloud and wet deposition processes as well as cloud chemistry are not included in the model. Again, this introduces uncertainties mainly for the winter simulation. It is impossible to estimate the overall effect of clouds on the aging process due to the competing effects of sulphate production in the aqueous phase and the removal by wet deposition as has been shown by Koch et al. (2003).

Despite these limitations the discussion of our case studies reveals the important issues that the time scale of soot aging is highly variable with respect to time and height and that nitrate can play an important role for the aging process.

5 Conclusions

We performed three-dimensional model simulations with the coupled mesoscale- γ model KAMM/DRAIS to derive the time scale τ that characterizes the transfer of diesel soot from external into internal mixture. From the two case studies that we considered we are not able to derive a parameterisation that will hold for the global scale. However, the method that we presented to derive the aging time scales are generally applicable and can be used as a tool in the future to improve the parameterisation for global models.

Generally, the aging time scales that we derived are smaller than most of the values that are currently used in global climate models and show a considerable variability in space and time. Hence, parameterising the aging process of soot with a fixed turnover rate represents an oversimplification.

During daytime in summer, condensation of sulphuric acid is the governing process for the aging of soot. In wintertime, the formation of ammonium nitrate gains in importance. To our knowledge the role of ammonium nitrate for the aging of soot has been highlighted for the first time. Overall, the time scales for daytime in summer and winter are comparable, about 2 h above 250 m and 8 h below during night time, condensation stops being the important process. Instead, coagulation becomes more significant, acting very slowly. This leads to a time scale during night of 10–40 h.

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