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Aerosol dynamics within and above forest in relation to turbulent transport and dry deposition

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Abstract. A 1-D atmospheric boundary layer (ABL) model coupled with a detailed atmospheric chemistry and aerosol dynamical model, the model SOSAA, was used to predict the ABL and detailed aerosol population (characterized by the number size distribution) time evolution. The model was applied over a period of 10 days in May 2013 to a pine forest site in southern Finland. The period was characterized by frequent new particle formation events and simultaneous intensive aerosol transformation. The aim of the study was to analyze and quantify the role of aerosol and ABL dynamics in the vertical transport of aerosols. It was of particular interest to what extent the fluxes above the canopy deviate from the particle dry deposition on the canopy foliage due to the above-mentioned processes. The model simulations revealed that the particle concentration change due to aerosol dynamics frequently exceeded the effect of particle deposition by even an order of magnitude or more. The impact was, however, strongly dependent on particle size and time. In spite of the fact that the timescale of turbulent transfer inside the canopy is much smaller than the timescales of aerosol dynamics and dry deposition, leading us to assume well-mixed properties of air, the fluxes at the canopy top frequently deviated from deposition inside the forest. This was due to transformation of aerosol concentration throughout the ABL and resulting complicated pattern of vertical transport. Therefore we argue that the comparison of timescales of aerosol dynamics and deposition defined for the processes below the measurement level do not unambiguously describe the importance of aerosol dynamics for vertical transport above the canopy. We conclude that under dynamical conditions reported in the current study the micrometeorological particle flux measurements can significantly deviate from the dry deposition into the canopy. The deviation can be systematic for certain size ranges so that the time-averaged particle fluxes can be also biased with respect to deposition sink.

1 Introduction

Turbulent fluxes of scalars are commonly measured by the eddy covariance (EC) technique above forests. From flux measurements the exchange of scalars between the ecosystem and the atmosphere is inferred by making simplifying assumptions, mainly horizontally homogeneous and stationary conditions, considering usually transport of passive scalars. From aerosol particle flux measurements deposition to ecosystem is inferred by neglecting all additional terms including the storage term. However, there are several mechanisms affecting the particle concentration, namely new particle formation, coagulation and source or sink terms for a particular size resulting from condensational growth. These processes, which we refer to as the aerosol dynamical processes throughout this study, govern the particle size distribution evolution. The significance of aerosol dynamical terms in comparison to dry deposition has been evaluated by comparing the respective timescales. The timescale for dry deposition for measurement level z has been estimated according to \( \tau_{dep}(z) = \frac{z}{V_d} \), where \( V_d = \frac{F(z)}{C(z)} \) denotes the bulk deposition velocity defined as the ratio of the total flux divided by the concentration at the same level (Pryor and Binkowski, 2004; Pryor et al., 2013). Such a definition of the timescale of dry deposition implies that frequently the aerosol dynamical terms have similar timescales to dry deposition and therefore affect the conservation of aerosol particle concentration.
during the transport pathway between the EC measurement level and the collecting surfaces. Depending on the particle size and the prevailing conditions, i.e., the nucleation rate, the availability of condensing vapors determining the condensational growth and the shape of the particle size spectrum, the aerosol dynamical terms can vary significantly. The timescale of aerosol dynamical processes varies typically between $10^3$ and $10^5$ s (Pryor and Binkowski, 2004; Pryor et al., 2013), i.e., being on the hourly timescale and more. This is a sufficient time to allow well-mixed conditions to establish within the unstable daytime atmospheric boundary layer (ABL), where the mixing timescale is estimated to be around 10 min (e.g., Stull, 1988). Under near-neutral and stable conditions such efficient mixing throughout atmospheric column cannot be assumed. Instead the characteristic timescales of turbulent transfer within and above forests have been estimated by different approaches (e.g., Zelger et al., 1997; Rinne et al., 2000, 2012; Rannik et al., 2009b). Such timescales of turbulent transfer depend on the observation conditions but typically remain on the order from a few tens of seconds to a few hundreds of seconds. In spite of different definitions used and large variation range of the timescales characterizing the scalar transport between the observation level and the collecting surfaces within forest, the turbulent transfer can be expected to occur much faster than the aerosol dynamical processes.

The aerosol particle dry deposition is strongly size dependent as different mechanisms operate at different particle sizes. The timescale of dry deposition depends on particle size and exhibits its maximum at around 100 nm. For small particles with a few nanometers in diameter this dry deposition timescale can be orders of magnitudes smaller due to efficient removal mechanism by Brownian diffusion. At particle sizes larger than 100 nm the particle collection is again enhanced due to interception and inertial impaction mechanisms (Petroff et al., 2008) and the respective timescale of dry deposition is smaller. In general, the dry deposition timescale has been frequently estimated to be in the same order of magnitude as the timescale for aerosol dynamics, leading to a conclusion that flux divergence may occur during transport due to aerosol dynamics (Pryor and Binkowski, 2004; Pryor et al., 2013).

The timescales of turbulent transfer and the timescale of dry deposition embed essentially different definitions and can lead also to different conclusions about the significance of aerosol dynamical terms during the transport between the underlying surfaces and the measurement level. The timescale of turbulent transfer is the characteristic time of the transfer within turbulent air layer. Dry deposition includes in addition the transport pathway within the laminar air layer surrounding the collecting surfaces. In the resistances framework (e.g., Monteith and Unsworth, 1990), the dry deposition includes the aerodynamic (corresponding to turbulent transport) as well as the leaf laminar sublayer resistances and under most conditions the dry deposition is limited by the laminar boundary layer transfer (e.g., Petroff and Zhang, 2010). Therefore comparison of the timescales of turbulent transport and dry deposition with that of aerosol dynamics leads us to the assumptions that (i) turbulent transport within and above forest is relatively fast and no significant transformation of aerosol population occurs within the respective timescale and (ii) depending on particle size the removal of aerosols via dry deposition occurs at the comparable timescale with aerosol dynamics and therefore the aerosol population can be modified during the removal process. Such modification occurs on hourly timescales and therefore is expected to occur throughout the ABL, where aerosol dynamical processes can depend strongly on height within the ABL via vertical profiles of condensing vapors.

The purpose of this study is to analyze the magnitude of different terms in the particle number conservation equation and to evaluate the timescales of particle turbulent transfer, aerosol dynamical processes and dry deposition over a wide range of particle sizes. Further, we evaluate the effect of these terms on inferring particle deposition velocities from flux measurements by micrometeorological techniques, in particular the influence on estimation of functional dependencies as well as systematic biasing effects. The study relies on the simulations by the model to Simulate the concentration of Organic vapors, Sulphuric Acid and Aerosols (SOSAA) and the measurements were used only to initialize the model (see Sect. 2.2 and Appendix B) or for evaluation of model outputs in terms of predicted particle size distributions and meteorological variables such as heat fluxes above canopy (see Sect. 3 below). Non-stationary conditions will be considered by simulating detailed ABL and aerosol dynamics inside and above the forest canopy during a period of 10 days, which includes highly dynamical conditions with new particle formation.

2 Materials and methods

The model was set up for a pine forest site in southern Finland and initialized with available measurements performed at the SMEAR II station. For description of the site and initialization of the model see Appendices A and B, respectively. The analysis relies on evaluation of the significance of different terms of the particle conservation equation.

2.1 Conservation equation for aerosol size distribution

In horizontally homogeneous conditions, when neglecting molecular diffusivity and applying the first-order closure to turbulent flux,

$$\overline{w' n'} = -D_i \frac{\partial \bar{\pi}}{\partial z},$$

(1)

the conservation equation for time-averaged particle number density $\bar{\pi} = \frac{\partial \bar{N}}{\partial \ln D_p}$ inside the canopy can be written as
where $N$ is the average particle number concentration, $D_p$ the particle diameter, $D_t$ the particle turbulent diffusivity, $w_s$ the settling velocity, $v_c$ the particle collection velocity by vegetation and $a$ the all-sided leaf area density. The source/sink term $S_{ad}$ incorporates all aerosol dynamical terms, consisting of nucleation $S_{nucl}$, condensational growth $S_{cond}$ and coagulation $S_{coag}$ terms. If the condensational growth rate is considered as $I_{cond}(\log_{10} D_p) = \frac{d \log_{10} D_p}{dt}$, then the respective source/sink term in Eq. (2) is expressed as $S_{cond} = \left[ \frac{d \pi}{dt} \right]_{cond} = - \frac{\delta (I_{cond})}{\delta (D_p)}$. For particle size range up to a few micrometers $D_t$ can be assumed to be equal to the eddy viscosity of the flow. The settling velocity $w_s$ is given as

$$w_s = \frac{C_c g \rho_p D_p^2}{18 \eta},$$

where $g$ is the acceleration due to the gravity, $\eta$ the dynamic viscosity of air, $\rho_p$ the particle density and $C_c$ the Cunningham slip correction factor (e.g., Hinds, 1982).

For the comparison of the significance of different terms of the conservation equation, the Eq. (2) was re-written so that the sum of all terms equaled 0, and the transport due to settling was merged with the particle collection by vegetation as

$$\left[ - \frac{\partial \pi}{\partial t} \right] + \left[ \frac{\partial}{\partial z} \left( D_t(z) \frac{\partial \pi}{\partial z} \right) \right]$$

$$+ \left[ -a(z) v_c \pi + \frac{\partial}{\partial z} (w_s \pi) \right] + [S_{ad}] = 0,$$

where the terms were called the storage, (vertical) transport, particle deposition and aerosol dynamical terms. Further, integration of Eq. (4) from the forest floor surface up to the canopy top $h$ was used to define the change velocities in analogy to deposition velocity. The change velocity due to particle deposition was defined as

$$V_{dep} = \frac{1}{\pi(h)} \int_0^h \left[ -a(z) v_c \pi + \frac{\partial}{\partial z} (w_s \pi) \right] dz$$

and the change velocity due to aerosol dynamics as

$$V_{ad} = \frac{1}{\pi(h)} \int_0^h S_{ad} dz.$$

In particular, for the transport term the respective change velocity was defined as

$$V_{transp} = \frac{1}{\pi(h)} \int_0^h \frac{\partial}{\partial z} \left( \frac{-\pi w'}{\pi} \right) dz$$

Note that in the modeling approach the vertical flux at the canopy top was obtained from the gradient diffusion approximation (Eq. 1) and the flux at the surface was defined by the ground deposition parameterization, which was applied as the sink term in the lowest model layer. Therefore in our model calculations $\frac{\pi w'}{\pi}(0) = 0$ and the transport velocity equaled to the exchange velocity defined at the canopy top by

$$V_e = - \frac{F(h)}{\pi(h)}.$$

The timescales of the processes affecting the particle concentration inside the canopy were defined by

$$\tau = \frac{h}{V_e},$$

with the change velocities $V_{dep}, V_{ad}$ and $V_e$ defining the timescales for deposition $\tau_{dep}$, aerosol dynamics $\tau_{ad}$ and exchange $\tau_e$, respectively. These timescales were calculated based on the numerical modeling results by SOSAA.

### 2.2 Simulation of aerosol transport and dynamics by model SOSAA

The model SOSAA is a 1.5-order RANS (Reynolds-averaged Navier–Stokes) model SCADIS (SCAlar DIStribution, 1-D version; Sogachev et al., 2002, 2012) coupled with detailed biogenic emissions, chemistry and aerosol dynamics. SCADIS describes the exchange between the vegetative canopy and atmosphere by considering the vegetation as a multi-layer medium and implementing parameterizations for radiation transfer, drag forces on leaves and stomatal conductance. The particle deposition processes in SOSAA are treated in the same manner as in the study by Lauros et al. (2011) based on the parameterization by Petroff et al. (2008). The parameterization considers Brownian diffusion and takes into account the influence of leaves on particle interception, impaction and settling. The model has been applied extensively in different forest sites for various studies concerning biogenic emissions, chemistry and aerosol formation (e.g., Kürten et al., 2011; Boy et al., 2013; Smolander et al., 2014; Mogensen et al., 2015; Zhou et al., 2015). Detailed model description is presented by Boy et al. (2011) and Zhou et al. (2014).

The model setup in this study was the same as in the study by Zhou et al. (2014) except that only kinetic nucleation
mechanism was employed in aerosol dynamics simulation (Weber et al., 1997; see also Sect. S2 in the Supplement). Zhou et al. (2014) presented the ability of SOSAA to reconstruct new particle formation events at Hyytiälä, which was the same site as in this study. The model was initialized with vertical profiles describing the initial atmospheric state (see Appendix B) and aerosol size spectrum observed at the surface and run for a 10-day time period similarly to Lauros et al. (2011). The aerosol size distribution was initialized each day at 00:00 LT based on the measurements at 2 m height. The first day the concentration profile was assumed constant (the same as at 2 m height) up to determined nighttime stable boundary layer (SBL) height (320 m) and 10% of the concentration values within the SBL above this level. During the next days the concentration profile was taken constant as per measurements at 2 m level up to the maximum ABL height occurring during the previous day and 10% of the within SBL values above that level. The initialization during the first day corresponded to the conditions of horizontal advection with very different properties of the air above the SBL, whereas during the other days the nighttime residual layer was assumed to retain the same properties as the SBL. The implications of these two contrasting assumptions for ABL mixing and vertical transport of aerosols will be discussed in Sect. 3.4. For meteorology simulations 10 sec time step was used along with the explicit forward in time integration method. The aerosol dynamics was simulated with 60 s time step.

2.3 Lagrangian estimation of turbulent transfer time

The Lagrangian stochastic (LS) simulations were used to estimate the turbulent transfer time. The conventional approach of using a LS model is to release particles at the surface point source and track their trajectories towards the point of interest forward in time (e.g., Wilson and Sawford, 1996). In case of horizontally homogeneous and stationary turbulence, the mean Lagrangian turbulent transfer time at the canopy top due to a sustained source located at height $z_0$ (near forest floor) can be described as

$$\tau_L(z) = \frac{1}{N} \sum_{i=1}^{N} \tau_i,$$  \hspace{1cm} (10)

where $\tau_i$ denotes the travel time of trajectory $i$ at the moment of intersection with observation height. For LS modeling the turbulence statistics such as the turbulent kinetic energy (TKE) and the vertical eddy diffusivity obtained from SOSAA were used to define the turbulent profiles of the dissipation rate of TKE and variances of the wind speed components.

Figure 1. Aerosol size distribution at 2 m height during a 10-day period in May 2013 as (a) measured by the DMPS system and (b) predicted by the model SOSAA.

3 Results

The selected time period consisted of 10 days in May 2013, days of year (DOYs) 121 (1 May) to 130 (10 May). On several days clear particle formation patterns were observed at the smallest particle sizes around midday, with subsequent growth to larger particle sizes (Fig. 1). In all days significant aerosol dynamics was taking place in terms of particle growth. The model simulations reproduced the observed particle size distributions qualitatively; however, it was unable to reproduce the exact particle size distribution patterns. In particular, during days with new particle formation the observed nucleation modes were not as clear; the particle growth was also overestimated, which can be observed clearly during the second half of the period. With respect to condensational growth of aerosols and resulting patterns of aerosol particle distributions a sensitivity analysis was performed (Fig. S2). The results of the sensitivity analysis are summarized in the end of Sect. 3.

The ABL height varied between about 600 (DOY 130) and 1400 m (DOY 123) as the peak height during different days (Fig. 2a). The heat fluxes were the primary drivers of the ABL growth and buoyancy-driven TKE. The simulated latent and sensible heat fluxes corresponded well to those measured at the site (Fig. 2b, c), but the simulated TKE had weak correlation with the values observed above the canopy (Fig. S1 in the Supplement). We ascribe this to the limited ability of the 1-D model to reproduce the actual flow field at the site. However, for the current study it is more important to reproduce diurnal variation and dynamics of the ABL, which is mainly driven by surface heating. The selected 10-day period showed significant variability in terms of aerosol and ABL dynamics and was therefore selected as the study case.
3.1 Aerosol dynamics and transport inside and above forest

The particle conservation terms were evaluated inside the forest on 7 May (DOY 127) at 12:00 and 21:00 LT (UTC +2). At noon the particle size spectrum was bi-modal, with nucleation and larger particle modes, by evening the nucleation mode had grown and almost merged into a single mode at around 200 nm (Fig. 3a). The rate of change by each term (as defined by the terms in Eq. 4) showed large particle sink due to deposition, which was compensated by the transport term at noon (Fig. 3b). The aerosol dynamical term was dominated by the condensational growth term, except at sizes smaller than a few tens of nanometers where coagulation was also important and at smallest sizes were particles due to nucleation appeared. The aerosol dynamics reduced the particle number of small particles less than about 10 nm in diameter, adding particle counts at larger sizes. The aerosol dynamical terms were reflected in relatively similar pattern in particle storage change (defined by the first term of Eq. 4). The positive value of the storage term implies a decrease of particle concentration and a negative value an increase. In the evening at 21:00 LT the change rates of small particles (less than 20 nm) were small due to low particle counts in this part of the size spectrum (Fig. 3a). The similarity (in magnitude, but opposite in sign) of aerosol deposition vs. transport and aerosol dynamical vs. storage change terms held also in the evening, leading us to conclude that particle loss due to deposition was mainly compensated by vertical transport and aerosol dynamical processes modified the concentration in time.

The aerosol concentration inside and above forest was homogeneous at noon and small vertical concentration gradients could not be observed from color presentation in Fig. 4a. The deposition pattern (dependence on particle size and height) was again similar to transport patterns (Fig. 4d and c). Aerosol dynamics affected the number concentration similarly throughout the column as presented in Fig. 4b and e. The same qualitative conclusions held also for the evening time, 21:00 LT (not shown).
When integrating the terms of the conservation equation (Eq. 4) from the surface up to the canopy top and normalizing with the concentration at the canopy top, one obtains change velocities as defined in Sect. 2.1. Such change velocities are comparable with the deposition velocity or the exchange velocity, which can be experimentally obtained from the flux measurements above the canopy. In terms of change velocities the deposition velocity (defined by Eq. 5) and the transport velocity (defined by Eq. 7 and equivalent to the exchange velocity in Eq. 8) appeared near symmetric for all particle sizes at noon (Fig. 5a). However, the correspondence was not exact, meaning that the flux defined at the canopy top did not correspond exactly to particle deposition. This was due to aerosol dynamics being responsible for additional sink inside the canopy for sizes up to 10 nm, creating concentration decrease as well as additional downward particle transport to compensate for the loss. Much larger differences in the respective patterns were observed in the evening at 21:00 LT, especially at small particle sizes (Fig. 5b). This implied a more complex relationship between particle source/sink/term (deposition and aerosol dynamics) and vertical mixing.

The vertical profiles of the aerosol dynamical term (normalized to simulated local concentrations, defining the local change rates) and the particle vertical fluxes (normalized with local concentration, defining the local exchange velocity) differed significantly for particle sizes and time of day (12:00 LT compared to 21:00 LT, 7 May), Fig. 6 upper and lower panels. The respective ABL heights were approximately 710 and 510 m according to the model results. At noon the particle deposition and aerosol dynamics led to vertical particle transport that depended on particle size and height. In the lower part of the ABL the small particles (3 and 10 nm) were transported downward to compensate for deposition sink inside the forest and particle loss through aerosol dynamics. The 100 nm particles were transported downward throughout the atmospheric column. For particles of 30 and 300 nm size it was predominantly the aerosol dynamics that drove the vertical transport, leading mostly to upward particle flux at heights above forest. The particle concentration gradients (Fig. 6a/u) were consistent...
3.2 Timescales of processes

The importance of aerosol dynamics for particle exchange measurements has been frequently assessed by comparing the timescales of aerosol dynamical and transport processes. Figure 7 presents the timescales defined in Sect. 2.1 and compares those with the Lagrangian turbulent transfer timescale determined according to Sect. 2.3. The time of turbulent transfer within forest (simulated as the time for an air parcel to travel between the surface and the forest height) was mostly much shorter than the timescales of deposition and aerosol dynamics. Only at smallest particle sizes and stable conditions did the turbulent timescale become comparable to the timescales of particle deposition and aerosol dynamics (Fig. 7b). The transport timescale, defined by Eqs. (9) and (7), accounts also for the effect of sources and sinks inside the canopy and is therefore very different from the turbulent transfer timescale \( \tau_L \). The transport timescale was determined mainly by deposition and modified by the impact of aerosol dynamics, reflecting the fact that particle vertical transport is mostly controlled by the sources and sinks and not limited by turbulent transfer speed.

The timescale of particle deposition strongly depended on particle size (resulting of respective dependence of particle collection on particle size), whereas the timescale of aerosol dynamics was occasionally shorter than the deposition timescale (even an order of magnitude, depending on particle size). Even though the turbulent transfer timescale \( \tau_L \) was much shorter than the other timescales, the flux at the canopy top deviated from the deposition to vegetation elements (can be inferred from the comparison of the deposition and the transport timescales). Note that even the sign of the flux at the canopy top differed for particles of about 100–300 nm in diameter; see the sign of the transport timescale in Fig. 7a. Although very short turbulent transfer time would suggest fast and efficient mixing (and therefore correspondence of flux to deposition), the difference can be explained by the importance of the aerosol dynamics which affect the concentrations throughout the atmospheric column and therefore drive the vertical redistribution of particles via vertical transport.

3.3 Time evolution and statistics of particle exchange

The idea behind micrometeorological particle flux measurements is to determine the particle dry deposition fluxes or equivalently the deposition velocities. Thus it is assumed that the fluxes observed above the forest represent the deposition...
Figure 7. The timescales of deposition, aerosol dynamics and transport (equivalent to vertical exchange) as defined by Eq. (9) together with Eqs. (5), (6) and (7) at (a) 12:00 LT (the values for the transport term are out of scale at about 200 and 450 nm, being about $+3.7 \times 10^6$ and $-1.05 \times 10^6$ s, respectively) and (b) 21:00 LT on 7 May 2013. In addition the Lagrangian timescale for turbulent transfer (corresponding to aerodynamic resistance only) as simulated according to Eq. (10), is presented as the median air parcel travel time between the forest floor and the canopy top with upper and lower quartiles. The “+” sign reflects the positive sign of the respective term (the source), whereas no such sign infers the negative (sink) term.

Figure 8. (a) Particle size spectrum and the change velocities (presented as the ratios to the absolute value of the deposition term) for selected particle sizes for (b) storage, (c) aerosol dynamics and (d) vertical exchange during 1 and 2 May (DOY 121 and 122) 2013.

Fluxes. Figure 8 compares the change velocities defined in Sect. 2.1 to the respective deposition change velocities during the first day of the simulations, 1 May (DOY 121), and the following nucleation day, 2 May 2013 (DOY 122). These two days differ in terms of initialization of vertical aerosol profiles at midnight (see Sect. 2.2). During the first day the aerosol dynamics barely affected the particle concentrations inside the forest, but 100 and 300 nm sizes were affected strongly by vertical transport occurring during the mixed layer (ML) growth period prior to noon. The initial concentration profile during this day corresponded to the conditions of horizontal advection. During the second day the aerosol dynamical term exceeded the deposition term several times (Fig. 8c). The storage change varied in approximately the same limits, and it is opposite in phase (Fig. 8b). The variation of the exchange velocity with respect to deposition was smaller (Fig. 8d), consistent with the analysis of Fig. 3 in which the vertical transport was the main mechanism compensating for aerosol loss due to deposition. Nevertheless, the magnitude of the exchange velocity can also differ several times compared to that of deposition. During the new particle formation and ABL growth period of the second day the vertical particle exchange showed downward transport of small particles (3, 10 and 30 nm) and upward transport of 100 nm particles. In particular during the first day (DOY 121), the upward particle transport was synchronous with the storage change, i.e., the concentration decrease (Fig. 8b), referring to the dilution of concentration within the canopy. Downward transport of 10 nm particles during the second day in turn exceeded significantly the particle deposition. This particle size range was affected then by the aerosol dynamical term changing (from negative to positive) during the morning hours due to particle growth (Fig. 8c), which was due to the fact that 10 nm size was on the lower edge of the dominant mode of the particle size spectrum (Fig. 8a). Note also that the storage change of 10 nm particles was similar to the aerosol dynamical term (opposite in sign) and not to the ex-
change velocity. Therefore the relatively large downward flux during the second day (DOY 122) was mainly driven by the aerosol dynamics occurring at night, whereas the growth of the ML initiated strong vertical mixing.

In order to understand overall trends and variability in aerosol dynamics and transport, the diurnal patterns of the averages together with the range of variation were presented in Fig. 9 for three particle sizes characterizing the nucleation (10 nm), Aitken (50 nm) and accumulation (300 nm) modes. For 50 nm particles the aerosol dynamics were a sink at night, whereas the condensational growth served as the source of 300 nm particles round the clock. The variation range of the aerosol dynamical term can be very large, indicating the role of ABL development during different days. Whereas the variation range of the aerosol dynamics and storage was large generally at night, the vertical exchange deviated from deposition mainly during the early morning SBL and further ABL growth period until noon (Fig. 9c). During this period the 50 nm particle fluxes were larger than induced by deposition, and during the ABL growth the 300 nm particle fluxes were lower than would have corresponded to deposition on average.

We further looked how different particle sizes were affected during different stages of the ABL state. At night the aerosol dynamics affected a wide range of particles and on average performed as the sink for particles less than 100 nm and the source for larger particles (Fig. 10c). The aerosol dynamical sink/source led primarily to particle concentration change. During the morning hours from sunrise until noon the ABL growth induced enhanced downward transport of about 30–200 nm particles, whereas vertical downward transport of larger particles was less than deposition sink (Fig. 10a). During the afternoon all the change velocities exhibited less variation compared to morning and night hours. Consequently deposition was also the best represented by the averages fluxes at the canopy top in the afternoon, with biggest deviation coinciding with the minimum in deposition velocity at around 100 nm (Fig. 10b). Figures 8, 9 and 10 (see also Fig. S3) illustrate that both the aerosol dynamics and ABL growth can strongly affect the vertical transport of aerosols and the fluxes above the canopy can deviate significantly from the deposition occurring within the canopy.

Due to instrumental limitations or by intention (frequently to obtain statistically significant particle counts in order to reduce particle flux random errors), a certain size interval of particles is measured. Figure 11 presents the vertical exchange velocity size-integrated values to represent the nucleation (3–30 nm), Aitken (30–100 nm) and accumulation (100–1000 nm) mode particles. During the first day with assumed conditions of horizontal advection the size-integrated particle fluxes showed clear upward transport during the morning hours for 30–200 and 100–1000 nm size ranges. The same has also been observed from the measurements and interpreted as the upward transport due to ABL growth and resulting dilution of relatively particle-rich air within forest with the particle-poor air transported down from aloft (e.g., Nilsson et al., 2001). The days with very large (both positive and negative) values of the exchange velocities compared to deposition velocities corresponded to the days with preceding very low ABL heights at nights (DOY 127, 129, 130). Therefore the ABL development can be identified as one of the main reasons for the large variation in vertical transport of particles. In case of experimental flux measurements the statistical uncertainty as well as natural variation originating from spatial heterogeneity and horizontal advection can additionally contribute to the variance of the calculated fluxes, leading to flux patterns with large variation, that are often difficult to interpret.

Table 1 presents the statistics of the fluxes at the canopy top (relative to deposition) for different particle sizes. Whereas for smaller particles 3–10 nm the time-averaged particle flux statistics converged to particle deposition within forest, for larger particles the fluxes (if measured by the micrometeorological technique) were biased in representing the particle deposition even on the average. The largest devia-
Table 1. Statistics of the ratio of the flux at the canopy top to deposition sink integrated over the canopy over a 10-day period in May 2013. The average statistics $\langle V_e \rangle$ and $\langle |V_{dep}| \rangle$ were averaged over the simulation period first and then the ratio was found, whereas the percentile statistics apply for the ratios $V_e / |V_{dep}|$ obtained from model simulations for each 10 min period.

<table>
<thead>
<tr>
<th>Particle size (nm)</th>
<th>3</th>
<th>10</th>
<th>30</th>
<th>100</th>
<th>300</th>
<th>850</th>
<th>3–30</th>
<th>30–100</th>
<th>100–1000</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\langle V_e \rangle$</td>
<td>0.90</td>
<td>0.99</td>
<td>1.36</td>
<td>2.09</td>
<td>0.53</td>
<td>0.82</td>
<td>1.11</td>
<td>1.99</td>
<td>0.66</td>
</tr>
<tr>
<td>$\langle</td>
<td>V_{dep}</td>
<td>\rangle$</td>
<td>$-0.24$</td>
<td>$-0.18$</td>
<td>$-0.32$</td>
<td>0.15</td>
<td>$-1.30$</td>
<td>0.20</td>
<td>0.33</td>
</tr>
<tr>
<td>$Q_5$</td>
<td>0.77</td>
<td>0.87</td>
<td>0.93</td>
<td>0.82</td>
<td>0.32</td>
<td>0.73</td>
<td>0.97</td>
<td>0.95</td>
<td>0.56</td>
</tr>
<tr>
<td>$Q_{25}$</td>
<td>0.97</td>
<td>1.00</td>
<td>1.06</td>
<td>0.94</td>
<td>0.85</td>
<td>0.92</td>
<td>1.06</td>
<td>1.04</td>
<td>0.86</td>
</tr>
<tr>
<td>Median</td>
<td>1.11</td>
<td>1.09</td>
<td>1.31</td>
<td>1.34</td>
<td>0.92</td>
<td>0.96</td>
<td>1.20</td>
<td>1.57</td>
<td>0.92</td>
</tr>
<tr>
<td>$Q_{75}$</td>
<td>1.81</td>
<td>1.70</td>
<td>3.36</td>
<td>9.59</td>
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<td>1.00</td>
<td>2.12</td>
<td>10.5</td>
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</tbody>
</table>

Table 2. Statistics of the ratio of the flux at the canopy top to deposition sink integrated over the canopy over a 10-day period in May 2013. For more details see Table 1. Morning refers to the time period from sunrise until noon, afternoon from noon until sunset and night from sunset until sunrise.

<table>
<thead>
<tr>
<th>Time</th>
<th>Morning</th>
<th>Afternoon</th>
<th>Night</th>
</tr>
</thead>
<tbody>
<tr>
<td>Particle size (nm)</td>
<td>3–30</td>
<td>30–100</td>
<td>100–1000</td>
</tr>
<tr>
<td>$\langle V_e \rangle$</td>
<td>1.25</td>
<td>2.92</td>
<td>0.48</td>
</tr>
<tr>
<td>$\langle</td>
<td>V_{dep}</td>
<td>\rangle$</td>
<td>$0.68$</td>
</tr>
<tr>
<td>$Q_5$</td>
<td>1.04</td>
<td>0.94</td>
<td>0.33</td>
</tr>
<tr>
<td>$Q_{25}$</td>
<td>1.17</td>
<td>1.17</td>
<td>0.84</td>
</tr>
<tr>
<td>Median</td>
<td>1.46</td>
<td>2.96</td>
<td>0.91</td>
</tr>
<tr>
<td>$Q_{75}$</td>
<td>2.21</td>
<td>29.6</td>
<td>0.98</td>
</tr>
</tbody>
</table>

3.4 Discussion of results

3.4.1 Aerosol dynamics and deposition

The simulations have shown that aerosol dynamics can have significant impact on aerosol population depending on particle sizes. It is mainly the condensational growth that can increase or decrease the particle numbers at certain sizes depending on the shape of the particle size spectrum. The aerosol dynamical impact on particle concentration at certain sizes can be equal to or even significantly exceed in magnitude the particle loss due to deposition within the canopy. This is in particular true for particle sizes at which deposition rate is minimal. Consistently with our result, Pryor and Binkowski (2004) and Pryor et al. (2013) have found that frequently the timescales corresponding to particle deposition and aerosol dynamical processes are on the same order of magnitude and therefore induce the concentration change with comparable magnitude. Pryor et al. (2013) evaluated these timescales to be on the order of 1–10 h during the daytime in summer over a pine forest. In the current study we presented that the aerosol dynamical timescale can be from approximately half an hour to tens of hours.
Figure 10. Variation of change velocities with particle size for (a) morning (from sunrise until noon), (b) afternoon (from noon until sunset) and (c) night (sun below horizon) periods: the blue line indicates the storage, magenta the aerosol dynamics and green the vertical exchange velocities. The lines present the ratios of the average change velocities to the average deposition term according to \( \langle V \rangle / \langle V_{\text{dep}} \rangle \) obtained from model simulations for each 10 min period and the shaded areas are the variation range as \( \pm \sigma V / \langle V_{\text{dep}} \rangle \) around the averages.

Figure 11. The exchange velocity \( V_e \) at the canopy top for selected particle size intervals during a 10-day period in May 2013 normalized with the absolute value of the deposition velocity \( |V_{\text{dep}}| \). Peak values for the size range 30–100 nm at DOYs 129 and 130 were about 30–35.

The timescales of turbulent transfer and vertical transport were determined to be essentially different. The vertical transport of aerosols was limited by the deposition and aerosol dynamical processes and only at stable conditions could the turbulent transfer become limiting to vertical transport of particles. The turbulent transfer timescales estimated in the current study by using the LS trajectory simulations were on the order of minutes during the daytime and could be up to a few tens of minutes under SBL conditions. Some other definitions of the timescales have been used in the analysis of the significance of chemical transformation of reactive scalars during transport pathway between the measurement level and sources or sinks located primarily at leaf surface. Rinne et al. (2000, 2012) used the ratio of the observation height to the friction velocity as the estimate for the mixing timescale. Zelger et al. (1997) used the definitions of Eulerian and Lagrangian turbulent timescales to characterize the turbulent transfer within and above forest. Holzinger et al. (2005) instead used the estimate of the residence time and obtained the value about 1.5 min for daytime conditions. The Lagrangian turbulent transfer times obtained in this study were consistent with the previous studies including the timescales obtained by the same approach by Rannik et al. (2009b).

### 3.4.2 Dynamics within ABL

The timescales of aerosol deposition and dynamics are much longer than the turbulent transfer times within the forest canopy. Therefore, one would expect a minor impact of aerosol dynamics on particle population during the vertical transfer within forest under most of the observation conditions and a relatively good vertical mixing of aerosols within and above forest. Nevertheless, we have seen in the current study that the vertical fluxes at the canopy top can deviate significantly from what would be expected from dry deposition only. From current model simulations we have seen that the aerosol dynamics is an important mechanism of aerosol transformation throughout the ABL, whereas the aerosol deposition occurs only inside the forest canopy. In addition, the impact of aerosol dynamics is height dependent. Within the canopy the emissions of the precursor gases for particle condensational growth (the volatile organic compounds) occur. The dominant condensing compounds are the OH oxidation products of monoterpenes, which form during the transport pathway from inside the forest to higher levels in the ABL. The concentrations of the condensing compounds are therefore larger within and immediately above the canopy and decrease with height. Such height dependence of the condensational growth of particles can lead to modification of concentration gradient and vertical flux profile. Even though the atmospheric mixing is fast compared to the above-discussed processes, we believe it is the extensive source/sink term by aerosol dynamics that operates throughout the atmospheric column (compared to the impact of deposition inside the canopy only) and can thus create significant vertical flux divergence and even upward particle transport.

The concentration time change, when summed up from the surface up to the measurement level, is called the stor-
age term and commonly accounted for in estimation of the net ecosystem exchange of carbon dioxide from the EC flux measurements (e.g., Foken et al., 2012). Such an approach inherently assumes that the storage change results from the source/sink activity below the observation level. Rannik et al. (2009a) studied the relevance of the storage term in estimation of the dry deposition from particle flux measurements. They concluded that in the case of aerosol particles the relevance of the storage term could not be established because of the different physical reasons for the concentration change during different phases of diurnal development of the ABL. This study supports the conclusion with the observation that the particle concentration change is primarily in correlation with the aerosol dynamics and the change occurs throughout the ABL. Therefore the particle storage change (which corresponds to accumulation or depletion) is not in general the sole component of the particle conservation equation that could help to improve particle deposition estimation from the flux measurements carried out above forest.

3.4.3 Upward particle fluxes

Particle fluxes determined by the micrometeorological techniques show typically large variability in magnitude as well as in sign. Occurrence of upward particle fluxes has been frequently reported in the literature (Pryor et al., 2007; Grönholm et al., 2007; Whitehead et al., 2010; Pryor et al., 2013). Even after careful classification of observations according to wind direction in order to remove the cases possibly affected by anthropogenic emissions, flux observation analyses by Pryor et al. (2008) revealed significant fraction of observations indicating emission. The upward particle flux values can be the result of large random uncertainty or caused by upward particle transport due to physical processes. Random flux errors of particle fluxes are due to stochastic nature of turbulence, instrumental noise and (limited) counting statistics of aerosol particles. The major source of the random uncertainty of particle flux estimates is the non-stationarity of particle concentration as well as its flux (for flux random uncertainties see Fairall, 1984). The particle fluxes have typically large statistical uncertainty, on the order of 100% and more (Pryor et al., 2008; Rannik et al., 2003); therefore, it is difficult to determine whether the calculated upward particle occurrence reflects the true transport or was obtained by chance. Pryor et al. (2008) investigated thoroughly the distribution and significance of upward fluxes as well as the relevance of several physical mechanisms that cause them by taking into account the error estimates of fluxes. They came to the conclusion that several possible physical mechanisms were responsible for upward particle transport including the entrainment of particle-free air from above during the intensive ABL growth periods. Whitehead et al. (2010) observed similar systematic pattern over a tropical rainforest in case of supermicron particles. Upward particle fluxes were also observed on seasonal average diurnal patterns by Rannik et al. (2009a) in the statistical analysis of long-term particle flux measurements over a pine forest, confirming that the phenomenon is common over a long period of time.

Nilsson et al. (2001) also associated the occurrence of upward particle fluxes to the solar radiation increase and boundary layer development. In addition, they studied the evolution of the Aitken and accumulation mode particle concentrations in the ML during the ABL growth and inferred the particle concentrations being entrained by using a simple ML growth model based on thermodynamical considerations. The model explained well the ML height as well as the particle concentration evolution. The entrained particle concentrations were determined to be virtually from 0 to 40% of the close-to-surface values, indicating that nighttime horizontal advection was a dominating process at the site affecting the vertical profiles of aerosols above the SBL. The initialization of the aerosol concentration profiles during the first day of simulations in the current study represent such advective conditions and resulted in strong upward particle transport during the early morning ML growth. Whereas the nighttime advection can be typical to SMEAR II site, it is certainly a site-specific phenomenon and therefore for the rest of the period we intended to use the initialization of profiles with uniform particle concentration up to the residual layer height. Therefore our simulation results for the first day represent the conditions characteristic to strong horizontal advection and are during the rest of the days expected to underestimate the vertical transport due to ML growth.

Gordon et al. (2011) observed major fraction (60%) of upward particle fluxes for size interval 18–450 nm above a mixed forest in Ontario, Canada, using the EC technique. The upward particle flux rate was highest for 75 nm particles. One of the mechanisms for upward fluxes was the entrainment of clean air from aloft as discussed previously. As an additional mechanism, the authors proposed the slowest growth rate of this particle size, suggesting that the authors referred to the aerosol dynamics as one of the reasons.

Pryor et al. (2013) also suggested the depletion mechanism as the most common cause of the upward fluxes above a sparse pine forest during the morning hours. Later in the day the authors attributed the upward fluxes of sub-30 nm particles to the growth of the newly formed particles by condensation of the biogenic volatile organic compounds (BVOCs). All the mechanisms as the reasons for upward particle fluxes discussed here appear to be the plausible reasons according to our model simulations and can dominate depending on location, emission rates of BVOCs, time of day, particle size and possibly some other factors. The results of the current study identified the aerosol dynamics as one of the main mechanisms causing upward transport of particles with 30 nm in diameter and larger.
3.4.4 Fluxes of above 100 nm particles

Our results have shown that the aerosol and ABL dynamics can introduce significant systematic deviation of the exchange velocities above the canopy from dry deposition on the average. For around 100 nm particles the fluxes above the canopy exceeded the dry deposition sink and for larger than 100 nm the deposition was poorly characterized by the fluxes above the canopy (see Table 1). The range of the flux to deposition ratio varied from negative to positive values, being especially large for particles about 100 nm, which coincides with the minimum of the particle deposition rate at this size. The median values presented in Table 1 were closer to unity than the averages. This implies that the averages are affected by extreme values corresponding to certain dynamical conditions occurring in the ABL. Such conditions certainly can take place in the real atmosphere. The fact that the median exchange velocities represent better deposition than the time average indicates that the median values are more robust statistics than the averages and should be perhaps used in representing the particle exchange instead of averages.

We note that the results based on model simulations were free of statistical uncertainty introduced by random errors to experimentally determined fluxes. Rannik et al. (2003) used a semi-empirical model to explain the size-integrated particle flux measurements performed at the same site with our model simulations. The model appeared to explain well the flux observation with particle population mainly consisting of below 100 nm particles. Deposition velocities for above 100 nm sizes were very uncertain. The authors proposed several reasons why the model was not able to explain the observations: presence of a mechanism controlling deposition of above 100 nm particles not described by the semi-empirical model as well as several other reasons such as temporary pollution sources in the measurement source area. The possible reasons of meteorological origin were suggested to be horizontal advection of particle concentration, boundary layer growth and concentration dilution, and roll circulation in the ABL (e.g., Buzorius et al., 2001). This study has shown that such apparent uncertainty in deposition pattern of above 100 nm particles could be the case even in horizontally homogeneous conditions due to aerosol dynamical and ABL development processes.

4 Conclusions

Simulations performed by the model SOSAA coupling turbulent exchange within the ABL with detailed atmospheric chemistry and aerosol dynamics indicated that the aerosol dynamics is strongly size dependent but a significant source/sink term to aerosol concentration throughout the atmospheric column. Whereas the vertical transport mostly compensates for particle loss inside the canopy due to the deposition, the aerosol dynamics leads to the concentration changes in the whole ABL. However, during the periods of intensive aerosol dynamics when new particle formation frequently occurs, the particle deposition and aerosol dynamics together with ABL development leads to complicated vertical transport patterns. For small particles (up to a few tens of nanometers) the deposition sink is relatively strong (compared to the aerosol dynamics) and the downward fluxes were predicted in the lower ABL. However, for some particle size ranges, depending on the aerosol dynamical processes, the stronger aerosol dynamical source inside and above forest (compared to higher ABL) can lead to upward particle transport such that the vertical fluxes above the canopy might not be coherent with dry deposition under such conditions. We have also observed that the ABL dynamics occasionally lead to upward particle transport which can be interpreted as the transport due to dilution of relatively particle-rich air within forest with the particle-poor air transported down from aloft during the active ABL growth phase.

The simulated turbulent transfer timescales inside the forest were much shorter than the timescales of deposition and aerosol dynamics for all sizes except the smallest at around 3 nm. In spite of efficient mixing inside the canopy, the particle fluxes at the canopy top can deviate from the deposition rates inside forest. This is due to the transformation of aerosol concentration throughout the atmospheric column resulting in the complicated pattern of particle vertical transport. Therefore, the within-canopy deposition and transformation processes do not determine solely the particle vertical transport within and above the canopy and the respective timescales are not sufficient to determine whether the aerosol dynamics can cause significant particle flux divergence below the measurement level.

We conclude that under dynamical conditions studied here the particle fluxes above the forest canopy occasionally deviated from the particle dry deposition sink inside the forest canopy. Such deviations can be very large and for certain particle sizes even systematic after performing diurnal averaging of results.
Appendix A: Description of measurements at SMEAR II

The SMEAR II (Station for Measuring Forest Ecosystem-Atmosphere Relations) field measurement station is located in Hyytiälä, Southern Finland (61°51’N, 24°17’E; 181 m a.s.l.). The station is located in the area covered mainly by pine-dominated forests. The dominant height of the stand (Launiainen et al., 2011). The forest floor vegetation is related low (mean height 1 m a.s.l.). The station is located in the area covered mainly by pine-dominated forests. The dominant height of the stand (Launiainen et al., 2011). The forest floor vegetation is relatively low (mean height ~0.2–0.3 m) but dense (total LAI ~ 1.5 m² m⁻²). However, in model setup a beta distribution of LAD was used that matched to observed turbulence statistics in and above the canopy and the forest floor vegetation as a separate layer was neglected (Boy et al., 2011). A more detailed description of the station and the measurements can be found in Hari and Kulmala (2005).

Turbulent fluxes of momentum, heat, CO₂ and H₂O were measured by means of the EC technique. The system, located at 23 m height above the ground on the top of a scaffolding tower, included an ultrasonic anemometer (Solent Research HS1199, Gill Ltd., Lymington, Hampshire, England) to measure the three wind velocity components and the sonic temperature, a closed-path infrared gas analyzer (LI-6262, LiCor Inc., Lincoln, NE, USA) that measured the CO₂ and H₂O concentrations. The data were sampled at 21 Hz, and a 2-D rotation of sonic anemometer wind components and filtering to eliminate spikes were performed according to standard methods (e.g., Aubinet et al., 2000). The high-frequency flux attenuation was corrected by using empirical transfer functions and co-spectral transfer characteristics (Mammarella et al., 2009).

Aerosol size distribution (from 3 nm to 1 µm) measurements were performed using a differential mobility particle sizer (DMPS) system. The aerosol was sampled from inside the forest at 2 m height. Details of the DMPS measurement system are presented in Aalto et al. (2001).

Appendix B: Initialization of model SOSAA

The chemistry scheme employed by the model for this study included the relevant Master Chemical Mechanism (MCM) chemical paths (Jenkins et al., 1997, 2003; Saunders et al., 2003) for the following parent molecules: methane, methanol, formaldehyde, acetone, acetaldehyde, MBO, isoprene, α-pinene, β-pinene, limonene and beta-caryophyllene. For other emitted organic compounds including myrcene, sabinene, 3-carene, ocimene, cineole and “other” monoterpenes, and farnesene and “other” sesquiterpenes, the MCM chemistry paths are not available and we thus included their first-order oxidations with OH, O₃ and NO₃. For the reactions of the stabilized Criegee intermediates (sCI) from α- and β-pinene and limonene, we used the rates from Mauldin III et al. (2012), similar to “Scenario C” in Boy et al. (2013). For the sCI from isoprene, we used the rates from Welz et al. (2012) as done in “Scenario D” in Boy et al. (2013). Sulfuric acid and nitric acid were removed from the gas phase based on the condensation sinks calculated from background aerosol loading.

There was no specific initialization of chemistry state for the model (all variables were initialized as 0 while created). Exceptions were the passive tracer concentrations (CO₂, NO, NO₂, SO₂ and O₃), which were initialized with measurements. The concentrations of these five passive tracers were always read in from measurements during the simulation. The time resolution of input data was half an hour and the data were linearly interpolated for each time step in the model run. The vertical profiles of the particle concentrations were initialized each night as described in Sect. 2.2.

Global short-wave radiation, top boundary temperature, humidity and wind speed were fixed to inputs throughout simulation. The global short-wave radiation was measured at SMEAR II. Temperature, humidity and wind speed at the top boundary were based on ECMWF reanalysis data.

The initial temperature profile was assumed linear, using the input top border temperature and input temperature gradient. The wind profile was set using the logarithmic wind law, the roughness and wind speed at the top boundary. Initial humidity was taken constant throughout the ABL and the heat fluxes and TKE were set to 0. Mixing length was initialized as $l = 0.40 \frac{z}{0.01z_0}$. At the lower boundary, soil humidity for the uppermost layer was set to 0.2 kg kg⁻¹. Soil temperature was set to −2°C from air temperature at the lowest level and leaf temperature was set equal to air temperature. Heat flux to the soil was based on the measurements from the SMEAR II station throughout simulations.
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