

One year boundary layer aerosol size distribution data from five nordic background stations

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Abstract. Size distribution measurements performed at five different stations have been investigated during a one-year period between 01 June 2000 and 31 May 2001 with focus on diurnal, seasonal and geographical differences of size distribution properties. The stations involved cover a large geographical area ranging from the Finnish Lapland (67° N) down to southern Sweden (56° N) in the order Värriö, Pallas, Hyytiälä, Aspvreten and Vavihill. The shape of the size distribution is typically bimodal during winter with a larger fraction of accumulation mode particles compared to the other seasons. Highest Aitken mode concentration is found during summer and spring during the year of study. The maximum of nucleation events occur during the spring months at all stations. Nucleation events occur during other months as well, although not as frequently. Large differences were found between different categories of stations. Northerly located stations such as Pallas and Värriö presented well-separated Aitken and accumulation modes, while the two modes often overlap significantly at the two southernmost stations Vavihill and Aspvreten.

A method to cluster trajectories was used to analyse the impact of long-range transport on the observed aerosol properties. Clusters of trajectories arriving from the continent were clearly associated with size distributions shifted towards the accumulation mode. This feature was more pronounced the further south the station was located. Marine- or Arctic-type clusters were associated with large variability in the nuclei size ranges.

A quasi-lagrangian approach was used to investigate transport related changes in the aerosol properties. Typically, an increase in especially Aitken mode concentrations was observed when advection from the north occurs, i.e. allowing

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more continental influence on the aerosol when comparing the different measurement sites. When trajectory clusters arrive to the stations from SW, a gradual decrease in number concentration is experienced in all modes as latitude of measurement site increases.

1 Introduction

Many processes in the atmosphere concerning aerosol particles are closely related to the size distribution, and a welldeveloped understanding of the size distribution is required in order to evaluate the effect of aerosols on both climate and human health. The effect on climate is generally divided into a direct and indirect effect. The direct effect comes from the ability of aerosols to scatter and absorb incoming solar radiation, while the indirect effect is a result of the ability of aerosols as acting as cloud condensation nuclei (CCN) and thereby altering the radiative properties of clouds (Twomey, 1974). Also, health problems arising from particulates have been given a greater amount of interest during last years (Künzli et al., 2000).

Most size distribution measurements so far are over a limited time period or only representative for a smaller area. Some work has been performed in this field, though. To be mentioned are recently published works by Birmili et al. (2001) that have sprung out with results that are useful for climatologically modelling. Birmili and co-workers (2001) found, when investigating the aerosol properties in Melpitz, typical size distribution attributable to the air mass where measured. Most pronounced differences were found between air masses of continental and marine origin, with a larger aerosol mass associated with air-masses of continental origin. Studies performed by Kulmala et al. (2000) have come up with similar results for the two Finnish stations Värriö and Hyytiälä. Winds from SW Russia and central Europe brought about far higher loads of accumulation particles as compared with those of arctic origin. Aitken mode was found to vary insignificantly when comparing the origin of air masses. Other long-term studies of the aerosol size distribution are reported from Jungfrauhoch (the Swiss Alps) with emphasis on air-mass origin (Nyeki et al., 1998).

The objective of this paper is to broaden the view of the Nordic aerosol in order to allow conclusions concerning the aerosol properties, i.e. find the seasonal variation for the whole Nordic region through the year of study and investigate how the aerosol varies over the region and establish a view on how long-distant transport effects different parts of the region. To achieve these goals 5 different stations observing size distribution data on timescales of minutes are utilized. The study will focus on data collected during one year at these five stations located in Finland and Sweden, namely Aspvreten (Institute for Applied Environmental Research, Air Pollution Laboratory), Vavihill (Lund University), Hyytiälä and Värriö (University of Helsinki) as well as Pallas (Finnish Meteorological Institute). In order to produce results relevant for modelling and process studies, the conceptual model of lognormal fitting has been applied to the individual scans in order to reveal seasonal as well air-mass history related effects on the aerosol size distribution during the year of study. To fulfil the goal of presenting a dataset with both seasonal and spatial considerations, a clustering model presented by Dorling et al. (1992a, b), have been used to treat a large number of trajectories arriving at the individual stations. This will allow conclusions concerning the effect of source area on the aerosol studied.

2 Methods

2.1 Clustering model

Cluster analysis is a multivariate technique aiming to maximize the variance between different groups and to minimize the variance within a group of variables. A cluster analysis applied to trajectory data may be useful in order to recognize and quantify the influence of synoptic meteorology on the aerosol climatology. The trajectory data itself relies on trajectories calculated with the HYSPLIT4 (HYbrid Single-Particle Lagrangian Integrated Trajectory) model (Draxler and Hess, 1998). For this study 96 h back trajectories where calculated throughout the year for all stations at four times a day (UTC: 05, 11, 17, 23).

Since each individual trajectory is constructed from wind fields, they must reflect the evolution of a synoptic pattern during the 4 days they reach back in time. Dorling et al. (1992a) indicate how a composite surface pressure pattern representing an ensemble of trajectories reveals typical atmospheric circulation features resulting in the transport patterns represented by the individual trajectories. When linking the cluster analysis to the actual size distribution measurements this will prove important. The evolution of the size distribution during a couple of days is strongly dependent on the weather situation and not only the source areas. Notably, cloud processing and washout by precipitation is of crucial importance. In order to make inter-comparisons between the different stations, it is therefore preferential to try to find and evaluate situations when similar meteorological history applies to the air mass measured in. Since the clustering procedure roughly will represent some typical weather situation we will be able to reduce the bias of the size distribution resulting from meteorological variations during the transport, as compared with cases if we choose only to look at the air mass origin in terms of sectors. This will allow us to investigate effects from transport in-between stations, finding the important differences resulting from factors acting upon the size distribution during transport.

The clustering method utilized for these purposes is an approach suggested and thoroughly described by Dorling et al. (1992a, b).

2.2 Model for aerosol size distribution

In order to allow comparisons between size distributions collected within different clusters as well as size distributions collected at different sites some general assumption has to be done concerning the size distribution. An often applied approach is the use of relatively simple equations to describe the aerosol size distribution as separated in one or more distinct modes, i.e. the lognormal aerosol size distribution (Whitby, 1978; Hoppel et al., 1994). This allows straightforward comparisons of the data between different stations by means of a set of well-defined parameters. This approach is well treated in the literature and the overall outcome has proved useful when parameterisations are required (e.g. Birmili et al., 2001; Mäkelä et al., 2000a).

In this study the aerosol size distribution has been interpreted in terms of three modes, nuclei (<30 nm), Aitken (30-110 nm) and accumulation mode (110-1000 nm). These modes are closely related to the processes leading to their appearance. Nuclei mode particles may be formed from condensing gases if some critical concentration is reached. This nucleation phenomenon has been extensively treated in the literature (e.g. Kulmala et al., 2001; Väkevä et al., 2000; Kulmala et al., 1998).

The lifetime of freshly formed particles is short, because of the growth by condensation but mostly due to removal by coagulation with larger particles. Coagulation between nucleation mode particles and condensation of gases onto nuclei mode particles constitute growth mechanisms of nuclei mode into the Aitken mode size range. While this growth by condensation and self-coagulation of nuclei mode particles leads to increased number concentration in the Aitken mode,

Table 1. Boundaries used as input for the automated fitting procedure for size distribution with three modes. Geometrical mean diameter (Dg), geometrical standard deviation (GSD) and number concentration (N) limits are used. Number concentrations between zero-infinity are allowed. Modal diameters may overlap.

	N1 (#cm ⁻³)	GSD1	Dg1 (nm)	N2 ($\#$ cm ⁻³)	GSD2	Dg2 (nm)	N3 ($\#$ cm ⁻³)	GSD3	Dg3(nm)
Lower boundaries	0	1.1	Smallest size of instrument	0	1.1	20	0	1.1	80
Upper boundaries	∞	1.8	60	∞	2.0	120	∞	2.0	Largest size of instrument

coagulation with larger particles might serve as a significant sink for newly formed nuclei mode particles. Transformation of nuclei mode particle mass into Aitken mode mass seems to largely be controlled by direct coagulation between nuclei mode particles and Aitken mode particles without observed growth of the actual nuclei mode.. Transformation of nuclei mode particle mass into Aitken mode mass seems to largely be controlled by direct coagulation between these particles without observed growth of the actual nuclei mode. This would only lead to an increased mass in the Aitken size ranges, leaving the number concentration unaffected. Only situations with favourable conditions (e.g. large nucleation rate and/or small coagulation sink) will lead to an observable growth of nuclei mode particles into the Aitken size ranges, thereby affecting their number concentration (Kerminen et al., 2001). Otherwise, in order to maintain the number concentration in the Aitken size ranges, direct emissions are required. In the literature it is argued that the primary emissions from anthropogenic activities largely contribute to aerosols in the Aitken size ranges (Birmili et al., 2001). Coarse mode particles may, if high enough concentration is present, serve as a significant sink of nucleation mode and Aitken mode particles. However, since coarse mode includes particles $>1 \,\mu m$ (Seinfeldt and Pandis, 1998), no evaluation of this mode can be done due to instrumental limitations. The instruments used typically cover a size range from <10 nm to 500 nm.

Subsequent growth into larger size classes requires a large amount of condensing gases. The most important mechanism transforming Aitken mode particles into accumulation mode particles is cloud processing of the aerosol. Theoretically, this will produce well-defined modes with a pronounced minimum in between, i.e. the Hoppel minima (Hoppel et al., 1994).

Since the goal was to fit each scan separately, some automated fitting procedure had to be developed, taking into account the huge amount of data produced by five stations during one year. In order to produce results with a physical relevance, some constrains had to be supplied to the fitting routine. For this purpose restrictions concerning the geometrical standard deviation (GSD) and mean modal diameter (Dg) given in Table 1 were used.

The fitting procedure is performed with a Matlab® routine involving two functions supplied with the Matlab® Optimization Toolbox: LSQNONLIN for a first crude fit and FMINCON for the final fitting. LSQNONLIN starts with a

first crude approximation concerning modal diameters and number concentration. The output from this first fit serves as first guesses in FMINCON which performs a constrained fit with boundaries as described in Table 1. However, since the number of mathematical solutions to the lognormal function is large, additional constraints have to be applied, especially to avoid super-positioning of modes, which is mathematically sound, but does not contribute with any useful information for this study. No modes are therefore allowed to have a spacing less than dlogDg<0.15, which roughly corresponds to a ratio of Dg's > 1.4. If a sound solution is not found within four iterations, a fitting with two modes is suggested. If the solution for two modes is not found within three attempts, fitting with one mode concludes the fitting procedure. This generally results in two modes, one in the Aitken and one in the accumulation size range, but appearance of a third mode occurs frequently, many times indicating new particle formation. The presence of this third mode is generally confined to midday, indicating the importance of photochemistry in production of nuclei mode particles. However, it should be pointed out that this third mode is not always a nuclei mode, but can be a second Aitken or accumulation mode.

In order to understand this convention we have to be aware of the fact that the nature of the processes contributing to the different modes is not to be considered as discrete, but rather a continuous process. For example we can consider the growth of the nuclei mode due to condensation into the Aitken size range. If an Aitken mode already is present, which of course is the normal case, this growth would lead to appearance of two modes in the Aitken size range. Since smaller particles grow faster (by size) than larger, the two modes would in due time appear as one in terms of physical size distribution properties. In order to take similar situations into account it is necessary to include a term for this growing mode. In the following we thus include the term Aitken 1 and Aitken 2, where Aitken 2 is referred to as the semi-persistent Aitken mode and Aitken 1 denotes the growing mode.

Only very seldom we were able to resolve more than one mode in the accumulation mode size range so this type of definition is limited to the Aitken mode only. Fitting of three modes to the aerosol size distribution is of course an oversimplification. Four or even five modes may also be present as reported by Birmili et al. (1998) and Birmili et al. (2001), and this is not considered in our analysis. However, as shown in Sect. 3.4, three modes are sufficient to reproduce the measured size distribution in this dataset.



Fig. 1. Location of the stations involved in the study.

2.3 Description of the stations

2.3.1 Hyytiälä

The background station Hyytiälä ($61^{\circ} 51' \text{ N} 24^{\circ} 17' \text{ E}$, 180 m a.s.l.) have been in focus in a number of large studies, especially concerning nucleation and new particle formation (e.g. Mäkelä et al., 2000; Nilsson et al., 2001a, b; Kulmala et al., 1998; Kulmala et al., 2001). The station, SMEAR II, (Station for Measuring forest Ecosystem-Atmosphere Relations) is characterized as a boreal forest site, with surroundings dominated by a flora of Scots Pine of about 30 years age. The station is located fairly far from urban pollution sites (Tampere at a distance of ~50 km SW and Jyväskylä ~100 km NE). The station has facilitated particle size distribution measurements since 1996, which have resulted in an extensive database including continuous measurement for over five years.

The instrument set-up consists of two differential mobility analyser systems, producing overlapping size distributions. The sample flow is split into two Vienna type differential mobility analysers (DMA). Two condensation particle counters (CPC) are used to detect the selected particles, one Model 3010 from TSI and one Model 3025 from TSI. The composite instrument set-up produces size distributions between 3– 25 nm overlapping with a size distribution spectrum covering the range 20–500 nm. A full size scan is produced every 10 minutes.

2.3.2 Värriö

The SMEAR I station $(67^{\circ} 46' \text{ N } 29^{\circ} 35' \text{ E}, 400 \text{ m a.s.l.})$ in Värriö is also classified as a background station and situated in the same vegetation as Hyytiälä (in this case a 40 year old Scots Pine forest). The station itself is located at a hill cap. The station is far from any pollution sources, although emissions on the Kola Peninsula give rather strong signals when winds are transporting air from this region (Kulmala et al.,

2000). Also, winds coming from the St. Petersburg area as well as Russia in general may bring elevated concentrations of acidifying gases as well as particulate pollution. The Differential Mobility Particle Sizer (DMPS) system set-up produces a complete size scan between 8 and 460 nm approximately every 10 min. Aerosol concentrations are generally low, on average somewhere in the order of 500 particles per cubic centimetre.

2.3.3 Pallas

The Matorova station at Pallas $(68^{\circ} 00' \text{ N} 24^{\circ} 14' \text{ E}, 340 \text{ m} \text{ a.s.l.})$ is located in the depths of the sub arctic pine forest in the Pallas – Ounastunturi National Park that spans over 50000 hectares. The particle measurements are made by Finnish Meteorological Institute (FMI) and the station itself is a part of the Global Atmospheric Watch (GAW) programme. The proximity to the Värriö station allows a good basis for comparison between these stations when simultaneous measurements are performed. DMPS-system set-up consists of one medium DMA separating the particles according to size, subsequently detected by a 3010 CPC from TSI. Size distributions between \sim 7–490 nm are observed with this set-up.

2.3.4 Aspvreten

The background station Aspvreten $(58^{\circ} 80' \text{ N}, 17^{\circ} 40' \text{ E}, 25 \text{ m a.s.l.})$, is located in Sörmland, some 70 km south west of Stockholm. The station is situated about 2 km from the coast in a rural area covered by mixed coniferous and deciduous forest with some meadows. The influence from anthropogenic activities is small, and the area around the station is sparsely populated. The station is operated by the Institute for Applied Environmental Research, Air Pollution Laboratory, and is a part of the European Monitoring and Evaluation Programme network (EMEP). The instrumental set up on the station, besides meteorology and PM 2.5/10 measurements, consist of a medium DMA along with a TSI 3010 CPC. The instrument set-up observes one size distribution spectrum every 6 min in the size ranges 10–452 nm.

2.3.5 Vavihill

Vavihill $(56^{\circ} 01' \text{ N}, 13^{\circ} 09' \text{ E}, 172 \text{ m a.s.l.})$ is a background station at the top of Söderåsen in Skåne. The surroundings are dominated by grasslands and deciduous forest. The station is situated about 10 km away from the closest small villages and about 20 km from the city Helsingborg. The area of Malmö and Copenhagen, with about 2 million inhabitants is situated about 60–70 km to the SSW. However it is not considered to be influenced by local anthropogenic activities and has facilitated background monitoring measurements since 1984. The station is mainly influenced by SW winds. The instrument set-up consists of two differential mobility analyser systems (Ultrafine Differential Mobility Analyser (UDMA)



Fig. 2. Example of typical nucleation events at Aspvreten (upper frame) and Värriö (lower frame).

and Differential Mobility Analyser (DMA), respectively). The separated particles are detected by a TSI CPC 3025 for sizes 3.1–22 nm and one TSI CPC 7610 detecting particles 22–~900 nm. During the year of study, measurements at Vavihill only covers a portion of the winter and spring period (February–April 2001). Therefore no seasonal variation can be evaluated with this data. Locations of the stations are displayed in Fig. 1.

3 Results

3.1 Diurnal variation

The diurnal behaviour of the aerosol at the different stations has been investigated. Mainly to reveal any local sources influence on the measurement. No indications of local anthropogenic influence were found. Nucleation events were observed at all stations. These occasions are typically characterized by a sharp increase of nuclei mode number concentration around noon. During this year of study, the frequency of the nucleation events has been shown to be largest around springtime, between March–May. This seasonal variation in nucleation frequency has been observed at Hyytiälä earlier (e.g. Kulmala et al 2001, Mäkelä et al., 2000b, Nilsson et al., 2001).

Figure 2 shows example of typical events at the measurement stations Aspvreten and Värriö, revealing the commonly

Table 2. Growth rate and peak concentration during class 1 events at the different stations. Number of events meeting criteria is also given in table during the year of investigation.

	GR(nm/h)	N _{max}	# class 1 events
Aspvreten	2.3	9661	17
Hyytiälä	1.9	7649	22
Pallas	1.6	3047	17
Värriö	2.0	2571	25

observed characteristics of the nucleation phenomenon. The fact that we are able to follow the growth for several hours indicates that the phenomenon occurs on a large spatial scale. Nucleation has earlier been reported to occur during sunny days (Kulmala et al., 2001). Further it was in the present study concluded that nucleation occur in air arriving from N/NW in most of the cases. Nucleation as shown in Fig. 2 is only very seldom observed in air advected from S/SE. Earlier reports stretch the importance of cold air advection and boundary layer height (e.g. Nilsson et al 2001). Also, high concentration of pre-existing aerosol is believed to quench the nucleation due to large condensational and coagulation sink, due to removal of the precursor gases and initially formed cluster respectively.

In Table 2 statistics of the observed nucleation events during the year of study is summarized. In this statistics we



Fig. 3. Time dependent evolution of the size distribution during March–May. Each plot represents the average during four hours intervals for the whole season. Upper frame shows days with nucleation and lower frame median of days with no nucleation.

only include the most pronounced events following a definition by Mäkelä et al. (2000b). The growth rate was found to be largest at Aspvreten, by 2.3 nm/h. Peak concentrations observed during the events was highest at Aspvreten (\sim 9600 cm⁻³ on average) and lowest at Värriö and Pallas. Thus both growth rate and number concentration of newly formed particles is highest at Aspvreten.

The diurnal variability in the shape of the size distribution was investigated. In Fig. 3 the composite time dependent median size distributions for Hyytiälä during March-May are depicted. The data have been divided into 4 h intervals and calculated as the median of the integral of scans during the period of the year.

The data was further divided into one sub-set with days with typical nucleation events and another with days when we did not observe nucleation events. Two features become obvious. First, a systematic diurnal variation is only observed for the sub-set with nucleation. In the case with nucleation events we experience this as an initial increase of small particle in the time interval between 08:00–12:00, even more pronounced during 12:00–16:00. Hereafter the size is shifted towards larger particles during time-steps 16:00–20:00 and 20:00–24:00. This behaviour could be interpreted as growth of the small, initially formed, particles. This since we believe that the nucleation phenomena can be observed over large areas simultaneously. We cannot observe a systematic diurnal variation in the sub-set with no nucleation. The sec-

ond feature comparing the behaviour of the two data sets is the fact that the nucleation sub-set has obviously much less mass associated with the aerosol as compared with the nonnucleating cases (e.g. much larger number concentration in the accumulation mode size range in non-nucleation cases as compared with nucleation cases). This probably reflects the fact that we observe nucleation when we have northerly advection, which is likely to bring rather clean air to the measurement sites.

This typical features comparing statistics concerning nonnucleation days and nucleation days are found at all stations. That is, nucleation events do affect the size distribution in a diurnal fashion. This feature is lacking for non-nucleation days.

The diurnal variation of the integral number per season has also been investigated. In Fig. 4 the results for Aspvreten are given. The data is presented as 1h median concentration for the season studied. Only small, if any, diurnal variation in integral number is found. The variations encountered are typically in the range $\pm 10\%$. 5–95 percentiles are indicated below each plot, showing occasions with maximum number concentration. The highest concentrations, shown by the 95 percentile, do occur during daytime.

Most probably, these peaks in integral number concentration can be attributed to nucleation events, treated in previous section. These peaks are furthermore confined to summer and spring periods of the year, which is in nice agreement



Fig. 4. Diurnal variation of median integral number concentration per season, June–August (JJA), September–November (SON), December–February (DJF) and March–May (MAM) at Aspvreten. Indicated in subplots are 5th-95th percentiles. T.O.D. denotes time of day. Note that the axis scale for the number density begins at 1500 cm^{-3} for the large frames.

with the nucleation inventory established for the different stations.

Figure 5 describes the diurnal variability of integral number recorded at Hyytiälä. In comparison with the Aspvreten dataset, even less pronounced variations for the Hyytiälä data was observed. Largest variations are encountered during summer and autumn, however still below $\pm 10\%$. Neither the winter nor summer median integral number data can be addressed with any diurnal variation. However, investigating the 5–95 percentile ranges, occasions with maximum concentrations are shown to occur during daytime. This especially applies during spring and summer, but same tendencies can be observed during autumn.

Performing the same analysis for Värriö we find only small tendencies to diurnal behaviour for all seasons (Fig. 6). The 95 percentile shows no or weak diurnal variation for all seasons except the spring period, when occasions with very high integral number concentration can be observed during noon. Smallest variations are encountered during winter months December, January and February (DJF). Overall lowest concentrations are found during autumn and winter during this year.

Turning the attention to Pallas (Fig. 7), similar behaviour as for Värriö is observed. Only weak diurnal variations seem to be present. No diurnal behaviour can be addressed to the winter season.

Conclusively, the diurnal variations of the integral number encountered at the stations are small, typically below $\pm 10\%$. This reduces the possibility that the aerosol is affected by local sources. The nature of industry and household activities in the vicinity of the stations typically follows a diurnal pattern (e.g. lack of large industries with 24 h production activity). Thus if local anthropogenic activities do affect the aerosol size distribution, this would be evident as diurnal variation of the aerosol number concentration. Furthermore, nucleation events seem to affect the integral number concentration for at least Aspvreten and Hyytiälä in a diurnal fashion when investigating the 95 percentile of the dataset. These trends are not equally obvious for the Värriö and Pallas datasets. However, the events occur as frequent at these stations as observed for e.g. Hyytiälä during this year of study. Although we do not see any large diurnal variations in the integral number we do observe typical diurnal changes concerning the shape of the size distribution. This is most obvious during spring and is attributed to new particle formation events.

3.2 Variation of aerosol properties between the seasons of the year of measurement

In Figs. 8–11 the seasonal features of the aerosol size distribution found during the year of study are depicted as median size distribution per season. The data have been cleaned from



Fig. 5. Diurnal variation of median integral number concentration per season, June–August (JJA), September–November (SON), December–February (DJF) and March–May (MAM) at Hyytiälä. Indicated in subplots are 5–95 percentiles. T.O.D. denotes time of day. Note that the axis scale for the number density begins at 500 cm^{-3} for the large frames.



Fig. 6. Diurnal variation of median integral number concentration per season, June–August (JJA), September–November (SON), December–February (DJF) and March–May (MAM) season at Värriö. Indicated in subplots are 5–95 percentiles. T.O.D. denotes time of day.

occasions indicating failure of the measurements. For each station and season, the resulting median has been lognormal fitted in three modes. The parameters are summarized in Table 3.

A nucleation day inventory has been established per season for all stations. Nucleation data is grouped following a scale from 0-3 where 1 indicates most pronounced event when growth can be observed for several hours. Event



Fig. 7. Diurnal variation of median integral number concentration per season, June–August (JJA), September–November (SON), December–February (DJF) and March–May (MAM) at Pallas. Indicated in subplots are 5–95 percentiles. T.O.D. denotes time of day.

Table 3. Modal parameters obtained from fitting of the median size distributions per season for each station. Also the frequency of nucleation events is presented. Here given as % days of the data set exhibiting nucleation as described under Sect. 3.1.

Station	Season	N1 ($\#$ cm ⁻³)	DSG1	Dg1 (µm)	N2 ($\#$ cm ⁻³)	DSG2	Dg2 (µm)	N3 (# cm ³)	DSG3	Dg3 (µm)	% Nucl.days
Pallas	Jun-Aug	93	1.71	0.029	282	1.49	0.053	115	1.44	0.182	10
	Sep-Nov	69	1.69	0.036	179	1.43	0.074	230	1.61	0.160	2
	Dec-Feb	_	-	-	152	1.85	0.053	123	1.48	0.212	2
	Mar–May	63	1.54	0.019	157	1.52	0.043	162	1.61	0.168	23
Värriö	Jun–Aug	103	1.51	0.043	541	1.60	0.085	141	1.47	0.191	6
	Sep-Nov	_	-	-	263	1.82	0.073	112	1.46	0.200	7
	Dec-Feb	_	-	-	108	1.54	0.048	201	1.63	0.175	12
	Mar–May	119	1.53	0.028	111	1.30	0.050	266	1.73	0.143	27
Hyytiälä	Jun–Aug	429	1.66	0.034	887	1.58	0.077	327	1.46	0.192	7
	Sep-Nov	227	1.63	0.025	776	1.62	0.068	295	1.44	0.200	7
	Dec-Feb	204	1.80	0.025	389	1.62	0.061	270	1.48	0.200	0
	Mar–May	431	1.74	0.025	849	1.69	0.067	284	1.44	0.200	25
Aspvreten	Jun–Aug	250	1.62	0.035	1178	1.49	0.078	423	1.51	0.205	10
	Sep-Nov	272	1.69	0.035	1429	1.61	0.071	494	1.47	0.220	7
	Dec-Feb	431	1.80	0.026	917	1.75	0.066	377	1.46	0.217	2
	Mar–May	507	1.73	0.033	1319	1.69	0.066	256	1.44	0.217	14

classes 2–3 do not show as pronounced growth as class 1. 0 represent occurrence of small particles, but no following growth is observed (Mäkelä, 2000b). In Table 3 class 1 & 2 events are considered. It should be pointed out that these events do not appear in the median values of size distribution since they are smoothed out by the large amount of data present.



Fig. 8. Seasonal variation of median size distribution at Aspvreten (58.8 N, 17.4 E). Error bars indicate 25–75 percentile range. Different modes from lognormal fitting appear in dashed, solid and dotted curves. % nucleation days indicated in upper left corner.



Fig. 9. Seasonal variation of median size distribution at Hyytiälä (61.51 N, 24.17 E). Error bars indicate 25–75 percentile range. Different modes from lognormal fitting appear in dashed, solid and dotted curves. % nucleation days indicated in upper left corner.

The seasonal characteristics for Aspvreten and Hyytiälä are quite similar as seen in Figs. 8 and 9. The median size distribution for the summer (June–August) and spring (March–May) period both show rather high concentration in the Aitken size ranges. The relative contribution of the accumulation mode is largest during winter. The overall lowest concentrations are encountered during the winter period, and the Aitken and accumulation mode are separated to a



Fig. 10. Seasonal variation of median size distribution at Värriö (67.46 N, 29.35 E). Error bars indicate 25–75 percentile range. Different modes from lognormal fitting appear in dashed, solid and dotted curves. % nucleation days indicated in upper left corner.

larger extent compared to the other seasons. The frequency of nucleation is largest during spring months, but some nucleation bursts where encountered during the summer and autumn months. At Aspvreten we observed the lowest frequency of nucleation events during winter and autumn (see Table 3, DJF period).

For Värriö, the overall largest concentrations are encountered during summer (see Fig. 10). The larger number concentration during the summer months is largely explained by an increase of particles in the Aitken size ranges. For all other seasons, the contribution from the accumulation and Aitken mode is found to be almost equal. A more or less pronounced minimum between the two modes is present during all seasons except summer. It is found as for the other stations that the maximum frequency of nucleation days is encountered during the spring period (March-May). However, a fairly high percentage of nucleation days were recorded at Värriö during the winter. This might be indicative of the closeness to smelters located at the Kola Peninsula. Easterly winds may bring elevated concentrations of nucleating gases or precursors thereof to the station from the smelters contributing to the observed nucleation. The seasonal pattern recorded at Pallas is similar to that at Värriö (see Fig. 11). Largest integral number concentrations are encountered during summer. However, there is a distinct minimum between accumulation and Aitken mode present during the summer The maximum in nucleation days occur during months. spring.

Unfortunately, the Vavihill station could not contribute with enough data to support a seasonal analysis. Data available from the station only covers February–April 2001.

Obviously there is a gradient in integral number when comparing the northernmost stations with the southerly located ones, with maximum aerosol concentrations found at Aspvreten. Although sparse in data, Vavihill tends to show equally high or higher aerosol concentrations as compared with Aspvreten. .

It is also obvious that the largest frequency of nucleation is confined to the spring period (MAM), although nucleation does occur both during summer and autumn, but more seldom (Table 3).

The maximum in integral number concentration is confined to the spring and summer months. This becomes obvious when investigating the daily mean concentration recorded for each station as presented if Fig. 12.

Potential explanations for this might be seasonal changes in air masses, lower rate of incoming solar radiation and thus less new particle formation during the winter period and/or a higher rate of precipitation and overall cloudiness during the winter.

3.3 Variation of trajectory orientation between the seasons of the year of measurement

In order to clarify the seasonal differences in typical advection schemes as well as inter-station related differences in air mass history we have chosen to interpret the seasonal



Fig. 11. Seasonal variation of median size distribution at Pallas (67.58 N, 24.07 E). Error bars indicate 25–75 percentile range. Different modes from lognormal fitting appear in dashed, solid and dotted curves. % Nucleation days indicated in upper left corner.



Fig. 12. Daily mean integral concentration for stations Värriö, Pallas, Hyytiälä and Aspvreten (blue bars) and 7 days running mean (red line).

variation of preferred transport pathways to the stations by means of cumulative trajectory plots, each one specific for both season and station. A selection of these results is presented in Fig. 13.

360 grid cells build up the plot. The colour of each cell represents the number of times a trajectory has passed through that specific grid cell during the season at hand.

Warm colours indicate that a large number for trajectories have crossed the grid cell. As an illustration we have chosen to compare the autumn and spring period between Aspvreten and Värriö.

Two features become obvious when studying the resulting plots. First, there is a strong seasonal variability in the orientation of the trajectories during the year of study. During the



Fig. 13. Seasonal variation in advection schemes. Above: The autumn and spring period for Aspvreten. Colour bar indicates cumulative number of times a grid cell has been crossed by a trajectory. Below: Same as above, but for Värriö.

autumn period a large number trajectories arrive from S/SW. Thus, both stations are likely to experience the sources on the continent, changing the aerosol properties accordingly." During spring the opposite applies, with trajectories arriving from the Arctic/marine environment dominating the transport to the different stations. The other feature is that for at least the spring period the two stations does not share the same advection pattern. The most dominant transport pathway to Aspvreten seems to be SE advection. Another important pathway is SW. The trajectories arriving Värriö are preferentially arrive from NE. These features of the transport to the different stations somewhat limits the possibilities of making straightforward comparisons between the stations. It should be mentioned that both Hyytiälä and Aspvreten compares quite well in terms of seasonal variation of trajectory orientation, and that the largest differences are encountered when comparing these southerly located stations with Värriö and Pallas.

3.4 Linking clustering model with model for aerosol size distribution

In order to investigate transport related effects on the aerosol it is important to find situations when similar large-scale advection situations apply to all stations studied. This will give us the opportunity to evaluate how the aerosol properties are changed, e.g. when the transport distance from the continent and its sources is increasing. Since the ensemble of stations provides reference points with a good spatial coverage such an approach will clearly constitute a good basis for future modelling exercises.

In the following section we present an example of how the cluster analysis and the mode fitting model are linked. We have made use of the Finnish station Hyytiälä, where the data set has been divided into four seasons, June–August, September–November, December–February and March–May. A new set of clusters is calculated for each season. The clustering procedure will separate out trajectories whit air of similar meteorological history arriving to all receptor sites. The size distribution is evaluated as the scans collected during a period of two hours before and two hours after each trajectory arrival (local time). In order to assure stable conditions concerning trajectory orientation and thus similar pressure situation, situations showing large deviations in trajectory orientation on time scales less than 12 h were filtered out.

The size distribution recorded around the arrival of all trajectories building the clusters is collected in this manner. Considering the timescale for the production of a new scan (minutes), this will give a large amount of size distribution data upon which the following analysis is built.

The resulting size distribution for three different cluster orientations arriving Hyytiälä during December to February 2000–2001 have been evaluated. The clusters resulting from the analysis is presented in Fig. 14.



Fig. 14. Representation of trajectory clusters arriving Hyytiälä for the period Sep–Nov. The clusters are here presented as the mean of the trajectories building the clusters. Numbers of trajectories in cluster are also indicated.

Table 4.	Modal parameters as	s derived from lognormal	aerosol model.	25-75th percentiles	indicated.	Data corresponds to	clusters A,	B and
C discus	sed above. Fractional	representation of the mo	de itself is also	indicated.				

	Nucleation mode Median	Nucleation 25–75% percentiles	Aitken 1 mode median	Aitken 1 25–75% percentiles	Aitken 2 mode median	Aitken 2 25–75% percentiles	Accumulation mode median	Accumulation 25–75% percentiles				
Cluster A												
$N (\# cm^{-3})$	586	282-834	938	803-1595	729	523-1080	268	220-340				
DSG	1.74	1.56 - 1.80	1.57	1.51-1.72	1.44	1.34-1.54	1.47	1.36-1.56				
$Dg (\mu m)$	0.021	0.016-0.025	0.033	0.031-0.035	0.053	0.049-0.058	0.176	0.159-0.190				
fraction	0.736	-	0.150	_	0.992	_	0.935	_				
				Cluster B	5							
$N (\# cm^{-3})$	241	139-386	153	97–266	828	535-1305	533	338-842				
DSG	1.80	1.63 - 1.80	1.61	1.45 - 1.70	1.61	1.51-1.73	1.52	1.44-1.61				
$Dg (\mu m)$	0.021	0.018-0.024	0.034	0.032-0.035	0.075	0.063-0.086	0.203	0.178-0.227				
fraction	0.369	-	0.125	_	0.936	_	0.989	_				
				Cluster C	1							
$N (\# cm^{-3})$	1310	428-1755	664	350-1035	733	336-1560	136	93-216				
DSG	1.48	1.38-1.63	1.39	1.28-1.47	1.45	1.31-1.60	1.41	1.31-1.52				
$Dg (\mu m)$	0.016	0.009-0.024	0.034	0.031-0.035	0.045	0.037 - 0.052	0.170	0.152-0.221				
fraction	0.540	-	0.155	-	0.868	-	0.675	_				

The general finding during December–February is that the most dominant transport pathways is southerly oriented. One cluster arriving from north (C) with a total number of 15 trajectories, one from south-west (A) with a total of 12 trajectories and one arriving from south (B) made up by 45 trajectories are presented here.

The numbers of size distributions in the different clusters are 306, 251 and 816 respectively. Each size distribution scan, collected around the arrival of each individual trajectory constituting the cluster, has been treated separately by fitting the lognormal aerosol model to the data. In Fig. 15 the median and 25th-75th percentile ranges of the size distributions observed in each cluster is shown. The median calculated on the lognormal mode fitting procedure is indicated with crosses in the figure. This result is derived from the median of the sum of model-reproduced scans. It is found that in most cases the lognormal mode model is able to reproduce the median size distribution with good accuracy. Only small deviations are experienced for the data resulting from cluster number A, where the fitting of the nuclei mode seems to be lacking in some cases.

In Table 4 the modal parameters obtained from fitting are presented. It is found that the largest nuclei mode concentration is present within the data collected in the marine



Fig. 15. Comparison of SW-cluster (left), S-cluster (middle) and N-cluster (right) resulting median size distribution. The red crosses show results derived from the fitting routine. Fraction of total number of scans accounted for by the model is indicated within brackets above.

cluster C. Further, the modal diameter is comparatively small. The Aitken mode shows largest differences when comparing the geometrical mean diameter, Dg. Typically, continental cluster (e.g. B) show an Aitken mode shifted towards larges sizes. Accumulation mode is found to be present more seldom in the marine case (C) as compared with clusters A and B that are influenced from continental sources to a larger extent. Concentrations found in this size range are also typically lower for the marine aerosol. Smallest accumulation mode diameters are found in the marine cluster (C). GSD's obtained are also typically larger for continental air masses.

This brief presentation of the lognormal size distribution model reveals the most pronounced differences when comparing clusters of different origin. The features presented above is found for all stations in most cases, putting confidence in both the cluster model as well as the simplified fitting methods used.

As a result of more large aerosols present in the continental clusters, these types of clusters also accommodate largest integral aerosol volume (3-450 nm). However, small variations in number concentration are found when comparing different clusters (Table 5). In this and several other cases we even find a lower integral number concentration to be associated with continental clusters as compared to those of marine origin. During the year of study we were able to identify situations when all or at least several stations had resulting clusters with similar orientations. As argued in the chapter dealing with the clustering model, situations with similar ensembles of trajectories arriving to different stations will allow us to evaluate the transport related changes with a reduced bias from meteorological variation. This since the trajectories roughly will represent the wind fields shaping the trajectories, and therefore the synoptic pressure situation. For this purpose a number of interesting cases have been selected to put light on how the aerosol is affected when transport over the Nordic countries occurs. The most dominant transport pathway was found to be SW trajectories. This orientation found representation for all stations during all seasons. However, also situations when other advection schemes applied have been investigated in detail.

In addition to the pre-classification delivered by the fitting routine as described earlier, the data was interpreted according to following conventions: the nuclei mode is confined to the size ranges smallest bin size of instrument -30 nm. Secondly, contribution to Aitken 2 is made by particles in the 30–110 nm size ranges. If three modes are present, although the first one could not actually be referred to as nuclei mode but falls in the Aitken size range, this mode is called Aitken 1. The accumulation mode is made up by particles >110 nm.

3.4.1 Case I, NE clusters

In Figs. 16a and b two occasions with NE cluster to the different station are shown. These situations occurred during the winter period (December–February, Fig. 16a) and spring period (Marc—May, Fig. 16b). During Dec-Feb Värriö, Pallas, Hyytiälä and Aspvreten are represented. During the spring period also data from Vavihill were available, giving an opportunity to follow the aerosol in-between receptor sites widely separated. The number of trajectories building each cluster is given in the figures.

As discussed in Sect. 2.1, clusters will represent a typical synoptic weather situation. A comparison of similar clusters arriving different stations presents similar average pressure fields. In Fig. 17 average pressure fields are presented for the four stations involved in this case study. The averages were calculated over the four day duration of the individual



Fig. 16. NE clusters arriving to the different stations during Dec–Feb (a) and Mar–May (b). Clusters are represented as means. The time spacing between each pair of endpoints (dots) in the figure is 5 h.

trajectories over the MAM-period (Fig. 16b). The different pressure fields present striking similarities as can be seen in the strong gradient over the North Sea and British Isles. The slight differences between the stations arise from the fact that the averages are calculated based on different days due to moving weather systems. This put confidence in the clustering procedure and actually confirms the idea that we will experience a reduced bias from station specific meteorology when comparing the different locations. Sea level pressure data was obtained from the NCEP/NCAR archives.

The resulting size distribution data for the NE-clusters is shown in Figs. 18a and b, both as median as well as the normalized distribution revealing the shape of the derived size distribution. When investigating the cluster data for the DJFperiod we find ensembles of trajectories arriving from northern Russia. The mean clusters have the same source area in the beginning, but do split up in height with the Kola Peninsula, where the air arriving to Aspvreten and Hyytiälä continue SW, while Värriö and Pallas clusters turn W. 27, 21, 13 and 17 trajectories each in the order Värriö, Pallas, Hyytiälä and Aspvreten build the clusters. The trajectories constituting these clusters are of preferentially continental origin.

Turning the attention to the median size distribution resulting from each cluster during December–February (Fig. 18a) several important features are exposed. First to be mentioned is the sharp increase in the Aitken mode observed for both Hyytiälä and Aspvreten as compared with Värriö and Pallas. This increase is most pronounced for Aspvreten. Concerning the accumulation mode median diameter there are obvious similarities between both modal diameter as well as number concentration when comparing Aspvreten, Värriö and Pallas. This feature is lacking for the Hyytiälä data. Investigating the individual trajectories it is found that some trajectories arriving to Aspvreten do cross over the Kola Peninsula as do trajectories arriving Pallas and Värriö, and this is not observed for the Hyytiälä dataset. This area is known for its heavy



Fig. 17. Composite surface pressure pattern for NE clusters presented in Fig. 16a. Surface pressure is presented as average. The averages were calculated over the four day duration of the individual trajectories over the period.

industry, giving rise to large emissions of particles as well as gaseous precursors (e.g. Virkkula et al., 1995 and Kulmala et al., 2000).

Trajectories arriving Aspvreten should further be treated with caution since they in most cases cross the region around Stockholm. Contribution from the city and its sources might affect the aerosol to a very large extent.

Only very small differences between Värriö and Pallas were found to be present when comparing those two clusters. The resulting size distributions are almost identical in both shape and number concentration. This indicates lack of important sources in-between the two stations.

For the March–May period we encountered clusters arriving to the different stations from NE. These clusters originate from the marine/Arctic environment. All clusters at hand use the same "corridor" of transport as shown in Fig. 16b. The clusters are fairly dense and compare quite well in-between stations. The Värriö, Pallas, Hyytiälä, Aspvreten and Vavihill clusters are made up by 22, 13, 33, 16 and 22 trajectories, respectively.

Starting with the resulting size distributions from Pallas and Värriö clusters, it is found that they compare very well both concerning resulting integral number as well as shape of the size distribution (Fig. 18b). The shape of the resulting size distribution is typically bimodal, reflecting the properties of the marine aerosol. Turning the attention to Aspvreten and Hyytiälä almost identical median size distributions is observed, and compared with Pallas and Värriö, a sharp increase in especially the Aitken size ranges are observed. Given the normalized distribution one also find that shape of the distribution is shifted from being dominated by accumulation mode particles to an aerosol with major part of the number concentration confined to Aitken mode. Even further south, at Vavihill, a slight increase in accumulation mode concentrations and a slight reduction in Aitken mode number concentration is observed.

Also, comparing the resulting clusters for Värriö and Pallas during March–May and December–February, clusters arriving Pallas and Värriö during DJF-periods are found to be associated with an aerosol exhibiting markedly different properties as compared with those arriving during March, April and May (MAM). If comparing the differences in mean trajectory orientation for the winter and spring period, the difference is probably a result of influence from the Kola Peninsula. The cases during the DJF-period are associated with an aerosol likely affected by industries on the Kola Peninsula.

Just comparing the median size distribution might be to generalizing and therefore the modal parameters derived from each scan have also been investigated. These parameters, shown in Table 6, are results from the lognormal fitting procedure described in the method part. Fitting these data often two modes in the Aitken range appeared, noted Aitken 1 and Aitken 2. When investigating these data several features



Fig. 18. Comparing resulting median size distribution and size distribution normalized to 1 between different stations with clusters of similar orientation. Dec–Feb (**a**) and Mar–May (**b**) are considered. Size distribution normalized to 1 are given in the right frame in order to highlight the changes in size distribution properties between the stations.

 Table 5. Volume and total number concentration median for clusters arriving in Sep–Nov. 25–75 percentiles and number of scans resulting in the calculated quantity are given.

Cluster orientation	Volume (μ g/cm ³)	Volume (25–75%)	Number $(\# \text{ cm}^{-3})$	Number (25–75%)	Number of scans
A (SW)	1.51	1.06-2.03	1833	1500-2250	251
B (S)	5.57	2.99-9.79	1680	1090-2660	816
C (N)	0.86	0.38-1.42	2159	1459–2980	306

become obvious. First an increase in nuclei mode concentrations is observed when going from north to south in conjunction with increased frequency of appearance of the nuclei mode itself. The maximum nuclei mode concentrations are found at Hyytiälä, although the mode it self tends to appear more seldom as compared with Aspvreten and Vavihill. For the two southernmost stations the fraction of individual scans exhibiting a nuclei mode is as large as 90% compared with 51% and 35% for Värriö and Pallas, respectively. However, a little caution has to be added since Värriö and Pallas do not measure sizes below 7 nm. The Aitken mode number concentration rises steeply when going from Pallas and Värriö to e.g. Hyytiälä and an almost tenfold increase in Aitken mode2 number concentrations can be observed. The frequency of occurrence does not vary as much as the nuclei mode. The Aitken mode diameter seems fairly stable throughout the southward journey. This also applies to the nuclei mode diameter, which is found in the sub 20 nm ranges most of the time. Accumulation mode parameters seem to be fairly stable, although a slight increase in number concentration is

	Nuclei median	Nuclei 25–75 percentile ranges	Aitken mode 1 median	Aitken 1 25–75 percentile ranges	Aitken mode 2 median	Aitken 2 25–75 percentile ranges	Accumulation mode median	Acc. mode 25–75 percentile ranges
Värriö								
$N (\# cm^{-3})$	202	66-682	68	45-132	75	40-233	151	116-205
GSD	1.34	1.25-1.46	1.54	1.41-1.73	1.46	1.35-1.63	1.55	1.47-1.64
$Dg(\mu m)$	0.015	0.012-0.021	0.034	0.032-0.037	0.04	0.035-0.046	0.215	0.191-0.235
Fraction	0.51	_	0.08	-	0.72	-	1	_
Pallas								
$N (\# cm^{-3})$	216	116-522	51	43-516	77	35-421	205	160-275
GSD	1.31	1.25-1.45	1.42	1.34-1.46	1.32	1.27-1.44	1.52	1.41-1.66
$Dg(\mu m)$	0.015	0.012-0.017	0.035	0.033-0.038	0.042	0.036-0.049	0.216	0.173-0.305
Fraction	0.35	-	0.11	-	0.74	-	0.99	_
Hyytiälä								
$N (\# cm^{-3})$	1029	628-2728	705	492-926	800	445-1266	321	208-364
GSD	1.48	1.36-1.77	1.56	1.51-1.68	1.58	1.42-1.76	1.44	1.32-1.67
$Dg(\mu m)$	0.0181	0.01-0.024	0.0327	0.032-0.034	0.0510	0.043-0.058	0.2006	0.163-0.223
Fraction	0.65	-	0.20	-	0.79	-	0.92	_
Aspvreten								
$N (\# cm^{-3})$	489	312-728	181	134-227	631	390-1012	235	171-464
GSD	1.55	1.47-1.72	1.28	1.22-1.49	1.56	1.39-1.81	1.42	1.35-1.6
$Dg(\mu m)$	0.0208	0.018-0.024	0.0365	0.033-0.037	0.0451	0.042-0.053	0.1950	0.161-0.238
Fraction	0.89	_	0.01	_	0.96	_	0.96	_
Vavihill								
$N (\# cm^{-3})$	474	325-786	743	507-888	646	380-964	375	308-461
GSD	1.70	1.38-1.80	1.80	1.72-2.0	1.56	1.41-1.78	1.43	1.36-1.52
$Dg(\mu m)$	0.017	0.01-0.02	0.032	0.031-0.033	0.048	0.041-0.058	0.164	0.149-0.174
Fraction	0.90	_	0.03	_	0.89	_	0.92	_

Table 6. Modal parameter from lognormal fitting procedure for NE-clusters arriving Värriö, Pallas, Hyytiälä, Aspvreten and Vavihill duringMarch–May. 25–75 percentile ranges are indicated.

observed. The fact that the modal geometrical mean diameter, Dg, seems to decrease when going from Pallas and Värriö to Vavihill could probably be explained by direct emissions contributing to the ~ 100 nm size range, reducing the fractional representation of the former and larger accumulation mode particles encountered at northerly located stations.

This data clearly points at the importance of both nucleation as well as direct emissions giving contributions in especially the nuclei-Aitken size ranges, shifting the size distribution when going south. It also becomes obvious that even with short transport distance as present between Värriö and Hyytiälä with sparsely populated areas in-between is enough to significantly alter the properties of the aerosol. This implies that rather fast processes are acting upon the aerosol during this southward transport.

3.4.2 Case II, SW clusters

SW trajectories ought to include transformation of aerosols when moving from extensively polluted areas in the conti-

nent up to clean background locations such as Pallas and Värriö. As mentioned earlier the SW clusters are the ones most commonly encountered. All seasons exhibit this advection situation. As an example of this kind of transport the winter period Dec- Feb has been chosen. For this season we found nice and distinct clusters, probably associated with low-pressure systems arriving from W. The clusters used in the analysis are presented in Fig. 19. All clusters are based on approximately 30 trajectories each. If comparing the clusters between each other one finds that the Pallas cluster is slightly different oriented as compared with the others arriving to Hyytiälä, Värriö and Aspvreten. In comparing the resulting size distributions associated with the clusters between the stations one has to be careful since the trajectories arriving Pallas comes from cleaner environment as compared with the others that most probably are affected by the sources in Europe and Great Britain. This will of course affect the aerosol measured.

Starting the analysis with a visual inspection of the resulting median size distribution and corresponding normalised



Fig. 19. SW clusters arriving to the different stations. Clusters are represented as means. The time spacing between each pair of endpoints (dots) in the figure is 5 h.

size distribution some properties become clear (Fig. 20). First a sharp decrease in all modes is noticed when going from southerly to northerly located stations.

Again, Värriö and Pallas size distribution data is found to agree quit well both concerning size and shape. However, a larger fraction of Aitken mode particles is observed in the Pallas dataset as compared with Värriö (~25%). Another important feature is the change in size distribution shape as the air moves northwards. This should be related to the actual transport time between the southerly and northerly stations, on average 2-3 days derived from the trajectory analysis. The aerosol alters from a size distribution dominated by particles by Aitken mode particles to an aerosol with the larger fraction of particles confined to the accumulation mode. This could have several explanations. First, Aitken mode particles do have a shorter lifetime than accumulation mode particles; therefore one would expect to observe a change in shape of the size distribution in this fashion due to coagulation. Another explanation could be cloud processing moving the Aitken mode particles into the accumulation mode and a subsequent wet deposition by precipitation of the larger accumulation mode particles of the aerosol. The influence from dilution and dry deposition can neither be neglected.

By complementing this first sectional data analysis with further investigations of the modal parameters the picture becomes more obvious (Table 7). Starting with the nuclei mode a decreasing concentration is observed as the distance from the continent increase. Also observed are the quite large modal diameters for the nuclei mode as compared with e.g. the NE clusters encountered during March-May. The concentrations in the mode are much smaller as well. If excluding Pallas from the analysis the fraction of scans associated with the nuclei mode is decreasing with increasing distance from the continent.

The Aitken 2 modal diameters are in the same size ranges for all stations, except for maybe Pallas. The concentration in the Aitken size range decrease sharply. Värriö exhibits approximately 10 times less Aitken mode particles as compared with Aspvreten. The accumulation mode concentration, in turn, decrease by less than 4 times which explains the shape of the median size distribution discussed previously. This indicates that there is a lack of sources to support the high number concentration encountered at Aspvreten when going northwards, and/or strong growth and deposition processes.

Investigating the changes of aerosol properties associated with northerly-southerly oriented airflow, a fast transition into an aerosol with much larger number concentrations associated with measurement sites far south as compared with the northerly-located stations was observed. This in spite of sparsely populated areas in-between the stations. Frequent occurrence of a nucleation mode associated with high number concentrations and increasing Aitken mode concentrations was noticed.

With SW cluster occasions the number concentration for all modes was observed to decrease when moving north. This might be puzzling since one would expect the same sources when going north as when going south. The direction of the airflow would not affect the source strength.



Fig. 20. Comparing resulting median size distribution between different stations with clusters of similar orientation. Resulting size distribution for period DEC-FEB. Size distribution normalized to 1 are given in the right frame in order to highlight the changes in size distribution properties between the stations.

Table 7. Modal parameter from lognormal fitting procedure for SW-clusters arriving Aspvreten, Hyytiälä, Pallas and Värriö from March-May. 25–75 percentile ranges are indicated.

	Nuclei mode median	Nuclei mode 25–75 percentile ranges	Aitken mode 1 median	Aitken 1 25–75 percentile ranges	Aitken mode 2 median	Aitken 2 25–75 percentile ranges	Acc. mode median	Acc. mode 25–75 percentile ranges
Aspvreten								
$N (\# cm^{-3})$	874	486-1803	346	203-851	1150	857-1703	455	386-524
GSD	1.740	1.62-1.90	1.690	1.54-1.90	1.560	1.49–1.67	1.420	1.37-1.52
$Dg (\mu m)$	0.023	0.02-0.026	0.036	0.033-0.052	0.060	0.052-0.069	0.214	0.177-0.240
Fraction	0.431	-	0.264	-	0.977	-	0.965	-
Hyytiälä								
N ($\# cm^{-3}$)	262	169-478	149	90-232	502	329-726	306	232-382
GSD	1.721	1.56-1.80	1.618	1.48-1.77	1.539	1.43-1.65	1.461	1.40-1.55
$Dg (\mu m)$	0.022	0.018-0.025	0.033	0.031-0.035	0.059	0.056-0.065	0.197	0.184-0.215
Fraction	0.614	_	0.159	_	0.988	-	0.997	_
Pallas								
$N (\# cm^{-3})$	84	47-158	45	22-103	152	79–275	161	60-271
GSD	1.465	1.33-1.62	1.404	1.30-1.57	1.533	1.39-1.75	1.505	1.39-1.63
$Dg (\mu m)$	0.019	0.015-0.024	0.038	0.033-0.050	0.049	0.044-0.054	0.182	0.163-0.196
fraction	0.403	-	0.051	-	0.922	-	0.936	-
Värriö								
$N (\# cm^{-3})$	84	31-155	55	29-102	106	56-173	141	82-206
GSD	1.510	1.42-1.58	1.370	1.27-1.52	1.450	1.33-1.66	1.520	1.4-1.62
$Dg (\mu m)$	0.024	0.02 - 0.027	0.044	0.039-0.048	0.056	0.047 - 0.078	0.183	0.161-0.212
fraction	0.141	-	0.395	-	0.767	-	0.984	-

With airflow from the continent, initially high concentrations of particles are present. This would constitute a large sink for newly formed particles as well as precursor gases. In situations with NE clusters the opposite applies. This would favour nucleation in trajectories associated with the NE-clusters as compared with SW-clusters. Another important suggestion that cannot be neglected is the weather situations leading to the formation of cluster specific trajectories in the two different cases. SW trajectories are most certainly associated with low-pressure systems arriving from W. This would probably include frequent precipitation for these trajectories. The fact that we actually have different types of weather situations associated with NE clusters compared with SW might explain the observed features, i.e. more frequent precipitation associated with SW clusters.

An analysis of the precipitation rate along the trajectories indicate up to four to five times more precipitation associated with SW clusters as compared with NE cluster, on average. In absolute numbers, the accumulated precipitation along SW trajectories between Hyytiälä and Värriö was found to be 2.6 mm on average. On the other hand, the accumulated precipitation along NE trajectories as in case 1 (3.4.1), was as low as 0.75. Wet deposition is well known to be a strong removal mechanism for especially accumulation mode, but also for Aitken mode particles.

4 Summary and conclusions

We have performed an analysis concerning one-year aerosol data during the period 01/06/00–31/05/01. The analysis has involved 5 stations ranging from the Finnish Lapland (Värriö & Pallas) down to Vavihill in southern Sweden. All stations have utilized similar DMPS-system set-up. The study has focused on seasonal, diurnal and geographical size distribution properties.

The diurnal variation in integral number concentration was found to be small. No larger variability than $\pm 10\%$ was encountered when investigating the hourly medians of integral number concentration. This reduces the possibility of local influence on the stations. Occasions with very high number concentrations were found to preferentially occur during midday hours though, especially during summer and spring. This was linked to new particle formation events. Nucleation events were observed at all stations, with the largest fraction of nucleation days during spring.

An investigation of the diurnal variation of the size distribution shape was performed. It was found that during nucleation days we observed a characteristic diurnal variation in the nuclei and Aitken size range, with nuclei mode particles appearing in the morning hours. The size distribution is later during the day shifted towards larger size classes. We interpret this as a growth of the freshly formed nuclei mode particles. No diurnal variation in the shape of the size distribution was found for non-nucleation days.

Large differences in aerosol properties were found to prevail when comparing southerly and northerly-located stations. These differences typically found in absolute number concentration and shape of size distribution. Smallest concentrations were encountered at the northernmost stations Pallas and Värriö. However, obvious similarities were observed when comparing these two stations. The southerly stations Hyytiälä and Aspvreten were found to be associated with especially high Aitken mode concentrations. Pallas and Värriö exhibit data with well-separated Aitken and accumulation modes, while we observe a more smoothed size distribution for the southern stations. Concerning the seasonal variation found through the year we experienced a minimum in Aitken mode during winter along with a better-separated Aitken and accumulation mode. Maximum in Aitken mode was experienced during summer and spring. The maximum in integral number concentration is further confined to the spring and summer months.

A trajectory clustering model has been applied to the data per season. Size distribution properties were related to the orientation of the different clusters. A larger fraction of accumulation mode particles was usually found in clusters of continental origin, whereas the marine cluster was found to be associated with a more pronounced Aitken mode as well as larger activity in the nuclei size ranges as compared with those of continental origin. This behaviour was most pronounced for the southerly-located stations.

The same clustering model was used to perform a semi-Lagrangian investigation of the effect on the aerosol from transport between the different stations. With a focus on situations when clusters of similar orientation arrived all stations, the aerosol could be assumed to be affected by similar meteorology at all stations and therefore giving an opportunity to examine the effect on the aerosol in terms of transport distance with a reduced bias from trajectory specific meteorology.

A focus was put on southerly and northerly airflow. It was shown that situations with SW-clusters were associated with a reduction of number concentration in all size ranges, whereas NE-clusters were typically associated with a dramatic increase of aerosols, especially in the Aitken size ranges, when going from one station to another. From this it is clear that strong deposition/dilution occurs when transport from the south applies. In turn, the anthropogenic and natural sources manifest their strength when we follow the transition from marine to continental air masses.

The measurement network seems ideal for evaluating the transport of suspended particulate matter. In the present study we have explored the possibilities of linking different measurement sites in space and time.

We conclude from the study that a constant field cannot describe the aerosol over the Nordic countries. Instead, we revealed typical differences between the stations. These differences were argued to be transport related. We did not performed an in depth study of the influence from different meteorological situations associated with the different types of advection. However, the influence from precipitation was argued important for the lifetime of the aerosol.

In the future this work of course has to be extended and a more detailed investigation of the role of meteorology and sources are called for. This investigation serves as a first step in this process of finding important factors affecting the aerosol over the Nordic countries and to find and supply parameterizations of aerosol processes to be included in regional models. Acknowledgements. This work was funded by the ASTA programme (International and National Abatement Strategies for Transboundary Air Pollution) of the MISTRA research foundation (Swedish Foundation for Strategic Environmental Research), Nordic Council of Ministers (NMR) and the Swedish Environmental Protection agency, Environmental Monitoring Programme. The valuable comments by two anonymous referees are greatly acknowledged.

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