Aerosol from Biomass Combustion in Northern Europe: Influence of Meteorological Conditions and Air Mass History

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Abstract: Alkali-containing submicron particles were measured continuously during three months, including late winter and spring seasons in Gothenburg, Sweden. The overall aims were to characterize the ambient concentrations of combustion-related aerosol particles and to address the importance of local emissions and long-range transport for the atmospheric concentrations in the urban background environment. K and Na concentrations in the PM1 size range were measured by an alkali aerosol mass spectrometer (Alkali-AMS) and a cluster analysis was conducted. Local meteorological conditions and some other data sets were obtained, and back trajectory analyses and chemical transport model (CTM) simulations were included for the evaluation. The Alkali-AMS cluster analysis indicated three major clusters: 1) biomass burning origin, 2) mixture of other combustion sources, and 3) marine origin. Low temperatures and low wind speed conditions correlated with high concentrations of K-containing particles, mainly due to regional emissions from residential biomass combustion; transport of air masses from continental Europe also contribute to cluster 1. The CTM results indicate that open biomass burning in the eastern parts of Europe may have contributed substantially to high PM2.5 concentrations (and to cluster 1) during an episode in late March. According to the CTM results the mixed cluster (2) is likely to include particles emitted from different source types and no single geographical source region seems to dominate for this cluster. The back trajectory analysis and meteorological conditions indicated that the marine origin cluster was correlated with westerly winds and high wind speed; this cluster had high concentrations of Na-containing particles, as expected for sea salt particles.

Keywords: biomass burning; residential wood combustion; aerosol mass spectrometry; potassium; chemical transport model

1. Introduction

Burning of wood and other biomass fuels for heating and cooking purposes is common around the globe [1]. Although biomass burning in fireplaces and wood stoves are such common processes, the knowledge about the influences on health is still fairly limited. Many epidemiological studies...
have demonstrated that ambient particulate matter (PM) may cause health problems [2,3]. Biomass burning and wood smoke from the combustion may pose considerable risk to respiratory and other aspects of human health, e.g., pneumotoxicity and other health problems [4,5]. It is important to characterize the PM exposure from biomass burning and other sources, to understand the health risk of PM. Viana et al. [6] reviewed studies dealing with source apportionment of atmospheric particulate matter in Europe between 1987 and 2007. They showed that studies throughout Europe agreed on the identification of four main source types: 1) a vehicular source, 2) a crustal source, 3) sea spray, and 4) a mixed industrial/fuel-oil combustion and secondary inorganic aerosol. Other sources, such as biomass combustion, were rarely identified with the methods traditionally employed, even though they may contribute significantly to PM levels in specific locations [7-12].

The most commonly used tracers for biomass burning are levoglucosan (LG) [13-19] and potassium [15,16,18]. LG is the product from pyrolysis of cellulose, and potassium, as one of the main nutrient elements, is released from the biomass during burning. A difference between the two tracers is that LG is only emitted during the ignition stage, as LG breaks down into smaller molecules at high temperatures [20,21] while the emission of alkali remain during the whole burning process. Aerosol mass spectrometers (AMS) provide the aerosol composition in ambient air and allow for studies on time-scales that are relevant for aerosol processes in the atmosphere [22-26]. We have developed an Alkali AMS technique based on surface ionization (SI) of particle-bound components [27,28]. Briefly, with this method, aerosol particles are vaporized on a hot metal surface, and the SI technique efficiently ionizes elements with low ionization potentials. Highly sensitive and selective detection can be achieved, and characterization of the alkali metal content in individual particles with diameters down to 14 nm has been demonstrated [27]. The Alkali AMS has previously been applied in field studies in urban air [29] and studies of large-scale biomass combustion [30,31]. The single-particle characterization significantly improves the chances of on-line identification of combustion aerosol and sea spray particles in ambient air. Long-time series of aerosol measurements from residential wood combustion and other solid fuel combustion sources are rare. In comparison to the use of other tracer substances, the alkali detection has the advantage that the compounds remain stable in the atmospheric environment — oxidation and UV irradiation will not change them. In contrast, the common marker compound, levoglucosan has the disadvantage that one needs to consider its reactions in the atmosphere [32,33].

This work describes the results from continuous AMS measurements of alkali-containing submicron particles in Gothenburg, Sweden, during three months period from February 16 to May 20, 2007. The overall aims are to characterize the ambient concentrations of combustion-related aerosol particles during a relatively long period, including the winter and spring seasons, and to address the importance of local emissions and long-range transport for the atmospheric concentrations in the urban background environment. The AMS results are compared to meteorological data, cluster analysis of back trajectories, and results from the EMEP MSC-W chemical transport model [34] to relate the observations to air mass origin and emission source categories.

2. Methods

2.1 Measurement site

The studies were performed in the city of Gothenburg, which is located on the west coast of Sweden and has a population of about 600,000 people. The AMS measurements were conducted at the atmospheric science center laboratory at the university campus (57.6911°N 11.9780°E). The campus is in close proximity to the downtown area and may be characterized as an urban background site. The laboratory is situated on the 4th floor and has access to a roof immediately outside the laboratory windows. It is facing a relatively wide street canyon with a two-lane paved local road. The sampled aerosol (1.0 L min⁻¹) was directed to the inlet of the AMS instrument through a 1 m long copper tube (i.d. 0.9 cm) on the outside of the wall and connected to 0.75 m of conducting tubing (i.d. 0.6 cm) inside the laboratory.
2.2 Alkali aerosol mass spectrometer measurements

The Alkali AMS used for on-line measurements of alkali-containing particles has been described in detail elsewhere [27,28]. Briefly, aerosol particles are drawn into the vacuum system of the AMS through a standard aerodynamic lens system [35,36], which produces a sharply focused particle beam with high transmission of particles into the detection unit [27,37]. The upper particle diameter limit is about 1 µm and the transmitted aerosol mass corresponds approximately to PM1 (total mass concentration of particles with a diameter < 1 µm) in the same way as for other widely used AMS systems [37,38]. The lower particle size limit is below 20 nm, which is lower than for the Aerodyne AMS because of a shorter distance (70 mm) from the outlet of the lens system to the ionizing unit in the present instrument.

The particle beam is directed onto a hot platinum (Pt) vaporizer in the detection chamber of the Alkali AMS [28]. The Pt vaporizer is designed as a small open box (3×3×5 mm), and the construction avoids problems caused by particle bouncing effects [27]. Individual particles decompose in contact with the hot surface, and the alkali metal content of the particles desorbs in ionic form from the metal surface, a phenomenon known as the surface ionization (SI) described in Ionov [39] and Zandberg [40]. The ionization probability is determined by the difference between the work function of the metal and the ionization potential of the desorbing compound. The alkali metals have unusually low ionization potentials, and the ionization probability approaches 100%, while desorption in neutral form dominates completely for other type of elements. During the experiments, the Pt surface was heated to a temperature of 1500 K, and the decomposition of an individual alkali salt particle on the hot surface typically resulted in the emission of alkali ions during 0.1–1 ms [27].

Particles consisting of alkali compounds that rapidly decompose and desorb at 1500 K are efficiently detected, including sea spray particles and typical aerosol particles produced by biomass and coal combustion. Combustion of biomass and coal results in a significant release of alkali-containing compounds, which subsequently re-condense into particles as the flue gases are cooled [27, 28, 31]. In contrast, strongly bound compounds including alkali silicates and aluminates that may be found in mineral dust and ash particles would require a considerably higher Pt vaporizer temperature to be efficiently detected by the Alkali AMS.

An orthogonal acceleration time-of-flight mass spectrometer (oa-TOFMS) was used to measure alkali ion mass spectra for individual nanometer-size aerosol particles [28]. One thousand consecutive 13.5 µs mass spectra scans were added and stored every 13.5 ms by a computer-controlled multi-channel scaler (FAST ComTec P7882, 200 MHz discriminator). A total of approximately 6-108 mass spectra were generated during the field campaign. The majority of the mass spectra only contained a low background signal, while occasionally a millisecond burst of ions was detected from a decomposing particle. The count rate of alkali-containing particles was typically low enough to make particle coincidence negligible, and the data set thus contains information about the Na and K content of single particles. The AMS was calibrated, and its operation was regularly controlled using laboratory-generated particles with known alkali salt content following the procedure described by Svane et al. [28].

2.3 Supporting measurements

The Environmental Office in Gothenburg provided trace gas, PM2.5, PM10, and meteorological data with a temporal resolution of 6 min. The measurements were performed 2 km from the campus at the urban background station “Femman” in the city center (57.7085° N; 11.9701° E), which is situated on a building with a flat roof and a height of 25 m. Additional meteorological measurements (Vaisala Inc., Model WXT510) performed at the AMS measurement site agreed well with the data from the Environmental Office.

2.4 Cluster analysis of mass spectra

The Na and K peaks in individual mass spectra were integrated, and mass spectra with zero signals for both ions were discarded. This reduced the data set to 9·10^7 mass spectra corresponding
to individual aerosol particles with a detectable Na and/or K content. The large size of the data set makes traditional clustering analysis techniques inapplicable, or at least highly impractical, and it was necessary to apply an alternative method. Three data clusters were initially qualitatively identified based on a density-based map of the complete data set indicated in Figure 1. In the next step, the density-based clustering approach DBSCAN (Density-Based Clustering of Applications with Noise) approach was used to extract the desired clusters and normalize them into templates [41,42]. The density-based clustering approach utilizes a local cluster criterion, and clusters are defined as regions in data space where the objects are dense and remain separated from one another by low-density regions [42]. DBSCAN can describe arbitrarily shaped clusters, and the method has been applied to AMS data and compared with the ART-2a method [41,60]. Finally, the boundaries of the three clusters were described by non-linear functions, as illustrated in Figure 1. A fraction with both low Na and K content was also discarded to avoid potential influence of noise in the data (Figure S1). A more detailed analysis of daily data sets shows that Cluster 1 and 3 are consistently described by the clusters indicated in Figure 1. On the other hand, Cluster 2 with both high K and Na content includes several overlapping sub-clusters (examples can be found in Figure S2), which suggests that this factor include contributions from a group of sources. As described below, Clusters 1 can be associated with residential wood combustion, Cluster 2 with a mixture of other combustion sources, and Cluster 3 with sea spray.

2.5 Air mass back-trajectories

Backward 72h trajectories ending 500 m above ground level in Gothenburg were calculated using the HYSPLIT4 model [43]. New trajectories were started every 6 h, giving a total of 376 trajectories for the entire measurement period. The trajectories were analyzed by cluster analysis using the built-in analysis module of the HYSPLIT program. The clustering process is an iterative process wherein each step, the two closest clusters (trajectories) are paired until all trajectories are collected in one cluster [43]. The change in total spatial variance is calculated for each clustering step, and a large change in spatial variance indicates that two relatively different clusters (trajectories) have been paired. Large changes in variance occurred in the formation of 5 and 10 clusters, which suggested that solutions with 6 and 11 clusters provided useful solutions in the clustering analysis. Results from the smaller six-cluster solution are used in the present study.

2.6 Chemical transport modelling

The EMEP MSC-W chemical transport model (CTM) [44,34,45] was used to track influences of different emission sources that may contribute to the observed K and Na concentrations in Gothenburg. The model is used in many different applications and includes treatment of the most important gases for describing photochemical ozone formation and secondary production of aerosol particles from anthropogenic and biogenic sources [46-50]. Primary emissions of particles from anthropogenic sources, sea salt, and wind-blown dust are also included in the model. Emissions of alkali compounds are not explicitly included in the model (except Na from sea salt). In the present work, regional-scale simulations were performed with the EMEP MSC-W CTM using a model domain that covers all of Europe and large parts of the North Atlantic Ocean and the Arctic area. The EMEP model results used in this study are mostly taken from the model simulations described by Bergström et al. [51]. The EMEP MSC-W model was run with a relatively coarse resolution; each grid point covered an area of about 50 km × 50 km, and the lowest model layer was ca. 90 m thick. This limits the ability to model the impact of local emission sources in detail since the emissions will be diluted over an area of ca 2500 km². The model will thus underestimate impacts from point (and small area) sources near the measurement site. The anthropogenic model emissions were handled as in Genberg et al. [48]; primary carbonaceous aerosol emissions from residential combustion were taken from the new emission inventory by TNO [8]; other primary carbonaceous aerosol emissions are taken from the EUCAARI inventory [8]; other anthropogenic emissions were taken from the standard EMEP emission inventory [34, 52]. Emissions from open biomass fires (both wildfires and agricultural burning) were taken from the Fire Inventory from NCAR version 1.0
(FINNv1) [53]. Natural emissions of biogenic volatile organic compounds from vegetation and sea salt from the oceans are calculated within the model [34].

The temporal distribution of the emissions is described in detail by Simpson et al. [34]; for small scale non-industrial combustion (SNAP-sector 2; dominated by residential combustion) the day-to-day variation of the emissions are based on the heating degree day concept, which means that the emissions depend on the daily temperatures in the respective emission areas [34].

3. Results and Discussion

3.1 Overview of measured K and Na concentrations

Figure 2 shows an overview of the Na and K concentrations measured during the period from February 16 to May 20, 2007, together with meteorological, PM10, and NOx data. The Na and K concentrations are typically in the range 1-50 ng m\(^{-3}\) and show large variability on the time scale of a few hours for both elements with peak concentrations above 100 ng m\(^{-3}\) for Na. The time series indicate that the two concentrations are correlated during some periods, but this is not generally true and the Na:K ratio usually varies in the range from 0.1 to 20. The alkali concentrations do not appear to show a general trend related to seasonal changes, and the reasons for the observed concentration variability are instead to be sought in the changes in atmospheric conditions and source strengths.

3.2 Influence of local meteorological conditions

The meteorological conditions changed from winter to spring during the measurement period, and the monthly average temperatures were 0.5, 6.0, 9.1, and 12.0 °C during February, March, April, and May, respectively. The temperatures were close to the normal monthly averages for February and May while March was 4 °C and April was 2 °C warmer than normal. Daily temperatures were below zero during several days in February, and peak day-time temperatures were in the range of 15 to 20 °C during some of the warmer periods. Ground-level temperature inversions are well known to have a strong influence on air quality in Gothenburg [54,55] and several strong inversions and associated periods of poor air quality occurred in March, while the effects of inversions on air quality were less important during other periods.

Figure 3 shows that the local wind speed has a relatively strong effect on the mean Na and K concentrations. Na concentrations increase considerably with wind speed above 3 m s\(^{-1}\), while the highest K concentrations are observed with wind speeds below 3 m s\(^{-1}\). Figure 4 illustrates that the wind direction also has an important effect on the concentrations of alkali-containing particles. High concentrations of Na-containing particles are observed with westerly winds from the nearby sea, while the highest K concentrations are observed for easterly winds from the inland region. Figure 5 shows that local temperature also has a substantial impact on the K concentrations. The concentrations of K (and Na) increase with decreasing temperature below 2 °C. The observed correlations between high Na concentrations and high wind speeds and westerly winds strongly indicate that the observed Na-rich particles mainly consist of sea salt particles. High K concentrations are, on the other hand, favored by low temperatures, low wind speeds and winds from the inland, which suggests that they may originate from biomass burning in the nearby region during cold periods. It appears that the local meteorological conditions have an important impact on the concentrations of Na and K in PM1. However, the observed concentrations show large variations under similar meteorological conditions, which indicates that regional production of particles may not be sufficient to explain the observations.

3.3 Influence of air mass history

To further evaluate the origin of the detected particles, back trajectories of air masses reaching Gothenburg were calculated and analyzed by group analysis. The results from a six-group solution are presented in Figure 6, where the traces for individual back trajectories are indicated. Figure 7 shows the resulting mean trajectories for the six group solutions and the relative contribution to each cluster in percent. Some of the clusters consist of trajectories with a complex pattern while some
were more directed from a certain region. The individual clusters are attributed to different regions of origin for the air masses: Central (C), Eastern (E), Northern (N), and Western Europe (W), as well as clusters with Atlantic (A) and Polar (P) origin.

Table 1 summarizes mean values for the alkali, PM10, and trace gas concentrations depending on the type of air mass reaching Gothenburg together with mean meteorological data. These data sets are only included from time periods where at least two consecutive air mass trajectories are attributed to the same group type. The highest mean Na concentration and the highest Na:K ratio are observed in air masses with an Atlantic origin (trajectory cluster No. 5). The highest mean K concentrations and lowest Na:K ratios are observed when the air masses have an Eastern or Central European origin. In these cases, the observed Na:K ratios are consistent with alkali-containing particles originating from solid fuel combustion processes [30,31]. NO, NOx and PM10 concentrations are also high for air mass types 1 and 2 compared to other types of air masses. The air masses originating from Northern and Western Europe, as well as from the Polar region appear to consist of local polluted air with different degrees of marine character.

To further address the importance of regional emissions and long-range transport, results from a short episode (March 24 to April 2, 2007) are illustrated in Figure 8 and 9. The period is characterized by large daily variations in temperature and relatively strong night-time temperature inversions. Figure 8a shows that the air masses reaching Gothenburg during this period originated in Eastern Europe, while the air mass origin shifted before and after this period. In addition to solid fuel combustion, fires are common in Eastern Europe during this period of the year due to agricultural burns and may potentially influence air quality in Northern Europe during short episodes. Figure 8b shows the distribution of fires in Europe (at 1 km resolution) during the short episode, where each dot on the map indicates a fire detected with the moderate resolution imaging spectroradiometer (MODIS) aboard the Terra satellite [56].

Figure 9a shows a close up of the measured Na and K concentrations together with wind direction (Figure 9b) and wind speed (Figure 9c) data during the same period as covered in Figure 8. The K concentration was relatively high during this period, while Na was low during most of the period. An interesting effect was that both K and Na appear to follow a diurnal pattern during this particular period, an effect that was not observed during other parts of the measurement campaign. For the K concentrations, often reached its highest values before noon, while Na peaked during the evening. The wind direction data (Figure 9b) indicates that the increase in Na concentration was related to a weak sea breeze that was observed daily in the afternoon and evening. The wind speed increased slightly during the sea breeze, but it was fairly low, and the Na signal may be attributed with limited formation of sea salt particles in the region near Gothenburg. The modulated K signal was likely due to a combination of long-range transport and regional emissions from biomass burning that were trapped near the ground during the relatively cold nights with strong night-time temperature inversions.

The results in Figure 10 indicate that the concentrations of Na and K are in general correlated in spite of occasional variations. Cluster 1 has a closer linkage in the winter season while Cluster 2 has no apparent seasonal trend coupling. This agrees with the temperature dependence of K mass concentrations illustrated in Figure 11a, which shows that low temperatures favor Cluster 1, while Cluster 2 shows no clear dependence on temperature. They are both favored by low wind speeds (Figure 11b).

3.4 Comparison between Alkali AMS and chemical transport modeling results

The experimental data are compared with results from model simulations with the EMEP MSC-W CTM, with special emphasis on the influence of biomass burning and fossil fuel combustion. Figure 12 shows the measured total concentration of PM2.5 at the urban background station Fenman and the modeled total concentration of carbonaceous PM from biomass burning (the sum of organic aerosol and elemental carbon from residential biomass combustion and open biomass fires). During the period 17 February – 20 May 2007, the measured hourly PM2.5 concentration varies from 0 to about 50 µg m⁻³, and the highest concentrations occur during a handful of major
peaks. The highest PM2.5 concentrations were observed during the period 23 March – 2 April (two concentration peaks) – at the same time the modeled concentration of particles from open fire and residential biomass burning is also high. There is a relatively high degree of correlation between the measured PM2.5 and the modeled biomass burning PM during the campaign period ($r = 0.81$, for hourly, moving 24h-mean concentrations; $r = 0.72$ for hourly concentrations). The modeled biomass combustion PM may constitute a considerable fraction of the total PM2.5 during some episodes. The average modeled biomass carbonaceous PM during the period is about 1.3 µg m$^{-3}$, which is about 12% of the average measured PM2.5 concentration (11 µg m$^{-3}$).

The measured K and particle number concentrations in the two combustion-related components (Clusters 1 and 2) are also compared to the results from the EMEP MSC-W model simulations. Since K is not explicitly modeled the comparison is based on model results for other particulate components that can be tracked to different source types. Model results for primary emitted fine particles (PPM2.5) from anthropogenic sources and open fires are used (Figures 13 and 14). The modeled PPM2.5 is split into PPM2.5 from (i) residential biomass combustion, (ii) open vegetation fires (including both wildfires and agricultural burning), and (iii) other anthropogenic sources (predominantly fossil fuel sources).

Figure 13 shows a comparison between modeled concentrations of PPM2.5 from biomass combustion (residential + open fires) and the measured K concentration in cluster 1. Overall the modeled concentrations show a high degree of correlation with experimental data. The main exception is a cold and windy 6-day period in February, where the model produces significantly lower values (for the first five days). The meteorological conditions during this period result in a high demand for local residential heating, and we attribute this discrepancy to an underestimation of contributions from local sources – the total PM2.5 concentrations at the urban background station Femman were low during the first four days of the “cluster 1 episode”, and this indicates that source(s) of the elevated K-concentrations measured with the AMS may have been located relatively close to the university site and possibly only impacting a relatively small area; the wind direction was rather stable during the whole episode. Local episodes of this type are difficult to model accurately with a large scale chemical transport model due to the low spatial resolution of the model simulations and emission data. The model simulations also confirm the importance of contributions from open fires, in particular, agricultural burns, in late March (compare with Figs. 8 and 9).

Figure 14 illustrates the modeled concentration of PPM2.5 from fossil fuel combustion and the measured K concentration in cluster 2. The correlation between the model and measurements is lower for cluster 2 than for cluster 1 — this is not surprising since the model PPM2.5 (from other sources than biomass burning) represents sums of many different sources, and the K emissions are likely to be different for different emission sources. However, several features in the experimental data are qualitatively represented by the model results.

A separate set of model simulations were performed to investigate the correlation between the AMS measurements and individual primary particle emission categories. The correlation coefficients between measured concentrations and modeled PPM2.5 from different emission sources are given in Table 2. The measured concentrations in cluster 1 are most strongly correlated with modeled biomass combustion particles, with a maximum $r = 0.76$ observed for total PPM2.5 from biomass combustion (residential combustion + open biomass burning). The correlation with modeled fossil fuel components is relatively low. In general, modeled concentrations are somewhat better correlated with measured number concentrations than with K concentrations. Although the measured Na concentrations are very low in cluster 1, the correlation (not shown) between these and the modeled concentrations are as good as for K for most components. Based on the relatively high correlations observed, we conclude that the cluster 1 particles observed with the Alkali AMS are mainly emitted from biomass combustion.

The correlation coefficients between the measured concentrations in Cluster 2 and various model components are also given in Table 2. Cluster 2 corresponds less well to any modeled compound than cluster 1 does; the highest correlations are found for emissions from the
Agricultural sector (SNAP-10; not including Open Burning of Biomass, \( r=0.65 \) when compared with the cluster 2 number concentration) and for the sum of all “non-biomass” emission sources (\( r=0.63 \)). Correlation is especially low for particles from residential biomass combustion (\( r<0.23 \)). This indicates that cluster 2 is at least not dominated by particles from residential biomass combustion in the region around Gothenburg. For cluster 2 the correlation coefficients are similar for particle number and K concentrations, while they are lower for Na concentrations (not shown; \( r<0.36 \) for all model components in Table 2 except for residential/non-industrial fossil fuel combustion, for which \( r=0.61 \) for Na in cluster 2). The lower correlations for Na have not been studied in detail, but one possible explanation may be interference from sea salt particles at the defined border between Cluster 2 and 3 (see Fig. 1).

To investigate if emissions from any single geographical region could be responsible for a large fraction of the cluster 2 particles, simplified model simulations, that track primary particle emissions from a number of areas in Europe (Sweden, Norway, Poland/Czech/Slovakia [PCS], Eastern Europe [EEU = Russia, Belarus, the Baltic states, Finland, Ukraine] and Western Europe [WEU = Denmark, Germany, BeNeLux, France], the British Isles [UK, Ireland]), were performed with the EMEP MSC-W model. These simulations indicated that different cluster 2 peaks correspond to different source regions; no single region seems to dominate cluster 2. The highest measured concentrations in cluster 2 (late April) seem to be associated with air masses from Western Europe. The broad peak at the end of March is associated with air from Eastern Europe and for the later part also with more southerly air that has passed Poland and other parts of Central Europe; Swedish emissions also influenced Gothenburg during this period. Several other peaks can be related to emissions in PCS, WEU, EEU, and Sweden.

We conclude that Cluster 2 seems to represent emissions from different emission sources and regions that contribute to the air quality in Gothenburg by long-range transport.

### 3.5 Comparison with earlier studies

The present results agree with measurements performed in Gothenburg during 2003 and 2004 with the Alkali AMS that only monitored one type of alkali ion at the time [29]. Mass concentrations of alkali in ambient air varied in the range 0.02–100 ng m\(^{-3}\) and the number of alkali-containing particles varied between 0.1 and 100 cm\(^{-3}\). The detected aerosol was concluded to be dominated by emissions from combustion of biomass and fossil fuels, with a significant contribution from sea-salt particles only during intrusion of marine air.

A long term measurement of Na and K in an urban and marine environment by Ooki et al. [57] reported an average K:Na ratio of 1.8 for the fine particles (\( D < 1.1 \mu m \)), the study concluded that the primary source of the fine particles was a domestic refuse incineration plant located within 15 km of the measurement site. This finding, K:Na ratio of 1.8, by Ooki et al. [57] agrees with the K:Na ratio of 1.2–1.7 reported by Mamane [58] from the direct household refuse incinerator aerosol measurements with the fine particles (\( D < 2.5 \mu m \)). In the present work, the air masses originating from western Europe (W) had an average K:Na value of 2.2, which may suggest the emissions were associated with refuse combustions. A slightly higher value of K:Na than Mamane [58] can be explained by different combusted source materials. In European countries, the usage of both small and large scale biomass burning systems are common; especially during the cold winter period, usage of domestic heating systems with biomass fuel may contribute to the higher K:Na ratio.

Stohl et al. [59] reported measurements of Arctic air pollution due to biomass burning associated with agricultural fire activities in Eastern Europe during the spring; this type of fire smoke can reach the Arctic region and potentially reduce the snow albedo and change the rate of snow/ice melting. The filter samples from the Zeppelin mountain station in Svalbard indicated elevated concentrations of levoglucosan, K, and CO for the biomass burning pollution events from April 27 to May 9, 2006. These results suggest that the monitoring of K and CO can be used as a good indicator for the potential contribution of air pollution from biomass burning in the Arctic region. Since the SI-AMS has single-particle analysis capability, it has further potential to obtain more detailed information.
4. Conclusions

Aerosol mass spectrometry has been used to measure the K and Na concentrations in submicron particles in Gothenburg, Sweden, during winter and spring seasons. Typical PM1 alkali concentrations were in the range from 1–50 ng m$^{-3}$, but with large variations in concentrations and Na:K ratio depending on the prevailing conditions. High concentrations of Na-containing particles were favored by westerly winds and high wind speeds and were concluded to be related to sea salt particles in marine air masses. High concentrations of K-containing particles correlated with low temperatures and low wind speeds and were mainly related to regional emissions and transport of air masses originating from continental Europe. Low Na:K ratios suggested that these particles originated from solid fuel combustion, including biomass burning. Diurnal variations in both K and Na concentrations were observed during a springtime period with daily temperature inversions. The K-rich particles were to a large extent transported from areas with extensive agricultural burns in Eastern Europe during the beginning of this period, during the later part residential biomass combustion emissions also made significant contributions. Further development and application of the methodology presented in this study, including analysis of single particle data may improve the ability to distinguish particles from different sources. To implement any regulatory approach to improve air quality, one needs to have a clear understanding of the emission sources. Alkali-AMS measurements can be a useful tool to identify contributions from biomass burning emissions to submicron particulate matter.
Figure 1. Cluster boundaries in logarithmic scale, where the boundaries were defined based on clustering analysis of Na and K content of individual aerosol particles.
Figure 2. Concentrations of Na and K in submicron particles as a function of time measured by the Alkali AMS, together with temperature, and PM10 and NOx concentrations measured at the Femman station. Displayed values are averages over 6 min intervals.
Figure 3. Concentrations of Na and K in Clusters 1–3 as a function of time. Displayed values are averages over 6 min intervals. Cluster 1 is associated with residential biomass combustion, Cluster 2 with multiple combustion sources, and Cluster 3 with sea salt in marine aerosol. See text for further details about the analysis and interpretation of Clusters 1–3.
**Figure 4.** a) Na and b) K concentrations as a function of wind direction (Femman station) for all data obtained during the measurement campaign.
Figure 5. a) Na and b) K concentrations as a function of temperature for all data obtained during the measurement campaign. Error bars correspond to ±1 standard error.
Figure 6. Back trajectory group analysis for six air mass types reaching Gothenburg during the period from 16 February 00:00 to 21 May 00:00 mean trajectories (uppermost panel), and individual trajectories attributed to the six groupings: C, E, N, W, A, and P. The traces correspond to 72 h backward trajectories calculated using the HYSPLIT4 model (Draxler and Hess, 2004). New trajectories were started every 6 h and ended 500 m above ground level in Gothenburg.
Figure 7. Mean back trajectories based on the group analysis illustrated in Figure 6. The six groups and percent of the total number of trajectories assigned to each group are indicated. See text for further details.
Figure 8. a) 72 h backward trajectories for the same period. A new trajectory was started every 12 h. b) Distribution of fires from March 24 to April 1, 2007. Each dot on the map indicates a fire detected with the moderate resolution imaging spectroradiometer (MODIS) aboard the Terra satellite (Roy et al., 2002).
Figure 9. Experimental data from an episode from March 24 to April 4 with high K concentrations: a) K and Na concentrations, b) wind direction, and c) wind speed.
Figure 10. K concentration in Clusters 1 and 2 as a function of a) temperature, b) wind speed, and c) wind direction.
Figure 11. Measured PM2.5 concentrations at the urban background station Femman in Gothenburg (black line, right scale), during the period 17 Feb. – 20 May 2007, and modeled concentrations of particulate carbonaceous matter (PCM) in PM2.5 from: residential biomass combustion (grey, dashed line), open biomass burning fires (including wild fires and agricultural fires; orange line), and total biomass burning PCM (red line). Unit: µg m⁻³. The data have been smoothed using moving 24 h mean concentrations.
Figure 12. Comparison of the modeled hourly concentrations of primary PM2.5 from residential biomass combustion (blue line) and open fires (red line) and measured (hourly average) concentrations of K in Cluster 1 (black line). Units, model results: ng m\(^{-3}\); measurements: ng K m\(^{-3}\).
Figure 13. Comparison of the modeled hourly concentrations of primary PM2.5 from fossil fuel combustion (red line) and measured (hourly average) concentrations of K in cluster 1 (black line). Units, model results: $\mu g m^{-3}$; measurements: ng K m$^{-3}$. 
Figure 14. Comparison of the modeled hourly concentrations of primary PM2.5 from fossil fuel combustion (red line) and measured (hourly average) concentrations of K in cluster 2 (black line). Units, model results: \( \mu g \text{ m}^{-3} \); measurements: ng K m\(^{-3}\).
Table 1. Back trajectory group analysis for six air mass types indicates the air mass category, number of 6 min.
samples, percent of the total number of trajectories assigned to each group, mean Na and K concentrations [ng
m$^{-3}$], Na:K concentration ratio, mean PM$_{10}$ and trace gas concentrations [$\mu$g m$^{-3}$], temperature (T) [$^\circ$C] and wind
speed (WS) [m s$^{-1}$].

<table>
<thead>
<tr>
<th>Origin group (Initial)</th>
<th>Samples</th>
<th>%</th>
<th>Na</th>
<th>K</th>
<th>Na/K</th>
<th>PM$_{10}$</th>
<th>O$_3$</th>
<th>NO</th>
<th>NOx</th>
<th>T</th>
<th>WS</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Central Eur. (C)</td>
<td>3238</td>
<td>19</td>
<td>3.9</td>
<td>14</td>
<td>0.27</td>
<td>34.8</td>
<td>46.2</td>
<td>11.5</td>
<td>51.4</td>
<td>7.0</td>
<td>3.5</td>
</tr>
<tr>
<td>2. Eastern Eur. (E)</td>
<td>2171</td>
<td>13</td>
<td>6.2</td>
<td>21</td>
<td>0.29</td>
<td>34.1</td>
<td>53.8</td>
<td>16.6</td>
<td>57.6</td>
<td>2.9</td>
<td>4.7</td>
</tr>
<tr>
<td>3. Northern Eur. (N)</td>
<td>5027</td>
<td>30</td>
<td>6.0</td>
<td>5.6</td>
<td>1.1</td>
<td>20.1</td>
<td>57.6</td>
<td>7.8</td>
<td>36.0</td>
<td>8.3</td>
<td>3.5</td>
</tr>
<tr>
<td>4. Western Eur. (W)</td>
<td>1393</td>
<td>8</td>
<td>4.9</td>
<td>11</td>
<td>0.45</td>
<td>25.3</td>
<td>49.3</td>
<td>4.6</td>
<td>32.0</td>
<td>8.0</td>
<td>3.6</td>
</tr>
<tr>
<td>5. Atlantic (A)</td>
<td>4188</td>
<td>25</td>
<td>3.6</td>
<td>3.6</td>
<td>9.2</td>
<td>27.2</td>
<td>66.6</td>
<td>2.7</td>
<td>23.3</td>
<td>7.2</td>
<td>5.4</td>
</tr>
<tr>
<td>6. Polar (P)</td>
<td>847</td>
<td>5</td>
<td>8.6</td>
<td>2.7</td>
<td>3.2</td>
<td>20.1</td>
<td>57.1</td>
<td>9.1</td>
<td>36.9</td>
<td>4.9</td>
<td>4.7</td>
</tr>
</tbody>
</table>
Table 2. Correlation coefficients ($r$) between measured concentrations in cluster 1 and 2 (particle number and K mass concentrations) and modeled concentrations of particulate matter from primary emissions (PM) from selected sources; both measured and modeled concentrations were smoothed using 24-h moving average concentrations. Correlation coefficients with values above 0.5 are marked in bold.

<table>
<thead>
<tr>
<th>Model tracer compound/source category</th>
<th>Cluster 1 Number</th>
<th>Cluster 1 K</th>
<th>Cluster 2 Number</th>
<th>Cluster 2 K</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total Biomass Burning (residential + fires)</td>
<td>0.76</td>
<td>0.64</td>
<td>0.24</td>
<td>0.31</td>
</tr>
<tr>
<td>Residential Biomass Combustion (SNAP-2)</td>
<td>0.71</td>
<td>0.62</td>
<td>0.15</td>
<td>0.23</td>
</tr>
<tr>
<td>Open Biomass Burning (Fires)</td>
<td>0.38</td>
<td>0.21</td>
<td>0.44</td>
<td>0.41</td>
</tr>
<tr>
<td>Total Other Anthropogenic PPM sources*</td>
<td>0.41</td>
<td>0.26</td>
<td>0.63</td>
<td>0.61</td>
</tr>
<tr>
<td>Non-industrial/residential combustion fossil fuels (coal, oil, gas) (SNAP-2)</td>
<td>0.51</td>
<td>0.54</td>
<td>0.27</td>
<td>0.37</td>
</tr>
<tr>
<td>Manufacturing industry (SNAP-3)</td>
<td>0.53</td>
<td>0.30</td>
<td>0.61</td>
<td>0.58</td>
</tr>
<tr>
<td>Road transport (SNAP-7)</td>
<td>0.39</td>
<td>0.19</td>
<td>0.57</td>
<td>0.52</td>
</tr>
<tr>
<td>International shipping</td>
<td>-0.04</td>
<td>-0.08</td>
<td>0.33</td>
<td>0.30</td>
</tr>
<tr>
<td>Other mobile sources/machinery</td>
<td>0.28</td>
<td>0.12</td>
<td>0.53</td>
<td>0.50</td>
</tr>
<tr>
<td>Agriculture (SNAP-10)</td>
<td>0.33</td>
<td>0.12</td>
<td>0.65</td>
<td>0.59</td>
</tr>
</tbody>
</table>

* Includes all anthropogenic primary PM sources, except residential biofuel combustion and Open Biomass Burning (Fires) – this is expected to be mainly from fossil fuel sources.

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References


