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Airborne measurements of the spatial distribution of aerosol chemical composition across Europe and evolution of the organic fraction

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Abstract. The spatial distribution of aerosol chemical composition and the evolution of the Organic Aerosol (OA) fraction is investigated based upon airborne measurements of aerosol chemical composition in the planetary boundary layer across Europe. Sub-micron aerosol chemical composition was measured using a compact Time-of-Flight Aerosol Mass Spectrometer (cToF-AMS). A range of sampling conditions were evaluated, including relatively clean background conditions, polluted conditions in North-Western Europe and the near-field to far-field outflow from such conditions. Ammonium nitrate and OA were found to be the dominant chemical components of the sub-micron aerosol burden, with mass fractions ranging from 20–50% each. Ammonium nitrate was found to dominate in North-Western Europe during episodes of high pollution, reflecting the enhanced NOₓ and ammonia sources in this region. OA was ubiquitous across Europe and concentrations generally exceeded sulphate by 30–160%. A factor analysis of the OA burden was performed in order to probe the evolution across this large range of spatial and temporal scales. Two separate Oxygenated Organic Aerosol (OOA) components were identified; one representing an aged-OOA, termed Low Volatility-OOA and another representing fresher-OOA, termed Semi Volatile-OOA on the basis of their mass spectral similarity to previous studies. The factors derived from different flights were not chemically the same but rather reflect the range of OA composition sampled during a particular flight. Significant chemical processing of the OA was observed downwind of major sources in North-Western Europe, with the LV-OOA component becoming increasingly dominant as the distance from source and photochemical processing increased. The measurements suggest that the aging of OA can be viewed as a continuum, with a progression from a less oxidised, semi-volatile component to a highly oxidised, less-volatile component. Substantial amounts of pollution were observed far downwind of continental Europe, with OA and ammonium nitrate being the major constituents of the sub-micron aerosol burden. Such anthropogenically perturbed air masses can significantly perturb regional climate far downwind of major source regions.

1 Introduction

The chemical composition of the atmospheric aerosol burden has significant implications for its climate impacts (e.g. Forster et al., 2007). Specifically, it plays a major role in determining the scattering or absorbing nature of the aerosol and has an important control upon its affinity for water uptake. Furthermore, the chemical composition of the particle phase is an important component of global and regional biogeochemical cycles (Andreae and Crutzen, 1997). These include the cycling of carbon, sulphur, nitrogen, oxygen and water. The spatial heterogeneity of the aerosol burden has significant implications for its subsequent impact. On regional scales, the direct effect of aerosols is capable of
substantially reducing the impact of greenhouse gas radiative forcing, due to their competing cooling and warming effects respectively (Charlson et al., 1992). This is particularly evident over industrialized and heavily populated regions of the Northern Hemisphere such as North America, Europe and South-East Asia. Additionally, aerosols can alter the microphysical properties of clouds (e.g. Haywood and Boucher, 2000), leading to changes in the radiation balance of the climate system and also regional meteorology (Denman et al., 2007).

Several recent intensive field studies have sought to elucidate aerosol chemical composition and the processes which change it, with a focus upon their regional impacts. These include studies in Asia such as the Indian Ocean Experiment (INDOEX, Ramanathan et al., 2001) and North America such as the New England Air Quality Study (NEAQS, Bates et al., 2005; Kleinman et al., 2007; Wang et al., 2007), the International Consortium for Atmospheric Research on Transport and Transformation (ICARTT, Quinn et al., 2006; Williams et al., 2007; Brock et al., 2008; de Gouw et al., 2008) and the Megacity Initiative: Local and Global Research Observations (MILAGRO, DeCarlo et al., 2006; Kleinman et al., 2008; Baumgardner et al., 2009; Fast et al., 2009). Recent airborne studies in Europe have focused upon polluted environments in the Adriatic and Black Seas (Crossier et al., 2007) and the UK region (Morgan et al., 2009). A major conclusion of these European studies was the significant contribution of ammonium nitrate to the sub-micron particulate burden, particularly during highly polluted conditions. Zhang et al. (2007) presented a summary of numerous field studies in the Northern Hemisphere from the Aerodyne Aerosol Mass Spectrometer (AMS, Jayne et al., 2000; Canagaratna et al., 2007). A major theme of such analyses was the high proportion of Organic Matter (OM) contributing to the sub-micron particulate burden. The study by Zhang et al. (2007) indicated that the OM component was dominated by oxygenated species relative to Primary Organic Aerosol (POA) and that the mass fraction of the more oxidised component compared to the total organic mass increases away from urban environments. Comparison of such ambient measurements of Secondary Organic Aerosol (SOA) with atmospheric chemistry models reveals significant discrepancies between them (Volkamer et al., 2006, and references therein). Furthermore, such discrepancies increase as a function of photochemical age.

Coupling of a thermodenuder system with an AMS indicated that the SOA component could be separated in terms of their volatility, with more aged-SOA being less volatile than fresher-SOA (Huffman et al., 2009). Furthermore, the POA component was shown to be semi-volatile. Such results are consistent with recent frameworks which have treated the entire OM component as semi-volatile (Donahue et al., 2006; Robinson et al., 2007). This is in contrast to traditional modelling approaches that prescribe the POA to be non-volatile and inert (Donahue et al., 2009). Recently, such a framework has been implemented to explain OM volatility and composition variations both downwind of a megacity source and across a global ground-based dataset (Jimenez et al., 2009).

The present study seeks to elucidate the spatial distribution and chemical evolution of the sub-micron particulate mass across a broad range of scales, with particular emphasis upon the OM component and its contrasting behaviour compared to inorganic species. We do so using measurements drawn from two related aircraft campaigns across Northern Europe, a region that plays an important role in the global aerosol budget.

The major part of the analysis consists of measurements made during May 2008 as part of the European Integrated Project on Aerosol Cloud Climate and Air Quality Interactions (EUCAARI, Kulmala et al., 2009) airborne intensive study, known as the EUCAARI-LONG Range EXperiment (EUCAARI-LONGREX, henceforth referred to as LON-GREX). These measurements are complemented by flight operations based out of the UK, which took place during April and September 2008 as part of the Appraising the Direct Impacts of Aerosol on Climate (ADIENT) project. A key aim of these projects is an evaluation of the relative contribution of particular chemical components to the aerosol burden. This includes characterisation of the multiple components which make up the OM burden, along with their subsequent evolution in the atmosphere. The present dataset is well placed to investigate processes governing the formation and transformation of atmospheric aerosol due to the broad range of spatial scales investigated.

2 Method

The UK Facility for Airborne Atmospheric Measurements (FAAM) BAE-146 research aircraft took part in the LONGREX campaign in conjunction with the Deutsches Zentrum für Luft-und Raumfahrt (DLR) Falcon 20-E5. The LON-GREX campaign was closely coordinated with the Intensive Observation Period at Cabauw Tower (IMPACT), the second major part of the EUCAARI airborne intensive operational period, although the analysis presented here is based solely on data obtained during LONGREX. Principally, the BAE-146 operated within the planetary boundary layer, while the Falcon operated at high-altitude in the free troposphere. The Falcon operated a LIDAR system which included a High Spectral Resolution LIDAR mode (HSRL, Esselborn et al., 2008) delivering aerosol backscatter and extinction coefficients. Real time LIDAR data were utilised during flight operations to identify the location of pollution plumes, which were subsequently sampled in-situ by the BAE-146. Flight operations were conducted across Northern Europe from the 6–22 May. A period of stable anticyclonic conditions characterised the first eight days of the project from the 6–14 May. Two more flights were conducted during LON-GREX over the 21–22 May period,
which was characterised by predominantly easterly conditions. Flights from LONGREX are considered along with the ADIENT flights from April and September 2008. The ADIENT flights were conducted in differing meteorological conditions, with the April flight conducted in southeast-easterly conditions, while the September flights took place during easterly conditions with similar transport patterns to the LONGREX period. The flight tracks are shown on Fig. 1 and are coloured according to the different flying periods and conditions. The meteorological conditions for the different periods are relatively consistent in terms of their transport patterns, with air masses advecting pollution from continental Europe downwind to either the UK region or the Eastern Atlantic Ocean (see Supplementary Materials Sect. 2, Figs. S1 and S2; http://www.atmos-chem-phys.net/10/4065/2010/acp-10-4065-2010-supplement.pdf). Some flights concentrated on instrument testing/calibration and are not included in the analysis. During some flights instrument performance was not optimal; data from these 3 flights have not been included. The flights included in this analysis are summarised in Table 1.

2.1 Instrumentation

The FAAM BAe-146 research aircraft houses a suite of instruments capable of resolving the chemical composition, microphysical, optical and hygroscopic properties of the in-situ aerosol population. Inboard aerosol instrumentation samples ambient air via stainless steel tubing from Rosemount inlets (Foltescu et al., 1995). Sub-micron particle losses have been shown to be negligible (Osborne et al., 2007). Number size distributions were measured using a wing-mounted Particle Measurement Systems (PMS) Passive Cavity Aerosol Spectrometer Probe 100X (PCASP, Liu et al., 1992; Strapp et al., 1992). The PCASP instrument optically counts and sizes particles between 0.1–3 µm diameter across 15 channels. Particle size is determined via experimental calibrations using Di-Ethyl-Hexyl-Sebacate (DEHS), which is converted to a Polystyrene Latex Sphere (PSL) equivalent size. Black Carbon (BC) mass and the number of particles containing a BC core were characterised as a function of particle size using a Droplet Measurement Technologies (DMT) Single Particle Soot Photometer (SP2, Stephens et al., 2003; Baumgardner et al., 2004). The SP2 was modified to include a two-element avalanche photodiode to improve determination of the optical size of absorbing
Table 1. Flight summary for ADIENT (B357, B401-B406) and LONGREX (B362-B380) operations included in this study. All flights were conducted during 2008. The predominant meteorological conditions present are summarised, where SE refers to south-easterly, E refers to easterly and HP refers to the high pressure system during the initial LONGREX flights described in the main text. Also included is an indication of which period each flight took place during, which are abbreviated as A1 (ADIENT-1) and A2 (ADIENT-2). The LONGREX flights refer to the 3 periods in the main text, which are referred to as LONGREX-1 (L1), LONGREX-2 (L2) and LONGREX-3 (L3).

<table>
<thead>
<tr>
<th>Flight</th>
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<td>B357</td>
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<td>B362</td>
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<tr>
<td>B370</td>
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<tr>
<td>B371</td>
<td>12 May</td>
<td>HP</td>
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<td>North-Western Europe &amp; North Sea</td>
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<tr>
<td>B373</td>
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<td>L2</td>
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<tr>
<td>B374</td>
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<td>L3</td>
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<tr>
<td>B380</td>
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<td>E</td>
<td>L3</td>
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<tr>
<td>B401</td>
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<td>E</td>
<td>A2</td>
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<tr>
<td>B402</td>
<td>19 September</td>
<td>E</td>
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<td>Eastern UK coast</td>
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<tr>
<td>B406</td>
<td>25 September</td>
<td>E</td>
<td>A2</td>
<td>Southern and Western UK coast</td>
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</table>

particles (Gao et al., 2007). Additionally, gas phase concentrations of CO (Carbon Monoxide), O$_3$ (Ozone) and NO$_x$ (defined as the sum of NO (Nitric Oxide) and NO$_2$ (Nitrogen Dioxide)) were measured. The facility also provides aircraft position information and measurements of standard atmospheric variables, such as temperature and relative humidity.

Measurements made by an Aerodyne compact Time-of-Flight Aerosol Mass Spectrometer (cToF-AMS, Drewnick et al., 2005; Canagaratna et al., 2007) form the major part of this study. The instrument provides the capability to quantitatively measure the size-resolved chemical composition of non-refractory particulate matter, including OM, sulphate, nitrate, ammonium and chloride. A major advantage of the cToF-AMS is the ability to provide high time resolution measurements with enhanced precision and sensitivity. Thus it is ideal for airborne deployments. Previous studies (Crosier et al., 2007; Capes et al., 2008; Morgan et al., 2009) have detailed the AMS sampling strategy onboard the BAe-146. The sampling losses for the cToF-AMS inlet system were estimated experimentally to be approximately 10% by number across the size range of the AMS. This was accomplished by comparing a Condensation Particle Counter (CPC) upstream of the inlet system with one situated immediately prior to the sampling orifice of the AMS while sampling monodisperse aerosol. Details specific to the cToF-AMS data analysis will be discussed in the following subsection.

### 2.2 AMS data quantification

The AMS data analysis was performed using the standard SQUIRREL (SeQUential Igor data RetRiEvaL) ToF-AMS software package. Mass spectrum deconvolution is accomplished using the fragmentation table approach described by Allan et al. (2004). Error estimates are generated according to the model documented by Allan et al. (2003). Mass concentrations derived from the AMS are reported as micrograms per standard cubic metre ($\mu$g sm$^{-3}$) i.e. at a temperature of 273.15 K and pressure of 1013.25 hPa. Power was unavailable to the AMS between flights due to operational constraints. Through the use of plug valves to isolate the AMS chamber, a vacuum of typically less than 0.5 Torr is maintained while the turbo-pumps are powered down. Ionisation efficiency (IE) calibrations were performed regularly before and after each flight during the flying periods. Values determined from both pre-flight and post-flight calibrations (i.e. taking place on the same day) exhibited little variability. Post-flight values were used as these were considered to be more reliable compared with pre-flight values due to the reduced instrument background post-flight. A faulty ground power unit during LONGREX caused a loss of power to the aircraft prior to flight B365 while the AMS was pumping down. This led to slightly enhanced background concentrations in the AMS vacuum chamber due to a filament failure associated with the power loss during flights B365–B369.

Recent laboratory evidence (Matthew et al., 2008) suggests that the AMS Collection Efficiency (CE, Huffman et al., 2005) is significantly modulated by particle phase. Previous AMS data collected in Europe on the BAe-146
has used a CE correction following the principle developed by Crosier et al. (2007). This was developed using a simple empirical treatment of the CE as a linear function of the nitrate content of the aerosol based upon comparison of AMS sulphate with filter measurements. Matthew et al. (2008) conducted a detailed laboratory study of the AMS CE across a range of compositions. Their study included a comparison of their laboratory derived correction with the Crosier et al. (2007) parameterisation. This indicated that the simpler linear treatment overestimated the CE at intermediate ammonium nitrate mass fractions. A comparison between the two techniques as part of this work revealed that the two approaches showed a maximum discrepancy at intermediate ammonium nitrate mass fractions that did not typically exceed 25% and was considerably less at high and low mass fractions. Consequently, we use the correction developed by Matthew et al. (2008) to estimate the CE of the AMS in this study. The AMS total mass concentrations were converted to total volume concentrations using the densities reported by Cross et al. (2007), which correspond to 1.27 g cm\(^{-3}\) for organics and 1.77 g cm\(^{-3}\) for inorganics. Comparison of the estimated AMS total volume with the PCASP indicates that campaign average agreement was within 30% (see Supplementary Materials Sect. 3, Fig. S4; http://www.atmos-chem-phys.net/10/4065/2010/acp-10-4065-2010-supplement.pdf). This is within the 30–50% uncertainty previously reported for PCASP volume concentration estimates (e.g. Moore et al., 2004; Hallar et al., 2006). All flights fall within the 50% agreement range except for B357. The uncertainties in the estimate are large due to the high sensitivity of the volume calculations to diameter (proportional to diameter\(^3\)) and the uncertainties in density required to convert the AMS mass to volume.

### 3 Spatial distribution of aerosol chemical composition across Europe

The differing spatial scales and meteorological contexts considered by this analysis provide a thorough examination of the aerosol chemical composition in Europe during anticyclonic conditions. The distribution of the chemical components measured by the AMS will be discussed here in order to set the context for the subsequent analysis and discussion. Both the broader scale and flight period/condition specific details will be summarised. The flight periods were split according to the period in which they were conducted and their general meteorological conditions. Their designations are summarised in Table 1 and Fig. 1.

The spatial distribution of the total OM for Straight and Level Runs (SLRs) below 3000 m is shown for the entire dataset in Fig. 2. The boundary layer height was predominantly below 2000 m but during some conditions, an elevated planetary boundary layer was observed so 3000 m was used as a threshold altitude. The distribution indicates that OM is a major contributor to the sub-micron aerosol burden, with typical mass fractions from 20–50%. Importantly, OM is a significant component during both background and highly polluted conditions. This is illustrated in Fig. 2, as the OM mass fraction almost always accounts for more than 20% of the total mass measured by the AMS.

The absolute and relative contributions of the chemical components measured by the AMS can be discerned from Fig. 1b. The flight data have been separated into different zones representing the main areas and emission fields sampled across Northern Europe and average composition data has been determined for each area. The OM concentration is typically 1.4–2.6 times greater than that of sulphate further east (zones 4–5) and is the dominant component mass in sub-micron aerosol in central and eastern Europe (zones 5 and 6), while in zones further west, the OM and sulphate mass concentrations are comparable. Sulphate contributes approximately 1.0–4.0 µg m\(^{-3}\) to the regional aerosol burden, with typical mass fractions of 10–30%. The spatial distribution of nitrate indicates large gradients in the concentration and relative contribution of this component. In particular, the nitrate mass concentration is enhanced in North-Western Europe compared to locations further east. In terms of its relative contribution, nitrate accounts for approximately 20–50% of the sub-micron mass across a wide range of locations and conditions. Furthermore, increases in the nitrate concentration are associated with enhanced total mass loadings, indicating that nitrate is a significant contributor under highly polluted conditions. Examination of the sulphate, nitrate and ammonium concentrations indicate that the aerosol was neutralised, thus sulphate is present in the form of ammonium sulphate and nitrate is in the form of ammonium nitrate.

The ADIENT-1 period focuses on one flight (B357, zone 2), which was conducted in the large-scale outflow
from continental Europe and the southern UK along the UK’s western edge. The flight included close to source sampling downwind of Manchester and Liverpool, in the north-west of England (identified on Fig. 1). During this flight, the secondary aerosol species’ mass concentrations were reduced when compared to the other periods. This is potentially a consequence of reduced photochemistry as the flight took place earlier in the year than the LONGREX and ADIENT-2 flying.

Analysis of the high pressure phase of LONGREX indicated significant differences in gas-phase and particulate loadings, which could be categorised into two distinct meteorological periods. During the initial phase of LONGREX from the 6–8 May (referred to as LONGREX-1), the air mass trajectories displayed a more zonal flow from east-to-west. The second phase from the 10–14 May (henceforth referred to as LONGREX-2) is characterised by the then well established high pressure system centred over northern Germany and Denmark, yielding more rotational anticyclonic flow. The LONGREX-1 period was characterised by enhanced CO and NO\textsubscript{x} levels compared to the second period. These strong signatures of urban pollution on the regional scale correlated with significant amounts of secondary material, with total mass concentrations reported by the AMS exceeding 25 µg sm\textsuperscript{−3}. Such instances were dominated by ammonium nitrate and OM. Such conditions were also prevalent during the LONGREX-3 and ADIENT-2 flying periods. These aforementioned periods concentrated upon sampling close to major anthropogenic sources and the immediate near-field downwind evolution of such sources.

The major distinction between LONGREX-2 and the operations conducted in each of the other identified periods was the large spatial scale sampled, where the measurements extended from the Baltic Sea region in Northern Europe, to the Eastern Atlantic Ocean, off the southern Irish coast. The operations represented approximately 3000 km and approximately 4–5 days of air mass transit based on air mass back trajectories. These were derived from European Centre for Medium-Range Weather Forecasts (ECMWF) wind fields, initialised from the SLRs in the east-Atlantic (51 N, 12 W). During the LONGREX-2 period, when NO\textsubscript{x} and background CO levels were reduced, the mass concentration of ammonium nitrate was diminished in North-Western Europe. In terms of the median AMS aerosol concentrations, LONGREX-2 was similar to LONGREX-1 in terms of sulphate and OM, while nitrate was reduced. The main difference between the periods was the absence of the more intense ammonium nitrate concentrations, which contributed to the significant pollution episodes encountered during LONGREX-1. The operations in the Eastern Atlantic probed total sub-micron mass loadings from the AMS exceeding 15 µg sm\textsuperscript{−3} with OM and ammonium nitrate being the dominant chemical components. The Falcon’s HSRL measured aerosol optical depth values from 0.3–0.5 in these conditions far downwind of the major source regions in Western Europe.

Vertical profiles of aerosol chemical composition are shown in Fig. 3 from the LONGREX-1 and LONGREX-2 periods. The location of each profile are shown in Fig. 1. The profiles represent the broad range of conditions sampled,
with the cleaner conditions in the Baltic region (panel C, zone 7), the immediate outflow from North-Western Europe into the North Sea (panel B, zone 4) and the outflow from continental Europe far downwind in the Eastern Atlantic (panel A, zone 1). The profiles show that the boundary layer typically extended to between 2000–3000 m, the actual height of which as determined by the location of the measurements in relation to the centre of the high pressure system. Also evident in panel B is the strong increase in concentrations close to the sea surface, which was coincident with a shallow moist layer where concentrations of both ammonium nitrate and OM increased significantly. Also discernible is the increase in sulphate at lower levels (below 1000 m) which is likely attributable to shipping activities. This was also observed in zone 3, which covers the English Channel region, where sulphate concentrations increased and the largest sulphate mass fractions were observed. Further details and observations regarding the vertical distribution of aerosol chemical composition are discussed in Morgan et al. (2010) in the EUCAARI special issue.

The following analysis and discussion will focus upon the evolution of the ubiquitous OM component across the full range of spatial scales investigated. This is accomplished using a factor analysis technique which will be presented and discussed in the following section. The results of this analysis will then be used to probe the evolution of the OM component and contrast it with the observed gradients in the ammonium nitrate and ammonium sulphate fields across Europe.

4 Factor analysis of organic aerosol: technique and discussion

Several recent studies have employed various factor analysis techniques in order to deconvolve the organic mass spectra derived from ambient AMS measurements (e.g. Zhang et al., 2005a,b; Lanz et al., 2007, 2008; Ulbrich et al., 2009). With the exception of biomass burning, wood combustion or urban environments with a prevalent cooking signature, such studies have attributed the OM component to be a combination of Hydrocarbon-like Organic Aerosol (HOA) and Oxygenated Organic Aerosol (OOA). These two factors commonly explain more than 90% of the variance in the ambient organic mass spectra (McFiggans et al., 2005; Zhang et al., 2005a; Rudich et al., 2007). Furthermore, such factors remain relatively constant across differing environments. Several studies (e.g. Lanz et al., 2007; Zhang et al., 2007; Ulbrich et al., 2009) have identified multiple OOA factors, which have been classified according to their level of oxidation.

Jimenez et al. (2009) characterised the evolution and volatility of these multiple OOA components using the terms Low-Volatility OOA (LV-OOA) and Semi-Volatile OOA (SV-OOA). These terms represent the OOA-1 and OOA-2 components identified in previous studies (e.g. Lanz et al., 2007; Ulbrich et al., 2009), though Jimenez et al. (2009) demonstrated that these factors were systematically correlated with lower and higher volatile fractions of the OM. LV-OOA is distinguished by the predominance of signal at m/z 44 (corresponding to the CO$_2^+$ ion arising from decarboxylation on the vapouriser surface). SV-OOA components typically exhibit enhanced signal at m/z 43 (C$_2$OH$_3^+$ and C$_3$H$_7^+$) and reduced signal at m/z 44, when compared to LV-OOA mass spectra. Examination of high-resolution AMS data has demonstrated the dominance of the C$_2$OH$_3^+$ ion at m/z 43 in ambient spectra when the contribution of OOA is high (Mohr et al., 2009). Thus this SV-OOA component represents a less oxidised OM fraction. Jimenez et al. (2009) argue that the atmospheric OOA evolves through a dynamic aging process of continual repartitioning between the particle and gas phases, which leads to a more oxidised, less volatile and more hygroscopic aerosol. Thus the initial OOA, which resembles SV-OOA, undergoes transformation processes that ultimately result in an OOA that exhibits LV-OOA like characteristics. Following this framework, the present analysis strives to simplify the OM burden into a limited number of key factors in order to probe their relative magnitudes, relations and evolution across Europe for the first time.

4.1 Positive Matrix Factorisation (PMF)

Positive Matrix Factorisation (PMF, Paatero and Tapper, 1994; Paatero, 1997) was utilised in order to accomplish some simplification of the OM burden. Several recent studies have detailed its application to AMS data (Lanz et al., 2007; Docherty et al., 2008; Ulbrich et al., 2009; Aiken et al., 2009). PMF employs a receptor-only factorisation model, which is based on mass conservation. The model assumes that a dataset matrix is comprised of a linear combination of factors with constant profiles, which have varying contributions across the dataset. The model employs the constraint of positive values upon the profiles and contributions. This work will follow the procedures identified by Ulbrich et al. (2009) in order to apply the PMF technique to AMS data. Version 4.2 of the PMF2 algorithm (provided by the University of Helsinki) is employed in robust mode to perform the factorisation.

The sampling and data collection strategy detailed in Sect. 2.2 resulted in the requirement that PMF be applied to each flight separately. Application of PMF to the whole dataset in one single matrix was found to be problematic due to the subtle changes in instrument performance across the 6 month flying period considered by this study coupled with the broad range of scales examined. Such changes impact the retrieved factors as instrument performance changes dominate the variability in the dataset, rather than the atmospheric processes and sources of interest here. Vertical profiles of temperature and AMS chemical composition indicated that the boundary layer top was typically between 2000–3000 m. Only data from below 3000 m was included in the analysis as we wish to consider boundary layer processes only and the
4.2 Results of the factor analysis

For the purposes of the subsequent discussions, the retrieved factor components are classified according to their level of oxidation, which is signified by their relative organic intensity at $m/z$ 44. Thus, factors with the greatest $m/z$ 44 signal are designated OOA-1 with subsequent factors designated as OOA-2, OOA-3 etc. This is conceptually similar to the nomenclature introduced by Lanz et al. (2007) and is used to initially compare the retrieved factors with reference mass spectra and external tracers. These will then be discussed in terms of their resemblance to LV-OOA, SV-OOA and HOA factors derived by Jimenez et al. (2009).

For all of the flights considered, two factor solutions were found to be the most appropriate; one that is highly oxidised (OOA-1) and a second that is less oxidised (OOA-2), which often represented a combination of HOA with fresher OOA components. The factors were chosen based upon comparison with reference mass spectra, external tracers and their numerical stability. Case study examples of a range of flights in differing conditions are presented in the Supplementary Materials (Sect. 5) (http://www.atmos-chem-phys.net/10/4065/2010/aqp-10-4065-2010-supplement.pdf). When more than two factors were chosen, the solutions were found to be numerically unstable based upon a bootstrapping analysis (Ulbrich et al., 2009, and references therein), where random resampling of the data matrix is performed in the time dimension.

The retrieved factors for the 2-factor solutions were compared with reference mass spectra in Fig. 4a. OOA-1 type mass spectra were highly correlated ($r>0.9$) with fulvic acid across the dataset and variability in the retrieved mass spectrum was low. Furthermore, the OOA-1 profiles had low correlation with the reference HOA mass spectrum from Pittsburgh ($r<0.45$ for all cases). The OOA-2 component typically exhibits enhanced signal at $m/z$ 43 relative to signal at $m/z$ 44, when compared to the OOA-1 mass spectra. The OOA-2 components had lower correlations with fulvic acid. The OOA-2 also exhibited greater correlation with the reference HOA spectrum with coefficients ranging from 0.5–0.9. This is unsurprising given that we have not separated the HOA contribution from the fresher-OOA component. The correlations of OOA-2 with fulvic acid and reference HOA reflect the chemical variation in the OOA component as they are anti-correlated with each other. Thus, as the OOA-2 becomes more oxidised it resembles HOA less and approaches a more LV-OOA or fulvic acid-like mass spectrum.

Previous studies (e.g. Lanz et al., 2007; Ulbrich et al., 2009) have reported that the time series of nitrate and OOA-2 show a close coupling, thus exhibiting an enhanced correlation coefficient. Similarly, OOA-1 and sulphate have been shown to exhibit enhanced correlation coefficients. Figure 4a displays the correlation coefficients for OOA-1 and OOA-2 with sulphate and nitrate in order to compare with previous results. The results from this study somewhat corroborate previous findings regarding such associations but some flights indicate substantial differences. Nitrate and OOA-2 are frequently found to be well correlated ($r>0.7$) but the same is also true for nitrate and OOA-1. The high correlations between nitrate and OOA-2 principally occur closer to source regions due to their semi-volatile nature. The examples of high correlation of nitrate and OOA-1 are further away from source due to the ammonium nitrate concentration being sustained downwind, while the OOA-2 has been processed to OOA-1. The most stark example of this is B374, where nitrate and OOA-2 are anti-correlated while nitrate and OOA-1 are highly correlated. B374 took place far downwind of the major sources in continental Europe in the Eastern Atlantic Ocean, where the air mass had aged significantly and little precipitation had occurred upward. Once formed, ammonium nitrate exists in a chemical equilibrium with ammonia and nitric acid, the losses of which are via dry and wet deposition only. This contrasts with OOA, which undergoes complex and continual processing involving repartitioning and oxidation. Thus the OM transforms from more volatile to less volatile condensable products as the air parcel ages downwind of source. Thus the OM is dominated by the more-oxidised form downwind of the major emission sources and is then well correlated with ammonium nitrate.
Sulphate is often well correlated with OOA-2 and OOA-1, which is likely due to the regional nature of these measurements and the covariance between the different chemical species. Thus factor interpretations where multiple OOA components are identified need to consider the meteorological, photochemical and geographical context at individual sampling locations when making suppositions based upon comparison with other secondary particulate species.

Given that the OOA-2 component generally contains a HOA contribution, it is desirable to estimate the relative importance of the primary versus secondary components. Some previous studies have included first order estimates of OM components based upon absolute intensities at specific mass spectral markers by comparing such markers with factor component solutions (e.g. Zhang et al., 2005a; Aiken et al., 2009). Zhang et al. (2005a) reported that the HOA mass concentration can be approximated based upon a linear scaling of the signal intensity at m/z 57. Aiken et al. (2009) reported a similar relationship but with an additional correction for oxidised fragments associated with m/z 57 based upon the organic signal intensity at m/z 44. In order to derive an approximate estimate of the contribution of HOA to the OM burden in this dataset, these first order estimates are used.

While such estimations are not fully quantitative, they do at least represent an approximate reference point which provides a level of justification for the decision to limit the analysis to the 2-factor solution sets. Furthermore, the estimated HOA is typically less than the OOA-2 concentration reported from the factor analysis. An important observation is that the enhanced HOA mass fractions are predominantly driven by the reduced contribution from secondary species, rather than a major increase in the absolute HOA mass loading.

The Aiken et al. (2009) estimate was found to be inappropriate for this dataset as the m/z 57 contribution was often close to zero, while the organic intensity at m/z 44 was typically an order of magnitude greater. This led to negative HOA concentrations frequently being estimated using this approximation. The solutions for the entire dataset are summarised in Fig. 4b by comparing the absolute mass concentrations for the OOA-1, OOA-2 and estimated HOA. The HOA contribution is calculated using the estimation from Zhang et al. (2005a) and indicates that the median concentration is typically less than 0.5 µg sm$^{-3}$, with concentrations rarely exceeding 1 µg sm$^{-3}$. Such a result indicates that HOA typically contributes 5–20% to the regional OM burden, which is in line with previous studies (Zhang et al., 2005a).
2007). Consequently, the dataset indicates that the OOA component dominates the OM burden across Europe on the regional scale, thus this will be the focus of the subsequent discussion.

### 4.3 Interpretation of the factor analysis

Typical example mass spectra from different flights from the 2 factor solutions are presented in Fig. 5a. Spectrum A is highly consistent with LV-OOA, with the spectrum being dominated by normalised organic signal intensity at m/z 44 and with a correlation coefficient of 0.99 with fulvic acid. This spectrum is taken from flight B366, as is spectrum B which is distinguished by the normalised m/z 43 and 44 peaks being almost equal. This is consistent with SV-OOA spectra (e.g. Ulbrich et al., 2009; Jimenez et al., 2009) and the relative concentration of this component versus the HOA estimated suggests it is dominated by OOA. Spectrum C is the B357 HOA-type factor which had a correlation coefficient of 0.91 with the reference HOA spectrum. This factor is likely dominated by HOA-like components rather than OOA. A key feature of these example spectra is the changing normalised signal intensities at m/z 43 and 44, with HOA being dominated by m/z 43, SV-OOA being closer to a 1:1 ratio and LV-OOA being dominated by m/z 44.

This is illustrated in Fig. 5b by the relationship between the organic signal intensity at m/z 43 and 44, which are both normalised to the total OM loading. Both individual data points from all of the flights and the normalised signal intensities from the resolved factor components for each flight are shown. This shows that as the m/z 43:OM ratio decreases, the m/z 44:OM ratio increases. This is suggestive of the OM burden aging as a continuum in terms of its oxygen content from freshly formed OOA, through to highly aged OOA which exhibits a high resemblance to fulvic acid-like mass spectra. The flight operations, given their transient nature, tend to probe this continuum which is a consequence of the constant evolution of the OM component on the regional scale. This is consistent with the difficulty in separating the OOA from the HOA in the factor analysis. The relative contribution of m/z 43 and 44 to the PMF factor components also vary from flight to flight. Clearly, while the generic terms OOA-1 and OOA-2 have been widely used to label different factors retrieved by PMF analysis, they are not chemically identical but vary from one dataset to another. Hence they also exhibit this continuum feature which can be seen in Fig. 5b. The mass spectra shown in Fig. 5a are identified on Fig. 5b in order to distinguish the general classification of the OM components. The factor components retrieved by this dataset indicate that the OOA-1 and OOA-2 are separated in terms...
of their $m/z$ 44:OM ratio, with OOA-2 being less than 0.10 and OOA-1 being greater than 0.14. This is reflected by the green dashed horizontal line on Fig. 5b between these two clusters. B357 is the only flight where the second factor is interpreted as being dominated by HOA rather than SV-OOA due to its high resemblance to reference spectra and good agreement with literature values for POA to primary emission tracers. The two factors (from B365 and B371) which also exhibit reduced signal at $m/z$ 44 do not agree as well with such literature values. Furthermore, they are partially oxidised ($m/z$ 44=0.01) and as shown in Fig. 4b, their absolute concentrations are significantly greater than the estimated HOA. Thus the second green dashed horizontal line is used to separate the B357 factor from the other OOA-2 components identified. The identified continuum is consistent with a PMF analysis of a worldwide AMS ground-based dataset by Ng et al. (2009). The black dotted lines in Fig. 5b show the general relationship between the $m/z$ 44:OM ratio and the $m/z$ 43:OM ratio from this worldwide dataset.

Based upon the framework presented by Jimenez et al. (2009), we classify the OOA-1 components as LV-OOA and the OOA-2 as SV-OOA except for the B357 OOA-2 factor, which is classified as HOA. Aiken et al. (2008) showed that the contribution of $m/z$ 44 to the OM is an excellent proxy for the Oxygen-to-Carbon (O:C) ratio of OM. Such an observation is attributed to SOA formation and photochemical aging. Thus the observed continuum of organic evolution is reflected by progressive aging from a SV-OOA dominated burden to a LV-OOA dominated burden.

The separation of the factor profiles in terms of their normalised $m/z$ 44 signal highlights that retrieved PMF profiles tend to be strongly determined by the extremes in the examined dataset as the actual data predominantly fall within a band of relatively oxidised OM between 0.10–0.20. The extremes on either side of this band are representative of very fresh and very aged OM respectively. The key question regarding this analysis technique is whether static PMF factor profiles are capable of reflecting changes in a continuum i.e. is it possible to simplify the observed evolution using such a limited number of factors? The variability in terms of the mass spectral fingerprints of the LV-OOA and SV-OOA components shown in Fig. 4b highlight this point. Such variations are likely a complex combination of both atmospheric/chemical processes and instrumental variability from flight to flight (and also within a single flight). Ng et al. (2009) show the same phenomenon in their multiple ground-based dataset. In order to test whether the mass fractions of LV-OOA and SV-OOA can replicate the evolution in the $m/z$ 44:OM ratio, correlation coefficients for each flight are calculated. The LV-OOA organic mass fraction and $m/z$ 44:OM have a correlation coefficient ranging from 0.55–0.98 across the dataset (see Supplementary Material Fig. S11; http://www.atmos-chem-phys.net/10/4065/2010/acp-10-4065-2010-supplement.pdf). This indicates that the dataset can reproduce the evolution in the OM by comparing the relative concentrations of the LV-OOA and SV-OOA components. Thus the retrieved factors reflect the continuity of processing by increased oxidation in the atmosphere, consistent with the processing paradigm proposed by Jimenez et al. (2009).

5 Evolution of the organic aerosol component

5.1 Transformation across Europe during a anticyclonic case study

The LONGREX-2 period presents an opportunity to probe the evolution of the chemical composition across the regional scale due to the consistent meteorological situation and reduced influence of wet deposition. Geopotential height fields and air mass back trajectories during the period are included in Supplementary Material Sect. 2, Fig. S3 (http://www.atmos-chem-phys.net/10/4065/2010/acp-10-4065-2010-supplement.pdf). The flight operations conducted during the period are summarised in Fig. 6a, where absolute mass concentrations between 250–2500 m are shown as a function of longitude. Points above 2500 m were generally outside of the boundary layer so are not included. Points lower than 250 m are omitted due to the formation of shallow layers over the sea surface decoupled from the atmosphere above. These were frequently encountered in marine regions sampled in this study. Such features, while interesting, are considered unrepresentative of the evolution of the chemical composition on the European scale. The aircraft flew on a roughly east-to-west transect across Northern Europe over the course of 4 days. At longitudes between 20 E and 25 E, concentrations were typically low (≈1 µg m$^{-3}$ for sulphate and organics) and constituted background conditions relative to the other locations sampled. Concentrations of both organic and inorganic species increased substantially to the west, with concentrations in the range of 3–7 µg m$^{-3}$ at the 75–95th percentiles, reflecting the increased density of anthropogenic sources encountered. The longitudinal gradients reveal that the median and the Inter Quartile Range (IQR) of the OM concentration is relatively constant west of 15 E, with concentrations typically between 3–5 µg m$^{-3}$. The larger IQR in the 15–20 E band is likely due to the transition from background conditions to polluted conditions. The OM concentrations contrast with the inorganic mass concentrations, which display somewhat differing trends and variability. The ammonium sulphate concentrations tend to build from east-to-west but with greater variability than the OM. This contrast is likely a consequence of the differing formation/processing time scales and source distributions for ammonium sulphate versus OM. The ammonium nitrate concentrations show enhanced variability throughout with no noticeable trend to the east of the meridian, although the median concentration increases to the west of this longitude. This is a reflection of the NO$_x$ and ammonia source fields across
Europe, with peak emissions of these emissions occurring in North-Western Europe (e.g. Reis et al., 2009). This is especially true in terms of ammonia, which is more readily available in Western Europe compared to further east due to intensive agricultural activities. The contrast with OM reflects this source distribution. Additionally, their differing formation and processing time scales discussed in Sect. 4.3 likely play a role.

In terms of the chemical evolution of the OM, this period highlights the highly dynamic nature of the OM burden and is shown in Fig. 6b. In the background conditions to the east, the LV-OOA dominates, with a median mass fraction of close to 90%. Between 15 E and 20 E, where OM concentrations sharply increase, the LV-OOA mass fraction is highly variable due to the transition from background to more polluted conditions. The enhanced OM concentrations are driven by the SV-OOA component, as more freshly formed material is encountered as the aircraft travels westwards. This input of fresher material continues further to the west as the SV-OOA fraction increases (decreasing LV-OOA fraction), which is consistent with the urban source distribution in continental Europe. The LV-OOA fraction is then relatively constant as far west as 10 W. West of this, over the Eastern Atlantic Ocean, LV-OOA undergoes a marked increase with the median and 75th percentiles being 65% and 80% respectively. This increase in the LV-OOA reflects the lack of fresh pollution contributing to new SV-OOA formation and is coincident with an increase in the median concentration of ozone beyond the zero meridian line, from 60 ppb to 80 ppb, which is suggestive of photochemical processing within the high pressure system. Based upon the aforementioned back trajectories, this represents approximately one day of air mass transit. A key observation during the LONGREX-2 period is the relationship between the LV-OOA mass fraction and the OM:ΔCO ratio shown in Fig. 6b (where ΔCO corresponds to CO minus the background value), with the enhancement in the OM:ΔCO ratio from a median value of approximately 20 µg sm$^{-3}$ ppm$^{-1}$ to over 100 µg sm$^{-3}$ ppm$^{-1}$ west of 15 E occurring upon the addition of SV-OOA mass which enhances the OM concentrations. These values then decrease west of the median to 60–100 µg sm$^{-3}$ ppm$^{-1}$ as the LV-OOA mass fraction becomes more dominant and OM concentrations decrease slightly.

De Gouw and Jimenez (2009) reported OM:ΔCO ratio’s for urban emissions containing large amounts of SOA in the range of approximately 50–90 µg sm$^{-3}$ ppm$^{-1}$ in North America and Tokyo. Thus the values reported here tend towards greater values or even exceed those reported previously. The measurements presented here are close to those reported for the highly polluted Po Valley region in Northern Italy described in Crosier et al. (2007).

The measurements are consistent with a rapidly formed OM close to source, which subsequently ages substantially downwind. This contrasts with the evolution of the inorganic

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**Fig. 6.** Boxplot summary statistics of aerosol chemical composition as a function of longitude during LONGREX-2, where the air mass transport was approximately east-to-west. (a) Summarises the concentrations for nitrate, sulphate and organics. (b) Presents the LV-OOA organic mass fraction (left) and the OM:ΔCO ratio (right). Crosses represent the mean value, while horizontal lines represent the 25th, 50th and 75th percentiles. The whiskers represent the 5th and 95th percentiles. Values are for altitudes between 250–2500 m.
constituents in the form of ammonium nitrate and ammonium sulphate, which evolve on differing time scales during this period.

5.2 Transformation with respect to distance from source and photochemical processing

The broadly similar transport patterns prevalent throughout the dataset (i.e. sampling of European air masses at distances both upwind, over continental Europe itself and at varying scales downwind) provides the potential to link the evolution of the aerosol chemical composition across this large range of spatial scales. The flight operations can be characterised according to the CO: NO\(_x\) and O\(_3\): NO\(_x\) ratios, which are qualitatively used as proxies for proximity to major sources and photochemical processing. The O\(_3\): NO\(_x\) ratio is shown versus the CO: NO\(_x\) ratio in Fig. 7c, indicating good first order agreement between their respective evolutions. In urban areas, CO concentrations are typically 5–15 times NO\(_x\) concentrations (e.g. Parrish et al., 2009) due to enhanced NO\(_x\) emissions associated with primary combustion sources. Furthermore, on urban-to-near-urban scales, NO\(_x\) levels are enhanced relative to O\(_3\) due to titration by NO. This yields an O\(_3\): NO\(_x\) ratio less than 1, thus this range in O\(_3\): NO\(_x\) and CO: NO\(_x\) is classified as near-urban. Subsequent dilution and photochemistry downwind of urban emissions will lead to an enhancement in the ratios, as NO\(_x\) is oxidised to form HNO\(_3\) and PAN on a time scale of a few hours (e.g. Neuman et al., 2009). This evolution is characterised by ratio values between 1 and 100, where O\(_3\) concentrations steadily increase (see Supplementary Material Sect. 4, Fig. S5; http://www.atmos-chem-phys.net/10/4065/2010/acpf-10-4065-2010-supplement.pdf). This increase in O\(_3\) and decrease in NO\(_x\) is characteristic of photochemical activity. Furthermore, across this range of O\(_3\): NO\(_x\), CO tends to steadily decrease, indicative of dilution downwind of its major sources. The 1–10 O\(_3\): NO\(_x\) is chosen as the near-source range as the CO concentrations are similar to those at less than 1, while O\(_3\) levels are steadily increasing from their minimum value close to urban sources. The 10–100 O\(_3\): NO\(_x\) range is then split between near-outflow and far-outflow regimes, principally based upon its gradient across the LONGREX-2 period (not shown) where it increased above 50 at longitudes to the west of 10W. Values greater than 100 are characteristic of background conditions, with reduced O\(_3\), CO and NO\(_x\) concentrations indicative of dilution.

The O\(_3\): NO\(_x\) ratio is used to characterise the flights as both measurements were available on all flights, whereas CO was absent during ADIENT-2. The results of this grouping are shown in Fig. 7 by contrasting the evolution from near-source conditions, predominantly over polluted regions of continental Europe and the associated near-field and far-field outflow from it. Also shown are the background conditions encountered, which were predominantly encountered in the Baltic Sea region, for a contrast with the more polluted regimes. The near-urban points are not included as too few were sampled to yield a statistically robust summary.

The analysis indicates that under polluted and highly photochemically active conditions at near-source locations, ammonium nitrate and OM are the dominant chemical components. Concentrations range from 4.5–10.0 µg sm\(^{-3}\) and 4.0–6.5 µg sm\(^{-3}\) at the 50–95th percentile levels respectively. The upper percentile dominance of nitrate diminishes with increasing distance from source but it still maintains a significant fraction of the sub-micron mass, with concentrations comparable to sulphate. OM is the dominant component at all scales outside of the most polluted conditions, where nitrate is dominant. The estimated HOA mass fraction is typically between 5–15% at the 25–75th percentile levels across the range of conditions sampled. The composition of the OOA evolves strongly as a function of the O\(_3\): NO\(_x\) ratio, with the LV-OOA fraction making up 50–65% of the OM at the 25–75th percentile levels close to source, through to 60–80% of the OM in the far-field outflow. In background conditions, LV-OOA makes up close to 90% of the OM at the 75th percentile and is at 100% at the 95th percentile. These features are reflected in the colouring of the Fig. 7c, with the LV-OOA increasing as a function of CO: NO\(_x\) and O\(_3\): NO\(_x\). The two apparent lines in the relationship are principally a consequence of the enhanced pollutant concentrations in LONGREX-1 (the lower line), during which NO\(_x\) and CO were enhanced thus the O\(_3\): NO\(_x\) is shifted downwards relative to the other periods shown. The LV-OOA organic mass fraction captures this change as the LONGREX-1 OM is more dominated by SV-OOA (reduced LV-OOA fraction).

The results show that significant OM concentrations are rapidly formed under polluted conditions in continental Europe and that these concentrations are maintained upon advection downwind of the major sources in Europe. Much of the enhanced mass in Europe is associated with an increase in the SV-OOA fraction but a significant fraction is associated with LV-OOA, which is indicative of rapid photochemical processing of the OM on the regional scale. The median LV-OOA fraction is always greater than the SV-OOA fraction and the LV-OOA mass fraction steadily increases upon advection downwind.

OOA across Europe was frequently well correlated with ammonium nitrate and ammonium sulphate, both of which likely originate from anthropogenic sources. Furthermore, OOA is correlated with O\(_3\): NO\(_x\), again suggestive of an anthropogenic influence. Potentially this may point to the OOA sources being anthropogenic in origin also, although this contrasts with several previous studies in Europe which have identified that SOA is predominantly derived from biogenic sources (e.g. studies in Zurich, (Szipat et al., 2006), rural/remote areas, Gelencsér et al., 2007 and Göteborg, Szipat et al., 2009). A similar picture is presented for the United States by Bench et al. (2007), who demonstrated that modern
carbon dominated in rural areas, which suggests a biogenic source. However, measurements in urban plumes and closer to major anthropogenic sources have indicated that anthropogenic Volatile Organic Compounds (VOCs) are the main source of OM (e.g. de Gouw et al., 2005, 2008; Bahreini et al., 2009). Furthermore, the presence of both significant anthropogenic and biogenic sources in Northern Europe could mean that the OOA results from a mixture of both anthropogenic and biogenic sources and/or that enhanced biogenic SOA is formed in the presence of anthropogenic pollution, which has previously been identified in the southeastern United States (Weber et al., 2007; Goldstein et al., 2009). However, the current study lacks suitable information regarding gas-phase organic precursors and/or additional OM chemical information to separate the different contributions of biogenic and anthropogenic to the total observed OM from the observations alone.

Enhanced SV-OOA concentrations in North-Western Europe are strongly coupled to ammonium nitrate concentrations, which are regularly observed to peak at the top of the boundary layer. This is prescribed to partitioning of semivolatile gas phase precursors to the particle phase at reduced temperature and enhanced relative humidity (e.g. Morino et al., 2006; Morgan et al., 2009). These observations are discussed in Morgan et al. (2010) in the EUCAARI special issue.

6 Conclusions

The spatial distribution of sub-micron aerosol chemical composition has been characterised based upon airborne measurements in the planetary boundary layer across Europe, north of the Alpine regions. Organic Matter (OM) and ammonium nitrate are the largest components, typically contributing 20–50% each to the non-refractory mass. Ammonium nitrate dominates the infrequent but highly polluted periods sampled with concentrations ranging from 4–8 µg sm\(^{-3}\) at the 75–95th percentile levels. This is consistent with the dominance of ammonium nitrate in Western Europe during periods of high pollutant concentrations identified previously (Putaud et al., 2004; Morgan et al., 2009). Ammonium nitrate dominates over sulphate over the whole of Europe, with OM concentrations typically 1.3–2.5 times greater than that of sulphate. Sulphate contributes 10–30% to the

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regional non-refractory particulate mass burden. Consideration of OM and ammonium nitrate in modelling assessments of the impact of atmospheric aerosol in Europe is evidently required. The measurements presented here provide a useful test for regional and global aerosol models, due to the contrasting distributions of the OM components and inorganic components across the significant spatial and temporal scales sampled.

A positive matrix factorisation analysis of the OM component was conducted, revealing the dominance of Oxidised Organic Aerosol (OOA) over Hydrocarbon-like Organic Aerosol (HOA), which is consistent with previous literature (Zhang et al., 2007). This dominance meant that the HOA component was difficult to separate from the OOA in a robust manner across the dataset. An empirical estimate based upon previous research indicated that HOA contributes less than 15% to the OM burden. Two factor solutions were found to be the most robust characterisation of the OM burden. Two separate OOA components were identified: one representing an aged-more oxidised organic aerosol and another representing fresher-less oxidised organic aerosol. These factors closely resemble those derived in ground-based global AMS datasets (Jimenez et al., 2009, Ng et al., 2009) where the less oxidised factor is associated with Semi-Volatile OOA (SV-OOA) and the more oxidised factor is found in the Low-Volatility OOA (LV-OOA) fraction. The OM chemical composition and the associated OOA factors derived for each flight were highly variable in terms of their oxygen content, based upon normalised organic signal intensities at m/z 44 and 43. When combined, the OM data and factors can be viewed as a continuum with a progression from a less oxidised, more-volatile component to a highly oxidised, less-volatile component. This progression was observed to occur as a function of the distance from source, with more oxidised components dominating further downwind. During highly polluted conditions in North-Western Europe, the SV-OOA displayed strong coupling to ammonium nitrate, indicative of gas-to-particle partitioning of semi-volatile components. Highly active photochemical conditions encountered during the study meant that LV-OOA was the largest OM component at all locations and this dominance increased with distance from source. In background conditions, LV-OOA made up more than 80% of the OM burden. This evolution is consistent with the recent paradigm proposed by Jimenez et al. (2009).

The processes identified in this work result in the build up of significant amounts of anthropogenically influenced aerosol downwind of major source regions with total sub-micron mass loadings from the AMS exceeding 15μg sm\(^{-3}\) with OM and ammonium nitrate being the dominant chemical components. Such concentrations have the capacity to significantly perturb regional weather and climate.

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References


