I Investigating the Annual Behaviour of Submicron

2 Secondary Inorganic and Organic Aerosols in London

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20 Abstract

For the first time, the behaviour of non-refractory inorganic and organic submicron 21 particulate through an entire annual cycle is investigated using measurements from an 22 Aerodyne compact time-of-flight aerosol mass spectrometer (cToF-AMS) located at a UK 23 urban background site in North Kensington, London. We show secondary aerosols account 24 for a significant fraction of the submicron aerosol burden and that high concentration events 25 are governed by different factors depending on season. Furthermore, we demonstrate that on 26 an annual basis there is no variability in the extent of secondary organic aerosol (SOA) 27 oxidation, as defined by the oxygen content, irrespective of amount. This result is surprising 28

given the changes in precursor emissions and contributions as well as photochemical activity
 throughout the year; however it may make the characterisation of SOA in urban
 environments more straightforward than previously supposed.

4 Organic species, nitrate, sulphate, ammonium, and chloride were measured during 2012 with average concentrations (\pm one standard deviation) of 4.32 (\pm 4.42), 2.74 (\pm 5.00), 1.39 (\pm 5 1.34), 1.30 (\pm 1.52) and 0.15 (\pm 0.24) µg m⁻³, contributing 43%, 28%, 14%, 13% and 2% to 6 7 the total non-refractory submicron mass (NR-PM₁), respectively. Components of the organic 8 aerosol fraction are determined using positive matrix factorisation (PMF) where five factors are identified and attributed as hydrocarbon-like OA (HOA), cooking OA (COA), solid fuel 9 OA (SFOA), type 1 oxygenated OA (OOA1), and type 2 oxygenated OA (OOA2). OOA1 10 and OOA2 represent more and less oxygenated OA with average concentrations of 1.27 (\pm 11 1.49) and 0.14 (± 0.29) μg m^-3, respectively, where OOA1 dominates the SOA fraction 12 (90%). 13

14 Diurnal, monthly, and seasonal trends are observed in all organic and inorganic species, due to meteorological conditions, specific nature of the aerosols, and availability of precursors. 15 16 Regional and transboundary pollution as well as other individual pollution events influence London's total submicron aerosol burden. High concentrations of non-refractory submicron 17 18 aerosols in London are governed by particulate emissions in winter, especially nitrate and 19 SFOA, whereas SOA formation drives the high concentrations during the summer. The findings from this work could have significant implications for modelling of urban air 20 pollution as well as for the effects of atmospheric aerosols on health and climate. 21

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23 1 Introduction

24 Atmospheric aerosols have adverse effects on human health (Pope and Dockery, 2006), air quality (AQEG, 2012), visibility (Watson, 2002), and climate (Boucher et al., 2013). 25 Pollution abatement is therefore important, especially in cities, when three-quarters of 26 Europe's population currently live in urban areas, a number that is expected to increase to 27 80% by 2020 (EEA, 2010). Regulations on air quality are based on PM₁₀ and, more recently, 28 PM_{2.5} (particulate matter with aerodynamic diameters less than 10 µm and 2.5 µm 29 respectively, European Union, 2008). A recent study (Aphekom Summary Report, 2011) 30 reported that life expectancy in London could increase by 2.5 months for persons 30 years of 31 32 age and older if average annual PM2.5 concentrations were decreased in line with the World

Health Organization's Air quality guidelines to 10 μ g m⁻³ (WHO, 2005). PM₁ (particulate matter with an aerodynamic diameter less than 1 μ m) is beginning to receive greater attention from the air quality community as they are associated with adverse health effects due to the depth within the lungs to which these particles can penetrate and can then enter the blood stream, and cause damage to other parts of the body (Oberdörster et al., 2005).

Primary and secondary aerosols have both natural and anthropogenic sources (Seinfeld and 6 Pandis, 2006), resulting in their diverse chemical composition, size, and concentration 7 (Pöschl, 2005). In urban areas, primary aerosols from transport, cooking, and solid fuel 8 burning are of great significance (Allan et al., 2010), particularly in the winter when 9 meteorological conditions are such that their concentrations are elevated resulting in pollution 10 events (Zhang et al., 2007; Huang et al., 2014). In addition, transported air masses frequently 11 12 influence the UK's atmosphere (Abdalmogith and Harrison, 2005) including polluted air masses from continental Europe and cleaner westerly conditions. Transported pollution 13 typically comprises secondary aerosols, with season having a strong influence on the 14 15 chemical composition and concentration (Charron et al., 2007). Previous studies highlight the variability in the contribution of both secondary inorganic and organic aerosol (SIA and 16 17 SOA, respectively) to the total mass depending on location (Jimenez et al., 2009). Furthermore, chemical composition varies with location due to a combination of local and 18 19 regional aerosol sources as well as daily and seasonal meteorological conditions.

The precursors and formation processes of SIA are relatively well understood, particularly as 20 anthropogenic emissions dominate although concentrations are significantly influenced by 21 regional and transboundary pollution. For example, Abdalmogith and Harrison (2006) 22 estimated that between 2002 and 2004, 88% of nitrate and 92% of sulphate in central London 23 originated from the regional background. Due to the non-linear response of SIA 24 25 concentrations from reductions in precursor emissions, the impacts on formation from changes in emissions are uncertain (AQEG, 2012). In contrast, the complexity of SOA 26 27 precursors, including the range of atmospheric processing they can undergo, lifetime, and 28 temporal and spatial variability presents a major challenge to understanding and 29 characterising SOA and its formation (Goldstein and Galbally, 2007). Additional variability of SOA sources and formation results from the long distances over which SOA precursors 30 31 and the resulting aerosols can be transported as well as dependency on meteorological conditions (Martin et al., 2011). Furthermore, SOA evolves in the atmosphere with properties 32

changing with age (Ng et al., 2010) meaning our ability to quantify and predict SOA remains
 limited.

The Aerodyne Aerosol Mass Spectrometer (AMS) measures size-resolved chemical 3 4 composition of non-refractory submicron particulates with high time resolution (Jayne et al., 2000; Canagaratna et al., 2007). The AMS has demonstrated its versatility in a range of 5 environments across the world (Zhang et al., 2007) and has been used to successfully 6 investigate SOA behaviour (e.g. Jimenez et al., 2009; Heald et al., 2010; Ng et al., 2010; 7 8 Kroll et al., 2011). However, despite its widespread use in such process studies, the instrument is infrequently used for long-term characterisation of aerosols. The related aerosol 9 chemical speciation monitor (ACSM, Ng et al., 2011a), however, is routinely used for long-10 term measurements of aerosol chemical composition. Here we present a year-long UK urban 11 12 background data set collected with a compact time-of-flight AMS (cToF-AMS) including results of positive matrix factorisation (PMF) analysis, which is the first time the AMS has 13 14 been used in this way in an urban environment. The temporal trends and contributions of 15 urban aerosols to PM₁ are evaluated and their sources are investigated. In this paper we will focus on the secondary aerosols; though primary organic aerosol sources are identified in this 16 17 paper, the behaviour of primary aerosols from these sources will be discussed in subsequent publications. 18

19 In Section 2 of the paper, the experimental site, instrumentation, and analysis methods utilised in this study are described. In Section 3, an overview of the bulk non-refractory PM₁ 20 (NR-PM₁) components including average mass, diurnal profiles, and seasonality is presented 21 along with a discussion on the factors governing concentrations and temporal trends. In 22 Section 4, the components of the organic fraction are investigated using receptor modelling. 23 In Section 4.3, we investigate two covarying factors derived from PMF analysis, with the 24 25 method used to estimate the concentrations of the two factors described in Section 4.4. In 26 Section 5, the organic components are identified and the results from the previous sections 27 are used to probe the behaviour of urban SOA including temporal trends (Section 5.1) and 28 state of oxidation (Section 5.2). In Section 6, the factors governing pollution events across the 29 year, as well as winter and summer, are assessed through identification of the dominant components of the high concentration events. Finally, Section 7 summarises the conclusions 30 31 from this study on secondary aerosols in London.

1 2 Experimental

2 2.1 Site and instrumentation

The measurements for this study were conducted as part of the Natural Environment 3 Air for London 4 Research Council (NERC) funded Clean (ClearfLo) Project (www.clearflo.ac.uk), a large, multi-institutional collaborative scientific project based in the 5 6 UK. A suite of state-of-the-art instrumentation, measuring aerosols, gases, radicals and meteorological parameters was deployed for two major intensive observation periods (IOPs) 7 8 during 2012, with long-term continuous measurements conducted between 2011 and 2013. 9 Measurements were conducted at the ClearfLo urban background site in the grounds of a 10 school in North Kensington (51.521055 N, 0.213432 W), where a permanent Department for Environment, Food and Rural Affairs (DEFRA) Automated Urban and Rural Network 11 12 (AURN, http://uk-air.defra.gov.uk/networks/network-info?view=aurn) monitoring station is located. A background site is defined by DEFRA as being located such that its pollution level 13 is not influenced significantly by any single source or street, but rather by the integrated 14 contribution from all sources upwind of the station" and "be representative for several square 15 kilometres". Situated in a residential area 7 km to the west of Central London, the sampling 16 site is not influenced by heavily trafficked roads and is representative of background air 17 quality (Bigi and Harrison, 2010). Along with the school buildings, a car park and a relatively 18 large playing field are also located at the site with several large trees both on site and lining 19 20 the surrounding pavements. Further details on the ClearfLo experimental campaigns and 21 locations are described in Bohnenstengel et al. (2014).

22 Aerosol chemical composition was measured by the cToF-AMS for a full calendar year (11 23 January 2012 – 23 January 2013) and by the high-resolution time-of-flight AMS (HR-ToF-24 AMS) during the two IOPs, which were conducted in the winter (January - February) and summer (July – August) of 2012. The cToF-AMS sampled through a PM_{2.5} inlet, with a 25 bypass flow of 16 l min⁻¹ and split using an asymmetric Y-piece. The HR-ToF-AMS was 26 located in a shipping container containing several other aerosol instruments, where aerosols 27 were sub-sampled from a sampling stack with a flow of 30 1 min⁻¹ via a 3.5 µm cut-off 28 cyclone. 29

Both AMS instruments operated in the standard configuration and took mass spectra (MS) and particle time of flight (pToF) data. An overview of the AMS can be found in Canagaratna et al. (2007) and detailed descriptions of both the cToF-AMS and HR-ToF-AMS can be

1 found in Drewnick et al. (2005) and DeCarlo et al. (2006) respectively. The instrument 2 operation and data analysis procedures pertinent to this study have been described elsewhere (e.g. Allan et al., 2010). The HR-ToF-AMS operated in both 'V' and 'W' ion path modes, 3 offering high sensitivity but low mass resolution, and low sensitivity but high mass 4 5 resolution, respectively. Only the V mode ambient data are analysed further here due to their better signal-to-noise ratio. The time-resolution of the cToF-AMS was 5 minutes throughout 6 7 the measurement period. As the HR-ToF-AMS sampled in an alternating sequence with other 8 black carbon and aerosol volatility measurements using a thermodenuder (Huffman et al., 9 2008) in the winter, 5-minute averaged ambient data in V mode were only obtained every 30 minutes. In the summer, there were no volatility measurements so average data were obtained 10 every 12 minutes. Both instruments were calibrated using 350nm mono-disperse ammonium 11 nitrate particles approximately once a month for the cToF-AMS and weekly during the IOPs 12 for the HR-ToF-AMS. Ammonium sulphate calibrations were also performed where possible. 13 The heater bias of the cToF-AMS was tuned to minimise the signal from surface ionised 14 15 potassium and the filament was run at a lower value than usual in order to prolong the life of the multi-channel plate (MCP). This configuration results in a reduced signal, which in turn 16 reduces the signal-to-noise ratio (Allan et al., 2003). 17

18 **2.2** Analysis and quality control of AMS data

CToF-AMS data were analysed within Igor Pro (Wavemetrics) using the standard TOF-AMS 19 analysis toolkit software package, SQUIRREL (SeQUential Igor data RetRiEval) v1.53. The 20 21 HR-ToF-AMS data were analysed using SQUIRREL v1.52J and PIKA v1.11J (Sueper, 2008). An average ionisation efficiency (IE) determined from all calibrations was applied to 22 23 the full data set. Relative ionisation efficiencies (RIEs) of ammonium, nitrate, and sulphate were estimated based on the molar ratios of each species from the ammonium nitrate 24 25 calibrations (see Table S1 in the Supplementary Information for the ammonium and sulphate RIE values for the cToF-AMS and HR-ToF-AMS). These were compared to particulate 26 27 sulphate measurements from the URG-9000B Ambient Ion Monitor (AIM) from North Kensington (AURN and Particle Numbers and Concentrations Network, http://uk-28 29 air.defra.gov.uk/networks/network-info?view=particle) where available (Fig. S1) and indicated 30 that the default RIE of sulphate of 1.2 may not be appropriate for either instrument. As a sulphate calibration was not performed on the HR-ToF-AMS during the winter IOP the RIE 31 32 was ambiguous, so concentrations were based on those reported by the cToF-AMS, which

was calibrated later during the campaign (see Section 2.1 in Supplementary Information for a 1 2 comparison of the concentrations between the two instruments for the winter IOP and Section 2.2 for the summer IOP). This approach, as opposed to using the default RIE of 1.2, was 3 deemed valid as it resulted in a more consistent volume concentration comparison with that 4 5 derived from a differential mobility particle sizer (DMPS) from the winter IOP (Fig. S2) where the volume concentration was estimated using the densities reported by Cross et al. 6 7 (2007). A time and composition dependent collection efficiency (CE) was applied to the data based on the algorithm by Middlebrook et al. (2012). This was also validated for both AMSs 8 9 by comparing the volume concentration with that derived from the DMPS measurements from the winter IOP (Figs. S2a and d). 10

11 Inspection of the data revealed step changes in cToF-AMS mass concentrations that coincided with changes in the flowrate, which were mostly due to partial blockages in the 12 13 pinhole (see Section 1.2 in the Supplementary Information). In each case, the pinhole was 14 either manually cleaned (through sonication in deionised water) or the flow returned to its 15 average rate of 1.3 cc/sec without intervention. Data were removed if clear mass changes were observed, with distinct start and end points (e.g. 2 June 2012, Fig. S3b). Other data were 16 17 flagged as suspect if the flow was significantly different from its normal rate (less than 1.2 cc/sec) but there were no distinct step changes in mass e.g. 4 September 2012 (Fig. S3c). The 18 19 final data set comprised 95% data that had not been removed or flagged as suspect.

20 2.3 Levoglucosan measurements

24-hour PM_{2.5} samples were collected on quartz fibre filters (Whatman QM-A) at NK during 21 22 the winter 2012 ClearfLo campaign using a high volume Digitel DHA-80 sampler at a flow 23 of 500 l/min. These samples were analysed for wood smoke marker levoglucosan using a 24 slightly modified version of the method of Yin et al. (2010) and Wagener et al. (2012). In brief, one portion of the Digitel filter sample was spiked with an internal standard (IS), 25 methyl-beta-D-xylopyranoside Sigma-Aldrich Ltd) and 26 (from extracted with dichloromethane and methanol under mild sonication at room temperature. The combined 27 extract was filtered and concentrated down to 50 µl. One aliquot of the extract was 28 29 evaporated dryness and derivatised by addition of N,Oto near bis(trimethylsilyl)trifluoroacetamide plus 1% trimethylchlorosilane (BSTFA + 1% TMCS) 30 and pyridine at 70°C for 1h, and finally cooled in a desiccator. Quantification was based on 31 the IS and a six point authentic standard calibration curve, using the selected ion monitoring 32

(SIM) mode on an Agilent GC-MS instrument. The ions monitored were 204 and 217 for the
 IS and 204, 217 and 333 for levoglucosan.

3

4 3 Results

5 The daily averaged time series of NR-PM₁ species, their diurnal patterns, and monthly average contributions to total submicron mass are shown in Fig. 1. On average, PM₁ 6 7 composition is dominated by the organic fraction (Org, 44%, Fig. 2) with the remainder of the total mass comprising SIA species. Nitrate (NO₃) is the largest SIA component, 8 9 comprising 28% of the total mass. Sulphate (SO₄) and ammonium (NH₄) contribute 14% and 10 13% respectively with a small contribution from non-refractory chloride (Chl, 1%). The 11 contribution of each species to the total mass varies with time; organics dominate in summer 12 and inorganics dominate in winter, with nitrate contributing up to 45% of the total mass in 13 spring (Fig. 1c).

Organic species constituted, on average, just under half of the non-refractory submicron mass 14 as measured by the AMS in 2012 (44%, Fig. 2), with a mean annual concentration (± one 15 standard deviation) of 4.32 (\pm 4.42) µg m⁻³. During the year, concentrations at times 16 increased up to, and over, an order of magnitude greater than this value with a maximum 5 17 minute concentration of 230 µg m⁻³, observed on 18 February. This is likely a locally sourced 18 event lasting approximately 6 hours as the maximum daily concentration was 16.97 μ g m⁻³, 19 observed on 25 July (Fig. 1a). The mean organic concentrations and diurnal patterns exhibit 20 21 little seasonality (Figs. S6a and b, respectively); a large evening peak is observed in all diurnal profiles but the number of peaks and their timing during the day vary slightly with 22 23 season. Although the total mass of organic species exhibits little seasonality, the organic 24 fraction of total PM₁ varies with season, being largest in summer and autumn.

The average annual PM1 nitrate concentration was 2.74 (± 5.00) $\mu g \text{ m}^{-3}$ with several high 25 26 concentration episodes occurring throughout the year (Fig. 1a). Peak events occurred mainly during the winter and spring, with a maximum 5 minute concentration of 48.35 μ g m⁻³ 27 measured on 23 March. Increases in the concentrations of all species are also observed 28 during these high nitrate events. Averaged across the year, nitrate exhibits a pronounced 29 diurnal pattern with an overnight increase in mass, peaking at 08:00 UTC, with a daytime 30 minimum at 16:00 UTC (Fig. 1b). The overall shape of the diurnal pattern varies little with 31 32 season although it becomes less pronounced in summer and autumn (Fig. S7b). In contrast, the total nitrate mass varies significantly with season (Fig. S7a), where the greatest concentrations are observed during the spring, which is also when the diurnal pattern is most pronounced due to a large range of concentrations. The lowest concentrations and smallest diurnal range occur during the summer months.

Submicron sulphate represents approximately 25% of the inorganic fraction, with a mean 5 concentration of 1.39 (\pm 1.34) µg m⁻³. The maximum 5 min sulphate concentration measured 6 in 2012 was 12.75 µg m⁻³ which occurred on 2 May. In general, increases in sulphate mass 7 are coincident with increases in concentration of other AMS measured species. In contrast to 8 nitrate, sulphate exhibits little seasonality although it dominates SIA mass in summer, with 9 10 higher mean concentrations occurring in spring and summer compared to autumn and winter (Fig. S8a). Furthermore, sulphate exhibits little diurnal variation for each season as well as 11 12 for the whole year (Figs. S8b and 1b).

Almost a quarter of the inorganic mass fraction is comprised of ammonium, with a mean 13 concentration of 1.30 (\pm 1.52) µg m⁻³. Averaged across the year, ammonium exhibits a weak 14 diurnal profile (Fig. 1b). However, this pattern varies with season, with a peak in 15 16 concentration between 08:00 and 10:00 UTC in all but the summer months (Fig. S9b). The most pronounced diurnal variability occurs in spring, which is also when there is the greatest 17 seasonal mass (Fig. S9a) and maximum concentration of the year (14.23 μ g m⁻³). The aerosol 18 was found to be neutral throughout the year as the balance between inorganic cationic and 19 anionic charge was maintained. 20

As the AMS does not detect chloride salts such as sodium chloride, the chloride measured here is primarily ammonium chloride. Although this represents a very small fraction of SIA, with an average concentration of 0.15 (\pm 0.24) µg m⁻³, some seasonal differences are apparent. Chloride exhibits a weak diurnal pattern with slightly higher concentrations at night compared to during the day (Fig. 1b), which changes with season (Fig. S10b). The highest chloride concentrations are in the winter with comparatively low concentrations in the summer (Fig. S10a).

3.1 Behaviour of bulk PM₁ components

29 3.1.1 Organic aerosols

Weak seasonality of organic aerosols in Paris has been previously suggested (e.g. Freutel et al., 2013) and observed in organic carbon (OC) measurements in Birmingham (Harrison and

Yin, 2008). The lack of seasonality arises because of the balance of sources that govern the 1 2 total concentration of organic species differently during each season (Zhang et al., 2007) rather than the constancy of any particular source. As well as differences in sources with 3 season, increased organic concentrations in winter are due to low temperatures and reduced 4 atmospheric mixing, whereas in summer similar concentrations are due to increased 5 photochemistry (Martin et al., 2011). In contrast to absolute mass, there are differences in the 6 7 organic fraction of total PM₁ with season, which have been observed in Paris (Crippa et al., 2013a; Freutel et al., 2013), Tokyo (Takegawa et al., 2006), and Zurich (Lanz et al., 2007), 8 9 and can be attributed to seasonal differences in concentrations of other species such as nitrate. Consistent with previous observations, organics in London exhibit little seasonality both in 10 terms of mass and diurnal profile. Any variations in diurnal pattern across the year are due to 11 both mixing layer height dynamics and the nature of the dominant source. The components of 12 the organic aerosol fraction are discussed in more detail in Section 4. 13

14 3.1.2 Nitrate

15 The annual cycle of nitrate mass is significantly influenced by season (Martin et al., 2011), driven by emissions of ammonia, which typically peak in the spring (Schaap et al., 2004), as 16 well as temperature and relative humidity (RH), which both control nitrate partitioning 17 (Stelson and Seinfeld, 1982). The diurnal pattern of nitrate in urban locations (e.g. Cork, 18 Dall'Osto et al., 2013; Paris, Freutel et al., 2013) is also largely governed by the semi-volatile 19 20 behaviour of ammonium nitrate. However, nitrate formation also strongly depends on availability of precursor gases (Ansari and Pandis, 1998) such as nitrogen oxides (NO_x) and, 21 22 in particular, ammonia, as emissions in urban environments are small compared to NO_x 23 (NAEI, 2013). Although some non-agricultural sources of ammonia are known (Sutton et al., 24 2000), their strengths and trends are not well understood. Pollution from continental Europe has also been identified as an important contributor to particulate concentrations in many 25 26 regions (e.g. Manchester, Martin et al., 2011; Paris, Freutel et al., 2013) with the highest nitrate concentrations occurring over North West Europe during pollution episodes (Morgan 27 et al., 2010). 28

29 Consistent with previous UK measurements (Harrison and Yin, 2008; AQEG, 2012), the 30 highest concentrations in this study occurred in spring. Although more pronounced in winter 31 and spring, the overall shape of the diurnal profile does not change with season, indicating 32 the strong semi-volatile behaviour of nitrate. Also consistent with previous studies (e.g. Abdalmogith and Harrison, 2005), increased nitrate concentrations occur in air masses influenced by continental North-Western Europe, indicating the importance of transboundary pollution. Nitrate concentrations are therefore governed by a combination of season, ambient conditions, availability of precursor emissions, and air mass trajectory rather than any one factor. Consequently, it was not possible to establish simple metrics that could be used to predict nitrate concentrations, highlighting the need for detailed modelling of aerosol chemistry and thermodynamics to accurately predict nitrate concentrations.

8 3.1.3 Sulphate

9 Sulphate concentrations have been decreasing at both urban and rural UK locations for at least the last 10 years (as summarised in Table 1) due to decreasing SO₂ emissions (Monks et 10 11 al., 2009). However, sulphate concentrations respond non-linearly to reductions in SO_2 emissions (Megaritis et al., 2013). The mean sulphate concentration (1.39 µg m^{-3}) measured 12 by the AMS in 2012 is comparable to the non-sea salt sulphate $(1.21 \ \mu g \ m^{-3})$ calculated from 13 AIM measurements also at North Kensington. The 2012 AMS measurements are therefore 14 consistent with the trend of decreasing sulphate concentrations observed at sites at North 15 Kensington and Harwell. Similar to the findings of Harrison et al. (2012) and Abdalmogith 16 and Harrison (2006), sulphate exhibits little seasonality and diurnal variation thus 17 emphasising the importance of regional pollution. 18

19 3.1.4 Ammonium

Changes in the diurnal profile and total mass of ammonium with season are very similar to those of nitrate and, to a lesser extent, sulphate (Morgan et al., 2009; Bressi et al., 2013). The springtime peak in concentrations is governed by the greater availability of ammonia and favourable meteorological conditions.

24 3.1.5 Chloride (non-refractory)

The seasonal variation and diurnal pattern of chloride is attributed to the semi-volatile nature of ammonium chloride as well as planetary boundary layer dynamics where low concentrations are expected during the summer due to increased mixing depth. The availability of ammonia will also govern the concentration of chloride. In addition, increased chloride concentrations in the winter could be attributable to increased coal combustion during this period (Sun et al., 2013).

4 Positive matrix factorisation analysis

2 To investigate the components and temporal trends of the organic fraction, positive matrix factorization (PMF) (Paatero and Tapper, 1994; Lanz et al., 2007) was applied to the organic 3 4 matrix from the year-long cToF-AMS data set, which is the first time PMF has been applied to a data set of this duration from an urban environment. Separating long-term data into 5 seasons before performing factorisation analysis may be used to reduce seasonal phenomena 6 affecting the retrieved factors, such as minimising the influence of variations in 7 8 photochemistry and also to address known PMF limitations such as mixing between factors. 9 However, splitting the data into seasons is subjective, resulting in a bias of the retrieved factors and loss of information on annual trends of potential OA components. 10

Identification of key organic components can also be achieved by using the related 11 12 multilinear engine ME-2, for which a protocol for the AMS is currently available (Canonaco et al., 2013) and has been found to produce more representative results in some circumstances 13 (e.g. Lanz et al., 2008), particularly when temporal co-variation of factors arises. However, a 14 15 priori knowledge by way of factor profiles and/or time series is required to utilise ME-2, so in principle it is preferable to obtain a factorisation without a priori assumption, which is 16 achieved in this study by way of unconstrained PMF analysis. Furthermore, several of the 17 18 factorisation problems that ME-2 overcomes when applied to data from the ACSM (Ng et al., 19 2011a) compared to the AMS are related to the fact that the ACSM has much lower signal-to-20 noise ratio (approximately by a factor of 40). We present the results from PMF analysis here to compare with earlier work and as a first stage in any further analysis. Furthermore, as we 21 will show in the data presented, temporal co-variation of factors can be overcome by careful 22 scrutiny of the data as well as from the use and support of associated measurements such as 23 24 from the HR-ToF-AMS.

25 4.1 Data preparation

PMF was performed on the organic data matrix for the year-long data set from the cToF-AMS and for the winter and summer periods when the HR-ToF-AMS was operating. The data preparation for all three data sets followed the recommended procedures as described by Ulbrich et al. (2009). However, for the final PMF solution, the summer period was removed from the cToF-AMS data set, as the mass spectrometer was mistakenly re-tuned for this period, which caused problems for the factorisation. The changes in the instrumental settings were evident in the data as the concentrations of several of the factors derived from PMF

analysis increased simultaneously with a step change in the heater bias. However, due to the 1 2 nature of the affected factors and the timing of the instrumental changes, it was not possible to calculate a reliable scaling factor to apply to the data from this period. The reader is 3 directed to Sects. 4.1 and 4.2 in the Supplement for more details regarding the data pre-4 5 treatment and quality assurance, including the identification and removal of problematic data 6 around the summer IOP. In addition to the standard methods, isotopes were not included in 7 the HR-ToF-AMS organic matrix. The peaks at m/z 30 and 46 were removed from the matrix, as they were not deemed to have been successfully retrieved using PIKA. APES light 8 9 v1.05 (Sueper, 2008) was used for the elemental analysis of the HR-PMF factors.

10 4.2 Factorisation results

A 5-factor solution to the PMF analysis was shown to be optimum for the cToF-AMS data set. The details of the choice of factors and solution criteria can be found in the Supplementary Information, Section 4.3. The reader is referred to Section 5 in the Supplementary Information for the HR-ToF-AMS PMF (HR-PMF) solution criteria, where 5factor solutions were chosen for both the winter and summer IOPs (Sections 5.1 and 5.2 in the Supplementary Information, respectively). The cToF-AMS PMF (cToF-PMF) solution criteria are briefly outlined here.

18 The 5-factor solution resulted in a better separation of the mass spectral profiles compared to the 4-factor solution, with improvements to diagnostics, such as Q/Q_{expected}, used to assess the 19 20 quality and suitability of a solution set. The 6-factor solution was discarded due to the similarity of several factors (spectra and time series). The 7-factor solution was also 21 22 discarded due to its significant dependency on the initialisation seed (unlike the solutions 23 with fewer factors) as well as the production of a factor that did not appear physically 24 meaningful. The 'fPeak' parameter was used to explore the rotational ambiguity of the 5-25 factor solution with the most central solution (fPeak=0) chosen for further analysis. Additional measurements were used to validate the chosen solution and for attribution of the 26 27 factors.

Three of the five PMF factors were clearly identifiable: hydrocarbon-like OA (HOA), cooking OA (COA), and type 1 oxygenated OA (OOA1). As the remaining two factors (labelled here as $SFOA_{PMF}$ and $OOA2_{PMF}$) exhibited similar temporal features, notably the diurnal pattern (Fig. 3) with an evening peak in concentration, they are investigated and addressed in detail in the following sections.

1 4.3 Identifying PMF limitations

2 The similarity of the diurnal patterns of SFOA_{PMF} and OOA2_{PMF} is likely due to the nature of 3 the aerosols where SFOA_{PMF} is likely emitted from domestic space heating, an activity that occurs in the evening. OOA2_{PMF} is typically thought to be semi-volatile oxygenated OA (SV-4 5 OOA) and will preferentially partition to the particle phase when temperatures are low and RH is high, again most likely in the evening. Conversely, the temporal co-variation of the 6 PMF solution could result in partial mixing of these two factors (Crippa et al., 2013b) leading 7 to the identification of an OOA2-BBOA factor (Crippa et al., 2013a). However, a clearer 8 separation of such factors was obtained through combined AMS and Proton Transfer 9 Reaction Mass Spectrometry (PTR-MS) PMF analysis (Crippa et al., 2013b). 10

The mass spectral profiles and time series of the cToF-PMF factors are compared to the 11 12 winter IOP HR-PMF factors, as factor retrieval from HR-ToF-AMS data is more robust with significantly reduced rotational ambiguity and improved separation of factors as individual 13 14 ion signals at the same nominal mass-to-charge ratio (m/z) are included (see Section 6 in the Supplementary Information for comparisons of the mass spectra and time series from the 15 winter and summer IOPs where available). In general, there is good correlation between most 16 factors from the two instruments (Pearson's r of 0.69-0.90, Table 2). However, the 17 concentration of the combined SFOA factors from the winter HR-PMF data set is 18 19 approximately double that of the cToF-PMF SFOA_{PMF} factor. A near equal concentration of SFOA from both AMSs is achieved when the cToF-PMF OOA2_{PMF} is combined with the 20 SFOA_{PMF} and correlated with the sum of HR-PMF SFOA factors. This suggests that most of 21 the SFOA_{PMF} mass measured by the cToF-AMS is being assigned to OOA2_{PMF} in PMF; the 22 total SFOA mass could therefore be up to a factor of two greater than previously estimated. 23

24 If SFOA represents all of levoglucosan and other similar species, we might expect good correlation between SFOA and levoglucosan to exist. As org60 (the organic fraction at m/z 25 26 60) has contributions from fatty acids arising from cooking POA emissions (Mohr et al., 2009) and carboxylic acids from SOA (e.g. DeCarlo et al., 2008), it is not expected that org60 27 28 and levoglucosan would correlate exactly when compared. **SFOA**_{PMF} and SFOA_{PMF}+OOA2_{PMF} are compared to 24-hour filter measurements of levoglucosan from the 29 30 winter IOP. SFOA_{PMF}+OOA2_{PMF} correlates better with levoglucosan than SFOA_{PMF} on its own (Pearson's r of 0.74 and 0.71 respectively), suggesting that some of the additional 31 32 variance is carried by a levoglucosan contribution to OOA2_{PMF}. Furthermore, org60 correlates slightly better with levoglucosan than SFOA_{PMF} (Pearson's r = 0.73), again 33

1 suggesting that SFOA_{PMF} is not capturing all the variability of levoglucosan. However, it is 2 unlikely that this is the full explanation as the m/z 60 signal of OOA2_{PMF} is relatively small. 3 The possibility that OOA2_{PMF} could be an overlap of what is OOA1 in the summer and an 4 aged SFOA in the winter has been explored but correlations between OOA2_{PMF} and nitrate 5 suggest that this is not likely to be the case.

This is not to suggest that all OOA2 factors contain some contribution of SFOA. However, if 6 SFOA is convolved with OOA2, as is the case in this study, it is possible to estimate the 7 proportion of SFOA convolved with OOA2 with the support of additional measurements. In 8 this study we have shown that comparisons of measurements from the two AMSs highlighted 9 10 a large difference in SFOA concentrations, which was further supported by levoglucosan measurements and those from the cToF-AMS itself such as org60. Improved separation of 11 OA factors may be achieved in the future, particularly in the absence of supporting 12 measurements, from the application of ME-2 to similar data sets such as those from the 13 14 ACSM. However, further work is therefore required to better resolve the issues arising from 15 PMF analysis regarding the separation of OA in to its primary and secondary constituents, particularly for long-term data sets. 16

17 4.4 Estimating concentrations of convolved factors

We infer from the correlations discussed in Section 4.3 that nearly all the SFOA_{PMF} is assigned to $OOA2_{PMF}$ during the winter IOP, where the proportion of SFOA_{PMF} that is convolved with OOA_{PMF} can be determined using the relationship between SFOA_{PMF} and OOA_{PMF} from the winter. Both factors have similar, strong diurnal profiles, the effect of which is reduced by using daily averages of each factor in the following equation:

$$23 \quad 00A2_{PMF} = a.SFOA_{PMF} + 00A2_{noSF} \tag{1}$$

where *a* is the gradient of an orthogonal distance regression fit, equal to 0.86, and $OOA2_{noSF}$ is the intercept which indicates the amount of $OOA2_{PMF}$ without a solid fuel signature. The remainder is the SFOA_{PMF} assigned to $OOA2_{PMF}$ during the PMF analysis and is estimated based on the gradient of the fit. The SFOA and OOA2 concentrations, SFOA_{mod} and $OOA2_{mod}$ respectively, can therefore be calculated using the following equations:

$$29 \quad SFOA_{mod} = SFOA_{PMF} + (a.SFOA_{PMF}) \tag{2}$$

$$30 \quad OOA2_{mod} = OOA2_{PMF} - (a.SFOA_{PMF}) \tag{3}$$

1 where $OOA2_{mod}$ in equation 3 is equivalent to $OOA2_{noSF}$ in equation 1. The relationship 2 based on the winter correlation does not hold true for the whole year and so the annual estimations are improved by using the relationship derived between the daily averages of 3 $OOA2_{PMF}$ and $SFOA_{PMF}$ from December 2012-January 2013, where a is equal to 0.52. 4 However, instrument maintenance (changes in instrument tuning during the summer and 5 change of the MCP in April) will likely add some variation to these estimates. Therefore, the 6 7 concentrations up until the summer period are estimated using the relationship from the 2012 winter IOP and the concentrations after the summer are estimated using the December 2012-8 9 January 2013 relationship. The estimated concentrations of OOA2_{mod} and SFOA_{mod} are used for further analyses. Figure 4 shows the retrieved OOA2_{mod} is dominated by noise, with an 10 average OOA2_{mod} concentration of 0.12 μ g m⁻³ and standard deviation of 0.46 μ g m⁻³ during 11 the winter IOP. The standard deviation provides a measure of uncertainty in our retrieval of 12 OOA2_{mod} and SFOA_{mod} using this approach. 13

14

15 5 Attribution and contributions of organic components

Attributing the PMF factors to different organic sources and components allows the organic 16 17 fraction to be split in to primary OA (POA) and secondary OA (SOA) and their contribution to total organic mass to be assessed. The behaviour of urban secondary OA (SOA) can then 18 19 be investigated. The HOA, COA, and SFOA_{mod} factors identified in the previous sections are grouped as primary OA (POA) and OOA1 and OOA2_{mod} are grouped as secondary OA 20 21 (SOA). The primary fraction has the greatest contribution from SFOA_{mod} 38% (Fig. 2), with smaller contributions from HOA (32%) and COA (30%). The secondary fraction is 22 23 dominated by OOA1 (90%) with only a small contribution from $OOA2_{mod}$ (10%).

The greatest contribution of the organic components to total OA mass, which does not 24 include some of the summer period (see Sect. 4.1), is from OOA1 (31%), followed by 25 SFOA_{mod} (25%), HOA (21%), and COA (19%). The remainder comprises OOA2_{mod} (4%). 26 During 2012, POA and SOA contributed 65% and 35% to total OA, respectively (Fig. 2). 27 28 However, the contribution of POA and SOA to total OA changes with season where SOA 29 contributes just over 50% on average during the spring and summer (Fig. 5). The smaller annual contribution from SOA could therefore be partly due to the omitted summer data, 30 31 where SOA dominates the mass fraction. However, the mean (and standard deviation) ozone 32 mixing ratio was not found to be statistically significantly different between the 5 week

- 1 period not included in PMF analysis and the whole June, July, and August period, suggesting
- 2 the data that is included in the analysis is representative of the data that were removed.

3 5.1 The behaviour of secondary organic aerosol in London background air

4 The average (\pm one standard deviation) OOA1 concentration observed was 1.27 (\pm 1.49) µg m^{-3} , with a maximum 5 minute concentration of 19.5 µg m^{-3} measured on 24 May 2012. 5 OOA1 does not exhibit a discernible diurnal pattern (Fig. 3), where the only change with 6 season is by way of concentration (Fig. 6a), suggestive of aged aerosol of a regional nature. 7 8 The peak in concentrations occurs in spring, where the average concentration is more than 9 double that of the autumn and winter and 1.7 times greater than the summer (Fig. 6b). This 10 spring time peak is consistent with secondary OC measurements in Birmingham (Harrison and Yin, 2008). 11

12 In comparison, the OOA2_{mod} concentration averaged 0.14 (\pm 0.29) µg m⁻³ over the year, with

maximum daily concentrations occurring in the summer. The seasonal trend of OOA2_{mod} is in
keeping with it being secondary in nature with concentrations increasing during the summer
(Fig. 6c) when photochemical processes and emissions of biogenic volatile organic
compounds (VOCs) (Holmes et al., 2014) are greatest.

Several high concentration events lasting 3-8 days are observed in both OOA1 and OOA2_{mod} time series (Fig. 4) such as in May (peaking on 27 May) and to a lesser extent September (peaking on 8-9 September). The event in May is associated mostly with Easterly conditions, likely the result of imported pollution. The September event is associated with a high-pressure system centred just off the SW UK coast with another high pressure system over continental Europe the following day. This resulted in an increase in concentrations in a stagnant air mass with additional imported pollution on the 9 September.

24 5.2 SOA chemistry and oxidation state

SOA forms in the atmosphere from the gas-phase oxidation of a number of VOCs (e.g. Goldstein and Galbally, 2007), which can be anthropogenic or biogenic in origin. SOA comprises a mixture of organic compounds with differing volatilities (Donahue et al., 2012) which partition between the gas and particle phases. SOA therefore exists across a variety of chemical states thus increasing its chemical complexity. As bulk chemical characterization of aerosols can be obtained from the AMS, several metrics and graphical representations of the data are used to investigate OA. The information gleaned from such metrics can be used to better inform models on SOA characteristics to improve the quantification and prediction of
 SOA.

One such metric for describing and explaining OA evolution in the atmosphere is the f_{44} vs. 3 4 f₄₃ space (Morgan et al., 2010; Ng et al., 2010), where f₄₄ and f₄₃ are the ratios of the organic signal at m/z 44 and 43 to the total organic signal in the component mass spectrum, 5 respectively. The degree of oxidation is inferred from the f_{44} value and the range of 6 precursors is suggested by the f_{43} values. Other metrics include the use of O:C and H:C ratios 7 8 in Van Krevelen space which reveals changes in functionality and therefore the likely degree of processing the aerosol has undergone (Heald et al., 2010). Kroll et al. (2011) combined 9 these ratios to derive the oxidation state of carbon and thus describe OA chemistry. 10 Furthermore, the chemical evolution of organic aerosol can be analysed by using the 11 oxidation state along with volatility in the two-dimensional volatility-oxidation space (2D-12 VBS, Donahue et al., 2012). Despite their differences, these metrics can all be used to 13 14 describe the evolution of gas-phase organic compounds through to semi-volatile OA and up 15 to the most oxidised OA with low volatility, concluding that atmospheric processing of fresh OA results in similarly aged and highly oxidised OA. 16

To characterise Northern Hemispheric OA and its evolution in the atmosphere, Ng et al. 17 (2010) compiled ambient AMS data from numerous urban and rural ground-based 18 19 measurement campaigns of varying duration (from a few days up to 5 weeks), occurring during different seasons. Morgan et al. (2010) investigated the organic chemical evolution 20 through various airborne measurement campaigns, tracking individual air masses and 21 crossing a range of European sites during different meteorological conditions. Data from both 22 studies exhibited a range of f₄₄ and f₄₃ values, reflecting the different photochemical ages and 23 sources. However, all data were found to lie within a very well defined triangular region in f_{44} 24 vs. f₄₃ space with OOA1 and OOA2 clustering in discrete regions of the triangular space. 25 26 This indicated that irrespective of source, atmospheric processes result in the convergence of all ambient OA to chemically similar, highly aged SOA. The most processed OA, with high 27 28 f₄₄ values, in both studies were from rural/remote locations and generally occurred during the 29 summer or during periods of elevated temperatures and greater photochemistry. To further investigate OA evolution and the corresponding changes in chemical composition Ng et al. 30 31 (2011b) transformed the f₄₄ vs. f₄₃ triangle into the Van Krevelen diagram (so called VKtriangle). The ambient data from Ng et al. (2010) fall within a narrow range within the VK-32

1 triangle whereas two urban data sets from different seasons fall in a narrow linear area (El

2 Haddad et al., 2013).

3 In contrast to the above studies, the measurements in this study are from a single location over the course of a year. In order to investigate trends within the secondary component of 4 OA, the PMF-derived primary components (SFOA_{mod}, HOA, and COA) are subtracted from 5 the total organic aerosol matrix with the remainder assumed to be secondary (hereafter 6 termed SOA_{calc}). Furthermore, to assess the degree of oxidation of the SOA in London the 7 contribution of the primary components to m/z 44 and 43 are subtracted, so the ratio of m/z 8 44:SOA_{calc} to m/z 43:SOA_{calc} can be determined. When plotted in f_{44} vs. f_{43} space, the 9 majority of the data in this study falls within the triangular space defined by Ng et al. (2010) 10 11 with the average value falling in the OOA1 region (Fig. 7). Along with the range of f_{44} values, correlations with temperature and time can elucidate the extent to which the aerosol is 12 13 oxidised, where in general, temperatures are elevated and photochemistry is greatest in the 14 summer. Figure 7 illustrates the f₄₄ vs. f₄₃ space for SOA_{calc} coloured as a function of time 15 (Fig. 7a) and temperature (Fig. 7b). SOA_{calc} exhibits little seasonality, where spring and summer averages have only a slightly higher ratio than autumn and winter, with this variation 16 17 occurring within a distinct area of the f₄₄ vs. f₄₃ space. There is also little evidence for a temperature trend (Fig. 7b). 18

19 OOA1 and OOA2 are thought to represent end members of OA aging from photochemical processing (Jimenez et al., 2009), where OOA1 is more oxygenated and highly aged 20 compared to the fresher and less-photochemically processed OOA2. The f₄₄ and f₄₃ for the 21 two OOA components identified from PMF analysis in this study, which are fixed factors 22 over the whole one year period, are therefore also plotted within the f44 vs. f43 space to further 23 24 constrain the degree of oxidation of what is hypothesised to be London SOA. The two subtypes are found to fall within their respective range of f₄₄ values expected for each of the 25 two subtypes, with higher f_{44} and f_{43} for OOA1 than OOA2. Using the relationship between 26 f₄₄ and O:C for unit mass resolution data from Aiken et al. (2008), the estimated O:C for the 27 long-term OOA2_{mod} and OOA1 factors are 0.36 and 0.79 respectively. In comparison, for the 28 29 summer IOP there is only a small difference between the O:C ratio of OOA2 and OOA1 (0.44 and 0.52 respectively) obtained from HR-PMF. Here, OOA1 has a much lower ratio 30 31 compared to other urban studies (e.g. Sun et al., 2011; Mohr et al., 2012).

While the concentration of SOA varies through the year, the extent of SOA oxidation shows no variability as a function of time of year or temperature and remains within a very narrow

1 range of values. This could be explained, in part, by the geographical position of London, 2 which leads to influences from both local sources and transported air masses from all directions. SOA in London forms from a variety of precursors across the year (Holmes et al., 3 2014) where the contributions of different precursors will change significantly with season. 4 5 In addition, the greater photochemistry during the summer results in an increase in SOA mass. However, the fraction of oxygen per molecule does not vary as the increase in locally 6 7 produced fresh SOA likely masks any increase in the oxidation of transported material resulting in chemically similar SOA throughout the year. Whether this extends to similar 8 9 urban background sites in other locations remains to be determined but if so, it makes a characterisation of SOA in urban environments more straightforward than may be previously 10 supposed, as the range of precursors and processes appears to lead to consistent average 11 characteristics. 12

13

14 6 Pollution events in London

15 Acute and short-term exposures to particulates have been associated with various adverse health effects including cardiovascular mortality as well as exacerbating existing illnesses 16 17 such as pulmonary disease (Pope and Dockery, 2006 and references therein). It is therefore important to investigate episodic pollution events to better understand their effects on human 18 health. During 2012, the average total NR-PM₁ concentration (\pm one standard deviation) was 19 9.91 (± 10.39) $\mu g m^{-3}$ in London, with slightly higher concentrations in the winter than 20 21 summer (Fig. 8a). Several pollution events occurred throughout the year where the contributions to the high concentrations differed for each of the NR-PM₁ components 22 23 depending on the time of year. To determine whether emissions or atmospheric processes are the controlling factor in driving such high concentration events, the contributions of the 24 different species to the top 10th percentile of the total annual concentration are assessed (Figs. 25 8a and b). Furthermore, the top 10th percentile of the winter and summer periods (Figs. 8c and 26 27 d, respectively) are also analysed to evaluate any seasonal changes in the dominant species 28 and sources.

Secondary aerosols are found to dominate throughout the year (Fig. 8b), irrespective of season, although the individual contributions from SIA and SOA change between winter and summer (Figs. 8c and d). High concentration events are dominated by nitrate in the winter (39%), with a greater contribution from POA than SOA to the organic fraction (79% and 21%, respectively). Furthermore, SFOA_{mod} is the greatest component of POA (43%) and total organic fraction (34%). In contrast, the high concentration events in summer are dominated
by organics (54%), with a significant contribution from SOA (47%), although POA is still the
dominant component of the organic fraction (53%). Nevertheless, the largest contribution to
the organic fraction is from OOA1 (42%).

Pollution events in the winter are therefore driven by particulate emissions, especially nitrate 5 6 and SFOA, whereas in the summer greater photochemistry results in higher concentrations predominantly comprised of SOA. Furthermore, the average mass of the pollution events in 7 8 the winter is greater than that of the summer, suggesting that the limits for daily average concentrations, set to improve air quality and protect human health, are more likely to be 9 10 exceeded in the winter than summer. Therefore, moderating sources of particulates is likely to be the most effective way of reducing particulates in the winter, although this does not 11 12 consider the refractory sources of aerosol, such as black carbon, which contribute to the total 13 PM mass in urban areas (e.g. Liu et al., 2014).

14

15 7 Conclusions

16 A full calendar year of NR-PM₁ chemical composition data were acquired using a cToF-AMS at an urban background site in North Kensington, London, where secondary aerosols 17 comprise approximately 71% of the total non-refractory submicron mass. Nitrate exhibited 18 strong seasonality, peaking in the spring as a result of favourable local meteorological 19 20 conditions and a peak in ammonia emissions. Several high nitrate concentration events occurred throughout the year, which were the result of a combination of ambient conditions, 21 22 availability of precursors, and air mass trajectory. Contrastingly, sulphate concentrations in London are predominantly influenced by regional pollution with few or no local sources and 23 ammonium concentrations are governed by the availability of precursor emissions and 24 25 meteorological conditions. Non-refractory chloride concentrations peak in the winter, governed by the lower temperatures favouring ammonium chloride partitioning to the aerosol 26 27 phase.

The organic fraction was separated into five factors using PMF analysis: HOA, COA, SFOA_{PMF}, OOA1 and OOA2_{PMF}. However, PMF was unable to account for the variance of two factors across the year, resulting in the assignment of some SFOA_{PMF} mass to OOA_{PMF} as indicated by comparison of the factors derived from cToF-PMF and HR-PMF during the winter IOP. Based on the relationship between SFOA_{PMF} and OOA2_{PMF} from the winter at the start and end of 2012, daily concentrations of SFOA_{mod} and OOA2_{mod} were calculated for
the year. OOA1 exhibited characteristics consistent with regional behaviour whereas
OOA2_{mod} exhibited a seasonal trend typical of SOA, peaking in the summer when VOC
emissions and photochemistry are greatest.

5 Although there is a substantial change in the concentration of SOA through the year, the 6 extent of oxidation of the SOA, as defined by the oxygen content of organic aerosol mass, shows no variability as a function of time of year, air mass history, or temperature at the site. 7 8 This suggests that in the urban background of London the range of precursors and chemical processing are insufficiently variable to yield secondary organic aerosol that has been 9 10 exposed to significantly different levels of chemical processing. This is surprising given the 11 variation in precursors throughout the year and the strong annual cycle in photochemical 12 activity. However, this could make characterisation of SOA in urban environments more 13 straightforward than may be previously supposed, as the range of precursors and processes 14 appears to lead to consistent average characteristics.

Several high concentration events occurred in London during 2012, driven by particulate 15 emissions in the winter and formation of SOA in the summer due to the greater 16 photochemistry. The limits for daily average concentrations set to improve air quality and 17 18 protect human health are more likely to be exceeded in the winter as the events had a greater 19 average mass than those in summer. Moderating sources of nitrate and POA is likely to be the most effective way of reducing particulates in the winter, and due to the dominance of this 20 season to the annual mean, for the whole year. SFOA, COA, and HOA all make a substantial 21 contribution to the POA fraction; however SFOA, along with COA, are less well 22 characterised than HOA so their variability requires further investigation. 23

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- 25

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- Table 1. Average annual sulphate concentrations (in μg m⁻³) from two UK locations
 measured between 2001 and 2012 as part of the AURN and Particulates networks. London
- 3 North Kensington is an urban background monitoring site and Harwell is a rural background
- 4 monitoring site.

	Location				
	London	Harwell			
Year	North Kensington				
2012	$1.39^{a}, 1.21^{b,*}$	-			
2011	2.2 ^b	-			
2010	2.3 ^c	1.6 ^c			
2009	1.7 ^c	1.3 ^c			
2008	2.6 ^c	2.4 ^c			
2007	2.8 ^c	2.4 ^c			
2006	3.5 ^c	3 ^c			
2005	3 ^c	2.4 ^c			
2004	3 ^c	2.3 ^c			
2003	2.6°	2.4 ^c			
2002	3.1 ^c	2.3 ^c			
2001	3.1 ^c	2.1 ^c			

 $5 \quad {}^{a}AMS (PM_{1})$, ClearfLo, this study.

⁶ ^bURG 9000B Ambient Ion Monitor (AIM) (PM₁₀), KCL (Courtesy of Dr. D. Green).

7 ^cThermo Scientific Partisol 2025 ion chromatography (PM₁₀), KCL (Courtesy of Dr. D.

8 Green).

9 ^{*}Calculated non-sea salt sulphate.

1	Table 2.	Time	series	comparison	of the	PMF	factors	from	the cT	oF-AMS	and	HR-	ΓoF-	AMS
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2 for the winter IOP.

cToF-PMF factor	HR-PMF factor	Slope	Pearson's r
HOA	HOA	0.95	0.90
COA	COA	0.58	0.89
SFOA _{PMF}	SFOA1	0.80	0.87
SFOA _{PMF}	SFOA2	0.85	0.72
SFOA _{PMF}	Combined SFOA	0.52	0.90
OOA2 _{PMF}	OOA	0.12	0.16
OOA1	OOA	0.90	0.91
$OOA2_{PMF} + OOA1$	OOA	1.02	0.69
$SFOA_{PMF} + OOA2_{PMF}$	Combined SFOA	0.93	0.89



Figure 1. (a) Daily averaged time series of all NR-PM₁ species. (b) Median diurnal profiles of
all NR-PM₁ species. (c) Average monthly fractional contribution of all species to total PM₁.
The months are grouped as seasons: January 2012, February, December, and January 2013
are in winter; March, April, and May are in spring; June, July, and August are in summer;
September, October, and November are in autumn.



Figure 2. Left: average annual fractional contribution of all species to total NR-PM₁. The 2 average annual PM_1 concentrations of SO₄, NH₄, and Chl were 1.39, 1.30, and 0.15 µg m⁻³, 3 respectively. Middle: Expansion of the organic fraction into its primary and secondary 4 components following PMF analysis. Right top: Expansion of the POA fraction into its three 5 components. Right bottom: Magnification of the SOA fraction showing its two subtypes. 6 SFOA and OOA2 refer to SFOA_{mod} and OOA2_{mod}, respectively. See text in Sect. 4.4 for 7 more details. Note that the organic data plotted in the middle and right bars do not include a 8 9 period from the summer as discussed in Sect. 4.1 as well as Sects. 4.1 and 4.2 in the Supplement. 10





2 Figure 3. Median diurnal profiles for each of the five PMF factors. Note that these data do not

3 include a period from the summer (see Sect. 4.1).



2 Figure 4. Time series of OOA1 and $OOA2_{mod}$, where $OOA2_{mod}$ is the daily average.



2 Figure 5. Seasonal fractional contributions of the PMF factors to total OA mass, with revised

- 3 masses (see Section 4.4). As discussed in Sect. 4.1, a period from the summer is not included
- 4 in these data.



Figure 6. (a) OOA1 seasonal median diurnal profiles. (b) Average seasonal concentration of OOA1, with the annual average denoted by the thick horizontal line. (c) Average seasonal concentration of OOA2_{mod}, with the annual average denoted by the thick horizontal line, both estimated in Sect. 4.4. Note that these data do not include the summer period (see Sect. 4.1).



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Figure 7. Daily averaged SOA within f_{44} vs. f_{43} space coloured by time (a) and temperature (b) where f_{44} and f_{43} refer to m/z 44:SOA_{calc} and m/z 43:SOA_{calc}, respectively. See text for more details. Daily averaged temperatures ranged from -0.5 to 26°C although are only coloured up to a maximum of 15°C here for clarity. Average annual and seasonal f_{44}/f_{43} values for SOA are denoted by the text. OOA1, OOA2 PMF factors are also plotted. The outline of the triangle as defined by Ng et al. (2010) is shown by the dashed black lines. Note that the organic data do not include the summer period (see Sect. 4.1)..



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Figure 8. (a) Mean concentrations of the full calendar year (2012-2013), winter, and summer 2 months. The average concentration of the top 10th percentile for the year as well as the top 3 10th percentile in the winter and summer are also shown. (b) Average fractional contributions 4 of all species to the top 10th percentile for the year, with an expansion of the organic fraction 5 into each of its primary and secondary components. (c) Average fractional contributions of all 6 species to the top 10th percentile in the winter, with an expansion of the organic fraction into 7 each of its primary and secondary components. (d) Average fractional contributions of all 8 species to the top 10th percentile in the summer, with an expansion of the organic fraction into 9 each of its primary and secondary components. In all figures, SFOA and OOA2 refer to 10 SFOA_{mod} and OOA2_{mod} respectively. Note that the PMF data (middle and right bars) do not 11 include the summer period as detailed in Sect. 4.1 as well as Sects. 4.1. and 4.2 in the 12 Supplement. 13