Aliphatic Carbonyl Compounds (C$_8$-C$_{26}$) in Wintertime Atmospheric Aerosol in London, UK

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ABSTRACT

Three groups of aliphatic carbonyl compounds, the n-alkanals (C8-C20), n-alkan-2-ones (C8-C26) and n-alkan-3-ones (C8-C19) were measured in air samples collected in London from January-April 2017. Four sites were sampled including two roof-top background sites, one ground-level urban background site and a street canyon location on Marylebone Road in central London. The n-alkanals showed the highest concentrations followed by the n-alkan-2-ones and the n-alkan-3-ones, the latter having appreciably lower concentrations. It seems likely that all compound groups have both primary and secondary sources and these are considered in the light of published laboratory work on the oxidation products of high molecular weight n-alkanes. All compound groups show relatively low correlation with black carbon and NOx in the background air of London, but in street canyon air heavily impacted by vehicle emissions, stronger correlations emerge especially for the n-alkanals. It appears that vehicle exhaust is likely to be a major contributor for concentrations of the n-alkanals whereas it is a much smaller contributor to the n-alkan-2-ones and n-alkan-3-ones. Other primary sources such as cooking may be significant but were not evaluated. It seems likely that there is also a significant contribution from photo-oxidation of n-alkanes and this would be consistent with the much higher abundance of the n-alkan-2-ones relative to the n-alkan-3-ones if the formation mechanism were to be through oxidation of condensed phase alkanes. Vapour-particle partitioning fitted the Pankow model well for the n-alkan-2-ones but less well for the other compound groups, although somewhat stronger relationships were seen at the Marylebone Road site than at the background sites.

Keywords: Carbonyl compounds; n-alkanals; n-alkan-2-ones; n-alkan-3-ones; organic aerosol; partitioning;
1. INTRODUCTION

Carbonyl compounds are classified as polar organic compounds, constituting a portion of the oxygenated organic compounds in atmospheric particulate matter (PM). Aliphatic carbonyl compounds are directly emitted into the atmosphere from primary biogenic and anthropogenic sources (Schauer et al., 2001, 2002a, b), as well as being secondary products of atmospheric oxidation of hydrocarbons (Chacon-Madrid et al., 2010; Zhang et al., 2015; Han et al., 2016).

The most abundant atmospheric carbonyls are methanal (formaldehyde) and ethanal (acetaldehyde), and many studies have described their emission sources and chemical formation in urban and rural samples (Duan et al., 2016). Long-chain aliphatic carbonyl compounds have been identified in PM and reported in few published papers (Gogou et al., 1996; Andreou and Rapsomanikis, 2009), and these compounds are considered to be formed from atmospheric oxidation processes affecting biogenic emissions of alkanes. Anthropogenic activity is also considered to be a significant contributor to the aliphatic carbonyls. Appreciable concentrations of aliphatic carbonyl compounds have been identified in emissions from road vehicles (Schauer et al., 1999), coal combustion (Oros and Simoneit, 2000), wood burning (Rogge et al., 1998) and cooking processes (Zhao et al., 2007b, a), spanning a wide range of molecular weights. Furthermore, chamber studies (Chacon-Madrid and Donahue, 2011; Algrim and Ziemann, 2016) have demonstrated that the aliphatic carbonyl compounds are very important precursors of secondary organic aerosol (SOA) when they react with OH radicals in the presence of NOx.
The oxidation of n-alkanes by hydroxyl radical is considered to be an important source of carbonyl compounds. It was believed that the n-alkanals with carbon atoms numbering less than 20 indicate oxidation of alkanes, whereas the higher compounds were usually considered to be of direct biogenic origin (Rogge et al., 1998). The homologues and isomers of n-alkanals and n-alkanones have been identified as OH oxidation products of n-alkanes in many chamber and flow tube studies (Zhang et al., 2015; Schilling Fahnestock et al., 2015; Ruehl et al., 2013). The commonly accepted oxidation pathways of n-alkanes generally divide into functionalization and fragmentation. Functionalization occurs when an oxygenated functional group (–ONO₂, –OH, –C=O, –C(O)O– and –OOH) is added to a molecule, leaving the carbon skeleton intact. Alternatively, fragmentation involves C–C bond cleavage and produces two oxidation products with smaller carbon numbers than the reactant. The chamber studies of dodecane oxidation have identified 1-undecanal, hexan-3-one, octan-3-one, heptan-2-one, nonan-2-one and decan-2-one as OH oxidation products (Schilling Fahnestock et al., 2015; Yee et al., 2012).

In London, with a high population density and a large number of diesel engine vehicles, the aliphatic hydrocarbons constitute an important fraction of ambient aerosols. Anthropogenic activities and secondary formation favour the emission and production of carbonyl compounds within the city. The objectives of the present study were the identification and quantification of aliphatic carbonyl compounds in particle and vapour samples collected in London from January to April 2017. This work has aided an understanding of the concentrations and secondary formation of carbonyls in the London atmosphere. Spatial and temporal variations of the studied carbonyl compounds were assessed and used to infer sources. One of the main objectives was to provide gas/particle partitioning
coefficients of identified carbonyls under realistic conditions. Diagnostic criteria were used to estimate the sources of identifiable atmospheric carbonyl compounds. Additionally, for the first time, concentrations of particulate and gaseous n-alkan-3-ones are reported.

2. MATERIALS AND METHODS

2.1 Sampling Method and Site Characteristics

Three sampling campaigns were carried out between 23 January and 18 April 2017 at four sampling sites (Figure 1) in London. The first campaign used two sampling sites, one located on the roof of a building (15 m above ground) of the Regent’s University (51°31′N, -0°9′W), hereafter referred to as RU, sampled from 23 January 2017 to 19 February 2017, the other located on the roof (20 m above ground) of a building which belongs to the University of Westminster on the southern side of Marylebone Road (hereafter referred to as WM), sampled from 24 January 2017 to 20 February 2017.

The third sampling site was located at ground level at Eltham (51°27′N, 0°4′E), hereafter referred to as EL, sampled from 23 February 2017 to 21 March 2017, which is located in suburban south London, and the fourth sampling site was located at ground level on the southern side of Marylebone Road (51°31′N, -0°9′W), hereafter referred to as MR, sampled from 22 March 2017 to 18 April 2017.

Marylebone Road is in London’s commercial centre, and is an important thoroughfare carrying 80-90,000 vehicles per day through central London. The Regent’s University site is within Regent’s Park to the north of Marylebone Road. The Eltham site is in a typical residential neighbourhood, 22 km from the MR site. Earlier work at the Marylebone Road and a separate Regent’s Park site is described by Harrison et al. (2012).
The particle samples were collected on polypropylene backed PTFE filters (47 mm, Whatman) which preceded stainless steel sorbent tubes packed with 1 cm quartz wool, 300 mg Carbograph 2TD 40/60 (Markes International, Llantrisant, UK) and sealed with stainless-steel caps before and after sampling. Sampling took place for sequential 24-hour periods at a flow rate of 1.5 L min\(^{-1}\) using an in-house developed automated sampler. Field blank filters and adsorption tubes were prepared for each site, and recovery efficiencies were evaluated. After the sampling, each filter was placed in a clean sealed petri dish, wrapped in aluminium foil and stored in the freezer at -18°C prior to analysis. Black carbon (BC) was simultaneously monitored during the sampling period at RU and WM sites using an aethalometer (Model AE22, Magee Science). Measurements of BC and NO\(_x\) at MR and NO\(_x\) at EL were provided by the national network sites of Marylebone Road, and Eltham (https://uk-air.defra.gov.uk/).

2.2 Analytical Instrumentation

The particle samples were analyzed using a 2D gas chromatograph (GC, 7890A, Agilent Technologies, Wilmington, DE, USA) equipped with a Zoex ZX2 cryogenic modulator (Houston, TX, USA). The first dimension was equipped with a SGE DBX5, non-polar capillary column (30.0 m, 0.25 mm ID, 0.25 mm – 5.00% phenyl polysilphenylene-siloxane), and the second-dimension column equipped with a SGE DBX50 (4.00 m, 0.10 mm ID, 0.10 mm – 50.0% phenyl polysilphenylene-siloxane). The GC × GC was interfaced with a Bench-ToF-Select, time-of-flight mass spectrometer (ToF-MS, Markes International, Llantrisant, UK). The acquisition speed was 50.0 Hz with a mass resolution of > 1200 fwhm at 70.0 eV and the mass range was 35.0 to 600 m/z. All data produced were processed using GC Image v2.5 (Zoex Corporation, Houston, US).
2.3 Analysis of Samples

Standards used in these experiments included 19 alkanes, C$_8$ to C$_{26}$ (Sigma-Aldrich, UK, purity >99.2%); 12 n-aldehydes, C$_8$ to C$_{13}$ (Sigma-Aldrich, UK, purity ≥95.0%), C$_{14}$ to C$_{18}$ (Tokyo Chemical Industry UK Ltd, purity >95.0%); and 10 2-ketones, C$_8$ to C$_{13}$ and C$_{15}$ to C$_{18}$ (Sigma-Aldrich, UK, purity ≥98.0%) and C$_{14}$ (Tokyo Chemical Industry UK Ltd, purity 97.0%).

The filters were spiked with 30.0 μL of 30.0 μg mL$^{-1}$ deuterated internal standards (dodecane-d$_{26}$, pentadecane-d$_{32}$, eicosane-d$_{42}$, pentacosane-d$_{52}$, butylbenzene-d$_{14}$, nonylbenzene-2,3,4,5,6-d$_5$, biphenyl-d$_{10}$, p-terphenyl-d$_{14}$; Sigma-Aldrich, UK) for quantification and then immersed in dichloromethane (DCM), and ultra-sonicated for 20.0 min at 20.0°C. The extract was filtered using a clean glass pipette column packed with glass wool and anhydrous Na$_2$SO$_4$, and concentrated to 50.0 μL under a gentle flow of nitrogen for analysis using GC × GC-ToF-MS. 1 μL of the extracted sample was injected in a split ratio 100:1 at 300°C. The initial temperature of the primary oven (80.0°C) was held for 2.0 min and then increased at 2.0 °C min$^{-1}$ to 210°C, followed by 1.5 °C min$^{-1}$ to 325 °C. The initial temperature of the secondary oven (120°C) was held for 2.0 min and then increased at 3.0°C min$^{-1}$ to 200°C, followed by 2.00°C min$^{-1}$ to 300°C and a final increase of 1.0°C min$^{-1}$ to 330 °C to ensure all species passed through the column. The transfer line temperature was 330 °C and the ion source temperature was 280°C. Helium was used as the carrier gas at a constant flow rate of 1.0 mL min$^{-1}$. Further details of the instrumentation and data processing methods is given by Alam et al. (2016a,b).
The sorbent tubes were analyzed by an injection port thermal desorption unit (Unity 2, Markes International, Llantrisant, UK) and subsequently analyzed using GC × GC-ToF-MS. Briefly, the tubes were spiked with 1 ng of deuterated internal standard for quantification and desorbed onto the cold trap at 350°C for 15.0 min (trap held at 20.0°C). The trap was then purged onto the column in a split ratio of 100:1 at 350°C and held for 4.0 min. The initial temperature of the primary oven (90.0°C) was held for 2.0 min and then increased to 2.0°C min⁻¹ to 240°C, followed by 3.0°C min⁻¹ to 310°C and held for 5.0 min. The initial temperature of the secondary oven (40.0°C) was held for 2.0 min and then increased at 3.0°C min⁻¹ to 250°C, followed by an increase of 1.5°C min⁻¹ to 315°C and held for 5.0 min. Helium was used as carrier gas for the thermally desorbed organic compounds, with a gas flow rate of 1.0 mL min⁻¹.

**Qualitative analysis**

Compound identification was based on the GC×GC-TOFMS spectra library, NIST mass spectral library and in conjunction with authentic standards. Compounds within the homologous series for which standards were not available were identified by comparing their retention time interval between their homologues, and by comparison of mass spectra to the standards for similar compounds within the series, by comparison to the NIST mass spectral library and by the analysis of fragmentation patterns.

**Quantitative analysis**

An internal standard solution (including dodecane-d₂₆, pentadecane-d₃₂, eicosane-d₄₂, pentacosane-d₅₂, triacontane-d₆₂, nonylbenzene-2,3,4,5,6-d₅, butylbenzene-d₁₄, biphenyl-d₁₀, p-terphenyl-d₁₄)
(Sigma-Aldrich, UK) was added to the samples to extract prior to instrumental analysis. Five internal standards (pentadecane-$d_{32}$, eicosane-$d_{42}$, pentacosane-$d_{52}$, triacontane-$d_{62}$, nonylbenzene-2,3,4,5,6-$d_5$) were used in the calculation of carbonyl compound concentrations.

The quantification for alkanes, aldehydes and 2-ketones was performed by the linear regression method using seven-point calibration curves (0.05, 0.10, 0.25, 0.50, 1.00, 2.00, 3.00 ng µL$^{-1}$) established between the authentic standards/internal standard concentration ratios and the corresponding peak area ratios. The calibration curves for all target compounds were highly linear ($r^2$>0.99, from 0.990 to 0.997), demonstrating the consistency and reproducibility of this method.

Limits of detection for individual compounds were typically in the range 0.04–0.12 ng m$^{-3}$. 3-ketones were quantified using the calibration curves for 2-ketones. This applicability of quantification of individual compounds using isomers of the same compound functionality (which have authentic standards) has been discussed elsewhere and has a reported uncertainty of 24% (Alam et al., 2018).

Alkan-2-ones and alkan-3-ones were not well separated by the chromatography. These were separated manually using the peak cutting tool, attributing fragments at m/z 58 and 71 to 2-ketones and m/z 72 and 85 to 3-ketones. The calibration for 2-ketones was applied to quantification of the 3-ketones.

Field and laboratory blanks were routinely analysed to evaluate analytical bias and precision. Blank levels of individual analytes were normally very low and in most cases not detectable. Recovery efficiencies were determined by analyzing the blank samples spiked with standard compounds. Mean
recoveries ranged between 78.0 and 102%. All quantities reported here have been corrected according to their recovery efficiencies.

3. RESULTS AND DISCUSSION

3.1 Mass Concentration of Particle-Bound Carbonyl Compounds

Fig. 2 shows the average total concentrations of particle-bound 1-alkanals, n-alkan-2-ones, and n-alkan-3-ones from January to April at four measurement sites, and the particle and gaseous phase concentrations are detailed in the Table S1 (Supporting Information). Total n-alkanals was defined as the sum of particle-bound n-alkanals ranging from C₈ to C₂₀. The particulate n-alkanals at the MR site accounted for 75.2% of the measured particle carbonyls with the average total concentration of 682 ng m⁻³, and concentrations at the other sites were 167 ng m⁻³ at EL, 117 ng m⁻³ at WM and 82.6 ng m⁻³ at RU, accounting for 57.0%, 57.9% and 56.3% of the measured particulate carbonyls, respectively. The n-alkanals identified in this study differed in some aspects from those previously reported in samples collected from Crete (Gogou et al., 1996) and Athens (Andreou and Rapsomanikis, 2009) in Greece. The n-alkanals from London presented narrower ranges of carbon numbers and a higher concentration than rural and urban samples from Crete. The concentrations of n-alkanal homologues (C₈-C₂₀) ranged from 5.50 to 141 ng m⁻³ (average 52.0 ng m⁻³) at MR which were far higher than 1.48-28.6 ng m⁻³ (average 6.44 ng m⁻³) at RU, 1.42-50.3 ng m⁻³ (average 9.03 ng m⁻³) at WM and 3.29-53.0 ng m⁻³ (average 13.0 ng m⁻³) at EL (Table S1), unlike Crete where the concentrations were 0.9-3.7 ng m⁻³ in rural (C₁₅-C₃₀) and 5.4-6.7 ng m⁻³ in urban (C₉-C₂₂) samples, and the average concentration of all four sites was much higher than the 0.91 ng m⁻³ measured in Athens (Andreou and Rapsomanikis, 2009) (C₁₃-C₂₀).
As part of the CARBOSOL project (Oliveira et al., 2007), air samples were collected in summer and winter at six rural sites across Europe. The particulate n-alkanals ranged from C\textsubscript{11} to C\textsubscript{30} with average total concentrations between 1.0 ng m\textsuperscript{-3} and 19.0 ng m\textsuperscript{-3}, with higher concentrations in summer than winter at all but one site. These concentrations fall well below those measured in the present study, although the range of compounds differed. Maximum concentrations at all sites were in compounds >C\textsubscript{22} indicating a source from leaf surface abrasion products and biomass burning. This far exceeds the C\textsubscript{max} values seen in the particulate fraction at our sites.

The n-alkan-2-one homologues measured in London ranged from C\textsubscript{8} to C\textsubscript{26}, and the average total particulate fraction concentration was 58.5 ng m\textsuperscript{-3} at RU, 75.1 ng m\textsuperscript{-3} at WM, 112 ng m\textsuperscript{-3} at EL and 186 ng m\textsuperscript{-3} at MR, approximately accounting for 39.9% (RU), 37.0% (WM), 38.1% (EL) and 20.5% (MR) of the total particulate carbonyls, respectively (Fig. 2). The published data from Greece indicated that the concentrations of n-alkan-2-ones were independent of the seasons, and an average of 5.40 ng m\textsuperscript{-3} (C\textsubscript{13}-C\textsubscript{29}) was measured in August and 5.44 ng m\textsuperscript{-3} in March at Athinas St, but 12.88 ng m\textsuperscript{-3} was measured in March at the elevated (20 m) AEDA site in Athens (Gogou et al. (1996)).

Concentrations in Crete for alkan-2-ones (C\textsubscript{10}-C\textsubscript{31}) were 0.4-2.1 ng m\textsuperscript{-3} at the rural site and 1.9-2.6 ng m\textsuperscript{-3} at the urban site (Andreou and Rapsomanikis, 2009).

The CARBOSOL project also determined concentrations of n-alkan-2-ones, between C\textsubscript{14} and C\textsubscript{31} with a C\textsubscript{max} at C\textsubscript{29} or C\textsubscript{29} at all but one site. Average concentrations ranged from 0.15 ng m\textsuperscript{-3} (C\textsubscript{17, 29}) to 3.35 (C\textsubscript{14}-C\textsubscript{31}), very much below the concentrations at our London sampling site. Cheng et al. (2006) measured concentrations of n-alkan-2-ones in the Lower Fraser Valley, Canada, in PM\textsubscript{2.5}.
Samples collected in a road tunnel showed the highest concentrations, total 1.8-12.6 ng m$^{-3}$ for C$_{10}$-C$_{31}$, and were higher in daytime than nighttime. Concentrations at a forest site were 1.1-7.2 ng m$^{-3}$ without a diurnal pattern. Values of C$_{\text{max}}$ ranged from C$_{16-17}$ at the road tunnel to C$_{27}$ (secondary maximum) at the forest site. Values of CPI averaged across sites from 1.00 to 1.34, giving little evidence for a substantial biogenic input from higher plant waxes.

Atmospheric concentrations of long-chain n-alkan-3-ones have not previously been reported in the literature. The n-alkan-2-one and n-alkan-3-one homologues with few carbon atoms are believed mainly to originate as the fragmental products of n-alkanes (Yee et al., 2012; Schilling Fahnestock et al., 2015), whereas the higher compounds are mainly generated from functional pathways (Zhang et al., 2015; Ruehl et al., 2013). The n-alkan-3-one homologues identified in the samples ranged from C$_8$ to C$_{19}$, and the average of individual compound concentrations was 0.52 ng m$^{-3}$ at RU, 0.94 ng m$^{-3}$ at WM, 1.37 ng m$^{-3}$ at EL and 3.34 ng m$^{-3}$ at MR. The concentrations of n-alkan-3-ones at the four sites were lower than the n-alkanals and n-alkan-2-ones, and MR had the highest average total mass concentrations 39.4 ng m$^{-3}$, followed by 14.3 ng m$^{-3}$ at EL, 10.4 ng m$^{-3}$ at WM and 5.65 ng m$^{-3}$ at RU, respectively.

Recently published studies have found that the isomeric distribution of first-generation oxidation products of n-alkanes depends strongly upon whether the reaction occurs in the gas phase or at the particle surface (Kwok and Atkinson, 1995; Ruehl et al., 2013). The homogeneous gas-phase oxidation occurs fast, and H-abstraction by OH radicals occurs at all carbon sites. The fractions of the OH radical reaction by H atom abstraction from n-decane at the 1-, 2-, 3-, 4- and 5-positions are
3.10%, 20.7%, 25.4%, 25.4%, and 25.4%, respectively, and the products from homogeneous reaction were generally in accord with structure-reactivity relationship (SRR) predictions (Kwok and Atkinson, 1995; Aschmann et al., 2001). Reaction of particulate n-alkanes is dominated by heterogeneous reactions with OH, and the H-abstraction occurs preferentially at the 2-position of the carbon chain (Zhang et al., 2015; Ruehl et al., 2013). The n-alkanes diffuse from the inner particle to the surface, where the OH will quickly attack the H atom of 1 and 2 position carbons. The intermediate products at the 2-position are relatively more stable than at the 1-position, and the products are dominated by oxidation of the 2-position. The isomeric carbonyls formed via OH-initiated heterogeneous reactions of n-octacosane (C_{28}) exhibit a pronounced preference at the 2-position of the molecule chain\textsuperscript{18}. The n-octacosan-2-ones have the highest relative yield (1.00), followed by n-octacosan-3-ones (0.50), while other isomeric carbonyl yields were lower than 0.20. The same results were found in the subsequent chamber studies of n-alkanes (Zhang et al., 2015) (C_{20}, C_{22}, C_{24}) but not C_{18}. The main reason was that OH oxidation of C_{18} was dominated by the homogeneous reaction as a large fraction of C_{18} evaporated into the gas phase.

During the field experiment, the 1-alkanal homologues were abundant in all samples, and this could be explained by a strong impact of anthropogenic activities. Thus, the n-alkanals are considered to arise mainly from primary emission sources. Furthermore, the particulate form of the n-alkane homologues (C_{14}-C_{36}) identified in the samples ranged from 50-100% in contrast to the low MW n-alkanes (C_{11}-C_{13}). The H-abstraction by OH radicals may therefore have been dominated by heterogeneous reactions generating the higher concentrations of n-alkan-2-ones than n-alkan-3-ones that were found in all samples. The ratio of n-alkan-2-ones/n-alkan-3-ones (C_{11}-C_{18}) with the same
carbon atom number ranged from 2.35-11.3 at four measurement sites. Surprisingly, although the n-alkane (C_{11}-C_{13}) oxidation was expected to be dominated by homogeneous reactions, the n-alkan-2-one/n-alkan-3-one ratios were still greater than 2.00. The probable reason was that the lower molecular weight n-alkan-2-ones were significantly impacted by primary emission sources. Another likely reason is that the n-alkan-2-one and n-alkan-3-one homologues with lower carbon atom numbers originated in part from the fragmental products of higher n-alkanes (Yee et al., 2012; Schilling Fahnestock et al., 2015).

The ratios of n-alkan-2-ones/n-alkanes, n-alkan-3-ones/n-alkanes (with same carbon numbers) were calculated and are reported in Table S2. The n-alkan-3-ones with carbon numbers higher than C_{20} were not identified in the samples, indicating that both the homogeneous and heterogeneous reactions of higher molecular weight n-alkanes were slow, the former probably due to the low vapour phase presence of n-alkanes. The ratios of n-alkan-3-ones/n-alkanes at four measurement sites gradually increased from C_{11}, and then decreased from C_{17}, while higher ratios of n-alkan-2-ones/n-alkanes were observed in the range from C_{17} to C_{22}, probably indicating a shift from homogeneous reactions to heterogeneous reactions with the increase of carbon numbers. The low ratios of n-alkan-2-ones/n-alkanes with carbon numbers from C_{23} to C_{26} were attributed to the low diffusion rate from the inner particle to the surface with the increasing carbon number of n-alkanes, even though heterogeneous reactions were the dominant pathway.
3.2 Temporal and Spatial Variations

The study of temporal and spatial variations of air pollutants can provide valuable information about their sources and atmospheric processing. The time series of particle-bound n-alkanals, n-alkan-2-ones, and n-alkan-3-ones are plotted in Fig. 3. It is clear that the concentrations of n-alkanals varied substantially with date, and were always higher than n-alkanones at four sites. It is also clear from Figure 2 that concentrations were broadly similar at the background sites, RU, WM and EL, but are elevated, especially for the n-alkanals, at MR. This is strongly indicative of a road traffic source.

3.3 Sources of Carbonyl Compounds

3.3.1 Homologue distribution and carbon preference index (CPI)

Fig. 4 shows the average concentrations, and molecular distributions of particle-bound carbonyl compounds at the four sites. The values of carbon preference index (CPI) were calculated to estimate the origin of carbonyl compounds, according to Bray and Evans (1961):

\[
\text{CPI} = \frac{1}{2} \left( \frac{\sum_{i=4}^{m} C_{2i} + 1 + \sum_{i=4}^{m} C_{2i+1}}{\sum_{i=4}^{m} C_{2i}} \right)
\]

For n-alkanals and n-alkan-3-ones (m=9):

\[
\text{CPI} = \frac{1}{2} \left( \frac{\sum \text{odd}(C_9-C_{19}) + \sum \text{even}(C_{10}-C_{20})}{\sum \text{even}(C_8-C_{18}) + \sum \text{even}(C_{10}-C_{20})} \right)
\]

For n-alkan-2-ones (m=12):

\[
\text{CPI} = \frac{1}{2} \left( \frac{\sum \text{odd}(C_9-C_{25}) + \sum \text{even}(C_{10}-C_{26})}{\sum \text{even}(C_8-C_{24}) + \sum \text{even}(C_{10}-C_{26})} \right)
\]

where \(i\) takes values between 4 and \(m\), and 5 and \(m\) as in the equation, and

\(m = 9\) for n-alkanal and n-alkan-3-ones

\(m = 12\) for n-alkan-2-ones
The carbon maximum number ($C_{\text{max}}$) was used to evaluate the relative contribution of the source and exhibit the homologue distribution of highest concentration. Table 1 presents the CPI and $C_{\text{max}}$ of particle-bound carbonyl compounds calculated in the current and other studies.

According to the low CPI (0.41-1.07) at four sites, the n-alkanal homologues with carbon number from $C_8$ to $C_{20}$ mainly originate from anthropogenic emissions or OH oxidation of anthropogenic hydrocarbons. The particle-bound n-alkanals exhibited a similar distribution of carbon number from January to April at four sites, and they had the same $C_{\text{max}}$ at $C_8$ with concentration 28.6 ng m$^{-3}$ at RU, 50.3 ng m$^{-3}$ at WM, 53.0 ng m$^{-3}$ at EL and 141 ng m$^{-3}$ at MR, respectively. This compound may be a fragmentation product, oxidation product or primary emission. In addition, the distribution of n-alkanals had a second concentration peak at $C_{15}$ (MR) and $C_{18}$ (RU, WM, and EL). The $C_{18}$ compound was observed accounting for the highest percentage of the total mass of n-alkanals in some rural aerosol samples (Gogou et al., 1996) in Crete. Andreou and Rapsomanikis reported the $C_{\text{max}}$ as $C_{15}$ or $C_{17}$ in Athens (Andreou and Rapsomanikis, 2009) and attributed this to the oxidation of n-alkanes. However, a $C_{\text{max}}$ at $C_{26}$ or $C_{28}$ in urban Crete (Gogou et al., 1996) was observed, suggestive of biogenic input. The homologue distribution and CPI of n-alkanals in this study differed from those previous reports, and demonstrated weak biogenic input and a strong impact of anthropogenic activities in the London samples.

In this study, n-alkan-2-ones have similar homologue distributions and $C_{\text{max}}$ ($C_{19}$ or $C_{20}$) (Table 2) at RU, WM and EL sites, and the total concentration from $C_{16}$ to $C_{23}$ accounts for 76.0%, 76.1% and 68.0% of $\Sigma n$-alkan-2-ones, respectively. The CPI values for n-alkan-2-ones ranged from 0.57 to 1.23 at the RU, MR and WM sites and were not indicative of biogenic input, and were considered
to mainly originate from anthropogenic activities and OH oxidation of anthropogenic n-alkanes. At EL, the CPI of 1.57 is probably indicative of a biogenic contribution in suburban south London. A difference was observed at the MR site, the n-alkan-2-ones with carbon atoms numbering from C_{12} to C_{18} accounting for 72.0% of $\sum$n-alkan-2-ones, with the C_{\text{max}} being at C_{16}. The C_{\text{max}} of n-alkan-3-ones was at C_{16} at the MR site, at EL, C_{\text{max}} = C_{16}, WM, C_{\text{max}} = C_{17} and at RU, C_{\text{max}} = C_{17}, respectively.

3.3.2 The ratios of n-alkanes/n-alkanals

Diesel engine emission studies have been conducted previously in our group; details of the engine set up and exhaust sampling system are given elsewhere (Alam et al., 2016b). Briefly, the steady-state diesel engine operating conditions were at a load of 5.90 bar mean effective pressure (BMEP) and a speed of 1800 revolutions per minute (RPM), and samples (n=14) were collected both before a diesel oxidation catalyst (DOC) and after a diesel particulate filter (DPF). The n-alkanes (C_{12} - C_{37}) and 1-alkanals (C_{9} - C_{18}) were quantified in the particle samples, while n-alkanones were not identified because their concentrations were lower than the limits of (detection 0.01–0.15 ng m$^{-3}$). The emission concentrations of n-alkanals ranged from 7.10 to 53.2 µg m$^{-3}$ (before DOC) and 1.20 to 11.5 µg m$^{-3}$ (after DPF), respectively, and the ratios of alkanes/alkanals (C_{12}-C_{18}) with the same carbon atom numbers ranged from 0.15 to 0.23 (before DOC) and 0.52 to 7.60 (after DPF). The n-alkane/n-alkanal (C_{12}-C_{18}) ratio at MR ranged from 0.92 to 5.03, while average ratios of 27.6 (RU), 22.1 (WM) and 15.1 (EL) were obtained, respectively. The similarity of the n-alkanes/n-alkanal ratio between MR and the engine studies (after DPF) strongly suggests that diesel vehicle emissions were the main source of 1-alkanals at MR.
The emission factors of total alkanes from diesel engines are reported to be 7 times greater than gasoline engines (Perrone et al., 2014), with n-alkanals with carbon atoms numbering lower than C_{11} being quantified in the exhaust from gasoline engines (Schauer et al., 2002b; Gentner et al., 2013). The n-alkane/n-alkanal (C_8-C_{10}) ratio with the same carbon numbers ranged from 5.60 to 14.3, suggesting that gasoline combustion may be another source of atmospheric n-alkanals.

Studies of n-alkanals showed that aldehydes have high reactivity when the OH radical attacks the aldehyde moiety (Chacon-Madrid and Donahue, 2011; Chacon-Madrid et al., 2010), and the rate constants are more than 3 times those of n-alkanes with the same carbon number. The mechanism and rate constants of H-abstraction by OH detailed in the Master Chemical Mechanism (MCM, v3.3.1), were obtained via [http://mcm.leeds.ac.uk/MCM](http://mcm.leeds.ac.uk/MCM) and used in the evaluation of our data.

### 3.3.3 Correlation analysis

Insights into the sources of carbonyls can be gained from correlation analysis with black carbon (BC) and NO_x. This has the advantage of comparing relative concentrations of pollutants, rather than absolute concentrations. The latter are strongly affected by weather conditions, making inter-site comparisons difficult when sampling did not occur simultaneously. In London, both black carbon and NO_x arise very substantially from diesel vehicle emissions (Liu et al., 2014; Harrison et al., 2012; Harrison and Beddows, 2017), and hence these are good measures of road traffic activity. The concentrations of BC were simultaneously determined by the online instruments during the sampling periods, with the average concentrations of 1.34, 1.94 and 3.58 µg m^{-3} at the RU, WM and MR sites, respectively. The data for NO_x were provided by the national network sites, with the average
concentrations of 23.4 and 202 µg m$^{-3}$ at the EL and MR sites, respectively. At the MR site, the
concentrations of BC and NO$_x$ averaged 5.00 µg m$^{-3}$ and 281 µg m$^{-3}$ when southerly winds were
dominant compared to 2.60 and 128 µg m$^{-3}$ for northerly winds. All correlations were carried out
with the sum of particle and vapour phases for the carbonyl compounds, and strong ($r^2 = 0.87$) and
weak ($r^2 = 0.12$) correlations between BC and NO$_x$ were obtained when the southerly and northerly
winds were prevalent at MR, respectively. Marylebone Road is a street canyon site where a vortex
circulation is established by the wind. The effect is that on northerly wind sectors the sampling site
on the southern side of the road samples near-background air, while on southerly wind sectors, the
traffic pollution is carried to the sampling site, leading to elevated pollution levels affected heavily
by the traffic emissions. The strong correlation between BC and NO$_x$ with southerly wind sectors is
a reflection of their emission from road traffic. In addition, the correlations between n-alkanals (C$_8$-
C$_{20}$) and BC, and between n-alkanals (C$_8$-C$_{20}$) and NO$_x$ were calculated to assess the contribution of
vehicular emission (Table S3). The results showed that the correlations ($r^2$) between n-alkanals and
BC gradually decreased from 0.61 (C$_9$) to 0.34 (C$_{20}$) at MR when the southerly winds were prevalent,
indicating that the distribution of n-alkanals, and especially the lower MW compounds, was
significantly impacted by the vehicular exhaust emissions. The average correlations at MR
(southerly winds) between n-alkanals and BC, and between n-alkanals and NO$_x$ were $r^2 = 0.47$ and
$r^2 = 0.32$, respectively. These moderate correlations demonstrated that the vehicular emissions were
a substantial source of n-alkanals at MR, and result in the high background concentrations of n-
alkanals in London. The other probable sources of n-alkanals include cooking emissions, wood
burning, photooxidation of hydrocarbons and industrial emissions. Poorer correlations between n-
alkanals and BC (average $r^2 = 0.15$), and between n-alkanals and NO$_x$ (average $r^2 = 0.15$) were
observed at MR in the north London background air sampled when northerly winds were prevalent.

There were very weak correlations (average $r^2 < 0.10$) between n-alkanals and BC, and between n-alkanals and NO$_x$ at the RU, WM and EL sites, which may be attributable to the high chemical reactivity of n-alkanals. High concentrations of furanones ($\gamma$-lactones) are generated via the photo-oxidation reaction of n-alkanals (Alves et al., 2001), and the total concentrations (particle and gas) were up to 376, 279, 347 and 318 ng m$^{-3}$ at RU, WM, WL, and MR, respectively for the sum of furanone homologues (from 5-propyldihydro-2(3H)-furanone to 5-tetradecyldihydro-2(3H)-furanone).

The relationships ($r^2$ values) between BC and NO$_x$ and the n-alkan-2-ones were low at all sites, but notably higher with southerly winds at MR (average $r^2 = 0.33$ and 0.35 for BC and NO$_x$ respectively) than for northerly winds ($r^2 = 0.16$ and 0.03 respectively). This is strongly suggestive of a contribution from vehicle exhaust to n-alkan-2-one concentrations, but smaller than that for n-alkanals. In the case of the n-alkan-3-ones, correlations averaged $r^2 = 0.25$ with BC and $r^2 = 0.21$ for NO$_x$ in southerly winds, compared to $r^2 = 0.08$ and $r^2 = 0.05$ respectively for northerly winds. This is also suggestive of a small, but not negligible contribution of vehicle emissions to n-alkan-3-ones.

The very low correlations observed in background air for both n-alkan-2-ones and n-alkan-3-ones with BC and NO$_x$ are suggestive of the importance of non-traffic sources, probably including oxidation of n-alkanes. The considerable predominance for n-alkan-2-one over n-alkan-3-one concentrations may be indicative of a formation pathway from oxidation of condensed phase n-alkanes, but this is speculative as primary emissions may be dominant.
3.4 The Partition Between Particle and Gas Phase

The partitioning coefficient $K_p$ between particles and vapour was calculated in this study according to the following equation defined by Pankow (1994):

$$K_p = \frac{C_p}{C_g TSP}$$

Where, $C_p$ and $C_g$ (µg m$^{-3}$) are the concentration of the compounds in the particulate phase and gaseous phase, respectively. TSP is the concentration of total suspended particulate matter (µg m$^{-3}$), which was estimated from the PM$_{10}$ concentration ($PM_{10}/TSP = 0.80$), and daily average PM$_{10}$ concentrations were taken from the national network sites. The partitioning coefficients $K_p$ calculated from our data and the percentages in the particulate form are presented in Table 2. For the three types of carbonyls, the n-alkanals $>$C$_{16}$, n-alkan-2-ones $>$C$_{19}$, and n-alkan-3-ones $>$ C$_{18}$ were assumed to have negligible vapour concentrations, and the partitioning into the particulate phase gradually increased from C$_8$ to high molecular weight compounds.

Log $K_p$ was regressed against vapour pressure ($VP_T$) for the relevant temperature derived from UManSysProp (http://umansysprop.seaes.manchester.ac.uk/) according to the following equation:

$$\log K_p = m \log (VP_T) + b$$

The calculated log $K_p$ versus log ($VP_T$) for the three types of carbonyls was calculated for each day, and the results appear in the Table S4. Data from four sites were over the temperature range 0.40–15.3 °C. A good fit to the data for n-alkan-2-ones ($r^2 = 0.54–0.94$ at RU, 0.64–0.93 at WM, 0.43–
0.95 EL and 0.45-0.89 at MR) was obtained. It is notable that the fit to the regression equation as indicated by the $r^2$ value is appreciably higher at the MR site than at the other sites, especially in the case of the alkan-3-ones. This is not easily explained, except perhaps by an increased particle surface area at the MR site which may enhance the kinetics of gas-particle exchange, leading to partitioning which is closer to equilibrium.

4. CONCLUSIONS

Three groups of carbonyl compounds were determined in the particle and gaseous phase in London and concentrations are reported for n-alkanals (C$_8$-C$_{20}$), n-alkan-2-ones (C$_8$-C$_{26}$) and n-alkan-3-ones (C$_8$-C$_{19}$). The Marylebone Road site has the highest concentration of particle-bound n-alkanals, and the average total concentration was up to 682 ng m$^{-3}$, followed by 167 ng m$^{-3}$ at EL, 117 ng m$^{-3}$ at WM and 82.6 ng m$^{-3}$ at RU. The particulate n-alkanals were abundant in all samples at all four measurement sites, accounting for more than 56.3% of total particle carbonyls. In addition, the average total particle concentrations of n-alkan-2-ones and n-alkan-3-ones at four measurement sites were in the range of 58.5-186 ng m$^{-3}$ and 5.65-39.4 ng m$^{-3}$, respectively. Diagnostic criteria, including molecular distribution, CPI, $C_{max}$, ratios and correlations, were used to assess the sources and their contributions to carbonyl compounds. The three groups of carbonyls have similar molecular distributions and $C_{max}$ values at the four measurement sites, and their low CPI values (0.41-1.57) at the four sites indicate a weak biogenic input during sampling campaigns. Heavily traffic-influenced air and urban background air were measured at the MR site when southerly and northerly winds were prevalent respectively; correlations of $r^2 = 0.47$ and $r^2=0.32$ were obtained between n-alkanals and BC, and between between n-alkanals and NO$_x$, respectively in southerly
winds. Vehicle emissions appear to be an important source of n-alkanals, which is confirmed by the similar ratios of n-alkanes/n-alkanals measured at MR (0.92-5.03) and in diesel engine exhaust studies (0.52-7.6), resulting in a high background concentration in London. In addition, the OH-initiated heterogeneous reactions of n-alkanes appear to be important sources of n-alkanones, even though weak contributions from vehicular exhaust emissions were suggested by correlation analysis with BC and NOx in southerly winds at MR. Anthropogenic primary sources appear to account for a large proportion of the alkan-2-one and alkan-3-one concentrations measured in London.

In addition, the partitioning coefficients of carbonyls were determined from the relative proportions of the particle and gaseous phases of individual compounds. The results of field measurements of partitioning between particle and vapour phases showed generally a better fit at MR than at the other three sites. The n-alkan-2-ones have a better fit at four sites than the n-alkanals and n-alkan-3-ones, with $r^2 = 0.78$ (0.54–0.94) at RU, 0.85 (0.64–0.93) at WM, 0.74 (0.43–0.95) EL and 0.70 (0.45–0.89) at MR, respectively in a regression of log $K_p$ versus the compound vapour pressure.

ACKNOWLEDGEMENTS

Primary collection of samples took place during the FASTER project which was funded by the European Research Council (ERC-2012-AdG, Proposal No. 320821). The authors would also like to thank the China Scholarship Council (CSC) for support to R.L., and the Natural Environment Research Council for support under the Air Pollution and Human Health (APHH) programme (NE/N007190/1).


TABLE LEGENDS

Table 1. The carbon preference index (CPI) and C_{max} for n-alkanals, n-alkan-2-ones, and n-alkan-3-ones in this study and published data.

Table 2. Percentages of particle phase form and the partitioning coefficient K_{p}.

FIGURE LEGENDS

Figure 1. Map of the sampling sites. RU-Regents University (15 m above ground); WM-University of Westminster (20 m above ground); EL-Eltham; MR-Marylebone Road (south side).

Figure 2. The average total concentration of particle-bound n-alkanals (C_{8}-C_{20}), n-alkan-2-ones (C_{8}-C_{26}), and n-alkan-3-ones (C_{8}-C_{19}), for each sampling period and site. The error bars indicate one standard deviation.

Figure 3. Time series of particle-bound Σ 1-alkanals, Σ n-alkan-2-ones and Σ n-alkan-3-ones at RU, WM, EL, and MR sites.

Figure 4. The molecular distribution of particle-bound carbonyl compounds at four sites (RU, WM, EL, and MR).
Table 1. The carbon preference index (CPI) and C\textsubscript{max} for n-alkanals, n-alkan-2-ones, and n-alkan-3-ones in this study and published data.

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<th>Reference</th>
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<th>C\textsubscript{max}</th>
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<th>Sampling period</th>
<th>Reference</th>
<th>CPI</th>
<th>C\textsubscript{max}</th>
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<td>RU, surrounded by Regent's Park, 15 m above ground</td>
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<td>0.52</td>
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Notes: All values are in ppm.
Table 2. Percentages of particle phase form and the partitioning coefficient Kp.

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Fig. 1. Map of the sampling sites. RU-Regents University (15 m above ground); WM-University of Westminster (20 m above ground); EL-Eltham; MR-Marylebone Road (south side).
Fig. 2. The average total concentration of particle-bound n-alkanals (C8-C20), n-alkan-2-ones (C8-C26), and n-alkan-3-ones (C8-C19), for each sampling period and site. The error bars indicate one standard deviation.
Fig. 3. Time series of particle-bound $\sum_1$-alkanals, $\sum_n$-alkan-2-ones and $\sum_n$-alkan-3-ones at RU, WM, EL, and MR sites.
Fig. 4. The molecular distribution of particle-bound carbonyl compounds at four sites (RU, WM, EL, and MR).