Interpretation of Particle Number Size Distributions Measured across an Urban Area during the FASTER Campaign
Interpretation of Particle Number Size Distributions Measured across an Urban Area during the FASTER Campaign
Distributions Measured across an Urban Area during the EASTER Campaign
Area during the FASTER Campaign
Area uuring the rabitles campaign
Day M. Hannigan ^{1*†} David C.S. Daddawal
Koy M. Harrison ', David C.S. Beddows
Mohammed S. Alam ¹ , Ajit Singh ¹ , James Brean ¹ ,
Ruixin Xu¹, Simone Kotthaus² and Sue Grimmond²
¹ Division of Environmental Health and Risk Management,
School of Geography, Earth and Environmental Sciences
University of Birmingham
Edgbaston, Birmingham B15 2TT
United Kingdom
8
² Department of Meteorology
University of Reading, Reading RG6 6BB
United Kingdom

^{*} To whom correspondence should be addressed.

Tele: +44 121 414 3494; Fax: +44 121 414 3709; Email: r.m.harrison@bham.ac.uk

[†]Also at: Department of Environmental Sciences / Center of Excellence in Environmental Studies, King Abdulaziz University, PO Box 80203, Jeddah, 21589, Saudi Arabia

24 ABSTRACT

Particle number size distributions have been measured simultaneously by Scanning Mobility 25 Particle Sizers (SMPS) at five sites in Central London for a one month campaign in January -26 27 February 2017. These measurements were accompanied by condensation particle counters (CPC) to measure total particle number count at four of the sites and aethalometers measuring Black 28 Carbon (BC) at five sites. The spatial distribution and inter-relationships of the particle size 29 30 distribution and SMPS total number counts with CPC total number counts and Black Carbon measurements have been analysed in detail as well as variations in the size distributions. One site 31 (Marylebone Road) was in a heavily-trafficked street canyon, one site (Westminster University) 32 33 was on a rooftop adjacent to the Marylebone Road sampler, a further sampler was located at Regent's University within a major park to the north of Marylebone Road. A fourth sampler was 34 located nearby at 160 m above ground level on the BT tower and a fifth sampler was located 4 km 35 to the west of the main sampling region at North Kensington. Consistent with earlier studies it was 36 found that the mode in the size distribution had shifted to smaller sizes at the Regent's University 37 (park) site, the mean particle shrinkage rate being 0.04 nm s⁻¹ with slightly lower values at low wind 38 speeds and some larger values at higher wind speeds. There was evidence of complete evaporation 39 of the semi-volatile nucleation mode under certain conditions at the elevated BT Tower site. 40 41 Whereas SMPS total count and Black Carbon showed typical traffic-dominated diurnal profiles, the CPC count data typically peaked during nighttime as did CPC/SMPS and CPC/BC ratios. This is 42 thought to be due to the presence of high concentrations of small particles (2.5 - 15 nm diameter)43 probably arising from condensational growth from traffic emissions during the cooler nighttime 44 conditions. Such behaviour was most marked at the Regent's University and Westminster 45 46 University sites and less so at Marylebone Road, while at the elevated BT Tower site the ratio of particle number (CPC) to Black Carbon peaked during the morning rush hour and not at nighttime, 47 unlike the other sites. An elevation in nucleation mode particles associated with winds from the 48

- 49 West and WSW sector was concluded to result from emissions from London Heathrow Airport,
- 50 despite a distance of 22 km from the Central London sites.

52 1. INTRODUCTION

The adverse health consequences of air polluted by particulate matter are now well recognised 53 (WHO, 2006). While the main focus has been on the public health impact of exposure to fine 54 particulate matter measured by mass (PM_{2.5}), there has also been concern over the possible 55 contribution of ultrafine particles of less than 100 nm diameter to adverse health outcomes. While 56 such particles contribute little to the total mass of particles in the atmosphere, they dominate particle 57 number (Harrison et al., 2000) and authoritative reviews have concluded that although evidence is 58 59 currently highly incomplete, they may contribute to the toxic hazard associated with ambient particulate matter (HEI, 2013; WHO, 2013). There have also been suggestions that particle surface 60 area plays a major role in health impacts and this resides largely in the accumulation mode which is 61 62 typically centred around 100-200 nm diameter (Harrison et al., 2000). Consequently, there is a strong interest from a health perspective in sub-micrometre particles and there are many reports of 63 their concentrations and size distributions within the atmosphere (Asmi et al., 2011; Kumar et al., 64 65 2010; 2014).

66

In addition to concerns over human health, there are other reasons for the study of the size 67 distribution of airborne particles. Not only does this strongly influence their location and efficiency 68 69 of deposition in the human lung, the particle size distribution can also be a strong indicator of particle source, with there being some clear differences between the modal diameter of particles 70 arising from different sources (Vu et al., 2015a). The clearest distinction is between particles 71 arising from combustion and other high temperature sources, which tend to be predominantly very 72 small, and particles generated by attrition processes which are typically far more coarse. However, 73 even within the particles generated from combustion and other high temperature sources, there may 74 well be different modal diameters associated with different sources or even multiple modes 75 associated with an individual source (Vu et al., 2015a). For example, exhaust emissions from diesel 76 engines typically comprise both a nucleation mode and an overlapping Aitken mode, reflecting in 77

78	the former case particles comprised mainly of condensed lubricating oil formed after the
79	combustion process, and in the latter case, solid carbonaceous particles formed within the
80	combustion process (Shi and Harrison, 1999; Alam et al., 2016).
81	

After their emission, particle size distributions are also liable to change through dynamic processes. These include evaporation which causes particles to shrink without changing the overall number, condensation which causes particles to grow without a change in total number, coagulation which also causes growth but reduces the total particle number, and deposition which causes a reduction in number and is a strong function of the particle size.

87

There are detailed assessments of the concentrations and size distributions of nanoparticles in the 88 rural atmosphere (Van Dingenen et al., 2004; Asmi et al., 2011), and of their dynamics during 89 90 atmospheric transport (Beddows et al., 2014), urban studies have been limited. There has been much research on emissions from road transport (Zhu et al., 2002a, b; Kumar et al., 2011), with 91 some attention given to shipping (Gonzalez et al., 2011) and to general modelling of sources 92 93 (Posser and Pandis, 2015). However, most urban measurement studies have been limited to a single 94 site (Morawska et al., 1998; Wang et al., 2011; Brines et al., 2015), although in a few instances more sites have been considered (Karl et al., 2016) but not as part of a concerted campaign. 95

96

Within this study, particle number size distributions were measured simultaneously by electrical
mobility spectrometers at five separate sites across London and the size distributions are compared
with a view to gaining a better understanding of the sources and processes affecting particles in the
urban atmosphere.

101

102

104 2. EXPERIMENTAL

105

Data were collected from 27 January 2017 to 16 February 2017 as part of the second campaign of
the FASTER project. Data recovery was high (100%, or close) at all sites except Westminster
University, where good SMPS data were collected on only three days, January 30 and 31 and
February 1, 2017.

110

111 **2.1** Sampling Sites

Data were collected at five sampling sites in total, three of which were established specifically for the FASTER campaign, Westminster University, Regent's University and BT Tower. The other two sites (London Marylebone Road and London North Kensington) collect data as part of the national Automatic Urban and Rural Network. The site locations (seen in Figure 1) and characteristics are as follows:

Marylebone Road. Air sampling equipment is housed in a large kerbside cabin on the sidewalk
 of a busy central London street canyon with an inlet approximately 2.5 m above ground-level
 (agl). The adjacent six-lane highway carries around 80,000 vehicles per day. The highway is
 relatively straight and runs almost due east-west (angle 80° from north). The buildings on
 either side of the highway are around six storeys in height giving a street canyon aspect ratio of
 approximately 1:1.

Westminster University. Air sampling instruments were located on the roof of the Westminster
 University building, almost directly above the Marylebone Road air sampling site on the
 southern side of the street. The instruments were housed in a temporary enclosure located
 approximately 26 m above street level and 4.5 m from the front edge of the roof where it
 overlooks the road, and with an inlet 1.5 above the roof.

Regent's University. A temporary enclosure for the instruments was located on the roof of
 Regent's University which is an isolated building within Regent's Park due north (i.e. 360°) of
 the Marylebone Road and Westminster University sites. The only highway lying between

Marylebone Road and the Regent's College site is a lightly trafficked road within Regent's 131 Park. The distance between the Westminster University and Regent's University sites is 132 estimated at 380 m. The instruments were located 16 m agl and 1 m from the edge of the roof. 133 London North Kensington. Instruments were sited in a permanent cabin located within the 134 • grounds of a high school in a lightly trafficked suburban area of central London, with an inlet 135 approximately 2.5 m agl. The air pollution climate at this site, often taken as representative of 136 the background air quality within central London, has been characterised in detail by Bigi and 137 Harrison (2010). 138

BT Tower. Instruments were sited on level T35 at approximately 160 m agl on a narrow tower
 which rises well above the surrounding buildings on a quietly trafficked street approximately
 380 m to the south of Marylebone Road. The site was used extensively in the REPARTEE
 experiment (Harrison et al., 2012a).

143

144 2.2 Sampling Instruments

The instruments (Table 1) were operated according to Wiedensohler et al. (2012) guidelines with 145 the omission of a dryer at three sites (discussed later), and calibrated and intercompared both before 146 and after the sampling campaign. Small correction factors (< 5%) were applied to CPC 147 148 (condensation particle counter) data as a result of the intercomparison. SMPS (scanning mobility particle sizers) data were analysed using the AIM9 and AIM10 software provided by TSI as 149 appropriate to the instrument. The national network sites (Marylebone Road and North Kensington) 150 are fitted with diffusion dryers according to EUSAAR/ACTRIS protocols (Wiedensohler et al., 151 2012), but the other sites were not. The particle size ranges measured were 14.9-615.3 nm at 152 Westminster University, Regent's University and BT Tower, 16.55-604.3 nm at Marylebone Road 153 and North Kensington, and a further system with a short DMA (differential mobility analyses) gave 154 4.96-145.9 nm at Regent's University. 155

156

It was not possible to use identical SMPS systems at each site. The variants used are shown in 157 Table 1. We expect little difference between the long column classifiers (TSI 3081) used at all sites 158 but with different platforms (TSI 3080 and TSI 3082) and CPCs (TSI 3775 and 3776). Differences 159 160 are expected to be minimal as platform-specific software was used to invert the data and both the CPC are butanol-based, with only slightly different lower cut-points which were well outside of the 161 range of measured particles. At the Regent's University site, both a long DMA (3081) and short 162 column DMA (3085) were utilised and the data were merged to give a single continuous size 163 distribution from 6 nm to 650 nm. A possible cause of divergence is the fact that two of the sites 164 (Marylebone Road and North Kensington) used diffusion dryers according to the EUSAAR/ 165 166 ACTRIS Protocol. The dryers were tested when installed and showed very low particle losses (less than 5%) and no significant change to particle size distributions (NPL, 2010). The dryer may, 167 however, affect the particle size distribution due to the hygroscopicity of certain kinds of particles. 168 Vu et al. (2015b) reviewed hygroscopic growth factors for submicron aerosols from different 169 sources. Their data are difficult to extrapolate to this study as measurements of hygroscopic growth 170 171 are typically made at very high relative humidities, normally around 90%. Even at 99.5% relative humidity, the growth of particles of less than 100 nm sampled from the atmosphere is relatively low 172 (Vu et al., 2015b). Consequently, a reduction in humidity from 88% typical of the campaign to the 173 values of 30-40% achieved in the dryer would be expected to have only a small effect on particle 174 sizes especially as fresh traffic-generated particles which comprise a large proportion of the sub-175 micrometre particulate matter in the urban atmosphere are hydrophobic and therefore undergo zero 176 or very limited growth in humid atmospheres. 177

178

179 2.

2.3 Weather Conditions During the Campaign

Wind speed and direction data were taken from Heathrow Airport to the west of London to reflect
the synoptic flow minimally affected by local building effects. At the start of the campaign (27
January 2017) the wind direction was easterly and moved to southerly by January 29th, briefly

183	passing through northerly before returning to a southerly circulation between January 31 and
184	February 3rd. During this time, wind speeds were typically around 4 m s ⁻¹ and temperatures mild
185	for the time of the year (mostly 6-10 °C). From February 4th to 8th there was a period of lower
186	wind speeds (1-4 m s ⁻¹) with variable wind directions and low nocturnal minima temperatures
187	(down to 1°C). From Feburary 8 – 12^{th} , a period of northerly winds (speeds of 3-5 m s ⁻¹) and lower
188	temperatures (1-3°C) without appreciable diurnal variation occurred. After February 12th, the
189	winds came from the east moving to south-westerly by February 17 th , with wind speeds variable
190	(between 0 and 6 m s ⁻¹) and temperatures steadily rising to daily maxima of 12°C.
191	

192 The mixed layer heights (MLH) were determined from Vaisala CL31 ceilometer data collected at the Marylebone Road site (Figure 1, Table 1). The observed 15 s (10 m gates) aerosol attenuated 193 backscatter profiles were pre-processed (Kotthaus et al., 2016) prior to using the CABAM 194 algorithm (Kotthaus and Grimmond, 2018) to determine 15 min intervals MLH. The multiple 195 196 aerosol layers (e.g. nocturnal residual layers) in the atmosphere are detected (Kotthaus and 197 Grimmond, 2018; Kotthaus et al., 2018). Here the lowest detected layer is analysed. At times the 198 MLH cannot be detected (e.g. during rain or very weak gradients in attenuated backscatter), but a residual layer might still be indicated. The ceilometer detects periods of precipitation, including 199 200 events that may not be recorded by ground-based stations (e.g. insufficient to trigger a tipping 201 bucket rain-gauge).

202

During the campaign the observed MLH varied from a daily minimum of 45 m agl to a daily maximum of 1312 m agl with an overall 15 min average (median) of 421 (382) m agl. The daily average (median) maximum MLH was 777 (695) and minimum was 194 (197) m agl. The daily range and the amount of data available per day are shown in Figure S1.

207

209 2.4 Modal Analysis of Size Distributions

Modes were fitted to the 15 min data obtained at Marylebone Road, Regent's and Westminster Universities using curve fitting and data analysis software "Fityk (version 1.3.1)" developed by Wojdyr (2010). In the present analysis, a standard peak function (equation 1) was used to disaggregate the size distributions into lognormal modes:

$$P_i = A_i \cdot exp\left[-\left(\frac{\ln(D/c_i)}{W_i}\right)^2\right] \tag{1}$$

By fitting linear a combination of n peaks $(P_1 + P_2 + ... + P_i + ... + P_n)$ to the number size distributions, the following information was calculated: 1) amplitude A_i and location of dN/dlogD at the mode of the distribution c_i , 2) area under the curve (nm cm⁻³), and 3) width of the lognormal curve W_i .

218

- 219 3. **RESULTS AND DISCUSSION**
- 220 **3.1** Particle Size Distributions

A time series of total particle number concentrations from the SMPS instruments appears in Figure
A strong diurnal variation is seen at all sites and is exemplified by the average daily variation
shown in Figure 3.

224

225 The data stratified by the wind direction measured at London Heathrow airport (LHR) (Figure 4) were used to perform the modal analysis. The log normal modes fit to the size distribution were 226 227 used to provide insights into the separate modes contributing to a measured size distribution. Although most measurements could be fit with three separate modes some distributions were best 228 fit with only two modes. An example of a three mode fit of a size distribution from North 229 Kensington appears in the data for the 270° wind sector at this site (Figure 5). It may be seen that 230 using three modes gives a very good overall fit to the data. The details of the modes fitted and their 231 relative magnitude and breadth appear in Table S1. 232

The Marylebone Road sampling site is located in a heavily trafficked (approx. 80,000 vehicles per 234 day) street canyon. The canyon is aligned almost east-west and the sampling site is at kerbside on 235 the southern side of the street. The canyon has a height to width ratio of ~1 consequently we expect 236 237 skimming flow when flow is perpendicular, with one or more vortices established in the canyon (Oke et al. 2017). When there is one vortex, the sampler is exposed to freshly emitted traffic 238 239 contaminants when the wind above the canyon is from the south (Figure 6). Particle number concentration on Marylebone Road is highest for the 225° and 270° wind sectors (Figure 4a) when 240 241 traffic-generated pollutants are carried efficiently to the sampler. When winds have a northerly component such as those for 0° and 45° in Figure 4a, the air reaching the sampler is typical of 242 243 background air from north London and peak concentrations fall by a substantial margin. The particle size data from Marylebone Road (Table S1) show no strong effect of wind direction on the 244 modal diameter for the first fitted mode in the distribution. The average diameter for the 180 and 245 225° wind sectors are 21.4 nm while for the 0 and 45° sectors they are 22.9 nm. The second and 246 third mode in the distribution are far more sensitive to wind direction, with the southerly traffic-247 248 dominated wind directions showing modes at around 32 and 76 nm as opposed to 56 nm and 263 nm for the northerly mode data. The former values compare well with modes in the number 249 distribution of around 20 nm and 50 nm previously attributed to the nucleation mode and Aitken 250 251 mode particles respectively from engine exhaust when sampled at Marylebone Road, with data analysed by Positive Matrix Factorization (Harrison et al., 2011). 252

253

The Westminster University sampling site is 26 m higher and slightly displaced (~8 m) horizontally from the Marylebone Road air sampling station. The observations at roof level are influenced by the flow separation over the roof, if the air is entering or exiting the canyon, and the background concentrations. The particle size data (Table S1) indicate a nucleation mode very similar in size to that observed within the street canyon at the Marylebone Road site. Concentrations are elevated for the 135 and 180° wind bearings suggesting that enhanced concentrations occurring within the

canyon on southerly winds are also elevated at the Westminster University sampler but the dataset
is very small and hence not included in Figure 4. The second mode appears to be broadly similar in
size to that at Marylebone Road and falls within the range of modal diameters measured at
Marylebone Road. Similarly, the third mode falls within the rather variable range also seen at
Marylebone Road.

265

The North Kensington site is widely taken as representative of the background air pollution climate 266 in central London (Bigi and Harrison, 2010; Bohnenstengel et al., 2015). At this site, the size of the 267 first mode in the size distributions is remarkably constant at 22-26 nm which is slightly larger than 268 269 that observed at Marylebone Road. The second mode is also less variable than at most other sites and broadly within the range of the second mode sizes at Marylebone Road (see Table S1). The 270 third mode is highly variable in size with wind direction but again broadly comparable to the data 271 from Marylebone Road. The Beddows et al. (2015) Positive Matrix Factorization of particle 272 number size distributions data from this site identified four factors contributing to the particle 273 274 number size distributions: a secondary component accounting for 4.4% of particle number with a mode at around 250 nm, an urban background factor (43% of particle number) peaking at around 50 275 nm, a traffic component (44.8% of particle number) peaking at around 30 nm and a regional 276 277 nucleation component (7.8% of particle number) peaking at 20 nm. The regional nucleation component showed a strong seasonality with greatest prevalence in the summer months and is 278 thought unlikely to have contributed significantly during the period of this campaign. This was a 279 winter campaign without clear evidence of nucleation leading to new particle formation at any of 280 the sites. A subsequent paper has investigated the factors influencing nucleation at three related 281 282 sites, including North Kensington and Marylebone Road (Bousiotis et al., 2018). Consequently, the first mode observed in our current study is very comparable to the traffic mode observed by 283 Beddows et al. (2015), and the second mode corresponds strongly to the urban background factor 284 285 identified by Beddows et al. (2015) who associated this factor with aged traffic emissions and wood

smoke, the latter of which is unlikely to have influenced the size distribution at Marylebone Roadsignificantly.

288

289 **3.2 Particle Shrinkage**

Previous London work has shown the tendency of nucleation mode traffic-generated particles sampled within Regent's Park to have shrunk by evaporation at rates of on average 0.13 nm s⁻¹ (Harrison et al., 2016) while particles in the regional atmosphere typically undergo condensational growth at a rate of about 0.6-0.9 nm h⁻¹ (Beddows et al., 2014). This reflects an initial local rapid loss of more volatile hydrocarbons, followed by a subsequent slower condensation of low volatility species formed by atmospheric oxidation in the regional atmosphere.

296

Under southerly flows the Regent's University site is downwind of Marylebone Road (Fig. 1). The 297 modal diameters measured at Regent's University in the nucleation mode (Table S1) are clearly 298 indicative of a shrinkage of particle diameter for the wind sectors 180°, 225° and 270°, 299 300 corresponding to air having passed over Marylebone Road. These data show that the nucleation mode is shrinking from a diameter in the range of 21-24 nm at Marylebone Road, and 22-24 nm at 301 Westminster University to a diameter of 14, 9 or 12 nm at the Regent's University site. In this case, 302 303 particle shrinkage seems to be limited to those three wind sectors, with possibly some shrinkage in the 45° wind sector, but particles in other wind sectors retain broadly similar diameters to those 304 measured at Marylebone Road and Westminster University. The second particle mode and third 305 particle mode (where identifiable) at Regent's University are broadly similar and considerably 306 larger than those measured at Marylebone Road or in the limited dataset at Westminster University. 307 308

In our earlier studies of the evolution of particle sizes between Marylebone Road and Regent's Park
(Harrison et al., 2016), the nucleation mode in the Marylebone Road size distributions lay between
20-24 nm (i.e. very similar to this study). In Regent's Park this had reduced to within the range of

6-11 nm with the largest sizes measured in the 0° wind sector and the smallest in the 180° wind 312 sector. The current data show a similar general pattern, although the extent of size reduction is 313 smaller. The travel distance to the Regent's University site is shorter, hence accounting in part for 314 less shrinkage, but the overall shrinkage rate in the current study (0.04 nm s⁻¹) was smaller than 315 previously (0.13 nm s⁻¹) (Harrison et al, 2016). This is probably explained by two factors. Firstly, 316 with warmer mean air temperatures (12-18°C) evaporation would be enhanced, and secondly, as the 317 site used for collection of the data described in the Harrison et al. (2016) study was in the centre of 318 319 the park and further from any major highways than the Regent's University site, it may have experienced lower vapour concentrations. Consequently, the two datasets appear highly consistent 320 321 with one another.

322

Previous BT Tower site observations have reported loss of < 20 nm particles (Dall'Osto et al. 323 2011). This loss was greatest when atmospheric turbulence levels were lowest and hence the time 324 for ground to sampling height (160 m) transport greatest. That analysis is not repeated in this study. 325 326 However, the nucleation mode size (Table S1) has grown slightly from the sizes measured at Marylebone Road for the nucleation mode. It is notable that unlike the earlier results, the amplitude 327 of this mode at the BT Tower was substantial and slightly larger than that observed at the ground-328 level background North Kensington site suggesting that there was generally good coupling between 329 ground-level and the Tower site. It is notable that the first mode diameter with greatest amplitude 330 was for the 270° sector (Figure 4d); this is discussed later. The particle size distribution associated 331 with the 225° wind sector had only one mode at 40 nm suggestive of the second solid particle mode 332 with complete evaporation of the semi-volatile nucleation mode. 333

334

Earlier studies have shown that particle number concentrations (< 100 nm) in a street canyon

(Olivares et al., 2007) and urban air (Hussein et al., 2006) increase with reducing temperature. This

is consistent with the semi-volatility of nucleation mode particles from road traffic (Harrison et al.,

2016), and consequently it would be expected that the particle size distribution as well as the 338 number concentration would be affected by ambient temperature. To investigate this, the size 339 distributions collected in the lowest quartile of air temperatures (1.1 to 3.8°C) were compared with 340 341 those in the highest quartile of temperature (9.1 to 11.8°C). This showed generally higher concentrations associated with the higher temperatures, and a clearer nucleation mode at higher 342 343 temperatures, at all sites, and most notably at Marylebone Road. Such behaviour is contrary to expectations, as greater evaporative losses would be expected at higher temperatures, reducing the 344 345 magnitude of the plot, or shifting the mode to smaller sizes. To understand this effect more clearly, wind directions with the coldest and hottest quartiles of temperature are analysed. The coldest 346 347 periods all occurred during northerly flows (270 to 90°) and >85% of highest quartile of temperatures occur during southerlies (90 to 270°). The behaviour, especially at Marylebone Road 348 and Regent's University therefore appears to be determined predominantly by synoptic wind 349 conditions. For Marylebone Road, the street canyon flow (Figure 6) is the dominant influence and 350 at Regent's University the traffic sources are most proximate with southerly flows. 351

352

353 **3.3** Particle Number Concentration (CPC) Data

Average diurnal variations of total particle number count derived from the Condensation Particle 354 355 Counters produced using the Openair Software Package (Carslaw and Ropkins, 2012) appear in Figure S2. At both Marylebone Road and Westminster University, these show a peak occurs 356 between midnight and 6 am before reducing and then rising to a second peak in the afternoon. CPC 357 concentrations at these sites far exceed those at Regent's University and the BT Tower, whereas 358 integrated counts from the SMPS instruments were considerably smaller and showed a diurnal 359 360 variation broadly similar to that expected for road traffic emissions (Figure 3). While it is quite normal for the CPC to give a higher count than the SMPS since it measures over a wider size range 361 and may have lower internal losses (although the SMPS data analysis software corrects for internal 362 363 losses), the ratio of CPC to SMPS is in our experience (e.g. Shi et al., 2001) typically around two,

but this value was significantly exceeded episodically, especially at Westminster University (Figure 364 S3). The overall pattern of CPC to SMPS ratios (Figure 7) shows that some of the highest ratios 365 were at Regent's University with two individual occasions exceeding 13. Some high peak values 366 367 were observed at Westminster University during the short SMPS time series. Wood burning is recognised as an influential source of particles in London (Harrison et al, 2012b; Crilley et al., 368 2015), and has a diurnal profile with higher concentrations typically at night. During the ClearfLo 369 370 winter campaign the BT Tower was influenced substantially by wood smoke irrespective of 371 boundary layer depth (Crilley et al, 2015). Since the BT Tower site was predominantly within the mixed layer during the 2017 campaign (Figure S1) and the CPC/SMPS average ratios at the Tower 372 373 show little nocturnal elevation, we consider it unlikely that wood smoke explains our observations. Furthermore, particle size distributions associated with biomass burning are typically larger than 374 those from road traffic, and outside of the sub-15 nm size range (Vu et al., 2015a). The occurrence 375 of the maximum in this behaviour at nighttime (3-4am) suggests that other heating-related 376 emissions (e.g. from natural gas combustion) are not the source. 377

378

To evaluate this phenomenon more closely, the Black Carbon data were examined. These are 379 typically taken as a good tracer of diesel exhaust which is expected to be the main source of the 380 381 particle number count. The diurnal variation in Black Carbon (Figure S4) conformed reasonably well to that expected for a traffic-generated pollutant with Marylebone Road concentrations far 382 exceeding those at the other sites and showing a typical traffic-associated pattern. Particle number 383 (derived from the CPC) to Black Carbon ratio (Figure S5) shows huge diurnal variability similar to 384 that seen in the ratio of particle number count from the CPC to that derived from the SMPS. We 385 386 infer from this behaviour that a large number of particles smaller than the lower limit of the SMPS and above the lower limit of the CPC (i.e. 2.5-14.9 nm for the 3776 instrument at Westminster 387 University and Regent's University; 4-14.9 nm for 3775 instrument at BT Tower; and 3-16.55 nm 388 for 3025 instrument at Marylebone Road) were present in the atmosphere. Both the mean ratio of 389

CPC to SMPS (Figure S3) and CPC to Black Carbon (Figure S6) have ratios that are greatest in the 390 early morning (midnight to 6 am). This is unexpected for the CPC/SMPS ratio, as the contribution 391 of traffic relative to regional aerosol is expected to be least and the coarser regional aerosol contains 392 393 few particles in the size range below the lower limit of the SMPS instrument. Similarly, for the Black Carbon data, one would expect that if traffic is the main source of particles measured by the 394 CPC, the latter would show a diurnal fluctuation like that of Black Carbon, which in London arises 395 396 mostly from traffic emissions. Consequently, it seems likely that nucleation processes favoured by 397 the cooler temperatures and lower condensation sink in the early hours of the morning are creating large numbers of particles in the range of 2.5-15 nm mobility diameter. These are forming as air 398 399 moves away from the traffic source and hence are greatest at the rooftop Westminster University site and have diminished to some extent by coagulation or re-evaporation by the time they reach the 400 Regent's University site which still shows a marked elevation in particle number to Black 401 Carbon ratio in the earlier hours of the morning compared to the Marylebone Road site. 402

403

404 Such behaviour is somewhat unexpected and a review of papers in which vertical gradients in particle number count have been measured above roadside sites showed no earlier evidence of such 405 behaviour (Lingard et al., 2006; Agus et al., 2007; Nikolova et al., 2011; Ketzel et al., 2003; 406 Longley et al., 2003; Kumar et al., 2008a, b; Kumar et al., 2009; Li et al., 2007; Vakeva et al., 407 1999; Zhu et al., 2002b; Wehner et al., 2002). However, evidence is seen in some of Villa et al.'s 408 (2017) observations, particle number count increased with height up to around 10 m above a multi-409 lane highway. The authors reported this unexpected pattern for some ascents/descents and 410 attributed it to exhaust tubes of heavy duty trucks tending to project vertically upwards and to be 411 412 located at a height of several metres above ground. They suggest this is not the case in urban canyons. 413

414

Another possibility arises from the report of Rönkkö et al. (2017) that large numbers of sub-4 nm 415 particles are observed in the exhaust of some diesel engines and the observation by Nosko et al. 416 (2017) of substantial numbers of similarly sized particles amongst emissions from brake wear. 417 418 Kontkanan et al. (2017) reported observations of sub-3 nm particles from many sites, the highest concentrations being in urban locations. The diurnal and regional variations did not relate clearly to 419 photochemistry and it was concluded that sub-3 nm particle concentrations are affected by 420 421 anthropogenic sources of precursor vapours. The correlation of sub-3 nm particle concentrations in 422 Helsinki with nitrogen oxides suggested a link with traffic emissions. Shi et al. (2001) measured particles of >9.5 nm by SMPS, >7 nm by CPC and >3 nm by ultrafine CPC, finding large numbers 423 424 of particles in urban air in the ranges 3-7 nm and 3-9.5 nm by differences of counts. Ratios of CPC (>3 nm):SMPS (>9.5 nm) were highly variable, but typically around 4. Clear links to road traffic 425 were seen, with drive-by experiments showing large numbers of particles in the 3-7 nm range in the 426 exhausts of both diesel and gasoline vehicles (Shi et al., 2001). Nanoparticles were also produced 427 in the plume downwind of a stationary combustion source (Shi et al., 2001). Herner et al. (2011) 428 429 measured the size distribution of particles emitted from vehicles equipped with diesel particle filters, and with diesel filters and selective catalytic reduction. The dominant mode in the size 430 distribution was at 10 nm diameter and comprised particles with a high fraction of sulphate. In 431 432 highway and roadside measurements in Helsinki, Enroth et al. (2016) measured particle size distributions with a dominant mode at 10 nm diameter. Such particles would be largely below the 433 lower threshold for counting by the SMPS but not the CPC. It is plausible that during the cooler 434 hours of the night a tail of <2.5 nm particles might be subject to condensational growth if the co-435 emitted vapour were to be supersaturated in the atmosphere within the street canyon. The 436 437 dominance of a 10 nm mode in the size distribution would appear to be the most plausible explanation for the high number concentration of particles observed at the Westminster University 438 rooftop location and the apparent transport of a substantial proportion of such particles to the 439 Regent's University measurement site. While this can explain the typically high CPC/SMPS ratios 440

observed, it does not explain their diurnal variation. This appears to require growth of sub-2.5 nm 441 particles into the range measured by CPC in the cooler, more humid nocturnal conditions. Rönkkö 442 et al. (2006) and Schneider et al. (2005) studied the formation of mechanisms and composition of 443 444 diesel exhaust nucleation particles in the laboratory and during car chasing. They conclude that formation of nucleation mode particles depends upon formation of sulphate nuclei upon which 445 hydrocarbons condense, consistent with earlier studies of Shi and Harrison (1999) and Shi et al. 446 (2000) conducted in our laboratory. Factors favouring nucleation mode particle formation were 447 found to be low temperature and high humidities, consistent with field measurements made on 448 Marylebone Road (Charron and Harrison, 2003). Both factors prevail at nighttime, probably 449 450 contributing to the relative increase in 2.5–15 nm diameter particles seen most notably between midnight and 6am (Figure S3). Salimi et al. (2017) reported nocturnal new particle formation 451 events in Brisbane, Australia, finding that air masses associated with nocturnal events were 452 typically transported over the ocean before reaching their sampling site, but the relevance to our 453 study is unclear, although the maritime air might sometimes be expected to show lower temperature 454 455 and higher humidity than that from the land.

456

Support for our observations also comes from the very detailed measurement and modelling study 457 of Choi and Paulson (2016). Measuring particle number size distribution downwind of a major 458 highway, they found a positive anomaly in particle number within the first 60 m of the plume peak, 459 as the peak for the small particles appeared further downwind than the peak in accumulation mode 460 particles. They attributed this to growth of unmeasured sub-5.6 nm particles into the smallest 461 measurable size range and suggested condensational growth or self-coagulation as the mechanism 462 463 (Choi and Paulson, 2016). Kerminen et al. (2007) measuring near a major road in Helsinki reported particle growth by condensation to be a dominant process during the road-to-ambient evolution 464 stage at nighttime in winter. They inferred that under such conditions (low wind speeds with a 465 temperature inversion), traffic-generated particle numbers were enhanced and could affect 466

submicron particle number concentrations over large areas around major roads. The distance scales
for such processes in both studies (Choi and Paulson, 2016; Kerminen et al., 2007) were within 100
m of source under the conditions of measurement but might conceivably extend over greater
distance scales. Similar processes of particle evolution within an aircraft exhaust plume have been
reported by Timko et al. (2013).

472

Pushpawela et al. (2018) report a phenomenon of hygroscopic particle growth at nighttime, which
can potentially be mistaken for new particle formation. This phenomenon was observed between
0.5-5.0 hours after sunset, peaking at 3.5 hours (Pushpawela et al., 2018). This would not appear to
explain our observations, where the peak in N/SMPS and N/BC plots (Figures S2 and S5) is
greatest at 3-4 am local time, which in London in winter is some 10-11 hours after sunset.
Additionally, such a phenomenon would be expected to be unrelated to local traffic emissions, and
hence more uniform across the various sites.

480

481 **3.5** Spatial Distribution of Particles – Horizontal and Vertical

Figure 2 shows the time series of particle concentrations from the SMPS instruments throughout the 482 campaign. Clearly, as expected, the Marylebone Road site shows the highest concentrations through 483 484 the campaign period due to its proximity to the road traffic source. The other sites tend to track one another quite closely with no consistent ranking of concentrations. There are periods such as 485 February 1st to 3rd when Regent's University well exceeds North Kensington, but at other times, 486 they are very similar (e.g. 10 - 12 February), or periods when North Kensington exceeds Regent's 487 University (e.g. 7 February) but these are few. In the former period (1 - 3 February), winds were 488 489 southerly and concentrations at Regent's University would be enhanced by passage of air across Central London, including Marylebone Road. In the situation where concentrations were similar (10 490 -12 February), winds were in the northerly sector, giving relatively low concentrations at all sites, 491 and rather little spatial variation. The temporal pattern at all sites showed substantial similarity 492

493 overall (Figure 2), including diurnal patterns (Figure 3), although the magnitude of concentrations494 varied.

495

496 A time series of CPC particle number concentrations (Figure 8) showed that under most conditions, the number count was lowest at the BT Tower site, and that the number count at Westminster 497 University frequently exceeded that at Marylebone Road, with Regent's University lower, but 498 499 above the concentration at the BT Tower (Figure 8). During the period of northerly winds (8 - 12)500 February), all sites showed low concentrations with Regent's University and BT Tower similar for much of the time, as for the SMPS data (Figure 2). The highest CPC count concentrations during 501 502 the latter were measured at Westminster University (Figure 9) which was downwind of Marylebone Road at those times. The similarity seen between Westminster University and Marylebone Road 503 for much of the campaign, with concentrations far in excess of those at BT Tower is strongly 504 suggestive of continuing particle growth into the size range 2.5–14.9 nm at Westminster University 505 with re-evaporation occurring before reaching the elevated BT Tower site, as previously observed 506 507 by Dall'Osto et al. (2011). Elevations in N/BC data were seen at the BT Tower site (Figure S4 and S6) but these occurred mainly during the morning rush hour period, presumably due to fresh traffic 508 emissions, rather than overnight as at the other sites (Figure S6). 509

510

Figure 2 suggests that vertical gradients between the proximate Regent's University and BT Tower 511 sites were small in SMPS count (Figure 2), but at certain times were substantial in the CPC count 512 (Figure 9). The particle size distributions measured at the BT Tower (Figure 4d) differ from 513 Marylebone Road and North Kensington (Figure 4a and b) in having no obvious mode in the 514 515 nucleation size range at 20 - 30 nm, a feature shared with Regent's University (Figure 4c). Only during westerly winds (270°) does the BT Tower show such a mode (Figure 4d), while at Regent's 516 University (Figure 5) the 270° wind direction also shows differences from the others with a mode at 517 518 below 20 nm. Anomalous behaviour in this wind sector is also observed at North Kensington

(Figure 4b), and at Marylebone Road. The most pronounced nucleation mode peak is associated 519 with the 270° and 225° wind directions. In the Marylebone Road case, these wind directions are 520 almost parallel to the highway, which might explain the high concentrations and pronounced 521 522 nucleation mode, but this explanation does not work for the other sites. A more likely explanation is that all sites are affected by emissions from Heathrow Airport which is to the west of London and 523 524 has been recognised as a major source of nucleation mode particles associated with aircraft and road traffic emissions (Masiol et al., 2017). At a site 1 km from the northern boundary of Heathrow 525 Airport, PMF factors attributed to aircraft (mode at <20 nm) and fresh road traffic emissions (mode 526 at 18–35 nm) accounted respectively for 31.6% and 27.9% of particle number count in the warm 527 528 season and 33.1% and 35.2% in the cold season (December 2014 – January 2015) data (Masiol et al., 2017). Heathrow Airport is located approximately 22 km from our Central London sites on a 529 bearing of 255°. Keuken et al. (2015) measured a large elevation in concentrations of particles of 530 10-20 nm diameter attributed to aircraft emissions (emission studies are reviewed by Masiol and 531 Harrison, 2014) at a site 7 km east of Schiphol Airport (Netherlands) and have shown by modelling 532 533 and measurement that concentrations are elevated to considerably greater downwind distances. Similarly, Hudda et al. (2014) reported PNC to have increased 4 to 5 fold at 8 – 10 km downwind 534 of Los Angeles International Airport (USA). 535

536

537 The size distributions have also been analysed according with mixed layer height (MLH),

determined by ceilometer (Kotthaus and Grimmond, 2018). Both Marylebone Road (Figure S7) and

539 Regent's University (Figure S8) have the highest concentrations associated with the deepest MLH

540 class (>1000 m). This seems likely to be due to an association with southerly winds and the street

541 canyon circulation. Whereas, North Kensington (Figure S9) has the highest concentrations during

shallow MLH (< 100 m and 100 - 200 m) when dispersion is limited for the low altitude emissions.

- 543 The most interesting behaviour is seen at the elevated (160 m) BT Tower site, which is consistent
- with Harrison et al. (2012a) and Dall'Osto et al. (2011). During the shallowest MLH (< 100 m) the

measurement site is above the inversion and the size distribution lacks an obvious nucleation mode 545 (Figure S10). As the MLH deepens, a nucleation mode appears which dominates the size 546 distribution for the deepest MLH categories (900 - 1000 m and > 1000 m) with a mode at 20 - 30547 548 nm, similar to that seen at Marylebone Road for the same MLH depths (Figure S7). The gradual transitioning of size distribution as the MLH deepens is consistent with the surface source (mainly 549 road traffic) of nucleation mode particles, and their evaporative loss which increases with the 550 551 timescale of vertical mixing to the height of the sampler, as reported by Dall'Osto et al. (2011), and the ultimate isolation of the sampler from ground-level emissions at the shallowest boundary layer 552 heights, as observed by Harrison et al. (2012a). 553

554

555 3.6 Detailed Comparison of Marylebone Road, Westminster University and Regent's 556 University

Unfortunately, a full dataset for the Westminster University site was only collected over the period 557 January 30th to February 1st due to a late set-up of the instrument and a malfunction after February 558 1st. This period however merits closer examination as it is the only period where SMPS data were 559 available for all three sites. For much of the time the SMPS data for the Westminster University 560 site looks surprisingly similar to that of the Marylebone Road site despite the former being on the 561 rooftop and the latter being within the street canyon. A detailed analysis hour by hour showed that 562 out of 51 hourly observations, in 23 the amplitude of the mode (dN/dlogD) at Westminster 563 University was within \pm 20% of that at Marylebone Road while in 25 cases the amplitude was 564 greater at Westminster University than at Marylebone Road, and in just two cases the amplitude 565 was smaller at Westminster University. In an attempt to explain this observation, the 566 567 meteorological data for the periods of similar magnitude and of different magnitudes were compared but no systematic difference was seen in wind direction, air temperature or relative 568 humidity between any of the periods. Wind directions were generally in a south-easterly to easterly 569

570 sector, mean temperatures around 8°C and relative humidity high (85 and 99%). The maximum

571 MLH were low and there was a lot of rain (Figure S1).

572

573 In order to gain further insight, the time series of observations were plotted for this period and appear in Figure 9. The SMPS integrated number counts shown in Figure 9(a) show a remarkable 574 similarity between Marylebone Road, Westminster University and Regent's University. For the 575 576 first two days, Regent's University concentrations are lower than those from the other two sites, 577 although on the third day they are very similar to those at Westminster University. On the first and last days, the peak concentrations at Marylebone Road exceed those at Westminster University but 578 579 on the middle day (January 31st) the differences between these two sites are very small. The CPC particle number counts shown in Figure 9(b) are very similar to those at Marylebone Road on the 580 first and last day but exceed those at Marylebone Road on January 31st. Concentrations at Regent's 581 University are typically only around half or less of those measured at Westminster University. The 582 magnitude of the CPC concentrations peaking at over 40,000 cm⁻³ is close to double the integrated 583 SMPS counts which peak at a little over 20,000 cm⁻³ indicating a large number of particles in the 584 size range below 14.9 nm. 585

586

However, the Black Carbon data (Figure 9c) have daytime concentrations at Marylebone Road that 587 far exceed those at Westminster University and Regent's University, the latter sites tracking each 588 other and having very similar concentrations. Since Black Carbon can be viewed as a conserved 589 tracer of vehicle emissions over these small time and distance scales, the inference is that particle 590 production must be continuing as the vehicle exhaust mixes upwards from the street canyon 591 592 Marylebone Road site to the Westminster University rooftop site. The southerly wind directions likely associated with upward flow on the Westminster University canyon wall (Fig. 6) would carry 593 vehicle exhaust past the Marylebone Road measurement station (south side of the road). 594

Air leaving the canyon and being entrained by the complex building roof flows could expose the 596 Westminster University sampler to air exiting the street canyon and to the general flow towards 597 Regent's University site (Fig. 6 and 1). Such behaviour is consistent with the observations of 598 599 particle growth in the sub-SMPS size ranges reported in the previous section extending into the SMPS size range. This is similar to behaviour observed by Kerminen et al. (2007) in Helsinki who 600 observed not only possibly evaporation of some particles in the 7-30 nm range, but also on apparent 601 602 growth of nucleation mode particles into the 30–63 nm size range between sampling points at 9 m 603 and 65 m downwind of a highway. The results in Figure 9 are suggestive of a substantial growth of nuclei into the range of the CPC at Westminster University. 604

605

606 4. CONCLUSIONS

The measurement of particle number size distributions in the atmosphere is resource intensive and 607 there have been rather few studies in which more than two samplers have been operated within a 608 city. Typically if there are two sites, one is a traffic-influenced site and the other urban background. 609 610 In this study, data have been collected at a total of five sites, although unfortunately the dataset from the Westminster University site is limited to only a few days. Nonetheless, the dataset allows 611 some deep insights into the spatial distribution of particle sizes and number counts not only 612 613 horizontally but in the vertical dimension. Not unexpectedly, concentrations of particles at the street canyon Marylebone Road site considerably exceed concentrations at other sites, but there are 614 nonetheless considerable similarities in diurnal profiles and the magnitude of concentrations at the 615 other, background sites. 616

617

One of the main motivating factors for this study was to confirm earlier observations of shrinkage of the nucleation mode particles between traffic emissions on Marylebone Road and the downwind site at Regent's University within Regent's Park. Particle shrinkage was observed within the current study although at a slower mean rate (0.04nm s⁻¹) than in the earlier study (Harrison et al.,

2016) in which the mean shrinkage rate was 0.13nm s⁻¹. However, temperatures in the current 622 study all fell below those in the earlier work of Harrison et al. (2016). Other factors may also have 623 been influential. There have been marked changes in the road vehicle fleet in London between the 624 625 two measurement campaigns. The earlier dataset as reported by Dall'Osto et al. (2011) and Harrison et al. (2016) was collected in 2006 at which time the sulphur content of diesel fuel was 626 regulated at below 50 ppm. Between the two campaigns, the sulphur content of both gasoline and 627 diesel motor fuels was reduced to below 10 ppm sulphur in order to facilitate the introduction of 628 629 diesel particle filters from 2011 onwards. The incorporation of a diesel particle filter on EURO 5 and EURO 6 vehicles leads to a substantial overall reduction in particulate matter emissions but 630 also a change in the hydrocarbon content of the particles. Secondly, the Regent's Park sampling 631 site used for the 2006 measurements was at about double the distance from Marylebone Road 632 compared to the Regent's University used in the latest study. This would allow for greater dilution 633 of the traffic plume from Marylebone Road and other adjacent highways, leading to a greater 634 reduction in vapour phase hydrocarbons at the more distant site causing an accelerated evaporation 635 636 process. The reduction in fuel sulphur content in 2007 was accompanied by a marked change in the size distribution of particles emitted from road traffic, including a reduction in the nucleation mode 637 particles (Jones et al., 2012). The work of Dall'Osto et al. (2011) also analysed data from the BT 638 Tower, showing increasing evaporative loss of nucleation mode particles as the travel time from 639 ground level to the sampling site on the Tower became longer with reduced atmospheric turbulence 640 levels. Although that phenomenon has not been studied in detail in the latest dataset, the results are 641 clearly consistent with such a process, and with an apparent total loss of the nucleation mode in 642 particles associated with regional pollution sampled when the boundary layer top was below the 643 644 sampling height on the tower.

645

Although the phenomenon of particle shrinkage had been seen in earlier work, there were twofurther major observations made in the current study which were not anticipated. The first, was the

clear influence of a major source to the west of London, almost certainly Heathrow Airport, upon 648 concentrations of nucleation mode particles. The association of an enhanced nucleation mode in the 649 270° or 225° sector is indicative of a major source of very fine particles, and the work of Masiol et 650 651 al. (2017) at a sampling site close to Heathrow Airport provides strong evidence for major emissions both from aircraft engines and the large volumes of road traffic attracted by the airport. 652 Earlier research by Keuken et al. (2015) and Hudda et al. (2014) gives a clear precedent for 653 measurement of strongly elevated concentrations of very fine particles several kilometres 654 downwind of a major airport, but to our knowledge this is the first observations of concentrations 655 above urban background at a distance of 22 km from the centre of the airport. 656

657

The other observation which was wholly unexpected was of the very poor relationship between total 658 particle numbers measured by the Scanning Mobility Particle Sizers and the total particle numbers 659 measured by co-located condensation particle counters. While both the SMPS counts and co-660 located Black Carbon measurements show a typical road traffic diurnal profile, the CPC data show 661 a quite different diurnal profile peaking at night. This is most evident in the ratios of CPC/SMPS 662 and CPC/BC seen at all sampling sites, with the exception of CPC/BC at the elevated BT Tower 663 site which does not show a nocturnal maximum, but peaks during the morning rush hour period. 664 Earlier studies such as that of Choi and Paulson (2016) and Kerminen et al. (2007) have reported 665 data consistent with such a phenomenon, but with very modest elevations in particle count 666 compared to those in the current data. The implication is of the presence of large numbers of 667 particles within the range of 2.5 - 15nm and hence observable with the CPC but below the lower 668 669 cut of the SMPS. It seems likely that such particles grow at night from vary small nuclei and it seems possible that the exceptional magnitude of this process within London results from the high 670 density of diesel traffic leading to substantial nocturnal concentrations of condensable vapours close 671 672 to the traffic source. A common feature to such observations appears to be its association with still conditions on winter nights which lead to poor dispersion of vehicle emissions and a pool of vapour 673

674 co-emitted with traffic particles which becomes supersaturated as it cools in the ambient

atmosphere, leading to condensation on small nuclei when the general particle concentrations andhence the condensation sink are relatively low in magnitude.

677

These very abundant particles within the 2.5 - 15 nm range are likely to prove ephemeral as they would be expected to re-evaporate as the air mass dilutes away from source. However, the health effects of exposure to particles within this range are poorly known and no recommendation can be given as to whether health-related studies would be best to measure the particle size range covered by the SMPS as is most typically performed at present, or whether CPC data going down to smaller particles sizes would be more appropriate.

684

There are some additional general conclusions from the work. Firstly the results demonstrate the dynamic behaviour of traffic-generated (and other) particles within the urban atmosphere. Our earlier paper (Dall'Osto et al., 2011) referred to "remarkable dynamics", and further remarkable dynamic processes have been observed in the current study. Secondly, as this work has revealed sources and processes that were not originally anticipated, although with the benefit of hindsight it might have been possible to predict them, there is clearly a need for further detailed observational studies of the behaviour of sub-100 nm particles within the urban atmosphere.

692

693 ACKNOWLEDGEMENTS

The authors are grateful to the management and staff of Westminster University, Regent's University and British Telecom for access to their buildings for air sampling. They also express gratitude to the National Centre for Atmospheric Science (NCAS) for the loan of sampling instruments, and to Dr Paul Williams (NCAS) for facilitating the instrument intercomparison. The operation of the ceilometers were supported by NERC ClearfLo, NERC AirPro, Newton Fund/Met Office CSSP (SG, SK) and University of Reading. We acknowledge the support of KCL LAQN for

700	the instrument sites and support and the Reading Urban Micromet group for maintaining the
701	instruments, notably in this period Elliott Warren and Kjell zum Berge. The work was funded by the
702	European Research Council (ERC-2012-AdG, Proposal No. 320821) and the UK Natural
703	Environment Research Council (R8/H12/83/011) and a NCAS studentship (to JB).
704	
705	AUTHOR CONTRIBUTIONS
706	DB, MA, JB and RX carried out the field measurements of particle size distributions, SK and SG
707	collected and interpreted the ceilometer data, and DB and AS carried out data analyses. RH led the
708	project and drafted the paper, with all co-authors contributing to subsequent enhancements.
709	

710 **REFERENCES**

- 711
- Agus, E. L., Young, D. T., Lingard, J. J. N., Smalley, R. J., Tate, J. E., Goodman, P. S., and Tomlin,
 A. S.: Factors influencing particle number concentrations, size distributions and modal parameters
 at a roof-level and roadside site in Leicester, UK, Sci. Tot. Environ., 386, 65-82, 2007.
- 715
- Alam, M. S., Rezaei, S. Z., Stark, C. P., Liang, Z., Xu, H. M., and Harrison R. M.: The
- characterisation of diesel exhaust particles composition, size distribution and partitioning, Faraday
 Discuss., 189, 69-84, 2016.
- 719

- Asmi, A., Wiedensohler, A., Laj, P., Fjaeraa, A.-M., Sellegri, K., Birmili, W., Weingartner, E.,
- Baltensperger, U., Zdimal, V., Zikova, N., Putaud, J.-P., Marinoni, A., Tunved, P., Hansson, H.-C.,
- Fiebig, M., Kivekas, N., Lihavainen, H., Asmi, E., Ulevicius, V., Aalto, P.P., Swietlicki, E.,
- 723 Kristensson, A., Mihalopoulos, N., Kalivitis, N., Kalapov, I., Kiss, G., Deleeuw, G., Henzing, B.,
- Harrison, R. M., Beddows, D., O'Dowd, C., Flentje, H., Weinhold, K., Meinhardt, F., Ries, L., and
- 725 Kulmala M.: Number size distributions and seasonality of submicron particles in Europe 2008-
- 726 2009, Atmos. Chem. Phys., 11, 5505-5538, 2011.
- Beddows, D. C. S., Dall'Osto, M., Harrison, R. M., Kulmala, M., Asmi, A., Wiedensohler, A., Laj,
- P., Fjaeraa, A.M., Sellegri, K., Birmili, W., Bukowiecki, N., Weingartner, E., Baltensperger, U.,
- Zdimal, V., Zikova, N., Putaud, J.-P., Marinoni, A., Tunved, P., Hansson, H.-C., Fiebig, M.,
- 731 Kivekäs, N., Swietlicki, E., Lihavainen, H., Asmi, E., Ulevicius, V., Aalto, P. P., Mihalopoulos, N.,
- 732 Kalivitis, N., Kalapov, I., Kiss, G., De Leeuw, G., Henzing, B., O'Dowd, C., Jennings, S. G.,
- Flentje, H., Meinhardt, F., Ries, L., Denier Van Der Gon, H. A. C., and Visschedijk, A.J.H.:
- Variations in tropospheric submicron particle size distributions across the European Continent
 2008-2009, Atmos. Chem. Phys., 14, 4327-4348, 2014.
- 736
- Beddows, D. C. S., and Harrison, R. M., Green, D., Fuller, G.: Receptor modelling of both particle
 composition and size distribution data from a background site in London UK, Atmos. Chem. Phys.,
 15, 10107-10125, 2015.
- 740
- Bigi, A., and Harrison R. M.: Analysis of the air pollution climate at a central urban background
 site, Atmos. Environ., 44, 2004-2012, 2010.
- 743
- Bohnenstengel, S. I., Belcher, S. E., Aiken, A., Allan, J. D., Allen, G., Bacak, A., Bannan, T. J.,
- Barlow, J. F., Beddows, D. C. S., Bloss, W. J., Booth, A. M., Chemel, C., Coceal, O., Di Marco, C.
- F., Dubey, M. K., Faloon, K. H., Fleming, Z. L., Furger, M., Geitl, J. K., Graves, R. R., Green, D.
- 747 C., Grimmond, C. S. B., Halios, C. H., Hamilton, J. F., Harrison, R. M., Heal, M. R., Heard, D. E.,
- Helfter, C., Herndon, S. C., Holmes, R. E., Hopkins, J. R., Jones, A. M., Kelly, F. J., Kotthaus, S.,
- Langford, B., Lee, J. D., Leigh, R. J., Lewis, A. C., Lidster, R. T., Lopez-Hilfiker, F. D., McQuaid,
- J. B., Mohr, C., Monks, P. S., Nemitz, E., Ng, N. L., Percival, C. J., Prévôt, A. S. H., Ricketts, H.
 M. A., Sokhi, R., Stone, D., Thornton, J. A., Tremper, A. H., Valach, A. C., Visser, S., Whalley, L.
- M. A., Sokhi, R., Stone, D., Thornton, J. A., Tremper, A. H., Valach, A. C., Visser, S., Whalley, L.
 K., Williams, L. R., Xu, L., Young, D. E., and Zotter, P.: Meteorology, air quality, and health in
- 752 K., Williams, E. K., Xu, E., Toung, D. E., and Zotter, T.: Meteorology, an 753 London: The ClearfLo project. Amer. Meteor. Soc., 779-804, 2015.
- 754
- Bousiotis, D., Dall'Osto, M., Beddows, D. C. S., Pope, F. D. and Harrison, R. M.: Analysis of new
 particle formation (NPF) events at nearby rural, urban background and urban roadside sites, Atmos.
 Chem. Phys. Discuss., <u>https://doi.org/10.5194/acp-2018-1057</u>, 2018.
- 758
- 759 Brines, M., Dall'Osto, M., Beddows, D. C. S., Harrison, R. M., Gómez-Moreno, F., Núñez, L.,
- 760 Artíñano, B., Costabile, F., Gobbi, G. P., Salimi, F., Morawska, L., Sioutas, C., and Querol, X.:

Traffic and nucleation events as main sources of ultrafine particles in high insolation developed 761 world cities, Atmos. Chem. Phys., 15, 5929-5945, 2015. 762 763 Carslaw, D. C., and Ropkins, K.: openair – An R package for air quality data analysis, Environ. 764 765 Model. Softw. 27-28, 52-61, doi:https://doi.org/10.1016/j.envsoft.2011.09.008, 2012. 766 767 Charron, A., and Harrison, R. M.: Primary particle formation from vehicle emissions during exhaust dilution in the roadside atmosphere, Atmos. Environ., 37, 4109-4119, 2003. 768 769 770 Choi W., and Paulson, S. E.: Closing ultrafine particle number concentration budget at road-toambient scale: Implications for particle dynamics, Aerosol Sci. Technol., 50, 5, 448-461, 2016. 771 772 773 Crilley, L. R., Bloss, W. J., Yin, J., Beddows, D. C. S., Harrison, R. M., Allan, J. D., Young, D. E., Flynn, M., Williams, P., Zotter, P., Prevot, A. S. H., Heal, M. R., Barlow, J. F., Hallios, C. H., Lee, 774 775 J. D., Szidat, S., and Mohr, C.: Sources and contributions of wood smoke during winter in London: 776 assessing local and regional influences, Atmos. Chem. Phys., 15, 3149-3171, 2015. 777 Dall'Osto, M., Thorpe, A., Beddows, D.C.S., Harrison, R.M., Barlow, J.F., Dunbar, T., Williams, 778 779 P.I., and Coe, H.: Remarkable dynamics of nanoparticles in the urban atmosphere, Atmos. Chem. Phys., 11, 6623-6637, 2011. 780 781 Enroth, J., Saarikoski, S., Niemi, J., Kouse, A., Jezek, I., Mocnik, G., Carbone, S., Kuulivainen, H., 782 Rönkkö, T., Hillamo, R., and Pirjola, L.: Chemical and physical characterization of traffic particles 783 in four different highway environments in the Helsinki metropolitan area, Atmos. Chem. Phys., 16, 784 5497-5512, 2016. 785 786 Gonzalez, Y., Rodriguez, S., Guerra Garcia, J. C., Trujillo, J. L., and Garcia, R.: Ultrafine particles 787 pollution in urban coastal air due to ship emissions, Atmos. Environ., 45, 4907-4914, 2011. 788 789 790 Harrison, R.M., Shi, J.P., Xi, S., Khan, A., Mark, D., Kinnersley, R., and Yin, J.: Measurement of 791 number, mass and size distribution of particles in the atmosphere, Phil. Trans. R. Soc. Lond., A, 792 358, 2567-2580, 2000. 793 794 Harrison, R.M., Beddows, D.C., and Dall'Osto, M.: PMF analysis of wide-range particle size 795 spectra collected on a major highway, Environ. Sci. Technol., 45, 5522-5528, 2011. 796 797 Harrison, R.M., Dall'Osto, M., Beddows, D.C.S., Thorpe, A.J., Bloss, W.J., Allan, J.D., Coe, H., 798 Dorsey, J.R., Gallagher, M., Martin, C., Whitehead, J., Williams, P.I., Jones, R.L., Langridge, J.M., Benton, A.K., Ball, S.M., Langford, B., Hewitt, C.N., Davison, B., Martin, D., Petersson, K., 799 Henshaw, S.J., White, I.R., Shallcross, D.E., Barlow, J.F., Dunbar, T., Davies, F., Nemitz, E., 800 Phillips, G.J., Helfter, C., Di Marco, C.F., and Smith, S.: Atmospheric chemistry and physics in the 801 802 atmosphere of a developed megacity (London): An overview of the REPARTEE experiment and its 803 conclusions, Atmos. Phys. Chem., 12, 3065-3114, 2012a. 804 805 Harrison, R.M., Beddows, D.C.S., Hu, L., and Yin, J.: Comparison of methods for evaluation of wood smoke and estimation of UK ambient concentrations, Atmos, Chem. Phys., 12, 8271-8283, 806 2012b. 807 808 Harrison, R.M., Jones, A.M., Beddows, D.C., and Dall'Osto, M.: Evaporation of traffic-generated 809 nanoparticles during advection from source, Atmos. Environ., 125, 1-7, 2016. 810 811

- HEI: Understanding the Health Effects of Ambient Ultrafine Particles, Health Effects Institute, HEI
 Perspectives 3, 2013.
- 814

830

834

839

- Herner, D.H., Hu, S., Robertson, W.H., Huai, T., Chang, M.C.O., Riger, P., and Ayala, A.: Effect
 of Advanced Aftertreatment for PM and NOx Reduction on Heavy-Duty Diesel Engine Ultrafine
- 817 Particle Emissions, Environ. Sci. Technol., 45, 2413-2419, 2011.818
- Hudda, N., Gould, T., Hartin, K., Larson, T.V., and Fruin, S.A.: Emissions from an international
 airport increase particle number concentrations 4-fold at 10km downwind, Environ. Sci. Technol.,
 48, 6628-6635, 2014.
- Hussein, T., Karppinen, A., Kukkonen, J., Harkonen, J., Aalto, P.P., Hameri, K., Kerminen, V.-M.,
 and Kulmala, M.: Meteorological dependence of size-fractionated number concentrations of urban
 aerosol particles, Atmos. Environ., 40, 1427-1440, 2006.
- Jones, A.M., Harrison, R.M., Barratt, B., and Fuller, G.: A large reduction in airborne particle
 number concentrations at the time of the introduction of "suphur free" diesel and the London Low
 Emission Zone, Atmos. Environ., 50, 129-138, 2012.
- Karl, M., Kukkonen, J., Keuken, M. P., Lutzenkirchen, S., Pirjola, L., and Hussein, T.: Modeling
 and measurements of urban aerosol processes on the neighborhood scale in Rotterdam, Oslo and
 Helsinki, Atmos. Chem. Phys., 16, 4817-4835, 2016.
- Kerminen, V.M., Pakkanen, T.A., Makela, T., Hillamo, R.E., Sillanpaa, M., Rönkkö, T., Virtanen,
 A., Keskinen, J., Pirjola, L., Hussein, T., and Hameri, K.: Development of particle number size
 distribution near a major road in Helsinki during an episodic inversion situation, Atmos. Environ.,
 41, 1759-1767, 2007.
- Ketzel, M., Wahlin, P., Berkowicz, R., and Palmgren, F.: Particle and trace gas emission factors
 under urban driving conditions in Copenhagen based on street and roof-level observations, Atmos.
 Environ., 37, 2735-2749, 2003.
- Keuken, M.P., Moerman, M., Zandveld, P., Henzing, J.S., and Hoek, G.: Total and size-resolved
 particle number and Black Carbon concentrations in urban areas near Schiphol airport (the
 Netherlands), Atmos. Environ., 104, 132-142, 2015.
- 847 848 Kontkanan, J., Lehtipalo, K., Ahonen, L., Kangasluma, J., Manninen, H.E., Hakala, J., Rose, C.,
- Sellegri, K., Xiao, S., Wang, L., Qi, X., Nie, W., Ding, A., Yu, H., Lee, S., Kerminen, V.-M.,
 Petaja, T., and Kulmal, M.: Measurements of sub-3 nm particles using a particle size magnifier in
 different environments: from clean mountian top to polluted megacities, Atmos. Chem. Phys., 17,
 2163-2187, 2017.
- Kotthaus S, and Grimmond, C.S.B.: Atmospheric boundary layer characteristics from ceilometer
 measurements Part 1: A new method to track mixed layer height and classify clouds, Q. J. R.
 Meteorol. Soc., https://doi.org/10.1002/qj.3299, 2018.
- 857
- Kotthaus, S., and Grimmond, C. S. B.: Atmospheric boundary layer characteristics from ceilometer
 measurements, Part 2: Application to London's urban boundary layer, Q. J. R. Meteorol. Soc.,
- 860 <u>https://doi.org/10.1002/qj.3298,</u> 2018.
- 861

Kotthaus, S., Halios, C. H., Barlow, J. F., and Grimmond, C.S.B.: Volume for pollution dispersion: 862 London's atmospheric boundary layer during ClearfLo observed with two ground-based lidar types 863 Atmos. Environ., 190, 401-414, 2018. 864 865 Kumar, P., Fennell, P., and Britter, R.: Measurements of particles in the 5-1000 nm range close to 866 road level in an urban street canyon, Sci. Tot. Environ., 390, 437-447, 2008a. 867 868 Kumar, P., Fennell, P., Langley, D., and Britter, R.: Pseudo-simultaneous measurements for the 869 vertical variation of coarse, fine and ultrafine particles in an urban street canyon, Atmos. Environ., 870 871 42, 4304-4319, 2008b. 872 873 Kumar, P., Garmory, A., Ketzel, M., Berkowicz, R., and Britter, R.: Comparative study of 874 measured and modelled number concentrations of nanoparticles in an urban street canyon, Atmos. Environ., 43, 949-958, 2009. 875 876 877 Kumar, P., Robins, A., Vardoulakis, S., and Britter, R.: A review of the characteristics of nanoparticles in the urban atmosphere and the prospects for developing regulatory controls, Atmos. 878 879 Environ., 44, 5035-5052, 2010. 880 Kumar, P., Ketzel, M., Vardoulakis, S., Pirjola, L., and Britter, R.: Dynamics and dispersion 881 modelling of nanoparticles from road traffic in the urban atmospheric environment - a review, J. 882 Aerosol Sci., 42, 580-602, 2011. 883 884 885 Kumar, P., Morawska, L., Birmili, W., Paasonen, P. H, M., Kulmala, M., Harriosn, R.M., Norford, L., and Britter, R.: Ultrafine particles in cites, Environ. Intl., 66, 1-10, 2014. 886 887 Li, X. L., Wang, J. S., Tu, X. D., Liu, W., and Huang, Z.: Vertical variations of particle number 888 concentration and size distribution in a street canyon in Shanghai, China, Sci. Tot. Environ., 378, 889 890 306-316, 2007. 891 Lingard, J. J. N., Agus, E.L., Young, D. T., Andrew, G. E., and Tomlin, A. S.: Observations of 892 893 urban airborne particle number concentrations during rush-hour conditions: analysis of the number based size distributions and modal parameters, J. Environ., Monitor., 8, 1203-1218, 2006. 894 895 896 Longley, I. D., Gallagher, M. W., Dorsey, J. R., Flynn, M., Allan, J. D., Alfarra, M. R., and Inglis, D.: A case study of aerosol (4.6 nm< $D_p < 10 \mu$ m) number and mass size distribution measurements 897 in a busy street canyon in Manchester, UK, Atmos. Environ., 37, 1563-1571, 2003. 898 899 900 Masiol, M., and Harrison, R. M.: Aircraft engine exhaust emissions and other airport-related contributions to ambient air pollution: A review, Atmos. Environ., 95, 409-455, 2014. 901 902 Masiol, M., Harrison, R. M., Tuan, V. V., and Beddows, D. C. S.: Sources of sub-micrometre 903 904 particles near a major international airport, Atmos. Chem. Phys., 17, 12379-12403, 2017. 905 906 Morawska, L., Thomas, S., Bofinger, N., Wainwright, D., and Neale, D.: Comprehensive characterization of aerosols in a subtropical urban atmosphere: Particle size distribution and 907 correlation with gaseous pollutants, Atmos. Environ., 32, 2467-2478, 1998. 908 909 Nikolova, I., Janssen, S., Vos, P., Vrancken, K., Mishra, V., and Berghmans, P.: Dispersion 910 911 modelling of traffic induced ultrafine particles in a street canyon in Antwerp, Belgium and 912 comparison with observations, Sci. Tot. Environ., 412, 336-343, 2011. 913

- Nosko, O., Vanhanen, J., and Olofsson, U.: Emission of 1.3-10 nm airborne particles from brake
 materials, Aerosol Sci. Technol., 51, 91-96, 2017.
- 916
 917 NPL: Design, construction and testing of a humidity management system for ultrafine particle field
 918 measurements, National Physical Laboratory, NPL Report AS 48, 2010.
- Oke, T., Mills, G., Christen, A., and Voogt, J.: Urban Climates, Cambridge University Press,
 doi:10.1017/9781139016476, 2017.
- Olivares, G., Johansson, C., Strom, J., and Hansson, H.-C.: The role of ambient temperature for
 particle number concentrations in a street canyon, Atmos. Environ., 41, 2145-2155, 2007.
- 925

947

950

956

960

919

- Posser, L. N., and Pandis, S. N.: Sources of ultrafine particles in the Eastern United States, Atmos.
 Environ., 111, 103-112, 2015.
- Pushpawela, B., Jayaratne, R., and Morawska, L.: Differentiating between particle formation and
 growth events in an urban environment, Atmos. Chem. Phys., 18, 11171-11183, 2018.
- 931
 932 Rönkkö, T., Virtanen, A., Vaaraslahti, K., Keskinen, J., Pirjola, L., and Lappi, M.: Effect of
 933 dilution conditions and driving parameters on nucleation mode particles in diesel exhaust:
 934 Laboratory and on-road study, Atmos. Environ., 40, 2893-2901, 2006.
- Rönkkö, T., Kuuluvainen, H., Karjalainen, P., Keskinen, J., Hillamo, R., Niemi, J. V., Pirjola,
 L., Timonen, H. J., Saarikoski, S., Saukko, E., Järvinen, A., Silvennoinen, H., Rostedt, A., Olin,
 M., Yli-Ojanperä, J., Nousiainen, P., Kousa, A., and Dal Maso, M.: Traffic is a major source of
 atmospheric nanocluster aerosol, PNAS, 114, 7549-7554, 2017.
- Salimi, F., Rahman, Md. M., Clifford, S., Ristovski, Z., and Morawska, L.: Nocturnal new particle
 formation events in urban environments, Atmos. Chem. Phys., 17, 521-530, 2017.
- 943
 944 Schneider, J., Hock, N., Weimer, S., Borrmann, S., Kirchner, U., Vogt, R., and Scheer, V.:
 945 Nucleation Particles in Diesel Exhaust: Composition Inferred from In Situ Mass Spectrometric
 946 Analysis, Environ. Sci. Technol., 39, 6153-6161, 2005.
- Shi, J. P., and Harrison, R. M.: Investigation of ultrafine particle formation during diesel exhaust
 dilution, Environ. Sci. Technol., 33, 3730-3736, 1999.
- Shi, J. P., Mark, D., and Harrison, R. M.: Characterization of Particles from a Current Technology
 Heavy-Duty Diesel Engine, Environ. Sci. Technol., 34, 748-755, 2000.
- Shi, J. P., Evans, D. E., Khan, A. A., and Harrison, R. M.: Sources and concentration of
 nanoparticles (< 10 nm diameter) in the urban atmosphere, Atmos. Environ., 35, 1193-1202, 2001.
- Timko, M. T., Fortner, E., Franklin J., Yu, Z., Wong, W., Onasch, T. B., Miake-Lye, R. C., and
 Herndon, S. C.: Atmospheric Measurements of the Physical Evolution of Aircraft Exhaust Plumes,
 Environ. Sci. Technol, 2013, 47, 3513-3520, 2013.
- Vakeva, M., Hameri, K., Kulmala, M., Lahdes, R., Ruuskanen, J., Laitinen, T.: Street level versus
 rooftop concentrations of submicron aerosol particles and gaseous pollutants in an urban street
 canyon, Atmos. Environ., 33, 1385-1397, 1999.
- 964

Van Dingenen, R., Raes, F., Putaud, J.-P., Baltensperger, U., Charron, A., Facchini, M.-C., 965 Decesari, S., Fuzzi, S., Gehrig, R., Hansson, H.-C., Harrison, R. M., Hüglin, C., Jones, A. M., Laj, 966 P., Lorbeer, G., Maenhaut, W., Palmgren, F., Querol, X., Rodriguez, S., Schneider, J., ten Brink, H., 967 Tunved, P., Tørseth, K., Wehner, B., Weingartner, E., Wiedensohler, A., and Wåhlin, P.: A 968 European aerosol phenomenology -1: Physical characteristics of particulate matter at kerbside, 969 970 urban, rural and background sites in Europe, Atmos. Environ., 38, 2561-2577, 2004. 971 Villa, T. F., Javaratne, E. R., Gonzalez, L. F., and Morawska, L.: Determination of the vertical 972 profile of particle number concentration adjacent to a motorway using an unmanned aerial vehicle, 973 974 Environ. Pollut., 230, 143-142, 2017. 975 976 Vu, T. V., Delgado-Saborit, J. M., and Harrison, R. M.: Review: Particle number size distributions 977 from seven major sources and implications for source apportionment studies, Atmos. Environ., 122, 114-132, 2015a. 978 979 980 Vu, T. V., Delgado-Saborit, J. M., Harrison, R. M.: A review of hygroscopic growth factors of submicron aerosols from different sources and its implication for calculation of lung deposition 981 efficiency of ambient aerosol, Air Qual. Atmos. Health, doi 10.1007/s11869-015-0365-0, 2015b. 982 983 Wang, Y., Hopke, P. K., Chalupa, D. C., and Utell, M. J.: Long-term study of urban ultrafine 984 particles and other pollutants, Atmos. Environ., 45, 7672-7680, 2011. 985 986 987 Wehner, B., Birmili, W., Gnauk, T., and Wiedensohler, A.: Particle number size distributions in a street canyon and their transformation into the urban-air background: measurements and a simple 988 model study, Atmos. Environ., 36, 2215-2223, 2002. 989 990 WHO: Air Quality Guidelines - Global Update 2005, World Health Organization, Copenhagen, 991 992 2006. 993 994 WHO: Review of Evidence on Health Aspects of Air Pollution - REVIHAAP Project, World 995 Health Organizatrion, Copenhagen, 2013. 996 Wiedensohler, A., Birmili, W., Nowak, A., Sonntag, A., Weinhold, K., Merkel, M., Wehner, B., 997 998 Tuch, T., Pfeifer, S., Fiebig, M., Fjaraa, A.M., Asmi, E., Sellegri, K., Depuy, R., Venzac, H., Villani, P., Laj, P., Aalto, P., Ogren, J.A., Swietlicki, E., Williams, P., Roldin, P., Quincey, P., 999 Huglin, C., Fierz-Schmidhauser, R., Gysel, M., Weingartner, E., Riccobono, F., Santos, S., 1000 Gruning, C., Faloon, K., Beddows, D., Harrison, R.M., Monahan, C., Jennings, S.G., O'Dowd, C. 1001 1002 D., Marinoni, A., Horn, H.-G., Keck, L., Jiang, J., Scheckman, J., McMurry, P. H., Deng, Z., Zhao, C. S., Moerman, M., Henzing, B., de Leeuw, G., Loschau, G., and Bastian, S.: Mobility particle 1003 size spectrometers: harmonization of technical standards and data structure to facilitate high quality 1004 long-term observations of atmospheric particle number size distributions, Atmos. Meas. Tech., 5, 1005 1006 657-685, 2012. 1007 Wojdyr M.: Fityk: A General purpose peak fitting program, J. Appl. Cryst., 43, 1126-1128, 2010. 1008 1009 Zhu, Y., Hinds, W. C., Kim, S., and Sioutas, C.: Concentration and size distribution of ultrafine 1010 particles near a major highway, JAWMA, 52, 1032-1042, 2002a. 1011 Zhu, Y., Hinds, W. C., Kim, S., Shen, S., and Sioutas, C.: Study of ultrafine particles near a major 1012 highway with heavy-duty diesel traffic, Atmos. Environ., 36, 4323-4335, 2002b. 1013 1014 1015 1016

1017 1018	TABLE LEGENDS				
1019 1020 1021 1022 1023	Table 1:	Location sites of instruments during the campaign. Mean sea level (msl), Above ground level (agl), Condensation particle counter (CPC), Scanning Mobility Particle Sizers (SMPS).			
1024	FIGURE LEGENDS				
1025 1026 1027 1028	Figure 1:	Study area locations (a) in central London (UK) and (b) more detail of the Marylebone Road (MR), Westminster University (WU) and Regent's University (RU) sites.			
1029 1030 1031	Figure 2:	Time series of total particle number count from the SMPS instruments at the five sites (Fig. 1, Table 1) over the campaign period.			
1032 1033 1034	Figure 3:	Campaign-average diurnal variation of particle number counts derived from the SMPS instruments with median (line) and inter-quartile range (shading) shown.			
1035 1036 1037 1038	Figure 4:	Average particle number size distributions stratified by 45° wind directions sectors (°, measured at LHR, value indicates mid-point of sector ers) for (a) Marylebone Road, (b) North Kensington (c) Regent's University, (d) BT Tower.			
1039 1040	Figure 5:	Lognormal modes fitted to the average particle size spectrum at North Kensington for wind direction sector 270°.			
1041 1042 1043 1044	Figure 6:	A schematic diagram of the wind flows in the street canyon of Marylebone Road (6 traffic lanes) during southerly and northerly winds. The orange marker represents the MR sampling site and red marker represents the WM sampling site.			
1045 1046 1047	Figure 7:	Time series (15 min) of ratio of total particle number counts, CPC/SMPS, for four sites over the campaign period.			
1048 1049 1050	Figure 8:	Time series (15 min) of total particle number count from the CPC instruments located at four sites over the campaign period.			
1051 1052 1053 1054 1055 1056	Figure 9:	Time series (15 min) of (a) SMPS integrated counts, (b) particle number counts (CPC) and (c) Black Carbon from Marylebone Road, Westminster University and Regent's University for 30 January to 1 February 2017.			

- **Table 1:** Location sites of instruments during the campaign. Mean sea level (msl), Above ground level (agl), Condensation particle counter (CPC),
 Scanning Mobility Particle Sizers (SMPS).
- 1059

Site Name	Marylebone Road	Westminster University	Regent's University	BT Tower	North Kensington
Lat (° N), Long	51.522530, 0.154611	51.522322.0.15515	51.525542, 0154570	51.521426, 0.138924	51.521082,
(°W)					0.213403
Height of ground	26	26	30	25	23
msl (m)					
Height of inlets agl	4	26	17	160	3
(m)					
Instruments	Long_DMA_SMPS/	Long_DMA_SMPS/CPC/	Long_DMA_SMPS/	Long_DMA_SMPS/CPC/	Long_DMA_SMPS
installed	CPC	(Micro)Aethalometer/Anemometer	Short_DMA_SMPS/	(Micro)	Vaisala CL31
	Vaisala CL31		CPC/Aethalometer/Anemometer	Aethalometer/Anemometer	
Particle	3080+3081+3775	3080+3081+3776	(3082+3081+3775)/(3082+3085+3776)	(3080+3081+3775)	(3080+3081+3775)
spectrometer type					
Aerosol dryer	Yes	No	No	No	Yes
CPC type	TSI 3025	TSI 3776	TSI 3776	TSI 3775	None

1061 Note: The SMPS size ranges are given in Section 2.2. The lower size cuts (D₅₀) of the CPCs are 3 nm (3025), 2.5 nm (3776) and 4 nm (3775).



Figure 1: Study area locations (a) in central London (UK) and (b) more detail of the Marylebone
 Road (MR), Westminster University (WU) and Regent's University (RU) sites.



Figure 2: Time series of total particle number count from the SMPS instruments at the five sites (Fig. 1, Table 1) over the campaign period.







Figure 3: Campaign-average diurnal variation of particle number counts derived from the SMPS
 instruments with median (line) and inter-quartile range (shading) shown.





(a)

Regent's University (Long+short column)





Figure 4: Average particle number size distributions stratified by 45° wind directions sectors (°,
measured at LHR, value indicates mid-point of sector ers) for (a) Marylebone Road, (b) North
Kensington (c) Regent's University, (d) BT Tower.





- 1102 for wind direction sector 270° .





Figure 6: A schematic diagram of the wind flows in the street canyon of Marylebone Road (6
traffic lanes) during southerly and northerly winds. The orange marker represents the MR sampling
site and red marker represents the WM sampling site.





Figure 8: Time series (15 min) of total particle number count from the CPC instruments located atfour sites over the campaign period.





Figure 9: Time series (15 min) of (a) SMPS integrated counts, (b) particle number counts (CPC) and (c) Black Carbon from Marylebone Road, Westminster University and Regent's University for

- 30 January to 1 February 2017.