

1 **Seasonal and diurnal trends in concentrations and fluxes**  
2 **of volatile organic compounds in central London**

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## 14 **Abstract**

15 Concentrations and fluxes of seven volatile organic compounds (VOCs) were measured  
16 between August and December 2012 at a roof-top site in central London as part of the ClearLo  
17 project (Clean Air for London). VOC concentrations were quantified using a proton transfer  
18 reaction-mass spectrometer (PTR-MS) and fluxes were calculated using a virtual disjunct  
19 eddy covariance technique (vDEC). The median VOC fluxes, including aromatics, oxygenated  
20 compounds and isoprene, ranged from 0.07 to 0.33 mg m<sup>-2</sup> h<sup>-1</sup>. Median mixing ratios were 7.3  
21 ppb for methanol and < 1 ppb for the other compounds. Strong relationships were observed  
22 between the fluxes and concentrations of some VOCs with traffic density, and between the  
23 fluxes and concentrations of isoprene and oxygenated compounds with photosynthetically  
24 active radiation (PAR) and temperature. An estimated 50-90 % of the fluxes of aromatic VOCs  
25 were attributable to traffic activity, which showed little seasonal variation, suggesting that  
26 boundary layer effects or possibly advected pollution may be the primary causes of increased  
27 concentrations of aromatics in winter. Isoprene, methanol and acetaldehyde fluxes and  
28 concentrations in August and September showed high correlations with PAR and temperature,  
29 when fluxes and concentrations were largest suggesting that biogenic sources contributed to  
30 their fluxes. Modelled biogenic isoprene fluxes from urban vegetation using the Guenther et  
31 al., (1995) algorithm agreed well with measured fluxes in August and September.  
32 Comparisons of estimated annual benzene emissions from the London and National  
33 Atmospheric Emissions Inventories agreed well with measured benzene fluxes. Flux footprint  
34 analysis indicated emission sources were localized and that boundary layer dynamics and  
35 source strengths were responsible for temporal and spatial VOC flux and concentration  
36 variability during the measurement period.

## 37 **1 Introduction**

38 Currently over 50 % of the global population lives in urban areas and with increasing migration  
39 to urban centres, air quality remains a high public health priority. In the European Union,  
40 including in the UK, volatile organic compound (VOC) emissions are subject to control under  
41 the European Commission Directive 2008/50/EC and emission reducing technologies have  
42 been implemented, yet urban air pollution continues to be a concern. VOCs from both  
43 anthropogenic and biogenic sources impact urban air quality and climate through their  
44 contribution to tropospheric ozone and aerosol particle formation. Some VOCs including  
45 benzene and 1,3-butadiene are also carcinogens, which can directly affect human health (Kim  
46 et al., 2001). Most VOCs in urban areas are assumed to come from fuel combustion or  
47 evaporative emissions (Kansal, 2009; Srivastava et al., 2005). However, in summer, urban  
48 vegetation may act as an additional source of VOCs such as methanol, isoprene and  
49 monoterpenes even in cities with a temperate climate and little green space, such as London  
50 or Manchester (Langford et al., 2009; 2010b).

51 Emission inventories such as the London and National Atmospheric Emissions Inventories  
52 (LAEI and NAEI) use a “bottom-up” approach based on activity data and emission factors to  
53 estimate emission rates from pollutant sources. Micrometeorologically-based eddy covariance  
54 techniques allow a “top-down” approach to quantify fluxes and these measurements can be  
55 compared with modelled emission inventory estimates. Such comparisons are essential as  
56 “bottom-up” emission inventories may inadvertently not include specific pollutant sources, or  
57 may use unrepresentative emission factors or activity profiles. “Top-down” approaches using  
58 Earth observation data from satellites are also available for a few chemicals (Lamsal et al.,  
59 2011), but not for primary VOCs. There have been few studies on VOC fluxes in urban areas,  
60 and these have been limited in spatial and temporal extent (Langford et al., 2009; 2010b; Park  
61 et al., 2010; 2011; Velasco et al., 2005; 2009). Due to the high technical demands of VOC flux  
62 measurements, it is difficult to increase spatial coverage or to make measurements for long

63 periods of time. Making further measurements of this kind is therefore a high priority in studies  
64 of air quality.

65 In this study we present flux and concentration measurements of seven selected volatile  
66 organic compounds made over five months in central London using the virtual disjunct eddy  
67 covariance method. The aims of this study were to i) quantify VOC fluxes above an urban  
68 canopy using proton transfer reaction-mass spectrometry and disjunct eddy covariance; ii)  
69 investigate seasonal, diurnal and spatial differences in VOC fluxes and concentrations; iii)  
70 examine possible major source contributions of speciated VOCs in central London; and iv)  
71 compare measured fluxes with those estimated by the London and National Atmospheric  
72 Emissions Inventories.

73 These observations were made as part of the ClearfLo (Clean air for London) project, which  
74 provided integrated short-term and long-term measurements of meteorology, gas phase and  
75 particulate pollutants over London and surrounding areas during 2011 and 2012  
76 (Bohnenstengel et al., 2015).

## 77 **2 Methods**

### 78 **2.1 Measurement site**

79 Micrometeorological flux measurements were made during the period 7<sup>th</sup> August - 19<sup>th</sup>  
80 December 2012 from a flux tower located on the roof of a building belonging to King's College,  
81 University of London (51.511667 N 0.116667 W, ground altitude 30 m a.s.l.) on the Strand in  
82 central London. Although the site is within the London Congestion Charge Zone (an area  
83 encompassing central London requiring road tolls to be paid and hence an area with reduced  
84 traffic density), surrounding roads supported a medium to high traffic volume (annual average  
85 of 50000-80000 vehicles per day, (Department for Transport, 2014)) with the River Thames  
86 situated 200 m to the south. This site is classified as Local Climate Zone (LCZ) Class 2  
87 Compact Midrise according to Stewart and Oke (2012) (i.e. dense mix of midrise buildings (3–

88 9 stories), few or no trees, land cover mostly paved, stone, brick, tile, and concrete  
89 construction materials). Land cover types (in %) were calculated based on the Ordinance  
90 Survey map for the 9 km<sup>2</sup> area (Figure 1) encompassing the site and are: roads (37 %),  
91 buildings (31 %), other paved areas (14 %), unpaved/ vegetation (11 %), and water bodies (7  
92 %).

93 The sampling inlet and sonic anemometer were mounted on a triangular mast (Aluma T45-H)  
94 at approx. 60.9 m (2.3 times mean building height,  $z_H$ ) above ground level (a.g.l.). The mean  
95 building height was around 25 m and the mast was located on an elevated area in the centre  
96 of the roof. A street canyon was located to the NW and an enclosed parking area to the SE,  
97 but generally surrounding buildings were of equal height. The sampling point (which we call  
98 KCL) is located 37 m west of a sampling point (KSS) that has been used for long-term energy  
99 and CO<sub>2</sub> flux measurements (Kotthaus and Grimmond, 2012). Although the site is not optimal  
100 for micrometeorological flux measurements due to the heterogeneity of the urban canopy, its  
101 suitability has been assessed in detail by Kotthaus and Grimmond (2014a; 2014b). This study  
102 describes in detail the measurement area and investigates the influence of source area  
103 characteristics on long-term radiation and turbulent heat fluxes for the KSS site. They conclude  
104 that the site can yield reasonable data on surface to atmosphere fluxes.

105 The weather in 2012 was somewhat cooler than the 1981 to 2010 long-term mean for London  
106 during summer and autumn, with several cold fronts bringing up to twice as much precipitation  
107 and associated winds as average, suppressing pollution levels. However, during the period of  
108 the Olympic and Paralympic Games (27<sup>th</sup> July – 12<sup>th</sup> August and 29<sup>th</sup> August – 9<sup>th</sup> September  
109 2012) the weather was hot and dry causing sustained pollution peaks. Winter 2012/2013 was  
110 generally warmer and drier in London than the 1981-2010 mean (Met Office, 2013).

## 111 **2.2 Instrumentation and data acquisition**

112 The CSAT3 sonic anemometer (Campbell Scientific) and inlet were faced toward the  
113 predominant wind direction (SW) to minimise flow distortion. Data from the sonic anemometer

114 were logged at a frequency of 10 Hz and flux measurements were calculated using 25 min  
115 averaging periods. The rotation angle  $\theta$ , used to correct measurements of the vertical  
116 wind velocity for minor misalignment of the sonic anemometer, showed no significant  
117 disturbance of the turbulence from interactions with the building when plotted against wind  
118 direction. Data were recorded in UTC (Universal Time Coordinated), which is one hour earlier  
119 than local time in summer and coincident with Greenwich Mean Time in winter. However, all  
120 analyses used local time.

121 VOC concentrations were measured using a high sensitivity proton transfer reaction-  
122 (quadrupole) mass spectrometer (PTR-MS) (Ionicon Analytik GmbH, Innsbruck, Austria) with  
123 three Varian turbo-molecular pumps (see for example de Gouw and Warneke, 2007; Hayward  
124 et al., 2002; Lindinger et al., 1998 for more detailed description of the instrument). Air was  
125 drawn through an inlet co-located with the sonic anemometer. Sample air was purged through  
126 a  $\sim 30$  m  $\frac{1}{2}$ " OD (3/8" ID) PTFE tube at a flow rate of 81 L  $\text{min}^{-1}$  to the PTR-MS, which was  
127 housed in a utility room below. The high flow rate ensured turbulent flow was maintained and  
128 signal attenuation minimised (Reynolds number,  $Re = 11177$ ). During the campaign, PTR-MS  
129 operating parameters were maintained at 1.95 mbar, 510 V and 50 °C for drift tube pressure,  
130 voltage and temperature respectively, to achieve an  $E/N$  ( $E$ : electric field strength,  $N$ : buffer  
131 gas number density) ratio of 123 Td ( $1 \text{ Td} = 10^{-17} \text{ V cm}^2$ ). This field strength forms a  
132 compromise between reagent ion clustering and fragmentation suppression (Hewitt et al.,  
133 2003). Further instrument parameters and meteorological conditions are summarized in Table  
134 1. The inlet flow rate into the instrument was 0.25-0.3 L  $\text{min}^{-1}$ .

135 The logging program was written in LabVIEW (National Instruments, Austin, Texas, USA) and  
136 operated the PTR-MS in multiple ion detection (MID) and SCAN modes for VOC  
137 concentrations of nine selected masses and a range of the protonated mass spectrum  $m/z$   
138 21-206 respectively. The sonic anemometer was not directly interfaced with the LabVIEW  
139 logging program, requiring the measurements to be synchronised during post-processing  
140 through the use of a cross-correlation function between the vertical wind velocity  $w$  and the

141 VOC ion counts *c.* A valve system controlled the measurement cycle, which consisted of 5  
142 min zero air (ZA), 25 min MID followed by further 5 min SCAN of sample air and 25 min MID  
143 mode. During the ZA cycle, air was pumped through a custom-made gas calibration unit  
144 (GCU) fitted with a platinum catalyst heated to 200°C to provide instrument background values  
145 at ambient humidity. In MID mode the quadrupole scanned nine predetermined protonated  
146 masses with a dwell time of 0.5 s each to which the following compounds were ascribed: *m/z*  
147 21 (indirectly quantified *m/z* 19 primary ion count via [H<sub>3</sub><sup>18</sup>O<sup>+</sup>]), *m/z* 33 (methanol), *m/z* 39  
148 (indirectly quantified *m/z* 37 first cluster [H<sub>3</sub>O<sup>+</sup> H<sub>2</sub>O<sup>+</sup>]), *m/z* 42 (acetonitrile, results not shown),  
149 *m/z* 45 (acetaldehyde), *m/z* 59 (acetone/propanal), *m/z* 69 (isoprene/furan), *m/z* 79 (benzene),  
150 *m/z* 93 (toluene), *m/z* 107 (C<sub>2</sub>-benzenes), and *m/z* 121 (C<sub>3</sub>-benzenes, results not shown). The  
151 total cycle time was 5.5 s. Secondary electron multiplier (SEM) voltage, as well as O<sub>2</sub><sup>+</sup> (*m/z*  
152 32) and photon “dark counts” (*m/z* 25) signals were monitored weekly.

153 The PTR-MS cannot distinguish between different compounds with the same integer mass,  
154 therefore isobaric interference can occur. For example, *m/z* 107 may result from several  
155 contributing C<sub>8</sub>-aromatics: ethyl benzene, (m+p)-xylene, o-xylene and some benzaldehyde  
156 (Warneke et al., 2003). Further interferences at measured *m/z* from additional compounds and  
157 fragmentation for this instrument in an urban environment are discussed in Valach et al.  
158 (2014). Although the O<sub>2</sub><sup>+</sup> and water cluster ions were kept < 2 % of the primary ion,  
159 interferences from <sup>17</sup>O<sup>+</sup> isotopes at *m/z* 33 were taken into account.

160 Single point calibrations were performed on-site once a month using a certified multiple  
161 component VOC gas standard (Ionimed, part of Ionicon Analytik GmbH, Austria), which was  
162 validated by cross-calibration with a second independent VOC standard (Apel Riemer  
163 Environmental Inc., CO, USA). Before and after the campaign, multistep calibrations were  
164 performed with both standards. Standards were diluted with catalytically converted zero air,  
165 since cylinder concentrations were approx. 1 ppm ± 5 % uncertainty (Ionimed Analytik) and  
166 0.5 ppm ± 10 % (Apel Riemer). Error propagation resulted in a total calibration uncertainty of  
167 < 20 %. Measured normalised instrument sensitivities (*S<sub>N</sub>*, Table 1) based on Taipale et al.

168 (2008) were used to convert normalised count rates (ncps) of protonated masses ( $RH^+$ ) to  
169 volume mixing ratios (Langford et al., 2010a). Only the o-xylene isomer was present in the  
170 Ionimed standard, which was used to determine instrument sensitivities for  $m/z$  107, but  
171 sensitivities agreed well when compared with sensitivities for p-xylene present in the Apel  
172 Riemer standard. Any remaining humidity effects on calibrations were previously investigated  
173 for this instrument and were found to be within the overall calibration uncertainty (Valach et  
174 al., 2014). Detection limits of VOC concentrations (Table 2) were calculated according to  
175 Taipale et al. (2008).

### 176 **2.3 Flux calculations and quality assessment**

177 Fluxes were calculated according to Karl et al. (2002) and Langford et al. (2009; 2010b) using:

$$178 \quad F = \frac{1}{n} \sum_{i=1}^n w' \left( \frac{i-t_{lag}}{\Delta_{tw}} \right) * c'(i), \quad (\text{Eq. 1})$$

179 where  $w'$  and  $c'$  are the instantaneous fluctuations around the mean vertical wind ( $w - \bar{w}$ ) and  
180 mean VOC concentration ( $c - \bar{c}$ ),  $n$  is the number of VOC concentration measurements per  
181 25 min averaging period ( $n = 273$ ),  $t_{lag}$  is the lag time between the wind and PTR-MS  
182 measurement due to the transit through the sampling line, and  $\Delta_{tw}$  is the sampling interval of  
183 the vertical wind speed measurements of the sonic anemometer (10 Hz = 0.1 s). Langford et  
184 al. (in review) recently demonstrated that the method used to determine the time lag becomes  
185 important where the signal-to-noise ratio of the analyser is poor, showing that methods that  
186 systematically search for a maximum in the cross-correlation function within a given window  
187 (MAX method) can bias the calculated fluxes towards more extreme (positive or negative)  
188 values. Their study recommends the use of a prescribed lag time determined either through  
189 the use of a monitored sample flow rate or by using the typical lag time derived by searching  
190 for a maximum. Here the prescribed lag times were determined by fitting a running mean to  
191 the time series of daytime lag times calculated using the MAX method for acetone, which had  
192 large fluxes and thus the clearest time-lags. Prescribed lag times for all other compounds were



193 set relative to that of acetone, accounting for the offset introduced by the sequential sampling  
194 of the PTR-MS.

195 Flux losses due to the attenuation of high and low frequency eddies were estimated for our  
196 measurement setup. High frequency flux attenuation was estimated to be on average 11 %  
197 using the method of Horst (1997), and a correction was applied. Attenuation from low  
198 frequency fluctuations for a 25 min flux period was investigated by re-analysing the sensible  
199 heat fluxes for longer averaging periods of 60, 90, 120 and 150 min. The coordinate rotation  
200 was applied to the joined files, which acted as a high pass filter to the three wind vectors,  
201 confirming that fluctuations of eddies with a longer time period than the averaging time did not  
202 contribute to the flux measurement (Moncrieff et al., 2004). The fluxes were compared back  
203 to the 25 min average fluxes, which had the coordinate rotation applied before joining, again  
204 to ensure only turbulent fluctuations of  $\leq 25$  min contributed to the flux (Supplementary  
205 information Figure A1). Flux losses due to low frequency attenuation were estimated to be <  
206 1.5 % and, therefore, no corrections were deemed necessary. The error due to the disjunct  
207 sampling was estimated by comparing the sensible heat fluxes calculated from the continuous  
208 data series with those calculated from a disjunct data series using a set sampling interval of  
209 5.5 s. The continuous data were averaged to match the sampling frequency of the disjunct  
210 data (i.e. 2 Hz). The difference between the eddy covariance and DEC sensible heat fluxes  
211 was minimal (0.01 %) and thus no additional corrections were applied.

212 Many of the 25 min resolved flux measurements were close to the limit of detection (LoD),  
213 based on 1 standard deviation using the method of Spirig et al. (2005), with an average fail  
214 rate of 82 %. Various techniques to statistically analyse or replace values below the LoD have  
215 been developed (Clarke, 1998). However, they often result in significant bias, either high or  
216 low depending on the value substituted, because values tend to be below the LoD when fluxes  
217 are indeed small (Helsel and Hirsch, 1992). In this study, our analysis focused on diurnally  
218 averaged flux profiles and we decided not to filter out individual flux values on the basis of  
219 being < LoD in order to avoid this bias. When averaging the 25 min flux data it is appropriate

220 to also average the LoD which, as shown by Langford et al. (in review), decreases with the  
221 square root of the number of samples averaged ( $N$ ). Therefore, although the majority of the  
222 individual 25 min flux measurements were below the LoD, their diurnal average profiles may  
223 exceed the LoD for the average and thus still yield important data on the net exchange of  
224 VOCs above the city.

$$225 \quad \overline{LoD} = \frac{1}{N} \sqrt{\sum_{i=1}^N LoD^2}. \quad (\text{Eq. 2})$$

226 The following describes the additionally applied filter criteria. 25 min flux values with a friction  
227 velocity ( $u^*$ )  $< 0.15 \text{ m s}^{-1}$  were rejected (3.4 % of total data) due to insufficient turbulence. The  
228 stationarity test and data quality rating methods of Foken and Wichura (1996) and Velasco et  
229 al. (2005) were used, and 47 % of the data files were rejected on this basis. The high number  
230 of files rejected in the stationarity test is to be expected for eddy covariance measurements  
231 over highly heterogeneous canopies, although horizontally averaged canopy morphology  
232 recovers some surface homogeneity. Furthermore, the low measurement height used can  
233 cause an increased sensitivity towards canopy roughness features resulting in non-  
234 stationarity. Since urban environments are inherently not ideal for micrometeorological flux  
235 measurements due to their heterogeneity, integral turbulence characteristics of this site were  
236 assessed by comparing the measured standard deviation of the vertical wind velocity ( $\sigma_w$ )  
237 normalised by  $u^*$  to the parameters of a modelled ideal turbulence (Foken et al., 2004). Results  
238 showed that 99.6 % of all the data were rated category six or better and 0.4 % were rejected  
239 using the criteria of Foken et al. (2004). This large pass rate gives further confidence that the  
240 measurements were not unduly affected by wake turbulence generated from the structure of  
241 the building. Erroneous meteorological data (2.6 % of total) were removed around wind  
242 directions of 14-15°, due to minor turbulence interferences from the presence of other sensors  
243 on the mast. Depending on compound between 40-61 % of flux data ( $N = 1934\text{--}2949$ ) passed  
244 all of the above quality controls. Exactly 2014 h of concentration data ( $N = 4834$ ) were  
245 obtained. For consistency regression coefficients ( $R^2$ ) were used throughout.

246 The traffic densities used for the analysis were obtained from a nearby site at Marylebone  
247 Road (approx. 3 km to the NW) and consisted of hourly vehicle counts covering the period 7<sup>th</sup>  
248 - 22<sup>nd</sup> August 2012. The major roads of the Strand and the Embankment surrounding the  
249 measurement site support a comparable traffic volume with an annual average of 50000 -  
250 80000 vehicles per day (Department for Transport, 2014) and diurnal patterns in traffic are  
251 likely to be similar across central London.

252 Photosynthetically active radiation (PAR) and CO<sub>2</sub> measurements used in the analysis were  
253 part of the long-term micrometeorological measurements at the same site and covered the  
254 period from August to September for PAR and from August to December for CO<sub>2</sub> respectively.  
255 Average diurnal profiles were calculated for the boundary layer mixing height, which was  
256 measured using three LiDARs located on rooftops within central London during an approx.  
257 two week period in summer and winter 2012 (Bohnenstengel et al., 2015).

### 258 **2.3.1 Flux footprint calculations**

259 Although there are no operational footprint models for urban environments which take the  
260 complex topography and spatial variability in building height and surface heat fluxes into  
261 account, the analytical footprint model of Kormann and Meixner (2001) has previously been  
262 applied in non-homogeneous terrain (Helfter et al., 2011; Neftel et al., 2008). The Kormann-  
263 Meixner (KM) model determines the 2D footprint density function explicitly from  
264 micrometeorological parameters, which are provided by the eddy covariance measurements,  
265 i.e., friction velocity ( $u^*$ ), measurement height ( $z_m$ ), Obukhov length ( $L$ ), horizontal wind velocity  
266 at the measurement height ( $u(z_m)$ ), and standard deviation of the lateral wind ( $\sigma_v$ ). The flux  
267 footprints were calculated for each 25 min flux period. Neftel et al. (2008) developed a  
268 Microsoft Excel based tool, which allows the footprint contributions (%) of user-defined spatial  
269 elements to be mapped. In this case we used a total of nine 1 km<sup>2</sup> grid squares to match the  
270 Ordnance Survey (OS) grid (Figure 1), centred on the measurement site. This grid resolution  
271 was validated using a simple parameterisation model (Kljun et al., 2004) with average diurnal  
272 cycle parameters for  $\sigma_w$ ,  $u^*$ , and boundary layer height ( $z_i$ ) during the campaign, which

273 calculated the distance of the maximum flux contribution ( $X_{max}$ ) and the extent of the 90 % flux  
274 footprint ( $X_{90}$ ).

275 The KM footprint calculation requires the Monin-Obukhov stability parameter ( $\zeta$ ) to be within  
276 the interval [-3, 3], where

$$277 \quad \zeta = \frac{z_m - d}{L}, \quad (\text{Eq. 3})$$

278 with  $d$  ( $d = \frac{2}{3}z_H = 16.7$  m) being the displacement height estimated as a fraction of the canopy  
279 height (Garrat, 1992). The footprint estimation for cases of extreme stability is of lower quality,  
280 but still provides useful information. The vertical turbulent flux  $F_c(0,0,z_m)$  measured at the  
281 height  $z_m$  is related to the corresponding surface flux area  $F_c(x,y,0)$  which is upwind of the  
282 measurement point, such that

$$283 \quad F_c(0,0,z_m - d) = \int_{-\infty}^{\infty} \int_0^{\infty} F_c(x,y,z_m - d)\Phi(x,y,z_m - d)dx dy, \quad (\text{Eq. 4})$$

284 where  $z_m$  is the measurement height and the x-axis is aligned with the mean horizontal wind  
285 direction.  $\Phi(x,y,z_m - d)$  is the footprint function and includes a weighting function to describe  
286 the influence of a unit point source on the flux from any surface location  $(x,y)$ . In order to  
287 compare VOC fluxes with estimated emissions from the London Atmospheric Emissions  
288 Inventory (LAEI), a 9 km<sup>2</sup> section of the 1 km<sup>2</sup> resolution OS grid system was used, which on  
289 average included 90 % of the footprint contribution to all measured fluxes. This area was  
290 limited to central London and partially included the following Boroughs: Westminster (squares  
291 1, 4, 5 and 7), Southwark (2, 3 and 6), Camden (8) and the City of London (9) (Figure 1).

## 292 **3 Results and Discussion**

### 293 **3.1 Diurnal profiles of VOC fluxes and concentrations**

294 Average diurnal cycles of measured VOC fluxes and mixing ratios are shown in Figure 2a and  
295 2b with descriptive statistics for all the data summarized in Table 2. Largest median

296 (interquartile range in parenthesis) fluxes per day were from C<sub>2</sub>-benzenes and toluene with  
297 7.86 (0.92-21.8) kg km<sup>-2</sup> d<sup>-1</sup> and 7.26 (1.83-15.3) kg km<sup>-2</sup> d<sup>-1</sup> respectively, followed by  
298 oxygenated compounds, i.e. methanol with 6.37 (2.99-10.0) kg km<sup>-2</sup> d<sup>-1</sup>, acetaldehyde 3.29  
299 (1.52-5.62) kg km<sup>-2</sup> d<sup>-1</sup>, and acetone 5.24 (2.33-9.62) kg km<sup>-2</sup> d<sup>-1</sup>. Isoprene and benzene  
300 showed smallest median fluxes with 2.14 (0.56-4.85) kg km<sup>-2</sup> d<sup>-1</sup> and 1.78 (0.06-4.34) kg km<sup>-2</sup>  
301 d<sup>-1</sup> respectively. The highest median mixing ratios were of the oxygenated compounds  
302 methanol (7.3 (6.8-7.9) ppb), acetone (0.95 (<LoD-1.36) ppb) and acetaldehyde (0.82 (0.59-  
303 1.13) ppb), followed by aromatics (C<sub>2</sub>-benzenes, toluene and benzene), and isoprene.

304 Oxygenated compounds commonly have relatively long atmospheric lifetimes and widespread  
305 origin including anthropogenic and biogenic sources and photochemistry, resulting in elevated  
306 concentrations and less pronounced diurnal profiles (Atkinson, 2000). Most VOC fluxes and  
307 concentrations were comparable to or lower than those previously observed in London  
308 (Langford et al., 2010b) and other UK cities (Langford et al., 2009), although C<sub>2</sub>-benzene  
309 fluxes and concentrations, as well as isoprene and benzene concentrations were slightly  
310 higher. The discrepancy in isoprene and benzene concentrations is consistent with  
311 photochemical loss during transport to the higher measurement height of the previous studies.  
312 Compared to other cities such as Houston Texas (Park et al., 2010) and Mexico City (Velasco  
313 et al., 2005), VOC fluxes and concentrations were lower, apart from C<sub>2</sub>-benzenes which were  
314 comparable or higher, although it must be noted that C<sub>2</sub>-benzenes in this study represent the  
315 sum of multiple VOC species. Unlike the other studies cited, Park et al. (2010) use relaxed  
316 eddy accumulation to measure VOC fluxes and hence the data obtained are not directly  
317 comparable with measurements made by EC-based methods.

318 Diurnal profiles of aromatic fluxes and concentrations presented clear double rush hour peaks  
319 during the morning and evening (07:00-10:00 and 17:00-20:00 local time). Concentration  
320 peaks are thought to be linked to additional advection of traffic-related pollution from larger  
321 commuter roads outside of the city centre, as well as boundary layer effects and  
322 photochemistry. VOC concentration measurements at canopy height can be affected by

323 boundary layer depth (Vilà-Guerau de Arellano et al., 2009). The rush hour emission peaks  
324 mostly coincide with the boundary layer expansion and collapse and therefore the effect of  
325 each factor cannot be separated. The morning concentration peak was slightly higher than the  
326 evening peak across traffic-related species even though fluxes tended to be larger during the  
327 evening rush hour. Morning emissions enter a shallow nocturnal boundary layer leading to  
328 relatively larger concentrations compared with higher afternoon emissions entering a  
329 developed boundary layer, leading to relatively lower concentrations. This enhanced dilution  
330 effect is found more often during summer when the boundary layer mixing height is higher  
331 (Figure 2). Therefore, the regression analyses below only refer to data from August (cf Section  
332 3.1.2 for comparisons with winter). Furthermore, increased photochemical degradation during  
333 the day removes VOCs, further contributing to the midday minimum in mixing ratios. The  
334 diurnal flux profiles of methanol, acetone, isoprene, and to a smaller extent acetaldehyde,  
335 showed one large peak just after midday (approx. 13:00 local time), which was only reflected  
336 in the concentration profiles of acetone and isoprene. Acetaldehyde concentrations presented  
337 a slight double peak similar to mixing ratios of aromatics. Methanol has a relatively long  
338 atmospheric lifetime and therefore high background concentrations, hence mixing ratios  
339 showed no distinct diurnal profile.

### 340 **3.1.1 Correlations with possible controlling variables of VOC fluxes and concentrations**

341 Aromatic compound fluxes closely followed the diurnal profile of traffic density with good  
342 correlations ( $R^2 = 0.51-0.92$ ,  $p < 0.05$ ) and slightly lower fluxes observed on the weekends. In  
343 central urban areas in the UK traffic densities — and therefore traffic-related VOC fluxes —  
344 increase steadily throughout the day, with discernible peaks during morning, midday and  
345 evening (Nemitz et al., 2002), which was also observed in this study. Previous studies have  
346 shown that the Marylebone Rd traffic count point can be used as a proxy representative of  
347 traffic flows throughout central London (Helfter et al., 2011).

348 The aforementioned concentration dilution due to boundary layer expansion resulted in  
349 negative correlations between boundary layer height and aromatic mixing ratios during August

350 ( $R^2 = 0.33-0.56$ ,  $p < 0.01$ ). As aromatic compound fluxes slightly dipped around midday, the  
351 mixing ratios were diluted by the deep boundary layer. The above evidence suggests that  
352 traffic-related emissions were the main contributors to fluxes and mixing ratios of aromatic  
353 compounds. Acetone and isoprene showed peak midday fluxes, which maintained daytime  
354 mixing ratios and produced positive correlations with boundary layer height ( $R^2 = 0.16$  and  
355  $0.59$  respectively,  $p < 0.01$ ). De Gouw et al. (2005) reported that changes in boundary layer  
356 meteorology could result in greater effects on observed concentrations of methanol and  
357 acetone due to their high background values. The mixing ratios of these compounds are,  
358 therefore, likely dominated by advected pollution rather than the local flux. Possibly a  
359 combination of boundary layer and photochemical effects were seen with methanol mixing  
360 ratios, wherein correlations with mixing height were negative ( $R^2 = 0.70$ ,  $p < 0.01$ ), whereas  
361 acetone and isoprene fluxes seemed to be sufficiently high during the day to maintain peak  
362 midday mixing ratios (Figure 3 example of isoprene). Vehicle emissions may have contributed  
363 to acetaldehyde and isoprene levels directly or indirectly (Figure 3 example of isoprene), since  
364 correlations of fluxes with traffic density were fairly high ( $R^2 = 0.60$  and  $0.46$  respectively,  
365  $p < 0.05$ ). The diurnal concentration profile of acetaldehyde to some degree mimicked those of  
366 traffic-related compounds reflecting a slight double peak.

367 VOC fluxes and concentrations plotted as a function of photosynthetically active radiation  
368 (PAR) showed strong daytime (defined as 06:00 to 18:00 local time) correlations for methanol,  
369 acetaldehyde and isoprene fluxes ( $R^2 = 0.71-0.78$ ,  $p < 0.001$ ) and concentrations ( $R^2 = 0.66-$   
370  $0.83$ ,  $p < 0.001$ ). Plotted as a function of temperature, high correlations with methanol,  
371 acetaldehyde and isoprene fluxes were seen ( $R^2 = 0.75$ ,  $0.63$ , and  $0.94$ ,  $p < 0.001$  respectively),  
372 whereas only methanol and acetone concentrations showed higher correlations with  
373 temperature ( $R^2 = 0.64$  and  $0.81$ ,  $p < 0.001$  respectively). Methanol fluxes correlated linearly  
374 with temperature ( $R^2 = 0.75$ ,  $p < 0.001$ ), but acetaldehyde and isoprene fluxes ( $R^2 = 0.64$  and  
375  $0.94$ ,  $p < 0.01$ ) and mixing ratios ( $R^2 = 0.45$  and  $0.55$ ,  $p < 0.01$ ) had exponential relationships  
376 with temperature (Figure 3 example of isoprene). The relationships of mixing ratios with PAR

377 and temperature for these compounds improved greatly when night time values were excluded  
378 (defined as PAR  $<100 \mu\text{mol m}^{-2} \text{s}^{-1}$ ) and when times of low temperature ( $< 5 \text{ }^\circ\text{C}$ ) were  
379 excluded. This indicates either separate source contributions or effects of boundary layer  
380 meteorology in these instances, whereby increased mixing ratios of these compounds with  
381 low PAR and temperature likely result from reduced dilution within a shallow boundary layer,  
382 e.g., at night or in winter, but also possible contributions of anthropogenic sources such as  
383 exhaust emissions, which are largely independent of light and temperature. Increases in  
384 concentrations due to high PAR and temperature suggest biogenic sources, increased  
385 evaporative emissions, and/or secondary atmospheric formation driven by oxidation of  
386 precursor hydrocarbons (Singh et al., 1994). Oxygenated compounds have a variety of  
387 different source contributions such as tailpipe emissions, evaporative emissions from fuel and  
388 solvents, direct emissions from plants, leaf decomposition, and secondary atmospheric  
389 production (Langford et al., 2009 and references therein).

390 Modelling studies have indicated that the contribution of secondary atmospheric formation to  
391 VOC concentrations could be more significant, especially in urban areas, during summer, i.e.  
392 with high PAR and temperatures (de Gouw et al., 2005; Harley and Cass, 1994). Acetone  
393 fluxes reached a maximum when PAR and temperature were around  $1000 \mu\text{mol m}^{-2} \text{s}^{-1}$  and  
394  $15\text{-}20 \text{ }^\circ\text{C}$  respectively, before declining, whereas mixing ratios increased exponentially with  
395 light and temperature. These observations resemble measurements over forest canopies (e.g.  
396 Schade and Goldstein, 2001). Aromatic compound concentrations and fluxes showed no  
397 correlations with PAR. Weak negative correlations were seen between aromatic  
398 concentrations and temperature and weakly positive correlations between fluxes and  
399 temperature, likely due to increased thermal mixing. The observed light and temperature  
400 responses associated with isoprene fluxes and mixing ratios in August and September can be  
401 explained by biogenic sources (cf Section 3.1.3).



### 402 3.1.2 Seasonal variability of VOC sources and meteorology

403 Most compounds showed larger fluxes in August and September than in October, November  
404 and December, with the exception of acetaldehyde which also showed increased fluxes in  
405 December (Figure 4 top). Increased acetaldehyde fluxes in December may have resulted from  
406 an additional source, such as domestic biomass burning (Andreae and Merlet, 2001; Lipari et  
407 al., 1984), although there are only few residential buildings in this area of London. Only toluene  
408 fluxes in September were significantly higher than in other months and benzene fluxes showed  
409 no significant seasonal differences. Seasonal variability in fluxes was likely due to increased  
410 emissions in summer, especially for compounds with biogenic and secondary atmospheric  
411 sources. Average monthly meteorological parameters are summarized in Table 3.

412 Mixing ratios of aromatics were generally lower in summer and highest in December (Figure  
413 4 bottom). This is likely due to less dilution effects in winter when the boundary layer is shallow  
414 or from advection of additional sources such as heating, since there was no increase in fluxes.  
415 Generally, in summer the boundary layer mixing height is higher and collapses later in the  
416 evening which maintains the dilution effect for VOC concentrations. In winter the average  
417 boundary layer mixing height is lower. It develops later in the morning and collapses earlier in  
418 the afternoon, which could increase overall VOC mixing ratios, but also individual maxima,  
419 e.g. during rush hours. Comparing average diurnal profiles of compound mixing ratios with  
420 boundary layer height during summer and winter showed that aromatic compound  
421 concentrations were associated with negative correlations in summer (cf Section 3.1.1) which  
422 became positive during winter ( $R^2 = 0.10-0.33$ ,  $p < 0.01$ ), while fluxes maintained positive  
423 correlations with boundary layer height regardless of season. This suggests boundary layer  
424 effects may be an important driver of increased concentrations in winter. Furthermore, traffic  
425 counts for the Congestion Charge Zone in central London indicate lower monthly average  
426 vehicle counts in December (Department for Transport, 2014). Oxygenated compounds and  
427 isoprene mixing ratios were highest in summer with the exception of acetone, which increased  
428 in December likely either from boundary layer effects, reduced photochemical degradation, or

429 advection. Correlations of mixing ratios and fluxes with boundary layer height were positive  
430 for acetone and isoprene during summer and winter, whereas methanol and acetaldehyde  
431 presented negative correlations during summer indicating stronger dilution effects (cf Section  
432 3.1.1).

433 Increased summer mixing ratios of oxygenated compounds and isoprene indicated a  
434 temperature dependent, possibly biogenic source contribution. While biogenic emissions may  
435 be advected from outside of the city, the concurrent increase in isoprene fluxes suggests the  
436 source to be largely local to the flux footprint. The temperature dependent fraction of observed  
437 isoprene mixing ratios, which may include advected pollution, was estimated using the  
438 isoprene temperature response function from Figure 9 in Langford et al. (2010b), which  
439 estimated a 30 % and 20 % contribution in August and September respectively. These values  
440 were significantly higher than for iso-pentane, a non-biogenic compound available from the  
441 Automatic Hydrocarbon Network, to which the same analysis was applied. The temperature  
442 dependent component of isoprene in October, November and December showed no  
443 significant difference to that of iso-pentane, suggesting the biogenic component was reduced  
444 or absent at lower temperatures. High correlations of  $m/z$  69 with light and temperature during  
445 August and September indicate that isoprene was the likely major component during these  
446 months, however the rest of the period the contribution of additional compounds such as furan  
447 and other alkenes at that mass may have increased, thereby overestimating the isoprene  
448 signal (Yuan et al., 2014).

### 449 **3.1.3 Modelling the biogenic isoprene contribution in London**

450 An attempt was made to model the biogenic isoprene component during August and  
451 September using the light and temperature algorithms of Guenther et al. (1995), hereafter  
452 termed G95. The foliar emissions based model calculates VOC fluxes as follows;

$$453 \quad F = D \times \varepsilon \times \gamma, \quad (\text{Eq. 5})$$

454 where  $D$  is the foliar density ( $\text{kg dry matter m}^{-2}$ ),  $\epsilon$  is an ecosystem dependent base emission  
455 rate ( $\mu\text{g C m}^{-2} \text{s}^{-1}$  normalised to a PAR flux of  $1000 \mu\text{mol m}^{-2} \text{s}^{-1}$  and leaf temperature of  $303.15$   
456 K), and  $\gamma$  is a dimensionless activity adjustment factor accounting for the effects of PAR and  
457 leaf temperature. Ambient air temperature and PAR measurements were used to calculate  
458 the light and temperature controlled parameters  $C_L$  and  $C_T$  for  $\gamma$ , where

$$459 \quad \gamma = C_L \times C_T. \quad (\text{Eq. 6})$$

460 The slope of the linear regression of the measured total isoprene flux and  $\gamma$  provided an  
461 emission factor in  $\text{mg m}^{-2} \text{h}^{-1}$ , which was converted to  $\mu\text{g g}^{-1} \text{h}^{-1}$  by dividing by the foliar density  
462 ( $D = 0.129 \text{ kg m}^{-2}$ ). The foliar density was estimated using the total tree leaf area as seen from  
463 visible satellite imagery within the flux footprint and tree leaf dry weight for representative  
464 species commonly planted in the area such as *Platanus x acerifolia* (City of Westminster,  
465 2009), which are also high isoprene emitters (Geron et al., 1994). The resulting base emission  
466 rate  $\epsilon$  from the measured fluxes was  $6.5 \mu\text{g g}^{-1} \text{h}^{-1}$  which compares well with the figure given  
467 in the literature ( $5 \mu\text{g g}^{-1} \text{h}^{-1}$ ) for cities in a cool climate (Guenther et al., 1995). For details of  
468 this calculation, see the Supplementary Information B. These estimates are representative of  
469 the biogenic isoprene fluxes from a highly heterogeneous canopy within the flux footprint,  
470 including both high and low isoprene emitting species as well as low average foliar density  
471 due to the sparse distribution of urban roadside and park trees. Green areas, as defined on  
472 the OS map, comprised 9 % of the total grid area and were evenly distributed across the 9  
473  $\text{km}^2$ . Only grid square 1 included a large green area of 23 ha (St. James' Park). The National  
474 Forest Inventory (NFI) England only included 4.4 % green areas within the grid selection (NFI,  
475 2012). The NFI excluded individual trees in parks and avenues, which can encompass up to  
476 50 % of trees maintained by the local authority in central London (City of Westminster, 2009).

477 Figure 5a and b shows that the modelled isoprene fluxes using the calculated base emission  
478 rate compared well with the measured fluxes by wind direction. Linear regressions from wind  
479 directions that have a strong anthropogenic component are lower, e.g. W ( $R^2 = 0.13$ ,  $p < 0.001$ ),

480 than from those areas dominated by biogenic sources, e.g. SE ( $R^2 = 0.81$ ,  $p < 0.001$ ) due to  
481 the nearby Temple Gardens. Modelled emissions seemingly underestimated observed  
482 isoprene fluxes since these included the traffic component, however it appears that biogenic  
483 isoprene represents a detectable source contribution in summer.

### 484 **3.2 VOC/VOC correlations and ratios**

485 Correlations of VOC/VOC fluxes ( $R^2 = 0.40-0.62$ ,  $p < 0.001$ ) indicated two groups of  
486 compounds with good correlations within each group, i.e. compounds related to traffic sources  
487 such as aromatics, and oxygenated and biogenic compounds, such as methanol, acetone and  
488 isoprene (Figure 6 top). Correlations of VOC/VOC concentrations ( $R^2 = 0.13-0.84$ ,  $p < 0.001$ )  
489 showed highest correlations between traffic-related compounds ( $R^2 = 0.45-0.84$ ,  $p < 0.001$ ) and  
490 good correlations between the oxygenated and biogenic compounds ( $R^2 = 0.55-0.69$ ,  
491  $p < 0.001$ ) (Figure 6 bottom). High correlations between oxygenated VOCs could indicate  
492 source commonality or formation mechanisms that depend on similar environmental factors.  
493 Scatterplots between aromatic compounds and isoprene/oxygenated compounds tend to  
494 show bimodal distributions indicating separate source contributions. Using temperature or, to  
495 a smaller extent, PAR as a third variable highlights a temperature or light dependency of the  
496 second source supporting the existence of additional biogenic and/or atmospheric sources. In  
497 the example of isoprene against benzene the relationship changes with temperature from 2:1  
498 to 1:2.

#### 499 **3.2.1 Benzene to toluene ratios**

500 Benzene to toluene (b/t) ratios can help identify source types and changes in ratios can  
501 indicate the photochemical age of an air mass as toluene reacts at a faster rate with OH in the  
502 atmosphere, assuming sufficient OH concentrations to drive the reaction (Warneke et al.,  
503 2007). Median (and interquartile range, IQR) b/t flux ratios were 0.21 (0.02-0.43) and median  
504 (IQR) b/t concentration ratios were 0.45 (0.39-0.48). Individual maxima and minima were seen  
505 in the b/t concentration ratios, examples of which are discussed below.

506 The observed ratios compared well with those of other European cities, which showed b/t  
507 concentration ratios of 0.35 in Zurich (Heeb et al., 2000), 0.57 in Manchester (Langford et al.,  
508 2009), 0.57-0.63 in London (Valach et al., 2014), and 0.1 at 190 m above London (Langford  
509 et al., 2010b). Traffic related emissions are considered to be an important source of benzene  
510 and toluene in London. B/t exhaust emission ratios based on derived yearly emissions in other  
511 megacities, such as Mexico City, were found to be 0.4 (Zavala et al., 2006), which agreed well  
512 with observed b/t concentration ratios in this study. Airborne flux measurements over Mexico  
513 City have shown average b/t flux ratios of 0.31 with lower ratios of 0.07 to 0.1 over industrial  
514 areas due to increased toluene emissions from industrial processes (Karl et al., 2009; Velasco  
515 et al., 2007). Evaporative emissions from gasoline or direct industrial toluene emissions may  
516 have contributed to the lower b/t flux ratios in London. Furthermore, low b/t concentration ratios  
517 of 0.26 from diesel emissions have been reported (Corrêa and Arbilla, 2006). The widespread  
518 use of diesel fuel in London (buses, taxis and some cars and trains) and diesel emissions from  
519 roads which exclude passenger cars, such as Oxford Street (approx. 1.3 km W from the  
520 measurement site) or central railway nodes, such as Waterloo Railway Station (1 km to the  
521 S), may have affected b/t ratios.

522 Wind speed and direction can play a role for b/t concentration ratios by transporting pollution  
523 over longer distances allowing more time to react with or exposure to higher OH  
524 concentrations, thus increasing the ratio. An example of this (Figure 7) was seen on the 12<sup>th</sup>  
525 August when median (IQR) b/t concentration ratios reached 0.5 (0.45-0.56) with stronger SE  
526 winds (mean 3.67 m s<sup>-1</sup>) possibly advecting pollution from Benelux/Northern Europe, whereas  
527 on the 9<sup>th</sup> August median b/t ratios were 0.34 (0.30-0.38) with low wind speeds (mean 1.28 m  
528 s<sup>-1</sup>) indicating higher contributions of local sources (i.e. 60 % London influence)  
529 (Bohnenstengel et al., 2015). On both days OH concentrations above London were around  
530  $1.25 \times 10^6$  molecules cm<sup>-3</sup> and b/t flux ratios were not significantly different making pollution  
531 advection a likely cause of the observed difference (L. Whalley, personal communication  
532 2014). Calculated back trajectories using the HYSPLIT trajectory model (Hybrid Single Particle

533 Lagrangian Integrated Trajectory Model (Draxler and Rolph, 2014)) were run at 3 h intervals  
534 starting at ground-level (10 m) from London and propagated 24 h backwards in time. These  
535 agreed with the changes in measured b/t ratios (Figure 7) and during episodes of high b/t  
536 ratios indicated that air masses passed over continental Europe within the past day which  
537 could have entrained pollution.

538 The median monthly b/t flux ratio during the measurement period stayed between 0.18 and  
539 0.26, which is to be expected since only local fluxes were detected, however the median (IQR)  
540 monthly b/t ratio for concentrations steadily increased from 0.41 (0.36-0.47) to 0.62 (0.55-  
541 0.70) from August to December. Advected pollution from mainland Europe may be common  
542 in winter or biomass burning may play a greater role in colder months, as this is associated  
543 with higher b/t ratios, e.g. 1.67 (Lemieux et al., 2004), due to the different fuel combustion  
544 emission profile. Furthermore, OH concentrations in London are often below the detection limit  
545 during winter (Bohnenstengel et al., 2015) resulting in less local photochemical removal during  
546 the winter months.

547 Median (IQR) concentration ratios for benzene to C<sub>2</sub>-benzenes were 0.31 (0.28-0.33) and  
548 toluene to C<sub>2</sub>-benenes were 0.72 (0.63-0.81), which both agree with previous values and  
549 suggest that these masses are indeed the ascribed traffic-related compounds (Heeb et al.,  
550 2000; Warneke et al, 2001).

### 551 **3.2.2 VOC to CO<sub>2</sub> correlations and ratios**

552 Generally, good correlations were found between averaged VOC and CO<sub>2</sub> fluxes measured  
553 concurrently at the site ( $R^2 = 0.03-0.81$ ,  $p < 0.001$ ). Traffic-related compounds were among the  
554 lowest correlations with CO<sub>2</sub> fluxes ( $R^2 = 0.03-0.48$ ,  $p < 0.01$ ). However, when points of peak  
555 CO<sub>2</sub> fluxes were removed the correlations with traffic-related VOC fluxes increased  
556 significantly to  $R^2 = 0.65-0.91$  ( $p < 0.001$ ). Presumably, the initial poor correlations resulted from  
557 an additional strong CO<sub>2</sub> source that is not a shared source of aromatic VOCs. The improved  
558 correlation is greater for traffic-related compounds due to the limited range of source

559 contributions to this group compared with oxygenated/biogenic compounds. The regression  
560 coefficient ( $R^2$ ) of benzene with  $\text{CO}_2$  fluxes increases from 0.48 to 0.91, whereas for isoprene  
561 fluxes the increase was small, i.e. 0.68 to 0.70 (Figure 8), as isoprene shares only few common  
562 sources with  $\text{CO}_2$ . Averaged VOC to  $\text{CO}_2$  concentration correlations were highest for traffic-  
563 related compounds ( $R^2 = 0.92-0.96$ ,  $p < 0.001$ ) and lower for the other compounds ( $R^2 = 0.71-$   
564  $0.90$ ,  $p < 0.05$ ).

565 Median VOC/ $\text{CO}_2$  flux ratios ranged from  $1.7 \times 10^{-5}$  to  $7.7 \times 10^{-5}$  ( $\text{mg m}^{-2} \text{ h}^{-1} / \text{mg m}^{-2} \text{ h}^{-1}$ ) with  
566 isoprene and benzene showing low ratios due to their low fluxes, and toluene and C<sub>2</sub>-  
567 benzenes high ratios. Highest flux ratios for all compounds were with W winds, whereas lowest  
568 for biogenic compounds with N and for traffic-related compounds S wind directions. Flux ratios  
569 declined towards December as  $\text{CO}_2$  fluxes increased and VOC fluxes decreased. Similarly,  
570 VOC/ $\text{CO}_2$  concentration ratios were between  $0.45 \times 10^{-6}$  and  $14.6 \times 10^{-6}$  (ppb/ppb) with  
571 isoprene and benzene representing the lowest and methanol and acetone the highest ratios.  
572 Highest concentration ratios were seen in August for oxygenated compounds/isoprene and  
573 December for traffic-related species.

### 574 **3.3 Wind direction and flux footprint analysis**

575 Polar Annulus and Polar plots were constructed for VOC fluxes and mixing ratios respectively  
576 and representative compounds are shown (Figure 9). Polar plots use a generalized additive  
577 model (GAM) to interpolate between wind direction and wind speed averaged data points  
578 within the OpenAir package in R (see Carslaw and Ropkins, 2012; Hastie and Tibshirani,  
579 1990; Wood, 2006). Polar Annulus plots averaged by time of day instead of wind speed show  
580 diurnal variability with wind direction. The majority of the time (83 %) unstable and near neutral  
581 conditions prevailed ( $\zeta < 0.2$ ), although the frequency varied between months with 87 %, 89  
582 %, 82 %, 84 % and 69 % during August, September, October, November and December,  
583 respectively. Wind directions with mostly unstable conditions were with W and S winds and  
584 near neutral with N or E winds. Mixing ratios were on average highest with low wind speeds

585 (showing a negative correlation) when pollutants accumulate due to reduced mixing, indicating  
586 local emissions (Figure 9, bottom).

587 Largest fluxes for all compounds were from the NW with either one daytime peak (e.g.  
588 isoprene) or two distinct rush hour peaks (e.g. benzene) (Figure 9, top). On average fluxes  
589 were largest from the W>E≥N>S (F-statistic = 60.37-227.06, p<0.001), because of increased  
590 emission rates of specific compound sources. Separated by month, fluxes were largest from  
591 W>N>E≥S in August and September, whereas during October, November and December  
592 fluxes followed the pattern W>E≥N>S. The flux footprint in this study was relatively small  
593 compared with that of measurements previously made at 190 m height from the BT Tower in  
594 central London (Langford et al., 2010b). Due to the relatively low measurement height in this  
595 study, flux measurements were always closely coupled with the surface layer, unlike  
596 measurements by Langford et al. (2010b), which were at times disconnected from the surface  
597 layer during stable night time conditions.

598 The average length of the maximum flux footprint contribution ( $X_{max}$ ) was around 330 m and  
599 90 % of all the fluxes ( $X_{90}$ ) originated from within 900 m. The median footprint area was 1.8  
600 km<sup>2</sup>. This established that the majority of emission sources contributing to the measured fluxes  
601 must have been local. Additionally, the selected emission grid (cf Section 2.3.1 above)  
602 encompassed 97 % of the footprint with S and W wind directions, but only 80 % and 84 %  
603 during E and N winds. Grid square 5 represented the maximum contribution area since it  
604 encompassed the measurement point. Average footprint contributions (mean ± SD) comprised  
605 of grid squares 1 (2 % ± 4 %), 2 (5 % ± 7 %), 4 (4 % ± 5 %) and 5 (52 % ± 31 %) during S and  
606 W wind conditions, squares 6 (4 % ± 9 %) and 9 (4 % ± 10 %) indicated E wind conditions,  
607 and square 8 (18 % ± 27 %) N wind conditions. During October contributions from square 9  
608 increased to 10 % and were more frequent at 30 % in December. Squares 3 (0.6 % ± 2 %)  
609 and 7 (0.9 % ± 2 %) provided minimal average contributions.



610 The River Thames to the S may have caused the low fluxes associated with S winds (i.e.  
611 squares 1, 2 and 3). Contributions of traffic-related compound fluxes were statistically  
612 significant from the W (i.e. squares 4, 5, and 7), followed by the N (square 8) and E (squares  
613 6 and 9) likely from the nearby heavily trafficked roads (Kingsway, Charing Cross, Strand and  
614 Blackfriars areas, respectively). Biogenic compound fluxes were highest from the W and E  
615 which coincides with significant nearby green areas within the flux footprint.

616 Correlations of fluxes with grid square contributions in the footprint can also give information  
617 on emission source strengths within the respective grid square (Figure 1). Generally positive  
618 correlations with fluxes across most compounds were seen from the W (squares 4, 5 and 7)  
619 confirming that high emission rates from sources within these grid squares were driving the  
620 large fluxes. Strongest correlations of fluxes with contributions from squares 4, 5, and 7 were  
621 seen during October and November ( $R^2 = 0.40-0.46$ ,  $p < 0.001$ ), especially for masses  
622 associated with biogenic sources ( $m/z$  33, 45, 59 and 69). Square 8 showed positive  
623 correlations for benzene and only in August for all compounds. Correlations of fluxes with  
624 contributions from squares 1, 2, 3, 6 and 9 were negative indicating weaker emission sources  
625 in these squares or increased VOC deposition.

626 Highest mixing ratios with wind direction were from  $E > N \geq W > S$  for traffic-related compounds,  
627 whereas oxygenated compounds/isoprene followed a similar pattern as the fluxes of  
628  $W \geq E > N \geq S$  (F-statistic = 47.49-86.95,  $p < 0.001$ ). Easterly winds in London are often associated  
629 with synoptic conditions that bring European Continental air masses to the UK, resulting in  
630 higher background concentrations. Furthermore, since the boundary layer was on average  
631 more stably stratified and mixing heights were lowest ( $640 \pm 80$  m) with E wind conditions, it  
632 is likely that pollutant concentrations were allowed to build up resulting in the observed higher  
633 concentrations to the E for the more ubiquitous compounds, whereas concentrations of  
634 compounds with biogenic contributions additionally had strong sources to the W, such as  
635 several green areas (St. James' Park, Hyde Park and Regents Park, total 331 ha).

## 636 **3.4 Comparisons with London and National Atmospheric Emissions**

### 637 **Inventories**

638 The London Atmospheric Emissions Inventory (LAEI) and National Atmospheric Emissions  
639 Inventory (NAEI) produce yearly emission estimates over the 1 km<sup>2</sup> OS grid for a range of  
640 pollutants and emission sources. Total VOC emission estimates are provided, but only  
641 benzene and 1,3-butadiene are estimated separately. Measured emissions were compared  
642 with annual estimated emissions for the above OS grid area selection from 2012 for benzene  
643 using the LAEI and indirectly speciated VOCs of the NAEI. Using the average flux footprint,  
644 the grid square estimates were compared with the scaled flux measurements from the  
645 equivalent area (Figure 10).

646 LAEI emission estimates included contributions from major (69 %) and minor roads (4 %), as  
647 well as evaporative emissions (27 %) (LAEI, 2012). No data were available on cold start  
648 emissions for benzene. The calculated standard errors provided some uncertainty  
649 approximation. Measured fluxes compared well with emission estimates, although the LAEI  
650 predicted slightly smaller benzene fluxes. Comparisons of fluxes with wind directions (Section  
651 3.3) agreed well with the LAEI emission estimates for the respective grid squares with highest  
652 emissions from squares 4, 5, 7 and 8 (i.e. W and N directions). This comparison assumes that  
653 the benzene fluxes during the measurement period were representative of annual emissions  
654 with any significant seasonal variation in benzene emission rates captured in this five month  
655 period. Section 3.1.2 confirmed that there was little month-to-month variability in the benzene  
656 flux.

657 Using speciated VOC emission contributions (% of total VOC emissions) for 2006 (Bush et al.,  
658 2006) and emission maps from 2012 for total non-methane VOC emissions, speciated  
659 estimates could be compared with observations (Figure 10). The NAEI includes a wide range  
660 of emission sources divided into 11 SNAP (Selected Nomenclature for sources of Air Pollution)  
661 sectors including industrial, commercial and residential processes, transport, waste treatment,

662 solvent use, point sources, agriculture and nature, although the latter two were unavailable for  
663 the London urban area. NAEI estimates for benzene exceed the LAEI due to the inclusion of  
664 a wider range of sources beyond traffic-related emissions. Total C<sub>2</sub>-benzene emission  
665 estimates consisted of ethyl benzene, (m+p)-xylene and o-xylene. Benzene and methanol  
666 emissions agreed very well, however for all the other compounds estimated emissions were  
667 significantly lower than the measured fluxes. Uncertainties related to the measurements, such  
668 as isobaric interferences within the PTR-MS could have contributed to measurement  
669 overestimation, whereas uncertainties within the modelled emissions and the use of older  
670 speciation values may have impacted the estimates. In the case of isoprene, only minimal  
671 emissions are assumed, which do not include the biogenic sources that contributed to the  
672 measured fluxes. It is also likely that some of the *m/z* 69 signal could be attributed to cyclic  
673 alkenes, but Section 3.1.3 showed that biogenic isoprene provided a significant contribution  
674 during August and September in central London.

## 675 **4 Conclusion**

676 Our measurements show that vehicle emissions are the dominant source of the fluxes and  
677 concentrations of VOCs in central London, although biogenic sources and secondary  
678 atmospheric formation may make a significant contribution, particularly in summer for some  
679 compounds. There were observable spatial variations in flux rates, which result from the  
680 varying spatial distribution of emission types and strengths of emission sources, such as  
681 vegetation and traffic. Temporal variations in relative source strengths can be seen in the  
682 diurnal and seasonal profiles, reflecting the diurnality and seasonality of some of the driving  
683 factors. The measured VOC fluxes mostly originated from an area within a 1 km radius around  
684 the measurement site but some instances of pollution advection were seen to affect  
685 concentrations at the site. However many of the spatio-temporal differences in the observed  
686 mixing ratios were attributable to changes in emission sources and strengths combined with  
687 effects of meteorological conditions. The diurnal and seasonal dynamics of the boundary layer  
688 mixing height are a significant driver of changes in observed VOC concentrations at the site.

689 The biogenic component of isoprene emissions was modelled using the G95 algorithm and  
690 the calculated base emission rate closely matched previous published values for urban areas.  
691 Even in this central urban area with a temperate climate there is a detectable biogenic  
692 component to isoprene emissions. Because of the relative importance of isoprene in  
693 atmospheric chemistry, its inclusion in photochemical pollution models is essential.

694 Close agreement between the flux footprint contributions and the LAEI for benzene emissions,  
695 a compound which is thought to be accurately estimated in the inventory but associated with  
696 high measurement uncertainty, gives confidence in the PTR-MS measurements. Good  
697 agreement was also seen with methanol estimated from the NAEI, but other compounds were  
698 all greatly underestimated in the emissions inventory.

699 This study provides further evidence for the successful implementation of VOC flux  
700 measurements in heterogeneous urban landscapes when measurement sites fulfil basic eddy

701 covariance criteria. Further VOC flux observations are essential for the validation of “bottom-  
702 up” emission inventories, especially as the latter are widely used for regulatory and compliance  
703 purposes.

## 704 **5 Author contributions**

705 E. Nemitz and B. Langford planned the measurement campaign, A. Valach with the help of B.  
706 Langford and E. Nemitz made the measurements, A. Valach with the help of B. Langford  
707 processed the data and completed the analyses. C. N. Hewitt designed the study, obtained  
708 funding and supervised the work. A. Valach prepared the manuscript with support from all the  
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929 **Tables**930 Table 1. Summary of instrument operating parameters and average meteorological conditions during the  
931 measurements in central London, August – December 2012.

Parameter	Unit	Mean (range)
Normalised sensitivity ( $S_N$ ) <sup>a</sup>	ncps ppb <sup>-1</sup>	11.5 ( <i>m/z</i> 33), 13.3 ( <i>m/z</i> 45), 10 ( <i>m/z</i> 59), 4 ( <i>m/z</i> 69), 3.6 ( <i>m/z</i> 79), 2.5 ( <i>m/z</i> 93), 1.5 ( <i>m/z</i> 107)
Primary ion ( <i>m/z</i> 19)	Cps	8.31×10 <sup>6</sup> (6.14 ×10 <sup>6</sup> – 1.15×10 <sup>7</sup> )
Water cluster ( <i>m/z</i> 37)	Cps	1.92×10 <sup>5</sup> (9.15 ×10 <sup>4</sup> – 3.86×10 <sup>5</sup> )
	% of <i>m/z</i> 19	2.3 (1.5-3.4)
O <sub>2</sub> <sup>+</sup>	% of <i>m/z</i> 19	<1.45 (1.11-2.01)
Temperature <sup>b</sup>	°C	14.0 (-1.81-30.39)
Relative humidity	%	76 (50-97)
Pressure	mbar	1004.27 (968.71-1023.27)
Wind speed <sup>b</sup>	m s <sup>-1</sup>	3.35 (0.12-14.96)
Friction velocity ( $u_*$ ) <sup>b</sup>	m s <sup>-1</sup>	0.5 (0.01-1.50)
SD of vertical wind speed ( $\sigma_w$ ) <sup>b</sup>	m s <sup>-1</sup>	0.65 (0.15-1.62)

932 <sup>a</sup> $S_N$ : Normalised sensitivity as calculated using Taipale et al. (2008).933 <sup>b</sup>Derived from measurements from the CSAT3 sonic anemometer (Campbell Scientific).

934 Table 2. Summary of 25 min VOC fluxes and mixing ratios above central London during August-December 2012.

Compound ( <i>m/z</i> )	Methanol ( <i>m/z</i> 33)	Acetaldehyde ( <i>m/z</i> 45)	Acetone ( <i>m/z</i> 59)	Isoprene ( <i>m/z</i> 69)	Benzene ( <i>m/z</i> 79)	Toluene ( <i>m/z</i> 93)	C <sub>2</sub> -benzenes ( <i>m/z</i> 107)
<i>Fluxes (mg m<sup>-2</sup> h<sup>-1</sup>)</i>							
Lifetime (OH <sup>a</sup> )	12 d	8.8 h	53 d	1.4 h	9.4 d	1.9 d	5.9 h
N	2920	2811	2945	2119	1908	2315	2053
Min.	-2.91	-0.28	-1.74	-0.35	-0.64	-2.31	-3.27
1. quartile	0.12	0.06	0.10	0.02	0.002	0.08	0.04
Median	0.27	0.14	0.22	0.09	0.07	0.30	0.33
Mean	0.29	0.16	0.31	0.13	0.09	0.41	0.54
3. quartile	0.42	0.23	0.40	0.20	0.18	0.64	0.91
Max.	3.36	1.09	2.85	1.16	0.59	4.86	8.63
SD	0.25	0.15	0.34	0.16	0.15	0.53	0.86
Skew	0.86	1.27	2.08	1.18	0.32	1.75	2.33
Kurtosis	20.37	2.85	7.57	2.81	0.76	8.04	14.48
<i>Mixing ratios (ppb)</i>							
N	4834	4834	4834	4834	4834	4834	4834
Min.	5.73	<LoD (0.14)	<LoD (0.02)	<LoD (0.03)	<LoD (0.04)	<LoD (0.05)	<LoD (0.14)
1. quartile	6.82	0.59	<LoD (0.65)	<LoD (0.16)	<LoD (0.18)	<LoD (0.38)	<LoD (0.57)
Median	7.27	0.82	0.95	<LoD (0.22)	<LoD (0.24)	<LoD (0.54)	0.75
Mean	7.53	0.94	1.10	0.25	0.29	<LoD (0.65)	0.87
3. quartile	7.90	1.13	1.36	0.30	0.34	0.77	1.03
Max.	17.06	5.17	6.07	1.86	1.71	5.30	4.96
SD	1.12	0.53	0.66	0.14	0.19	0.45	0.50
Skew	2.21	2.14	1.65	1.97	2.80	3.07	2.79
Kurtosis	7.22	7.83	4.06	7.27	12.37	15.89	12.99
LoD <sup>b</sup>	0.96	0.45	0.66	0.25	0.28	0.66	0.71

935 <sup>a</sup>Atmospheric lifetimes with regard to OH for a 12 h daytime average OH concentration of 2.0 x 10<sup>6</sup> molecules  
936 cm<sup>-3</sup> (Atkinson, 2000).937 <sup>b</sup>LoD: Limit of detection calculated using Taipale et al. (2008).  
938

939 Table 3. Summary of site meteorology by month in central London during 2012.

Parameter	Data coverage (%)	Median stability ( $\zeta$ )	Wind speed ( $\text{m s}^{-1}$ )	Dominant wind direction (%)	Footprint <sup>a</sup> length (m)	Footprint width (m)
Aug	67	-0.0086	3.3	S (54)	2417	1355
Sep	83	-0.0154	3.2	W (48)	1285	880
Oct	89	-0.0006	3.5	S (29)	2624	1327
Nov	51	-0.0037	3.4	S (53)	2329	1156
Dec	40	0.0047	3.4	N (32)	1804	990

940 <sup>a</sup>Calculated two-dimensional description of the oval footprint according to the KM model. Length  
 941 parameter is the length between the point nearest to the sensor where the crosswind-integrated  
 942 footprint function reaches 1 % of its maximum value to the point where it drops below 1 % of the  
 943 maximum value.

944 **Figure captions**

945 **Figure 1.** Map of central London overlaid with the Ordinance Survey grid including the  
946 measurement site (KCL) at King's College (green point) with references to the geography of  
947 Greater London and Great Britain. Outlines of the areas that contribute the maximum ( $X_{max}$ ),  
948 as well as 75 %, 90 %, and 99 % to the flux footprint using overall median meteorological  
949 values are shown as black contour lines with their respective labels laid out according to the  
950 median wind direction.

951 **Figure 2a.** Average diurnal profiles in local time for selected VOC fluxes ( $\text{mg m}^{-2} \text{h}^{-1}$ )  
952 separated into all days, weekdays (red dashed line) and weekends (blue dotted line) with  
953 traffic density (vehicles  $\text{h}^{-1}$ ), detection limit (patterned area), and upper and lower confidence  
954 intervals (shaded area). Traffic density (with weekday and weekend) and boundary layer  
955 mixing height (for summer and winter) are shown in separate panels. Compounds are:  $m/z$   
956 33 (methanol),  $m/z$  45 (acetaldehyde),  $m/z$  59 (acetone/propanal),  $m/z$  69 (isoprene/furan),  
957  $m/z$  79 (benzene),  $m/z$  93 (toluene), and  $m/z$  107 ( $\text{C}_2$ -benzenes).

958 **Figure 2b.** Average diurnal profiles in local time for selected VOC mixing ratios (ppb)  
959 separated into all days, weekdays (red dashed line) and weekends (blue dotted line) with  
960 detection limit (dotted line), and upper and lower confidence intervals (shaded area). Traffic  
961 density (with weekday and weekend) and boundary layer mixing height (for summer and  
962 winter) are shown in separate panels. Compounds are:  $m/z$  33 (methanol),  $m/z$  45  
963 (acetaldehyde),  $m/z$  59 (acetone/propanal),  $m/z$  69 (isoprene/furan),  $m/z$  79 (benzene),  $m/z$   
964 93 (toluene), and  $m/z$  107 ( $\text{C}_2$ -benzenes). The mixing ratio axes start from zero apart from  
965 that of methanol, which begins at 6.4 ppb due to the high atmospheric background.

966 **Figure 3.** Examples, using isoprene, of averaged VOC fluxes (left) and mixing ratios (right)  
967 as a function of photosynthetically active radiation (PAR) ( $\mu\text{mol m}^{-2} \text{s}^{-1}$ ), temperature ( $^{\circ}\text{C}$ ),  
968 traffic density (vehicles  $\text{h}^{-1}$ ) and boundary layer mixing height (m) based on 25 min VOC  
969 means with linear or exponential regressions, formulae,  $R^2$ -values and detection limit  
970 (shaded area for fluxes and dashed line for mixing ratios).

971 **Figure 4.** Diurnal profiles by month with confidence intervals and bar charts showing hourly  
972 averages for the respective month and representative compound (top) fluxes ( $\text{mg m}^{-2} \text{h}^{-1}$ )  
973 ( $m/z$  45, 69 and 79) and (bottom) mixing ratios (ppb) ( $m/z$  59, 69 and 79). Letters (a-d)  
974 indicate statistically significant subgroups using Tukey's HSD (Honest Significant Difference)  
975 post hoc test.

976 **Figure 5a.** Time series of both measured (grey) and modelled (black) fluxes, as well as PAR  
977 and temperature measurements for August and September 2012.

978 **Figure 5b.** Correlation between modelled and measured isoprene fluxes ( $\text{mg m}^{-2} \text{h}^{-1}$ ) by  
979 wind direction using the G95 algorithm with temperature as a third variable, Ordinary Least  
980 Squares (OLS) regression lines, 99<sup>th</sup> confidence intervals, formulae, and  $R^2$ -value.

981 **Figure 6.** Selected scatterplots of representative correlations of VOC/VOC fluxes (top) and  
982 mixing ratio (bottom) with temperature as a third variable showing an example of bimodal,  
983 strong linear and medium linear correlations as commonly seen in the mixing ratio  
984 correlations with  $R^2$ -values, 1:1 line, 1:2 and 2:1 lines for the bimodal example in the bottom  
985 left panel.

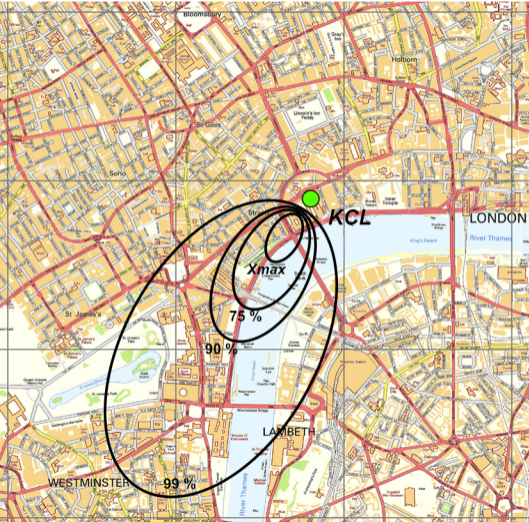
986 **Figure 7.** Top: 24h back trajectories from the NOAA HYSPLIT trajectory model during  
987 selected days in August 2012 corresponding to periods of low (left) and high (right)  
988 benzene/toluene concentration ratios. Daily release in 3 h intervals (10 m height) for 24 h  
989 prior. Bottom: Scatterplots showing benzene to toluene concentration ratios during the 9<sup>th</sup>  
990 August 2012 (left) and 12<sup>th</sup> August 2012 (right) with linear regression with 95<sup>th</sup> confidence  
991 interval, regression equation and coefficient ( $R^2$ ).

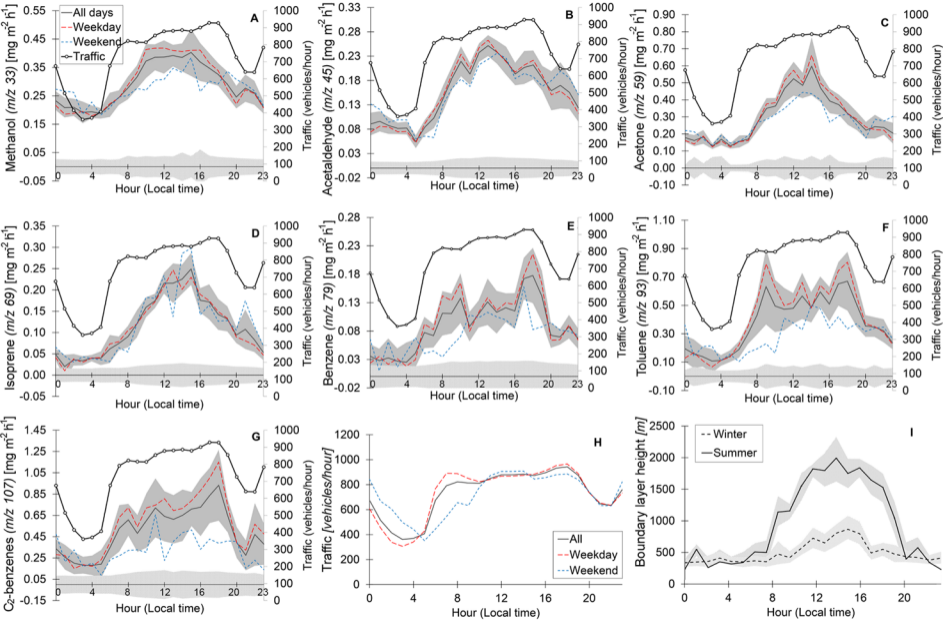
992 **Figure 8.** Scatterplots showing averaged flux and concentration regressions of isoprene and  
993 benzene as a function of CO<sub>2</sub> fluxes and concentrations based on 25 min VOC means with  
994 linear regressions, formulae,  $R^2$ -values and detection limit (shaded area for fluxes and  
995 dashed line for mixing ratios).

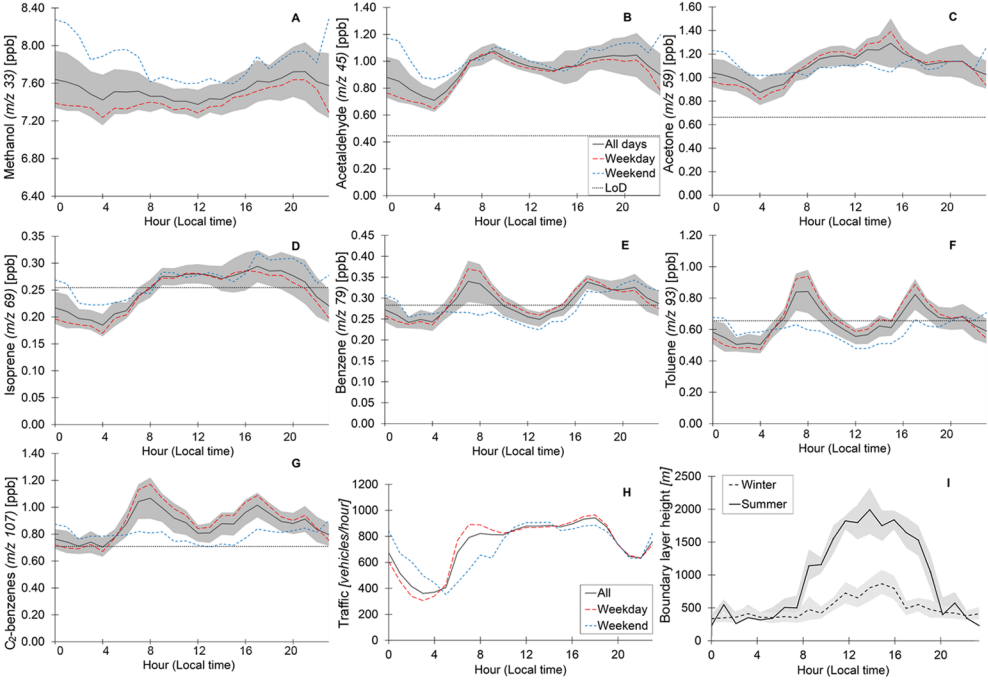
996 **Figure 9.** Polar Annulus and Polar plots for isoprene ( $m/z$  69) and benzene ( $m/z$  79) VOC  
997 fluxes (top) and mixing ratios (bottom) (colour scale) by time of day (top), wind speed  
998 (bottom) and wind direction.

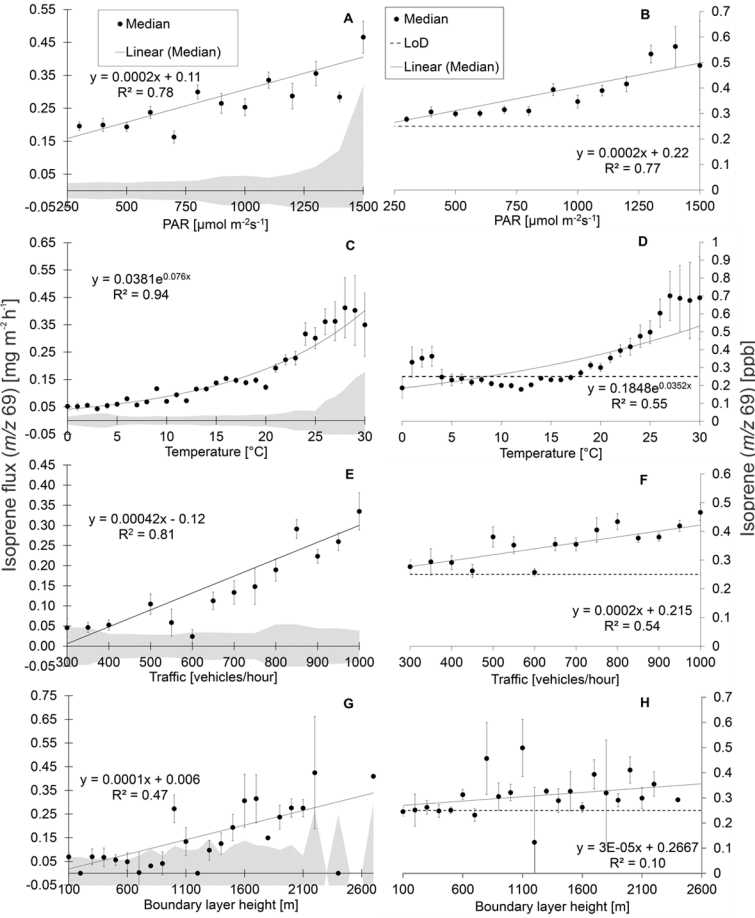
999 **Figure 10.** Bar chart showing scaled comparisons of LAEI and NAEI estimates against  
1000 measured fluxes in  $\text{t km}^{-2} \text{a}^{-1}$  for speciated VOCs with error bars.

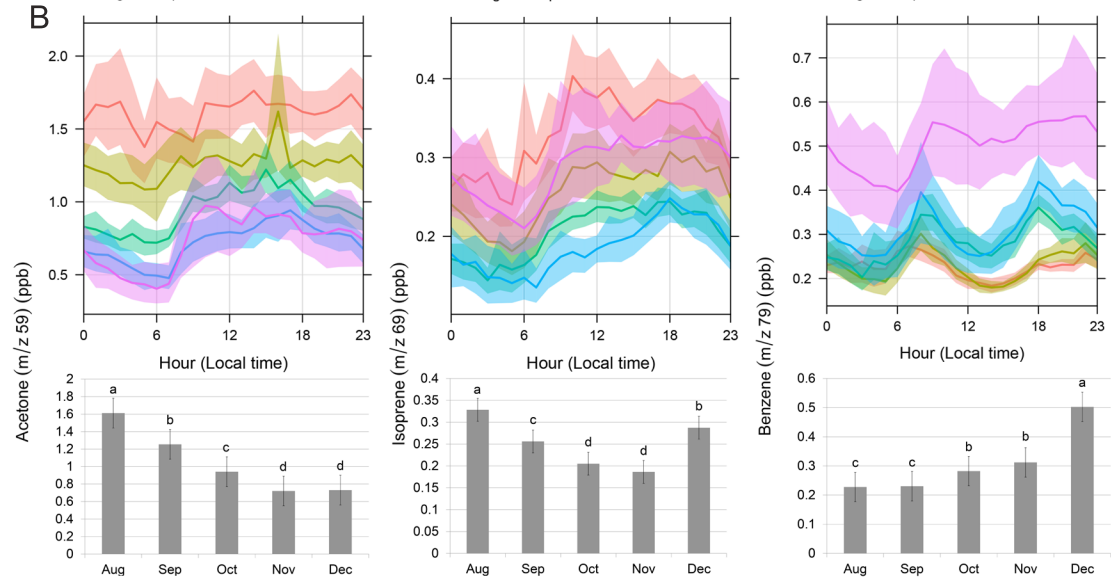
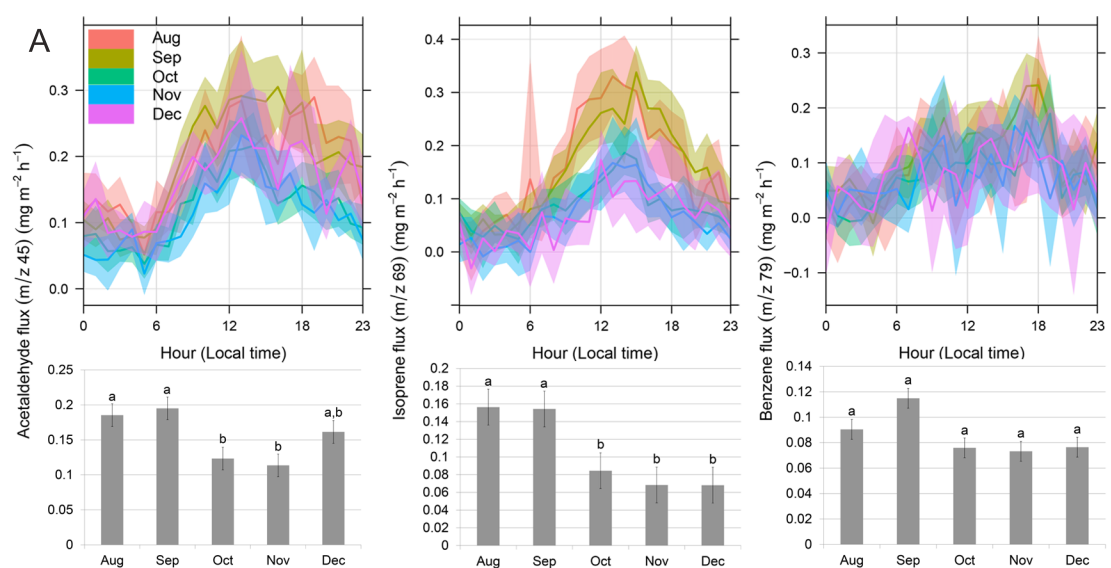


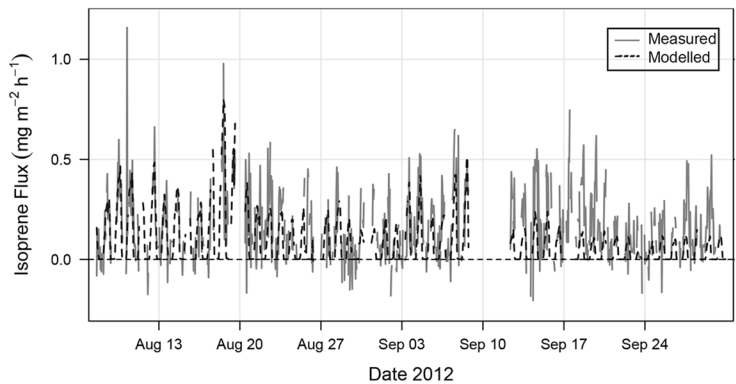
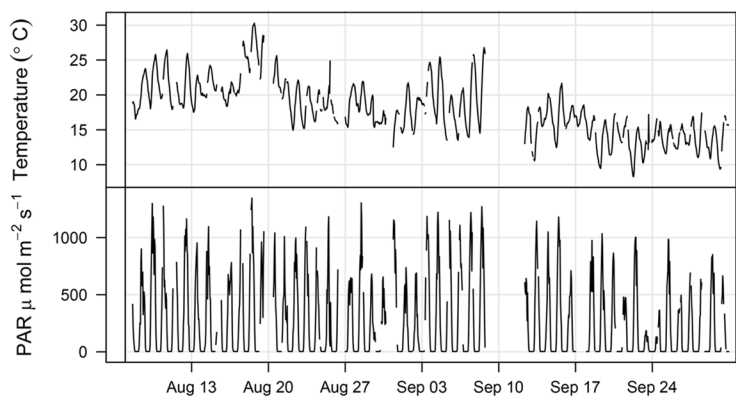












Temperature (°C)

10

15

20

25

30



0.0

0.5

1.0

NW

Mod=0.32[Meas]+0.025 R<sup>2</sup>=0.23

N

Mod=0.6[Meas]+0.02 R<sup>2</sup>=0.48

NE

Mod=0.48[Meas]+0.094 R<sup>2</sup>=0.4

0.8

0.6

0.4

0.2

0.0

W

Mod=0.23[Meas]+0.036 R<sup>2</sup>=0.13

E

Mod=0.7[Meas]+0.016 R<sup>2</sup>=0.53

0.8

0.6

0.4

0.2

0.0

SW

Mod=0.43[Meas]+0.037 R<sup>2</sup>=0.31

S

Mod=0.5[Meas]+0.076 R<sup>2</sup>=0.31

SE

Mod=0.83[Meas]-0.016 R<sup>2</sup>=0.81

0.8

0.6

0.4

0.2

0.0

0.0

0.5

1.0

0.0

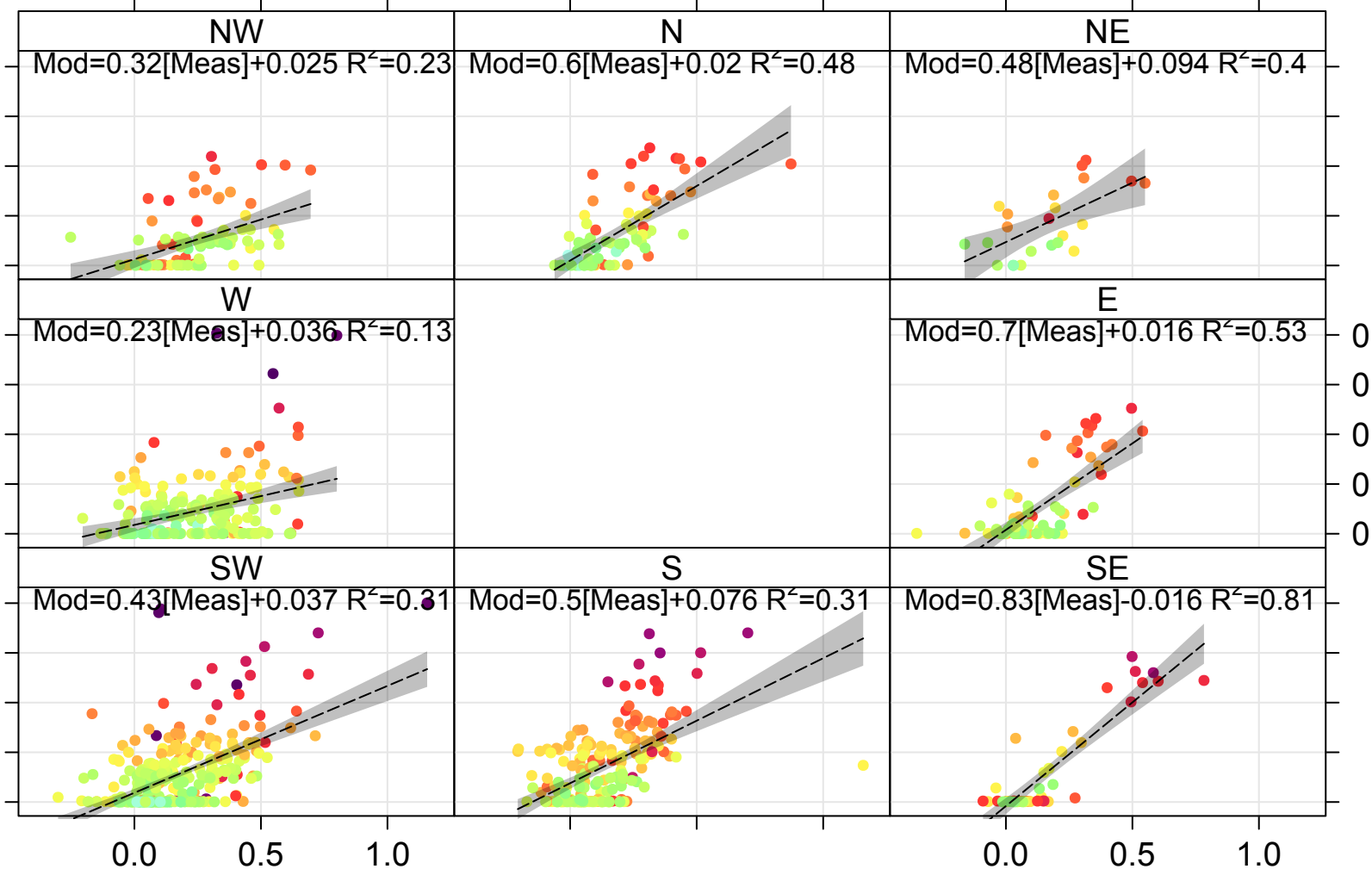
0.5

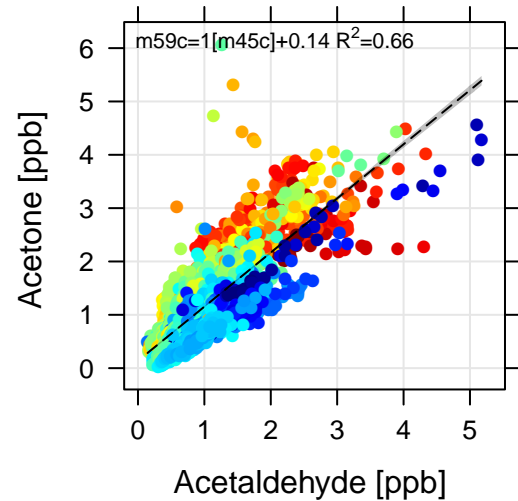
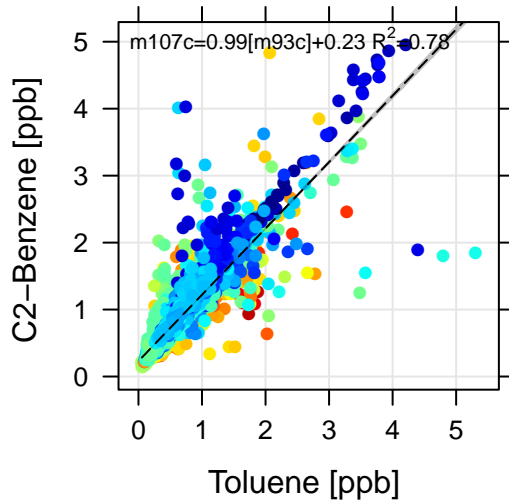
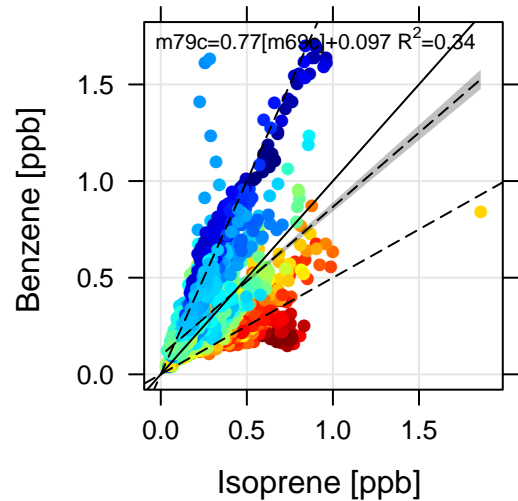
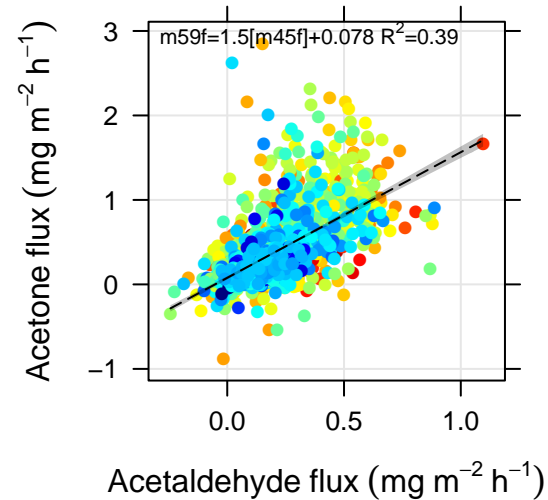
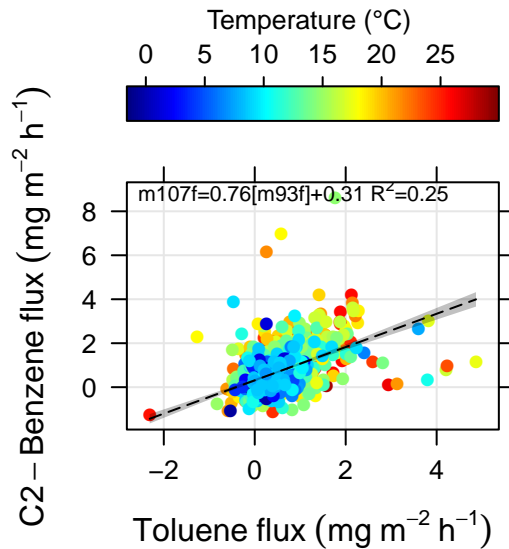
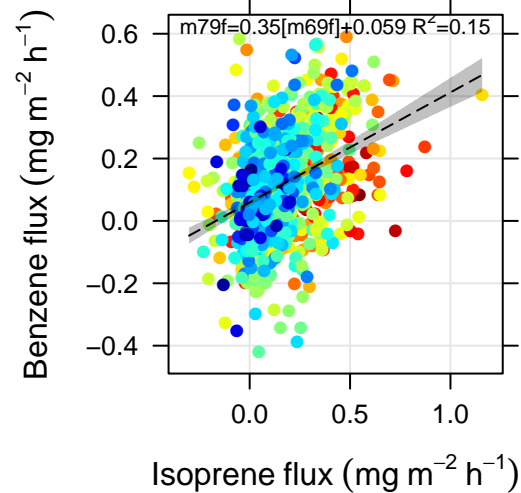
1.0

Modelled Flux (mg m<sup>-2</sup> h<sup>-1</sup>)

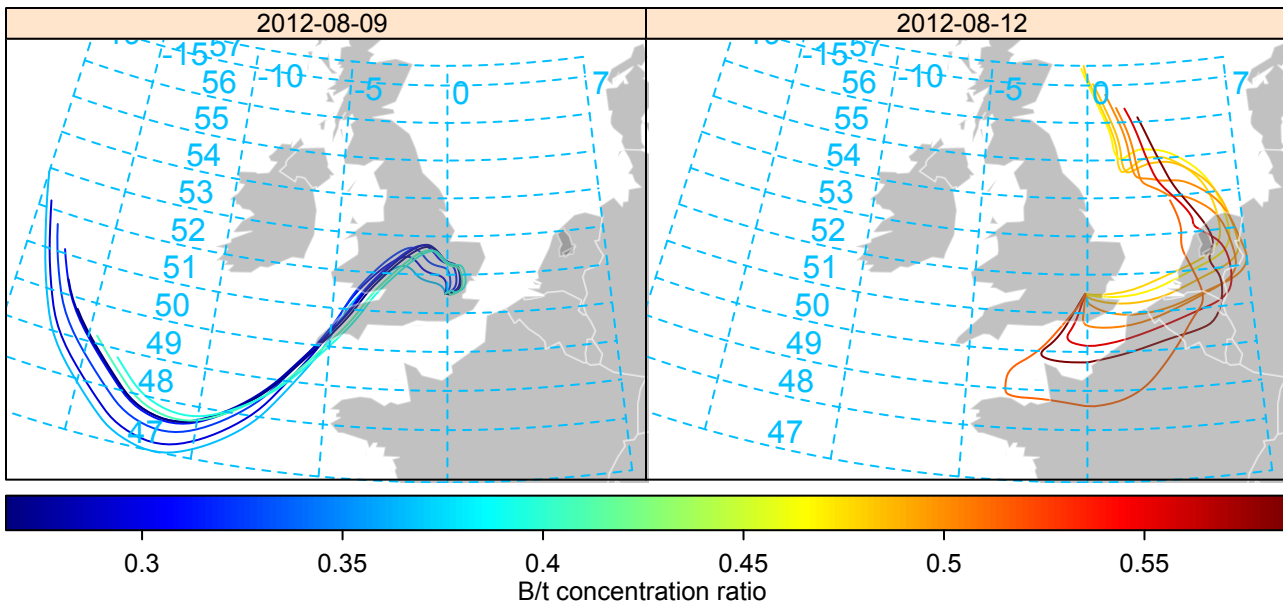
Measured Flux (mg m<sup>-2</sup> h<sup>-1</sup>)

Modelled Flux (mg m<sup>-2</sup> h<sup>-1</sup>)

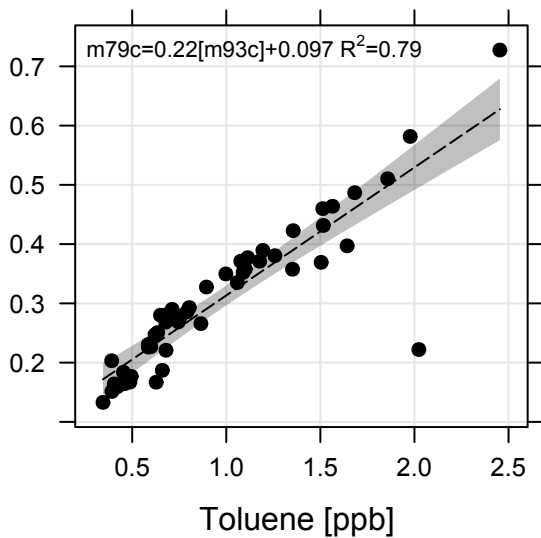




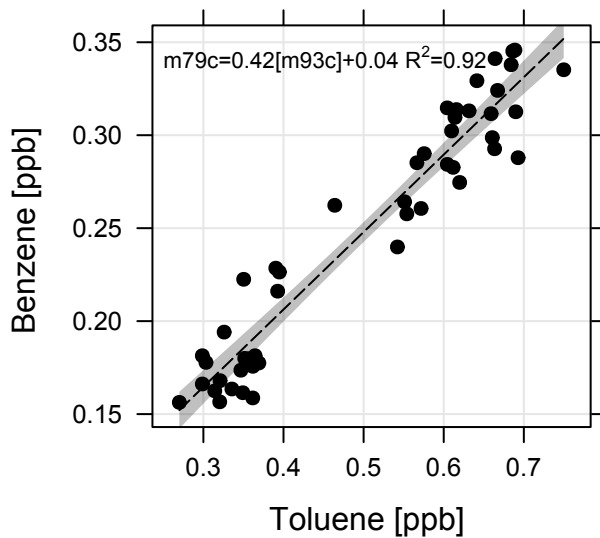


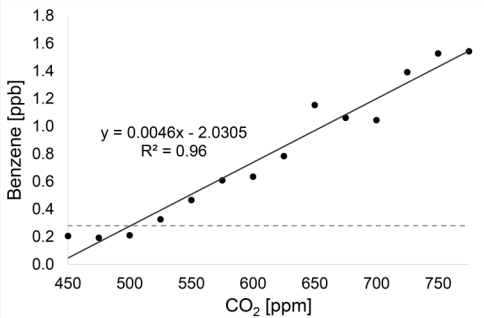
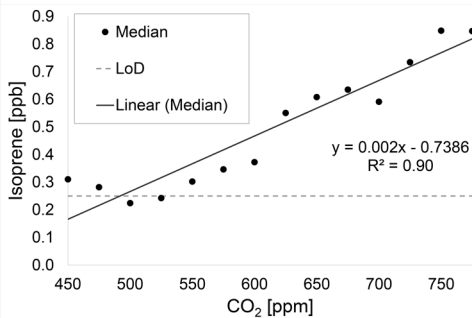
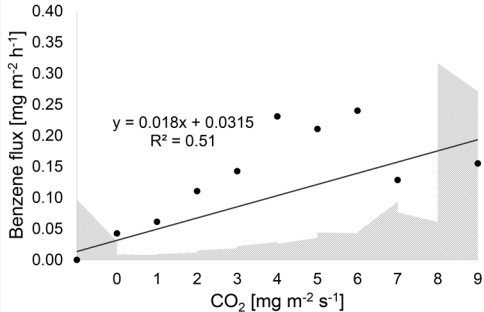
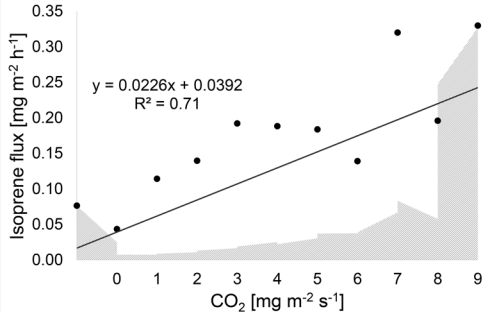


B/t ratio 9th August

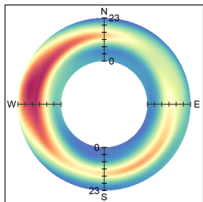


B/t ratio 12th August





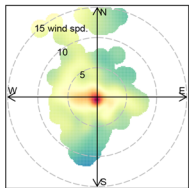
Isoprene (m/z 69)



0.05 0.1 0.15 0.2 0.25

Flux ( $\text{mg m}^{-2}\text{h}^{-1}$ )

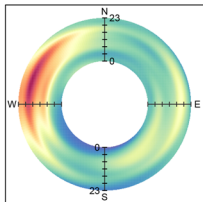
Isoprene (m/z 69)



0 0.1 0.2 0.3 0.4 &gt;0.5

Mixing ratio (ppb)

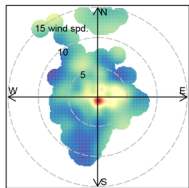
Benzene (m/z 79)



0.05 0.1 0.15 0.2 0.25

Flux ( $\text{mg m}^{-2}\text{h}^{-1}$ )

Benzene (m/z 79)



0.2 0.3 0.4 0.5 0.6

Mixing ratio (ppb)

