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1	Measurement of NO_x fluxes from a tall tower in			
2	central London, UK and comparison with			
3	emissions inventories.			
4				
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13				
14				
15	Abstract			
16	Direct measurements of NO _x concentration and flux were made from a tall tower in central			

17 London, UK as part of the Clean Air for London (ClearfLo) project. Fast time resolution (10

18 Hz) NO and NO₂ concentrations were measured and combined with fast vertical wind

19 measurements to provide top-down flux estimates using the eddy covariance technique. 20 Measured NO_x fluxes were usually positive and ranged from close to zero at night to 2000 – 8000 ng m₋₂ s₋₁ during the day. Peak fluxes were usually observed in the morning, coincident 21 22 with the maximum traffic flow. Measurements of the NO_x flux have been scaled and 23 compared to the UK National Atmospheric Emissions Inventory (NAEI) estimate of NO_x emission for the measurement footprint. The measurements are on average 80% higher than 24 the NAEI emission inventory for all of London. Observations made in westerly airflow (from 25 parts of London where traffic is a smaller fraction of the NO_x source) showed a better 26 27 agreement on average with the inventory. The observations suggest that the emissions 28 inventory is poorest at estimating NOx when traffic is the dominant source, in this case from 29 an Easterly direction from the BT tower. Agreement between the measurements and the 30 London Atmospheric Emissions Inventory (LAEI) are better, due to the more explicit 31 treatment of traffic flow by this more detailed inventory. The flux observations support previous tailpipe observations of higher NO_x emitted from the London vehicle diesel fleet 32 33 than is represented in the NAEI or predicted for several EURO emission control technologies. 34 Higher than anticipated vehicle NO_x is likely responsible for the significant discrepancies that 35 exist in London between observed NO_x and long-term NO_x projections.

36

37 Introduction

The oxides of nitrogen NO_x (defined as the sum of NO and NO_2), are emitted as a consequence of most combustion processes. The majority of NO_x is emitted as NO, which is rapidly oxidised to NO_2 upon reaction with ozone (O₃), with the reverse of this process being caused by the action of sunlight on NO_2 to form NO and O_3 . NO_2 is known to have significant direct health effects on humans. At high concentrations it causes inflammation of the airways and long-term exposure may affect lung function and enhance the response to

allergens^{1, 2}. In addition, NO_x contributes to the formation of O_3 and secondary particles 44 through a series of photochemical reactions³. As a result of this, NO₂ is included in a series 45 of air pollutants identified as part of the EU Air Quality Directive (AQD, 2008)⁴ which sets 46 limit values for hourly and annual mean exposure. It has been shown by measurements and 47 models that the annual mean limit value of 40 μ g m⁻³ continues to be exceeded in many urban 48 centres throughout the UK⁵, including London. Measures are in place to control the 49 50 emissions of nitrogen oxides and UK emissions are projected to decline by about 35 % between 2010 and 2020⁶. However, it is known that ambient NO₂ concentrations do not 51 52 respond linearly to reductions in the concentration of NO_x (e.g. Derwent et al., 1995⁷), mainly because of the chemical coupling of ozone (O_3) and NO_x under ambient conditions⁸. In 53 addition, changes in diesel emission control technology have led to increases in directly 54 emitted NO_2^9 . Trends in ambient concentrations of NO_x and NO_2 in the UK have generally 55 56 shown a decrease in concentration from 1996 to 2002, followed by a period of more stable concentrations from 2004–2012¹⁰. This is not in line with the expected decrease suggested 57 by the UK emission factors¹¹. 58

59 Air pollutant emission inventories provide input data for air pollution models, which in turn are used for predicting current and future air pollution. This is typically done using a 'bottom' 60 up' approach involving estimated emissions from different source sectors to produce yearly 61 emission estimates. However it is known that this method can contain large uncertainties, 62 with the errors propagating through into errors in air pollution models¹². Evaluation of 63 emission inventories can be carried out by comparing air quality model predictions (using 64 inputs from the inventory) to observed concentrations^{13, 14}, however this method does not 65 provide a direct comparison with the emission rate as it requires knowledge of other 66 67 paramters such as chemistry and meteorology. The eddy covariance technique provides a 68 direct measurement of the flux to the atmosphere of a particular pollutant, thus providing a

'top down' approach to quantifying emissions¹⁵. Flux measurements also provide 69 information on both spatial and temporal change in emissions from a calculated flux 70 footprint, giving insight into controls and sources. The majority of eddy covariance 71 measurements made to date have concentrated on fluxes of greenhouse gases (CO₂, CH₄ and 72 N₂O)^{16, 17} and volatile organic compounds (VOCs)¹⁸⁻²⁰, largely from biogenic sources. Some 73 eddy covariance NO_x flux measurements have been made and have typically focused on 74 emissions from soils²¹, forests²²⁻²⁴ or snow^{25, 26}. Recently however, it has been shown that 75 this method can be extended to the urban canopy for CO_2^{27-29} and $VOCs^{30-32}$, with one study 76 of urban NO_x^{33} . 77

In this study, we use the eddy covariance technique to directly measure the flux of NO and NO₂ from a tall tower (190 m) in central London as part of the Clean Air for London (ClearfLo) project³⁴. The results are compared to local traffic flow and a flux footprint is calculated to allow comparison with two emission inventories, one for the whole of the UK and one specific to London.

83

84

85 Experimental

86 *Measurement site*

Measurements were made during June – August 2012 and March – April 2013 from the top of the BT tower, a 190m tall telecommunications tower situated in central London, UK ($51^{\circ}31'17.4''N 0^{\circ}8'20.04''W$). Mean building height is 8.8 ± 3.0 m within 1 - 10 km of the tower and 5.6 ± 1.8 m for suburban London beyond this^{30, 35}. The area surrounding the tower is dominated by roads and commercial / residential buildings, but also includes some urban parkland and pervious ground. A map of the location of the tower within London is shown in the supplementary information (Figure S1). The gas inlet and ultrasonic anemometer were attached to a mast which extended ~ 3 m above the top of the tower. Air was pumped down a ~40 m long Teflon tube (1/2" OD) at a flow rate of ~ 30 L min⁻¹ to the gas instruments which were housed in a room inside the tower.

97 The most prevalent wind direction during the summer 2012 measurement period was the 98 SW sector (~50 % of the time), with other wind sectors split approximately equally. Wind speed was 6.7 ms⁻¹ on average, with the highest wind speeds measured when the wind was 99 100 from a NW direction. Average temperature was 15.1 ± 4.3 °C. During the March and April 2013 measurement period, the most prevalent wind direction was between 0 - 90° (50 %), 101 again with other directions split approximately equally. Wind speed was higher than summer 102 2012, being 8.8 m s⁻¹ on average with the highest wind speed when the wind was from the 103 104 SW direction. As expected, average temperature was lower than the summer 2012 period, being $9.7 \pm 2.4^{\circ}$ C. 105

106

107 *NO_x measurements*

108 Measurements of NO were made using an Ecophysics 780TR instrument, which uses the chemiluminescence technique^{36, 37}. NO₂ was quantified in a second identical NO instrument 109 110 by initial photolytic conversion to NO using blue light LED diodes centred at 395 nm. The 111 395 nm wavelength has a specific affinity for NO₂ photolytic conversion to NO, giving high analyte selectivity within the channel³⁸ and there is a low probability of other species such as 112 nitrous acid (HONO) being photolysed. The diode based converter also has a very low 113 114 residence time for the air sample (< 0.1s) which allows 10 Hz measurements of NO₂ to be 115 made. The NO instruments were calibrated every 36 hours by addition of a known amount of 116 NO to the sample line, made by diluting a gas standard (5 ppm NO in N_2 , BOC – traceable to NPL scale) in NO_x free air (Ecophysics PAG003). The conversion efficiency of the NO_2 117 converter was also measured during each calibration by gas phase titration of the known NO 118

119 upon addition of O_3 , with typical conversion efficiencies being 30 - 35%. It is estimated that 120 the total error (including accuracy and precision) is around 10 % for NO and 15 % for NO₂ at 121 10 ppbv.

122

123 Meteorology measurements

Fast (20 Hz), 3 dimensional wind vectors and sonic temperature were measured from next to the sample line inlet by a Gill Instruments R3-50 ultrasonic anemometer. The data was logged, along with that from the NO_x instrument, using a custom National Instruments LabViewTM program. The boundary layer height was measured using a HALO Photonic Doppler LiDAR instrument³⁹.

129

130 Flux calculations and uncertainties

131 NO and NO₂ fluxes (F_{NO} and F_{NO_2}) were calculated using eq. 1 and 2 below.

132
$$F_{NO} = \frac{\overline{w'C'_{NO}}}{S_{NO}V_{mol}} \qquad (1)$$

133

134
$$F_{NO2} = \frac{1}{\alpha V_{mol}} \left\{ \frac{\overline{w'C'_{NO2}}}{S_{NO2}} - \frac{\overline{w'C'_{NO}}}{S_{NO}} \right\}$$
 (2)

135

136 C_i is the number of instrument counts (in Hz) and S_i is the associated instrument sensitivity 137 (in Hz ppb⁻¹) for species *i* (*NO and NO*₂). V_{mol} is the molar volume (calculated for each 138 individual point), α is the photolytic conversion efficiency of NO₂ to NO and *w* is the vertical 139 wind component measured by the ultrasonic anemometer. A "prime" symbol represents an 140 instantaneous deviation from the mean and a horizontal bar denotes the covariance of 2 141 scalars. Processed data were filtered using a three-step quality assurance algorithm whereby datawere deemed of satisfactory quality if:

144 The level of turbulence was sufficient, i.e. locally-derived friction velocity $u_* \ge 0.2 \text{ m s}^{-1}$ 145 (<5% of the data is rejected due to this parameter)

146 The number of spikes in w, NO and NO_2 did not exceed 1% of total in each half-hourly 147 averaging period.

The stationarity test described by Foken et al.^{40, 41}, which requires the flux for the complete averaging interval (here 30 min) to be within 30 % of the fluxes calculated for the subintervals (6×5 minutes), was satisfied.

Total measurement uncertainty, i.e. the sum of total random and systematic uncertainties, was estimated using the 24-hour differencing method⁴² whose assumption is that the difference between pairs of observations taken exactly 24 hours apart under similar meteorological conditions (air temperature, wind speed and direction) is mainly attributable to stochastic factors. Using multiple pairs of observations, the standard deviation of the random error can be calculated from eq. (3).

157
$$\sigma = \frac{\sigma(x_{1,t} - x_{2,t})}{\sqrt{2}}$$
 (3)

158

159 The environmental conditions were deemed similar if:

160 Air temperatures diverged by less than 3 °C.

161 Wind speed diverged by less than 2 m s^{-1} .

162 Wind directions originated from the same quadrant.

163

164 Causes of systematic uncertainties are varied and include calibration procedures, 165 instrumentation limitations or data processing artefacts. Unlike random uncertainties, 166 systematic errors can be minimized by careful data processing and correction. 167 Successive calibration events were linearly interpolated over time cancelling out errors due 168 to calibration drifts provided that the drift is linear over time.

169 To estimate potential turbulence attenuation in the sampling line, which can lead to underestimation of the actual flux, fluxes of CO₂ measured using by a Picarro G2301-f 170 171 sampling off the same line as the NO and NO₂ analysers were compared with fluxes measured by a Licor 7500 open-path analyser mounted near the ultrasonic anemometer. The 172 underlying assumption is that turbulence attenuation and molecular interactions with the 173 174 sampling tube are comparable for CO₂, NO and NO₂ molecules. Rather than correct for 175 attenuation, this systematic uncertainty was added to the estimated stochastic component and 176 presented as confidence interval in what follows.

177

178 Flux footprint

179 In order to carry out meaningful interpretation of the data, it is necessary to calculate the flux footprint of the measurement. It is not possible to get footprint models to fully account 180 for the spatial variability of building heights, topography and surface heat flux from an urban 181 environment. In this case, the Kormann and Meixner (2001)⁴³ footprint model (K-M model) 182 183 was applied, which accounts for non-neutral stratification but assumes homogeneous 184 surfaces. The aerodynamic roughness length for momentum was assumed to be 1 m as used in previous BT Tower flux studies³⁵. The sample height for the BT Tower was 190 m. The K 185 - M model was used to estimate the flux footprint on a half hourly time base. A Microsoft 186 187 Excel tool (based on the K - M model) calculates the distance from the measurement point at 188 which a set percentage of the measured flux is emitted from. Figure S4 in the supplementary information shows a histogram of the calculated footprints for 50 %, 70 % and 90 % of the 189 flux for the measurement period. The analysis here uses the footprint where 90 % of the flux 190

191 is predicted to originate from, which shows a range of 150 m - 19980 m with a median of 192 4695 m.

193

194 **Results and discussion**

Measurements of the NO_x flux were made during two time periods, June – August 2012 (36 days) and March – April 2013 (28 days). Downtime was due mainly to instrument failure of both the fast NO_x instrument and 3-D sonic anemometer, as well as a failure in the sample pump. Despite this, data coverage on the days when measurements was taken 61 %), meaning the dataset provides a unique opportunity to examine the diurnal and seasonal behavior of NO_x fluxes from central London.

201 The full time series of data is shown in the supplementary information Figure S2, with NO_x concentrations averaged to the 30 minute flux averaging time. Typically NO concentrations 202 vary from close to zero at night to a maximum of $10 - 100 \ \mu g \ m^{-3}$ during the day, whereas 203 NO₂ ranges from $5 - 80 \mu \text{g m}^{-3}$. Also shown in Figure S2 is the time series of NO and NO₂ 204 205 from an urban background site in at North Kensington, London, which is approximately 5 km west of the BT tower⁴⁴. These data show similar trend to the BT tower for most of the time, 206 207 although at generally higher levels. A regression analysis of the two datasets (BT tower and 208 North Kensington, shown in supplementary information Figure S3), shows North Kensington data being on average 10 % higher for NO and 6 % higher for total NO_x (R^2 of 0.65 and 0.58 209 respectively). This result gives confidence that, at least for total NO_x , the BT tower site is 210 211 representative of the wider London area.

Random uncertainties (1 σ) obtained by 24-hour differencing were 441 ng m⁻² s⁻¹ for F_{NO}, 475 ng m⁻² s⁻¹ for F_{NO₂} and 510 ng m⁻² s⁻¹ for F_{NO_x} (F_{NO} + F_{NO₂}); residual systematic uncertainties, were estimated at 15% of the measured flux. Maximum NO_x fluxes are measured during the daytime, with values from 2000 ± 741 to 5000 ± 1191 ng m⁻² s⁻¹ for NO

and 2000 ± 775 to 12000 ± 2275 ng m⁻² s⁻¹ for NO₂. Measured fluxes are usually positive, 216 217 demonstrating, as expected, that NO_x emission dominates over deposition in this urban 218 environment and that it is likely to be dominated by anthropogenic emissions. NO_x can be lost to the surface by dry deposition⁴⁵, and assuming a deposition velocity of 0.1 cm⁻¹ and a 219 NO_x concentration of 50 µg m⁻³, then the downward flux can be estimated to be in the region 220 of 100 ng $m^{-2} s^{-1}$, which is more than an order of magnitude smaller than the observed values. 221 222 NO and NO₂ fluxes show a distinct diurnal profile. NO flux is close to zero at night (although still positive), with a rise starting at 05:00 to a peak of 1800 - 1900 ng m⁻² s⁻¹ between 08:00 223 224 and 12:00. The NO flux then usually starts to decrease throughout the rest of the day and into the night, reaching the nighttime value of 100 - 200 ng m^{-2} s⁻¹ at around 20:00. NO₂ flux also 225 typically shows a diurnal profile with 500 - 1000 ng m^{-2} s⁻¹ measured at night followed by a 226 rise to 2200 - 2300 ng m⁻² s⁻¹ from 05:00 until 12:00, with levels then remaining constant 227 until around 16:00. There follows a steady decrease in NO₂ flux throughout the rest of the 228 day and into the night, with levels reaching around 1200 ng m^{-2} s⁻¹ at midnight. 229

230 Very few direct flux measurements of NO and NO₂ have been made in an urban 231 environment, however the values measured in this study are comparable to a study in the 232 urban area of Norfolk, Virginia, USA, which reported total NO_x fluxes in the range 5000 -8000 ng m⁻² s^{-1 33}. Direct measurements of NO_x fluxes have been made previously over 233 234 forested and snow pack environments, with the measured fluxes still positive, but typically an order of magnitude smaller than measured here^{22, 24, 25}. Because of the close coupling of NO 235 and NO₂, it is the sum NO_x that is typically reported in emission inventories, and so the rest 236 237 of this work will concentrate on measurements of total NO_x. This also allows us to discount the chemistry associated with the inter-conversion of NO and NO₂, which can happen on a 238 239 very fast timescale., Total NO_x is likely to be conserved between emission and sampling on 240 the BT tower, as formation of NO_x reservoir species such as PAN and HNO₃ takes place on a

241 much longer timescale than the time between emission from street level and sampling at the 242 tower (estimated as 3 - 8 minutes).

Analysis of the wind sector dependence of the flux can help to identify the sources of the 243 244 species in question. Figure 1 shows bivariate polar plots with the joint flux footprint-wind direction of the NO_x flux, created using the Openair package⁴⁶. The flux footprint used was 245 calculated using the method described above. Two plots are shown to reflect daytime (05:00 246 247 -19:00) and night time fluxes. During the daytime, there are clearly higher fluxes measured when the calculated footprint is smaller, in particular when the wind is from an E / NE 248 249 direction from the tower. Fluxes then get smaller as the footprint gets larger in all directions. 250 This is a reflection of the reduced traffic density (and hence traffic emissions), further away 251 from central London. At night the fluxes are lower in all directions and for all footprints (as 252 expected), however there is much less of a reduction in flux as the footprint gets larger. An explanation for this behavior is likely that traffic emissions are much less important for the 253 254 total night time NO_x emission, with the majority of the emissions from commercial, industrial 255 and domestic combustion. Hence there is more homogeneity over London during the night 256 compared to the daytime. There are still greater fluxes measured when the wind was from the NE – SE sector, which is probably due to the area to the east of the tower being more 257 258 urban in nature than that to the west.

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260



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- 262

Figure 1. Wind sector dependence of the NO_x flux for all data averaged during (a) daytime (05:00 - 19:00) and (b) nighttime (20:00 - 04:00). The radial axis shows the calculated flux footprint in metres for each measurement.

268 Concentrations of a given pollutant in the atmosphere are largely dependent on its emission 269 rate, meteorology and chemical processing. It is useful to consider diurnal profiles in all these 270 quantities because it can help understand the processes leading to what is observed. For 271 diurnal averages, systematic uncertainties greatly outweigh random uncertainties which decrease as $1/\sqrt{n}$, with *n* the sample size. Average diurnal cycles have been calculated for 272 the entire measurement period, for NO_x flux, average traffic volume at 20 traffic counting 273 sites within the flux footprint of the site, boundary layer height and NO_x concentration and 274 this data is shown in Figure 2 (all times local time). Standard deviations of the average 275 276 diurnals are also shown, demonstrating the relatively small day to day variability of the 277 measurements. The traffic data used can be thought of as a proxy for total traffic flow across 278 the entire flux footprint area and a map of the location of the traffic counting sites used is 279 shown in the supplementary information (Figure S5). Data from each day is binned into 280 hourly time periods (UTC = local time -1 hour) and averaged, with the time stamp being the 281 mid-time of the averaging period. NO_x flux shows a diurnal cycle with positive fluxes seen 282 throughout the day. From 00:00 to 04:00 fluxes are slightly decreasing from 1400 ± 210 ng $m^{-2} s^{-1}$ to $450 \pm 67 ng m^{-2} s^{-1}$, with a rise starting at around 04:30, consistent with the onset of 283 the morning rush hour in London (at 05:30 local time). There follows a steady increase in the 284 NO_x flux to around 4000 \pm 600 ng m⁻² s⁻¹ at 10:00, levels that remain until 17:00 (with a 285 286 slight second peak at 16:00). This is broadly similar to the average traffic count data, providing more evidence that the majority of the NO_x emissions sampled at the BT tower are 287 from road traffic emissions. There then follows a steady decrease in the NO_x flux throughout 288 the rest of the day, to around 1200 ± 180 ng m⁻² s⁻¹ at 00:00. This is again broadly in line with 289 the traffic flow. NOx concentrations are reasonably stable at ~ 18 - 20 $\mu g~m^{\text{-3}}$ throughout the 290 night, followed by a rapid rise starting at 04:30 (at similar time to the rise in NO_x flux). This 291

292 rapid rise is due to a combination of the increase in fluxes, and the fact that the boundary 293 layer height does not increase until around 06:30. Once the boundary layer starts to grow (from ~ 300 m at 08:00 to 1700 m at 12:00), the rise in NO_x concentrations is less rapid, and 294 in fact they start to fall after a peak of 22 μ g m⁻³ at 08:00 until 16:00. This is likely due to 295 dilution effects caused by the increasing height of the boundary layer meaning the NO_x is 296 297 emitted into a larger volume. After 15:30, the NO_x concentrations start to rise again, despite a 298 decrease in flux. This is again likely due to the meteorology, with a decreasing boundary 299 layer height into the night.

300 Also, plotted in Figure 2 is the weekday and weekend diurnal average for the data. During 301 the day, traffic counts are on average lower during the weekend, particularly during the 302 morning where the difference is up to 50 %. This is reflected in the NO_x flux data, although it 303 does not show as pronounced a difference between weekend and weekday. This is potentially 304 due to the type of traffic at the weekend, which is likely to be predominantly buses and larger 305 vehicles (mainly powered by diesel engines), whereas during the week, private cars and taxis 306 maybe more prevalent. During the night, traffic levels are actually higher at the weekend 307 than during the day, also likely to be a result of public transport and the large nighttime 308 weekend economy of London. This is also reflected in the NO_x flux measurements showing 309 higher values from midnight to 06:30 for weekends compared to weekdays.





Figure 2. Average diurnal profiles for 36 days of data during Jun – Aug 2012 and 28 days during March and April 2013. Data shown are average traffic count (see text for further details), NO_x flux, boundary layer depth and NO_x mass mixing ratio. All times local time with the time stamp the mid-point of an hour averaging period. Error bars reflect the 95 % confidence intervals in the mean of the different measurements used to calculate the diurnal average. The red dotted line shows weekday data and the blue dashed line show weekend data.

318 The flux data was binned into 4 different regimes according to the calculated footprint (0 -2.5, > 2.5 - 5, > 5 - 10 and > 10 - 20 km radial distance from the BT tower) and average 319 diurnal profiles for each are plotted in Figure 3. The shaded regions represent the 95 % 320 confidence of the day to day variability of the flux measurements. All regimes show a 321 322 similar diurnal profile, with the flux starting to rise at around 04:30, with a peak between 10:00 and 14:00. The highest fluxes are seen in the two smallest footprint regimes, with both 323 showing similar values during daytime of around 4500 ± 675 ng m⁻² s⁻¹. The 5 - 10 km 324 regime shows lower daytime peak fluxes of 3200 ± 480 ng m⁻² s⁻¹, with the 10 - 20 km 325 regime lower still, with a peak of 2950 \pm 442 ng m⁻² s⁻¹ at 10:00 and then a decline 326 327 throughout the day. All 4 regimes show similar NO_x fluxes at night of around 1000 ± 150 ng $m^{-2} s^{-1}$, the exception being the 0 - 2.5 km, which does exhibit some elevated flux levels up to 328 1500 ng m⁻² s⁻¹, and appears to start to rise slightly earlier than the other regimes. All this 329 330 behavior is consistent with traffic emissions being the dominant source of NO_x, especially in 331 central London. It is expected that traffic volume will be higher closer to central London and 332 this is shown by the average traffic counts also plotted in the different footprint bins in figure 4. As a result of this, the smaller footprint regimes from the BT tower show the largest 333 daytime fluxes. At night, it is likely that a smaller proportion of the NO_x will come from 334 335 traffic sources, meaning the measured flux will be similar in all flux regimes out to 20 km 336 from the measurements site.

337



338

Figure 3. Average diurnal profiles for NO_x flux in 4 different footprint regimes (red trace). The error bars reflect the 95 % confidence intervals in the mean of the different measurements used to calculate the diurnal average. All times local time with the time stamp the mid-point of an hourly averaging period. Also shown is the average traffic flow at 6 sites within each of the individual footprint areas (blue trace).

344 *Emissions inventories*

In order to put the measured data in some context, a comparison has been carried out against inventories of NO_x emissions for London. The UK National Atmospheric Emissions Inventory (NAEI) shows official annual, spatially disaggregated 1 x 1 km gridded emission maps for a wide range of atmospheric pollutants, including NO_x . A detailed description on how the emissions maps are produced is given in Bush et al. 2008⁴⁷. Briefly, annual emission estimates are generated from 11 source sectors, according to those laid out by the United 351 Nations Economic Commission for Europe (UNECE). For each sector a national total 352 emission estimate is produced from a combination of reported emissions and estimates based on modelling. The UK National Atmospheric Emission Inventory (NAEI) gives an estimate 353 of the NO_x emissions in 1 km² grids over the UK, including a breakdown of the different 354 sources. The NAEI estimate for NO_x emissions for London is shown in the supplementary 355 information (Figure S6). The map is centered on the BT tower and features of London 356 357 characterized by large NO_x emissions can clearly be seen (e.g. Heathrow airport to the West 358 and the M25 orbital motorway circling the city). Four maps are shown, with the contribution 359 from 3 of the most important sectors (road transport, domestic, industrial and commerical 360 combustion and other transport (rail and shipping), as well as the total emissions. Also shown 361 on the maps are 5 km and 10 km radius circles from the tower, indicative of the flux footprint 362 bins described above. It suggests that around 65 % of NO_x emissions from central London are 363 from road and other transport, with the majority of the remainder from commercial, domestic and industrial combustion. 364

365 The London Atmospheric Emissions Inventory (LAEI) provides emissions estimates of 9 air pollutants (including NO_x), on a 20 m x 20 m grid square scale. The inventory reflects the 366 geography of the roads in London, enabling an accurate assessment of population exposure 367 368 and health impacts. Two versions of the LAEI are used in this study. The standard LAEI 369 (LAEI base) is the 2012 inventory based on methods set out in the Greater London Authority datastore⁴⁸, but updated for 2012 emission data. Also, we use an enhanced version of the 370 371 LAEI, which uses measured roadside emissions based on extensive vehicle emission remote sensing⁴⁹. Both emissions inventories discussed are purely annual averages with no seasonal 372 373 or finer temporal detail.

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375



377 *Comparison with measurements*

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Figure 4. Comparison of the averaged measured fluxes, scaled to give an annual emission rate, with the estimate of the National Atmospheric Emission Inventory (NAEI) and two versions of the London Atmospheric Emissions Inventory (LAEI) – see text for details. The different colours in the columns represent the estimates from different source sectors.

384

385

Figure 4 shows estimated emissions of NO_x taken from the NAEI and LAEI for 2.5 km, 5 km and 10 km radial distance from the tower, along with the estimates for sections in easterly $(30 - 150^{\circ})$ and westerly $(210 - 330^{\circ})$ directions and the source sector estimate divided into road transport, commercial and residential combustion and other transport (which is mainly rail in London). For the NAEI data for a 20 km radial distance is also plotted however this data is not available for the LAEI. Also plotted is the averaged measured NO_x flux for the different footprint regimes, also divided into periods of easterly and westerly wind directions and scaled to give a yearly emission rate.

394 The measurements are seen to be significantly higher than the NAEI (outside the estimated 395 experimental flux systematic error of 15 %) under all regimes. The agreement between the 396 measurement and the inventory tends to get worse for the larger footprint regimes, with the 397 measurement being 2.2 times higher than the inventory for the 10 - 20 km regime, and only 398 1.6 times higher for the 0 - 2.5 km regime. There is much more scope for error when 399 considering a comparison between larger flux footprints and the inventory as the further the 400 air has travelled, the more different emission inventory grid squares it could have passed 401 over, making a comparison with the inventory more difficult. In general the agreement is 402 better for the westerly flow conditions, with the measurement being 1.36 and 1.38 times 403 higher than the inventory for the 2.5 km and 5 km footprints respectively, whereas for the 404 easterly flow, the agreement is worse (1.6 and 1.9 times higher for 2.5 km and 5 km). The 405 difference in source sector between the 2.5 km and 5 km radius is small. Road transport 406 dominates (62 % and 60% for 2.5 km and 5 km respectively), with the remainder from commercial and domestic combustion (29 % and 27 %) and other transport (4 % and 10 %). 407 408 There is a lower contribution from road transport for the westerly flow conditions (48 % for 409 both 2.5 km and 5 km radius), giving a potential reason for the better agreement here. It is likely that road transport is the most poorly constrained part of the NAEI, and hence when 410 411 this is less important to the total emission rate, the agreement with the measurement is better.

412 For the base LAEI, the comparison shows a much closer agreement of the measurements 413 with the inventory compared to that with the NAEI discussed above. The inventory is within the measurement error for the average of all wind directions, with the measurement 1.03 and 414 415 1.1 times higher than the inventory in the 2.5 km and 5 km regimes respectively. The 416 agreement is similarly good in westerly flow, and although in Easterly flow the measurement 417 is now 1.07 times lower for the 2.5 km footprint and 1.03 times higher for the 5 km footprint, 418 these are still well within measurement error. For the 10 km footprint, the LAEI falls outside 419 the systematic error of the measurements for all the data separated into easterly and westerly flow regimes, the measurements being 1.16 and 1.48 times higher than the inventory for 420 421 westerly and easterly flow respectively. A comparison of the measurements to the enhanced 422 LAEI (which has generally increased road transport NO_x emissions), shows the 423 measurements being slightly lower than the inventory for data from the 2.5 km and 5 km flux 424 footprints, although again these is still within the systematic error of the measurements for all 425 the data and the westerly flow. It is in the easterly flow conditions where the measurements 426 are now significantly lower than the inventory, with the underestimation of 20 % and 17 % 427 for the 2.5 km and 5 km regimes falling outside the flux measurement error. For the 10 km 428 flux footprint regime, the enhanced LAEI brings the emission estimates into much better 429 agreement with the measurements than the base case, with the data from both easterly and 430 westerly flows showing agreement within 10 %.

In general both the LAEIs seem to be doing a reasonable job of estimating NO_x emissions in central London, and certainly better than the NAEI estimations. The LAEI, particularly in its enhanced form with measured road traffic emissions, has a much more explicit treatment of road transport emission than the NAEI, thus potentially providing a more accurate estimate of NO_x emissions in London. It uses vehicle speed and vehicle flow data from each road link using GPS based vehicle speed, as well as automatic number plate recognition data to 437 enhance vehicle stock information. The inventory also makes predictions of primary NO₂ 438 emissions, something that is potentially important in London due to the high proportion of diesel fuelled vehicles, which are likely to have a higher direct primary NO₂ emission 439 compared to petrol vehicles⁵⁰. The LAEI containing the enhanced treatment of traffic 440 441 emissions actually overestimates the NO_x emission in the central London footprint regimes (0 - 5 km from the BT tower), with greater overestimation outside the error of the measurements 442 443 under easterly flow conditions. This suggests potential extra errors in the treatment of traffic flow in the center of London to the east of the BT tower within the LAEI. The LAEI has a 444 significant contribution from other sources, which are mainly from non-road mobile 445 446 machinery (e.g. cranes, construction vehicles). These are virtually zero in the NAEI and it 447 could be errors in these sources that are contributing to the overestimation of the inventory in 448 central London. The better comparison with the LAEI compared to the NAEI support 449 previous tailpipe observations of higher NO_x emitted from the London vehicle diesel fleet 450 than is represented in the NAEI or predicted for several EURO emission control technologies 451 and show that a detailed treatment of traffic emissions is required to properly predict the NO_x emissions¹¹. There are no studies to our knowledge that specifically evaluate the London or 452 453 national inventories. However, it is clear from recent remote sensing measurements in 454 London during 2012 that emissions of NO_x have not decreased as expected through emissions legislation⁴⁹. This higher than anticipated vehicle NO_x is likely responsible for the significant 455 discrepancies that exist in London between observed NO_x and long-term NO_x projections, 456 457 and show that a detailed representation of traffic emissions is required to accurately represent 458 NO_x in London.

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461 Associated content

Figures S1 – S6 show a location map of the site, the time series of NO_x levels and fluxes from the BT tower, regression between BT tower NO_x and NO_x measured at a nearby urban background site, flux footprint statistics, a map of the location of the traffic count sites and maps of the 1km x 1km National Atmospheric Emission Inventory (NAEI) source specific emissions for NO_x . This information is available free of charge via the Internet at http://pubs.acs.org/

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