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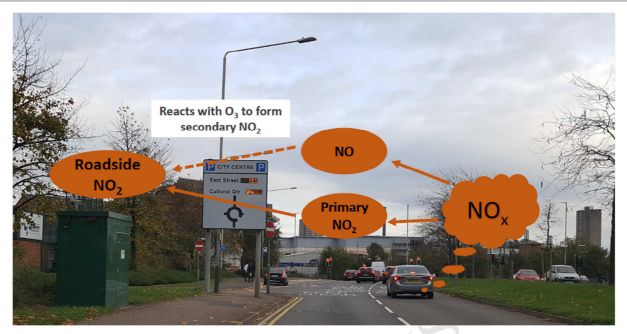
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10	Abstract
11	Nitrogen oxides (NO and NO ₂ , collectively NO_x) derived from vehicle exhausts are critical
12	pollutants with significant implications for urban air quality and human and environmental
13	health. In this study, we investigate trends in measured ambient nitrogen dioxide (NO $_2$) and NO $_x$

mixing ratios at urban traffic - dominated monitoring sites in the UK for the period 2009-2016. 14 15 We apply an oxidant analysis approach alongside a number of assumptions to the ambient data to determine trends in the inferred primary NO₂/NO_x ratio, and examine evidence for enhanced 16 17 vehicle "cold start" effects upon these inferred emissions. Ambient NO₂ and NO_x mixing ratios have experienced an overall decrease of 17.2% and 11.3% respectively for the locations 18 considered over this time period. The inferred primary NO₂/NO_x ratio for the majority of the 19 study locations is found to have fallen by 29% (from 0.175 to 0.125) as a monthly mean from 20 21 2009 to 2016, with a statistically significant median decrease of 0.32 percentage points per year. However, during cold weather (temperatures less than or equal to 5 °C), the inferred primary 22 NO₂/NO_x ratio averaged across all locations, when compared with normal conditions 23 (temperatures higher than 5 $^{\circ}$ C) increased from 0.062 (±0.004) to 0.102 (±0.001) (64.5% higher) 24 and from 0.056 (±0.004) to 0.098 (±0.001) (75% higher) for cold morning and evening rush 25 26 hours, with substantially greater increases at some sites. This "cold start" result suggests that the 27 combination of recent vehicle driving history and ambient weather conditions, in conjunction with technological constraints on the operating temperature range of emission control systems 28 29 in some vehicles, affects NO_x emissions and hence has a detrimental impact upon air quality in 30 urban environments. Increased cold start emissions imply an increased NO₂ - derived health 31 burden from air pollution, under certain conditions, assessment of which should consider changes in vehicle use as a result of weather, and hence altered personal exposure. 32

33

Key words: Urban air quality, Nitrogen oxides - primary NO₂, vehicle cold starts, trend analysis,
 vehicle emissions, low temperature emissions

36

37 **1. Introduction**

38 1.1 Background and objectives

39 Vehicle emissions are a dominant source of air pollution in urban environments in 40 developed nations. Direct emissions include nitrogen oxides (NO_x, the sum of nitrogen monoxide (NO) and nitrogen dioxide (NO₂)) and particulate matter (PM). NO_x emissions are of particular 41 importance to urban air quality, since they not only contribute to local and regional scale air 42 43 pollution, but also to the formation of secondary pollutants such as ozone (O_3) and secondary 44 PM. Vehicular pollutants have been reported to have substantial adverse impacts on human health (Papapostolou et al., 2011), with exposure to NO₂ known to lead to reduced lung function 45 and increased risk of cancer (Adam et al., 2015; Hamra et al., 2015; WHO, 2013), and is 46 47 responsible for tens of thousands of premature deaths each year across Europe (EEA, 2016; RCP, 48 2016; COMEAP, 2010). The International Agency for Research on Cancer has recently classified 49 diesel engine exhaust as a Group I carcinogen, based on its association with lung cancer 50 incidence (Attfield et al., 2012; Silverman et al., 2012).

Within the EU, a series of legislative measures have been introduced to address this 51 challenge. The first directive 96/62/EC (EC, 1996), which is commonly referred as the Air Quality 52 53 Framework Directive, and its daughter Directives, established standards in the period up to 2004 for a range of pollutants including NO₂ in ambient air. The three first daughter directives were 54 55 consolidated into a single ambient air quality Directive adopted as 2008/50/EC (EC, 2008a) and together with the fourth daughter directive 2004/107/EC they provide the current framework 56 57 for the control of ambient concentrations of air pollution in the UK. However, the need for 58 greater reductions in NO₂ mixing ratios has led to further directives, which gradually aimed to 59 decrease the emission levels directly from vehicles (EC, 2008b; 2011). In the UK, evidence indicates that new Euro limits (Euro 5 and Euro 6), in conjunction with new after-treatment 60 technologies (see section 1.2) have had a positive impact in the reduction of roadside PM_{2.5} 61 (Harrison and Beddows, 2017) and NO_x (DEFRA, 2017), while factors such as aging of catalysts 62

leading to reduced oxidative capacity via thermal deactivation might also play a role in affecting
primary NO₂ emissions from traffic (Carslaw et al., 2016b).

65 Here, we analyse trends in ambient NO_x and NO₂ mixing ratios between 2009 and 2016 66 measured at urban monitoring stations in the UK, where NO_x mixing ratios are dominated by 67 local traffic emissions. We assess changes in the ambient NO₂/NO_x ratio, and analyse the 68 ambient data to infer potential trends in the primary (directly emitted) NO_2/NO_x ratio from 69 vehicles. The NO_x data are coupled with meteorological observations and a methodology is 70 introduced to assess potential changes in vehicle emissions associated with "cold start" 71 operation, as indicated by ambient temperature, and the impact of this behaviour upon ambient 72 air quality.

73

74 1.2 Controls of vehicle NO_X and NO₂ emissions

75 Internal combustion vehicle technologies have developed to reduce fuel consumption, improve engineNitrogen oxides (NO and NO2, collectively NOX) derived from vehicle exhausts, 76 are critical pollutants with significant implications for urban air quality and human and 77 environmental health. In this study, we investigate trends in measured ambient nitrogen dioxide 78 79 (NO2) and NOX mixing ratios at urban traffic - dominated monitoring sites in the UK for the 80 period 2009-2016. We apply an oxidant analysis approach alongside a number of assumptions to the ambient data to determine trends in the inferred primary NO2/NOx ratio, and examine 81 82 evidence for enhanced vehicle "cold start" effects upon these inferred emissions. Ambient NO2 83 and NOX mixing ratios have experienced an overall decrease of 17.2% and 11.3% respectively for 84 the locations considered over this time period. The inferred primary NO2/NOx ratio for the 85 majority of the study locations is found to have fallen by 29% (from 0.175 to 0.125) as a monthly 86 mean from 2009 to 2016, with a statistically significant median decrease of 0.32 percentage 87 points per year. However, during cold weather (temperatures less than or equal to 5 OC), the 88 inferred primary NO2/NOx ratio averaged across all locations, when compared with normal 89 conditions (temperatures higher than 5 OC) increased from 0.062 (±0.004) to 0.102 (±0.001) 90 (64.5% higher) and from 0.056 (±0.004) to 0.098 (±0.001) (75% higher) for cold morning and 91 evening rush hours, with substantially greater increases at some sites. This "cold start" result 92 suggests that the combination of recent vehicle driving history and ambient weather conditions, 93 in conjunction with technological constraints on the operating temperature range of emission

94 control systems in some vehicles, affects NOx emissions and hence has a detrimental impact 95 upon air quality in urban environments. Increased cold start emissions imply an increased NO2 -96 derived health burden from air pollution, under certain conditions, assessment of which should 97 consider changes in vehicle use as a result of weather, and hence altered personal exposure. 98 performance, and (primarily through the addition of after-treatment systems) reduce emissions 99 of air pollutants such as NO_x. Of the two principal categories of internal combustion engine, the 100 emission problem is more straightforward to address for gasoline (spark ignition) vehicles in 101 comparison to diesel (compression ignition) power units. Most gasoline vehicles now employ a 102 three-way catalyst (TWC), a key abatement technology introduced in the 1980s and applied 103 widely from the 1990s to reduce vehicle tailpipe emissions. TWC are designed to simultaneously 104 convert carbon monoxide (CO) to carbon dioxide (CO₂), hydrocarbons (HC) to water and NO_x 105 species to nitrogen. Theoretically, TWC can cut CO, HC and NO_x emissions by over 99% (under 106 stoichiometric conditions) if the air to fuel ratio in the exhaust stream is accurately controlled, 107 although they have been associated with emissions of the greenhouse gas nitrous oxide (N_2O) 108 (Berges et al., 1993; Jimenez et al., 2000).

Diesel after-treatment systems face more challenges for the reduction of emissions. Diesel vehicles (Euro 3 and later) use diesel oxidation catalysts (DOC), which normally contain palladium, platinum and aluminium oxide, all of which serve as catalysts to oxidize HC and CO to CO₂ and H₂O. However, this oxidation can lead to increased NO₂ emissions when no further after-treatment technology is applied.

Exhaust gas recirculation (EGR) was introduced as a further emission control technology in larger size Euro 3 engines, and became the standard in Euro 4 and later diesel passenger cars and light duty vehicles (LDV). The EGR system channels and recirculates a portion of the exhaust gas into the filtered, high-pressure, fresh combustion air at the engine intake. The higher the engine load the better the EGR performance and the greater the NO_x reduction (Yokomura et al., 2003).

Diesel particle filters (DPF), introduced in 2009 to achieve Euro 5 limits, physically capture diesel particles and prevent their release to the atmosphere. However, the stored PM must be oxidized in order to avoid blocking the filter. This is achieved via reaction with O_2 at high temperatures (600 °C) and via reaction with NO_2 at low temperatures (250 – 450 °C). Owing to the amount of NO_2 needed in the DPF to burn the soot, the DPF is attached after the DOC system, forming a continuously regenerated trap (CRT) or catalyzed continuously regenerated trap (CCRT), which reduces PM, CO and HC. The secondary NO_2 formation as a by-product of the catalysis, however, is a major issue in these systems and can lead to increased NO_2 emissions from the vehicle.

129 The latest after-treatment technologies involve Lean NO_x trap (LNT) and Selective 130 catalytic reduction (SCR) technologies, which can achieve Euro 6 limits. LNT technology achieves 131 NO_x storage during lean engine operation, and NO_x reduction during rich operation phases. 132 During lean engine operation, NO_x is retained in the storage components in the form of nitrates 133 and nitrites. In a subsequent short fuel-rich period, the NO_x trap is regenerated by NO_x release 134 and reaction with HCs achieves reduction to N₂. Since fuel consumption for NO_x trap 135 regeneration depends on the regeneration frequency, which is a function of NO_x trap 136 performance parameters, the impact of catalyst aging must be considered in the operation 137 strategy.

Selective catalytic reduction (SCR) is an after-treatment system which catalyzes NO_x (NO + NO₂) reduction by addition of reactive nitrogen compounds, such as ammonia or urea-based mixtures. In SCR, the mixture of ammonia/urea reacts with NO_x to form N₂, CO₂, and H₂O. The SCR process requires precise control of the ammonia injection rate, because an insufficient injection rate may result in unacceptably low NO_x conversion, while an excessive injection rate results in releases of ammonia (undesirable "ammonia slip") to the atmosphere and increased SCR reductant consumption.

145 All these after-treatment systems need to reach a certain temperature threshold (i.e. regular operating conditions) in order to effectively reduce NO_x emissions. This difference in the 146 147 temperature, for both the engine and the catalytic converter, under which the vehicle is initially operated comparing to regular operating conditions can be expressed as a "cold start" when the 148 engine is operated with the temperature of the oil, coolant and all elements of the engine at the 149 150 ambient temperature. At lower ambient temperatures, the engine and catalyst warm up period 151 is prolonged and this can have an adverse effect on vehicle emissions. For modern gasoline and 152 diesel vehicles equipped with TWC and DOC after-treatment systems, under cold operation, this 153 implies dis-proportionally higher release of gaseous pollutants, since the temperature of the catalyst is not sufficient to ensure efficient NOx conversion. Currently in the EU, only the 154 155 emissions from gasoline Euro 3/4 vehicles under cold-start in low ambient temperature are 156 regulated under the directive 98/69/EC (EC, 1998), as identified elsewhere (e.g. Dardiotis et al., 157 2013; Bielaczyc et al., 2011; 2012).

159 1.3 NO₂ and NO_x emission measurement approaches

158

160 There are several different approaches to estimate vehicle combustion (tailpipe) 161 emissions - see e.g. the review of Franco et al., (2013). Here, we briefly introduce the key 162 approaches to provide context for the methodology used in this work. Most widely known are approved bench (chassis dynamometer) tests, where the tested vehicle follows a certain driving 163 cycle (sequence of speed/acceleration/deceleration over a pre-defined period of time that 164 165 corresponds to notional urban driving or extra-urban driving behaviour) and relevant analyzers 166 sample directly from the exhaust to determine (e.g.) NO_x emissions (Nine et al., 1999; Yanowitz 167 et al, 2000). However, such tests are very short (approximately 20 min each test) and, as has 168 been widely reported, fail to capture the real world operation NO_x emissions either by accident 169 or design (Andersson et al., 2014; Degraeuwe and Weiss, 2017). Thus, it is thought that historical 170 emission factors based solely upon such tests may not be representative of real-world on-road 171 vehicle behaviour.

172 Secondly, portable emission measurement systems (PEMS) are devices mounted to 173 individual test vehicles which measure directly from the exhaust/tailpipe during on-road driving 174 (Weiss et al, 2011). PEMS capture individual vehicle emissions under real driving conditions, but are expensive, can only be fitted in one car at a time, normally require tailpipe adaptations to 175 176 sample directly, and may suffer power limitations if powered by the vehicle on-board low voltage 177 power supply system. A related approach is the chase measurement method. In this approach, 178 instruments are mounted in a second, monitoring vehicle and an inlet is used in order to sample 179 from ambient air, while following individual (or groups of) target vehicles (Brantley et al., 2014). 180 Chase approaches can, compared with test bench measurements and PEMS approaches, more 181 readily give accurate information about fleet emissions from a number of vehicles and variations 182 with driving behaviour (e.g. urban, rural, motorway), but suffer from limitations of the need to 183 account for mixing with background air (commonly achieved via use of CO_2 as an exhaust tracer), 184 and overlap of multiple vehicle plumes.

Thirdly, remote sensing technologies (Bishop et al, 1989), where a light beam passes through the exhaust plume of an individual or series of target vehicles prior to measurement at a detector, wherein the amount of light absorbed is proportional to the concentration of gases in the plume (Carslaw and Rhys-Tyler, 2013). Those approaches are commonly coupled with vehicle 189 sensing and number plate recognition tools to identify speed, acceleration and vehicle/engine 190 characteristics. Although remote sensing methodologies are in many senses the "gold standard" 191 approach, in that they can measure the real-world on-road emissions from large numbers of 192 individual vehicles under favourable operating conditions, they suffer from constraints of cost, 193 complexity and potentially perturbations to traffic flow/behaviour. Furthermore, such 194 techniques have issues when located on multiple lane roads as emissions from vehicles 195 alongside the target vehicle can interfere with the result, in particular when the wind is in an 196 unfavourable direction - an alternative top-down geometry, which can sample individual, parallel 197 vehicle lanes, has recently been demonstrated (Ropkins et al., 2017).

198 Lastly, vehicle emissions may be inferred from analysis of ambient air quality 199 measurements, from monitoring sites located in suitable proximity to the road. This method has 200 the advantage of utilizing the data that in most cases are used to report ambient air pollution 201 levels, so no additional experiments are needed, and is able to exploit long term observations to 202 assess the efficacy of policy measures. However, it cannot give detailed information for the 203 vehicle fleet or driving behaviour. In this study we adopt this last approach, exploiting the 204 extensive UK air quality network to infer trends in cumulative UK fleet NO_x emissions and their 205 response to temperature, building upon previous, related work applying a similar approach to 206 infer changes in relative vehicle emissions (e.g. Carslaw et al., 2016b; Carslaw and Beevers, 2005; 207 Jenkin, 2004), and potential signatures of increased diesel vehicle penetration and tightening of emission standards (e.g. Carslaw et al., 2016b; Grange et al., 2017). 208

209

210 2. Data and Methodology

211 2.1 Experimental data

Atmospheric mixing ratios of NO_x and O_3 are monitored across the UK by the Automatic Urban and Rural Network (AURN), which is operated and maintained by the Department for Environment, Food and Rural Affairs (DEFRA), the Scottish Executive, the National Assembly for Wales and the Department of the Environment for Northern Ireland. Ambient mixing ratios are reported at hourly resolution, with measurements made by well-established conventional techniques, UV photometry for O_3 and chemiluminescence for NO_x with a molybdenum converter (see comments below, section 2.2, re NO_2 selectivity). The measurement uncertainty

219 for NO_x and O_3 from the AURN stations is <15%, following the European Committee for 220 Standardisation (documents BS EN14211:2012 -NO_x and BS EN14625:2012 -O₃).

221 The analysis presented here uses hourly data from every urban-traffic monitoring station 222 (defined as stations that are located at roadside and kerbside locations) in the UK AURN 223 network. For each urban-traffic station, background data were obtained from a nearby 224 background site, defined as the urban or rural background site (applying the AURN classification) 225 closest to the location of the urban traffic monitoring station. Further data availability 226 constraints of coverage from the start of 2009 to the end of 2016, and overall data capture 227 greater than 75% were applied (75 % selected as this is the minimum requirement for a valid 228 aggregated value over a day and a year according to the EU air quality reports (EEA, 2016; Font 229 and Fuller, 2016)). Additionally, for each urban traffic monitoring site, meteorological 230 measurements (UK Met-Office, downloaded from the British Atmospheric Data Centre (BADC, 231 https://badc.nerc.ac.uk/data/)) were used, in order to couple the air pollution measurements 232 with local meteorology. In total, we included 17 urban traffic stations, paired with 10 urban 233 background sites and 5 rural sites from the AURN network, alongside the 14 meteorological 234 sites. The rural stations were required as not all urban traffic sites have a proximate urban background station. The difference in the numbers of urban background and meteorological 235 236 stations arises as for London two urban background sites were used for four urban traffic 237 stations, and one meteorological station for the whole area. The map of the stations used is 238 shown in Fig 1, and their details are listed in the supplementary materials (Table S1).

239 2.2 Data processing methods and assumptions

240 Monitored data were processed and displayed using ArcGIS and R version 2.15.1 (R Core 241 Team, 2016) along with the R packages ggplot2 (Wickham, 2009), openair (Carslaw and Ropkins, 242 2012), and mcgv (Wood, 2003). The data were de-seasonalised by applying the smoothing LOESS 243 function, which is a non-parametric regression method that combines multiple regression 244 models in a k-nearest-neighbor-based meta-model (Cleveland et al., 1990). The statistical 245 approach that we followed for trend calculation is a non-parametric Mann-Kendall approach, while the trend slope is calculated with the Theil-Sen method (Sen, 1968; Theil, 1950) available 246 247 in the R-openair package. In this method, for a given set of n x, y pairs, the slopes between all pairs of points are calculated and the median is taken as an estimate of the most probable slope 248 249 (trend). This method is robust to outliers and can be used in both non-normal and heteroscedastic (non-constant error variance) data series. Bootstrap re-sampling was used for the calculation of confidence intervals at the 95% level and p-values. A statistically significant trend was assumed when p < 0.1 (represented with a '+' symbol), meaning that the trend was not random at a 90% chance; p-values of: p < 0.05, p < 0.01 and p < 0.001, marked by '*', '**' and '***', respectively, indicate very highly significant trends, while p > 0.1 shows insignificant trends.

To derive the inferred primary NO_2/NO_x ratio from ambient monitoring data we used the total oxidant (Ox = $NO_2 + O_3$) approach developed by Clapp and Jenkin (2001). It is well known that in the daytime atmosphere the interconversion of O_3 to NO_2 and vice versa is generally dominated by the following reactions:

(1)

(2)

(3)

 $260 \qquad NO+O_3 \rightarrow NO_2+O_2$

261 $NO_2 + hv \rightarrow NO + O$

which constitute a null cycle. Reactions (1) - (3) result in the cycling of NO_x between NO and NO_2 263 264 and total Ox between O_3 and NO_2 , however the total mixing ratios of both NO_x and Ox remain 265 constant. During daylight, the equilibrium that occurs in the above reactions determines the 266 photo-stationary steady state (PSS). Clapp and Jenkin, (2001) illustrated that when in PSS there is 267 a linear relation between Ox and NO_x for roadside station data, which may be interpreted as a 268 NO_x - independent and a NO_x - dependent contribution to Ox. The former is the regional 269 contribution which corresponds to the regional background O_3 level, whereas the latter is 270 effectively a local contribution that is associated with additional NO₂ (i.e. primary NO₂ emissions, 271 under the assumption that no other sources are significant). While this analysis only holds in 272 volume mixing ratio space, it should be noted that the AURN data-series are reported in $\mu g/m^3$, 273 therefore before we apply the total oxidant approach, data were converted from $\mu g/m^3$ to ppb by applying EU/DEFRA conversion factors (20 °C temperature and 1013 hPa pressure). For the 274 275 analysis presented here, "Daytime" was defined as all whole hours between sunrise and sunset, 276 based on the local time in London.

277 In the case of vehicle emissions from ambient urban traffic monitoring data, Jenkin, 278 (2004), demonstrated that by considering the 'total oxidant' slope, $(NO_2 + O_3) / NO_x$, estimates 279 could be drawn for the primary (direct) NO_2/NO_x ratio emitted from vehicles. However, most of 280 the AURN urban traffic monitoring sites in the UK do not have O_3 measurements which poses a 281 limitation to the total oxidant approach and to the estimation of the primary NO_2/NO_x ratio. In 282 such cases, Carslaw and Beevers, (2005) suggested a methodology that calculates the NO_2/NO_X 283 ratio without the need of O_3 observations from the urban traffic site, but with additional 284 observations (of NO_2 , NO_x and O_3) from a nearby urban background site. In general, this 285 approach assumes that the increment in NO₂ mixing ratio between a given urban traffic 286 monitoring site and a nearby urban background site is partitioned into NO_2 that is chemically 287 derived through the reaction between NO and O₃, and NO₂ which is emitted directly by road 288 vehicles. Therefore, data from paired sites (roadside and background/rural site) are used. 289 Additionally, in the calculation of the primary NO_2/NO_x ratio, the difference in NO_x between the 290 urban traffic and urban-background/rural stations is considered; therefore impacts from non-291 traffic related sources are essentially removed. The method uses a simple constrained model 292 and basic set of chemical reactions to explain the time-dependent variation in NO, NO₂ and O_3 as 293 vehicle plumes mix with background air. Carslaw and Beevers (2005), showed that the primary 294 NO_2/NO_x ratio may then be obtained from fitting the observed NO_2-NO_x data to that predicted, 295 whilst optimizing the primary NO₂ fraction and the mixing time between emission and observation at the urban site. Here, we follow this methodology, but note some limitations. 296 297 Specifically, the approach assumes that the increment in NO₂ mixing ratio above a local 298 background site is controlled by the availability of O_3 and directly emitted NO and NO₂ only and 299 does not include additional chemical reactions (e.g. of peroxy radicals with NO). Furthermore, it 300 should be mentioned that because the AURN analysers employ a heated molybdenum converter 301 to detect NO_2 (as NO), other NOy species will be detected as NO_2 . The influence of those 302 interferences might lead to overestimation of NO₂ in urban areas (Dunlea et al., 2007; Harrison 303 et al., 2012). In addition, direct emissions of HONO, thought to comprise a small, but still 304 significant and uncertain, component of vehicle exhaust (Crilley et al., 2016; Jenkin et al., 2008; 305 Kurtenbach et al., 2001), is also a potential interferent in NO₂ measurements. However, in our 306 analysis (see section 2.3 below) all the above potential interferences would be expected, to a 307 first approximation, to co-vary with NO_2 and hence have limited effect upon the inferred primary 308 NO_2/NO_x ratio.

309 2.3 Investigation of cold-start/emissions under low ambient temperature conditions.

310	To assess evidence for altered emissions under conditions of low ambient temperatures,
311	we applied the total oxidant approach for rush hour periods during the morning (06:00-10:00)
312	and afternoon (16:00-19:00) during winter time (November, December, January, February), with
313	a temperature condition of 5 ^{o}C (i.e., T \leq 5 ^{o}C). We select rush hours because the vehicle fleet
314	experiences less (and known) variations during those hours with respect to the rest of daytime.
315	It should be noted here that, although we used the total oxidant approach (notionally daylight
316	only, Clapp and Jenkin, 2001), by including morning pre-sunrise and evening post-sunset periods,
317	sensitivity tests showed that this introduced a very small difference in the overall results (less
318	than a 2% change in the inferred primary NO_2/NO_x ratio; less than 2 ppb difference in the ozone
319	background). According to European regulations, the bench/laboratory tests for standard vehicle
320	emissions are made under an (air) temperature range of 20 – 30 $^{\circ}$ C (EC, 91/441/ECC) and
321	normally at the fixed temperature value of 23 $^{\circ}$ C (DfT, 2016). The vehicle is given a standard pre-
322	conditioning in a temperature control room so that the whole vehicle including engine oil and
323	coolant is 'soaked' to the regulated temperature range. As a result, the selected temperature
324	threshold method (T \leq 5 $^{\rm o}C$), is well below that for the tested/regulated range, and very close to
325	the winter mean temperature in the UK (4.4 $^{ m o}$ C). Results for days that satisfied the temperature
326	criterion were compared with those that did not (Matthaios et al., 2017), in order to examine
327	any potential difference in emissions-driven air quality.

328

329 3. Results and Discussion

330 3.1 Ambient NO_x and NO₂ trends and relationships

331 The annual averaged ambient NO_x and NO_2 values for each urban traffic station are shown in Fig 2, normalised to their mean value (2009-2016). A clear decline in the levels of both 332 333 NO_x and NO_2 is evident, which based on the overall average from all sites is 11.3% for NO_x (black 334 dashed line, Fig 2a) and 17.2% for NO2 (black dashed line, Fig 2b) across the time period 335 considered (2009 - 2016). The larger NO_2 than NO_x decline in the UK urban traffic sites might be due to an alteration in PSS, emissions or in the background ozone that contributes to the 336 337 photochemical cycle. Fig 3 shows how the monthly mean rush hour NO2 mixing ratios (normalized to 100 on 1/1/2009) varies for the examined period. An overall mean reduction of 338 339 approximately 20% (see black line) in the NO2 mixing ratios is apparent. This sharper NO2 340 decrease cannot be explained by the changes in background ozone, since the background ozone 341 has not changed significantly during the examined years (0.5% increase - see Fig S1 in the 342 supplementary material). This decrease in NO₂ is somewhat counter-intuitive to that expected 343 when considering the increase in the number of licensed diesel vehicles in the UK, which, 344 according to the Department for Transport, have increased from about 39% (2009) to 52% (2016) in urban areas of the UK – and which are associated with higher NO_2 emissions than 345 346 petrol vehicles. However, it should be highlighted that during the period studied here, new Euro 347 limits came into force for the reduction of NO_x and NO₂ vehicle emissions. Briefly, Euro 5 limits, 348 first introduced in 2009, introduced a limit value of 0.18 g/km for NO_x in passenger cars, while 349 the tighter limits of Euro 6 for passenger cars (0.08 g/km for NO_x) came into force around 2014, 350 driving the introduction of new after-treatment technologies such as SCR and LNT. Depending 351 upon fleet penetration, these changes are beginning to play an important role in the (fleet 352 averaged) primary NO_2/NO_x ratio from vehicles (Carslaw et al., 2016b).

Figure 4 illustrates the variation in the ambient NO_2/NO_x ratio over time, as a function of 353 354 NO_x abundance, for the ambient monitoring data from urban traffic locations. Fig 4a (top) shows the distribution of hourly mean NO_x mixing ratios for 2009 and 2016; Fig 4b (right) the 355 356 distribution in hourly mean NO_2/NO_x ratios for the same years and Fig 4c (centre) the hourly 357 mean NO_2/NO_x ratio as a function of NO_x abundance. The values of NO_2/NO_x as a function of NO_x (Fig 4c) tend towards a minimum value, which approximates the primary NO_2/NO_x emission ratio 358 (this asymptote can be used as an alternative to the total oxidant approach described above in 359 360 order to estimate the primary NO_2/NO_x emission fraction; Itano et al, (2007) used a similar 361 approach to estimate the primary NO₂/NO_x ratio from an urban traffic site in Osaka, Japan). From 362 the distribution in Fig 4b (right) it can be seen that the NO₂/NO_x ratio has reduced slightly between 2009 and 2016 (median falling from 0.46 to 0.43), while from the asymptotes in Fig 4c 363 364 (centre) the primary NO_2/NO_x emission can be seen to have fallen from 0.175 to 0.125.

Figure 4d shows the absolute changes in NO_2/NO_x ratio as a function of the change in total NO_x mixing ratio (hourly mean for corresponding date / time / location) between 2009 and 2016, for each urban traffic location. The greatest reduction in NO_2/NO_x ratio has occurred in the locations where NO_x mixing ratio has increased (i.e. sites which have become more polluted, or busy, with positive values on the x axis), while for the majority of sites where NO_x abundance has decreased (negative x axis values), the NO_2/NO_x ratio has increased. To assess whether the 371 observed changes in NO_2/NO_x could be explained by NO_x - O_3 PSS chemistry alone a simple PSS 372 model, based on reactions 1-3, was used. In the first instance the model was initiated with 373 baseline conditions of 30 ppb of O_3 , 100 ppb of NO_x (75:25% NO:NO₂). In subsequent runs only 374 the initial NO_x mixing ratio was adjusted. In each case the model was run until NO, NO₂ and O_3 375 reached PSS and the difference in the final NO₂/NO_x ratio between the baseline run and all other 376 runs was plotted against the change in initial NO_x (see Fig 4b – blue line). It is observed that as 377 the NO_x abundance grows (positive x axis values), the PSS-chemistry-derived changes in the 378 NO_2/NO_x ratio (blue line) deviate from the observed ambient behavior (black points). Further 379 model runs (not shown here) demonstrated that if the PSS model was initiated with a higher 380 NO_2 fraction of 0.45 (i.e. similar to the mean NO_2/NO_x ratio observed in 2013 from Fig 4a) then 381 the difference between PSS-only predicted and ambient observed NO₂/NO_x differences becomes 382 even larger. Additional model PSS simulations for different initial ozone values $O_3 = 100$ ppb (extreme case) also showed deviation from observations. The results of the model runs suggest 383 384 that the response of the PSS-chemistry to changes in overall NO_x emissions cannot, in isolation, 385 explain the variation in the NO_2/NO_x ratio observed between 2009 and 2016.

386 The observed changes in NO_2/NO_x therefore are likely to be influenced primarily by 387 changes in vehicle emissions, which might have resulted from the combined changes in: (1) 388 policy, where new Euro limits came into force for new car passenger models (Euro 5 in 2009 and 389 Euro 6 in 2014), and for London (4 stations included here), a low emission zone came into effect, and daily charges for the London congestion zone increased. (2) Diesel cars, where in 2009, 90% 390 391 of diesel cars in the UK were Euro 3 and Euro 4, while in 2016, 69% of UK diesel cars were Euro 5 392 and Euro 6, according to the DfT / National Atmospheric Emissions Inventory NAEI, 2016). (3) 393 After-treatment technologies, where new technologies such as LNT and SCR have been 394 introduced (see section 1.2) and factors such as 'catalyst thrifting', where catalyst developers and 395 manufacturers reduce the amount of platinum group metal (which will potentially affect the 396 amount of NO₂ formed), or catalyst deactivation over time (where aged DOC catalyst 397 technologies may have reduced oxidative capacity through thermal deactivation or poisoning 398 and therefore less efficient conversion of NO to NO₂ (Carslaw et al., 2016b). Our results are in 399 accordance with other monitoring observations which show trends from eight long-running 400 urban traffic stations (1995 - 2015), and highlight that six out of eight stations show a downward 401 trend in the ambient NO₂ concentrations (DEFRA, 2017). In a recent study across Europe, Grange 402 et al. (2017) conclude that the ambient NO_2 levels measured at roadside monitoring sites have

403 fallen, and that the primary NO_2/NO_x emission ratio is lower than assumed by some key emission 404 inventories. This result is also in agreement with the findings of Font and Fuller (2016), who 405 analysed trends from ambient measurements in London and found that NO₂ and PM show a 406 declining trend from 2010 to 2015, attributed to a variety of factors, including a reduction of the 407 primary NO_2/NO_x emission ratio due to the absolute and relative reduction in numbers of older 408 diesel passenger cars in the overall fleet. These studies represent a growing body of evidence 409 that there has been a recent reduction in the ambient and primary emitted NO_2/NO_x ratio in UK 410 urban environments.

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412 3.2 Trends in inferred primary NO₂/NO_x ratio

The trend in inferred primary NO_2/NO_x ratio is presented in Fig 5 (monthly daytime 413 414 values from all urban traffic sites, calculated using the methodology of Palmgren et al., (1996) / 415 Carslaw and Beevers, (2005), after removing the observed seasonality (see section 2.1; Carslaw, 416 2016a)). Despite some individual monthly increases (mainly in 2011), the inferred primary 417 NO₂/NO_x ratio, shows a clear downward trend, with a change (median value) of -0.32%/year with 95% confidence interval from -0.45% to -0.2% per year, significant to the 0.001 level (p < 0.001). 418 419 Fig 5 shows an overall reduction in the inferred primary NO_2/NO_x ratio from 17.5% to 12.5% 420 between 2009 and 2016. Despite the increases reported for the urban traffic monitoring sites in 421 the UK before 2009 (AQEG, 2004, 2007), the inferred primary emitted NO_2/NO_x experiences an 422 overall reduction of 5 percentage points over the 2009 - 2016 period. In detail, from 2009 until 423 2011, a small reduction in the emissions is evident (from 17.5% to 15%). From 2011 until mid-2013 no significant changes are observed and the mean inferred primary NO_2/NO_x ratio is steady. 424 425 At that period the Euro 5a and Euro 5b limits for passenger and light duty vehicles, as well as the 426 Euro IV standards for heavy duty vehicles came into force for sales of new vehicles in September 427 2011 and January 2013 for light and heavy-duty vehicles respectively. Standards, which set a 428 prerequisite for diesel vehicles to have a DPF after-treatment system (Euro 5 diesel cars 429 constituted 20% of the UK diesel fleet in 2011 and they reached 52% by 2014 according to NAEI, 430 (2016)). This technology, as discussed in section 1.2, uses NO_2 in order to burn the collected 431 soot/PM in the filters, thus can also impact NO₂ emissions in the exhaust. In September 2014, 432 the most recent and stricter limits of Euro 6 came into force. The introduction of new Euro-limit-433 driven emissions control technologies and their penetration through the fleet (Euro 5 and Euro 6

434 passenger cars) may therefore have contributed to this decrease of 2.5 percentage points in the 435 mean primary NO_2/NO_x ratio from early 2014, and the overall decrease of 5 percentage points 436 that is apparent from early 2009. This 29% relative decrease in inferred primary NO_2/NO_x ratio in 437 the last period has to be taken into account in the emission inventories for future modelling 438 estimates of exposure and policy making which currently they estimate an increase until 2015 439 for UK (NAEI, 2016) and Europe (Kiesewetter et al., 2014; Grange et al., 2017).

440 The change in inferred primary NO₂/NO_x ratio between 2009 and 2016 shows some 441 variability with geographic region across the UK (Fig 6). With few exceptions, the primary 442 NO_2/NO_x ratio has fallen in the majority of the regions, by a range from 0.5 to 10 % (Table 1). The 443 observed increases in the ratio in some regions (e.g York, Swansea) are difficult to explain 444 individually - they may be associated with other local factors such as changing road layouts and 445 urban infrastructure. A decline in the median inferred primary NO_2/NO_x emission for London of 5% is apparent, in agreement with Carslaw et al., (2016b), who found a decrease of about 7% in 446 447 the primary NO₂/NO₂ ratio for inner London from 2009/10 to 2014/15. However, these variations 448 were not always consistent with the monthly primary NO_2/NO_x trend (calculated for the whole 449 data series) (Table 1), indicating that the trends are not significant in every area and more data 450 (i.e. a longer time series) are needed in order for a clear conclusion to be drawn. This potentially 451 reflects that the oxidant analysis assumptions (essentially, that the NO_{3}/O_{3} PSS represents a fully 452 closed system) may be less valid for some locations and highlights the complexity and the 453 heterogeneity in the implementation and outcome of policy interventions to control air pollution. 454

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456 3.3 Low temperature primary NO₂ emissions – evidence for "cold start" effects

The inferred NO₂/NO_x emission ratios for cold (T \leq 5°C) and non-cold/normal (T > 5°C) 457 458 conditions during winter (November, December, January, February) morning and evening rush 459 hours are depicted in Fig 7. The change in gradient between cold (Fig 7b, 7d) and non-460 cold/normal (Figs 7a, 7c) conditions is clearly apparent, with the O_x -NO_x analysis indicating an 461 increase of 64.5% (mornings) and 75% (evenings) in the inferred primary NO₂/NO_x ratio for the 462 former compared with the latter. The increase in overall NO_x abundance (greater x-axis range, most likely arising primarily from lower and more stable atmospheric boundary layer/mixing 463 464 heights under cold conditions, alongside reduced winter month NO₂ photolysis) is also apparent.

The 64.5% increase in inferred primary NO₂/NO_x ratio is statistically significant (p < 0.001), changing by over 4 percentage points from 6.2 (\pm 0.4) %, for normal winter mornings to 10.2 (\pm 0.1) %, for cold winter mornings. Similar behaviour is observed for evenings (75% increase from normal to cold conditions), where the difference is slightly higher at 4.2 percentage points, from 5.6 (\pm 0.4) to 9.8(\pm 0.1) % (values in the parenthesis indicate the standard error). For individual locations the difference between cold and normal conditions ranges over a factor of 1.6 to 3.8 (Table 2).

472 Atmospheric photochemical processes (i.e., chemical reaction rate constants and to a 473 lesser extent photolysis frequencies) are dependent upon temperature, which can introduce 474 changes in concentrations, as a function of temperature, under otherwise constant conditions 475 i.e. without any emission change. To assess the extent to which this may contribute to the 476 observed trends, a series of tests were performed using a simple chemical box-model including a 477 full inorganic chemical mechanism (taken from the MCM, http://mcm.leeds.ac.uk/, Saunders et 478 al., 2003) (described in detail in the supplementary material). The model was run for a 24-hour period for a notional winter day (15thDecember), for a range of temperatures (-10 to +30 °C), to 479 assess the impact of the temperature dependence of chemical processes (alone) upon the 480 ambient NO_2/NO_x ratio. The results showed that the daytime and nighttime average primary 481 482 NO_2/NO_x ratios during the rush hour period (i.e., the period examined here for the cold starts) 483 varied by between 0.5% and 8.4%, with a mean relative change of 2.4%, due to the effect of 484 temperature-dependent chemical kinetics (see supplementary material Fig S4, S5). The model 485 results indicate that the observed increments in the inferred primary NO₂/NO_x ratio with low 486 temperatures, of 64.5% and 75%, are substantially larger than can be accounted for by the 487 temperature-dependence of the atmospheric photochemistry. We suggest therefore that 488 changed vehicle emissions with operating temperature are contributing towards the observed 489 difference in inferred NO_2/NO_x emission ratio with temperature. While the ambient data will 490 contain contributions from vehicles which have been operating for some time and hence are not 491 under "cold start" conditions, statistics on mode of transport from the UK DfT National Travel 492 Survey in 2016 indicate that 20% of all trips under 1 mile (1.6 km), were by car (either as driver or passenger), increasing to 77% for trips between 1 and 5 miles (1.6 - 8 km) (DfT, 2017). For 493 494 urban journeys, distances tend to be shorter than these UK mean values suggest. Short urban 495 trip duration alongside the reduced efficiency or designed deactivation of emission control systems at temperatures below 250 °C may contribute to the substantial increase in the inferred 496

497 primary NO_2/NO_x ratio, and hence reduced attainment of NO_2 air quality standards, under cold 498 weather conditions. Recent testing of vehicles in Germany conducted in 2016 found that over 499 half of ca. 50 vehicles tested had their EGR system inoperative under low ambient temperatures, 500 for engine protection purposes (BMVI, 2016). Such differences in the vehicle NO_x emissions under low temperatures are observed in Euro 6 passenger cars (Suarez-Bertoa and Astorga, 501 502 2018). During regulated laboratory tests Suarez-Bertoa and Astorga (2018) observed that Euro 6 503 gasoline and diesel vehicles emitted two to more than three times more NO_x at pre-conditioned initial engine temperatures of - 7 °C, compared to emissions at 23 °C. These results underline 504 the lower effectiveness in the performance of the after-treatment systems under the low initial 505 506 operating temperatures found in some real driving conditions. However, the relative 507 contribution of gasoline and diesel vehicles, or passenger cars vs light- and heavy-goods vehicles, to the change in inferred primary NO_2/NO_x ratio cannot be readily determined from these data. 508 At intermediate temperatures (5 – 15 $^{\circ}$ C) reduced changes in the inferred primary NO₂/NO_x ratio 509 510 were found (see Fig S2, S3 in supplementary material).

511 Fig 8 shows the relationship between inferred mean primary NO_2/NO_x ratio and mean ambient temperature, for the cold months (November, December, January and February). The 512 primary NO₂/NO_x ratio is significantly negatively correlated with temperature ($R^2 = 0.54$), 513 according to the inverse relationship fNO₂ = $15\% - [0.77 \pm 0.15] \times (T / {}^{\circ}C)$. Similar results to those 514 515 presented here were reported by Degraeuwe and Weiss (2017), who found that for on-road 516 tests, Euro 5 and Euro 6 cars emit four and three times higher NO_x at low ambient temperatures than the legislative limits, respectively. For Euro 6 cars, similar results were reported by Kwon et 517 al. (2017), who compared NO_x emissions at lower ambient temperatures (0 – 5 $^{\circ}$ C) with higher 518 ambient temperatures $(15 - 20 \degree C)$ and found a difference of 82 – 192%. 519

These literature results, along with the analysis presented here, support the finding that higher vehicle emissions at lower ambient temperatures may have a measurable impact upon ambient air quality. Vehicle emissions effects are likely exacerbated by meteorology during wintertime (lower boundary layer height, and increasing tendency to stable conditions (temperature inversions) during the worst pollution episodes and coldest weather); reduced photolysis frequencies (shifting the NO_x PSS towards NO₂), and potentially greater vehicle use during low temperature conditions in the UK, relative to other travel options.

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528 4. Conclusions

529 This study presents an analysis of ambient NO_x and NO_2 mixing ratios, to derive inferred 530 NO_2/NO_x emission ratios from urban traffic monitoring stations in the UK, for the period from 531 2009 until 2016. The analysis showed:

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 In keeping with other recent results, a decrease in the mean ambient NO₂ mixing ratio at selected urban roadside measurement sites between 2009 and 2016, alongside a smaller reduction in NO_x mixing ratios.

- 535 Direct emissions are not measured in this work, and a number of assumptions are made 536 in deducing emitted NO_x ratios from ambient data for a finite number of measurement 537 stations. The mean inferred primary NO_2/NO_x ratio shows an overall reduction of 5 538 percentage points from 17.5% to 12.5%, with a calculated reduction of -0.32 percentage 539 points/year (95% confidence interval of -0.2 to -0.45 percentage points/year), over this 540 time period. The period of this reduction corresponds to the combined development of 541 policy (new Euro limits), changes in fleet composition and, to some extent, to the initial penetration of new after-treatment technologies introduced in the last eight years. 542 However, this reduction is not evident in every location, underling the complexity of the 543 problem and implying that more stringent measures may be needed to further reduce 544 ambient NO₂ in urban environments, a challenge that will be heightened by increasing 545 vehicle numbers. 546
- Enhanced "cold-start" emissions, inferred from ambient monitoring data under low temperature conditions. The results imply that the overall vehicle primary NO₂/NO_x ratios increase on average by 64.5% and 75% for morning and evening rush hours under cold (\leq 5 °C) conditions, compared with (for the UK) normal conditions (> 5 °C), while this difference can be 1.6 - 3.8 times higher, when examining individual urban traffic monitoring stations. An inverse relationship is found between ambient temperature and primary NO₂/NO_x ratio.

These results suggest that the combination of (short duration) driving patterns and the temperature dependence of the current after-treatment systems under low ambient temperatures leads to measurable impacts upon ambient air quality. They also highlight the importance of consideration of cold-start emissions within vehicle test cycles, as these may cause a measurable deterioration in air quality in urban areas. Assessment of the related health burden – which must integrate personal exposure considerations – should combine the impact
of meteorological factors on both vehicle performance, and public behaviour vs transport
choices.

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Region	Median 2009 primary	Median 2016 primary	Difference primary	Monthly trend with 95%
	NO ₂ /NO _x (%)	NO ₂ /NO _x (%)	NO ₂ /NO _x (%)	confidence limits (%/year)
Aberdeen	19	10	-9	-1.33 (-1.52, -1.1) ***
Birmingham	20	10	-10	0.24 (-0.29, 0.78)
Cambridge	12	8	-4	-0.83 (-1.04, -0.6) ***
Carlisle	21.5	17	-4.5	-0.16 (-0.51, 0.18)
Chepstow	14	16	2	0.06 (-0.28, 0.34)
Exeter	18	15	-3	-0.45 (-0.75, -0.13) **
Leeds	16	13	-3	-0.17 (-0.5, 0.2)
Liverpool	21	19	-2	0.42 (-0.13, 1.01)
London	15	10	-5	-0.78 (-1.08, -0.38) **
Newcastle	11	10	-1	-0.5 (-1.26, 0.37)
Sandy	10	15	5	0.68 (0.39, 0.96) ***
Stanford le hope	20	17.5	-2.5	-1.79 (-2.45, -1.13) ***
Swansea	27	26.5	-0.5	0.16 (-0.12, 0.43)
York	8	15	7	-0.8 (-1.61, -0.06) *

Table 1. Median inferred primary NO₂/NO_x ratios, difference and monthly trend (percentage points/year) for individual urban traffic monitoring sites in the UK from 2009 to 2016. Symbols (***, ***, *, +) indicate the level of significance p (0.001, 0.01, 0.05, 0.1). Note that the median primary NO₂/NO_x values were acquired with a localised fitted regression model, while the monthly trend was calculated with a fitted linear regression model.

Region	Cold start primary NO ₂ emissions (%)		Primary NO ₂ emissions (%)		Relative increment (%)	
	Morning	Evening	Morning	Evening	Morning	Evening
Aberdeen	$12.3 \pm 0.17^{***}$	$13.1 \pm 0.18^{***}$	12.0±0.3	13.0±0.3	2.5	0.1
Birmingham	10±0.14***	10.4±0.23***	7.2±0.4	8.2±0.3	38.8	26.8
Cambridge	8.5±0.36***	7.2±0.36***	3.6±0.7	4.7±0.5	136.1	53.2
Carlisle	12.6±0.43***	13.2±0.47***	10.1±0.8	7.0±0.5	24.8	88.5
Chepstow	9.7±0.62***	11.9±0.1***	9.9±0.9	12.0±0.8	-	0.1
Exeter	12.1±0.19***	11.7±0.33***	6.5±0.3	6.8±0.4	86.2	72.1
Leeds	8.5±0.2***	8.0±0.3***	4.6±0.4	3.2±0.3	84.8	150
Liverpool	9.4±0.3***	11.3±0.48***	7.8±0.7	11.2±0.6	20.5	0.1
London Camden	15±0.2***	14.2±0.4***	12.2±0.3	11.1±0.3	22.9	27.9
London Haringey	6.8±0.2***	6.9±0.04***	1.4±0.5	3.3±0.3	385.7	109.1
London Marylebone	12.3±0.1***	13.7±0.2***	12±0.2	13.7±0.2	2.5	-
London tower hamlet	13.5±0.3***	13.8±0.7***	12.6±0.6	13.8±0.5	7.1	-
Newcastle	10.7±0.1***	11.4±0.2***	7.9±0.4	8.3±.0.3	35.4	25.3
Sandy	16.3±0.3***	15.1±0.3***	15±0.7	13.2±0.6	8.7	14.3
Stanford le hope	7.0±0.3***	8.2±0.4***	4.8±0.9	7.8±0.6	45.8	12.8
Swansea	24.4±0.3***	23.4±0.4***	15±0.4	13.2±0.4	62.7	77.2
York	13.8±0.3***	13.5±0.4***	5.7±0.7	5.3±0.4	142.1	158.4

Table 2. Primary NO_2/NO_x ratio during cold weather rush hours (representing cold start emissions) and normal weather rush hours (representing normal emissions) and the corresponding increment for individual areas across the UK. Symbols (***, **, *, +) indicate the level of significance of the slope p (0.001, 0.01, 0.05, 0.1).

FIGURES

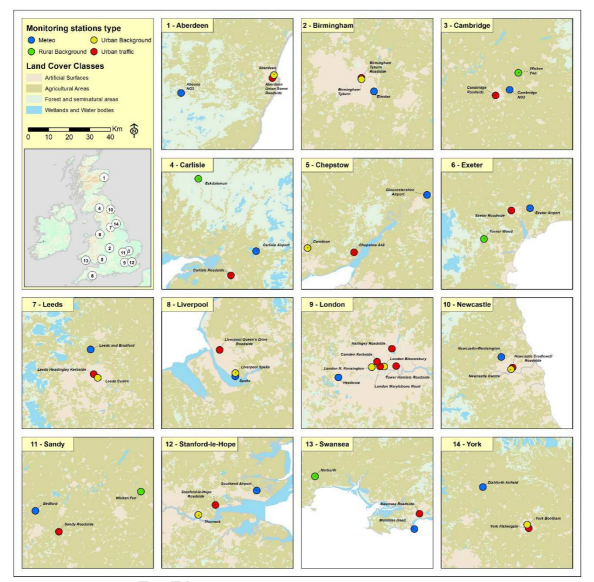


Fig 1. Monitoring sites used in this study. Blue indicates the meteorological stations, red the urban traffic sites, yellow the urban background and green the rural sites.

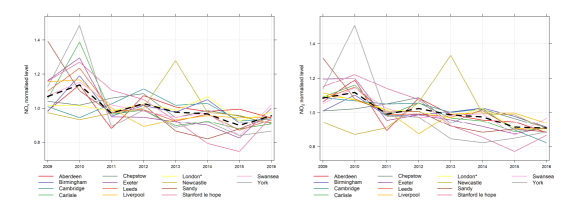


Fig 2. Annual average NO_x (left) and NO_2 (right) values for urban traffic stations in the UK normalized to their mean value. Black dashed line is the average from all sites used.

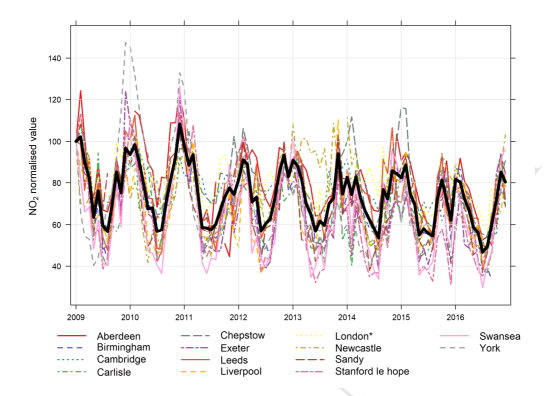


Fig 3. Monthly NO₂ mixing ratios during rush hour periods (06:00-10:00, 16:00-19:00), normalized to their 1/1/2009 value, for urban traffic monitoring sites across the UK. Black line indicates the mean value across all stations used.

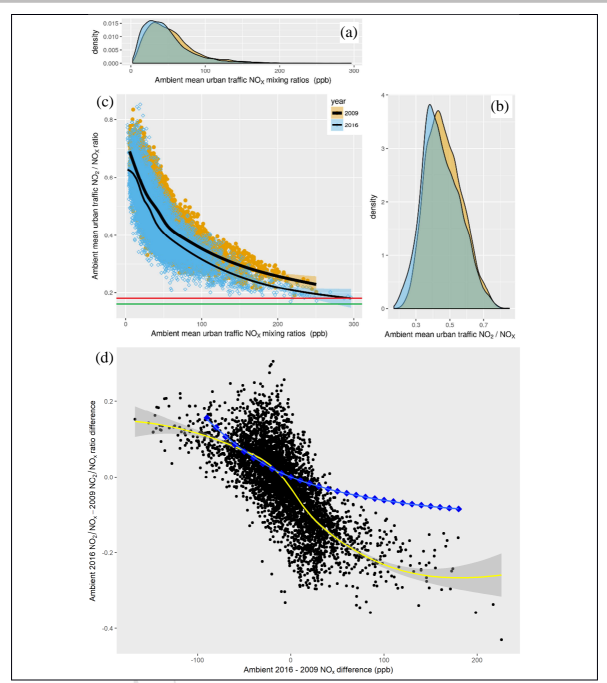


Fig 4. Daytime hourly mean ambient (a) NO_x and (b) NO_2/NO_x distributions for 2009 and 2016 from all urban traffic monitoring sites in the UK. (c) NO_2/NO_x ratio vs NO_x : the green (2016) and red (2009) lines indicate the baselines of the distributions and can be used to estimate the primary NO_2 (see text). (d) ambient NO_2/NO_x ratio as a function of the difference in NO_x mixing ratio for each location, between 2009 and 2016. The yellow line indicates the locally fitted regression (LOESS) line with associated 95% confidence interval, while the blue line indicates the variation in NO_2/NO_x expected, for a change in overall NO_x abundance, on the basis of the NO_x - O_3 PSS chemistry alone (see text for details).

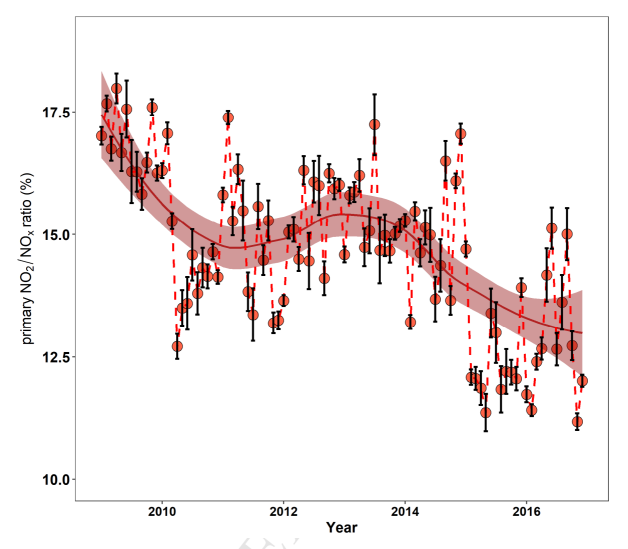


Fig 5. Mean overall trend of the inferred primary NO_2/NO_x ratio averaged over all UK urban traffic monitoring sites. The error bars indicate the standard error of the slope estimates. The local regression fitted line (red solid line) is weighted by taking into account the standard errors of the individual slopes. The shaded area indicates the 95% confidence intervals.

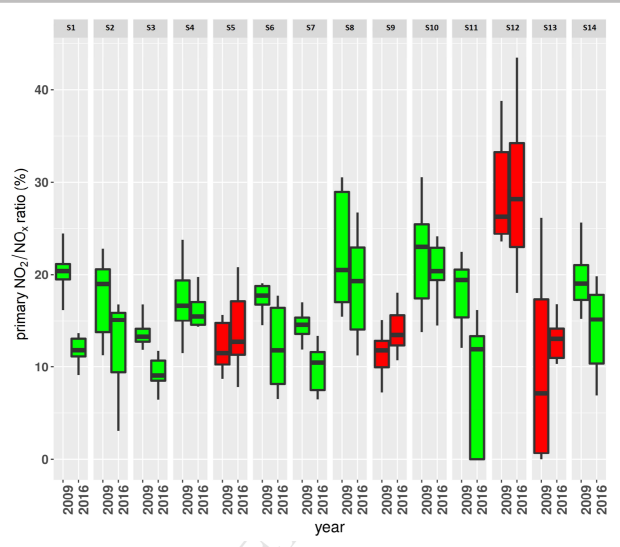


Fig 6. Box-whisker plots for the inferred primary NO_2/NO_x ratio for the individual urban traffic monitoring sites considered in this study, during 2009 and 2016. Green indicates a reduction in the median value between the two dates for each location, red an increase, between 2009 and 2016. S1: Aberdeen, S2: Birmingham, S3: Cambridge, S4: Carlisle, S5: Chepstow, S6: Exeter, S7: Leeds, S8: Liverpool, S9: Newcastle, S10: Sandy, S11: Stanford le Hope, S12: Swansea, S13: York, S14: London.

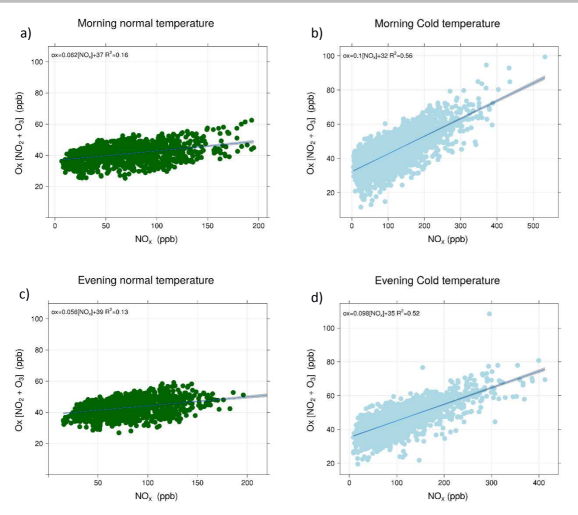


Fig 7. Ox -NO_x plots during normal winter morning and evening rush hours (Fig 7a, Fig 7c, green) and during cold (T \leq 5 °C) morning and evening rush hours (Fig 7b, Fig 7d, blue), representing normal and potential cold start emissions. The shaded areas indicate the 95% confidence intervals of the linear regression fit.

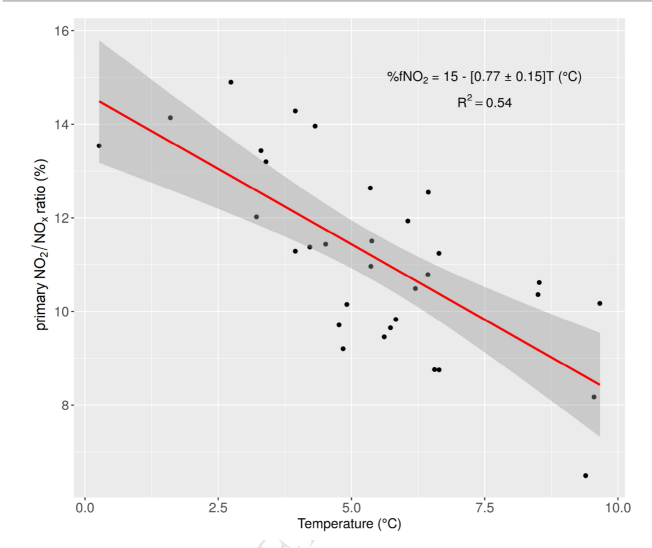


Fig 8. Inferred mean monthly primary NO_2/NO_x ratio (%fNO₂) and its dependence on mean ambient temperature in the UK. The grey shaded areas indicate the 95% confidence intervals of the regression relationship.

Highlights

- A methodology is developed to assess enhanced "cold-start" primary NO₂ emissions from vehicles.
- Ambient mixing ratios of NO₂ and NO_x measured at selected UK urban traffic sites have fallen over the period from 2009-2016.
- Overall inferred primary NO₂/NO_x ratio experiences a statistically significant decrease from 17.5% to 12.5% between 2009 and 2016.
- Inferred cold-start morning and evening primary NO₂ vehicle emissions are significantly higher than those found under warmer conditions
- Inferred cold-start primary NO₂ vehicle emissions have measurable impacts upon urban air quality in the UK.

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