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Characterization of gas and particulate phase organic emissions (Č9-C37) from a diesel engine and the effect of abatement devices

Alam, Mohammed S.; Zeraati-Rezaei, Soheil; Xu, Hongming; Harrison, Roy M.

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2 3 4 5	Characterisation of Gas and Particulate Phase Organic Emissions (C9-C37) from a Diesel Engine and the Effect of Abatement Devices
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7	Mohammed S. Alam ¹ , Soheil Zeraati-Rezaei ² ,
8	Hongming Xu ² and Roy M. Harrison ^{1*†}
9	
10	¹ School of Geography, Earth & Environmental Sciences
12	University of Birmingham, Edgbaston, Birmingham, B15 2TT
13	United Kingdom
14	
15	
16	² Department of Mechanical Engineering
17	School of Engineering, University of Birmingham, Edgbaston
18	Birmingham, B15 211
19 20	United Kingdom
20	

^{*} To whom correspondence should be addressed. Tele: +44 121 414 3494; Fax: +44 121 414 3708; Email: r.m.harrison@bham.ac.uk

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



24 ABSTRACT

Particulate and vapour phase emissions in the diluted exhaust of a light duty diesel engine designed 25 for Euro 5 application have been sampled. The engine was operated in three modes, and samples 26 27 were collected from exhaust without aftertreatment but also with aftertreatment by an exhaust oxidation catalyst and particle filter. The samples were analysed by 2-dimensional gas 28 chromatography with Time-of-Flight mass spectral detection. The results show overall removal 29 30 efficiencies for organic compound mass by the combination of oxidation catalyst and particle filter 31 of 50%, 56% and 74% for the high speed/high load, low speed/low load and high speed/low load conditions respectively. The results are clearly indicative of substantial repartitioning of the 32 33 particulate and vapour components within the abatement devices and show an apparently reduced efficiency for removal of high molecular weight alkanes under high speed/high load conditions 34 relative to lower molecular weight compounds, although this may be due to alkane formation by 35 thermocracking of other species. A notable feature is the presence of oxygenated compounds in the 36 emissions which are not present in the fuel. These are increased under high speed/high load 37 38 conditions and the results suggest formation in the aftertreatment devices as well as in the combustion 39 process.

40 INTRODUCTION

Diesel vehicles are an important source of atmospheric pollutants, particularly in urban environments¹. Numerous studies have investigated diesel exhaust generated particulate matter (PM) due to the associated adverse health and environmental effects^{2,3,4,5}. Diesel exhaust PM consists mainly of agglomerated solid carbonaceous material and ash, together with organic and sulphur containing compounds⁶. The organic fraction is mainly sourced from unburned/partially-oxidised fuel and lubricating oil while their contribution to emitted particles varies with engine design, operating conditions and fuel formulation, and can be between <10 to >90% by mass^{6,7}.

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49 Traditional studies of organic species emitted in diesel engine exhaust have focused on volatile organic compounds (VOC) and primary organic aerosol (POA)^{8,9,10,11}. More recently, the focus has 50 been on the measurement of intermediate-volatility organic compounds (IVOC)^{12,13,14,15,16} and semi-51 volatile organic compounds (SVOC)^{17,18,19} which have effective saturation concentrations (C*) of 52 10^3 - 10^6 and 1- $10^3 \mu g m^{-3}$, respectively²⁰. This typically corresponds to hydrocarbons in the range C₁₂-53 C₂₂ and C₁₅-C₃₅ for IVOC and SVOC in engine exhaust, respectively. SVOC include any compound 54 with >1% of its mass in both the condensed and vapour phases, while IVOC are found almost 55 exclusively in the vapour phase but have a high potential to generate lower volatility products after 56 reaction²¹. Henceforth, in this study, S/IVOC will be taken to include both SVOC and IVOC. The 57 emissions of S/IVOC can lead to the formation of ozone and secondary organic aerosol 58 (SOA)^{13,22,23,24}. A full understanding of the composition of POA from diesel emissions and its 59 contribution to SOA formation still remain elusive, as many uncertainties exist relating to the semi-60 volatile component of the particles¹⁸. 61

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Emissions inventories such as the National Atmospheric Emissions Inventory (NAEI) in the UK (or NEI in the US) report many fuel based emissions factors (EF), including EFs of non-volatile particulate matter and volatile organic compounds $<C_{12}$ in the vapour phase. EFs of S/IVOC from diesel vehicles, however are scarce and are usually reported only for individual species such as PAH
or n-alkanes^{25,26,27}. Recent literature has reported IVOC and SVOC emissions from diesel
vehicles^{11,14,15,28}.

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At present relatively little is known about the chemical composition and magnitude of IVOC and 70 SVOC from diesel engine emissions, largely because of the scarcity of analytical techniques that 71 72 accurately resolve the complexity and allow quantification of speciated organic emissions. 73 Conventional gas chromatography (GC) techniques have been used to identify S/IVOC in diesel exhaust^{14,26,27}. The species identified/quantified in these studies are typically polycyclic aromatic 74 75 hydrocarbons (PAH), n-alkanes, hopanes and steranes; and 80-90% of the S/IVOC composition remains uncharacterised because it is unresolved by traditional GC, forming a large hump in the 76 chromatogram referred to as 'unresolved complex mixture' (UCM)²⁹. With significant advancements 77 in analytical techniques, studies have reported more comprehensive S/IVOC compositional 78 information for diesel fuel^{19,22,30,31}, lubricating oil^{19,32,33,34,35} and S/IVOC from diesel 79 emissions^{15,16,19,36}. Recently, Alam et al.¹⁹ comprehensively characterised diesel fuel, lubricating oil 80 and diesel exhaust emissions in the gas and particulate phases. The majority of diesel and lubricating 81 oil is comprised of acyclic and monocyclic alkanes (76% and 59%, respectively). This is reflected in 82 83 the gas and particulate phase emissions which were also largely comprised of these compounds. Diesel exhaust gas phase emissions were remarkably similar to diesel fuel (with the exception of an 84 increased number of monocyclic aromatics in the emissions), whereas particulate emissions had an 85 organic composition similar to lubricating oil. Emissions in the particulate phase, however, were 86 comprised of few compound groups that were not identified in the lubricating oil including, PAH, 87 biphenyls, oxygenates and FAMEs (see Table S1 in Alam et al.¹⁹). 88

89

90 The majority of studies have investigated diesel emissions using a dynamometer rig, averaging
91 emissions over a particular vehicle driving cycle (e.g. NEDC, WLTP, FTP-75, etc.) and thus

attempting to simulate real world driving conditions^{10,37}. Although the use of an engine or chassis 92 dynamometer permits tight control of engine and abatement device parameters, these studies offer 93 limited information with regard to combustion processes and specific engine conditions that play a 94 95 crucial role in the emissions of S/IVOC, particularly when averaging data over an entire cycle. A limited number of studies have therefore investigated S/IVOC emissions under steady-state engine 96 conditions (constant engine torque, speed, etc.) with and without emission control technologies^{15,16,17}. 97 The fraction of vehicles with diesel oxidation catalysts (DOC) and diesel particulate filters (DPF) is 98 relatively smaller in developing countries³⁸, and so it is important to comprehensively characterise 99 the composition of S/IVOC emissions from diesel engines both with and without abatement devices. 100 101 The global on-road light duty diesel fleet includes older vehicles with no exhaust aftertreatment, vehicles of intermediate age with a DOC, and in countries where the fuel quality is adequate, vehicles 102 with both a DOC and DPF. This study therefore includes measurements of all such configurations. 103

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In this study we use thermal desorption coupled to comprehensive two dimensional gas 105 chromatography time-of-flight mass spectrometry (TD-GC×GC-ToFMS) to characterise and 106 quantify the composition of S/IVOC, both in the gas and particulate phases, from a light duty diesel 107 engine under different steady-state operating conditions with and without exhaust aftertreatment 108 systems. We investigate compositional changes and emission factors with varying engine speed/load, 109 with and without DOC and DPF. Detailed chemical speciation based on structure and functionality is 110 reported using an established mapping and quantification technique for isomer sets of S/IVOC¹⁹. The 111 results aim to shed light ultimately on the atmospheric fate of emitted S/IVOC and to better 112 understand the sources of emission which include unburned and partially burned fuel and lubricant, 113 as well as products of pyrolysis of fuel/lubricant or products of catalysis. 114

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118 EXPERIMENTAL

119 Sampling

Experiments were conducted with a light-duty 2.2 L, 4-cylinder inline compression ignition diesel 120 121 engine (designed for Euro 5 applications); equipped with a common rail direct injection system, a variable-nozzle-turbine turbocharger, a diesel oxidation catalyst (DOC) and a diesel particulate filter 122 123 (DPF). The open engine control unit (Open-ECU) allowed full control over the engine operating parameters. 30 min samples were collected at steady-state engine operating conditions at three 124 125 different load/speed combinations (see Table S1). Each experiment type was performed before, and in some experiments after, the DOC, and after the DPF to assess the performance of the abatement 126 127 devices installed. The DPF fitted to the test engine is believed to be catalysed as it oxidised unconverted exhaust CO and HC leaving the upstream DOC during the experiments. It also depends 128 upon the production of nitrogen dioxide in the upstream DOC which is able to oxidise the deposited 129 carbonaceous particles at a much reduced temperature compared to oxidation by molecular oxygen. 130 Hence the DPF is continuously regenerating but exhaust pressure before and after the DPF is 131 132 monitored to use an active regeneration strategy if the exhaust back pressure becomes excessive. This did not occur during our experiments and therefore the results represent the conditions occurring only 133 during continuous regeneration. 134

135

A pilot+main fuel injection strategy was used for all of the engine operating conditions. Exhaust gas 136 recirculation (EGR) and fuel injection pressure were fixed at approximately 55% and 500 bar, 137 respectively, for 1.4 bar BMEP and 1000 RPM, 33% and 600 bar for 1.4 bar BMEP and 1800 RPM, 138 17% and 1000 bar for 5.9 bar BMEP and 1800 RPM. Sampling was started after ensuring that the 139 140 engine was fully warmed-up and thermally stable. Standard European, EN590 specifications, ultralow sulphur diesel (ULSD, S < 10 ppm) and 5W-30 part synthetic engine lubricating oil were used 141 for the experiments. Comprehensive details regarding the engine test rig is described in a companion 142 paper³⁹. For S/IVOC characterisation, engine exhaust was diluted (1:50) using a modified TSI 3302A 143

diluter described in detail elsewhere¹⁸. Briefly, the undiluted hot exhaust sample was delivered via a 144 heated line maintained at 191°C (minimising sampling wall losses) and mixed with cleaned (moisture 145 free) compressed air, at ambient temperature, by utilising a high flow rate moisture trap and two 146 147 HEPA filters. The diluted sample was divided into two streams. Stream 1 was fixed to a flow rate of 9 L/min and was used to collect samples on 47mm quartz fibre filters (Whatman, Maidstone, UK). 148 Stream 2 demonstrated a fixed flow rate of 2 L/min and was utilised to collect gas phase constituents 149 150 using stainless steel thermal adsorption tubes packed with 1cm quartz wool, 300mg Carbograph 2TD 40/60 (Markes International) downstream of a polypropylene backed PTFE 47 mm filter (Whatman, 151 Maidstone, UK) to remove any particulates. The temperatures at the sampling points were $25 \pm 5^{\circ}$ C. 152 153 A schematic representation of the experimental setup is presented in Figure S1. The sample collected via stream 1 (onto the quartz fibre filters) was classed as particulate, while samples collected via 154 stream 2 (on the adsorption tubes) were classed as gas phase. The deposit collected on the PTFE filter 155 upstream of the adsorption tube was also analysed for particulates and compared to the results 156 obtained from the quartz filters. The results obtained between the two types of filters in these 157 158 experiments were in good agreement and within $\pm 10\%$ across the entire carbon range (C₉-C₃₇), apart from C₁₃ compounds where quartz filters exhibited up to 14% larger concentrations than PTFE filters. 159

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161 **Quantitative and Qualitative Analysis**

Sample deposits were extracted from the filter after addition of deuterated standards by using 162 dichloromethane (DCM), ultrasonicated and concentrated to 50 µL under a gentle flow of nitrogen 163 for subsequent analysis using a 2-dimensional gas chromatograph (GC, 7890A, Agilent 164 Technologies, Wilmington, DE, USA) equipped with a Zoex ZX2 cryogenic modulator (Houston, 165 TX, USA). Samples collected on adsorption tubes were desorbed using thermal desorption (Unity 2, 166 Markes International, Llantrisant, UK) and subsequently analysed using the GC×GC which was 167 interfaced with a BenchTOF-Select, time-of-flight mass spectrometer (ToF-MS, Markes 168 International, Llantrisant, UK) with a scan speed of 50 Hz and a mass range of 30 to 600 m/z. Electron 169

impact ionisation energies were tuned between 10 and 70 eV, the former retaining the molecular ion 170 and the latter causing extensive fragmentation and allowing comparison with standard library 171 spectra³⁴. All data produced were processed using GC Image v2.6 (Zoex Corporation, Houston, US). 172 173 Individual compounds were identified on the basis of retention times (in two dimensions) and mass spectra, and were aggregated into compound groups according to their location in the 2D 174 chromatogram. Quantification was on the basis of ion current after calibration with appropriate 175 standards. Further details are provided in the Supplementary Information. In depth details regarding 176 sample preparation, instrumental parameters, identification and quantification methodologies are 177 presented in detail elsewhere^{18.19}. 178

179

180 RESULTS AND DISCUSSION

Over 500 compounds were identified and quantified, and aggregated into eight groups in the vapour 181 phase and six groups in the particulate phase. On average 83% and 72% of the total ion current was 182 accounted for in the gas and particulate phase sample chromatograms, respectively; in agreement 183 with our previous work¹⁹. The EFs calculated in this study for the three steady state conditions with 184 and without after-treatment devices are shown in Table S2, for the gas and particulate phases of all 185 measured groups of compounds in the range C_9 - C_{37} . The calculation method for the reported EFs in 186 this study is described in S3 and further details are available in a companion paper³⁹. Briefly, the EFs 187 are emission concentrations normalised to the actual (measured) engine fuel consumption (kg.fuel) 188 at each engine operating condition. 189

190

191 Gas Phase Emission Factors – Speeds and Loads

The eight groups with their relative contributions to the total identified organic compounds for all the conducted experiments included; normal (n-) and branched (i-) alkanes (40 - 63%), monocyclic alkanes (17 - 33%), bicyclic alkanes (5 - 11%), monocyclic aromatics (7 - 24%), bicyclic aromatics (2 - 7%), tetralins (1 - 7%), aldehydes and ketones (<1%) and PAH (<1%). Figure 1 illustrates the

EF of compounds from the diesel engine in the gas phase before the diesel oxidation catalyst (DOC) 196 and after the diesel particulate filter (DPF) at different speeds and loads. The most abundant 197 compound class was the n- and i-alkanes accounting for 37-60% and 46-68% of the total identified 198 199 organic compounds between C₁₁-C₁₈ before DOC and after DOC+DPF, respectively. The majority of C_{19} compounds (90%) were n- and i-alkanes and C_{20} - C_{26} compounds were exclusively n-alkanes, 200 although present in relatively low concentrations. This is because the majority of compounds in this 201 volatility range would be expected to be in the particulate phase. The largest total EF of $C_{11} - C_{26}$ gas 202 203 phase compounds were in the order of HS/LL > LS/LL > HS/HL (for both before DOC and after DOC+DPF). This general trend was only observed for the n- and i-alkanes and monocyclic alkanes 204 205 (the two most abundant compound classes). For all other compound classes the largest EF were observed in the order LS/LL > HS/LL > HS/HL. Some bicyclic alkanes (C₁₁ and C₁₅) were 206 significantly larger in LS/LL conditions, factors of 1.8–10.2 larger than other conditions before DOC 207 and after DOC+DPF. Bicyclic alkanes have been shown to be both thermally stable and endothermic 208 fuel components for advanced jet fuels which may give rise to the larger EF for bicyclic alkanes for 209 LS/LL⁴⁰. The presence of bicyclic alkanes in diesel fuel can reduce smoke emissions without 210 sacrifices in brake specific energy consumption⁴¹. The longer ignition-delay of some bicyclic alkanes 211 makes it possible to complete fuel injection early in the combustion stage and thus decreases the 212 213 quantity of fuel directly injected into the hot flame zone which leads to reduced smoke emissions⁴¹.

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As previously reported the gas phase emissions are remarkably similar to diesel fuel while the particulate phase emissions resemble that of the composition of lubricating oil^{18,19}. In this study the S/IVOC gas phase composition between different speeds/loads does not differ much, in contrast to previous studies of IVOC emissions by Cross et al.¹⁵ who reported IVOC EF and chemical compositional changes over a wider range of engine conditions, using a medium duty diesel engine. At low loads, IVOC emissions were dominated by saturated hydrocarbon species similar to unburned fuel, but at higher loads the IVOC emissions became increasingly characterised by unsaturated

hydrocarbons and oxygenated species, possibly formed from incomplete combustion processes¹⁵. In 222 this study a larger relative proportion of classes of compounds other than alkanes contributed to the 223 overall EF for LS/LL in comparison to the HS/HL and HS/LL conditions. The largest gas phase EF 224 225 were present in low load conditions (LS/LL and HS/LL), but did not seem to vary greatly with speed. This may suggest that the contribution of unburned fuel is larger for low load conditions, and 226 improved combustion efficiency is observed for the fuel for higher loads. In a diesel engine, for a 227 given speed, the equivalence ratio, defined as the ratio of the actual fuel/air ratio to the stoichiometric 228 229 fuel/air ratio, normally increases with load. CO, total hydrocarbon (THC) and organic gas phase compounds would expect to typically increase with equivalence ratio; however, other factors such as 230 231 the in-cylinder gas turbulence and temperature, fuel injection timing and pressure calibration and/or number of injection events can also play vital roles and differ depending on a particular engine. 232

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234 Effect of Abatement Devices on Gas Phase Constituents

Figure 1 shows the effect of the presence of a DOC+DPF on the gas phase composition for three 235 236 steady state conditions. In all instances the presence of the DOC+DPF shows a decrease in the EF of gas phase constituents. For HS/LL conditions, measurements were also made after the DOC to assess 237 the efficiency of the DOC and DPF separately. The DOC is expected predominantly to adsorb and/or 238 239 oxidise gas phase constituents, whereas the catalysed DPF is expected to mainly filter particulate phase species and oxidise gas phase species at lower rates. The percentage decreases in the EF for the 240 sum of all species for the three conditions are shown in Figure S3 (before DOC to after DOC+DPF 241 for all conditions; and before DOC to after DOC for HS/LL only). For HS/HL, the low molecular 242 weight (LMW) monocyclic and bicyclic alkanes, tetralins and monocyclic aromatics compounds in 243 the vapour phase are more efficiently removed than high molecular weight (HMW) compounds; 244 however, as the acyclic alkanes represent the majority of the mass, this is not reflected in Figure S3. 245 Alam et al.¹⁸ reported a greater efficiency of the DOC for HS/HL engine steady state conditions for 246 LMW acyclic alkanes in the gas phase. This is not the case for HS/LL (after DOC), or for LS/LL and 247

HS/LL (both after DOC+DPF) in this study where a more uniform decrease in EF as a function of 248 carbon number is observed for acyclic alkanes. The average decreases in the total organic EF for 249 HS/HL, LS/LL and HS/LL are 50%, 56% and 74% using the DOC+DPF, respectively. DOCs have 250 251 been reported as typically reducing emissions of PM by 20-40% or more and gaseous emissions by 50-70%⁴². Figure S3 also shows the percentage decrease of organic vapour phase compounds for 252 HS/LL conditions after the DOC and after the DPF. The EFs of gas phase C_{11} - C_{18} compounds were 253 254 reduced on average by 40% after the DOC and a further 34% after the DPF. The HMW gas phase 255 species (C₁₈-C₂₆) showed a modest 20% average decrease after the DOC and a further 53% after the DPF. These results appear to reflect oxidation of vapour on the catalytic surfaces of the DPF. 256

257

258 Particulate Phase Emission Factors – Speeds and Loads

The six groups with their relative contributions to the total identified organic compounds for all the 259 conducted experiments included; normal (n-) and branched (i-) alkanes (37 - 68%), monocyclic 260 alkanes (5 - 19%), bicyclic alkanes (3 - 10%), monocyclic aromatics (3 - 12%), oxygenated 261 compounds, including alkyl-3(2H) furanones, aldehydes and ketones (13 - 37%) and PAH (<1%). 262 Figure 2 shows the EF of compounds emitted in the particulate phase before the DOC and after the 263 DOC+DPF at the three studied steady state conditions. As for the gas phase, the most abundant 264 compound class was the n- and i-alkanes accounting for 63-65% (before DOC) and 37-68% (after 265 DOC+DPF) of the total identified organic compounds between C₉-C₃₇. The lower average abundance 266 of acyclic alkanes after the DOC+DPF must reflect the efficiency of the DOC+DPF in the removal 267 of these compounds. The majority (>83%) of particulate phase compounds $\geq C_{28}$ were n- and i-alkanes 268 and all measureable compounds $\geq C_{33}$ were exclusively n-alkanes. LMW particulate species with C₉-269 C₁₃ were exclusively oxygenated compounds, present in modest concentrations relative to the 270 aliphatic alkanes for HS/HL (before DOC and after DOC+DPF) and HS/LL (after DOC+DPF), see 271 Figure 2. Some of the most abundant identified oxygenated compounds are presented in 272 Supplementary Information S4. In our previously published work, bimodal peaks were observed for 273

the particulate phase for HS/HL steady state conditions and 2D-chromatography of the particulate 274 phase indicated the presence of two unresolved complex mixtures¹⁸. The LMW UCM was attributed 275 to a diesel fuel component while the HMW UCM indicated emissions from the lubricating oil. 276 277 Previously published measurements using the same techniques have shown that the composition of the same diesel fuel used for this study did not contain oxygenated compounds¹⁹. Thus, the origin of 278 the oxygenated compounds in this study is not from unburned fuel or lubricating oil, but more likely 279 from combustion products. Wagner and Wyszyński⁴³ showed how hydrocarbons from unburned fuel 280 are oxidised to aldehydes and ketones during in-cylinder mixing and in the exhaust manifold. The 281 formation of these oxygenated compounds, however, is dependent upon numerous engine parameters, 282 283 including speed and load, mixture composition, compression ratio, air-fuel ratio and heat losses to the wall/in the exhaust. The larger EF of these oxygenated compounds for HS/HL and HS/LL suggest 284 that the engine parameters are such that they favour their formation in comparison to LS/LL, which 285 is a factor of 3-6 smaller (for before DOC). There is much debate whether diesel fuel composition 286 has a significant effect on the oxygenated hydrocarbon emissions. Some studies suggest low density 287 and low aromatic content diesel fuel produces minimal aldehyde emissions⁴⁴, and others show no 288 notable difference⁴⁵. In this study we observe a notable difference in the emissions of oxygenated 289 compounds with varying speed and load, but do not test the effect of different fuel formulation. 290

291

292 Effect of Abatement Devices on Particulate Phase Constituents

Figure 2 shows the effect of the presence of a DOC+DPF on the particulate phase composition for three steady state conditions. In all instances the presence of the DOC+DPF shows a decrease in the EF of particulate phase constituents and the largest decrease is observed for HS/LL. The percentage decreases in EF of all species for the three conditions are shown in Figure S3 for before DOC to after DOC+DPF and for before DOC, after DOC and after DOC+DPF for HS/LL. The average decrease observed after DOC+DPF for the total organic EF, as a function of carbon number, for LS/LL, HS/HL and HS/LL are 68%, 44% and 89%, respectively. The higher efficiency of the abatement devices for

HS/LL may reflect the composition and concentration of organics, as total EF for HS/LL were a factor 300 of 3.0 and 2.3 larger than HS/HL and LS/LL, respectively. The acyclic alkanes are the main 301 302 constituent in the organic particulate fraction and so the overall efficiency of the DOC+DPF is 303 dependent upon the removal of these species. For LS/LL the decrease in the total EFs of organic species $C_9 - C_{18}$ (which are predominantly in the vapour phase, see Figure S4) was 54%, significantly 304 lower than compounds C₁₉-C₃₇ which was 75%. Although this observation is similar to HS/LL (albeit 305 306 to a lesser extent, see C_9-C_{16} HS/LL, Figure S3), it is in contrast to HS/HL, where reductions in EFs of species after the DOC+DPF was lower (38%) for HMW compound $C_{19} - C_{37}$, see Figure S3. The 307 reduced efficiency of the abatement devices for removing the particulate phase HMW aliphatic 308 309 alkanes for HS/HL may reflect a shift in the volatility distribution or n-alkyl hydrocarbon thermocracking. At elevated temperatures (~250°C) during engine operation n-alkyl substituted 310 naphthenic and naphthalene-aromatic compounds have been reported to undergo mild 311 thermocracking, preferably at the tertiary carbon atom, leading to the production of HMW alkanes, 312 even when the fuels/lubricants were originally deficient in those compounds^{8,46}. The EF of LMW 313 alkanes (C₉- C₁₈) for HS/HL are reduced by 56% and the overall EFs after the DOC+DPF are within 314 25% of LS/LL and HS/LL, indicating similar emissions for LMW alkanes for all steady state 315 conditions. This suggests that the efficiency of the DOC+DPF for removing HMW alkanes is 316 dependent upon driving conditions and is less effective for HS/HL conditions. However, it may also 317 indicate a shift in volatility distribution of the acyclic alkane emissions. Cross et al.¹⁵ did not observe 318 significant differences in IVOC volatility profiles when investigating different loads, and this was 319 attributed to the possibility of different types of IVOC being emitted at differing loads but with the 320 same volatility distributions. In this study, for HS/HL we observe a possibility of a volatility shift for 321 the acyclic alkanes, but in the presence of a DOC+DPF, which may most likely be the inefficiency of 322 DOC+DPF removal of these compounds. Williams et al.⁴⁷ reported that the survival rate of fuel 323 aliphatic components increases with molecular weight, when investigating the role of unburned fuel, 324 which gives rise to the larger relative EFs after DOC+DPF. However, unburned fuel is expected to 325

be a larger fraction of the emissions for lower exhaust gas temperatures and for conditions of low 326 speed/load¹⁷; thus the HMW aliphatic alkanes $(C_{19} - C_{37})$ probably originate from unburned 327 lubricating oil rather than fuel^{18,32,35}. Alam et al.¹⁸ reported <25% particulate phase concentration 328 329 reductions of n-alkanes for HS/HL conditions in the presence of a DOC only, in agreement with the small reductions in EFs observed in this study for the same conditions. Huang et al.¹⁷ reported large 330 reductions in concentrations (>99%) in PM including PAH, NPAH, hopane and sterane emission 331 rates, but also reported reductions of 83-99% during regeneration of the DPF in comparison to 332 emissions without a DPF. During regeneration, exhaust temperatures increase burning off PM 333 accumulated in the DPF, therefore it can possibly increase the emissions of PM and S/IVOC. We do 334 335 not believe that the reduced reduction of HMW alkanes is due to active regeneration in this study, as exhaust back pressure was insufficient to trigger active regeneration. 336

337

The overall reduction in EF of monocyclic alkanes and aromatics for LS/LL, HS/LL and HS/LL are 338 between 60-94% using DOC+DPF. Although the overall decrease in EF for oxygenated compounds 339 340 is 59%, 61% and 82% for LS/LL, HS/HL and HS/LL, respectively, with DOC+DPF, there is an inefficiency of the removal of this compound class by the DOC. An average of 15% reduction in EF 341 was observed for oxygenated compounds C_9 - C_{26} , after the DOC, compared to >81% reduction for 342 343 other compound classes. This increased to 82% removal after the DPF for HS/LL. Thus 40% of the total particulate organic composition, between C₉-C₂₆, after the DOC are oxygenated compounds. 344 This indicates a contribution of oxygenated compounds to primary organic aerosol (POA) from direct 345 vehicular emissions in the absence of a catalysed DPF. Gentner et al.³⁶ suggest that it is unlikely that 346 alkenes, aromatics and oxygenates contribute to POA and although this may be true for vehicles fitted 347 348 with DOC+DPF, the results of this study show that in the absence of a catalysed DPF, oxygenated compounds comprise a significant amount of the overall emitted composition. May et al.¹¹ reported 349 DPF-equipped diesel vehicles with gravimetric PM mass emissions over two orders of magnitude 350 lower than vehicles with no DPF and attributed the mass reduction to EC, which reduced from 351

approximately 200 mg/kg-fuel to less than 1 mg/kg-fuel. Although particulate OC was not specifically targeted in their study, reductions of approximately 70% were observed in OC between vehicles manufactured pre-1994 and in 1994-2003. Zhao et al.¹⁴ reported total-IVOC emissions from on- and off-road diesel vehicles and showed that the magnitude of total-IVOC emissions was substantially lower for vehicles with exhaust after-treatment devices but there was little variation in the composition. This is in broad agreement with the results from this study apart from after DPF for HS/HL conditions where we see a shift from LMW to HMW compound emissions (see Figure 2).

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360 Gas and Particulate Phase Partitioning

Figure S5 illustrates the gas-particle phase distribution of the total characterised organic compounds 361 identified in this study. All compound classes exhibited the same characteristics in all speed/load 362 (with/without abatement devices) conditions investigated. Compounds $\geq C_{21}$ were >90% in the 363 particulate phase and species $\leq C_{16}$ were >90% in the vapour phase. Observations of organic 364 compounds between C_{17} - C_{20} showed varying ratios between the gas and particulate phases, apart 365 from monocyclic alkanes where compounds $\leq C_{18}$ were exclusively in the gas phase and $\geq C_{19}$ were in 366 the particulate phase. Up to 15% of C₁₃ and C₁₄ monocyclic aromatic compounds were also present 367 in the particulate phase. A slight increase in the vapour phase EF was observed for HS/HL between 368 C_{22} - C_{29} peaking at C_{26} , see Figure S4; in agreement with a previous report¹⁸. Tetralins and bicyclic 369 aromatics were only observed in the vapour phase and were below the detection limit of the 370 instrument in the particulate samples. Oxygenated compounds were only measured in the particulate 371 phase and not observed in the vapour phase, most probably due to their lower volatilities. The species 372 identified in the gas phase for all speeds/loads studied are very similar to the diesel fuel composition 373 374 suggesting unburned fuel emissions. However, when looking at the HS/HL particulate phase emissions, the larger relative contribution of oxygenated compounds to the total concentration 375 indicates products of oxidation which partition into the particulate phase (see Figure 2, HS/HL). 376

The total gas and particulate phase EF for the three steady state conditions are shown in Table 1. The 378 EF ratios of total gas to total particulate phase identified for HS/HL, LS/LL and HS/LL are 1.5, 2 and 379 1 (before DOC); and 1.6, 6 and 7 (after DOC+DPF), respectively. This suggests that there is very 380 381 active partitioning between the gas and particulate phases within the DOC+DPF abatement devices, and that the DOC+DPF are relatively ineffective in reducing gas phase emissions of hydrocarbons in 382 the molecular weight range studied. The average decreases in the total gas phase emissions are 42-383 59%, and 44-89% for particulate phase emissions. Zhao et al.¹⁴ reported IVOC composition and EF 384 from on- and off-road diesel vehicles, evaluating the effects of driving cycles and exhaust abatement 385 devices. Little variation in IVOC composition was observed over different cycle tests but the 386 magnitude of the emissions varied significantly for vehicles with or without abatement devices (a 387 factor of 7 to 28 lower with DPF equipped vehicles, depending on driving cycle). In this study the EF 388 are reduced by a factor of 2-4 with DOC+DPF under the steady-state testing conditions. 389

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This study observed a decrease in EF values from 305 mg kg⁻¹ to 85 mg kg⁻¹ for HS/LL to HS/HL, respectively for experiments performed without emission control devices, which is consistent with differences in combustion efficiency at different loads and with previous studies^{14,15,24}. Our results also indicate that speed affects the EF (see Table 1), as the EF for LS/LL are a factor of 2 smaller than HS/LL for engine runs without abatement devices, and 1.2 times smaller in the presence of DOC+DPF.

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The measured partitioning is likely to represent that in the engine exhaust system, and will not reflect that in the atmosphere where the conditions are cooler and more dilute, and the matrix of other absorbent/adsorbent particles is different. Some changes in partitioning may have occurred during sampling, but these are expected to be small due to the very short residence times in the sampling system (ca. 0.5s).

404 Comparison of Emission Factors with Other Studies

In order to conduct a comparison, the emission factors for n-alkanes have been converted from µg/kg 405 fuel to µg/km travelled using a vehicle fuel consumption of 0.06 kg.fuel/km and assuming that the 406 407 engine runs for 33% of time at low load and the remainder at high load. The estimated emission factors appear in Table S3, and can be compared with particle phase data for n-alkanes of C_{19} - C_{26} 408 reported by Charron et al.⁴⁸ for Euro 3 (E3) and Euro 4 (E4) diesel passenger cars with a DOC and 409 Euro 4 with a DOC and DPF. The two vehicles with DOC only tested by Charron et al.⁴⁸ showed 410 markedly different concentrations both peaking a C₂₁ with emission factors (EF) of 31.5 (E3) and 411 5.84 (E4) μ g/km for the C₂₁ n-alkane. This compares with EFs of 29.2 (before DOC) and 4.0 μ g/km 412 413 (after DOC and DPF) in our data (Table S3), suggesting high comparability. Peak abundances in our data (Table 4) are at C₂₂ (before DOC) and C₂₃ (after DOC), the small differences from Charron et 414 al.⁴⁶ probably explicable in terms of engine oil composition. The vehicle fitted with a DPF sampled 415 by Charron et al.⁴⁸ peaked at C_{20} with an EF of 2.8 µg/km with concentrations of all compounds well 416 below those in our data (Table S3) indicating a more efficient emissions control system. Our EF data 417 may also be compared with EF values inferred for mixed vehicle fleets from atmospheric 418 concentration data, as reported by Xu et al.⁴⁹ who measured concentrations on Marylebone Road, 419 London and reviewed data from a Chinese road tunnel⁵⁰ and the Grenoble Ring Road, France⁴⁸. All 420 field-derived EF data show peak abundance at C_{21} - C_{23} and broadly similar concentrations across all 421 field studies, comparable with our lab-derived data for a light duty diesel (Table S3). The on-road 422 vehicle fleet contains light duty gasoline and heavy duty diesel vehicles in addition to light duty 423 diesels and it appears the emissions from the heavy duty diesels counteract the beneficial effects of 424 gasoline vehicles for high molecular weight n-alkane emissions. The close comparison with some 425 426 published studies may be fortuitous due to the limited operating conditions and single engine used in our study. This emphasises the need for studies of further engines or vehicles over full operating 427 cycles. 428

430 DATA ACCESSIBILITY

- 431 Data supporting this publication are openly available from the UBIRA eData repository at
 432 https://doi.org/10.25500/edata.bham.00000328
- 433

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- 439

440 SUPPORTING INFORMATION

- 441 Supporting Information provides further details of analytical protocols, experimental conditions for
- the engine, measured emission factors and the efficiency of abatement devices.
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444 CONFLICT OF INTERESTS

445 The authors declare no competing financial interest.

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641 TABLE LEGENDS

642 Table 1.643

Emission factors of total identified compounds (as OC mass) in the gas and particulate phases for three studied conditions.

644

645 FIGURE LEGENDS

- 646 Figure 1. Gas phase emission factors from the 2.2 L light-duty diesel engine under high speed / high load (top), low speed / low load (middle) and high speed / low load (bottom).
 648 Different coloured bars represent different compound classes identified in the engine exhaust emissions. Black: n-+i-alkanes, red: monocyclic alkanes, green: bicyclic alkanes, purple: monocyclic aromatics, blue: tetralins, orange: bicyclic aromatics. The lighter shades of each colour represent emission factors with DOC and DOC+DPF emission control devices.
- Figure 2. Particulate phase emission factors from the 2.2 L light-duty diesel engine under high speed / high load (top), low speed / low load (middle) and high speed / low load (bottom). Different coloured bars represent different compound classes identified in the engine exhaust emissions. Black: n-+i-alkanes, red: monocyclic alkanes, green:
 bicyclic alkanes, orange: monocyclic aromatics, blue: oxygenated compounds. The lighter shades of each colour represent emission factors with DOC and DOC+DPF emission control devices.

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Table 1. Emission factors of total identified compounds (as OC mass) in the gas and particulatephases for three studied conditions.

	HS / HL			LS /LL			HS / LL		
TOTAL OC (mg/kg-fuel)	Gas	Particulate	Total	Gas	Particulate	Total	Gas	Particulate	Total
None	51.2	33.9	85.1	97.9	49.5	147	154	151	305
DOC							101	33.7	135
DOC+DPF	25.9	16.3	42.2	55.3	9.31	64.6	68.2	9.84	78.1
% Decrease							34	78%	56%
(DOC)							%	7070	5070
% Decrease	400/	520/	500/	4.4.04	Q10/	560/	56	020/	7404
(DOC+DPF)	49%	3270	50%	44 %	01%	50%	%	75%	/ 4 %



Figure 1. Gas phase emission factors from the 2.2 L light-duty diesel engine under high speed / high load (top), low speed / low load (middle) and high speed / low load (bottom). Different coloured bars represent different compound classes identified in the engine exhaust emissions. Black: n-+i-alkanes, red: monocyclic alkanes, green: bicyclic alkanes, purple: monocyclic aromatics, blue: tetralins, orange: bicyclic aromatics. The lighter shades of each colour represent emission factors with DOC and DOC+DPF emission control devices.



Figure 2. Particulate phase emission factors from the 2.2 L light-duty diesel engine under high speed / high load (top), low speed / low load (middle) and high speed / low load (bottom). Different coloured bars represent different compound classes identified in the engine exhaust emissions. Black: n-+i-alkanes, red: monocyclic alkanes, green: bicyclic alkanes, orange: monocyclic aromatics, blue: oxygenated compounds. The lighter shades of each colour represent emission factors with DOC and DOC+DPF emission control devices.