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Fine iron aerosols are internally mixed with nitrate in the urban european atmosphere

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Article

FINE IRON AEROSOLS ARE INTERNALLY MIXED WITH NITRATE IN THE URBAN EUROPEAN ATMOSHPERE

Manuel Dall'Osto, David C.S. Beddows, Roy Michael Harrison, and Burcu Onat

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33 **TOC**



37 ABSTRACT

38 Atmospheric iron aerosol is a bioavailable essential nutrient playing a role in oceanic productivity. 39 Using Aerosol Time-Of-Flight Mass Spectrometry (ATOFMS), the particle size (0.3-1.5 µm), 40 chemical composition and mixing state of Fe-containing particles collected at two European urban 41 sites (London and Barcelona) were characterised. Out of the six particle types accounting for the 42 entire Fe-aerosol population, that arising from long range transport (LRT) of fine Fe-containing 43 particles (Fe-LRT, 54-82% across the two sites) was predominant. This particle type was found to 44 be internally mixed with nitrate and not with sulphate, and likely mostly associated with urban 45 traffic activities. This is in profound contrast with previous studies carried out in Asia, where the 46 majority of iron-containing particles are mixed with sulphate and are of coal combustion origin. 47 Other minor fine iron aerosol sources included mineral dust (8-11%), traffic brake wear material (1-48 17%), shipping/oil (1-6%), biomass combustion (4-13%) and vegetative debris (1-3%). Overall -49 relative to anthropogenic Asian Fe-sulphate dust - anthropogenic European dust internally mixed 50 with additional key nutrients such as nitrate is likely to play a different role in ocean global 51 biogeochemical cycles.

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- 54

55 Keywords: Iron particles; mineral dust; non-exhaust; nitrate; ATOFMS

57 1. INTRODUCTION

Atmospheric particulate matter (PM) has well known adverse effects on human health¹ and 58 atmospheric visibility; and partly compensates climate forcing by greenhouse gases.² Iron (Fe) is a 59 60 major element found in the atmospheric aerosol. Atmospheric Fe has both natural (e.g. aeolian desert dust transport) and anthropogenic (e.g. iron/steel and general industrial activities, traffic) 61 origins.^{3,4} Coarse particles ($PM_{2,5,10}$) are often considered to be related to crustal elements and 62 resuspension of road dust.⁴ However - within the urban environment - Harrison et al.⁵ found that the 63 64 iron concentration within coarse dusts was much greater at roadside sites than at urban background 65 sites and this was considered indicative of road traffic and most probably the vehicles themselves as a source. Gietl et al.⁶ also showed that Fe and other brake wear elements (Cu, Ba, Sb) have very 66 similar particle size distributions at a roadside site, implying a common source. Dall'Osto et al.⁷ 67 found that Fe and Cu together can also be used as a tracer of brake wear also in PM_{2.5} aerosol 68 69 sampled in the urban agglomerate of Barcelona (Spain).

70

In general little is known about aerosol sources of Fe-containing particles in the finer mode $(PM_1)^{8-1}$ 71 ⁹⁻¹⁰ in the urban atmosphere. Although fine aerosol modes contribute little Fe to the total aerosol 72 73 mass, it has much greater potential for respiratory intake, and its involvement in health effects. Fe is 74 a transition metal which is considered to exert toxic effects, which may release free radicals in lung tissues via the Fenton reaction, and are hypothesized to cause cellular inflammation.^{11,12} Iron 75 aerosol particle size influences their reactivity, toxicity and solubility.¹³ Since particle-bound metals 76 77 need to dissolve and become free ions in the lung fluid, particle solubility is considered a major criterion for the bioavailability and therefore toxicity.^{12,14} Atmospheric Fe may be associated with 78 adverse health effects due to mechanisms such as DNA strand breakage and tissue or cell damage.¹⁵ 79 80

81 Fine Fe plays a role also in ocean/climate biogeochemical cycles, and it might actually be more 82 important than coarse Fe-containing aerosol^{16,17}. Briefly, iron is an essential nutrient for all

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83 organisms, used in a variety of enzyme systems, including those for photosynthesis, respiration, and nitrogen fixation.¹⁶ Several studies have argued that the greater solubility at lower dust mass 84 concentration could be due to a larger surface area to volume ratio of the finer dust particles.¹⁷⁻¹⁹ 85 86 However, the size dependence of the fractional Fe solubility (FFS) can at most explain a small part 87 of the measured variability in FFS in atmospheric aerosol samples. The high values of the FFS of > 88 1% in atmospheric aerosol samples are related to a number of atmospheric processes and natural 89 and anthropogenic sources and are clearly not well understood.¹⁷ It is also important to keep in 90 mind that although combustion sources are not important for the total Fe mass compared to natural sources, they can contribute up to 50% of soluble Fe deposited into oceanic regions²⁰⁻²¹. Overall, the 91 92 relationships between iron solubility, oxidation state, and bioavailability are complex and poorly 93 understood.

94

95 Evidence for the importance of such a dust-ocean-climate feedback process is still limited. Whilst the role of desert dust is currently being studied¹⁷⁻²⁰, far less information is available on Fe-96 containing urban anthropogenic dust²². Urban influenced air masses can travel long distances. 97 98 Continental polar air masses originating in Europe travel southwards across the Mediterranean, North Africa and finally reach the tropical Atlantic Ocean²³. Continental polar air masses 99 originating in Portugal and Ireland also travel southwards to the Canary Island waters²⁴. Northern 100 101 European air masses originating in the continent of Europe can travel across the British Isles and 102 further to the North Atlantic Ocean. For example, results from Mace Head (Ireland) show that these continental polar air masses travel to the Atlantic Ocean fairly often, about a third of the time²⁵⁻²⁶. 103 104 Once urban anthropogenic dust is deposited in the ocean, its impact on marine productivity will 105 depend on a number of factors, including the bioavailability of the chemical components present in the dust 3,27,28 . 106

107

108 In order to characterise the sources, transport and transformations of Fe aerosols in the atmosphere, 109 a comprehensive approach that simultaneously characterises single particles for aerodynamic diameter and chemical composition is needed²⁹. The use of single-particle analysis is an important 110 111 analytical tool because it allows determination of how the chemical constituents are distributed 112 between individual particles (mixing state). Specifically, in this work we are interested in 113 apportioning which secondary chemical components (i.e. nitrate, sulphate) are mixed within Fe 114 containing aerosols; because this can have implications for solubility and additional nutrient capability³⁰⁻³². We use Aerosol Time-of-Flight Mass Spectrometry (ATOFMS), which provides 115 116 information on the abundance of different types of aerosol particles as a function of particle size with a much high time resolution and better count statistics²⁹. For the purpose of our work, the 117 118 ATOFMS is more suitable than other single particle techniques. For example, Transmission 119 electron microscopy (TEM) is a powerful technique capable of simultaneously obtaining unique 120 information on individual particles regarding metal speciation, size, mixing state, and morphology $^{33-34}$. However, semi-volatile particle types (i.e. nitrate) are liable to vaporise in the high 121 vacuum environment of the TEM and therefore not be observed³⁵. 122

123

124 A few specific studies reporting the sources and mixing state of iron-containing particles can be found in the literature. Furutani et al.³⁶ utilized single particle mass spectrometry to classify Fe-125 126 containing particles based on their chemical associations. Fe-containing particles have numerous 127 sources, especially anthropogenic sources such as coal combustion. Indeed, mineral dust was found 128 to be a minor contributor to the total iron concentration, whereas the importance of the coal combustion source in East Asia was emphasised. Complementary measurements^{21,36} performed at 129 the same time and location¹⁸ showed that field-collected atmospheric Fe-containing particles were 130 131 found to be internally mixed with secondary species such as sulfate, soot, and organic carbon. Recently, Zhang et al.³⁷ showed ATOFMS results collected in Shanghai (China) for nearly 22 d 132 133 during the winter of 2011. Fe-containing particles were mainly clustered into four chemical groups,

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134 comprising Fe-rich, K-rich, Dust and V-containing particle types. It was concluded that major 135 sources of Fe-containing particles correspond to various anthropogenic sources including iron/steel 136 industrial activities, and fly ash from both biomass burning and coal combustion, accounting for 55% and 18%, respectively. Hu et al.³⁸ also stressed that sources of Fe-bearing particles in the 137 138 Beijing atmosphere (China) are mainly local steel industries and oil fuel combustion. As a 139 consquence, the majority of the Fe-containing particles collected in Asia are found to be mainly internally mixed with sulphate ^{21,36-38}. With regard to other study areas, to our knowledge there are 140 141 not specific ATOFMS studies on the matter.

142

143 In summary, whilst coarse Fe-containing particles $(PM_{2,5-10})$ are often associated with mineral dust, 144 road dust and non-exhaust traffic material, little is known of the sources of finer Fe-containing 145 particles. Furthermore, to our knowledge all aerosol mixing state studies on Fe-containing particle 146 studies presented in the literature discuss ambient data collected in Asia. This study reports 147 information on sources and mixing state of Fe-containing particles sampled over the European 148 continent. Both meteorology and emissions show large variations across Europe, and conclusions 149 from a certain region cannot necessarily be extrapolated to other parts of the continent. Latitude variations within Europe influence mineral matter³⁹. Saharan dust intrusion⁴⁰, secondary inorganic 150 aerosol⁴¹ and sea salt concentrations⁴². For this reason, we consider two different ambient ATOFMS 151 152 datasets from Barcelona (Spain) and London (England). Whilst both monitoring sites are classified 153 as "roadside" sites, the Spanish one presents a minor road in an environment subject to more Saharan dust intrusion in a drier climate.²⁶ By contrast, the monitoring site in London is near a 154 major road under typical North European weather conditions.⁴⁴ We use two road side sites cause we 155 want to apportion both local road traffic²² and regional long range transport sources. Further 156 157 ATOFMS studies previously published in other background European locations (Mace Head, 158 Ireland; Po Valley, Italy and Port Talbot, Wales) are also compared with the presented results. 159 Different aerosol sources are inferred and discussed, including their average aerosol sizes and mixing state. An important comparison with existing ATOFMS data collected in Asia is alsopresented, and its implication for marine biogeochemistry discussed.

162

163 **2. METHODS**

164 **2.1** Locations

165 This study aimed to elucidate the Fe-containing aerosol sources and their mixing state in two 166 different European cities. The first was London - which has a population of 7.6 million, whilst the 167 Greater London metropolitan area has between 12.3 and 13.9 million, making it the largest in the European Union.⁴⁵ The sampling campaign was conducted at Marylebone Road (51.52°N, 0.15°W), 168 169 a six-lane heavily trafficked road within an urban canyon, between 22nd May and 11th June 2009. 170 Instruments were installed in a permanent monitoring station located at kerbside, 1m from the busy 171 six-lane highway, Marylebone Road, carrying ca 80,000 vehicles per day. Inlets were sited about 3 172 m above ground level (a.g.l.) at differing distances from the road, mostly 1-3 m distant. The second 173 city was Barcelona, capital of Catalonia, a large coastal urban agglomerate located in the Western 174 Mediterranean Basin in the north-east (NE) part of Spain. The sampling campaign (SAPUSS, Solving Aerosol Problems by Using Synergistic Strategies)⁴² took part in Barcelona between 20th 175 176 September and 20th October 2010 (local time, UTC+2). The Road site (lat 41.388423; lon 177 2.150213) was situated in a car park next to a major road (Carrer Urgell). The road, which cuts the 178 city from South East to North West, is a street canyon composed by a two-way cycle path and a 179 one-way four lane vehicle road. Vehicle intensity for the month of measurements was about 17,000 180 vehicles per day. Air was drawn from outside the instruments throughout a number of home-181 designed inlets, situated on the vans' roofs at a height of about 4m above ground with 1/4 inch 182 stainless steel tube led inside each trailer (length about 2 m) to a manifold.

183 With regard to air mass back trajectory analysis run for the two ambient field campaigns herein 184 presented, it was shown in previous studies that air masses travelling over the study areas continue to travel and reach nearby and far oceans and seas about a quarter of the ambient field study
 time^{43,46}

187

188 2.2 Instrumentation

189 Since its introduction in the late 1990s, the ATOFMS has given valuable insights into the size and composition of individual airborne particles.^{47,48} The ATOFMS (Model 3800-100, TSI, Inc.) 190 191 collects bipolar mass spectra of individual aerosol particles. Ambient aerosol is focused into a narrow particle beam for sizes between 100 nm and 3 µm. Using a 2-laser velocimeter, particle 192 193 sizes are determined from particle velocity after acceleration into the vacuum. In addition, the light 194 scattered by the particles is used to trigger a pulsed high power desorption and ionization laser 195 which evaporates and ionizes the particle in the centre of the ion source of a bipolar reflectron ToF-196 MS. Thus, a positive and a negative ion spectrum of a single particle are obtained. The mass 197 spectrum is qualitative in that the intensities of the mass spectral peaks are not directly proportional 198 to the component mass but are dependent on the particle matrix, the coupling between the laser and 199 the particle and the shot to shot variability of the laser. However, the ATOFMS can provide 200 quantitative information on particle number as a function of composition; providing a measure of all 201 particle components and can be used to assess mixing state.

202

Depending on the scientific objectives, past studies have typically used the aerodynamic lens system to study ultrafine and fine anthropogenic emissions⁴⁹ or the nozzle orifice system to study dust particles.²¹ In this study we focus on fine aerosol, so we chose the aerodynamic lens system configuration for better count statistics. Additionally, it is important to note that this study uses the same ATOFMS aerodynamic lens configuration used in past ATOFMS studies characterising Fecontaining particles in Asia.^{21,36,37}

209

210 Fe-containing particles commonly produce an intense ion peak at mass-to-charge ratio (m/z) 56 in the positive ion mass spectrum. As discussed in Zhang et al.³⁷, simply using the ion peak at m/z 56 211 as a unique marker for Fe is not suitable, since it might also be caused by the presence of CaO^+ , 212 KOH^+ , or $C_3H_4O^+$ from other components. There are also other contributors (mainly organic) at 213 m/z = 54. Therefore, as previously described^{36,37} the natural isotopic composition of Fe (⁵⁴Fe/⁵⁶Fe) 214 was adopted as a basis for the Fe screening process to minimize the interference from other species. 215 216 A peak-area ratio 56 Fe/ 54 Fe > 10 was finally applied as recommended by Zhang et al.³⁷ to strengthen 217 the screening through excluding more ambiguous assignments. The screened Fe-containing 218 particles were subsequently imported into YAADA (Yet Another ATOFMS Data Analyzer) and 219 single-particle mass spectra were grouped with adaptive resonance theory neural network, ART-2a.⁵⁰ The parameters used for ART-2a in this experiment were: learning rate 0.05, vigilance factor 220 0.85, and 20 iterations. Further details of the parameters can be found elsewhere.³⁷ An ART-2a area 221 222 matrix (AM) of a particle cluster represents the average intensity for each m/z for all particles 223 within a group. An ART-2a AM therefore reflects the typical mass spectra of the particles within a 224 group.

225

226 **3. RESULTS**

Overall, 890,873 and 693,462 ATOFMS mass spectra were apportioned in Barcelona⁴³ and 227 London⁴⁶, respectively. The two datasets were merged and the resulting total number of particles 228 229 (1,584,335) were queried for Fe following the methodology explained in section 2.2, and a total 230 number of 10,549 particles (6,375 in Barcelona, 4,174 in London) were found. By running ART-2a 231 on the combined dataset, more than 300 clusters were found initially but many were merged if they presented similar temporal trends, size distributions and similar mass spectra.⁵¹ By merging similar 232 233 clusters, the total number of clusters describing the whole database was reduced to six, which are 234 described in the following sub-sections.

3.1 Regional Fe-Containing Particles from Long Range Transport (Fe-LRT)

237 This was found to be the most frequently occurring cluster (54-82%, Table 1). The average ATOFMS Art-2a cluster (Fig. 1a) shows strong signals at m/z 54 and 56 (⁵⁴Fe and ⁵⁶Fe. 238 239 respectively) and iron oxide ([FeOH]⁺, m/z 73). It is found internally mixed with nitrate (m/z -46 240 and -62 in the negative spectra). This particle type was mainly distributed in the fine mode peaking 241 at around 0.4-0.7 µm in aerodynamic diameter (Fig. 2). In Barcelona, Fe-rich particles were found 242 to correlate with ATOFMS Long Range Transport (LRT) nitrate associated with long range 243 transport of pollutants⁴³. The small mode of this particle type reflects the fact that only fine particles 244 were likely to travel long distances relative to the coarser ones which were lost during transport. 245 Likewise, in London they were associated mainly with air masses arriving from the European 246 continent.

247

248

8 **3.2** Fe-containing Particles from Mineral Dust (Fe-DUST)

249 A significant Fe contributor to global tropospheric particulate matter is suspension of soil dusts or 250 other crustal material. The typical ATOFMS mass spectra of a dust particle type rich in Fe is shown 251 in Fig. 1b. Peaks associated with sodium (m/z 23), magnesium (m/z 24) and potassium (m/z 39) can 252 be seen. In addition, peaks for calcium (m/z 40) and aluminium (m/z 27) are present. The negative 253 spectra also show a dominant set of peaks occurring at m/z = -16, -17, -63 and -79. These indicate 254 the presence of O⁻, OH⁻, PO₂⁻ and PO₃⁻, respectively. Peaks of silicate (SiO₂⁻ and SiO₃⁻) at m/z 60 255 and 76 (respectively) are also detected. This particle type was found to be the second most abundant 256 (8-11% of the total Fe-containing particles). It presented a size mode in the largest ATOFMS size 257 range at about 1.4 µm, and these were found to be the most coarse particles detected in both cities. 258 A number of studies have focused on the long-range transport of Saharan air masses over Western and Eastern North Atlantic areas⁵² and across the Mediterranean.⁵³ For example, titanium rich 259 260 particles (along with aluminosilicate) originated from the Azores high pressure region which draws air from North Africa were already identified by the ATOFMS.⁵⁴ However, during this analysis we 261

were not able to separate a specific mineral dust attributable to Saharan mineral dust, leaving this

263 ATOFMS particle type classified as general mineral dust.

264

265 **3.3 Fe-containing Particles from Brake Wear (Fe-Brake)**

266 Fig. 1c shows the typical mass spectra of single particles associated with brake wear material originating from non exhaust traffic brake wear material.⁵⁵ Many brake linings contain high metal 267 268 concentrations, although there is also a large variation between different types and different brands. 269 However, the list of ingredients contained in European brake pads, notably include metallic copper, 270 steel and iron oxide, and barites, which is highly consistent with the observed correlation between Cu, Ba, and Fe.¹² The characteristic features of brake wear particle types include intense [Fe]⁺ and 271 272 $[FeO_2]^-$ (m/z 56 and -88) signals, along with $[Ba]^+$ and $[BaO]^+$ (m/z 138 and 154). Moreover, it was noted that FeO_2^- (m/z -88) is a good indicator of brake dust particles.⁵⁵ Peaks due to copper ([Cu]⁺, 273 274 m/z 63 and 65) were seen in about 30% of the particles analysed. This particle type was mainly 275 found in the supermicron mode (>1 μ m). Additionally, it was found to be a major contributor near 276 the major road in London (17%) but not in Barcelona (1%, Table 1). This is likely due to the much higher number of cars passing at the London site (80,000 day⁻¹) than the Barcelona site (17,000 day⁻¹) 277 ¹). 278

279

280 **3.4** Fe-Containing Particles from Oil Combustion (Fe-V)

Field observations have confirmed that ships produce significant amounts of soot, vanadium, nickel, and sulfate.⁵⁶ ATOFMS studies often report a very specific particle type containing vanadium (m/z $51 [V]^+$ and m/z 67 $[VO]^+$)⁵⁷⁻⁵⁸, as shown in the Art-2a cluster detected in this study (Fig. 1d). This particle type was found mainly internally mixed with nitrate (m/z -46, m/z -62) and in minor amount with sulphate (HSO₄⁻, m/z -97). This class accounted for only 1-6% of the particles characterised, with a fine aerosol unimodal distribution peaking at about 500 nm. V-containing particles have been primarily attributed to residual fuel oil combustion, associated with sources such

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as ships and refineries in the USA.⁵⁷ Likewise, the recent study of Zhang et al.³⁷ also reported this
particle type in Shanghai, which is a coastal mega-city with relatively high density of shipping.
Using single - particle online chemical analysis data, Ault et al.⁵⁷⁻⁵⁸ showed that V-containing
aerosols emitted from ships could contribute a large fraction of atmospheric aerosols in the
California basin.

293

3.5. Fe-containing Particles from Combustion Mixed with Sulphate (Fe-SUL)

295

296 The mass spectrum of this particle type (Fig. 1e) shows the presence of iron (m/z 56) and potassium (m/z 39). The unique feature in the positive mass spectra is the strong peaks at m/z 27 $[C_2H_3]^+$ and 297 m/z 43 $[(CH_3)CO]^+$ usually associated with oxidized secondary organic aerosol.⁵⁹ Additionally, the 298 299 signal of sulphate (m/z - 97) is the strongest of all the ATOFMS particle types described in this 300 study, hence the name characterising this particle type. This particle type contributed only 4-13% of 301 the total Fe-containing particles analysed. A key feature of particle type Fe-SUL is that it is 302 distributed in the smallest size range of the Fe-containing particles. As seen in Fig. 2, a strong mode 303 at about 350 nm can be seen, representing more than 80% of the total Fe-containing particles of this 304 size detected in this study. Overall, the finest size mode and the sulphate mixing state suggests this is a combustion type particle. This particle type was similar to the ones detected in Asia^{36,37} and 305 306 associated with fly ash, coal and biomass burning. The frequent association of Asian dust single 307 particle mass spectra with peaks due to lithium (m/z 7, Li^+) and to potassium (m/z 39: K^+) was 308 indicative of a large contribution from coal combustion since biomass burning aerosols produce negligible Li ion peak^{36,60}. In contrast, the absence of signal from Li in single particle mass spectra 309 310 reported in Fig. 1e suggests that the Fe-SUL particle type is more related to biomass burning than 311 coal combustion.

312

313 **3.6** Fe-containing Particles from Leaves/Vegetative Debris (Fe-VEG)

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314 The ATOFMS has already proved to be a good tool able to separate dust (mainly Ca-rich or Al-Si rich) from biological particles⁶¹. The Art-2a cluster of Fe-VEG is shown in Fig. 1f. In addition to 315 Fe, a very strong potassium signal (K^+ at m/z 39 and 41) is attributed to biogenic sources.⁶² 316 317 Additionally, the biogenic plant debris origin is also supported by the presence of phosphate (m/z -63, -79), as suggested in previous studies.⁵¹ This cluster was found in minor abundance (1-3%, 318 319 Table 1), and presented a broad size distribution (Fig. 2, 51% super-µm, 49% sub-µm). K-rich particles have previously been attributed to biomass burning^{60,63} and have been found to correlate 320 321 with gas-phase measurements of acetonitrile, a good biomass-burning tracer. However, the presence of peaks at m/z 39 $[K]^+$, m/z 113 $[K_2CI]^+$ and m/z -35 $[CI]^-$ suggesting a combustion source related 322 323 to biomass burning are not seen in the particle mass spectra, excluding such a source.

324

325 4. **DISCUSSION**

326

327 This work aims to better understand the aerosol sources of fine Fe-containing particles and their 328 aerosol mixing state in the European atmosphere. Single particle mass spectrometry (ATOFMS) 329 was used to acquire more than 1,500,000 positive and negative mass spectra of single particle of 330 known size. It is important to stress the fact that whilst the ATOFMS provides important 331 information on the mixing state of the aerosols and temporal trends of unique particle types over 332 time, the total number of particles sampled by the instrument does not have quantitative meaning 333 without extensive calibrations as it is biased by transmission efficiency⁶⁴. Nevertheless, the 334 ATOFMS can precisely size particles within the $0.3-1.5 \,\mu m$ range. The six particle types identified 335 were mostly distributed in the submicron mode (<1 μ m): Fe-LRT (72% of the particles were 336 smaller than 1 µm), Fe-SUL (99%) and Fe-V (76%). Vegetative dust (Fe-P) was equally distributed (51% submicron, 49% supermicron). By contrast, mineral dust and brake dust were found more 337 338 abundant in the supermicron (>1 µm) size range: 82% and 73%, respectively, as found in previous studies^{5,6}. 339

340 This study - unique in the aspect that it analyses submicron Fe-containing particles at very high time 341 resolution at two European sites - suggests that the main contribution of submicron iron particles by 342 particle number is related to regional and long range transport of aerosol. The most abundant Fe-343 containing particle type (54-82%) was found to be associated with nitrate, and the absence of 344 mineral dust peaks (Al, Ti, Ca, Si) suggests an anthropogenic urban-industrial source. Fe-containing 345 particles from long range transport have a size distribution peaking at about 600 nm. Previous 346 studies in the same location in London support our findings. Charron et al.⁶⁵ found that the regional 347 background aerosol was the largest contributor measured at Marylebone Road between January 348 2002 and December 2004. Very often these episodes were associated with air masses coming from 349 mainland Europe which carried large concentrations of secondary aerosol adding particulate mass to 350 primary emissions from local urban sources around London.

351

352 By contrast, local sources were found to contribute only in small percentages to Fe aerosol particle 353 concentrations. In urban areas, local road traffic is recognised as an important source of particulate 354 matter (PM). Numerous studies have been conducted regarding traffic-emitted pollutants, with most 355 investigations focusing on exhaust emissions. The most important direct emission sources are the 356 wear of tyres, brakes, and road surfaces. None of the six Fe-containing particle types detected at the 357 two road sites used in this study correlated temporally with the traffic counts. Meteorological data 358 was examined for correlation with aerosol types. However, only a weak correlation between brake wear (Fe-Brake) and wind speed ($R^2 = 0.25$ and 0.41 at LND and BCN, respectively) was observed. 359 360 As expected, mineral dust was found to be a major source of Fe-particles in both Barcelona and 361 London. However, an interesting finding of this work is that at the heavily trafficked London road 362 site, brake wear was found to contribute a higher percentage by number (17%) than mineral dust 363 (8%). Another minor anthropogenic source containing iron was associated with the marine 364 transport sector (ATOFMS Fe-V), which is especially polluting due to the use of low-quality 365 residual fuels containing high amounts of sulphur and heavy metals. Concentrations of V are

366 elevated in the Mediterranean region owing to increased consumption of fuel oil for power generation, shipping and industrial emissions^{66,67}. In this study the ATOFMS Fe-V particle type 367 was found to be highly correlated with Fe-LRT (temporal $R^2 = 0.85$), suggesting that this particle 368 369 type is also primarily related to long range transport rather than to local shipping from the nearby 370 ports of Barcelona and London. With regards to other combustion sources, it is important to 371 recognise that particles associated with biomass burning/coal combustion contribute only 4-13% of 372 the particles sampled at the two European sites. This is in profound contrast to studies of the same 373 type conducted in Asia, where comparing the chemical composition and spatial distribution of iron 374 within ambient particles and standard Asian mineral dust, it was determined that the main source of field collected atmospheric Fe-containing particles was coal combustion.^{36,37} 375

376

377 One limitation of this study is that the ATOFMS uses a laser desorption/ionization (LDI). Such 378 process is heavily influenced by particle size, morphology and matrix composition, since these will 379 influence energy transfer from the laser beam to the particle, vaporization of the particle and ion 380 formation in the vaporization plume. As a result, whilst the ATOFMS can provide the precise 381 aerodynamic diameter of individual particles, the chemical information obtained by this type of 382 single particle mass spectrometry can only be treated as qualitative, or semi-quantitative after a complex calibration⁶⁸. Luckily, there are a number of filter-based off line studies that have focused 383 384 on the aerosol mass source apportionment of Fe in the PM1 and PM2.5 aerosol size fraction. 385 Interestingly, in the coarse fraction (PM > $2.5 \,\mu$ m) Fe is usually associated with mineral dust, road 386 dust and traffic related matter. However, the source distribution of Fe in the finer size fraction is not clear at all. In the city of Barcelona, Amato et al.⁶⁷ reported that major sources of PM_{2.5} Fe are due 387 to mineral and industrial activity. However, about 15-30% of the PM1 and PM2.5 Fe was attributed 388 to secondary more aged material not of local origin. Cusack et al.⁸ reported a detailed source 389 390 apportionment of PM_1 at a regional background site in the western Mediterranean. It was found that 391 about 45% of PM₁ iron was associated with secondary sulphate and secondary organic aerosol.

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Perrone et al.9 reported almost two years of data of PM2.5 and PM1 samples collected at a central 392 393 Mediterranean site. It was found that 60% of the aerosol mass PM_{2.5} occurred in the PM₁ range. 394 The PM source identification showed that Fe in the PM_{2.5} was distributed almost equally between 395 four sources: reacted dust, heavy oil combustion, secondary marine and traffic. Within the PM₁ mass region, Perrone et al.⁹ reported that about 60% of the Fe was related to long range transport 396 397 sources: nitrate with reacted dust, secondary marine and ammonium sulphate. During the same 398 SAPUSS field campaign used in this work, Dall'Osto⁷ reported hourly metal concentrations. PM_{2.5} 399 Fe was higher at the road site than the urban background site, stressing the importance of traffic 400 activities. About 70% was related to non-exhaust traffic activity, and the rest to industry (about 401 10%) and dust (about 20%). However, nitrate was not detectable by the analytical technique used 402 (PIXE), so the results from this study cannot readily be compared. Fewer studies are available in London, although recently Visser et al.¹⁰ reported a very detailed study of the kerb and urban 403 404 increment of highly time-resolved trace elements in PM10, PM2.5 and PM1 aerosol in London. While 405 the $PM_{2.5-10}/PM_{0.3-1}$ Fe aerosol mass ratio at the road site was found to be 7.7, it was less at the 406 background sites (5.5 and 5.6), suggesting the higher importance of fine Fe-containing aerosols 407 away from traffic sources. Indeed - regardless of the local impact of vehicles at each monitoring site 408 - this study shows that most of the fine Fe-containing particles detected are related to long range 409 transport.

410

Another interesting finding is that the major particle type characterised in this study - related to regional and long range transport - is internally mixed with nitrate. Our results are in profound contrast with previous important work carried out in Asia, where Fe-containing particles were found to be internally mixed with secondary species such as sulphate, soot, and organic carbon, but not nitrate.^{21,36,37}. It is relevant that within the European Union (EU), in the last decades much lower SO₂ levels are being recorded following strict implementation of industrial pollution controls⁶⁹. EU abatement of traffic related NO_x levels is still required to maintain levels within international

standards⁶⁹. Control actions in Asia have also been taken, although their effectiveness is outweighed 418 by increased energy demand, with larger amount of SO_2 from coal relative to Europe⁷⁰⁻⁷¹. The fact 419 420 that our detected long range transport Fe-containing aerosols are internally mixed with nitrate is not 421 surprising given the higher amount of NO_x relative to SO_2 in the urban atmosphere, whereas the 422 latter has much higher concentrations in Asia. As regards of the seasonality of the studied main Fe 423 particle type (Fe-LRT), the two field studies herein reported were carried out in spring (London) 424 and fall (Barcelona). Whilst previous analysis show a reduction due to evaporation of nitrate during 425 warm summer months, the evolution of nitrate can be heavily influenced by meteorological factors⁷². Given the likely urban anthropogenic source of fine iron-nitrate particles, we speculate the 426 427 mixing state may not change dramatically across most of the year.

428 We looked at iron containing particles sampled by ATOFMS at other European sites. A previous 429 study of the size distributions and the chemical characterization of airborne particles in the vicinity of a large integrated European steelworks⁷³ found that Fe-rich particles were mainly related to 430 431 emissions from the blast furnace as a result of furnace tapping and slag quenching operations. It was 432 found that Fe-rich particles are internally mixed with nitrate (and very little sulphate), reporting 433 mass spectra similar to those shown in Figure 1a, suggesting a contribution of this source to the 434 ATOFMS Fe-LRT particle type. Data sets from two other studies carried out at Mace Head, Ireland²⁶ and in the Po Valley, Italy⁷⁴ were queried for fine Fe-containing particles. It was found 435 436 that most of the fine Fe containing particles are of anthropogenic origin and internally mixed with 437 nitrate - consistent with this study. However, Fe-containing particles represented less than 0.5% of 438 the total sampled particles.

439

We did not study the solubility of the detected particles. Generally, Fe-nitrate chemical species are usually more soluble than Fe-sulphate species. Baker and Jickells¹⁸ reported an inverse relationship between dust mass concentration and FFS, arguing that the greater solubility at lower dust mass concentration could be due to a larger surface area to volume ratio of the finer dust particles.

However, later studies¹⁷⁻¹⁹ reported that the size dependence of the FFS can at most explain a small 444 445 part of the measured variability in FFS in atmospheric aerosol samples. Overall, natural Fe-dust is generally found in the coarser mode than anthropogenic urban related Fe-dust²². While no 446 447 conclusions can be drawn on aerosol solubility, this study shows that the main type of fine Fe-448 containing particles in the urban atmosphere are internally mixed with nitrate, hence of different 449 chemical composition of natural Fe-dust. Additionally, it is important to remember that while 450 sulphate is not considered a nutrient for marine microorganisms, nitrate is a key one, alongside iron. 451 Atmospheric reactive nitrogen is mainly derived from anthropogenic combustion or agricultural sources in densely populated regions throughout the world.³⁰ The supply of new nutrients to the 452 453 ocean from external sources such as atmospheric deposition has been extensively addressed in ironlimited High Nutrient-Low Chlorophyll (HNLC) regions,³¹ most of which receive low atmospheric 454 inputs at the present time.³² How the ocean responds to pulses of deposition - as either transient or 455 long-term impacts on diversity of the natural assemblage and/or carbon export - is not obvious. 456 Guieu et al.⁵⁸ recently reviewed the significance of the episodic nature of atmospheric deposition to 457 458 low nutrient LNLC regions. Most of the experimental studies presented used desert dust but few 459 considered its solubility and hence bioavailability. This is particularly important because desert 460 dusts derived from soils contain very little soluble nitrogen whereas "atmospherically processed 461 dust" is rich in nitrogen. It was stressed that differences in the source of the material used (e.g., soil 462 vs. locally collected aerosols), related composition, and solubility may partially explain the 463 observed variability in response. Deposition of iron, nitrogen, phosphorus and organic matter has increased considerably since preindustrial times³⁰ and nitrogen could further slightly increase in the 464 future (Guieu et al.³² and ref. included). The present study shows that anthropogenic Fe-containing 465 466 particles also contain nitrate, and the interactive effects and the possible synergy of both nutrients is 467 not currently accounted for in ocean global biogeochemical cycles. The solubility of Fe in 468 atmospheric aerosols, model soils, and desert samples has been investigated thoroughly in various ocean and atmospheric environments during the past decade,¹⁷⁻¹⁹ illustrating the complex nature of 469

470 iron solubility in atmospheric aerosols. This study shows that the majority of fine Fe-containing 471 particles collected in urban European environment are internally mixed with nitrate, and it is 472 important to keep this in mind when studying both the health effects of urban aerosol and the 473 impact of iron in LNLC-HNLC oceanic regions impacted by long range transport of European 474 anthropogenic aerosol.

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789	TABLE LEGEND					
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791 702	Table 1:	Major types of Fe-containing particles identified by the ATOFMS, and their abundance in the Baraglana (BCN) and London (LND) studies				
792		abundance in the Barcelona (BCIV) and London (LIVD) studies.				
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795	FIGURE L	EGENDS				
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797 798	Figure 1:	Positive and Negative ART-2a area vectors attributed to (a) Fe-LRT, (b) Fe-Dust, (c) Fe Brake (d) Fe V (e) Fe SUL and (f) Fe P				
790		(c) 1^{-1} 1^{-1} 1^{-1} 1^{-1} 1^{-1} 1^{-1} 1^{-1} 1^{-1} 1^{-1} 1^{-1}				
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801	Figure 2:	Size distributions of the six different particle types detected at the two European				
802	8	sites. It is clear that combustion Fe-SUL are in the fine mode, the nitrate-Fe in the				
803		accumulation mode, and the Fe-Brake and Fe-Dust in the coarser mode. Fe-V in the				
804		fine combustion mode, Fe-P (vegetative debris) is bimodal in both fine and coarse				
805		modes.				
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- 820 Table 1: Major types of Fe-containing particles identified by the ATOFMS, and their abundance in
- the Barcelona (BCN) and London (LND) studies. In the text the interval of the % are given as %
- across the two monitoring sites (BCN and LND, i.e. Fe LRT is 54-82%).

	Total particle		%	
ATOFMS particle type	BCN	LND	BCN	LND
Fe_LRT	5221	2254	82	54
Fe_Dust	664	344	11	8
Fe_Brake	41	695	1	17
Fe_V	61	244	1	6
Fe_SUL	282	524	4	13
Fe_P	106	113	1	3
Total	6375	4174	100	100







Figure 2: Size distributions of the six different particle types detected at the two European sites. It is clear that combustion Fe-SUL are in the fine mode, the nitrate-Fe in the accumulation mode, and the Fe-Brake and Fe-Dust in the coarser mode. Fe-V in the fine combustion mode, Fe-P (vegetative debris) is bimodal in both fine and coarse modes.

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