CHARACTERISTICS OF TYRE DUST IN POLLUTED AIR: STUDIES BY SINGLE PARTICLE MASS SPECTROMETRY (ATOFMS)

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Abstract

There is a paucity of quantitative knowledge on the contributions of non-exhaust (abrasion and re-suspension) sources to traffic emissions. Abrasive emissions can be broadly categorised as tyre wear, brake wear and road dust/road surface wear. Current research often considers road dust and tyre dust as externally mixed particles, the former mainly composed of mineral matter and the latter solely composed of mainly organic matter and some trace elements. The aim of this work was to characterise tyre wear from both laboratory and field studies by using Aerosol Time-Of-Flight Mass Spectrometry (ATOFMS). Real-time single particle chemical composition was obtained from a set of rubber tyres rotating on a metal surface. Bimodal particle number size distributions peaking at 35 nm and 85 nm were obtained from SMPS/APS measurements over the range 6-20,000 nm. ATOFMS mass spectra of tyre wear in the particle size range 200-3000 nm diameter show peaks due to exo-sulphur compounds, nitrate, Zn and ions of high molecular weight (m/z > 100) attributed to organic polymers. Two large ATOFMS datasets collected from a number of outdoor studies were examined. The former was constituted of 48 road dust samples collected on the roads of London. The latter consisted of ATOFMS ambient air field studies from Europe, overall composed of more than 2,000,000 single particle mass spectra. The majority (95%) of tyre wear particles present in the road dust samples and atmospheric samples are internally mixed with metals (Li, Na, Ca, Fe, Ti), as well as phosphate. It is concluded that the interaction of tyres with the road surface creates particles internally mixed from two sources: tyre rubber and road surface materials. Measurements of the tyre rubber component alone may underestimate the contribution of tyre wear to concentrations of airborne particulate matter. The results presented are especially relevant for urban aerosol source apportionment and PM$_{2.5}$ exposure assessment.
Keywords: Tyre dust; road surface wear; non-exhaust emissions; road traffic; ATOFMS

1. INTRODUCTION

Air quality management is an essential aspect of environmental policy. In 2005, the European Commission published its Thematic Strategy on Air Pollution under the 6th Environmental Action Programme: The Clean Air For Europe (CAFE, 2005) programme, aiming to establish a long-term policy strategy targeting the adverse effects of air pollution on human health and environment. Vehicular traffic is known to be one of the major, if not the largest, source of primary atmospheric particulate matter (PM) in the urban environment (EEA, 2009).

Road traffic emissions comprise not only tailpipe exhaust emissions but also non-exhaust emissions derived from the vehicle-induced resuspension of dust deposited on the road, and from the direct emissions from vehicle wear (brakes, tyres, discs etc.). Road dust is comprised of soil and materials that arise from mechanical abrasion including wear of pavement, tyres, brakes and clutches, as well as corrosion of chassis, bodywork, and other vehicle components; and corrosion of street signs, guiderails, and fencing (Luhana et al., 2004). Road dust is a chemically complicated mixture with a multitude of sources, including natural sources (e.g., soil, plant fragments and other biological materials) and traffic-related sources.

Traffic related non-exhaust emissions are currently not subject to legislative controls, and consequently these emissions are increasingly significant contributors to the environmental burden of particles as exhaust particle emissions decrease (Denier van der
Gon et al., 2013). Following the recent review of Pant and Harrison (2013), the four biggest components of non-exhaust particles are:

**Brake Wear**
Frictional contact between brake system components during forced deceleration is an important source of particulate matter emissions from motor vehicles. Gietl et al. (2010) found that barium comprises 1.1% of brake wear (PM$_{10}$) particles from the traffic fleet as a whole, allowing its use as a quantitative tracer of brake wear emissions at other traffic influenced sites. Dall’Osto et al. (2013) found that Fe-Cu can also be used as a tracer of brake wear. These elements are typical traffic markers, as the abrasion of brakes produces particles characterised by high concentrations of Cu, Ba, Zn and Fe (Sanders et al., 2003; Johansson et al., 2008).

**Mineral Dust**
The composition of road dust has been found to be dominated by elements and compounds typically associated with crustal material (Ca, Ti, and minor trace elements such as La, Ce, Ga and Sr.). Road dusts and roadside soils often also contain significant concentrations of a range of metals, including Pb, Cu, Cd, and Zn, indicative of contamination by road vehicle emissions. The road dust consists primarily of soil derived minerals (60%) with quartz averaging 40-50% and remainder being clay minerals such as albite, microcline, chlorite and muscovite originating from surrounding soils. Organic matter primarily originating from plant matter is also present (Gunawardana, 2012).

**Tyre Wear**
Tyre wear particles are produced from the interaction of tyres with the roadway surface. Generally, tyres comprise a rubber compound reinforced with steel wires and textile. At
present, very few reliable estimates of tyre wear in the ambient air are available, primarily due to limitations in the chemical markers used to quantify this type of PM. Previous research has shown tyre dust particles to be elongated particles with mineral incrustations from the road surface, which are quite different from tread particles produced by grinding or abrading a tyre tread (Kreider et al., 2010).

Road Surface Wear

Particles abraded from the road surface also become suspended, either directly or as resuspension from road surface dust. Their composition reflects that of the road surfacing materials: typically minerals with a bitumen binder.

While the brake wear is directly generated from brake components, and road dust comprises more general mineral components, the research on tyre dust has been less exhaustive and far less is known of its characteristics (Pant and Harrison, 2013). Several studies have been conducted to characterise tyre wear particles (Camatini et al., 2001; Dahl et al., 2006; Gustafsson et al., 2008; Sjödin et al., 2010; Lee et al., 2013). As regards the contribution to urban PM concentrations, results indicated that tyre concentrations in the PM$_{10}$ fraction are low with averages ranging from 0.05 to 0.70 µg m$^{-3}$, representing an average PM$_{10}$ contribution of 0.84% (Panko et al., 2013). Kwak et al. (2013) also reported a minor amount of tyre wear in urban dust, whereas the main proportion consisted of crustal matter generated from road surface wear and road dust. This finding is consistent with previous conclusions of Boulter (2005), who reported that less than 10% of car tyre wear materials are emitted as PM$_{10}$ under ‘typical’ driving conditions and by Gehrig et al. (2004), who concluded that tyre wear accounted for 7.5% of PM$_{10}$ at a kerbside site and 2% of PM$_{10}$ at an urban background site. Recently, Harrison et al. (2012a) reported separate estimation of the contributions of brake dust, tyre dust, and resuspension to...
particle mass in the range 0.9-11.5 μm aerodynamic diameter of 55%, 11% and 38%, respectively. As regards the tyre dust chemical composition, automobile tyre rubber is composed of 40-60% rubber polymers, 20-35% reinforcing agents/fillers such as carbon black or silica, and 15-20% mineral oils such as extender oils/softeners. The composition of the rubber depends on where in the tyre the rubber is used. Zinc typically comprises about 1% of tyre mass and Zn in road dust is often derived primarily from tyre wear (Adachi and Tainosho, 2004), although many other sources of Zn exist in urban areas (Pant and Harrison, 2013) and this element should not be regarded as a specific tyre wear marker. The key tracer components of tyre wear include n-alkanes, n-alkanoic acids, PAHs, benzothiazoles and trace metals. Overall, Pant and Harrison (2013) reported that tyre dust may have significant difference in composition among different countries.

The objective of this study was to reduce the uncertainty regarding tyre wear contributions in urban air. It used single particle mass spectrometry (SPMS), which gives direct insights into the mixing state of aerosols. Single particle analysis, as in SPMS, allows determination of whether a population of particles is “internally” (each single particle belonging to the aerosol population presents the same chemical composition as the bulk composition) or “externally” (single particles of different chemical compositions) mixed. In order to do so:

- a tyre wear experiment was run in the laboratory with a rubber tyre moving on a metal surface in order to evaluate the chemical composition of aerosolised tyre fragments;
- a large number of road dust samples were collected in order to estimate the tyre dust content;
- tyre dust mass spectra were sought in data from a number of ambient field studies carried out in different locations in which an Aerosol Time-of-Flight Mass Spectrometer (ATOFMS) was used for acquisition of compositional data on individual particles.
One of the major problems in the analysis of non-exhaust PM using field data has been the difficulty in distinguishing between wear and tear emissions and road dust since the chemical composition is often very similar (Bukowiecki et al., 2010; Pant and Harrison, 2013). The ATOFMS cannot provide quantitative aerosol mass loading concentrations, but its unique strength relies in the fact that it can monitor in real time variations in the single particle composition. Previous single particle studies have identified rubber particles from tyre wear experiments in the laboratory and found Zn using SEM-EDX (Scanning Electron Microscope-Energy Dispersive X-ray Spectroscopy) analysis (Camatini et al., 2001). Adachi and Tainosho (2004) found single particle tyre tread being associated with ZnO. In this study we use single particle mass spectrometry which gives more statistically robust results through characterisation of the many thousands of single particle mass spectra obtained. Nevertheless, it is worth pointing out that a limitation of this study is that it is a laboratory study which does not fully represents real world conditions. However, this unique single particle study allows us to study the mixing state of tyre dust, a source of urban aerosol not well characterised to date.

2. METHODOLOGY

2.1 Instrumentation

During the experiment instruments deployed were:

ATOFMS (Model 3800-100, TSI, Inc.) 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 (vacuum aerodynamic diameter). Thus, a positive and a negative ion spectrum of a single particle are obtained. The mass spectrum is qualitative in that the intensities of the mass spectral peaks are not directly proportional to the component mass
but are dependent on the particle matrix, the coupling between the laser and the particle and the shot to shot variability of the laser. However, the ATOFMS can provide quantitative information on particle number as a function of composition; providing a measure of all particle components and can be used to assess mixing state. During this study, the ART-2a neural network algorithm was used for classifying the large number of particles collected. The TSI ATOFMS datasets were imported into YAADA (Yet Another ATOFMS Data Analyzer) and single particle mass spectra were grouped with Adaptive Resonance Theory neural network, ART-2a (Song et al., 1999). The parameters used for ART-2a in this experiment were: learning rate 0.05, vigilance factor 0.85, and iterations 20. Further details of the parameters can be found elsewhere (Dall’Osto and Harrison, 2006).

An ART-2a area matrix (AM) of a particle cluster represents the average intensity for each m/z for all particles within a group and therefore reflects the typical mass spectrum of the particles within a group. The ART-2a algorithm generated a number of clusters used to describe the dataset herein presented.

Aerodynamic Particle Sizer (APS TSI 3321, size range 523-20,000 nm, aerodynamic diameter) and two Scanning Mobility Particle Sizer (SMPS TSI3071/CPC 3022A size range 15 nm-800 nm, and SMPS TSI3085/CPC3025 size range 5 nm-160 nm, mobility diameter) were used to provide the aerosol size distributions of the atmospheric dust particles herein described.

2.2 Laboratory Tyre Dust Experiment

The tyre wear rig (Fig. 1) comprises a vehicle wheel and tyre pressed under load against a large rotating metal wheel. Friction was generated by setting an angle between the vehicle
wheel and the metal wheel, simulating a cornering manoeuvre. The experiment was run for a total number of 25 hours. The first part ran on 18th April 2011 from 11:30 till 20:00. Experiments were run on different tyres (HOMO 88H 185/65/R15 and Michelin 185/70/14) and blanks were obtained by sampling the background and also with the metal wheel rotating with no wheel attached. During night-time the aerosol instruments were left sampling the background air. The morning after, on 19th of April 2011, the experiment was repeated again between 0900 and 1230. Figure 2 shows the data collected in the lab experiment. The sampling inlets were placed at distance of about 15 cm from the tyre/wheel interface and connected to the instruments via a 1 m long 1/4 inch conductive plastic tubing.

It is important to stress that the limitation of this study is the flat metal surface used to run the rubber tyre. This does not represent real world conditions, as road wear is generated from the friction between the tyre and the road, which occurs far more easily on asphalt/concrete roads than on a smooth metal surface. However, this gives a unique opportunity to study the chemical composition of tyre dust and then to see if such particles can be found under real world ambient conditions.

2.3 Single Particle Measurements of Road Dust Samples

A number of road dust samples were collected at four different locations within the city of London during the REPARTEE experiment (Harrison et al., 2012b). Samples were collected between 8th October and 30th October 2006 from four different locations around London. The four sites were very different from each other, in particular in relation to their volumes of traffic (see map of the four sites in Harrison et al., 2012b). The four sites were within 1 km of each other:
A: Marylebone Road site (MR). This road is spread on 6 lanes with frequent congestion, with traffic flows of over 80,000 vehicles per day.

B: BT tower (BT). A side road situated on the corner between Cleveland Street and Maple Street, within the congestion charge zone of London.

C: Inner Circle of Regents Park (RR). Road situated within Regent's Park, with leaves and soil often seen on the ground.

D: Footpath in Regents Park (RP). This little pathway is made of concrete/stones and no cars are allowed, although bicycles are.

Road samples were taken by sweeping the road surface with a brush into a small plastic bag. A different brush was used for each location, and the brush was washed before the collection. Forty-eight samples (12 each monitoring site) were taken during the month of October 2006. Mass spectra from each dust sample were obtained by placing each dust sample into a flask in a sonicator to create a suspension of dust particles under filtered lab air flow. The aerosol was then directed into the ATOFMS inlet for analysis and about 500 single-particle mass spectra were collected per sample (2791 particles in total).

2.4 Field Studies

A number of field studies have been run by the University of Birmingham in the last decade. The ATOFMS can be used with two different inlets. The ATOFMS TSI model 3800-100 uses an aerodynamic lens to direct particles to the sizing region of the instrument (Su et al. 2004). By contrast, the ATOFMS TSI Model 3800 utilizes a simple nozzle/skimmer inlet (Gard et al. 1997). In terms of particle collection efficiency, the aerodynamic lens provides a major improvement toward smaller particle sizes (<1 µm).
the following studies, the aerodynamic lens ATOFMS TSI 3800-100 system (200-3000 nm), was used:

- A remote marine site in the coastal area of Mace Head, Ireland (Dall’Osto et al., 2010a).
- A cruise in the Tropical Atlantic Ocean in 2006 (Dall’Osto et al., 2010b).
- A cruise in the North Atlantic Ocean during MAP (Decesari et al., 2012).
- A regional site in the Po Valley, Italy in 2009 (Decesari et al., 2013).
- A regional site in Harwell, England in 2010 (Giorio et al., 2012).
- A roadside site on Marylebone Road, London, UK in 2009.
- A roadside site in Barcelona (Spain) during the SAPUSS field study (Dall’Osto et al., 2013).

3. RESULTS

3.1 Laboratory Tyre Dust Experiment

3.1.1 Size distributions of tyre dust

Figure 2 shows the plot of aerosol size distributions (both SMPSs and APS) taken during the day of the tyre experiment. When the mean aerosol size distributions are plotted (Fig. 3), it is clear that a bimodal particle distribution is seen below 100 nm with modes at about 35 nm and 85 nm. Unimodal (70-90 nm) and bimodal (<10 and 30-60 nm) number size distributions in the nano size range have been reported for tyre particles under low and high speed conditions respectively (Mathissen et al., 2011). Dahl et al. (2006) found an enrichment of tyre particles between 15-50 nm using a road simulator. They identified two types of particle: one comprising mineral oils from the softening filler, the other of soot-like agglomerates from the carbon-reinforcing filler material. Our APS results showed little enhancement of coarse particles above background across the whole size range during
experiments 1 and 2. In summary, few particles above 1 µm were detected, although those may significantly contribute to mass.

Kreider et al. (2010) characterized the physical and chemical properties of particles generated from the interaction of tyres and road surfaces. However, these particles were distributed in the very large coarse mode, spanning from 4-6 µm to 265-280 µm with the mode centred at approximately 50-75 µm.

3.1.2 Chemical speciation of tyre dust

Art-2a was run on the 25,000 ATOFMS single particle mass spectra detected during the lab studies. A number of Art-2a clusters were detected, most of which were associated with background conditions. One cluster was found to peak during the experiment when the tyre was running on the metal surface. The spectrum obtained is shown in Figure 4, and is the average of the two different tyres used. Given that the two tyres generated the same single particle mass spectra, the results are presented as representing tyre dust as generated in the laboratory. In the positive mass spectrum there is a main peak at \( m/z \) 39, which could be due to potassium or an organic fragment. Other organic peaks can be seen at \( m/z \) 41, 43 and 69. The peaks at \( m/z \) 69, 81 and 95 (particularly the strong peak at \( m/z \) 95) may represent the exo-sulphur aromatic series (sulphur attached to an aromatic ring) (McLafferty, 1993). Sulphur is used as vulcaniser in the tyre, so S internally mixed with organic compounds is expected in tyre dust. Zinc (Zn) is added to tyre tread as zinc oxide and organozinc compounds to facilitate the vulcanization process. However, Zn is emitted from brake wear, motor oil and other sources and cannot be used as a unique tracer for tyre wear (Lough et al., 2005). Zn can be detected in ATOFMS single particle mass spectra. For example, a Zn particle type detected in a steel industry study showed the presence of ions at \( m/z \) 64, \([\text{Zn}]^+\); 97, \([\text{ZnCl}]^+\) and –169, \([\text{ZnCl}_2]^–\) (Dall’Osto et al., 2008).
During this study Zn (query search $m/z$ 64 without the presence of $m/z$ 48 [Ti], Dall’Osto et al., 2010b) was detected in about 16% of the particles. The negative mass spectrum shows the presence of peaks due to CN ($m/z$ -26, -42) as well as nitrate ($m/z$ -46, -62) which are not expected. Nitrogen compounds in tyre dust are usually not seen and the mass spectra from the ATOFMS may be simply a representation of the extreme sensitivity to K and nitrate of the instrument (Gross et al., 2000). A number of unique peaks can be seen in the high molecular weight ($m/z$ > 100) range. The peaks at $m/z$ 140 and $m/z$ -163 may be due to potassium nitrate ((K$_2$NO$_3$) and (K(NO$_3$)$_2$, respectively (McLafferty, 1993), but also to unidentified organic compounds. Other unique peaks at $m/z$ 173, $m/z$ -127, $m/z$ -146 can be seen. These are not characterised and are likely due to rubber copolymer components.

3.2 Single Particle Experiment with Road Dust Samples

The ATOFMS spectrum obtained from the tyre experiment above was compared with the 48 road dust samples using YAADA. Only few single particle mass spectra (< 10) identical with the laboratory generated tyre dust were found. Overall, less than 0.1% of the road dust mass spectra matched the tyre road dust sample. This implies that tyre dust was not found in the same form that was generated in the laboratory. However, by searching the unique peaks associated with the tyre material ($m/z$ 140, $m/z$ 173, $m/z$ -163), it was found that about 4% of the particles were associated with such unique markers. This implies that the mixing state of the particles found in the real world is different from the laboratory study. An average spectrum of the particles containing specific tyre dust $m/z$ peaks is shown in Fig. 5. It is clear other peaks are present. Fig. 6 shows the difference, taken as:

Difference (Fig. 6) = real world road dust with tyre markers (Fig. 5) minus laboratory tyre dust (Fig. 4), and scaled relative to the two biggest peaks ($m/z$ 39 and $m/z$ -62).
The difference comprises mainly ions due to crustal elements (particularly Li, Na, Al, Fe and Si). The positive mass spectrum clearly shows the peaks associated with Ca (m/z 40) and Fe (m/z 56). Additionally minor peaks are Al (m/z 27), and this is confirmed clearly by the negative peak at m/z -43 (AlO). Peaks with m/z of 48 and 64 are due to Ti and TiO, another common marker seen in dust, and at m/z 7 is attributed to Li. The negative peaks include -26 (organic peak), EC (-48) and phosphate (-63, -79). The presence of the phosphate may be due to lubricating oil - a marker that the ATOFMS is sensitive to (Spencer et al., 2006). The signal at m/z 64 could be due to Zn, but the signal could also be associated with titanium oxide (TiO, m/z 64).

The results suggest that the tyre, when rotating on a real road surface, creates particles internally mixed with the minerals contained in road surface materials. Hence, the tyre is degrading the road surface, creating a particle internally mixed between tyre material and road surface material. This is not a new observation, as a number of studies have already pointed out this mechanism (Kreider et al., 2010). However, this study shows for the first time that most, if not all, tyre dust particles are internally mixed with minerals coming from the abrasion of the road surface. Additionally, whilst the study of Kreider et al. (2010) focused on particles in the size range 4-280 µm, our study mainly characterise PM smaller than 3 µm and more relevant to PM$_{2.5}$ exposures.

3.3 Field Studies

To provide further evidence, ATOFMS road dust markers (sum of m/z 140, m/z 173 and m/z -163, peak area >100) were sought in a number of ambient field studies run in the past. This very large ATOFMS dataset is an ensemble of field studies carried out by the University of Birmingham (UK) in more than a decade, spanning from very remote open ocean areas to heavily trafficked road sites (please see references reported in section 2.4 for
further details). Table 1 clearly shows that tyre related dust was not found at sites away from urban agglomerates. Furthermore, it can be seen that the highest percentages were found at roadside sites (London, Birmingham and Barcelona - about 2%) and in the road dust samples collected (4%). Whilst Table 1 may appear simple, it comprises more than 2,000,000 single particle mass spectra collected across the European Continent.

4. DISCUSSION AND CONCLUSION

Road dust, of which crustal dust is a key component, consists primarily of coarse-sized particles derived from different sources such as traffic, industrial emissions, mineralogical dust etc. The composition of road dust shows spatial as well as temporal variation and it is often difficult to classify dust unequivocally into crustal/re-suspended/direct emission etc. Very few reliable estimates of tyre wear in the ambient air are available, primarily due to limitations in the elemental markers which have been used to quantify this type of PM.

This study suggests that pure tyre wear particles are not often found in real world road dust samples and ambient aerosol. By contrast, most of them are found internally mixed with minerals and other substances, possibly including contributions from lubricating oil and vegetative debris. It is important to note that, within the road dust samples, all the tyre road dust particles were found in the samples taken on the roads (Marylebone Road and the road next the BT tower), whereas no tyre dust-like spectra were found in the Inner Circle of Regent’s Park (RR) or in the Footpath in Regent’s Park (RP) in the city of London. This gives confidence in our ATOFMS markers and the results herein presented. Our results are quantitatively very consistent with those of Kreider et al. (2010) and quantitatively broadly consistent with those of Panko et al. (2013) in showing an association of tyre rubber with mineral material.
The ATOFMS cannot provide quantitative measurements without appropriate scaling factors (Dall’Osto et al., 2006). However, when looking at the particles containing ATOFMS tyre wear markers (4% by number), we find that most of them by volume were distributed in the size range 1-3 µm.

There is clearly a discrepancy in the size distributions of tyre particles measured in the laboratory (mainly sub-100 nm) and those in the field studies (1-3 µm). If present in the atmospheric samples, the sub-100 nm particles would not be detected by the ATOFMS. The particle counting instruments showed only modest production of 1-3 µm particles in the laboratory study, although sufficient to be detected by the ATOFMS, suggesting that the mechanism of particle generation in the laboratory from friction between the tyre and a smooth metal surface may be different from that on the road where the surface is rough. However, earlier studies using rougher surfaces have also shown a predominance of nano-sized particles (Dahl et al., 2006; Mathissen et al., 2011).

One consequence of our findings is that tyre wear obtained by abrading a tyre, e.g. by using a coarse stainless steel file at between two and four different locations on each tyre (Sadiktsis et al. 2012) may not be appropriate for toxicological studies. Our study shows that the large majority of tyre dust particles in urban air are internally mixed with minerals. This implies that field studies using chemical tracers for tyre wear and resuspension particles may under-estimate those arising from tyre-road interaction and over-estimate the resuspension component.
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Table 1. Percentages of tyre particles in ambient field studies run with the ATOFMS. Twelve field studies were examined, with a total of more than 2 millions particle mass spectra.

Figure 1. Photo of the tyre rig at the University of Birmingham. The arrow points towards the point of contact between the tyre (on the left) and the metal wheel (to the right).

Figure 2. The tyre dust lab experiment. Top chart shows the Art-2a ATOFMS particle type associated to tyre dust (ATOFMS counts/hour), middle charts show aerosol size distributions (APS, SMPS, nano-SMPS) and bottom chart shows the experimental mode used.

Figure 3. SMPS size distributions of the tyre experiment (and relative background). The difference (average of diff exp 1 and diff exp 2) shows an enhancement of particles at 35 and 85 nm.

Figure 4. ATOFMS art2a cluster associated to tyre dust in the laboratory.

Figure 5. Average spectra of road dust single particles containing tyre dust key m/z peaks.

Figure 6. Difference between tyre dust (from lab study) and road dust with tyre markers.
Table 1. Percentages of tyre particles in ambient field studies run with the ATOFMS. Twelve field studies were examined, with a total of more than 2 millions particle mass spectra.

<table>
<thead>
<tr>
<th>Monitoring site</th>
<th>Characteristic</th>
<th>% of tyre dust (by number of particles)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mace Head</td>
<td>Marine</td>
<td>0%</td>
</tr>
<tr>
<td>North Atlantic</td>
<td>Marine</td>
<td>0%</td>
</tr>
<tr>
<td>Tropical Atlantic</td>
<td>Marine</td>
<td>0%</td>
</tr>
<tr>
<td>Po Valley</td>
<td>Regional background</td>
<td>0.5%</td>
</tr>
<tr>
<td>Harwell</td>
<td>Regional background</td>
<td>0.5%</td>
</tr>
<tr>
<td>Marylebone road</td>
<td>Roadside</td>
<td>2.5%</td>
</tr>
<tr>
<td>SAPUSS Barcelona</td>
<td>Roadside</td>
<td>2%</td>
</tr>
<tr>
<td>London streets</td>
<td>Road dust samples</td>
<td>4%</td>
</tr>
</tbody>
</table>
Figure 1. Photo of the tyre rig at the University of Birmingham. The arrow points towards the point of contact between the tyre (on the left) and the metal wheel (to the right).
Figure 2. The tyre dust lab experiment. Top chart shows the Art-2a ATOFMS particle type associated to tyre dust (ATOFMS counts/hour), middle charts show aerosol size distributions (APS, SMPS, nano-SMPS) and bottom chart shows the experimental mode used.
Figure 3. SMPS size distributions of the tyre experiment (and relative background). The difference (average of diff exp 1 and diff exp 2) shows an enhancement of particles at 35 and 85 nm.

Figure 4. ATOFMS art2a cluster associated with tyre dust in the laboratory.
Figure 5. Average spectra of road dust single particles containing tyre dust key m/z peaks.
Figure 6. Difference between tyre dust (from lab study) and road dust with tyre markers.