

Partitioning of trace elements and metals between quasi-ultrafine, accumulation and coarse aerosols in indoor and outdoor air in schools

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HIGHLIGHTS

- Quasi-ultrafine particles (q-UF) are measured indoors and outdoors in 39 schools.
- Particle size distribution patterns of trace elements and metals are described.
- High proportions of typically coarse elements are found in q-UF particles.
- Potentially health-hazardous metals are found in q-UF in indoor air.
- The location of schools in trafficked areas increases children exposure to q-UF.

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ABSTRACT

Particle size distribution patterns of trace elements and metals across three size fractions (<0.25 μm, quasi-ultrafine particles, q-UF; 0.25–2.5 μm, accumulation particles; 2.5–10 μm, coarse particles) were analysed in indoor and outdoor air at 39 primary schools across Barcelona (Spain). Special attention was paid to emission sources in each particle size range. Results evidenced the presence in q-UF particles of high proportions of elements typically found in coarse PM (Ca, Al, Fe, Mn or Na), as well as several potentially health-hazardous metals (Mn, Cu, Sn, V, Pb). Modal shifts (e.g., from accumulation to coarse or q-UF particles) were detected when particles infiltrated indoors, mainly for secondary inorganic aerosols. Our results indicate that the location of schools in heavily trafficked areas increases the abundance of q-UF particles, which infiltrate indoors quite effectively, and thus may impact children exposure to these health-hazardous particles.

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1. Introduction

Most of the concern over atmospheric particles derives from their impact on a number of aspects, ranging from climate (Shindell et al., 2012) or visibility (Cheung et al., 2005), to human health (WHO, 2013). In regard to the latter, aerosol effects are dependent on particle size, with different health effects being associated to

different particle diameters (WHO, 2013). Aerosols may be classified based on particle diameter as coarse (aerodynamic diameter > 1 μm), fine (1 μm > aerodynamic diameter > 100 nm), ultrafine particles (<100 nm) and nanoparticles (<50 nm) (Seinfeld and Pandis, 1998). Some of these definitions may vary, especially for the finer size fractions (Kumar et al., 2010). Recent studies have extended the use of the term ultrafine particles to particles up to 250 nm in diameter, referring to this size fraction as “quasi-ultrafine” (Saffari et al., 2013).

Numerous studies have evidenced a wide range of adverse health outcomes on the human cardiovascular and respiratory

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systems linked to the exposure to airborne coarse, fine and ultrafine particles (Cassee et al., 2013). Whereas coarse aerosols impact mostly on respiratory system (Pérez et al., 2008), ultrafine particles have the ability to translocate to the circulatory system and induce inflammation (Oberdorster, 2001).

The partitioning of elements among various sizes depends on their sources and affects their chemical composition and therefore will impact the potential toxicity induced on different regions of the body (Cassee et al., 2013). Based on their chemical composition different types of particles generate reactive oxidative species, with metallic particles such as Cu and Fe being a well-known example (Fu et al., 2014) albeit not the only one (e.g., soluble and mineral particles; Hetland et al., 2001). Because literature results evidence that health effects from airborne particles depend on their chemical composition (e.g., transition metals, combustion-derived organic particles) but also on physical properties (size, particle number; Cassee et al., 2013), it is essential to assess the link between the two by evaluating the concentration of major and trace aerosol components across particle size fractions. This assessment could be used as input data for the assessment of health effects of particles reaching different regions of the body.

In addition to health effects studies, source apportionment analyses of ultrafine particles and nanoparticles by receptor modeling tools also require chemical size distribution data. For coarse and fine aerosols, tracer elements of specific emission sources (e.g., vehicular traffic, mineral dust, sea salt, among others) are well established and have been frequently used in the literature (Bruinen de Bruin et al., 2006). However, studies have indicated that such tracers may not be common across particle size fractions (Lin et al., 2005; Miller et al., 2007; Gietl et al., 2010; Buonanno et al., 2011; Liati et al., 2012; Patel et al., 2012). As an example, whereas Al and Ca are known to be tracers of mineral dust in coarse aerosols, they may be derived from local-scale combustion processes when found in ultrafine particles (Lin et al., 2005). As a result, interpreting Al and Ca as tracers of mineral matter in ultrafine particles may be misleading for specific cases (Sanderson et al., 2014). Similar cases are reported for Fe, Ca, P and Mg in ultrafine particles from lubricating oils (Miller et al., 2007; Patel et al., 2012) and for Ca, Mg, Al and Fe in ultrafine particles from diesel soot (Liati et al., 2012; Sanderson et al., 2014). Consequently, relatively few source apportionment studies of ultrafine particles are currently available (Karanasiou et al., 2007; Sanderson et al., 2014).

Finally, understanding the way in which elements partition across particle size fractions also allows us to interpret changes in such particle size distributions, especially those induced by physical and/or chemical processes, such as particle infiltration from outdoor to indoor air. Infiltration processes may alter particle size distribution and aerosol composition for specific aerosols, e.g., volatilization of ammonium nitrate (Hering et al., 2007). The assessment of the chemical size distribution data of aerosol components should enhance our understanding of the extent to which infiltration processes affect indoor aerosols, and provide better insights on indoor exposures. In the present study, this analysis targets children and their exposure to indoor airborne pollutants during school hours. Previous studies on indoor air quality in schools (Blondeau et al., 2005; Diapouli et al., 2007; Weichenthal et al., 2008; Morawska et al., 2009; Mejía et al., 2011; Mazaheri et al., 2014) concluded that research on indoor air quality in typical schools is vital to students' health and their performance, given that numerous outdoor airborne pollutants are present in schools.

This work aims to quantify the distribution of major metals and trace elements across three particle size fractions in urban outdoor and indoor air. This assessment was carried out in primary schools in Barcelona, both indoors and outdoors, and at different distances from traffic emissions. Our results provide qualitative and

quantitative descriptions of particle size distribution patterns for metals and trace elements, with a special focus on their emission sources in each particle size range.

2. Methodology

2.1. Sampling locations

Measurements were carried out in 39 schools across Barcelona (Spain), selected in the framework of the Advanced ERC Grant BREATHE. Schools were divided into those representative of the city's urban background pollution levels and of high traffic emissions (Rivas et al., 2014), and then grouped into pairs (one school of each type) for concurrent sampling. In each school, indoor and outdoor sampling was carried out simultaneously. Further details of the sampling locations are provided elsewhere (Rivas et al., 2014; Viana et al., 2014). Two six-month sampling campaigns took place between January 2012 and February 2013, at a rate of 2 schools per week. Sampling was carried out at each school twice.

2.2. Sample collection and analysis

Filter samples were collected using Personal Cascade Impactor Samplers (PCIS; Misra et al., 2002) connected to SKC Leland Legacy pumps, operating at 9 l pm. Four PCIS were placed simultaneously (2 indoors, 2 outdoors) in each school. Samples were collected only during school hours (9:00–17:00 local time, 8 h) over a period of 4 consecutive days (Monday to Thursday). Thus, samples accumulated 32 sampling hours on the same filters and were representative of weekly (4-day) school-hour concentrations.

Each PCIS collected size-fractionated particles in the size ranges <0.25 μm (quasi-ultrafine particles, q-UF), 0.25–2.5 μm (accumulation mode particles), and 2.5–10 μm (coarse PM). The collection substrates were Pall quartz-fibre filters (Pall Tissuquartz 2500QAT-UP), 25 mm in diameter for the coarse and accumulation stages, and 37 mm for q-UF particles.

After collection, mass concentrations were gravimetrically determined by weighing the filter substrates on a Mettler-Toledo Microbalance. All samples were then analysed for major and trace elements. For each school, week and environment (indoor and outdoor), 2 parallel samples (with 3 filter substrates each) were available: A and B. The mass concentration on samples A and B was comparable. The filters from PCIS A were acid digested ($\text{HNO}_3\text{:HF:HClO}_4$) to determine major and trace elements by ICP-MS and ICP-AES (Querol et al., 2001). The filters from PCIS B were cut in two halves: one half was leached in deionised water to extract the soluble fraction, which was analysed by ion chromatography and ion selective electrode to determine concentrations of SO_4^{2-} , NO_3^- , Cl^- and NH_4^+ . The remaining half filter was used for determination of OC and EC by thermal–optical transmission (TOT) with a Sunset OCEC Analyzer following the temperature protocol NIOSH. This methodology was applied equally to each of the size ranges (q-UF, accumulation and coarse).

In total, the number of valid samples for outdoor air was 84 and 144 for sampling campaigns 1 and 2, respectively, and 75 and 126 for indoor air for sampling campaigns 1 and 2.

3. Results and discussion

3.1. Mean indoor and outdoor concentrations

The size-fractionated mean concentrations of major and trace components, as well as the gravimetrically determined mass, for indoor and outdoor air samples, are shown in Fig. 1. The data shown are the mean concentrations for both sampling campaigns,

and thus representative of cold and warm periods separately, and of the annual mean when combined. The measurements were taken during the school year (while children were inside the classrooms). The mass closure of indoor and outdoor aerosols is not described in detail, as it is not the main focus of this work. The mass closure for the first sampling campaign was discussed in detail in Viana et al. (2014), whereas this work focusses on particle size distribution patterns and across both sampling campaigns.

An initial assessment demonstrates larger differences between the mass of the three size fractions in indoor than in outdoor air, with a mass ratio between coarse and accumulation mode particles of 2.3 indoors and of 1.5 outdoors, respectively. This difference is mainly induced by the higher indoor organic carbon (OC) mass. In indoor air, OC was the major contributor to all size fractions, accounting for 74% of coarse, 80% of accumulation and 39% of q-UF particles. As described by Rivas et al. (2014) and Viana et al. (2014), the main origin of organic aerosols in indoor air in the schools under study was human activity (textiles, skin flakes, etc.). Also in indoor air, the second largest contributor to all size fractions were major elements Al₂O₃, Ca, Fe, K and Mg, most of them tracers of mineral matter in coarse and accumulation particles. In q-UF particles, the source origin of these major metals is most probably also mineral. However, in trace concentrations much lower than the ones obtained in this study, these elements may also originate from road dust (Dahl et al., 2006), traffic fuels (diesel or gasoline) and/or to lubricants, among others (Sanderson et al., 2014). It is also possible that the high contribution of mineral elements to q-UF particles is, to a certain extent, an effect of bounce-off from previous stages in the impactors, or to a slight deviation from the expected cut-off diameter due to flow control. This is a known limitation of impaction techniques (Gomes et al., 1990). However, Misra et al. (2002) showed that bounce is negligible with PCIS impactors due to the relatively low jet velocity (645 cm/s) in the 2.5 μm stage. This potential artefact was not quantified in the present work.

Mineral elements accounted for 15% of coarse, 6% of accumulation and 9% of q-UF particles indoors. In coarse particles, the main source of mineral dust was re-suspension of indoor- and outdoor-originated dust within the classrooms (Amato et al., 2014; Rivas et al., 2014; Viana et al., 2014). The high contribution of mineral elements to q-UF particle mass (2.3 μg/m³) was an unexpected result, given that mineral contributions are generally considered to be minimal in this size fraction (Seinfeld and Pandis, 1998). However, Rivas et al. (2014) showed that the levels of mineral matter in PM_{2.5} at the Barcelona schools are much higher than those measured in the Barcelona urban background. Tracers of marine aerosols (Na and Cl⁻) contributed with 4–5% of particle mass across size fractions, although it should be stated that traces of Na could originate from other (anthropogenic) sources in q-UF particles, as

in the case of other major elements (based on the low R^2 between Na and Cl⁻, <0.3).

Relatively larger similarities were observed in the chemical composition of the different particle size fractions outdoors, when compared to indoor air. Organic matter was again the main contributor to all size fractions outdoors, with contributions ranging from 57% of coarse particles to 74% of accumulation and 30% of q-UF particles. The main source of outdoor carbonaceous aerosols (including elemental carbon, contributing with 1–5% of the PM mass) in the Barcelona schools under study was vehicular traffic (Amato et al., 2014). Mineral elements accounted for 13% of coarse, 5% of accumulation and 10% of q-UF particle mass, similarly to indoor air, even if in terms of mass concentration the contribution was higher indoors for coarse particles (5.6 μg/m³ indoors vs. 2.7 μg/m³ outdoors). As in the case of indoor air, the mineral mass in q-UF particles found outdoors (2.3 μg/m³) was high, as observed also by Rivas et al. (2014) for PM_{2.5}. The mineral mass in q-UF particles was higher than that reported by Daher et al. (2013) for traffic-influenced sites in central Los Angeles (0.84 μg/m³), suggesting the possibility of an additional source of q-UF mineral particles in the Barcelona schools. Saffari et al. (2013) also reported mineral matter concentrations lower than in Barcelona. These results, together with the higher organic mass indoors also in coarse particles (27.1 μg/m³ indoors vs. 11.9 μg/μm³ outdoors) confirm the high exposure of schoolchildren in Barcelona to mineral and organic aerosols reported in detail elsewhere (Rivas et al., 2014; Viana et al., 2014).

3.2. Partitioning of chemical components across particle size fractions

The particle size distribution patterns of major and trace elements are discussed in the following sections. Because only three size fractions are available, the terms “patterns” or “particle size distribution patterns” are used throughout the text, as opposed to “particle size distributions” which is more appropriate when a larger number of size channels is available. The terms “unimodal” or “bimodal” are only used on specific occasions.

a) Outdoor air

The mean distribution of major and trace elements in outdoor air are shown in Fig. 2. The behaviour of each individual element and component was assessed in order to evaluate how they partition between size fractions, attempting to identify common patterns and to infer potential emission sources. Initially, the prevalence of mass concentrations in one or two modes was identified, but the distinction between them was at times not straightforward. In order to apply a systematic approach, an arbitrary threshold of 20% was defined as the minimum difference

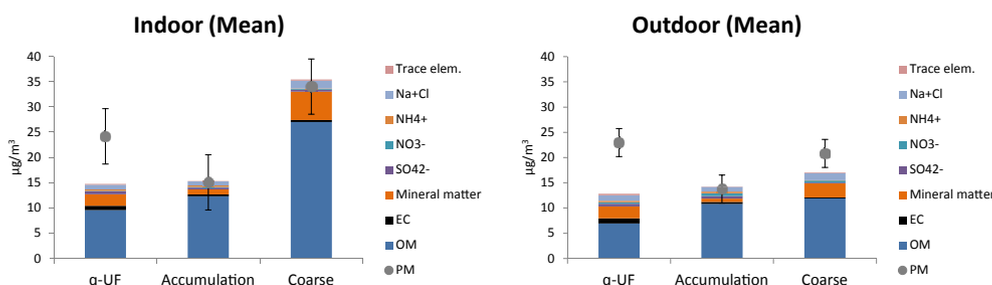


Fig. 1. Size-fractionated mean concentrations of major species, metals and trace components, as well as the gravimetrically determined mass, for indoor and outdoor air. Number of valid samples: 228 for outdoor air and 201 for indoor air.

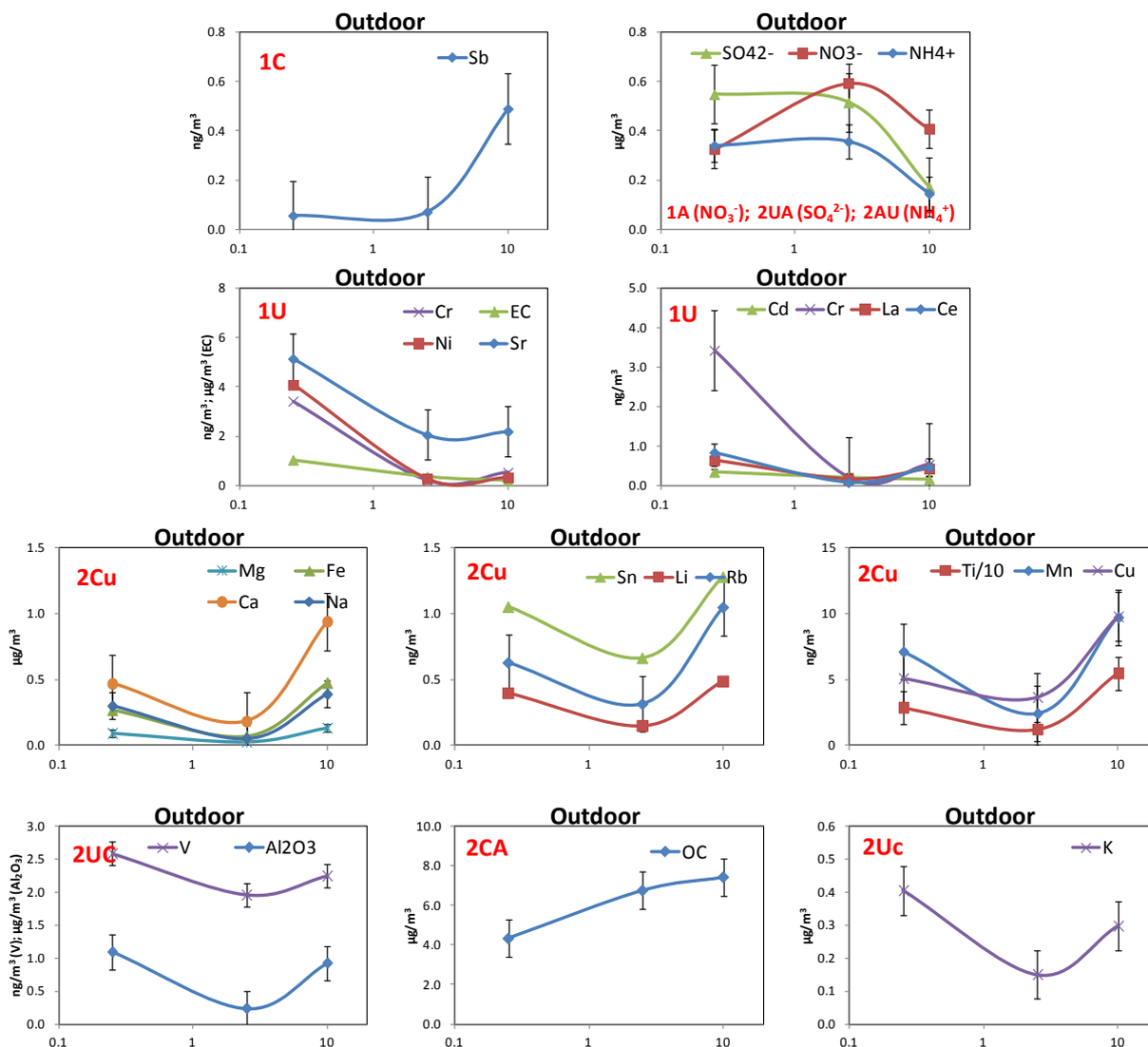


Fig. 2. Size distribution patterns identified for outdoor air. The patterns (in red, top left corner) are defined as follows: one mode, with prevalence for q-UF particles (1U); one mode, with prevalence for accumulation particles (1A); one mode, with prevalence for coarse particles (1C); two modes (coarse and q-UF), with prevalence for coarse particles (2Cu); two modes (coarse and q-UF), with prevalence for q-UF particles (2UC); two modes (coarse and q-UF), with similar mass in both modes (2UC); two modes (q-UF and accumulation), with similar mass in both modes (2CA). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

between mass concentrations to define a mode. As an example, for a given element, the ratio between the mass concentration in the coarse and accumulation modes was calculated, and also the ratio between q-UF and accumulation modes. If both ratios indicated mass differences between modes >20%, then the distribution was classified as having 2 modes, because there was a >20% mass difference between the coarse and accumulation modes, and between the q-UF and accumulation modes (“U” shaped distribution). Conversely, if only one ratio showed a mass difference >20% then the distribution was classified as having only 1 mode.

Applying this classification scheme, the following patterns were identified (Fig. 2 and Table 1):

One mode, with prevalence for accumulation particles (1A): as expected (Seinfeld and Pandis, 1998), this was the characteristic size distribution of NO_3^- aerosols in outdoor air. Because NO_3^- may

be present in the form of NH_4NO_3 but also forming salts with coarser cations (Ca, Na, etc.), 46% of the mean NO_3^- mass was found in accumulation particles followed by 31% of the mass in coarse and 23% of the mass in q-UF particles. Despite showing 2 modes (Table 1), SO_4^{2-} and NH_4^+ are also described here due to their prevalence in the accumulation mode. Sulphate was found mostly in the form of $(\text{NH}_4)_2\text{SO}_4$ and therefore showed a strong contribution in the accumulation mode but also in q-UF particles (q-UF, 44%, and accumulation, 42%, classified as 2UA). The same pattern was followed by NH_4^+ , with prevalence for accumulation (45% of the mass) and q-UF (43%) particles (pattern 2UA).

One mode, with prevalence for coarse particles (1C): components with this size distribution were Sb and Ga. This distribution suggests a single source for each of these species, or different sources emitting particles in the same size range. In the case of Sb, the emission sources are mostly linked to vehicular emissions (road dust re-suspension, Bruinen de Bruin et al., 2006), which generate coarse particles.

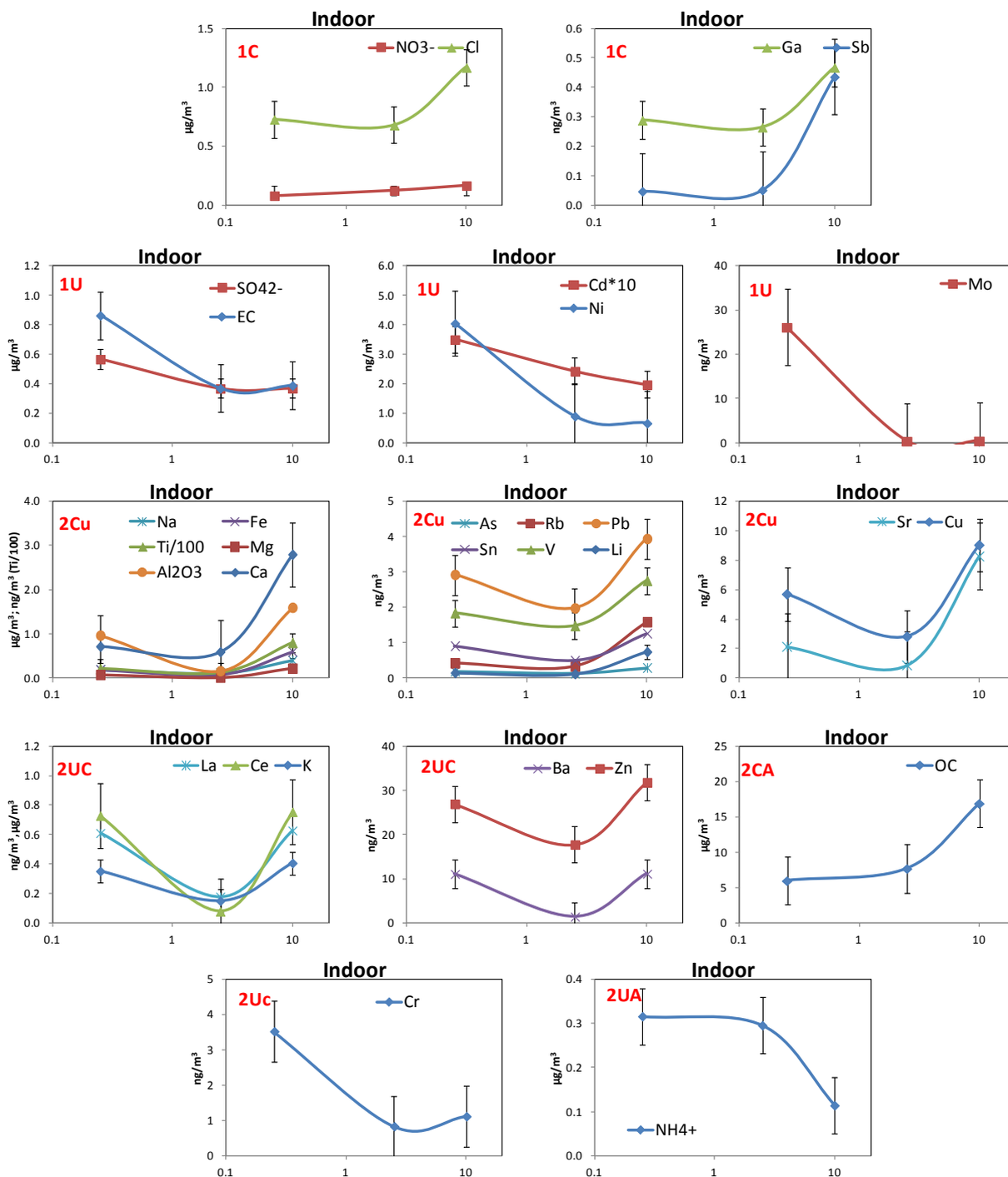


Fig. 3. Size distribution patterns identified for indoor air.

Two modes (coarse and q-UF), with prevalence for coarse particles (2Cu): components with prevalence for coarse particles but with a secondary q-UF particle mode in indoor air were Fe, Ca, Mg, Na, Li, Ti, Mn, Cu, Rb, Sn, Al₂O₃, V, As, Sr and Pb. The first (Fe, Ca, Mg, Na, Li, Ti, Mn, Cu, Rb, Sn) showed the same pattern outdoors, suggesting that at least one indoor source of these elements was infiltration from outdoor air. Given their mostly mineral origin, another indoor source would be re-suspension of indoor dust (originated indoors or entrained from playgrounds by the children). In the case of Ti, concentrations in the coarse mode were 80 ng/m³ indoors and 55 ng/m³ outdoors, whereas they were fairly similar in the q-UF

and accumulation modes. This may indicate the presence of a specific indoor source of coarse Ti particles, which could be related to the re-suspension of settled particles of outdoor origin or to TiO₂ used as white pigment in paint (Uhde and Salthammer, 2007). For Ca, an additional enrichment in coarse particles indoors was detected, which results from the contribution from chalk dust (Rivas et al., 2014). Absolute indoor Ca concentrations were higher than outdoors by a factor of 2.6, especially in the coarse mode (I/O factor of 3.0). Mineral dust re-suspension and the subsequent enrichment in coarse particles was also the most probable cause for the modal shift observed for the remaining components in this

group (Al_2O_3 , V, As, Sr, Pb). Al_2O_3 followed a 2UC pattern outdoors and shifted to a coarse pattern (2Cu) indoors, due to an increase in the coarse Al_2O_3 mass. V followed a similar pattern, as it may substitute Al in crystalline structures. A similar increase in coarse Sr was detected in indoor air, together with a marked decrease in q-UF Sr with respect to outdoor air (where Sr followed a 1U pattern). Elements which may be less related to mineral matter (e.g., As) also showed increases in coarse mode mass and decreases in the masses of accumulation and q-UF particles in indoor with respect to outdoor air. Absolute concentrations were always similar or lower indoors.

As in the case of outdoor air, these results indicate the presence in q-UF particles in indoor air of typical tracers of mineral matter (both of indoor and outdoor origin), and in relatively high proportions (e.g., 35% of Al_2O_3 , 17% of Ca, 21% of Fe, 39% of K, 24% of Mg or 27% of Na, Table S2). These results also show the presence in indoor air of trace and potentially toxic metals in q-UF particles, such as Mn, Cu, Sn, V or Pb. The origin of these metals is probably infiltration from outdoor air. As a result, metallic q-UF particles are detected indoors and thus may impact exposure of schoolchildren.

Two modes (coarse and q-UF), with prevalence for q-UF particles (2UC): only one element (Cr) was identified with a consistent and strong prevalence for q-UF particles and a smaller contribution in coarse particles.

Two modes (coarse and q-UF), with similar mass in both modes (2UC): a similar case to the 2Cu pattern, but with an even stronger enrichment in coarse particles, was detected for K, Zn, Ba, La and Ce. These elements showed a prevalence for q-UF particles in outdoor air (1U or 2Uc patterns), whereas indoors an increase in the absolute coarse mass concentration of all of them was detected. This suggests the presence of indoor sources (possibly, mineral), and the increased exposure of children to these elements in coarse aerosols.

Two modes (q-UF and accumulation), with similar mass in both modes (2UA): in indoor air, NH_4^+ was the only component to show this mass size distribution pattern. A similar pattern was expected for SO_4^{2-} , but however absolute concentrations were similar ($1.2 \mu\text{g}/\text{m}^3$ outdoors vs. $1.3 \mu\text{g}/\text{m}^3$ indoors) but a shift toward q-UF particles was detected in indoor air (pattern 1U, Fig. 3). This shift was relative and linked not with an increase of q-UF SO_4^{2-} but with a reduction in accumulation mode SO_4^{2-} (ratio I/O for accumulation particles = 0.7). This could be related to infiltration losses due to particle size, even though infiltration is known to be most efficient for particles in the accumulation mode (Long et al., 2001). In addition, an increase in coarse SO_4^{2-} particles was detected (ratio I/O for coarse particles = 2.2), which probably originate from indoor-specific sources such as chalk from blackboards (Rivas et al., 2014; Viana et al., 2014).

Two modes (accumulation and coarse), with similar mass in both modes (2CA): indoor OC followed the same pattern as in outdoor air, as a result of infiltration of outdoor particles indoors. In addition to the outdoor sources of OC described above, one additional source of coarse indoor OC (human activity, Rivas et al., 2014) contributed to an enhanced coarse OC mode, with a larger coarse/accumulation ratio (2.2 indoors vs. 1.1 outdoors).

In sum, typically mineral elements (e.g., Ca, Fe, Mg, Al, Ti, Sr) prevailed in coarse aerosols indoors due to the influence of indoor sources (dust re-suspension). In addition, exposure to metallic particles in the q-UF size mode was also evidenced (EC, Ni, Mo, Cd), which resulted mainly from outdoor infiltration. The presence of typically mineral elements in q-UF particles (e.g., Fe, Mg, Na, Li, Ti) was also shown in indoor air, probably sourcing from finely ground mineral matter from playgrounds (Rivas et al., 2014). During infiltration, secondary inorganic aerosols underwent a modal shift, towards coarse particles in the case of NO_3^- and towards q-UF particles in the case of SO_4^{2-} .

c) Influence of road traffic

A detailed analysis was carried out for schools located in high- and low-traffic areas of the city, as described in the Methodology section and by Rivas et al. (2014). This analysis was only carried out for outdoor air, aiming to avoid the influence of other variables such as infiltration.

When compared to the mean size distributions obtained for outdoor air, the results from the high traffic schools show a marked shift towards the finer grain-sized fractions, with most of the components evidencing at least one mode in the q-UF particle range (Table 1). This likely reflects the influence of fresh vehicular emissions, characterised by a higher fraction of ultrafine particles. On the other hand, at low traffic schools the prevailing size distributions were dominated by the accumulation mode, suggesting the transport and atmospheric ageing of fresh vehicular emissions towards urban background areas and the subsequent increase in particle size. Thus, the particle size of elements such as Cu, Fe or Rb which showed prevalence for coarse and q-UF particles (pattern 2Cu) at high traffic schools, shifted towards coarser particles only at low traffic schools (pattern 1C for Cu, and 1A for Fe and Rb). Similar cases were observed for elements such as Ca, Mg, Al or Pb. The number of elements showing prevalence for the coarse mode was higher at low traffic with respect to high traffic schools (1C patterns for OC, NO_3^- , Cu, Se, Sn and Sb at low traffic schools vs. for Cl^- and Sb at high traffic schools). Results show that NO_3^- prevailed in the accumulation mode after oxidation from NO_x emission in high traffic areas, and increased towards a coarse distribution after transport toward low traffic areas. Sulphate aerosols, on the other hand, showed maximum mass concentrations in q-UF and accumulation particles at high traffic schools, and shifted toward a unimodal q-UF pattern after ageing by means of a decrease in accumulation mode particles which probably shifted towards the coarse mode, as shown by the relative increase in coarse SO_4^{2-} particles ($0.22 \mu\text{g}/\text{m}^3$ in high-traffic vs. 0.35 in low traffic schools). Because the size distribution of particles outdoors influences indoor air quality through infiltration processes, our results evidence that the location of schools in heavily trafficked areas of the city has an impact on the amount of q-UF particles with the potential to infiltrate indoors, and thus on the increased exposure of schoolchildren to health-hazardous q-UF particles. This result supports the implementation of actions to reduce traffic emissions around schools.

d) Influence of seasonality

The variability in particle size of secondary organic and inorganic species (OC, SO_4^{2-} , NO_3^-) was assessed as a function of seasonality. This variability, induced by outdoor ambient temperature, should have an impact on indoor particles of outdoor origin through infiltration and thus on indoor exposure. To this end, mean outdoor ambient temperatures were calculated for each school and for the period Monday through Thursday, from 9 to 17 h (school hours). Two types of scenarios were then identified: warm periods, with mean outdoor ambient temperature equal to or $>20^\circ\text{C}$ (classified as “summer”; mean daily temperature 23.1°C with 2.2°C standard deviation), and cold periods with ambient temperatures $<20^\circ\text{C}$ (“winter”; mean daily temperature 13.5°C with 3.5°C standard deviation). In total, 23 summer and 53 winter scenarios were analysed.

In outdoor air, the particle size patterns obtained for secondary inorganic aerosols followed expectations (Seinfeld and Pandis, 1998). In winter, under lower ambient temperatures, which favour the thermal stability of NH_4NO_3 , NO_3^- concentrations peaked in the accumulation mode and thus NO_3^- was mostly present in this

form (Fig. 4). Under higher summer temperatures, NH_4NO_3 (accumulation mode) decomposed and coarser NO_3^- species (e.g., Ca or NaNO_3) were formed. Sulphate, on the other hand, did not undergo such a modal shift but the influence of higher summer temperatures was evidenced as an increase in absolute accumulation mode SO_4^{2-} concentrations (mostly $(\text{NH}_4)_2\text{SO}_4$) due to the higher oxidation rate of SO_2 (Hidy, 1994).

The particle sizes and concentrations of outdoor secondary inorganic aerosols influenced indoor air (Fig. 4). Indoor NO_3^- followed a similar although less pronounced pattern than outdoors, with prevalence for accumulation particles (NH_4NO_3) in winter and for coarse particles (e.g., calcium/sodium NO_3^-) in summer. As expected, concentrations were lower indoors than outdoors in winter, when outdoor concentrations reached their annual maxima, with the largest reduction being observed for q-UF and accumulation particles in indoor air (outdoor/indoor ratios of 5.1–5.7 for q-UF and accumulation particles vs. of 3.1 for coarse particles). Therefore outdoor ambient temperatures seem to modify the particle size distribution patterns for NO_3^- aerosols outdoors, and that the same patterns were followed by NO_3^- indoors.

Sulphate aerosols, on the other hand, evidenced different patterns indoors as a function of outdoor temperatures. In outdoor air throughout the year, and indoors in summer, SO_4^{2-} was mostly present in accumulation and q-UF particles as expected. However, in indoor air and in winter, a decrease in accumulation SO_4^{2-} particles was consistently detected, which remains unexplained with the data available. This effect seems to be related with seasonality,

as it was only detected in winter, and it is possibly related with infiltration given that the windows remain closed in winter. I/O ratios >1 were obtained in summer and winter for coarse SO_4^{2-} (CaSO_4), as a result of indoor emissions from blackboard chalk and the high mineral dust re-suspension (Rivas et al., 2014).

Finally, organic aerosols were also assessed, showing prevalence for accumulation and coarse particles in outdoor air (ratios >1.6 for accumulation and coarse over q-UF particles). In indoor air, particle size distribution patterns seemed to be more strongly influenced by indoor sources than by seasonality. Both in winter and summer, indoor OC prevailed in coarse particles originating from human activity as described in the previous sections. Indoor concentrations were higher in winter due to the fact that windows were kept closed. In summer the absolute concentrations of q-UF and accumulation OC particles were similar to outdoor air because windows remained open most of the time, while coarse OC particles were higher indoors (ratio I/O = 1.4). This evidences that, as expected, OC emissions from human activity are mainly coarse.

4. Conclusions

This work aimed to describe particle size distribution patterns of major metals and trace elements in indoor and outdoor air in Barcelona, with a special focus on their emission sources in each particle size range. The main conclusions extracted may be summarised as follows:

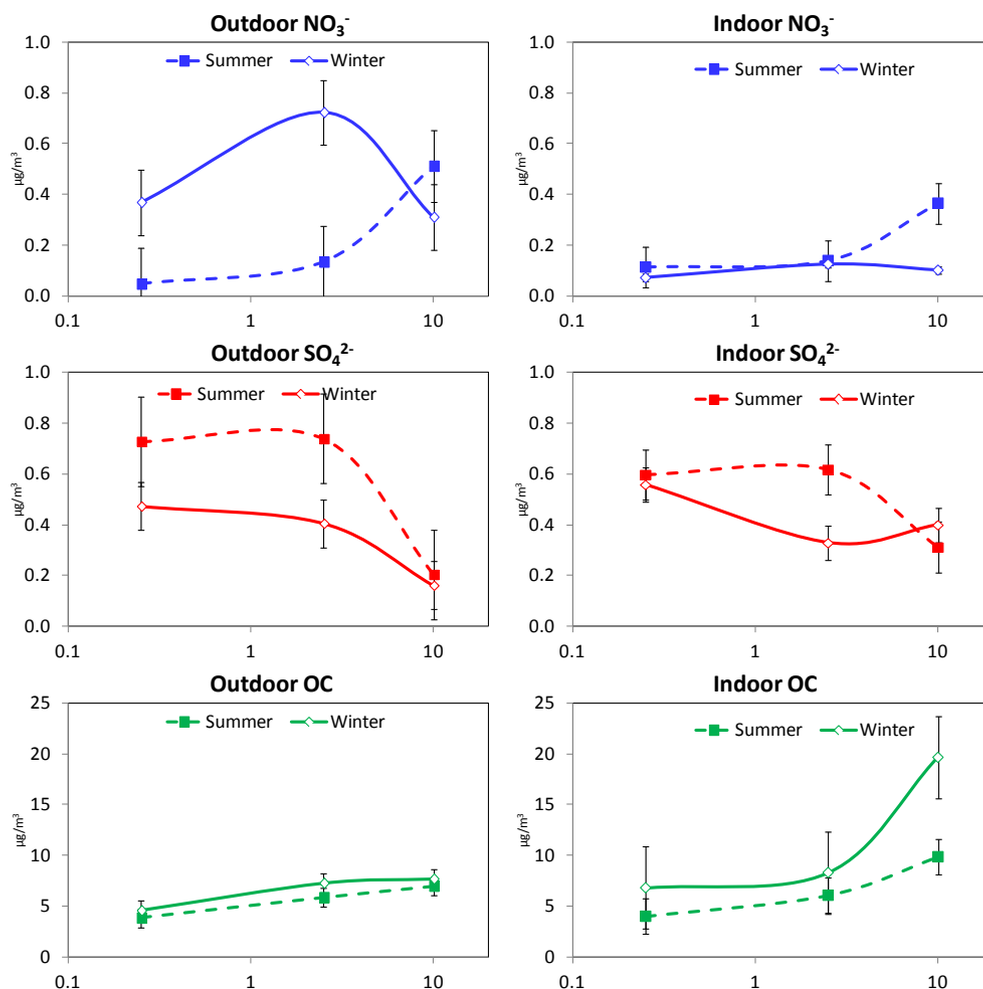


Fig. 4. Particle size distribution patterns identified for secondary organic and inorganic species as a function of seasonality.

- Mineral and typically coarse elements (e.g., Ca, Fe, Al, Mg, Na) were detected in relatively high proportions in indoor and outdoor q-UF particles (20–40% of their mass in q-UF). These elements originate mainly from mineral matter (anthropogenic and natural) and sea salt, and they may also source in trace concentrations from anthropogenic emissions in particles <100 nm. The presence in indoor air of potentially toxic metals in q-UF particles (Mn, Cu, Sn, V, Pb) was also detected, with a potential impact on child exposure.
- Secondary inorganic aerosols, which prevailed in accumulation particles outdoors, evidenced modal shifts (NO_3^- towards coarse particles due to evaporation losses, and SO_4^{2-} towards q-UF particles due to the loss of accumulation particles) after they infiltrated indoors. The influence of an indoor-specific source of coarse sulphate was detected.
- At high traffic schools, size distribution patterns were dominated by q-UF particles reflecting the influence of fresh vehicular emissions. At low traffic schools, accumulation particles prevailed suggesting the transport of fresh vehicular emissions from high traffic areas. Our results demonstrate that the location of schools in heavily trafficked areas increases the amount of q-UF particles with the potential to infiltrate indoors, and thus impacts child exposure to health-hazardous q-UF particles. It is recommended that actions to reduce traffic emissions around schools should be undertaken.
- Outdoor ambient temperatures modified the particle size distribution patterns for NO_3^- aerosols outdoors, and the same patterns were followed by NO_3^- indoors. Sulphate, on the other hand, displayed different patterns indoors with respect to outdoors as a function of seasonality. The particle size distribution patterns for organic carbon in indoor air seemed to be more strongly influenced by emission sources than by seasonality.

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Appendix A. Supplementary data

Supplementary data related to this article can be found at <http://dx.doi.org/10.1016/j.atmosenv.2014.07.027>.

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