Elemental Composition of Suspended Particulate Matter Collected at Two Different Heights above the Ground in A Sub-Urban Site in Kenya

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Abstract
Suspended particulate matter samples were collected in a sub-urban area in Nairobi over a 12 month period at two different heights above ground using a “Gent” SFU sampler. A total of 126 sets of duplicate fine and coarse particulate matter samples were collected. The samples were analysed by energy dispersive x-ray fluorescence (EDXRF) and atomic absorption spectroscopy (AAS) for up to 10 elements. It was found that 66% of the samples collected at two metres and 50% of the samples collected at four metres height exceeded the WHO 24 guideline of 70 μg m⁻³. Reduction in concentration of between 30 to 74 % for Ca, Ti, Zr and Fe were observed both in coarse and fine particulate matter fractions at the higher height. The elements Cu, Zn, Pb and Br represented 0.5 to 1.1 % of the total coarse particulate matter at both heights. Higher proportions of 1.5 to 3.5 % were observed at both heights in the fine particulate matter fraction. High enrichment factors were observed for Cu (10.8 - 228.3), Zn (12.4 - 124.6), Pb (59.4 - 1967) and Br (152.5 - 3038.7) at both heights suggesting anthropogenic activities such as industrial, urban refuse burning, residential and vehicular emissions could be the major contributors. Pb and Br were mainly from the vehicular emissions as indicated by the strong correlations (r > 0.593) and the Br/Pb ratios (0.307 to 0.339).

Keywords: Sub-urban, EDXRF, AAS, Enrichment factors, Heavy metals, particulate matter.

Introduction
Elevated levels of heavy metals have been observed in soils, vegetation and air in the vicinity of certain industries like smelters, mines, battery manufacturing and near busy roads and within urban area in general. The measured levels of heavy metals serve as an index of the extent of pollution (Nguyen et al; 1997; Khandekar et al; 1980). Although some metals such as Mn, Fe, Cu and Zn are essential micronutrients are toxic if exposure levels are sufficiently high (Prasad, 1988).

Numerous studies (Mahadevan et al; 1989; Karue et al; 1992; Noll et al; 1990) have demonstrated that metal contaminated environments are reflected by metal concentrations in suspended particulate matter. In Kenya several studies on elemental characterisation of suspended particulate matter have been done in the city of Nairobi (Karue et al; 1992; Gatebe et al; 1994) but non has ever concentrated on determination of in a slum set-up. This study was designed with an aim of assessing the load of heavy metals in dust on these paths could be at higher risk from inhaling the contaminated dust.

Materials and Methods
Description of the study area
The study was carried out in a slum area about seven (7) kilometers southwest of city centre and about five (5) kilometres southwest of industrial area. The village covers an area of about 3 km², with a population density of approximately 40,881 people km⁻² with a total of 42,722 households (CBS, 1989). Houses are constructed of mud, wood and roofed with corrugated iron sheets. The area is dotted with many cottage industries such as metal works, “Posho” milling, and timber works. There were a total of five sampling sites(A-E)(Fig 1) located on the small open spaces between the residential housing units. The area is criss-crossed with unpaved roads, a railway line, vehicle movements and burning of urban refuse in the open is a common phenomena.

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Sampling procedures

Sampling was done using the “Gent” stacked filter unit (SFU) (Maenhaut et al; 1992; Thomas et al; 1994). The sampler is fitted with an impaction plate which produces a threshold of roughly 10\(\mu\)m for the size of incoming particles. The impaction plate was coated with vaseline oil to reduce particles bouncing off. The PM\(_{10}\) inlet was placed at a height of two metres above the ground for the first seven months and four metres for five months. Two polycarbonate nucpore filters in sequence were used, both of 47mm diameter. These were coarse (8\(\mu\)m pore) and fine (0.4\(\mu\)m pore) diameter to separate the coarse and fine fractions of the particulate matter respectively. The sampling was done over an eight hour period starting at between 8.00 AM and ending at around 5.00 PM in the evening. A total of 126 duplicate coarse and fine particulate matter samples were collected in the five sites.

Weighing of filters

Filters were weighed in an air conditioned laboratory using 1\(\mu\)g sensitivity (Ainsworth Type 24N) weighing balance before and after sampling. Prior to weighing all filters were left to equilibrate for 24 hours in the room. They were irradiated with \(^{241}\)Am \(\alpha\)-emitting source (5\(\mu\)Ci) to remove the static buildup before and after sampling (IAEA, 1996). All filter handling was done using plastic tweezers to avoid contamination by grease from fingers and by heavy metals from metallic tweezers.

Spectral data analysis

A Canberra S-100 PC based multi-channel analyser (MCA) was used for spectral data acquisition and storage. Both the coarse and fine loaded filters were irradiated for 60,000 seconds using \(^{109}\)Cd source. The analysis of the x-ray spectra was done by AXIL (Espen et al; 1985) and quantitative analysis of environmental samples (QAES) for spectral deconvolution and quantitative measurements (Kump, 1993).

Sample analysis by AAS

The loaded filters were digested by boiling gently in 20ml of high purity 70% nitric acid for 2 hours, cooled to room temperature and 10ml of 70% perchloric acid added. This solution was boiled gently for 30 minutes, cooled to room temperature and filled to 50ml mark in 1% nitric acid solution. Quantitative analysis was done by atomic absorption spectrometer, Model AA-680 (Shimadzu Corporation, 1991).

Quality control

The reliability of the analytical procedures was checked by analysis of standard reference materials (SRM-1648). Results of analysis by both Energy Dispersive X-ray Fluorescence (EDXRF) and Atomic Absorption
Spectroscopy (AAS) showed good agreement with the certified values. Values obtained were within 10% range of the certified values. Selected fine and coarse particulate matter samples analysed by EDXRF were reanalysed by AAS. T-tests did not show any significance difference between the values obtained by the two techniques.

**Results and discussions**

**Coarse and fine particulate matter concentrations**

Coarse and fine particulate matter levels (µg m⁻³) variation at two metres over the sampling period for the five sites are summarised graphically in Figures 2 and 3. The levels of coarse particulate matter recorded for each site reflected the location and kind of activities near the site. Site D nearest to unpaved road and several children’s playground the highest mean monthly levels (123.6 to 233.9 µg m⁻³). Site E far away from any road/footpath, recorded low levels with most of the samples collected having a mean monthly values of less than 100 µg m⁻³. This is also noted for the fine suspended particulate matter samples. It should be noted that the ground cover around this site was mainly hence less soil dust. This points to dust blown off unpaved roads to be a major contributor to the load of suspended particulate matter in this area.

Site C located on an open field away from the slum houses recorded values comparable to site B. The levels observed (23.1 to 152.2 µg m⁻³) for this site could be due to several factors among them dust from unpaved roads/footpaths, vehicular emissions and burning refuse nearby.

For the fine particulate matter levels, sites A, B and C recorded comparable values for most of the months (Fig 3). Site D recorded the highest mean monthly levels (20.6 to 36.1 µg m⁻³) strongly suggesting the location and activities near the site were the main determinant of levels observed.

Trends similar to particulate matter loading obtained at two metres height was observed at four metres for both coarse and fine particulate matter samples (Figures 4 and 5). The highest value obtained at four metres was 184 µg m⁻³ at site D as compared to 233.9 µg m⁻³ at two metres height for coarse particulate matter samples.

For the fine particulate matter, sites B, C and D recorded values above 20 µg m⁻³ for most of the months but at four metres values below 20 µg m⁻³ were observed. The maximum mean monthly value observed at four metres was 29 µg m⁻³ observed at two metres. Reduction in particulate loading of between 17 to 45% was observed for the coarse particulate fraction while the fine particulate fraction registered reduction of between 18 to 40%. Resuspended soil dust at the lower height was observed to contribute to the high levels of suspended particulate matter observed.

**Elemental concentrations**

The mean elemental concentration ranges (µg m⁻³) in the coarse and fine particulate matter fractions are given in Tables 1 and 2. The elements K, Ca, Ti, Mn and Fe associated with crustal dust had the highest concentrations at both heights for both coarse and fine particulate matter fractions. Reduction in concentration of between 30 to 72% was observed in the coarse particulate fraction as compared to 25% for Cu, Zn and Br at four metres height.

In the fine particulate matter fraction higher reductions of between 70 to 74% were observed for Ca and 26 to 61% for Fe. However, slight reduction of between 10 to 23% were observed for Ti, Cu, Zn and Pb. Elements K, Ca, Ti, Mn and Fe represented 13.6 to 21.2% of the total coarse particulate mass at two metres and 10 to 18.2% at four metres height. A higher proportion (33.1 to 49.5%) was observed for these elements in the fine particulate fraction at two metres than at four metres (11.2 to 17.3%). This could indicate the effect of resuspended soil dust is more pronounced in the fine particulate range. The elements Cu, Zn, Pb and Br associated with anthropogenic activities represented 0.5 to 1.1% of the total coarse particulate mass at both heights. Higher proportions were observed in the fine particulate fraction at both two metres (1.5 to 3.5%) and four metres (2.3 to 3.1%) height. This could be attributed to the reduced levels of resuspended soil dust, and influence of incoming air masses, especially with the north easterlies prevalent in the area.
Figure 2: Variation of suspended coarse particulate matter levels with time for the various sampling sites at two meters height

Figure 3: Variation of suspended fine particulate matter levels with time for the various sampling sites at two meters height
Figure 4: Variation of suspended course particulate matter levels with time for the various sampling sites at four meters height

Figure 5: Variation of suspended fine particulate matter levels with time for the various sampling sites at four meters height
Correlation coefficients

At two metres height

Correlations between the various crustal elements K, Ca, Ti, Mn, Fe, Zr and SPM levels ranged between 0.612 to 0.984 in the coarse particulate matter fraction. The same elements correlate well (0.515 to 0.927) with SPM levels in the fine particulate fraction for all the sites. However, high correlations (0.658 to 0.773) between Zn, Pb and SPM were observed at site D. This was attributed to probable surface dust contamination by fine particle fallout over time form metal works nearby and lead particles from vehicular emissions in near road. Pairwise correlations between the various crustal elements were > 0.851 in the coarse particulate matter fraction, while in the fine particulate matter fraction they ranged from 0.552 to 0.942. The correlations between Zn, Pb, Cu and Br were much weaker for most of the sites. However, strong correlations between Pb and Br (0.593 to 0.971) were observed at sites B, C, and E in the fine particulate matter fraction.

At four metres height

A similar trend as obtained at two metres height is observed. Strong correlations between Ca, Ti, Mn, Fe, Zr and SPM ranging from 0.514 to 0.970 were observed for both coarse and fine particulate matter fractions for all the sites. However, an exception is Cu which shows high correlations (0.945 to 0.963) in the fine particulate matter fraction for all the sites. This could probably be attributed to transport from industrial area about seven kilometres away. The pairwise correlation coefficients between the various crustal elements Ca, Ti, Mn, Fe and Zr ranged from 0.501 to 0.993 for both coarse and fine particulate matter fractions for all the sites. The correlations between the other elements are much weaker, except for Br, Pb and Cu where the pairwise correlations ranged from 0.514 to 0.932 in the coarse particulate fraction in sites B, C and E.

Br/Pb Ratios

Site D had some of the highest Br/Pb ratios (0.354 to 0.387) at both heights in both the coarse and fine particulate matter fractions. This could be due to the fact that this site was nearest to a busy road (Fig 1). At both heights higher Br/Pb ratios were observed in the coarse particulate matter fraction. Higher Br/Pb ratios were also observed at the lower height for both particulate matter fractions. This could be due to the vaporization of Br at the lower height (i.e., by the warm surface) hence more of the Br could be in the vapor phase at the higher height.

Automotive emissions are the most likely sources for Br and Pb as suggested by the good correlation between these two elements ($r = 0.593\pm 0.971$) at some of the sites and the average Br/Pb ratios of (0.339 ± 0.023, n=50) for coarse particulate matter samples and (0.307 ± 0.035, n=50) for the fine particulate matter samples at two metres height. These Br/Pb ratios are similar to those observed in other sub-urban and urban areas in the world, 0.35 to 0.39 in cities and 0.25 in suburbs, 0.309 to 0.535 in a city (Karue et al; 1992; Gatebe et al; 1996).

Enrichment factors

At two metres height

Crustal enrichment factors (Efs) relative to Fe and Mason’s average crustal rock composition (Mason, 1966) were calculated. In both the coarse and fine particulate matter fractions the Efs for elements K, Ca, Ti, Mn and Zr were less than 6 (Table 3 and 4). These elements apart from Zr registered high concentrations in the coarse particulate matter fraction. However, Mn shows marginal enrichment (6.2 to 7.67) in the coarse particulate matter fraction and Ca (6.28 to 9.64) in the fine particulate matter fraction respectively. This points to probable local anthropogenic sources for these elements. These probably includes the frequent wood/charcoal/kerosine fires and burning of urban refuse. The other elements Cu, Zn, Pb and Br have high Efs values (12.4 to 1812) particularly in the fine particulate matter fraction and various anthropogenic sources are probably responsible for their atmospheric concentrations. Cu and Zn could mostly be attributed to local anthropogenic activities such as the metal welding works but dust transport from the industrial area about 3 kms away by the prevalent north easterlies cannot be ruled out. Pb and Br are probably due to vehicular emissions.

At four metres height

A similar trend as observed for samples collected at two metres is evident. K, Ca, Ti, Mn and Zr showed low enrichment factors both in the coarse and fine particulate matter fractions. Cu, Zn, Pb and Br showed high enrichment factors. However marginal enrichment was observed for Mn (10.5 to 17) in the fine particulate matter fraction indicating possible anthropogenic sources. Higher enrichment factors were observed for Cu, Zn, Pb and Br at four metres height than at two
metres in the fine particulate matter fraction. This could be due to the longer residence time of the fine particles in the air.

**Conclusion**

Similar suspended particulate matter (SPM) concentration profiles were observed at both heights for all the sites sampled. Site D recorded the highest levels while site E had the lowest at both heights. Reduction in particulate loading of between 17 to 45 % was observed for the coarse particulate matter fraction while a reduction of between 18 to 40 % were registered for the fine particulate matter fraction.

Elements K, Ca, Ti, Mn and Fe had the highest concentrations at both heights for both coarse and fine particulate matter fractions. Reduction in concentration of between 30 to 72 % for Ca, Ti, and Zr was observed in the coarse particulate fraction, while Cu, Zn and Br registered a reduction of 25 % in the coarse particulate matter fraction. For the fine particulate fraction higher reductions of between 70 to 74 % were observed for Ca and 26 to 61 % for Fe. Elements Cu, Zn, Pb and Br associated with anthropogenic activities represented 0.5 to 1.1 % of the total coarse particulate mass at both heights. Higher proportions were observed in the fine particulate matter fraction at both two and four metres height. At both heights higher Br/Pb ratios were observed in the coarse particulate matter fraction. Higher Br/Pb ratios were also observed at the lower height for both particulate matter fractions.

Automotive emissions are the most likely sources of Br and Pb as indicated by the good correlation ($r > 0.593$) observed at some sites and the Br/Pb ratios (0.339 ± 0.023) for coarse particulate matter and (0.307 ± 0.035) for fine particulate matter samples. Cu and Zn were probably from anthropogenic activities within and outside the area.

**References**


Maenhaunt W; Filip, F and Jan C. (1992). Institute for nuclear sciences, University of Gent, Belgium. The “Gent” Stack Filter Unit (SFU) sampler for the collection of atmospheric aerosols in two size fractions. Description and instructions for installation and use. IAEA Co-ordinated Research programme, CRP E4-10.08.


Acknowledgements
This work could not have been possible without the cooperation of the owners of the five houses in the study area who provided electricity outlet to power the sampling pump and the Institute of Nuclear Science (University of Nairobi) for providing funds and the EDXRF instrument for analysis of the samples.

Table 1: Mean elemental concentration (µg m⁻³) range at heights of two and four metres for the suspended coarse particulate matter samples

<table>
<thead>
<tr>
<th>Element</th>
<th>Two (2) metres</th>
<th>Four (4) metres</th>
</tr>
</thead>
<tbody>
<tr>
<td>K</td>
<td>1.435 –4.533 (1.804)</td>
<td>--------</td>
</tr>
<tr>
<td>Ca</td>
<td>5.185 –7.104 (7.054)</td>
<td>1.55 - 1.97 (1.745)</td>
</tr>
<tr>
<td>Ti</td>
<td>0.387 –0.809 (0.419)</td>
<td>0.233 -0.541 (0.447)</td>
</tr>
<tr>
<td>Mn</td>
<td>0.381 –0.790 (0.526)</td>
<td>0.277 - 0.656 (0.455)</td>
</tr>
<tr>
<td>Fe</td>
<td>3.485 –6.025 (5.148)</td>
<td>2.31 - 6.29 (4.125)</td>
</tr>
<tr>
<td>Cu</td>
<td>0.119 –0.169 (0.135)</td>
<td>0.095 - 0.117 (0.102)</td>
</tr>
<tr>
<td>Zn</td>
<td>0.143 –0.247 (0.195)</td>
<td>0.118 - 0.165 (0.157)</td>
</tr>
<tr>
<td>Pb</td>
<td>0.168 –0.337 (0.243)</td>
<td>0.120 - 0.222 (0.156)</td>
</tr>
<tr>
<td>Br</td>
<td>0.029 –0.064 (0.059)</td>
<td>0.045 - 0.054 (0.049)</td>
</tr>
<tr>
<td>Zr</td>
<td>0.046 –0.152 (0.096)</td>
<td>0.036 - 0.095 (0.071)</td>
</tr>
</tbody>
</table>

(Median value shown in parenthesis) *K at four metres could not be used for intercomparison because it was only detected in two of the five months sampled at only two of the four sites.

Table 2: Mean elemental concentration (µg m⁻³) range at heights of two and four metres for the suspended fine particulate matter samples

<table>
<thead>
<tr>
<th>Element</th>
<th>Two (2) metres</th>
<th>Four (4) metres</th>
</tr>
</thead>
<tbody>
<tr>
<td>K</td>
<td>1.503 -2.25 (1.78)</td>
<td>LDL</td>
</tr>
<tr>
<td>Ca</td>
<td>4.77 -6.298 (5.481)</td>
<td>1.45 -1.65 (1.550)</td>
</tr>
<tr>
<td>Ti</td>
<td>0.133 -0.280 (0.238)</td>
<td>0.222 -0.245 (0.234)</td>
</tr>
<tr>
<td>Mn</td>
<td>0.079 -0.110 (0.095)</td>
<td>0.107 -0.192 (0.123)</td>
</tr>
<tr>
<td>Fe</td>
<td>0.862 -1.102 (0.982)</td>
<td>0.326 -0.818 (0.518)</td>
</tr>
<tr>
<td>Cu</td>
<td>0.075 -0.135 (0.088)</td>
<td>0.088 -0.115 (0.099)</td>
</tr>
<tr>
<td>Zn</td>
<td>0.106 -0.150 (0.132)</td>
<td>0.081 -0.116 (0.110)</td>
</tr>
<tr>
<td>Pb</td>
<td>0.165 -0.250 (0.233)</td>
<td>0.162 -0.225 (0.187)</td>
</tr>
<tr>
<td>Br</td>
<td>0.031 -0.055 (0.052)</td>
<td>0.031 -0.034 (0.033)</td>
</tr>
<tr>
<td>Zr</td>
<td>0.011 -0.027 (0.016)</td>
<td>0.012 -0.022 (0.016)</td>
</tr>
</tbody>
</table>

(Median value shown in parenthesis)(LDL-low detection limits)