



1 **Atmospheric trace metals measured at a regional**
2 **background site (Welgegund) in South Africa**

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10

11 **Abstract**

12 Atmospheric trace metals can cause a variety of health-related and environmental problems.
13 Only a few studies on atmospheric trace metal concentrations have been conducted in South
14 Africa. Therefore the aim of this study was to determine trace metals concentrations in aerosols
15 collected at Welgegund, South Africa. PM₁, PM_{1-2.5} and PM_{2.5-10} samples were collected for 13
16 months and 31 atmospheric trace metal species were detected. Atmospheric iron (Fe) had the
17 highest concentrations in all three size fractions, while calcium (Ca) was the second most
18 abundant species. Chromium (Cr) and sodium (Na) concentrations were the third and fourth
19 most abundant species, respectively. The concentrations of the trace metal species in all three
20 size ranges were similar, with the exception of Fe that had higher concentrations in the PM₁
21 size fraction. With the exception of titanium (Ti), aluminium (Al) and manganese (Mg), 70%
22 or more of the trace metal species detected were in the smaller size fractions, which indicated
23 the influence of industrial activities. However, the large influence of wind-blown dust was
24 reflected by 30% and more of trace metals being present in the PM_{2.5-10} size fraction.
25 Comparison of trace metals determined at Welgegund to those in the western Bushveld Igneous
26 Complex indicated that at both locations similar species were observed with Fe being the most
27 abundant. However, concentrations of these trace metal species were significantly higher in the
28 western Bushveld Igneous Complex. Fe concentrations at the Vaal Triangle were similar to
29 levels thereof at Welgegund, while concentrations of species associated pyrometallurgical



1 smelting were lower. Annual average Ni was four times higher and annual average As was
2 marginally higher than their respective European standards limit values, which could be
3 attributed to regional influence of pyrometallurgical industries in the western Bushveld Igneous
4 Complex. All three size fractions indicated elevated trace metal concentrations coinciding with
5 the end of the dry season, which could partially be attributed to decreased wet removal and
6 increases in wind generation of particulates. Principal component factor analysis (PCFA)
7 revealed four meaningful factors in the PM_1 size fraction, i.e. crustal, pyrometallurgical-related
8 and Au slimes dams. No meaningful factors were determined for the $PM_{1-2.5}$ and $PM_{2.5-10}$ size
9 fractions, which was attributed to the large influence of wind-blown dust on atmospheric trace
10 metals determined at Welgegund. Pollution roses confirmed the influence of wind-blown dust
11 on trace metal concentrations measured at Welgegund, while the impact of industrial activities
12 was also substantiated.

13

14 **1 Introduction**

15 Atmospheric aerosols are either directly emitted into the atmosphere (primary aerosols) from
16 natural and/or anthropogenic sources, or are formed through gaseous reactions and gas-to-
17 particle conversions (secondary aerosols). Aerosols have high temporal and spatial variability,
18 which increases the need and importance for detailed physical and chemical characterisation
19 on a regional scale in order to assess the impacts of aerosols (Pöschl, 2005). Particulate matter
20 (PM) is classified according to its aerodynamic diameter, as PM_{10} , $PM_{2.5}$, PM_1 and $PM_{0.1}$,
21 which relates to aerodynamic diameters being smaller than 10, 2.5, 1 and 0.1 μm , respectively.
22 Larger particulates have shorter lifetimes in the atmosphere compared to smaller particles,
23 while the impacts of these species are also determined, to a large degree, by their size (Tiwari
24 et al., 2012, Colbeck et al., 2011). The largest uncertainties in the estimation of direct and
25 indirect radiative forcing from aerosols are related to the insufficient knowledge of the high
26 spatial and temporal variability of aerosol concentrations, as well as their microphysical,
27 chemical and radiative properties (IPCC 2014). Aerosols consist of a large number of organic
28 and inorganic compounds, of which typical inorganic species include ionic species and trace
29 metals.

30 Natural sources of atmospheric trace metals include mineral dust, crustal species, oceans and
31 biomass burning (wild fires), while major anthropogenic sources are pyrometallurgical



1 processes, fossil fuel combustion and incineration (Pacyna and Pacyna, 2001). Larger aerosol
2 particles ($>2.5 \mu\text{m}$) are usually associated with natural emissions through processes such as
3 rock weathering and soil erosion (Nriagu et al., 1989). Trace metal species usually associated
4 with natural emissions include sodium (Na), silicon (Si), magnesium (Mg), aluminium (Al),
5 potassium (K), calcium (Ca), titanium (Ti), chromium (Cr), manganese (Mn) and iron (Fe)
6 (Adgate et al., 2007). Arsenic (As), barium (Ba), cadmium (Cd), copper (Cu), nickel (Ni), zinc
7 (Zn), vanadium (V), molybdenum (Mo), mercury (Hg) and lead (Pb) are mostly related to
8 anthropogenic activities (Pacyna (1998); Polidori et al., 2009). One of the most significant
9 sources of anthropogenic trace metal emissions is the industrial smelting of metals. Industrial
10 pyrometallurgical processes produce the largest emissions of As, Cd, Cu, Ni and Zn (Zahn et
11 al., 2014). Cr, Ba, Mo, Zn, Pb and Cu are typically associated with motor-vehicle emissions
12 and oil combustion, while Fe, Pb and Zn are emitted from municipal waste incinerators (Adgate
13 et al., 2007). However, most of these atmospheric trace metals are emitted through a
14 combination of different anthropogenic sources (Polidori et al., 2009).

15 Although trace heavy metals, i.e. metals $> \text{Ca}$, represent a relatively small fraction of
16 atmospheric aerosols (with the exception of Fe that could contribute a few percent) (Colbeck,
17 2008), these species can cause a variety of health-related and environmental problems, which
18 depends on the aerosol composition, extent and time of exposure (Pöschl, 2005). The potential
19 hazard of several toxic species is well documented as discussed, for instance, by Polidori et al.
20 (2009), indicating that trace metals such as As, Cd, Co, Cr, Ni, Pb and Se are considered human
21 and animal carcinogens even in trace amounts (CDC, 2015). It has also been shown that Cu,
22 Cr and V can generate reactive oxygenated species that can contribute to oxidative DNA
23 damage (Nel, 2005). Furthermore, trace metals such as Cr, Fe and V have several oxidation
24 states that can participate in many atmospheric redox reactions (Seigneur & Constantinou,
25 1995), which can catalyse the generation of reactive oxygenated species (ROS) that have been
26 associated with direct molecular damage and with the induction of biochemical synthesis
27 pathways (Rubasinghege et al., 2010). Guidelines for atmospheric levels of many trace metals
28 are provided by the World Health Organization (WHO) (WHO 2005). In addition, lighter
29 metals such as Si, Al and K are the most abundant crustal elements (next to oxygen), which
30 can typically constitute up to 50% of remote continental aerosols. These species are usually
31 associated with the impacts of aerosols on respiratory diseases and climate.



1 South Africa has the largest industrialised economy in Africa, with significant mining and
2 metallurgical activities. South Africa is a well-known source region of atmospheric pollutants,
3 which is signified by three regions being classified through legislation as air pollution priority
4 areas, i.e. Vaal Triangle Airshed Priority Area (Government Gazette, 2006), Highveld Priority
5 Area (Government Gazette, 2007) and Waterberg-Bojanala Priority Area (Government
6 Gazette, 2012). Air quality outside these priority areas is often adversely affected due to
7 regional transport and the general climatic conditions, such as low precipitation and poor
8 atmospheric mixing in winter. Only a few studies on the concentrations of atmospheric trace
9 metals in South Africa have been conducted (Van Zyl et al., 2014; Kgabi, 2006; Kleynhans,
10 2008). In addition, these studies were also conducted within two of these priority areas
11 containing a significant number of large point sources, and regional impacts of atmospheric
12 trace metals could therefore not be assessed.

13 In this study, trace metals were determined in three size ranges in aerosol samples collected for
14 one year at the Welgegund atmospheric measurement station in South Africa. Welgegund is a
15 comprehensively equipped regional background atmospheric measurement station that is ~100
16 km downwind of the most important source regions in the interior of South Africa (e.g. Tiitta
17 et al., 2014). In an effort to determine major sources of trace metals on a regional scale, source
18 apportionment was also performed by applying principal component factor analysis (PCFA).

19

20 **2 Experimental**

21 **2.1 Site description**

22 Aerosol sampling was performed at Welgegund (www.welgegund.org, 26°34'11.23"S,
23 26°56'21.44"E, 1480 m a.s.l. (above sea level)) in South Africa. As indicated in Figure 1,
24 Welgegund is situated in the interior of South Africa and is frequently affected by air masses
25 moving over the most important anthropogenic/industrial source regions in the interior (Beukes
26 et al., 2013, Tiitta, et al., 2014, Jaars, et al., 2014, Vakkari et al., 2015; Booyens et al., 2015).
27 Also indicated in Figure 1 are the major industrial point sources, i.e. coal-fired power plants,
28 petrochemical industries and pyrometallurgical smelters. In Beukes et al. (2013) Tiitta, et al.,
29 2014 and Jaars, et al., 2014, reasons for the site selection, prevailing biomes and pollution
30 sectors are discussed in detail. In summary, air masses affecting the site from the west, between



1 north- and south-west, are considered to be representative of the regional background, since
2 they move over a sparsely populated region without any large point sources. In the sector
3 between north and north-east from Welgegund lays the western limb of the Bushveld Igneous
4 Complex, which holds eleven pyrometallurgical smelters (most commonly related to the
5 production of Cr, Fe, V and Ni) within a ~55 km radius, in addition to other industrial, mining
6 and residential sources. In the north-east to eastern sector, the Johannesburg-Pretoria (Jhb-Pta)
7 conurbation is situated, which is inhabited by more than 10 million people, making it one of
8 the forty largest metropolitan areas in the world. In the sector between east and south-east from
9 Welgegund is the Vaal Triangle region, where most of the South African petrochemical and
10 petrochemically-related industries are located, together with other large point sources, such as
11 two coal-fired power stations (without de-SO_x and de-NO_x) and large pyrometallurgical
12 smelters. Welgegund is also affected by the Mpumalanga Highveld in the eastern sector
13 (indicated by MP in Figure 1). In this region, there are 11 coal-fired power stations (without
14 de-SO_x and de-NO_x technologies) with a combined installed generation capacity of ca. 46 GW,
15 as well as a very large petrochemical plant, several pyrometallurgical smelters and numerous
16 coal mines, all within a ca. 60 km radius. Furthermore, Welgegund is also affected by air
17 masses passing over the pyrometallurgical smelters in the eastern limb of the Bushveld Igneous
18 Complex situated north-east from Welgegund in the Limpopo Province (indicated by LP in
19 Figure 1).

20

21 **Insert Figure 1**

22

23 **2.2 Sampling and analysis**

24 Aerosol samples were collected for one year from 24 November 2010 until 28 December 2011.
25 A Dekati (Dekati Ltd., Finland) PM₁₀ cascade impactor (ISO23210) equipped with PTFE filters
26 was used to collect different particulate size ranges, i.e. PM_{2.5-10} (aerodynamic diameter ranging
27 between 2.5 and 10 μm), PM_{1-2.5} (aerodynamic diameter ranging between 1 and 2.5 μm) and
28 PM₁ (aerodynamic diameter <1 μm). The pump flow rate was set at 30 L min⁻¹. Samples were
29 collected continuously for one week, after which filters were changed. A total of 54 samples
30 were collected for the 54-weeks sampling period for each of the three size ranges. The trace



1 metals in the PM collected on the 216 PTFE filters were extracted by hot acid leaching (20 ml
2 HNO₃ and 5 ml HCl) and diluted in deionised water (18.2 MΩ) up to 100 mL for subsequent
3 analysis with an inductively coupled plasma mass spectrometer (ICP-MS). In total, 31 trace
4 metals could be detected with ICP-MS analysis, which included Na, Mg, Al, K, Ca, Ti, Cr, Mg,
5 Fe, As, Ba, Cd, Cu, Ni, Zn, V, Mo, Hg, Pb, manganese (Mn), cobalt (Co), platinum (Pt),
6 beryllium (Be), boron (B), selenium (Se), palladium (Pd), barium (Ba), gold (Au), thallium
7 (Tl), antimony (Sb) and uranium (U). Trace metal concentrations below the detection limit of
8 the ICP-MS were considered to have concentrations half the detection limit of the species
9 considered. This is a precautionary assumption that is frequently used in health-related
10 environmental studies (e.g. Van Zyl et al., 2014).

11 **2.3 Statistical analysis**

12 In an attempt to identify possible sources of trace metals detected, PCFA with Varimax rotation
13 (v. 13.0 SPSS Inc., Chicago, IL, USA) was performed on the dataset. PCFA has been used
14 widely in receptor modelling to identify major source sectors. The technique operates on
15 sample-to-sample fluctuations of the normalised concentrations. It does not directly yield
16 concentrations of species from various sources, but identifies a minimum number of common
17 factors for which the variance often accounts for most of the variance of species (e.g. Van Zyl
18 et al., 2014 and references therein). The trace metal concentrations determined for the 32
19 species in all three size fractions were subjected to multivariate analysis of Box-Cox
20 transformation and Varimax rotation, followed by subsequent PCFA. In addition, Spearman
21 correlations were also performed in order to establish correlations between trace metals in order
22 to substantiate results obtained with PCFA.

23

24 **3 Results**

25 **3.1 Size-resolved trace metal concentrations**

26 Although nitric digestion is commonly used to extract and dissolve metals for ICP-MS analysis,
27 it is unable to dissolve and extract silicate minerals. Therefore Si could not be quantified in this
28 study. In addition, this limitation of the nitric digestion could also result in determining lower
29 concentrations of metals associated with the silicate component such as Al and K. It is



1 estimated that approximately only 7 % Si and 30 % Al is extracted by nitric acid leaching (Ahn
2 et al., 2011). Therefore, since Si and Al are considered to be the most abundant crustal elements
3 after oxygen, the trace metal concentrations presented in this paper should be related to the
4 limitation of nitric digestion, i.e. Si-Al-K components missing from the digestions phase.
5 Silicate minerals can be dissolved in a mixture of aqua regia and hydrofluoric acid. However,
6 this is a very difficult procedure, which results in the formation of gaseous SiF_3 that is not
7 determinable by ICP-MS.

8 In Figure 2, the combined trace metal concentrations in all three size fractions (Figure 2 (a)),
9 as well as concentrations of the trace metals determined in each of the size fractions are
10 presented (Figure 2 (b), (c) and (d)). Hg and Ag concentrations were below the detection limit
11 of the analytical technique for the entire sampling period in all three size fractions and the
12 concentrations of these species are therefore excluded from Figure 2.

13

14 **Insert Figure 2**

15

16 The highest median concentration was determined for atmospheric Fe, i.e. $1.4 \mu\text{g m}^{-3}$, while
17 Ca was the second most abundant species with a median concentration of $1.1 \mu\text{g m}^{-3}$. Fe
18 concentrations were significantly higher compared to the other trace metal species determined
19 at Welgegund. Cr and Na concentrations were the third and fourth most abundant species,
20 respectively. The median Cr concentration was $0.54 \mu\text{g m}^{-3}$, while the median Na level was
21 $0.39 \mu\text{g m}^{-3}$. Relatively higher concentrations were also determined for Al, B, Mg, Ni and K
22 with median concentrations of $0.20 \mu\text{g m}^{-3}$, $0.30 \mu\text{g m}^{-3}$, $0.18 \mu\text{g m}^{-3}$, $0.02 \mu\text{g m}^{-3}$ and $0.18 \mu\text{g m}^{-3}$,
23 respectively. The combined atmospheric concentrations of the other trace metals in all the
24 size fractions were clearly lower.

25 A comparison of the trace metal concentrations in the three size fractions indicates that Fe and
26 Ca were the most abundant species in all three size fractions. Fe had the highest median
27 concentration in the PM_1 size fraction, i.e. $0.63 \mu\text{g m}^{-3}$, while Ca had the highest median
28 concentrations in the $\text{PM}_{1-2.5}$ and $\text{PM}_{2.5-10}$ size fractions, i.e. $0.39 \mu\text{g m}^{-3}$ and $0.29 \mu\text{g m}^{-3}$,
29 respectively. The median concentration of Fe in the PM_1 was significantly higher compared to
30 the median concentrations thereof in the $\text{PM}_{1-2.5}$ and $\text{PM}_{2.5-10}$ size fractions. The third and fourth



1 most abundant species in all three size fractions were Cr and Na, respectively. Relatively higher
2 concentrations were also determined for Al, B, Mg, Ni and K in all three size fractions. With
3 the exception of Fe concentrations in the PM₁ size fraction, the concentrations of each of the
4 trace metal species were similar in all size fractions.

5 A major source of the trace metal species with elevated levels in all three size fractions can be
6 considered to be wind-blown dust. Trace metal species typically associated with wind-blown
7 dust include Fe, Ca, Mg, Al and K. As mentioned, Welgegund is a regional background
8 location affected by air masses passing over large pollutant source regions and a relatively
9 clean background area (Figure 1). It is therefore expected that wind-blown dust could have a
10 major impact on atmospheric trace metal concentrations. In addition, the western Bushveld
11 Igneous Complex is a major source region affecting Welgegund, with a large number of
12 pyrometallurgical smelters and mining activities (Tiitta et al., 2014; Jaars et al., 2014). This
13 source region could contribute to regional elevated levels of Fe, Cr, Ni, Zn, Mn and V measured
14 at Welgegund. The possible sources of trace metal species measured at Welgegund will be
15 further explored in section 3.5.

16

17 3.2 Size distribution of trace metals

18 In Figure 3, the mean size distributions of each of the trace metal species identified above the
19 detection limit in the three size fractions are presented. Ti had significantly higher contribution
20 (80%) in the PM_{2.5-10} size fraction, while Al and Mg also had relatively higher contributions
21 (~50 and 45%, respectively) in the PM_{2.5-10} size fraction. The PM_{2.5-10} size fraction is usually
22 associated with wind-blown dust typically comprising Al, Fe, Na, Mg and Ti (Polidori et al.,
23 2009). 70% or more of all the other trace metal species detected were in the two smaller size
24 fractions, with approximately 35 to 60% occurring in the PM₁ size fraction. The presence of
25 these trace metal species predominantly in the smaller size fractions, especially considering the
26 relatively large contribution in the PM₁ size fractions, indicates the influence of industrial (high
27 temperature) activities on air masses measured at Welgegund. However, the large influence of
28 wind-blown dust on trace metal concentrations determined at Welgegund is also reflected with
29 approximately 30% of most of these trace metals being present in the PM_{2.5-10} size fraction.
30 Trace metal concentrations measured at Marikana (van Zyl et al., 2014) indicated that Cr, Mn,



1 V, Zn and Ni occurred almost exclusively in the PM_{2.5} size fraction, with no contribution by
2 coarser particles.

3

4 **Insert Figure 3**

5

6 **3.3 Comparison to previous studies and ambient air quality standards**

7 In Table 1, the annual average PM₁₀ trace metal concentrations determined in this study are
8 compared to trace metal concentrations determined in other studies. Although the aerosol
9 sampling periods and frequencies for most of these previous trace metal studies were not
10 similar to the aerosol sampling period and frequency in this investigation, these results could
11 be utilised to contextualise the trace metal concentrations. As mentioned previously, Hg and
12 Ag concentrations were below the detection limit of the analytical technique for the entire
13 sampling period in all three size fractions. Therefore, concentrations presented for these species
14 are most likely to be an over estimate due to the precautionary assumption.

15

16 **Insert Table 1**

17

18 The annual mean PM₁₀ trace metal concentrations at Welgegund (Table 1) were typically lower
19 than previous studies conducted in South Africa (Kgabi, 2006; Kleynhans, 2008; Van Zyl et
20 al., 2014). This is expected, as Welgegund is a regional background location and the previous
21 studies were conducted at sites within two priority areas, as mentioned previously. These sites
22 were also located in two of the major source regions influencing air masses arriving at
23 Welgegund. Marikana (Van Zyl et al., 2014) and Rustenburg (Kgabi, 2006) are situated
24 approximately 100 km north-north-west from Welgegund within the western Bushveld Igneous
25 Complex source region, while the site in the Vaal Triangle (Kleynhans, 2008) source region is
26 situated approximately 90 km east from Welgegund.

27 Fe was also the most abundant species at Marikana and Rustenburg, with significantly higher
28 concentrations compared to Welgegund. Mg was the second most abundant species at
29 Marikana, with Mg concentrations being an order of magnitude higher than levels thereof at



1 Welgegund, while Mn and Cr concentrations were the second and third highest, respectively
2 at Rustenburg. Cr levels at Rustenburg were approximately 2.5 times higher than levels thereof
3 at Welgegund. However, Cr concentrations measured at Welgegund were approximately two
4 times higher compared to Cr levels determined at Marikana, which could be attributed to the
5 contribution of Cr units from wind-blown mineral dust at Welgegund. Ni and Zn concentrations
6 at Welgegund were an order of magnitude lower compared to levels thereof at Marikana and
7 Rustenburg. Mn and V concentrations determined at Welgegund were significantly lower
8 compared levels thereof measured at Rustenburg. V levels measured at Marikana were similar
9 to concentrations at Marikana, while Mn levels were two times higher at Marikana. Similar to
10 Welgegund, Na, B and Al were also relatively abundant at Marikana with concentrations of
11 these species an order of magnitude higher at Marikana. Ca concentrations determined at
12 Welgegund were similar to the levels thereof determined at Marikana, while K levels were
13 three times higher at Marikana.

14 Atmospheric Na had the highest concentrations in the Vaal Triangle, while Fe and K were the
15 second and third most abundant species, respectively. Fe concentrations were similar at Vaal
16 Triangle than levels thereof at Welgegund, while the annual average Na concentration was
17 seven times higher and the annual average K level was an order of magnitude higher at the
18 Vaal Triangle. In addition, Mg concentrations were approximately five times higher in the Vaal
19 Triangle. Cr, Ni and Zn that are typically associated with pyrometallurgical industries were
20 significantly lower in the Vaal Triangle compared to levels thereof at Welgegund. However,
21 Mn concentrations at the Vaal Triangle were higher compared to levels thereof at Welgegund
22 and Marikana. This can be attributed to the presence of a ferromanganese (FeMn) smelter in
23 the Vaal Triangle region, as indicated in Figure 1.

24 The atmospheric trace metal concentrations determined at Welgegund were also compared to
25 measurements at regional background sites near Beijing, China (Duan et al., 2012), the west
26 coast of Portugal (Pio et al., 1996) and Spain (Querol et al. 2007). Al concentrations near
27 Beijing were significantly higher compared to other trace metal species, while Na was the
28 second most abundant species. Elevated levels of K, Fe and Ca were also determined near
29 Beijing. Al, Na and K concentrations were an order of magnitude higher compared to levels of
30 these species determined at Welgegund. Fe levels were twice as low near Beijing, while Mg
31 concentrations were three times higher. Ca, Pb and Mn concentrations at Welgegund were
32 similar to levels thereof near Beijing. All the other trace metal species measured near Beijing



1 were an order or two orders of magnitude lower compared to concentrations of these species
2 at Welgegund. Annual average trace metal concentrations determined at the two European
3 regional background sites were an order or two orders of magnitude lower compared to trace
4 metal levels determined at Welgegund. The generally lower trace metal concentration
5 determined at these sites in China and Europe compared to Welgegund can be attributed to the
6 sites in China and Europe being more removed from a conglomeration of metal sources.

7 Also indicated in Table 1 are the existing ambient air quality guidelines and standard limit
8 values for trace metal species prescribed by the WHO air quality guidelines for Europe (WHO,
9 2005), the European Commission Air Quality Standards (ECAQ, 2008) and the South African
10 National Air Quality Standards of the South African Department of Environmental Affairs
11 (DEA) (Government Gazette, 2009). There are currently only guidelines and standards for
12 seven trace metal species, of which each of the above-mentioned institutions only prescribe
13 limit values for some of these trace metal species. Comparison of the annual average trace
14 metal concentrations determined at Welgegund with the annual average standard limit values
15 indicates that Ni and As exceeded standard limits set by the European Commission of Air
16 Quality Standards. The annual average Ni concentration of $0.079 \mu\text{g m}^{-3}$ were approximately
17 four times higher than the European standard limit value of $0.02 \mu\text{g m}^{-3}$, while the annual
18 average As level of $0.0084 \mu\text{g m}^{-3}$ marginally exceeded the annual standard limit of $0.006 \mu\text{g m}^{-3}$.
19 These exceedances can most probably be ascribed to the regional impacts of
20 pyrometallurgical activities in the Bushveld Igneous Complex. Van Zyl et al. (2014) indicated
21 that the exceedance of Ni at Marikana situated within the western Bushveld Igneous Complex
22 could be attributed to base metal refining.

23 The WHO guideline of $2.5 \times 10^4 \mu\text{g m}^{-3}$ listed for Cr is only for atmospheric concentrations of
24 Cr(VI) with a lifetime risk of 1:1 000 000. The $0.50 \mu\text{g m}^{-3}$ annual average Cr concentration
25 determined can therefore not be compared to the guideline, since this value represents the total
26 atmospheric Cr concentrations in all the oxidation states. V only has a 24-hour standard limit
27 value. Therefore, V concentrations determined in this study cannot directly be compared to this
28 standard limit. However, the 24-hour average calculated from the highest weekly V
29 concentration ($0.084 \mu\text{g m}^{-3}$) was $0.012 \mu\text{g m}^{-3}$, which was two orders of magnitude lower than
30 the 24-hour V standard limit of the European Commission Air Quality Standards.



1 Since Pb is the only trace metal for which a South African ambient air quality standard limit
2 exists, it must also be noted that Pb concentrations did not exceed any standard limit. The
3 annual average Pb concentrations determined at Welgegund ($0.0078 \mu\text{g m}^{-3}$) were an order of
4 magnitude lower than levels thereof at Marikana and Vaal Triangle, and three orders of
5 magnitude lower than Pb levels determined at Rustenburg. However, the annual average Pb
6 concentrations at Vaal Triangle, Marikana and Rustenburg were below the standard limit
7 (Kleynhans, 2008; Van Zyl et al., 2014; Kgabi, 2006). These low Pb concentrations can be
8 partially ascribed to de-leading of petrol in South Africa. Furthermore, Pb concentrations
9 determined at Beijing were similar to levels thereof determined at Welgegund.

10 Since the measurement of the ambient Hg concentrations is receiving increasing attention in
11 South Africa and it is foreseen that a standard limit value for Hg levels will be prescribed in
12 the near future, it is also important to refer to the Hg concentrations that were below the
13 detection limit of the analytical instrument for the entire sampling period. Van Zyl et al. (2014)
14 also indicated that Hg was below the detection limit of the analytical technique for aerosol
15 samples collected at Marikana. This can be expected, since particulate Hg only forms a small
16 fraction of the total atmospheric Hg, with Hg being predominantly present in the atmosphere
17 as gaseous elemental Hg (GEM) (Venter et al., 2015; Slemr et al., 2011).

18

19 **3.4 Seasonal trends**

20 The climate and weather of South Africa is characterised by its distinctive wet and dry seasons,
21 which have an influence on concentrations of atmospheric species (Tyson and Preston-Whyte,
22 2000). Therefore, in Figure 4, the total concentrations of the trace metal species in the PM_1 (a),
23 $\text{PM}_{1-2.5}$ (b) and $\text{PM}_{2.5-10}$ (c) size fractions measured at Welgegund for each month are presented,
24 with the contributing concentrations of each of the trace metals indicated. In the $\text{PM}_{1-2.5}$ and
25 $\text{PM}_{2.5-10}$ size fractions relatively higher total trace metal concentrations are observed from
26 August to December. These periods coincided with the end of the dry season, which occurs in
27 this part of South Africa typically from mid-May to mid-October (e.g. Tyson and Preston-
28 Whyte, 2000). The end of the dry season is typically characterised by increases in wind speed
29 in August (e.g. Tyson and Preston-Whyte, 2000). Therefore, these elevated trace metal
30 concentrations determined in the $\text{PM}_{1-2.5}$ and $\text{PM}_{2.5-10}$ size fractions can partially be attributed
31 to decreased wet removal in conjunction with increases in wind generation thereof. The PM_1



1 size fractions also had relatively higher during the end of dry season period, especially during
2 September and October. However, slightly higher trace metal concentrations are also observed
3 in the PM_1 size fraction in the austral winter months from June to August. This can be ascribed
4 to the presence of more pronounced inversion layers during this time of the year (e.g. Tyson
5 and Preston-Whyte, 2000) that trap pollutants near the surface, which signifies the contribution
6 of industrial sources to PM_1 species.

7

8 **Insert Figure 4**

9

10 The monthly concentrations of each of the trace metal species determined in the PM_1 and $PM_{1-2.5}$
11 size fractions reveal the highest contributions from Fe and Ca in both these size fractions for
12 each of the months. The concentrations of Na and Cr that were the third and fourth most
13 abundant species, respectively, as well as the elevated levels of Al, B, Mg, Ni and K are also
14 reflected in the monthly distributions in the PM_1 and $PM_{1-2.5}$ size fractions. However, although
15 Fe and Ca were slightly higher in the $PM_{2.5-10}$ size fraction, a more even contribution from the
16 concentrations of Fe, Ca, Na, Cr, Al, B, Mg, Ni and K is observed (with the exception of
17 November as mentioned previously). This can be attributed to species in this larger size fraction
18 consisting predominantly of wind-blown dust (Adgate et al., 2007) with no additional industrial
19 sources of these species.

20

21 **3.5 Source apportionment**

22 As a first approach in the source apportionment investigation, Spearman correlation diagrams
23 were prepared for each size fraction. In Figure 5, Spearman correlations of the PM_1 , $PM_{1-2.5}$
24 and $PM_{2.5-10}$ size fractions are presented, i.e. Figures 5a, 5b and 5c, respectively. From Figure
25 5 relatively good correlations is observed between trace metals associated with
26 pyrometallurgical activities, i.e. Fe, Cr, Zn, Mn and V in all three size fractions. Na, Mg and
27 Ca also correlate with each other in all three size fractions, indicating the crustal (earth)
28 influence. Relatively good correlations are also observed between Ti and crustal species in the
29 $PM_{2.5-10}$ size fraction. In addition, these crustal species (Na, Mg, and Ca) also correlate with
30 species associated with pyrometallurgical activities (Fe, Cr, Zn, Mn and V). As mentioned in



1 Sections 3.1 and 3.2, although the influence of the pyrometallurgical smelters in the western
2 Bushveld Complex is evident, the large influence of wind-blown dust on trace metal
3 concentrations determined at Welgegund is also reflected with approximately 30% of most of
4 the trace metals being present in the $PM_{2.5-10}$ size fraction.

5

6 **Insert Figure 5**

7

8 In an effort to determine sources of trace metals, PCFA was applied as an exploratory tool,
9 since much larger datasets are required for definitive source apportionment with PCFA.
10 Therefore, only the most apparent groupings of metal species relating to expected sources in
11 the region were identified. PCFA of the $PM_{1-2.5}$ and $PM_{2.5-10}$ size fractions did not reveal any
12 meaningful factors. This was attributed to the large influence of wind-blown dust on trace
13 metals measured at Welgegund with all the factors obtained for the $PM_{1-2.5}$ and $PM_{2.5-10}$ size
14 fractions containing mostly crustal species loadings. In Figure 6, the factor loadings obtained
15 for the PM_1 size fraction are presented indicating four statistically significant factors with
16 eigenvalues equal to or greater than one (Pollisar et al., 1998). These four factors obtained
17 explained 88% of the variance.

18

19 **Insert Figure 6**

20

21 Factor 1 explained 59.6 % of the total system variance and was mainly loaded with trace metal
22 species that are typically associated with wind-blown dust, i.e. Ca, Fe, Na, Mg and Al (Adgate
23 et al., 2007). Therefore, this factor was identified as the crustal factor. The contribution of small
24 metal ore units from wind-blown dust is also reflected in this factor with a relatively high
25 loadings of species such as V, Mn, Zn and Cr. Mn is present in most of the ores from which
26 metals are produced in the western Bushveld Igneous Complex. The smaller contribution from
27 Mn compared to Fe in this factor is also indicative of wind-blown dust, since Mn is more
28 volatile than Fe (Kemink, 2000). Therefore, a higher contribution is expected from Mn
29 compared to Fe from pyrometallurgical sources.



1 Factor 2 and 3 explained 16.5 and 4.3 % of the variance in the data, which was identified as
2 pyrometallurgical-related factors. Factor 2 revealed higher loadings of Cr, Fe Mn, Ni and Cu,
3 while Factor 3 was predominantly loaded with Cr, Fe and V. Fe and Cr are associated with the
4 large number of ferrochromium smelters in the Bushveld Igneous Complex, while Ni related
5 to base metal smelters that refine base metals extracted from the PGM production processes.
6 In addition, Al present in Factor 2 is may be associated with fly ash formed during high
7 temperature processes, which include coal combustion. It must be noted that coal fly ash has a
8 composition, which is rather similar to that of crustal material (Mouli, et al., 2006). Mn has a
9 substantially lower vapour pressure than most of the heavy metals produced in this region.
10 Therefore, the coincidental influence of the pyrometallurgical industries is reflected by the high
11 loadings of Mn and Ni in Factor 2.

12 Factor 4 was considered to be indicative of trace metal species associated with slimes dams
13 from Au mining and recovery in the region, which is especially signified by the U and Au
14 loadings in this factor. In addition, this factor is mostly loaded with the metal species for which
15 significantly lower concentrations were measured. This factor explained 7.6 % of the total
16 system variance.

17 Pollution roses of each of the trace metal species detected were also compiled in an effort to
18 substantiate the sources identified with PCFA for the PM_{10} size fraction, as well as to verify the
19 influence of wind-blown dust that contributed to obtaining no meaningful factors for $PM_{1-2.5}$
20 and $PM_{10-2.5}$. In Figure 7, these pollution roses are presented, which indicate higher trace metal
21 concentrations associated with wind directions from the north to western sector from
22 Welgegund for all the trace metal species. As mentioned previously, the north to south-western
23 sector from Welgegund is considered to be a relatively clean region without any large pollutant
24 sources. Therefore, the most significant source of atmospheric trace metal species originating
25 from this sector can be considered to be wind-blown dust (e.g. from the Karoo and Kalahari).
26 This is also indicated by the higher atmospheric concentrations of specifically Ca, Fe, Na, Mg,
27 Al and Ti associated with the north-western sector. Furthermore, the concentrations of trace
28 metal species originating from the north can also be associated with pyrometallurgical
29 industries in the western Bushveld Igneous Complex. The influence of these activities is
30 reflected by the relatively higher concentrations of Cr, Ni, Mn, V and As associated with winds
31 originating in the north. It is also evident from these pollution roses that atmospheric Fe



1 concentrations have contributions from wind-blown dust from the north-western sector, as well
2 as from pyrometallurgical activities in the north.

3

4 **Insert Figure 7**

5

6 **4 Conclusions**

7 Of the elements analysed in the aerosol samples, atmospheric Fe had the highest concentrations
8 in all three size fractions, while Ca was the second most abundant species. Cr and Na
9 concentrations were the third and fourth most abundant species, respectively, while relatively
10 higher concentrations were also determined for Al, B, Mg, Ni and K. With the exception of Fe
11 that had higher concentrations in the PM₁ size fraction, the concentrations of the trace metal
12 species in all three size ranges were similar. With the exception of Ti, Al and Mg, 70% or more
13 of the trace metal species detected were in the two smaller size fractions, which indicated the
14 influence of industrial activities on trace metals measured at Welgegund. However, the large
15 influence of wind-blown dust on trace metal concentrations determined at Welgegund is
16 reflected by 30% and more of trace metals being present in the PM_{2.5-10} size fraction

17 A comparison of trace metal concentrations determined at Welgegund with trace metal
18 measurements conducted in the western Bushveld Igneous Complex (Kgabi, 2006; van Zyl et
19 al., 2014) indicated that Fe was also the most abundant species, while other trace metals
20 determined at Welgegund were also measured in the western Bushveld Igneous Complex.
21 However, concentrations of these trace metal species were significantly higher in the western
22 Bushveld Igneous Complex. Trace metal concentrations were also compared to levels thereof
23 in the Vaal Triangle (Kleynhans, 2008) where. Fe concentrations were similar to levels thereof
24 at Welgegund, while concentrations of species associated with pyrometallurgical smelting
25 were lower. Comparison to atmospheric trace metal species measured at international
26 background sites indicated that trace metal concentrations at Welgegund were generally lower,
27 with the exception of Al, Na and K concentrations measured at Beijing, China (Duan et al.,
28 2012) that were an order of magnitude higher. Annual average Ni ($0.079 \mu\text{g m}^{-3}$) were four
29 times higher than the European Commission Air Quality Standards limit value, which could
30 possibly be attributed to the influence of base metal refining in the western Bushveld Igneous



1 Complex. As marginally exceeded the European Commission Air Quality Standards limit
2 value, which also reflects the regional impacts of pyrometallurgical industries.

3 All three size fractions indicated elevated trace metal concentrations coinciding with the end of
4 the dry season. This could partially be attributed to decreased wet removal and increases in
5 wind generation of particulates.

6 PCFA analysis revealed four statistically significant factors in the PM₁ size fraction, i.e. crustal,
7 pyrometallurgical-related and Au slimes dams. No meaningful factors were determined for the
8 PM_{1-2.5} and PM_{2.5-10} size fractions, which were attributed to the large influence of wind-blown
9 dust on atmospheric trace metals determined at Welgegund. Pollution roses confirmed this
10 influence of wind-blown dust on trace metal concentrations, while the impact of industrial
11 activities was also substantiated.

12

13 **5 Acknowledgements**

14 The financial assistance of the National Research Foundation (NRF) towards this research is
15 hereby acknowledged. Opinions expressed and conclusions arrived at are those of the author
16 and are not necessarily to be attributed to the NRF. V. Vakkari wishes to acknowledge financial
17 support by the Academy of Finland Center of Excellence program (grant no. 272041).

18

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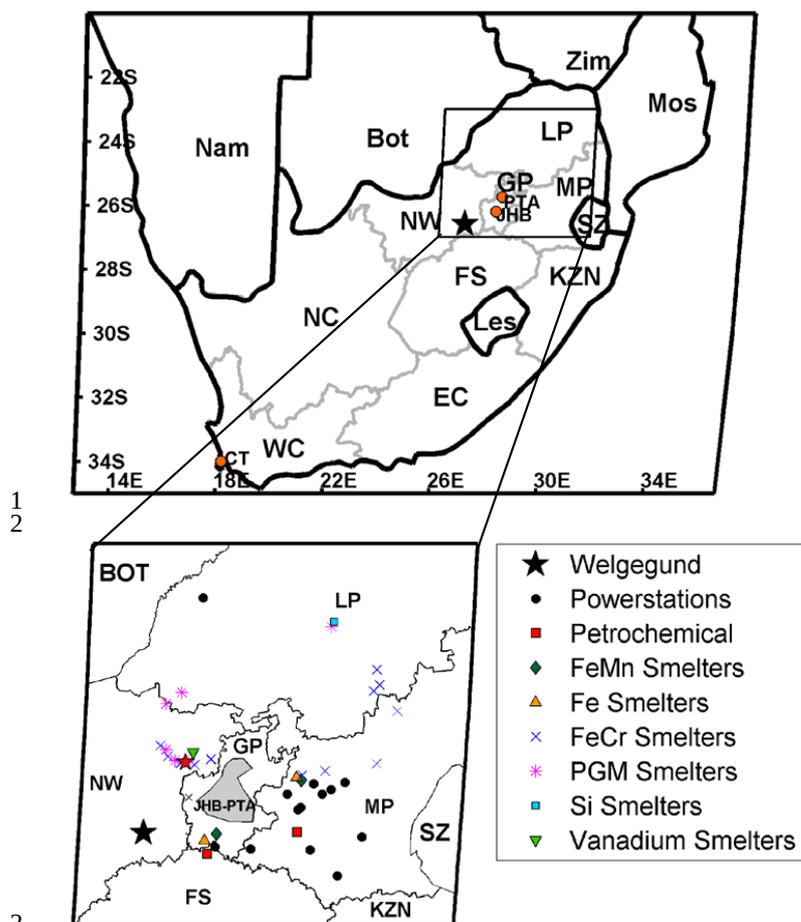
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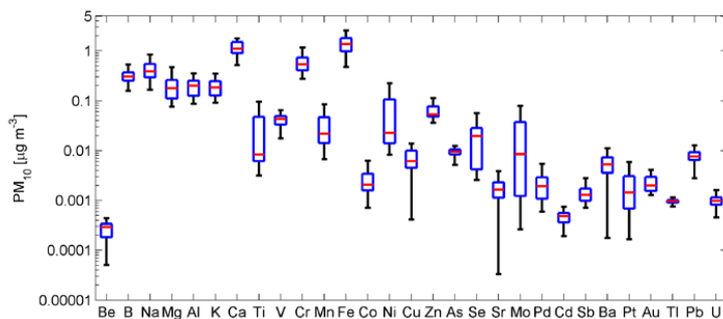
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5 Figure 1: Geographical map indicating Welgegund (black star), as well as the major point
 6 sources and the Johannesburg-Pretoria (JHB-PTA) conurbation. Neighbouring countries to
 7 South Africa (Nam = Namibia, Bot = Botswana, Zim = Zimbabwe, Mos = Mozambique, SZ
 8 = Swaziland, Les = Lesotho) as well as South African provinces (LP = Limpopo, NW =
 9 North-West, FS = Free State, KZN = Kwa-Zulu Natal, MP = Mpumalanga, NC = Northern
 10 Cape, EC = Eastern Cape and WC = Western Cape) are also indicated.
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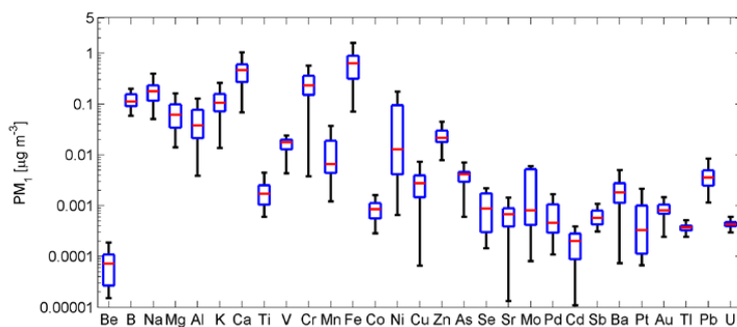


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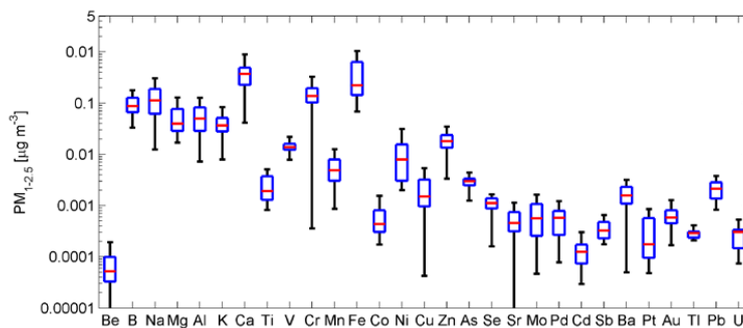
(a)

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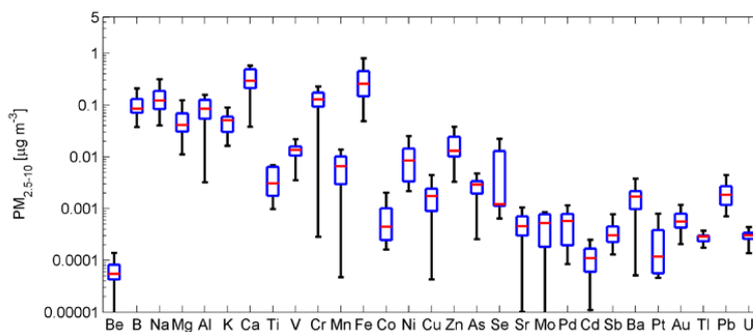


(b)

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(c)



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(d)

2 Figure 2: Box and whisker plots of trace metal concentrations in the (a) PM_{10} (sum of trace
3 metal concentrations in the three size fractions), (b) PM_1 , (c) $PM_{1-2.5}$, and (d) $PM_{2.5-10}$ size
4 fractions. The red line indicates the median concentrations, the blue rectangle of the boxplot
5 represents the 25th and 75th percentiles, while the whiskers indicate ± 2.7 times the standard
6 deviation

7



1 Table 1: Annual mean PM₁₀ trace metal concentrations measured at Welgegund, annual
 2 average standard limits, as well as annual average trace metal levels determined in other
 3 studies in South Africa, China and Europe. Concentration values are presented in $\mu\text{g m}^{-3}$

PM ₁₀ annual average	ICP detection limits ($\times 10^{-5}$)	Welgegund (This study)	Annual standard limit	South Africa			Beijing, China (Duan et al., 2012)	West coast of Portugal (Pio et al., 1996)	Spain (Querol et al., 2007)
				Marikana (Van Zyl et al., 2014)	Rustenburg (Kgabi, 2006)	Vaal Triangle (Kleynhans, 2008)			
Be	0.293	0.0002		0.020			0.100		<0.001
B	4.415	0.28		1.300					
Na	8.515	0.38		1.410		2.800	1.450		
Mg	3.504	0.23		2.040		1.000	0.637		
Al	6.960	0.17		1.280			2.180	0.200	
K	12.98	0.14		0.680		1.300	1.170		
Ca	19.88	1.1		1.080			0.996		
Ti	5.729	0.072		0.120	0.180	0.020	0.069		0.019
V	1.736	0.037	1.000 ^{(b)#}	0.040	0.160			<0.001	0.005
Cr	0.233	0.50	2.5 $\times 10^4$ ^{(a)*}	0.240	1.370	0.050	0.022	<0.001	0.001
Mn	2.064	0.026	0.15 ^(a)	0.060	4.390	0.120	0.036	0.002	0.005
Fe	15.86	1.2		2.540	9.760	1.280	1.090	0.028	
Co	0.8146	0.0035		0.140			<0.001		<0.001
Ni	4.000	0.079	0.020 ^(b)	0.330	0.770	0.040	0.020	<0.001	0.003
Cu	3.529	0.0069		0.180	0.210	0.050	0.010	0.003	0.008
Zn	14.13	0.053		0.490	0.340	0.090	0.027	0.003	0.026
As	4.730	0.0084	0.006 ^(b)	0.260			0.003	0.002	<0.001
Se	10.51	0.0074		0.580			0.001	<0.001	0.001<

4 Table 1: continued...



Sr	0.819	0.0017					0.010		0.005
Mo	0.421	0.015					0.007		0.004
Pd	7.394	0.0018		0.410					
Ag	1.030	0.0005					<0.001		
Cd	0.637	0.0004	0.005 ^{(a)(b)}	0.030			<0.001	<0.001	<0.001
Sb	0.444	0.0013					<0.001		<0.001
Ba	3.194	0.0040		0.140			0.018		<0.008
Pt	6.962	0.0016		0.350					
Au	7.340	0.0031		0.380					
Hg	9.971	0.0002	1.000 ^(a)	0.550					
Tl	4.917	0.0007		0.270					<0.001
Pb	2.592	0.0078	0.5 ^{(a)(b)(c)}	0.080	0.420	0.040	0.053	0.003	0.009
U	8.527	0.0009							

1

2

* WHO guideline for Cr(VI) concentrations associated with an excess lifetime risk of 1:1 000 000

3

24-h limit value

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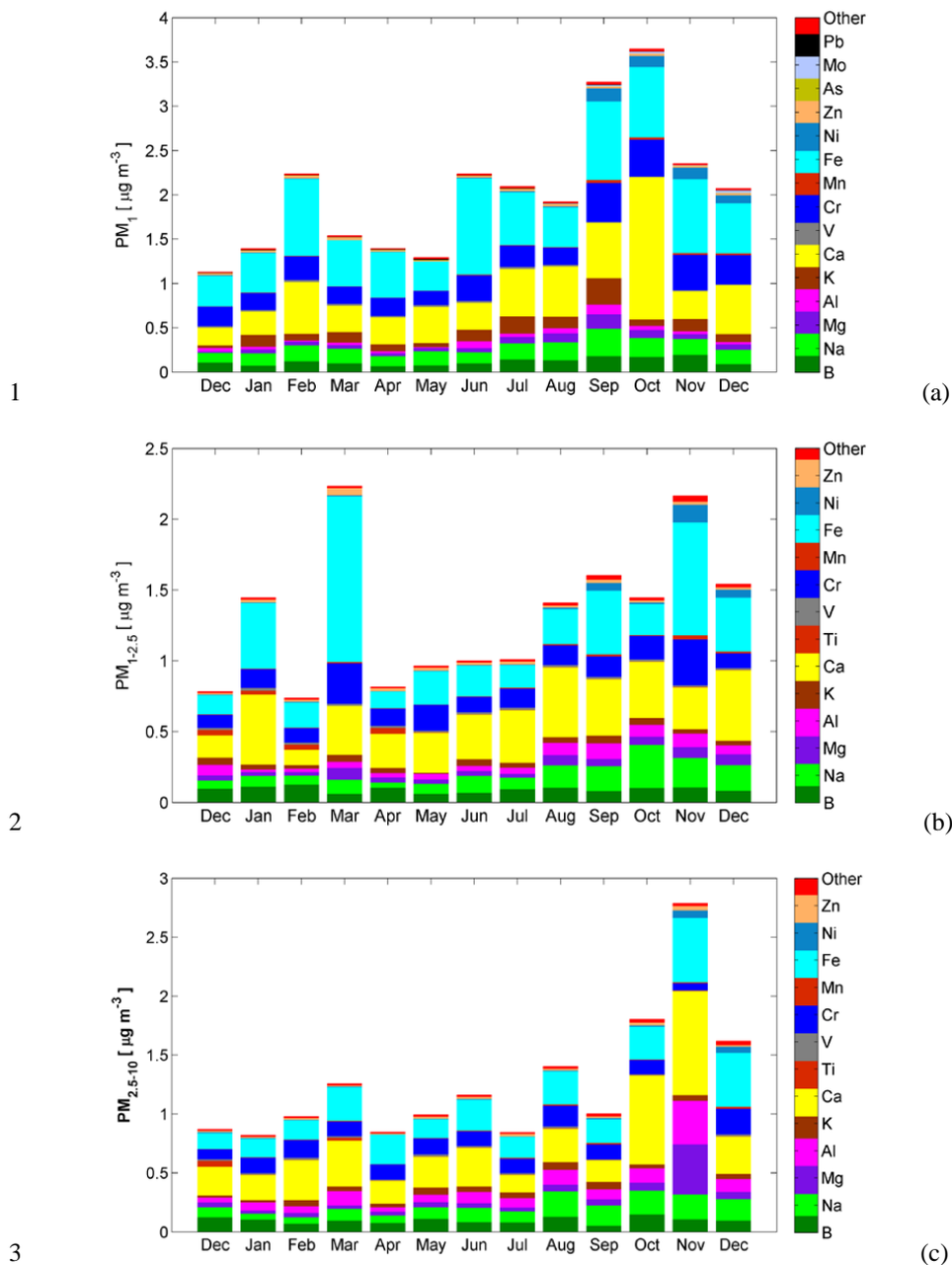
a) WHO air quality guidelines for Europe, b) European Commission Air Quality Standards, c) National Air

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Quality Act of the South African Department of Environmental Affairs

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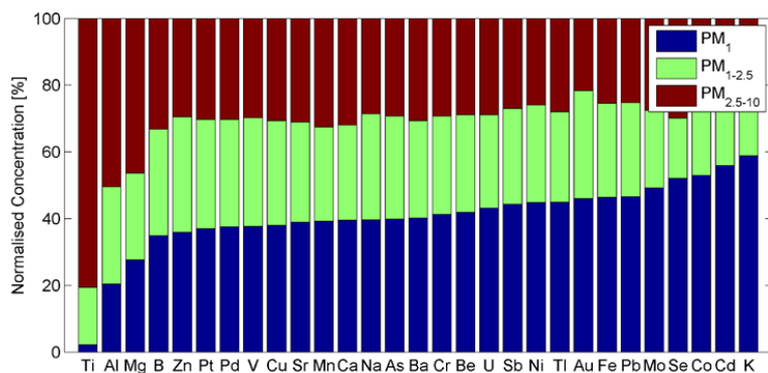


4 Figure 4: The monthly median trace metal concentrations in the PM₁ (a), PM_{1-2.5} (b) and
 5 PM_{2.5-10} (c) size fractions

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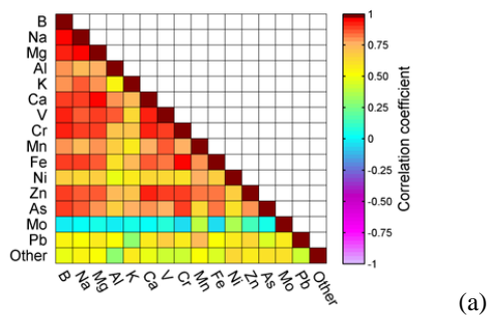
3 Figure 3: Mean size distributions of individual trace metal species detected. Species are
4 arranged by increasing concentration in the PM₁ size fraction

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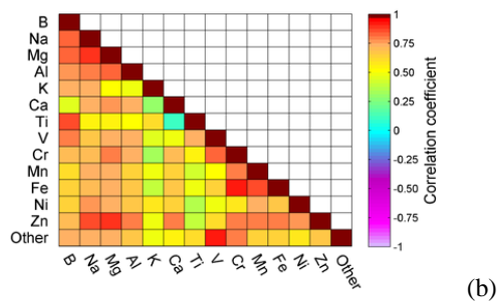
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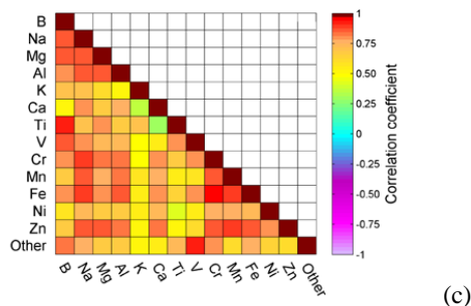
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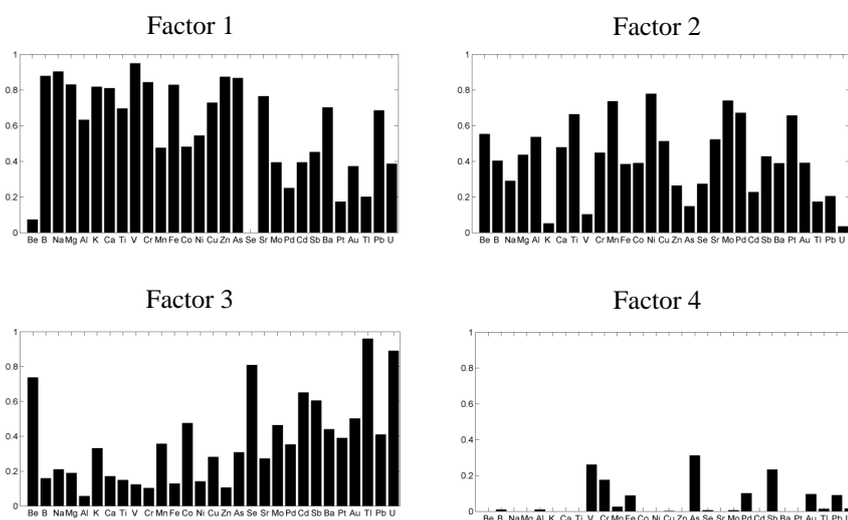


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4 Figure 5: Spearman correlations of trace metal species in the PM₁ (a), PM_{1-2.5} (b) and PM_{2.5-10}
5 (c) size fractions

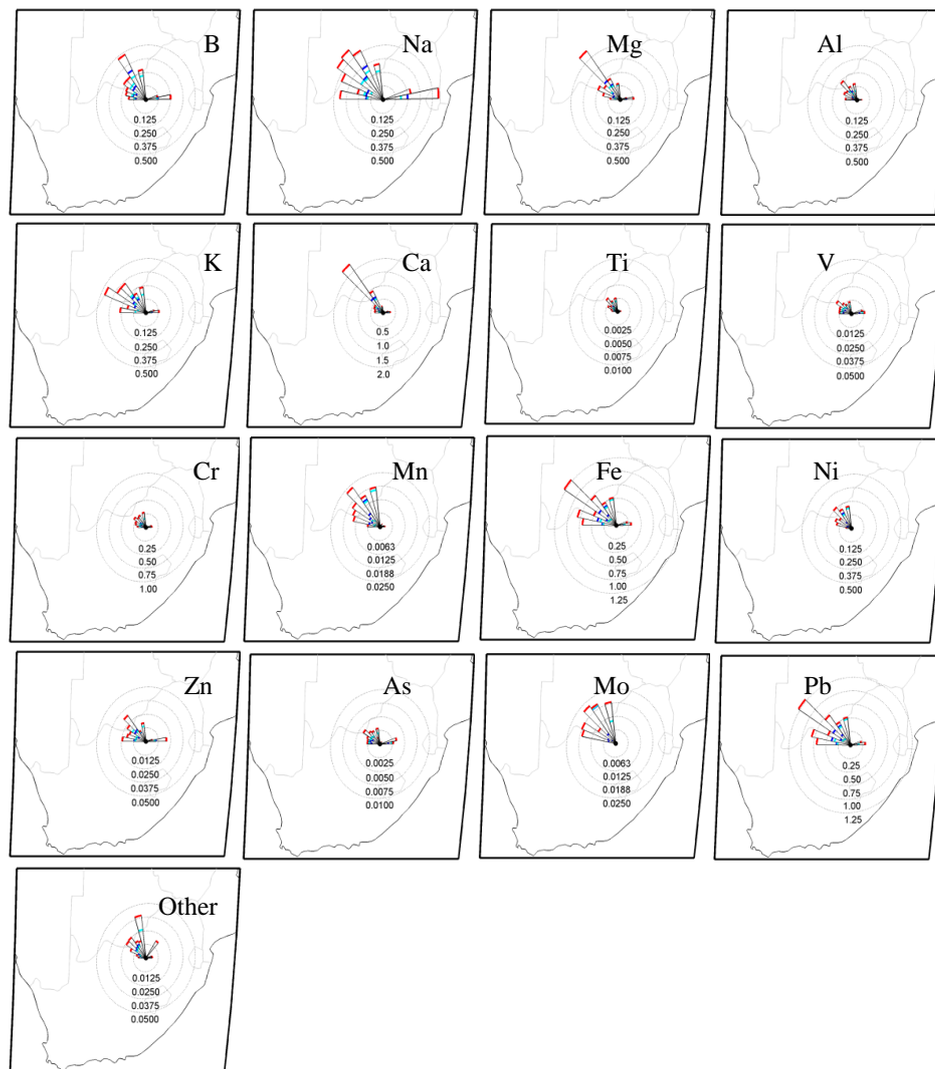
6



1

2

3 Figure 6: PCA/FA of the trace metal concentration in the PM₁ size fraction. Four dominant factors are identified.



1

2

3 Figure 7: Pollution roses of trace metal species that were 25% or more of the time detected with

4 the analytical technique