Basic characteristics of atmospheric particles, trace gases and meteorology in a relatively clean Southern African Savannah environment

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Abstract

We have analyzed one year (July 2006–July 2007) of measurement data from a relatively clean background site located in dry savannah in South Africa. The annual-median trace gas concentrations were equal to 0.7 ppb for $\text{SO}_2$, 1.4 ppb for NOx, 36 ppb for $\text{O}_3$ and 105 ppb for CO. The corresponding PM$_{1}$, PM$_{2.5}$ and PM$_{10}$ concentrations were 9.0, 10.5 and 18.8 $\mu$g m$^{-3}$, and the annual median total particle number concentration in the size range 10–840 nm was 2340 cm$^{-3}$. Gases and particles had a clear seasonal and diurnal variation, which was associated with field fires and biological activity together with local meteorology. Atmospheric new-particle formation was observed to take place in more than 90% of the analyzed days. The days with no new particle formation were cloudy or rainy days. The formation rate of 10 nm particles varied in the range of 0.1–28 cm$^{-3}$ s$^{-1}$ (median 1.9 cm$^{-3}$ s$^{-1}$) and nucleation mode particle growth rates were in the range 3–21 nm h$^{-1}$ (median 8.5 nm h$^{-1}$). Due to high formation and growth rates, observed new particle formation gives a significant contribute to the number of cloud condensation nuclei budget, having a potential to affect the regional climate forcing patterns.

1 Introduction

Air quality and interactions between land ecosystems and atmosphere in a changing climate are issues that influence most of the developing world. In Africa, for example, changing climate affect vegetation and may thereby cause emigration (Boko et al., 2007). Emigration, on the other hand, increase urban population and air pollution. Although most of the population accumulates into urban areas, air quality is affected also by the surrounding landscape. One spectacular example of this are the intensive dust episodes related to the Kosa desert in Asia, increasing occasionally PM$_{10}$ concentrations up to values of about 1000 $\mu$g m$^{-3}$ (Mori et al., 2002). For that reason, studying the air quality and anthropogenic influences in urban areas requires also knowledge
about the natural conditions, and in contrast, urban and industrial plumes may have an adverse effect on vegetation even far away from the source of emissions. Furthermore, vast areas of even sparse vegetation may contribute significantly to the global atmospheric chemistry (Guenther et al., 1996) and, via the formation of new aerosol particles, to the properties of clouds and global radiation balance (Went, 1960; Kurten et al., 2003; Kerminen et al., 2005).

Africa is one of the least studied continents in the World with respect to air quality (Laakso et al., 2006). Aerosol measurements have been made mainly in equatorial Africa (Andreae et al., 1992; Afeti et al., 1998; Gatari et al., 2005), with the main information coming from the SAFARI-92 and SAFARI 2000 campaigns conducted in the southern parts of Africa (Swap et al., 2003 and references therein). The main findings from the SAFARI-92 campaign were (1) that the anticyclonic circulation links all the environments in southern Africa, (2) that in addition to biomass burning, there are several other sources of trace gases and aerosol particles in Africa, and (3) that a large fraction of smoke and haze is transported southward from lower latitudes. In the SAFARI2000 campaign the focus was in biomass burning plumes due to the exceptional weather conditions. The published results concentrate on the optical and radiative effects of such plumes, including aerosol-cloud interactions (Ross et al., 2003), as well as pollutant emissions from vegetation and fires. In addition to results from the SAFARI campaigns, there is a great deal of publications about VOC emissions from entire savanna, and individual tree species (e.g. Guenther et al., 1996; Otter et al., 2003). Currently, there are also several different studies in the framework of African Monsoon Multidisciplinary Analyses (AMMA) from Western and Central Africa (e.g. Caminade and Terray, 2006) but, to our knowledge, no articles based on aerosol observations has been published yet.

Despite some previous observations, combined long-term measurements of trace gas concentrations, aerosol particle mass concentrations and number size distributions (especially in ultrafine size range), air ion number size distributions and meteorological variables are practically non-existent. Also the exposure of people in Africa to air pollu-
tants in informal settlements, heavily-industrialized areas and mining regions is poorly known (Pikketh et al., 2005; Kgabi, 2006).

In order to get a more detailed picture on various aerosol and related trace gas properties in southern Africa, we built a transportable measurement trailer (Petäjä et al., 2007). The trailer is currently located in a game reserve in a savannah environment away from local pollution sources. Later we will place the trailer in a populated mining region, in an industrial area and in a suburban residential background for one year in each location. Although the meteorological variability from year to year hampers the comparison of measurement results between the different environments, we aim to characterize air quality in these environments and to investigate reasons for observed differences. In this paper, we will describe the measurement site and observational procedures and present a general overview of the measurement results from the period 23 July 2006 to 23 July 2007.

2 The measurement site and characteristics

2.1 The location

The measurement site considered here is the Botsalano game reserve in North-West Province, South Africa (Fig. 1). The reserve is located in a clean background site about 50 km north of the nearest city, Mafikeng, with approximately 260 000 inhabitants. One of the large regional pollution sources in North-West Province, the Rustenburg mining region, is located approximately 150 km east of Botsalano (Pikketh et al., 2005). Rustenburg is one of the main platinum group metal mining and metallurgical extraction regions in South Africa. Such activities create significant sulphate emissions because platinum-group metals are often connected with sulphides. The capital of Botswana, Gaborone, is approximately 100 km north of Botsalano. Other potential nearby sources affecting the site are shown in Fig. 1.
The neighborhood of the measurement trailer is shown in Fig. 2. All the buildings in the park are heated by electricity. The apartments of rangers and that of managers accommodate eight and one person, respectively. The workshop of the game reserve is used mainly during the mornings between 06:00 and 08:00 local time, whereas the slaughter house is used only few times per year during the hunting season. The waste burning site is used approximately twice a week for a short period, mainly during the afternoon. Game, mainly antelopes, giraffes, rhinoceros and other herbivores, go freely throughout the reserve. On the savanna grazing ground outside the game reserve, there are sporadically cows and other livestock.

The traffic on the gravel road outside the fence is sparse, totaling a few tens of cars per day. The traffic rate between the management gate, workshop and head quarters is less than 20 cars per day. The road next to the trailer is used a few times per day in maximum.

2.2 Synoptic scale meteorology in Botsalano

The Botsalano game reserve is dry bushveld. The annual precipitation amount, of which approximately 60% comes during the summer, is 540 mm year\(^{-1}\). Typical winter temperatures vary between 4 and 20\(^\circ\)C, the summer temperatures being between 17 and 31\(^\circ\)C (South African weather service, 2007).

The seasons in the region are clearly defined and each of them lasts for a period of approximately three months. The autumn, characterized by relatively high temperatures and relative humidity, lasts from February until April. The winter (May–July) has low temperatures, relative humidities and wind speeds. The spring (August–October) is characterized mainly by low relative humidities, whereas during the summer (November–January) high relative humidities and temperatures together with frequent precipitation are typically encountered.

The large-scale meteorology in the region is characterized by a high degree of stability and anticyclonic circulation (Tyson et al., 1996). Due to the limited vertical mixing, the atmosphere is layered, containing clean and polluted horizontal cells (Hobbs,
2003). The limited vertical mixing, together with high stock heights and anticyclonic circulation, results frequently to a situation in which air masses are contaminated at least to a certain degree, either by industrial sources or by biomass burning. In some cases, air can re-circulate over the sub-continent up to 20 days (Tyson et al., 1996).

2.3 Vegetation

The vegetation of the reserve is fairly homogeneous, consisting of the species of Acacia, Rhus, Ziziphus, Vitex and Grewia as the dominant trees or shrubs typical of a savanna biome. Other tree or shrubby species include Euclea undulata, Ozoroa paniculosa, Ximenia caffra and Tarchonanthes comphoroides.

The typical grass species found in this mixed bushveld vegetation include for example Themeda triandra, Cymbopogon plurinoides, Aristida spp., Eragrostis spp., Heteropogon contortus, Schmiditia pappophoroides, Echinochloa sp., Cynodon dactylon, Melinis repens, Elionurus argenteus and Panicum spp.

The herbaceous layer include Hypoxis hemerocallidea, Alternanthera pungens, Nidorella resedifolia, Lippia scaberrina, Commelina livingstonii, Jatropha zeyheri, Hermbstaedtia adorata, Aerva leucura, as well as species of Ophrestia, Cucumis, Solanum and Hibiscus.

Based on the earlier studies, savanna trees like Acacia emits significant amount of isoprene and monoterpenes (e.g. Güenther et al., 1996). Biogenic emissions from other types of vegetation in savannah, such as grass and herbs, are not well known.

3 Measurements and methods

Air quality monitoring instruments were mounted into an air-conditioned Eurowagon 4500 U trailer (length 4.5 m, width 2.1 m, height 2.3 m, weight 2500 kg). The aim of the trailer is to be a self-sufficient and transportable monitoring station. Only prerequisites are three-phase power and periodical maintenance of the instruments. Data
is downloaded automatically every day to a server via GRPS-modem. This enables intermittent supervision of the performance of the trailer instruments remotely.

The instrumentation, technical solutions and calibrations are discussed in detail by Petäjä et al. (2007), so here we summarize them only briefly and present only prime information in Table 1.

Sub-micron aerosol number size distribution was measured with a Differential Mobility Particle Sizer (DMPS, Aalto et al., 2001) in the size range from 10 to 840 nm. The sample was drawn through a Digitel PM2.5 inlet. Prior sizing the particles were dried with a Nafion-drier and then brought to a known charging state with a Ni-63 beta-active neutralizer. The particles were classified with a Vienna-type (length 0.28 m) Differential Mobility Analyzer (Winklmayr et al., 1991) and counted with a TSI Condensation Particle Counter (CPC) model 3010. Time resolution of the system is 7.5 min.

Concentration of air ions and charged aerosol particles were measured with a Air Ion Spectrometer (AIS, Airel Ltd, Estonia, Mirme et al., 2007). The instrument is a a multi-channel, parallel-principle device, measuring simultaneously ion concentrations in 27 mobility fractions of both positive and negative ions. It has two identical cylindrical aspiration-type DMAs, one for each polarity. A radial electrical field separates naturally charged particles (cluster ions and aerosol particles) which are then deposited on different electrodes on the outer core of the DMA depending on their electrical mobility. There are 21 insulated collector electrodes on each of the DMAs. The current carried by the collected ions are amplified and measured. The concentration of ions with mobilities between 0.0013 to 3.2 cm² V⁻¹ s⁻¹ are monitored. The corresponding diameter range of singly-charged particles is approximately from 0.4 to 40 nm using modified Millikan formula (Tammet, 1995, 1998). Time resolution is 5 min.

Suspended aerosol mass concentrations were monitored with a Tapered Element Oscillating Microbalance (TEOM) model 1400a (Rupprecht and Patashnick R&P, Co. Inc.). The sample was drawn to the system via a Thermo Andersen PM₁₀ inlet through a custom made inlet switcher. This switcher alternated between three operation modes enabling consecutive measurements of mass concentrations of PM₁₀, PM₂.₅ and PM₁.
In practice a linear motor changed between a straight tube (for PM$_{10}$), a PM$_{2.5}$ cyclone or a PM$_1$ cyclone in the sampling line. This configuration enabled us to monitor all relevant mass fractions with a single TEOM-instrument with a three hour time resolution. Values for each mass corresponds 20 last min (or 4 last measurement points) for each inlet position, since instrument stabilization takes approximately 40 min after each inlet mode change.

Gaseous pollutants (SO$_2$, NO$_x$, CO and O$_3$) were monitored in one minute time resolutions with a set of gas analyzers sharing a PTFE-sampling line. Sulfur dioxide was measured with Thermo-Electron 43S, NOx with a Teledyne 200AU, CO using a Horiba APMA-360 and ozone with a Environnement s.a. 41A gas analyzer. The gas data was corrected based on on-site calibrations done in April 2006, May 2007 and October 2007.

Local meteorological parameters (temperature, relative humidity, wind speed and direction, photosynthetically available radiation (PAR) and amount of precipitation were logged in one minute time resolution. The meteorological instruments were mounted on a mast located on the roof of the trailer.

All the instruments were checked and maintained weekly, and a full service was made approximately every three months.

Power in Botsalano game reserve was taken with a 100 m long on-ground cable from the workshop.

The data analyses were carried out using MATLAB scripting language with a trailer_plot_period.m, which automatically filters out questionable data. Such values were recorded quite often after frequent electricity breaks. For the same reason, we used median values instead of means in our subsequent data-analysis. Furthermore, all the gathered data were checked visually to make sure that the bad data points were excluded.

The number size distributions acquired with the DMPS-system were the basis of this classification yielding separate datasets for new particle formation events, non-events and undefined days depending on whether new particle formation was observed or not.
in the DMPS measurement data. The classification was done according to Dal Maso et al. (2005). Also ion measurement data from the AIS were classified in a similar way, and we used the AIS data to support the event classification when the DMPS data not available. We calculated formation rate of the smallest detectable particles of 10 nm diameter (J10) and the particle growth rate (GR) in size range 10–25 nm from DMPS size distributions using the method of Dal Maso et al. (2005).

In addition to the continuous measurements described in Table 1, we measured sporadically volatile organic compounds (VOC’s) and the chemical composition of aerosol particles in PM$_{10}$, PM$_{2.5}$ and PM$_{1}$ samples. In August 2007, a new DEBITS-site was started next to the trailer (http://www.igac.noaa.gov/DEBITS.php). DEBITS measurements cover deposition of several biologically important compounds.

4 Results and discussions

4.1 Annual behaviour of meteorology, trace gases and aerosol particles

The monthly statistics of local meteorological variables are shown in Fig. 4. As can be seen, the monthly-median temperature varied by less than 15°C over a course of a year. Freezing temperatures below 0°C were encountered only during a couple of nights during the measurement period. In summertime temperatures were relatively frequently in the range 25–30°C and reached occasionally values in excess of 35°C. The relative humidity was quite low with typical monthly-median values in the range 30–60%. Highest average relative humidities were typically detected during the months with most frequent precipitation. The winds speed had a clear seasonal trend, being it’s highest during the summer months. Most of the precipitation came during the summer.

The seasonal variation of wind direction is shown in Fig. 5. During all seasons, wind was coming most often from the Eastern (NE, E, and SE) sector. As discussed earlier in this paper, this is the sector where we can assume most of the anthropogenic pollution originates from.
The monthly-median SO$_2$ concentrations were typically in the range 0.5–2 pbb with an annual median equal to 0.7 pbb (Fig. 6a). The highest SO$_2$ concentrations exceeded 10 pbb. Sinha et al. (2003) measured SO$_2$ using an aircraft in August–September during the SAFARI2000 campaign. Their SO$_2$ concentrations were higher than those obtained by us, possibly because the SAFARI flights were closer to the mining and industrial area of Rustenburg and Johannesburg. SO$_2$ concentrations over the full annual cycle have been measured in only a few regional locations in Africa, including the sites Elandsfontein (annual median equal to 7 pbb) and Cape point (0.3 pbb) in South-Africa (Carmichael et al., 2003). The annual median SO$_2$ concentration measured in equatorial and northern Africa are substantially lower (<0.1 pbb). Overall, most regional background sites in the World show annual median SO$_2$ concentrations less than 1 pbb (Carmichael et al., 2003).

The monthly-median NO$_x$ concentration varied between 1 and 2.5 pbb with an annual median equal to 1.4 pbb (Fig. 6b). The highest values were measured during the local autumn (March–April). These concentrations are comparable to those observed over mid-latitude continental areas and substantially larger than those found typically over the southern oceans (Levy et al., 1999).

The carbon monoxide concentration had a clear seasonal trend with elevated values observed during the late winter and spring (Fig. 6c). The most probable reason for the elevated CO concentration during the driest months were the regional bush fires and possibly more distant biomass burning sources. The monthly-median CO concentrations varied from about 90 ppb in April up to about 140 ppb in September, the annual median value being equal to 104 ppb. These values are similar to those measured by Sinha et al. (2003) during the SAFARI2000 campaign but roughly twice those measured annually at Cape Point, South Africa (Duncan et al., 2007). Overall, CO concentrations comparable to our measurements have been reported for continental Europe, North America and Asia (Holloway et al., 2000; Duncan et al., 2007).

The monthly-median ozone concentrations shown varied between 30 and 42 ppb and had an annual median of 36 ppb (Fig. 6d). These values are similar to those
reported by Sinha et al. (2003) and comparable to those measured in various continental surface sites at mid and low latitudes (Scheel et al., 1997; Solomon et al., 2000; Carmichael et al., 2003). Noteworthy in our data is the rather weak seasonal variation of the ozone concentration. In most surface sites, the annual cycle of monthly-median ozone concentration is above 15 ppb (e.g. Scheel et al., 1997; Helmig et al., 2007).

Figure 7 shows the measured trace gas concentrations as a function of the local wind direction. We may instantly see that the highest SO₂ concentrations were always associated with easterly winds bringing heavily-polluted air from the Rustenburg mining region approximately 150 km east of Botsalano. During westerly winds SO₂ concentrations were very low, typically around 0.5 ppb or less. The NOx concentrations were highest when the wind blew from the direction of Mafikeng. The CO concentrations were more homogeneously distributed between the different wind directions, even though a minor increase in the CO concentrations due in NE to E directions could be addressed to the nearby villages of Khunotswane and Dinokana. In contrast to SO₂ and NOx, ozone did not have any significant dependence on the wind direction.

The relatively even distributions of the CO and O₃ concentrations between the different wind sectors is probably attributed to the fact that these pollutants are mostly of secondary origin and their formation is due to a combination of several primary pollutants and meteorological conditions. O₃, for example, results from VOC and NO₂, whereas CO originates from the oxidation of biogenic hydrocarbons, soil, termites and vegetation (Finlayson-Pitts and Pitts, 2000; Holloway et al., 2000). The North-East sector was somewhat elevated for CO probably due to fires at lower latitudes.

Figure 8 shows the monthly statistics of measured PM₁, PM₂.₅ and PM₁₀ concentrations. Similar to the CO data, the highest concentrations were observed during the late winter, especially for PM₁ and PM₂.₅. The PM₁₀ concentration increased earlier during the winter (May–July 2007), being indicative of the presence of coarse dust particles from deserted areas (e.g. Kalahari) during that period of the year. Annual median concentrations of PM₁, PM₂.₅ and PM₁₀ were equal to 9.0, 10.5 and 18.8 µg m⁻³, respectively. These values are comparable to those observed in rural or continental environments.
Figure 9 shows the relation between the particle mass concentrations and wind direction. We see that the 95% percentile of PM$_{10}$ is somewhat elevated in the North-West and South-East sectors, the former being probably due to the dust from Kalahari and the latter due to activities in the direction of game park management gate and workshop (Fig. 2). Measured PM$_{2.5}$ and PM$_{1}$ concentrations were relatively equally distributed between the different wind sectors.

The total particle number concentration in the size range 10–840 nm (Fig. 10) had an annual median of 2340 cm$^{-3}$ with no clear seasonal trend. The monthly-median values were typically between 2000 and 3500 cm$^{-3}$. The highest particle number concentrations were due to daytime new-particle formation events discussed in more detail in Sect. 4.3. The values shown here compare relatively well to those by Sinha et al. (2003), the differences being probably due to the combined effects different measured size range and new-particle formation. The measurement flights by Sinha et al. (2003) were carried out in the middle of the day at the time when we observed most intensive new-particle formation. The highest particle number concentrations were observed in the same wind sector as the highest SO$_2$ concentrations (Fig. 11). The reason for this is discussed shortly in Sect. 4.3.

4.2 Diurnal cycles

In addition to seasonal variations, we also investigated diurnal variation of the measured quantities during the different seasons. Of temperature, relative humidity, wind speed and global radiation (Fig. 12), it is interesting to note the very clear diurnal behavior of the wind speed, with highest values observed typically during the mornings and relatively low values during the nights. The wind direction had also a diurnal pattern. Winds coming from East-North-East sector were frequent in the morning, turning more toward the West during the afternoon. A detailed trajectory analysis concerning the influences of air mass origin and local topography on measured aerosol and trace
gas concentrations will be carried out after we have completed the measurements in Botsalano in January 2008.

The SO$_2$ concentration had a clear diurnal cycle in all the seasons with peak values observed typically before the noon (Fig. 14a). The high values during the winter compared with other seasons can be attributed to less effective mixing, which affects the dilution of pollution originating from the Rustenburg mining area. Because of strong night-time inversions and lack of local sources, SO$_2$ concentrations decreased during the nights due to deposition. The concentrations of NOx and O$_3$ were strongly anticorrelated over the diurnal cycles as one would expect (Fig. 14b and d) (e.g. Finlayson-Pitts and Finlayson, 2000). The night-time behavior of the NOx concentration during winter and spring differed significantly from that during summer and autumn. The reason for the winter anomaly is biological activity in the ground related to the availability of the water during the humid season, which increases NO emissions (Otter et al., 1999). The CO concentrations had a weak late afternoon minimum potentially due to the rapid reaction of CO with the OH radical. The high spring values for CO are attributed to biomass burning during the dry period as mentioned earlier in Sect. 4.1.

Figures 15 and 16d depict the diurnal cycles of PM$_1$, PM$_{2.5}$ and PM$_{10}$ concentrations and ratio of PM$_1$ to PM$_{10}$ concentrations during the different seasons. During some periods a bimodal structure with a morning and late afternoon/evening maximum can be distinguished. In urban areas such a structure is attributed usually to the combined effect of traffic emissions and dilution, but this cannot be the case here due to the very low traffic amounts. The average decrease of the PM$_1$ to PM$_{10}$ concentration ratios throughout the morning until late afternoon co-insides with the period of high wind speeds and might therefore be indicative of the influence of desert dust. In summer and autumn, the high PM$_1$ to PM$_{10}$ concentration ratios during the late afternoon could be due to secondary aerosol formation associated with biogenic precursor emissions. It appears that multiple sources together with variable mixing conditions, rather than a single source or factor, is needed to explain the diurnal cycle of observed PM concentrations.
The total particle number concentration was found to peak very clearly in the afternoon, which is be related to new-particle formation taking place almost every day. During the winter, total particle number concentrations decreased much less rapidly toward the night and morning than during other seasons. Potential reasons for this might be the low winter-time wind speeds resulting in less efficient particle dry deposition and dilution, as well as the influence of smoke from the nearby villages and settlements (e.g. Khunotswane and Dinokana).

4.3 New-particle formation episodes

New particle formation was frequent in Botsalano. Figure 18 shows a typical new-particle formation day observed in Botsalano. During this day, nucleation started soon after sunrise and lasted for approximately three hours. The average formation rate of 10 nm particles was $6.7 \text{ cm}^{-3} \text{s}^{-1}$ and the average particle growth rate was $9.7 \text{ nm h}^{-1}$. About 30 000 new particles cm$^{-3}$ were formed into the size range $>10$ nm during the day.

Figure 19 shows the frequency of the observed new-particle formation events in each month. The events were analyzed separately from the AIS and DMPS data. We had several periods when only one of the devices worked properly but, due to the overlapping size range of 10–40 nm, the analyzed days are well comparable. The event probability was very high, taking place on more than 90% of the days. Days without new-particle formation (non-events), as well as undefined days, corresponded mainly to cloudy or rainy days.

Figure 20 shows the formation rate of 10 nm particles for those days when DMPS data was available (approximately 60% of the time). The observed formation rates varied between about 0.1 and $28 \text{ cm}^{-3} \text{s}^{-1}$ with no clear seasonal cycle. The corresponding particle growth rates varied between 3 and 21 nm h$^{-1}$, again with no clear seasonal cycle.
If we compare the observations with measurements carried out in other sites (see Kulmala et al., 2004, and references therein) as well as recent study from Po-valley, Italy (Hameed et al., 2007) we notice that the frequency of the new-particles formation events, as also the particle formation and growth rates, are among the highest observed in continental areas. Other important difference is the any lack of seasonal variability – clearly, we have enough nucleating and condensing vapors as well as solar radiation for nucleation to take place throughout the year.

5 Summary and conclusions

In this paper, we have introduced a new atmospheric measurement site in Southern African savannah and presented the main results from a full year of aerosol and trace gas measurements. Based on the conducted measurements, we may state that our site is in general relatively clean but influenced occasionally by anthropogenic plumes. As a result, the site provides an optimal location between polluted industrial regions to the East and clean, sparsely-populated regions to the West of the measurement site. The location allows us to investigate the influences of sources in different regions with the help of air mass trajectories.

Our observations have provided information on trace gas concentration, aerosol particle properties and meteorological variables. When comparing the measurement results with those obtained from earlier studies, we found that the concentrations of trace gases were in the level reported in literature, with seasonal and diurnal variations influenced by both biological activity and local meteorology.

Since there is no prior measurements of aerosol particle number size distributions from the regions, we observed for the first time that new particle formation takes place almost every day in such an environment. Due to their high growth rates, the fresh-formed particles reach easily sizes at which they can act as cloud condensation nuclei and affect thereby the regional radiation balance.
The first analysis of the data set raised also several questions, such as what is the effect of volatile organic compounds on new-particle formation and growth and how different sources influence trace gas and aerosol concentrations. These issues will be discussed in more detail in our future publications.

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References


### Table 1. Measurements and devices.

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Fig. 1. The measurement site. Location of the detailed map (red square) on a larger scale is indicated on the small map in the lower right hand corner. Detailed map: 1. Botsalano game reserve (measurement site: 25.32°28′ S, 25.45°16′ E, 1424 m a.s.l.), 2. Mafikeng (population 260 000), 3. Slurry (cement factory), 4. Ramathlabama (small village and chicken farm), 5. Zeerust (pop. 140 000), 6. Khunotswane (pop. few thousands), 7. Dinokana (pop. >10 000), 8. Lobatse municipal waste burning area. In addition, directions and distances from the measurement site to some regional major pollution sources are shown.
Fig. 2. Measurement site. All roads on the map are gravel roads.
Fig. 3. Surroundings of the trailer (trailer itself is located in the middle of the picture). Picture is taken approximately 300 m North-East of the trailer on 16 October 2007, during the local spring.
Fig. 4. Monthly variation of (a) temperature (b) relative humidity (c) wind speed (d) monthly accumulated precipitation. Red line in the middle is 50% percentile, boxes represent 25 and 75% percentiles and bars 5 and 95% percentiles. Numbers at the top of each month show the coverage of good quality data for each month in percent.
Fig. 5. Frequency distribution of wind direction based on 1-min data.
Fig. 6. Annual variation of gaseous pollutants (a) SO$_2$; (b) NOx; (c) CO; (d) O$_3$. Red line in the middle is 50% percentile, boxes represent 25 and 75% percentiles and bars 5 and 95% percentiles. Numbers at the top of each month show the coverage of good quality data for each month in percent.
Fig. 7. Gas concentrations as a function of wind direction, (a) SO$_2$; (b) NOx; (c) CO; (d) O$_3$. Different colors represent percentiles: green: 5%, blue 25%, black 50%, magenta 75% and red 95%.
Fig. 8. Particulate mass concentrations. Red line in the middle is 50% percentile, boxes represent 25 and 75% percentiles and bars 5 and 95% percentiles. Numbers at the top of each month show the coverage of good quality data for each month in percent.
Fig. 9. Particulate mass as a function of wind direction. Wind directions is the mean wind direction corresponding each mass measurement period.
Fig. 10. Annual variability of particle number concentration. Red line in the middle is 50% percentile, boxes represent 25 and 75% percentiles and bars 5 and 95% percentiles. Numbers at the top of each month show the coverage of good quality data for each month in percent.
Fig. 11. Total number concentration of particles as a function of wind direction.
Fig. 12. Diurnal variation of temperature, Relative humidity, Wind speed and Global radiation during different seasons. Spring: August–October; Summer: November–January; Autumn: February–April; Winter: May–July.
Fig. 13. Diurnal frequency distribution of wind direction in Botsalano game reserve, South Africa for the period 23 July 2006–23 July 2007.
Fig. 14. Diurnal variation of trace gases for each season.
Fig. 15. Diurnal variation of particulate mass during different seasons.
Fig. 16. PM$_1$/PM$_{10}$ in Botsalano game reserve as a function of time of the day for different seasons.
Fig. 17. Diurnal variation of 10–840 nm particles.
Fig. 18. New particle formation event on 2 February 2007. Figure is merged from negative AIS and DMPS data and it is shown for illustrative purposes only.
Fig. 19. Fraction of new particle formation days observed with DMPS and/or AIS. Numbers at the top of each month show percentage of data available for each month.
Fig. 20. Upper panel: formation rate of >10 nm particles in nucleation burst. Lower panel: Nucleation mode particle growth rates. Note that data have gaps during the periods when DMPS has not worked, longest gaps are in September and November 2006.