Study of Some Aerosol Features in Buenos Aires

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Abstract We present the analysis of in-situ measurements obtained in an experimental campaign in Buenos Aires. Observations of total condensation nuclei (CN), black carbon (BC), polycyclic aromatic hydrocarbon concentration in particulate phase (PPAH), vertical profiles of aerosol light backscattering and meteorological parameters, were obtained at a coastal site during a one-year period. The study provides detailed and novel information about the contributions to the aerosol load in the atmosphere of Buenos Aires due to the combined action of a variety of sources and dispersion mechanisms. Furthermore, they provide a unique dataset to evaluate modeling tools.

Keywords Urban Air Pollution, Black Carbon, Polycyclic Aromatic Hydrocarbon

1. Introduction

Buenos Aires (34°38'S, 58°28'W) is a megacity situated on the right coast of the Rio de la Plata. With a surface of nearly 3800 km², it has a population of 11 million inhabitants and ranks as the third in Latin America. The air pollution in the city is due both to the local sources and regional or remote contributions. In particular, the aerosol load in the urban atmosphere might be an issue of concern not only in the short term as regards public health or visibility, but also by the possible impacts on climate and the hydrological cycle.

Among the studies that reported measurements of particles, Arkouli et al. (2010) performed a one year campaign and the obtained PM10 and PM2.5 values were higher from July to September. The PM10 daily values were greater than the European Union limit on 36 occasions. The World Health Organization (WHO) guideline value for daily PM2.5 was exceeded 21 times. As regards the annual averages, the PM10 mean was almost 70% of the Buenos Aires limit and the PM2.5 annual mean was similar to the limit. The annual averages were above the WHO air quality guidelines. Another study reported twenty-four hour averages of PM10 and PM2.5 measured (not simultaneous) at a site near downtown Buenos Aires. The values of PM2.5 concentrations correlated well with the concentrations of carbon monoxide during winter, indicating that direct traffic emissions have an important contribution to PM2.5.

The data were less correlated for PM10, indicating that coarse fraction has an important contribution of other sources, for example re-suspended material[2].

These works represent the impact of the local sources, mainly the traffic as well as the thermo electrical plants and industries. In addition, the regional pollution is related to biomass burning, sea salt and dust. On the side of natural aerosols like sea salt, Dos Santos et al. (2012) found robust evidence that the marine aerosol from the South Atlantic Ocean reaches the city of Buenos Aires. Eventually, the western region of South America has also an important number of active volcanoes that can erupt and release to the atmosphere a variety of pollutants, which can travel far away transported by the free atmosphere winds.

The contribution of biomass burning is more noticeable from August to October. These months correspond to the main burning activities in central South America (northern Argentina, Paraguay, Bolivia and central Brazil[4,5,6]. However, as the practice is common in the agricultural sector, there are also contributions from nearby locations in Argentina[7,8]. The latter are most frequent during the austral autumn.
Light absorbing carbon, namely black carbon (BC), is mainly due to incomplete combustion in a variety of sources. Epidemiological studies have suggested that it is associated with increased cardiovascular and respiratory effects. As BC is chemically inert and dry deposition does have little effect on its removal, airborne BC is found not only close to combustion sources but also in areas remote from these sources. Similar to BC, particle-bound polycyclic aromatic hydrocarbons (PPAHs) are products of incomplete combustion. Polycyclic aromatic hydrocarbons (PAHs) are semi-volatile organic contaminants that may cause a wide range of effects due to their carcinogenic and mutagenic properties. Atmospheric PAHs are in particulate and gaseous phases. However, the carcinogenic species are predominantly associated with particles, especially those in the accumulation mode. PPAHs have also been associated with induction of oxidative stress and acute respiratory responses. Understanding the short-term temporal behavior between these two components of particulate matter may help to advance in the knowledge of their contribution to local air quality and the mitigation actions.

A research team from Universidad de Buenos Aires and Universidad Nacional Autónoma de México conducted a collaborative effort during 2011. A suite of sensors was installed at the site of Ciudad Universitaria, Buenos Aires (34°35' S, 58°22'W), and continuous measurements of some properties of aerosols were made. Figure 1 shows the location of the building and the most important nearby sources.

![Figure 1. Map of Buenos Aires city and location of the measurement site](image)

A highway that links the downtown region with the suburban zone passes at about 500 m from the building. It is heavily used during the day by cars and buses and the diurnal variation is in accordance to the office hours on workdays. There are also an important number of trucks related to the transportation of goods to/from the dock. The domestic/regional airport is at 3 km from the measurement site. Farther south are situated the thermal power plants (at about 7 km).

The aims of the experimental campaign were: (1) to provide the first comprehensive assessment of optical and physical properties of particle aerosols in the city, (2) to study the evolution of these properties in relation with the synoptic situation, atmospheric boundary layer structure and emission sources and (3) to evaluate direct and indirect modelling systems. The results will help to estimate the possible impacts on health, visibility, hydrological cycle and climate system.

### 2. Data and Methodology

Table 1 details the equipment that provided the data reported in this paper. The suite was fully operative from April to December 2011.

The measurements from the condensation nuclei counter, nephelometer and particle soot absorption photometer were recorded every second. The mass concentration of particle bound polycyclic aromatic hydrocarbons was recorded every 5 s. The weather meteorological station measured the state parameters and wind and the recording rate was every minute.

Black carbon (BC) mass concentration was derived from the absorption coefficient, $B_{\text{abs}}$, using the factory recommended specific absorption coefficient of 10 m $\mu$g$^{-1}$. 
This conversion factor was applied after the absorption coefficients had been corrected to account for differences from factory specifications in deposit area and flow rate and for the effects of light scattering. The reported concentrations from the PPAH analyzer are discussed with respect to relative changes because they were not compared with an analytical method, as recommended by the manufacturer.

Table 1. Equipment and measured variables

| Variable                        | Equipment             | Range                  |
|--------------------------------|-----------------------|------------------------|
| Condensation nuclei (CN)       | TSI 3010              | 0.05 – 3 μm            |
| Particle-Bound Polycyclic Aromatic Hydrocarbons (PPAH) | Ecochem PAS 1000 | 1 – 100 μgm$^{-3}$ |
| Absorption Coefficient (Babs)  | Radiance Research Particle Soot Absorption Photometer | 1 – 1000 Mm$^{-1}$ |
| Scattering Coefficient (Bscat) | Radiance Research Nephelometer | 1 – 1000 Mm$^{-1}$ |
| Backscattering Vertical Profile | Vaisala Ceilometer    | 20 m to 7000 m         |

3. Results and Discussion

Figure 2 depicts the monthly values obtained from a basic statistical analysis of the derived concentrations. The CN mean values ranged from 4880 cm$^{-3}$ to 13500 cm$^{-3}$ and the median from 3400 cm$^{-3}$ to 11000 cm$^{-3}$. Maximum values varied from 24000 to 64000 cm$^{-3}$, BC median values were between 0.04 μgm$^{-3}$ and 0.07 μgm$^{-3}$, the mean values ranged from 0.06 to 0.1 μgm$^{-3}$ and the maxima from 0.8 to 2 μgm$^{-3}$. PPAH mean resulted between 0.02 and 0.06 μgm$^{-3}$, the median values varied from 0.06 to 0.3 μgm$^{-3}$, and the maxima between 0.3 and 0.6 μgm$^{-3}$. The obtained monthly values are within the order of magnitude than similar locations worldwide[9].

In general, the measurements show the higher values during winter suggesting that the health risk due to exposure in winter is higher than in summer. During the cold period, the concentrations exhibited a high variability. The seasonal behaviour is similar to the variations in aerosol properties observed elsewhere[9]. These variations are likely due to a combination of changes in emissions rates and meteorology. In winter the increase in anthropogenic emissions associated with domestic heating combine with unfavourable dilution ability mainly due to shallow atmospheric boundary layer[10]. The variability is mainly related to the mid-latitude transient systems that reach the city more frequently during the cold season. Another source of variability was the arrival of the ash plume from the Puyehue-Cordon Caulle in Chile that erupted on 4 June and reached the city several times till October.

Figure 3 shows the relationships between the concentrations of BC, PPAH and CN for different wind sectors: NE (from the de La Plata River), SE (from the power plants and industries), SW (from the city and suburbs) and NW (from the northern region of the city). In the NW sector the variables are well correlated and have the largest concentrations, in the SE sector relatively lower BC and PPAH concentrations are registered; the SW sector has a bifurcation for high CN values and the NE sector corresponds to clean air masses. High concentrations of CN, PPAH, BC are associated with winds from W and NW, which points to urban common sources. Intermediate BC concentrations, low CN concentrations and very low PPAH concentrations are present with variable wind directions. Strong easterly winds that bring clean air from the La Plata River are related to intermediate BC, very low CN and PPAH concentrations.
Figure 2. Monthly CN, BC and PPAH concentrations at Buenos Aires. Line within box is the median value; square symbol is the mean value; top and bottom of box are the 25th and 75th percentiles, respectively; ends of the whiskers are the 5th and 95th percentiles, respectively.

Figure 3. BC and PPAH concentrations (ng m$^{-3}$) versus CN concentrations (cm$^{-3}$) by wind sector.
The time evolution of CN, PPAH and BC concentrations and the wind is in Figure 4. The period shown is June. High values of CN, PPAH, BC are associated with winds from W and NW (urban sources). Intermediate BC, low CN and very low PPAH values are present with variable wind directions. Strong easterly winds (clean air from the de La Plata River) are related to intermediate BC, very low CN and PPAH concentrations. The CN reached 60000 cm⁻³, BC peaked at 800 ng m⁻³ and PPAH exceeded 500 ng m⁻³.

The ceilometer measurements provided the vertical distribution of aerosols and the boundary layer diurnal variation (Figure 5). An interesting feature is that the intrusion of the volcanic ash plume from the Puyehue-Cordon Caulle was detected (days 159 and 165). There is also excellent agreement with the aerosol optical thickness derived from the AERONET site (AÉrosol RObotic NETwork) from NASA (National Atmospheric and Science Administration) at CEILAP-BA (34.5° S, 58° W).
4. Conclusions

The condensation nuclei (CN) reached 60000 cm$^{-3}$, black carbon (BC) peaked at 800 ng m$^{-3}$ and polycyclic aromatic hydrocarbon (PPAH) concentration in particulate phase exceeded 500 ng m$^{-3}$. These values are as large as those often found in other urban areas with populations much larger than in Buenos Aires. The different relationships between pollutants according to the wind sector clearly show the influence of sources. To the knowledge of the authors this is the first observational campaign that reports the concentrations of CN, BC and PPAH in Buenos Aires, for nearly a year period. The adverse effects of BC and PPAH demand this type of characterization to contribute to manage local, regional as well as global air pollution control strategies.

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