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To cite this version:
A. Marinoni, P. Cristofanelli, Paolo Laj, R. Duchi, F. Calzolari, et al.. Aerosol mass and black carbon concentrations, a two year record at NCO-P (5079 m, Southern Himalayas). Atmospheric Chemistry and Physics, European Geosciences Union, 2010, 10 (17), pp.8551-8562. 10.5194/ACP-10-8551-2010.

HAL Id: insu-00562250
https://hal-insu.archives-ouvertes.fr/insu-00562250
Submitted on 24 Feb 2012

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Aerosol mass and black carbon concentrations, a two year record at NCO-P (5079 m, Southern Himalayas)

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Received: 16 March 2010 – Published in Atmos. Chem. Phys. Discuss.: 31 March 2010
Revised: 8 July 2010 – Accepted: 10 August 2010 – Published: 10 September 2010

Abstract. Aerosol mass and the absorbing fraction are important variables, needed to constrain the role of atmospheric particles in the Earth radiation budget, both directly and indirectly through CCN activation. In particular, their monitoring in remote areas and mountain sites is essential for determining source regions, elucidating the mechanisms of long range transport of anthropogenic pollutants, and validating regional and global models. Since March 2006, aerosol mass and black carbon concentration have been monitored at the Nepal Climate Observatory-Pyramid, a permanent high-altitude research station located in the Khumbu valley at 5079 m a.s.l. below Mt. Everest. The first two-year averages of PM¹ and PM¹−10 mass were 1.94 µg m⁻³ and 1.88 µg m⁻³, with standard deviations of 3.90 µg m⁻³ and 4.45 µg m⁻³, respectively, while the black carbon concentration average is 160.5 ng m⁻³, with a standard deviation of 296.1 ng m⁻³. Both aerosol mass and black carbon show well defined annual cycles, with a maximum during the pre-monsoon season and a minimum during the monsoon. They also display a typical diurnal cycle during all the seasons, with the lowest particle concentration recorded during the night, and a considerable increase during the afternoon, revealing the major role played by thermal winds in influencing the behaviour of atmospheric compounds over the high Himalayas. The aerosol concentration is subject to high variability: in fact, as well as frequent “background conditions” (55% of the time) when BC concentrations are mainly below 100 ng m⁻³, concentrations up to 5 µg m⁻³ are reached during some episodes (a few days every year) in the pre-monsoon seasons. The variability of PM and BC is the result of both short-term changes due to thermal wind development in the valley, and long-range transport/synoptic circulation. At NCO-P, higher concentrations of PM¹ and BC are mostly associated with regional circulation and westerly air masses from the Middle East, while the strongest contributions of mineral dust arrive from the Middle East and regional circulation, with a special contribution from North Africa and South-West Arabian Peninsula in post-monsoon and winter season.

1 Introduction

Atmospheric aerosol particles are the focus of increasing attention, due to their role in many atmospheric processes, in particular those involved in the Earth’s radiative balance. The uncertainty linked to aerosol forcing is a major obstacle to the accurate prediction of future anthropogenic-induced climate changes (Forster et al., 2007). Aerosol radiative forcing depends either directly or indirectly on several intensive properties of particles, including mass concentration, presence of absorbing material in their composition and size distribution.

While the variability of aerosol mass (PM) concentration is well documented in urban and peri-urban areas due to air quality regulations, uneven global and regional coverage of PM observations remains a limitation for assessing global aerosol atmospheric loadings and source identification. Access to PM information is limited in many parts of the world, especially in remote regions and the free troposphere (Laj et al., 2009). In many studies on climate and air quality, knowledge of PM variability needs to be integrated with information on the main constituents of fine atmospheric aerosol.
Aerosol particles may be transported over long distances and reach remote areas, especially in dry climates, because of the slow rate of their removal from the atmosphere in the absence of precipitation. Long range transport of aerosol to clean background areas may have a substantial effect on regional climate change, but also on public health issues (Chung and Seinfeld, 2005; Kanakidou et al., 2005; Sun and Ariya, 2006).

Ubiquitous throughout the atmosphere, BC is present in low levels in remote areas, like the South Pole (Andreae et al., 1984; Hansen et al., 1984, 1988) and Manua Loa (Bodhaine, 1995) but higher concentrations (1–30 µg m$^{-3}$) are observed near sources, in urban areas and biomass burning regions (Cachier et al., 1986; Ruellan et al., 1999). Jacobson (2001) pointed out that the magnitude of direct radiative forcing due to BC may exceed that due to methane, thereby making it one of the main species contributing to global warming. INDOEX measurements have reported 14% of BC in fine aerosol dry mass in the marine boundary layer (Lelieveld et al., 2001; Ramanathan et al., 2001). Apportionment of BC is a major concern when modelling aerosol radiative properties. Such estimates are extremely rare for Indian locations, where surprisingly high pollution levels have also been observed over the surrounding ocean. The Indian subcontinent is a region where atmospheric pollution is rapidly on the rise, causing extensive air quality degradation, with local, regional and global implications, including an impact on the oxidizing power of the atmosphere. (Lelieveld 2001). In fact, recently surface and satellite observations have pointed to the persistence of so-called “Brown Clouds”, i.e. wide polluted tropospheric layers characterised by anthropogenic aerosol optical depth (AOD) greater than 0.3 and absorbing AOD greater than 0.03 (Ramanathan et al., 2007)

A knowledge of aerosol mass and black carbon concentrations in remote regions is therefore necessary to determine emission sources, quantify long range transport and validate both regional and global models. High-elevation sites are well-suited for documenting components of the free troposphere, in order to characterize its typical background conditions and investigate the influence of human activity on its composition. For these reasons, aerosol mass and black carbon concentrations have been monitored at the Nepal Climate Observatory-Pyramid (NCO-P) since March 2006 in the framework of ABC (Atmospheric Brown Clouds) UNEP and SHARE (Station at High Altitude for Environmental Research) Ev-K2-CNR projects. Located in the Khumbu valley at 5079 m a.s.l. below Mt. Everest, NCO-P is one of the very few monitoring stations in this area. The two-year continuous record (March 2006–February 2008) of fine and coarse aerosol mass concentrations and black carbon content completes the very rare records published for this area (Shrestha et al., 2000; Carrico et al., 2003).

The present paper describes observations of PMs and BC at the site and discusses factors influencing aerosol diurnal and seasonal variability. In particular, the role of valley breezes and synoptic circulation on the aerosol transport up to such high altitude is also investigated. Because NCO-P is located within the area of the Himalayan glaciers, a knowledge of concentrations and sources of BC, the most efficient light-absorber in atmosphere and snow, is a key step towards a better assessment of anthropogenic pressures on snow-pack and glacier dynamics.

With the presentation of an unique data set referring to aerosol mass and black carbon observations carried out in a crucial area that plays a key role in climate change, the main objectives of this paper are to present and discuss the principal factors influencing the aerosol variability, in order to provide important information for the characterization of tropospheric background conditions and precious input to improve the performance of climate and atmospheric chemistry models at different scales.

2 Measurement site and experimental set up

A complete description of the NCO-P station (Nepal, 27.95 N, 86.82 E, 5079 m a.s.l.) is provided in Bonasoni et al. (2008) and Bonasoni et al. (2010). At NCO-P, the seasonal variation of atmospheric conditions is influenced both by the local mountain wind system (with a strong diurnal valley wind and a weaker mountain night-time breeze), and by the large-scale Asian monsoon circulation. In particular, besides determining the seasonal variations of meteorological parameters, the annual variations of the main synoptic circulation can also modulate the diurnal cycles characterizing the local mountain weather regime (Bonasoni et al., 2010).

The instrumentation includes a Multi-Angle Absorption Photometer (MAAP 5012, Thermo Electron Corporation) and an optical particle counter (GRIMM#190), both devices sampling air from a TSP (Total Suspended Particle) head. On-site technical support is provided year-round by the Ev-K2-CNR Committee, while a field campaign on site is organized once a year for extra maintenance work, instrument calibration, and possible technical upgrades. The energy required for NCO-P operations is provided by 98 photovoltaic cells and 120 battery storage, other than those supplied by Pyramid panels and cells. A diesel generator (located 100 m below NCO-P) is used to supply electricity in the event of solar-energy shortage, although this occurs rarely (less than 6% of total period), and tests have been performed to insure that no contamination of NCO-P measurements occurs due to diesel use.

PM is measured using the optical particle counter (OPC), which measures the size distributions for particle with diameters greater than 0.25 up to 32 µm by means of 90° laser light scattering. The optical diameters determined by the OPC (classified in 31 bins) are converted into geometric diameters
assuming spherical particles. According to the manufacturer, the accuracy of the Grimm 190 OPC in particle counting is ±2% over the entire measurement size range. The OPC particle sizing is based on a calibration with latex spheres (1 µm Duke Scientific Corp.) performed by the manufacturer prior to starting measurements at NCO-P. Additional calibration was performed in February 2008 using latex spheres (400 nm Duke Scientific Corp.), showing a slight, but acceptable, diameter underestimation of less than 5%. To calculate the apparent particulate mass within the different size ranges (PM$_{2.5}$, PM$_{10}$ and PM$_{10}$), the particle density was chosen by the manufacturer during the first calibration using Micro Dolomit DR80 (Basserman & Co). Particles smaller than 0.3 µm were assumed not to significantly contribute to the particulate mass: the error associated to PM determination was lower than 2% in each size fraction. The OPC was placed in the NCO-P shelter with temperature almost constant (between 20 and 25 °C), at least 10 °C higher with respect to the external one, thanks to heating produced by the instrument energy consumption. The particles were thus measured at a relative humidity slightly lower than ambient RH. However above 70% RH (measured in the sample head), a dehumidification system becomes active, removing humidity over a Nafton ® membrane-tube from the sample pipe to avoid condensation during measurement. The detection limit was determined as 3σ of 12 h of measurement with an absolute filter, corresponding to 5.8 pL$^{-1}$, i.e. equivalent to 0.06 µg m$^{-3}$.

The measurement of the aerosol absorption coefficient was obtained by MAAP: it measures the transmission and the back scattering of a light beam (Petzold and Scholinner, 2004) incident on a fibre filter where aerosol particles are deposited by sampling flow. As determined during an intercomparison experiment (March 2007) at the Leibniz Institute for Tropospheric Research (Leipzig, Germany), the wavelength of the light source employed in the MAAP is 635 nm (T. Müller, personal communication, 2008). The reduction of light transmission, multiple reflection intensities, air sample volume are continuously integrated over the sample run period to provide real-time data output of the particle light absorption coefficient $σ_{abs}$, and thus black carbon concentration.

In the visible spectral range, black carbon is the most efficient light-absorbing aerosol species (Horvath, 1993). The relationship between the aerosol absorption coefficient $σ_{abs}$ (m$^{-1}$) and the corresponding black carbon mass concentration BC (ng m$^{-3}$) is established by a mass absorption efficiency $σ_{me,B}$ (m$^2$ g$^{-1}$) via the relationship:

$$BC = \frac{σ_{abs}}{σ_{me,B}}$$

where $σ_{me,B}$ and $σ_{abs}$ are wavelength dependent. $σ_{me,B}$ depends on the type of aerosol, the ageing and the size of the BC particles (Lioussse et al., 1993). A thermo-optical method (OC/EC analyzer, Sunset laboratory Inc.) was used to determine the mass absorption efficiency ($σ_{me,B}$), assuming that elemental carbon (EC) is equivalent to BC (details of analyses in Decesari et al., 2009). Here, the value of $σ_{me,B}$ adopted was 6.5 m$^2$ g$^{-1}$, as recommended by Petzold et al. (2002) and set by default on MAAPs. The very good correlation between EC and equivalent BC ($R^2=0.94$), obtained by totally independent methods confirms the fact that, despite the presence of high levels of dust (Duchi et al., 2010), MAAP absorption measurements are primarily influenced by BC. This justifies the use, in the remaining part of the paper, of the term “equivalent BC” for MAAP measurements.

In order to check the MAAP noise and detection limit, a 12 h measurement with an absolute filter was performed during the calibration campaign in February 2008, together with the calibration of flow and temperature-pressure sensors. The detection limit (3σ of blank measurements) was calculated as 11 ng m$^{-3}$, with an integration time of a 30 min basis.

Both instruments worked continuously without major failures, and interruptions were extremely rare and of limited duration. Measurements (available in real-time at the web address http://evk2.isac.cnr.it/realtime.html) were performed with a time resolution of 1 min and data were averaged to a common time base of 30 min. All measurements presented in this study refer to STP conditions (1 atm and 0 °C) and Nepal Standard Time (NST, UTC+5.45).

3 Results and discussion

3.1 Variability of PMs and BC concentrations

Mass concentrations of accumulation (0.25–1 µm) and coarse (1–10 µm) particulate matter calculated by OPC, as well as equivalent BC mass concentration, are reported in Fig. 1, while their seasonal statistics are given in Table 1. The accumulation fraction mass will be referred hereinafter as PM$_1$, since the sum of chemical species analysed on PM$_1$ filters (Decesari et al., 2009) accounts on average for 91% of the accumulation fraction mass calculated by OPC: the fraction of particles smaller than 0.25 µm (not detected by OPC) are thus assumed as negligible.

The two-year average of PM$_1$ and PM$_{10}$ mass were 1.94 µg m$^{-3}$ and 1.88 µg m$^{-3}$, with a standard deviation of 3.90 µg m$^{-3}$ and 4.45 µg m$^{-3}$, respectively. Thus, the coarse fraction accounts for half of the aerosol mass measured by OPC, with a greater variability with respect to the accumulation fraction. The black carbon concentration averaged over two years of measurements was 160.5 ng m$^{-3}$, with a standard deviation of 296.1 ng m$^{-3}$, accounting for 10.8% of PM$_1$ on average.

3.1.1 Seasonal variations of PMs and BC

Both fine and coarse fractions of aerosol mass exhibit a well defined diurnal cycle, as well as significant annual variations. In fact, as shown in Figs. 1 and 2, large mass concentrations...
occur during the pre-monsoon season, while the minima appear during the monsoon season (prolonged to the post monsoon for the coarse fraction). PM$_1$ shows a more enhanced annual variation than PM$_{1-10}$: the pre-monsoon average is larger by a factor of 5.6 with respect to the monsoon mean value, while the coarse fraction mass (characterised by a larger year-to-year variation) shows a pre-monsoon average value larger than the monsoon average by a factor of 3.0. The year 2007 was characterized by a more peaked variation, especially for coarse fraction. The ratio between the seasonal maximum and minimum of aerosol mass in 2007 was 11.4 for the coarse fraction and 5.6 for the fine fraction, while in 2006 they were 1.7 and 5.7, respectively. The origin of the inter-annual difference may be associated with the inter-annual variations in meteorological and dynamic conditions already observed in this area, such as the Tropospheric Biennial Oscillation, leading to warmer and wetter even years than odd years (Bertolani et al., 2000; Pillai and Mohankumar, 2007).

The annual behaviour of BC concentration is very similar to the PM$_1$ annual cycle, with a maximum during the pre-monsoon season, a minimum during the monsoon and a similar cycle amplitude (pre-monsoon average value is larger than the monsoon value by a factor of 5.0 over the 2 years, 4.5 in 2006 and 6.6 in 2007). A weak inter-annual variation characterised the BC behaviour at NCO-P. In fact, except for the post-monsoon and winter season, the year 2006 appears slightly more “polluted” than 2007 (see Table 1 and Fig. 2). The very high BC standard deviation reflects the high variability of BC concentrations: frequent “background conditions” (55% of the time), with concentrations mainly below 100 ng m$^{-3}$ are observed, while during some episodes (a few days every year) in pre-monsoon seasons, BC concentrations up to 5 µg m$^{-3}$ (30 min averaged values) are reached (Fig. 1).

The aerosol mass and BC seasonality observed at NCO-P is similar to the variations observed at other Indian sites, with lower values during the monsoon season, mostly due to wet scavenging, and larger ones in the winter and pre-monsoon periods (Chandra Mouli et al., 2006; Ganguly et al., 2006). In the Indogangetic plain, at Gadanki station (India, 375 m a.s.l., 13.5$^\circ$ N, 79.2$^\circ$ E), Krishnan and Kunhikrishnan (2004) observed that the atmospheric boundary layer height was maximum during the pre-monsoon period, and minimum during the winter months. It can therefore be supposed that the high pre-monsoon values observed at NCO-P are influenced by the larger vertical extension of the regional PBL (indicative of the ABC vertical extension), which together with the valley breezes, can favour the transport of regional pollution up to the 5000 m a.s.l. of NCO-P.

### Variations of BC/PM$_1$

The contribution of BC to fine aerosol mass at NCO-P was found to be 10.8%, with a very large variability (standard

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**Table 1.** Seasonal average and standard deviation for BC, PM$_1$, PM$_{1-10}$ (30-min averaged values). In brackets, the annual average values for 2006 (Y06) and 2007 (Y07) are also reported. $N$ represents the number of averaged values.

| Seasons   | PM$_1$ (µg m$^{-3}$) | PM$_{1-10}$ (µg m$^{-3}$) | BC (ng m$^{-3}$) | BC/PM$_1$ (%) |
|-----------|----------------------|---------------------------|------------------|---------------|
| Pre-monsoon | 3.9±5.5 (Y06: 4.4; Y07: 3.5) | 3.4±6.7 (Y06: 2.5; Y07: 3.6) | 320±469 (Y06: 376; Y07: 279) | 10.5±28.0 |
| N = 10045 | 0.6±1.1 (Y06: 0.8; Y07: 0.3) | 1.1±2.7 (Y06: 1.4; Y07: 0.6) | 56±75 (Y06: 85; Y07:41) | 40.0±141 |
| Monsoon   | 1.5±1.4 (Y06: 1.5; Y07: 1.5) | 0.5±1.4 (Y06: 0.5; Y07: 0.6) | 137±126 (Y06: 132; Y07: 144) | 10.0±5.0 |
| N = 4189 | 1.4±3.6 (Y06: 1.0; Y07: 2.3) | 1.6±4.3 (Y06: 1.4; Y07: 2.2) | 125±147 (Y06: 124; Y07: 126) | 12.4±7.4 |
| Winter    | 1.6±4.3 | 28.0 | 7.4 |
| N = 4777 | 12.4±7.4 | 12.8; Y07: 11.7 |

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**Fig. 1.** Time series of 30-min averaged BC (black), PM$_1$ (cyan) and PM$_{1-10}$ (brown) at NCO-P.
deviation: 9.5%). As shown in Fig. 3, the BC/PM$_{1}$ ratio shows a maximum contribution of carbonaceous absorbing material to the fine particulate matter during the monsoon season (about 18% in July-August), while the minimum values characterize the pre-monsoon season (less than 9% in April-May). This is plausibly due to a preferential scavenging by monsoon precipitation of the inorganic fraction of aerosol with respect to the less hygroscopic black carbon (Sellegri et al., 2003). Moreover, this may also be addressed to the influence of local BC emissions due to incomplete combustion processes from domestic heating and cooking activities. In fact, during the monsoon season highly efficient scavenging processes clean air masses originating over Indian plains, while local/regional sources, even when weak, could influence the background composition of Himalayas atmosphere. Even though BC contributes only a few percent to the total aerosol mass, it produces significant radiative effects. The apportionment of BC is therefore very important in assessing the aerosol radiative properties, as confirmed by Marcq et al. (2010) who report lower single scattering albedo values during the monsoon season.

The BC/PM$_{1}$ ratio observed at NCO-P is higher than values observed at lower altitude Himalayan sites in India. In fact, Saha et al. (2005) found a winter average value of ~5%, at Manora Peak (29.4° N; 79.5° E; 1951 m a.s.l., India) during winter 2004, while Hyvärinen et al. (2009) pointed out a ~4% average BC/PM$_{2.5}$ ratio at Mukteshwar (29.4°N; 79.6° E, 2180 m a.s.l., India). This indicates the existence of significant differences in aerosol composition along Himalayas ridge. In fact, for NCO-P, the BC/PM$_{1}$ is similar to those observed within brown clouds over Indian Ocean and populated urban areas. Moreover the BC/PM$_{1}$ ratio seems to be higher with cleaner atmospheric conditions, suggesting a preferential removing of non-BC aerosol. INDOEX measurements (January to March 1999) reported surprisingly high pollution levels (comparable to urban air pollution in North America and Europe) over the entire northern Indian Ocean, with 14% of BC in fine aerosol dry mass in the marine boundary layer (Lelieveld et al., 2001; Ramanathan et al., 2001b). Novakov et al. (2000) found a BC content similar to the NCO-P one in the Indian Ocean haze, as high as 17% of the total fine particle mass. This very high fraction correlated very well with total carbon and sulphate, suggesting the predominantly primary nature of aerosol carbon and the significant influence of fossil fuel combustion sources. As reported by Babu and Moorthy (2002), BC/PM$_{1}$ values ranging from ~3–4% (during monsoon) to 9% (during winter) were observed at Trivandrum (~1 million inhabitants, South India). Except for the remote site of Mukteshwar (2180 m, North India), where on average only 4% of PM$_{2.5}$ mass consisted of BC, the ensemble of Asian observations shows values generally higher than the European continental background: Cozic et al. (2006) found a BC contribution to PM$_{1}$ equal to 2.5% in summer and 4.7% in winter at the free troposphere station of Jungfraujoch (Switzerland, 3850 m a.s.l.), while Putaud et al. (2004) found a contribution in the range of 5–10% to PM$_{2.5}$ at a European natural background sites. As indicated
by Babu et al. (2004), the BC/PM$_1$ ratio is a key parameters to define the atmospheric forcing efficiency of aerosols. Thus, the high BC/PM$_1$ ratio observed at the NCO-P suggests that BC plays a major role in driving atmosphere energy budget over High Himalayas, as also indicated by Marcq et al. (2010).

### 3.1.2 Diurnal variations

The NCO-P average seasonal diurnal variations of PM$_1$, PM$_{1-10}$ and black carbon, together with specific humidity (SH), often used as a tracer of PBL-influenced air masses for mountain sites (e.g. Henne et al., 2008), are shown in Fig. 4. Alongside day-to-day variability, PM$_1$ and black carbon also show a typical diurnal cycle in all the seasons with the lowest particle concentrations at night and a considerable increase during the afternoon. As shown by Venzac et al. (2008) and Bonasoni et al. (2010), such behaviours suggest an important influence of thermal winds in determining the variation of atmospheric compounds over the high Himalayas. Similar phenomena have been observed at many mountain stations in the European Alps (Baltensperger et al., 1997), Italian Apennines (Marinoni et al., 2008), Massif Central in France (Venzac et al., 2009), and Hawaii (Bodhaine, 1995). The maximum of pollutant (PM$_1$ and BC) concentrations is reached between 16:00 and 18:00 NST during the pre-monsoon season and about one or two hours earlier in the other seasons, while the minimum appears between 06:00 and 08:00 NST in all the seasons, except for the winter one, when it is reached around 04:00 NST and persists longer.

It is interesting to note that the SH diurnal cycle shows the same behaviour as BC and PM$_1$ variations, except in the monsoon season, when the BC and PM peaks and minima are delayed by about two hours with respect to SH. Here, it is hypothesized that the early peak of SH in the monsoon season is due to an additional water vapour source from evapotranspiration along the valley in central hours of the day, while outside the monsoon season, humidity reaches NCO-P simultaneously with pollutants from low levels of the Khumbu Valley and Pahar region.

Pollutants reaching the high altitude site, pumped up by thermal winds or transported by synoptic circulation, can be injected into the free troposphere, where they attain a longer life time, and/or accumulate in high residual layers. As shown in Table 2, the amplitude of the BC and PM$_1$ diurnal cycle is very close to that of SH during the monsoon season, and higher in other seasons. Additionally, the nighttime BC contribution to the fine aerosol mass is, on average, between 2% and 10% higher than the day-time contribution. The higher BC fraction in night-time aerosol sampled at the station may reflect an ageing of the aerosol population; in fact the polymerisation of organic material in aerosol particles can lead to increasingly absorbing properties (Andreae and Gelenscer, 2006).

Following the approach of Baltensperger et al. (1997), the fraction of days showing a strong influence of valley-induced thermal convection was calculated, on the basis of the black carbon concentration. Applying this criterion a “strong” diurnal variation was found in 38% of all days (55% in the pre-monsoon and only 7% in the monsoon season).

The most “polluted” days (characterized by a daily BC average larger than the 90th percentile) were characterized by a larger amplitude of pollutant diurnal variation: for the total average the 15:00–21:00 (NST) period shows a mean BC value 3.5 higher than the 03:00–09:00 period, while considering only days with a daily BC average over the 90th percentile (345 ng m$^{-3}$), the 15:00–21:00 period became 8.2 times higher than the 03:00–09:00. Nevertheless, among the “polluted” days, more than 30% showed limited amplitude of the diurnal cycle and therefore not classified as days with “strong” diurnal variation. The recirculation and transport of pollutants possibly stored in reservoir layers can partially explain the limited amplitude of the diurnal cycle. However, these findings seem to confirm that two main mechanisms are involved in the transport of pollutants to NCO-P: thermal winds, leading to an enhanced diurnal cycle, and synoptic circulation, which constrains and limits the thermal wind influence.

### Table 2. Diurnal cycle amplitudes of the typical day in different seasons for specific humidity (SH), black carbon (BC), PM$_1$, and PM$_{1-10}$. Amplitudes are expressed as the ratio between the highest and the lowest 30-min averaged values.

| Parameter | Pre-monsoon | Monsoon | Post-monsoon | Winter |
|-----------|-------------|---------|--------------|--------|
| SH        | 1.7         | 1.2     | 1.8          | 2.5    |
| BC        | 4.8         | 1.4     | 3.5          | 3.8    |
| PPM$_1$   | 2.1         | 1.9     | 4.2          | 7.2    |
| PM$_{1-10}$ | 1.2     | 0.7     | 1.9          | 2.0    |
Typical events with marked diurnal variations, with the highest concentration of PM$_1$ (around 60 µg m$^{-3}$) and BC (attaining 5 µg m$^{-3}$), were recorded in March–April of both 2006 and 2007: the strongest of such events are recognizable also in 30-min averaged time series shown in Fig. 1 (i.e., 30–31 March, 1–3 and 15–17 April 2006, 30–31 March, 1 and 3–6 April 2007). Conversely, when the rise in pollution concentration shows a very limited diurnal cycle, it is most likely that the air mass is transported by synoptic circulation, which superimposes and masks locally originated thermal winds. A typical prolonged long range transport is recorded both in 2006 and 2007 in the monsoon onset period (12–22 June 2006 and 22 May–7 June 2007).

The aerosol coarse mass has a more irregular diurnal variation than PM$_1$ and BC, with daytime concentrations (10:00–18:00 NST) higher than night-time ones in all seasons except for the monsoon. The diurnal peaks are not linked with the lifting of air masses by valley winds, but seem to be linked with wind strength, with a maximum at midday (4–5 h before pollution maximum), suggesting a limited contribution from the local re-suspension of coarse particles. Furthermore, the coarse mass variations are weaker than the seasonal and diurnal cycles of the fine fraction (see Table 2), while PM$_{1-10}$ and the respective diurnal cycle amplitude are higher in the post monsoon and winter seasons, when the soil is drier (Fig. 4).

### 3.1.3 Comparison with other sites

At NCO-P, the two-year average of PM$_1$ and PM$_{1-10}$ mass were 1.94 µg m$^{-3}$ and 1.88 µg m$^{-3}$, with a standard deviation of 3.90 µg m$^{-3}$ and 4.45 µg m$^{-3}$, respectively, while the BC mean concentration was 160.5 ng m$^{-3}$, with a standard deviation of 296.1 ng m$^{-3}$, accounting for 10.8% of PM$_1$ on average.

As expected, the aerosol mass values observed at NCO-P were much lower (at least one order of magnitude) than the corresponding ones measured everywhere else in the Indian Subcontinent: the annual average mass of PM$_{10}$ was 33 µg m$^{-3}$ (annual average, with similar seasonality as NCO-P) in Tirupati, South of India (Chandra Mouli et al., 2006), 74 µg m$^{-3}$ in Agra (Khemani et al., 1982), 66 µg m$^{-3}$ in Ahmedabad (Ganguly et al., 2006) both cities in the Indo-Gangetic Plain. This reflects the influence of the brown clouds, as well as the larger transcontinental pollution over South and East Asia (Ramanathan, 2001).

The PM$_{10}$ mean value recorded at NCO-P is comparable with TSP concentrations measured at Nam Co (4730 m a.s.l., Tibetan Plateau) for the period August 2006 to July 2007 showing an annual average of 7 µg m$^{-3}$ and falling in the range 0.5–36 µg m$^{-3}$ (Cong et al., 2009), while in the remote and rural area of Tengchong (1960 m a.s.l., Tibetan Plateau), PM$_{10}$ was measured as 34 µg m$^{-3}$ and PM$_{2.5}$ as 28 µg m$^{-3}$ (Chan et al., 2006). The aerosol mass values observed at NCO-P are also lower than the European natural background defined by Putaud et al. (2004), within the range 5–10 µg m$^{-3}$.

The BC concentration levels at NCO-P appear very similar to values at Nam Co. For the period when BC measurements are available at both sites (July 2006–January 2007), BC was on average 82±70 ng m$^{-3}$ at Nam Co (Ming et al., 2010) and 91±113 ng m$^{-3}$ at NCO-P, suggesting that both stations are representatives of Himalayan background conditions. The monthly mean BC concentration for December 2004, measured at the mountain site of Nainital (Manora Peak, 1950 m a.s.l., located in the Shivalik ranges of the central Himalayas) was 1.36±0.99 µg m$^{-3}$ (Pant et al. 2006), thus significantly higher than NCO-P levels for the same season, probably because of the lower altitude and greater proximity to the strong sources of the Indo-Gangetic plain. In fact, on the Indian plains, BC daily levels range from few µg m$^{-3}$ up to 27 µg m$^{-3}$ measured in Delhi by Beegum et al. (2009) and Babu et al. (2002). Even on the southern Indian Ocean, BC concentrations are mostly higher than those observed at NCO-P, especially during winter, testifying to the widely extending atmospheric brown cloud over this area. At NCO-P, monsoon averaged BC values fall mainly in the range of the European continental background, measured between 8 and 104 ng m$^{-3}$ (monthly median) at Jungfraujoch (Alps, 3450 m a.s.l.) by Nyeki et al. (1998) and at Monte Cimone (winter average below 100 ng m$^{-3}$) by Marini et al. (2008). The NCO-P monsoon values are approximately two orders of magnitude higher than Antarctic background concentrations (~0.65 ng m$^{-3}$) at the South Pole, one order of magnitude higher with respect to a remote marine boundary layer on the island of Mauna Loa (~5.8 ng m$^{-3}$), and on the same order than the baseline atmospheric station of Barrow, on the northernmost tip of Alaska (~41 ng m$^{-3}$) (Bodhaine, 1995).

### 3.2 Source origin: thermal wind regime versus long range circulation

As shown by Bonasoni et al. (2010), NCO-P atmospheric compound concentrations are the result of local/regional scale dynamics along the Khumbu valley and long-range transport and synoptic circulation. During the pre-monsoon season in particular, the Khumbu valley is an efficient and persistent channel for transporting large amounts of particles to NCO-P during the day-time up-valley breeze, as also suggested by the analysis of diurnal variations investigated in Sect. 3.1.2. Figure 5 shows the seasonal average BC, PM$_1$ and PM$_{1-10}$ split into clusters with respect to wind direction (anabatic or catabatic) and typically clean (night-time) or polluted (afternoon) periods. It is clear that valley breezes play an important role in pollution transport from the low level of Khumbu valley and Himalayan foothills during all seasons (blue bars), while the lowest pollutant concentrations are recorded during mountain breeze conditions (brown and yellow bars). Nevertheless, the effect of seasonality strongly

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In Fig. 5, it is interesting to note that pre-monsoon lowest concentrations of PM$_1$ and BC are associated with mountain winds. This is particularly evident during the afternoon, when the day-time valley winds from South-West are suppressed by synoptic scale forcing, transporting air-masses from North, often characterised by high levels of mineral dust aerosol (Duchi et al., 2010). In fact, as shown in Fig. 5 the largest PM$_{1-10}$ concentrations (about one order of magnitude larger than the overall average) observed in winter and pre-monsoon season, are associated with mountain wind during the afternoon, suggesting a not-negligible role of mineral dust mobilisation and recirculation from the Tibetan Plateau and Taklimakan desert. However, a significant contribution of PM$_{1-10}$ is carried to NCO-P by afternoon valley winds, especially in the pre-monsoon seasons, when the Indo-Gangetic Plains are strongly influenced by the transport of dust outbreaks originating in the north-western desert in India (Thar desert, see Gautam et al., 2009), and perhaps even from far off regions (North Africa, see Decesari et al., 2009).

Besides local/regional circulation, synoptic air-mass circulation can also influence the atmospheric composition at NCO-P. To evaluate the possible influence of synoptic-scale air-mass transport on atmospheric composition observed at NCO-P, BC, PM$_1$ and PM$_{1-10}$ data were analysed as a function of the air-mass cluster classification provided by Bonasoni et al. (2010). In particular, depending on geographical origin and paths of the analysed back-trajectories, seven clusters were identified and classified into three main classes: SW south-westerly (SW-AP: Arabian Peninsula, SW-AS: Arabian Sea, SW-BG: Bengal Gulf); W westerly (W-NA: North Africa, W-EU: Europe, W-ME: Middle East) and REG regional. Following the same methodology presented by Bonasoni et al. (2010), only night time data (00:00–06:00) were considered, in order to avoid interaction with air masses carried up with valley winds, and data over the 95th percentile (6 h average night-time values) were neglected (5.0 µg m$^{-3}$ for PM$_{1-10}$, 3.4 µg m$^{-3}$ for PM$_1$ and 26.7 µg m$^{-3}$ for BC) to avoid the influence of outliers.

Figure 6 shows the averaged concentrations of aerosol mass (PM$_1$ and PM$_{1-10}$) and BC split into the different back-trajectory clusters for each season. Even if seasonality plays an important role in defining atmospheric composition at NCO-P, as is evident looking at scales of the graphs, significant differences are observed within each season between aerosol concentration in air masses of different origin.

During the pre-monsoon season, the largest content of aerosol is associated with westerly air masses from W-ME: such air masses are characterised by a large mineral dust load, together with a high pollution level, and they represents 19.3% of air masses in this season. To a lesser extent, REG air masses also bring a significant amount of dust and pollution to NCO-P. This cluster represents the second main class of air masses for the pre-monsoon season (23.9%) and differs from others because of its lower back-trajectory.
altitude, tracing the influence of regional sources (Bonasoni et al., 2010) and explaining the association of dust and pollution. The lower amount of mineral dust is connected to other W and SW air masses, while pollution shows a minimum with W-EU air masses. A possible role of the Arabian peninsula and Middle-East/Persian Gulf (Lelieveld et al., 2009) as sources of pollution for South Himalayas is suggested by the high fraction of BC/PM$_1$ from SW-AP and SW-AS.

During the monsoon season, low BC average values characterised all the analysed air-mass circulation. However, REG and W-ME air masses are associated with the highest amount of PM$_1$ and BC, the highest accumulation mode concentrations were also observed for aerosol size distribution in these air masses (Sellegri et al., 2010). It is important to specify that REG air masses accounted for 73.8% of air masses in this season, while W-ME accounted for only 3.9%. The SW-BG air masses (20% of monsoon back-trajectories) brought to NCO-P the cleanest air over the whole year, with only 0.14 µg m$^{-3}$ of PM$_1$ and 38 ng m$^{-3}$ of BC. In fact, SW air masses are often associated with low pressure over the Gulf of Bengal, causing intense precipitation on the whole Indian Subcontinent up to the Himalayas foothills (Barros and Long, 2003). Only one event from SW-AS (not shown in Fig. 6) brought very high levels of aerosol in concentrations mostly above the seasonal averages (1.7 µg m$^{-3}$ PM$_{1-10}$, 5.5 µg m$^{-3}$ PM$_1$ and 316 ng m$^{-3}$ BC).

The most frequent post-monsoon back-trajectories (W-ME: 37.1%, REG: 26.3% and W-NA: 18.6%) were also the most polluted of the season, with high average values of PM$_1$ and BC but a limited contribution of mineral dust as deduced by the low PM$_{1-10}$ values. Conversely, SW-AS (7.8%) air-masses were characterised by the highest content of mineral dust.

During the winter season, the highest PM$_1$ and BC averaged value was related with W-EU air-masses (frequency: 19.8%), for which also relatively high PM$_{1-10}$ amount were observed at NCO-P. Relatively high pollution levels were traced back to SW-AS air-masses (frequency: 8%), while high mineral dust contents were also detected with the most frequent W-NA cluster (frequency: 47.2%).

The analysis of cluster concentrations and their frequency shows that the long distance air mass transport significantly impacts aerosol concentration and composition: W-ME and REG air-masses were significantly affected by polluted aerosols from pre-monsoon to post-monsoon seasons, while during winter the highest BC and PM$_1$ average values were related to W-EU air-masses. Concerning mineral dust, the strongest contributions are mainly from W-ME and REG during pre-monsoon, with possible contributions from SW-AS during the post-monsoon and far westerly regions during winter.

4 Summary and conclusions

The present paper addresses the diurnal and seasonal variability of aerosol mass and black carbon concentration observed over a two-year period at Nepal Climate Observatory-Pyramid (5079 m a.s.l.).

The annual mean aerosol mass concentration is 3.8±8.4 µg m$^{-3}$. The coarse fraction accounts for half of aerosol mass measured by OPC (1.88±4.45 µg m$^{-3}$), with a larger variability with respect to the accumulation fraction. Black carbon average concentration over two years of measurements is 160.5 ng m$^{-3}$, with a standard deviation of 296.1 ng m$^{-3}$.

All the parameters investigated in the paper show a well defined seasonal variation, as well as significant inter-annual variations. The highest concentrations occur during the
pre-monsoon season, while the minima appear during the monsoon season (prolonged to the post-monsoon period for the coarse mass). The monsoon minimum is attributed to wet scavenging due to frequent cloud and precipitation in this season. The pre-monsoon maximum is linked to an increase of anthropogenic emissions in Indian Subcontinent coupled with the dry meteorological conditions, and to the larger vertical extent of the regional PBL on the Indo-Gangetic plains, associated to the efficient “chimney” effect of the Himalayan valleys.

In fact, PMs and BC concentrations show a strong diurnal variation: the daytime maxima of PM$_1$ and BC are linked to up-valley thermal winds with a maximum in the afternoon, while PM$_{1-10}$ is linked to wind strength, with a maximum peaked at noon, suggesting a limited contribution from local re-suspension of coarse particles. Night-time mass and BC concentrations are relatively low throughout the year, providing information on free tropospheric background levels at high altitudes in the Himalayas. However, although lower than the daytime values, the night-time levels follow the same seasonality as the diurnal values, suggesting that pollutants reaching the high altitude, pumped by thermal winds or transported by synoptic circulation, can be injected into the free troposphere, where they attain a longer lifetime, and/or accumulate in high residual layers. This is particularly evident in the pre-monsoon season, when the night-time/early morning concentrations remain still high with respect to the rest of the year.

The mean values of PM reported in this study are definitely lower than the Asian and Europe background. However, as shown by the high mean BC values, at NCO-P the non-monsoon seasons are more affected by regional/continental emissions than European high altitude sites, indicating the influence of the brown cloud up to 5000 m a.s.l.

The contribution of BC to the total aerosol mass at NCO-P is, on average, higher than 10% (standard deviation: 9.5%). The BC/PM$_1$ ratio shows a maximum contribution of carbonaceous absorbing material to the fine particulate matter during the monsoon season (about 18% in July-August), while the minimum values characterize the pre-monsoon season (less than 9% in April-May). Such values are much higher than those measured at other mountain sites, and more similar to the values reported for urban polluted sites. The monsoon maximum can be attributed to a preferential scavenging by monsoon precipitation of the inorganic aerosol fraction with respect to the less hygroscopic black carbon. However, during this season, a possible role of local/regional sources cannot be completely ruled out.

Besides valley breeze regime, synoptic air-mass circulation can also influence the atmospheric composition at NCO-P. The analysis of BC, PM$_1$ and PM$_{1-10}$ night-time observations as a function of the air-mass cluster classification provided by Bonasoni et al. (2010), confirms that long distance transport of air masses significantly impacts aerosol concentration and composition. Significant differences are observed within each season between aerosol concentration in air masses of different origin/path: polluted aerosols are mainly related to W-ME, REG and W-EU air masses, while the strongest contributions of mineral dust are mainly from W-ME and REG during pre-monsoon and monsoon, with a special contribution from W-AS during post-monsoon and far westerly regions during winter.

Acknowledgements. This work was carried out in the framework of the UNEP-ABC ( Atmospheric Brown Clouds) and Ev-K2-CNR-SHARE ( Stations at High Altitude for Research on the Environment) projects. The contribution of CNRS through the PICS bilateral program between CNR and CNRS and through the LEFE-INSU program is gratefully acknowledged. The authors also thank Tenzing C. Sherpa, Kaji Bista, Laxman Adhikary, Pema Sherpa, Lhakpa T Sherpa, Lakpa T Sherpa, Chhimi T Sherpa and Hari Shrestha for their support at the Nepal Climate Observatory-Pyramid, and Michael Sprenger (ETHZ) for providing LAGRANTO back-trajectories.

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