Carbonaceous and inorganic species in atmospheric aerosols during wintertime over urban and high-altitude sites in North India

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1 Synchronous sampling of bulk-aerosols, carried out during wintertime from the two strategically located sites in North India, reveals that total suspended particulates (TSP) over an urban site (Hisar: 29.2°N 75.7°E; 219 m asl) ranged from 67 to 396 μg m⁻³; in contrast, TSP at Manora Peak (a high-altitude station: 29.4°N 79.5°E; 1950 m asl) was relatively low (range: 13.7 to 42.7 μg m⁻³). At Hisar, on average, water-soluble ionic species (WSIS, range: 14.1 to 78.3 μg m⁻³) contribute nearly one-fourth by weight to TSP, with dominant contribution from SO₄²⁻, NO₃⁻ and NH₄⁺. The time series analysis over a span of 30 days shows somewhat uniform distribution of organic carbon/elemental carbon (OC/EC) ratio centering around 8.5 ± 2.2 at this urban site; and the water-soluble organic components (WSOC range: 6.7 to 42.0 μg m⁻³) account for 11.5 % to the TSP concentration. Both WSOC and OC exhibit significant positive correlation with water-soluble K⁺ (r = 0.88 and 0.79 respectively), suggesting their dominant contribution from biomass burning. At Manora Peak, the chemical composition of ambient aerosols show characteristically lower abundances of WSIS (range: 2.0 to 9.9 μg m⁻³) and WSOC (range: 1.4 to 6.0 μg m⁻³); together they account for one-third of the TSP. The characteristic low abundances of OC (range: 2.8 to 6.9 μg C m⁻³) and EC (range: 0.34 to 1.4 μg C m⁻³) at this high-altitude site and their significant correlation with K⁺ and SO₄²⁻ suggest contribution from long-range transport of anthropogenic species. This study represents a first comprehensive data set for documenting the chemical characteristics of ambient aerosols and source apportionment of EC, OC, WSIS and mineral dust over urban and high-altitude sites in north India, an important data set required for the south Asian region. If the observed OC/EC ratios for urban and high-altitude sites in north India, almost 50% of its global emissions are estimated to arise from the latter source [Cooke and Wilson, 1996]. The primary sources of OC are either anthropogenic or biogenic in nature; whereas its secondary source results from the condensation of semi-volatile organic vapors to particle surfaces and via atmospheric photo-oxidation reactions of precursor gases such as terpene [Odum et al., 1996; Griffin et al., 1999; Kanakidou et al., 2005]. The role of black carbon and organic aerosols is getting increasingly recognized in the atmospheric chemistry and radiative forcing models [IPCC, 2007]. The organic aerosols have potential to scatter sun’s radiation and to reduce the hygroscopicity of inorganic species, thus causing variations in the light scattering property of aerosols with change in relative humidity [Sjogren et al., 2007]. In particular, the water-soluble organic compounds (WSOC) can alter the surface properties of aerosols from hydrophobic to hydrophilic and thus influence their optical properties as well as atmospheric cycles of several important trace gases (such as ozone); and eventually the number

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