



# Road traffic related influences upon PM<sub>2.5</sub> load in Beijing, China

Klaus Schäfer<sup>1\*</sup>, Rong-rong Shen<sup>1</sup>, Jürgen Schnelle-Kreis<sup>2</sup>, Yuesi Wang<sup>3</sup>, Longyi Shao<sup>4</sup>, Stefan Emeis<sup>1</sup>

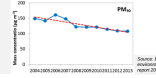
- <sup>1</sup> Institute of Meteorology and Climate Research, Atmospheric Environmental Research (IMK-IFU) at the Karlsruhe Institute of Technology (KIT), 82467 Garmisch-Partenkirchen, Germany
- <sup>2</sup>Joint Mass Spectrometry Centre, Cooperation group "Comprehensive Molecular Analytics", Helmholtz Zentrum München, 85764 Neuherberg, Germany
- <sup>3</sup> Institute of Atmospheric Physics (IAP), Chinese Academy of Sciences (CAS), 100029 Beijing, P.R. China
- <sup>4</sup> School of Geoscience and Surveying Engineering, China University of Mining and Technology (CUMTB), 100083 Beijing, P. R. China

#### **OBJECTIVES**

Emission reduction measures were performed to improve air quality during the Olympic Summer Games in 2008: mainly  $PM_{10}$  reduced, haze became a problem - why?

Work program:

- Chemical composition of PM
- Characteristics of chemical compounds
- Special case studies during haze events for source apportionment

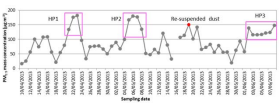


### METHODOLOGY

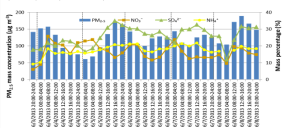
Particulate sampling on quartz fibre filters with 2 High-Volume Samplers DHA80 (Digitel) from 10 April till 8 June 2013. Sampling time 24 hours (00:00 - 24:00); 4 hours during some haze episodes. Backward trajectories calculated with HYSPLIT4. Meteorological data: ZBAA (http://weather.uwyo.edu/upperair/sounding.html) and IAP

Particle composition: Inorganic elements by ICP-MS and PM<sub>2.5</sub> mass concentrations by filter weighing (sampler A). Inorganic water-soluble ions, EC / OC and organic speciation by IC with continuous flow analyzer, thermal/optical carbon analyzer and GC-MS (sampler B)

Source apportionment: Positive matrix factorization (PMF3.0, EPA)

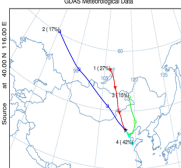


Temporal variation of  $\rm PM_{2.5}$  mass concentration from 10 April till 8 June 2013 in Beijing with indication of three haze episodes (HP1, HP2 and HP3) and a dust episode (re-suspended dust)



Secondary inorganic pollutant mass concentrations ( $NO_3$ ',  $SO_4$ <sup>2</sup> and  $NH_4$ \*) of  $PM_{2.5}$  during high-temporal sampling (4 hours) from 3 till 8 June 2013

Cluster means - Standard 60 backward trajectories GDAS Meteorological Data



Cluster means of backward trajectories from 10 April till 8 June 2013

## RESULTS

#### PM<sub>2.5</sub>/PM<sub>10</sub> ratios:

higher values during haze days (0.68) than during clear days (0.38); fine particles and high transformation efficiency into secondary aerosol dominant factors for haze pollution

#### PM<sub>2.5</sub> characteristics:

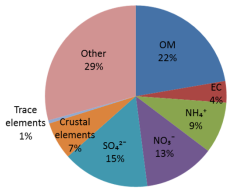
EC, OM (organic matter), Cl-,  $SO_4^{2-}$ ,  $NO_3^-$ ,  $NH_4^+$ , K+, Cu, Ni, Zn, As, Cd, Tl and Pb mass concentrations **increased during haze days** in comparison to clear days (see mass balance right).  $SO_4^{2-}$ ,  $NO_3^-$ ,  $NH_4^+$  mass concentrations 6 times higher.

4 h sampling: average  $PM_{2.5}$  mass concentration highest 16:00 - 20:00.  $NO_3$ -mass percentage high 0:00 - 8:00 and low 12:00 - 20:00.  $SO_4$ <sup>2</sup> mass percentage varied with  $PM_{2.5}$  mass concentration, highest values 12:00 - 20:00 (see left).

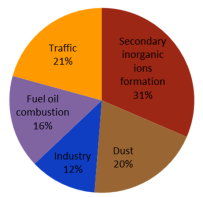
Fe, Ca and Ba (crustal elements) represent dust particles, **Zn**, **As and Pb indicate haze** particles.

High amount of secondary inorganics: major chemical species of PM<sub>2.5</sub> during haze originated from anthropogenic sources.

 $NO_3$ -/  $NO_3$ - mass ratio during haze days (0.84) higher than during 2001 - 2003 (0.80) (Wang et al., 2005) and 1999-2000 (0.58) (Yao et al., 2002): vehicle exhaust emissions are a rational for haze days in Beijing.



Mass balance of PM<sub>2.5</sub> compounds from 10 April till 8 June 2013



Result of source apportionment by PMF from 10 April till 8 June 2013

#### PM<sub>2.5</sub> emission sources:

The result of PMF analyses are (see right):

- Main sources: secondary inorganic ion formation, traffic, dust, fuel oil combustion and industry.
- Dust and traffic emissions main sources during clear days, secondary inorganic ion formation dominant sources during haze days.

Based on PAH diagnostic ratios: coal and liquid fossil fuel combustion dominant sources of PAHs.

Some ratios of compounds as the Hopane and Homohopane index: high amount of **vehicle exhaust emissions** and coal combustion emissions.

Enrichment factor analyses: aerosols originated from

- geogenic sources are most likely re-suspended road dust and dust storm and
- anthropogenic sources are vehicle exhaust, chemical industry, coal combustion and fertilizer application.

Backward trajectories from the South, calculated for 72 hours in 500 m altitude at the target point, were always found to be short and with high PM mass concentration (see above): southerly flows were the main source of haze particles in Beijing - industrial companies and big cities are in a distance of some 100 km.

#### Acknowledgement

This work was partly funded by Chinese Scholarship Council (CSC) and KIT Centre of Climate and Environment

#### CONCLUSIONS

Haze becomes more frequent because an increasing number of fine and anthropogenic particles is emitted.

Control of emissions of precursor gases of SO<sub>4</sub><sup>2-</sup>, NO<sub>3</sub><sup>-</sup> and NH<sub>4</sub><sup>+</sup> in the local and regional scale and thus of **road traffic emissions** is necessary for reducing haze.

Installation of **cleaning equipment in** industrial and **mobile exhaust vents**, **improvement of road cleaning standards** and reduction of construction dust also becomes important for improving air quality.

There is a need to do further research on formation mechanisms and sources of haze episodes during different seasons in order to improve the urban air quality.

#### REFERENCES

Sun, Y.L., G.S. Zhuang, A. Tang, Y. Wang and Z.S. An (2006), Chemical characteristics of PM2.5 and PM10 in haze-fog episodes in Beijing, *Environmental Science & Technology*, 40, 3148-3155. Wang, Y., G.S. Zhuang, A. Tang, H. Yuan, Y. Sun, S. Chen and A. Zheng (2005), The ion chemistry and the source of PM2.5 aerosol in Beijing. *Atmos. Environ.*, 39, 3771-3784.

Wang, Y., G.S. Zhuang, Y. Sun and Z.S. An (2006), The variation of characteristics and formation mechanisms of aerosols in dust, haze, and clear days in Beijing, Atmos. Environ, 40, 6579–6591. Yao, X., C.K. Chan, M. Fang, S. Cadle, T. Chan, P. Mullawa, K. He and B. Ye (2002). The water-soluble ionic composition of  $PM_{2.5}$  in Shanghai and Beijing, China. Atmos. Environ, 36, 4223–4234.

Zhang, J.K., Y. Sun, Z.R. Liu, D.S. Ji, B. Hu, Q. Liu, and Y. S. Wang (2014), Characterization of submicron aerosols during a month of serious pollution in Beijing, 2013, *Atmos. Chem. Phys.*, 14, 2887–2903.

<sup>\*</sup> Contact: Klaus Schäfer – klaus.schaefer@kit.edu