1	Evaluation of tropospheric SO ₂ retrieved from MAX-DOAS	
2	measurements in Xianghe, China	
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12	Abstract	
13	Ground-based Multi-Axis Differential Optical Absorption Spectroscopy (MAX-DOAS)	
14	measurements of sulfur dioxide (SO ₂) have been performed at the Xianghe station (39.8°N,	
15	117.0°E) located at ~50 km southeast of Beijing from March 2010 to February 2013.	
16	Tropospheric SO ₂ vertical profiles and corresponding vertical column densities (VCDs),	
17	retrieved by applying the Optimal Estimation Method to the MAX-DOAS observations, have	
18	been used to study the seasonal and diurnal cycles of SO2, in combination to correlative	
19	measurements from in situ instruments, as well as meteorological data. A marked seasonality	
20	was observed in both SO_2 VCD and surface concentration, with a maximum in winter	
21	(February) and a minimum in summer (July). This can be explained by the larger emissions in	
22	winter due to the domestic heating and, in case of surface concentration, by more favorable	
23	meteorological conditions for the accumulation of SO ₂ close to the ground during this period.	
24	Wind speed and direction are also found to be two key factors in controlling the level of the	
25	SO ₂ -related pollution at Xianghe. In the case of east or southwest wind, the SO ₂ concentration	

does not change significantly with the wind speed, since the city of Tangshan and heavy 1 2 polluting industries are located to the east and southwest of the station, respectively. In contrast, when wind comes from other directions, the stronger the wind, the less SO₂ is 3 observed due to a more effective dispersion. Regarding the diurnal cycle, the SO₂ amount is 4 larger in the early morning and late evening and lower at noon, in line with the diurnal 5 variation of pollutant emissions and atmospheric stability. A strong correlation with 6 7 correlation coefficients between 0.6 and 0.9 is also found between SO₂ and aerosols in winter, 8 suggesting that anthropogenic SO_2 , through the formation of sulfate aerosols, contributes significantly to the total aerosol content during this season. The observed diurnal cycles of 9 MAX-DOAS SO2 surface concentration are also in very good agreement (correlation 10 coefficient close to 0.9) with those from collocated in-situ data, indicating the good reliability 11 and robustness of our retrieval. 12

13 **1 Introduction**

14 Sulfur dioxide (SO_2) , one of the most common air pollutants, is of major concern in pollution 15 control acts (Gauderman et al., 2000). In China, the Ministry of Environmental Protection 16 (MEP) lists SO₂ as one of the three conventional pollutants, together with NO₂ and PM10, and daily averaged SO₂ concentrations were used as an indicator to quantify the level of pollution 17 18 (Yan et al., 2010). This trace gas is predominantly produced by the burning of fossil fuels 19 including oil and coal, and the smelting of mineral ores that contain sulfur (Yan et al., 2005; 20 Zhao et al., 2012). SO₂ contributes to a large extent to the process of acidification resulting in 21 acid rain and to the formation of sulfate aerosols, both of which cause human health damages, 22 building surface corrosion, and visibility reduction. In particular, the secondary pollutant 23 sulfate aerosols generated by SO_2 are the primary source of fine solid particles in cities, which 24 are also responsible for severe air pollution issues (Meng et al., 2009). In addition, the 25 on-going industrial development, population growth, and heavy traffic contribute to higher energy consumption and therefore, to an increase in SO₂ emissions into the atmosphere (Wu 26 27 et al., 2013). Consequently, in order to meet the urgent demand to improve and control air quality in China, as well as to promote sustainable development, it is of the greatest
 importance to study the evolution of a pollutant like SO₂ and to identify its possible origins.

3 So far, the SO₂ surface concentration has been monitored using in-situ and long-path DOAS 4 (Differential Optical Absorption Spectroscopy) instruments (Meng et al., 2009), while satellite 5 sensors like GOME, SCIAMACHY, GOME-2, OMI, OMPS, and IASI have shown their ability to measure the SO₂ vertical column density (VCD) over polluted areas (see e.g. 6 Eisinger and Burrows, 1998; Krotkov et al., 2006; Lee et al., 2009; Nowlan et al., 2011; 7 Fioletov et al., 2013; Yang et al., 2013; Boynard et al., 2014). During the last decade, a new 8 remote sensing technique called MAX-DOAS (Multi-Axis Differential Optical Absorption 9 Spectroscopy) has been developed, providing information on both VCD and vertical 10 11 distribution of trace gases in the troposphere (Hönninger et al., 2004; Platt and Stutz, 2008). It 12 is based on the measurement of sunlight scattered at multiple elevation angles towards the 13 horizon, thus increasing the sensitivity to absorbers present close to the ground compared to 14 the zenith viewing geometry (Hönninger et al., 2004). MAX-DOAS studies published so far 15 have been mainly focused on the retrieval of NO₂ (e.g. Wittrock et al., 2004; Vlemmix et al., 2010; Frins et al., 2012; Hendrick et al., 2014; Ma et al., 2013; Wang et al., 2014), halogen 16 17 oxides like BrO and IO (e.g. Frieß et al., 2011; Großmann et al., 2013), formaldehyde (e.g. 18 Heckel et al., 2005; Wagner et al., 2011), and aerosols (e.g. Wagner et al., 2004; Frieß et al., 19 2006; Clémer et al., 2010). A lot of work has been done on MAX-DOAS measurements of 20 volcanic SO₂ (e.g. Bobrowski et al., 2007a; Galle et al., 2010), but so far, only a few studies 21 deal with MAX-DOAS observations of this species in polluted area (e.g. Irie et al., 2011; Lee 22 et al., 2008; Wu et al., 2013), despite the fact that as for other trace gases like NO₂, HCHO, and BrO, the combination of both surface concentration and VCD retrievals makes 23 MAX-DOAS a useful technique for validating SO₂ satellite data. 24

Here we present three years (March 2010-February 2013) of continuous MAX-DOAS SO₂ observations at the Xianghe Observatory, China (39.75°N, 116.96°E), located at about 50 km southeast of Beijing, at the borders among Beijing, Tangshan and Tianjin (see Fig. 1). The station is operated by the Institute of Atmospheric Physics (IAP)/ Chinese Academy of

Sciences (CAS) while the MAX-DOAS instrument was developed by the Belgian Institute for 1 2 Space Aeronomy (BIRA-IASB) and validated in several intercomparison exercises, in particular as part of the international Cabauw Intercomparison of Nitrogen Dioxide measuring 3 Instruments (CINDI, Roscoe et al., 2010) and more recently a national Chinese MAX-DOAS 4 instruments intercomparison campaign held in Xianghe (Wang et al., 2013). SO₂ 5 MAX-DOAS observations are used here in combination with in-situ measurements as well as 6 7 conventional meteorological data (temperature, humidity, wind direction and speed) to 8 investigate the seasonal and diurnal cycles of SO₂ vertical profiles and VCDs. The paper is divided into three main Sections. In Sect. 2, the SO₂ measurements are described, including 9 10 the DOAS analysis, vertical profile retrieval, and retrieval verification through comparison with in situ data. The seasonal and diurnal cycles of SO₂, and the relationship between SO₂ 11 12 and aerosols are investigated in Sect. 3. Finally, conclusions are given Sect. 4.

13 **2 Data**

14 **2.1 Instrument**

The MAX-DOAS instrument operated at the Xianghe Observatory consists of three 15 components: a thermo-regulated box containing two spectrometers, an optical head mounted 16 17 on a sun tracker, and two computers for instrument control and data storage (Clémer et al., 18 2010). The optical head and the two spectrometers are linked by two-way splitter optical 19 fibers (Clémer et al., 2010; Wang et al., 2013). This setup is capable of measuring scattered as 20 well as direct sunlight. One spectrometer works in the UV region (300 to 390 nm) and its 21 instrumental function is close to a Gaussian with a full width at half maximum (FWHM) of 22 0.4 nm. The other spectrometer covers the visible wavelength range from 400 to 720 nm with 23 a FWHM equal to 0.9 nm. During the observation, the azimuth direction of the telescope is 24 fixed to the North. A full MAX-DOAS scan consists of 9 elevation viewing angles (2°, 4°, 6°, 8°, 10°, 12°, 15°, 30°, and 90°) and lasts about 15 minutes (Clémer et al., 2010). The 3-year 25 data set investigated in this study covers the March 2010 to February 2013 period. 26

1 2.2 DOAS analysis

Scattered-sunlight spectra measured at different elevation angles (EVAs) are analyzed using
the DOAS technique (Platt and Stutz, 2008) where high-frequency molecular absorption
structures in the UV and visible regions of the spectrum are exploited to detect and quantify a
number of key atmospheric gases such as SO₂.

6 In this work, the spectra obtained from MAX-DOAS observations are analyzed using the **BIRA-IASB** 7 **QDOAS** spectral-fitting software suite developed at 8 (http://uv-vis.aeronomie.be/software/QDOAS/). QDOAS calculates the SO₂ differential slant 9 column densities (DSCDs), which are defined as the difference between the trace-gas 10 concentration integrated along the effective light path and the amount of the absorber in a 11 measured reference spectrum. (MAX-)DOAS is recognized as a "self-calibrating" technique 12 because differential absorptions are measured and therefore the impact of possible 13 instrumental degradations can be largely removed by using appropriate reference spectra. In 14 contrast, in-situ instruments need to be optically and/or chemically calibrated on a regular 15 basis, especially when performing long-term measurements. For tropospheric studies, a zenith 16 spectrum is frequently chosen as reference, in this way also removing the contribution of the 17 stratosphere in off-axis DSCDs.

The SO₂ DOAS settings have been investigated through sensitivity tests on several key parameters, such as wavelength interval, choice of absorption cross sections, polynomial order, and intensity off-set terms. The selected settings are summarized in Table 1 and described below.

SO₂ fitting windows ranging between 303 and 325 nm have generally been used in previous studies (Bobrowski and Platt, 2007b; Lee et al., 2008; Galle et al., 2010; Irie et al., 2011). At wavelengths shorter than 303 nm, the limiting factor is the strong ozone absorption which interferes with SO₂, leading to lower signal to noise ratio. At wavelengths longer than 325 nm, the SO₂ differential absorption signal becomes too weak. In order to identify the wavelength interval which minimizes both random and systematic uncertainties on SO₂ retrieval, 6

1 wavelength intervals have been investigated. The results of these sensitivity tests for two example days are presented in Figs. 2 and 3. On the first day (1st October 2011), the SO₂ 2 content is minimum and stable in time. On the second day (4th October 2011), large variations 3 of the SO₂ content occur, so the ability of the different intervals to give consistent and stable 4 values can be verified. As can be seen, the 305-317.5 nm interval provides the lowest fitting 5 6 errors throughout the day and the smallest dependence on the solar zenith angle (SZA) for 7 both days. Due to the larger absorption and therefore interference by O_3 at large SZAs, it has been decided to exclude measurements taken at SZAs larger than 75°. For these tests, the 8 following spectral signatures have been included: SO₂, O₃, NO₂, and the Ring effect (Grainger 9 and Ring, 1962; Chance and Spurr, 1997). Daily zenith-sky radiance spectra recorded around 10 11 local noon have been selected as reference. To account for the temperature dependence of the ozone absorption, cross sections at 2 different temperatures (223°K and 243°K) were used 12 13 according to Van Roozendael et al. (2006). A fifth-order polynomial is applied to fit the 14 low-frequency spectral structure due to Rayleigh and Mie scattering and instrumental effects. 15 Attempts to further adjust these settings, e.g. by adding BrO cross-section or by including 16 additional ozone correction terms according to Pukite et al. (2010) were not successful (less 17 stable retrievals with larger noise on the SO₂ DSCDs).

Fig. 4 shows a typical example of a DOAS fit for SO₂ at 43° SZA. We see that fitting 18 residuals range in between -2×10^{-3} and 2×10^{-3} , corresponding to a root-mean-squares (RMS) 19 of 9×10^{-4} , which appears to be small in comparison to the SO₂ differential structures presented 20 in the lowest panel of the figure. The typical fitting uncertainty on SO₂ DSCDs is of about 21 $1-6 \times 10^{15}$ molec cm⁻² (less than 10%), and for the case illustrated here, corresponds to 2%. For 22 near-noon conditions, the detection limit on the SO₂ DSCD can be conservatively estimated 23 as 3 times the one-sigma uncertainty on the slant column, which means approximately 3×10^{15} 24 molec \cdot cm⁻². This detection limit is similar for the vertical columns estimated using the 25 geometrical approximation at 30° elevation (see Sect. 2.3). Vertical columns derived from the 26 full inversion generally have a smaller detection limit, owing to the gain in sensitivity 27 obtained when including near horizontal viewing measurements. 28

1 2.3 Profile retrieval

2 SO₂ vertical profiles are retrieved for each MAX-DOAS scan by using the bePRO profiling 3 tool developed at BIRA-IASB (Clémer et al., 2010; see also Hendrick et al., 2014). It is based 4 on the Optimal Estimation Method (Rodgers, 2000) and includes the LIDORT radiative 5 transfer model (RTM) as a forward model. A two-step approach is implemented in bePRO: First, aerosol extinction profiles are retrieved from measured O₄ DSCDs. This step is needed 6 7 because the aerosols strongly influence the effective light path in the atmosphere and therefore the optical density of trace gases like SO₂. Secondly, bePRO is applied to measured 8 9 trace-gas DSCDs using the retrieved aerosol extinction profiles for the radiative transfer 10 calculations (see below). Since the DOAS analysis is performed using daily zenith radiance 11 spectra around noon as reference, bePRO is feeded for each scan with SO₂ and O₄ DSCDs 12 obtained by taking the difference between off-axis DSCDs and the zenith DSCD interpolated 13 at the time of each off-axis measurement using the zenith DSCDs of two consecutive scans. Proceeding this way allows to properly remove the contributions of the stratosphere from the 14 measurements and is similar, at least for SZA < 75°, as taking the zenith spectrum of each scan 15 16 as reference for the DOAS analysis.

Both linear and non-linear iterative approaches have been implemented in our profiling algorithm. For weak absorbers like NO_2 , HCHO and SO_2 , the linear method is selected (see e.g. Hendrick et al., 2004). In case of strong absorbers like O_4 , the non-linear iterative approach is used:

21
$$\mathbf{x}_{i+1} = \mathbf{x}_i + (\mathbf{S}_a^{-1} + \mathbf{K}_i^{\mathrm{T}} \mathbf{S}_{\varepsilon}^{-1} \mathbf{K}_i)^{-1} \cdot [\mathbf{K}_i^{\mathrm{T}} \mathbf{S}_{\varepsilon}^{-1} (\mathbf{y} - \mathbf{F}(\mathbf{x}_i)) - \mathbf{S}_a^{-1} (\mathbf{x}_i - \mathbf{x}_a)]$$
(1)

where *y* is the observation vector with the DSCDs at the different EVAs, **F** is the forward model describing the physics of the measurements, **K** is the weighting function, expressing the sensitivity of the measurements to changes in the aerosol extinction or SO₂ vertical profile and calculated on-line by the LIDORT RTM, S_{ε} is the measurement uncertainty covariance matrix, x_a and S_a are the a priori vertical profile and its corresponding error covariance matrix.

1 A priori information is needed in the OEM method in order to indirectly reject unrealistic 2 solutions compatible with the measurements. Another important quantity in the OEM is the averaging kernel matrix A, which represents the sensitivity of the retrieval to the true state. 3 More specifically, each element A_{ij} in the matrix A describes the sensitivity of the retrieval at 4 i^{th} level to the true states at the different altitude levels *j*. Furthermore, the trace of the matrix 5 A gives the degrees of freedom of signal (DFS), which corresponds to the number of 6 7 independent pieces of information contained in the measurements. Due to the nonlinearity of 8 the inverse problem in case of aerosols, the solution to equation (1) must be iterated until satisfactory convergence is achieved between measured DSCDs and those calculated using 9 10 the retrieved aerosol extinction vertical profile.

11 Regarding the choice of the a priori profile x_a , exponentially decreasing a priori SO₂ and 12 aerosol extinction profiles with a fixed scaling height of 0.5km have been constructed 13 according to the following expression:

14
$$X_{a}(Z) = \frac{\text{VCD}_{a}}{\text{SH}} e^{-\frac{Z}{\text{SH}}}$$
(2)

where $x_a(z)$ is the a priori profile, SH the scaling height (0.5 km), and VCD_a (AOD_a) is the a 15 16 priori vertical column density (aerosol optical depth). For each scan, VCD_a is derived using the geometrical approximation method, i.e. the SO_2 layer is assumed to be located below the 17 18 scattering altitude at 30° EVA, so that tropospheric SO₂ VCDs can be derived by applying a geometrical air mass factor (AMF) to measured 30° EVA DSCDs (Hönninger et al., 2004; 19 Brinksma et al., 2008; see also Hendrick et al., 2014). In case of aerosols, a fixed AOD of 0.2 20 is used. Since the DOAS fitting intervals are different for SO₂ and aerosols, the aerosol 21 extinction profiles utilized as input for the calculation of SO₂ weighting functions have been 22 23 derived by directly converting the aerosol profiles retrieved in the 338-370 nm wavelength range to the 305-317.5 nm interval using the Ångström exponents (Cachorro et al., 2000) 24 25 retrieved from collocated CIMEL/AERONET supplotometer measurements (Holben et al., 26 1998; see http://aeronet.gsfc.nasa.gov):

1 Extinction(z, 313 nm) = Extinction(z, 360 nm) x $(313/360)^{-\alpha}$ (3)

2 where z is the altitude and α is the Ångström exponent.

17

3 The 340-440 nm exponents are used in a first approximation since values for a wavelength 4 range closer to the SO_2 fitting interval (305-317.5 nm) are not available so far. The corresponding mean scaling factor for the March $2010 - \text{February } 2013 \text{ period is of } 1.16 \pm 0.06.$ 5 6 The single scattering albedo and phase function of aerosols at 360 nm required by bePRO for 7 retrieving aerosol extinction profiles are calculated off-line based on the aerosol size distribution and refractive index retrieved from the same CIMEL/AERONET sunphotometer 8 9 measurements as above. The temperature-pressure profiles are obtained from the US standard atmosphere. S_{ε} and S_a matrices are similar as in Clémer et al. (2010) and Hendrick et al. 10 11 (2014). S_{ε} is a diagonal matrix, with variances equal to the square of the DOAS fitting error. 12 For S_a , the diagonal element corresponding to the lowest layer, $S_a(1,1)$, is set equal to the 13 square of a scaling factor β times the maximum partial VCD (AOD) of the profiles. Here 14 β =0.4 for SO₂ and 0.2 for aerosol. The other diagonal elements decrease linearly with altitude down to $0.2 \times S_a(1,1)$. The off-diagonal terms in S_a , were set using Gaussian functions as 15 16 follows:

$$\mathbf{S}_{a}(i,j) = \sqrt{\mathbf{S}_{a}(i,i)\mathbf{S}_{a}(j,j)\exp(-\ln(2)(\frac{z_{i}-z_{j}}{\gamma})^{2})}$$
(4)

where z_i and z_j are the altitudes of ith and jth levels, respectively. The correlation length is set to 0.1 km for SO₂ and 0.05km for aerosol in order to optimize the DFS.

The retrieval altitude grid is also the same as in Clémer et al. (2010) and Hendrick et al. (2014), i.e. ten layers of 200 m thickness between 0 and 2 km, two layers of 500 m between 2 and 3 km and 1 layer between 3 and 4 km.

Fig. 5 shows an example of a SO₂ profile retrieval (Xianghe, 29 September 2010, 10:15 LT).
Fig. 5(a) compares the a priori and retrieved profiles; Fig. 5(b) shows an example of fit results,
i.e. the comparison between measured DSCDs and those calculated from the retrieved profile.
The quality of the profile retrieval is checked for each scan by calculating the relative Root

Mean Square Error (RMS) between observed and calculated DSCDs. This RMS corresponds to the standard RMS expressed in molec/cm² divided by the mean DSCD of the scan. All retrievals based on the following selection criteria have been selected: RMS < 15%, DFS >0.7, and negative values not allowed. For each year, the number of selected retrievals using these criteria reaches ~70% of the total number of scans.

Also shown in Fig. 5 are the smoothing and noise errors (c) and the averaging kernels (d). 6 7 Regarding the errors, the smoothing error limits the ability of the retrieval to obtain solutions far from the a priori, while the noise error is related to the propagation of the noise in the 8 measurements into the retrieval (Rodgers, 2000). From Fig. 5(c), we see that the smoothing 9 error is significantly larger than the noise error, except in the 0-200m layer. The averaging 10 11 kernels show that the retrieval is mainly sensitive to the layer close to the surface in addition to the total vertical column. In this example, the DFS is about 2.4, suggesting that two 12 independent pieces of information can be determined from the measurements. 13

14 The error budget is presented in Table 2. Uncertainty related to aerosols is estimated by retrieving SO₂ profiles using wavelength-converted retrieved aerosol profiles plus their 15 16 corresponding error (i.e. the sum of smoothing and noise errors plus a 20% error due to the 17 uncertainty on the O_4 cross sections (Clémer et al., 2010)) as input and comparing the results to the standard retrievals. The uncertainty on the SO₂ cross sections is set to 5%, as suggested 18 19 by Vandaele et al. (1994). The uncertainty on the a priori profiles is estimated by taking SH =20 1 km in Eq. (2) instead of 0.5 km in the standard retrieval. The total uncertainty is calculated by adding the different terms in Gaussian quadrature. 21

Monthly-mean SO_2 profiles are shown in Fig. 6. There is a maximum SO_2 concentration in the 200-400m layer for each profile, except in summer where the maximum is located near the surface. The largest vertical gradient is observed in February and November, the minimum in July and August. This is mainly due to the fact that the SO_2 emissions are the highest in February and November. This will be discussed in detail below.

27 Fig. 7 shows the seasonal mean of diurnal cycle of DFS. The diurnal distribution in any

season shows a single peak at mid-day due to the fact that the retrieval error at late evening or early morning overweights that at noon. If we compare the DFS around noon among the different seasons, values in summer are lower compared to the other seasons due to the lower SO₂ amounts associated with larger uncertainties observed during this period.

5

6 2.4 SO₂ surface concentration retrieval verification

For verification purpose, our retrieved SO_2 surface concentrations have been compared to measurements from a modified commercial in-situ instrument, based on pulsed UV fluorescence technology (Thermo Environmental Instruments Model 43C) (Li et al., 2007). Comparison results for December 2011 when the in-situ instrument was freshly calibrated are shown in Fig. 8. Hourly and daily averages of SO_2 concentration are plotted in Fig. 8(a) and (b), respectively. A good agreement is obtained with a correlation coefficient of 0.86 and a slope of 0.95.

In Fig. 9, the daytime variations of the MAX-DOAS and in-situ SO₂ surface concentration are compared for 9 continuous days. A very good agreement is found between both data sets, indicating the good overall reliability and the robustness of our MAX-DOAS retrievals.

17 **3 Results and discussion**

Based on the SO₂ profiles retrieved for the period from March 2010 to February 2013, we have investigated the daily and seasonal variations of the SO₂ VCD and surface concentration and the possible influence of meteorological conditions, including atmospheric stability, wind direction and speed. We have adopted the following convention for the seasons: MAM, JJA, SON, and NJF for spring, summer, autumn, and winter, respectively.

23 **3.1 Seasonal variation of SO₂**

Fig. 10(a) shows that the SO₂ VCD is highly correlated with concentration close to the ground

(correlation coefficient of 0.85). From Fig. 10(b), we see that the temporal evolutions of SO₂
 VCD and concentration are very similar, consistent with the fact that the SO₂ emission
 sources are located near the ground.

4 The monthly averaged SO₂ VCD and surface concentrations are shown in Fig. 11. Both show 5 a marked seasonal signature with a maximum in winter and a minimum in summer, implying 6 that SO₂ originates mainly from human sources rather than natural ones (Lin et al., 2011). 7 Generally, the fluctuations of any atmospheric pollutant in a region of interest can be mainly attributed to three factors: emission level, residence time, and atmospheric transport (Wang et 8 9 al., 2010; Lin et al., 2011). From the perspective of emission level, firstly, owing to enhanced 10 domestic heating and associated coal and oil consumption in winter, the heating-related 11 emissions of SO₂ are much larger during this period than in summer. Secondly, the residence 12 time, defined as the rate of removal mechanisms, also plays an important role in determining the seasonal variation of SO₂ concentrations (Lee et al., 2011). Processes responsible for the 13 14 removal of SO₂ involve dry and wet deposition and homogeneous or inhomogeneous 15 gas-phase reactions leading to the production of H₂SO₄ or sulfate (Tu et al., 2004). As shown 16 in Fig. 12, the relative humidity is lower in winter, so that the removal of SO_2 through wet 17 deposition is not as substantial as in summer. Thirdly, the transport can also influence the 18 evolution of SO_2 at a given location. Although in winter the wind is stronger at Xianghe, the 19 emissions also increase during the same period. In addition, the reduced atmospheric 20 boundary layer height and frequent temperature inversion events result in larger surface 21 concentrations due to an accumulation of SO₂ in the lower troposphere (Meng et al., 2009). In 22 summary, the aforementioned three factors jointly lead to the observed seasonal pattern of 23 SO₂ concentration in Xianghe.

From Fig. 11, we see that the amount of SO₂ strongly increases in November with respect to October, as a consequence of increasing domestic heating (November is the beginning of the domestic heating season). Moreover, the higher wind speed observed in December (see Fig. 12) leads to a decrease of SO₂ during this month due to more efficient diffusion and dilution effects. Finally, it is also noticeable that SO₂ in January 2011 is remarkably lower than that in 1 other years. This will be further discussed below.

2 **3.2** Impact of meteorological conditions

3 Because of the high correlation coefficient and similar seasonal variations of the SO₂ VCD 4 and concentration, we decided to investigate the impact of meteorological conditions on 5 VCDs only. The variation of the SO₂ VCD is closely linked not only to the spatial distribution 6 of emission sources but also to meteorological conditions including wind (speed and direction) and precipitation. As shown in Fig. 12, in general, the variations of temperature and humidity 7 8 appear to exhibit similar behavior from year to year. This suggests that the contribution of the 9 wind speed and direction as driver for the SO₂ VCD variation is probably different over the different years investigated here. We further explore the relationship between SO₂ and wind 10 (speed and direction), as displayed in Fig. 13. It can be seen that the amount of SO_2 is strongly 11 12 dependent on the wind direction (Fig. 13a): high VCDs are prominent when the winds blow 13 from the east, because Tangshan, a heavy industrial city releasing large amounts of SO₂, is 14 situated to the east of Xianghe (see Fig. 1); in contrast, the northwest direction corresponds to 15 a minimum in SO₂ VCD, since it is a mountain area, characterized by much less emissions 16 than in Xianghe. The wind therefore contributes significantly to the dispersion of the 17 pollutants, as expected. Regarding the dependence of the SO₂ VCD on wind speed, Fig. 13(b) shows that the VCD is almost constant with wind speed for the E and SW, which means that 18 19 no good dispersion happens with the wind from these directions, since high-emission 20 industrial areas and Tangshan are located to the southwest and east of Xianghe, respectively. 21 In contrast, an anti-correlation is observed for NE/NNE, NW, and SE, which means that the 22 wind from these directions corresponding to less polluted areas can efficiently disperse 23 pollutants. In addition, the SO₂ content at Xianghe is more sensitive to the emission sources in 24 Tangshan (E) than in Beijing (WNW), which is consistent with the fact that Beijing has taken 25 regulatory actions to reduce air pollution through traffic-control measures and the closure of 26 heavy polluting industries initiated before the 2008 Olympic Games (Yu et al., 2010).

27 The annual cycles of SO_2 are generally in good agreement among the different years.

However, the SO₂ VCD in January 2011 drastically deviates by up to 30% from the values 1 2 during the same month in 2012 and 2013, which is also the case in May 2012. Wind roses in Fig. 14 reveal that the inter-annual variability of wind speed and direction is responsible for 3 the significantly different SO₂ VCD in January 2011. During that month, the frequency of 4 north-west winds reaches 70% and wind speed predominantly exceed 5m.s⁻¹. As mentioned 5 above, the strong northwesterly wind favors the atmospheric dispersion of pollutants. 6 Consequently, the SO₂ VCDs are generally lower than 4×10^{16} molec.cm⁻². For January 2012 7 and 2013, uniformly distributed wind on each side and low velocity (<5 m.s⁻¹, 8 frequency>50%) jointly result in relatively high SO₂ VCDs compared to January 2011. 9 Similar features can explain the May 2012 case. 10

11 **3.3 Diurnal Cycle**

12 In Fig. 15, we further compare the diurnal cycles of SO_2 VCDs for the different seasons. 13 Since the sunshine duration is different in the four seasons, the available time period for MAX-DOAS observations also differs: 7:30-17:30 in spring and autumn, 6:30-18:30 in 14 15 summer, and 8:30—16:30 in winter. As can be seen, the diurnal cycles for all years are very consistent, especially in summer. The retrieved SO₂ VCDs in autumn 2011 and spring 2012 16 17 are significantly higher than those during the same period of the other years due to the 18 anomalous VCD values in November 2011 and May 2012. Furthermore, the amplitude of the SO₂ VCD diurnal cycle, which shows a minimum at noon and a maximum in the morning and 19 20 late afternoon, is larger in winter. This can be explained a strengthened diurnal variation of 21 emission sources during this period (Meng et al., 2009).

It should be noted that similar investigations have been done for NO_2 (Wang et al., 2014). One can conclude that both NO_2 and SO_2 display a similar seasonal variation and are impacted in the same way by meteorological conditions. However, SO_2 abundances are always higher than NO_2 ones and their diurnal cycles are different, especially in winter and summer: SO_2 has a more pronounced diurnal cycle than NO_2 in winter which is in line with the known diurnal cycle of burning of fossil fuels for heating and atmospheric stability, and the photochemical reaction activity leads to an obvious decrease of NO₂ during daytime in summer (Wang et al.,
 2008; Meng et al., 2009; Lin et al., 2011).

3 3.4 Relationship between SO₂ and aerosols

SO₂ is known as a major aerosol precursor in the Beijing area through its conversion into sulfates and sulfuric acid by reaction with OH (see e.g. Ma et al., 2012 and Zhang et al., 2013). Since aerosol extinction profiles are retrieved in the first step of the SO₂ retrieval (see Sect. 2.3), our data set offers a unique opportunity to investigate the relationship between SO₂ emission and aerosol production in the suburban Beijing. This will be done through a correlation study as in Lu et al., 2010 and Veefkind et al. (2011).

10 Fig. 16 shows monthly scatter plots of the SO₂ concentration versus aerosol extinction coefficient retrieved in the 0-200m layer for the March 2010 – February 2013 period. In all 11 12 plots, data points correspond to MAX-DOAS scans satisfying the selection criteria based on 13 the quality of the retrievals (see Sect. 2.3). A strong correlation (Pearson correlation 14 coefficients in the 0.6-0.9 range) is obtained in J, F, M and O, N, D while a significantly lower 15 correlation is observed in late spring/summer with correlation coefficients around 0.3 in J, J, A. Similar features are found from the scatter plots of SO₂ VCD versus AOD but also when 16 outliers outside the 95% confidence interval are removed and/or the uncertainties on both SO₂ 17 and aerosol data are taken into account (not shown here). The marked seasonality of the 18 19 correlation between SO_2 and aerosols is further illustrated in Fig. 17 where monthly 20 correlation coefficients for both surface concentration and integrated column are reported. The 21 positive correlation (>0.2) observed throughout the year indicates that in most cases, high 22 pollution events in Xianghe are associated with enhanced SO₂ and aerosol levels (Chan and 23 Yao, 2008; Li et al., 2007). The higher correlation coefficients obtained in winter (>0.6)suggest that anthropogenic SO₂, through the formation of sulfate aerosols, is a major 24 25 contributor to the total aerosol content during this period of the year. In late spring/summer, 26 the Beijing area is strongly influenced by other sources of aerosols, especially particles 27 emitted from massive agricultural fires in the surrounding region (Xia et al., 2013) as well as

dust particles transported from the Kumutage and Taklimakan deserts in western China and 1 from the Mongolian deserts (Yu et al., 2009). These perturbations by other aerosol sources 2 combined to lower SO_2 emissions, shorter lifetime of SO_2 due to a more efficient oxidation, 3 and different meteorological conditions, could likely explain the significantly weaker 4 correlation between anthropogenic SO₂ and aerosols obtained in J. J. A. The intercept values 5 much larger than zero found in summer scatter plots (see Fig. 16) further support the fact that 6 aerosol sources other than anthropogenic ones play a significant role in summer, as also 7 suggested by Lu et al. (2010) from a correlation study between SO₂ emission inventories and 8 9 AODs measured by the MODIS satellite instrument. It is however important to note that co-located measurements of the chemical composition of aerosols in Xianghe as well as 10 additional investigations on the type and photochemical age of the air masses probed by the 11 12 MAX-DOAS instrument would be needed to confirm our findings.

13

14 **4 Summary and conclusions**

Tropospheric SO₂ vertical profiles and corresponding column densities at the Xianghe station 15 16 have been retrieved by applying an OEM-based profiling tool to continuous ground-based MAX-DOAS observations from March 2010 to February 2013. The 305-317.5 nm 17 wavelength range was found to be the most suitable fitting window for near-noon DOAS 18 analysis of SO_2 . For verification purpose, retrieved SO_2 surface concentrations have been 19 20 compared to collocated in-situ data. An excellent agreement was found, with correlation 21 coefficient and slope close to 0.9, indicating the good reliability and robustness of our retrievals. 22

These MAX-DOAS measurements have been used to investigate the seasonal and diurnal cycles of SO₂ vertical columns and surface concentrations, in combination with conventional meteorological data (temperature, humidity, and wind speed and direction). Regarding the seasonal variation, both VCD and surface concentrations exhibit the same patterns, with a maximum in winter (February) and a minimum in summer (July), in accordance with the large

1 emissions due to domestic heating in winter. The high levels of SO₂ during the cold season are 2 further enhanced by the weakness of the wet deposition mechanism and, in case of surface concentration, by the frequent temperature-inversion events occurring during this period, 3 favoring the accumulation of SO₂ in the atmospheric layers close to the ground. The variation 4 of the SO₂ amount in Xianghe is also found to be largely driven by wind speed and direction. 5 In the case of east or southwest wind, the VCD at the station remains almost constant with the 6 7 increase of wind speed, since the city of Tangshan and heavy polluting industries are located 8 to the east and southwest of Xianghe, respectively. In contrast, an anti-correlation between SO₂ VCD and wind speed is observed for NE/NNE, NW, and SE directions, which means the 9 10 wind from these directions can efficiently disperse the pollution in Xianghe. With respect to the diurnal cycle, larger SO_2 amounts are obtained in the early morning and late evening with 11 a minimum around noon, in line with the diurnal variation of pollutant emission and 12 atmospheric state. Moreover, the diurnal cycle is more pronounced during wintertime, mainly 13 due to the more marked diurnal variation of emission sources during this season. The 14 15 relationship between SO₂ and aerosols has been also investigated. A strong correlation 16 between both is found in winter but not in summer. This seasonality could be related to the 17 fact that in the Beijing area in winter, the aerosol content depends significantly on anthropogenic SO₂ through the formation of sulfate aerosols while in spring/summer, dust and 18 biomass burning particles, which are much less SO₂-dependent, are also important aerosol 19 sources. It is however worth noting that such kind of correlation analysis should be combined 20 to aerosol composition measurements in order to definitely conclude whether the conversion 21 22 of SO₂ to sulfate is a dominant aerosol source or not.

These three-year MAX-DOAS SO_2 measurements in Xianghe constitute a unique data set for validating and improving space-borne observations over China, which is the region in the world where anthropogenic SO_2 emissions are the largest (Yang et al., 2013; Boynard et al., 2014). In particular, retrieved SO_2 vertical profiles can be used as a priori information for the AMF calculation in satellite retrievals. Moreover, the combination of both integrated columns and surface concentrations could provide useful information to make explicitly the link

- 1 between measured satellite columns and surface concentrations.
- 2

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Parameter	Data source	Fitting interval (nm)	
		338-370 (O ₄)	305-317.5(SO ₂)
NO ₂	Vandaele et al. (1998) 220K, 294K	x	x(only 294K)
SO_2	Vandaele et al. (1994) 294K		Х
O ₃	Bogumil et al. (2003) 223K, 243K	x(only 223K)	Х
O_4	Hermans et al. (2003) 296K	Х	
BrO	Fleischmann et al. (2004) 223K	Х	
H ₂ CO	Meller and Moortgat (2000) 293K	х	
Ring	Chance and Spurr (1997)	Х	Х
Polynomial		5	5
degree			

1 Table 1: Settings used for the SO_2 and O_4 DOAS analysis.

Uncertainty (%)	Concentration (0-200m)	VCD
Smoothing + noise errors	16	11
Uncertainty related to aerosols	16	5
Uncertainty related to the a priori	8	19
Uncertainty on SO ₂ cross section	5	5
Total uncertainty	24	23

1 Table 2: Error budget of retrieved SO₂ concentration (0-200m) and VCD.





2 Figure 1: Location of the Xianghe Observatory (red star) and major neighborhood cities.



Figure 2: SO₂ DSCDs (1st column) and corresponding fitting uncertainties (2nd column) retrieved at 4°
(upper plots), 30° (lower plots) elevation for different wavelength intervals on 1st October 2011. Local
time (h) and corresponding SZA (°) are given on the x-axis.



2 Figure 3: Same as Figure 2, but for 4th October, 2011.



Figure 4: Example of DOAS fit result for SO₂. It corresponds to 29 September 2010 at ~11:20
LT. SZA and EVA values are 43° and 30°, respectively.



Figure 5: Example of SO₂ vertical profile retrieval from MAX-DOAS measurements at
Xianghe (29 September, 2010 at 10:15 LT). (a) a priori (blue) and retrieved profile (red); (b)
observed (red) and calculated (blue) DSCD (c) smoothing error (red), noise error (green) and
sum of these two (blue); (d) averaging kernels.



2 Figure 6: Monthly-averaged SO_2 concentration vertical profiles for the March 2010 -

- 3 February 2013 period.
- 4





2 Figure 7: Seasonally-averaged DFS diurnal cycles corresponding to the SO₂ profile retrievals.



Figure 8: (a) Scatter plot of in situ SO₂ surface concentrations (0-200m layer) against MAX-DOAS data for December 2011 (hourly-averaged concentrations). The red line denotes the linear least-squares fit to the data. (b) Temporal evolution of daily averaged MAX-DOAS and in situ SO₂ concentrations during December 2011. Gaps in the data series are due to missing MAX-DOAS measurements.



Figure 9: Comparison between in situ (blue, hourly means) and MAX-DOAS SO₂ surface
concentrations (red, each point represents the retrieval from one scan) for the December 15-23,
2011 period (upper plots are for December 15-17, middle plots are for December 18-20, lower
plots are for December 21-23).



Figure 10: (a) Scatter plot of SO₂ VCD against surface concentration. The red line represents
the linear least-squares fit to the data. (b) Temporal evolutions of monthly mean VCD and
concentration from March 2010 to February 2013.



- 2 Figure 11: Monthly mean SO₂ VCD (a) and surface concentration (b) for the March 2010 -
- 3 February 2013 period.





Figure 12: Seasonal cycles (monthly means) of temperature, humidity, and wind speed in
2010 (marker: star), 2011 (plus), 2012 (circle), and 2013 (square).



Figure 13: (a) Wind rose showing the SO₂ VCD (10¹⁶ molec.cm⁻²) as a function of the wind
direction (average for all wind speed). (b) Dependence of SO₂ VCD (10¹⁶ molec.cm⁻²) on
wind direction for different wind speeds.



Figure 14: Wind rose for wind speed $(1^{st} row; m.s^{-1})$ and SO₂ VCD $(2^{nd} row; 10^{16} molec.cm^{-2})$

3 for January 2011 (1^{st} column), 2012 (2^{nd} column), and 2013 (3^{rd} column).

4



2 Figure 15: (a) Seasonally-averaged SO_2 VCD diurnal cycles, and (b) corresponding errors.

3 Data points represent hourly means.

4



Figure 16: Scatter plots of aerosol extinction coefficient versus SO₂ concentration in the O-200m layer for months 1-12 of the March 2010 – February 2013 period (first row from left to right is for J, F, M, respectively; second row for A, M, J; third row for J, A, S; fourth row for O, N, D). The data points correspond to the different MAX-DOAS scans. The red line denotes the linear least-squares fit to the data.



Figure 17: Seasonal variation of the correlation coefficient between SO₂ and aerosols over the
March 2010-February 2013 period. The red curve corresponds to VCD versus AOD and the
blue curve to SO₂ concentration versus aerosol extinction coefficient in the 0-200m layer.