

Supplement of

Ground-based MAX-DOAS observations of tropospheric aerosols, NO2, SO² and HCHO in Wuxi, China, from 2011 to 2014

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Supplement for "Ground-based MAX-DOAS observations of tropospheric aerosols, NO2, SO² and HCHO in Wuxi, China, from 2011 to 2014"

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1 Meteorological conditions

The ground-based weather station near the MAX-DOAS instrument records the ambient temperature, wind speed and direction, and relative humidity during the whole observation period. Figure S1 shows their seasonally mean diurnal variations. A large seasonal difference occurs only for the ambient temperature, but not for the wind speed and relative

20 humidity. Similar diurnal variations for the three meteorology parameters are found for the different seasons. The ambient temperature and the relative humidity reach the maximum and minimum values around noon, respectively. The wind speed has the maximum value around 16:00 LT. The wind directions recorded by the same weather station are shown by the wind roses for the individual seasons in Fig. S2, indicating that the dominant wind is from the northeast in all seasons. In spring and summer the non-dominant wind directions occur more frequently than in winter and autumn.

25 **2 DOAS analysis and data screening**

In the DOAS analysis, the slant column densities (SCDs) of the trace gases (TGs) are retrieved from the off-axis spectra using a zenith measurement from the same elevation sequence as the Fraunhofer reference spectrum (FRS). As the latter also contains (usually small) absorptions features, the resulting SCD actually represents the differences between the SCDs of the measured spectrum and the FRS. This difference is usually referred to as the differential SCD (dSCD). The use of a FRS

30 from the same elevation sequence can minimise any effects caused by changes of the properties of the instrument (relevant for long term analyses) and the stratospheric absorptions (relevant for measurements at high solar zenith angle (SZA)). The

effect of rotational Raman scattering is considered by including a Ring spectrum (Shefov 1959; Grainger and Ring, 1962; Solomon et al., 1987; Chance and Spurr, 1997; Wagner et al., 2009) computed by the DOASIS software (Kraus, 2006, using a routine from Bussemer 1993). To account for the different wavelength dependencies of the filling-in in clear and cloudy skies, an additional Ring spectrum as described in Wagner et al. (2009) is also included.

- 5 For the retrieval of O_4 and NO₂, the wavelength range of 351 to 390 nm is selected, covering two O_4 absorption bands and several NO₂ absorption bands. A 3rd order polynomial is used. Besides the NO₂ cross section at the temperature of 294 K, another cross section at 220 K is also included in the fit to account for the temperature dependence of the $NO₂$ absorptions. The detailed DOAS settings for the retrieval are listed in Table 1 of the main manuscript. In Fig. S3a and b, the O_4 and NO_2 $dSCDs$ from all measurements are plotted against SZA. NO₂ and O₄ dSCDs show an obvious systematic increase or decrease
- 10 with increasing SZA, respectively, for SZA larger than 75°. For NO² this behaviour can be explained by the larger differences of the stratospheric light paths between the measurement and the FRS for large SZA. The opposite dependencies in the morning and evening (indicated by the different solar azimuth angles) are due to the decrease or increase of the stratospheric light path with time in the morning and evening, respectively. The $O₄$ behaviour might be related to the interference of the so called intensity offset (see below) and the O₄ absorption. But this hypothesis is still not clearly
- 15 confirmed.

A quite large relative intensity offset (RIO) is found for measurements at large SZAs as indicated in Fig. S3c, which implies a possible interference of the offset corrections and the derived TG dSCDs (see also Coburn et al., 2011). Thus we skip the data for the SZA larger than 75° to avoid the interference with the stratospheric contributions and RIO on the retrieved tropospheric dSCDs.

- $20\quad$ SO₂ dSCDs are retrieved in the wavelength interval from 307.8 nm to 330 nm including O₃, SO₂, HCHO cross sections and Ring spectra shown in Table 1 of the main manuscript. Wang et. al (2014) performed sensitivity studies to find the optimum wavelength interval which minimizes both random and systematic uncertainties on the $SO₂$ retrieval. They found that the wavelength range of 305 to 317.8 nm provides the lowest fitting errors. Below 305 nm, interference with the strong ozone absorption can affect the SO_2 retrieval. At small wavelengths also the signal to noise ratio decreases. Considering the rather
- 25 low sensitivity of the miniature spectrometer in the UV range used in our study compared to scientific grade spectrometer used in the study of Wang et. al (2014), here we limit the lower wavelength range to 307.8 nm. We also changed the upper wavelength range to 330 nm to minimise the possible interference with other species. The SO_2 , O_3 and Ring dSCDs as well as the intensity offset are plotted against SZAs in Fig. S4. At large SZAs strong changes of all quantities are found indicating the possible interference of the stratospheric ozone absorptions and the intensity offset on the SO_2 retrieval. To avoid these
- 30 interferences, we screen the SO_2 dSCD data for SZA larger than 75 $^{\circ}$.

HCHO dSCDs are retrieved in the wavelength interval from 324.6 to 359 nm including O_3 , O_4 , SO_2 , HCHO cross sections and Ring spectra shown in Table 1 of the main manuscript. Pinardi et. al (2013) found that the interferences between BrO, Ring spectrum and HCHO can strongly affect the retrieved HCHO dSCDs and they recommended the wavelength range of 336.5 to 359 nm, to minimise the uncertainties of the HCHO retrieval. In the wavelength range below 336.5 nm, the ozone

absorption interferes with the HCHO retrieval, like for SO_2 . However this conclusion is only appropriate for the retrieval with the daily noon zenith spectrum as the FRS. In this study, the sequential FRS is used. Because BrO is mostly located in the stratosphere, the difference of the BrO absorptions between the measurement and the FRS is negligible (the differential optical depth of the BrO absorption is typical only 1×10^{-4}) and much lower than using a daily noon FRS (typical 6×10^{-4}).

- 5 Considering that the BrO absorption signal is too weak to impact the HCHO retrieval, the BrO cross section is not included in the HCHO DOAS fit in this study. Tropospheric BrO is not expected to be found due to large $NO₂$ concentrations (e.g. Holla et al 2015). Moreover similar to the BrO interference, the effective stratospheric ozone absorption is also much smaller if a sequential FRS is used compared to a daily noon FRS. Thus the wavelength interval can be extended to a shorter wavelength to cover more and stronger absorption bands of HCHO. Moreover a wider wavelength range usually makes the
- 10 fit more stable, but at shorter wavelengths the interference of the ozone absorption is also stronger. To find the optimal retrieval wavelength interval, the examples of the HCHO retrieval in three different wavelength ranges of 310 to 359 nm, 324.6 to 359 nm and 336.5 to 359 nm are shown in Fig. S5. The measured structure from the DOAS fit in the wavelength range 310 to 359 nm indicates the strong interference of the ozone absorption. In addition, the HCHO dSCDs and the fitting errors in the three wavelength ranges on two days with low and high HCHO load are shown in Fig. S6. We find that the
- 15 HCHO dSCDs in the wavelength range of 324.6 to 359 nm are consistent with those in 336.5 to 359 nm, which is recommended by Pinardi et. al (2013). And both of them are quite different from the values in the wavelength range of 310 to 359 nm, especially on the day with the low HCHO load. The reverse "U" diurnal variation of the HCHO dSCDs in the wavelength range of 310 to 359 nm is an indication for the strong interference of the stratospheric ozone absorption. Moreover the wavelength range of 324.6 to 359 nm has much smaller fitting errors than the wavelength range of 336.5 to
- 20 359 nm. Thus we conclude that in general the wavelength range of 324.6 to 359 nm is the optimal wavelength range in which the ozone interference is weak and the fitting error is small. To avoid remaining interferences of the HCHO results with the stratospheric ozone absorption and intensity offset we exclude the HCHO dSCD for SZA > 75° (see Fig. S7). After applying these filters, the mean RMS of the residual is 6×10^{-4} for NO₂, O₄ and HCHO, and 1.3×10^{-3} for SO₂. The detection limit of the dSCDs (assumed as two times of the mean RMS) is 3×10^{15} molecules cm⁻² for NO₂, 5×10^{41}
- 25 molecules² cm⁻⁵ for O₄, 5×10^{15} molecules cm⁻² for SO₂, 5×10^{15} molecules cm⁻² for HCHO. Only 0.7%, 0.4%, 3.3%, 6.6% of the filtered measurements have results below the respective detection limits for $NO₂, O₄, SO₂$ and HCHO, respectively.

3 PriAM inversion algorithm

The profile inversion is based on the fact that the vertical distribution of the light paths depends on the elevation angle of the observation. The vertical trace gas profiles are assumed to be constant for the duration of the elevation angle sequence and 30 also in horizontal dimensions. If the light paths are well-known, vertical trace gas profiles can be derived from a set of dSCDs for the different elevation angles. Besides the observation geometry and sun position, scattering on air molecules

(Rayleigh scattering), aerosols and cloud particles (often referred to as Mie scattering) determine the atmospheric light paths. Opposed to the well-known Rayleigh scattering, scattering on aerosols and cloud particles depends on their respective optical properties, which are diverse and depend on a size, shape and composition. Vertical profiles of AEs can be retrieved from a set of O_4 dSCDs for individual elevation sequences using the well-known vertical profile of the O_4 concentration,

- 5 which is proportional to the square of the concentration of molecular oxygen and thus only depends on temperature and pressure (Hönninger et al., 2004; Wagner et al., 2004; Frieß et al., 2006). Like for other algorithms, a two-step inversion procedure is used in the PriAM algorithm: in the first step the aerosol extinction (AE) profiles and in the second step the profiles of the trace gas VMRs are retrieved. In PriAM we applied the Levenberg-Marquardt modified Gauss-Newton procedure (Rodgers, 2000) to solve the ill-posed inversion problem for AEs (Frieß, et al., 2006 and Yilmaz, 2012) through 10 the numerical iteration:
	- $x_{i+1} = x_i + \left((1 + \gamma_i) S_a^{-1} + K_i^T S_{\varepsilon}^{-1} K_i \right)^{-1} (K_i^T S_{\varepsilon}^{-1} (y F(x_i)) S_a^{-1} (x_i x_a))$ (s1)

with x_{i+1} and x_i the solutions of atmospheric state at the i and i+1 step. x_a is the a-priori profile and y the measurement vector. γ_i is the Levenberg-Marquardt factor, which is multiplied or divided by two to make the minimization of the cost function faster and more stable than for the normal Gauss-Newton algorithm. S_a is the covariance of the error of the a-priori

15 profile and S_{ε} is the covariance of the errors of the measurements. K_i and $F(x_i)$, which are calculated for each iteration step, are the weighting function and the forward model value at the state of x_i , respectively.

The set of O_4 dSCDs for the m non-zenith elevation angles in each scan (in this study 5° , 10° , 20° and 30°) is the measurement vector to retrieve the AE (σ) in n atmospheric layers. In this study 20 atmospheric layers from the surface to 4 km with height intervals of 0.2 km are used (the same layers are used for the retrievals of the trace gas profiles). Considering

20 the frequent variation of aerosols, very little is known about the expected AE profile. Thus a fixed smoothed box-shaped apriori AE profile (Boltzmann distribution) is used, as introduced by Yilmaz (2012):

$$
\sigma(z) = \frac{\sigma(0)}{z - \frac{\tau}{\sigma(0)}}\tag{S2}
$$

Here $\sigma(z)$ and $\sigma(0)$ denote the extinction coefficient at the altitude z (km) and at the surface, respectively. τ is the optical depth. In this study, $\sigma(0)$ and τ are 0.15 km⁻¹ and 0.3, respectively. The covariance matrix S_a is constructed as follows:

$$
25 \quad S_{a_{ik}} = \sigma_{a_i}^2 \times e^{-\frac{|z_i - z_k|}{\eta}}
$$
\n
$$
\tag{S3}
$$

With σ_{a_i} the a-priori AE at the atmospheric layer i. z_i and z_k are the heights of the atmospheric layer i and k, respectively. The smoothing factor η is 0.5 km. The covariance matrix of the measurement uncertainties S_{ε} contains diagonal elements representing the square of the fitting errors of the $O₄$ dSCDs and off-diagonal elements of zero.

In most previous studies, the optimal linear inverse method (Rodgers, 2000; Frieß, 2011) is used to retrieve the vertical 30 profiles of the trace gas VMRs. In PriAM, we use the Gauss-Newton numerical procedure as in Eq. (s1) because the use of the safe state of AEs and trace gas VMRs (see below) converts the linear problem into a nonlinear one.

Similar to the retrieval of the AE profiles, the diagonal element of S_{ε} is the square of the fitting errors of the respective trace gas dSCDs and the off-diagonal elements are zero. The elements of S_a are calculated from Eq. (s3) but the σ_{a_i} are replaced by the a-priori VMR of the respective trace gas (ρ_{a_i}) . One fixed a-priori profile of the VMRs for each trace gas is used. The a-priori profiles of $NO₂$ and $SO₂$ are described as an exponential function (similar to Yilmaz, 2012 and Hendrick et al 2014):

$$
5 \quad \rho(z) = \rho(0) \times e^{-\frac{z}{H}}
$$
\n
$$
(s4)
$$

Here $\rho(z)$ and $\rho(0)$ are the VMR of the trace gases at altitude z (km) and near the surface, respectively. H is the scaling height (in this study fixed to 1km). The ground VMR $p(0)$ is set to 4 ppb for NO₂ and 8 ppb for SO₂.

MAX-DOAS and aircraft measurements in Milano during summer of 2003 indicated that the layer of high HCHO concentration often extends to 1 km or even higher altitudes (Wagner et al., 2011, Junkermann, 2009). Thus for HCHO the 10 same a-priori profile (Boltzmann distribution) as for the AE is used. The surface mixing ratio ρ(0) is set to 4ppb and the VCD to 1.7×10^{16} molecules/cm².

During the profile inversion for aerosols and trace gases, negative values can occur, which are physically invalid. To avoid them, the original atmospheric state vector x is transformed to its corresponding 'safe state' *^x* (Yilmaz, 2012):

$$
x' = \ln(x) \tag{s5}
$$

15 After finishing the calculation of x' , x' is transformed back to the original format

$$
x = e^{x'} \tag{86}
$$

In this way it is ensured that x is always positive.

The averaging kernel (AK) is an important quantity to characterize the vertical resolution of the measurement and the sensitivity of the retrieved state \hat{x} to the true state as a function of altitude. The trace of the AK matrix yields the degree of

20 freedom (DoF) of the signal, which represents the number of independent pieces of information that can be retrieved. The error of the retrieved state S consists of the smoothing error S_s (due to the limited vertical resolution of the retrieval) and the retrieval noise S_m (due to measurement errors).

3.1 Influence of the choice of the a-priori profiles on the retrieved profiles

- We investigate the impact of the choice of the a-priori profiles on the retrieved profiles, VCD (AOD) and near-surface VMR 25 (AE) (from the ground to 200 meters) for two months (July 2011 and February 2012) by either varying the VCD (AOD) by 0.5 or 2, or changing the profile shape by replacing the Boltzmann distribution with the exponential distribution (for aerosols and HCHO) or the other way around (for $NO₂$ and $SO₂$) (see Fig. S8). We compared the respective differences of the measured dSCDs and modeled dSCDs (results of the forward model) and the retrieved profiles (see Fig. S9), VCDs (AODs) and the near-surface VMRs (AEs) (see Table S1). We found a stronger influence of the a-priori profile for aerosols than for
- 30 the trace gases. By changing the a-priori profiles, the maximum change of the retrieved VCDs and AODs is on average about 10% and 20%, respectively. The retrieved near-surface VMRs and AEs change by around 2% and 10%, respectively. For

both aerosols and trace gases, typically the smallest differences between the measured and modelled dSCDs are found for the standard a-priori profiles (see Fig. S9) indicating that the standard a-priori profiles are the preferred assumptions.

3.2 Evaluation of the internal consistency of the inversion algorithm

The retrieval quality is evaluated for favourable measurement conditions, namely cloud-free sky with relatively low aerosols

5 (the sky condition is directly identified by MAX-DDOAS observations as described in section 2.2.5 in the main manuscript), and the performance of the retrievals in different seasons is discussed.

Comparing the measured TG dSCDs to the modelled dSCDs (the results of the forward model corresponding to the retrieved AE and TG profiles) is a direct way to evaluate how close to the real profile the retrieved profile is. Ideally, the differences between measured and modelled dSCDs are minimized by the inversion. However because of measurement errors,

- 10 deviations of the forward model from reality (e.g. for cloudy skies, shown in section 2.2.5 in the main manuscript) and the not always realistic assumption of the Gauss-Newton Algorithm in Eq. (1) in the main manuscript (especially under the condition with strong aerosol load, also shown in section 2.2.5 in the main manuscript), the derived profiles might strongly deviate from the real profiles. The mean differences (and standard deviations denoted by error bars) between the measured and modelled dSCDs for the four species are plotted against the elevation angles during the whole measurement period in
- 15 Fig. S10. For the aerosol retrieval, a larger negative difference of the O_4 dSCD of 2.9×10^{41} molecules² cm⁻⁵ is found for 5°elevation angle, indicating an underestimation of the aerosol extinction in the layer close to the surface; however the magnitude of the underestimation is only about 2% based on the mean O_4 dSCD of about 1.6×10^{43} molecules² cm⁻⁵ for 5°elevation angle. For the TG retrievals, in general the differences for high elevation angles are slightly larger than those for low elevation angles. This finding probably indicates the higher sensitivity of the inversion algorithm to lower altitudes. This
- 20 is also indicated by the AKs in Figs. S12c, S13c and S14c. Even so, the mean deviations of the dSCDs for the 30° elevation angle are only -0.28 $\times 10^{15}$ molecules cm⁻² for NO₂ (mean dSCD of 2.6×10¹⁶ molecules cm⁻²), -0.07× 10¹⁵ molecules cm⁻² for SO₂ (mean dSCD of 3.3×10^{16} molecules cm⁻²) and 0.65×10^{15} molecules cm⁻² for HCHO (mean dSCD of 1.6×10^{16} molecules cm^{-2}).
- The mean AKs for retrievals of AE, $NO₂$, $SO₂$ and HCHO are shown in Figs. S11c S14c, respectively. They indicate that 25 the inversions are sensitive to the layers from the surface up to 1.5 km. The degrees of freedom (DoF) are about 1.5 for aerosols (similar to Frieß et al., 2006), 2 for NO₂ and 2.3 for SO₂ and HCHO. The total AKs in different seasons are shown in Figs. S11d - S14d for the four species, respectively. The generally similar total AKs in different seasons indicate the consistent response of the measurements to the true atmospheric state. The slight seasonality is probably related to the variation of the SZA. The same reason probably causes the weak diurnal variation of the DoF of the inversions as shown in
- 30 Fig. S15. The averaged profiles retrieved from the measurements during the whole period and in different seasons are shown in Figs. S11a - S14a for the four species together with the corresponding a-priori profiles. The retrieved profiles below 1.5 km are quite different from the a-priori profiles, indicating that the measurements contain sufficient information for the altitude below 1.5 km. The mean contributions of the noise and the smoothing error (this error originates from the limited

resolution of the inversion) of the retrievals are shown in Fig. S13b - S16b. The total (absolute) retrieval errors have a maximum around 1 km and decrease towards the surface. The relative errors are minimal close to the surface (10% for AE, NO₂ and SO₂, and 30% for HCHO). Most of the errors originate from the smoothing error, which largely contributes to the total error at high altitudes.

5 **3.3 Deriving O⁴ VCDs from the measured surface temperature and pressure**

To derive O_4 VCDs from the measured surface temperature and pressure (TP), we first fit $6th$ order polynomials to the seasonal variations of surface TP (see Fig. 3 in the main manuscript). Second, we calculate height profiles of the temperature assuming a lapse rate of 0.645K / 100m:

$$
T_{i+1} = \begin{cases} T_i - 0.645 \times \frac{z_{i+1} - z_i}{0.1} & (0 < z < 12km) \\ T(z = 12km) & (z > 12km) \end{cases}
$$
 (s7)

¹⁰ Here T_i (in unit of kelvin, K) and z_i indicate the temperature and height of the atmospheric layer i, respectively. Above 12 km the temperature is kept constant to represent the temperature inversion around the tropopause. Here it should be noted that this simplification has only negligible influence on the derived O_4 VCD, because most of the O_4 is present at lower altitudes. For the same reason, the TP profiles are only calculated up to an altitude of 20 km.

Based on the calculated temperature profile and the surface pressure we calculate the corresponding pressure profile:

15
$$
D_i = 28.9 \times 10^{-3} \times \frac{P_i}{(T_i \times R)}
$$
 (s8)
 $P_{i+1} = P_i - (g \times 100 \times D_i)$ (s9)

Here D (in units of kg/m³) and P (in unit of hPa) indicate the air density and pressure, respectively. R is the ideal gas constant (8.31 J/mol·K), and g is the gravitational constant (9.8 N/kg). Because the O_4 concentration is proportional to the square of the oxygen concentration (which represents 21% of the air density), the profile and VCD of O_4 can be calculated

20 from the assumed air density profile. Fig. 3 in the main manuscript shows the seasonal variation of the $O₄ VCD$ calculated from the measured surface TP in 2012. The O₄ VCD in summer is systematically lower than in winter (by about 2×10^{42} molecules² cm⁻⁵, which is about 15% of the annually mean O_4 VCD).

4 Dependencies of the errors of the VCD derived by the geometric approximation and the profile inversion on the aerosol load

25 In Fig S25, $Diff_{total}$, $Diff_{inversion}$ and $Diff_{geometry}$ for the different TGs are plotted against the AODs for elevation angles of 20° and 30° (for a range of the SAA between 100° to 110). We found linear relations of $Diff_{geometry}$ against AOD for the three species, especially for 20° elevation angle. The weaker dependence of $Diff_{geometry}$ on AODs for an elevation angle of 30° is due to the lower sensitivity of MAX-DOAS observations on aerosols than for an elevation angle of 20°. Correlation coefficients of the linear regressions of $Diff_{geometry}$ and AODs are largest for HCHO due to its higher layer height compared to the other species. The same reason causes the better correlation for SO_2 than for NO_2 . For relatively large AODs, the relation of $Diff_{total}$ and AOD follows a linear relation of $Diff_{geometry}$ and AODs, but for low AODs, $Diff_{\text{inversion}}$ contributes most to $Diff_{\text{total}}$. $Diff_{\text{inversion}}$ is mostly between $\pm 20\%$ and is randomly distributed around zero. Thus $Diff_{\text{inversion}}$ can not be the reason for the systematic bias between the VCD_{geo} and the VCD_{pro} .

Figures

5

Figure S1: Seasonally mean diurnal variations (2011 to 2014) of ambient temperature (a), wind speed (b) and relative humidity (c) obtained from the observations of the weather station nearby the MAX-DOAS instrument.

5 **Figure S2: Wind rose diagrams based on all hourly averaged observations of the weather station for winter (a), spring (b), summer (c) and autumn (d) from 2011 to 2014.**

Figure S3: SZA dependence of the NO² dSCDs (a), O⁴ dSCDs (b) and relative intensity offset (c) derived from the NO² DOAS fits for all measured spectra during the whole observation period. The blue vertical lines indicate a SZA of 75°. The colours indicate the solar azimuth angle (SAA) with north as zero. Small (large) SAA indicate measurements in the morning (evening).

Figure S4: SZA dependence of the SO² dSCDs (a), O³ dSCDs (b), relative intensity offset (c) and Ring optical depth (d) derived from the SO² DOAS fits for all measured spectra during the whole observation period. The blue lines indicate a SZA of 75°. The colours indicate the solar azimuth angle (SAA) with north as zero. Small (large) SAA indicate measurements in the morning 10 **(evening).**

15 **Figure S5: Examples of DOAS fits of HCHO in the wavelength ranges of 310 to 359 nm (a), 324.6 to 359 nm (b) and 336.5 to 359**

Figure S6: HCHO dSCDs derived from the DOAS fits in the three wavelength ranges on 10 January (a) and 7 August 2014 (b) as well as the fit errors in the wavelength ranges of 324.6-359 nm and 336.5-359 nm on the both days in (c) and (d).

Figure S7: SZA dependence of the HCHO dSCDs (a), relative intensity offset (b), O³ dSCDs (c), and Ring optical depths (d) 5 **derived from the HCHO DOAS fits of all measured spectra during the whole observation period. The blue lines flag the position of SZA of 75°. The blue lines indicate a SZA of 75°. The colours indicate the solar azimuth angle (SAA) with north as zero. Small (large) SAA indicate measurements in the morning (evening).**

10 **Figure S8: Four a-priori profiles for aerosol extinction (a), NO² VMRs (b), SO² VMRs (c) and HCHO VMRs (d) used for the sensitivity tests of the MAX-DOAS profile retrievals. The baseline a-priori profile is used for the standard retrieval of the whole measurements.**

 (g) (h)

Figure S9: Left: elevation angle dependencies of the differences between measured and modelled dSCDs of O4, NO2, SO2 and HCHO for the different a-priori profiles shown in Fig. S8. Right: the average profiles of AE, NO² VMR, SO² VMR and HCHO 5 **VMR derived for the different a-priori profiles. The results are obtained for measurements in July 2011 and February 2012.**

Figure S10: Mean differences and the standard deviations (error bars) between the measured and modelled dSCDs of O⁴ (a), NO² (b), SO² (c) and HCHO (d) for clear sky conditions with low aerosols plotted against the elevation angles of MAX-DOAS measurements.

Figure S11: (a) Mean AE profiles for the different seasons derived from observations under clear sky conditions with low aerosols. Also shown is the a-priori profile; (b) total, noise and smoothing errors of the averaged aerosol extinction profile; (c) the corresponding mean averaging kernels for the different height layers (DoF is the degree of freedom); (d) the total averaging kernel 5 **for the different seasons.**

Figure S12: Same as Fig. S11, but for the NO² retrieval.

Figure S13: Same as Fig. S11, but for the SO² retrieval.

Figure S14: Same as Fig. S11, but for the HCHO retrieval.

Figure S15: Seasonally averaged diurnal variations of the degrees of freedom (DoF) of the inversions for aerosols (a), NO² (b), SO² (c) and HCHO (d). For all four species, the DoF in the morning and afternoon are smaller than around noon mainly due to lower scattering probability in the boundary layer.

Figure S16: Time series of near-surface temperature and pressure obtained from the ground-based weather station near the MAX-DOAS instrument from 2011 to 2014.

Figure S17: Dependence of the difference of the AOD (left) and AE (right) for January 2011 retrieved using TP profiles for either winter or summer versus the AOD and AE (retrieved using TP profiles for winter). TP profiles for winter are calculated assuming 10 **a surface pressure of 1020 hPa and surface temperature of 280 K; those for summer are calculated assuming a surface pressure of 993 hPa and surface temperature of 299 K. The retrievals based on the wrong (summer) TP profiles underestimate the AODs and near-surface AEs by about 20% and 27%, respectively.**

15

5

 Figure S18: Histograms of the differences between MAX-DOAS results and independent techniques for different seasons for measurements under clear sky conditions with low aerosols. (a) difference of the MAX-DOAS AOD compared with the Taihu AERONET level 1.5 data, (b) difference of the MAX-DOAS AE compared with the results from the visibility meter; (c, d) VMRs of NO² and SO² derived from MAX-DOAS compared with the results of the nearby long path DOAS instrument. The mean differences, standard deviations and total numbers of observations are given in brackets for each season.

Figure S19: Scatter plots of the near-surface NO² VMR derived from MAX-DOAS versus those from LP-DOAS in summer (a) and winter (b) for clear sky conditions with low aerosols. The left and right subfigures show results for morning (before 12:00 local time (LT)) and afternoon (after 12:00 LT), respectively. Results of the linear regression are shown in the individual subfigures.

(b)

(d)

Figure S20: Mean results of all MAX-DOAS retrievals under different sky conditions. Besides the AOD and AE (a) and the trace gas mixing ratios and VCDs (b to d), also the cost functions and degrees of freedom are shown. The cost functions of all species are higher under cloudy conditions compared to clear sky conditions. The effect of clouds is stronger for aerosols compared to TGs. This is consistent with the larger discrepancy between modelled dSCDs and the measured dSCDs shown in Fig. 8 of the main 10 **manuscript. The reason is that clouds are not included in the forward model. The DoF of the inversions strongly depend on the cloud and aerosol load. A large aerosol load leads to an increase of the DoF of the aerosol inversion, but to a decrease of the DoF for the TG inversions. The column densities and near surface TG mixing ratios and AE are found to be quite different for the different sky conditions, probably due to cloud effects on the inversions and the different atmospheric chemistry conditions (photolysis) and dynamics for different cloud conditions.**

Figure S21: Mean absolute differences, standard deviations as well as correlation coefficients (R), slopes and intercepts derived from linear regressions of the AODs from MAX-DOAS and the Taihu AERONET level 1.5 data sets for different seasons (different colours) and sky conditions. Numbers of data in each comparison are shown in the bottom panel.

Figure S22: Same as Fig. S21 but for the near-surface AEs compared with the results from visibility meters.

(a)

Figure S25: Diff_{total}, Diff_{inversion} and Diff_{geometry} derived for MAX-DOAS observations under clear sky conditions at SAA **between 100° and 110° for NO² (a), SO² (c), HCHO (e) for elevation angle of 20°; the corresponding results for an elevation angle of 30° are shown in sub-figures (b), (d), (f).**

Figure S26: SO² VCDs (a) and VMRs near surface (b) plotted against NO² VCDs and VMRs, respectively. The linear regressions are shown in each subfigure.

(c)

 Figure S27: Monthly mean profiles of NO² (a), SO² (b), HCHO (c) VMRs under clear sky conditions for May 2011 to November 2014.

Figure S28: Wind roses for March 2012 (a), 2013 (b) and 2014(c).

Figure S29: Average profiles of aerosol extinction (a), NO² VMR (b), SO² VMR (c) and HCHO VMR (d) in the morning and afternoon for winter and summer, respectively.

Figure S30: Diurnal variations of VCDs (left) and surface VMR (right) of NO² (a), SO² (b) and HCHO (c) for different days of the week, averaged over the period of May 2011 to November 2014.

Table S1 Absolute and relative differences of the retrieved VCDs (and AOD) and near-surface VMRs (AEs) of NO₂, SO₂ and HCHO between either using the three test a-priori profiles or the standard a-priori profile (shown in Fig. S6)

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