Effect of Atmospheric Interfering Absorption on Measurement of BTX by DOAS

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It was reported on the elimination of interfering absorption of BTX, the absorption of \( \text{O}_2 \) includes different absorption bands, which change differently when the partial pressure of oxygen is varied. The cause of the nonlinear absorption of \( \text{O}_2 \) and the observed band shape to vary with the column density of \( \text{O}_2 \). The absorption ratio of molecular absorption in each of the Herzberg bands and dimer absorptions, as well as the contribution to the correction error of molecular absorption, are studied based on the characteristic of these absorption bands. The optimized way to eliminate the interfering absorption is obtained in the end and the effectiveness of using interpolation proposed by Volkamer et al. to remove \( \text{O}_2 \) absorption is proved again.

II. THEORIES AND INSTRUMENTS

The analysis of DOAS spectra is based on the Lambert-Beer’s law [14]:

\[ D = \ln \frac{I_0(\lambda)}{I(\lambda)} = L \sum_{i=1}^{n} \sigma_i(\lambda)C_i + \mu_R(\lambda) + \mu_M(\lambda) \]  

A differential optical density \( D \) is defined in Eq.(1), \( \lambda \) is wavelength, \( I_0(\lambda) \) is intensity of the light source, \( I(\lambda) \) is intensity at the detector.
is light intensity after pass through the atmosphere, \( \mu_R(\lambda) \) is Rayleigh-scattering coefficient, \( \mu_M(\lambda) \) is Mie-scattering coefficient, \( \sigma_i(\lambda) \) is differential absorption cross section of absorber \( i \), \( C_i \) is the concentration of absorber \( i \), \( L \) is total light path length, \( n \) is the number of absorbers. Numerical filters and special algorithms remove broadband variation of the cross section as well as the wavelength dependency of the measured spectra due to Rayleigh- and Mie-scattering and the lamp. The species concentration can then be calculated according to Eq.(2):

\[
C_i = \frac{D_i(\lambda)}{\sigma_i(\lambda)L}
\]

In theory, after removing the absorptions of different species, the residual is constituted mainly by short noise. Usually, the minimum detectable optical density is indicated as the difference between the maximum and minimum values in the residual spectrum.

DOAS system: a Cassegrain telescope with a high pressure Xenon short-arc lamp (150 W, Osram, Germany) as the light source; thirteen retro reflectors, a spectrometer (Acton500, focal length is 500 mm, grating are 1800, 1200, and 600 L/mm blazed at 300, 300, and 500 nm, respectively), a fiber bundle, 1024 pixels photo diode array (PDA) (Hoffman Messtechnik GmbH, Germany) kept at \(-30^\circ C\). The data produced by the detector was digitized by a 16-bit analog-to-digital converter (ADC) and then transmitted to the Hoffman controller. After being processed, the signals were transferred to the computer by RS232. The measurement of three different types of spectra is required: the atmospheric spectra, a lamp spectra and spectra of the background light. The DOAS spectra evaluation algorithm is well described in Ref.[14].

III. ANALYSIS OF INTERFERING ABSORPTION

A. Interfering absorption of \( \text{O}_2 \)

In the UV spectral region below 300 nm \( \text{O}_2 \) molecules show weak absorption due to electronic excitation in the Herzberg I \((A^3\Sigma_u^-\rightarrow X^3\Sigma_g^-)\), Herzberg II \((A^1\Sigma_u^-\rightarrow X^3\Sigma_g^-)\), and Herzberg III \((A^3\Delta_u\rightarrow X^3\Sigma_g^-)\) transitions (shown in Fig.1). At the same time, Herzberg III \((A^3\Delta_u\rightarrow X^3\Sigma_g^-)\) give rise to additional diffuse absorption bands due to \( \text{O}_2 \)-dimers [15]. All four band-systems strongly overlap, extending from the joint dissociation limit of \( \text{O}_2 \) at 242.9 nm towards longer wavelengths with the absorption becoming negligible above 287 nm. Regarding the transparency of the ground level atmosphere in the UV (below 300 nm), Ditchburn and Young concluded that attenuation by \( \text{O}_2 \) in the Herzberg systems was comparable to extinction by tropospheric ozone and consequently has to be taken into account in transparency calculations [16]. \( \text{O}_2 \) is the most important absorber in the UV, and its absorption is constituted mainly by Herzberg band-systems and \( \text{O}_2 \)-dimers’ absorption-bands. In 1915, Warburg first mentioned a quadratic pressure dependence of the absorption of gaseous oxygen [18] and Wulf first ascribed this behavior to the formation of oxygen-dimers [19]. At a pressure of about 400 kPa of pure \( \text{O}_2 \), dimers’ absorption begins to dominate \( \text{O}_2 \)-absorption [17]. The important dimers in the atmosphere are \( \text{O}_2-\text{O}_2 \) and \( \text{O}_2-\text{N}_2 \), overlapped with Herzberg band-systems. In any case, dimer-formation forces the cross section to increase linearly with pressure, which for dimer-absorption in the Herzberg band’s wavelength range was observed in pure oxygen gas [20,21] as well as in mixtures with other gases, e.g. nitrogen, argon [17,18]. Moreover, the \( \text{O}_2-\text{N}_2 \) complex is only half as strong an absorber as the \( \text{O}_2-\text{O}_2 \) complex, leading to a predominant fraction (66%) of the dimer-absorption to be due to \( \text{O}_2-\text{N}_2 \) complexes under atmospheric conditions [17].

The interfering absorption of \( \text{O}_2 \) poses several problems for the measurements of BTX by DOAS, which were well researched by Volkamer et al. [5]. (i) The Herzberg bands consist of a series of narrow ro-vibronic lines with a width of the order of 1 pm. In order to fully resolve these lines a spectral resolution well below 1 pm (\( \Gamma \): indicated as full width at half maximum, FWHM) would be required. Spectral resolution used in DOAS system is usually much larger than 1 pm, so the individual lines can not be separated and the apparent optical density (OD) will not increase linearly with the \( \text{O}_2 \) concentration (or \( L \)). As a consequence, the shape of the spectral features will depend on the \( \text{O}_2 \) column density. These effects of unresolved bands generally limit the linear range of DOAS measurements [17], but are of special importance for the correction of...
the Herzberg bands since at atmospheric path lengths of a few hundred meters, the stronger lines of the Herzberg I bands \( (σ≤1.6×10^{-22} \text{ cm}^2) \) become saturated. \( σ \) of Herzberg absorption-bands depends on \( λ \), and great difference exists in the intensity of absorption lines, so the nonlinearity effect of \( O_2 \) absorption spectrum is different from this wavelength range to the other. A single reference spectrum taken at a different column density then cannot be scaled linearly to eliminate both “weak” and “strong” absorptions in a measured spectrum at the same time. As a consequence, residual structures remain.

(ii) When the \( O_2 \) partial pressure is changed, the different absorption-bands of \( O_2 \) scale differently. Namely the Herzberg bands scale in a way different from that of dimer-absorptions and even three Herzberg bands and the absorption of two dimers \( (O_2-O_2 \text{ and } O_2-N_2) \) scale in a way different from each other. In addition, all these absorption-bands overlap with each other and can not be separated, so it is impossible to correct these absorption-bands in a measured spectrum separately. The relative strength absorption of \( O_2-O_2 \text{ and } O_2-N_2 \) scales differently from the \( O_2 \) partial pressure. Even though this effect can reduce the change of the ratio of molecular absorption in the Herzberg bands and dimer-absorptions, it needs to be accounted for, if the DOAS is used to measure BTX in the atmosphere.

(iii) The absorptions of \( O_2 \) and BTX overlap with each other, and the absorption features of \( O_2 \) may influence the apparent absorption features of BTX at long path lengths if spectral features are measured at “low” resolution. Hence, further residual structures may result from an incomplete correction of the absorption features of the atmospheric absorber [21].

B. Interfering absorptions of \( O_3 \) and \( SO_2 \)

The absorptions of BTX overlap with those of \( O_3 \) and \( SO_2 \) besides \( O_2 \) (shown in Fig.2 and Fig.3). Though the intensities of two gases’ differential cross sections are less than that of most BTX, their concentrations are ten times or even one hundred times higher than BTX, and their differential absorption ODs in the atmospheric spectra are much higher than BTX. Especially in China, \( SO_2 \) concentration is usually high in most places. As a consequence, the slight elimination error of \( O_3 \) and \( SO_2 \), which is negligible for the quantification of \( O_3 \) and \( SO_2 \) by DOAS, will introduce additional features to the residual and interfere the retrieval of BTX concentrations.

IV. EFFECT OF ABSORPTION ON MEASUREMENT RESULTS

A. Interfering evaluation of \( O_2 \) absorption

To characterize the magnitude of the \( O_2 \) interference with DOAS measurements of BTX, the signal of benzene in a measured spectrum can be compared to that of the nearby \( O_2 \) band at 259 nm. When \( L=682 \text{ m} \) and \( Γ=0.15 \text{ nm} \), \( O_2 \) bands in atmospheric spectra have an OD of \( \sim 0.06 \). Assuming a mixing ratio for benzene of \( 1.5×10^{-9} \) (a typical level in the urban air), benzene will have an OD of \( 3.3×10^{-3} \), 5.5\% of \( O_2 \) absorption. That means that the \( O_2 \) absorption will dominate the measured spectrum and even when the elimination error of \( O_2 \) absorption is kept as low as 1\%, it will be difficult to accurately detect benzene for most of time in urban sites. Fortunately, most BTX show their strongest absorption bands towards longer wavelengths, where the \( O_2 \) absorption is weaker (Fig.4) [5], but still the \( O_2 \) absorption bands are likely to be stronger than the aromatic absorption, since most aromatics are present at lower concentrations than benzene.
B. Interfering evaluation of O$_3$ and SO$_2$ absorption

The elimination error of O$_3$ and SO$_2$ absorption mainly originates from the temperature effect of absorption cross section. However the atmospheric temperature changes inevitably and periodically. Taking a mixing ratio of 6×10$^{-8}$ (a typical level in the urban air) as an example, the differential absorption spectra of O$_3$ at different temperatures are shown in Fig.2 ($L=682$ m, $\Gamma=0.15$ nm) and OD increases with the decrease of temperature. In order to study if the changes of O$_3$ differential absorption spectra at different wavelength range are synchronous, the differences of relative intensity are calculated at different wavelength, as shown in Fig.5. With the increase of temperature, the change magnitude of differential spectra depends on $\lambda$. If a single reference spectrum taken at a different temperature cannot be scaled linearly to eliminate both, “weak” and “strong” absorptions in a measured spectrum at the same time. As shown in Fig.2, the differential absorption OD of O$_3$ (280 K) at 259 nm is $\sim$4.3%, thirteen times higher than that of benzene detected at the same time. When a reference spectrum taken at 273 or 295 K is used to fit this spectrum, 0.8% and 1.0% residual noise will remain respectively, both of which are much larger than the differential absorption OD of benzene (1.5×10$^{-9}$). Though O$_3$ absorption is weaker towards longer wavelengths, it is still likely to be stronger than the BTX absorption.

In the same way, as shown in Fig.3 and Fig.5, the absorption cross section of SO$_2$ also suffers from temperature effect and the change magnitude of differential spectra depends on $\lambda$ with the increase of temperature. SO$_2$ is usually higher in China and taking a mixing ratio of 10$^{-7}$ (a typical level in the urban air) as an example, the differential absorption OD of SO$_2$ (280 K) at $\sim$270 nm is $\sim$3.0% (273 K, $L=682$ m, $\Gamma=0.15$ nm), much higher than that of BTX in this wavelength range. As shown in Fig.3 [5], the difference between the SO$_2$ differential spectra at different temperatures can lead to 0.8% and 1.0% additional residual respectively, which are much larger than the differential absorption ODs. The SO$_2$ absorption in the detected atmospheric spectra is usually much higher than BTX, so the slight change of absorption feature will make the retrieval of BTX difficult.

V. ELIMINATION OF THE INTERFERENCES

The magnitude of the O$_2$, SO$_2$, and O$_3$ interference with DOAS measurements of BTX is all determined by their elimination error, so the accurate elimination of interfering absorptions is critical to DOAS measurements of BTX.

A. Elimination of O$_2$ interference

Axelsson et al. and Trost used a “zero” reference spectrum taken in “clean” air to correct O$_2$ absorption [26,27]. However, this method will introduce an unknown negative offset for all species present in the “zero” spectrum, as this spectrum is subtracted from each measured spectrum. Volkamer et al. proposed a
solution to this problem—using interpolation to remove $O_2$ absorption and proved it [5]. Here we used more details to prove the effectiveness of this method.

\[ O_2 + O_2(N_2) \rightleftharpoons O_2-O_2(N_2) \]

As said above, absorption features of $O_2-O_2$ and $O_2-N_2$ overlapped with Herzberg bands should be separated first, in order to remove $O_2$ absorption from the measured spectrum. As shown in Eq.(3), the formation of $O_2-O_2$ depends on the pressure, which causes a quadratic pressure dependence of the absorption of gaseous oxygen and a linear pressure dependence of the absorption of dimer [21,28]. With regard to the dimer’s nature, the classification of oxygen dimers under atmospheric conditions as mixtures of metastable Van der Waals dimers, metastable form and colliding pairs are most possible. Only bound Van der Waals dimers should exhibit structured absorption features, metastable form and colliding pairs would result in broadened absorptions [29].

By adjusting a spectrum taken in different oxygen/nitrogen ratios, band positions and bandwidths of the dimer absorption below 267 nm could be determined. These spectra cannot be used as dimer reference spectra since residual structures due to the unresolved molecular absorption (Fig.6) superimpose with the dimer absorption. Figure 6(a) shows the absorption cross section of $O_2$ at different pressures and temperatures [30]. It is clear that the band positions and bandwidths of the dimer absorption depend on pressures and temperatures, and the bandwidths of the dimer absorption are much wider than the BTX absorption. As shown in Fig.6(b), by the application of an optimized Savitzky Golay high pass filter [31,32], the wide band absorption can be almost eliminated. Seen from filtered spectra $A'$, $B'$, $C'$, and $D'$, the resulting spectra of $A$, $B$, $C$, and $D$ respectively after being high filtered.

After the elimination of the absorption of $O_2$ dimers, the overlapped Herzberg bands need to be separated in order to eliminate the oxygen absorption present in the atmospheric spectra. As shown in Table I, the absolute oscillator strengths ($f$) for these three systems are $8.49\times10^{-10}$, $0.44\times10^{-10}$, and $0.51\times10^{-10}$ [23-25] and the Herzberg I transition be more than one order of magnitude stronger than the Herzberg II and Herzberg III systems, it also can be seen from Fig.1 that the strong lines of the Herzberg I band dominate the spectrum and the absorption features of the Herzberg II and Herzberg III systems can be negligible. Therefore, the absorption features of Herzberg I determine the nonlinearity absorption features of Herzberg bands systems. In addition, as seen from the line intensity of Herzberg I transition.
TABLE I. Spectroscopic properties of the Herzberg transitions

<table>
<thead>
<tr>
<th>System</th>
<th>Spectroscopic notation</th>
<th>Selection rule violated</th>
<th>$f$</th>
<th>Relative strength</th>
</tr>
</thead>
<tbody>
<tr>
<td>Herzberg I</td>
<td>$A^3\Sigma_u^+ \leftarrow X^3\Sigma_g^-$</td>
<td>$\Sigma^+ \neq \Sigma^-$</td>
<td>$\approx 8.49 \times 10^{-10}$ [22]</td>
<td>19.3</td>
</tr>
<tr>
<td>Herzberg II</td>
<td>$C^1\Sigma_u^+ \leftarrow X^3\Sigma_g^-$</td>
<td>$\Delta \Sigma \neq 0$</td>
<td>$\approx 0.44 \times 10^{-10}$ [23]</td>
<td>1.0</td>
</tr>
<tr>
<td>Herzberg III</td>
<td>$A^3\Delta_u \leftarrow X^3\Sigma_g^-$</td>
<td>$\Delta \Delta = 0, \pm 1$</td>
<td>$\approx 0.51 \times 10^{-10}$ [24]</td>
<td>1.2</td>
</tr>
</tbody>
</table>

FIG. 7 Oxygen reference spectrum of the Herzberg II and III ($\Gamma=0.15$ nm).

II and III, very few lines can become saturated at atmospheric path lengths of a few hundred meters.

As shown in Fig.7, cross sections for Herzberg II and Herzberg III are calculated using the integrated absorption cross sections [24,25]. The resulting cross section spectrum is used to calculate absorption spectra for oxygen at three column densities, which are chosen to differ by a constant value and are equivalent to a ground level path length in the atmosphere of 400, 800, and 1200 m. Subsequently, these spectra are convoluted with a Gaussian-shape instrument function ($\Gamma=0.15$ nm) in order to adjust them to a resolution comparable to atmospheric DOAS measurements. When $L=800$ m, the differential absorption ODs of Herzberg II and III are $3.8 \times 10^{-2}$ (260 nm) and $2.5 \times 10^{-2}$ (258 nm) respectively. The change of $L$ has little effect on the absorption features and when the differential spectra at different column densities are used to fit each other, the residual structure is in a level less than $10^{-6}$. So this slight difference has almost no effect on the absorption feature changes of $O_2$.

After removing the interference of Herzberg II and Herzberg III absorption, the last step is the accurate elimination of nonlinearity absorption of Herzberg I absorption. In the same way as said above, absorption spectra of Herzberg I at path lengths of 400, 800, and 1200 m are calculated ($\Gamma=0.15$ nm), as shown in Fig.8. The individual ro-vibronic transitions are not resolved and appear as Q-branches in the spectrum. The nonlinearity of the increasing absorption features can already be seen in Fig.8, as the vertical spacing between the spectra A and B is larger than that between the spectra B and C. The change in the relative strength of the bands is clearly visible. This becomes more obvious when seen in the residual structures shown in Fig.8(d).

B. Elimination of $SO_2$ and $O_3$ interference

The temperature effect on $O_3$ absorption features is relatively large. In order to characterize the magnitude of temperature effect on the change of absorption features, the absorption cross sections of $O_3$ at different temperatures are used to calculate absorption spectra with $6 \times 10^{-8}$ concentration at a light path of 682 m. Subsequently, these spectra are convoluted with a Gaussian-shape instrument function ($\Gamma=0.15$ nm) and are used to calculate differential absorption spectra. The fitting errors between these spectra are indicated as the remaining noise after using one spectrum to correct for another. Using one differential absorption spectrum at different temperatures separately or simultaneously with the spectrum at 295 K to correct for the differential absorption spectrum $x$ (where was the $O_2$ absorption spectrum at 280 K), the fitting results are shown in Table II. It can be seen that when only one spectrum is used, as the temperature of used spectrum become closer and closer to 280 K, the fitting error becomes less and less. However, it is still at the level of $10^{-3}$; but when two
FIG. 8 The 4-0 and 5-0 vibronic band of the Herzberg I band system of O2 [23]. (a) High resolution absorption cross section. (b) Two part of cross section shown in layer (a). (c) The shown spectra correspond to \( L = 400 \) m (A), 800 m (B), and 1200 m (C) (\( \Gamma = 0.15 \) nm). (d) The remaining residual structures (D, E, and F) after B are fitted by A and C separately and simultaneously.

TABLE II The difference among the differential absorption spectra at different temperatures

<table>
<thead>
<tr>
<th>Temperature/K</th>
<th>228</th>
<th>230</th>
<th>246</th>
<th>273</th>
<th>293</th>
<th>295</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fitting error/0.1%</td>
<td>1</td>
<td>9.25</td>
<td>7.18</td>
<td>5.01</td>
<td>2.45</td>
<td>3.08</td>
</tr>
<tr>
<td>2</td>
<td>0.74</td>
<td>0.67</td>
<td>0.31</td>
<td>0.07</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

1: The fitting error of differential absorption spectra at different temperature to \( x \) (the O\(_3\) absorption spectrum at 280 K).
2: The fitting error of differential absorption spectra at different temperature and differential absorption spectra at 295 K to \( x \) (the O\(_3\) absorption spectrum at 280 K); wavelength range: 255-270 nm.

spectra with different temperatures are used, the fitting error can be reduced to the level of \( 10^{-4} \) to \( 10^{-5} \), much less than the detection limit of benzene. That means that when two spectra with the temperature closer to the detected spectrum are used, the fitting error can be reduced as small as possible.

As shown in Fig.3, the fitting error resulting from the temperature effect of SO\(_2\) can be reduced to the level of \( 10^{-5} \), much less than the detection limit OD of xylene.

VI. MEASUREMENT RESULTS AND CONCLUSION

In July of 2007, long time continuous monitoring of BTX was carried out at Environmental Monitoring Department of Guangzhou with a time resolution of about 3-8 min. A ventilative measurement site with some small emission source distributed equally around was selected. The results are shown in Fig.9 and are compared with the detected results of gas chromatography (GC) (shown in Fig.10). It can be seen from the results that the same trend of BTX diurnal variation can be obtained from these two methods with high correlation. Correlation coefficient for toluene is 0.9209, benzene 0.8864, and \( m&p\)-xylene 0.8118. Only the BTX absolute concentrations of two methods have some difference. Certainly this difference may come from different principles: line measurements are used in DOAS where measurements are performed along an optical path, and point measurements are used in GC where measurements are carried out by taking samples at one spot. The detection limits of BTX are calculated based on the residual noise (\( 10^{-3} \)) remaining after removing

FIG. 9 The diurnal fluctuation of BTX concentration in the atmosphere. a, c and e: The results of DOAS. b, d and f: The results of GC. a and b: Toluene; c and d: Benzene; e and f: \( m&p\)-xylene.
Featuring excellent response characteristics and detection sensitivity and with much lower operational cost, differential optical absorption spectroscopy (DOAS) can be a powerful tool to trace concentration variation of BTX (the main aromatic compounds in the atmosphere). This work focuses on the elimination of interfering absorption of BTX: the absorption of O$_2$ includes different absorption bands, which change differently when the partial pressure of oxygen is varied. Namely the Herzberg bands change in the way different from that of dimer absorption and even three Herzberg bands and the absorption of two dimers (O$_2$-O$_2$ and O$_2$-N$_2$) change in ways different from each other. These cause the nonlinear absorption of O$_2$ and the observed band shape to vary with the column density of O$_2$. The absorption ratios of molecular absorption in each of the Herzberg bands and dimer absorptions, as well as the contribution to the correction error of molecular absorption, were studied based on the characteristic of these absorption bands. The optimized way to eliminate the interfering absorption is obtained in the end and the effectiveness of using interpolation proposed by Volkamer et al. to remove O$_2$ absorption is proved again. As to O$_3$ and SO$_2$, the effect of the thermal effect of characteristic spectra on the elimination error of their absorption is studied. Solutions to these problems are discussed and demonstrated together with methods to optimize the interpolation of spectra. As a sample application, DOAS measurements of BTX were carried out and a low detection limit and good correlation with point instruments was achieved. These evidences prove the feasibility of the methods.

VII. ACKNOWLEDGMENT

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