The influence of instrumental line shape degradation on NDACC gas retrievals: total column and profile

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Abstract:

We simulated Instrumental line shape (ILS) degradations with respect to typical types of misalignment, and compared their influence on each NDACC (Network for Detection of Atmospheric Composition Change) gas. The sensitivities of total column, root mean square of fitting residual (RMS), total random uncertainty, total systematic uncertainty, total uncertainty, degrees of freedom for signal (DOFs), and profile with respect to different levels of ILS degradation for all current standard NDACC gases, i.e., O\textsubscript{3}, HNO\textsubscript{3}, HCl, HF, ClONO\textsubscript{2}, CH\textsubscript{4}, CO, N\textsubscript{2}O, C\textsubscript{2}H\textsubscript{6}, and HCN, were investigated. The influence of an imperfect ILS on NDACC gases retrieval were assessed, and the consistency under different meteorological conditions and solar zenith angles (SZA) were examined. The study concluded that the influence of ILS degradation can be approximated by the linear sum of individual modulation efficiency (ME) amplitude influence and phase error (PE) influence. The PE influence is of secondary importance compared with the ME amplitude. Generally, the stratospheric gases are

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more sensitive to ILS degradation than the tropospheric gases, and the positive ME influence is larger than the negative ME. For a typical ILS degradation (10%), the total columns of stratospheric gases O₃, HNO₃, HCl, HF, and ClONO₂ changed by 1.9%, 0.7%, 4%, 3%, and 23%, respectively. While the columns of tropospheric gases CH₄, CO, N₂O, C₂H₆, and HCN changed by 0.04%, 2.1%, 0.2%, 1.1%, and 0.75%, respectively. In order to suppress the fractional difference in total column for ClONO₂ and other NDACC gases within 10% and 1%, respectively, the maximum positive ME degradations for O₃, HNO₃, HCl, HF, ClONO₂, CO, C₂H₆, and HCN should be less than 6%, 15%, 5%, 5%, 5%, 5%, 9%, and 13%, respectively; the maximum negative ME degradations for O₃, HCl, and HF should be less than 6%, 12%, and 12%, respectively; the influence of ILS degradation on CH₄ and N₂O can be regarded as negligible.

Key words: NDACC, FTIR, Instrumental line shape, Profile retrieval

1 Introduction

In order to achieve consistent results between different FTIR (Fourier transform infrared) sites, the TCCON (Total Carbon Column Observing Network, http://www.tccon.caltech.edu/) and NDACC (Network for Detection of Atmospheric Composition Change, http://www.ndacc.org/) have developed strict data acquisition and retrieval methods to minimize site to site differences (Hase et al., 2012; Wunch et al., 2010 and 2011; Washenfelder, 2006; Messerschmidt et al., 2010; Kurylo, 1991; Davis et al., 2001; Schneider, et al., 2008; Kohlhepp et al., 2011; Hannigan et al., 2009; Vigouroux et al., 2008 and 2015). Interferograms are acquired with similar instruments operated with common detectors, acquisition electronics and/or optical filters. These interferograms are first converted to spectra and then these spectra are analyzed using dedicated processing algorithms, i.e., GFIT, PROFFIT or SFIT (Wunch et al., 2010 and 2015; Hase et al., 2006; Hannigan and Coffey, 2009). Typically, the TCCON network only uses the Bruker 125HR instruments (http://www.tccon.caltech.edu/; https://www.bruker.com/) with specified settings.
In the NDACC network, other instruments are used as well, e.g., the Bruker M series, a BOMEM DA8 in Toronto, Canada and a self-built spectrometer in Pasadena, USA (http://www.ndacc.org/; https://www.bruker.com/). FTIR spectrometers are highly precise and stable devices, and if carefully aligned, the instrumental line shape (ILS) might not be far from the theoretical limit. However, their alignment can change abruptly as a consequence of operator intervention or drift slowly due to mechanical degradation over time (Olsen et al., 2004; Duchatelet et al., 2010; Hase et al., 2012; Feist et al., 2016). Moreover, the NDACC observation may change the entrance field stop size if incident radiation changes. This practice may introduce a dependency of the instrument alignment status on the optical settings because the mechanical errors between different field stops may be non-negligible and inconsistent (Sun et al., 2017). Biases between sites would arise if all these misalignments are not properly characterized.

The TCCON network only operates in near infrared (NIR) region and aims at column of fewer gases. While the NDACC network operates in both NIR and mid-infrared (MIR) regions and aims at both columns and profile of many gases. The TCCON assumes an ideal ILS in spectra retrieval, and the maximum ILS degradation is prescribed as 5% for the modulation efficiency (ME) amplitude (Wunch et al., 2011 and 2015). This assumption still holds within the required accuracy of the results. In the NDACC gases retrieval, the ILS can be assumed as ideal if spectrometer is well aligned, or if misalignment exists, described by LINEFIT results derived from dedicated cell measurements or retrieved together with the gas profile from an atmospheric spectrum using a polynomial (Vigouroux et al., 2008 and Vigouroux et al., 2015). How these ILS treatments influence the NDACC gases retrieval and how much ILS deviation from unity is acceptable for each NDACC gas if an ideal line shape is assumed are still not fully quantified, and it may be better to assume an ideal ILS. The practice of co-retrieving ILS parameters from atmospheric spectra without dedicated cell measurements is not to be recommended because the observed shapes of spectral lines are exploited primarily for inferring the vertical distribution of the
trace gases, the ILS and the trace gas profiles have similar effects on the line shape, i.e., changing the shape and width of the line. Overlapping lines, i.e., due to interfering gases may introduce an asymmetry in the absorption lines which may be undistinguishable from an ILS phase deviation.

This paper investigates the influence of ILS degradation on total column and profile of current standard NDACC gas retrievals and deduces the maximum ILS deviations allowable for suppressing the influence within a specified acceptable ranges.

2 Characteristics of ideal and imperfect ILSs

The ILS is the Fourier transform of the weighting applied to the interferogram. This weighting consists of two parts: an artificially applied part to change the calculated spectrum and an unavoidable part which is due to the fact that the interferogram is finite in length (box car function), the divergence of the beam is non-zero (due to the non-zero entrance aperture), and several other effects which are due to misalignment (Davis et al., 2001, chapter 9). The ILS consisting of only the unavoidable parts of the line shape is called the ideal line shape.

The theoretical ideal ILS as defined in equation (3), when the instrument is well aligned, is a convolution of sinc and rectangular functions (defined in equations (1) and (2)), representing the finite length of the interferogram and the finite circular field of view (FOV) of the spectrometer (Davis et al., 2001).

\[ SINC(\sigma, L) = 2L \frac{\sin(2\pi \sigma L)}{2\pi \sigma L} \]  

\[ RECT(\sigma, \sigma_0, \theta) = \begin{cases} \frac{2}{\sigma_0 \theta^2} & \text{if } -0.5\sigma_0 \theta^2 \leq \sigma \leq 0 \\ 0 & \text{otherwise} \end{cases} \]  

\[ ILS(\sigma, \sigma_0, L, \theta) = SINC(\sigma, L) \ast RECT(\sigma, \sigma_0, \theta) \]  

where \( \sigma \) is the wavenumber, \( \sigma_0 \) is the central wavenumber, \( L \) is the optical path difference (OPD) and \( \theta \) is the angular radius of the circular internal FOV of the spectrometer. For standard NDACC measuring conditions, \( L \geq 180 \) cm and \( \theta \) defined
by the entrance field stop size in the light path.

The LINEFIT software calculates the deviation of the measured ILS from the ideal ILS (Hase et al., 2001 and 2012). It retrieves a complex ME as a function of OPD, which is represented by a ME amplitude and a phase error (PE) (Hase et al., 1999). The ME amplitude is connected to the width of the ILS while the PE quantifies the degree of ILS asymmetry. For a perfectly aligned spectrometer, it would meet the ideal nominal ILS characteristics if smear and vignetting effects were neglected, and thus have an ME amplitude of unity and a PE of zero along the whole interferogram. However, if a FTIR spectrometer is subject to misalignment, the ME amplitude would deviate from unity and the PE deviate from zero (Hase et al., 2012). This results in an imperfect ILS.

3 Simulation of ILS degradation

We use the program ALIGN60 to simulate ILS degradation in a high resolution FTIR spectrometer typically used in the NDACC network. As an auxiliary tool of LINEFIT, ALIGN60 is a raytracing model for FTIR spectrometers following the classical Michelson design, assuming one fixed and one movable arm, and using cube corners instead of plane mirrors. It calculates the resulting phase distortions in the recombined beam and from these deduces the variable intensity observed by the detector. ALIGN60 takes into account the lateral shear error of the movable retro-reflector as function of OPD, a decenter of the field stop with respect to the optical axis, an unsharp boundary line or deformation of the field stop image (as possibly caused by a defocused collimator), and vignetting effects with increasing OPD. It can generate trustworthy results with respect to all types of misalignment (Hase et al., 1999). In this simulation, the entrance beam section was assumed to be circular with a diameter of 8.0 cm. The ILS was only calculated from positive side of interferogram. The smear and vignetting effects were not taken into account. The misalignment of a FTIR spectrometer can be expressed via two perpendicular axes perpendicular to the beam direction. For a circular entrance beam, the same misalignment in either direction results in a similar ILS. Thus, this work only
The misalignments as inputs of ALIGN60 are listed in Table 1, the resulting ILSs are shown in Fig. 1, and the corresponding Haidinger fringes at the maximum OPD are shown in Fig. 2. The ME deviation, decenter of Haidinger fringes and ILS deterioration varying over misalignment are evident. All types of misalignment cause nonlinear ME deviations except decentering of measuring laser (c) and the constant shear (d) which mainly affect PE and result in linear PE deviation. Two types of ILS degradation are evident, one is referred to as positive ME and has a ME amplitude of larger than unity. The other one is referred to as negative ME and has a ME amplitude of less than unity. Typically, the increasing misalignment with increasing OPD (b, f, h or i) causes negative ME amplitude and the decreasing misalignment with increasing OPD (e, g or j) causes positive ME amplitude. For the same misalignment amplitude, the decreasing misalignment causes more ME deviation than the increasing misalignment. Regardless of positive or negative ME, the ME deviation shape depends on misalignment type and the same misalignment amplitude causes the same deviation in ME amplitude. The decentering of the entrance filed stop is equivalent to the linear increasing misalignment.

4 NDACC gases retrieval

4.1 Retrieval strategy

The influence of ILS degradation on all current standard NDACC gases, i.e., O₃, HNO₃, HCl, HF, ClONO₂, CH₄, CO, N₂O, C₂H₆, and HCN, is investigated. Typical atmospheric vertical profiles of these gases are shown in Fig.3. There are five stratospheric gases and five tropospheric gases. The retrieval settings for all these gases as recommended by the NDACC are listed in Table 2 (https://www2.acom.ucar.edu/irwg/links). The latest version of profile retrieval algorithm SFIT4 v 0.9.4.4 is used (http://www.ndacc.org/). The basic principle of SFIT4 is using an optimal estimation technique for fitting calculated-to-observed spectra (Rodgers, 2000; Hannigan and Coffey, 2009). All spectroscopic line parameters are adopted from HITRAN 2008 (Rothman et al., 2009). This might not be ideal, but we keep it to achieve consistent results. A priori profiles of pressure,
temperature and water vapor for the measurement days are interpolated from the National Centers for Environmental Protection and National Center for Atmospheric Research (NCEP/NCAR) reanalysis (Kalnay et al., 1996). A priori profiles of the target gases and the interfering gases except H$_2$O use the WACCM4 (Whole Atmosphere Community Climate Model) model data. We follow the NDACC standard convention with respect to micro windows (MWs) selection and the interfering gases consideration (https://www2.acom.ucar.edu/irwg/links). For the interfering molecules that affect the target gas retrieval, H$_2$O should be treated with care as it is almost always present in all MWs, to varying degrees. It has been dealt with differently for different gas. For HNO$_3$ and ClONO$_2$, H$_2$O is treated as the other interfering species: only a scaling of a single a priori profile is made. For other gases, the H$_2$O profile is retrieved simultaneously with the target gas profile. No de-weighting signal to noise ratios (SNR) are used except for CO and HCl which utilize a de-weighting SNR of 500 and 300, respectively.

The selection of the regularization (a priori covariance matrix $S_a$ and SNR) cannot be easily standardised because it depends on the real variability for each gas. In optimal estimation, the selection of $S_a$ is very important in the inversion process and, together with the measurement noise error covariance matrix $S_\epsilon$, will lead to the following averaging kernel matrix $A$ (Rodgers, 2000):

$$ A = G_y K_x = (K_x^T S_\epsilon^{-1} K_x^T + S_a^{-1})^{-1} K_x^T S_\epsilon^{-1} K_x $$

(4)

where $G_y$ is the sensitivity of the retrieval to the measurement. $K_x$ is weighting function matrix or Jacobian matrix that links the measurement vector $y$ to the state vector $x$ : $\Delta y = K_x \Delta x$. $A$ characterizes the vertical information contained in the FTIR retrievals. In this study, we assume $S_\epsilon$ to be diagonal and its diagonal elements are the inverse square of the SNR. The vertical information content of the retrieved target gas profile can be quantified by the number of degrees of freedom for signal (DOFs), which is the trace of $A$, defined in Rodgers (2000) by:

$$ d_s = tr(A) = tr((K_x^T S_\epsilon^{-1} K_x^T + S_a^{-1})^{-1} K_x^T S_\epsilon^{-1} K_x) $$

(5)

The diagonal elements of $S_a$ represent the assumed variability of the target gas volume mixing ratio (VMR) at a given altitude, and the off diagonal elements represent the correlation between the VMR at different altitudes. We can see in Table 3 that, except CO and HCN, the target gases are using an a priori covariance matrix.
with diagonal elements constant with altitude corresponding to 10, 20, 50 or 100 % variability; the largest variability are for HNO₃, HCl and ClONO₂. For CO, the diagonal elements of Sₘ correspond to 27% from ground to 34 km and decrease down to 11% at the top of atmosphere. For HCN, the diagonal elements of Sₘ correspond to 79% from ground to 5 km and decrease down to 21% at the top of atmosphere. No correlation of off diagonal matrix elements is used in all retrievals except for ClONO₂ which uses exponential correlation with a HWHM (half with at half-maximum) of 8 km. The SNR values for all retrievals are the real values taken from each individual spectrum. The ILSs for all retrievals are using the simulations in section 3.

4.2 Averaging kernels

The rows of A are the so called averaging kernels and they represent the sensitivity of the retrieved profile to the real profile. Their FWHM is a measure of the vertical resolution of the retrieval at a given altitude. The area of averaging kernels represents sensitivity of the retrievals to the measurement. This sensitivity at altitude k is calculated as the sum of the elements of the corresponding averaging kernels, \(\sum A_{k,i}\). It indicates the fraction of the retrieval at each altitude that comes from the measurement rather than from the a priori information (Rodgers, 2000). A value close to zero at a certain altitude indicates that the retrieved profile at that altitude is nearly independent of measurement and is therefore approaching the a priori profile.

The averaging kernels and their areas for these ten NDACC gases are shown in Fig. 4. The altitude ranges with sensitivity larger than 0.5 and the corresponding total DOFs are summarized in Table 3. These sensitive ranges indicate that the retrieved profile information comes by more than 50% from measurement, or, in other words, that the a priori information influences the retrieval by less than 50%. Each gas has different sensitive range. The sensitive range for HCN, CO and C₂H₆ is mainly tropospheric, and for ClONO₂, HCl and HF is mainly stratospheric. O₃, CH₄ and N₂O have high retrieval sensitivity in both troposphere and stratosphere. The HNO₃ has high retrieval sensitivity in stratosphere and in atmospheric boundary layer below 1.5 km.

4.3 Error analysis

As listed in Table 2, we classified errors as systematic or random according to whether they are constant between consecutive measurements, or vary randomly. For
comparison, the error items considered in error analysis are the same for the retrieval of all gases. The smoothing error $E_s$ is calculated via equation (6), the measurement error $E_m$ is calculated via equation (7), and all other error items $E_{var}$ are calculated via equation (8) (Rodgers, 2000).

$$E_s = (A - I)S_s(A - I)^T$$ (6)

$$E_m = G_yS_cG_y^T$$ (7)

$$E_{var} = G_yK_{var}S_{var}K_{var}^T G_y^T$$ (8)

where $S_{var}$ is the error covariance matrix of $var$. $K_{var}$ is weighting function matrix of $var$. Here $var$ refers to one of the error items in Table 2 except smoothing error and measurement error. In this study, the a priori error covariance for all non-retrieval parameters are set the same for all gases retrieval.

5 ILS influence study

This section presents the ILS influence study, whereby the degraded ILSs that simulated by ALIGN60 are used in the SFIT forward model, and the fractional difference (D%) in various quantities for each gas relative to the retrieval with an ideal ILS are computed. For each gas, sections 5.1 and 5.2 only select one typical spectrum for study. In order to retrieve these ten gases, five spectra with different wavenumber coverage are used. All of them are randomly selected from the routine measurements on a clear day at Hefei on February 16, 2016. The consistency of the resulting deduction is evaluated in section 5.3 where one year of measurements from August 2015 to August 2016 are used. The Hefei site has run NDACC observations with the Bruker 125HR for more than three years. We regularly use a low-pressure HBr cell to diagnose the misalignment of the spectrometer and to realign the instrument when indicated. As shown in Fig.5, all actual ILS degradations of the FTIR spectrometer within this selected period are less than 2% and can be regarded as ideal. For all spectra used in this study, the retrievals with all levels of ILS degradation fulfill the following filter criteria:

1) The root mean squares (RMSs) of the residual (difference between measured and calculated spectra after the fit) in all fitting windows has to be less than 3%.

2) The retrievals should converge for all levels of ILS degradation.
3) The concentrations of the target and interfering gases at each sub layer should be positive.

4) The solar intensity variation (SIV) should be less than 10%. The SIV within the duration of a spectrum is the ratio of the standard deviation to the average of the measured solar intensities.

These criteria are used to remove those spectra that have sampling errors or contaminated by aerosols, clouds, hazes or other unpredictable objects which cause a low SNR or a large detecting intensity variation. In following calculations, we have taken the retrievals with an ideal ILS as the reference. The fractional difference is defined here as,

$$D\% = \frac{X - X_{ref}}{X_{ref}} \times 100 \quad (9)$$

where $X$ is a vector which can include multiple elements such as gas profile or only one element such as DOFs, RMS, total column, total random uncertainty, total systematic uncertainty, or total uncertainty. The total random uncertainty and systematic uncertainty are the sum in quadrature of each individual uncertainty listed in Table 2, and the total uncertainty is the sum in quadrature of total random uncertainty and total systematic uncertainty. $X_{ref}$ is the same as $X$ but for the nominal ideal ILS.

### 5.1 ME amplitude and PE influence

In order to determine how the ILS degradation affects the NDACC gas retrievals, the results deduced from ILS considering both ME amplitude and PE are compared to those only considering ME amplitude or PE. All types of ILS degradation in section 3 are used in this study. Fig.6 exemplifies the case of ILS $j$, where the differences in total column, RMS, random uncertainty, systematic uncertainty, total uncertainty, and DOFs for each gas relative to the retrieval with an ideal ILS are compared. Fig.7 shows the fractional difference in profile of each gas for ILS $j$. The results show that the influence of ILS degradation on the total column, RMS, random uncertainty, systematic uncertainty, total uncertainty, DOFs, and profile can be approximated by
the linear sum of individual ME amplitude influence and PE influence. The PE influence is of secondary importance compared with the ME amplitude influence. The comparisons for the results retrieved with ILS a to i come to the same conclusions.

Figs. 8 and 9 show the influence of ILS a to j on total column and profile of all NDACC gases. The resulting influence amounts depend on deviation amount and deviation shape of ME. For positive MEs, in most cases, the ILS j causes the maximum influence, and for negative MEs, the ILS i causes the maximum influence. In a real instrument, the misalignment is a combination of misalignment a to j. In principle, for the same misalignment amplitude, it should not cause influence exceeding misalignment i or j. In the following, misalignment i and j are selected on behalf of negative and positive ME respectively to investigate how the ILS degradation influence the NDACC gas retrievals.

5.2 Sensitivity study

We simulated seven levels of negative ME i and positive ME j with ALIGN60, and incorporated them in the SFIT forward model, and then calculated the fractional difference in various quantities for each gas relative to the retrieval with an ideal ILS. The misalignments as inputs of ALIGN60 and the resulting ILSs are shown in Figs. 10 and 12. The corresponding Haidinger fringes at the maximum misalignment position are shown in Figs. 11 and 13. The ME deviation, decenter of Haidinger fringes and ILS deterioration varying over misalignment are evident. Fig. 14 is the sensitivity of total column with respect to different levels of ILS degradation. Figs. 15 ~ 18 are the same as Fig. 14 but for DOFs, RMS, uncertainty and profile. The results show that the ILS degradation affected total column, RMS, DOFs, retrieval uncertainty, and profile. Generally, the larger the ME deviation, the larger the influence. The positive and negative ME have opposite influence on total column, DOFs, total uncertainty and profile.

With respect to total column, the influence of ILS degradation on stratospheric gases is generally larger than the tropospheric gases. For a typical ILS degradation (10%), the total columns of stratospheric gases O₃, HNO₃, HCl, HF, and ClONO₂...
changed by 1.9%, 0.7%, 4%, 3%, and 23%, respectively. While the total columns of tropospheric gases \text{CH}_4, \text{CO}, \text{N}_2\text{O}, \text{C}_2\text{H}_6, \text{and HCN} changed by 0.04%, 2.1%, 0.2%, 1.1%, and 0.75%, respectively. For \text{O}_3 and \text{HNO}_3, positive ME causes an overestimated total column and negative ME causes an underestimated total column. For other gases, negative ME causes an overestimated total column and positive ME causes an underestimated total column. For all gases except \text{O}_3 and \text{CH}_4, the positive ME influence is larger than the negative ME influence. For \text{CH}_4, the negative ME influence is larger than the positive ME influence. For \text{O}_3, the level of the positive ME influence and the negative ME influence is very close.

For all gases, positive ME increases the DOFs and negative ME decreases DOFs. For all gases except HF and \text{CH}_4, both positive ME and negative ME increase RMS. For HF, positive ME increases RMS while negative ME decreases RMS. For \text{CH}_4, positive ME decreases RMS and negative ME increases RMS.

The influence on systematic uncertainty and random uncertainty depends on ME deviation type and gas type. The influence on total uncertainty is the combination of the influence on total systematic uncertainty and total random uncertainty. For all gases except \text{O}_3, positive ME decreases total uncertainty and negative ME increases total uncertainty. For \text{O}_3, positive ME increases total uncertainty and negative ME decreases total uncertainty.

The ILS degradation causes an evident difference in profile within the altitude ranges that show high retrieval sensitivity in Fig.4, or in other words, the sensitive ranges listed in Table 3. Generally, the profile is more sensitive to positive ME than negative PE, and the influence of ILS degradation on stratospheric gases is larger than the tropospheric gases.

5.3 consistency evaluation

This section uses the spectra recorded at Hefei from August 2015 to August 2016 to evaluate the consistency of above study. These spectra span a large difference in atmospheric water vapor, SZAs, surface pressures, surface temperatures, wind speeds, and wind directions (Fig. 19). All retrievals fulfill the above filter criteria are included.
in this study. A simulated ILS $j$ with maximum ME amplitude deviation of 5% is used in the retrieval. The results are compared to the retrievals deduced from an ideal ILS.

Fig. 20 exemplifies the fractional difference in total column, RMS, total uncertainty, and DOFs for each gas as a function of SZA. The results show that the fractional difference in total column, total uncertainty, and DOFs for all gases are consistent under different SZAs. For most gases, the fractional difference in RMS exhibits more scatters than the total column, total uncertainty, and DOFs. However, they are independent of SZA, and most of them are less than 10%. In general, the influence of ILS degradation on NDACC gases retrieval shows good consistency under different SZAs. The fractional difference as functions of humidity, pressure, SZA, temperature, wind direction, and wind speed come to the same conclusions.

6 Discussion and recommendation

For each gas, the $a$ priori covariance matrices of $S_a$, $S_e$, and $S_{var}$ are the same in the aforementioned study. According to equations 6 ~ 8, we conclude that the ILS degradation altered the weighting function matrix $K_x$ and eventually altered the quantities such as the total column, RMS, random uncertainty, systematic uncertainty, total uncertainty, DOFs, and profile. The change of $K_x$ is attributed to the fact that the ILS degradation alters gas absorption line shape and hence alters the structure of calculated spectra, and aggravates the mismatch between the calculated spectra and the measured spectra.

The stratospheric gases are more sensitive to ILS degradation than the tropospheric gases, and the ClONO$_2$ exhibits the largest sensitivity. This is because the absorption structure in stratosphere is narrower than that in troposphere, and is more easily affected by ILS degradation. We set the acceptable fractional difference in total column for ClONO$_2$ and other NDACC gases as 10% and 1%, respectively. Considering an excessively large of ME degradation (e.g., > 20%) seldom occurred within NDACC network because of the regular alignment at each site, the permitted maximum ILS degradation for each gas is deduced in Table 5 as:

1) The influence of ILS degradation on CH$_4$ and N$_2$O can be regarded as
negligible.

2) If a misalignment causes positive ME degradation, the maximum degradations for O₃, HNO₃, HCl, HF, ClONO₂, CO, C₂H₆, and HCN should be less than 6%, 15%, 5%, 5%, 5%, 9%, and 13%, respectively.

3) If a misalignment causes negative ME degradation, the maximum degradations for O₃, HCl, and HF should be less than 6%, 12%, and 12%, respectively.

Note that the retrievals of certain gases, e.g., O₃, CH₄, CO, and N₂O, can be divided into multiple independent sub layers depending on total DOFs. The above deductions don’t apply to partial column integrated over each sub layer because, as Figs. 17 and 18 show, the sensitivity of profile to ILS degradation is altitude dependent. How ILS degradation influences partial column of each NDACC gas and how much ILS deviation from unity is acceptable if an ideal line shape is assumed beyond the scope of this paper and will be published elsewhere.

7 Conclusion

We assessed the influence of instrumental line shape degradation on all current NDACC gases retrieval via investigation of sensitivities of total column, root mean square of fitting residual, total random uncertainty, total systematic uncertainty, total uncertainty, degrees of freedom, and profile with respect to modulation efficiency degradations. The study concluded that the influence of instrumental line shape degradation can be approximated by the linear sum of individual modulation efficiency amplitude influence and phase error influence. The phase error influence is of secondary importance compared with the modulation efficiency amplitude influence. The influence amounts depend on deviation amount and deviation shape of the modulation efficiency.

The stratospheric gases are more sensitive to instrumental line shape degradation than the tropospheric gases, and the positive modulation efficiency has more influence on total column or profile than the negative modulation efficiency. For a typical ILS degradation (10%), the columns of stratospheric gases O₃, HNO₃, HCl, HF, and ClONO₂ changed by 1.9%, 0.7%, 4%, 3%, and 23%, respectively. While the columns
of tropospheric gases \( \text{CH}_4 \), \( \text{CO} \), \( \text{N}_2 \text{O} \), \( \text{C}_2 \text{H}_6 \), and \( \text{HCN} \) changed by 0.04\%, 2.1\%, 0.2\%, 1.1\%, and 0.75\%, respectively. The influence of instrumental line shape degradation on NDACC gas retrievals shows good consistency under different meteorological conditions and solar zenith angle. In order to suppress the fractional difference in total column for \( \text{ClONO}_2 \) and other NDACC gases within 10\% and 1\%, respectively, the maximum positive modulation efficiency degradations for \( \text{O}_3 \), \( \text{HNO}_3 \), \( \text{HCl} \), \( \text{HF} \), \( \text{ClONO}_2 \), \( \text{CO} \), \( \text{C}_2 \text{H}_6 \), and \( \text{HCN} \) should be less than 6\%, 15\%, 5\%, 5\%, 5\%, 9\%, and 13\%, respectively; the maximum negative modulation efficiency degradations for \( \text{O}_3 \), \( \text{HCl} \), and \( \text{HF} \) should be less than 6\%, 12\%, and 12\%, respectively; the influence of ILS degradation on \( \text{CH}_4 \) and \( \text{N}_2 \text{O} \) can be regarded as negligible.

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**9 Figs**

Fig. 1. Simulated ILS degradation with respect to different types of misalignment. The results are derived from ALIGN60. Top left demonstrates different types of misalignment (a to j) used in the simulation, top right is the resulting ILS, bottom left is the resulting ME amplitude, and bottom right is the resulting PE. Descriptions for the misalignment a to j are listed in Table 1.

Fig. 2. The Haidinger fringes at maximum OPD for misalignment a to j shown in Fig. 1.
Fig. 3. Typical profiles of ten NDACC gases. Bottom panels are five tropospheric gases, i.e., CH₄, CO, N₂O, C₂H₆, and HCN. Top panels are five stratospheric gases, i.e., O₃, HNO₃, HCl, HF, and ClONO₂. Although the CO concentration above 60 km is much higher than that in the troposphere, it is regarded as tropospheric gas because it is an anthropologic pollution gas and shows large variation in troposphere.

Fig. 4. Averaging kernels of ten NDACC gases (color fine lines), and their area scaled by a factor of 0.2 (black bold line). They are deduced from the spectra recorded at Hefei on February 16, 2016 with an ideal ILS.
Fig. 5. ME amplitudes (left) and phase errors (right) along with OPD deduced from HBr cell measurements at Hefei.

Fig. 6. Fractional difference in total column, RMS, total random uncertainty, total systematic uncertainty, total uncertainty, and DOFs for misalignment $j$. “ME amplitude” represents the ILS only taken ME amplitude deviation into account. “PE” represents the ILS only taken PE deviation into account. “ME amplitude & PE” represents the ILS taken both ME amplitude and PE deviations into account. “Linear sum” represents the fractional difference of each item is linear sum of “ME amplitude” and “PE”. The ME amplitude and PE are obtained from ALIGN60 with misalignment $j$ in Fig. 1. The results are deduced from the spectra recorded at Hefei on February 16, 2016.
Fig. 7. Fractional difference in profile for misalignment $j$. The nomenclature in the plot legend is same as Fig. 6. The results are deduced from the spectra recorded at Hefei on February 16, 2016.

Fig. 8. Sensitivity of total column to different types of ILS degradation. The ILS $a$ to $j$ correspond to misalignment $a$ to $j$ in Table 1. The results are deduced from the spectra recorded at Hefei on February 16, 2016.
Fig. 9. Sensitivity of profile to different types of ILS degradation. The ILS $a$ to $j$ correspond to misalignment $a$ to $j$ in Table 1. The results are deduced from the spectra recorded at Hefei on February 16, 2016.

Fig. 10. Simulated positive ME deviations along with OPD. Top left demonstrates the misalignment, top right is the resulting ILS, bottom left is the resulting ME amplitude, and bottom right is the resulting PE.
Fig. 11. The Haidinger fringes at maximum OPD (the maximum misalignment position) for Fig. 10

Fig. 12. Simulated negative ME deviations along with OPD. Top left demonstrates the misalignment, top right is the resulting ILS, bottom left is the resulting ME amplitude, and bottom right is the resulting PE.
Fig. 13. The Haidinger fringes at 1/2 maximum OPD (the maximum misalignment position) for Fig. 12.

Fig. 14. Sensitivity of total column with respect to ME deviation. "P_Tclmn" represents the sensitivity of total column with respect to positive ME deviation and "N_Tclmn" represents the sensitivity of total column with respect to negative ME deviation. The results are deduced from the spectra recorded at Hefei on February 16, 2016.
Fig. 15. The same as Fig. 14 but for DOFs and fitting RMS. The acronyms in the legend are similar to those in Fig. 14.

Fig. 16. The same as Fig. 14 but for total random uncertainty, total systematic uncertainty and total uncertainty. The acronyms in the legend are similar to those in Fig. 14. “Trnd”, “Tsys” and “Tstd” represent total random uncertainty, total systematic uncertainty and total uncertainty, respectively.
Fig. 17. Sensitivity of profile with respect to ME deviation. “4%” represents the ME amplitude deviation is 4%. The nomenclature for other plot labels is straightforward. The results are deduced from the spectra recorded at Hefei on February 16, 2016.

Fig. 18. The same as Fig. 17 but for negative ME deviation.
Fig. 19. The meteorological data and SZA’s record at Hefei. Large span of all these parameters are shown within the period from August 2015 to August 2016 (black dotted square).

Fig. 20. Fractional difference in total column, RMS, total uncertainty, and DOFs as a function of SZA from August 2015 to August 2016 where ILS j with a maximum ME deviation of 5% is used.
<table>
<thead>
<tr>
<th>Type</th>
<th>Description</th>
<th>Input</th>
<th>Output in maximum</th>
</tr>
</thead>
<tbody>
<tr>
<td>a</td>
<td>No misalignment occurs: interferometer in ideal condition</td>
<td>none</td>
<td>ME amplitude: 1.00 PE: 0.000 rad.</td>
</tr>
<tr>
<td>b</td>
<td>Decenter of entrance field stop defining FOV: causes a linear increase in misalignment along OPD</td>
<td>0.33 [mrad] field stop error</td>
<td>ME amplitude: 0.86 PE: -0.056 rad.</td>
</tr>
<tr>
<td>c</td>
<td>Decenter of path measuring laser: causes a linear increase in phase error along OPD</td>
<td>0.33 [mrad] laser error</td>
<td>ME amplitude: 1.00 PE: -0.152 rad.</td>
</tr>
<tr>
<td>d</td>
<td>Constant shear: causes a constant shear offset of fixed retro-reflector</td>
<td>0.03 [cm]</td>
<td>ME amplitude: 1.00 PE: -0.056 rad.</td>
</tr>
<tr>
<td>e</td>
<td>Decreasing linear shear: causes a linear decrease in misalignment along OPD</td>
<td>0.03-0.00017*OPD [cm]</td>
<td>ME amplitude: 1.16 PE: -0.007 rad.</td>
</tr>
<tr>
<td>f</td>
<td>Increasing linear shear: causes a linear increase in misalignment along OPD</td>
<td>0.00017*OPD [cm]</td>
<td>ME amplitude: 0.86 PE: -0.056 rad.</td>
</tr>
<tr>
<td>g</td>
<td>Cosine bending of scanner bar: causes a cosine decrease in misalignment along OPD</td>
<td>0.03<em>cos(π</em>OPD/360) [cm]</td>
<td>ME amplitude: 1.16 PE: -0.013 rad.</td>
</tr>
<tr>
<td>h</td>
<td>Sine bending of scanner bar: causes a sine increase in misalignment along OPD</td>
<td>0.03<em>sin(π</em>OPD/360) [cm]</td>
<td>ME amplitude: 0.86 PE: -0.056 rad.</td>
</tr>
<tr>
<td>i</td>
<td>Cosine &amp; sine bending of scanner bar: causes a chord increase in misalignment before 1/2 maximum OPD and causes a chord decrease in misalignment after 1/2 maximum OPD</td>
<td>0.073*(sin(π<em>OPD/360)+cos(π</em>OPD/360))-0.073 [cm]</td>
<td>ME amplitude: 0.86 PE: -0.029 rad.</td>
</tr>
<tr>
<td>j</td>
<td>Constant shear plus cosine &amp; sine bending of scanner bar: causes a chordal decrease in misalignment before 1/2 maximum OPD and causes a chordal increase in misalignment after 1/2 maximum OPD</td>
<td>-0.073*(sin(π<em>OPD/360)+cos(π</em>OPD/360))+0.103 [cm]</td>
<td>ME amplitude: 1.16 PE: -0.056 rad.</td>
</tr>
</tbody>
</table>
The $b, f, h$, and $i$ are referred to increasing misalignment, the $e, g$, and $j$ are referred to decreasing misalignment.

Table 2. Summary of the retrieval parameters used for all NDACC gases. All micro windows (MW) are given in cm$^{-1}$

<table>
<thead>
<tr>
<th>Gases</th>
<th>O$_3$</th>
<th>HNO$_3$</th>
<th>HCl</th>
<th>HF</th>
<th>ClONO$_2$</th>
<th>CH$_4$</th>
<th>CO</th>
<th>N$_2$O</th>
<th>C$_2$H$_6$</th>
<th>HCN</th>
</tr>
</thead>
<tbody>
<tr>
<td>MW for profile retrievals</td>
<td>1000-1004.5</td>
<td>867.5-870</td>
<td>2727.73-2727.83</td>
<td>2775.7-2775.8</td>
<td>2925.8-2926.0</td>
<td>4109.4-4110.2</td>
<td>779.85-780.45</td>
<td>782.55-782.87</td>
<td>2613.7-2615.4</td>
<td>2835.5-2835.8</td>
</tr>
<tr>
<td>Retrieved interfering gases</td>
<td>H$_2$O, CO$_2$, C$_2$H$_4$, O$_3$668, O$_3$686</td>
<td>H$_2$O, OCS, NH$_3$, CH$_4$, NO$_2$, O$_3$, N$_2$O, HDO</td>
<td>H$_2$O, HDO, CH$_4$</td>
<td>O$_3$, HNO$_3$, H$_2$O, CO$_2$</td>
<td>CO$_2$, NO$_2$, H$_2$O, HDO</td>
<td>O$_3$, N$_2$O, CO$_2$, OCS, H$_2$O</td>
<td>CO$_2$, CH$_4$</td>
<td>H$_2$O, CH$_4$, O$_3$</td>
<td>H$_2$O, OCS, C$_2$H$_4$, CH$_4$</td>
<td></td>
</tr>
<tr>
<td>H$_2$O treatment</td>
<td>Profile retrieval</td>
<td>Scaling retrieval</td>
<td>Profile retrieval</td>
<td>Profile retrieval</td>
<td>Profile retrieval</td>
<td>Profile retrieval</td>
<td>Profile retrieval</td>
<td>Profile retrieval</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SNR for de-weighting</td>
<td>None</td>
<td>None</td>
<td>300</td>
<td>None</td>
<td>None</td>
<td>None</td>
<td>500</td>
<td>None</td>
<td>None</td>
<td>None</td>
</tr>
<tr>
<td>$S_a$</td>
<td>Diagonal: 20%</td>
<td>Diagonal: 50%</td>
<td>Diagonal: 50%</td>
<td>Diagonal: 10%</td>
<td>Diagonal: 100%</td>
<td>Diagonal: 10%</td>
<td>Diagonal: 10%</td>
<td>Diagonal: 11% ~ 27%</td>
<td>Diagonal: 10%</td>
<td>Diagonal: 10%</td>
</tr>
<tr>
<td>Error analysis</td>
<td>Systematic error:</td>
<td>-Smoothing error</td>
<td>-Errors from parameters not retrieved by sfit4</td>
<td>Background curvature, Optical path difference, Field of view, Solar line strength, Background slope, Solar line shift, Phase, Solar zenith angle, Line temperature broadening, Line pressure broadening, Line intensity</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Random error:</td>
<td>-Interference errors: Retrieval parameters, Interfering species</td>
<td>-Measurement error</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*The input uncertainties of all these items are the same and are included into error analysis if they are not retrieved. Otherwise, the corresponding uncertainties wouldn’t be included.
Table 3. Altitude ranges with sensitivity larger than 0.5 for all NDACC gases

<table>
<thead>
<tr>
<th>Items</th>
<th>O₃</th>
<th>HNO₃</th>
<th>HCl</th>
<th>HF</th>
<th>ClONO₂</th>
<th>CH₄</th>
<th>CO</th>
<th>N₂O</th>
<th>C₂H₆</th>
<th>HCN</th>
</tr>
</thead>
<tbody>
<tr>
<td>Altitude ranges (km)</td>
<td>Ground</td>
<td>17 - 28</td>
<td>18 - 42</td>
<td>18-44</td>
<td>20 - 28</td>
<td>Ground</td>
<td>Ground</td>
<td>Ground</td>
<td>Ground</td>
<td>4.5-18</td>
</tr>
<tr>
<td>Total DOFs</td>
<td>5.2</td>
<td>1.4</td>
<td>1.5</td>
<td>1.3</td>
<td>0.55</td>
<td>3.5</td>
<td>3.8</td>
<td>4.0</td>
<td>1.2</td>
<td>1.1</td>
</tr>
</tbody>
</table>

Table 4. Recommendation for suppressing fractional difference in total column for ClONO₂ and other NDACC gases within 10% and 1%, respectively

<table>
<thead>
<tr>
<th>Items</th>
<th>O₃</th>
<th>HNO₃</th>
<th>HCl</th>
<th>HF</th>
<th>ClONO₂</th>
<th>CH₄</th>
<th>CO</th>
<th>N₂O</th>
<th>C₂H₆</th>
<th>HCN</th>
</tr>
</thead>
<tbody>
<tr>
<td>Positive ME</td>
<td>&lt; 6%</td>
<td>&lt;15%</td>
<td>&lt;5%</td>
<td>&lt;5%</td>
<td>&lt;5%</td>
<td>*</td>
<td>&lt;5%</td>
<td>*</td>
<td>&lt;9%</td>
<td>&lt;13%</td>
</tr>
<tr>
<td>Negative ME</td>
<td>&lt; 6%</td>
<td>*</td>
<td>&lt;12%</td>
<td>&lt;12%</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td>*</td>
<td>*</td>
</tr>
</tbody>
</table>

*The influence on ClONO₂ is less than 10% and on all other NDACC gases are less than 1% even the ILS degrade by an excessively large of 28%, and thus can normally be regarded as negligible.