- Using a Speed-Dependent Voigt Line Shape to Retrieve O<sub>2</sub> from
- 2 Total Carbon Column Observing Network Solar Spectra to Improve
- 3 Measurements of XCO<sub>2</sub> Improving the Retrieval of XCO<sub>2</sub> from
- 4 Total Carbon Column Network Solar Spectra
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- 12 **Abstract**. High-resolution, laboratory, absorption spectra of the  $a^1\Delta_q \leftarrow X^3\Sigma_q^-$  oxygen(O<sub>2</sub>) band measured using
- 13 cavity ring-down spectroscopy were fitted using the Voigt and speed-dependent Voigt line shapes. We found that
- the speed-dependent Voigt line shape was better able to model the measured absorption coefficients than the Voigt
- line shape. We used these line shape models to calculate absorption coefficients to retrieve atmospheric #total
- 16 columns abundances of O<sub>2</sub> were retrieved from ground-based high resolution absorption spectra from four Fourier
- 17 transform spectrometers that are apart of the Total Carbon Column Observing Network (TCCON) sites using both
- 18 Voigt and speed dependent Voigt line shapes to calculate absorption coefficients. A lLower O2 concentration total
- columns was were retrieved with the speed-dependent Voigt line shape, with and the difference between the total
- 20 columns retrieved using the Voigt and speed-dependent Voigt line shapes increasing increased as a function of solar
- 21 zenith angle. Previous work has shown that carbon dioxide (CO<sub>2</sub>) total columns were also retrieved from the same
- 22 spectra using a Voigt line shape and are better retrieved using a speed-dependent Voigt line shape with line mixing.
- The column-averaged dry-air mole fraction of CO<sub>2</sub> (XCO<sub>2</sub>) was calculated using the ratio between the columns of
- 24 CO<sub>2</sub> and O<sub>2</sub> columns retrieved (from the same spectra) with both line shapes from measurements made over a one-
- year period at the four sites, and compared. The inclusion of speed dependence in the O<sub>2</sub> retrievals significantly
- 26 reduces the airmass dependence of XCO<sub>2</sub>. The TCCON empirical airmass correction factor for XCO<sub>2</sub> derived from a
- 27 year of measurements from TCCON sites at Darwin, Lamont, and Park Falls for XCO2 improved from
- 28 0.0071±0.0057 to 0.0012±0.0054 when speed dependence was included. XCO2 retrieved with the Voigt and speed
- 29 dependent Voigt line shapes was compared to aircraft profiles measured at 13 TCCON sites. T and the bias between
- the TCCON measurements and the calibrated integrated aircraft profile measurements was reduced from
- 31 0.9897±0.0005 1% to 1.0041±0.0005 0.4%, for XCO2 retrieved with the Voigt and speed dependent Voigt line
- 32 shapes respectively. These results suggest that speed dependence should be included in the forward model when
- fitting near-infrared CO<sub>2</sub> and O<sub>2</sub> spectra to improve the accuracy of XCO<sub>2</sub> measurements.

#### 1. Introduction

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35 Accurate remote sensing of greenhouse gases (GHGs), such as CO<sub>2</sub>, in Earth's atmosphere is important for studying 36 the carbon cycle  $\frac{1}{100}$  in order to better understand and predict climate change. The absorption of solar radiation by  $O_2$  in 37 the Earth's atmosphere is important because it can be used to study the properties of clouds and aerosols, and to 38 determine vertical profiles of temperature and surface pressure. Wallace and Livingston (1990) were the first to retrieve total columns of  $O_2$  from some of the discrete lines of the  $a^1\Delta_q \leftarrow X^3\Sigma_q^-$  band of  $O_2$  centered at 1.27  $\mu$ m 39 40 (which will be referred to bellow as the 1.27 µm band) using atmospheric solar absorption spectra from the Kitt 41 Peak observatory. Mlawer et al. (1998) recorded solar absorption spectra in the near-infrared (NIR) region to study collision-induced absorption (CIA) in the  $a^1\Delta_g \leftarrow X^3\Sigma_g^-$  band of  $O_2$  centered at 1.27  $\mu$ m (which will be referred to as 42 43 the 1.27 µm band) as well as two other O<sub>2</sub> bands. The spectra were compared to a line-by-line radiative transfer 44 model and the differences between the measured and calculated spectra showed the need for better absorption 45 coefficients in order to accurately model the 1.27 µm band (Mlawer et al., 1998). Subsequently, spectroscopic 46 parameters needed to calculate the absorption coefficients from discrete transitions of the 1.27 µm band were 47 measured in multiple studies (Cheah et al., 2000; Newman et al., 1999, 2000; Smith and Newnham, 2000), as was 48 collision-induced absorption (CIA) (Maté et al., 1999; Smith and Newnham, 2000), while Smith et al. (2001) 49 validated the work done in Smith and Newnham (2000) using solar absorption spectra. 50 The 1.27 µm band is of particular importance to the Total Carbon Column Observing Network (TCCON) (Wunch 51 et al., 2011). TCCON is a ground-based remote sensing network that makes accurate and precise measurements of 52 GHGs for satellite validation and carbon cycle studies. Using the O<sub>2</sub> column retrieved from solar absorption spectra, 53 the column-averaged dry-air mole fraction of CO<sub>2</sub> (XCO<sub>2</sub>) has been shown to provide better precision than using the 54 surface pressure to calculate XCO<sub>2</sub> (Yang et al., 2002) (Washenfelder et al., 2006). The O<sub>2</sub> column is retrieved from 55 the 1.27 µm band because of its close proximity to the spectral lines used to retrieve CO<sub>2</sub>, thereby reducing the 56 impact of solar tracker mis-pointing and an imperfect instrument line shape (ILS) (Washenfelder et al., 2006). To 57 improve the retrievals of O<sub>2</sub> from the 1.27 µm band, Washenfelder et al. (2006) found that adjusting the 58 spectroscopic parameters in HITRAN 2004 (Rothman et al., 2005) decreased the airmass and temperature 59 dependence of the O2 column. These revised spectroscopic parameters were included in HITRAN 2008 (Rothman et 60 al., 2009). Atmospheric solar absorption measurements from this band made at the Park Falls TCCON site by 61 Washenfelder et al. (2006) were the first measurements to observe the electric-quadrupole transitions (Gordon et al., 62 2010). Leshchishina et al. (2011, 2010) subsequently used cavity-ring-down spectra to retrieve spectroscopic 63 parameters for the 1.27 µm band using a Voigt spectral line shape and these parameters were included in HITRAN 64 2012 (Rothman et al., 2013). Spectroscopic parameters for the discrete spectral lines of the O<sub>2</sub> 1.27 μm band from HITRAN 2016 (Gordon et al., 2017) are very similar to HITRAN 2012 except that HITRAN 2016 includes 65 66 improved line positions reported by Yu et al. (2014). 67 Extensive spectral line shape studies have been performed for the O<sub>2</sub> A-band, which is centered at 762 nm and used 68 by the Greenhouse Gases Observing Satellite (GOSAT) (Yokota et al., 2009) and the Orbiting Carbon Observing

69 Observatory-2 (OCO-2) satellite (Crisp et al., 2004) to determine surface pressure. These studies Studies have 70 showed shown that the Voigt line shape was is inadequate to describe the spectral line shape of the discrete O<sub>2</sub> lines 71 in the A-band, Dicke narrowing occurs when the motion of the molecule is diffusive due to collisions changing the 72 velocity and direction of the molecule during the time that it is excited. This diffusive motion is taken into account 73 by averaging over many different Doppler states resulting in a line width that is narrower than the Doppler width 74 (Dicke, 1953). The need to take into account Dicke narrowing was shown in Long et al. (2010) and Predoi-Cross et 75 al. (2008) found it necessary to use a spectral Line shape model that accounted for Dicke narrowing when fitting the 76 discrete lines of the O<sub>2</sub> A-band. and Line mixing, which occurs when collisions transfer intensity from one part of 77 the spectral band to another (Lévy et al., 1992), was shown to be prevalent in multiple studies (Predoi-Cross et al., 78 2008; Tran et al., 2006; Tran and Hartmann, 2008). Tran and Hartmann (2008) showed that including line mixing 79 when calculating the O<sub>2</sub> A-band absorption coefficients reduced the airmass dependence of the O<sub>2</sub> column retrieved 80 from TCCON spectra. When fitting cavity ring-down spectra of the O<sub>2</sub> A-band, Drouin et al. (2017) found it 81 necessary to use a speed-dependence Voigt line shape, which takes into account different speeds at the time of 82 collision (Shannon et al., 1986), with line mixing to properly fit the discrete spectral lines of the O<sub>2</sub> A-band. 83 The need to include non-Voigt effects when calculating absorption coefficients for the O<sub>2</sub> 1.27 µm band was first 84 shown in Hartmann et al. (2013) and Lamouroux et al. (2014). In Hartmann et al. (2013) and Lamouroux et al. 85 (2014), Lorentzian widths were calculated using the re-quantized classical molecular-dynamics simulations 86 (rCMDSs) and used to fit cavity-ring-down spectra with a Voigt line shape for some isolated transitions in the O<sub>2</sub> 87 1.27 µm band. The studies concluded that a Voigt line shape is insufficient for modeling the spectral lines of the O<sub>2</sub> 88 1.27 µm band and that effects such as speed dependence and Dicke narrowing should be included in the line shape 89 calculation. 90 In this study, air-broadened laboratory cavity-ring-down spectra of the O<sub>2</sub> 1.27 µm band were fitted using a spectral 91 line shape that takes into account speed dependence. The corresponding derived spectroscopic parameters for the 92 speed-dependent Voigt line shape were used to calculate absorption coefficients when fitting high-resolution solar absorption spectra. Using these new O<sub>2</sub> total columns, and the simultaneously measured were combined with CO<sub>2</sub> 93 94 total columns, from using the updated line shape model described by Mendonca et al. (2016), to calculate XCO<sub>2</sub> and 95 compared these results with XCO<sub>2</sub> retrieved using a Voigt line shape. Section 2 details the formulas used to 96 calculate absorption coefficients using different spectral line shapes. In Section 3, we describe the retrieval of 97 spectroscopic parameters from three air-broadened cavity-ring-down spectra fitted with a speed-dependent Voigt 98 line shape. For Section 4, the speed-dependent line shape along with the retrieved spectroscopic parameters is used 99 to fit solar absorption spectra from four TCCON sites and retrieve total columns of O2, which is compared to O2 100 retrieved using a Voigt line shape. In Section 5, we investigate the change in the airmass dependence of XCO<sub>2</sub> with 101 the new O<sub>2</sub> parameters retrievals. In Section 6, we discuss our results and their implications for remote sensing of 102 greenhouse gases.

## 2. Absorption Coefficient Calculations

### 104 2.1 Voigt Line Shape

- The Voigt line shape is the convolution of the Lorentz and the Gaussian Doppler line shapes profiles, which model
- pressure and Doppler broadening of the spectral line respectively. The corresponding absorption coefficient, k, at a
- given wavenumber v becomes:

$$k(v) = N \sum_{j} S_{j} \left(\frac{1}{\gamma_{D_{j}}}\right) \left(\frac{\ln(2)}{\pi}\right)^{1/2} \left(Re\left[c(v, x_{j}, y_{j})\right]\right)$$
(1)

- where N is the number density,  $S_j$  is the line intensity of spectral line j,  $\gamma_{D_j}$  is the Doppler half-width (HWHM), c is
- the complex error function, and

$$x_j = \frac{\left(v - v_j^0 - P\delta_j^0\right)}{\gamma_{D_j}} (\ln(2))^{1/2}, \ y_j = \frac{\gamma_{L_j}}{\gamma_{D_j}} (\ln(2))^{1/2}.$$
 (2)

- Here,  $v_j^o$  is the position of the spectral line j, P is the pressure, and  $\delta_j^o$  is the pressure-shift coefficient. The Lorentz
- 111 half-width,  $\gamma_{L_i}$ , is calculated using:

$$\gamma_{L_j}(T) = P \gamma_{L_j}^o \left(\frac{296}{T}\right)^n \tag{3}$$

- where  $\gamma_{L_i}^o$  is the air-broadened Lorentz half-width coefficient (at reference temperature 296 K) and n is the exponent
- of temperature dependence. The Voigt line shape assumes that pressure broadening is accurately represented by a
- Lorentz profile calculated for the stastical average velocity at the time of collission.
- 115 2.2 Speed-Dependent Voigt Line Shape
- The speed-dependent Voigt line shape refines the pressure broadening component of the Voigt by calculating
- multiple Lorentz profiles for different speeds at the time of collision. The final contribution from pressure
- broadening to the speed-dependent Voigt is the weighted sum of Lorentz profiles (weighted by the Maxwell-
- Boltzmann speed-distribution) calculated for different speeds at the time of collision. To take speed dependence into
- 120 account, we use tThe speed-dependent Voigt line shape (Ciuryło, 1998) with the quadratic representation of the
- Lorentz width and pressure shift (Rohart et al., 1994) is:

$$k(v) = N\left(\frac{2}{\pi^{\frac{3}{2}}}\right) \sum_{j} S_{j} \int_{-\infty}^{\infty} e^{-V^{2}} V\left(tan^{-1} \left[\frac{x_{j} - Ba_{\delta_{j}}((V)V^{2} - 1.5) + V}{y_{j}(1 + a_{\gamma_{L_{j}}}((V)V^{2} - 1.5))}\right]\right) dV$$
 (4)

- where  $a_{\gamma_{L_j}}$  is the speed-dependent Lorentz width parameter (unitless) for line j,  $a_{\delta_j}$  is the speed-dependent pressure-
- shift parameter (unitless), B is  $\frac{(\ln(2))^{1/2}}{\gamma_{D_i}}$ , V is the ratio of the absorbing molecule's speed to the most probable speed
- of the absorbing molecule, and all other variables are defined before.

#### 3. Fitting Laboratory Spectra

- O<sub>2</sub>, unlike CO<sub>2</sub> and CH<sub>4</sub>, cannot produce an electric dipole moment and therefore should not be infrared active.
- However, O<sub>2</sub> has two unpaired electrons in the ground state that produce a magnetic dipole moment. Due to the
- unpaired electrons in the ground state  $(X^3\Sigma_q^-)$  the rotational state (N) is split into three components which are given
- by J = N-1, J = N, and J = N+1, while in the upper state  $(a^1\Delta_a)$ , J = N. When labeling a transition, the following
- nomenclature is used  $\Delta N(N'')\Delta I(I'')$  (Leshchishina et al., 2010), where  $\Delta N$  is the difference between N' in the upper
- state and N'' in the lower state,  $\Delta J$  is the difference between J' in the upper state and J'' in the lower state. The
- magnetic transitions of  $a^1\Delta_g \leftarrow X^3\Sigma_g^-$  allow for  $\Delta J=0, \pm 1$ . This leads to 9 branches observed: P(N'')Q(J''),
- 133 R(N'')Q(J''), and Q(N'')Q(J''), for  $\Delta J = 0$ , O(N'')P(J''), P(N'')P(J''), and Q(N'')P(J''), for  $\Delta J = -1$ , and Q(N'')P(J''),
- 134 R(N'')R(J''), and Q(N'')R(J''), for  $\Delta I = 1$ .
- Absorption coefficients for three room temperature air-broadened (NIST Standard reference materal® 2659a
- 136 containing 79.28 % N<sub>2</sub>, 20.720(43) % O<sub>2</sub>, 0.0029 % Ar, 0.00015 % H<sub>2</sub>O, and 0.001 % other compounds) spectra
- were measured at the National Institute of Standards and Technology (NIST) using the frequency-stabilized cavity-
- ring-down spectroscopy (FS-CRDS) ) technique (Hodges et al., 2004; Hodges, 2005). The absorption spectra were
- acquired at pressures of 131 kPa, 99.3 kPa, and 66.9 kPa, at temperatures of 296.28 K, 296.34 K, and 296.30 K
- respectively. Figure 1a shows the three measured absorption spectra. A more detailed discussion of the present FS-
- 141 CRDS spectrometer can be found in Lin et al. (2015).
- The spectra were fitted individually using a Voigt line shape (Eq. 1), with  $S_j$ ,  $\gamma_{L_j}^o$ , and  $\delta_j^o$  for the main isotope of the
- magnetic dipole lines of the O<sub>2</sub> 1.27 μm band for lines with an intensity greater than 7.0x10<sup>-28</sup> cm<sup>-1</sup>/(molecule cm<sup>-2</sup>).
- 144 The spectroscopic parameters measured in Leshchishina et al. (2011) for the spectral lines of interest were used as
- the a priori for the retrieved spectroscopic parameters. The line positions were left fixed to the values measured in
- Leshchishina et al. (2011), and all other O<sub>2</sub> spectral lines (intensity less 7.0x10<sup>-28</sup> cm<sup>-1</sup>/(molecule cm<sup>-2</sup>)) were
- calculated using a Voigt line shape with spectroscopic parameters from HITRAN 2012 (Rothman et al., 2013).
- Spectral fits were done using the Isqnonlin function in Matlab, with a user-defined Jacobian matrix. The Jacobian
- was constructed by taking the derivative of the absorption coefficients with respect to the parameters of interest.
- Using an analytical Jacobian instead of the finite difference method is both computationally faster and more
- accurate. The Voigt line shape was calculated using the Matlab code created by Abrarov and Quine (2011) to
- 152 calculate the complex error function and its derivatives. To take collision-induced absorption (CIA) into account, a
- set of 50 Legendre polynomials were added together by retrieving the weighting coefficients needed to add the
- polynomials to fit the CIA for each spectrum. Figure 1b shows the residual (measured minus calculated absorption
- coefficients) when using a Voigt line shape with the retrieved spectroscopic parameters. The plot shows that residual
- structure still remains for all three spectra. The Root Mean Square (RMS) residual values for the spectra are given
- by the legend at the side of the plot.
- Figure 2 is the same plot as Figure 1 but for the P(11)P(11), P(11)Q(10), P(9)P(9), and P(9)Q(8) spectral lines only.
- Figure 2b shows that for all four spectral lines there is a "W" shaped residual at the line center. The P(11)P(11) line

160 was also measured by Hartmann et al. (2013) at pressures ranging from 6.7 to 107 kPa. Figure 5 of Hartmann et al. 161 (2013) shows the P(11)P(11) line at a pressure of 66.7 kPa, which is approximately the pressure of the 66.9 kPa 162 spectrum (blue spectrum in Figure 1 and 2). When one compares the blue residual of the P(11)P(11) line in Figure 163 2b to that of the residual of the left panel of Figure 5 of Hartmann et al. (2013), one can see that the residuals are the 164 same. Figure 6 of Hartmann et al. (2013) show that the amplitude of the residual increases with decreasing pressure, 165 which is also seen in Figure 2b. Figure 3 of Lamouroux et al. (2014) shows the same "W" residual for the P(9)P(9) lines and that the amplitude of the residual increases with decreasing pressure (although for lower pressures) 166 167 consistent with the results for the P(9)P(9) line in Figure 2b. 168 Figure 1c shows the residual when using the speed-dependent Voigt (Eq. 4) to fit each spectrum individually. To use 169 Eq. (4) requires integration over all possible speeds, which is not computationally practical, so we employ the simple numerical integration scheme as was done by Wehr (2005). When fitting the spectra, parameters  $S_i$ ,  $\gamma_{L_i}^o$ ,  $\delta_i^o$ , 170  $a_{\gamma_{L_i}}$  and  $a_{\delta_i}$  were retrieved for lines of intensity greater than  $7.0 \times 10^{-28}$  cm<sup>-1</sup>/(molecule cm<sup>-2</sup>), while all other O<sub>2</sub> lines 171 172 were calculated using a Voigt line shape and spectroscopic parameters from HITRAN 2012 (Rothman et al., 2013b). 173 The Jacobian matrix was created by taking the derivative with respect to each parameter of interest, as was done 174 with the Voigt fits. By taking speed-dependent effects into account, the residuals were reduced to 25 times smaller 175 than those for the Voigt fit and the RMS residuals (given in the legend of Figure 1c) are 10 times smaller. However, 176 some residual structure still remains, which is more evident in the Q and R branches than the P branch. Figure 177 2c shows the four lines in the P branch, as discussed when analyzing the Voigt fits. A small residual "W" remains at 178 line center, as well as residuals from weak O2 lines. 179 Figure 3 shows the averaged intensity, Lorentz width coefficient, pressure shift coefficient, and speed-dependent 180 shift coefficient of the 1.27  $\mu$ m O<sub>2</sub> band, retrieved from the three spectra, plotted as a function of  $\frac{f''}{f''}$  quantum number m which is m=-J (where J is the lower state rotational quantum number) for the P-branch lines, m=J for the 181 O-branch lines, and m=J+1 for the R-branch lines. The intensity, Lorentz widths, and pressure shifts show a \( \frac{\mu}{\text{m}} \) m 182 183 dependence for these parameters for the P and R sub-branches. The measured Lorentz widths and pressure shifts for the Q sub-branches show a #" m dependence but are not as strong as the P and R sub-branches. This is because the 184 185 Q branch lines are broadened enough to blend with each other since they are spaced closer together than the P or R 186 branch lines. Figure 1c shows that some of the residual structure in the Q branch increases with pressure and is 187 partly due to the blending of these transitions as the pressure increases. The weak O<sub>2</sub> absorption lines also blend in 188 with the Q branch, contributing to the residual structure in Figure 1c. We tried retrieving the spectroscopic 189 parameters for the weak O<sub>2</sub> absorption lines, but since they were overlapping with the strong O<sub>2</sub> lines, it was not 190 possible. Figure 4a shows the retrieved speed-dependent width parameter averaged over the three spectra, plotted as a function of  $\frac{\mu}{m}$ , showing that it increases with  $\frac{\mu}{m}$ . Error bars correspond to the  $2\sigma$  standard deviation and are 191 192 large regardless of sub-branch. Figure 4b shows the retrieved speed-dependent width for the PQ sub-branch for the 193 different pressures. The speed-dependent width shows the same  $\frac{J''}{m}$  dependence regardless of pressure, but also 194 increases with decreasing pressure as is the case for sub-branches. It should be noted that the speed-dependent width 195 parameter should be independent of pressure.

# 4. Fitting Solar Spectra

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197 High-resolution solar absorption spectra were measured at four TCCON sites using a Bruker IFS 125HR FTIR 198 spectrometer with a room temperature InGaAs detector at a spectral resolution of 0.02 cm<sup>-1</sup> (45 cm maximum 199 optical path difference). The raw interferograms recorded by the instrument were processed into spectra using the 200 I2S software package (Wunch, D. et al., 2015) that corrects them for solar intensity variations (Keppel-Aleks et al., 201 2007), phase errors (Mertz, 1967), and laser sampling errors (Wunch, D. et al., 2015), and then preforms a fast 202 Fourier transforms to convert the interferograms into spectra (Bergland, 1969). The GGG software package (Wunch, 203 D. et al., 2015) is used to retrieve total columns of atmospheric trace gases. GFIT is the main code that contains the 204 forward model, which calculates a solar absorption spectrum using a line-by-line radiative transfer model and an 205 iterative non-linear least square fitting algorithm that scales an a priori gas profile to obtain the best fit to the 206 measured spectrum. A priori profiles for GHGs are created by an empirical model in GGG that is based on 207 measurements from the balloon-borne JPL MkIV Fourier Transform Spectrometer (FTS) (Toon, 1991), the 208 Atmospheric Chemistry Experiment (ACE) FTS instrument aboard SCISAT (Bernath et al., 2005), and in situ 209 GLOBALVIEW data (Wunch et al., 2011). Temperature and pressure profiles, as well as H<sub>2</sub>O a priori profiles are 210 generated from the National Centers for Environmental Prediction (NCEP) data. The calculations are performed for 211 71 atmospheric layers (0 km to 70 km), so all a priori profiles are generated on a vertical grid of 1 km. 212 In the current GGG software package (Wunch, D. et al., 2015), the forward model of GFIT calculates absorption 213 coefficients for the discrete lines of the O<sub>2</sub> 1.27 µm band using a Voigt line shape and spectroscopic parameters 214 from Washenfelder et al. (2006a) and Gordon et al. (2010). To take CIA into account, absorption coefficients are 215 calculated using a Voigt line shape and spectroscopic parameters from the foreign-collision-induced absorption 216 (FCIA) and self-collision-induced absorption (SCIA) spectral line lists provided with the GGG software package 217 (Wunch, D. et al., 2015). Spectroscopic parameters in the FCIA and SCIA line lists were retrieved by Geoff Toon by 218 fitting the laboratory spectra of Smith and Newnham (2000). This was done by retrieving the integrated absorption 219 at every 1 cm<sup>-1</sup> of the spectrum and using a Voigt line shape, with fixed Lorentz width and no pressure shift. In 220 GFIT, a volume scale factor is retrieved for the CIA and discrete lines separately so that the O<sub>2</sub> column is derived 221 from the discrete lines of the 1.27 µm band only. Airglow is not considered when fitting the 1.27 µm band since the 222 spectrometer views the sun directly, and airglow is overwhelmed by such a bright source. The continuum level and 223 tilt of the 100% transmission level is fitted using a weighted combination of the first two Legendre polynomials. 224 Absorption coefficient for all other trace gases are calculated using a Voigt line shape and spectroscopic parameters 225 from the atm. 101 line list (Toon, G. C., 2014a) (Wunch, D. et al., 2015) and solar lines are fitted using the solar line 226 list (Toon, G. C., 2014b)(Wunch, D. et al., 2015). 227 Figure 5 shows the spectral fit to a solar absorption spectrum recorded at Eureka on March 27, 2015, at a solar 228 zenith angle (SZA) of 81.32° (airmass of 6.3). This spectrum is an average of 5 Eureka scans. The TCCON standard 229 is single scan but 5 scans were averaged to decrease the noise. The measured spectrum (red circles), calculated 230 spectrum (black circles) and transitions from all gases in the window (colored lines, refer to the legend for different 231 gases) are shown in Figure 5b. The residual obtained using a Voigt line shape to calculate the discrete lines of the O<sub>2</sub>

232 1.27 µm band is shown in red in Figure 5a. The blue residual is the result of using a speed-dependent Voigt line 233 shape with the spectroscopic parameters retrieved from fitting the absorption coefficients in Section 3. To decrease 234 the amount of time it takes to calculate the absorption coefficients, the quadratic-Speed Dependent Voigt (qSDV) 235 computational approach of Ngo et al. (2013) and Tran et al. (2013) was used instead of Eq. (4) since it requires the 236 Voigt calculation only twice, while Eq. (4) requires numerical integration scheme with 33 iterations. The 237 temperature-dependent parameter of the Lorentz width of the discrete lines of the O2 1.27 µm band reported in HITRAN 2012 was used to take temperature dependence into account for  $\gamma_{L_i}(T)$ . There was only a slight 238 239 improvement in the fit residuals with the new absorption coefficients (using the qSDV), as seen in Figure 5a. 240 Absorption coefficients calculated with the qSDV were used to retrieve total columns of O<sub>2</sub> from solar spectra 241 recorded over a one year period at TCCON sites in Eureka (eu) (Nunavut, Canada) (Batchelor et al., 2009; Strong et al., 2017)(Batchelor et al., 2009), Park Falls (pa) (Wisconsin, U.S.A) (Washenfelder et al., 2006; Wennberg et al., 242 243 2017)(Washenfelder et al., 2006b), Lamont (oc) (Oklahoma, U.S.A) (Wennberg et al., 2017b), and Darwin (db) (Australia) (Deutscher et al., 2010; Griffith et al., 2017) (Deutscher et al., 2010). In total 131 124 spectra were fitted 244 245 using the qSDV and the average root mean square (RMS) residual of the fit only decreased by 0.5 %.

#### 5. Impact of O<sub>2</sub> Columns on XCO<sub>2</sub> Measurements

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The O<sub>2</sub> column retrieved from the 1.27 μm band with a Voigt line shape and spectroscopic parameters from the atm.101 line list (Toon, G. C., 2014a) (Wunch, D. et al., 2015) has an airmass dependence such that the O<sub>2</sub> column retrieved increases as a function of solar zenith angle (or airmass). Using spectra recorded from Eureka, Park Falls, Lamont, and Darwin over one-year periods, total columns of O<sub>2</sub> were retrieved using (1) a Voigt spectral line shape with spectroscopic parameters from the atm.101 line list and (2) the qSDV with the spectroscopic parameters determined in Section 3. Figure 6 shows the percent difference calculated as the column from the qSDV retrieval minus the column from the Voigt retrieval, which was then divided by the latter and multiplied by 100, plotted as a function of solar zenith angle (SZA). At the smallest SZA, the qSDV retrieves 0.75% less O<sub>2</sub> than the Voigt, with the difference increasing to approximately 1.8% as the SZA approaches 90°. Figure 7 shows XAIR from Park Falls on June 18, 2013. XAIR is the column of air (determined using surface pressure recorded at the site) divided by the column of O<sub>2</sub> retrieved from the spectra and multiplied by 0.2095, which is the dry air mole fraction of O<sub>2</sub> in Earth's atmosphere. Ideally XAIR should be 1 but when using O2 retrieved with a Voigt line shape (red points) it is closer to 0.98 near noon (small SZA) and lower near the start and end of the day (large SZA). When using O2 retrieved with the qSDV, XAIR is closer to 0.988 near noon and a bit higher near the start and end of the day. This means the O<sub>2</sub> column, retrieved with the qSDV, decreases as a function of SZA, while previously the column increased as a function of SZA when the Voigt line shape is used.

### 5.1 Airmass Dependence of XCO<sub>2</sub>

Since the standard TCCON XCO<sub>2</sub> (and all other XGas) is calculated using the column of O<sub>2</sub> instead of the surface pressure, errors associated with the retrieval of O<sub>2</sub>, such as the airmass dependence of the O<sub>2</sub> column, will affect XCO<sub>2</sub>. Figure 8 is XCO<sub>2</sub> calculated for four different combinations pertaining to the two CO<sub>2</sub> column retrievals and

267 the O<sub>2</sub> column retrievals. The CO<sub>2</sub> columns were retrieved with either a Voigt line shape (the standard GGG2014 268 approach) or the qSDV with line mixing as done in Mendonca et al. (2016) while the O<sub>2</sub> columns were retrieved 269 with either a Voigt (the standard GGG2014 approach) or the new qSDV approach developed here. Figure 8 shows a 270 spurious symmetric component to XCO<sub>2</sub> when the total column of O<sub>2</sub> is retrieved with the Voigt line shape, 271 regardless of line shape used to retrieve CO<sub>2</sub>. When the qSDV is used to retrieve total columns of O<sub>2</sub>, the symmetric 272 component of XCO<sub>2</sub> is dismissed regardless of line shape used to retrieve CO<sub>2</sub>. This is because the airmass 273 dependence of the column of O<sub>2</sub> retrieved using the qSDV is more consistent with the airmass dependence of the 274 column of CO<sub>2</sub> (for both line shapes used to retrieve CO<sub>2</sub>). Mendonca et al. (2016) showed that using the qSDV with 275 line mixing results in better fits to the CO<sub>2</sub> windows and impacts the airmass dependence of the retrieved column of 276 CO<sub>2</sub>. When using a Voigt line shape the retrieved column amount of CO<sub>2</sub> decreases as airmass increases until the 277 airmass is large (SZA of about 82°) at which point the retrieved column of CO<sub>2</sub> increases as the airmass increases, 278 changing the shape of the airmass dependence of the CO<sub>2</sub> column. When the qSDV with line mixing is used, the 279 retrieved column of CO<sub>2</sub> decreases as a function of airmass (up until the sun is above the horizon).

In order to To correct for this, an empirical correction is applied to all TCCON XCO<sub>2</sub> (and XGas). The empirical correction determines the antisymmetrical component of the day's XCO<sub>2</sub>, which is assumed to be the true variation of XCO<sub>2</sub> throughout the day, as well as the symmetrical component, which is caused by the airmass dependence of the retrieved column of the gas of interest and O<sub>2</sub>. We can, therefore, represent a measurement as (Wunch et al., 2011):

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$$y_i = \hat{y}[1 + \alpha S(\theta_i) + \beta A(t_i)] \tag{5}$$

where  $\hat{y}$  is the mean value of XCO<sub>2</sub> measured that day,  $\beta$  is the fitted coefficient of the antisymmetric function  $A(t_i)$  and  $\alpha$  is the fitted coefficient of the symmetric function  $S(\theta_i)$ . The antisymmetric function is calculated by (Wunch et al., 2011):

$$A(t_i) = \sin(2\pi(t_i - t_{noon})) \tag{6}$$

288 where  $t_i$  is the time of the measurement and  $t_{noon}$  is the time at solar noon, both in units of days. The symmetric function is calculated by (Wunch et al., 2011):

$$S(\theta_i) = \left(\frac{\theta_i + 13^o}{90^o + 13^o}\right)^3 - \left(\frac{\theta_i 45^o + 13^o}{90^o + 13^o}\right)^3 \tag{7}$$

where θ<sub>i</sub> is the SZA in degrees. To determine α for the different line shapes, total columns of CO<sub>2</sub> were retrieved
 using the Voigt line shape (Wunch, D. et al., 2015) and the qSDV with line mixing (Mendonca et al., 2016).
 Henceforth, we will refer to XCO<sub>2</sub> calculated from O<sub>2</sub> and CO<sub>2</sub> using the Voigt line shape as XCO<sub>2</sub> Voigt and the
 qSDV line shape as XCO<sub>2</sub> qSDV.

Figure  $\frac{97}{7}$  shows the average  $\alpha$  calculated for each season at Darwin, Lamont, and Park Falls. Eureka XCO<sub>2</sub> cannot be used to determine  $\alpha$  because Eureka measurements do not go through the same range of SZAs as the other three

296 sites due to its geolocation. The average  $\alpha$  values derived from XCO<sub>2</sub> Voigt are represented by stars in Figure 97, 297 while the squares indicate XCO<sub>2</sub> qSDV. At all three sites,  $\alpha$  is closer to 0 when the qSDV line shape is used in the 298 retrieval compared to the Voigt retrieval, regardless of the season. The average  $\alpha$  for XCO<sub>2</sub> Voigt calculated from a 299 year of measurements from Darwin, Park Falls, and Lamont is -0.0071±0.0057 and that for XCO2 qSDV is -300  $0.0012 \pm 0.0054$ . 301 For all four sites,  $\alpha = -0.0071$  is used to correct XCO<sub>2</sub> Voigt measurements. Figure  $\frac{10}{8}$ a shows the XCO<sub>2</sub> Voigt 302 anomalies plotted as a function of SZA. The data is expressed as the daily XCO<sub>2</sub> anomaly, which is the difference 303 between the XCO<sub>2</sub> value and the daily median value, in order to remove the seasonal cycle. When XCO<sub>2</sub> is left 304 uncorrected for airmass dependencies, there is a clear airmass dependence where the amount of XCO<sub>2</sub> decreases as a function of SZA up to a SZA of approximately 82°, at which point XCO2 and increases as a function of SZA at 305 306 andgles greater than 82°. Figure 108 b shows XCO2 Voigt corrected for the airmass dependence. This airmass 307 correction works well only up to a SZA of approximately 82°, after which the correction only serves to increase the 308 airmass dependence. Figure 108c is the same as 108a but for the uncorrected XCO2 qSDV measurements, while 309 Figure 108d is the same as 108b but for the corrected XCO<sub>2</sub> qSDV measurements. When the airmass correction is 310 applied to XCO<sub>2</sub> qSDV there is a small difference between the corrected and uncorrected XCO<sub>2</sub> qSDV 311 measurements, with the difference only noticeable for the Darwin measurements recorded at SZA > 60°. For XCO<sub>2</sub> 312 qSDV measurements made at SZA > 82° XCO<sub>2</sub> does not increase with SZA as it does with the Voigt. 313 5.2 Accuracy of XCO<sub>2</sub> 314 To assess the accuracy of TCCON XCO<sub>2</sub> measurements, they are compared to aircraft XCO<sub>2</sub> profile measurements 315 using the method described in Wunch et al. (2010). Figure 119a shows the comparison between the aircraft XCO<sub>2</sub> 316 measurements (legend on at the top details the different aircraft) and TCCON XCO<sub>2</sub> Voigt measurements for 13 317 TCCON sites (given by the color-coded legend on at the bottom right). The gray line indicates the one-to-one line 318 and the dashed line is the line of best fit. There is a bias of 0.9897±0.0005, as given by the slope of the line of best 319 fit in Figure 119a, for the XCO<sub>2</sub> Voigt measurements. Figure 119b is the same as 119a but for the XCO<sub>2</sub> qSDV 320 measurements. The bias between the aircraft XCO2 measurements and the XCO2 qSDV measurements is 321 1.0041±0.0005 as given by the slope of the line of best fit in Figure 119b. This increase in the slope can be 322 explained by an increase in the retrieved column of CO<sub>2</sub> when using the qSDV with line mixing as shown in 323 Mendonca et al. (2016) as well as combined with a decrease in the retrieved O<sub>2</sub> column due to using the qSDV. As 324 discussed previously (section 5) the decrease in the retrieved O<sub>2</sub> column is an improvement but the expected column 325 of O<sub>2</sub> is still approximately 1.2% higher (at the smallest SZA) than it should be. Therefore, the retrieved column of 326 CO<sub>2</sub> is higher than it should be, and the slope would be greater if the retrieved column of O<sub>2</sub> was 1.2% lower. Never 327

TCCON XCO<sub>2</sub> measurements are divided by the scale factors (or bias determined in Figure 119) to calibrate to the WMO scale. For all TCCON XCO<sub>2</sub> measurements retrieved with a Voigt line shape, the airmass correction is first

XCO<sub>2</sub> and aircraft XCO<sub>2</sub> measurements by 0.62 %.

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the less Using using the qSDV to retrieve total columns of CO<sub>2</sub> and O<sub>2</sub> thus reduces the difference between TCCON

331 applied to the data and the result is divided by the determined bias factor, 0.9897. Figure 1240a to 1240d shows 332 XCO<sub>2</sub> Voigt (for Eureka, Park Falls, Lamont, and Darwin respectively) indicated by red square boxes in the plots. 333 XCO<sub>2</sub> Voigt measurements made at SZA > 82° have been filtered out because they cannot be corrected for the 334 airmass dependence. The blue boxes are XCO<sub>2</sub> qSDV corrected for airmass dependence and scaled by 1.0041. No 335 filter was applied to the XCO<sub>2</sub> qSDV measurements for SZA since the airmass dependence correction works at all 336 SZA. Figure 1240e to 1240h shows the difference between XCO<sub>2</sub> Voigt and XCO<sub>2</sub> qSDV for Eureka, Park Falls, 337 Lamont, and Darwin respectively. The mean differences for the data shown in Figures 1210e to 1210h are 338 0.113±0.082, -0.102±0.223, -0.132±0.241, and -0.059±0.231 µmol/mol (ppm) for Eureka, Park Falls, Lamont, and 339 Darwin respectively. The difference throughout the day at Park Falls, Lamont, and Darwin varies between -0.6 to 340 0.2 µmol/mol and is SZA dependent. 341 Figure 1344a shows XCO<sub>2</sub> Voigt corrected for the airmass dependence, as well as XCO<sub>2</sub> qSDV, uncorrected and 342 corrected for the airmass dependence. These XCO<sub>2</sub> measurements were retrieved from Park Falls spectra recorded 343 on June 18, 2013. For all three XCO<sub>2</sub> measurements, the amount of XCO<sub>2</sub> decreases throughout the day. Figure 344 1311b shows the difference between the corrected Voigt XCO2 and the uncorrected qSDV XCO2, as well as the 345 difference between the corrected Voigt XCO2 and the corrected qSDV XCO2. The difference between the Voigt and 346 the qSDV (corrected and uncorrected) shows that at the start and end of the day, more XCO2 is retrieved with the 347 qSDV, while at midday less is retrieved with the qSDV. The range in the differences seen in Figure 1240e to 1240h 348 varies with SZA throughout the day as shown in Figure 1311b. 349 **6. Discussion and Conclusions** 350 Using cavity ring-down spectra measured in the lab, we have shown that the Voigt line shape is insufficient to 351 model the line shape of O<sub>2</sub> for the 1.27 µm band, consistent with the results of (Hartmann et al. (2013) and 352 Lamouroux et al. (2014). By using the speed-dependent Voigt line shape when calculating the absorption 353 coefficients, we were better able to reproduce the measured absorption coefficients than using the Voigt line shape. 354 However, some residual structure still remains as seen Figures 1 and 2. This is partly due to the blending of spectral 355 lines (i.e., line mixing) and the inability to retrieve the spectroscopic parameters for weak O<sub>2</sub> transitions. Fitting low-356 pressure spectra would help with isolating spectral lines and decreasing the uncertainty on the retrieved 357 spectroscopic parameters for the Q branch lines. 358 Accurate measurements of the pressure shifts in the 1.27 µm band have been hard to obtain as shown in Newman et 359 al. (1999) and Hill et al., (2003). While the retrieved pressure shifts show a dependence on quantum number m 360 (Figure 3c) as one would expect, this dependence is not as strong as the m dependence of the Lorentz widths (Figure 361 3b). This can be explained by the fact that line mixing, which is shown to be important for the O<sub>2</sub> A-band, was not 362 considered when fitting the cavity-ringdown spectra. Neglecting line mixing usually produces an asymmetric residual in the discrete lines as well as a broad residual feature associated with the fact that collisions are transferring 363 364 intensity from one part of the spectrum to another. By fitting a set of Legendre polynomials for CIA we could be 365 simultaneously fitting the broader band feature associated with line mixing while the retrieved pressure shifts, and

| 366 | speed-dependent pressure shifts could be compensating for the asymmetric structure one would see in the discrete                     |
|-----|--|
| 367 | lines when neglecting line mixing. The remaining structure, as seen in Figure 1c, could be due to neglecting line                    |
| 368 | mixing especially in the Q-branch where the spacing between spectral lines is small (in comparison to the P and R                    |
| 369 | branches) and line mixing is most likely prevalent.  |
| 370 | The pressure dependence of the retrieved speed-dependent width parameter is an indication that Dicke narrowing                       |
| 371 | needs to be taken into account, as shown by Bui et al. (2014) for CO <sub>2</sub> . However, when When dealing with both             |
| 372 | speed dependence and Dicke narrowing are present, a multi-spectrum fit needs to be used due to the correlation                       |
| 373 | between the parameters (Bui et al., 2014). Domysławska et al. (2016) recommend using the qSDV to model the line                      |
| 374 | shape of O2 based on multiple line shape studies of the O2 B-band. In these studies, a multi-spectrum fit to low                     |
| 375 | pressure (0.27-5.87 kPa) cavity-ring down spectra was preformed testing multiple line shapes that took speed-                        |
| 376 | dependence and Dicke narrowing into account both separately and simultaneously. They found that the line shapes                      |
| 377 | that only used Dicke narrowing were not good enough to model the line shape of the O2 B-band lines, but a line                       |
| 378 | shape that included either speed-dependence or both speed-dependence and Dicke narrowing produced similar                            |
| 379 | quality fits, ultimately concluding that speed-dependence has a larger effect than Dicke narrowing. It was noted in                  |
| 380 | the study by Wójtewicz et al., (2014) that both Dicke narrowing and speed-dependent effects might simultaneously                     |
| 381 | play an important role in modeling the line shape of the O2 B-band lines. However, the speed-dependent and Dicke                     |
| 382 | narrowing parameters are highly correlated at low pressures. To reduce the correlation requires either a multi-                      |
| 383 | spectrum fit of spectra at low pressures with high enough signal to nosie ratio or spectra that cover a wide range of                |
| 384 | pressure (Wójtewicz et al., 2014). So, by combining the high-pressure spectra used in this study with low pressure                   |
| 385 | spectra in a multipspectrum fit both the speed-dependence and Dicke narrowing parameters could be retrieved. The                     |
| 386 | temperature dependence of the Lorentz width coefficients of this band has never been measured before, which could                    |
| 387 | have an impact on the airmass dependence of O2. Combining high-pressure cavity-ring-down absorption coefficient                      |
| 388 | measurements with those for low pressures and different temperatures as done in Devi et al. (2015 and 2016) for                      |
| 389 | CH <sub>4</sub> in would lead to more accurate line shape parameters for O <sub>2</sub> .  |
| 390 | By taking speed dependence into account for both CO <sub>2</sub> (in the work of Mendonca et al., 2016) and O <sub>2</sub> (the work |
| 391 | presented here), we were able to significantly decrease the airmass dependence of TCCON XCO <sub>2</sub> and the bias                |
| 392 | between TCCON and aircraft XCO $_2$ . With the qSDV line shape, XCO $_2$ measurements made at SZA $> 82^{\rm o}$ no longer           |
| 393 | have to be discarded, resulting in more XCO2 measurement available from all TCCON sites. This is particularly                        |
| 394 | important for high-latitude TCCON sites, such as Eureka, because measurements made from late February to late                        |
| 395 | March and from late September until mid-October are made at SZA > 82°. Filtering out these large SZA                                 |
| 396 | measurements thus limits the knowledge of the seasonal cycle of XCO2 at high latitudes. The airmass dependence of                    |
| 397 | the O2 column not only effects XCO2 but all trace gases measured by TCCON and in the future the airmass                              |
| 398 | dependence of all XGas will be determined with these new O <sub>2</sub> columns.   |

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## 652 Figures

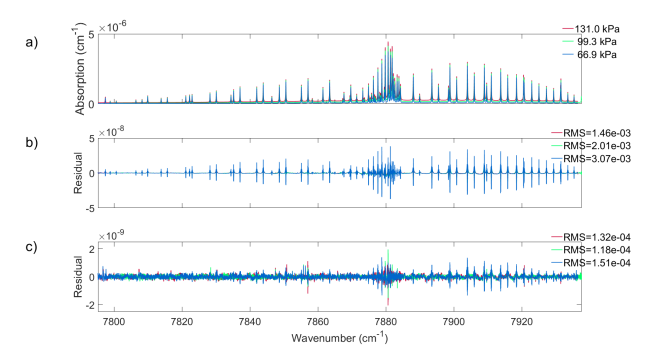
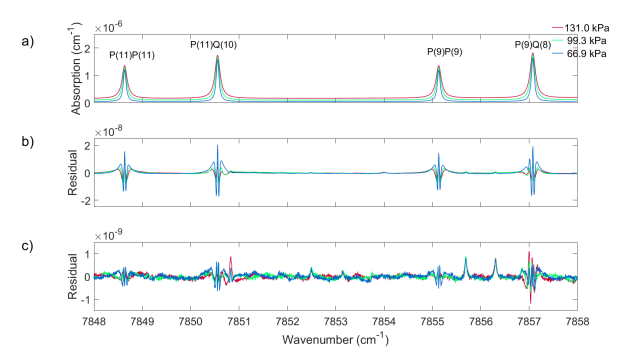


Figure 1: (a) Cavity-ring-down absorption coefficients for  $O_2$  measured at the three pressures indicated in the legend at approximately room temperature and a volume mixing ratio of 0.20720(43) for  $O_2$ . The difference between measured absorption coefficients and those calculated using (b) a Voigt line shape, and (c) the speed-dependent Voigt line shape. Note the difference in scale between panels (b) and (c).



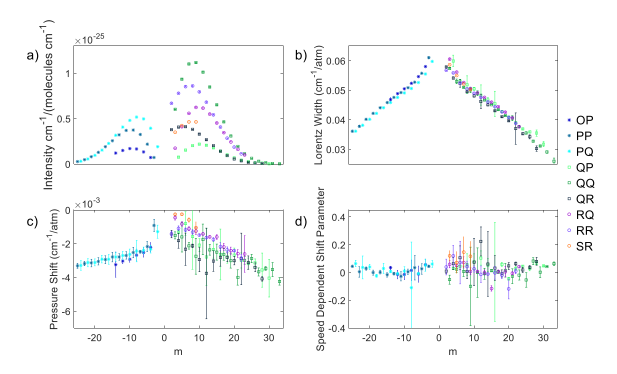


Figure 3: The averaged measured (a) intensity, (b) Lorentz line width, (c) pressure shift, and (d) speed-dependent pressure shift retrieved from the three cavity ring-down spectra of the 1.27  $\mu$ m band of O<sub>2</sub>. All data are plotted as a function of lower state rotational quantum number J m which is m =-J for the P-branch lines, m=J for the Q-branch, and m=J+1 for the R-branch (where J is the lower state rotational quantum number) and the uncertainties shown are 2 $\sigma$ .

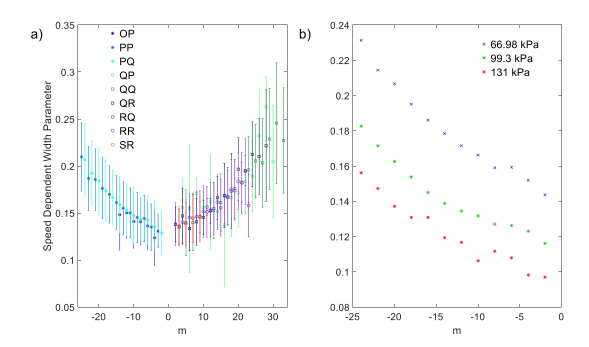
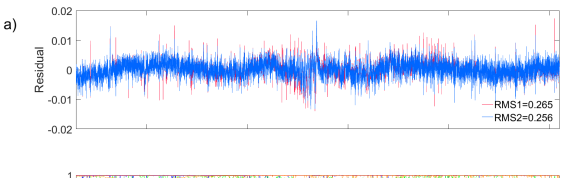


Figure 4: (a) The averaged measured speed-dependent width parameter of the 1.27  $\mu$ m band of O<sub>2</sub> plotted as a function of m. lower state rotational quantum number J''. (b) The measured speed-dependent width parameter for spectral lines that belong to the PQ sub-branch plotted as a function of m. lower state rotational quantum number J''.



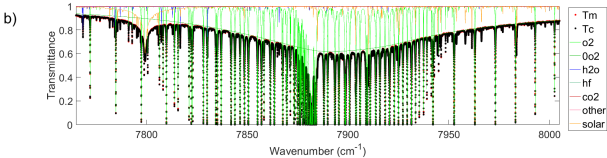


Figure 5: (a) The residuals (measured minus calculated) for a spectrum measured at Eureka on March 27, 2015 at a SZA of 81.32°. The red residual is the result of using the Voigt line shape and the blue is from using the qSDV. (b) The measured (red dots) and calculated (black dots), with the qSDV, spectrum, along with the gases included in the fit (refer to the legend to the right) in the spectral window.

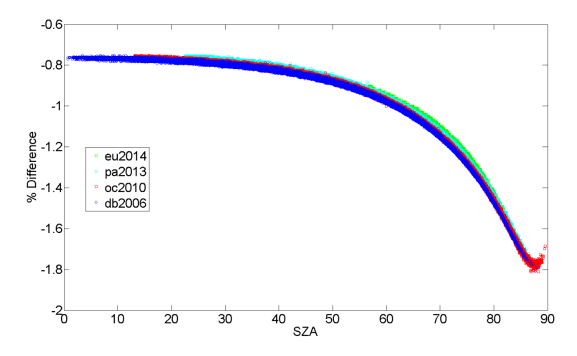


Figure 6: The percent difference between the  $O_2$  column retrieved with the Voigt and qSDV line shapes for a year of measurements from Eureka (eu), Park Falls (pa), Lamont (oc), and Darwin (db).

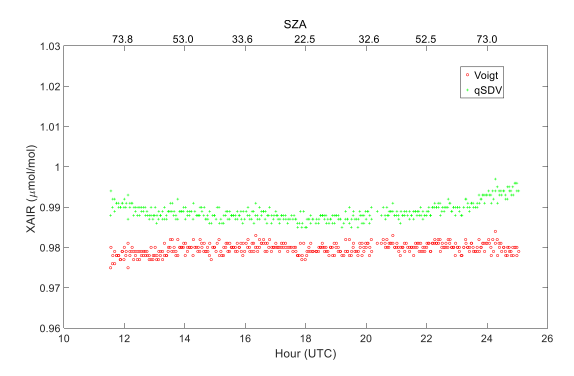


Figure 7: XAIR from Park Falls retrieved from spectra recorded on June 18, 2013. XAIR is calculated using O<sub>2</sub> columns retrieved using a Voigt (red) and qSDV (green) line shapes. The top x-axis is the SZA that corresponds to the hour on the bottom x-axis.

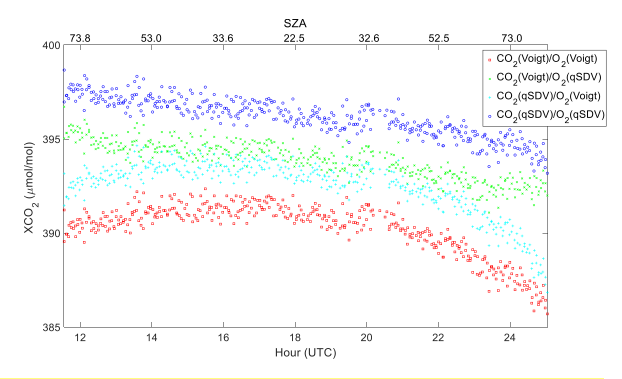


Figure 8: XCO<sub>2</sub> calculated from the CO<sub>2</sub> and O<sub>2</sub> columns retrieved from Park Falls spectra recorded on June 18, 2013. The CO<sub>2</sub> columns were retrieved using either the Voigt line shape or the qSDV with line mixing, while the O<sub>2</sub> columns were retrieved using either the Voigt or qSDV line shapes. XCO<sub>2</sub> was calculated in four ways: 1) Both CO<sub>2</sub> and O<sub>2</sub> columns retrieved using the Voigt line shape (red), 2) CO<sub>2</sub> columns retrieved with the Voigt and O<sub>2</sub> columns retrieved with the qSDV (green), 3) CO<sub>2</sub> columns retrieved with the qSDV and line mixing and O<sub>2</sub> columns retrieved with the Voigt (cyan), and 4) CO<sub>2</sub> columns retrieved with the qSDV and line mixing and O<sub>2</sub> columns retrieved with the qSDV (blue). The top x-axis is the SZA that corresponds to the hour on the bottom x-axis.

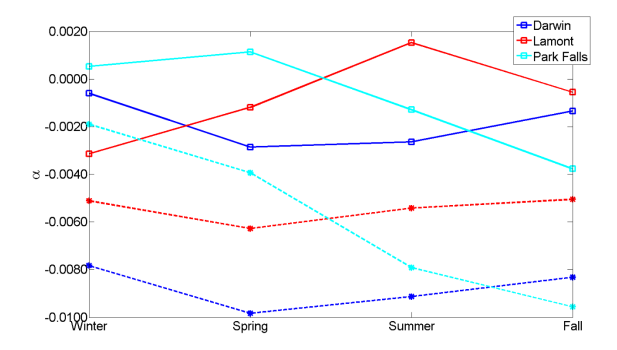


Figure 97: The average airmass-dependent correction factor for  $XCO_2$  derived from a year of spectra measured at Darwin, Lamont, and Park Falls for different seasons. The dashed lines with stars are the  $\alpha$  for  $XCO_2$  retrieved using a Voigt line shape for both  $CO_2$  and  $O_2$  columns. The solid lines with squares are from  $XCO_2$  retrieved using the qSDV for both  $CO_2$  and  $O_2$  columns.

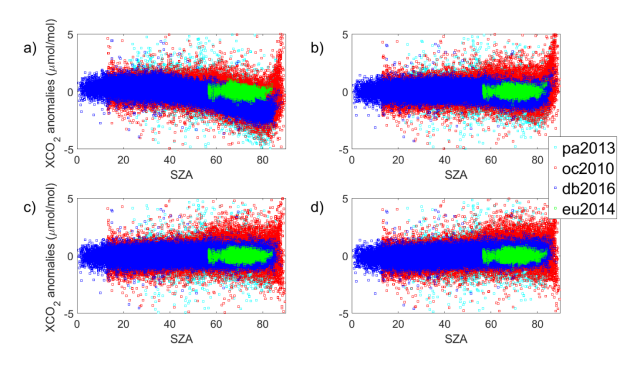


Figure 108: (a) XCO<sub>2</sub> Voigt anomaly for a year of measurements from the four TCCON sites. The XCO<sub>2</sub> anomaly is the difference between each XCO<sub>2</sub> value and the daily median XCO<sub>2</sub>. (b) The XCO<sub>2</sub> Voigt anomaly after the airmass dependence correction is applied to the XCO<sub>2</sub> Voigt data. (c) XCO<sub>2</sub> qSDV anomaly. (d) XCO<sub>2</sub> qSDV anomaly after correction for the airmass dependence.

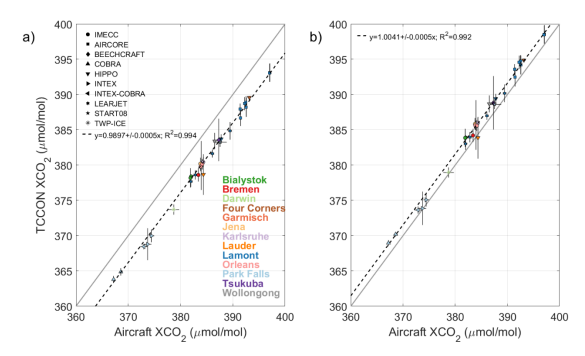


Figure 119: (a) Correlation between TCCON and aircraft XCO<sub>2</sub> Voigt measurements for 13 TCCON sites. Each aircraft type is indicated by a different symbol given by the legend in the top left corner. Each site is represented by a different colour given by the legend in the bottom right corner. The grey line indicates the one-to-one line and the dashed line is the line of best fit for the data. The slope of the line of best fit as well as the error on the slope are given in the plot. (b) the same as (a) but for XCO<sub>2</sub> qSDV.

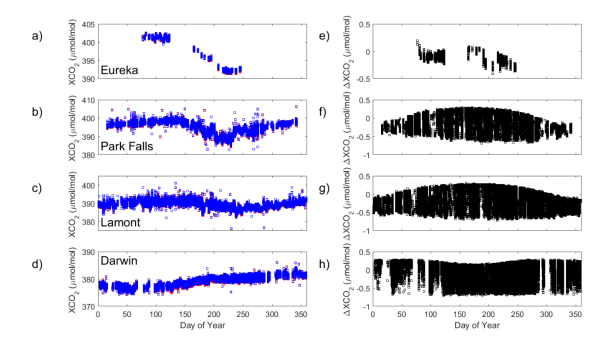


Figure 1210: (a) to (d) XCO<sub>2</sub> plotted as a function of day of the year for Eureka (2014), Park Falls (2013), Lamont (2010), and Darwin (2006) respectively. The mostly-hidden red boxes are XCO<sub>2</sub> calculated from using a Voigt line shape in the retrieval and the blue boxes are from using the qSDV. (e) to (h) the difference between XCO<sub>2</sub> Voigt and XCO<sub>2</sub> qSDV.



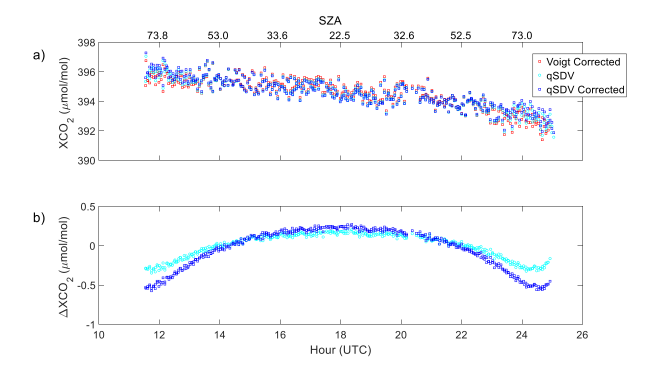


Figure 1311: (a) XCO<sub>2</sub> from Park Falls retrieved from spectra recorded on June 18, 2013. Plotted is XCO<sub>2</sub> retrieved: (1) with a Voigt line shape and corrected for the airmass dependence (red squares), (2) with the qSDV (cyan circles), and (3) with the qSDV and corrected for the airmass dependence (blue squares). (b) the difference between the Voigt XCO<sub>2</sub> corrected and the qSDV XCO<sub>2</sub> (cyan circles), and the difference between the Voigt XCO<sub>2</sub> corrected and the qSDV XCO<sub>2</sub> corrected (blue squares). The top x-axis is the SZA that corresponds to the hour on the bottom x-axis.