New and improved infra-red absorption cross sections and ACE-FTS retrievals of carbon tetrachloride (CCl₄)

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A B S T R A C T

Carbon tetrachloride (CCl₄) is one of the species regulated by the Montreal Protocol on account of its ability to deplete stratospheric ozone. As such, the inconsistency between observations of its abundance and estimated sources and sinks is an important problem requiring urgent attention (Carpenter et al., 2014) [5]. Satellite remote-sensing has a role to play, particularly limb sounders which can provide vertical profiles into the stratosphere and therefore validate stratospheric loss rates in atmospheric models.

This work is in two parts. The first describes new and improved high-resolution infra-red absorption cross sections of carbon tetrachloride/dry synthetic air over the spectral range 700–860 cm⁻¹ for a range of temperatures and pressures (7.5–760 Torr and 208–296 K) appropriate for atmospheric conditions. This new cross-section dataset improves upon the one currently available in the HITRAN and GEISA databases. The second describes a new, preliminary ACE-FTS carbon tetrachloride retrieval that improves upon the v3.0/v3.5 data products, which are biased high by up to ~20–30% relative to ground measurements. Making use of the new spectroscopic data, this retrieval also improves the microwindow selection, contains additional interfering species, and utilizes a new instrumental lineshape; it will form the basis for the upcoming v4.0 CCl₄ data product.

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1. Introduction

A clear, colourless, sweet-smelling liquid at room temperature, carbon tetrachloride (CCl₄; also known as tetrachloromethane or CFC-10) was first synthesised in 1839 from chloroform and chlorine gas by Henry Victor Regnault [1]. As carbon tetrachloride does not occur naturally in the Earth system, its presence in the atmosphere, soil and ocean arises through its anthropogenic uses.

In the early twentieth century, carbon tetrachloride found many uses, for example in dry cleaning (excellent solvent properties), fire extinguishers, postage stamp collecting (to reveal watermarks without damaging the paper), lava lamps, and even as a refrigerant in early refrigerators. The discovery that carbon tetrachloride is harmful to human health – a potential human carcinogen, it has been shown to depress the central nervous system, inhibit liver and kidney function, and even kill – led to a decline in these uses. However, this had no effect on the largest industrial application of carbon tetrachloride in the twentieth century, its use as a feedstock in the production of trichlorofluoromethane (CFC-11) and dichlorodifluoromethane (CFC-12), which were commercialised in the 1930s as nonflammable and non-toxic refrigerants [2].
It was not until the realisation that long-lived chlorine-containing species such as CCl4 and the CFCs (chlorofluorocarbons) could destroy stratospheric ozone that the use of CCl4 in industry began to decline, as regulated by the 1987 Montreal Protocol (and its later amendments). Under the terms of the Protocol production and consumption of CCl4 were eliminated for developed countries in 1996 and for developing countries in 2010 [3]. There are no limits on the use of CCl4 as a feedstock, however, so it is still used, for example, in the production of hydrofluorocarbons. Although an ozone-depleting substance, with an ozone depletion potential of 0.72 [4], CCl4 is also a very strong greenhouse gas with a 100-year global warming potential of 1730 [4].

As one of the species controlled by the Montreal Protocol, there is much work carried out in monitoring CCl4 atmospheric concentrations by global networks such as AGAGE (Advanced Global Atmospheric Gases Experiment), NOAA (National Oceanic and Atmospheric Administration), and UCI (University of California, Irvine). The latest ozone assessment report from 2014 [5] indicates that the global surface mean mole fraction of CCl4 has continued to decline between 2008 and 2012, with the AGAGE and UCI networks reporting a rate of decline of 1.2–1.3% from 2011 to 2012, and 1.6% reported by the NOAA network. These rates of decline are comparable with those determined by remote sensing instruments of 1.1–1.2% yr⁻¹ from the Atmospheric Chemistry Experiment-Fourier transform spectrometer (ACE-FTS) on SCISAT and the ground-based FTS at Jungfraujoch [6].

Recently there has been particular interest in carbon tetrachloride, on account of the inconsistency between observations of its abundance and estimated sources and sinks [3]. A recent study [3] has suggested that the observed inter-hemispheric gradient for CCl4 (1.5 ± 0.2 ppt for 2000–2012) is primarily caused by ongoing emissions, however the 2007–2012 emissions estimate based on the UNEP reported production and feedstock usage is near-zero. Additionally, the decline of atmospheric concentrations is slower than can be accounted for by our current best estimates of the total CCl4 lifetime and its uncertainties.

In addition to the ground-based networks of CCl4 measurements, remote-sensing measurements which provide vertical profiles into the stratosphere are particularly useful in validating stratospheric loss rates in atmospheric models (the atmospheric loss of CCl4 is essentially all due to photolysis in the stratosphere). The literature reports a number of remote-sensing instruments capable of measuring CCl4 in the Earth’s atmosphere: ATMOS (Atmospheric Trace Molecule Experiment) [7] and CIRRIS 1A (Cryogenic Infra-Red Radiance Instrumentation for Shuttle) [8], both deployed on the space shuttle, the JPL balloon-borne MkIV interferometer [1], MIPAS (Michelson Interferometer for Passive Atmospheric Sounding) on ENVISAT (ENVironmental SATEllite) (e.g. [9]), and the ACE-FTS [1,10].

In the literature, the only global distribution of CCl4 extending up to the mid-stratosphere is derived from ACE-FTS measurements. Covering the spectral region 750–4400 cm⁻¹ with a resolution of 0.02 cm⁻¹, the ACE-FTS instrument uses the sun as a light source to record limb transmission through the Earth’s atmosphere (~300 km effective length) during sunrise and sunset (‘solar occultation’). The measured spectra, with high signal-to-noise ratios and through long atmospheric path lengths, provide a low detection threshold for retrieving the profiles of trace species. In fact, the ACE-FTS can detect more of these species than any other satellite instrument, although it only records spectra for at most 30 occultation events per day [11,12 (this issue)]. Unfortunately, the ACE-FTS v3.0/v3.5 CCl4 retrieval is biased high by up to ~20–30% relative to ground measurements; the reasons for this have been attributed to errors in the CCl4 spectroscopy, bad line parameters from other absorbing species in the microwave, and several absorbing species missing from the forward model. The aim of the present work was to improve the ACE-FTS CCl4 retrieval and minimise this high bias. This problem has been attacked on two fronts: (1) through the utilisation of new laboratory spectroscopic measurements of air-broadened CCl4 samples over a range of atmospheric pressure-temperature (PT) combinations and; (2) through improvements in the retrieval itself, in particular improved microwindow selection (the avoidance of spectral regions associated with poor or inadequate line parameters of interfering species), the inclusion of new interfering species, and a new instrumental line-shape (ILS). This new scheme will form the basis for the upcoming processing version 4.0 of ACE-FTS data. Section 2 of this manuscript presents some spectroscopic background to CCl4, and a discussion of previous infra-red (IR) absorption cross section datasets. Section 3 provides details on the new measurements, the derivation of cross sections, and a discussion of the results and comparison with previous measurements. Section 4 provides details of the new ACE-FTS CCl4 retrieval scheme.

2. Infra-red spectroscopy of carbon tetrachloride

2.1. Spectroscopic background

Since carbon and chlorine each have two stable isotopes, there are ten stable isotopologues of carbon tetrachloride, namely 12/13C35Cl4, 12/13C35Cl37Cl, 12/13C35Cl37Cl2, 12/13C35Cl37Cl3, and 12/13C37Cl4; these belong to the point groups Td, C3v, C3v, C3v, and Td, respectively. With natural abundances of ~99% and ~1% for 12C and 13C, and ~76% and ~24% for 35Cl and 37Cl, it turns out that the four isotopologues 12C35Cl4, 12C35Cl37Cl, 12C35Cl37Cl2, and 12/13C35Cl37Cl3 account for ~99% of the isotopologues found, that is 33%, 42%, 20% and 4%, respectively [13].

CCl4 has nine normal vibrational modes; in the Td point group these are labelled v1 (non-degenerate, A1), v2 (doubly-degenerate, E), v3 (triply degenerate, T2), and v4 (triply degenerate, T2). For the lower-symmetry isotopologues, the degeneracy can be lifted; v2 splits into A1 and A2 for C2v, and v3 and v4 split into A1 and E under C3v, A1, B1, and B2 under C2v. In the literature, however, it is more common to label bands under the assumption of Td symmetry, as the splittings are likely small to a first approximation.
The strong $\nu_3$ fundamental and $\nu_1+\nu_4$ combination bands of CCl$_4$ are situated in the 700–860 cm$^{-1}$ spectral region, with Fermi resonance between these levels resulting in both bands having mixed character and the latter having enhanced intensity [13]. The extremely dense rotation–vibration spectrum is complicated by the presence of isotopologues, which contribute to the spectrum in proportion with their natural abundances. As such, it is a difficult task to derive spectroscopic line parameters for CCl$_4$. For this reason, remote-sensing of the Earth’s atmosphere requires the use of absorption cross sections derived from air-broadened spectra recorded in the laboratory. Fig. 1 provides a plot of the new absorption cross section at 208.0 K and 7.501 Torr with these main band systems labelled. Full details on the measurement conditions and derivation of this cross section are given in Section 3.

2.2. A brief history of carbon tetrachloride absorption cross sections

The HITRAN 1986 compilation [14] included for the first time high resolution (0.03 cm$^{-1}$) cross sections for the simulation of the spectra of a number of important atmospheric molecules, including CCl$_4$, for which no line parameters were available. These cross sections, with a quoted accuracy of 10–25%, were derived from laboratory absorption spectra of pure samples recorded at 296 K at the University of Denver [15]. The first published temperature-dependent CCl$_4$ cross sections appeared in 1992 [16]. Derived from measurements of pure CCl$_4$ at 0.05 cm$^{-1}$ resolution and 223, 248, 273 and 298 K, these data were later included in the HITRAN 1996 compilation [17].

N$_2$-broadened CCl$_4$ absorption cross sections between 750 and 812 cm$^{-1}$, derived from measurements over a range of temperatures down to 208 K at a spectral resolution of 0.03 cm$^{-1}$ [18], henceforth referred to as the Nemtchinov dataset, were subsequently included in HITRAN 2000 [19]. This dataset has been used extensively for remote-sensing applications over the last decade and a half. This dataset has remained unchanged for subsequent HITRAN compilations, including the most recent HITRAN 2012 [20], and has additionally been included in the most recent GEISA 2003 [21] and 2009 [22] compilations. Despite its widespread use, the Nemtchinov dataset has a number of deficiencies which will be fully discussed in Section 3. Such deficiencies in the underlying laboratory spectroscopy compromise the accuracy of retrieved quantities, such as volume mixing ratios (VMRs), in remote-sensing applications. The new spectroscopic dataset described in the present work provides an improvement over previous available datasets.

3. New absorption cross sections of air-broadened carbon tetrachloride

3.1. Experimental

The experimental setup and procedures have been used previously for related measurements (e.g. [23,24]), so only brief details are provided here. All measurements were performed at the Molecular Spectroscopy Facility (MSF), Rutherford Appleton Laboratory (RAL), using a Bruker Optics IFS 125 HR FTS, and an internally mounted 26-cm-pathlength sample cell connected to a Julabo F95-SL Ultra-Low Refrigerated Circulator filled with ethanol. The cell temperature was monitored by four platinum resistance thermometers (PRTs) in thermal contact at different points on the exterior surface of the cell. Mixture pressures were measured using Baratron capacitance manometers (MKS). Samples were provided by Fluka (CCl$_4$; analytical standard, Z$_{99.95\%}$ purity, natural-abundance isotopic mixture) and BOC Gases (dry synthetic air or ‘Air Zero’, total hydrocarbons < 3 ppm, H$_2$O < 2 ppm, CO$_2$ < 1 ppm, CO < 1 ppm). The carbon tetrachloride sample (liquid at room temperature) was freeze-pump-thaw purified multiple times prior to use, whereas the synthetic air was used ‘as is’. Air-broadened CCl$_4$ IR spectra were recorded over a range of pressures and temperatures, and at resolutions between 0.01 and 0.03 cm$^{-1}$ (defined as the Bruker instrument resolution of 0.9/MOPD; MOPD = maximum optical path difference), 0.01 cm$^{-1}$ for the lowest...
pressures (in the Doppler-limited regime), and 0.03 cm$^{-1}$ for the highest pressures. Pure nitrous oxide (N$_2$O) spectra were additionally recorded at each temperature to calibrate the wavenumber scale of the air-broadened CCl$_4$ spectra. The FTS instrumental parameters and settings are summarised in Table 1, with sample pressures, temperatures, and their experimental uncertainties and associated spectral resolutions listed in Table 2.

### 3.2. Generation of absorption cross sections

The procedure for generating absorption cross sections from experimental data has been reported previously (e.g. [23,24]). After generation of transmittance spectra from measured interferograms, the wavenumber scale is calibrated against the positions of isolated N$_2$O absorption lines taken from the HITRAN 2012 database [20]. Next, using a successful approach developed previously (e.g. [23,24]), absorption cross sections are derived and calibrated against a “calibration standard” integrated band strength. This is necessary to counter problems with carbon tetrachloride adsorption in the vacuum line and on the cell walls, resulting in its partial pressure during each measurement differing from the initial, measured value.

Previous absorption cross sections created for ACE-FTS retrievals have had their integrated band strengths calibrated using data from the Pacific Northwest National Laboratory (PNNL) IR database (http://nwir.pnl.gov) [25]. However, data from the National Institute of Standards and Technology (NIST) Quantitative Infra-red Database (http://webbook.nist.gov/chemistry/quant-ir/) [26] are also suitable as calibration standards. Surprisingly, it was found that for carbon tetrachloride, the NIST integrated band strength (between 700 and 860 cm$^{-1}$) is 5.4% higher than that for PNNL. The main difference between the CCl$_4$ spectra in the two databases is that the NIST measurements were made using flow-through samples in a White cell, with the PNNL measurements employing a static sample cell. For the purposes of the ACE-FTS retrievals in Section 4, it was found that calibrating to the NIST intensities resulted in a better agreement with ground-based measurements. Therefore, the absolute intensity of the absorption cross sections, $\sigma(\nu, P_{\text{air}}, T)$, at wavenumber $\nu$ (cm$^{-1}$), temperature $T$ (K) and synthetic air pressure $P_{\text{air}}$, was calibrated according to

$$\int_{700 \text{ cm}^{-1}}^{860 \text{ cm}^{-1}} \sigma(\nu, P_{\text{air}}, T)d\nu = 6.7163 \times 10^{-17} \text{ cm molecule}^{-1},$$

where the value on the right hand side is the integrated band intensity over the spectral range 700–860 cm$^{-1}$ for the composite 760-Torr-N$_2$-broadened CCl$_4$ NIST spectrum at 296 K.

### 3.3. Absorption cross section uncertainties

The wavenumber accuracy, i.e. uncertainty in $\nu$, of the new absorption cross sections is comparable to the accuracy of the N$_2$O lines used in the calibration; HITRAN error codes indicate this is between 0.001 and 0.0001 cm$^{-1}$. Since only one spectrum is recorded at each PT combination, due to time constraints, it is not possible to obtain an estimate of the $y$-axis random errors. However, systematic errors are believed to make the dominant contribution to the uncertainty in $y$. Maximum uncertainties in the sample temperatures ($\mu_T$) and total pressures ($\mu_P$) are 0.2% and 0.8%, respectively (Table 2). The photometric uncertainty ($\mu_{\text{phot}}$) is estimated to be $\sim 2\%$. The pathlength uncertainty ($\mu_{\text{path}}$) is estimated to be negligibly small, lower than 0.1%. The systematic error, $\mu_{\text{NIST}}$, in the NIST CCl$_4$ spectrum used for intensity calibration is 2.1%. Assuming that the error estimates for all quantities are uncorrelated, the overall systematic error in the dataset
can be calculated from:

\[ \mu_{\text{systematic}} = \mu_{\text{NIST}}^2 + \mu_T^2 + \mu_P^2 + \mu_{\text{phot}}^2. \]  

(2)

Note that using a NIST spectrum for intensity calibration effectively nullifies the errors in the \( \text{CCl}_4 \) partial pressures and cell pathlength, so these do not have to be included in Eq. (2). According to Eq. (2), the systematic error contribution, \( \mu_{\text{systematic}} \), to the new cross sections is \( \sim 3\% \).

3.4. Comparison between absorption cross-section datasets

This section provides a comparison between the new dataset presented as part of this work and the older Nemtchinov dataset. This comparison will focus on the quality of the datasets in terms of their spectral resolution, signal-to-noise ratio, integrated band strength, wavenumber scale, and PT coverage. This new dataset is available electronically from the first author, and will be made available to the community via the HITRAN and GEISA databases. Preliminary retrievals from the ACE-FTS instrument using the new spectroscopic dataset are presented in Section 4.

3.4.1. Spectral resolution

All spectra used to create the Nemtchinov cross-section dataset were recorded at 0.03 cm\(^{-1}\) spectral resolution. While such a resolution is appropriate for high-pressure measurements, it leaves much of the fine structure underresolved when the total pressure of the sample mixture is low. This is well illustrated in Figs. 2 and 3, which compare selected spectral ranges of the new cross section at 208.0 K and 7.501 Torr with the Nemtchinov cross section at 207.9 K and 8.22 Torr. For the new measurements, resolutions between 0.01 and 0.03 cm\(^{-1}\) were used, depending on the total pressure of the mixture gas.

3.4.2. Signal-to-noise ratios (SNRs)

The SNRs of the new transmittance spectra, calculated using Bruker’s OPUS software at \( \sim 840 \text{ cm}^{-1} \) where the transmittance is close to 1, range from 800–1600 (rms). A direct comparison between the SNRs of the two absorption cross-section datasets is somewhat misleading because of the incorrectly chosen spectral resolution used for a number of Nemtchinov measurements. Whereas the SNRs are comparable for all the 0.03 cm\(^{-1}\) measurements, the new measurements at 0.01 cm\(^{-1}\) are noisier than the Nemtchinov measurements at 0.03 cm\(^{-1}\) by at most a factor of two. This has minimal impact for remote-sensing applications, however, given the overall high SNR of the new dataset.

3.4.3. Integrated band strengths

In order to compare integrated band strengths between datasets, integrals have been calculated over the spectral range of the Nemtchinov cross-section files, 750–812 cm\(^{-1}\). This wavenumber range, however, does not extend far enough to obtain a true measure of the baseline position for comparison with the new dataset. Fig. 4 is a plot of integrated band strength (without error bars for clarity) against temperature for each dataset. The Nemtchinov integrated band strengths display a spread in values, likely resulting from baseline inconsistencies in these cross sections. The apparent temperature dependence in the new measurements is an artefact caused by the narrower spectral range of the integrals. When calculated over the range 700–860 cm\(^{-1}\), integrated band strengths for the new cross sections correspond to \( 6.7163 \times 10^{-17} \text{ cm molecule}^{-1} \), as given in Eq. (1), for all temperatures.

3.4.4. Wavenumber scale

As with other absorption cross-section datasets produced from the same laboratory, e.g. HFC-134a [27], CFC-12 [24], and HCFC-22 [28], the wavenumber scale for the Nemtchinov dataset is not accurate; it appears not to have been calibrated. The dataset presented in this work has been calibrated against the positions of isolated \( \text{N}_2\text{O} \) absorption lines in the HITRAN 2012 database [20]. The difference in wavenumber scales is illustrated in Figs. 2 and 3, which compare the new cross section at 208.0 K and 7.501 Torr with the Nemtchinov cross section at 207.9 K and 8.22 Torr. In this case the Nemtchinov cross section is shifted too low by \( \sim 0.010 \text{ cm}^{-1} \) (a correction factor of \( \sim 1.000013 \)).

3.4.5. Pressure–temperature coverage

An absorption cross-section dataset used in remote sensing should cover all possible combinations of pressure and temperature appropriate for the region of the atmosphere being observed; it is more accurate to interpolate between cross sections rather than extrapolate beyond them. Fig. 5 provides a graphical representation of the PT combinations for both datasets. It shows clearly that the

![Fig. 2. The new IR absorption cross section of carbon tetrachloride/dry synthetic air (part of the \( \nu_1 + \nu_4 \) band) at 208.0 K and 7.501 Torr, with that of Nemtchinov [18] at 207.9 K and 8.22 Torr overlaid.](image-url)
new dataset presented in this work greatly extends the PT coverage (26 PT combinations in total). Ideally, the dataset should cover temperatures down to 190 K; however, as noted by Nemtchinov and Varanasi [18], due to the sharp drop-off in CCl4 vapour pressure below 208 K, it is not possible to record a spectrum below this temperature using a 26-cm-pathlength sample cell.

4. New ACE-FTS retrieval

As touched on in the Introduction, the ACE-FTS v3.0/v3.5 CCl4 retrieval is biased high by up to ~20–30% relative to ground measurements. Note that the CCl4 retrieval scheme is identical for the v3.0 and v3.5 datasets, the difference being in the meteorological data used as input for the pressure and temperature retrievals (the lowest ACE-FTS levels use these data directly). Due to an error in these inputs, v3.0 data should only be used for measurements taken until the end of September 2010, while v3.5 is valid for all ACE-FTS measurements. This section outlines an improved, preliminary ACE-FTS CCl4 retrieval, which addresses the problem of the high bias. The retrieval is based on version 3.5 of the ACE-FTS retrieval software [29], with new CCl4 spectroscopy (this work), updated microwindows, additional interfering species, and a new ILS [30] designed for the upcoming v4.0 processing. (The use of this new ILS, instead of that for v3.0/v3.5, reduces systematic errors in the retrieved VMRs arising from the ILS contributions to the residuals.)

The new microwindow set (listed in Table 3) consists of 35 microwindow 'slices' across the broad CCl4 spectral feature, with a common set of baseline parameters (a scaling factor and a slope for the baseline are included in the least-squares analysis). These slices were chosen so as to avoid regions of bad spectral residuals, for instance near the 792 cm\(^{-1}\) Q branch of CO2 (11101-10002 transition), which suffers from strong line mixing effects that the software is not currently equipped to calculate [6], and near numerous H2O lines, where current spectroscopic line parameters are unable to provide the required accuracy. The altitude limits of the microwindows vary with latitude according to the phenomenological expression \(\sin^2(\text{latitude})\), to reflect the latitude dependence at lower altitudes of the saturation of spectral features in the microwindows, and at upper altitudes of the variation in CCl4 abundance. An additional eight microwindows are included to improve the retrievals of various interferers, because the least-squares process can prematurely converge before reaching the optimal solution if the information content for one or more of the fitting parameters is too low.

The ACE-FTS v3.5 retrieval procedure has been explained previously [29]. In general, vertical VMR profiles of trace gases (along with temperature and pressure) are derived from the recorded transmittance spectra via a nonlinear least-squares global fit to the selected spectral region(s) for all measurements within the altitude range of
interest. For the CCl₄ retrievals carried out for this work, the atmospheric pressure and temperature profiles and the tangent heights of the measurements were taken from the v3.5 processing of the ACE-FTS data. The abundances of most molecules with absorption features in the microwindows (see Table 4) were adjusted simultaneously with the CCl₄ VMR. A weak contribution for CFC-13 (CCIF3), which is not listed in Table 4 as its VMR was not adjusted during the retrieval, was calculated based on abundances measured via in-situ studies. Spectroscopic line parameters and absorption cross sections for most molecules were taken from the HITRAN 2012 database [20], with the exception of HCFC-22 [28], CFC-113 [31], and CHCl₃ (PNL). An ACE-FTS transmittance spectrum (from occultation ss8706) over the range 772–812 cm⁻¹ is plotted in Fig. 6, along with several residual plots which provide an indication of the retrieval quality.

For the purposes of this work, a subset of 527 ACE-FTS occultations, covering March and April 2005 and providing a good coverage over all latitudes, was processed by the retrieval software. Fig. 7 is a plot of median VMR profiles in each of nine 20° latitude bins (from south to north, each bin contains 63, 64, 37, 45, 63, 74, 55, 69 and 57 profiles respectively); errors at each altitude are calculated as ± 1 MAD (median absolute deviation). Overlaid on each plot is the corresponding v3.0/v3.5 median profile (without error
Table 4
Summary of the molecules included in the microwindows for the new ACE-FTS CCl₄ retrieval.

<table>
<thead>
<tr>
<th>Molecule</th>
<th>Lower altitude limit (km)</th>
<th>Upper altitude limit (km)</th>
<th>Nature of spectroscopic data</th>
</tr>
</thead>
<tbody>
<tr>
<td>ClONO₂</td>
<td>8 – 2sin²(latitude°)</td>
<td>30 – 5sin²(latitude°)</td>
<td>Cross sections</td>
</tr>
<tr>
<td>CHClF₂ (HCFC-22)</td>
<td>8 – 2sin²(latitude°)</td>
<td>20</td>
<td>Cross sections</td>
</tr>
<tr>
<td>C₂Cl₃F₃ (CFC-113)</td>
<td>8 – 2sin²(latitude°)</td>
<td>20</td>
<td>Cross sections</td>
</tr>
<tr>
<td>H₂¹⁷O</td>
<td>8 – 2sin²(latitude°)</td>
<td>21</td>
<td>Line parameters</td>
</tr>
<tr>
<td>C₃H₆ONO₂ (PAN)</td>
<td>8 – 2sin²(latitude°)</td>
<td>20</td>
<td>Cross sections</td>
</tr>
<tr>
<td>HO₂NO₂</td>
<td>8 – 2sin²(latitude°)</td>
<td>30 – 5sin²(latitude°)</td>
<td>Cross sections</td>
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<td>8 – 2sin²(latitude°)</td>
<td>22</td>
<td>Line parameters</td>
</tr>
<tr>
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<td>30 – 5sin²(latitude°)</td>
<td>Line parameters</td>
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</tr>
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<td>Line parameters</td>
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<tr>
<td>CH₂₄</td>
<td>8 – 2sin²(latitude°)</td>
<td>21</td>
<td>Line parameters</td>
</tr>
</tbody>
</table>

Fig. 6. Top panel: an ACE-FTS transmittance spectrum over the 772–812 cm⁻¹ region for occultation ss8706 (recorded on 25 March 2005, off the coast of northern Scotland) at a tangent height of 8.60 km. The features in red represent the microwindows used in the new CCl₄ retrieval. Middle panel: the total observed - calculated residuals for the CCl₄ retrieval. Bottom panel: Observed/calculated ratio (without the inclusion of CCl₄ in the forward model), with the calculated CCl₄ transmittance contribution to the measurement overlaid. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
bars for clarity) for the same occultations. Unavoidably, by reducing the total wavenumber coverage of combined microwindows (the v3.0/v3.5 retrieval utilises a single CCl4 microwindow of 787.50–805.50 cm⁻¹), the VMRs retrieved using the new scheme are noisier than was observed with v3.0/v3.5, however they are certainly more accurate; this accuracy can be assessed by comparison with ground measurements from early 2005. Based on data from the 2010 ozone assessment report [32], in particular Figure 1–1, mean global surface mixing ratios of CCl4 at the time of the ACE-FTS measurements range from 92–95 ppt. With an atmospheric lifetime of 44 years [33] due to stratospheric loss, meaning that CCl4 is well-mixed in the troposphere, these surface VMRs should be representative of those in the free troposphere. As shown by Fig. 7, these values compare well with retrieved ACE-FTS VMRs in the upper tropical troposphere (30°S and 30°N), where the air is relatively young. (Note that if the cross sections had been scaled using PNNL spectra, the retrieved VMRs would be on average 5.4% higher, and agreement with ground measurements worse.) However, there is an unphysical ‘kink’ in the upper tropospheric profile between 10°S and 10°N. It is possible that this is caused by strong water absorption contaminating the selected retrieval microwindows; water is typically much more abundant in the tropics compared to other regions. This will be investigated prior to the upcoming v4.0 processing.

The improvement in retrieval accuracy relative to v3.0/v3.5 can be attributed directly to the new cross sections and the changes made to the retrieval scheme. Disentangling the individual contributions to the systematic differences between profiles for the two schemes is beyond the scope of this work, however based purely on integrated band strengths the new CCl4 cross sections lead to 8% lower VMRs. It is likely that the new v4.0 ILS makes only a relatively small contribution of several percent to the improvement. The sizes of these systematic differences vary for different profiles, depending on, for example, the concentrations of interferers that are either missing from the forward model or represented by inadequate spectroscopy (in v3.0/v3.5), as well as the differences between the interpolated CCl4 cross sections used for the two schemes. Since a number of the interferers inadequately characterised in the v3.0/v3.5 retrieval have larger concentrations in the troposphere, e.g. PAN which was not included in v3.0/v3.5, the systematic differences are larger in the troposphere.
Median profiles for the 527 ACE-FTS CCl4 VMR profiles assigned into 10° latitude bins have been used to construct a CCl4 zonal median VMR cross-section plot for March/April 2005 (Fig. 8); this provides an alternative means of viewing the data from Fig. 7. Those data points within the tropical troposphere (fewer than ten points in total) which deviate significantly from neighbouring values were removed and replaced by averages of these neighbouring VMRs. The plot in Fig. 8 illustrates clearly that CCl4 VMRs decrease more rapidly with increasing altitude in the high-latitude/polar regions, reflecting the large-scale circulation of air around the globe, in which tropospheric air ascends into the stratosphere over the tropics and descends into the troposphere towards the poles.

The new retrieval scheme described in this work will form the basis for the CCl4 retrieval in the upcoming processing version 4.0 of ACE-FTS data. This processing version will feature a new pressure and temperature retrieval, which will further improve the CCl4 data product and provide more accurate trends.

5. Conclusions

New high-resolution IR absorption cross sections for air-broadened carbon tetrachloride have been determined over the spectral range 700–860 cm⁻¹, with an estimated uncertainty of ~3%. Spectra were recorded for mixtures of CCl4 with dry synthetic air in a 26-cm-pathlength cell at spectral resolutions between 0.01 and 0.03 cm⁻¹ (calculated as 0.9/MOPD) over a range of temperatures and pressures appropriate for upper troposphere – lower stratosphere conditions (7.5–760 Torr and 208–296 K). Intensities were calibrated against a CCl4 spectrum in the NIST IR database. These new cross sections improve upon those currently available in the HITRAN and GEISA databases; namely they cover a wider range of pressures and temperatures, they have a more accurately calibrated wavenumber scale, they have more consistent integrated band intensities, and they were derived from spectra recorded at appropriate spectral resolutions.

It has been demonstrated that the new absorption cross sections, coupled with improvements in the microwindow selection, the addition of new interfering species, and a new ILS, lead to a more accurate ACE-FTS retrieval than v3.0/v3.5. It is important to note that the improvements made to the CCl4 retrieval scheme arise predominantly from spectroscopic considerations. The revised CCl4 absorption cross sections contribute about an 8% reduction in the bias relative to v3.0/v3.5 (based on a simple comparison of integrated band strengths). The rest of the bias has been reduced by the careful selection of microwindows based on sound spectroscopic judgement, and ensuring that spectral regions in which the line parameters of interfering species do not adequately calculate the measured ACE-FTS spectra are not included in the retrieval. A similar approach can be extended to CCl4 retrievals from atmospheric IR spectra recorded by other remote-sensing instruments, e.g. MIPAS. The preliminary scheme outlined in this work will form the basis for the upcoming processing version 4.0 of ACE-FTS data.

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