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The rotationally-resolved absorption spectrum of formal dehyde from 6547 to 6804 $\rm cm^{-1}$

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Abstract

The room temperature absorption spectrum of formaldehyde, H₂CO, from 6547 to 6804 cm⁻¹ (1527 to 1470 nm) is reported with a spectral resolution of 0.001 cm⁻¹. The spectrum was measured using cavity-enhanced absorption spectroscopy (CEAS) and absorption cross-sections were calculated after calibrating the system using known absorption lines of H₂O and CO₂. Several vibrational combination bands occur in this region and give rise to a congested spectrum with over 8000 lines observed. Pressure broadening coefficients in N₂, O₂, and H₂CO are reported for an absorption line at 6780.871 cm⁻¹, and in N₂ for an absorption line at 6684.053 cm⁻¹.

Key words: Formaldehyde, H_2CO , near infrared spectrum, combination bands, cavity enhanced absorption spectroscopy (CEAS), pressure broadening, high resolution

1 Introduction

Formaldehyde has a rich chemistry in the atmosphere, where it occurs as an intermediate species in the oxidation of alkanes and alkenes, nitric oxide, and dimethyl sulphoxide [1]. The photolysis of formaldehyde, H_2CO :

 $H_2CO + h\nu \rightarrow H + HCO$

and the subsequent formation of the hydroperoxyl radical, HO_2 , in the presence of oxygen:

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Number	Symmetry	$\tilde{\nu} \; [{\rm cm}^{-1}]$	Mode
$ u_1 $	a_1	2782.5	CH_2 symmetric stretch
$ u_2 $	a_1	1746.0	CO stretch
$ u_3$	a_1	1500.2	CH_2 scissor
$ u_4$	b_1	1167.3	CH_2 wag
$ u_5 $	b_2	2843.3	CH_2 asymmetric stretch
$ u_6 $	b_2	1249.1	CH ₂ rock

Table 1 Transition energies for the six normal vibrational modes of H_2CO [12].

 $\mathrm{HCO} + \mathrm{O}_2 \mathop{\rightarrow} \mathrm{CO} + \mathrm{HO}_2$

 $\mathrm{H} + \mathrm{O}_2 \rightarrow \mathrm{HO}_2$

play a key role in oxidation chain reactions. The importance of these reactions has encouraged efforts to improve the spectroscopic data on H_2CO and HO_2 for use in field studies and laboratory experiments [2–8].

Various spectroscopic methods have been employed to measure the gas-phase absorption spectrum of H₂CO. To increase the sensitivity of the measurement, either a multipass cell, or cavity-enhanced spectroscopic methods such as cavity ring-down spectroscopy (CRDS) and cavity enhanced absorption spectroscopy (CEAS) [9,10], have typically been used. Barry et al. reported the $2\nu_5$ overtone band (around 5676 cm⁻¹) using CEAS [2], and have shown that the results from CEAS closely match those from CRDS [11]. For applying absorption spectroscopy to trace measurements of H₂CO, the stronger fundamental vibration bands have usually been used. For instance, Fried and co-workers used a tunable diode laser spectrometer (TDLAS) system to carry out airborne measurements of H₂CO using the 2831 cm⁻¹ absorption line [3]. They demonstrated a detection limit of 55 ppt with an integration time of 20 s. Dahnke et al. have achieved a 2 ppb sensitivity to H₂CO with cavity leak-out spectroscopy at 2853 cm⁻¹ [8].

In this paper we report the H_2CO absorption spectrum from 6547 to 6804 cm⁻¹. The H_2CO molecule has C_{2v} symmetry and six normal modes (Table 1), with several combination bands between 6500 and 6830 cm⁻¹ (Table 2). These transitions have been observed with a 1 cm⁻¹ resolution by Bouwens et al. in a jet experiment using dispersed fluorescence spectroscopy from a number of vibrational bands in the S_1 excited state [13]. The H_2CO combination bands consist of three to five normal modes. A sensitive spectroscopic technique is therefore necessary to measure the absorption of these combination bands' rovibrational lines. In this work we used CEAS to record the H_2CO spectrum.

Although the weak H_2CO absorption lines from 6547 to 6804 cm⁻¹ are not

Table 2

Combination	Symmetry	$\tilde{\nu}$
band		$[\mathrm{cm}^{-1}]$
$3_24_26_1$	b_2	6508.8
$1_1 3_1 4_2$	a_1	6562.7
$2_14_26_2$	a_1	6578.8
$1_2 4_1$	b_1	6611.6
$3_1 4_2 5_1$	b_2	6635.7*
$2_{3}3_{1}$	a_1	6652.2
$2_1 3_1 4_3$	b_1	6693.4*
$3_14_15_16_1$	b_1	6710.0
$3_15_16_2$	b_2	6759.0
$2_1 3_1 4_2 6_1$	b_2	6777.6*
$4_{1}5_{2}$	\mathbf{b}_1	6795.1
$3_{3}4_{2}$	a_1	6815.2
$1_1 2_1 4_2$	b_1	6825.5

Combination bands between 6500 and 6830 cm^{-1} from the dispersed fluorescence spectra of Bouwens et al. [13]. Tentatively identified combination bands observed in our spectrum are indicated with asterisks.

suited for field observations of H_2CO , they should be useful in laboratory studies and database development. We note that recent efforts to calculate the vibrational and rotational constants of HO_2 from its $2\nu_1$ spectrum (between 6603 and 6686 cm⁻¹) have been hampered by ambiguities in the line positions, possibly arising from interfering H_2CO lines [4]. Pressure broadening coefficients in N₂, O₂, and H₂CO are reported for an absorption line at 6780.871 cm⁻¹, and in N₂ for an absorption line at 6684.053 cm⁻¹. The integrated cross-sections of these lines are also reported.

2 Experimental

CEAS is based on measuring the intensity of light transmitted through a high finesse optical cavity as a function of wavelength [10,14]. In comparison, in CRDS the decay time of light in an optically stable resonator is the desired parameter. For small losses per pass, the absorption coefficient, α , of the sam-

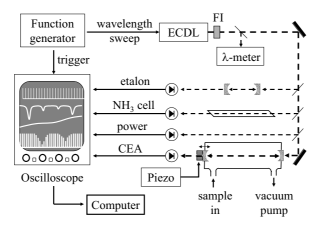


Fig. 1. Diagram of the CEAS system showing the beam path (dashed) from the external cavity diode laser (ECDL), through a Faraday isolator (FI), and through the sample cavity to give the cavity-enhanced absorption (CEA) signal. Before the sample cavity, small fractions of the beam intensity are split off to record the relative power and to calibrate the wavelength scale.

ple is adequately described by [15]:

$$\alpha \approx \frac{1}{d} \left(\frac{I_0}{I} - 1 \right) (1 - R) \tag{1}$$

where d is the cavity length, R is the mirror reflectivity, and I and I_0 are, respectively, the intensity transmitted through the cavity with and without a sample present. CEAS is $(1 - R)^{-1}$ times more sensitive than a single-pass absorption method assuming that the achievable signal-to-noise ratio remains the same for both approaches [16]. CEAS does not yield absorption crosssections directly; the wavelength-dependent mirror reflectivities must be either known or calibrated against a spectrum of known absorption cross-section.

Our experimental setup is shown schematically in Fig. 1. The light source was a temperature-stabilized external cavity diode laser (Sacher Lasertechnik, TEC 500) in a Littman configuration. The laser power was ≈ 3 mW and the bandwidth was ~ 1 MHz. The total tuning range of the system was between 6547 and 6804 cm⁻¹ (corresponding to 1527 and 1470 nm). The spectrum we present here covers this entire spectral range, except for two gaps near the red edge of the range (from 6549.7 to 6551.3 cm⁻¹ and 6553.5 to 6554.5 cm⁻¹) where stable laser operation was not obtained.

The laser was linearly scanned over a range of 2 cm^{-1} (referred to as a "spectral segment") at approximately 40 Hz and synchronized with the oscilloscope. The laser beam was coupled into an 85 cm cavity formed by two spherical mirrors of 5 m radius of curvature and a maximum reflectivity of ~0.996. An optical isolator was placed after the laser output aperture to reduce feedback

into the laser and associated laser instability. The distance from the laser to the cavity was also made as large as possible (2.5 m) for the same reason. The light transmitted through the cavity was detected with an InGaAs photodiode (NewFocus Nirvana 2017). For each 2 cm⁻¹ segment of the spectrum, 1000 sweeps were collected and averaged with a digital oscilloscope (LeCroy Waverunner LT 264). The exposure time was 10 s at a sampling rate of 4 Hz, limited by our oscilloscope. The sampling interval of 0.0008 cm⁻¹ was enough to resolve the H₂CO Doppler-broadened FWHM of about 0.015 cm⁻¹.

A key consideration in CEAS is to minimize the influence of the cavity's mode structure on the wavelength-dependent intensity of the transmitted light. This is achieved by continuously modulating the cavity length in order to shift the mode structure between sweeps. A ring-shaped piezoelectric transducer attached to the cavity's exit mirror was used for this purpose. We also adopted an off-axis configuration in order to excite a large number of high order transverse modes [17]. The laser was scanned at a rapid rate of 6 MHz μ s⁻¹ (0.2 cm⁻¹s⁻¹), which was significantly larger than the ratio of the cavity mode linewidth to the ring-down time (0.3 MHz μ s⁻¹ for the TEM₀₀ mode). Consequently, the light intensity in a given mode was far away from saturation. These strategies avoided the build up of significant light in any single cavity mode. The transmitted intensity per frequency interval (which is determined by the integration time) is then averaged over the number of sweeps per measurement. Discrete cavity modes were not observed in the transmitted light.

Small fractions of the beam intensity were split off before the cavity to calibrate the wavenumber scale and to measure the laser power. For the wavenumber calibration, the spectrum of NH_3 was recorded by passing a beam through a 60 cm cell filled with NH₃ at 10 mbar. Absorption peaks were matched to the NH_3 spectrum reported by Lundsberg-Nielsen et al. [18]. The spectral density of NH_3 lines is high enough over the entire spectrum presented here in order to obtain several absolute wavenumbers per spectral segment. To compensate for small deviations from linearly scanning the laser over a $\sim 2 \text{ cm}^{-1}$ spectral segment, the relative wavenumber spacing was also determined from a Fabry-Perot etalon with a free-spectral range of about 0.03 cm^{-1} . We estimate the uncertainty in our wavenumber scale to be approximately 0.001 cm^{-1} [19]. The etalon also served to identify mode-hops within the laser scanning range. When mode-hops were observed, the temperature of the laser diode was adjusted until the scanning region was mode-hop free. A third beam was used to normalize the laser power over the tuning range. These three signals were measured with InGaAs photodiodes (Thorlabs DET410) and recorded simultaneously with the cavity-enhanced absorption signal on the digital oscilloscope (see Fig 1).

 H_2CO was prepared by pyrolysis of paraformaldehyde (Aldrich, 95% pure) under vacuum. The gaseous H_2CO was first passed through a cooling trap below

200 K to remove water vapour and polymerization products of H₂CO. Water vapour catalyses the polymerization of H₂CO. The monomeric H₂CO was trapped and stored at 77 K under vacuum. The sample cavity was evacuated to approximately 10^{-6} mbar, ensuring that it was virtually free of gaseous water. H₂CO gas was introduced into the cavity by slowly heating the solid H₂CO in the cooling trap with the valve to the cavity opened. When the vapour pressure of the H₂CO reached the required value of 2 mbar the valve was closed and the solid H₂CO was cooled again for further use. The pressure in the cavity was measured with a capacitance manometer operating in the range of 10^{-2} to 120 mbar with a precision of 0.01 mbar. With 2 mbar of H₂CO in the sample cavity, the pressure decreased by about 0.1 mbar before stabilizing, whereas at pressures above 10 mbar the pressure dropped by 50% over a period of a few hours. Polymerization of H₂CO therefore appears to be negligible at 2 mbar. The temperature of the system was 291 ± 2 K.

3 Results and Discussion

3.1 Absorption spectrum

 I_0 , I, and the reflectivity spectrum of the mirrors, R, must be known to calculate the absorption coefficient of the sample with Eq. (1). Because the measurement of I_0 for every corresponding 2 cm⁻¹ segment of the spectrum is prohibitively time-consuming for a 250 cm⁻¹ spectral range, we adopted a pragmatic approach to estimate I_0 . In areas of the H₂CO spectrum where no absorption occurs, the transmitted intensities with and without a sample should be the same. Therefore, an estimated I_0 was obtained from a fourthorder polynomial fit to well-spaced maxima in the spectrum of I. The absorption cross-section, $\sigma = \alpha N^{-1}$, where N is the number density of the absorbing species, was calculated by using this estimated I_0 in Eq. (1).

The system was calibrated using the integrated cross-sections of H₂O and CO₂ lines from the HITRAN database to calculate R as a function of wavelength [20]. The corresponding calibration curves in Fig. 2 show that the integrated absorption of a transition varies linearly with the pressure, as expected. Furthermore, there is a minimal offset at the lowest pressures, suggesting that our approximation for I_0 does not introduce a large systematic error. From the H₂O and CO₂ absorption lines the mirror reflectivity, R, was determined as a function of wavelength as shown in Fig. 3. Points with large error bars were determined from single measurements, and those with small error bars from linear regressions of pressure dependent measurements as in Fig. 2. Because the reflectivity decreases towards the blue end of the spectrum, a second-order polynomial was fitted to (1 - R) as a function of wavelength. In the fit the

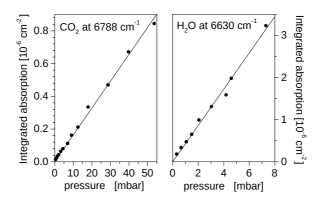


Fig. 2. Integrated absorption coefficient as a function of pressure. The left panel shows a CO_2 transition at 6788 cm⁻¹ and the right panel an H₂O transition at 6630 cm⁻¹. The slopes of these graphs were used to calculate the mirror reflectivities at these wavenumbers.

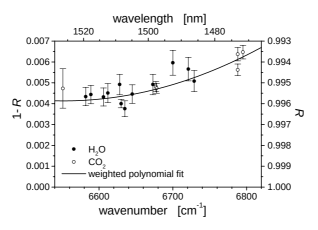


Fig. 3. Mirror reflectivity obtained using the integrated cross-sections of H_2O and CO_2 lines from HITRAN [20]. The curve is a weighted second-order polynomial fit to the experimental points and was used for the reflectivity.

different uncertainties of calibration measurements were accounted for. The overall systematic error of this calibration of the wavelength-dependent reflectivity was established to be $\pm 5\%$. The calculated values of R agree well with the manufacturer's specified maximum mirror reflectivity of 0.996 around 6500 cm⁻¹.

The individual segments of the spectrum were joined together to obtain the complete spectrum. The overlap of two typical spectral segments is shown in Fig. 4 and the general agreement between the absorption coefficients of such segments is very satisfactory. For the final spectrum, the overlap regions of adjacent spectral segments were averaged, or where one spectral segment was to be preferred for experimental reasons, the best measurement was chosen. The overall statistical uncertainty in the H₂CO absorption cross-section is estimated at $\approx 10\%$ over most of the spectrum. However, where the laser is at its tuning limit for a given laser mode, the absorption coefficients of adjacent

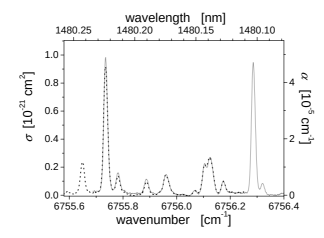


Fig. 4. Typical overlap of two adjacent spectral segments. The agreement in the wavenumber between two adjacent segments is usually within 0.001 cm⁻¹, and the difference in the absorption maxima is usually less than 10%.

spectral segments show poorer agreement. In these cases, which constitute less than 3% of the entire spectral range, we estimate the uncertainty in the absorption coefficients to be $\approx 25\%$. These regions are identified in the supplementary material [21]. The signal-to-noise ratio of the spectrum is better than 500. The minimum detectable absorption of our system is approximately 1.5×10^{-7} cm⁻¹ with a 10 s exposure time.

ν	σ	$\tilde{\nu}$	σ	$\tilde{\nu}$	σ
$[\mathrm{cm}^{-1}]$	$[10^{-21} \text{ cm}^2]$	$[\mathrm{cm}^{-1}]$	$[10^{-21} \text{ cm}^2]$	$[\mathrm{cm}^{-1}]$	$[10^{-21} \text{ cm}^2]$
6567.402	0.71	6652.368	0.70	 6739.425	1.04
6567.418	0.60	6653.443	1.07	6741.337	1.08
6576.402	0.59	6654.459	0.76	6741.819	1.00
6576.425	0.77	6654.802	0.55	6743.692	1.26
6592.086	0.57	6654.868	0.57	6744.231	1.22
6595.268	0.63	6655.834	1.11	6746.066	0.99
6596.641	0.53	6656.654	0.83	6746.640	1.38
6596.731	0.53	6657.316	0.64	6748.460	1.16
6598.324	0.85	6657.355	0.62	6749.057	1.09
6598.359	1.00	6658.252	0.84	6750.868	1.32
6600.261	0.97	6658.718	0.69	6751.471	1.14
6601.348	0.89	6659.809	0.69	6753.294	1.19
6602.345	1.00	6659.828	0.69	6753.884	1.10
6604.211	1.18	6660.637	0.52	6754.619	0.56
6604.509	1.00	6660.711	0.89	6755.735	0.97
6606.318	0.54	6662.285	0.84	6756.286	0.95
6606.489	0.51	6663.242	0.74	6758.194	0.79
6606.723	1.13	6664.739	1.10	6758.682	1.19

Table 3: H₂CO absorption lines between 6547 cm⁻¹ and 6804 cm⁻¹ with absorption cross-sections, σ , greater than 0.5×10^{-21} cm².

			- C	ontinued from	previous page
$\tilde{\nu}$	σ	$\tilde{\nu}$	σ	$\tilde{\nu}$	σ
$[cm^{-1}]$	$[10^{-21} \text{ cm}^2]$	$[cm^{-1}]$	$[10^{-21} \text{ cm}^2]$	$[cm^{-1}]$	$[10^{-21} \text{ cm}^2]$
6606.964	1.12	6667.186	0.95	6760.665	0.54
6607.740	0.61	6667.433	0.53	6761.070	0.56
6608.975	1.10	6669.390	0.55	6763.774	0.53
6609.622	1.35	6669.625	0.67	6766.298	0.64
6610.538	0.55	6670.321	0.51	6767.336	0.69
6611.256	1.39	6672.197	0.52	6775.121	0.59
6612.212	1.40	6681.776	1.67	6775.757	0.67
6613.559	0.83	6684.691	0.54	6777.411	0.87
6614.757	1.18	6685.278	0.53	6778.311	0.83
6615.879	1.06	6686.209	0.54	6779.693	1.04
6617.276	1.13	6693.920	0.77	6780.872	1.01
6618.209	1.06	6695.400	0.93	6781.956	1.29
6619.780	0.92	6696.342	0.74	6782.003	0.73
6620.550	0.89	6696.805	0.51	6783.441	0.97
6622.281	0.77	6697.739	0.58	6784.213	1.24
6622.896	0.76	6698.758	1.02	6786.020	1.09
6624.779	0.93	6701.165	1.15	6786.457	1.15
6625.248	0.53	6703.558	0.85	6788.603	1.37
6638.777	0.51	6705.933	0.59	6788.699	1.24
6638.804	0.52	6705.953	0.59	6790.930	0.91
6639.331	0.61	6708.654	0.55	6791.191	1.27
6639.845	0.62	6709.633	0.68	6793.169	1.23
6641.674	0.78	6710.621	0.51	6793.778	1.25
6642.355	0.72	6710.678	0.65	6795.409	1.32
6644.015	0.66	6712.921	0.53	6795.959	0.97
6644.854	0.91	6717.946	0.56	6796.368	1.23
6646.361	0.92	6727.917	0.55	6797.668	1.09
6647.331	0.83	6729.629	0.65	6798.804	0.54
6648.712	1.13	6730.104	0.80	6798.952	1.12
6649.264	0.51	6732.042	0.97	6799.771	0.67
6649.675	0.58	6732.369	0.82	6799.955	1.08
6649.769	0.94	6734.362	1.05	6801.522	0.83
6649.839	0.58	6734.689	1.06	6802.281	1.19
6651.069	0.94	6736.676	1.11	6803.447	0.53
6652.157	0.96	6737.044	0.99	6804.058	0.87
6652.260	0.63	6738.999	1.23	6804.362	0.57

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The complete absorption spectrum of H_2CO from 6547 to 6804 cm⁻¹ is shown in Fig. 5. The multiple transitions and overlapping rovibrational combination bands produce a congested spectrum, making it difficult to assign individual absorption lines. The rotational envelopes of the fundamental modes are

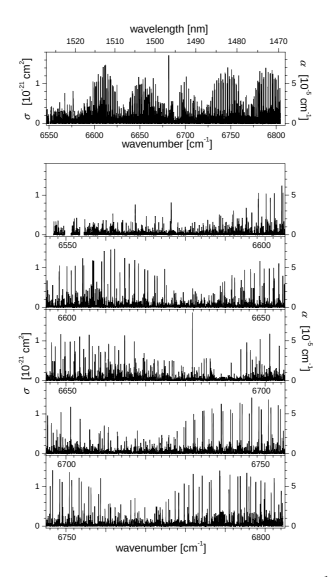


Fig. 5. The absorption spectrum of H_2CO from 6550 to 6800 cm⁻¹. The top panel shows the complete spectrum; the spectrum is broken into 50 cm⁻¹ sections in the lower panels.

slightly greater than 100 cm^{-1} wide at room temperature [22]; the width should be similar for the combination bands. Apparent rovibrational bands occur from 6590 to 6680 cm⁻¹, from 6720 to above 6800 cm⁻¹, and possibly from 6650 to 6730 cm⁻¹. These lines correspond most closely to the $3_14_25_1$ (6636 cm⁻¹), $2_13_14_3$ (6693 cm⁻¹), and $2_13_14_26_1$ (6778 cm⁻¹) combination bands reported by Bouwens et al. (also see Table 2). However, a comprehensive spectral analysis will be necessary to deconvolute the overlapping combination bands and to assign the vibrational frequencies with precision. Such an analysis is beyond the scope of this study. The strongest absorption lines in the spectrum (those with absorption cross-sections greater than 5×10^{-22} cm²) are listed in Table 3. This cut-off criterion results in 162 lines; the entire spectrum, in which over 8000 lines have been identified, is available in the supplementary

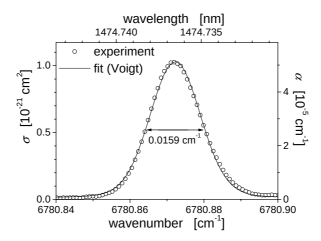


Fig. 6. Fit of a Voigt profile to the 6780 cm⁻¹ absorption line of pure H₂CO at 2 mbar. Fit parameters for Voigt: Gaussian FWHM=0.00151 cm⁻¹, Lorentzian FWHM=0.00015 cm⁻¹. Empirical FWHM=0.0159 cm⁻¹.

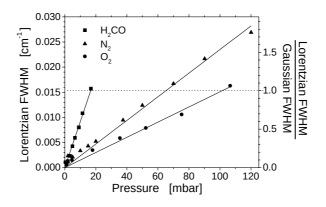


Fig. 7. Broadening of the 6780.871 cm⁻¹ line in N₂, O₂, and H₂CO, showing the Lorentzian FHWM (left axis) and the ratio of Lorentzian to Gaussian FWHM (right axis).

material [21]. The integrated absorption cross-section of the 6780.871 cm⁻¹ transition is 1.6×10^{-23} cm/molecule and that of the 6684.053 cm⁻¹ transition is 7.2×10^{-24} cm/molecule. The former transition is about 35 times weaker than the 5676.21 cm⁻¹ transition of the $2\nu_5$ overtone, and over 4000 times weaker than the ν_1 C–H stretch [2].

3.2 Pressure broadening

The pressure broadening of the 6780.871 cm⁻¹ line in N₂, O₂, and H₂CO, and of the 6684.053 cm⁻¹ line in N₂, was studied. For different pressures of pure H₂CO, Voigt profiles were fitted to the absorption lines using a non-linear least square method (Levenberg-Marquardt) [23]. From the Voigt fits the Lorentzian and Gaussian contributions to the overall FWHM were de-

Table 4

Transition	Gas	FWHM pressure
		broadening
$[\mathrm{cm}^{-1}]$		$[10^{-4} \text{cm}^{-1} \text{mbar}^{-1}]$
6780.871	$\rm H_2CO$	9.18 ± 0.18
6780.871	O_2	1.51 ± 0.05
6780.871	N_2	2.35 ± 0.05
6684.058	N_2	2.01 ± 0.10

Pressure broadening of H₂CO absorption lines at ≈ 291 K for the transitions at 6780.871 cm⁻¹ and 6684.053 cm⁻¹. The Gaussian contribution to the FWHM of 0.015 cm⁻¹ (determined at 2 mbar H₂CO) was fixed for each gas investigated.

termined. The measurements showed that at pressures <2 mbar the FWHM was entirely dominated by Doppler broadening. The corresponding value of 0.0151 cm^{-1} for the Gaussian width agreed with the laboratory temperature within the uncertainty of the Voigt fit $(8 \times 10^{-4} \text{ cm}^{-1})$. A Voigt fit to the experimental data for H_2CO at 2 mbar is shown in Fig. 6. The quality of this fit was slightly affected by an adjacent absorption line; however, the resulting additional uncertainty of the FWHM was negligible. For each series of pressure dependent measurements in N_2 and O_2 the first measurement was performed with pure H_2CO before adding the respective buffer gas. The Gaussian FWHM (0.015 cm^{-1}) was then fixed in Voigt fits of absorption measurements in N₂ and O_2 at higher pressures. The Lorentzian pressure broadening contribution to the FWHM increases linearly with pressure for all the gases, as shown in Fig. 7 for the line at 6780.871 cm⁻¹. The effect of the H_2CO pressure is much more marked than that of N_2 and O_2 . Table 4 lists the corresponding broadening parameters, which were obtained by linear regression of the Lorentzian widths versus pressure. The broadening parameters determined in N_2 for the two different absorption lines agree reasonably well. The small discrepancy can result from the fact that the 6780.871 cm^{-1} line is well isolated, whereas the line at 6684.058 cm^{-1} may have an underlying additional absorption feature, which can be attributed to the small asymmetry of the line. The errors stated in the table only refer to the quality of the linear fit. Finally, no pressure shift in the peak positions was observed within our experimental uncertainty.

4 Summary

We have measured the formaldehyde spectrum from 6547 to 6804 cm⁻¹ (1527 to 1470 nm) with a resolution of about 0.001 cm⁻¹ using CEAS. The absorption cross-section for the spectrum was estimated by comparison to the known

linestrengths of CO_2 and H_2O . The pressure broadening coefficients in N_2 have been determined for two absorption lines.

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Table 1 Transition energies for the six normal vibrational modes of H_2CO [12].

Number	Symmetry	$\tilde{\nu} \ [\mathrm{cm}^{-1}]$	Mode
$ u_1 $	a_1	2782.5	CH ₂ symmetric stretch
ν_2	a_1	1746.0	CO stretch
$ u_3$	a_1	1500.2	CH_2 scissor
$ u_4$	b_1	1167.3	CH_2 wag
$ u_5 $	b_2	2843.3	CH_2 asymmetric stretch
$ u_6 $	b_2	1249.1	CH ₂ rock

Table 2 $\,$

Combination	Symmetry	$\tilde{ u}$
band		$[\mathrm{cm}^{-1}]$
$3_24_26_1$	b_2	6508.8
$1_1 3_1 4_2$	a_1	6562.7
$2_14_26_2$	a_1	6578.8
1_24_1	b_1	6611.6
$3_1 4_2 5_1$	b_2	6635.7*
$2_{3}3_{1}$	a_1	6652.2
$2_1 3_1 4_3$	b_1	6693.4^{*}
$3_14_15_16_1$	b_1	6710.0
$3_1 5_1 6_2$	b_2	6759.0
$2_1 3_1 4_2 6_1$	b_2	6777.6*
$4_{1}5_{2}$	\mathbf{b}_1	6795.1
$3_{3}4_{2}$	a_1	6815.2
$1_1 2_1 4_2$	b_1	6825.5

Combination bands between 6500 and 6830 cm^{-1} from the dispersed fluorescence spectra of Bouwens et al. [13]. Tentatively identified combination bands observed in our spectrum are indicated with asterisks.

$\tilde{\nu}$	σ	$\tilde{ u}$	σ	$\tilde{ u}$	σ
$[\mathrm{cm}^{-1}]$	$[10^{-21} \text{ cm}^2]$	$[\mathrm{cm}^{-1}]$	$[10^{-21} \text{ cm}^2]$	$[cm^{-1}]$	$[10^{-21} \text{ cm}^2]$
6567.402	0.71	6652.368	0.70	6739.425	1.04
6567.418	0.60	6653.443	1.07	6741.337	1.08
6576.402	0.59	6654.459	0.76	6741.819	1.00
6576.425	0.77	6654.802	0.55	6743.692	1.26
6592.086	0.57	6654.868	0.57	6744.231	1.22
6595.268	0.63	6655.834	1.11	6746.066	0.99
6596.641	0.53	6656.654	0.83	6746.640	1.38
6596.731	0.53	6657.316	0.64	6748.460	1.16
6598.324	0.85	6657.355	0.62	6749.057	1.09
6598.359	1.00	6658.252	0.84	6750.868	1.32
6600.261	0.97	6658.718	0.69	6751.471	1.14
6601.348	0.89	6659.809	0.69	6753.294	1.19
6602.345	1.00	6659.828	0.69	6753.884	1.10
6604.211	1.18	6660.637	0.52	6754.619	0.56
6604.509	1.00	6660.711	0.89	6755.735	0.97
6606.318	0.54	6662.285	0.84	6756.286	0.95
6606.489	0.51	6663.242	0.74	6758.194	0.79
6606.723	1.13	6664.739	1.10	6758.682	1.19
6606.964	1.12	6667.186	0.95	6760.665	0.54
6607.740	0.61	6667.433	0.53	6761.070	0.56
6608.975	1.10	6669.390	0.55	6763.774	0.53
6609.622	1.35	6669.625	0.67	6766.298	0.64
6610.538	0.55	6670.321	0.51	6767.336	0.69
6611.256	1.39	6672.197	0.52	6775.121	0.59
6612.212	1.40	6681.776	1.67	6775.757	0.67
6613.559	0.83	6684.691	0.54	6777.411	0.87
6614.757	1.18	6685.278	0.53	6778.311	0.83
6615.879	1.06	6686.209	0.54	6779.693	1.04
6617.276	1.13	6693.920	0.77	6780.872	1.01
6618.209	1.06	6695.400	0.93	6781.956	1.29
6619.780	0.92	6696.342	0.74	6782.003	0.73
6620.550	0.89	6696.805	0.51	6783.441	0.97
6622.281	0.77	6697.739	0.58	6784.213	1.24
6622.896	0.76	6698.758	1.02	6786.020	1.09
6624.779	0.93	6701.165	1.15	6786.457	1.15
6625.248	0.53	6703.558	0.85	6788.603	1.37
6638.777	0.51	6705.933	0.59	6788.699	1.24
6638.804	0.52	6705.953	0.59	6790.930	0.91
6639.331	0.61	6708.654	0.55	6791.191	1.27
6639.845	0.62	6709.633	0.68	6793.169	1.23
	0.02	0.00.000	0.00	0.00.100	1.20

Table 3: H₂CO absorption lines between 6547 cm⁻¹ and 6804 cm⁻¹ with absorption cross-sections, σ , greater than 0.5×10^{-21} cm².

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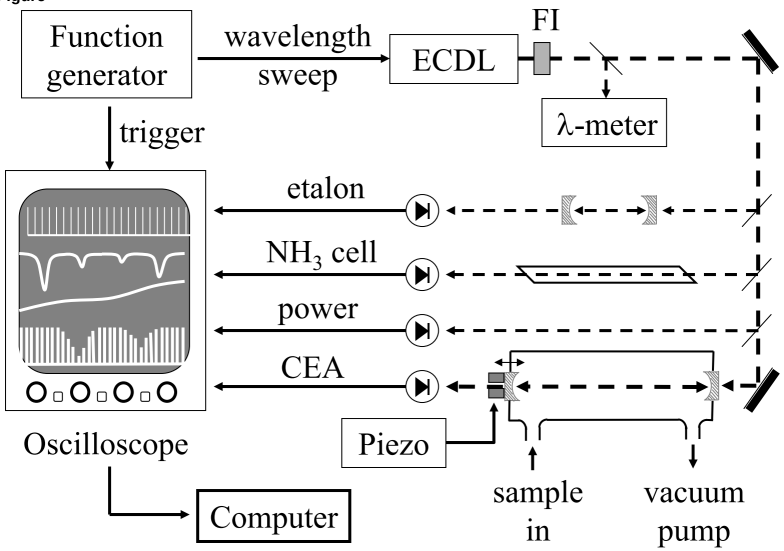
		- continued from previous page			
$\tilde{\nu}$	σ	$\tilde{ u}$	σ	$\tilde{\nu}$	σ
$[\mathrm{cm}^{-1}]$	$[10^{-21} \text{ cm}^2]$	$[\mathrm{cm}^{-1}]$	$[10^{-21} \text{ cm}^2]$	$[\mathrm{cm}^{-1}]$	$[10^{-21}~{\rm cm}^2]$
6642.355	0.72	6710.678	0.65	6795.409	1.32
6644.015	0.66	6712.921	0.53	6795.959	0.97
6644.854	0.91	6717.946	0.56	6796.368	1.23
6646.361	0.92	6727.917	0.55	6797.668	1.09
6647.331	0.83	6729.629	0.65	6798.804	0.54
6648.712	1.13	6730.104	0.80	6798.952	1.12
6649.264	0.51	6732.042	0.97	6799.771	0.67
6649.675	0.58	6732.369	0.82	6799.955	1.08
6649.769	0.94	6734.362	1.05	6801.522	0.83
6649.839	0.58	6734.689	1.06	6802.281	1.19
6651.069	0.94	6736.676	1.11	6803.447	0.53
6652.157	0.96	6737.044	0.99	6804.058	0.87
6652.260	0.63	6738.999	1.23	6804.362	0.57

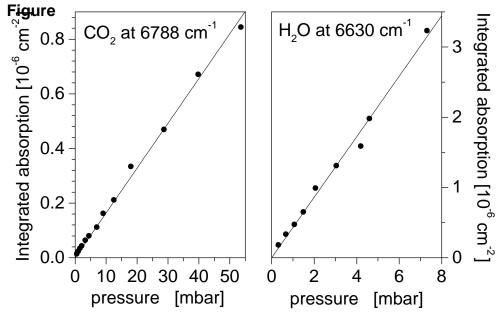
Table 4

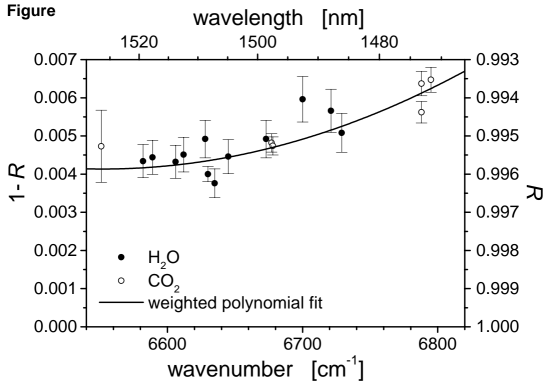
Pressure broadening of H₂CO absorption lines at ≈ 291 K for the transitions at 6780.871 cm⁻¹ and 6684.053 cm⁻¹. The Gaussian contribution to the FWHM of 0.015 cm⁻¹ (determined at 2 mbar H₂CO) was fixed for each gas investigated.

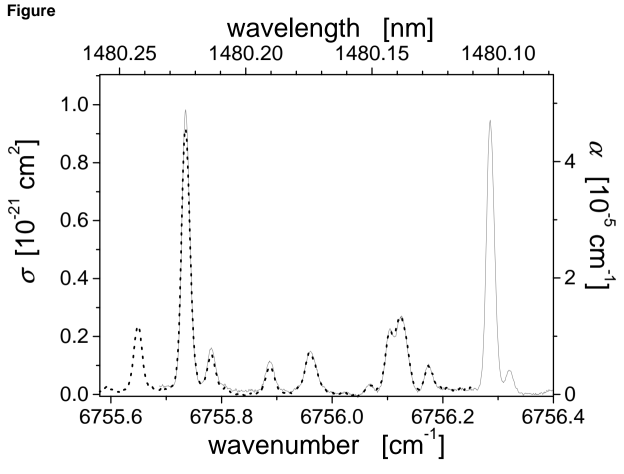
Transition	Gas	FWHM pressure
		broadening
$[\mathrm{cm}^{-1}]$		$[10^{-4} {\rm cm}^{-1} {\rm mbar}^{-1}]$
6780.871	$\mathrm{H}_{2}\mathrm{CO}$	9.18 ± 0.18
6780.871	O_2	1.51 ± 0.05
6780.871	N_2	2.35 ± 0.05
6684.058	N_2	2.01 ± 0.10

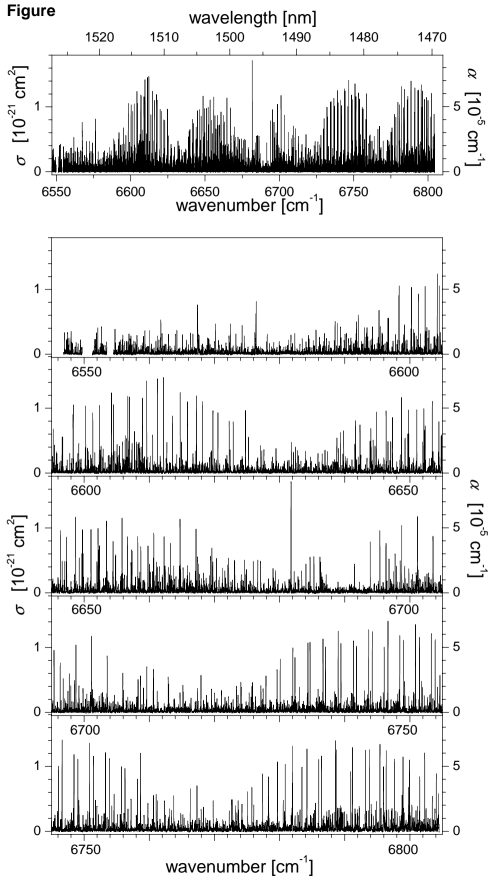


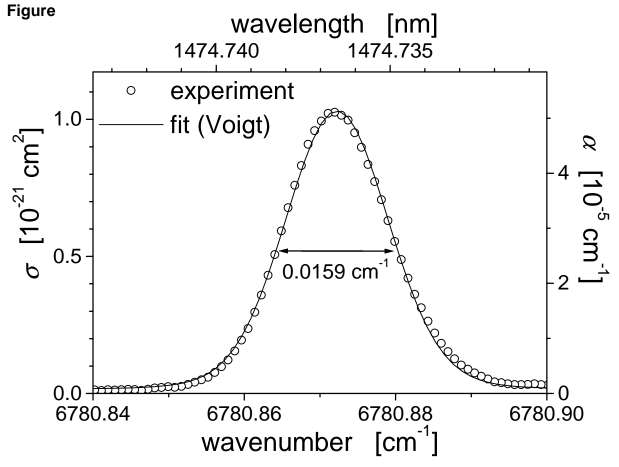


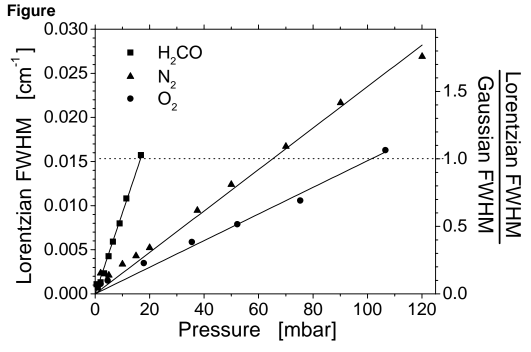












The supplementary material for the web consists of two additional files - the second file could not be uploaded on submission.

(A) Supplementary text: Staak_et_al_Supplement1.doc (Word file)Outlines regions (3%) of the published spectrum where errors are somewhat larger than the general uncertainties stated in the main manuscript. Reference is made in the main text - Ref. [21].

(B) Original data of measured spectrum: We intended to provide the full spectrum, comprising over 8000 absorption features, as supplementary material, since it is of interest for database development and atmospheric modelling. However, the online submission system does not allow the uploading of ordinary ASCII data or zipped files. The original ASCII data file has a size of 6.4 MB. The zipped data (using WinZip) has a size of 1.9 MB. If the data were embedded in a WORD document it would be 27 MB which is unacceptable.

Eventually we would like to provide the spectrum on the web in zipped ASCII form, if that is possible - please advice how to proceed with the submission of these data. E-mail: a.ruth@ucc.ie

The rotationally-resolved absorption spectrum of formaldehyde from 6547 to 6804 cm⁻¹

- Supplementary Material -

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Absorption spectrum

The full spectrum is available as supplementary material in two data columns; 1^{st} column: \tilde{v} [cm⁻¹], 2^{nd} column: σ [10⁻²¹ cm² / molecule].

Less accurate spectral regions

Based on the accurate of the mirror reflectivity as a function of wavelength, a maximum systematic uncertainty of the absorption cross-section of $\pm 5\%$ was established. An additional statistical uncertainty of less than 10% was established for the general measurement error of the absorption – both estimates are conservative.

Less than 3% of the overall spectrum have a larger statistical uncertainty of 25% in the absorption due to an insufficient overlap or inadequate correspondency of spectral segments to satisfactorily verify the absorption in these regions.

Spectral region $[cm^{-1}]$ 6546.86 - 6547.20 6549.40 - 6549.70 6551.26 - 6551.92 6553.20 - 6553.51 6558.10 - 6559.00 6559.30 - 6559.80 6564.80 - 6565.10 6575.60 - 6575.20 6598.00 - 6598.30 6612.15 - 6612.25 6613.50 - 6614.80 6643.60 - 6644.00 6655.20 - 6655.60

In the spectral regions listed below a shift in the wavenumber position was observed for the same absorption lines in different overlapping spectral segments. The shifts may be due to (a) unnoticed mode-hops of the laser or (b) discrepancies in the calibration data of NH₃ absorption lines. The region and the wavenumber difference between the corresponding lines is reported.

Spectral region [cm ⁻¹]	Wavenumber shift [cm ⁻¹]
6673.90 - 6674.74	0.005
6677.70 - 6678.20	0.004
6695.35 - 6695.45	0.004
6782.90 - 6782.10	0.007
6753.40 - 6754.10	0.007
6783.00 - 6783.60	0.005
6794.30 - 6794.50	0.005
6630.75 - 6630.85	0.003
6681.75 - 6681.85	0.004
6800.65 - 6801.20	0.010

We regard the following spectral region as rather uncertain in both wavenumber and absorption cross-section:

Spectral region [cm⁻¹] 6801.20 – 6802.16