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The quantitative infrared and NIR spectrum of CH₂I₂ vapor: vibrational assignments and potential for atmospheric monitoring

T. J. Johnson, T. Masiello, and S. W. Sharpe

Pacific Northwest National Laboratory, Richland, WA 99354, USA

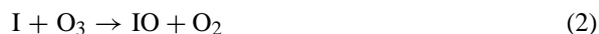
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Abstract. Diiodomethane (CH₂I₂) has recently become a molecule of significant atmospheric interest as it can contribute to coastal IO formation. As part of the PNNL database of gas-phase infrared spectra, the quantitative absorption spectrum of CH₂I₂ has been acquired at 0.1 cm⁻¹ resolution. Two strong b₂ symmetry A-type bands at 584 and 1114 cm⁻¹ are observed, but are not resolved when broadened to 760 Torr with nitrogen and appear as B-type. In contrast, the b₁ symmetry C-type bands near 5953, 4426 and 3073 cm⁻¹ are resolved with rotational structure, including Q-branches with widths ≤1 cm⁻¹. The quantitative infrared and near-infrared vapor-phase spectra (600–10 000 cm⁻¹) are reported for the first time. Some bands are discussed in terms of their potential for atmospheric monitoring and theoretical detection limits on a selected basis. FT-Raman spectra and ab initio calculations are used to complete vibrational assignments in the C_{2v} point group.

1 Introduction

Twenty-five years ago Chameides and Davis (1980) suggested the potential importance of iodine in tropospheric chemistry. They recognized that certain species such as methyl iodide, CH₃I, are easily photolyzed to liberate atomic iodine, which can react with O₃ and then NO_x or HO_x to form other inorganic species such as IO, HOI or IONO₂. The analogous reactions for diiodomethane, CH₂I₂, are:



Correspondence to: T. J. Johnson
(timothy.johnson@pnl.gov)

Similar to the ClO_x and BrO_x systems, the various iodine species can be cycled back to a radical I atom with catalytic O₃ destruction and an increase in the NO₂/NO ratio. Subsequent works continue to emphasize the importance of iodine in atmospheric chemistry (Solomon et al., 1994). Although typical concentrations are low (sub-ppb_v), most organic iodides are readily photolyzed in the visible and near-UV and the ensuing kinetics are typically fast, thus leading to the importance of the iodine cycles.

In addition to CH₃I, the photolysis of CH₂I₂ was also investigated some years ago where it was suggested that I₂ can appear as an end photoproduct and at high concentrations there is regeneration of the parent compound (Schmitt and Comes, 1980; Kasper et al., 1964, 1965). More recently and more importantly, the photolysis of CH₂I₂ has had renewed interest since laboratory experiments have shown that the photolysis of certain iodine-containing species in the presence of ozone (O₃) plays a direct role in the formation of aerosols (Jimenez et al., 2003; Kolb, 2002; Burkholder et al., 2004). It has been shown that the photolysis of CH₂I₂ in the presence of ozone results (via Reaction 2) in IO, and CH₂I₂ is thus a major contributor to coastal IO formation. Field experiments have subsequently shown that the seas are a direct source of several gas-phase halogenated species, (McFiggans et al., 2004; Carpenter et al., 1999) including CH₂I₂. Diiodomethane and other alkyl iodides have been identified with maritime biogenic sources, (Carpenter et al., 1999) in particular *Laminaria* Macroalgae (McFiggans et al., 2004) or the brown algae *Fucus* (Mäkelä et al., 2002). The existence of the iodine dioxide molecule, OIO, was first observed in the laboratory by Himmelmann et al. (1996). Ravishankara and co-workers (Harwood et al., 1997; Gilles et al., 1997) first investigated the kinetics of the IO radical reacting either with BrO or with itself, while Cox and colleagues (Bloss et al., 2001; Rowley et al., 2001; Cox et al., 1999) first observed OIO as a primary reaction product:



Tucceri et al. (2006) have recently measured the OIO absorption cross-sections using pulsed laser photolysis. O'Dowd and co-workers have also shown that nucleation was observed over several orders of magnitude in CH₂I₂ and suggested that the gas-phase OIO is responsible for the new particle growth (Jimenez et al., 2003; Mäkelä et al., 2002). CH₂I₂ photolysis in the presence of ozone thus plays a role in marine boundary layer ozone loss (Vogt et al., 1999).

In terms of spectroscopy, relatively little has been reported regarding gas-phase CH₂I₂. The early photolysis work of Pimentel and co-workers, as well as Kroger et al. (1976) and Schmitt and Comes (1980) showed that photolysis liberates an I atom and an internally excited CH₂I radical. Vibrational spectra of CH₂I₂ liquid have been reported previously by both Ford (1975) and Voelz et al. (1953), with the fundamentals being assigned in the C_{2v} point group, as well as a few combination and overtone bands. Although both the liquid- and vapor-phase spectra have been recorded in the IR, (Ford, 1975; Voelz et al., 1953), it appears the resolution was not sufficient to resolve any ro-vibronic structure in the vapor. There have also been assignments made of the $\Delta\nu_{\text{CH}}=3, 4, 5$ and 6 overtones in the near-infrared spectrum, (Henry and Hung, 1978) but these were also in the liquid phase. The anharmonicity was used to extrapolate a fundamental frequency of 3059 cm⁻¹ for the liquid, which would correspond to the C-H stretch ν_6 fundamental (vide infra). While Marshall et al. (2005) have calculated frequencies, but not intensities, of CH₂I₂ using ab initio methods, to date there have been no quantitative vapor spectra of this moderately volatile species (1.2 Torr at 298 K).

The Pacific Northwest National Laboratory (PNNL) continues to construct a reference library of quantitative high-resolution infrared spectra designed for tropospheric monitoring (Sharpe et al., 2004). This Northwest Infrared (NWIR) database, is optimized for best signal-to-noise ratio (SNR) in the long-wave infrared (LWIR), 800–1300 cm⁻¹, but each spectrum has a minimal spectral range of ≤ 600 to ≥ 6500 cm⁻¹. When certain bands are recognized as having sufficient SNR outside this minimal spectral range, these data are also included. The reference spectra are intended primarily for open path monitoring; each laboratory measurement has the analyte back-pressurized with ultra-high purity (UHP) nitrogen to 760±5 Torr. The data are nevertheless recorded at 0.112 cm⁻¹ resolution in the desire to bring out all possible spectral features, not only the resolved ro-vibronic lines of lighter species, but also the Q-branches of many polyatomics that display halfwidths of 1 cm⁻¹ or less. Recognizing that most species have typical pressure broadening coefficients of ~0.1 cm⁻¹/atm, additional resolution was deemed unnecessary for these reference data.

While constructing the NWIR database, it was recognized that CH₂I₂ has recently garnered great interest among the atmospheric community, and it would be beneficial to point

out alternate method for monitoring this trace gas. While GC/MS offers excellent sensitivity for CH₂I₂ analysis, it is limited to point measurements (Wevill and Carpenter, 2004). Since estimated maritime boundary layer concentrations are at most 100's of ppt_v, it was recognized that the only optical methods with sufficient sensitivity to see CH₂I₂ would be, for example, long path DOAS (Platt, 1994) or narrow-band IR laser techniques (Johnson et al., 1991a; Sharpe et al., 1998): Special attention was thus paid to narrow spectral features: e.g., transitions over which a probing laser frequency can easily be tuned.

2 Experimental

Using the methods of Chu et al. (1999) and Sharpe et al. (2004) a series of measurements is made, each corresponding to a different concentration-path length burden. Each fitted spectrum, in fact, represents a series of (typically >10) individual measurements at 298.1 K, with the different measurements covering a large range (often >2 orders of magnitude) of analyte burdens, each burden pressurized with UHP N₂ to one atmosphere. The fitted spectrum is derived by fitting a Beer's law plot at each wavelength channel to each of the individual burdens. In order to account for any of several different nonlinearity phenomena in the A=f(P) fit, the individual burdens are in addition weighted according to T² (where T=I/I₀). All values with T<0.025 (i.e. decadic absorbance A>1.6) are weighted with zero. This multiple burden with weighted data approach retains several advantages over any single measurement or few measurements: First, multiple measurements greatly enhance the SNR ratio. Second, the high burden measurements enhance the SNR for the weak bands that might not exceed the noise floor in any measurement designed to keep the strong bands on scale. Third, for the strong bands, the weighting mechanism brings out a better fidelity to account for Beer's law saturation effects, photoconductive detector nonlinearity effects or other phenomena that cause the absorbance values to not scale with burden. Finally, analyzing the residual vector has proven helpful at recognizing chemical impurities as their signatures typically do not scale with the fit.

Allen et al. (2005) have also recently published the advantages of fitting the absorbance as a function of partial pressure in order to calculate A(λ), as first discussed by Chu et al. (1999). The Chu et al. methods go further, however, that in addition to the absorbance data linear fit at each wavelength bin, the raw data also have a weighting factor to account for nonlinearity mechanisms. In this method the residual fit vector is, moreover, carefully analyzed since any chemical impurities, including the uncommon ones, readily manifest themselves in the residual. The PNNL method typically measures twelve to fifteen separate burdens ranging over two orders of magnitude. The resulting error analysis (Sharpe et al., 2003) shows that the maximal expected

systematic errors in absorbance are 3% for the PNNL static system measurements and 7% for flow system measurements, including the CH₂I₂ values reported here. The measured random values are less than this, and these results have been vetted against NIST for a host of different molecules as described by Sharpe (2003, 2004), which contain further details of the error analysis. The above discussion is valid for each of the 400+ molecules in the PNNL database. In addition, for CH₂I₂ and a handful of other species, high resolution (0.0015 cm⁻¹) studies have assigned all observed bands to the analyte, increasing confidence as to chemical purity. (Maki et al., 2001; Masiello et al., 2005).

The CH₂I₂ sample itself was from Aldrich Chemicals and its purity was monitored by comparing to known infrared spectra. A Bruker IFS 66v/S vacuum spectrometer (Johnson et al., 2005) was used over the 600 to 6500 cm⁻¹ range. The spectrometer hardware characteristics have been previously documented (Johnson et al., 2002) according to IUPAC guidelines, (Bertie et al., 1998) taking care to avoid ghosting, “warm aperture” (Johnson et al., 2002) and detector nonlinearity (Chase, 1984; Birk et al., 1996) artifacts. Due to the modest vapor pressure of CH₂I₂ (1.2 Torr at 298 K, Yaws, 1999) these measurements were made on a long path flowed system whereby the liquid analyte is quantitatively delivered into a heated stream of ultra-high purity N₂ carrier gas, (Sharpe et al., 2003) using a specially constructed vapor dissemination system (Johnson et al., 2006¹). The measurement at 298.1 K is made in a specially designed White cell with a circulating liquid jacket which can provide more precise temperature control (Ballard et al., 1994) with the optical path set to 7.96 m (±0.5%). The temperature is measured by placing a NIST-traceable temperature probe directly into the gas in the White cell; the stated accuracy is ±0.2 K. For each of the 12 burdens, 256 interferograms were averaged to reduce noise. Gain ranging improved the signal-to-noise by amplifying the wings of the interferogram before digitization. The single-sided interferograms were zero-filled 2× and phase corrected using Mertz’s method (Mertz, 1967). Decadic absorbance spectra were calculated in the usual manner [−log(I/I₀)] using the spectrum of the cell with just N₂ carrier gas for I₀. The final data product is a spectrum that corresponds to an optical path of 1 m through a mixing ratio of 1 ppm_v of analyte at 296 K.

For this particular molecule, several bands of interest were noted in the near-infrared (NIR) range, so seventeen additional burdens were measured with alternate hardware optimized to improve the NIR signal/noise. These included a tungsten halide source, a Si/CaF₂ beamsplitter, and a photovoltaic InSb detector. The mirror scan speed was adjusted to 20 kHz, but all other collection parameters were as for the mid-infrared measurements (Sharpe et al., 2004). The

NIR spectra covered the 1900 to 10 000 cm⁻¹ range, for a total range of 550–10 000 cm⁻¹. Data are only plotted to 6500 cm⁻¹ as the few bands at higher frequencies are relatively weak. The FT-Raman spectra were collected over the range from 50 to 3600 cm⁻¹ Stokes shift and from −100 to −1000 cm⁻¹ anti-Stokes shift using a previously described instrument (Williams et al., 2006). Raman frequencies are calibrated using the interferometer HeNe laser and the Raman Nd:YAG laser. For a given FT-Raman spectrum the frequency accuracy has been shown to be better than 1 cm⁻¹. Data collection parameters and methods for the low pressure, high resolution IR spectra (0.0015 cm⁻¹ on a Bruker IFS 125), including for peaks reported in the far-IR, have already been reported by Maki et al. (2001). Theoretical ab initio calculations were performed using the NWChem software algorithm. The vibrational frequencies and infrared line intensities (peak maxima) were calculated at the MP2 level with the largest basis set (6–311 g**) possible that delivers both frequencies and intensities; the frequencies are typically too large, and are customarily scaled by a factor of 0.955 (Marshall et al., 2005).

3 Results

Figure 1 reports the quantitative gas-phase infrared spectrum² of methylene iodide (diiodomethane) from 550 to 6500 cm⁻¹. The y-axis is quantitative such that the absorbance values correspond to an optical path of 1 m through a concentration of 1 ppm_v of analyte at 296 K and 1 atm total pressure. The reported spectrum represents the weighted average of 12 individual measurements for the mid-infrared and 17 measurements for the near-infrared. The concatenated spectrum has been broken into four separate plots for visual clarity. Note that the plot in Fig. 1a has a different ordinate scale as the ν₉ band at 584.21 cm⁻¹ and the ν₈ band at 1113.87 cm⁻¹ are both IR-allowed fundamentals and are very intense bands. Plots 1b, 1c and 1d are all on the same scale, and all plots represent an average from multiple burdens of CH₂I₂, each burden pressure-broadened to 760 Torr. The peak positions and band types of the fundamentals for the present vapor-phase data are given in Table 1. Also in Table 1 are the corresponding liquid-phase values, both infrared and Raman, as well as the vibrational assignments. Table 2 presents the integrated vapor-phase band strengths (in units of cm⁻¹ cm² molecule⁻¹, or cm molecule⁻¹) for the fundamentals. While the data in the NWIR database use base 10 logarithms, the band integrals in Table 2 use Napierian (natural) logarithms at 296 K for the peak height as this is the more common unit for *S*. Also in Table 2 are the frequencies and intensities as calculated by the ab initio program NWChem. As these are the first Fourier-transform measurements of vapor-phase CH₂I₂ the accuracy along the

¹Johnson, T. J., Sharpe, S. W., and Covert, M. A.: A Method for Quantitative Dissemination of Vapors, Review of Scientific Instruments, in review, 2006.

²The CH₂I₂ spectrum and other NWIR data can be found at <http://nwir.pnl.gov>.

Table 1. Observed Raman (liquid), Infrared (liquid and gas) vibrational frequencies, Raman polarizations and (for vapor measurements) band types of CH₂I₂, together with symmetries and vibrational assignments. Frequencies are in cm⁻¹. Fundamental frequencies are printed in bold. Details can be found in text. NS = not seen, OOR = out of range.

Ram Liq	Ram str	Ram Polz	IR Liq	Liq Str	IR Gas	Band Type Rel. Intens.	Assm't	SYM
121	vs	pol	122	w	NS		ν_4	a₁
197	w	pol	NS	NS	NS			
238	w	pol	NS	NS	NS		2 ν_4	A ₁
365	w	pol	NS	NS	NS		$\nu_3 - \nu_4$	A ₁
448	vw		448		463.79		$\nu_9 - \nu_4$	
486	vs	pol	485	w	493.01	B-vw	ν_3	a₁
571	m	dp	571	s	584.21	A-vs	ν_9	b₂
604	w	pol	NS	NS			$\nu_3 + \nu_4$	A ₁
717	w	dp	717	s	718.08	C	ν_7	b₁
851	w		851	vw	NS			
969	w	pol	NS	NS	NS		2 ν_3	A ₁
1032	w	dp	1033	w	1041.99	vw	ν_5	a₂
					1075.04	vw	$\nu_3 + \nu_9$	B ₂
1109	m	dp	1106	vs	1113.87	A-vs	ν_8	b₂
1134	m	pol	NS	NS	NS	NS	2 ν_9	A ₁
1181	vw		1181	w	1189.1	w	$\nu_3 + \nu_7$	B ₁
NS			1226	m	1229.42	A-s	$\nu_4 + \nu_8$	B ₂
1353	m	pol	1352	m	1373.61	B-w	ν_2	a₁
1434	m	pol	NS	NS	NS		2 ν_7	A ₁
			1595	vw	1603.39	w	$\nu_3 + \nu_8$	B ₂
1672	w	pol	NS	NS			$\nu_8 + \nu_9$	A ₁
			NS	NS	1714.86	C-w	$\nu_4 + \nu_5 + \nu_9$	B ₁
1755	w	dp	1754	vw	1759.38	A	$\nu_5 + \nu_7$	B ₂
NS			1837		1864.97	w	$\nu_7 + 2\nu_9$	B ₁
NS			1921	w	1953.86	w	$\nu_2 + \nu_9$	B ₂
1995	vw	pol	NS		NS			
2058	w	dp	2056		2075.67	C	$\nu_2 + \nu_7$	B ₁
NS			2132		2142.42	C-s	$\nu_5 + \nu_8$	B ₁
2209	w	pol	2209		2222.42	B	2 ν_8	A ₁
2238	w	pol	NS	NS	NS			A ₁
2360	w	pol	NS	NS	NS			A ₁
2453	w	dp	2454		2482.14	A	$\nu_2 + \nu_8$	B ₂
2504	vw		NS		NS		$\nu_2 + 2\nu_9$	A ₁
NS		NS	2549	vw	2560.12	w	$\nu_2 + \nu_3 + \nu_7$	B ₁
2602	m	pol	NS	NS	2631.70	vw	2 $\nu_5 + \nu_9$	B ₂
2967	vs	pol	2968	s	3002.00	B-s	ν_1	a₁
3046	s	dp	3046	vs	3073.01	C-s	ν_6	b₁
3332	w	pol	NS	NS	3323.23	B-w		
3573	w	pol	NS	NS	3585.93	B-w	$\nu_1 + \nu_9$	B ₂
OOR			3757	m	3785.26	B-w	$\nu_6 + \nu_7$	A ₁
OOR			4061	m	4097.60	A/B-s	$\nu_5 + \nu_6$	B ₂
OOR			4377	m	4426.52	C-s	$\nu_2 + \nu_6$	B ₁
OOR			5852	w	5921.62	B-m	2 ν_1	A ₁
OOR			5892	w	5953.31	C-m	$\nu_1 + \nu_6$	B ₁
OOR			6094	w	6102.68	B-m	2 ν_6	A ₁

wavelength axis is much higher (Sharpe et al., 2004; Maki and Wells, 1991). High resolution vapor-phase infrared spectra confirm our low resolution estimates for many of the band

origins given in Table 1. However, precise determination of the band origins can only be achieved through analysis of the high resolution data. Consequently, an error of ± 0.1 cm⁻¹

should be assumed for all values given in Table 1 for the vapor-phase spectra.

The vibrational assignments largely agree with those of Voelz and co-workers (1953) as well as Ford (1975). The nine fundamentals in the C_{2v} point group are printed in bold in Table 1. The long path gas cell and greater instrumental sensitivity allow for observation of even very weak fundamentals including ν_3 (a₁ symmetry, 493.01 cm⁻¹) ν_7 (b₁, 718.08 cm⁻¹) and ν_2 (a₁, 1373.61 cm⁻¹). The Raman spectra aided in the assignment and confirmation of many vibrational bands, especially the polarized Raman spectra which helped to confirm assignment of vibrational modes with a₁ symmetry. The symmetric ν_1 (3002.00), ν_2 (1373.61), ν_3 (493.01) and ν_4 (122 cm⁻¹) a₁ modes are easily assigned due to their medium to very strong Raman signals which are all strongly polarized (i.e. a₁ modes). We note that the ν_4 (122 cm⁻¹) fundamental was not observed in any of the IR gas-phase spectra, including path lengths >35 m. This diminished intensity does agree with that predicted by the ab initio calculations (Table 2). The ν_8 (1113.87) and ν_9 (584.21) IR bands are both b₂ modes and dominate the IR spectrum. Although both appear to be B-type bands in the pressure-broadened 0.1 cm⁻¹ gas-phase measurements, high resolution (0.0015 cm⁻¹) measurements at low pressure do show A-type contours, and centralized Q-branch features as expected.

The ν_5 (a₂) fundamental is seen in both the liquid Raman and liquid IR spectra at 1033 cm⁻¹. As an a₂ mode, it is IR forbidden in the C_{2v} point group, but we can tentatively assign the very weak 1041.99 cm⁻¹ band as the corresponding gas phase fundamental which has modest intensity in the liquid due to condensed phase interactions. The depolarized Raman signal helps assign the symmetry. The $\nu_3 + \nu_9$ combination band is seen only in the gas phase at 1075.04 cm⁻¹, while the band at 2075.67 is the $\nu_2 + \nu_7$ combination. Many of the overtone, combination and difference bands are assigned for the first time. In particular, we assign for the first time many of the bands at frequencies >3500 cm⁻¹. Although these frequencies are outside the Raman spectrum, the vapor-phase band profiles (Fig. 2), along with the NWChem ab initio calculations make the assignments straightforward in most cases.

In terms of band strengths, the b₂ ν_9 (584.21 cm⁻¹) and ν_8 (1113.87 cm⁻¹) bands are clearly the most intense IR bands and dominate the spectrum. Both are pressure broadened at 760 Torr (Nitrogen) to form unresolved bands that are B-type in appearance, but in fact are A-type. Of the two, ν_8 is the more intense with a width (FWHM) of 11.8 cm⁻¹ and a peak amplitude 1.2×10^{-3} absorbance for 1 ppm-m. The spacing of the CH₂I₂ rotational lines in the ν_8 and ν_9 bands is quite small and these do not resolve at atmospheric pressure. They do resolve at low pressure, however, and a high resolution study assigning both these bands is currently under way in our laboratory using previously described high resolution instrumentation (Maki et al., 2001).

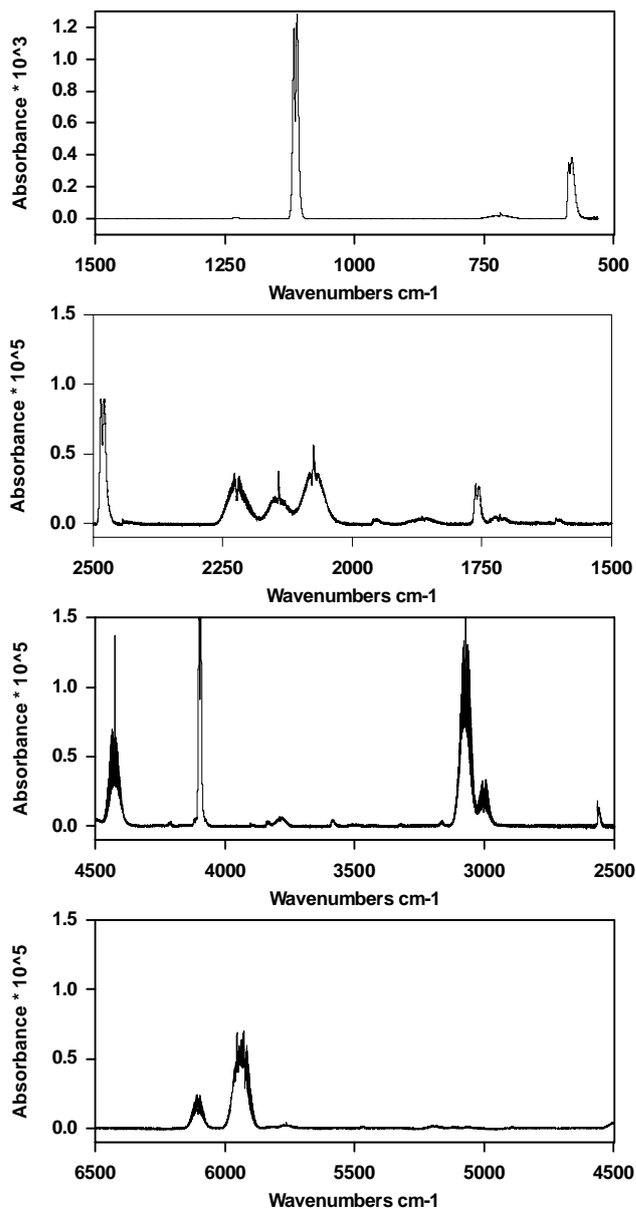


Fig. 1. Mid- and near-infrared quantitative spectrum of diiodomethane (CH₂I₂) from 6500 to 550 cm⁻¹, recorded at 0.1 cm⁻¹ resolution. The spectrum is actually a composite derived from multiple spectra that corresponds to 1 ppm-m of gas at 296 K pressure-broadened to 760 Torr. Note that due to the strong bands at 584 and 1114 cm⁻¹ panel (a) has been multiplied by 10³, while panels (b–d) have been multiplied by 10⁵ on the y-axis. See text for details.

For the remaining pressure-broadened bands shown in Fig. 1, many are of approximately the same band strength, and of these several are C-type bands that exhibit resolved Q-branches. We focus on these in particular, since the resolved structure better lends itself to open path monitoring, either via methods such as DOAS (Platt, 1994; Volkamer et al., 2005) or laser-based techniques. Three of these

Table 2. Observed Raman (liquid) and infrared (liquid and gas) vibrational frequencies along with gas-phase band integrals for the fundamental vibrations of CH₂I₂. Frequencies are in cm⁻¹ and band integrals are in units of cm-molecule⁻¹, using Napierian (natural) logarithms at 296 K for the peak heights, and have been multiplied by 10¹⁸. Also listed are the frequencies (scaled by 0.955) and IR intensities relative to ν_8 as predicted by NWChem. Details in text.

Assm't	SYM	Ram Liq	IR Liq	IR Gas	Gas Bnd <i>f</i> limits	Band <i>f</i> cm/molec	NWC Freq'y	NWC Ints'y
ν_4	a ₁	121	122	NS	NS	296 K × 1e18 NS	Adjusted 117	Rel to ν_8 0.02
ν_3	a ₁	486	485	493.01	NC	NC	476	0.15
ν_9	b ₂	571	571	584.21	600–550	5.1240	579	28.60
ν_7	b ₁	717	717	718.08	775–665	0.8785	705	9.02
ν_5	a ₂	1032	1033	1041.99	1048–1031	0.0008	1045	–
ν_8	b ₂	1109	1106	1113.87	1135–1090	13.0800	1135	100.00
ν_2	a ₁	1353	1352	1373.61	1420–1330	0.0159	1366	0.16
ν_1	a ₁	2967	2968	3002.00	3030–2950	0.0756	3030	0.26
ν_6	b ₁	3046	3046	3073.01	3140–3030	0.4147	3111	2.80

bands with the C-type profile will be considered in more detail for pressure-broadened (ambient) monitoring. Figure 2's top trace displays the ν_1 fundamental at 3002.00 cm⁻¹ along with the C-type ν_6 band at 3073.01 cm⁻¹, where the plot again represents an optical depth of 1 ppm-m. For the ν_6 band the Q-branch has a peak height of 2.65×10^{-5} OD with a Q-branch differential peak height of 1.98×10^{-5} . The middle trace of Fig. 2 plots the $\nu_2 + \nu_6$ combination band at 4426.52 cm⁻¹, and the lower trace shows the $2\nu_1$ overtone at 5921.62 cm⁻¹ along with the $(\nu_1 + \nu_6)$ combination C-type band at 5953.31 cm⁻¹. Note that the y-axis has been held the same in the three plots to show the relative line strengths, most of which are comparable, though all these bands are approximately two orders of magnitude weaker than ν_8 or ν_9 bands in the long-wave infrared.

4 Discussion

Both the intense IR fundamentals, ν_9 at 584.21 cm⁻¹ and ν_8 at 1113.87 cm⁻¹, shown in Fig. 1 are in fact unresolved A-type bands and fall in an “atmospheric window”. For low resolution monitoring, the ν_8 band at 1113.87 cm⁻¹ is the most promising as it lies in the center of the LWIR window. We note that both the ν_8 and ν_9 fundamentals are of b₂ symmetry in the C_{2v} point group. At 0.1 cm⁻¹ resolution and pressure broadened by N₂, both appear as B-type bands, but very high resolution spectra recorded at low pressure have shown the

expected A-type structure. The ν_8 band has a width (FWHM) of 11.8 cm⁻¹, and a peak amplitude of 1.2×10^{-3} OD for 1 ppm-m. At lower resolution, typically 4 or 8 cm⁻¹, used in most ambient monitoring situations with an active or semi-active FTIR system (Johnson et al., 2004), the one minute detection limits are of the order 10⁻⁴ OD (Griffith and Jamie, 2000). This suggests that for a 500 m (open) path, an optimistic detection limit for CH₂I₂ is 500 ppt_v. Consequently, current FT-methods would be challenged to observe the anticipated <100 ppt_v biogenic levels of CH₂I₂ (Carpenter et al., 1999). Although they are the strongest bands, the ν_8 and ν_9 bands are unresolved at 760 Torr and may not be useful for open-path laser monitoring. As mentioned above, however, these bands do resolve at low pressure, and their assignments and potential for monitoring will be discussed in a separate work. We are optimistic that the line strengths are sufficient to be useful for extractive monitoring, using e.g. Pb-salt or QC-laser systems.

Of the laser methods, Pb-salt systems have been widely used in the mid-IR ($\lambda=3$ to 20 μ m), primarily as point sensors using long path cells at low pressure (Werle et al., 2004). Several near-IR laser systems have been deployed using techniques such as FM modulation (Johnson et al., 1991b) or cavity ring-down methods (Bitter et al., 2005). Until recently, the greatest limitations appear to be the tunability and wavelength availability of the lasers, predominantly limited to the wavelengths used by the telecommunications industry

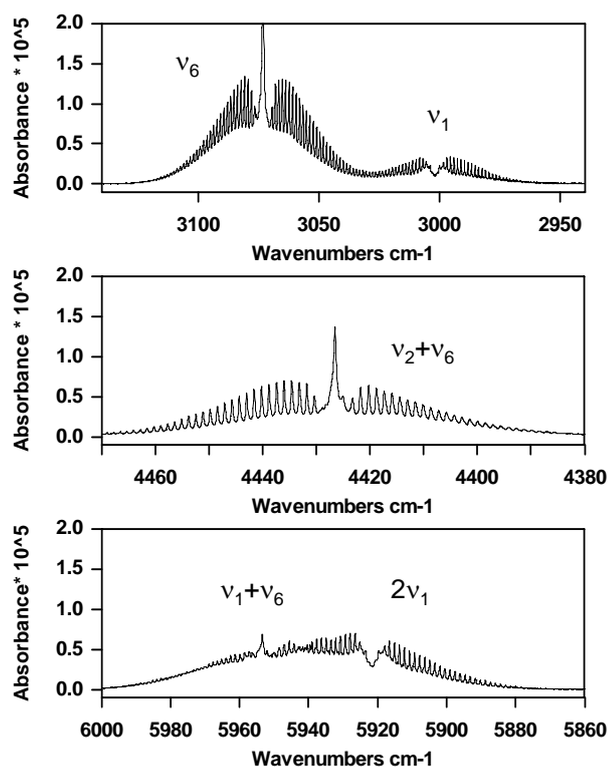


Fig. 2. Details of three spectral regions of interest for monitoring. The spectra are all pressure broadened to 760 Torr and are composite spectra that correspond to 1 ppm-m optical depth. The y-axis is on the same scale in all three panels. Panel (a) shows the ν_1 band (3002 cm^{-1}) along with the ν_6 band (3073 cm^{-1}), panel (b) shows the $(\nu_2+\nu_6)$ combination band at 4427 cm^{-1} , while panel (c) shows the $2\nu_1$ overtone band (5921 cm^{-1}) along with the $(\nu_1+\nu_6)$ combination band at 5953 cm^{-1} .

(e.g. 1.3 or $1.55\text{ }\mu\text{m}$). The recent advent of quantum cascade (QC) lasers shows excellent promise as these are available from the near-IR to the terahertz (THz) wavelengths. These lasers have also demonstrated an important feature in that they have significant tuning ranges, typically $>0.5\%$ or more of the central wavelength, depending on the nature of the device fabrication. This allows, for example, tuning across a 1 cm^{-1} wide Q-branch for atmospheric monitoring purposes.

Figure 3 demonstrates one possibility for monitoring using the Q-branch of the 760 Torr ν_6 band at 3073.01 cm^{-1} ; this CH₂I₂ spectrum (1 ppm-m) is plotted in pink. Although 1 ppm is far greater than expected ambient concentrations, this may simulate a chamber experiment. Also plotted in Fig. 3 are 1 m paths for typical ambient concentrations (Warneck, 1988) of 1.8 ppm for methane (green) and the optical density corresponding to 10 Torr of H₂O (blue). The water data are from the HITRAN database (Rothman et al., 2005) since the HITRAN calculated data have no noise that can obfuscate the signal when expanded thousands of times. The database was examined for other potential interferents with high background concentrations (e.g. N₂O, CO₂, CO), but

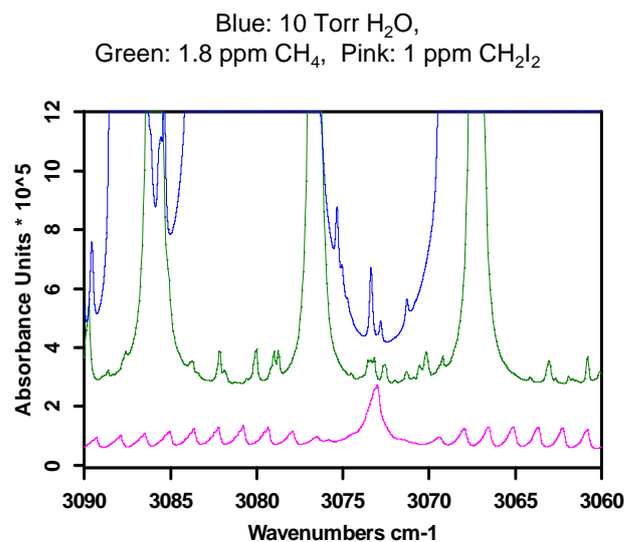


Fig. 3. Comparison plot (all at 0.1 cm^{-1} resolution) showing potential interferents to the CH₂I₂ ν_6 band at 3073.01 cm^{-1} . The traces correspond to 1 ppm-m of CH₂I₂ (lower trace – pink), 1.8 ppm-m of CH₄ (middle trace – green), and 10 Torr of H₂O (upper trace – blue). The H₂O data are from the HITRAN data base. The spectra have been vertically offset for clarity. See text for details.

these did not show any interfering absorptions at these wavelengths. The Q-branch at 3073 cm^{-1} corresponds to a differential peak height of 1.98×10^{-5} OD for 1 ppm-m, which is also equivalent to 1.98×10^{-5} for 1 ppb-km. The $3.25\text{ }\mu\text{m}$ wavelength can be accessed by both Pb-salt and quantum cascade diode lasers. Assuming an open path laser-based system with a 10 km path length, in order to achieve a 50 ppt_v detection limit, a spectrometer would need a detection sensitivity corresponding to $\sim 1\times 10^{-5}$ OD or better. For an extractive laser-based system with a 200 m path length, in order to achieve a 50 ppt_v detection limit, a spectrometer would need a detection sensitivity corresponding to $\sim 2\times 10^{-7}$ OD or better at these wavelengths. Although such a measurement would pose a challenge, the best extractive measurements at these wavelengths are now achieving such detection limits using difference frequency generation (Richter et al., 2002) or improved Pb-salt diode laser systems (Wert et al., 2003; Schiller et al., 2001). The weak methane lines near 3073 cm^{-1} pose a concern, but the CH₄ mixing ratios can be expected to be low and constant in marine atmospheres. It is also possible that using an extractive system with a near-infrared detector (e.g. InSb) and a long path cell that some of the lines in the Q-branch resolve and separate from those of methane, also possibly increasing the depth of any one of the CH₂I₂ lines. Orphal and Ibrahim (2005) have announced recent results showing that the ν_1 and ν_6 bands near $3.3\text{ }\mu\text{m}$ do resolve at low pressure and thus may be useful for extractive monitoring.

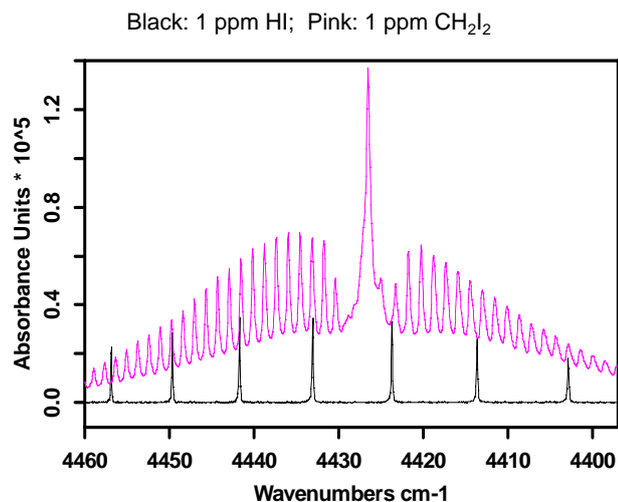


Fig. 4. Comparison plot (all at 0.1 cm^{-1} resolution) showing potential laser combination measurements using the CH₂I₂ ($\nu_2+\nu_6$) combination band at 4426.52 cm^{-1} . The traces correspond to 1 ppm-m of CH₂I₂ (upper trace – pink), and 1 ppm-m of hydrogen iodide, HI (lower trace – black). The spectra have been vertically offset for clarity.

Figure 4 plots the ($\nu_2+\nu_6$) combination band at 4426.52 cm^{-1} which we report for the first time. Figure 4's bottom trace also shows some of the sharp lines of the $2\nu_1$ overtone band of hydrogen iodide gas, HI. It was hoped that using a single laser it may be possible to simultaneously monitor both HI and CH₂I₂ by slight tuning of the wavelength. However, closer inspection of the database immediately showed that expected concentrations of 1.8 ppm_v of CH₄ or 316 ppbv of nitrous oxide (Evans and Puckrin, 1997) would interfere with such a measurement, particularly the N₂O. However, extractive measurements may still afford a practical separation of the HI/CH₂I₂ pair from the interfering signals, particularly at wavenumbers $>4440\text{ cm}^{-1}$, although wall adhesion mechanisms are of concern for species such as HI, HCl, or HNO₃ (Chackerian et al., 2003; Johnson et al., 2003). Nevertheless, such bands could be of use in e.g. a chamber kinetics study without CH₄ or N₂O.

The traces in Fig. 5 display two new bands of CH₂I₂, namely the $2\nu_1$ overtone (A₁ symmetry, B-type) at 5921.62 cm^{-1} , as well as the ($\nu_1+\nu_6$) combination C-type band at 5953.31 cm^{-1} . Both have resolved rotational structure even when pressure broadened to one atmosphere. The structure in these bands suggests the possibility of remote sensing as the Q-branch at 5953 cm^{-1} as well as several of the individual rotational lines, e.g. in the $2\nu_1$ R branch near 5930 cm^{-1} have differential absorbances of $\sim 3\times 10^{-6}$. Such structure is very helpful not only for laser monitoring, but also for DOAS methods, which have been used to monitor trace gases such as HONO at the (sub-) ppbv levels (Platt et al., 1980). Detection limits of 10^{-4} are achievable, and path lengths of $>10\text{ km}$ are now common. In a much related work

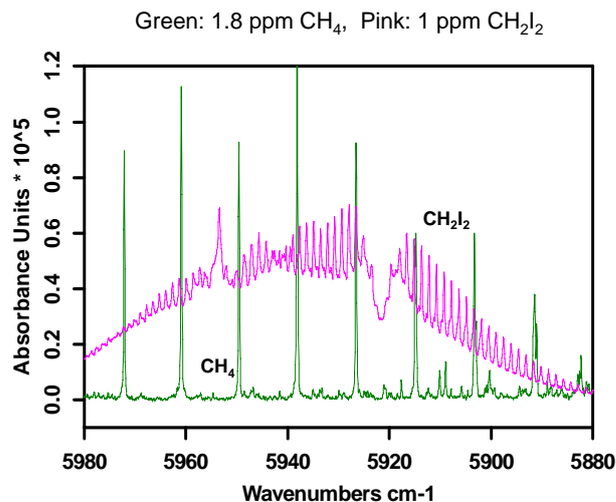


Fig. 5. Comparison plot (0.1 cm^{-1} resolution) showing potential laser combination measurements using the CH₂I₂ $2\nu_1$ overtone band (5921 cm^{-1}) along with the $\nu_1+\nu_6$ combination band at 5953 cm^{-1} along with the potential interference from CH₄. The traces correspond to 1 ppm-m of CH₂I₂ (upper trace – pink), and 1.8 ppm-m of methane, CH₄ (lower trace – green). The spectra have been vertically offset for clarity. See text for details.

Alicke and co-workers (1999) have used such a set-up in the UV-visible with a 14.6 km path to measure IO at levels of $<10\text{ ppt}_v$. For the case of CH₂I₂, if one assumes a 15 km (e.g. estuary) path and a differential cross section of $3\times 10^{-6}\text{ ppm-m}$ (equivalent to $2.8\times 10^{-21}\text{ cm}^2/\text{molecule base-e}$) in order to observe 100 ppt_v of CH₂I₂ it would be necessary to detect an absorbance of 4.5×10^{-6} at these wavelengths. This is certainly a challenging detection limit, but DOAS detection for methane has recently been reported at these wavelengths (Frankenberg et al., 2005).

Of the CH₂I₂ near-infrared bands discussed above, several fortuitously display resolved structure, even when broadened to 1 atmosphere. Although the telecommunications diode lasers are limited in wavelength selection, several recent technologies offer methods for sensitive trace gas detection using such lines. These include tunable optical parametric oscillators, difference frequency generation (Richter et al., 2002) and, recently, red-shifting the emission wavelength of near-IR lasers via high pressures (Adamiec et al., 2004). In terms of the CH₂I₂ photochemistry in the presence of ozone, it may be possible to monitor the CH₂I₂, HI and IO simultaneously in a chamber type experiment as suggested by Mäkelä et al. (2002). The ν_6 3073 cm^{-1} band may be useful in laboratory experiments where CH₄ and H₂O concentrations are reduced. Also, monitoring at $3\text{ }\mu\text{m}$ offers the advantage of less light scattering compared to visible or NIR regions.

5 Summary

The quantitative infrared and near-infrared vapor-phase spectra of diiodomethane, CH₂I₂, have been reported for the first time. Using FT-IR and FT-Raman spectroscopy as well as ab initio computational methods we have completed a vibrational assignment. The atmospherically broadened spectra exhibit C-type bands near 5953, 4426 and 3073 cm⁻¹ all have resolved rotational structure, including Q-branches with widths of ~1 cm⁻¹. While GC-MS currently offers better sensitivity for point monitoring, some of the quantitative bands discussed here could be potentially useful for (remote) monitoring of this important species. The ν₆ Q-branch at 3073 cm⁻¹ shows promise for extractive monitoring by Pb-salt or QC diode laser systems and the band strengths suggest estimated CH₂I₂ mixing ratios are near (or close to) the detection limit of current technology's best systems. The two very strong b₂ A-type bands (appearing as B-type, ν₉ at 584.21 cm⁻¹ and ν₈ at 1113.87 cm⁻¹) do not resolve at atmospheric pressure, but show great promise for low pressure extractive laser studies. However, high resolution studies will first be needed.

The ν₁+ν₆ band shows a resolved Q-branch at 4426 cm⁻¹. While this band would be of little use for field measurements due to N₂O interferences, it could be used e.g. in smog chambers (synthetic air) with FTIR/laser spectrometers. As Fig. 4 depicts, it has the added benefit of being able to simultaneously measure the adjacent HI lines in a kinetics study. The two bands shown in Fig. 5 with strong structure near 5950 cm⁻¹ may be of interest to e.g. the satellite-based DOAS systems as the long optical paths increase the burden for greater optical depth, and the wavelengths are just accessible on the red end of many current spectrometers.

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