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# New analysis of the $\nu_6$ and $2\nu_3$ bands of methyl iodide ( $\text{CH}_3\text{I}$ )

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## Abstract

A new rovibrational study of the  $\nu_6 = 1$  band of methyl iodide was conducted to obtain a rather complete line list. Therefore, we accomplished a new analysis in line position. The spectrum of this band has been first recorded using the FTIR Bruker HR125 at the AILES beamline of the SOLEIL Synchrotron facility and later with the Bruker FTIR IFS125HR located at the LISA facility in Créteil. The final analysis led to the assignment of about 10000 infrared lines up to  $J = 85$  and  $|K| = 20$  quantum numbers, 6000 lines of which are due to the  $^{127}\text{I}$  nuclear quadrupole coupling hyperfine structure revealed in the spectrum owing to the iodine nuclei. A global fit based on the Watson model of these new infrared data combined with previously assigned microwave lines [1] helped improving the last set of published parameters for this band [2]. The  $\nu_3 = 2$  band was also analysed and around 1500 lines were assigned. Due to the high values of quantum numbers achieved for the  $\nu_6 = 1$  energy levels, it proved to be crucial to take into account an  $\alpha(\Delta\ell = \pm 1; \Delta K = \pm 2)$  and a  $C_x(\Delta\ell = \pm 1; \Delta K = \pm 1)$  types of Coriolis interactions with the levels of  $\nu_3 = 2$  and of  $\nu_2 = 1$  respectively.  
*Keywords:* methyl iodide –  $\text{CH}_3\text{I}$  – Line List – High-resolution Fourier

## 1. Introduction

Methyl iodide, ( $\text{CH}_3\text{I}$ ) is emitted to the atmosphere by the oceans and photolyzes with a lifetime of the order of a week. It is a source for the ozone destruction in the upper troposphere and in the lower stratosphere [3]. It is  
5 as well of nuclear interest. In the case of a sever nuclear accident, the iodine fission products represent the major part of the released radioactivity and are of high concern due to their affinity to the thyroid. Methyl iodide is produced in the primary circuit by the reaction, under ionising radiation, of iodine with the organic coating of the enclosed containment. Because, in the case of an ac-  
10 cident, organic iodine is not efficiently trapped on filters, it is crucial to monitor its release into the atmosphere, as an important goal of the nuclear safety and radio-protection. Hence, the present work concentrates on acquiring a database for a future spectroscopic remote sensing detection of methyl iodide.

To our knowledge, there is up to now no infrared detection of this specie  
15 in the atmosphere from satellite instrument. The rather strong  $\nu_6$  band of  $\text{CH}_3\text{I}$ , located at  $892.916\text{ cm}^{-1}$ , which coincides with the  $11\ \mu\text{m}$  window of transparency in the atmosphere, could therefore be a good candidate for such detection. In addition it is relatively isolated from the absorption of the atmospheric molecules, especially water, as shown in figure 1. However, there exist  
20 no spectroscopic parameters for this molecule in the common access spectroscopic databases HITRAN<sup>1</sup> or GEISA<sup>2</sup>. This work, which deals with the line positions, is the very first contribution to fill this gap.

This molecule is also one of the most intensively studied symmetric top molecules by means of high resolution spectroscopy for both the for the  $^{12}\text{CH}_3\text{I}$   
25 and  $^{13}\text{CH}_3\text{I}$  isotopologues. It has six vibrational modes which are described

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<sup>1</sup><http://hitran.org/>

<sup>2</sup><http://ether.ipsl.jussieu.fr/etherTypo/?id=950>

shortly in Table 1. Methyl iodide is a prolate symmetric top molecule with an equilibrium configuration belonging to the  $C_{3v}$  point group. It has nine fundamental modes, three of which,  $\nu_1$ ,  $\nu_2$ ,  $\nu_3$ , being non-degenerate modes and belong to the  $A_1$  symmetry representation in addition to three degenerate  
30 modes  $\nu_4$ ,  $\nu_5$ ,  $\nu_6$  having the  $E$  symmetry. The infrared spectrum of methyl iodide is dense in the infrared region due to the iodine nuclei which, on one hand, makes of the methyl iodide a heavy molecule with small values of the rotational constants, and on the other hand, complicates the spectrum by the nuclear hyperfine effect which splits each rovibrational level into six sub-levels labelled by  
35 the  $F$  quantum number. Its rotational spectrum in the ground and in the first excited vibrational states was investigated in detail by microwave [4, 5, 1], and by Doppler-free double resonance techniques [2, 6]. This molecule was also the subject of numerous infrared studies which concern the  $\nu_1$  [7],  $\nu_3$  [8],  $\nu_4$  [9],  $\nu_5$  [10],  $\nu_2$ ,  $\nu_5$  and  $\nu_3+\nu_6$  band system, [11],  $\nu_6$  [12, 13, 14, 15, 16, 17], and  $2\nu_3$ [18]  
40 bands and several combination bands in the  $2800\text{ cm}^{-1}$  spectral region [19]. In this way, vibrational band centers, rotational and coupling constants, together with, eventually, hyperfine parameters could be obtained for the ground state and for several vibrational states of  $^{12}\text{CH}_3\text{I}$  and  $^{13}\text{CH}_3\text{I}$ . Indeed, for  $\text{CH}_3\text{I}$ , hyperfine splittings are easily observable in the microwave region, and eventually also  
45 in the infrared region, due to the large values of the iodine nuclear quadrupole moment and spin of the iodine nucleus ( $I=5/2$ ).

As far as the  $\nu_6 = 1$  and  $\nu_3 = 2$  vibrational states are concerned, the most recent papers differs in strategy from the present one. The analysis of the  $\nu_6$  band of  $^{12}\text{CH}_3\text{I}$  performed by Paso and Alanko [16] by Fourier transform technique  
50 is, up to now, the most complete one existing in the infrared literature. Indeed, about 4000 transitions of the  $\nu_6$  band were identified for lines involving  $J$  up to 70 and  $K$  up to 18. However, the identification of this band was complicated by the existence of hyperfine structures which are easily observable for rotational transitions involving rather low  $J$  values as compared to the  $K$  values ( $J \leq 2$   
55  $\times K$ ). By correcting perturbed lines from their hyperfine structure, they could increase the number of rotational transitions to be included in the fit. During

the energy level calculation, the weak first order Coriolis resonance coupling together the  $v_6 = 1$  and  $v_3 = 2$  energy levels was accounted for explicitly. One has to mention that this  $v_6 = 1 \Leftrightarrow v_3 = 2$  resonance which was also considered  
60 during the investigation of the  $v_6$  band for the  $^{13}\text{CH}_3\text{I}$  isotopologue [15] was not accounted for during the investigation of the  $2v_3$  overtone band of  $^{12}\text{CH}_3\text{I}$  and  $^{13}\text{CH}_3\text{I}$  performed by Alanko et al. [18]. Later on, a new investigation of the rotational structure of the ground and  $v_6 = 1$  vibrational states of  $^{12}\text{CH}_3\text{I}$  was performed using a Doppler free double resonance technique by Carocci et al. [2].  
65 The results of these rotational measurements were combined with a large set of literature data achieved by microwave techniques [20, 21, 4, 22, 23, 1] together with infrared investigations of the  $v_6$  band. In this way a complete set of rotational and of hyperfine (quadrupole experimental rotational frequencies) was obtained both for the ground and the  $v_6 = 1$  excited states of  $^{12}\text{CH}_3\text{I}$ . However,  
70 the existence of a  $v_6 = 1 \Leftrightarrow v_3 = 2$  Coriolis resonance was not mentioned and therefore not accounted for explicitly.

Finally, a new investigation of the  $v_6$  band of  $^{12}\text{CH}_3\text{I}$  was performed very recently by Haykal et al. [17]. This new study aimed at testing the  $\text{C}_{3v}\text{TDS}$  package of the XTDS suite of programs [24] and to demonstrate its capabilities  
75 [25]. However, this preliminary study involves a rather restricted set of the  $v_6$  transitions as compared to the one used during the investigation of Paso and Alanko [16]. Indeed, as compared to reference [16] the Fourier transform spectrum used during Haykal's study [17] was recorded for a better resolution, but for a (pressure  $\times$  path length) product which is about 20% weaker than  
80 in the Paso and Alanko's study. In addition, the hyperfine structure was not accounted for, and the corresponding perturbed lines were not considered in the calculation.

The goal of the present work is to generate a set of accurate line positions for the  $v_6$  band of  $^{12}\text{CH}_3\text{I}$ . As the existence of a resonance was mentioned previously,  
85 the analysis of the  $2v_3$  band was performed in parallel. The infrared data delivered during the present analysis were combined with all existing literature data for rotational transitions within the  $v_6 = 1$  and  $v_3 = 2$  states. The

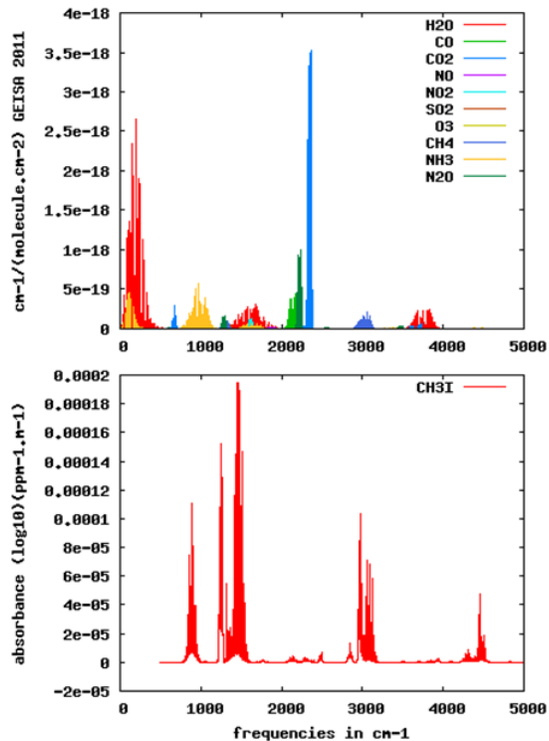


Figure 1: A comparison between a low resolution spectrum of methyl iodide, in red, issued from the PNNL (Pacific Northwest National Laboratory) with another spectrum (the upper spectrum), which combines the GEISA 2011 database for some atmospheric molecules.

theoretical model used during this study accounts for the Coriolis resonances coupling together the  $v_6 = 1 \Leftrightarrow v_3 = 2$  energy levels. Surprisingly enough, it proved necessary also to account also for the Coriolis resonance coupling the  $v_6 = 1 \Leftrightarrow v_3 = 2$  resonating levels.

Section 2 describes the experimental conditions in which five spectra were recorded. Details on the analysis procedure are listed in section 3 and of the calculations in section 4. Finally, section 5 gives a general conclusion to the present work in addition to some future perspectives. Finally, the theory of the rovibrational Hamiltonian for symmetric molecules presenting a nuclear hyperfine structure effect is recalled in Appendix A.

Table 1: Vibrational fundamental bands of  $^{12}\text{CH}_3\text{I}$ 

Mode	Symmetry	Description	$E_v^*$	Reference
$v_1$	$A_1$	$\text{CH}_3$ stretch	2971.8375	[26]
$v_2$	$A_1$	$\text{CH}_3$ deformation	1251.192826	[11]
$v_3$	$A_1$	C-I stretch	533.216836	[8]
$v_4$	$E$	$\text{CH}_3$ stretch	3060.07890	[9]
$v_5$	$E$	$\text{CH}_3$ deformation	1435.013447	[11]
$v_6$	$E$	C-I Bending	892.918	[16]
$v_6$	$E$	C-I Bending	882.91833100(427)	this work
$2v_3$		Overtone $v_3$	1059.993378	[18]
$2v_3$		Overtone $v_3$	1059.99357(1)	this work

\* Vibrational energies ( $E_v$ ) and band center ( $v$ ) in  $\text{cm}^{-1}$ .

## 2. Experimental details

100 The sample was purchased from Sigma Aldrich (Fluka Chemie GmbH, 99 %  
purity) and used without further purification. Two absorption spectra of methyl  
iodide in the mid-infrared spectral region were successively recorded. The first  
spectrum (spectrum n° 1) was recorded using the Bruker IFS125HR Fourier  
transform spectrometer of the AILES beamline at the SOLEIL Synchrotron  
105 facility. It presents a high signal to noise ratio thanks to the combination of the  
synchrotron light and of a new home-made detector developed at the AILES  
beamline of the synchrotron SOLEIL [27]. A multi-pass cell was used which  
provided a path length of 4 m. The pressure in the cell was maintained at  
0.206 mbar. The spectrum n° 1 was recorded at room temperature and with  
110 a resolution of  $0.00102 \text{ cm}^{-1}$  (according to the Bruker definition, resolution is  
 $0.9/882 \text{ cm}$ , 882 cm being the maximum optical path difference). However, the  
pressure  $\times$  path length ( $P \times L$ ) was not sufficiently high and a second spectrum  
(n° 2) was recorded with the Bruker IFS125HR spectrometer [28] located at the

LISA facility in Créteil at a resolution of  $0.0019\text{ cm}^{-1}$ . As for the spectrum n° 1,  
115 the home-made detector [27] was implemented. An optical path of 19.249 m was  
provided by a multi-pass cell. The pressure in the cell was maintained at 0.934  
mbar. This spectrum was recorded at 295 K. For both spectra the instrument  
was equipped with a KBr/Ge beam splitter and an empty cell spectrum was  
used as the reference. No apodization was applied (Boxcar function in the  
120 Bruker software). Peak positions were determined after zerofilling thanks to the  
Bruker OPUS software using the standard method for well separated peaks and  
the second derivative for partially overlapping lines.

Figures 2 and 5 represent an overview of the  $\nu_6$  band of methyl iodide, in  
the 750 - 1070  $\text{cm}^{-1}$  frequency range, recorded during this study at a low and a  
125 high  $P \times L$  products (Spectrum n° 1 and n° 2, respectively). One can see that  
the  $2\nu_3$  band is detected for the spectrum n° 2 in the figure 5, while in the case  
of the spectrum n° 1 in the figure 2, this band is too weak to be observable.  
Figures 3 and 4 with figures 6 and 7 give detailed views of the spectra n° 1  
and n° 2, respectively. It is clear when comparing figures 3 and 4 with the  
130 figures (1) and (2) from reference [16], that the high resolution of the spectrum  
n° 1 provides more information on the hyperfine patterns in comparison to this  
previous investigation.

### 3. Rovibrational analysis

According to symmetry considerations for a  $C_{3v}$ -type molecule,  $\nu_6$  is a per-  
135 pendicular band, while  $2\nu_3$  is a parallel band. The analysis proved to be straight-  
forward in general for the rather unperturbed  $\nu_6$  and  $2\nu_3$  bands. However,  
the hyperfine structure which is hardly observable for the  $2\nu_3$  parallel band, is  
a problem for  $\nu_6$  transitions involving rather weak values of the  $J$  rotational  
quantum number as compared to  $K$  ( $J \leq 2 \times K$ ). Also, for the  $P$  branches of  
140 each band, lines from several hot bands, like  $3\nu_3-\nu_3$  centered at  $1016.114\text{ cm}^{-1}$   
[18] and  $\nu_3 + \nu_6 - \nu_3$  at  $877.581\text{ cm}^{-1}$  [11], are clearly observable in the spectra  
which lead, sometimes, to complications in the analysis procedure. The first  $\nu_6$

and  $2\nu_3$  assignments were performed using the calculated predictions both for line positions and relative line intensities. As the Coriolis resonances perturbing  $\nu_6$  and  $2\nu_3$  are rather weak, these effects were not accounted for during the preliminary identification process, and the symmetric-top Hamiltonian used for the energy level calculation is described in Appendix A. As usual it includes diagonal z-Coriolis terms together with  $(\Delta\ell ; \Delta K) = (\pm 2; \pm 2)$  - type operators for the degenerated ( $\ell = 1$ )  $\nu_6 = 1$  state. In addition, the hyperfine structure was considered explicitly and Appendix A details the nuclear quadrupole and the spin-rotation operators which were considered during our calculations. However, in similarity with the references [5] and [1], the  $(\Delta J = \pm 1)$  and  $(\Delta J = \pm 2)$  matrix elements (see Appendix A) were considered through a second order perturbation treatment.

For the computation of the ground state energy levels we used the rotational and centrifugal distortion constants quoted in reference [2], and this all along the present study. Indeed, the recent ground state rotation constants quoted in table 2 of reference [17] lead to computed ground state combination differences which are in significant disagreement with the observed ones (up to  $\sim 0.005$   $\text{cm}^{-1}$  for  $J \geq 75$ ).

For the upper states, the sets of vibrational energies and rotational constants from reference [2, 18] were used as starting values for the calculation of the  $\nu_6 = 1$  and  $\nu_3 = 2$  upper state energies, respectively.

In table 4 are listed the nuclear quadrupole and spin rotational constants originating from Reference [2] and this for the ground and the  $\nu_6 = 1$  vibrational states, as for the  $\nu_3 = 2$  state they were taken from reference [1]. These hyperfine parameters are the ones used along the present study since it proved not to be faithful to refine them. In fact, the present study does not bring any new microwave data detailing the rotational structure in the ground,  $\nu_6 = 1$ , or  $\nu_3 = 2$  states. In addition, the relative vibration- rotation intensities were computed using the method described in Reference [29, 30]. More explicitly, the  $\nu_6$  and  $2\nu_3$  transition moment operators were expanded up to the first order

using the following expressions:

$${}^{33}\mu_Z^{\Delta\ell=0} = {}^{33}\mu_0^{\Delta\ell=0} \times \Phi_z \quad (1)$$

and;

$${}^6\mu_Z^{\Delta\ell=0} = {}^6\mu_0^{\Delta\ell=1} \times \Phi_x \quad (2)$$

175 Where,  $\Phi_z$  and  $\Phi_x$  are the  $Z_z$  and  $Z_x$  components of the cosines direction respectively between the  $Z$  laboratory fixed axis and the  $z$  and  $x$  molecular axes. The estimated ratio  ${}^{33}\mu_0^{\Delta\ell=0}$  and  ${}^6\mu_0^{\Delta\ell=1}$  first order terms are:

$$\frac{{}^6\mu_0^{\Delta\ell=1}}{{}^{33}\mu_0^{\Delta\ell=0}} \approx 9 \quad (3)$$

In addition, the relative contribution to the intensities of the hyperfine sub-components were computed using the standard  $6j$ - method [31, 32].

180 The assignments of the  $\nu_6$  and  $2\nu_3$  bands were performed using these predictions and accounting for the perturbations due to the hyperfine structure. As pointed out by Paso and Alanko [16], the individual vibration rotation transitions may be split into different hyperfine subcomponents, or look unsymmetrical or broadened, or, for large  $J$  values, be unaffected by such perturbation. This is observable on Figures 5 and 3 where we give, respectively, a detailed  
185 views of the band-heads for the  ${}^PQ_9$  and  ${}^RQ_9$  branches of the  $\nu_6$  band. So, we adopted the same strategy as the one applied by Paso and Alanko in reference [16]. Thus, two types of assignments were performed during the  $\nu_6$  and  $2\nu_3$  analyses: For the majority of the  $\nu_6$  lines and almost all the  $2\nu_3$  transitions, the  
190 hyperfine structure is not observable in the infrared. Examples of such “pure vibrational-rotational” transitions are given in the bottom part of Figures 5, 3 and 4. These transitions were included in our list of assignments at their estimated accuracy and without further correction. However a significant portion of the assignments for the  $\nu_6$  band, and some lines of the  $2\nu_3$  band, are lines  
195 for which the effects of hyperfine structure cannot be ignored. In this case, the observed structures correspond usually to clusters of several hyperfine subcomponents. These “hyperfine perturbed transitions” are shifted in position relative

to the corresponding “pure vibration-rotation” lines. These “perturbed lines” were only introduced in our line list after a correction of the calculated hyperfine shift. The uncertainty associated to each of these additional “perturbed” transitions accounts for their relative intensity, in the hyperfine cluster, and for the quality of the associated measured peak position in the Fourier transform spectrum. The final results of the assignments of the  $\nu_6$  and the  $2\nu_3$  bands are described in table 3. For the  $\nu_6$  band (resp. the  $\nu_3$  band), the present work represent a significant (resp. reasonable) progress in comparison to the previous infrared investigation [16] (resp. Reference [18]) where the assignments were restricted to  $J$  and  $K$  values with  $J \leq 70$  and to (resp. ( $J \leq 70$  and  $|K| \leq 12$ )). The set of the experimental vibrational-rotational energy levels for  $\nu_6 = 1$  and for  $\nu_3 = 2$  were generated by adding the line positions to the computed ground state energy levels [2]. It is important to mention that when accounting for the lines perturbed by the hyperfine structure in our list of assignments we increased the number of measured rotational energy levels which grows from 1797 (resp. 678) for the and the  $\nu_6 = 1$  state (resp. and the  $\nu_3 = 2$  state) to 2177 (resp. 692).

Table 2: Statistical analysis of energy level calculation

	$\nu_6$	$2 \nu_3$
(A) Range of observed energy levels during the analysis of the 12 $\mu\text{m}$ bands of $\text{CH}_3\text{I}$ :		
Total number of lines*	9248	1495
“Pure vibration-rotation” lines*	3976	1370
Max $J, K$	$J \leq 85, \ell \times K = -19$ to $\ell \times K = +20$	$J \leq 73,  K  \leq 15$
	$\nu_6 = 1$	$\nu_3 = 2$
Total number of rotational levels	2177	692
(B) Statistical analysis of the results of the energy level calculation:		
Calculation n <sup>o</sup>	1 <sup>▲</sup>	2 <sup>▼</sup>
	( $\nu_3 = 2, \ell = 0$ )	( $\nu_6 = 1, \ell = 1$ )
Number of levels	692	2177
$0.0 \leq \delta^* \leq 2 \times 10^{-4} \text{cm}^{-1}$	86.2 %	85.2 %
$2 \times 10^{-4} \leq \delta^* \leq 4 \times 10^{-4} \text{cm}^{-1}$	11.4 %	11.8 %
$4 \times 10^{-4} \leq \delta^* \leq 7 \times 10^{-4} \text{cm}^{-1}$	2.4 %	2.8 %
$8 \times 10^{-4} \leq \delta^* \leq 1.8 \times 10^{-3} \text{cm}^{-1}$		0.2 %
Standard deviation (in $\text{cm}^{-1}$ )	$0.12 \times 10^{-3}$	$0.17 \times 10^{-3}$
Calculation n <sup>o</sup>		3 <sup>◆</sup>
	( $\nu_3 = 2, \ell = 0$ )	( $\nu_6 = 1, \ell = 1$ )
Number of levels	692	2177
$0.0 \leq \delta^* \leq 2 \times 10^{-4} \text{cm}^{-1}$	86.8 %	86.2 %
$2 \times 10^{-4} \leq \delta^* \leq 4 \times 10^{-4} \text{cm}^{-1}$	12.7 %	11.4 %
$4 \times 10^{-4} \leq \delta^* \leq 6.2 \times 10^{-4} \text{cm}^{-1}$	0.4 %	2.4 %
Standard deviation (in $\text{cm}^{-1}$ )		$0.14 \times 10^{-3}$
(C) Microwave rotational transitions within the ( $\nu_6 = 1, \ell = 1$ ) and ( $\nu_3 = 2, \ell = 0$ ) vibrational states:		
$\nu_3 = 2$	$\nu_6 = 1$	both levels
95	160	255
$0.0 \text{ MHz} \leq 92.63\% \leq 1.0 \text{ MHz}$	$0.0 \text{ MHz} \leq 92.50\% \leq 1.0 \text{ MHz}$	$0.0 \text{ MHz} \leq 92.55\% \leq 1.0 \text{ MHz}$
$1.0 \text{ MHz} \leq 6.32\% \leq 2.0 \text{ MHz}$	$1.0 \text{ MHz} \leq 7.50\% \leq 2.0 \text{ MHz}$	$1.0 \text{ MHz} \leq 7.06\% \leq 2.0 \text{ MHz}$
$2.0 \leq 1.05\% \leq 4.0 \text{ MHz}$	$2.0 \text{ MHz} \leq 0.00\% \leq 4.0 \text{ MHz}$	$2.0 \text{ MHz} \leq 0.39\% \leq 4.0 \text{ MHz}$

\* Pure “vibration-rotation” transitions and “hyperfine perturbed” subcomponents (see text).

\* Pure “vibration-rotation” transitions (see text).

\*  $\delta = |E_{obs} - E_{calc}|$ .

▲ As an isolated state.

▼ As an isolated state.

◆ ( $\nu_6 = 1, \ell = 1$ ) accounting for  $\alpha(\Delta\ell = \pm 1; \Delta K = \pm 2)$  and for  $C_x(\Delta\ell = \pm 1; \Delta K = \pm 1)$  Coriolis resonance with ( $\nu_2 = 1, \ell = 0$ ) and ( $\nu_3 = 2, \ell = 0$ ) respectively.

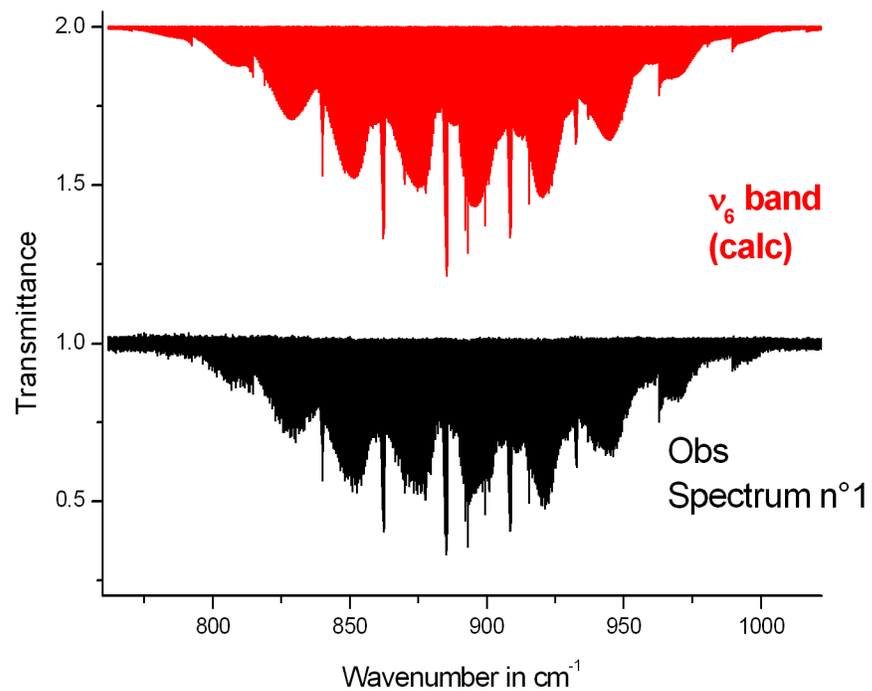


Figure 2: Overview of the absorption spectrum of  $^{12}\text{CH}_3\text{I}$  (spectrum n°1) in the 740-1055  $\text{cm}^{-1}$  spectral region. "Obs" trace: observed "spectrum n°1" recorded with a resolution of  $1.02 \times 10^{-3} \text{ cm}^{-1}$ , an optical path length of 4 m, a pressure of 0.206 mbar and a stabilized temperature of 295 K. "Calc" trace: synthetic spectra, calculated for the  $\nu_6$  and  $2\nu_3$  bands at the same experimental conditions using the line lists generated in this work. The observed and synthetic spectra are in transmission and shifted for clarity.

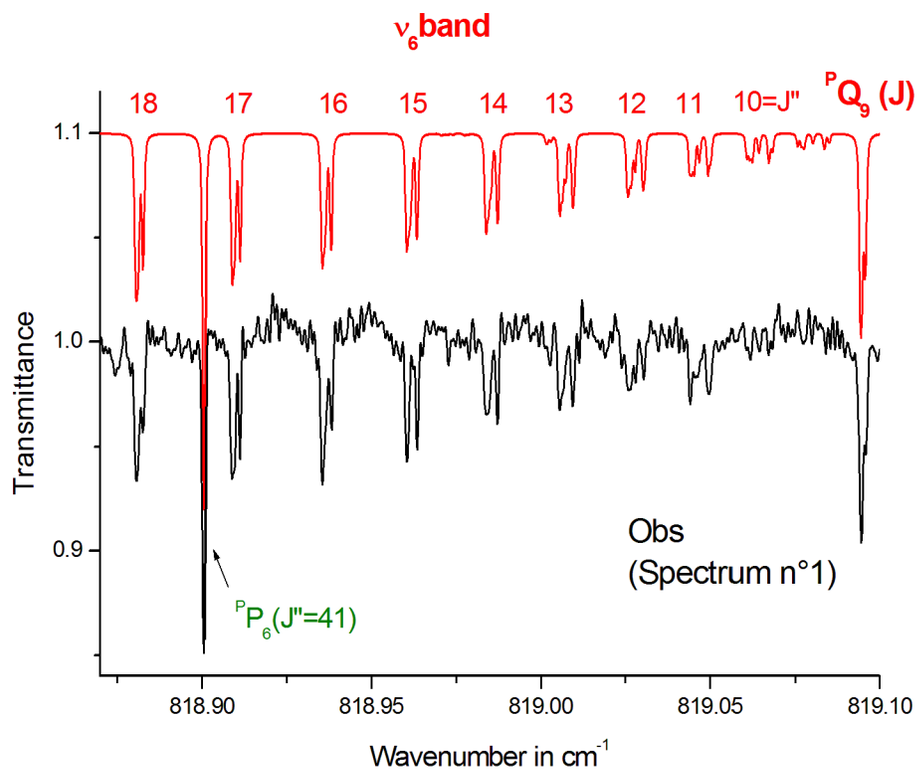


Figure 3: Portion of the absorption spectrum of  $^{12}\text{CH}_3\text{I}$  (spectrum n°1) in the  $818\text{ cm}^{-1}$  spectral region. The observed and synthetic spectra are in transmission and are shifted for the sake of clarity. Some assignments are given. For the  $^P Q_9$  band-head, the hyperfine structure has to be accounted for.

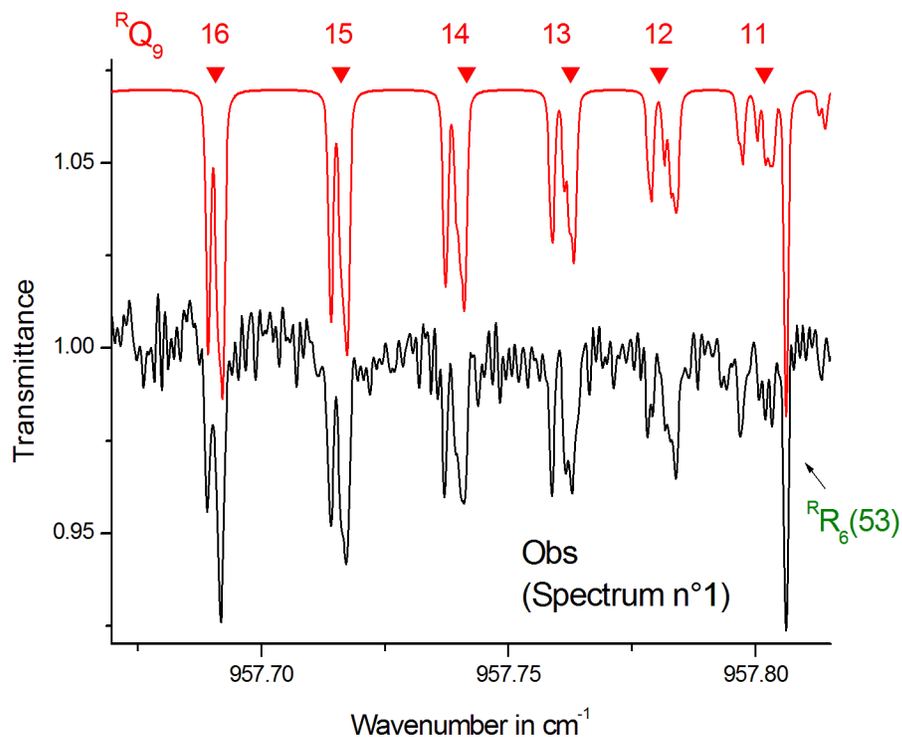


Figure 4: Portion of the absorption spectrum of  $^{12}\text{CH}_3\text{I}$  (spectrum n°1) in the  $957.7\text{ cm}^{-1}$  spectral region. The observed and synthetic spectra are in transmission and are shifted for the sake of clarity. Some assignments are given. For the  $^R\text{Q}_9$  band-head, the hyperfine structure has to be accounted for.

#### 4. Energy levels calculations

The infrared energy levels, achieved during the present study, were included  
 220 in a least squares fit calculation together with the rotational transitions available  
 in the literature, within the  $\nu_6 = 1$  [4, 5, 1, 2] and the  $\nu_3 = 2$  [1] vibrational  
 states, which were achieved through different microwave techniques. Each set  
 of data were accounted for at their estimated accuracy in a process which is

described in details in Reference [2]. The model used during this calculation is  
225 described in Appendix A. The hyperfine structure is accounted for during the  
computation of the rotational transitions within  $v_6 = 1$  and  $v_3 = 2$  states. As  
pointed out previously, it proved unnecessary to refine the electric quadrupole  
and spin-rotation parameters which were fixed to the values given in references  
[2] and [1] for  $v_6 = 1$  and  $v_3 = 2$ , respectively.

230 Two types of energy levels calculations were performed. During our first two  
calculations, calculation n° 1 and calculation n° 2, we assumed that the  $v_3 = 2$   
and  $v_6 = 1$  states can be considered as isolated ones, with no significant pertur-  
bation. As far as the  $v_3 = 2$  state is concerned, it proved that we were not able  
to improve significantly the quality of the calculation, for the energy levels and  
235 for the microwave transitions, and this in comparison to the calculations previ-  
ously performed by Alanko et al. [18] and by Wlodarczak et al. [1]. Therefore,  
the parameters quoted in table 4 are those given by Alanko et al. [18], and we  
accomplished a simple readjustment of the  $v_3 = 2$  rotational parameters, and  
this is due to the fact that we used a different set of ground state parameters  
240 [33] in respect to the one used by Alanko et al. [18]. On the other hand, during  
the calculation n° 2, we had to refine significantly the  $v_6 = 1$  parameters in  
order to reproduce the observed energy levels. The resulting parameters to-  
gether with their associated uncertainties are collected in table 4 in the case  
 $v_6 = 1$  state. This table compares the values obtained during the present work  
245 with the parameters achieved during the study conducted by Carocci et al.  
[2]. In comparison to this previous investigations [2] we considered one addi-  
tional flexible parameter ( $\eta_{JKK}$ ) during the energy level calculation. This was  
expected, since the present range of rotational energy levels exceeds the one  
achieved during these previous investigations. In the case of the infrared levels,  
250 the table 3 gives a statistical analysis of the (observed - calculated) values. It  
seems relevant to compare the quality of the present energy calculation with  
the one achieved when using the parameters (band centers and rotational con-  
stants) from references [2, 18]. This is not the case for the  $v_6 = 1$  state, and  
figure 8 shows a plot of the Obs-Calc differences for the  $v_6 = 1$  energy levels

255 as a function of the  $K$  and  $J$  rotational quantum numbers ( $K + (J/100)$  in our case). It is clear that the centrifugal distortion constants are more reliable in the present work than in the reference [2] because we used a significantly larger set of energy levels. However, it is clear also that some series of energy levels could not be reproduce correctly. For example for the energy levels involving  
 260  $(\ell \times k) = -17$ , the disagreement grows up with  $J$  to reach  $-1.7 \times 10^{-3} \text{ cm}^{-1}$  for  $J = 37$ . Therefore, it was clear that resonances due to other vibrational states may occur. According to the recommendation given by Paso and Alanko [16] we first performed a new calculation accounting for the  $(\Delta \ell ; \Delta K) = (\pm 1; \pm 1)$  Coriolis resonance coupling the energy levels from  $v_6 = 1$  with those of the  
 265  $v_3 = 2$  state. We observe some progress in the quality of our fit, but still, the series of energy levels in  $(\ell \times k) = -17$  were not reproduced satisfactorily. We presumed that some series belonging to the  $v_2 = 1$ ,  $(v_3 = 1; v_6 = 1)$  or  $v_5 = 1$  states of  $^{12}\text{CH}_3\text{I}$  [11] could possibly be involved in this resonance. After trials and errors, it appeared that the  $v_2 = 1$  state was responsible for this resonance  
 270 through an  $\alpha(\Delta l = \pm 1; \Delta K = \pm 2)$  type Coriolis resonance. Therefore, in the final performed calculation we took into account the resonances involving the  $v_6 = 1$  energy levels with those from both  $v_3 = 2$  and  $v_2 = 1$  levels. For the  $v_2 = 1$  vibrational state, the band center and the  $(v=2\text{X}-v=0\text{X})$  vibrational dependences of the rotational constants were maintained to the values proposed by  
 275 Alanko [11]. The final set of band centers, rotational and interaction constants are given in table 4 together with their associated uncertainties. It is important to mention that for the  $\alpha(\Delta l = \pm 1; \Delta K = \pm 2)$  resonance, only the second order term,  ${}^6,2\alpha_K$ , could be determined. We presume that this is due to the fact that our calculation does not account for the resonances which couple together the  
 280  $v_2 = 1$  energy levels with those of the  $(v_3 = 1; v_6 = 1)$  and  $v_5 = 1$  states. These resonances are rather weak. To give an order of magnitude, the percentage of mixing of the wave-functions for the following resonating series are, for  $J = 75$ :

- $v_6 = 1, (\ell \times K) = -17 \Leftrightarrow v_2 = 1, K = 15$  : mixing of 0.5 %
- $v_6 = 1, (\ell \times K) = +18 \Leftrightarrow v_2 = 1, K = 16$  : mixing of 0.5 %

- 285 •  $v_6 = 1, (\ell \times K) = +20 \Leftrightarrow v_3 = 2, K = 19$  : mixing of 0.6 %
- $v_6 = 1, (\ell \times K) = +19 \Leftrightarrow v_3 = 2, K = 18$  : mixing of 0.1 %

However, provided that the analysis is pursued up to rather high values of the rotational quantum numbers, even for the lower energy vibrational state, it proved that the  $v_6 = 1$  level of  $^{12}\text{CH}_3\text{I}$  cannot be considered as a true "isolated" state because its higher energy levels are coupled with those from other states,  $v_3 = 2$  and  $v_2 = 1$ . Such situation occurs for many molecules, particularly in the case of  $^{12}\text{CH}_3\text{F}$  [34] and HDCO [35], for example. Table 3 gives a statistical analysis on the results of this energy level calculation (Calculation n° 3). In figure 8, the Obs-Calc differences, in energy, are plotted as a function of the rotational quantum numbers ( $K + (J/100)$ ). It is clear that the progress is significant, especially for the series in  $(\ell \times K)=15$  and in  $(\ell \times K)=-17$ . Finally, the (C) part of table 3 describes the results of the present calculation concerning the data existing in the literature for the rotational transitions within the  $v_6 = 1$  and  $v_3 = 2$  vibrational states. There is no significant differences in term of quality of the fit between the two types of computations (calculations n° 1 and n° 2 and calculation n° 3). Furthermore, these results in case of the microwave data are also quite similar in quality to those achieved previously in the literature when using the previous set of vibrational energies, rotational and hyperfine constants [2, 18, 1].

305 To check the quality of our calculation, a line list in positions and relative intensities was generated for the  $v_6$  and  $2v_3$  bands. Therefore, we used the vibration-rotation parameters quoted in table 4 for the computation of the  $v_6 = 1$  and  $v_3 = 2$  rotation energy levels, and the rotation constants of reference [2] in the case of the ground state. The hyperfine structure of the lines was accounted for by using the hyperfine parameters quoted in table 4 [2, 1]. The relative line intensities were computed as described previously.

Figures 2 and 5 give overviews of the  $v_6$  and  $2v_3$  bands while figures 3, 4, 6 and 7 compare the observed and calculated spectra in several regions of these bands. It is clear that the agreement is very satisfactorily for  $v_6$  lines even when

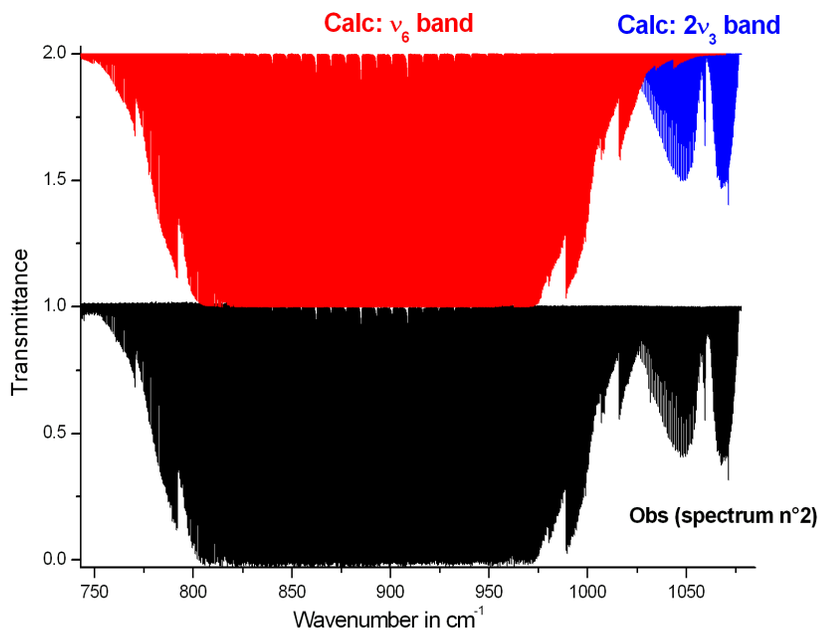


Figure 5: Overview of the absorption spectrum of  $^{12}\text{CH}_3\text{I}$  (spectrum n°2) in the 740-1020  $\text{cm}^{-1}$  spectral region. “Obs” trace: observed “spectrum n°2” recorded with a resolution of  $0.0019 \text{ cm}^{-1}$ , an optical path length of 4 m, a pressure of  $0.34 \text{ mbar}$  and a stabilized temperature of 293 K. “Calc ” trace: synthetic spectra, calculated for the  $\nu_6$  band at the same experimental conditions using the line lists generated in this work. The observed and synthetic spectra are in transmission and shifted for clarity.

315 the hyperfine structure is quite observable (figures 3,4 and 6). The agreement is as well satisfactory for  $\nu_6$  transitions involving high  $K$  values and for  $2\nu_3$  transitions (figure 7).

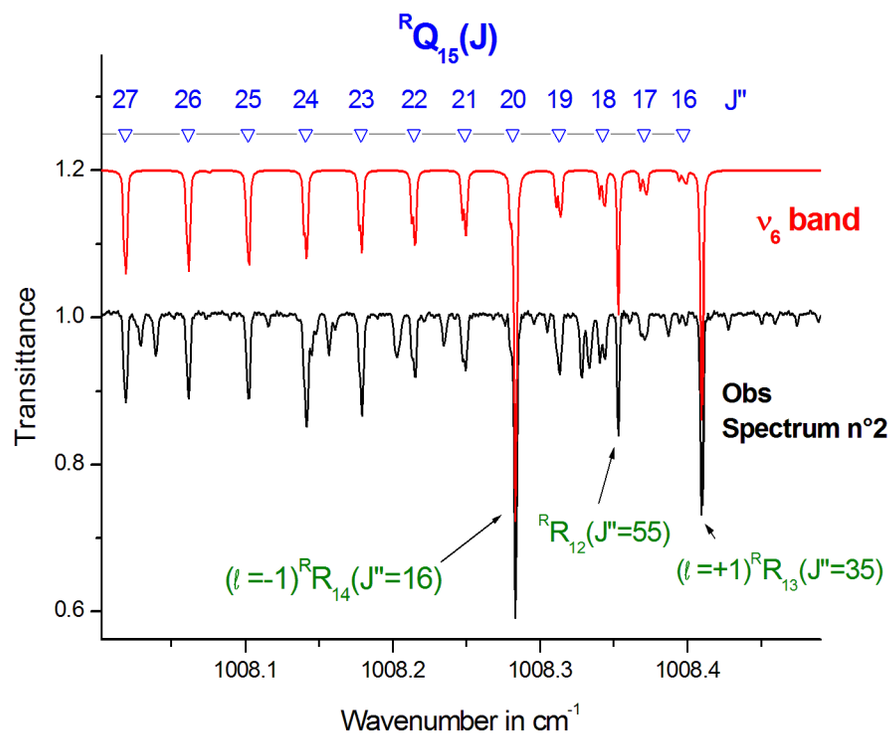


Figure 6: Portion of the absorption spectrum of  $^{12}CH_3I$  (spectrum n°2) in the  $1008.3\text{ cm}^{-1}$  spectral region (spectrum "2"). The observed and synthetic spectra are in transmission and shifted for clarity. Some assignments are given. For the  $RQ_{15}$  band-head, the hyperfine structure has to be accounted for.

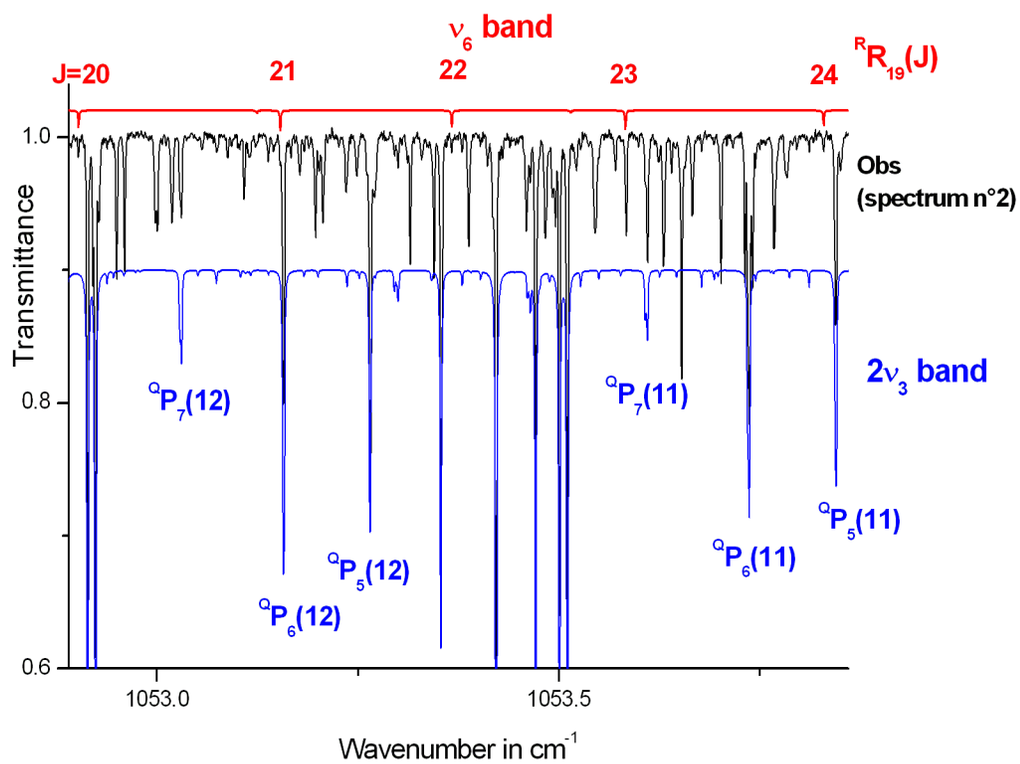


Figure 7: Portion of the absorption spectrum of  $^{12}\text{CH}_3\text{I}$  (spectrum n°2) in the  $1053.5 \text{ cm}^{-1}$  spectral region (spectrum "2"). The observed and synthetic spectra are in transmission and shifted for clarity. Some assignments are given for the  $2\nu_3$  bands, together with lines from the  $\nu_6$  band involving high  $K = 20$  in the upper energy levels.

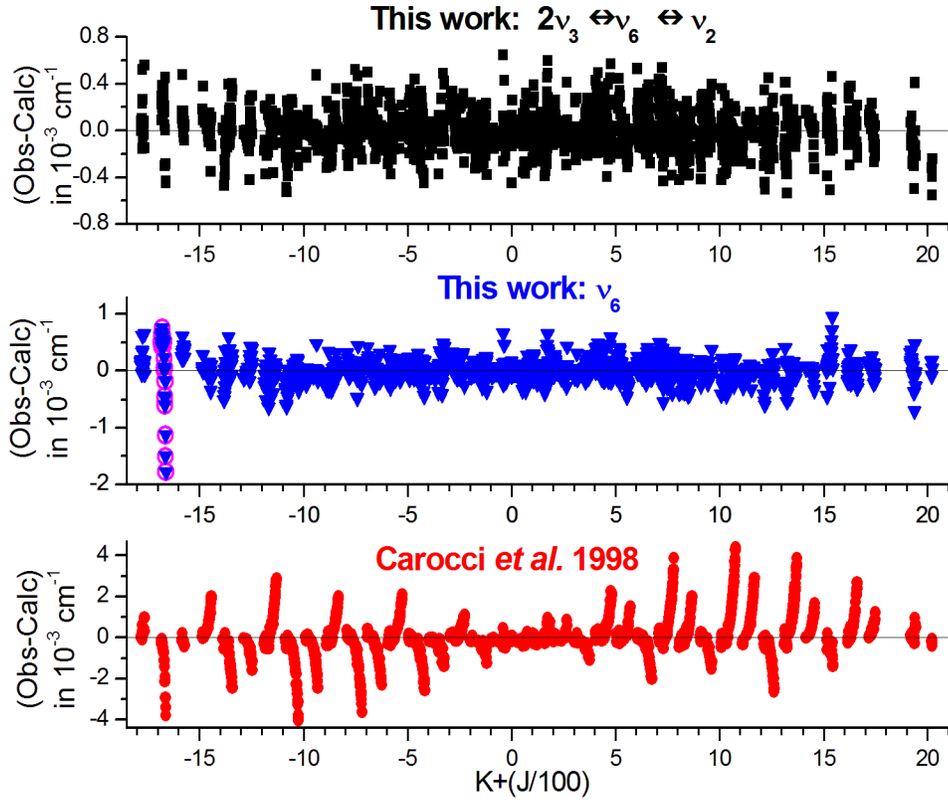


Figure 8: Plots of the (observed- calculated) energy levels for the  $v_6 = 1$  as a function of  $[(\ell \times K) + J/100]$  and this for different sets of  $v_6 = 1$  parameters (band centers, rotational and coupling constants). Note the difference of scale in the y-axes for the three traces. Lower trace:  $v_6 = 1$  as an isolated state and the parameters of reference [2]. Medium trace :  $v_6 = 1$  as an isolated state and the parameters are taken from the table 4 of this work (calculation n° 1). Noticeable disagreements are obtained for energy levels involving  $K = -17$  (open circles). Upper trace:  $v_6 = 1$  in resonance with  $v_2 = 1$  and  $v_3 = 2$  and the parameters from table 4 of this work (calculation n°3).

Table 3: Vibrational and rotational constants for the ground and the  $v_3 = 2$  vibrational states (Calculation n° 1).

	Calculation n° 1	
	Ground	( $v_3 = 2, \ell = 0$ )
	[2]	[18] <sup>*</sup>
$E_v$ ( $\text{cm}^{-1}$ )		1059.993378
A (GHz)	155.1106938	154.703709
B (GHz)	7.501275753	7.39248335
$D_J$ (kHz)	6.30762	6.392491
$D_{JK}$ (kHz)	98.7657	98.6572
$D_K$ (MHz)	2.62714	2.594741
$H_J$ (mHz)	-3.431	<sup>a</sup>
$H_{JK}$ (mHz)	58.8	<sup>a</sup>
$H_{KJ}$ (Hz)	4.602	<sup>a</sup>
$H_K$ (Hz)	135.6	<sup>a</sup>

<sup>a</sup> Fixed to the ground state value [2].

<sup>\*</sup> Calculated from the ground state parameters of reference [2] and the ( $X''$ - $X'$ )  $2\nu_3$  vibrational dependence of the rotational constants.

Table 4: Hyperfine constants for the ground and the ( $v_6 = 1, \ell = 1$ ) and ( $v_3 = 2, \ell = 0$ ) vibrational states of CH<sub>3</sub>I.

	Ground <sup>a</sup>	( $v_6 = 1, \ell = 1$ ) <sup>a</sup>	( $v_3 = 2, \ell = 0$ ) <sup>b</sup>
$eqQ$ (GHz)	-1.9341306	-1.94034	1.93478
$X_J$ (kHz)	-1.55	-1.81	
$X_K$ (kHz)	-33.36	-37.6	
$X_d$ (kHz)	22.45	20.83	
$\eta$ (kHz)		-1.19	
$C_N$ ( $\times 10^{-3}$ )	-17.398	17.62	-13.92
$C_K$ (kHz)	-17.10	-28.53	

<sup>a</sup> Reference [2].

<sup>b</sup> Reference [1].

Table 5: Vibrational energies, rotational and centrifugal distortion constants for the ( $v_6 = 1, \ell = 1$ ) Considered as an isolated state.

	$v_6 = 1$	
	This work (calc. n° 2)	Reference [2]
$E_v$ ( $\text{cm}^{-1}$ )	882.918 38 (1)	882.918 206 4 (6)
$(A_\zeta)$ (GHz)	32.852 43 (32)	32.852 482 (22)
$\eta_J$ (kHz)	200.899 (41)	200.933 (33)
$\eta_K$ (MHz)	4.153 6 (11)	4.161 92 (76)
$\eta_{JJ}$ (mHz)	159.4 (68)	157.4 (86)
$\eta_{JK}$ (Hz)	9.30 (63)	9.04 (24)
$\eta_{KK}$ (Hz)	-760.4 (40)	-792.8 (32)
$\eta_{JKK}$ (mHz)	6.8 (26)	–
$q_{22}^6$ (MHz)	-1.480 52 (14)	-1.480 5 (13)
$q_{22}^{6,J}$ (Hz)	4.515 (42)	4.513 (32)
A (GHz)	156.150 901 1 (11)	156.150 936 9(53)
B (GHz)	7.477 662 25 (24)	7.477 663 12 (22)
$D_J$ (kHz)	6.349 61 (14)	6.349 91 (18)
$D_{JK}$ (kHz)	98.799 7 (61)	98.825 6 (45)
$D_K$ (MHz)	2.729 30 (10)	2.729 748 (55)
$H_J$ (mHz)	-3.715 (18)	-3.697 (34)
$H_{JK}$ (mHz)	68.33 (98)	69.2 (28)
$H_{KJ}$ (Hz)	3.824 (26)	4.010 (27)
$H_K$ (Hz)	141.58 (25)	143.14 (18)

Table 6: Vibrational and rotational constants for the  $v_2 = 1$ ,  $v_3 = 2$ , and  $v_6 = 1$  interacting vibrational states of  $^{12}\text{CH}_3\text{I}$ .

	calculation n° 3		
	$v_2 = 1$	$v_6 = 1$	$v_3 = 2$
$E_v$ ( $\text{cm}^{-1}$ )	125 1.192 826*	882.918 37 (1)	105 9.993 58 (3)
$(A_\zeta)$ (GHz)		32.852 519 (41)	
$\eta_J$ (kHz)		201.009 (46)	
$\eta_K$ (MHz)		4.159 6 (14)	
$\eta_{JJ}$ (mHz)		166.9( 79)	
$\eta_{JK}$ (Hz)		8.87 (34)	
$\eta_{KK}$ (Hz)		-775.97 (177)	
$\eta_{JKK}$ (mHz)			
$q_{22}^6$ (MHz)		-1.479 68 (17)	
$q_{22}^{6,J}$ (Hz)		4.521 (48)	
A (GHz)	155.725 385**	156.150 947 (12)	154.703 776 (29)
B (GHz)	7.475 513 96**	7.477 664 07 (25)	7.392 479 45 (65)
$D_J$ (kHz)	6.356 622**	6.349 66 (14)	6.392 48 (16)
$D_{JK}$ (kHz)	99.387 5**	98.821 8 (61)	98.668 (10)
$D_K$ (MHz)	2.712 53**	2.729 84 (11)	2.595 24 (19)
$H_J$ (mHz)	<sup>a</sup>	-3.706 (19)	<sup>a</sup>
$H_{JK}$ (mHz)	<sup>a</sup>	71.6 (10)	<sup>a</sup>
$H_{KJ}$ (Hz)	<sup>a</sup>	4.109 (29)	<sup>a</sup>
$H_K$ (Hz)	<sup>a</sup>	142.97 (28)	<sup>a</sup>
$^{33,6}C_x$ (MHz)		92.98(16)	
$^{2,6}\alpha_K$ (Hz)		9.01(41)	

<sup>a</sup> Fixed to the ground state value.

\* Reference [11].

\*\* Calculated from the ground state parameters of reference [2] and the  $(X''-X')$   $\nu_2$  vibrational dependence of the rotational constants [11].

## 5. conclusion

During this work a new analysis of the  $\nu_6$  and  $2\nu_3$  bands was performed using high resolution Fourier transform spectra. Along the assignment process, the hyperfine structure was accounted for during the analysis procedure as well as during the energy level calculation. In consequence, a large set of experimental energy levels was accurately measured for the  $\nu_6 = 1$  and the  $\nu_3 = 2$  states. The energy levels calculation was performed by considering, in a preliminary step, the  $\nu_6 = 1$  and the  $\nu_3 = 2$  as an isolated states. In an advanced stage of the study, it proved necessary to account for the resonances which were found to couple the  $\nu_6 = 1$  energy levels with those of  $\nu_3 = 2$  and  $\nu_2 = 1$ .

## 6. Acknowledgements

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## A. Appendix.

Table 7: Hamiltonian matrix used to describe the  $\{(\nu_6 = 1; \ell = \pm 1), (\nu_3 = 2; \ell = 0), (\nu_2 = 1; \ell = 0)\}$  resonating states of  $\text{CH}_3\text{I}$ .

	$(\nu_6 = 1; \ell = \pm 1)$	$(\nu_3 = 2; \ell = 0)$	$(\nu_2 = 1; \ell = 0)$
$(\nu_6 = 1; \ell = \pm 1)$	${}^6W + {}^6Hyp$	c.c.	c.c.
$(\nu_3 = 2; \ell = 0)$	$C_x(\Delta\ell = \pm 1; \Delta K = \pm 1)$	${}^{33}W + {}^{33}Hyp$	
$(\nu_2 = 1; \ell = 0)$	$\alpha(\Delta\ell = \pm 1; \Delta K = \pm 2)$		${}^2W + {}^2Hyp$

(A) Vibrational rotational operators : The  $W(v; \ell = \pm 1)$  are rotational diagonal in v-operators, including both diagonal and non diagonal  $\ell$  terms. The  $(\Delta\ell; \Delta K) = (0; 0)$  z-type Coriolis and Anharmonic rotational operator.

(1)  $W(v; \ell = \pm 1)$  v-diagonal operators :

$$\begin{aligned} \langle v; \ell, JK | \mathcal{H} | v; \ell, JK \rangle &= E_v + B_v J(J+1) + (A_v - B_v) K^2 \\ &- D_J^v J^2(J+1)^2 - D_{JK}^v K^2(J+1) - D_K^v K^4 + H_J^v J^3(J+1)^3 \\ &+ H_{JK}^v J^2(J+1)^2 K^2 + H_{KJ}^v J(J+1) K^4 + H_K^v K^6 + L_{KJ} K^4 J^2(J+1)^2 \\ &+ [-2A_{\zeta v} + \eta_J^v J(J+1) + \eta_K^v K^2 + \eta_{JJ}^v J^2(J+1)^2 + \eta_{JK}^v K^2(J+1) + \eta_{JKK}^v K^4 J(J+1)] K\ell \end{aligned}$$

345 The diagonal z-Coriolis terms ( $A_{\zeta v}$ ) and their expansion ( $\eta^v$  ect..) vanish for all  $\ell = 0$  vibrational states ( $\nu_2 = 1$  and  $\nu_3 = 2$ ).

For the ( $v_6 = 1; \ell = \pm 1$ ) vibrational state, the  $(\Delta\ell; \Delta K) = (\pm 2; \pm 2)$  - type of operator was taken into account. <sup>3</sup>

$$\begin{aligned} \langle v; \ell = \pm 1, JK | \mathbf{H} | v; \ell = \pm 1, JK \pm 2 \rangle &= \\ 2 ( q_{22}^v + q_{22}^{v,J} J(J+1) ) \times F_2(J, K, K \pm 1) \end{aligned}$$

(2)  $C(\pm 1; \pm 1)$  Coriolis off diagonal in v operator.<sup>4</sup>

$$\begin{aligned} \langle v_6 = 1; \ell = \pm 1, JK | \mathbf{H} | v_2 = 1; \ell = 0, JK \pm 2 \rangle &= \\ \pm ( {}^{6,2}\alpha_0 + {}^{6,2}\alpha_K (K^2 + (K \pm 2)^2) ) F_2(J, K, K \pm 2) \end{aligned}$$

$$\begin{aligned} 350 \langle v_6 = 1; \ell = \pm 1, JK | \mathbf{H} | v_3 = 2; \ell = 0, JK \pm 1 \rangle &= {}^{6,33} C_0 F_1(J, K, K \pm 1) \\ F_2(J, K, K \pm 2) &= \sqrt{J(J+1) - (K(K \pm 1))(J(J+1)) - (K \pm 1)(K \pm 2)} \\ F_1(J, K, K \pm 2) &= \sqrt{J(J+1) - K(K \pm 1)} \end{aligned}$$

(B) Hyperfine operators ( $Hyp(v)$ ): For  $\text{CH}_3\text{I}$ , two type of hyperfine operators are to be considered in order to describe the observed hyperfine structure:

355 1. The nuclear quadrupole operator  $H_{Quadr}$ :

- Diagonal term :

$$\langle J, K, F | H_{Quadr} | J, K, F \rangle =$$

<sup>3</sup> Note the different convention used in reference [2] for the  $(\Delta\ell; \Delta K) = (\pm 2; \pm 2)$  off-diagonal operator and as a consequence for the  $q_{22}^{WW}$  parameter :

$$\langle v; \ell = \pm 1, JK | \mathbf{H} | v; \ell = \pm 1, JK \pm 2 \rangle = 1/2 F_2(J, K, K \pm 2) \times (q_{22}^{WW} + \dots)$$

<sup>4</sup>The  $\alpha(\Delta\ell = \pm 1; \Delta K = \pm 2)$  Coriolis operator is set at zero when the Coriolis resonances are not considered explicitly.

$$(eqQ'' + X_J''J(J+1) + X_K''K^2) \times \left( \frac{3K^2}{J(J+1)} - 1 \right) + X_d'' \frac{K^2(4K^2-1)}{J(J+1)} Y(I, J, F)$$

where;

$$Y(I, J, F) = \frac{3C(C+1) - I(I+1)J(J+1)}{8I(2I-1)(2J-1)(2J+3)}$$

360 where;

$$C = F(F+1) - I(I+1) - J(J+1)$$

- Non diagonal terms :

$$\langle J, K, F | H_{Quadr} | J+1, K, F \rangle = \frac{3eqQ'' K [F(F+1) - I(I+1) - J(J+2)]}{8I(2I-1)J(J+2)}$$

$$\times \left[ \left( 1 - \frac{K^2}{(J+1)^2} \right) (I+J+F+2) \times \frac{(J+F-I+1)(I+F-J)(I+J-F+1)}{(2J+1)(2J+3)} \right]^{\frac{1}{2}}$$

and;

$$\langle J, K, F | H_{Quadr} | J+2, K, F \rangle = \frac{3eqQ''}{16I(2I-1)(2J+3)}$$

$$\times \left[ \left( 1 - \frac{K^2}{J^2(J+1)^2} \right) \left( 1 - \frac{K^2}{J^2(J+2)^2} \right) (I+J+F+3) \right]$$

$$\times (I+J+F+2)(I+J-F+2)(I+J-F+1)(J+F-I+2)$$

$$\times \frac{(J+F-I+1)(I-J+F)(I+F-J-1)}{(2J+1)(2J+5)}^{\frac{1}{2}}$$

365

2. The spin-rotation operator  $H_{SR}$ :

$$H_{SR} = -\frac{1}{2} \left[ C_N'' + (C_K'' - C_N'') \frac{K^2}{J(J+1)} \right] \times F(F+1) - I(I+1) - J(J+1)$$

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