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Conformational landscape and internal dynamics of limona ketone, a key oxidation product of limonene



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ABSTRACT

The rotational spectrum of limona ketone (4-acetyl-1-methyl-1-cyclohexene), a key oxidation product of limonene, was recorded in the gas phase using a Fourier transform microwave spectrometer coupled to a supersonic jet expansion over the 4–20 GHz range. To display a complete view of the conformational landscape of this relatively flexible system, the relative stability and inter-conversion barriers between conformers were explored by electronic structure calculations. The lowest energy equatorial conformer was detected, and its spectroscopic molecular parameters were determined from the analysis of the spectrum. The internal rotation of the methyl group attached to the aldehyde moiety was taken into account in the analysis. The internal dynamics of the methyl group within the acetyl moiety was quantitatively described. Experimental lines of both A and E symmetry were fitted to experimental accuracy, and the experimental barrier height of the methyl group internal rotation is discussed with the support of quantum chemical calculations.

1. Introduction

Biogenic volatile organic compounds (BVOCs) are abundant in the Earth's atmosphere with a global emission by vegetation estimated at 760 Tg(C)yr⁻¹ [1]. Many atmospheric BVOCs are terpenes, that primarily consist of two bonded isoprene subunits (C5H8). Limonene is the second major terpene released naturally in the atmosphere. It is mostly emitted from lemon trees, orange and citrus fruits. Its ozonolysis reaction is considered as one of the primary sources of the formation of secondary organic aerosols (SOA) [2-5]. Because of its two chemically different double bonds, an endocyclic and an exocyclic double bond, the limonene ozonolysis has been the subject of many investigations [6–11], using a variety of techniques, in order to identify the products of reaction and evaluate the thermodynamic and kinetic parameters. In particular, the first-generation products of the limonene ozonolysis may be unsaturated and exhibit high reactivity for further oxidation. Baptista et al. [6], using density functional methods (DFT), theoretically investigated the reaction steps of the ozonolysis with a focus on primary ozonide formation. The oxidation regime and SOA composition in limonene ozonolysis was also investigated [8]. Despite its importance as precursor of SOAs, only a few studies concerning the structure of limonene in the gas phase at the microwave accuracy are reported in the literature. Avilés-Moreno et al. [12] identified different conformations of limonene in the gas phase. Loru et al. [13] reported conformational flexibility of limonene oxide and characterized up to five conformers from the analysis of the pure rotation spectrum. More

recently, Loru et al. [14] characterized axial conformers of limonene, carvone and perillaldehyde.

Limona ketone (C9H14O, 4-acetyl-1-methyl-1-cyclohexene, LK), referred also as keto-limonene, is one of the products of limonene oxidation [15]. Arey et al. [16] investigated products formed from the OH radical-initiated reactions of β -pinene and limonene in the presence of NO_x. By using gas chromatography with flame ionization detection (GC-FID), they analyzed the GC-FID traces and found two significant product peaks from d-limonene, one of them was identified as LK. Gallimore et al. [17] investigated the oxidation of limonene in the Cambridge Atmospheric Simulation Chamber (CASC) and observed that LK is a short-lived oxidation product produced at low concentration and it is removed by further ozonolysis. LK is also identified as a product of ozonolysis of limonene. In 2005, Leungsakul et al. [18] proposed and tested experimentally a kinetic mechanism for predicting the formation of SOA from the reaction of limonene with O3. Limononaldehyde was the major identified product followed by LK, keto-limonaldehyde, limononic acid and keto-limononic acid. Ham et al. [19] were able to detect five carbonyl compounds from limonene ozonolysis: LK, 7-hydroxy-6-oxo-3-(prop-1-en-2yl) heptanal, 3-isopropenyl-6-oxo-heptanal, 2-acetyl-5-oxohexanal and 3-acetyl-6-oxoheptanal. As for the firstgeneration products of limonene ozonolysis, LK exhibits high reactivity. It was found that it reacts with ozone in the Hyytiälä forest in spring

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Fig. 1. Scan in low resolution mode ($250 \,\text{kHz}$ steps) between 8000 and 9000 MHz of limona ketone, with insets at high resolution ($1.8 \,\text{kHz}$) for assigned peaks. Rotational lines are identified with $J_{K,K}$ label and the split components associated with internal rotation are labeled with A/E symmetry. The star symbol indicates unassigned peaks.

time [20]. Moreover, Donahue et al. [21] showed that LK itself generates SOA with an efficiency that is statistically indistinguishable from α pinene, probably due to the fact that they have a 1-methylcyclohexene moiety in common.

Fourier transform microwave (FTMW) spectroscopy in jet-cooled conditions is a powerful technique for investigating terpenoids, which are oxygenated derivatives of isoprene units. The first investigation of a group of terpenes using this technique was reported in 1978 [22]. The characterization of camphor using FTMW has been published more recently, in 2003 [23]. Since then, jet-cooled FTMW spectroscopy has been used to characterize a number of monoterpenes such as α -pinene [24], β -pinene [25], camphene [26], verbenone [27] and some of their oxidation products, namely perillaldehyde [28], nopinone [25], myrtenal [29] and fenchol [30]. Other monoterpenes obtained from oxydation and studied at microwave accuracy are fenchone [31], menthol and (iso)menthone [32], thymol and carvacrol [33].

In this paper, we employed FTMW spectrometers to record the rotational spectrum of LK in jet-cooled conditions. The lowest energy conformer was detected and its molecular parameters are determined from the analysis of the spectrum. The internal rotation of the methyl group attached to the aldehyde group is taken into account in the analysis. Experimental lines for both A and E states are fitted to instrumental accuracy and the experimental barrier height of the methyl group internal rotation is discussed with the support of quantum chemical calculations. In addition, the relative stability and inter-conversion barriers between conformers are explored by electronic structure calculations in order to display a complete view of the conformational landscape of this relatively flexible system.

2. Methods

2.1. Microwave spectroscopy

LK (referred as 4-acetyl-1-methyl-1-cyclohexene, 90%) was purchased from Fluorochem and used without further purification. The pure rotation spectra of LK were recorded in the gas phase in the microwave region (3.8–19.3 GHz) using the Fabry–Perot Fourier-Transform microwave (FP-FTMW) technique coupled to a pulsed supersonic jet [34]. A heated nozzle [35] allowed to mix LK vapor with the carrier gas (neon) at an absolute backing pressure of about 0.4 MPa. A temperature of 343 K was found to optimize the signal-to-noise ratio (SNR) of recorded lines. The mixture was introduced into a Fabry–Perot cavity through a series 9 General Valve pin hole nozzle (1 mm) at a

repetition rate of 1 Hz. Jet-cooled molecules (T_{rot} of a few K) were polarized within the supersonic expansion by a 2 µs microwave pulse. The free-induction decay (FID) signals were recorded using a heterodyne detection at 30 MHz and were digitized at a sampling rate of 120 MHz. After Fourier transformation of the average time domain signals, lines of the amplitude spectrum were observed as Doppler doublets due to the coaxial arrangement of the jet and the Fabry-Perot cavity. Each resonance frequency was measured as the average frequency of the two Doppler components. The number of recorded points was chosen so that the frequency grid was set to 1.8 kHz. Scans over several hundred of MHz were performed with a step of 250 kHz, each one consisting of 30 FIDs averaged and Fourier transformed, in order to roughly locate LK rotational lines. The experimental accuracy of the MW measured lines was estimated to 2kHz. Examples of observed signals, both in low-resolution scan mode and high-resolution mode (insets), are shown in Fig. 1. In particular, all lines were found split in two components, signature of an internal motion.

2.2. Theoretical calculations

Before the initial search of the rotational transitions, we performed quantum chemical calculations in order to obtain information on the relative energies of possible conformers and on their equilibrium geometries. All calculations were performed using the Gaussian 16 rev B.01 software package [36] on the high-performance computing cluster of PhLAM Laboratory. Møller-Plesset second order perturbation theory (MP2) [37-40] as well as DFT methods, including Becke98 functional (B98) [41], Becke 3-parameter Lee-Yang-Parr functional (B3LYP) [42-45] and Minnesota Functional (M06-2X) [46] were used. Different basis sets were used throughout including: Dunning and coworkers augmented correlation consistent triple zeta (aug-cc-pVTZ) [47], Ahlrichs and coworkers split valence and triple zeta (def2TZVP) [48,49] and Pople's 6-311++G(d,p) [50]. The quadratic synchronous transit (QST2) method [51], as implemented in Gaussian 16, was used at the MP2, B98, B3LYP and M06-2X levels of theory to calculate the energy of the transition state, and consequently the barriers of the methyl group internal rotation and the ring puckering motion (interconversion between equatorial and axial conformers, see hereafter). Concerning the acetyl group torsion, relaxed potential energy surface (PES) scans at the B98/aug-cc-pVTZ level were performed to characterize possible stable conformations. The geometry was optimized at each point of the scan (i.e. at each step of rotation of the dihedral angle containing the acetyl group). Geometries of the most stable conformers were



Fig. 2. Optimized geometries of all calculated stable equatorial and axial conformers of limona ketone at the MP2/aug-cc-pVTZ level of theory; τ represents the value of the dihedral angle C5C4C8O10.

subsequently fully optimized at the MP2, B98, B3LYP, M06-2X levels of theory. Vibrational frequencies (harmonic) and rotational constants (equilibrium) were calculated for each of the optimized geometries. A full list of the calculations, for all the predicted conformers, is given in the supplemental file.

3. Results and analyses

3.1. Conformational analysis

LK is a cyclic ketone having an endocyclic C–C double bond between C1 and C2 carbon atoms, an acetyl (COCH₃) group attached to a cyclohexene ring at carbon C4 and a methyl (CH₃) group attached to the ring at carbon C1. At C4, the acetyl group can be oriented in either equatorial (along the plane of cyclohexene ring) or axial (perpendicular to the plane of the ring) positions. Interconversion barrier between equatorial and axial is expected to be high (several tens of kJ mol⁻¹, see *e.g.* [52]), since it implies the acetyl group torsion in addition to the ring puckering motion. Indeed, QST2 calculations at the MP2/augcc-pVTZ result in a barrier height between most stable conformations at equatorial and axial orientations of 20.9 kJ mol⁻¹ (ZPE corrected at the harmonic level).

The torsion around the C4-C8 bond (acetyl group torsion), in both equatorial and axial orientations, can lead to different conformations displayed in Fig. 2. To explore the conformational landscape, we performed a relaxed PES scan along the dihedral angle C5C4C8O10, with

a step of 5°. Upon exploring the 1D potential energy surface (PES), we can see from Fig. 3 that four equatorial conformers and two axial conformers are possible.

We named the equatorial conformers EQ1, EQ2, EQ3 and EQ4 and the axial conformers AX1 and AX2, from lowest to highest value of calculated energy. The calculated dihedral angle C5C4C8O10 value, at the MP2/aug-cc-pVTZ level of the theory, is equal to 272°, 325°, 141° and 103° for EQ1, EQ2, EQ3 and EQ4 and 19° and 236° for AX1 and AX2, respectively. The relative equilibrium structure energies, calculated with different methods and taking into account the zero point energy (ZPE) corrections at the harmonic level, along with rotation–vibration constants, are summarized in Fig. 4.

The global minimum corresponds to the EQ1 conformer. Its optimized geometry at the MP2/aug-cc-pVTZ level of theory is presented in Fig. 2(a). Conformational relaxation within the supersonic expansion is expected when the conformational energy is above the barrier height. In our experimental conditions, the conformational energy of LK is equal to $2.8 \text{ kJ} \text{ mol}^{-1}$ (energy corresponding to a reservoir temperature of 343 K). A bare theoretical estimation of the barrier height is given by energy differences between minima and maxima from the relaxed scan of the PES along the acetyl group torsion (see Fig. 3(a)). As shown in Fig. 4, for EQ4 and EQ3, this difference ranges from 0.6 to $0.87 \text{ kJ} \text{ mol}^{-1}$ according to the different calculations performed. The barrier height from EQ4 to EQ3 is estimated to $0.53 \text{ kJ} \text{ mol}^{-1}$, far below conformational energy and this barrier is a disputable point (see discussion hereafter), thus the EQ4 conformer is expected to relax to



Fig. 3. Potential energy scan for the equatorial (a) and axial (b) configurations of limona ketone at the B98/aug-cc-pVTZ level of theory, with respect to the dihedral angle of the acetyl group COCH₃. In panel (a), surprising non-smooth points at certain angles are marked by an asterisk (*); the dash line represents the lowest energy path extracted from a 2-dimensional exploration of the potential energy surface (see in text section 4.2 conformational landscape for details).



Fig. 4. Relative energy (ΔE (kJ mol⁻¹)) ordering of all calculated conformers of limona ketone from different methods and basis sets. Zero represents the lowest energy value. Values are ZPE corrected.

EQ3 conformer. Similarly, EQ2 is expected to relax to EQ1 since the energy difference ranges from 0.45 to $0.92 \text{ kJ} \text{ mol}^{-1}$ and the barrier height from EQ2 to EQ1 is estimated to $0.24 \text{ kJ} \text{ mol}^{-1}$. The barrier height from EQ3 to EQ2 is estimated to 2.08 kJ mol⁻¹, so that another

possible relaxation route goes from EQ4 to EQ1 passing by EQ2. Finally, the barrier height from EQ3 to EQ1 is estimated to 3.11 kJ mol^{-1} , slightly higher than conformational energy, thus relaxation from EQ3 to EQ1 is not expected, although possible. Such assumptions were verified for comparable systems: perillaldehyde, carvone and limonene [12,28]. For axial conformers, the minimum energy structure corresponds to the AX1. Its optimized geometry at the MP2/aug-cc-pVTZ level is shown in Fig. 2(e). The AX2 conformer is not expected to relax to AX1 conformer because the barrier height is above the conformational energy value. In summary, based on energetic considerations, four conformers: EQ1, EQ3, AX1 and AX2 could be observed. A table provided in the supplemental file summarizes relevant molecular constants from all calculations methods used for the six conformers.

A noticeable feature in the experimental spectrum was observed. As we can see in Fig. 1, each experimental line had a neighboring line with similar intensity, splittings ranging from few kHz to few MHz. This splitting pattern is characteristic of an internal rotation motion, such as the torsion of a methyl group which changes the degeneracy of the rotational levels and splits them into two symmetry states: A and E. Thus, each transition will be split into two lines of equal intensity. This splitting is further discussed hereafter. Table 1

Experimental and calculated rotational and quartic centrifugal distortion constants, of limona ketone lowest energy equatorial conformer (EQ1), along with other spectroscopic parameters.

Parameters	Exp. value (SPFIT)	Exp.value (XIAM)	MP2 ^a	MP2 ^b	B3LYP ^c	b98 ^b	M06-2X ^a
A (MHz)	3159.78866(11)	3159.509939(292)	3151.24	3173.01	3184.07	3169.26	3175.23
B (MHz)	735.407909(45)	735.390027 (96)	732.36	741.18	734.05	727.51	738.52
C (MHz)	653.518570(52)	653.512896 (91)	660.44	662.23	656.74	651.66	656.66
Δ_J (kHz)	0.03016(18)	0.030581 (395)	0.03	0.03	0.03	0.03	0.02
Δ_{JK} (kHz)	0.0571(24)	0.05208 (312)	0.06	0.07	0.07	0.07	0.06
Δ_K (kHz)	0.3659(86)	0.3362 (158)	0.44	0.40	0.38	0.39	0.37
δ_J (kHz)	0.001516(74)	0.002012 (238)	0.003	0.001	0.002	0.002	0.001
δ_K (kHz)	0.834(21)	0.8315 (260)	0.63	0.56	0.63	0.60	0.51
V ₃ (cm ⁻¹)		298.39(25)	207.12	183.24	178.27	238.89	125.52
e (rad)		2.615051 (959)	2.36	2.32	2.35	2.34	2.39
δ (rad)		2.230887 (506)	2.13	2.08	2.09	2.08	2.09
Dpi2- (kHz)		-317.622 (807)				•••	
F ₀ (GHz)		163.561 (115)				•••	
RMS ^d (kHz)	1.32	3.36				•••	
N ^e	101	214				•••	
μ_a^{f} (D)		•••	1.2	1.2	1.2	1.2	1.3
μ_b^{f} (D)			1.9	1.9	2.0	2.0	2.1
$\mu_c^{\rm f}$ (D)			1.8	1.7	1.8	1.8	1.7

^a6-311++G(d,p) basis set.

^bdef2TZVP basis set.

^caug-cc-pVTZ basis set.

^dRoot mean square value of observed - calculated line frequencies.

eNumber of lines used in the fit: only A symmetry components with SPFIT, both A and E symmetry components with XIAM.

^fDipole moment component in the principal axes system.

3.2. Spectral analysis

LK in EQ1 conformation is a prolate asymmetric rotor with Ray's asymmetric parameter having a value of $\kappa = -0.935$. Preliminary predictions of the spectrum were made based on a semi-rigid rotor Hamiltonian (Watson's A-reduction) from the rotational constants calculated at different levels of theory, using SPFIT/SPCAT program suite [53]. The assignment started by fitting intense lines observed in the scan to predicted transitions, and then including gradually series of lines with increasing quantum numbers and different variations of K_a and K_c . Lines of different transition types were added, as well as Q-branch transitions. In the following, the notations used include information about the assigned lines. The a,b,c exponents represent the transition type with respect to dipole components in the principal axes. The R or O symbols correspond to R- and O-branch transitions ($\Delta J = 1$ and $\Delta J = 0$), respectively. The two subscript components represent the ΔK_a and ΔK_c values. A set of 107 lines including 45 of a-type (all of them being ${}^{a}R_{01}$), 37 b-type (17 ${}^{b}R_{11}$, 1 ${}^{b}R_{01}$, 14 ${}^{b}R_{1-1}$, 4 ${}^{b}R_{-11}$, 1 ${}^{b}R_{10}$) and 25 c-type (19 °R₁₀, 1 °R₁₋₁, 5 °R₁₋₂) up to J = 13 and $K_a = 4$ was finally obtained, corresponding to a consistent set of A symmetry transitions. The line intensities of the different transition types were in accordance with the calculated dipole moment components ($|\mu_a|$ < $|\mu_c| < |\mu_b|$, see Table 1 for calculated values). The root-mean-square (RMS) value of the fit is 1.32 kHz, in agreement with the instrumental accuracy. The fitted values of the rotational and quartic distortion constants are summarized in Table 1. The values are in good agreement with calculated constants, with the difference ranging between 0.3 and 1.1 %. This magnitude of differences is expected when comparing ground state experimental values with equilibrium theoretical values.

A more detailed look shows that the A constant is best calculated using the MP2 method, whereas the B and C constants are better predicted with the DFT (B98 and B3LYP) methods. Concerning quartic centrifugal distortion constants, and because of the limited range of values of J, K_a and K_c accessible in the microwave region, their deviations from theoretical calculations are relatively more important, but their sign and order of magnitude are consistent as usually expected.

In order to address the observed splitting, we should consider the two methyl groups that are present in LK: one attached to C1 atom and the other attached to the carbonyl moiety of the acetyl group. We named them M1 and M2, respectively. The magnitude of the splitting

depends on the barrier height of the methyl torsion. Thus, the barrier height (V_3) associated to M1 and M2 internal rotations was calculated at different levels of theory using the QST2 method. For M1, we obtained a value of 790 cm⁻¹ at the MP2/6-311++G(d,p) level of theory. From experience, such barrier height higher than roughly 500 cm⁻¹ is expected to cause splitting on the lowest energy levels ranging from several kHz to few tens of kHz which are too small to be observed at our instrumental resolution. For M2, V_3 varies in a wide range from 125 cm⁻¹ to 240 cm⁻¹ depending on the method used. The values obtained are summarized in Table 1. A barrier height of such magnitude is considered to be intermediate, and a splitting into two components of A and E symmetry of each observed transition expected to be observed, ranging from few kHz to few hundreds of MHz. Hence, the splitting due to internal rotation of M2 was considered in the spectral analysis.

The XIAM program [54], which treats the internal rotation motion, was used to perform an effective fit of A and E transitions observed in the spectrum. XIAM was chosen because it uses a simple Hamiltonian based on Watson's 1st order Hamiltonian, and it usually results in reliable fits in the case of medium-to-high internal rotation barriers as in the present case of LK EQ1. Starting from predictions based on the fitted parameters with SPFIT, and the ZPE corrected barrier height calculated at the MP2/aug-cc-pVTZ level, experimental frequencies of the E components have been assigned and fitted in addition to the A components that have been previously assigned. The final XIAM fit includes 214 lines (78 a-type including 22 aR01 and 56 R0-1, 82 btype including 16 $^{b}R_{-1-1},$ 10 $^{b}R_{1-1},$ 22 $^{b}R_{-11}$ and 34 $^{b}Q_{-11}$ as well as 54 c-type including 40 $^{\rm c}R_{-10},$ 6 $^{\rm c}Q_{10},$ 2 $^{\rm c}Q_{-1-2}$ and 6 $^{\rm c}Q_{-10})$ up to J = 13 and $K_a = 4$ and shows a RMS deviation of 3.36 kHz. A complete list of assigned transitions is available in the supplemental file. The V_3 potential energy parameter, the angles δ (angles between the internal rotation axis and the principal axis z) and ϵ (angle between the principal axis x and the projection of the internal rotation axis onto xy-plane) and the internal rotation constant F_0 are fitted. The use of the "Dpi2-" empirical parameter that is programmed in XIAM [55] (internal rotation-overall rotation distortion parameter) was necessary to reach instrumental accuracy. Values from the best fit are listed in Table 1.

The experimentally deduced rotational constants are compared to the calculated ones in Table 1. Noticeably, the rotational constants obtained with XIAM, by fitting both A and E symmetry components, are close to those obtained from the semi-rigid rotor fit, by fitting A symmetry components only, and they are still in good agreement with calculations. Individual errors also depend on the method used.

4. Discussions

4.1. Internal rotation motion

The V_3 value is theoretically underestimated whatever the method used (see Table 1). In addition, the use of the Dpi2- parameter is unexpected when fitting centimeter-wave data only. Such discrepancy might be explained accounting for the fact that XIAM program refers to an isolated system where only the methyl group rotates, whereas all the system is relaxed and optimized in theoretical calculations. In this context, the barrier height was then calculated at the MP2/aug-ccpVTZ level of theory with an "adiabatic" path where only the methyl group was rotated and all other atoms were kept fixed at equilibrium structure. The obtained "adiabatic" value V_3^{ad} is 442 cm⁻¹. As a result, the experimental barrier height $V_3^{exp} = 298 \text{ cm}^{-1}$ is almost at the center of the range obtained from the "relaxed" value $V_3^{rel} = 183 \text{ cm}^{-1}$ to the "adiabatic" value $(V_3^{rel} < V_3^{exp} < V_3^{ad})$. This result indicates that on interaction exists have $V_3^{rel} = 1.000 \text{ cm}^{-1}$. an interaction exists between M2 internal rotation and the skeleton of LK. Such effect was already evidenced, for example in the case of methacrolein [56], methylglyoxal [57] or 2-nitrotoluene [58]. Another noticeable example of similar situation was observed in 2-Acetyl-5methylfuran [59]. As a consequence, in the absence of a model taking into account an interaction between large amplitude motions, only an effective fit can be performed to reach instrumental accuracy, by using second or higher order terms (Dpi2- in the present case) in the isolated internal rotation hamiltonian. Other arguments pointing toward an interaction between internal motions are developed in the following.

4.2. Conformational landscape

The study of relaxed potential energy scans (Fig. 3) was the first step in exploring the conformational landscape of LK. The scans, for both axial and equatorial conformers, showed some non-smooth points at certain angles, that can be seen as small bumps in the curve (as marked by a * in Fig. 3). This was also pointed out in the case of 2-Acetyl-5-methylfuran [59]. A recheck of these points was necessary to understand whether they are artifacts from the calculations or they signify a physical interaction. Focusing on the equatorial configuration, one of these points leads to the assumption that we might have two minima around 120°, thus expecting EQ2 and EQ3 conformers, when in fact we might have just one minimum. A primary check was to calculate the PES scans using different combinations of method and basis sets and particularly have a deeper look around these points with smaller steps (1°). The bumps occurred through all calculations (see figure provided in supplemental file). Second, we followed the optimized geometry at each point of the scan to check any sudden change in geometry. We noticed that, for example, when the dihedral angle of the acetyl group is around 119°, M2 unexpectedly jumps of about 40°. This movement is shown in Fig. 5. When we compared the two geometries, the hydrogen atoms of M2 seem to mirror themselves through this rotation by a "symmetry" plane of LK skeleton, suggesting that the optimization process tries to lower the steric hindrance by suddenly switching to an almost iso-energetic geometry. This movement can be observed similarly at the other bumps for both axial and equatorial conformers.

As this observation is not enough to understand whether these bumps have a physical significance or not, and driven by the fact that the experimental barrier height of M2 does not match the calculated value with two approaches (see previous subsection), a twodimensional PES exploration has been performed. Indeed, as pointed out in a recent study [58], an iso-surface can be used to visualize such effects. The PES was calculated across two relevant dihedral angles, C5C4C8C10 and C4C8C9H22 corresponding to the rotation of



Fig. 5. Geometry of the methyl group M2 (CH₃) of equatorial conformers of limona ketone at an acetyl group (COCH₃) orientation of 119° (gray bonds) and 120° (green dashed bonds). The calculated angle of sudden rotation is about 40° .

the acetvl and M2 groups, respectively, by means of relaxed scans of 5° steps. The PES and its iso-surface are shown in Figs. 6(a) and 6(b). respectively. The overall PES displays a smooth behavior without any break from one point to the next. Along M2 rotation (i.e. when acetyl dihedral angle is fixed), curves look like a standard 3-fold potential as expected. Along acetyl rotation, curves qualitatively agree with the fully relaxed scan (Fig. 3(a)). Without interactions between the two movements, one would expect a PES with horizontal and vertical orientations of valleys which means that one group rotates without influencing the other. However, the present iso-surface clearly displays a diagonal orientation of valleys (see Fig. 6(b)) which is, on the contrary, an evidence of an interaction between these two rotations. Indeed, starting from one minimum, only a concerted torsional motion is possible to reach another relative minimum (i.e. by roughly following dark blue points on the iso-surface). Noticeably, complete rotations of M2 to reach an equivalent position (i.e. rotations of 120°) are achieved at the acetyl group angles where bumps are observed in the relaxed scan. Therefore, we followed quantitatively the minimum energy path in the 2D PES: starting from a minimum position (acetyl=275°; M2=60°), we took the closest lowest energy point on the next PES slice (at acetyl angle $+5^{\circ}$) and so on until complete rotation of the acetyl group. The obtained PES cut is shown in Fig. 3(a) where it is compared to the fully relaxed scan. First, one can clearly see that bumps are no longer present in the cut, leading to the conclusion that these bumps have no significant physical meaning and they are due to calculation "artifacts" correlated with the relaxed scan procedure. Second, EQ3 no longer exists in the cut, suggesting that EQ3 and EQ4 are rather "mirror images" of themselves than two different stable conformations (see Figs. 2(c)and 2(d)). Third, the well around EQ1 and EQ2 appears very flat in the minimum energy path curve. One explanation could come from the fact that following a minimum path in a 2D surface only takes into account constraints between the two followed elements, and does not take into account the constraints with the rest of the molecule, whereas a fully relaxed scan does. For example, when rotating the acetyl group at fixed steps, some electrostatic repulsion or perturbations of the overlapping molecular orbitals might be compensated by an "over-rotation" of the methyl group which would not have occurred in a system fully relaxed. Finally, the most realistic potential energy curve of the internal rotation of the acetyl group should be somewhere in between, probably closer to the fully relaxed scan but without bumps. In any case, this behavior can be qualitatively used to confirm the presence of interactions between M2 internal rotation and a LK skeleton movement such as acetyl rotation, which explains the disagreement between the experimental barrier height (V₃ parameter) and its theoretical estimation using two different approaches ("adiabatic" and "relaxed" energy curves), as well as the unexpected use of the Dpi2- parameter.

4.3. Search for higher energy conformers

After analysis of LK EQ1 spectrum, taking into account line splitting due to the internal rotation of the methyl group M2, some observed



(b) Isosurface view.

Fig. 6. 2-dimensions potential energy surface of limona ketone in equatorial configuration with respect to $COCH_3$ and CH_3 rotations, displayed in (a) 3D view and (b) isosurface plot. In panel (b), The red line represents the following of the path with the lowest energy value.

lines remain unassigned. Although many of them are very weak thus most probably belong to ¹³C isotopic species in natural abundance (whose analysis is out of the scope of the present study), few of them are strong enough to deserve further investigation (weak lines of EQ1 have SNR from 5 to 99, whereas one unassigned line has a SNR of 750, the few others having SNR from 10 to 90). The most reasonable guess is that they might arise from the other conformers of LK, in particular AX1, EQ2 and AX2 which are relatively close in energy to EQ1 (see Fig. 4). Indeed, the Boltzmann distribution at sample temperature (343 K) over all five conformers based on MP2 relative energies would give an initial sample (before expansion in the jet) composed of about 28 % of EQ1, 22 % of AX1, 18 % of AX2, 18 % of EQ2 and 14 % of EQ3/EQ4. However, all attempts to perform a fit at instrumental accuracy with a set of molecular parameters corresponding to LK conformers were unsuccessful. This tends to correlate with our initial assumption of a strong relaxation to EQ1 within our jet expansion.

Then, a search for the presence of precursor molecules and/or impurities in the sample was undertaken, starting from the LK synthesis process. LK is synthesized using a Diels–Alder reaction of isoprene and methyl vinyl ketone on dry silica gel [21]. The rotational constants of isoprene [60,61] and methyl vinyl ketone [62] being well known, we tried to assign unidentified lines to isoprene and/or methyl vinyl ketone, but here again without success.

During the synthesis of LK, the 4-acetyl-2-methylcycloxene, a meta substituted isomer, was observed [21]. Therefore, calculations at the MP2/6-311++G(d,p) level were performed to identify the possible conformers and to obtain the rotational constants of the most stable one. One more time, attempts to attribute unidentified signals to this molecule were unsuccessful.

Finally, searches into splatalogue spectroscopic database [63] were undertaken focusing on spectra of known molecules similar to LK, with respect to the C8O10 double bond, such as acetone or acetaldehyde, unsuccessfully. As a consequence of all these numerous attempts, we believe that the assignment of these few lines is much more challenging than expected and falls beyond the scope of the work presented here.

5. Conclusion

In this paper, a microwave study of limona ketone is presented through a combined experimental and theoretical approach. We use a Fourier transform microwave spectrometer coupled to a supersonic jet expansion over the 3.8-19.3 GHz range. Considering the theoretical relative energy values, four conformers were expected, but only one, the most stable from calculated energies, is observed. This is most probably due to conformational relaxation upon our supersonic expansion conditions. The spectrum clearly shows line splittings characteristic of an internal rotation motion, identified to the methyl group attached to the carbonyl moiety of the acetyl group. Both SPCAT/SPFIT and XIAM program suites are employed in a sequential method: a preliminary fit to the A symmetry transitions using SPFIT followed by a fit of both A and E symmetry transitions using XIAM. The experimentally deduced value of the barrier to internal rotation is $V_3^{exp} = 298 \text{ cm}^{-1}$. Besides the barrier height value, the analysis of the internal rotation provided information about the interaction of the methyl group within the molecular framework. Indeed, although the agreement between theoretical and experimental internal rotation parameters is only qualitative, a two-dimensional exploration of the potential energy surface clearly evidences an interaction between the methyl group internal rotation and a skeleton movement, that is the acetyl group torsion, which explains the discrepancy. Moreover, a standard unidimensional scan of the PES displayed some surprising features (discontinuities and an unexpected conformer EQ3), which have been identified as computational artifacts thanks to a more detailed 2D exploration. The methodology proposed in the present work should be of interest for other compounds containing acetyl moiety. Attempts to identify the higher energy conformers in the spectrum were unsuccessful, confirming the assumption of a strong relaxation to the most stable conformer EQ1 within the jet expansion. Finally, from the atmospheric application point of view, the present data can provide the fundamentals for future studies of microsolvation effect and/or aggregation of limona ketone.

CRediT authorship contribution statement

Noureddin Osseiran: Formal analysis, Investigation, Writing – review & editing, Visualization. Annunziata Savoia: Formal analysis, Investigation, Writing – original draft, Visualization. Pascal Dréan: Conceptualization, Formal analysis, Investigation, Project administration, Writing – review & editing. Thérèse R. Huet: Conceptualization, Formal analysis, Supervision, Project administration, Validation, Writing – review & editing. Manuel Goubet: Formal analysis, Project administration, Methodology, Validation, Writing – review & editing.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A. Supplementary data

Supplementary material related to this article can be found online at https://doi.org/10.1016/j.jms.2022.111643.

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