Vibrational spectra of pilocarpine hydrochloride crystals

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Pilocarpine is a natural substance with potential application in the treatment of several diseases. In this work Fourier Transform (FT)-Raman spectrum and the Fourier Transform infra red (FT-IR) spectrum of pilocarpine hydrochloride $C_{11}H_{17}N_2O_2^+$. Cl⁻ were investigated at 300 K. Vibrational wavenumber and wave vector have been predicted using density functional theory (B3LYP) calculations with the 6-31 G(d,p) basis set. A comparison with experiment allowed us to assign most of the normal modes of the crystal.

Keywords: Raman scattering, infrared spectroscopy, normal modes, pilocarpine hydrochloride

1. INTRODUCTION

In recent years, there has been a growing interest in the study of spectroscopic properties of plant cells in order to identify their chemical constituents through non-destructive analysis [1]. The main researches deal with primary metabolites, i.e., substances essential for their growth, surviving and reproduction (among them, amino acids, proteins, carbohydrates, lipids and fatty acids). On the contrary, the investigation of the vibrational property of isolated secondary metabolites from plants (used as defense against parasite and diseases as well as used to reinforce reproductive processes) is still poorly explored, although many of them have potential application as therapeutic drugs [2,3].

Secondary metabolity pilocarpine ($C_{11}H_{17}N_2O_2$), an alkaloid extracted from the leaves of the South American shrubs *Pilocarpus jaborandi*, *Pilocarpus microphyllus* and other *Pilocarpus* species [4], is an imidazole derivative that exhibits some pharmacological activities. These activities include diaphoretic effects, stimulation of parasympathetic system [5], miotic action [6], being also used in ophthalmology [6,7]. The action of pilocarpine on the parasymphatetic nervous system has been extensively investigated and it is known that the substance act mainly as a cholinergic drug [8]. Despite of several therapeutic effects, pilocarpine is used clinically only to treat glaucoma [8].

Pilocarpine molecule, which contains both imidazole and γ - lactone rings forms two semi-organic compounds in the solid state phase, trichlorogermanate hermihydrate [9] (C₁₁H₁₇N₂O₂, GeCl₃,1/2H₂O) and hydrochloride (C₁₁H₁₇N₂O₂⁺.Cl⁻). [10] For both compounds it was discovered that the crystal structures are monoclinic, space group *P*2₁, although the conformation of the pilocarpine molecule itself differs significantly from one structure to the other [9,10].

From the biological point of view, pilocarpine hydrochloride has been used in certain eye diseases, as for exemple, in the treatment of intraocular hemorrhages, opacities of the vitreous and aqueous fluids [8], while trichlorogermanate hermihydrate pilocarpine presents a weak activity of muscarinic stimulants [11].

In this work an infrared analysis and a Raman scattering study in the spectral range 40 cm⁻¹ to 4000 cm⁻¹ of pilocarpine hydrochloride crystal obtained from *Pilocarpus trachyllophys* [4] is reported. In order to assign the normal modes of vibrations of the material a Density Functional Theory (DFT) calculation was performed.

2. EXPERIMENTAL

FT-Raman spectrum was taken using a Bruker RFS100/S FTR system and a D418-T detector, with the sample excited by means of the 1064 nm line of a Nd:YAG laser. Infrared spectrum was obtained by using an Equinox/55 (Bruker) Fourier Transformed Infrared (FTIR) spectrometer. FT-Raman and

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FT-IR spectra were collected from samples confined in screw cap standard chromatographic glass vials, at a nominal resolution of 4 cm^{-1} accumulating 60 scans per spectra and using a laser power of 150 mW.

3. COMPUTATIONAL METHOD

Density functional theory (DFT) calculations were carried out using the Gaussian 98 programme package [12]. The B3LYP functional was used with the 6-31 G(d,p) basis set. The calculations were performed using an isolated molecule of pilocarpine cation: $C_{11}H_{17}N_2O_2^+$. The structure obtained from the X-ray analyses of pilocarpine hydrochloride at 77 K was used as starting structure [10]. This structure was optimized and the vibrational wavenumbers were then calculated. The output file contained the optimized structure, the vibrational frequencies in the harmonic approximation, and the atomic displacements for each mode. At the optimized structure of the molecule, no imaginary frequency was obtained, proving that a true minimum of the potential energy surface was found. The calculated vibrational wave numbers were adjusted to compare with experimental Raman and IR frequencies.

4. RESULTS AND DISCUSSION

The crystal of pilocarpine hydrochloride at room temperature belongs to the monoclinic structure with $P2_1(C_2^2)$ space group, with Z=2, and lattice parameters a = 11.057 [10.965] Å, b = 9.212 [9.177] Å, c =6.697 [6.507] Å and β = 110.05 [109.19]^o o (where the values in brackets hold for the 77 K determination) [10].

Figure 1 shows the molecular structure of the pilocarpine hydrochloride ($C_{11}H_{17}N_2O_2^+.Cl^-$). The numbering of the atoms in Fig. 1 follows that of Codding [10] in which the *N*-methylated nitrogens (N_1) are separated from the ether oxigen by four carbon atoms and from the carbonyl oxygen by five atoms. This labelling will be used in describing the parameters for optimized structure and the molecular wave vectors. The distribution of the two molecules of pilocarpine hydrochloride in the unit cell is showed in Fig. 2.

Tables 1, 2 and 3 show respectively, bond distances, bond angles and some selected torsion angles for pilocarpine cation, $C_{11}H_{17}N_2O_2^+$, for optimized structure of the molecule (Cal) and that obtained from X-ray analysis (Exp) [10]. The results show that optimized structure was observed to reproduce the experiments with good agreement.

FT-Raman spectrum and the FT-infrared (IR) spectrum of pilocarpine hydrocholride C_{11} H₁₇ N₂O₂⁺.*Cl*⁻ are presented in Figs. 3(a) and 3(b), respectively.

The molecule of pilocarpine hydrochloride has C_1 site symmetry, and therefore, 93 molecular vibrations among all 99 are allowed in the Raman spectrum. The couplings of vibrations due to the presence of two molecules in the unit cell give rise to twice greater number of modes in the crystal. The number of normal modes expected for the crystal is then, 198, distributed into irreducible representations of C_2 factor group as 99 (A + B); from these modes 99 A + 99 B modes are Raman active. Assuming that the weakness of the intermolecular coupling causes negligible factor group splitting, the task is



FIG. 1: The molecular structure of an isolated molecule of pilocarpine hydrochoride: $C_{11} H_{17} N_2 O_2^+ Cl^-$.



FIG. 2: Unit cell of pilocarpine hydrochloride; the Z plane is perpendicular to the figure.

simplified to the assignment of the 93 molecular modes. Table 4 lists a detailed description of assignments for vibrational wavenumbers of pilocarpine hydrochloride. In the first column the calculated values for the wavenumbers are given. We also present the experimental wavenumber values for the crystal obtained by FT-Raman and FT-IR spectroscopies (the second and third columns, respectively); the fourth column in Table 4 gives the assignment of the bands. In order to better visualise the vibrations, we refer to the two rings in the assignments of Table 4, as follows: imidazole ring type or 1-methylimidazole as **R1** and γ - lactone ring type or γ -butyrolactone as **R2**. The nomenclature employed in the classification of normal modes is given below the Table 4.

The assignment for pilocarpine hydrochloride shows that most of the bands observed through FT-Raman and FT-IR spectroscopies correspond to a mixture of vibrational modes. The mixture of modes is common in molecules of C_1 site sym-

	Exp.	Calc.
N(1)–C(2)	1.328	1.337
N(1)-C(5)	1.384	1.399
N(1)-C(15)	1.459	1.469
C(2)-N(3)	1.327	1.335
N(3)-C(4)	1.383	1.381
C(4)-C(5)	1.362	1.369
C(5)-C(6)	1.488	1.498
C(6)-C(7)	1.523	1.535
C(7)-C(8)	1.530	1.543
C(7)-C(11)	1.544	1.549
C(8)-O(9)	1.446	1.431
O(9)-C(10)	1.354	1.368
C(10)-C(11)	1.513	1.537
C(10)-O(12)	1.206	1.200
C(11)- C(13)	1.537	1.548
C(13) C(14)	1.526	1.533

TABLE 1: Bond distances (Å) for Pilocarpine cation: $C_{11} H_{17} N_2 O_2^+$.

TABLE 2: Bond angles (°) for Pilocarpine cation: $C_{11} H_{17} N_2 O_2^+$.

	Exp.	Calc.
C(2)-N(1)-C(5)	109.3	109.4
C(2)-N(1)-C(15)	124.7	124.9
C(5)-N(1)-C(15)	126.0	125.7
N(1)-C(2)-N(3)	108.5	107.9
C(2)-N(3)-C(4)	108.9	109.5
N(3)-C(4)-C(5)	107.0	107.2
N(1)-C(5)-C(4)	106.3	105.9
N(1)-C(5)-C(6)	121.8	122.5
C(4)-C(5)-C(6)	131.7	131.6
C(5)-C(6)-C(7)	113.4	114.2
C(6)-C(7)-C(8)	110.0	111.9
C(6)-C(7)-C(11)	118.3	118.7
C(8)-C(7)-C(11)	102.3	101.9
C(7)-C(8)-O(9)	105.0	105.3
C(8)-O(9)-C(10)	110.2	111.0
O(9)-C(10)-C(11)	110.9	109.7
O(9)-C(10)-O(12)	120.4	122.5
C(11)-C(10)-O(12)	128.6	127.9
C(7)-C(11)-C(10)	101.6	101.4
C(7)-C(11)-C(13)	114.6	116.2
C(10)-C(11)-C(13)	109.6	110.4
C(11)-C(13)-C(14)	114.7	113.5

metry. The superposition of modes precludes a direct identification of the bands. However, an effort was carried out through this work to make a detailed description of assignments of vibrational modes of the crystal. In order to illustrate the assignment, atomic displacements corresponding to selected normal modes from the isolated molecular structure of pilocarpine hydrochloride are shown in Fig. 4.

Now we discuss the main calculated and observed vibrations of pilocarpine hydrochloride. Two fundamental units of the pilocarpine molecule are the rings. Imidazole ring, which is a characteristic part of pilocarpine, is present in several substances of biological interest, as for example, L-histidine amino acid [13-15], and in other substances [16,17]. Their vibrations spread over a large spectral range of wavenumbers. Lactone, the other ring, is also found in several different substances of biological interest [18-20]; for some of them spectroscopic studies have revealed the wavenumber of the main vibrations [19]. Calculations show that at low wavenumber ($\omega < 150 \text{ cm}^{-1}$) where it is expected to be observed bands associated to lattice vibrations, some internal modes are also present. For example, torsional vibrations of the two rings are observed together with lattice modes at very low wavenumber. This should be expected because the rings are very large structures; so, we assign the bands in this spectral region as a mixture of lattice modes and torsional vibrations of the rings R1 and R2. Fig. 4(a) shows atomic displacements associated to deformations { γ_{oop} (R1), δ_{oop} (R2) [τ (C10O9O12)], r(C13H₂), r(C14H₃)} corresponding to the strong Raman bands observed at 96 cm⁻¹ ($\omega_{cal} = 97$ cm⁻¹).

Another class of vibrations is related to deformation of rings. In plane ring deformation vibration appears in a large spectral region (690 – 1900 cm⁻¹) and out of plane ring deformation vibration appears for 546 < ω < 1140 cm⁻¹. However, most of them are mixed with other kind of vibrations such as rocking and bending of CH, torsion of CH₂ and stretching of CC. Fig. 4(b) represents the mixtures of vibrational modes { δ_{ip} (R2) [sc(C8C7C11); v_s (C11C1009)], r(C8H₂; C13H₂), r(C14H₃), δ (C7H)} giving rise to the strong Raman peak ob-

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	Exp.	Calc.	
C(2)-N(1)-C(5) –C(6)	176.7	179.1	
C(3)-C(4)-C(5) –C(6)	-176.3	-178.9	
N(1)-C(5)-C(6) –C(7)	179.8	177.2	
C(4)-C(5)-C(6)-C(7)	-4.2	-4.1	
C(5)-C(6)-C(7)-C(8)	168.4	171.6	
C(5)-C(6)-C(7) –C(11)	-74.5	-70.1	
C(6)-C(7)-C(8) –O(9)	157.0	159.2	
C(6)-C(7)-C(11) –C(10)	-149.7	-153.3	
C(6)-C(7)-C(11) –C(13)	-31.6	-33.5	
C(7)-C(8)-O(9) -C(10)	-20.4	-20.2	
O(9)-C(10)-C(11) –C(7)	18.2	19.8	
O(12)-C(10)-C(11) -C(7)	-162.8	-161.2	
C(7)-C(11)-C(13) –C(14)	171.4	179.4	

TABLE 3: Some selected torsion angles(°) for Pilocarpine cation: $C_{11} H_{17} N_2 O_2^+$.



FIG. 3: (a) Pilocarpine hydrochloride FT-Raman spectrum. (b) Pilocarpine hydrochloride FT-IR spectrum.



FIG. 4: Some selected representations of atomic vibrations corresponding to the strongest Raman bands of pilocarpine cation: C_{11} H₁₇ N₂O₂⁺.

served at 766 cm⁻¹ ($\omega_{cal.}$ = 765 cm⁻¹).

The whole structure presents deformation vibrations $\delta(all$ structure) at $\sim 650 \text{ cm}^{-1}$ as well as at $\sim 1080 \text{ cm}^{-1}$ and at 1300 - 1340 cm⁻¹. A calculated δ (all structure) vibrational is also expected at 770 cm⁻¹ but, possibly, is mixed with the 765 cm⁻¹ complex vibration. Four strong IR bands are associated to ring deformation. Fig. 4(c) illustrates one of them observed at 1027 cm⁻¹ ($\omega_{cal} = 1026 \text{ cm}^{-1}$), corresponding to { δ_{oop} (R2) [v_{as} (C11C13C14)], wag (C13H₂), r (C6H₂; $C8H_2$), r (C14H₃), δ (C7H; C11H)}. Another band is observed at 1181 cm⁻¹ ($\omega_{cal} = 1185$ cm⁻¹), corresponding to $\{\delta_{iv} (R2) [sc (C8C7C11), v (C10O9)], r (C13H_2), r (C14H_3),$ wag (C6H₂; C8H₂), δ (C7H; C11H)}. The other two strong IR bands are associated with the deformations { δ_{ip} (R1) [v (N1C15; N3C2), ν (C4C5)], wag (C15H₃), δ (C2H; C4H), δ (N3H)} and { δ_{ip} (R1) [v (C4C5; C5C6), v (C2N1)], wag (C6H₂), r (C15H₃), δ (C2H; C4H), δ (N3H)}, corresponding to the peaks observed at 1752 cm⁻¹ ($\omega_{cal} = 1598 \text{ cm}^{-1}$) and $1767 \text{ cm}^{-1} (\omega_{cal} = 1655 \text{ cm}^{-1})$, respectively.

It is also interesting to note that the rocking vibrations of the three CH₂ units (C6H₂, C8H₂ and C13H₂) are observed at similar wavenumbers. On the contrary, although CO₂ vibrations can be expected for pilocarpine molecule, rocking vibration of CO₂ (a well characteristic vibration in amino acid crystal at 500 - 540 cm⁻¹) is absent. This is because the only CO₂ possibility is C10O9O12, but O9 is held at lactone ring; as a consequence, it is impossible to have a rocking C10O9O12 vibration.

Because there are many C-C bonds in pilocarpine molecules the CC stretching vibrations are observed in a large range of wavenumbers. The lowest wavenumber value is for C11-C13 stretching, which was observed at 716 cm⁻¹ while the highest wavenumber value corresponding to a v(CC) is calculated at 1140 cm⁻¹ · v(NC) vibrations contributes with bands observed at 939 and 1754 cm⁻¹.

It is possible to note a marked localization of the scissoring vibrations (CH₂ and CH₃) in the range 1382 cm⁻¹ < ω_{cal} < 1537 cm⁻¹. As an example, the Raman band observed at 1492 cm⁻¹ (ω_{cal} = 1493 cm⁻¹) corresponds to the scissoring vibration sc(C15H₃).

A large number of bands associated with overtones and combination tones may be found in the region about 2800 cm^{-1} due to the large number of bands in the region between 84 cm^{-1} and 1800 cm^{-1} . The bands 2819, 2834 and 2844

ω_{calc}	$\omega_{FT-Raman}$	ω_{FT-IR}	Assignment
29			γ _{oop} (R1), r (C11C13C14)
43			γ_{oop} (R1), γ_{oop} (R2)
64	84m		γ_{oop} (R1), γ_{oop} (R2)
97	96s		γ_{oop} (R1), δ_{oop} (R2) [τ (C10O9O12)], r (C13H ₂), r (C14H ₃)
123	114s		γ_{ip} (R1), δ_{oop} (R2) [τ (C10O9O12)], r (C13H ₂), r (C14H ₃)
132			τ (C15H ₃)
177	159w		γ _{oop} (R1), r (C11C13C14)
192	199w		γ _{oop} (R1), δ _{oop} (R2) [δ (C8O9)], r (C8H ₂), r (C14H ₃)
212			γ_{oop} (R1), γ_{oop} (R2), r (C11C13C14), δ (C15N1)
214			γ_{oop} (R1), δ_{oop} (R2) [δ (C10O12)], r (C8H ₂), r (C14H ₃ ; C15H ₃)
228	229m		γ_{oop} (R1), δ_{oop} (N1C15), r (C6H ₂), r (C14H ₃)
247	256w		γ_{oop} (R1), γ_{ip} (R2), δ_{oop} (N1C15), r (C6H ₂), r (C14H ₃)
288			γ_{oop} (R1), δ_{oop} (N1C15), r (C6H ₂ ; C8H ₂ ; C13H ₂); r (C14H ₃)
298	306w		γ_{oop} (R1), γ_{oop} (R2), r (C6H ₂ ; C8H ₂ ; C13H ₂); r (C14H ₃)
366	374w		sc (C11C13C14), γ _{oop} (R2), r (C6H ₂)
412	414w	415vw	γ_{ip} (R1), δ_{ip} (N1C15), δ_{oop} (R2)
479		480m	δ (all structure)
548	546w	546w	δ_{oop} (R2) [sc (C6C7C11); sc (O9C10O12)], r (C6H ₂ ; C8H ₂)
621			δ_{oop} (R1) [δ (C4N3C2)]
627	625vw	628m	δ (all structure)
641	643m	643m	δ (all structure)
654			δ_{oop} (R1) [δ (C4C5N1)], δ_{oop} (C2H; C4H), δ_{oop} (N3H), τ (C6H ₂),
			r (C8H ₂ ; C13H ₂), r (C15H ₃)
685			δ_{oop} (N3H)
690	691vw	690w	δ_{ip} (R2) [v _s (C8O9C10)], r (C6H ₂ ; C8H ₂ ; C13H ₂),
			r (C14H ₃)
710	716vw	716w	δ_{oop} (R2) [δ_{oop} (C10O12)], v (C11C13), δ (C11H), r (C13H ₂),
	_ / /		r (C14H ₃)
765	766s	762w	δ_{ip} (R2) [sc (C8C7C11); v _s (C11C10O9)], r (C8H ₂ ; C13H ₂),
			r (C14H ₃), δ (C7H)
770			δ (all structure)
787	824vw	822m	δ_{oop} (R1) $[\delta_{oop}$ (C2H; C4H)], r (C6H ₂)
834	850w	843m	$\frac{\delta_{oop} (\text{C2H}; \text{C4H}), \delta_{oop} (\text{N3H})}{(\text{C2H}; \text{C4H}), \delta_{oop} (\text{N3H})}$
852	861vw	866m	δ_{oop} (R2) [n (C11C13); v _s (C8C/C11)], τ (C13H ₂), r (C14H ₃),
000		000	$\frac{0(C/H;CI1H)}{(C10012)} = \frac{(C11C1000)}{(C11L)} = \frac{(C11L)}{(C11L)} = \frac{(C11L)}{(C1$
880		880m	o_{ip} (K2) [o (C10012); v_s (C11C1009)], t (C8H ₂ ; C13H ₂), f (C14H ₃),
017	018m	005m	$\frac{0_{000} (C4H)}{\delta} = \frac{(C11C12C14) \times (C7C8)}{\delta} = \frac{\delta}{\delta} = \frac{C4H}{\delta} + \frac{\delta}{\delta} + \frac{C7H}{\delta}$
030	030m	903W	$\frac{\mathbf{b}_{oop}}{\mathbf{k}_{2}} = \frac{\mathbf{p}_{s}}{\mathbf{k}_{1}} \left[\mathbf{p}_{s} \left(\mathbf{C}_{11} \mathbf{C}_{12} \mathbf{C}_{14} \right), \mathbf{p}_{s} \left(\mathbf{C}_{12} \mathbf{N}_{24} \right), \mathbf{p}_{s} \left(\mathbf{C}_{12} \mathbf{N}_{14} \right), p$
959	939W	921W	$\frac{\delta_{ip}(R1)[v(C5N1), sc(C2N5C4)], t(C15H3)}{\delta_{ip}(R2)[v(C11C13C14)], v(C7C8)] \delta_{ip}(C7H; C11H)}$
950			σ_{00p} (K2) [v_s (C11C13C14), v (C7C8)], v (C7C8)], v (C7C8), v
1026	1031m	1027s	$\frac{1}{8}$ (R2) [v (C11C13C14)] wag (C13H ₂) r (C6H ₂ : C8H ₂)
1020	105111	10275	$r_{000}(R2)[v_{as}(C11C15C14)], wag(C15112), r(C0112, C0112), r(C14H_2)]$
1039			$\frac{1}{8}$ (R2) [v(C7C8)] v C11C13C14) wag (C6H ₂) r (C8H ₂)
1057			δ_{ip} (K2) [v(eveo)], v_{as} errerserv), wag (corr_2), i (corr_2), δ (C11H)
1068	1063w	1063m	δ_{con} (B2) [v(C7C8: C10O9)] r (C6H ₂ : C13H ₂): r (C14H ₂)
1000	10001	1000111	δ (C2H: C4H: C11H). δ (N3H)
1078			δ (all structure)
1083	1081w	1085w	δ (all structure)
1106	1102w	1103w	δ_{aaa} (R2) [r (C7C11C13); v (C8O9; C10O9)], v (C13C14),
			r (C14H ₃), τ (C13H ₂), δ (C7H; C11H)
1124	1121w		δ_{in} (R1) [v (C4N3)], r (C15H ₃), v (C7C11), δ (C2H; C4H)
1126			δ _{in} (R1) [v (C5N1; C4N3)], r (C14H ₃ ; C15H ₃), r (C8H ₂), v (C7C11)
			d (N3H)
1140			δ _{oon} (R2) [n (C11C13)], r (C14H ₃ ; C15H ₃), r (C8H ₂), δ (C2H; C4H),
			δ (N3H)
1152	1151vw	1150w	r (C15H ₃), δ (C2H; C4H), δ (N3H)
1153			δ (all structure)
1185		1181s	δ_{ip} (R2) [sc (C8C7C11), v (C10O9)], r (C13H ₂), r (C14H ₃),
			wag (C6H ₂ ; C8H ₂), δ (C7H; C11H)
1207	1196m		δ_{ip} (R1) [sc (C5C4N3)], wag (C6H ₂), r (C8H ₂),
			δ (C2H; C4H; C7H; C11H), δ (N3H)
1235	1218w	1228m	r (C14H ₃), τ (C6H ₂ ; C8H ₂), δ (C2H; C7H; C11H)
1253	1245vw		r (C14H ₃), τ (C6H ₂ ; C8H ₂), δ (C4H; C7H; C11H), δ (N3H)
1267		1272vw	δip (R1) [v (C4N3)], r (C14H ₃), τ (C8H ₂ ; C13H ₂),
			δ (C2H; C4H; C7H; C11H), δ (N3H)

TABLE 4: Calculed vibrational wavenumbers unscaled, Raman band positions in units of cm^{-1} and assignments for vibrational modes of pilocarpine hydrochloride cation: $C_{11} H_{17} N_2 O_2^+$.

cm⁻¹ were assigned as combination tone. The intensity of

bands at 2844 $\rm cm^{-1}$ suggests that is a combination involving

Continued)		
$\omega_{FT-Raman}$	ω_{FT-IR}	Assignment
1279w	1281vw	$δ_{ip}$ (R1) [v (C4N3)], r (C14H ₃ ; C15H ₃), τ (C6H ₂ ; C13H ₂), δ (C2H; C4H; C7H; C11H), δ (N3H)
	1292vw	δ (all structure)
1315w		δ (all structure)
1337m	1331m	δ (all structure)
		τ (C6H ₂ ; C8H ₂ ; C13H ₂), δ (C11H), r (C14H ₃)
1369s	1367w	wag (C14H ₃), wag (C13H ₂), τ (C6H ₂ ; C8H ₂), ν _{as} (C11C13C14), r (C5C6C7), δ (C2H; C7H; C11H), δ (N3H)
1387vw	1387w	sc (C14H ₃), τ (C6H ₂ ; C8H ₂), wag (C13H ₂), ν _{as} (C11C13C14), δ (C7H; C11H)
		δ_{in} (R1) [sc (C4C5N3); v (N1C2; N1C5)], sc (C15H ₃),
		wag (C6H ₂ ; C8H ₂ ; C13H ₂), ν _{as} (C5C6C7), δ (C7H)
		wag (C14H ₃), wag (C6H ₂ ; C8H ₂ ; C13H ₂), sc (C15H ₃) δ (C7H)
1422vw		wag (C14H ₃), wag (C6H ₂ ; C8H ₂ ; C13H ₂), δ (C7H; C11H)
1433w	1432w	wag (C14H ₃)
1449m	1452m	wag (C15H ₃), sc (C6H ₂), δ (C2H), δ (N3H)
1468m		sc (C14H ₃ ; C15H ₃), sc (C6H ₂ ; C13H ₂)
1492s	1492w	sc (C15H ₃)
		sc (C14H ₃ ; C15H ₃), sc (C6H ₂ ; C13H ₂)
		sc (C14H ₃ ; C15H ₃), sc (C13H ₂), δ (N3H)
		sc (C14H ₃ ; C15H ₃), sc (C13H ₂), δ (N3H), δ _{ip} (R1) [v (C2N3; C4N3)]
		sc (C14H ₃ ; C15H ₃), sc (C6H ₂ ; C13H ₂), δ (N3H)
1559vw	1558m	sc (C14H ₃), sc (C13H ₂)
1611w	1613m	sc (C8H ₂)
1754vw	1752s	$δ_{ip}$ (R1) [v (N1C15; N3C2), v (C4C5)], wag (C15H ₃), d (C2H; C4H), δ (N3H)
1767w	1767s	δ_{ip} (R1) [v (C4C5; C5C6), v (C2N1)], wag (C6H ₂), r (C15H ₃), δ (C2H; C4H), δ (N3H)
	1863w	δ_{ip} (R2) [v_s (C10=O12)]
2819w		combination
	2834vw	combination
2844w		combination
2908vs	2911vw	v _s (C6H ₂ ; C13H ₂), v (C7H)
2938vs	2935vw	v _s (C6H ₂ ; C8H ₂ ; C13H ₂), v (C7H)
2972s	2970w	v _s (C6H ₂ ; C8H ₂ ; C13H ₂), v (C7H; C11H)
3004s	3007w	v _{as} (C6H ₂), v _s (C8H ₂), v (C7H; C11H)
3025m	3030w	v _s (C14H ₃), v _{as} (C13H ₂), v (C11H)
		v _s (C14H ₃), v _{as} (C6H ₂ ; C13H ₂), v _s (C8H ₂), v (C11H)
3079s		v _s (C15H ₃), v _{as} (C6H ₂ ; C13H ₂), v (C7H; C11H)
		v _s (C15H ₃), v _{as} (C13H ₂), v (C7H; C11H)
		v _s (C15H ₃), v _{as} (C6H ₂ ; C13H ₂), v (C7H; C11H)
3116vw		ν _{as} (C14H ₃), ν _{as} (C6H ₂ ; C13H ₂), ν (C7H; C11H)
		ν _{as} (C14H ₃), ν _{as} (C8H ₂ ; C13H ₂), ν (C7H; C11H)
		v_{as} (C14H ₃), v_{as} (C13H ₂)
		v_{as} (C15H ₃)
		v_{as} (C15H ₃)
	Continued) ω _{FT-Raman} 1279w 1315w 1337m 1369s 1387vw 1422vw 1433w 1449m 1468m 1492s 1559vw 1611w 1754vw 1767w 2819w 2844w 2908vs 2938vs 2972s 3004s 3025m 3079s	Continued) $\omega_{FT-Raman}$ ω_{FT-IR} 1279w 1281vw 1292vw 1315w 1337m 1331m 1369s 1367w 1387vw 1387w 1387vw 1387w 1422vw 1432w 1449m 1452m 1468m 1492s 1492s 1492w 1559vw 1558m 1611w 1613m 1754vw 1752s 1767w 1767s 1863w 2819w 2844w 2908vs 2972s 2970w 3004s 3007w 3025m 3030w

 $[\]tau$ = twisting; γ = torsion; sc= scissoring; wag = wagging; δ = deformation; δ_{ip} = deformation in plane; δ_{oop} = deformation out of plane; r = rock; ν = stretching; ν_{as} = asymmetric stretching; ν_s = symmetric stretching. vs=very strong ; s = strong; m = medium; w = weak; vw = very weak.

at least one strong Raman band. Thus, the modes at 96, 114, 766, 1369, and 1492 cm^{-1} can be involved.

The spectral region between 2800 and 3150 cm⁻¹ of the Raman spectrum of pilocarpine hydrochloride crystal consists of a series of very intense Raman bands, and a series of less intense IR bands. However, all bands are well resolved, allowing for their identification as listed in Table 4. For organic crystals the region about 3000 cm⁻¹, in general, contains the bands originated from C-H, CH₂, CH₃, and N-H vibrations [21, 22]. For some materials this region condenses very important informations, being a tool to understand conformation of the molecules in the unit cell or even interactions such as hydrogen bonds. For example, a study on L-methionine crystal have shown that the behaviour of Raman bands under pressure in this spectral region can be understood as consequence of

structural changes instead of simple conformational changes of molecules in the unit cell [23]. So, the understand of the origin of these bands can be fundamental to understand the behaviour of pilocarpine hydrochloride under different conditions, in particular, related to the conditions found in drug artefacts. The scheme of Fig. 4(d) shows, as an example, the mixtures of stretching modes { v_s (C6H₂; C13H₂), v (C7H)} corresponding to a very strong Raman band observed at 2908 cm⁻¹ (ω_{cal} = 3024 cm⁻¹).

5. CONCLUSIONS

The phonon spectrum of the pilocarpine hydrochloride, a potential pharmaceutical substance to be used in several dis-

ease treatments, was measured at room temperature through FT-Raman and FT-IR techniques. Density functional theory calculations were carried out by using the Gaussian 98 package and the B3LYP functional with the 6-31 G(d,p) basis set. The calculations were observed to reproduce the experiments with good agreement. This agreement allowed us the assignment of the observed wavenumbers to atomic motions in the molecules. In particular, it was observed that most bands are associated to mixing of vibrational modes, even in the low wavenumber region where, generally, the lattice modes are found. The absence of stretching vibrations of water molecule which can be observed at ~ 3400 cm⁻¹ in this region indicates

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that the crystal is free of water molecules.

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