

Study of the 3-(3,3-Dimethylbutanoyl)-4-hydroxy-6-neopentyl-2H-pyran-2-one by IR, Raman spectroscopy, and DFT

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Abstract

The heterocyclic structure of pyrones has a variety of biological activities and plays an important role in the creation of new drugs. Therefore, the study of the structure and spectra of pyrones is of considerable interest. In this work, the IR and Raman spectra of 3-(3,3-Dimethylbutanoyl)-4-hydroxy-6-neopentyl-2H-pyran-2-one (1) in its crystalline state was studied. The tautomerization of 1 was followed by a quantum-chemical method at the DFT/B3LYP/6-311G** level. The calculation for the 4-hydroxy enol tautomer (A) reproduces the experimental IR and Raman spectra of compound 1. The classification of the bands in the experimental vibrational spectra of 1 has been carried out. The intramolecular H-bond was characterized by IR spectroscopy. The free energies of the tautomers and their populations were calculated for two different solvents. It appears from our data that type A dominates. The content of tautomer B increases in the nonpolar solvent but does not exceed 13%. As evident from the calculations and experimental X-ray data, the pyran ring of the molecule is flat. HOMO and LUMO of molecule 1 are located on the pyran ring. During tautomeric transformations, there is a significant delocalization of charge and a change in the reactivity of the molecule. The reactivity of pyrone 1 was characterized using descriptors. The form B was found to have higher ionization energy, electron affinity, chemical potential, and electrophilic index than the A form. The dipole moment is higher for form A, and the softness of the two molecules is the same.

Keywords:

acids; infrared spectra; raman spectra; hydrogen bonds; normal vibrations; density functional theory

I. Introduction

The study of heterocyclic pyrone derivatives is interesting and important because they are used in the pharmaceutical, cosmetic, and food industries [1–5]. Pyrones are biologically active substances and are used for the manufacture of analgesics, anti-cancer drugs, and to fight against HIV [1-10]. Pyrones are the initial reagents in the synthesis of many organic compounds [1–10]. Studies on NMR spectra have shown that among the five tautomeric forms of the pyrones, two enolic forms predominate [11,12]. The IR and NMR spectra of pyrones have been studied [13–18].

This particular work for the first time characterized the two low-energy tautomeric forms of the 3-(3,3dimethylbutanoyl)-4-hydroxy-6-neopentyl-2H-pyran-2-one (1) using the methods of IR and Raman spectroscopy and quantum chemistry. The choice of compound 1 is linked to the tautomerism of the central structural fragment of pyrandione, which can modify certain features characteristic of the functional groups. The enamine derivative of synthesized pyrone 1 has inhibitory activity against the human carcinoma cell line HeLa and the herpes virus VPG [19]. We attempted to trace the change in the structure of the acid, the strength of the hydrogen bond and

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its vibrational spectra during tautomeric transformations. The comparison of the free energies of the tautomers allows for an estimation of their population.

It was important to follow the evolution of the geometry, electronic structure, and spectra of pyrone 1 during tautomeric transformations. Active centers of the molecule for nucleophilic and electrophilic attacks have been determined. The calculation of the charges on the acid atoms made it possible to estimate the capacity of the atoms to form hydrogen bonds and attract ions and metal atoms. The electrophilicity index characterizes the biological activity of compound 1.

2. Materials and Methods

2.1. Experimental

The neopentyl derivative of dehydroacetic acid 3-(3,3-dimethylbutanoyl)-4-hydroxy-6-neopentyl-2H-pyran-2-one (1) has been obtained by $CF_3SO_3H/(CF_3CO)_2O$ activated acylation of carboxylic acids according to [19,20]. The white product has the crystalline powder form (melting point 77–78°C). Compound 1 can exist in two tautomeric enol forms, A and B (Figure 1).

IR spectra were recorded by accumulating 64 scans in the region of 4000–400 cm⁻¹ with a resolution of 4 cm⁻¹. A Bruker Vector 22 spectrometer was used [21]. The samples were compressed into KBr pellets.

Raman spectra of the pyrone were recorded in the 3500–50 cm⁻¹ region via the FTIR spectrometer VER-TEX 70 and the Bruker FT-Raman RAM II module [21]. The 1064 nm excitation line provided by an Nd:YAG laser with a power of 50 mW was used.

2.2. Computational Details

The calculation of the vibrational spectra of compound 1 was performed using the B3LYP functional [22,23] and the basis set 6-311G**. The calculations were performed using the Gaussian09 program [24]. As a first approximation, the experimental coordinates of the atoms obtained through the X-ray diffraction method were used (Supplementary Information S1). Standard optimization methods were used to find minima on the potential surface. Full geometry optimization was performed without any restrictions. The Hessian analysis made it possible to determine the minima of potential energy.

Optimized geometrical parameters of the tautomers were used to calculate the harmonic vibration frequencies. Theoretical structural and spectral data were obtained for both tautomers at 298 K, 1 atm. The potential energy distribution was calculated to attribute the vibra-

tions [25]. Calculated frequencies were scaled using a multiplier of 0.96. The theoretical spectral curves were constructed, taking the Lorentz band shape and a half-width of 10 cm⁻¹.

The calculation of natural bonding orbitals (NBO) has been performed to characterize the electronic properties of molecules [26]. The chemical potential, hardness, softness, and electrophilicity index are related to the first vertical ionization energy and electron affinity by the following formulas: $\mu \approx -(IE + EA)/2$, $\eta \approx (IE - EA)$, $S = 1/\eta$, and $\omega = \mu^2/2\eta$ [27]. The Fukui functions for nucleophilic $f_k^+(r) = [q_k(N+1) - q_k(N)]$ and electrophilic $f_k^-(r) = [q_k(N) - q_k(N-1)]$ attacks were calculated using the natural atomic charges on atoms q_k and the number of electrons N in a molecule. The local softness of atoms has also been calculated $s_k^+ = Sf_k^+, s_k^- = Sf_k^-$ [27].

Using the difference in free energies of the tautomers, their populations at 298.15 K can be calculated $p = \frac{\exp(-\Delta G_i/RT)}{\sum_{j} \exp(-\Delta G_j/RT)}$ [28]. A polarizable continuum model was used to assess the influence of the polar environment on the tautomeric equilibrium [28].

3. Results and Discussion

3.1. Structural Analysis

As can be seen from the X-ray diffraction data in the crystalline state at room temperature, the A tautomer of compound 1 is realized (Supplementary Information S1). In the more stable tautomer A, an intramolecular hydrogen bond is realized. The measured distance between the O3 and O4 atoms is 2.437 Å. Supplementary Information S2 lists the measured bond lengths and angles of tautomer A.

The results of geometry optimization of tautomers A and B are shown in Figure 2 and in Supplementary Information S2. Gibbs's free energy and Boltzmann weights of tautomers are shown in Table 1. It appears from our data that type A dominates. The content of tautomer B increases in the less polar chloroform but does not exceed 13%. Vibrational spectra were calculated for tautomers A and B.

As can be seen from the calculations and experimental X-ray data, the pyran ring of the molecule is flat. A satisfactory agreement is observed between the calculated geometrical parameters of tautomer *A* and the experimental X-ray data.

Bond lengths change during tautomeric transformation. In tautomeric form A, the calculated bond lengths are (Å) 1.313 (O(3)–C(7)), 1.250 (O(4)–C(27)), 1.413 (C(6)–



Figure 1: The structure of tautomeric forms A and B of compound 1.

C(7)), and 1.467 (C(6)–C(27)), and for the *B* tautomer, the length of these bonds changes to 1.259, 1.302, 1.458, and 1.411, respectively. Such changes in bond lengths are consistent with the change in their properties during the tautomeric transformation. The H-bond lengths in the *A* and *B* tautomers are also different. The calculated $O(3)\cdots O(4)$ distances for tautomers *A* and *B* are 2.473 and 2.418 Å, respectively.

3.2. Frontier Orbitals and Descriptors

The HOMO and LUMO molecular orbitals for the *A* and *B* tautomers are located on the pyran ring (Figure 3). Conjugation leads to a flat structure for this ring. It is interesting to see how the charge distribution changes during tautomeric transformations. In the tautomeric transformation from form *A* to *B*, the negative charge on the O3 atom increases, and that on the O4 atom decreases (Supplementary Information S3). In the *B* form, the charges on the atoms C5, C6, and C7 increase, and on the C27, H44 atoms decrease. It follows from these data that the chemical properties change during the tautomeric transition.

The pyrone 1 molecule contains several functional groups. Their reactions can be described using descriptors. Form B was found to have higher ionization energy, electron affinity, chemical potential, electrophilic index, and energy band gap than Form A (Table 2). The dipole moment is higher for form A, and the softness of both molecules is the same.

The calculation of the local electrophilic indices makes it possible to estimate the reactivity of the atoms for two tautomeric forms (Supplementary Information S3). The electrophilic index of oxygen atoms is higher for the *A* form. For the carbon atoms C5, C6, C7, and C27, the electrophilicity index is higher for form *A*. It should be emphasized that the C6 atom is highly reactive. Detailed analysis of chemical descriptors allows finding new ways to obtain drugs with desired properties.

3.3. NBO Analysis

The molecular orbitals are similar for the two tautomeric forms, but there is an orbital $\sigma(2)_{C6-C7} = 0.8137 (\text{sp}^{1.00} \text{d}^{0.00})_{C6} + 0.5812 (\text{sp}^{1.00} \text{d}^{0.00})_{C7}$ in form A, which is not in form B. In form B there is an orbital $\sigma(2)_{O3-C7} = 0.8605 (\text{sp}^{1.00} \text{d}^{0.00})_{O3} + 0.5095 (\text{sp}^{1.00} \text{d}^{0.00})_{C7}$, which is not in form A. These extra orbitals are of π character and indicate an increase in bond order.

In form *A*, significant interactions of C6–C7, C8–C10 bond orbitals with antibonding orbitals of the O2–C5, O4–C27, C6–C7 σ_2 (C6–C7) $\rightarrow \sigma^*_2$ (O2–C5), σ_2 (C6–C7) $\rightarrow \sigma^*_2$ (O4–C27), σ_2 (C8–C10) $\rightarrow \sigma^*_2$ (C6–C7) with stabilization energies 34.18, 31.20, and 24.96 kcal/mol (Supplementary Information S4). Additionally, the molecule has lone electron pairs of oxygen atoms interactions with O2–C5, C5–C6, C6–C7, and O3–H44 bonds, n(LP₂O1) $\rightarrow \sigma_1^*$ (O2–C5), n(LP₂O2) $\rightarrow \sigma_1^*$ (O1–C5), n(LP₂O2) $\rightarrow \sigma_2^*$ (C5–C6), and n(LP₂O4) $\rightarrow \sigma_1^*$ (O3–H44) with energies 29.25, 40.10, 15.56, and 42.96 kcal/mol.

In the tautomeric form B, the delocalization of electrons is maximal for the C8–C10 bond and is distributed over the antibonding orbitals O3–C7 $\sigma_2(C8-C10) \rightarrow \sigma^*_2(O3-C7)$ with the stabilization energies 27.83 kcal/mol. Additionally, the molecule has lone electron pairs of oxygen atom interactions $n(LP_2O1) \rightarrow \sigma_2^*(O2-C5)$, $n(LP_2O1) \rightarrow \sigma_1^*(C8-C10)$, $n(LP_2O2) \rightarrow \sigma_1^*(O1-C5)$, $n(LP_2O2) \rightarrow \sigma_1^*(C5-C6)$, and $n(LP_2O3) \rightarrow \sigma_1^*(O4-H44)$ with energies 31.36, 34.08, 37.96, 15.85, and 63.18 kcal/mol.

3.4. Vibrational Analysis

The determination of the type of vibration in the experimental spectra was carried out by analyzing the potential energy, the atomic displacements, and the comparison with related compounds [29,30]. The experimental and calculated vibrational spectra for two tautomers A and B of compound 1 are shown in Figures 4 and 5 and Table 3.

The sharp medium-intensity peak in the IR spectrum at 3107 cm^{-1} and the frequency of 3108 cm^{-1} in the



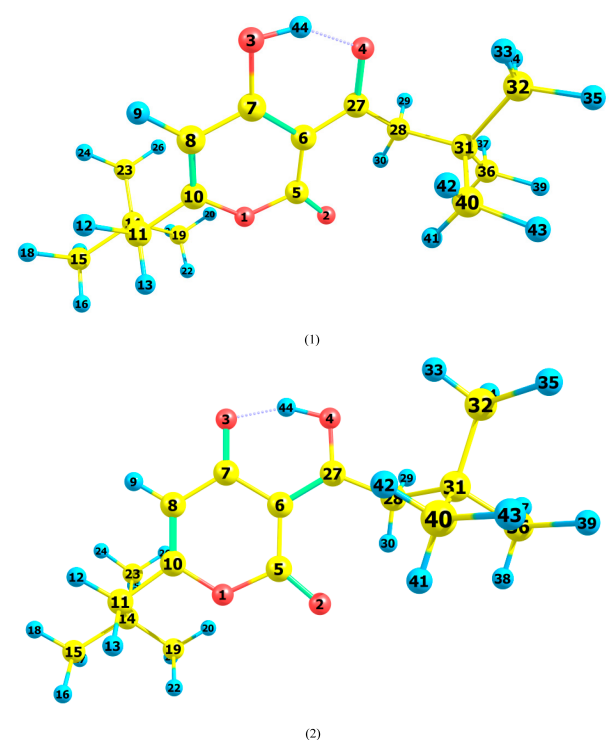


Figure 2: Geometry and atom numbering for tautomeric forms A(1) and B(2) of compound 1.

Raman spectrum refer to CH stretching vibrations (C8–H9 bond). Experimental frequencies in the region 3100–2960 cm⁻¹ in vibrational spectra refer to $\nu_{as}(CH_2)$ and $\nu_{as}(CH_3)$ stretching vibrations. The symmetrical stretching vibrations of the methyl and methylene groups cause

frequencies in the range of 2910–2860 ${\rm cm}^{-1}$ in the experimental spectra.

Stretching of carbonyl groups without H-bonds (C5=O bond) causes a band at 1719 cm⁻¹ in the experimental IR and Raman spectra (Figures 4 and 5). The stretch-



Table I: Gibbs relative free energy ΔG (kcal/mol), Boltzman weighting factor p (%) of low energy tautomers of 1 calculated in the B3LYP/6-311**G level.

Gas			Chlorofori	n	Dimethylsi	Dimethylsulfoxide		
Tautomer	ΔG	p	ΔG	р	ΔG	p		
A	0	90	0	87	0	90		
В	1.35	10	1.12	13	1.32	10		

Table 2: Calculated ionization energy (I), electron affinity (A), energy band gap (|GAP|), chemical potential (μ), global softness (S), global electrophilicity index (ω), and dipole moment (M) for tautomeric forms A and B of compound $\mathbf{1}$.

Tautomer	I, eV	A, eV	GAP	μ , eV	S, eV	ω , eV	M, D	
A	8.661	0.803	4.618	-4.732	0.127	2.850	3.414	
В	8.795	0.883	4.693	-4.839	0.126	2.960	1.935	

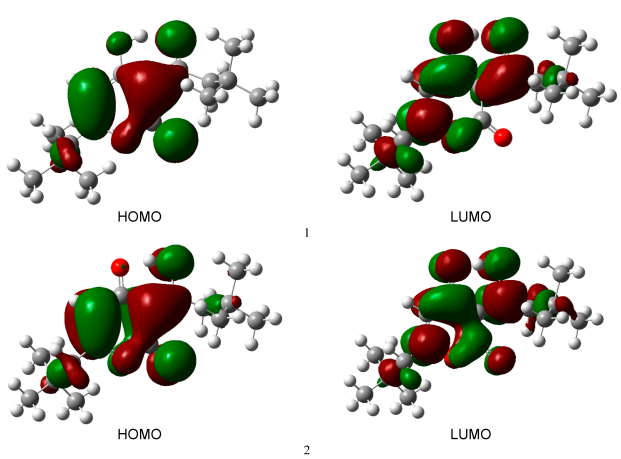


Figure 3: Molecular orbital surfaces for tautomeric forms A(1) and B(2) of compound 1.

ing vibrations of the carbonyl group forming the H-bond (C3=O7 bond) are shifted to the low-frequency region of 1636 cm⁻¹.

The bands in the region 1460–1400 cm $^{-1}$ in the experimental spectra are due to the $\delta_{as}(CH_3)$ and $\delta_{as}(CH_2)$ bending vibrations. Symmetrical bending vibrations $\delta_s(CH_3)$ and $\delta_s(CH_2)$ cause bands in the region 1400–1340 cm $^{-1}$. The frequency at 1315 cm $^{-1}$ in the exper-

imental IR spectrum and the band at 1318 cm⁻¹ in the experimental Raman spectrum are due to the wagging vibrations of the methylene groups.

Stretching vibrations of the *CO* and *CC* bonds of the pyran ring (C5–O1 and C5–C6 bonds) cause frequencies between 1290 and 1150 cm⁻¹ in the experimental spectra. The bands in the region of 1060 to 1000 cm⁻¹ in the ex-



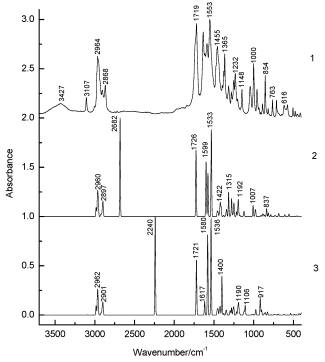


Figure 4: Experimental IR spectra of crystalline compound 1 (1) and calculated IR spectra of tautomeric forms A(2) and B (3). The theoretical spectra were modeled using Lorentz functions centered on the calculated frequencies, scaled by a factor of 0.96 and an FWHM of 10 cm^{-1} .

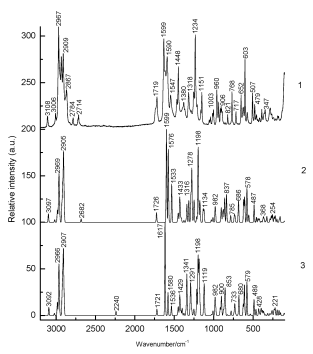


Figure 5: Experimental Raman spectra of crystalline compound 1 (1) and theoretical Raman spectra of tautomeric forms A(2) and B (3). The theoretical spectra were modeled using Lorentz functions centered on the calculated frequencies, scaled by a factor of 0.96 and an FWHM of 10 cm^{-1} .

perimental spectra were attributed to the deformation of the *CCH* angles and stretching of the *CC* bonds.

The rocking vibrations of the methyl groups $\rho(CH_3)$ cause bands between 960 and 880 cm⁻¹ in the



experimental IR and Raman spectra. Bands of average intensity in the region of 860–760 cm⁻¹ in the experimental spectra refer to *CO* and *CC* bonds (C5–O1 and C5–C6 bonds) stretching vibrations.

The band at 713 cm⁻¹ in the experimental spectra refers to the stretching vibrations of the CC bonds (C10–C11 bond). Bending vibrations of the pyran ring cause bands in the 620–470 cm⁻¹ region in the experimental spectra. Bending and torsional vibrations of the pyran ring cause bands in the 500–100 cm⁻¹ region of IR and Raman spectra.

It is important to understand the changes that occur in the vibrational spectra of compound 1 during the tautomeric transformation. The spectra of tautomeric forms A and B are similar (Figures 4 and 5). For tautomers A and B, the frequencies of most bands remain unchanged, but their intensity changes. Bands 1726, 1599, 1576, 1533, 1422, 1315, and 1192 cm⁻¹ of the form A IR spectrum are shifted to frequencies 1721, 1617, 1580, 1536, 1400, 1341, and 1190 cm⁻¹ of the form B IR spectrum (Figure 4). Bands 1726, 1599, 1576, 1533, 1433, 1340, 1316, 1278, 1249, 1198, 837, 686, 607, 578, 487 cm⁻¹ in the Raman spectrum of form A are shifted to frequencies 1721, 1617, 1580, 1536, 1429, 1341, 1291, 1198, 853, 680, 579, 489 cm⁻¹ in the Raman spectrum of form B (Figure 5).

The theoretical spectra align with the experimental vibrational spectra of tautomer A across a wide frequency range (Figures 4 and 5). Thus, the use of the DFT approximation for the considered molecular system is correct.

3.5. Hydrogen Bond

Compound 1 has a strong H-bond with a cyclic chelate structure. The enol form of β -diketones is a well-known case of a 6-membered ring (chelate) with a strong resonance-assisted H-bond [31–34]. The characteristic spectral features observed for such structures include strong absorption bands in the region of 1580–1630 cm⁻¹, instead of pronounced carbonyl stretching bands [16]. The stretching of the OH bond band shifts up to 2200 cm⁻¹ [16].

Additionally, due to the resonance of two enol tautomers (both are chelate enol forms), this structural fragment can be considered a quasisymmetric O···H···O hydrogen bond. This bond has a broad and shallow potential well with two minima, which leads to a series of diffuse absorption bands scattered over a wide wavenumber region.

The analysis of the observed IR spectrum 1 in the region of 2700–2000 cm⁻¹ shows that there are several weak bands, $\nu(OH)$ (Figure 6). The calculated $\nu(OH)$ frequencies after scaling are 2682 and 2240 cm⁻¹ for tau-

tomers A and B, respectively, and they are in the range between 2700 and 2200 cm $^{-1}$. An empirical formula is proposed that establishes a correlation between the frequencies $\nu(OH)$ observed and those calculated in the harmonic approximation: $\nu_{\rm obs} = -757 + 1.173 \ \nu_{\rm harm}$ [33]. The calculation by this formula yields $\nu(OH)$ frequencies of 2520 and 1980 cm $^{-1}$ for tautomers A and B, respectively. In the experimental IR spectrum of compound 1, the band 2524 cm $^{-1}$ is observed, which can be attributed to $\nu(OH)$ vibrations. The shift of this band to low frequencies depends on the strength of the intramolecular H-bond. We see that in the most energy-stable tautomer A, the H-bond is weaker.

This conclusion is consistent with the fact that the H-bond length for A and B tautomers is 2.473 and 2.418 Å, respectively. The strength of the H-bond can be described using Wiberg bond indices 0.158 (tautomer A) and 0.204 (tautomer B) [26]. These values of the Wiberg indices indicate that strong H-bonds are formed in compound 1 for both tautomers. It appears from our data that there is a correlation between the frequencies $\nu(OH)$ calculated in the harmonic approximation and the Wiberg indices. The stronger the H-bond, the lower the $\nu(OH)$ frequency and the higher the Wiberg index.

The strength of the intramolecular H-bond in compound 1 can be estimated as the interaction $n(LP_2O4) \rightarrow \sigma_1*(O3-H44)$ with the energy 42.96 kcal/mol of the tautomer A and $n(LP_2O3) \rightarrow \sigma_1*(O4-H44)$ with energy 63.18 kcal/mol of tautomer B. The energy of the donor-acceptor interaction of the H-bond is larger for the B tautomer; therefore, it is stronger. Interactions $\sigma_1(O3-H44) \rightarrow \sigma^*_1(C7-C8)$, $\sigma_2(C6-C7) \rightarrow \sigma^*_2(O4-C27)$, $\sigma_2(C6-C7) \rightarrow \sigma^*_2(C8-C10)$, $\sigma_2(C8-C10) \rightarrow \sigma^*_2(C6-C7)$ with energies 5.83, 31.20, 7.49, 24.96 kcal/mol (tautomer A), and $\sigma_2(C8-C10) \rightarrow \sigma^*_2(O3-C7)$ with energy 27.83 kcal/mol (tautomer B) are realized due to the conjugation of bonds in a six-membered ring.

4. Conclusions

In conclusion, the correlation between the structure and H-bonding was established in 3-(3,3-Dimethylbutanoyl)-4-hydroxy-6-neopentyl-2H-pyran-2-one. For pyrone 1, the X-ray diffraction, DFT-calculations and IR, Raman spectroscopy revealed the most favourable tautomeric form *A*. The content of tautomer *B* increases in the nonpolar solvent but does not exceed 13%.

As can be seen from the calculations and experimental X-ray data, the pyran ring of the molecule is flat. A satisfactory agreement is observed between the calculated geometrical parameters of tautomer A and the experimental X-ray data.



Table 3: Observed and calculated wavenumbers ν (cm⁻¹), the intensity of the bands in the IR spectra I (km/mol) and relative intensity of the bands in the Raman spectra J (a.u.) and assignments for the tautomeric forms A and B of compound 1 in the gas phase by using the B3LYP/6–311++G** method.

Experimental		B3LYP/6-311++G** Method							
R	Raman	A		В		Assignments			
/	ν	ν	I	J	ν	I	J		
427w									
107w	3108w	3097	1.1	9.2	3092	1.1	8.8	νC8–H9	
004vw	3006w	3018	1.0	2.7	3021	0.6	2.2	νC28–H29 ν _{as} CH ₂	
		2985	30.6	5.7	2984	26.8	5.5	νC19–H20 ν _{as} CH ₃	
		2984	26.3	4.7	2984	33.3	4.8	νC32–H33 ν _{as} CH ₃	
		2980	17.6	4.8	2983	16.2	4.2	νC40–H41 ν _{as} CH ₃	
		2969	64.4	14.8	2969	55.8	10.8	νC23–H24 ν _{as} CH ₃	
		2966	33.2	11.1	2966	43.1	13.4	νC15–H16 ν _{as} CH ₃	
964s	2967s	2964	43.8	12.8	2965	30.9	7.2	νC11–H12 ν _{as} CH ₂	
		2963	26.0	6.0	2963	45.5	12.3	νC36–H37 ν _{as} CH ₃	
		2961	3.1	2.1	2962	70.0	18.6	ν C19–H21 ν_{as} CH ₃	
		2960	79.0	21.3	2961	1.9	1.8	ν C32–H34 ν _{as} CH ₃	
956sh		2957	8.0	1.8	2957	6.7	1.8	ν C23–H25 ν _{as} CH ₃	
		2956	3.8	1.5	2956	4.4	1.2	ν C25 H25 ν as CH3 ν C15–H17 ν as CH3	
		2954	4.7	1.9	2956	22.0	2.7	ν C32–H34 ν _{as} CH ₃	
		2953	26.1	2.9	2955	3.2	1.2	$vC40$ –H42 v_{as} CH ₃	
	2933m	2926	15.8	18.8	2923	14.8	20.7	$vC40=H42 v_{as} CH_3$ $vC28=H29 v_{as} CH_2$	
	2733111	2926	13.8	37.1	2923	14.8	35.8	ν C26–H29 ν _{as} CH ₂ ν C11–H12 ν _{as} CH ₂	
	2909m	2908	19.5	22.9	2918	20.3	21.8	ν C11–H12 ν _{as} CH ₂ ν C19–H21 ν _s CH ₃	
906m	2707111	2905	26.0	50.4	2908	23.9	42.3	ν C19=H21 ν_s CH3 ν C32=H33 ν_s CH3	
JUUIII		2905 2900	25.1	30. 4 12.2	2907	26.5	42.3 1.4	$vC32-H33 v_s CH_3$ $vC23-H24 v_s CH_3$	
		2900 2899	28.5						
			28.5	1.0	2900	26.0	9.6	ν C40–H41 ν_s CH ₃	
868m	2867w	2897 2895		2.2	2896	28.7	3.4	ν C15–H16 ν_s CH ₃	
			31.1	4.6	2896	29.3	4.5	ν C36–H37 ν_s CH ₃	
675vw	2784vw	2682	709.1	3.7	2240	787.7	4.4	νO3–H44	
666vw	2714vw								
666vw									
631vw									
608vw									
568vw									
524vw									
372vw									
364vw									
355vw									
2338vw	4=00.4								
737sh	1738sh	1-0-	4=0 =		1-0:	440 =		05.00	
719vs	1719m	1726	479.7	10.4	1721	440.3	6.8	νC5=O2	
683w									
636s	1633vs	1599	394.4	100.0	1617	110.3	100.0	νC3=O7	
615w	1617sh								
588m	1590m	1576	311.0	78.6	1580	641.7	24.6	βΟ3–Η44	
553vs	1547m	1533	626.8	41.2	1536	775.5	10.3	νC6–C7	
174sh									
462sh	1467m	1460	18.0	0.7	1460	21.1	0.4	β C32–H33 δ_{as} CH ₃	
455m		1458	17.0	0.9	1458	19.6	0.6	βC19–H20 δ _{as} CH ₃	
		1452	12.7	1.5	1451	19.4	1.6	βC36–H39 δ _{as} CH ₃	
		1451	4.9	5.1	1450	14.0	3.7	βC15–H16 δ _{as} CH ₃	
		1449	6.7	1.5	1449	6.6	1.4	βC11–H12 δ _{as} CH ₂	
445sh	1448m	1447	11.3	2.6	1447	10.0	3.4	β C28–H29 δ _{as} CH ₂	
	1438sh	1433	7.8	11.7	1432	20.9	5.3	β C40–H41 δ _{as} CH ₃	
437sh	1420811			11./	1734	20.7	٥.٥		



		1429	3.4	7.4	1429	2.0	7.5	β C15–H16 δ_{as} CH ₃
		1428	6.4	4.2	1428	3.5	5.0	β C19–H20 δ_{as} CH ₃
		1425	0.4	0.2	1425	0.4	0.2	β C23–H25 δ_{as} CH ₃
		1424	2.4	0.3	1424	2.0	0.0	β C36–H37 δ_{as} CH ₃
		1422	77.7	4.5	1422	63.1	1.3	β C28–H29 δ_s CH ₂
		1417	32.8	2.2	1415	6.4	4.6	β C11–H12 δ_s CH ₂
1394vw		1409	64.0	4.0	1400	312.5	7.8	β C28–H29 δ_s CH ₂
1376w	1380m	1379	4.3	1.4	1379	5.8	1.1	β C19–H21 δ_s CH ₃
1365m	1369sh	1376	4.7	0.8	1377	6.8	1.3	β C32–H35 δ_s CH ₃
	1365sh	1351	7.6	1.7	1351	6.2	3.7	β C19–H20 δ_s CH ₃
		1349	6.0	0.3	1350	14.3	0.3	βC32–H33 δ _s CH ₃
		1347	7.3	0.3	1346	8.3	1.2	βC23–H24 δ _s CH ₃
		1344	7.5	0.6	1345	8.6	0.6	βC36–H37 δ _s CH ₃
		1340	44.1	18.9	1341	23.0	43.3	νC5-O1
1315w	1318m	1316	176.4	23.6	1298	30.1	13.8	βC28–H29 wag CH ₂
		1289	8.9	29.5	1291	13.8	29.5	βC11–H12 wag CH ₂
1282w	1285vw	1278	129.0	57.2	1279	58.2	5.9	νC6–C7
- - ··		1261	16.8	2.1	1260	39.7	1.7	β C28–H29 wag CH ₂
1251w	1253m	1249	97.4	23.9	1251	63.1	6.6	vC6–C27
1232m	1234vs	1216	34.5	6.1	1215	26.9	28.2	vC6–C27
1203w	1208w	1198	57.3	72.3	1198	60.9	61.8	βC23–H24 wag CH ₃
1203 W	1200W	1192	98.5	21.7	1190	90.7	22.0	βC32–H35 wag CH ₃
		1177	4.7	10.8	1184	8.2	49.9	vC5–C6
		1174	2.9	12.4	1176	7.5	3.0	βC23–H24 τw CH ₃
1161sh	1163w	1170	4.2	3.2	1174	3.4	9.7	νC5–O1
1148m	1151m	1134	2.0	14.2	1128	11.0	8.1	βC8–H9
1140111	1131111	1123	25.1	11.1	1122	3.4	16.2	βC28–H29 τw CH ₂
	1056vw	1119	19.1	3.6	1119	29.0	18.1	βC11-H12 τw CH ₂
1044m	1041w	1030	0.7	2.7	1106	78.5	0.4	βC40–H41
1044111	1041w 1014w	1017	0.6	0.8	1007	2.2	1.0	βC23–H24
	1014W	1017	1.2	1.7	1018	4.0	0.3	βC36–H39
		1009	15.2	2.4	1010	1.7	2.7	βC23–H24 τw CH ₃
1000m	1003w	1007	64.9	1.3	1008	4.1	2.3	δO3–H44
1000111	1005W	982	22.5	13.9	982	1.2	18.8	vC7–C8
959m	960m	976	41.1	7.2	973	44.3	7.2	νO1–C10
935w	937w	928	0.1	0.1	928	0.5	0.1	γC32–H33 ρ CH ₃
929sh	93 / W	926	0.1	0.1	926	0.3	0.1	γC23–H24 ρ CH ₃
912vw	914w	910	3.7	2.7	917	135.1	8.4	γC28–H29 ρ CH ₃
912VW	906w	907	4.9	2.7	907	1.4	1.3	γC19–H20 ρ CH ₃
	900W	907	2.1	8.9	907	2.9	7.7	νC14–C23
	899w	903	0.4	4.4	903	3.7	3.1	νC31–C36
888w	891w	902 887	17.5	9.3	902	35.3	10.5	
ooow	891W	877	17.3					νC5–C6, νC5–O1
	0.61 -1-			12.9	876	1.0	6.3	vC14-C19
0.5.4	861sh	862	3.2	3.9	865	12.8	11.0	vC14-C15
854m	856vw	861	6.9	22.6	857	5.1	14.3	νC10-C11
017	844sh	837	57.4 26.2	30.0	853	16.6	18.5	νO1–C5
817w	821w	815	26.3	0.3	826	25.0	0.7	γC8–H9
779sh	768m	785 728	13.7	7.4	779 742	2.5	4.8	ρC27=O4
763w	754sh	738	2.6	7.6	742	4.5	2.4	vC8–C10
712	717	729 720	7.2	5.7	733	2.1	12.5	ρC7-O3
713w	717w	720	1.4	0.2	721	2.8	1.9	vC10-C11
697sh	(52	686	18.3	21.3	691	7.4	0.6	vC27–C28
686sh	652m	680	6.8	3.9	680	0.3	21.8	ρC7–O3
650sh	625w	620	1.1	24.4	625	2.2	12.2	νC10-O1
616w	603m	607	12.3	27.0	611	2.8	32.7	ρC7–O3
574w	575w	578	0.4	36.4	579	0.8	35.6	νC10–C11
544sh		555	17.8	8.0	535	12.8	1.5	$\beta_{ m R}$
536sh								



529sh								
505w	507m							
477vw	479w	487	2.6	18.8	489	6.0	17.1	β_R
453vw	456w	458	4.1	5.4	464	2.0	6.6	β_R
445vw		438	3.1	4.3	430	10.3	2.6	$\beta_{\rm R}$
436vs	439vw	426	1.7	1.9	428	0.0	4.8	$\beta_{ m R}$
412vw	418vw	419	0.2	3.1	417	8.3	2.4	$ au_{ m R}$
		396	0.1	0.7	400	14.6	5.2	$ au_{ m R}$
	389w	391	3.1	0.5	395	4.8	3.9	βC27–O4
	381sh	375	3.5	2.4	383	11.1	5.5	ρC5=O2
		368	0.3	5.3	369	0.4	4.4	βC14_C19
	359sh	359	0.6	2.9	360	3.0	1.8	, βC27–C28
	347w	340	2.1	1.6	340	0.9	0.8	γC10–C11
		326	6.6	6.4	315	3.0	2.1	βC27–O4
	288sh	287	0.4	3.0	288	0.2	1.7	τC14-C15
	281w	283	1.5	0.6	283	0.2	0.3	τC31–C32
		280	0.3	0.4	280	0.2	0.2	τC14–C15
		276	0.1	1.1	277	0.1	0.9	τC14–C23
	264vw	269	2.2	6.5	265	2.1	2.8	$ au_{ m R}$
		258	1.0	2.6	260	0.3	2.2	τC14–C23
		254	0.9	6.5	255	0.5	3.5	β_R
	241vw	235	1.3	4.5	236	0.4	4.3	τC14–C23
		230	0.2	1.9	230	1.5	3.7	τC31-C40
		220	0.6	6.0	221	0.3	6.1	τC14-C15
	211vw	219	0.3	1.1	219	0.6	2.0	$ au_{ m R}$
	188vw	199	3.6	3.6	195	0.7	5.9	$ au_{ m R}$
	178vw	167	0.3	3.1	171	1.0	5.4	$ au_{ m R}$
	168vw	157	2.7	6.2	152	2.5	3.0	$ au_{ m R}$
	105vw	103	0.2	1.3	105	0.9	2.4	$ au_{ m R}$

Abbreviations: vs, very strong; s, strong; m, medium; w, weak; vw, very weak; γ , stretching; β , deformation in plane; γ , deformation out of plane; wag, wagging; τ , torsion; β_R , deformation ring; τ_R , torsion ring; ρ , rocking; τ_R , twisting; δ , deformation; a, antisymmetric; s, symmetric.

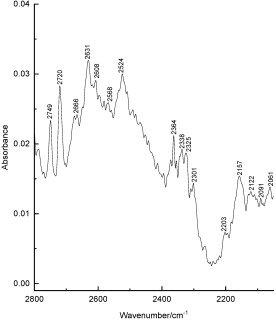


Figure 6: The experimental IR spectrum of compound 1 in the region $3600-2700 \text{ cm}^{-1}$.



The calculation of the normal vibrations by the DFT method gives a detailed description of the dynamics of pyrone 1. The intensities of the bands in the IR spectra show high sensitivity to the H-bond in compound 1.

The HOMO and LUMO orbitals of the acid molecule are located on the pyran ring. During tautomeric transformations, there is a significant delocalization of the charge, which modifies the reactivity of the molecule.

The reactivity of compound 1 was characterized using descriptors. Form B was found to have higher ionization energy, electron affinity, chemical potential, and electrophilic index than Form A. The dipole moment is higher for Form A, and the softness of the two molecules is the same.

The obtained results provide the opportunity better understand the interplay between the tautomeric flexibility of the pyrone ring and its H-bonding, providing a new approach for rational design of drugs with desired properties.

Abbreviations

IR Infrared

DFT Density Functional Theory
NMR Nuclear Magnetic Resonance
FTIR Fourier Transform Infrared
NBO Natural Bond Orbital

HOMO Highest Occupied Molecular Orbital LUMO Lowest Unoccupied Molecular Orbital

IE Ionization Energy EA Electron Affinity

FWHM Full Width at Half Maximum

Author Contributions

V.F.: conceptualization, methodology, software, writingoriginal draft preparation and editing. A.V.: investigation of IR and Raman spectra. E.N., M.M.: synthesis of pyran. V.K.: conceptualization, methodology, reviewing and editing. V.K.: conceptualization, methodology, reviewing and editing.

Availability of Data and Materials

No datasets were generated or analyzed during the current study.

Consent for Publication

Not applicable.

Conflicts of Interest

The authors declare no competing interests.

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Supplementary Materials

Supplementary materials are available for download here.

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