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FRONTIER MOLECULAR, NATURAL BOND ORBITAL, UV-VIS SPECTRAL STUDY, SOLVENT INFLUENCE ON GEOMETRIC PARAMETERS, VIBRATIONAL FREQUENCIES AND SOLVATION ENERGIES OF 8-HYDROXYQUINOLINE

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ABSTRACT: *N*-heterocyclic compounds have extensive biological and pharmaceutical applications. 8-Hydroxyquinoline (8-HQ) also plays a significant role in many fields of life. The excellent biological significance of the 8-HQ prompted us to extend the DFT based studies. The frontier molecular orbitals (FMOs), UV-VIS and solvation model based studies remained unknown. Therefore, we intended to study the natural bond orbital, FMOs, UV-VIS, thermodynamic properties and medium influence on solvation energies, dipole moment, FT-IR and FT-Raman using polarizable continuum model (PCM) and density-based solvation model (SMD). The electronic properties of molecule were calculated by M06-2X/6-31G (d,p) and B3LYP/6-31G (d,p) level of theories. The solvent influence on the geometric parameters, FT-IR and FT-Raman were studied by B3LYP /6-31G(d) method. A good correspondence is found between the optimized parameters and the reported X-ray data. Natural bond orbital reveals that the maximum stabilization energy reached up to 39.64kJ/mol which is responsible for extra stability of the molecule. In solvated 8-HQ, a significant medium effects on FT-IR and FT-Raman intensities is observed. The intensities enhanced from gas to solvent phase. The solvation free energies are found to be -28.710 and -39.456 kJ/mol in PCM and SMD models respectively. FMOs suggested that this molecule contain less hardness and larger softness values. These findings reveal that the molecule might be bioactive.

INTRODUCTION: *N*-heterocyclic compounds are most versatile organic compounds contain broad spectrum biological and pharmaceutical significance.¹

In this context, 8-hydroxyquinoline (1-azanaphthalene) is also one of the *N*-heterocyclic compounds which play a fundamental role in many fields of life. The compound 8-hydroxyquinoline (8-HQ) has been used as preservative in textile, wood and paper industries.² It is used in the chemicals which help to kill the crop insects and pests in an agricultural field.^{1, 2} It is also used as antimalarial,³ antibacterial,⁴ anticancer drugs,⁵ anthelmintic, amebicidal, anti-HIV⁶ and neuroprotective.³

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It is used as a powerful chelator for the treatment of diseases caused by metals.^{6, 2} A basic component of organic light emitting diode (OLEDs) is Alq3 which is produced by the reaction of 1-azanaphthalene and aluminum (III).⁴ 8-HQ and its derivatives have multiple applications that range from pharmacological and pharmaceutical agents to carries the electron in OLEDs and fluorescent chemo sensor for metal ions as well as very useful chelating agent in analytical and radioactive chemistry.^{2, 7} 8-HQ has strong fluorescence in concentrated acids, water dearth of fluorescence or clusters of hydrogen-bonded with water, alcohol and ammonium molecules.

Due to these outstanding applications, various DFT based studies have been reported on this molecule as; Arıcı *et al.* (2005) have reported IR in gas phase.⁸ Krishnakumar *et al.* (2005) also reported FT-IR and FT-Raman spectra in gas phase.⁹ R. P. Gangadharan *et al.* (2014) reported NBO using B3LYP with 6-31G (d, p) basis set,¹⁰ however, some discrepancies have been observed regarding the assignment of NBO analysis.

Usually, FT-IR vibrational spectra based on computational analysis is performed in the gas phase as carried out regarding this molecule.^{6, 7} However, the experimental studies mostly include in solvent phase. Hence, it is important to consider solvent effects in computational approaches. The chemistry of the molecule under investigation can be changed due to synergic effect of the solvent and the solute. The solvent-solute association can modify stability, energy (i.e. vibrational frequency, spectrum) and molecular arrangement as well as play a vital role in the pharmaceutical industry for drug design.¹¹

As In 1890, W. Ostwald wrote:

“Almost all the chemical processes, which occur in nature, whether in animal or vegetable organisms or in non-living surface of the Earth ... take place between substances in solution”

To the best of our knowledge, neither frontier molecular orbitals (FMOs), global reactivity parameters, UV-VIS analysis nor the solvation models based DFT regarding title molecule are reported so for.

Therefore, the objective of our study is to present the frontier molecular orbitals (FMOs), global reactivity parameters, UV-VIS analysis and also to explore the solvent effects on geometry, FT-IR, FT-Raman spectra and solvation energy of title molecule using Polarizable Continuum Model (PCM) and Density-based Solvation Model (SMD). Moreover, one of the main objectives of our study is also to improve the scientific contribution by elucidating the right assignment in NBO analysis using same level of theory as reported [B3LYP/6-31G (d,p)] and more accurate M06-2X/6-31G (d,P) method.

MATERIALS AND METHODS: Complete calculations were performed using Gaussian 09 program package¹² with density functional theory (DFT)¹³⁻¹⁵ at the M06-2X/6-31G (d,P) and B3LYP/6-31G (d,p)/ 6-31G (d) levels of theory. The geometry optimization and frequency calculations were performed for title compound. Consequently, a nonattendance of negative eigen value demonstrated that the geometry of the compound is in the globe minima of the potential energy surface.

All input files were prepared using Gauss View 5.0.16 Chemcraft,¹⁷ Gauss View 5.0 and Avogadro software¹⁸ were used to analyze the output files.

RESULTS AND DISCUSSION: Equilibrium geometry: The 8-HQ geometry is optimized by using M06-2X and B3LYP level of theories with basis set as 6-31G(d,p). However, the molecule is also optimized at B3LYP/6-31G(d) method in gas phase and dimethyl sulfoxide as a solvent to study the solvent influence on optimized geometry. The obtained parameters are tabulated in **Table 1**.

The geometry of a compound is very important to study the vibrational frequencies since slight alteration in geometry can significantly alter the frequency. The intermolecular forces are more essential in this context, when any molecule goes from gaseous to solution phase. Subsequently solute-solvent interactions take place and might be geometry of solute changes.⁸ Keeping in view these solute-solvent interactions, we studied our system in gas and solvent phase. **Table 1** shows a variation in bond lengths and angles.

The bond lengths such as R(1-2), R(3-4), R(4-18), R(5-16), R(10-13), R(16-17) are slightly increased in solvent phase than gas phase. However, some bond lengths like that R(1-8), R(2-9), R(4-5), R(6-11), R(7-10), R(12-13) and R(13-15) are slightly decreased in solvent phase as compared to gas phase. These variations are occurred only in third decimal. Interestingly, some bond lengths i.e. R(1-6), R(2-3), R(3-10), R(12-14), R(12-18) are found exactly similar in both phases. Moreover, the obtained bond lengths show good agreement to the X-ray data ¹⁹ and DFT based observed bond lengths. ⁶

Furthermore, the bond angles like A(1-2-9), A(1-6-11), A(4-3-10), A(3-10-7), A(5-4-18), A(4-5-6),

A(5-16-17), A(13-12-18) and A(14-12-18) are slightly increased in solvent phase than gas phase. While an opposite trend is observed in A(1-6-5), A(2-3-4), A(2-3-10), A(3-4-18), A(6-5-16) and A(12-13-15) where bond angles are decreased in solvent phase to a little extent than gas phase. However, bond angles such as A(2-1-6), A(2-1-8), A(1-2-3), A(6-1-8), A(3-2-9), A(3-4-5), A(3-10-13), A(4-5-16), A(4-18-12), A(10-13-12) and A(10-13-15) exhibited exactly similar trend in both phases. Hence, the straightforward relation is not observed between solvent and gas phase.

The found bond angles show correspondence with the X-ray data ¹¹ and DFT based calculated bond angles. ¹²

TABLE 1: COMPARISON OF SELECTED BOND LENGTHS (Å) AND ANGLES (°) FOR 8-HQ.

Bond length	Gas Phase	Solvent Phase ^a	Bond Angle	Gas Phase	Solvent Phase ^a
R(1-2)	1.375	1.377	A(2-1-6)	120.9	120.9
R(1-6)	1.414	1.414	A(2-1-8)	120.3	120.3
R(1-8)	1.087	1.086	A(1-2-3)	119.7	119.7
R(2-3)	1.42	1.42	A(1-2-9)	121	121.1
R(2-9)	1.087	1.086	A(6-1-8)	118.8	118.8
R(3-4)	1.431	1.432	A(1-6-5)	120.9	120.7
R(3-10)	1.42	1.42	A(1-6-11)	119.6	119.7
R(4-5)	1.434	1.433	A(3-2-9)	119.3	119.3
R(4-18)	1.361	1.364	A(2-3-4)	120.3	120.2
R(5-6)	1.383	1.383	A(2-3-10)	122.9	122.8
R(5-16)	1.358	1.362	A(4-3-10)	116.8	117
R(6-11)	1.089	1.088	A(3-4-5)	118.4	118.4
R(7-10)	1.088	1.087	A(3-4-18)	123	122.8
R(10-13)	1.374	1.375	A(3-10-7)	119.1	119.2
R(12-13)	1.418	1.416	A(3-10-13)	119.7	119.7
R(12-14)	1.09	1.09	A(5-4-18)	118.6	118.8
R(12-18)	1.32	1.32	A(4-5-6)	119.9	120
R(13-15)	1.086	1.085	A(4-5-16)	116.9	116.9
R(16-17)	0.971	0.972	A(4-18-12)	117.8	117.8

B3LYP /6-31G(d), ^asolvent=DMSO

TABLE 1: Continued.....

Bond Angle	Gas Phase	Solvent Phase ^a
A(6-5-16)	123.2	123
A(5-6-11)	119.5	119.6
A(5-16-17)	108.4	108.7
A(7-10-13)	121.2	121.1
A(10-13-12)	118.6	118.6
A(10-13-15)	121.5	121.5
A(13-12-14)	119.6	119.5
A(13-12-18)	124.1	124.2
A(12-13-15)	120	119.9
A(14-12-18)	116.3	116.4

Natural Bond Orbital (NBO) Analysis: The natural bond orbital (NBO) analysis is a proficient method to study the intra- and inter-molecular bonding, the charge transfer and conjugative interaction between the electron donor and acceptor.^{9, 2} In the presence of large stabilization energy electron donors have more rigorous interactions with electron acceptors. NBO analysis is useful to transmit of electron density form filled electron orbital to unoccupied orbital.²⁰ According

to second order perturbation approach, the stabilization energy formula can be shown by Equations 1.

$$E^{(2)} = q_i \frac{(F_{i,j})^2}{\varepsilon_j - \varepsilon_i} \quad \text{Equations 1}$$

Where $E^{(2)}$ is stabilization energy, q_i donor orbital occupancy, $F(i,j)$ is the diagonal and ε_j and ε_i are the off-diagonal NBO Fock matrix elements.

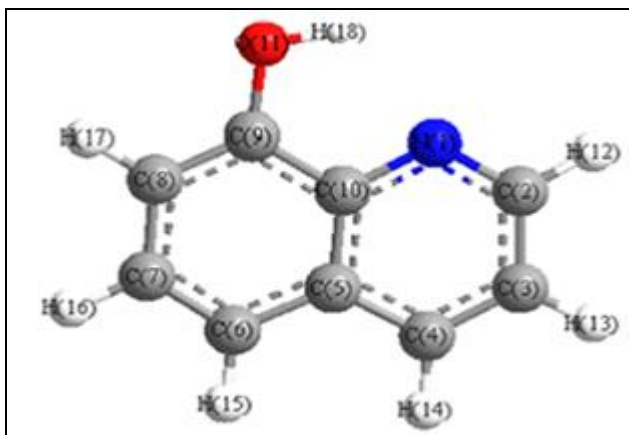


FIG. 1: STRUCTURAL REPRESENTATION OF 8-HUDROXYQUINOLINE

TABLE 2: NATURAL BOND ORBITAL (NBO) ANALYSIS USING B3LYP/6-31G(d,P)

Donor(i)	Type	Acceptor (j)	Type	$E(2)^a$	$E(j) - E(i)^b$ [a.u.]	$F(i; j)^c$ [a.u.]
C1 - C2	π	C3 - C4	π^*	17.81	0.28	0.067
C1 - C2	π	C5 - C6	π^*	14.91	0.29	0.060
C3 - C4	π	C1 - C2	π^*	14.81	0.28	0.060
C3 - C4	π	C5 - C6	π^*	18.33	0.27	0.065
C3 - C4	π	C10 - C13	π^*	17.28	0.27	0.065
C3 - C4	π	C12 - N18	π^*	15.88	0.26	0.060
C5 - C6	π	C1 - C2	π^*	19.93	0.30	0.070
C5 - C6	π	C3 - C4	π^*	13.68	0.29	0.059
C10 - C13	π	C3 - C4	π^*	15.21	0.29	0.063
C10 - C13	π	C12 - N18	π^*	23.11	0.28	0.073
C12 - N18	π	C3 - C4	π^*	19.34	0.33	0.076
C12 - N18	π	C10 - C13	π^*	12.18	0.33	0.057
O16	LP(2)	C5 - C6	π^*	32.52	0.35	0.099
N18	LP(1)	C3 - C4	π^*	10.57	0.86	0.086
N18	LP(1)	C3 - C4	π^*	2.20	0.85	0.039
N18	LP(1)	C4 - C5	π^*	10.65	0.87	0.087

^a $E^{(2)}$ means energy of hyper conjugative interaction (stabilization energy in kJ/mol).

^bEnergy difference between donor and acceptor i and j NBO orbitals.

^c $F(i; j)$ is the Fock matrix element between i and j NBO orbitals.

TABLE 3: NATURAL BOND ORBITAL (NBO) ANALYSIS USING B3LYP/6-31G(d,P)

Donor(i)	Type	Acceptor (j)	Type	$E(2)^a$	$E(j) - E(i)^b$ [a.u.]	$F(i; j)^c$ [a.u.]
C1 - C2	π	C3 - C4	π^*	23.29	0.36	0.087
C1 - C2	π	C5 - C6	π^*	19.12	0.37	0.077
C3 - C4	π	C1 - C2	π^*	18.80	0.36	0.077
C3 - C4	π	C5 - C6	π^*	22.60	0.35	0.082
C3 - C4	π	C10 - C13	π^*	22.67	0.35	0.084
C3 - C4	π	C12 - N18	π^*	21.48	0.35	0.080
C5 - C6	π	C1 - C2	π^*	27.10	0.39	0.092

C 5 - C6	π	C3 - C4	π^*	18.06	0.37	0.076
C10 - C13	π	C3 - C4	π^*	19.84	0.37	0.080
C10 - C13	π	C12 - N18	π^*	29.97	0.37	0.095
C12 - N18	π	C3 - C4	π^*	25.39	0.41	0.097
C12 - N18	π	C10 - C13	π^*	16.10	0.42	0.074
O16	LP(2)	C5 - C6	π^*	39.64	0.46	0.126
N18	LP	C3 - C4	π^*	11.89	1.01	0.099
N18	LP	C12 - C13	π^*	12.30	1.00	0.100

See **Table 2** foot note for the definition of superscripted alphabets (^{a, b, c})

The stabilization energies vary among the different orbitals as exhibited in **Table 2** and **3**. Presented data in **Table 2** has determined using B3LYP/6-31G(d,P) level of theory. The high probable transition takes place in 8-HQ as (C10-C13)→ $\pi^*(C12-N18)$ with 23.11 kJ/mol. It is the enormous value among all the stabilization energies. Other transitions such as $\pi(C12-C18)$ → $\pi^*(C3-C4)$ and $\pi(C3-C4)$ → $\pi^*(C5-C6)$ also show a very high stabilization energies as 19.34 kJ/mol and 18.33 kJ/mol respectively. While transition as $\pi(C12-N18)$ → $\pi^*(C10-C13)$ contains 12.18 kJ/mol. This is the least stabilization energy among all. This lowest energy results in the weak interaction between the electron donor and acceptor.

The same trend of interaction is also noticed in relation to resonance in the molecule. For example LP(O16)→ $\pi^*(C5-C6)$ produces 32.52 kJ/mol. This is the highest value. While LP(N18)→ $\pi^*(C3-C4)$ produces 2.20 kJ/mol which exhibits a very low electron donating interaction energy.

Presented data in **Table 3** has been determined using M06-2X/6-31G(d,P) level of theory. The interaction as $\pi(C10-C13)$ → $\pi^*(C12-N18)$ has the highest value (29.97 kJ/mol) among all the energies. Some other stabilization energies are 27.10 kJ/mol and 25.39 kJ/mol due to $\pi(C5-C6)$ → $\pi^*(C1-C2)$ and $\pi(C12-C18)$ → $\pi^*(C3-C4)$ respectively. These values also show the strong stabilization energies. However $\pi(C12-C18)$ → $\pi^*(C10-C13)$ and $\pi(C5-C6)$ → $\pi^*(C3-C4)$ have the interaction energies 16.10kJ/mol and 18.06 kJ/mol respectively, which are the least values among the whole data of donor (π) and acceptor (π^*). In case of the resonance, the transition as LP(O16)→ $\pi^*(C5-C6)$ has energy value 39.64 kJ/mol which is enormous energy value. While a least value 11.89 kJ/mol was exhibited by LP(N18)→ $\pi^*(C3-C4)$. By comparing the data of

both **Tables 2** and **3**, it is concluded that the highest energy values are calculated at M06-2X/6-31G(d,P) as compared to B3LYP/6-31G(d,p) level.

For example, the highest stabilization energies such as 39.64 and 29.97 kJ/mol are represented at M06-2X/6-31G(d,P) level (**Table 3**). In contrast, the maximum stabilization energies are found to be 32.52 and 23.11 kJ/mol at the B3LYP/6-31G(d,p) level (**Table 2**).

Some conflicts are detected with the assignment of the C10 reported by R. P. Gangadharan et al. (2014) in which the C10 containing LP(1) with high stabilization energy as 81.91 kJ/mol was reported.⁷ In the recent study, same reported method as B3LYP/6-31G(d,p) and new M06-2X/6-31G(d,P) method are used to clear this confusion from the literature. Anyway, there is no carbon of the molecule showed any lone pair [LP(1)] see **Table 2** and **3**. As it is obvious that carbon never contains any lone pair [LP(1)] in neutral molecule. Moreover, in the present study, the maximum stabilization energies reached up to 32.52 and 39.64 kJ/mol due to lone pairs [LP(2)] of oxygen in B3LYP/6-31G(d,p) and M06-2X/6-31G(d,P) methods respectively. These energies are around about 2 times less than reported values. Although, the M06-2X level is considered most authentic method to calculate the non covalent interactions (NCIs), stabilization energy, high accuracy in thermochemistry and excitation electronic energies for valence and rydberg states.²¹

FT-IR and FT-Raman analysis: The quantity of atoms in 8-HQ are 48 atoms with symmetry point group C1. The active vibrational normal modes are also found to be 48 in FT-IR and FT-Raman spectra. The vibrational modes were assigned with the help of animation option of Avogadro software. The calculated frequencies are summarized in **Table 4**.

TABLE 4: CALCULATED VIBRATIONAL FREQUENCIES OF 8-HQ BY M06-2X/6-31G(d,P)

^a Freq	^a I _{IR}	^a I _{Raman}	Vibrational assignments
416	80	5	$\nu(\tau)$ O-H
443	30	3	$\nu(\gamma)$ O-H+ $\nu(\gamma)$ C-H _{Ben}
474	0.56	7	$\nu(\delta)$ H-C=C-H
478	0.86	0.28	$\nu(\tau)$ C-H _{Ben}
492	0.48	5	$\nu(s)$ C-N=C-H
549	5	3	$\nu(\delta)$ O-H + $\nu(\delta)$ C-H _{Ben}
583	5	6	$\nu(\delta)$ C=C-C=N
598	0.43	3	$\nu(w)$ C-H _{Ben}
659	2.1	1.08	$\nu(\tau)$ C-H _{Ben}
729	6	27	$\nu(s)$ C-C=C
770	32	5	$\nu(w)$ C-H _{Ben}
813	14	2.34	$\nu(w)$ C-H _{Ben}
818	10	1.71	$\nu(s)$ H-C=C-H
839	53	0.04	$\nu(w)$ C-H _{Ben}
890	0.01	4	$\nu(\tau)$ C-H _{Ben}
902	3.4	2	$\nu(as)$ H-C=C-H
978	0.29	4	$\nu(\tau)$ C-H _{Ben}
982	0.96	0.19	$\nu(\tau)$ C-H _{Ben}
1020	0.9	0.07	$\nu(w)$ C-H _{Ben}
1060	6.8	5	$\nu(\tau)$ C-H _{Ben} + $\nu(s)$ C-C
1091	55	3	$\nu(\delta)$ C-H _{Ben}
1120	42	7	$\nu(\delta)$ C-H _{Ben}
1162	0.33	10	$\nu(\delta)$ C-H _{Ben}
1187	25	5	$\nu(\delta)$ C-H _{Ben} + $\nu(\rho)$ O-H
1227	21	2.28	$\nu(\rho)$ C-H _{Ben}
1259	22	1.38	$\nu(\delta)$ C-H _{Ben}
1296	43	3	$\nu(\rho)$ C-H _{Ben}
1326	21	5	$\nu(\rho)$ C-H _{Ben}
1397	82	10	$\nu(\rho)$ C-H _{Ben} + $\nu(\delta)$ O-H
1434	44	64	$\nu(s)$ C-H _{Ben} + $\nu(s)$ C-C
1453	4.6	35	$\nu(\rho)$ C-H _{Ben} + $\nu(s)$ C-C-N
1486	40	132	$\nu(s)$ C-H _{Ben} + $\nu(s)$ C-C
1541	16	4	$\nu(\rho)$ C-H _{Ben} + $\nu(as)$ C-C
1590	78	0.25	$\nu(\rho)$ C-H _{Ben} + $\nu(s)$ O-H+ $\nu(s)$ C-C
1671	39	40	$\nu(s)$ C-C=C+ β C-C=C
1698	25	4	$\nu(s)$ C-C=C+ β C-C=C
1721	20	3	$\nu(as)$ C-C=C+ β C-C=C
3176	24	154	$\nu(as)$ C-H _{Ben}
3196	10	74	$\nu(as)$ C-H _{Ben}
3209	10	128	$\nu(s)$ C-H _{Ben} + $\nu(as)$ C-H _{Ben}
3211	5.0	72	$\nu(s)$ C-H _{Ben}
3226	11	150	$\nu(s)$ C-H _{Ben}
3231	12	231	$\nu(s)$ C-H _{Ben}
3897	71	145	ν O-H

Frequencies are given in cm^{-1} , ν =stretching, β =in-plane bending, γ =out-plane bending δ =scissoring, ρ =rocking, w =wagging, s =symmetric, as =asymmetric, τ =twisting, Ben=benzene ring.

O-H vibrations: The FT-IR and FT-Raman modes of hydroxyl group (OH) in 8-hydroxyquinoline have been located at 3418, 1456 and 896 cm^{-1} . **Error! Bookmark not defined.** In the current study, the vibrations of C-H and O-H are mostly observed at 443, 549, 1187 and 1397 cm^{-1} . The vibrational frequency for O-H is located at 3897 cm^{-1} and their assignments can be seen in **Table 4**.

C-H stretching vibration: The C-H vibrations of heteroaromatic organic compounds and their derivatives are found very close to C-H vibrations in benzene ring. They consist of multiple weak bands in range of $3100\text{-}3000 \text{ cm}^{-1}$ due to stretching vibrations of C-H. ²² In current study, the aromatic ring carbon-hydrogen (C-H) stretching frequencies occurred at a range $3231\text{-}3176 \text{ cm}^{-1}$ in FT-IR and FT-Raman.

However, the Raman intensities are found stronger than FT-IR intensities for above mentioned vibrational frequencies.

The assignments of mentioned vibrational frequencies are observed: the aromatic ring carbon-hydrogen (C-H) stretching showed symmetric modes at 3231 and 3226 cm^{-1} . While anti-symmetric stretching modes of C-H are observed at 3196 and 3176 cm^{-1} . However 3209 cm^{-1} is the only frequency mode in this range which showed both symmetric and anti-symmetric stretching of C-H.

In hydroxyquinoline derivatives, C-H vibrations were also studied in range 1247-1087 cm^{-1} . **Error! Bookmark not defined.** In the current study, the C-H stretching modes and their intensities are also observed at low frequencies as shown in **Table 4**. The rocking stretching modes are found to be 1326, 1296 and 1227 cm^{-1} . While, the scissoring

frequencies are ranged in 1162-1091 cm^{-1} . The wagging bands are observed at 1020, 839, 813, 770 and 598 cm^{-1} . The twisting vibrations of C-H are observed at 982, 978, 890, 659 and 478 cm^{-1} as can be seen in **Table 4**.

C-C stretching vibration: The C-C stretching vibrations in the benzene derivatives appear in range of 1650-1400 cm^{-1} .²³ In this study, a similar trend of C-C vibrations is observed at 1721-1486 cm^{-1} and their assignments can be seen in **Table 4**.

C-N bands: The evaluation of C-N bands is difficult task due to the possibility of overlapping with several other vibrations. The C-N stretching modes are observed at 1600, 1450, 1286 and 1273 cm^{-1} as can be seen in **Table 4**. The FT-IR and Raman frequencies were calculated by using method B3LYP/6-31G(d) in gas and solvent phase. The results are tabulated in **Table 5**.

TABLE 5: CALCULATED VIBRATIONAL FREQUENCIES USING B3LYP/6-31G(d) IN GAS AND SOLVENT PHASE

Gas Phase				Solvent Phase ^b			
Frequency Unscaled	Frequency Scaled ^a	I _{IR}	I _{Raman}	Frequency Unscaled	Frequency Scaled ^a	I _{IR}	I _{Raman}
137	132	0.8782	1.5121	134	129	2.7372	3.5805
177	170	6.5759	0.1195	176	169	9.4653	0.2082
267	257	4.1632	0.9354	263	253	11.7116	1.8121
286	275	7.8046	1.3711	288	277	13.0914	2.7918
405	389	88.6766	4.9427	378	363	135.1242	8.4871
440	423	16.9382	2.8406	436	419	6.9541	6.7659
470	453	0.5153	6.8084	469	451	1.032	14.0655
480	462	0.4826	0.3724	480	461	0.0051	0.4244
491	472	0.3334	5.0577	491	472	0.3685	12.7386
546	525	5.4075	4.0793	544	523	8.9243	8.7544
585	562	3.7901	7.3314	584	561	5.451	17.3027
593	570	0.206	2.5227	593	570	0.4699	4.1147
651	626	1.7612	0.8915	650	625	2.4071	1.1882
718	690.	5.8373	30.4289	716	688	13.0399	77.2163
763	733	27.637	3.977	768	738	41.8657	3.9044
805	774	21.912	1.9765	804	773	20.9449	4.3067
819	787	10.7307	1.122	816	785	17.0746	3.3164
831	799	37.1822	0.391	834	801	60.1606	0.752
866	832	1.155	3.2693	878	845	0.3944	3.9965
898	863	4.5371	1.7607	897	863	8.9497	6.9381
960	923	0.3468	0.0223	969	932	0.7135	3.3205
964	927	0.1897	3.4266	974	936	0.2248	1.7089
995	956	0.8516	0.1491	1006	967	1.2269	0.5807
1058	1017	7.2201	6.6479	1059	1018	27.7016	16.1963
1084	1041	63.8909	4.624	1079	1038	104.9651	17.6372
1114	1071	34.8322	9.516	1110	1067	70.8533	30.9298
1169	1124	0.6704	8.1774	1168	1123	1.7045	19.0348
1197	1150.4	17.0417	1.9885	1193	1147	24.0337	1.7467
1230	1182	25.1357	2.9646	1229	1182	39.9485	8.6426
1259	1211	8.191	2.2252	1256	1208	8.4728	10.7766
1297	1247	52.724	4.2669	1293	1243	99.3074	8.396

1321	1270	29.2186	8.3763	1315	1264	46.4538	32.8852
1386	1333	80.9085	23.0063	1378	1324	143.7059	46.1463
1408	1353	55.2562	87.2255	1405	1355	79.1135	403.9155
1449	1393	0.2306	17.621	1447	1391	1.3109	58.2746
1480	1423	42.909	75.8039	1477	1420	67.847	219.1444
1525	1466	11.2768	5.2207	1521	1463	26.6199	13.8429
1566	1505	43.9668	0.2358	1563	1502	95.3223	1.3084
1624	1561	39.7479	40.1334	1620	1558	44.1799	155.3946
1657	1593	15.3705	6.2409	1653	1589	24.6183	20.3393
1674	1609	19.0884	2.198	1671	1607	25.656	4.665
3160	3038	31.4188	150.2075	3166	3042	56.8655	351.9797
3167	3044	20.1376	106.4854	3183	3060	20.4974	225.9812
3183	3060	5.7772	25.7498	3194	3070	2.6982	19.519
3187	3064	11.0221	106.0116	3196	3073	23.3264	309.8756
3206	3082	28.599	172.6484	3212	3088	38.5252	469.8366
3210	3086	22.2084	266.8348	3220	3096	29.8359	529.5724
3741	3596	44.098	154.9709	3729	3585	109.1334	310.4658

Frequency units are given in cm^{-1} , ^ascaling factor= 0.9613, ^bsolvent=DMSO.

According to this data the values of the FT-IR and Raman frequencies are increased in solvent phase than gas phase such as 288,768, 834, 878, 969, 974, 1006, 1059 and 3166-3220 cm^{-1} . While in gas phase frequencies like 137,177, 267, 405, 440, 470, 546, 585, 651,718, 805, 819, 898, 1084-1674 and 3741 are shifted upward as compared to solvent phase. Hence, the straightforward relation is not observed between solvent and gas phase.

However, a significant straightforward relation is observed in intensities of IR and Raman of both solvent and gas phase. The intensities of FT-IR and FT-Raman are remarkably increased in solvent than gas phase. For example the frequencies as 378, 716,1079,1378,1405, 3166, 3220 and 3790 cm^{-1} showed the maximum intensities in solvent phase. In current study, a good agreement is observed in

FT-IR and Raman frequencies computed by both M06-2X/6-31G (d,P) and B3LYP/6-31G (d,p) method. However, these vibrational frequencies are slightly greater than the reported frequencies. **Error! Bookmark not defined. Error! Bookmark not defined.** The solute-solvent interactions can be well explained with the help of free energy difference.²⁴ The solvation free energy is calculated through PCM model²⁵ by pursuing electrostatic, dispersions, repulsions, and cavitations terms. These terms can be shown by following relationship.

$$G_{\text{solvation}} = G_{\text{electrostatic}} + G_{\text{dispersion-repulsion}} + G_{\text{cavitation}}$$

Nevertheless, in SMD (Density-based Solvation Model),²⁶ the process of solvation free energy is completely different than PCM model. This model involves complete solute electron density without considering partial atomic charges.

TABLE 6: DIPOLE MOMENT, FREE ENERGY AND SOLVATION FREE ENERGY USING PCM AND SMD.

	Dipole Moment ^a	Free Energy ^b	Salvation Free Energy ^b
Gas Phase	2.21	-1252456	
PCM model	3.10	-1252484	-28.7
SMD model	3.21	-1252495	-39.5

^aunits in Debye, ^bunits in kJ/mol

In the present study, the PCM and SMD models have been compared, and the calculated dipole moment solvation free energies are summarized in **Table 6**. The dipole moment and free energies are obtained to be greater in solution than the gas phase. The dipole moment and solvation free energies are found to be 3.10 μ and -28.7 kJ/mol in PCM model. While, SMD model showed the dipole moment and solvation free energies are 3.21 μ and -39.5 kJ/mol.

By comparing PCM and SMD models, the solvation free energy and dipole moment is increased in SMD model than PCM model.

UV-Visible analysis: The ultraviolet spectral analysis was calculated by M06-2X/6-31G(d,P) method. The outcomes of ultraviolet spectrum are represented in **Table 7**.

TABLE 7: WAVE LENGTH, EXCITATION ENERGY, AND OSCILLATOR STRENGTH OF 8-HQ

λ (nm)	E (eV)	f	MO contributions
273.97	4.5254	0.0028	H-2 \rightarrow L (90%), H-2 \rightarrow L+2(6%)
269.58	4.5991	0.0763	H-1 \rightarrow L+1(3%), H \rightarrow L(96%)
260.70	4.7559	0.0051	H-1 \rightarrow L (40%), H \rightarrow L+1(57%)
218.09	5.6851	0.0000	H-2 \rightarrow L+1 (96%)
207.53	5.9744	0.7542	H-3 \rightarrow L+1 (5%), H-1 \rightarrow L(51%), H-1 \rightarrow L+2(4%) H L+1 \rightarrow (38%)
191.64	6.4696	0.0416	H-3 \rightarrow L (15%), H-1 \rightarrow L+1 (62%), H \rightarrow L+2 (20%)

MO=molecular orbital, H=Homo, L=LUMO, f = oscillator strength

Table 7 shows the wavelengths as 191.67, 269.58 and 207.59 nm with oscillator strengths i.e 0.0763, 0.0416 and 0.7542 respectively which indicate relatively stronger allowed transitions. While the wavelengths as 260.70 and 269.58 nm with low magnitude of oscillator strength show weak transitions. Frontier molecular orbitals (FMOs). The frontier molecular orbital (FMOs) theory has been recognized as a most outstanding theory in elucidating the chemical stability of species.²⁷ The highest occupied molecular orbital is denoted by

HOMO and lowest unoccupied molecular orbital is denoted by LUMO, which are considered substantial orbitals of frontier molecular orbitals (FMOs). Usually, the HOMO vitality demonstrates the capacity of donating an electron, though, the LUMO vitality speaks to the capacity of picking up an electron.^{28, 29} The frontier orbital energy gap is a valuable parameter in order to get knowledge regarding the dynamic stability and chemical reactivity of species.³⁰

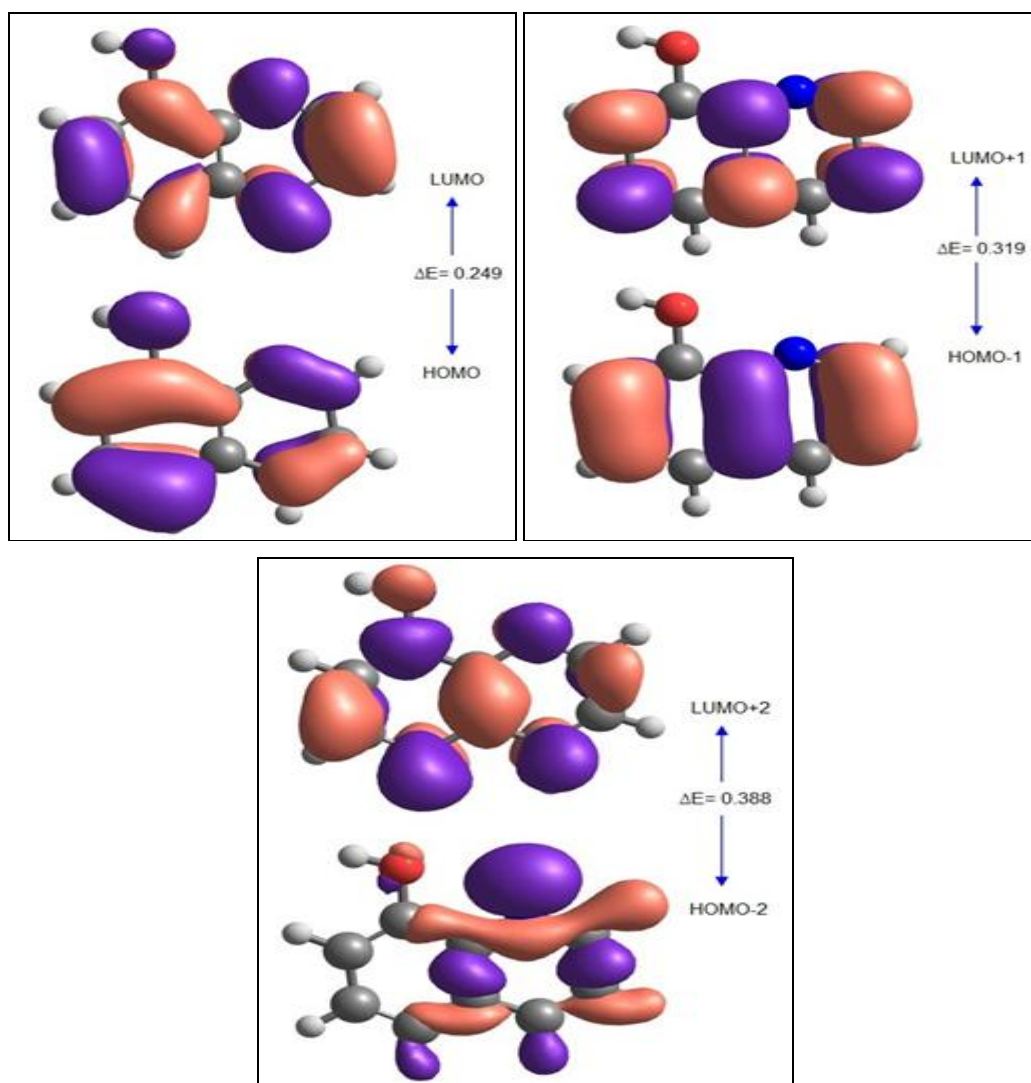


FIG. 2: FRONTIER MOLECULAR ORBITALS OF 8-HQ AND UNITS IN HARTREE (a.u.).

The orbital analysis of 8-HQ has been computed by using M06-2X/6-31G(d,p) method. The calculated energy of HOMO, LUMO, HOMO-1, LUMO+1, HOMO-2, LUMO+2 has been obtained around -0.2596, -0.0106, -0.2999, 0.0189, -0.314 and 0.074 a.u respectively. Subsequently, the energy gap such as $E_{\text{HOMO}}-E_{\text{LUMO}}$, $E_{\text{HOMO-1}}-E_{\text{LUMO+1}}$, $E_{\text{HOMO-2}}-E_{\text{LUMO+2}}$ were obtained 0.249, 0.319 and 0.388 respectively, see **Fig. 2**. The energies of FMOs are helpful to determine the global reactivity descriptors, such as global softness (S), electron affinity (EA), ionization potential (IP)³¹, electro negativity (X)³², global hardness (η), global electrophilicity index (ω) and the chemical potential (μ)³³⁻³⁶ calculated utilizing Equations 2-8.

$$IP = -E_{\text{HOMO}} \quad \text{Equations 2}$$

$$EA = -E_{\text{LUMO}} \quad \text{Equations 3}$$

Where; IP defines Ionization potential (a.u), EA defines electron affinity (a.u).

TABLE 8: IONIZATION POTENTIAL (IP), ELECTRON AFFINITY(EA), ELECTRO NEGATIVITY (X) CHEMICAL POTENTIAL (μ) GLOBAL HARDNESS (η) GLOBAL SOFTNESS (S) AND GLOBAL ELECTROPHILICITY (ω).

	A	B	C
IP	-0.260	-0.299	-0.314
EA	-0.011	0.019	0.074
X	0.135	0.141	0.120
μ	-0.135	-0.141	-0.120
η	0.249	0.319	0.388
S	4.016	3.137	2.577
ω	0.073	0.062	0.037

A= HOMO & LUMO, B= HOMO-1 & LUMO+1, C= HOMO-2 & LUMO+2, units in Hartree (E_h).

The stability, selectivity and reactivity of specie can be described by global reactivity parameters.³⁹⁻⁴² The most important chemical property is electronegativity which described the capacity of specie to attract electrons towards itself. The global softness (S) of this molecule is found to be 4.016 Eh. It is 16 times greater than the magnitude of global hardness (η). These findings suggest that this molecule contain less hardness and larger softness values because of the low HOMO-LUMO energy gap. The softness value reveals that the molecule may be biological activate. This indication has good agreement to reported biological activities of 8-HQ.

Molecular electrostatic potential (MEP): The three dimensional plot of total electron density is

The chemical potential (μ), electro negativity (X) and chemical hardness has been determined by using Koopmans's theorem²⁸ and as following Equations.

$$X = \frac{[IP + EA]}{2} = -\frac{[E_{\text{LUMO}} + E_{\text{HOMO}}]}{2} \quad \text{Equations 4}$$

$$\eta = \frac{[IP - EA]}{2} = -\frac{[E_{\text{LUMO}} - E_{\text{HOMO}}]}{2} \quad \text{Equations 5}$$

$$\mu = \frac{E_{\text{HOMO}} + E_{\text{LUMO}}}{2} \quad \text{Equations 6}$$

The global softness (σ) is defined by Equations 7³⁷

$$\sigma = \frac{1}{2\eta} \quad \text{Equations 7}$$

Parr et al.³⁸ introduced an electrophilicity index (ω) and it can be described Equations 8.

$$\omega = \frac{\mu^2}{2\eta} \quad \text{Equations 8}$$

This parameter was proposed to describe the range of the electrophilic strength for specie

known as molecular electrostatic potential (MEP). MEP surface is developed utilizing the optimized geometry of title compound at the M06-2X/6-31G(d,p) level of theory. MEP can be significant descriptor for explanation of non-covalent interactions (NCIs), attack of electrophilic and nucleophilic at appropriate zone of investigated system.⁴³ The magnitude of electrostatic potential at the MEP map are addressed by standard colors i.e. red color defines the most negative zones, blue displays the most positive and green assume the role of the zero electrostatic potential, therefore, the charge contribution can be ordered i.e. red > orange > yellow > green > blue. Accordingly, the negative (red and yellow) zones on MEP have been associated to the chance of electrophilic attack and the positive zones (blue) to encounter nucleophilic.

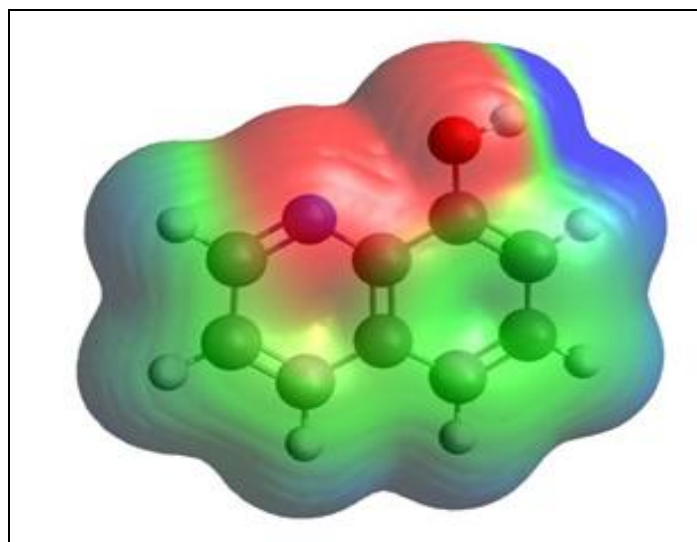


FIG. 3: MOLECULAR ELECTROSTATIC POTENTIAL MAP FOR TITLE COMPOUND

Fig. 3 shows, negative potential which is confined over the oxygen atoms and the N atoms, whereas positive region is confined around the hydrogen atoms.

Thermodynamic parameters: The thermodynamics parameters of 8-HQ were calculated using the M06-2X/6-31G(d,p) and B3LYP/6-31G(d) methods and presented in Table 9.

TABLE 9: THERMODYNAMIC PARAMETERS OF 8-HQ WITH VARIOUS TEMPERATURES

T (K)	S_m^0 (J/mol.K) ^a	$C_{p,m}^0$ (J/mol.K) ^a	H_m^0 (kJ/mol) ^a	S_m^0 (J/mol.K) ^b	$C_{p,m}^0$ (J/mol.K) ^b	H_m^0 (kJ/mol) ^b
25.00	212.50	33.55	0.83	212.59	33.54	0.83
100.00	267.97	54.26	4.01	267.95	54.08	4.00
200.00	318.92	100.11	11.62	318.63	99.47	11.57
298.15	368.27	150.34	23.92	367.67	149.39	23.79
300.00	369.21	151.27	24.19	368.60	150.32	24.07
400.00	419.27	197.76	41.71	418.37	196.72	41.48
500.00	467.65	235.81	63.46	466.52	234.82	63.13
600.00	513.41	265.87	88.60	512.11	264.99	88.18
700.00	556.25	289.72	116.43	554.82	288.93	115.92
800.00	596.24	308.95	146.39	594.71	308.25	145.81
900.00	633.57	324.72	178.10	631.96	324.10	177.45
1000.0	668.48	337.85	211.25	666.81	337.30	210.54

^aCalculated via M06-2X/6-31G(d,p), ^bCalculated via B3LYP/6-31G(d).

The thermodynamics parameters calculated by two methods namely M06-2X and B3LYP with same basis set as 6-31G (d). The magnitude of thermodynamics parameters is appeared to be similar in both methods. The rise in temperature (25-1000K) enhanced the molecular vibration which in turn increases the entropy (S), heat capacity (C) and enthalpy (H).

CONCLUSION: The present study indicates that the optimized bond lengths and bond angles show good correspondence with the reported X-ray and DFT based data. As reported in literature that the C10 contain LP(1) with high stabilization energy as

81.91 kJ/mol. However, the findings of both B3LYP/6-31G(d,p) and M06-2X/6-31G(d,p) methods indicated that there is no lone pair on carbon of the title compound. It is obvious that carbon never contains any lone pair in neutral molecule. The maximum stabilization energy reached up to 39.64kJ/mol which is 2 times less than reported value. The FT- IR and Raman frequencies have increased as compared to reported values in all methods, but good consistency is found from among investigated methods. In solvated 8-HQ, the FT- IR and Raman frequencies are almost insensitive to medium effects.

However, significant medium effect on FT-IR and FT-Raman intensities is observed. The intensities enhanced from gas to solvent phase. The dipole moments are found to be 3.10 and 3.21 and solvation free energies are -28.710 and -39.456 kJ/mol in PCM and SMD models respectively. By comparing both models, the dipole moment and solvation free energy are increased in SMD model than PCM model. FMOs suggest that this molecule contain less hardness and larger softness values. These findings reveal that the molecule might be bioactive.

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