

VIBRATIONAL SPECTRA, NBO, HOMO-LUMO ANALYSES OF FURFURAL SEMICARBAZONE

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ABSTRACT

The molecular structure and vibrational spectra of Furfural Semicarbazone (FSC) have been calculated with the help of density fuctional theory (DFT) using 6-31++G(d,p) and 6-311++G(d,p) basis sets. The solid phase FT-IR and FT-Raman spectra of furfural semicarbazone have been recorded in the range 4000-400cm⁻¹ and 4000-100 cm⁻¹, respectively. On the basis of B3LYP calculations, the normal coordinate analysis has been performed to assign the vibrational fundamental frequencies according to potential energy distribution. The optimum molecular geometry, harmonic vibrational frequencies, infrared intensities and Raman Scattering activities were calculated by density functional theory (DFT/B3LYP) method with 6-31++G(d,p) and 6-311++G(d,p) basis sets. The difference between the observed and scaled wave number values of most of the fundamentals is very small. Stability of the molecule arising from hyper conjugative interactions, charge delocalization have been analyzed using Natural Bond Orbital (NBO) analysis. The electro dipole moment (μ) and first hyperpolarizability (β) values of the investigated molecule are computed using Scaled Quantum Mechanical Force field (SQMF) method. Besides, HOMO and LUMO analysis and Fukui functions of the title molecule have also been calculated using DFT method.

Indexing terms/Keywords

DFT, Natural Bond Orbital (NBO) analysis, HOMO and LUMO, Fukui functions

Academic Discipline And Sub-Disciplines

Applied Physics

SUBJECT CLASSIFICATION

Physics Subject Classification; Spectroscopy

TYPE (METHOD/

Quantum Chemical Calculation – DFT studies

INTRODUCTION

A number of aliphatic and aromatic semicarbazone derivatives have been prepared by the condensation of a variety of ketones with semicarbazide. The products have been characterized by analytical and spectral methods. The semicarbazone derivatives were screened for anti-bacterial activity by diffusion method, using nutrient agar medium and some of the products have shown biological activities against common test organisms. Recently some workers had reviewed the bioactivity Rouse et al (2003) activities. Accordingly and by considering the biological potential of semicarbazones.

Extensive work has been done with the structure analogues of semicarbazone, and may have been found to exhibit interesting biological and chemotherapeutic properties. The spectroscopic studies of semicarbozone and its derivatives have been motivated by their occurrence in natural biological systems Kupchan et al (1974), Georgieva et al (2006) although free molecules do not occur in biological systems. The understanding of the vibrational spectra of free base molecules would be very useful for the understanding of specific biological processes and in the analysis of relatively complex system

SYNTHESIS

1 gram of semicarbazide hydrochloride and 1.5 gram of sodium acetate were dissolved in 10 ml of water. 0.5 gram of the furfural was added to the above mixture. The mixture was stirred for a few minutes and allowed to stand. Furfural semicarbazone crystallized out from solution on standing. The crystals were filtered, washed with cold water and recrystallised from water. The scheme of the synthesis is shown in Figure 1.





Figure 1 The scheme of the synthesis of FSC

EXPERIMENTAL DETAILS

The fourier transform infrared spectra of the title compound was recorded in the region 4000-400 cm⁻¹, using Bracker IFS 66V model FT-IR Spectrometer equipped with an MCT detector, a KBR beam splitter and globar source. The spectral resolution is \pm cm⁻¹.

The FT-Raman Spectrum of FSC was recorded on a computer interfaced BRUCKER IFS 66V model interferometer equipped with FRA-106 FT-Raman accessories. The spectrum was measured in the region 4000-100 cm⁻¹ using Nd:YAG laser operating at 200 mW power continuously with 1064 nm excitation. The spectra were recorded with scanning speed of 30 cm⁻¹ min⁻¹ of spectral width 2 cm⁻¹. The frequencies of all sharp bands are accurate to ± 1 cm⁻¹.

COMPUTATIONAL DETAILS

In the frame work of DFT approach, different exchange and correlation functions were routinely used. Among these, the B3LYP combination (Becke 1993) is the most used since it proved its ability in reproducing various molecular properities, including vibrational spectra. In order to provide information with regard to the structural characteristics and the normal vibrational modes of FSC, the DFT-B3LYP correlation functional calculations have been carried out. The molecular geometry optimizations, energy and vibrational frequency calculations were carried out for FSC with the GAUSSIAN 03W software package. The DFT level calculations, employing the B3LYP keyword, which invokes Becke's three parameter hybrid method Pulay et al (1983), Fogarasi & Pulay (1985), (Sundius 2002) has been computed using the correlation function of Lee et al (1998), implemented with 6-31++G(d,p) and 6-311++G(d,p) basis sets. All the parameters were allowed to relax and all the calculations converged to an optimized geometry which corresponds to a true minimum, as revealed by the lack of imaginary values in the wavenumber calculations. The Cartesian representation of the theoretical force constants have been computed at optimized geometry by assuming C1 point group symmetry. Scaling of the force field was performed according to SQMF procedure Pulay et al (1983), Fogarasi & Pulay (1985) using selective scaling in the natural internal coordinate representation Fogarasi & Pulay (1985). Normal coordinate analysis has been performed to obtain full description of the molecular motion pertaining to the normal modes with MOLVIB program version 7.0 written by Sandius (2002). For plots of simulated IR spectra, pure Lorentizian band shapes were analysed with a band width (FWHM) of 10 cm⁻¹. The Raman intensities (δ_i) calculated by Gaussian 03W program have been suitably adjusted by the scaling procedure with MOLVIB and subsequently converted to relative Raman intensities (Ii) using the following relationship derived from the basis theory of Raman scattering.

 $I_{i} = f (v_{o}-v_{i})^{4\delta i} / v_{i} (1-exp (hcv_{i}/k_{b}T))$

(1)

Where is the exciting frequency in cm⁻¹, v_i the vibration wave number of the ith normal mode, h, c and k_b are the fundamental constants and f is a suitably chosen common normalization factor for the entire peak intensities Olszak et al (1995). The simulated IR and Raman spectra have been plotted using Lorentizian shapes with Full Width at Half Maximum (FWHM) of 10 cm⁻¹. Natural Bond Orbital (NBO) analysis Molvib (v.7.0), Keresztery et al (2002), Glendering et al (1998) has been performed using NBO 3.1 program as implemented in the Gaussian 03W package at the DFT/B3LYP/6-31++G(d,p) level of theory.

RESULT AND DISCUSSION

Molecular Geometry

The optimized molecular structure with the numbering of atoms of the title compound is shown in Figure 1.2. The most optimized structural parameters are also calculated by B3LYP method with 6-31++G(d,p) and 6-311++G(d,p) as basis sets and they were presented in Table 1 they were presented in Table 1. The calculated values were compared with X-ray diffraction results. From the structural data given in Table 1. Table compares with those of experimental data Klots et al (1994), one can find that most of the optimized bond lengths are larger than experimental values. This is due to the fact that the theoretical calculations result from isolated molecules in gaseous phase while the experimental results are from molecules in solid state. Comparing bond angles and bond lengths of B3LYP/ 6-31++G (d,p) and 6-311++G (d,p) basis set for FSC, it is also observed that the geometrical parameters were found to be almost same at both levelsThe calculated C-C bond length of the Furan ring vary from 1.36 to 1.42 Å calculated by B3LYP/6-311++G (d, p) method. The C(1)-C(2) bond length is 1.442 Å. This is larger than the C-C bond distance of the furan ring. This clearly shows that there is no delocalization of the π - electrons of N=C towards the ring of FCS. The calculated furan ring bond angle by B3LYP/6-31++G (d, p) and 6-311++G(d,p) basis sets lies in the range 106.55° to 110.64°. The bond angle of C(1)-C (2)-O(6) and C(3)-C(2)-O(6) are equal to 119.59° and 109.61°, respectively. This variation of the bond angle depends on the electro negativity of the central atom. If the electro negativity of the central atom increases, bond angle increases.





Figure 2 Molecular Structure of FSC with numbering of atoms

Detailed description of vibrational modes can be given by means of normal coordinate's analysis. For this purpose the full set of 48 standard internal coordinates were analysed. For the title compound are defined in Table 2. From these, a non-redundant set of local symmetry coordinates were constructed by suitable linear combinations of internal coordinates following the recommendations of Fogarasi & Pulay (1985) which are summarized in Table 3

Parameters	B3LYP/ 6-31++G(d p)	B3LYP/ 6-311G++(d p)	Experimental								
	BOND I FI	NGTH									
C(1) C(2)	1 4417	1 4402	1 462								
C(1)-C(2)	1.4417	1.4402	1.462								
C(1)-N(7)	1.2882	1.2836	1.2599								
C(1)-H(12)	1.0958	1.0942	1.080								
C(2)-C(3)	1.3754	1.3714	1.369								
C(2)-O(6)	1.3682	1.3657	1.373								
C(3)-C(4)	1.4292	1.4279	1.4259								
C(3)-H(13)	1.0809	1.0787	1.081								
C(4)-C(5)	1.3648	1.3605	1.368								
C(4)-H(14)	1.0804	1.0781	1.1								
C(5)-O(6)	1.3629	1.3606	1.353								
C(5)-H(15)	1.0788	1.0769	1.1								
N(7)-N(8)	1.3523	1.3497	1.352								
N(8)-C(9)	1.3957	1.3971	1.215								
N(8)-H(16)	1.0172	1.0159	1.012								
C(9)-O(10)	1.2276	1.2194	1.208								
C(9)-N(11)	1.36	1.3588	1.309								
N(11)-H(17)	1.0062	1.0047	1.0119								
N(11)-H(18)	1.0087	1.0074	1.012								
	BOND ANGLE										
C(2)-C(1)-N(7)	122.50	120.55	119.99								
C(2)-C(1)-H(12)	115.51	115.43	119.99								
N(7)-C(1)-H(12)	121.98	122.01	125.91								
C(1)-C(2)-C(3)	130.80	130.70	128.49								

Table 1	Optimized	geometrical	parameters	for FSC	with	different	basis	sets



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C(1)-C(2)-O(6)	119.59	119.64	111.04
C(3)-C(2)-O(6)	109.61	109.64	105.85
C(2)-C(3)-C(4)	106.55	106.52	109.97
C(2)-C(3)-H(13)	126.04	126.06	126.58
C(4)-C(3)-H(13)	127.41	127.42	127.561
C(3)-C(4)-C(5)	106.03	106.03	106.33
C(3)-C(4)-H(14)	127.49	127.52	133.39
C(5)-C(4)-H(14)	126.49	126.44	126.02
C(4)-C(5)-O(6)	110.64	110.66	110.93
C(4)-C(5)-H(15)	133.62	133.54	133.39
O(6)-C(5)-H(15)	115.74	115.80	116.87
C(2)-O(6)-C(5)	107.18	107.15	106.90
C(1)-N(7)-N(8)	118.21	118.42	119.09
N(7)-N(8)-C(9)	121.97	122.07	123.03
N(7)-N(8)-H(16)	122.41	122.31	124.05
C(9)-N(8)-H(16)	115.62	115.62	115.66
N(8)-C(9)-O(10)	119.90	120.02	120.00
N(8)-C(9)-N(11)	114.86	114.51	114.56
O(10)-C(9)-N(1)	125.23	125.48	125.78
C(9)-N(11)-N(17)	117.68	117.80	117.99
C(9)-N(11)-H(18)	120.93	120.77	120.23
H(17)-N(11)-H(18)	121.39	121.43	120.53

Table 2 Definition of internal coordinates of FSC

No.	Туре	Definition								
	Stretching (r _i)									
1-3	0-C	C2-O6, C5-O5, C9-C10								
4-7	C-C	C1-C1, C2-C3, C3-C4, C4-C5								
8-10	C-N	C9-N8, C1-N7, C9-N11								
11-14	C-H	C3-H13, C4-H14, C5-H15, C1-H12								
15-17	N-H	N8-H16								
	NH ₂	NH ₂ (sym), NH ₂ (asym)								
		In-Plane Bending(β _i)								
18-22	Ring	O6-C2-C3, C2-C3-C4, C3-C4-C5, C4-C5-O6, C5-O6-C2								
23	C-C-C	C3-C2-C1								
24	O-C-C	C3-C2-O6								
25	C-C-N	C2-C1-N7								
26-28	C-N-N	C1-N7-N8, C9-N8-N7, N8-C9-N11								
29-30	N-C-O	N8-C9-O10, N11-C9-O10								
31	N-N-H	N7-N8-H16								
32-33	C-N-H	C9-N8-H16, C1-N7-H12								



34-39	С-С-Н	C1-C2-H12, C2-C3-H13, C4-C3-H13, C3-C4-H14, C4-C5-H14, C4-C5-H15
40	О-С-Н	O6-C5-H15
41	N-H-H	N11-H17-N18
	I	Out-of- Plane Bending (v _i)
42	C-C	C1-C2-C3-O6
43-46	C-H	C4-C5-O6-H15, C3-C4-C5-H14, C1-C2-C3-H13, C1-C2-C3-H12
47-48	N-H	C1-N7-N8-H16, N7-N8-C9-H16
49	N-O	N8-C9-N11-O10
50	C-N	C9-N11-H17-H18
	I	Torsion(T)
51-55	T _{ring}	O6-C2-C3, C2-C3-C4, C3-C4-C5, C4-C5-C6, C5-O6-C2
56	TC-NH2	C9-N11-H17-H18

Table 3 Definition of local symmetry coordinates in B3LYP/6-31++G (d,p) for FSC

S.No	Symmetry Coordinates	Description
1	S ₁ = r ₁₂	vC ₁ C ₂
2.	$S_2 = r_{17}$	v C ₁ N ₇
3.	$S_3 = r_{112}$	vC ₁ H ₁₂
4.	$S_4 = r_{23}$	vC ₂ C ₃
5.	$S_5 = r_{26}$	vC ₂ O ₆
6.	$S_6 = r_{34}$	vC ₃ C ₄
7.	S ₇ = r ₃₁₃	vC ₃ H ₁₃
8.	$S_8 = r_{45}$	v C ₄ C ₅
9.	$S_9 = r_{56}$	v C ₅ O ₆
10.	$S_{10} = r_{515}$	v C ₅ H ₁₅
11.	$S_{11} = r_{78}$	v N ₇ N ₈
12.	$S_{12} = r_{89}$	v N ₈ C ₉
13.	$S_{13} = r_{816}$	v N ₈ H ₁₆
14.	$S_{14} = r_{910}$	v C ₉ O ₁₀
15.	$S_{15} = r_{911}$	v C ₉ N ₁₁
16.	$S_{16} = r_{1117}$	v N ₁₁ H ₁₇
17.	$S_{17} = r_{1118}$	v N ₁₁ H ₁₈
18.	$S_{18} = \beta_{2313} - \beta_{4313}$	βC ₃ H ₁₃
19.	$S_{19} = \beta_{3414} \beta_{5414}$	$\beta C_4 H_{14}$
20.	$S_{20} = \beta_{4515} + \beta_{6515}$	$\beta C_4 H_{15}$
21.	$S_{21} = \beta_{7816} \cdot \beta_{9816}$	$\beta N_8 H_{16}$
22.	$S_{22} = \beta_{8911} \cdot \beta_{10911}$	βC ₉ N ₁₁
23.	S ₂₃ = β ₉₁₁₁₈ - β ₁₇₁₁₁₈	βN ₁₁ H ₁₈
24.	$S_{24} = \beta_{2112} \beta_{7112}$	βC1H12
25.	S25 = β 18 11 9 - β 7 11 9	βN11C9
26.	S26 = β 2 6 5 + a(β 6 5 4 + β 3 2 6)+ b(β 2 3 4 + β 4 5 6)	βring



27.	S27 = u1 2 6 3	u C1C2
28.	S28 = u13 3 2 4	u C3H13
29.	S29 = u14 4 5 3	u C4H14
30.	S30 = u15 5 4 6	u C5H15
31.	S31 = u12 1 7 2	u C1H12
32.	S32 = u10 8 9 7	u N8H16
33.	S33 = u9 11 17 18	u C9N11
34.	S34 = b(u5 6 2 3+ u2 3 4 5)+a(u6 5 4 3+ u4 3 2 6)+ u3 2 6 5	u ring
35.	S35 = T 13 3 4 14	т Н13С3
36.	S36 = T 14 4 5 15	т Н14С4
37.	S37 = T 12 1 2 3	т Н12С1
38.	S38 = T 16 8 7 1	т Н16N8
39.	S39 = T 10 9 8 16	тО10С9
40.	S40 = T 11 17 9 10	тN11H17
41.	S41 = T 1 2 3 13	т С1С2
42.	S42 = T 7 1 2 6	т N7С1
43.	S43 = T 8 9 11 18	т N8С9
44.	S44 = T 2 6 5 15	т С2О6
45.	S45 = T 5 4 3 13	т С5С4
46.	S46 = T 2 3 4 14	т С2С3
47.	S47 = т 7 8 9 10	т N7N8
48.	S48 = T 14 4 5 6 + T 13 3 4 14 + T 15 5 6 2	т СН

Vibrational Assignments

The molecule FSC belongs to C_1 point group symmetry and its 48 fundamental modes of vibrations. All these modes are found to be active both in the Raman scattering and infrared absorption.





Figure 3 Comparison of experimental and computed FT-IR spectra of FSC

Figure 4 Comparison of experimental and computed FT-Raman spectra of FSC

The detailed vibrational assignments of fundamental modes of FSC along with the calculated IR and Raman frequencies and normal mode descriptions are reported in Table 4.



	Experimental		Theoretical Values cm ⁻¹		B3LYP/6-31++G(d,p)		B3LYP/6-31		
Mod e N.	FT-IR	FT- RAMAN	B3LYP/ 6-31++G(d,p)	B3LYP/ 6-311+G(d,p)	^a IR Intensity	[⊳] Raman Intensity	^a IR Intensity	[⊳] Raman Intensity	Assignments
1	3456(s)	-	3476	3459	15.08	17.06	14.77	20.47	vNH ₂ (asy)(98)
2	3342(vs)	3366(s)	3351	3352	5.34	42.82	5.08	52.28	vNH ₂ (sym)(99)
3	3280(m)	3171(ms)	3295	3281	1.91	82.82	1.95	100.00	vNH(99)
4	3095(s)	3077(w)	3070	3048	0.01	51.65	0.02	63.29	vCH(96)
5	2998(w)	-	3045	3024	0.05	37.36	0.02	42.07	vCH(92)
6	2996(s)	-	3033	3013	0.65	36.07	0.61	43.52	vCH(98)
7	2977(ms)	2962(ms)	2959	2938	4.74	26.81	4.44	32.94	vCH(99)
8	1857(vs)	1843(ms)	1814	1809	100.00	41.07	100.00	49.16	vOC(69), vNC(10)
9	1689(vs)	-	1624	1611	2.02	68.68	2.14	88.87	vN=C(57), vCC(14),βHCN(12)
10	1658(m)	1628(vw)	1646	1636	5.76	0.46	6.42	0.65	vN=C(15), vCC(18),vCC(3 2)
11	1601(vw)	1611(w)	1584	1588	45.49	11.33	39.44	14.84	βNH ₂ (85)
12	1519(ms)	1557(w)	1473	1464	8.24	57.32	7.39	57.98	vCC(56), βHNN(13), βHCO(10)
13	1473(vs)	1473(ms)	1421	1415	24.92	100.00	18.00	130.66	vCC(40), βHNN(37)
14	1456(vs)	-	1376	1365	1.39	68.98	2.01	85.89	βCOO(10), βHCC(22), βHCO(14), βCCO(21)
15	1418(vw)	1392(s)	1349	1340	47.13	1.97	50.74	1.30	vCC(32), vNC(16),βHNC(14), βOCN(13)
16	1332(vs)	1332(ms)	1299	1294	4.09	17.96	5.78	27.26	vCN(58)
17	1278(vs)	1279(s)	1255	1243	0.49	22.77	0.44	25.22	vCC(20), vOC(28),βCCO (11)
18	1220(s)	1223(s)	1202	1194	3.47	6.62	3.27	6.71	βHCC(41), βHCO(18)
19	1156(ms)	1154(ms)	1143	1134	6.59	7.77	12.13	16.59	βHCO(32),vOC (17), vNN(20),
20	1139(w)	1139(w)	1129	1120	24.89	22.28	19.19	23.04	vOC(16), vNN(39)
21	1120(s)	1118(w)	1065	1058	0.52	4.54	0.47	5.39	vCC(10), vOC(38),βCCO (16)
22	1098(vw)	1073(s)	1044	1044	6.07	1.73	8.37	1.51	δNH ₂ (73),βHC N(34)
23	1010(vs)	1026(w)	1003	996	3.94	16.05	3.84	21.65	βHCC(31), βCOC(16)

Table 4 Experimental and B3LYP levels computed vibrational assignments of FSC



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24	998(ms)	1012(vs)	957	951	1.94	2.34	2.01	2.66	γHCC(38),vNC(24),βNCC(19)
25	947(w)	-	914	911	3.38	3.87	3.51	5.47	γHCC(34),vCC(12), βCCO(11)
26	930(s)	931(w)	895	896	2.16	2.46	1.93	2.90	γHCN(46)
27	-	908(vw)	863	867	1.85	3.70	1.66	4.53	βCCO(61), βCOC(21)
28	883(ms)	884(ms)	847	847	0.00	0.72	0.00	0.93	тНССС(22), тНССО(17), тНСОС(15)
29	812(vw)	-	783	783	1.78	0.56	1.66	0.43	THCOC(19)
30	803(ms)	789(w)	765	765	1.01	3.18	1.06	4.38	βNCC(38), βCCO(15)
31	751(s)	-	722	724	9.97	1.03	1.70	0.07	ωNH ₂ (66)
32	739(vs)	724(vw)	715	720	1.33	0.03	9.17	0.89	THCCO(34)
33	656(ms)	-	654	655	0.04	0.40	0.05	0.53	τCCOC(26), ωCCOC(31)
34	644(s)	623(w)	611	611	0.01	0.78	0.01	0.93	vNC(50), βNCC(40), βNCN(23)
35	595(vs)	598(vw)	578	579	1.23	0.11	1.25	0.11	тНСОС(11),тС СОС(15)
36	575(vw)	573(w)	563	560	0.74	0.11	0.48	0.10	THNCN(38)
37	531(ms)	526(vw)	529	532	5.05	1.16	4.90	1.41	βOCN(65)
38	461(s)	412(ms)	449	444	13.66	0.23	13.70	0.33	THNNC(27)
39	418(vw)	403(vs)	423	424	4.44	1.22	4.25	1.53	τNH₂(48),βCCO (25), βNNC(11),
40	347(vs)	372(vw)	352	350	0.56	0.32	0.60	0.50	TNNCC(27),τC COC(16),ωCC OC(12)
41	321(w)	-	305	304	0.25	0.99	0.27	1.20	vCC(20), vNC(13),βNNC(22)
42	-	269(ms)	270	272	10.07	0.55	9.52	0.64	TNCCC(17),τN CNN(27),τCNN C(13),ωCCOC(13)
43	202(w)	201(ms)	214	215	1.88	0.70	1.82	0.85	βNCN(16), βCCO(34), βCNN(30),
44	-	152(ms)	129	125	0.01	0.14	0.07	0.17	TNCCC(24), TCNNC(21), ωCCOC(24)
45	111(w)	109(w)	99	93	15.40	0.07	11.72	0.07	тHNCN(25), тHCCC(18), тNNCC(15)
46	92(m)	90(w)	75	74	0.10	0.64	0.09	0.78	βNCC(35), βCCO(18).
L									//



									βNNC(24)
47	56(w)	-	52	50	0.04	0.50	0.01	0.65	тNCCC(38), тCNNC(21)
48	-	-	37	41	6.54	0.16	8.63	0.28	тHNCN(25), тNCNN(12)

Abbreviations: w-weak, vw- very weak, m- medium, s- strong, vs- very strong, u- stretching, u_{as} - asymmetric stretching, u_{s} - symmetric stretching, β - in-plane bending, γ - out-of plane bending, ω -wagging and τ - torsion.

NH₂/NH vibrations

The NH₂ group gives rise to six internal modes, namely symmetric stretching-v_s(NH₂), anti-symmetric stretching- $v_{as}(NH_2)$, scissoring (or) symmetric deformation (or) simply deformation- $\beta(NH_2)$, anti-symmetric deformation (or) rocking $\delta(NH_2)$, wagging- $\omega(NH_2)$, and torsion or twist- τ (NH₂) modes. For FSC the v_s, v_{as}, τ and ω modes are usually localized and are pure group modes, whereas β and $\overline{\delta}$ modes mix up with other ring modes Arivazhagan et al (2010). The calculated NH₂ asymmetric stretching vibrations give rise to the strong band at 3463, 3446 cm⁻¹ compared to the experimental result at 3456 cm⁻¹ Bambagiooti et al (1974). The theoretically observed NH₂ symmetric vibration give rise to the strong band at 3312,302 cm⁻¹ compared to the experimental values at 3342 and 3366 cm⁻¹ in FT-IR and FT Raman spectrum of FSC. The N-H groups show their stretching vibration in the region 3500-3220 cm⁻¹. The position of the absorption in this region depends upon the degree of hydrogen bonding and, upon the physical state of the sample (or) the polarity of the solvent. In the present work, the bands appear at 3280 cm⁻¹ in the FT-IR and 3271 cm⁻¹ in the FT-Raman. The theoretically observed NH symmetric stretching vibration gives rise to 3233 and 3222 cm⁻¹. The scissoring mode (β) of the NH₂ groups gives rise to its characteristic frequencies in the region 1600-1700 cm⁻¹ which contains a broad and strong IR band with peak at 1601 cm⁻¹ in FT-IR and 1611 cm⁻¹ in FT Raman for FSC. The rocking δ (NH₂) mode usually appears in the region 900-1150 cm⁻¹ for nucleic acid bases. In the present case the IR band at 1098 cm⁻¹ is associated with this mode. In the assignment the $\omega(NH_2)$ and $\tau(NH_2)$ modes have been assigned at 751 cm⁻¹ and 418 cm⁻¹.

C-H vibration

The heteroaromatic organic compounds and its derivatives are structurally very close to benzene and commonly exhibit multiple weak bands in the region 3100-3000 cm⁻¹ Zinczuk et al (2009) due to C-H stretching vibration Rastogi et al (2002), Arivazhagan et al (2010). In our study, the FT-IR vibrational frequencies found at 3095, 2998, 2996 and 2977 cm⁻¹ are assigned to C-H stretching vibration of FSC. The theoretically computed C-H vibrations by B3LYP/6-31++G(d,p) and 6-311++G(d,p) methods show good agreement with recorded spectrum as well as the literature data. The C-H in-plane and out-of-plane bending vibrations generally lie in the range1300-1000 cm⁻¹ and 1000-750 cm⁻¹, respectively. The FT-IR band at 1220, 1156, 1098 and 1010 cm⁻¹ and FT-Raman band 1223, 1154, 1073 and 1026 cm⁻¹ are assigned to C-H in-plane bending vibration of FSC. The peaks at 998,947 and 930 cm⁻¹ in FT-IR and 931 and 908 cm⁻¹ confirm the C-H out- of- plane bending vibration which agrees with the above said literature values. In general, the C-H vibrations (stretching, in plane and out of plane bending vibrations) calculated theoretically are in good agreement with experimental values Klots et al (1994), Zhang et al (2011).

C-N vibrations

The identification of C-N vibrations is a very difficult tasks, since mixing of several bands are possible in this region. Silverstein &Webster (1981) assigned C-N stretching absorption in this region 1382-1266 cm⁻¹, for aromatic amines. In this study the bands observed at 1332 cm⁻¹ in IR have been assigned to C-N stretching vibration of FSC. The slight shift in wave number is due to the fact that force constants of the C-N bond increases due to resonance with the ring. Pyrimidine absorbs strongly in the region 1600-1500 cm⁻¹ due to the C=C and C=N ring stretching vibrations (Dani 1995), Arivazhagan et al (2010). In FSC, the band observed at 1689 cm⁻¹ in IR spectrum is assigned to C=N stretching vibration and the corresponding force constants contribute 87% to the Total Energy Distribution (TED). The C-N in- plane and out-of-plane bending vibrations have also been identified and are listed in Table 3.

Furan ring vibrations

Generally the C-C stretching vibrations in aromatic compounds from the band in the region of 1430-1650 cm⁻¹ Zinczuk et al (2009). According to (Socrates 2001), the presence of conjugate substituent such as C=C causes a heavy doublet formation around the region 1625-1575 cm⁻¹. The six ring carbon atoms undergo coupled vibrations, called skeletal vibrations and give a maximum of four bands in the region 1660-1420 cm⁻¹ (Kalsi 1993). Five membered ring hetero aromatic compounds with two double bond in the ring, generally shows three ring stretching bands near 1590,1490 and 1400 cm⁻¹ Klots et al (1994), Zhang et al (2011). As predicted in the earlier references the prominent peaks at 1519/1557, 1473/1473, and 1418/1392 cm⁻¹ in the FT-IR/ Raman spectra are assigned to C=C and C-C stretching vibrations for the title compound and these modes are supported by their TED values. Most of the ring vibrational modes are affected by the substitutions in the aromatic ring of the title compound. In the present study, the mode with major contribution from furan ring deformation is attributed at 883 and 812 cm⁻¹ in Infrared spectrum and corresponds well with the calculated wave number. The peak observed at 526 cm⁻¹ in Raman and 531 cm⁻¹ in IR Spectrum are assigned to the furan ring torsional modes by careful consideration of their quantitative descriptions. The reductions in the frequencies of



these modes are due to the change in force constants and the vibrations of the functional groups present in the molecule. The theoretically computed values by B3LYP/6-311++G(d,p) method gives excellent agreement with experimental data.

Natural Bond Analysis (NBO)

NBO analysis is an efficient method for studying intra and intermolecular bonding and interaction among bond and also provides a convenient basis for investigating charge transfer or conjugate interactions in molecular systems. It was performed by using NBO 3.1 program implemented in the Gaussian 03W package at the DFT/B3LYP level in order to understand various second order interactions between the filled orbitals of one subsystem and vacant orbitals of another subsystem Read et al (1988), Choo et al (2002), which is a measure of the delocalization or hyperconjugation. NBO method gives useful information about the interactions in both filled and virtual orbital spaces. In this analysis large E ⁽²⁾ value shows the intensive interaction between electron donors and electron acceptors a greater extent of conjugation of the whole system. For each donor (i) and acceptor NBO (j) the stabilization energy E ⁽²⁾ associated with an electron delocalization between donor and acceptor is estimated as Luque et al (2000), Okulik & Jubert (2005):

$$E^{(2)} = \Delta E_{ij} = q_i \frac{F(i,j)^2}{\varepsilon_j - \varepsilon_i}$$

(2)

qⁱ = orbital occupancy

 $\varepsilon_j - \varepsilon_i$ = diagonal elements

F(i,j) = off diagonal NBO Fock matrix element

Donor(i)	Туре	ED(e)	Acceptor(j)	Туре	ED(e)	E ^{(2)a} (kJ/mol)	E(j)– E(i) ^b (a.u.)	F(i, j) ^c (a.u.)	
			C ₁ -N ₇	σ*	0.0107	4.08	1.14	0.068	
			C ₁ -H ₁₂	σ*	0.0368	0.75	1.12	0.026	
			C ₂ -C ₃	σ*	0.0277	7.18	1.39	0.089	
C ₁ -C ₂	σ	1.9575	C ₂ -O ₆	σ*	0.0326	3.28	1.29	0.058	
			C ₃ -C ₄	σ*	0.0124	1.06	1.39	0.034	
			C ₅ -O ₆	σ*	0.0154	2.97	1.29	0.055	
			N ₇ -N ₈	σ*	0.0238	4.88	1.15	0.067	
C ₁ -N ₇			C ₁ -C ₂	σ*	0.0346	3.32	1.55	0.064	
	G	1 9861	C ₁ -H ₁₂	σ*	0.0368	0.82	1.29	0.029	
	0	1.9001	C ₂ -C ₃	σ*	0.0277	2.23	1.55	0.053	
			N ₈ -C ₉	σ*	0.0717	1.83	1.38	0.045	
C ₄ -N-	π	π	1 0282	C ₁ -N ₇	π*	0.02763	0.50	0.36	0.013
		1.3202	C ₂ -C ₃	π*	0.3697	13.37	0.37	0.069	
			C ₁ -C ₂	σ*	0.0346	8.15	1.36	0.094	
			C ₁ -N ₇	σ*	0.0107	1.85	1.39	0.045	
	G	1 9697	C ₂ -O ₆	σ*	0.0326	1.80	1.27	0.043	
02-03	Ŭ	1.5057	C ₃ -C ₄	σ*	0.0124	3.20	1.37	0.059	
			C ₃ -H ₁₃	σ*	0.0210	1.36	1.14	0.035	
			C ₄ -H ₁₄	σ*	0.0126	1.69	1.13	0.065	
C ₉ -O ₁₀	σ	1.9910	N ₇ -N ₈	σ*	0.0238	1.89	1.45	0.047	
			N ₈ -C ₉	σ*	0.0717	1.17	1.50	0.038	
N ₁₁ -H ₁₇	σ	1.9897	N ₈ -C ₉	σ*	0.0717	4.33	1.11	0.063	
			C ₉ -O ₁₀	σ*	0.0112	0.54	1.32	0.024	
O ₆	LP ₁	1.9495	C ₁ -C ₂	σ*	0.0346	1.94	1.20	0.043	
			C ₂ -C ₃	σ*	0.0277	5.74	1.21	0.075	

Fable 5 Second orde	r perturbation theo	ry analysis of Fock	matrix in NBO	basis of FSC
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			C4-C5	σ*	0.0209	6.66	1.12	0.078
			C ₂	RY [*] (1)	0.00818	3.01	2.10	0.072
			C ₂	RY [*] (2)	0.00586	1.65	2.84	0.062
			C ₅	RY (1)	0.00644	4.81	1.76	0.083
O ₆	LP ₂	1.6370	C ₂ -C ₃	π*	0.3967	43.07	0.41	0.119
			C ₂	RY [*] (3)	0.00379	4.56	1.49	0.081
			C ₁ -C ₂	σ*	0.03466	1.94	1.20	0.043
			C ₄ -C ₅	π*	0.31739	46.65	0.38	0.120
N ₇	LP ₁	1.9135	C ₁ -H ₁₂	σ*	0.0368	10.17	0.75	0.079
			N ₈ -C ₉	σ*	0.0717	1.01	0.84	0.026
			C ₁	RY [*] (1)	0.00689	5.14	1.47	0.079
			C ₁ -C ₂	σ*	0.03466	3.24	1.01	0.052
			N ₈ -H ₁₆	σ*	0.03387	8.68	0.75	0.073
N ₈	LP ₁	1.70147	C ₁ -N ₇	π*	0.27638	31.59	0.28	0.085
			C ₉ -O ₁₀	π*	0.37858	52.61	0.30	0.115
O ₁₀	LP ₁	1.98006	N ₈ -C ₉	σ*	0.07177	1.29	1.12	0.034
			C ₉ -N ₁₁	σ*	0.06431	1.47	1.12	0.037
O ₁₀	LP ₂	1.8563	N ₇ -N ₈	σ*	0.0238	0.90	0.63	0.022
			N ₈ -C ₉	σ*	0.0717	24.22	0.68	0.117
			C ₉ -N ₁₁	σ*	0.0643	23.54	0.69	0.116
			N ₁₁ -H ₁₈	σ*	0.0116	0.64	0.66	0.019

ED Electron Density

^a E⁽²⁾ means energy of hyper conjugative interaction (stabilization energy).

^b Energy 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.

The results of second order perturbation theory analysis of Fock matrix at B3LYP/6-31++G(d,p) level of theory are collected in Table 5. The strong intramolecular hyperconjugative interaction of the σ and π electrons of the C-C, C-O, C-H and C-N to the anti C-C, C-O, C-H and N-H bond of the furan ring as well as the semicarbazone group leads to stabilization of some part of the ring. The intermolecular hyper conjugation interaction of the $\sigma(C_1-C_2)$, distributed to $\sigma^*(C_2-O_6)$ (C₁-N₇), (C₂-C₃) leads to stabilization of approximately 3.3. 4.1 and 7.2 kJ/mol. furthermore, $\pi(C_1-N_7)$ NBO conjugates with the anti-bonding orbitals of $\pi^*(C_2-C_3)$ which lead to strong delocalization of 13.37 kJ/mol. The LP₂(O₆) of NBO conjugated with the $\pi^*(C_2-C_3)$ results in a high stabilization of 43.07 kJ/mol. These interactions are observed as increase in Electron Density (ED) in C-C, C-N, anti-bonding orbital that weakens the respective bonds. The energies for the interaction LP₁(O₆) $\rightarrow \sigma^*(C_4-C_5)$ and LP₁(N₇) $\rightarrow \sigma^*(C_1-H_{12})$ are 6.66 and 10.17 kJ/mol, respectively, which clearly demonstrate that the intermolecular hyper conjugative interaction between the semi carbazone group is strong in the ground state. There occurs a strong intermolecular hyperconjugative interaction of (C₂-C₃) from O₆ (LP₂ (O₆) $\rightarrow \pi^*(C_2-C_3)$ which increases ED (0.3967 eV) that weakens the respective bonds (C₂-C₃) leading to stabilization of 43.07 kJ/mol. These charge transfer interactions of FSC are responsible for pharmaceutical and biological properties. Hence, the FSC structure is stabilized by these orbital interactions.

Frontier Molecular Orbital Analysis

Molecular orbitals (HOMO and LUMO) and their properties such as energy are very useful for physicists and chemists and are very important parameters for quantum chemistry. This is also used by the frontier electron density for predicting the most reactive position in π - electron systems and also explains several types of reaction in conjugated system.



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S.no	Molecular orbital (eV)	B3LYP/ 6- 31++G(d,p)	B3LYP/ 6- 311++G(d,p)
1.	Еномо	-6.0402	-6.0913
2.	Еномо-1	-7.5218	-7.5825
3.	E _{HOMO-2}	-7.5256	-7.5887
4.	ELUMO	-1.7307	-1.7627
5.	ELUMO+1	-0.3679	-0.3689
6.	E _{LUMO+2}	-0.0822	-0.0849
7.	ΔE(homo-lumo)	4.3095	4.3285
8.	ΔE(homo-1-lumo)	5.7911	5.8197
9.	ΔЕ(номо-2-lumo)	5.7949	5.8260
10.	ΔE(homo-lumo+1)	5.6722	5.7223
11.	ΔЕ(номо-1-цимо+1)	7.1539	7.2135
12.	ΔЕ(номо-2-цимо+1)	7.1577	7.2197
13.	ΔE(homo-lumo+2)	5.9579	6.0064
14.	ΔE(_{HOMO-1-LUMO+2})	7.4396	7.4976
15.	ΔE(_{HOMO-2-LUMO+2})	7.4434	7.5038

The electronic absorption corresponds to the transition from the ground to the first excited state and is mainly described by one electron excitation from the highest occupied molecular orbital (HOMO) to the lowest unoccupied molecular orbital (LUMO). Commonly the atom occupied by more densities of HOMO should have stronger ability for detaching electrons, whereas the atom with more occupation of LUMO should be easier to gain electrons Ravi Kumar et al (2008), Dixon et al (2003). The atomic orbital compositions of the frontier molecular orbital are shown in Figure 5.



LUMO PLOT (First Excited State)

Figure 5 Molecular orbital surface Figure for the HOMO and LUMO of FSC

In order to evaluate the energetic behaviour of the title molecule, the energies of HOMO, LUMO and their orbital energy gaps using B3LYP/6-31++G(d,p) and 6-311++G(d,p) basis sets are calculated and presented in Table 6. The HOMO lying at -6.0402 eV is delocalized over π orbital and includes mixing of lone pairs of electrons on all the oxygen as well as nitrogen atoms. The H-1 at-7.5218 eV below the HOMOs is localized over nitro group through the furan ring. The H-2 at -7.5256 eV below the HOMO and has mainly of π bonding character along with some signa bond character, whereas the LUMO lying at-1.7307 eV is a π orbital, delocalized over the entire molecule with large anti-bonding character from Table 1.6. It is evident that the absorption band centered around arises mainly due to the four electronic transitions given by $H\rightarrow L$, $H-1\rightarrow L$, $H-2\rightarrow L$, $H\rightarrow L+1$, $H-1\rightarrow L+1$, $H-2\rightarrow L+1$, $H\rightarrow L+2$, $H-1\rightarrow L+2$, $H-2\rightarrow L+2$. Are assigned as $\pi-\pi^{\dagger}$ type whereas the more intense band around 312 nm, having maximum oscillator strength (0.567), corresponds to H→L transition and is



mainly characterized to the $LP_{2} \rightarrow \pi^{\hat{}}$ type. The higher intensity of band corresponding to the $LP_{2} \rightarrow \pi^{\hat{}}$ type transition is attributed to the presence of large number of free lone pairs of electrons available on two oxygen and three nitrogen atoms. It can be seen from the plots that the HOMO is spread over the entire molecule and shows appreciable amount of π bonding character. The LUMO is also found to be uniformly distributed over the molecule and reflects a lot of antibonding π character. The energy gap of HOMO-LUMO explains the eventual charge transfer interaction within the molecule, and the frontier orbital energy gap in case of FSC is found to be 4.3095 eV obtained at B3LYP/6-31++G(d,p) method.

Fukui Function

The hard and soft acids and bases principle has been used in a local sense in terms of DFT concepts such as the Fukui function of f(r). Pared Yang showed that sites in chemical species with the largest values of Fukui function f(r) are those with higher reactivity. The Fukui function is defined as Huizar et al (2011):

$$F(r) = [\delta \rho(r) / \delta N]_{u}$$

(3)

 $\label{eq:product} Where \qquad \rho(r) \to electronic \ density,$

 $N \rightarrow$ number of electrons.

The Fukui function is a local reactivity descriptor that indicates the preferred regions where a chemical species will change its density when the number of electrons is modified.

Therefore, it indicates the property of the electronic density to deform at a given position upon accepting (or) donating electrons Roy et al (1999) and also it is possible to define the corresponding condenses or atomic fukui functions on the jth atom site as;

$f_{k} = q_{k} (N) - q_{k} (N-1)$	(4)
$f_{k}^{+} = q_{k} (N+1) - q_{k} (N)$	(5)
$f_k^o = \frac{1}{2}[q_k (N+1) - q_v (N-1)]$	(6)

For an electrophilic $f_k^{-}(r)$, nucleophilic $f_k^{+}(r)$ and free radical attack $f_k^{\circ}(r)$, on the references molecule, respectively. In these equation, q_k is the atomic charge (evaluated from Mulliken population analysis, electrostatic derives charge, etc.) at the kth atomic site in the neutral (N), anionic (N+1) or cationic (N-1) chemical species. Hence, it is important to mention q_k independently of the approximations used for calculation.

$$f(r) dr = 1$$
 (7)

The condensed Fukui functions and condensed local softness indices allow us to distinguish each part of the molecule on the basis of its distinct chemical behaviour due to the different substituent functional groups. Thus, the site for nucleophilic attack will be the place where the value of $\int f(r) dr = 1$ for nucleophilic attack in the most reactive site of FSC is on the O_{10} and C_3 atoms respectively. For electrophilic attack the most reactive site of FSC is on the C_5 and C_1 respectively. These results are shown in Table 7. The condensed local softness indices Sf_k^- and Sf_k^+ are related to the condensed Fukui functions. The local softness also follows the same trend to fukui functions.

Atoms	f _k ⁺	f _k	f _k ⁰	Sf _k ⁺	Sf _k	Sf _k ⁰
C ₁	0.0081	0.1964	0.1022	0.0019	0.0456	0.0237
C ₂	-0.0047	-0.1356	-0.0701	-0.0011	-0.0315	-0.0163
C ₃	0.1180	0.1436	0.1308	0.0274	0.0333	0.0304
C ₄	0.0246	-0.0661	-0.0207	0.0057	-0.0153	-0.0048
C ₅	0.0868	0.2383	0.1626	0.0201	0.0553	0.0377
O ₆	0.0430	0.0107	0.0269	0.0100	0.0025	0.0062
N ₇	0.0815	0.1157	0.0986	0.0189	0.0268	0.0229
N ₈	0.0893	-0.0290	0.0302	0.0207	-0.0067	0.0070
C ₉	-0.0175	0.0014	-0.0081	-0.0041	0.0003	-0.0019
O ₁₀	0.1299	0.0899	0.1099	0.0301	0.0209	0.0255
N ₁₁	0.0275	0.0196	0.0235	0.0064	0.0046	0.0055
H ₁₂	0.0695	0.0843	0.0769	0.0161	0.0196	0.0178
H ₁₃	0.0786	0.0816	0.0801	0.0182	0.0189	0.0186



H ₁₄	0.0762	0.0744	0.0753	0.0177	0.0173	0.0175
H ₁₅	0.0830	0.0765	0.0797	0.0193	0.0178	0.0185
H ₁₆	0.0567	0.0567	0.0567	0.0132	0.0132	0.0132
H ₁₇	0.0389	0.0411	0.0400	0.0090	0.0095	0.0093
H ₁₈	0.0105	0.0005	0.0055	0.0024	0.0001	0.0013

CONCLUSION

In this study the spectroscopic properties such as molecular parameters, frequency assignments and structural confirmation of title compound are examined by using FT-IR, FT-Raman and tools derived from the DFT. The optimized geometric parameters (bond length and bond angles) were theoretically determined B3LYP/6-31++G(d,p) and 6-311++(d,p) levels of theory and compared with the structurally similar compounds. The vibrational FT-IR and FT-Raman spectra of FSC were recorded and computed vibrational wave numbers and their PED were calculated. When all theoretical results are scanned, they are showing good correlation with experimental data. The differences between the observed and scaled wave number values of most of the fundamentals are very small. Therefore, the assignments made at DFT level of theory with only reasonable deviations from the experimental values seem to be correct. NBO analysis is made and it is indicating the intra-molecular charge transfer between bonding and antibonding orbitals. The bond order and atomic charges of the title molecule have been studied by DFT methods. Charge transfer occurs in the molecule between HOMO and LUMO energies, and Frontier energy gap is calculated and presented. The lowering of the HOMO-LUMO energy gap value has substantial influence on the intra-molecular charge transfer and bioactivity of the molecule. Fukui function helps to identify the electrophilic/ nucleophilic nature of a specific site within a molecule.

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