



FT-IR, FT-Raman and ab-initio studies of 1,3-diphenyl thiourea

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ABSTRACT

The FT-IR and FT-Raman spectra of 1,3-diphenyl thiourea were recorded and analyzed. The vibrational wavenumbers were examined theoretically with the aid of the Gaussian03 package of programs using the HF/6-311++G(d,p) and B3LYP/6-311++G(d,p) levels of theory. The data obtained from vibrational wavenumber calculations are used to assign vibrational bands obtained in IR and Raman spectroscopy of the studied molecule. The first hyperpolarizability, infrared intensities and Raman activities are reported. The calculated first hyperpolarizability is comparable with the reported values of similar derivatives and is an attractive object for future studies of non-linear optics. The geometrical parameters of the title compound are in agreement with the values of similar structures. The changes in the C-N bond lengths suggest an extended π -electron delocalization over the thiourea moiety which is responsible for the non-linearity of the molecule.

1. Introduction

Thiourea and substituted thioureas are versatile precursor units in the synthesis of many useful heterocyclic compounds [1]. Theoretical study of the Hg^{2+} recognition by 1,3-diphenylthiourea has been reported [2]. The mechanism of enantioselective Michael addition of acetylacetone to a nitroolefin catalyzed by a thiourea based chiral bifunctional organocatalyst is investigated using density functional theory calculations [3]. Yang *et al.* [4] reported the structural and spectroscopic study of *N*-2-fluorobenzoyl-*N'*-4-methoxyphenyl thiourea. Badawi reported the structural stability, C-N internal rotations and vibrational spectral analysis of non-planar phenylurea and phenylthiourea [5]. Metal organic coordination compounds as non linear optical (NLO) materials have attracted much more attention for their high NLO coefficients, stable physico-chemical properties and better mechanical intension. Many metal organic coordination complexes of thiourea materials with good NLO effects have been designed and synthesized [6-11]. Pfeffer *et al.* [12] reported the anion recognition using pre-organized thiourea functionalized polynorbormance receptors. There is currently great interest in the development of supramolecular systems that have the ability to bind and signal the presence of anions, as well as transporting such species across vesicle and cell membranes [13,14]. Molecules that possess functional groups such as amides, ureas and thioureas [15-17] have proven to be particularly effective in this regard as they are able to bind anions using directional hydrogen bonding interactions. Thiourea and substituted thioureas are especially important for their non linear optical properties. Hence the theoretical study of a symmetrically substituted aryl thiourea, 1,3-diphenyl thiourea, has been attempted.

2. Experimental

1,3-Diphenyl thiourea were prepared by reported method [18]. Briefly, 20 g sodium hydroxide (0.5 mol), 30 mL carbon disulphide (0.5 mol), 94 mL aniline (1 mol) and 5 mL ethyl alcohol were refluxed for 30 minutes in an RB flask. The solid thiourea separated is filtered at the pump, washed with water and ether and dried. White flake with a melting point of 153 °C was obtained. The purity was ascertained by TLC. The FT-IR (Figure 1) was recorded using a Perkin Elmer FTIR Spectrometer Spectrum RX1 with KBr pellets, number of scans 16, resolution 2 cm^{-1} . The FT-Raman spectrum (Figure 2) was obtained on a Bruker Equinox 55/s spectrometer with FRA Raman socket, 106/s. For excitation of the spectrum the emission of Nd:YAG laser was used, excitation wavelength 1064 nm, laser power 250 mW, resolution 2 cm^{-1} , number of scans 128, measurement on solid sample.

3. Computational details

Calculations of the title compound were carried out with Gaussian03 software program [19] using the HF/6-311++G(d,p) and B3LYP/6-311++G(d,p) basis sets to predict the molecular structure and vibrational wavenumbers and also with Becke's three parameter hybrid model using the Lee-Yang-Parr correlation functional (B3LYP) method. Molecular geometries were fully optimized by Berny's optimization algorithm using redundant internal coordinates. Harmonic vibrational wavenumbers were calculated using analytic second derivatives to confirm the convergence to minima on the potential surface. At the optimized structure (Figure 3) of the examined species, no imaginary wavenumber modes were obtained, proving that a true minimum on the potential surface

was found. The DFT hybrid B3LYP functional tends to overestimate the fundamental modes; therefore scaling factors have to be used for obtaining a considerably better agreement with experimental data. Therefore, a scaling factor of 0.9613 and 0.8929 were uniformly applied to the DFT and HF calculated wavenumbers [20]. The observed disagreement between theory and experiment could be a consequence of the anharmonicity and of the general tendency of the quantum chemical methods to overestimate the force constants at the exact equilibrium geometry. Table 1 gives the calculated (DFT) geometrical parameters of the title compound.

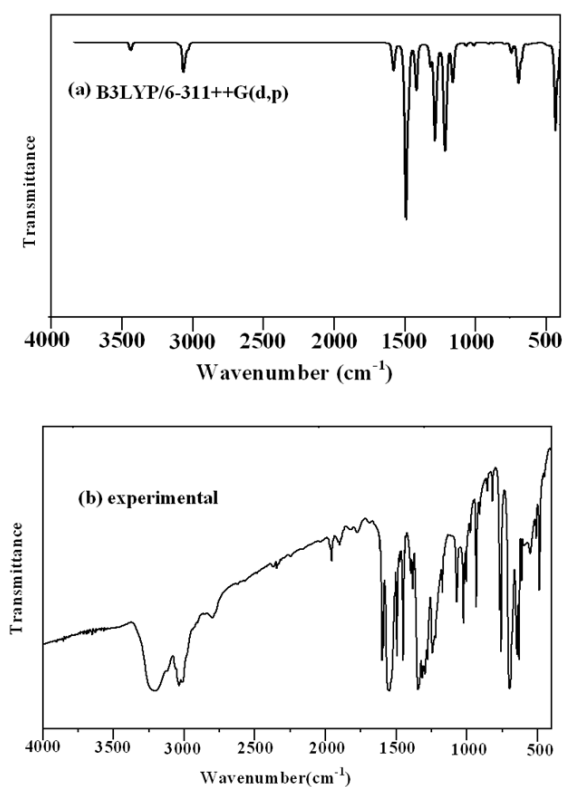


Figure 1. FT-IR spectrum of 1,3-diphenyl thiourea (a) theoretical (b) experimental.

4. Results and Discussion

4.1. IR and Raman spectra

The observed IR, Raman bands with the relative intensities, scaled wavenumbers and assignments are given in Table 2. The assignments of the benzene ring vibrations of the title compound in Wilson notation [21] is made by referring the case of benzene derivatives with mono substitution as summarized by Roeges [22]. The NH stretching vibration [22] appears as a strong broad band in the region $3390 \pm 60 \text{ cm}^{-1}$. In the present study, the NH stretching band is observed at 3217 cm^{-1} in the IR spectrum and at 3207 cm^{-1} in the Raman spectrum. The corresponding calculated values (DFT) are 3438 and 3432 cm^{-1} . The NH stretching wavenumber is redshifted in the IR spectrum with a strong intensity from the computational wavenumber which indicates the weakening of the N-H bond resulting in proton transfer to the neighbouring sulfur [23]. El-Asmy and Al-Hazmi [24] reported ν_{NH} in the region $3138\text{--}3323 \text{ cm}^{-1}$. The CNH vibrations in which N and H atoms move in opposite directions of the carbon atom in the amide moiety appear at $1507, 1494 \text{ cm}^{-1}$ in the IR spectrum, $1517, 1490 \text{ cm}^{-1}$ in the Raman spectrum and at $1507, 1491 \text{ cm}^{-1}$ theoretically (DFT) and the CNH vibrations in which N and H atoms move in the same direction of the carbon atom in the amide group

appear at $1298, 1278 \text{ (IR)}, 1249 \text{ (Raman)}$ and $1292, 1285 \text{ cm}^{-1}$ (DFT) [25-27]. The NH rock in the plane is not seen in the IR spectrum but the DFT calculations give this mode at 1139 cm^{-1} [27]. The out-of-plane wagging [22] of NH is moderately active with a broad band in the region $790 \pm 70 \text{ cm}^{-1}$ and the band at 858 cm^{-1} in the IR spectrum, 858 cm^{-1} in the Raman spectrum and $872, 868 \text{ cm}^{-1}$ (DFT) are assigned as this mode. El-Shahawy *et al.* [27] reported a value 710 cm^{-1} for this mode. Badawi reported [5] the NH vibrational modes at $3315, 1498, 1268 \text{ cm}^{-1}$ in the IR spectrum, $3320, 1508, 1264 \text{ cm}^{-1}$ in the Raman spectrum and at $3617, 1502, 1256 \text{ cm}^{-1}$ theoretically. Panicker *et al.* [28] reported the NH bending modes at $1538, 1220 \text{ cm}^{-1}$ in the IR spectrum and at $1558, 1223 \text{ cm}^{-1}$ theoretically (DFT). The C-N stretching vibration [22] coupled with δ_{NH} , is moderately to strongly active in the region $1275 \pm 55 \text{ cm}^{-1}$. El-Shahawy *et al.* [27] observed a band at 1320 cm^{-1} in the IR spectrum as the ν_{CN} mode. Yang *et al.* [4] reported aromatic C-N stretching band at 1359 cm^{-1} . In the present case, the bands at $1326, 1312 \text{ cm}^{-1}$ in the IR spectrum and at $1345, 1316 \text{ cm}^{-1}$ in the Raman spectrum are assigned as this mode. The DFT calculations give the corresponding bands at 1318 and 1307 cm^{-1} which are not pure but contain significant contributions from other modes also.

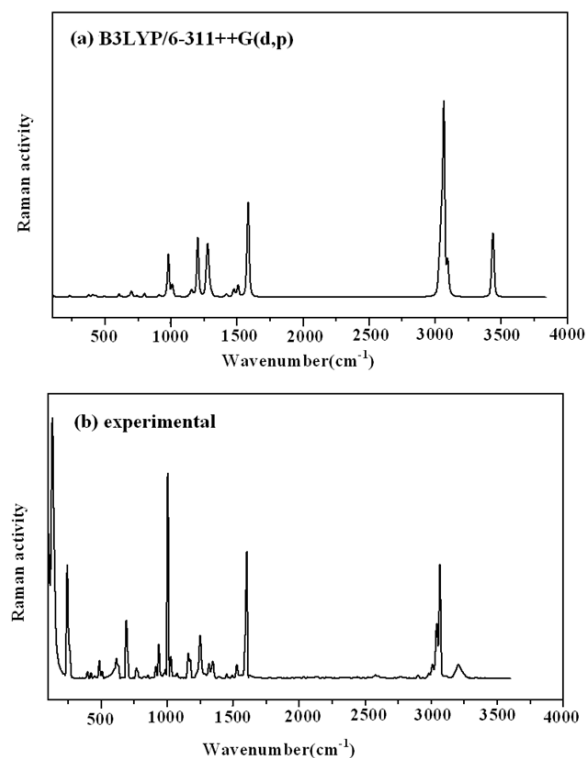


Figure 2. FT-Raman Spectrum of 1,3-diphenyl thiourea (a) theoretical (b) experimental.

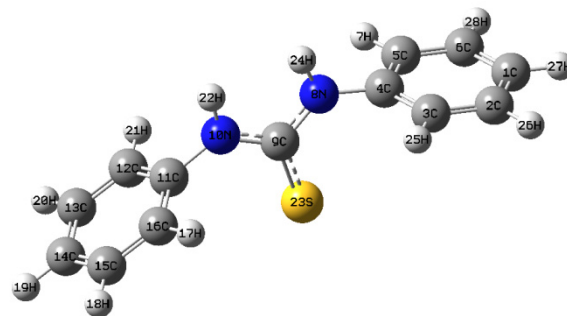


Figure 3. Optimized geometry of 1,3-diphenyl thiourea.

Table 1. Geometrical parameters (DFT) of 1,3-diphenyl thiourea, atom labeling according to Figure 3.

Bond lengths(Å)		Bond angles(°)		Dihedral angles(°)	
C1-C2	1.3949	C2-C1-C6	119.4	C6-C1-C2-C3	-0.1
C1-C6	1.3928	C1-C2-C3	120.8	C2-C1-C6-C5	-0.6
C2-C3	1.3915	C2-C3-C4	119.7	C1-C2-C3-C4	0.8
C3-C4	1.3976	C3-C4-C5	119.6	C2-C3-C4-C5	-0.8
C4-C5	1.3981	C3-C4-N8	122.1	C2-C3-C4-N8	-177.0
C4-N8	1.4188	C5-C4-N8	118.2	C3-C4-C5-C6	0.0
C5-C6	1.3928	C4-C5-C6	120.3	N8-C4-C5-C6	176.4
C9-N8	1.3814	C1-C6-C5	120.1	C3-C4-N8-C9	-47.3
C9-N10	1.3814	C4-N8-C9	128.6	C5-C4-N8-C9	136.4
C9-S23	1.6636	N8-C9-N10	110.1	C4-C5-C6-C1	0.7
C11-N10	1.4188	N8-C9-S23	124.9	C4-N8-C9-N10	173.0
C11-C12	1.3975	N10-C9-S23	124.9	C4-N8-C9-S23	-7.0
C11-C16	1.3981	C9-N10-C11	128.6	N8-C9-N10-C11	173.1
C12-C13	1.3915	N10-C11-C12	122.1	S23-C9-N10-C11	-6.9
C13-C14	1.3949	N10-C11-C16	118.2	C9-N10-C11-C12	-47.4
C14-C15	1.3928	C12-C11-C16	119.6	C9-N10-C11-C16	136.3
C15-C16	1.3928	C11-C12-C13	119.7	N10-C11-C12-C13	-177.0
		C12-C13-C14	120.8	C16-C11-C12-C13	-0.8
		C13-C14-C15	119.4	N10-C11-C16-C15	176.4
		C14-C15-C16	120.1	C12-C11-C16-C15	0.0
		C11-C16-C15	120.3	C11-C12-C13-C14	0.8
				C12-C13-C14-C15	-0.2
				C13-C14-C15-C16	-0.6
				C14-C15-C16-C11	0.7

The C=S stretching bands [22,25,29] are observed in the range $800 \pm 130 \text{ cm}^{-1}$, usually with a moderate intensity. Since the S atom is less electronegative than the O atom, the C=S group is not as polar as the C=O group and less prone to form bridges. The C=S deformation bands are expected in the regions [22] 510 ± 90 and $420 \pm 100 \text{ cm}^{-1}$. Nigam *et al.* [30] reported the C=S stretching mode at 847 and 835 cm^{-1} . For the title compound, the C=S stretching mode is observed at 908 cm^{-1} in IR, 911 cm^{-1} in Raman and at 908 cm^{-1} theoretically (DFT).

The existence of one or more aromatic ring in a structure is normally readily determined from the C-H and C=C related vibrations. The C-H stretching occurs above 3000 cm^{-1} and is typically exhibited as a multiplicity of weak to moderate bands, compared with the aliphatic C-H stretch [31]. For mono substituted benzenes, the CH stretching modes are expected in the region 3105-3000 cm^{-1} [22]. There are five CH stretching modes 20a, 20b, 2, 13, 7b for mono substituted benzenes. According to selection rules all five bands are allowed in the IR spectrum [26]. The calculated values are in the range 3097-3032 cm^{-1} . The bands observed at 3113, 3062, 3042, 3020 cm^{-1} in the IR spectrum and at 3113, 3068, 3040, 3010 cm^{-1} in the Raman spectrum were assigned as νCH modes of the benzene rings.

The benzene ring possesses six ring stretching vibrations, of which the four with the highest wavenumbers (8a, 8b, 19a and 19b, occurring respectively near 1600, 1580, 1490 and 1440 cm^{-1}) are good group vibrations. In the absence of ring conjugation, the band near 1580 cm^{-1} is usually weaker than that at 1600 cm^{-1} . The fifth ring stretching vibration νPh 14 is active near $1335 \pm 35 \text{ cm}^{-1}$, a region which overlaps strongly with that of the CH in-plane deformation and the intensity is in general, low or medium high [22,26]. The sixth ring stretching vibration or ring breathing mode νPh 1 appears as a weak band near 1000 cm^{-1} in mono substituted benzenes [22]. The bands observed at 1600, 1589, 1562, 1475, 1460, 1434, 1326, 1312 cm^{-1} in the IR spectrum and at 1602, 1465, 1345, 1316 cm^{-1} in the Raman spectrum are assigned as νPh ring stretching modes. As seen from the Table 2, the DFT calculations give these modes in the range 1584- 1307 cm^{-1} . These vibrations are expected in the region 1620-1300 cm^{-1} [22]. The frequency of the vibrational pair 8, 19a in mono substituted benzenes is rather insensitive of substitution [26]. The frequency interval for vibration 19a is 1470-1515 cm^{-1} [26,32] and for all mono substituted benzene derivatives 19a have a band between 1477 and 1511 cm^{-1} in the IR spectrum. When derivatives have substituents of donor character, the frequency is found

between 1493 and 1511 cm^{-1} and in this group of derivatives the band has higher intensity when a carbon atom is attached directly to the ring. For the title compound the ring breathing modes are assigned at 979, 978 cm^{-1} theoretically. The strong band at 1004 cm^{-1} in the Raman spectrum is assigned with confidence to the characteristic ring breathing mode of the title compound. The ring breathing mode for mono substituted benzene is reported at 983 cm^{-1} by Panicker *et al.* [33] and at 1010 cm^{-1} in the Raman spectrum by Badawi *et al.* [5] for phenyl urea.

In mono substituted benzenes, there should be five CH in-plane bending vibrations 3, 9a, 9b, 15 and 18a. The mode 9a appears between 1170 and 1181 cm^{-1} in the spectra of all mono substituted benzene derivatives, mode 15 is found between 1060 and 1080 cm^{-1} and the intensity of 9a and 18a are the most intense in the IR spectrum [26,32]. The mode 3 is weak in general for both IR and Raman in the range 1253-1331 cm^{-1} [26]. In the present case the bands observed in the range 1003-1266 cm^{-1} in the IR spectrum, 1165 – 1024 cm^{-1} in the Raman spectrum and in the range 1009-1275 cm^{-1} (DFT) are assigned as δCH modes of the phenyl rings. Some modes are not pure, but contain significant contributions from other modes also. The in-plane CH vibrations are expected in the range 1015-1300 cm^{-1} for mono substituted benzenes [22].

The out-of-plane CH deformations γCH of mono substituted benzene derivatives have the modes 5, 10a, 11, 17a and 17b [22,26]. These modes are expected in the range 1000-700 cm^{-1} [22]. Generally, the CH out-of-plane deformations with the highest wavenumbers have a weaker intensity than those absorbing at lower wavenumbers. The stronger γCH band occurring in the region $775 \pm 45 \text{ cm}^{-1}$ (γCH 11 or umbrella mode) tends to shift to lower (higher) wavenumbers with increasing electron donating (attracting) power of the substituent, but seems to be more sensitive to mechanical interaction effects. The lowest wavenumbers for this umbrella mode are found in the spectra of benzenes substituted with a saturated carbon or heavy atoms such as halogen, sulfur or phosphorus [26,34-36]. The bands at 938, 809, 789, 752, 698 cm^{-1} in the IR spectrum, 936, 765, 690 cm^{-1} in the Raman spectrum and the calculated values (DFT) in the range 945-699 cm^{-1} are assigned as the out-of-plane CH modes of the phenyl rings. The out-of-plane CH deformation γCH 11 at 698 cm^{-1} and the out-of-plane ring deformation γPh 4 at 655, 632 cm^{-1} in the IR spectrum form a pair of strong bands characteristic of mono substituted benzene derivatives, which find support from computational results [22,36].

Table 2. Calculated (scaled) vibrational wavenumbers, measured infrared and Raman band positions and assignments for 1,3-diphenyl thiourea*.

HF/6-311++G(d,p)			B3LYP/6-311++G(d,p)			IR		Raman	Assignments
ν (cm ⁻¹)	IR intensity	Raman activity	ν (cm ⁻¹)	IR intensity	Raman activity	ν (cm ⁻¹)	ν (cm ⁻¹)		
3437	54.13	207.48	3438	18.55	273.43	3217 sbr	3207 sbr	vNH	
3429	8.21	0.22	3432	10.57	28.07			vNH	
3009	0.21	521.89	3097	0.26	90.23	3113 sh	3113 w	vCH 20a	
3009	7.66	27.09	3097	4.36	21.65			vCH 20a	
3005	54.10	0.71	3066	0.07	516.99		3068 s	vCH 20b	
3005	0.00	0.04	3066	56.25	63.04	3062 sh		vCH 20b	
2997	5.70	103.11	3054	25.16	104.16			vCH 2	
2996	70.71	13.68	3054	28.19	87.46			vCH 2	
2987	14.18	26.86	3044	0.27	118.88		3040 s	vCH 13	
2987	0.79	198.91	3044	1.17	72.08	3042 m		vCH 13	
2976	0.02	70.85	3032	8.47	42.84			vCH 7b	
2976	0.32	6.60	3032	12.46	41.17	3020 w	3010 w	vCH 7b	
1605	0.01	69.97	1584	0.68	378.46	1600 s	1602 vs	vPh 8a	
1601	37.17	3.50	1583	27.14	4.37	1589 s		vPh 8a	
1584	4.32	3.81	1578	1.36	7.12			vPh 8b	
1583	0.00	19.94	1574	69.67	6.01	1562 sbr		vPh 8b	
1529	0.75	5.15	1507	427.45	44.96	1507 m	1517 m	δ NH	
1514	694.37	14.95	1491	427.45	0.74	1494 s	1490 w	δ NH	
1478	0.08	1.81	1475	0.41	38.50	1475 w		vPh 19a	
1478	3.31	2.65	1474	172.86	1.01	1460 s	1465 w	vPh 19a	
1431	10.43	0.19	1426	2.85	8.28	1434 sh		vPh 19b	
1431	0.00	2.05	1416	152.88	7.10			vPh 19b	
1300	0.41	0.01	1318	48.36	2.73	1326 s	1345 m	vCN, vPh 14	
1299	0.00	1.29	1307	2.35	4.15	1312 m	1316 w	vCN, vPh 14	
1283	286.93	5.00	1292	26.02	47.13	1298 m		δ NH	
1234	405.06	2.81	1285	243.79	0.76	1278 m	1249 s	δ NH	
1224	97.96	32.87	1275	106.34	240.58	1266 m		δ CH 3	
1181	3.91	0.55	1216	368.10	0.35	1218 w		δ CH 3	
1181	0.00	0.77	1202	75.85	198.71			δ CH 9a	
1169	28.65	0.05	1161	75.94	0.28	1168 w	1165 m	ρ CH 9a	
1145	1.62	5.29	1159	0.52	20.29		1159 m	ρ CH 9b	
1144	6.15	0.83	1156	51.81	9.34	1156 w		δ CH 9b	
1071	14.49	0.00	1139	0.34	6.78			ρ NH	
1071	0.00	4.42	1139	1.01	4.21			ρ NH	
1039	1.96	0.46	1065	10.56	0.38	1070 s	1070 w	vCX(X) 7a, δ CH 15	
1039	0.00	0.36	1063	0.70	1.60			vCX(X) 7a, δ CH 15	
1001	0.73	11.18	1011	0.46	41.57	1018 m	1024 w	δ CH 18a	
998	13.77	0.27	1009	10.19	1.01	1003 w		δ CH 18a	
992	0.00	0.43	979	0.26	156.74		1004 vs	vPh 1	
992	0.01	0.43	978	0.06	2.62	975 w		vPh 1	
975	0.55	0.00	945	0.01	0.12			γ CH 5	
975	0.00	0.00	945	0.19	0.14	938 s		γ CH 5	
972	0.62	76.75	920	0.53	1.78		936 w	γ CH 17a	
970	4.41	0.20	920	0.16	0.34			γ CH 17a	
931	4.33	0.06	908	6.07	9.76	908 w	911 w	vCS	
930	1.64	34.93	872	4.01	0.41			ω NH	
898	10.46	24.68	868	0.22	0.40	858 w	858 w	ω NH	
839	2.26	0.71	800	0.73	0.56	809 w		γ CH 17b	
838	0.00	0.12	798	1.19	12.03			γ CH 17b	
807	10.40	1.43	792	3.90	1.21	789 s		γ CH 10a	
759	26.25	13.32	747	20.68	1.21	752 s	765 w	γ CH 10a	
756	12.66	2.89	742	13.22	4.86			γ CH 11	
684	7.79	0.20	699	64.56	27.49	698 s	690 s	γ CH 11	
683	145.77	6.87	691	70.30	0.66	690 s		δ Ph(X) 12	
669	10.40	14.71	673	29.56	3.08			δ Ph(X) 12	
637	0.51	0.00	667	10.06	0.12	655 s		γ Ph 4	
635	14.26	0.37	629	6.45	2.33	632 s	628 w	γ Ph 4	
619	16.00	3.96	608	2.50	4.14	611 m	615 m	δ CS	
604	0.00	8.57	607	0.74	8.56	599 w		δ Ph 6b	
602	1.27	1.10	571	2.86	1.60	563 w		δ Ph 6b	
490	9.63	2.85	497	5.95	3.54	505 w	506 w	γ CS	
476	13.50	2.01	488	0.75	1.74	486 s	485 w	γ Ph(X) 16b	
436	0.01	0.23	478	4.96	0.02			γ Ph(X) 16b	
425	102.84	0.30	433	216.39	5.80	439 w	440 w	ρ CSN	
410	0.00	0.02	415	74.15	6.74		419 w	γ Ph(X)	
403	39.72	0.16	402	0.07	5.98			γ Ph(X)	
400	0.00	0.17	401	0.15	0.42		398 w	δ Ph(X) 6a	
380	1.73	0.29	375	2.79	8.72			δ Ph(X) 6a	
238	0.83	3.53	321	0.32	0.21			δ CX(X) 18b	
222	4.407	2.09	235	0.75	7.78		242 s	δ CX(X) 18b	
218	0.35	4.50	228	2.54	0.77			ρ C(=S)NH	
175	0.00	0.02	204	1.03	0.79			ρ CSN	
103	4.98	4.25	130	8.04	1.88		129 s	γ CX(X) 10b	
71	0.00	0.75	106	0.03	8.58			γ CX(X) 10b	
52	0.84	1.94	54	0.31	4.35			tPh	
27	2.95	0.15	40	1.80	4.71			tPh	
20	3.68	8.51	32	0.05	12.78			tCSN	
7	0.00	1.26	28	0.13	2.00			tCSN	

* ν -stretching; δ -in-plane deformation; γ -out-of-plane deformation; ρ -rocking; ω -wagging; t-torsion; X-substituent sensitive; Ph-phenyl ring.

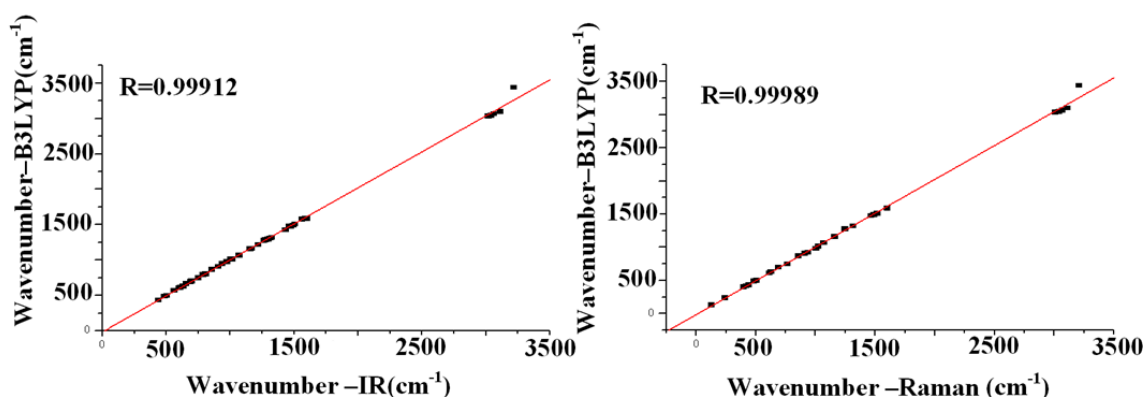


Figure 4. Correlation graph between experimental and calculated wavenumbers.

In mono substituted benzenes, six modes of vibrations are substituent sensitive, which means that their wavenumbers shift systematically with mass or inductive or mesomeric effects of the substituent. The highest substituent sensitive mode $\nu_{CX(X)}$ mode 7a appears in the region $1195 \pm 90 \text{ cm}^{-1}$ for mono substituted benzenes [22,37]. The band corresponding to mode 7a is generally strong both in IR and Raman spectra. For the title compound the $\nu_{CX(X)}$ modes are observed in the IR spectrum at 1070 cm^{-1} , and at 1070 cm^{-1} in Raman spectrum. The DFT calculations give values at $1065, 1063 \text{ cm}^{-1}$. Moreover, the other substituent sensitive modes are identified (Table 2). The correlation graph between experimental and calculated wavenumbers is shown in Figure 4. The IR bands in the $1729\text{--}2770 \text{ cm}^{-1}$ region and their large broadening support the intramolecular hydrogen bonding [33].

4.2. Geometrical parameters and first hyperpolarizability

The C-C bond length in the phenyl rings lie between 1.3915 and 1.3981 \AA . Here for the title compound, benzene is a regular hexagon with bond lengths somewhere between the normal values for a single (1.54 \AA) and a double (1.33 \AA) bond [38]. The $C_{11}\text{--}C_{12}$, $C_{11}\text{--}C_{16}$, $C_4\text{--}C_3$ and $C_4\text{--}C_5$ bond lengths are longer than all the other C-C bonds. This is due to the presence of the electronegative nitrogen atom, which can pull electrons from the ring elongating the bond. Since the molecule is symmetric, all the structural parameters also exhibit the symmetry on both sides. Both the N-H bonds are oriented away from the C-S bond so as to reduce the repulsive forces. Another interesting feature of the molecule is that, the planes of both the benzene rings are perpendicular to each other. Yang *et al.* [4] reported bond lengths $N_{10}\text{--}C_9=1.38$, $N_8\text{--}C_9=1.404 \text{ \AA}$, whereas in the present case, the corresponding value is 1.3814 \AA and shorter than C-N single bond of 1.4725 \AA . The partial double bond character of the structure is presumed as a result of the intra-molecular H-bond "locking" the molecule into a planar six-numbered ring structure.

The C-S bond length in the present study 1.6636 \AA lies between the values of C-S single and double bonds. The $C_4\text{--}N_8$ bond length for the title compound is 1.4188 \AA , formally a single bond, which is remarkably shortened. The reported values of the bond lengths are, $C_9\text{--}S_{23}=1.663$ [4] and $C_4\text{--}N_8=1.421 \text{ \AA}$ [4]. Dilovic *et al.* [39] reported the bond lengths, $C_9\text{--}S_{23}=1.6792$, $C_9\text{--}N_8=1.3422$, $C_9\text{--}N_{10}=1.3462$. $C_4\text{--}N_8=1.4122 \text{ \AA}$, bond angles $C_{11}\text{--}N_{10}\text{--}C_9=129.7$, $N_{10}\text{--}C_9\text{--}N_8=114.3$, $N_{10}\text{--}C_9\text{--}S_{23}=116.8$, $S_{23}\text{--}C_9\text{--}N_8=128.8$, $C_9\text{--}N_8\text{--}C_4=130.7$, $N_8\text{--}C_4\text{--}C_5=124.7^\circ$ and the dihedral angles $C_{11}\text{--}N_{10}\text{--}C_9\text{--}S_{23}=-173.4$, $C_4\text{--}N_8\text{--}C_9\text{--}S_{23}=-1.8$, $C_4\text{--}N_8\text{--}C_9\text{--}N_{10}=179.1^\circ$, which are in agreement with the corresponding values for the title compound. Ramnathan *et al.* [40] reported $C_9\text{--}S_{23}=1.7073$, $C_4\text{--}N_8=1.445$, $C_{11}\text{--}N_{10}=1.4375 \text{ \AA}$, $C_9\text{--}N_8\text{--}C_4=127.8^\circ$, $C_9\text{--}N_{10}\text{--}C_{11}=125.5^\circ$, $N_8\text{--}C_9\text{--}N_{10}=117.3^\circ$, $N_8\text{--}C_9\text{--}S_{23}=$

120.4° , $N_{10}\text{--}C_9\text{--}S_{23}=122.2^\circ$, whereas the corresponding values in the present case are, 1.6636 , 1.4188 , 1.4188 \AA , 128.6 , 128.6 , 110.1 , 124.9 , and 124.9° .

The first hyperpolarizability (β_0) of this novel molecular system is calculated using B3LYP/6-311++G(d,p) method, based on the finite field approach. In the presence of an applied electric field, the energy of a system is a function of the electric field. First hyperpolarizability is a third rank tensor that can be described by a $3 \times 3 \times 3$ matrix. The 27 components of the 3D matrix can be reduced to 10 components due to the Kleinman symmetry [41]. The components of β are defined as the coefficients in the Taylor series expansion of the energy in the external electric field. When the electric field is weak and homogeneous, this expansion becomes

$$E = E_0 - \sum_i \mu_i F^i - \frac{1}{2} \sum_{ij} \alpha_{ij} F^i F^j - \frac{1}{6} \sum_{ijk} \beta_{ijk} F^i F^j F^k - \frac{1}{24} \sum_{ijkl} \gamma_{ijkl} F^i F^j F^k F^l + \dots \quad (1)$$

where, E_0 is the energy of the unperturbed molecule, F^i is the field at the origin, μ_i , α_{ij} , β_{ijk} and γ_{ijkl} are the components of dipole moment, polarizability, the first hyper polarizabilities, and second hyperpolarizabilities, respectively. The calculated first hyperpolarizability of the title compound is 1.86×10^{-30} esu, which is 14.36 times that of urea (0.13×10^{-30} esu) [42]. The C-N distances in the calculated molecular structure vary from 1.3814 to 1.4188 \AA which are intermediate between those of a C-N single bond (1.48 \AA) and a C=N double bond (1.28 \AA). Therefore, the calculated data suggest an extended π -electron delocalization over the thiourea moiety [42,43] which is responsible for the nonlinearity of the molecule. We conclude that the title compound is an attractive molecule in future for non linear optical applications.

5. Conclusion

The FT-IR and FT-Raman spectra of 1,3-diphenyl thiourea were recorded and analyzed. The wavenumbers are calculated theoretically using Gaussian03 software package. The normal modes of the phenyl ring are assigned by comparing experimental wavenumbers and theoretically calculated wavenumbers. The NH stretching wavenumber is redshifted in the IR spectrum with a strong intensity from the computational wavenumber which indicates the weakening of the N-H bond resulting in proton transfer to the neighboring sulfur atom. The small differences between experimental and calculated vibrational modes are observed. It must be due to the fact that hydrogen bond vibrations present in the crystal lead to strong perturbation of the infrared wavenumbers and intensities of many other modes. Furthermore, we state that experimental results belong to solid phase and theoretical calculations belong to gaseous phase. The first hyperpolarizability, infrared

intensities and Raman activities are reported. The calculated first hyperpolarizability is comparable with the reported values of similar derivatives and is an attractive object for future studies of non-linear optics.

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