

Molecular Dynamic Simulation Studies of 4-(Trifluoromethyl)phenylacetonitrile

D. Gnanasoundar Iraiyan¹, S. Joseph Vedhagiri²,
M. Govindarajan³, K. Parimala^{4*}

¹Faculty of Allied Health Sciences, Vinnayakka Mission University,
Puducherry-607402, India

²PG & Research Department of Physics, TBML College
(Affiliated to Bharathidasan University), Porayar-609307, India

³Department of Physics, Auvaigar Govt. College for Women
(Affiliated to Pondicherry University), Karaikal-609602, India

⁴PG&Research Department of Physics, Nehru Memorial College
(Affiliated to Bharathidasan University), Trichy-621007, India

*Corresponding author E-mail: kparimala79@yahoo.co.in

Received 04 August 2021, Revised 27 September 2021

Abstract. In this present look at on 4-(Trifluoromethyl)phenylacetonitrile (4TFMNB) has been investigated with the aid of FT-IR, FT-Raman and NMR (¹H and ¹³C) spectral works. Most of the vibrational assignments of FT-IR and FT-Raman spectra predicted using density functional theory (DFT) method match well with experimental findings. The ¹³C and ¹H NMR chemical shifts of the title compound is obtained by the gauge independent atomic orbital (GIAO) method and compared with experimental results.

KEY WORDS: FT-IR, FT-Raman, NMR, DFT.

1 Introduction

Phenylacetonitrile molecules having regular structure and the identically stabilized with the aid of using dipole-dipole moment interactions, while alcohols are self-associated through the hydrogen bonding in their hydroxyl groups, developing multimers of various degrees. Alcohols are used as hydraulic fluids for machine, in medications for animals, in manufacturing of perfumes, flavours and dyestuffs, paint removers, as defrosting and as an antiseptic agent. Phenylacetonitrile is used in organic synthesis of dyes, perfumes, pesticides, pharmaceuticals and especially penicillin precursors. Recently, substantial research work has been reported on the excess properties of acetonitrile+alkanols [1], acrylonitrile+alkanols [2, 3] and benzonitrile+alkanols [4], while that on phenylacetonitrile+alkanol is relatively very rare.

Literature survey reveals that no detailed B3LYP with 6-311+G(d,p) basis set of FT-IR, FT-Raman and NMR (^{13}C and ^1H) chemical shifts calculation of 4TFMNB have been reported so far. It is, therefore thought worth to make the theoretical and experimental vibrational spectral details based on optimized molecular structure to give the correct assignment of fundamental bands in the experimentally observed FT-IR and FT-Raman spectra. In this study, molecular geometry and vibrational frequencies are predicted by hybrid density functional method. This method is used to give relatively correct molecular structure and vibrational spectra with moderate computational effort.

2 Experimental Details

The fine sample of 4-(Trifluoromethyl)phenylacetonitrile (4TFMNB) is purchased from Sigma-Aldrich chemicals, USA and it was used as such without any further purification. The FT-IR spectrum of selected compound has been recorded using Perkin-Elmer 180 spectrometer in the range of $4000\text{--}400\text{ cm}^{-1}$. The spectral resolution is $\pm 2\text{ cm}^{-1}$. The FT-Raman spectrum of the compound was also recorded in same instrument with FRA 106 Raman module equipped with Nd:YAG laser source operating in the region $3500\text{--}100\text{ cm}^{-1}$ at 1064 nm line width with 200 MW powers. ^{13}C and ^1H NMR spectra were taken in CDCl_3 solutions and all signals were referenced to TMS on a BRUKER TPX-400 FT-NMR spectrometer.

3 Computational Details

The molecular structure of the 4TFMNB molecule in the ground state is computed through B3LYP/6-311+G(d,p) method. The optimized structural parameters are used with in the vibrational wavenumber calculations at B3LYP method. The DFT calculations had been done for 4TFMNB with GAUSSIAN 09W software package [5]. Initial geometry generated from the standard geometrical parameters become minimized without any constraint, which invokes Becke's three parameter hybrid method [6] with Lee-Yang-Parr correlation functional (LYP) [7], implemented the same basis set for higher description of the bonding properties. All the parameters had been allowed to loosen up and all the calculations converged to an optimized geometry which corresponds to a real minimum, as revealed by the lack of imaginary values in the wave number calculations. The Cartesian representation of the theoretical force constants had been computed on the completely optimized geometry.

Transformation of force field, the subsequent evaluation of normal coordinate along with the least square refinement algorithm of the scale factors and calculation of the full strength energy distribution (TED) were achieved on a PC with the MOLVIB program (model V7.0-G77) written by Sundius [8, 9]. The calculated frequencies are scaled using 0.9689 scale factor [10, 11]. As a result,

the unscaled frequencies, reduced masses, force constants, infrared intensities and Raman had been obtained. The systematic comparison of the consequences from DFT theory with results of experiments has proven that the method usage of B3LYP functional is the most promising in providing accurate vibrational wave numbers.

The Raman activities (S_i) calculated by the Gaussian 09W software become transformed to relative Raman intensities (I_i) by the following relationship derived from the intensity theory of Raman scattering [12, 13]:

$$I_i = \frac{f(\nu_0 - \nu_i)^4 S_i}{\nu_i [1 - \exp(-hc\nu_i/KT)]}, \quad (1)$$

where ν_0 is the laser exciting wavenumber in cm^{-1} (in this work, we have used the excitation wavenumber $\nu_0 = 9398.5 \text{ cm}^{-1}$, which corresponds to the wavelength of 1064 nm of a Nd:YAG laser), ν_i is the vibrational wavenumber of the i^{th} normal mode (cm^{-1}), while S_i is the Raman scattering activity of the normal mode ν_i , f (is a constant equal to 10^{-12}) is a suitably chosen common normalization factor for all peak intensities h , k , c and T are Planck and Boltzmann constants, speed of light and temperature in Kelvin, respectively.

In this study, the ^{13}C and ^1H NMR chemical shifts and nuclear magnetic shielding tensor values of DFNB in the ground state for B3LYP/6-311+G(d,p) method were calculated with the gauge independent atomic orbital (GIAO) method [14, 15]. All calculations were performed using Gaussian 09W program package [5] employing B3LYP method with 6-311+G(d,p) basis set.

4 Results and Discussion

4.1 Vibrational assignments

The optimized molecular structure of 4TFMNB is shown in Figure 1. From the structural point of view the compound is assumed to have C1 point group sym-

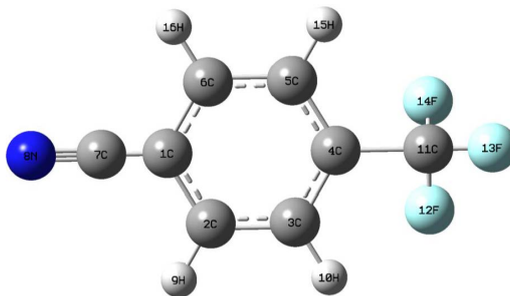


Figure 1. The molecular structure with atom numbering of 4TFMNB.

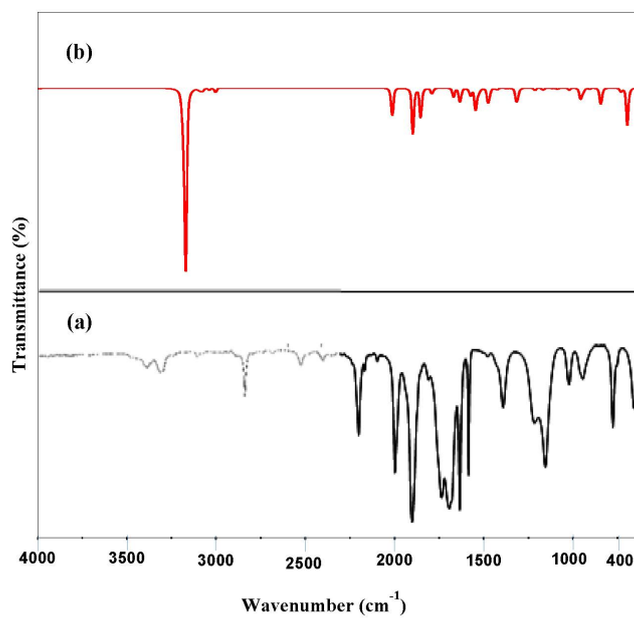


Figure 2. FT-IR spectra of 4TFMNB

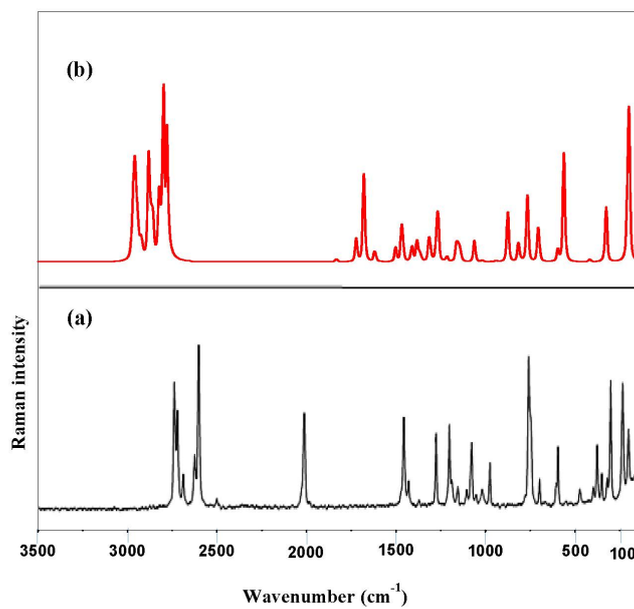


Figure 3. FT-Raman spectra of 4TFMNB

metry and hence, all the calculated frequency transforming to the same symmetry species (A). The 4TFMNB molecule consists of 16 atoms and expected 42 normal modes of vibrations. The detailed vibrational assignments of fundamental modes of 4TFMNB along with the TED are reported in Table 1. The observed FT-IR and FT-Raman spectra of 4TFMNB are shown in Figures 2 and 3, respectively.

4.1.1 C–H vibrations

The C–H stretching vibrations are occurred in the region 3100–3000 cm^{-1} and these vibrations are not found to be affected due to the nature and position of the substituent [16, 17]. Accordingly, in title compound the C–H stretching vibrations are observed at 3415 and 3300 cm^{-1} in the FT-IR spectrum.

The C–H in-plane bending vibrations usually occurs in the region 1430–990 cm^{-1} [18, 19]. The FT-IR peaks observed at 1700 and 1560 cm^{-1} and FT-Raman peaks observed at 1425 cm^{-1} are assigned to C–H in-plane bending vibrations. C–H out-of-plane bending vibrations are occur in the region 900–667 cm^{-1} . In 4TFMNB peaks obtained at 980, 915 and 820 cm^{-1} in both FT-IR and FT-Raman spectra confirms the C–H out-of-plane bending vibrations which agrees with the calculated values.

4.1.2 C–C vibrations

The bands observed at 1430–1670 cm^{-1} are assigned to C–C stretching modes [20]. Socrates [21] mentioned that the presence of conjugate substituent such as C–C vibrations 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} [22]. As predicted in the earlier references, in 4TFMNB compound also there are prominent peaks at 2780, 2000, 1880, and 1750 cm^{-1} in FT-IR spectrum and peaks observed at 2750 and 2000 cm^{-1} in FT-Raman spectrum are assigned to C–C stretching vibrations. The C–C in-plane and out-of-plane bending vibrations are appeared at 1480, 1310, 1115 cm^{-1} and 750, 220, 160 cm^{-1} in FT-Raman spectrum are listed in Table 1. The same assignments are in good agreement with the literature [23].

4.1.3 C–N vibrations

The identification of C–N vibrations are a very difficult task, since the mixing of several modes are possible in the characteristic region. Silverstein *et. al.*, [24] have assigned C–N stretching vibrations in the region 1382–1266 cm^{-1} for aromatic compounds. In the present study, observed band at 1380 cm^{-1} in

Table 1. Vibrational assignments of fundamental frequencies are obtained for 4TFMNB using B3LYP/6-311+G(d,p) level of theory. Experimental relative intensities are abbreviated as follows: vs – very strong; s – strong; m –medium; w – weak; vw – very weak. Abbreviations; ν – stretching; sym – symmetric stretching; asym – asymmetric stretching; b – in-plane bending; ω – out-of-plane bending; ipb – in-plane bending; opb – out-of-plane bending; sb – symmetric bending; ipr – in-plane rock; opr – out-of-plane rock; t – torsion.

Sl. No.	Observed frequencies frequencies (cm^{-1})		Calculated frequencies frequencies (cm^{-1})		Assignments with > 10% of TED
	FT-IR	FT-Raman	B3LYP		
			Unscaled	Scaled	
1	3415(vw)		3400	3378	ν C-H (100)
2	3300(vw)		3243	3231	ν C-H (99)
3			3125	3178	ν C-H (98)
4			3092	3081	ν C-H (98)
5			3050	3031	ν C-C (95)
6			3037	3018	ν C-C (93)
7			3006	2918	ν C-C (91)
8			2987	2963	ν C-C (91)
9	2780(m)	2750(s)	2932	2908	ν C-C (90)
10	2000(s)	2000(s)	1936	1905	ν C-C (85)
11	1880(vs)		1816	1793	ν C-C (84)
12	1750(vw)		1771	1614	ν C-C (82)
13	1700(w)		1706	1607	bC-H (81)
14	1560(vs)		1583	1576	bC-H (83)
15			1545	1523	bC-C (78)
16		1480(s)	1487	1462	bC-C (73)
17			1455	1424	bC-H (71)
18		1425(vw)	1436	1408	bC-H (73)
19	1380(m)		1398	1369	ν C-N (76)
20		1310(vw)	1385	1360	bC-C (69)
21		1230(s)	1280	1243	CF3 sym (89)
22	1215(vw)		1219	1207	CF3 asym (83)
23	1180(m)	1200(w)	1202	1193	CF3 asym (84)
24		1115(m)	1115	1085	bC-C (78)
25			1068	1047	bC-C (67)
26		1010(vw)	1015	980	bC-N (61)
27		980(m)	986	963	ω C-H (59)
28	915(m)		917	909	ω C-H (53)
29			855	835	ω C-N (60)
30			851	784	ω C-H (62)
31	820(m)		836	761	ω C-H (65)
32			802	723	ω C-C (57)
33		750(vs)	735	708	ω C-C (55)

Table 1. Continued

Sl. No.	Observed frequencies (cm ⁻¹)		Calculated frequencies (cm ⁻¹)		Assignments with > 10% of TED
	FT-IR	FT-Raman	B3LYP		
			Unscaled	Scaled	
34	450(s)	590(m)	620	608	bCF3 ipb (58)
35			583	571	bCF3 opb (53)
36			429	411	bCF3 sb (54)
37		250(s)	362	345	bCF3 ipr (52)
38			334	321	bCF3 opr (50)
39			200	168	ω C-C (47)
40		220(m)	193	172	ω C-C (43)
41			185	163	ω C-C (49)
42			162	145	tCF3 (44)

FT-IR spectrum is assigned to C–N stretching vibration for 4TFMNB. The in-plane and out-of-plane vibrations of the title compound are occurred within the characteristics region.

4.1.4 CF3 vibrations

Usually symmetric and antisymmetric CF3 stretching vibrations are in the ranges 1290–1235 and 1226–1200 cm⁻¹, respectively [25, 26]. Therefore the band located at 1230 cm⁻¹ in FT-Raman spectrum is assigned to symmetric stretching vibrations. CH₃ asymmetric frequencies are assigned at 1215, 1180 cm⁻¹ and 1200 cm⁻¹ in FT-IR and FT-Raman spectra for title compound. CF3 deformations usually occur in regions 690–631 cm⁻¹, 640–580 cm⁻¹ and 570–510 cm⁻¹ [21]. Accordingly CF3 opb is identified at 590 cm⁻¹ in FT-Raman spectrum. CF3 rocking vibrational frequency ranges at 460–350 cm⁻¹ [27]. In the present study, the CF3 opr band observed at 250 cm⁻¹ in FT-Raman spectrum.

5 NMR Spectral Analysis

The ¹³C and ¹H NMR chemical shifts are calculated within gauge independent atomic orbital (GIAO) method using B3LYP /6-311+G(d,p) method. A comparison of the experimental and theoretical NMR spectra can be very useful in making correct assignments and understanding the basic chemical shift molecular structure relationship. The experimental ¹³C and ¹H NMR spectra of the title compound are given in Figure 4 ((a) ¹³C and (b) ¹H)). In Table 2, reported the detailed experimental and theoretical ¹³C and ¹H isotropic chemical shifts values in ppm for the present compound are given. The ¹³C chemical shift statistics

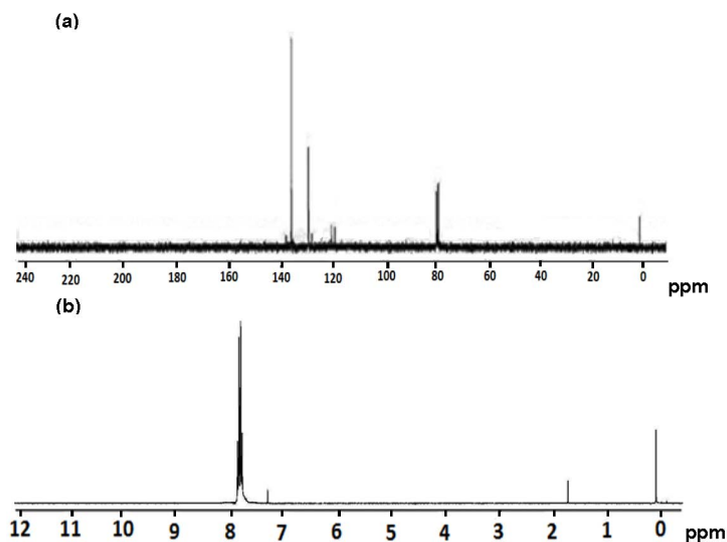


Figure 4. The experimental (a) ^{13}C and (b) ^1H NMR spectra of 4TFMNB.

for all calculations have the range from 121 to 72 ppm at B3LYP/6-311+G(d,p) level in average for 4TFMNB. Downfield and upfield chemical shift values observed at 118 and 68 ppm of carbon atoms C2 and C1 are due to electron donating effect of nitrile group. The carbon atoms C3, C4, C5, C7 and C11 are significantly observed in the upfield with chemical shift values of 114, 108, 110, 102 and 112 ppm, respectively, which well-known show that the influences of the electronegative nitrile and trifluoromethyl group atoms are negligibly small and their signal are observed in the normal region.

The ^1H chemical shift values for all calculations have 7.53 from 4.32 ppm at B3LYP/6-311+G(d,p) method in the average for 4TFMNB molecule. As can be seen from Table 2, theoretical ^{13}C and ^1H chemical shift results of the title com-

Table 2. Experimental and theoretical ^{13}C and ^1H chemical shift (ppm) of 4TFMNB.

Atoms	Theoretical shift (ppm)	Experimental shift (ppm)	Atoms	Theoretical shift (ppm)	Experimental shift (ppm)
C1	72	68	C7	103	102
C2	121	118	C11	105	112
C3	115	114	H9	7.53	6.8
C4	109	108	H10	6.65	6.0
C5	111	110	H15	7.23	6.4
C6	93	—	H16	4.32	3.4

pound are generally closer to the corresponding experimental chemical shift data except for C6 atom. The small shifts can be explained in detail as a consequence of the change in the molecular environment.

6 Conclusions

The compound 4TFMNB was studied by various analytical techniques such as FT-IR, FT-Raman and NMR (^{13}C and ^1H). The specific interpretations of the vibrational spectra had been accomplished for title molecule. The observed wavenumbers are found to be in good agreement with the theoretically calculated values. The ^{13}C and ^1H NMR chemical shifts results seemed to be in a good agreement with experimental data.

References

- [1] P.S. Nikam, L. Shirsat, M. Hasan (1998) *J. Chem. Eng. Data* **43** 732.
- [2] J.S. Sandhu, A. Singh (1992) *J. Chem. Thermodyn.* **24** 81.
- [3] M.I. Aralguppi, C.V. Jadar, T.M. Aminabhavi (1999) *J. Chem. Eng. Data* **44** 216.
- [4] P.S. Nikam, B.S. Jagdale, A.B. Sawant, M.J. Hasan (2000) *Chem. Eng. Data* **45** 214.
- [5] M.J. Frisch, A.B. Nielsen, A.J. Holder (2009) “*Gaussview user’s manual*” (Gaussian Inc., Pittsburgh, PA).
- [6] A.D. Becke (1993) *J. Chem. Phys.* **98** 5648.
- [7] C. Lee, W. Yang, R.C. Parr (1998) *Phys. Rev. B* **37** 785.
- [8] T. Sundius (2002) *Vib. Spectrosc.* **29** 89.
- [9] T. Sundius (2002) MOLVIB (V.7.0): Calculation of Harmonic Force Fields and Vibrational Modes of Molecules, QCPE Program No. 807.
- [10] D.C. Young (2001) “*Computational Chemistry: A Practical guide for applying Techniques to Real world Problems (Electronic)* (John Wiley and Sons Inc., New York).
- [11] N. Sundaraganesan, S. Illakiamani, H. Saleem, P.M. Wojciechowski, D. Michalska (2005) *Spectrochim. Acta A* **61** 2995.
- [12] G. Keresztury, J.M. Chalmers, P.R. Griffiths (2002) “*Raman Spectroscopy: Theory in Hand Book of Vibrational Spectroscopy* (John Wiley & Sons Ltd., New York).
- [13] G. Keresztury, S. Holly, G. Besenyeyi, J. Varga, A.Y. Wang, J.R. Durig (1993) *Spectrochim. Acta A* **49** 2007.
- [14] M. Karabacak, D. Karagoz, M. Kurt (2009) *Spectrochim. Acta A* **61** 2995.
- [15] N. Sundaraganesan, S. Kalaichelvan, C. Meganathan, B. Dominic Joshua, J. Cornard (2008) *Spectrochim. Acta A* **71** 898.
- [16] R. Senthil Kumar, N. Marudhaiveeran (2015) *Elixir Vib. Spec.* **87** 35682.
- [17] Y.R. Sharma (1994) “*Elementary Organic Spectroscopy – Principles and Chemical Applications*” (S. Chand & Company Ltd., New Delhi 92).
- [18] G. Vijayakumar, N. Saravanan (2016) *Elixir Comp. Chem.* **93** 39522.

D. Gnanasoundar Iraiyan, S. Joseph Vedhagiri, M. Govindarajan, K. Parimala

- [19] L. Vien, N.B. Colthup, W.G. Fateley, J.G. Grassell (1991) “*The Handbook of Infrared Raman Characteristic Frequencies of Organic Molecules*” (Academic Press, Boston, MA).
- [20] E. Gladis Anitha, S. Joseph Vedhagiri, K. Parimala (2015) *Spectrochim. Acta A* **136** 1557-1568.
- [21] D.N. Sathyanarayana (2004) “*Vibrational Spectroscopy – Theory and Applications*”, Second ed. (New Age International (P) Limited Publishers, New Delhi).
- [22] G. Socrates (2001) “*Infrared and Raman, Characteristics Group Frequencies*”, 3rd edition (Wiley, New York).
- [23] V.R. Dani (1995) “*Organic Spectroscopy*” (Tata-McGraw Hill Publishing Company, New Delhi).
- [24] R.M. Silverstein, G. Clayton Basseler, T.C. Morrill (1991) “*Spectrometric identification of Organic Compounds*” (John Wiley and Sons, New York).
- [25] V. Balachandran, K. Parimala (2013) *Spectrochim. Acta A* **102** 30-51.
- [26] L.E. Fernandez, A. Benaltabef, A. Navarro, M. Fernandez Gomez, E.L. Varetti (2000) *Spectrochim. Acta A* **56** 1101.
- [27] E.L. Fernandez, A. Ben Altabet, E.L. Varetti (2002) *J. Mol. Struct.* **785** 612.