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Structural, electronic and optical properties of 2,5-dichloro-p-xylene: experimental and theoretical calculations using DFT method†

G. Venkatesh,*a M. Govindaraju,b C. Kamal,c P. Vennilad and S. Kayae

The vibrational spectra including FT-IR and FT-Raman for 2,5-dichloro-p-xylene (DCPX) have been recorded. All the energies, the geometry of the compound and vibrational frequencies of (DCPX) have been calculated using density functional theory (DFT/B3LYP) method coupled with 6-31G* and 6-311+G** functionals. The occurrence of charge transfer was known from the HOMO-LUMO energy gap of DCPX. In order to find out the electronic excitation energies, oscillator strength and nature of the respective excited states, the closed-shell singlet calculation method has been utilized for DCPX. The values of the total dipole moment (μ) and the first order hyperpolarizability (β) of DCPX have been investigated. The non-linear optical (NLO) behavior of DCPX has been identified from the results of hyperpolarizability values.

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1 Introduction

Xylene is an aromatic hydrocarbon which has two methyl groups substituted in benzene at various positions. Xylene is a major petrochemical produced by the carbonisation of coal in the manufacturing of coke fuel, which represents about 0.5–1% of crude oil and is established in a small amount in gasoline and aircraft fuels. p-Xylene is a benzene based aromatic hydrocarbon having two substituted methyl groups with the molecular formula C_8H_{10} or $C_6H_4(CH_3)_2$. The position of the two methyl groups in the aromatic ring has been identified from the symbol "p" which revealed it as para position. Xylene is used on a large scale for the manufacture of terephthalic acid for polyester.¹⁻⁴

Xylene is the principal precursor to terephthalic acid and dimethyl terephthalate, both are used in the production of polyethylene terephthalate (PET) plastic bottles and polyester clothing. It is widely used as a solvent in the leather, rubber and printing industries. Xylene is used as a sealer/carrier solvent in acrylic concrete which is a widely used concrete protector.⁵⁻⁹ Arjunan *et al.*, examined the vibrational properties of 4-nitro

and 5-nitro-m-xylene.10 Govindarajan and his co-workers have

True to the best of our knowledge, the structural features and vibrational analysis of 2,5-dichloro xylene by DFT methods have not been reported, so far. Thus, considering the industrial importance of xylene derivatives, the experimental and theoretical calculation of DCPX using DFT method with the B3LYP (Becke's three-parameter (B3) Lee-Yang-Parr's (LYP) correlation exchange in functional) functional has been reported in the present study with a view to explore the structural characteristics.14,15 Quantum chemical computational methods have been proved to be very significant in interpreting and assuming the vibrational spectra, recent past. For instance, a complete vibrational and molecular structure analysis of 2-methoxy-5nitrophenol and 2-methoxy-4-methylphenol isomers has been evaluated using quantum mechanical approach by HF and DFT calculations by Meenakshi.16 Spectroscopic, single crystal XRD structure, DFT and molecular dynamics investigation of Sheena mary et al., examined 1-(3-chloro-4-fluorophenyl)-3-[3-(trifluoromethyl)phenyl]thiourea using DFT for it molecular dynamics.17 DFT has also been applied to examine the structural features, NLO properties of heterocyclic 5-nitro-1,3benzodioxole and its applications on solar cells.18

A significant development has been achieved by combining the empirical method with DFT information in the scaled quantum mechanical (SQM) force field method. In SQM approach, the deviations of the computed harmonic force field are rectified using some scale factors which are found to be well

reported vibrational, spectral and NLO properties of 2-chloroxylene. Chemical properties and anticancer activity of $\alpha, \alpha, \alpha', \alpha'$ tetra bromo-p-xylene were reported by M. Manzoor ali and his co-workers. Arivazhagan *et al.*, reported the vibrational spectroscopic studies of 4-bromo-p-xylene.

^aR&D Centre, Bharathiar University, Coimbatore-641 046, India. E-mail: venkateshindhuja@gmail.com; Tel: +91 9965285541

^bDepartment of Chemistry, Arignar Anna Government Arts College, Namakkal-638 052,

Department of Chemistry, VSA Group of Institutions, Salem, Tamil Nadu-636010, India

^dDepartment of Chemistry, Thiruvalluvar Government Arts College, Rasipuram-638 052, India

Department of Chemistry, Cumhuriyet University, Sivas-58140, Turkey

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transferable between chemically related molecules and they were preferred to use. The calculated IR and Raman vibrational spectral assessment of DCPX were also simulated using the computed dipole moments for IR intensities and polarizabilities for Raman intensities.

2 Experimental details

DCPX as finely powdered solid was obtained from Lancaster Chemical Company of UK. The FT-IR spectra of DCPX have been recorded using KBr pellet technique in the region 4000–400 cm $^{-1}$ with a resolution of 1 cm $^{-1}$ using BRUKER IFS 66 V spectrophotometer coupled with He–Ne laser source at room temperature. The FT-Raman spectra has also been recorded on a same instrument equipped with FRA-106 FT-Raman accessory in 4000–100 cm $^{-1}$ stoke region with 1064 nm line of a Nd:YAG Laser for excitation which is operating at 200 mW power. The minimal accuracy of wave numbers was assumed to be $\pm 1~{\rm cm}^{-1}$.

3 Computational details

To provide the complete information regarding the molecular structural characteristics and the fundamental vibrational modes of DCPX, the restricted DFT-B3LYP correlation functional calculations have been carried out.14,15 The calculations of geometrical parameters in the ground state were performed using the GAUSSIAN 09W programs. The geometry optimization was carried out using the initial geometry generated from standard geometrical parameters and B3LYP methods adopting 6-31G*/6-311+G** functional to characterize all stationary points as minima. The optimized structural parameters of the compound were used for harmonic vibrational frequency calculations resulting in IR and Raman frequencies together with intensities and Raman depolarization ratios. In DFT methods, Becke's three parameter exchange-functional (B3) jointed with gradientcorrected correlation functional of Lee, Yang and Parr (LYP) by employ the split-valence polarized 6-311+G** functional have been developed for the computation of molecular structure optimization, vibrational frequencies and energies of the molecular optimized structures.

The vibrational modes were assigned by means of visual inspection using the GAUSSVIEW program. The symmetry of the vibrational modes was determined by standard procedure of decomposition of the traces of the symmetry operations into the irreducible representations. The derivation (given below) obtained from the theory of Raman scattering has been utilized to convert the Raman activities (S_i) calculated by GAUSSIAN 09W program into relative Raman intensities (I_i)

$$I_{i} = \frac{f(v_{0} - v_{i})^{4} S_{i}}{v_{i} [1 - \exp(-hcv/KT)]}$$
(1)

where, v_0 and v_i are exciting frequency (in cm⁻¹) and vibrational wavenumber of the normal mode, respectively; h, c & k represent universal constants and f stands for common normalization factor chosen for all peak intensities.

4 Results and discussion

4.1. Geometrical parameters

DCPX contains two methyl groups at para position and two chlorine atoms in aromatic benzene ring. The optimized molecular structure has been obtained from Gaussion09W program.19 The non linear molecular structure of DCPX illustrates C_s symmetry (Fig. 1). The global minimum energy has been obtained by the DFT molecular optimization structure and is given in Table S1 ESI.† The obtained energies are as follows -1229.95 (Hartee) B3LYP/6-31G*, -1230.07 (Hartee) by B3LYP/6-311G** for gaseous phase and -1229.54 (Hartee) by B3LYP/6-311G** -1230.09 (Hartee) aqueous phase. Table 1 compares the theoretically calculated bond lengths, bond angles and dihedral angles with X-ray data values. The optimized bond lengths are in the order as follows C2-C3 < C5-C6 < C1-C2 < C4-C5 < C1-C6 < C3-C4. It is clear from Table 1 that, the carbons substituted by methyl groups have larger bond lengths and the carbons substituted by halogens possess lower bond lengths when compared to all other ring atoms. 10,12,20 This could be due to the inductive effect of methyl group and chlorine atoms as well as resonance effect of phenyl ring (substitution effect). +I effect of methyl group pushed out the electrons towards the phenyl ring. But, the -I effect of chlorine atoms and resonance effect of phenyl ring competed with each other to pull the electrons towards themselves. These effects resulted in larger bond lengths of carbons substituted by methyl group meanwhile shorter bond lengths of carbons substituted by halogen atoms.21,22 From the bond lengths, it is clearly known that, the aromatic ring is slightly distorted from perfect hexagonal structure. It could be due to the

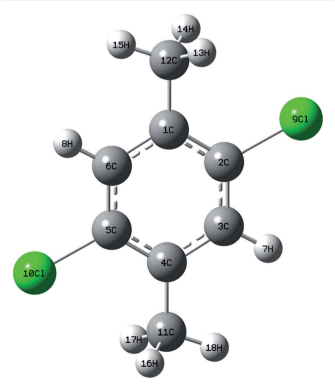


Fig. 1 Optimized molecular structure of DCPX.

Table 1 Optimized geometrical parameters of DCPX obtained by B3LYP/6-31G*, B3LYP/6-311G** methods in gaseous phase and B3LYP/6-311G** method in aqueous phase using DFT calculations

Gaseous phase			Aqueous phase			
B3LYP/6-31G*		B3LYP/6-311G**		B3LYP/6-311G**		V was d-4
Bond length	Value (Å)	Bond length	Value (Å)	Bond length	Value (Å)	X-ray da values (Å
C_1 – C_2	1.389	C_1 – C_2	1.396	C ₁ -C ₂	1.396	1.399
C ₁ -C ₆	1.398	C_1 – C_6	1.403	C_1 – C_6	1.402	1.399
C ₁ -C ₁₁	1.508	C_1 - C_{11}	1.507	C_1-C_{11}	1.507	1.511
C_2 – C_3	1.381	C_2 – C_3	1.387	C_2 – C_3	1.386	1.399
C_2 - Cl_9	1.831	C_2 – Cl_9	1.835	C_2 - Cl_9	1.836	1.771
C ₃ -C ₄	1.406	C_3 - C_4	1.405	C_3 – C_4	1.405	1.399
C_3 – H_7	1.083	C_3 – H_7	1.080	C_3 - H_7	1.080	1.098
C_4 - C_5	1.391	C_4 - C_5	1.398	C_4 – C_5	1.397	1.399
C_4 - C_{12}	1.508	C_4 - C_{12}	1.507	C_4 - C_{12}	1.507	1.511
$C_5 - C_6$	1.383	$C_5 - C_6$	1.389	C_5 - C_6	1.388	1.399
C ₅ -Cl ₁₀	1.831	C_5 - Cl_{10}	1.835	C_5 – Cl_{10}	1.836	1.771
C ₆ -H ₈	1.083	C_6 - H_8	1.080	C_6-H_8	1.080	1.098
C ₆ 11 ₈ C ₁₁ -H ₁₃	1.096	C ₁₁ -H ₁₃	1.092	C_{11} - H_{13}	1.092	1.120
	1.096		1.092	C_{11} - H_{13} C_{11} - H_{14}	1.092	1.120
C ₁₁ -H ₁₄		C ₁₁ -H ₁₄				
C ₁₁ -H ₁₅	1.093	C ₁₁ -H ₁₅	1.090	C ₁₁ -H ₁₅	1.088	1.120
C ₁₂ -H ₁₆	1.096	C ₁₂ -H ₁₆	1.092	C ₁₂ -H ₁₆	1.092	1.120
C ₁₂ -H ₁₇	1.093	C ₁₂ -H ₁₇	1.090	C ₁₂ -H ₁₇	1.088	1.120
C ₁₂ -H ₁₈	1.096	C ₁₂ -H ₁₈	1.092	C ₁₂ -H ₁₈	1.092	1.120
Dond anala	Value (°)	Bond angle	Value (°)	Dand angle	Value (°)	X-ray da values (
Bond angle	Value (°)	Bond angle	value ()	Bond angle	value ()	values (
C ₂ -C ₁ -C ₆	117.130	C_2 - C_1 - C_6	117.059	$C_2-C_1-C_6$	117.124	118.6
C_2 - C_1 - C_{11}	120.945	C_2 – C_1 – C_{11}	120.008	C_2 - C_1 - C_{11}	120.346	_
$C_6 - C_1 - C_{11}$	120.925	C_6 - C_1 - C_{11}	120.983	C_6 – C_1 – C_{11}	120.930	_
$C_1 - C_2 - C_3$	121.574	$C_1 - C_2 - C_3$	121.691	C_1 - C_2 - C_3	121.683	120.0
C_1 – C_2 – Cl_9	119.333	C_1 – C_2 – Cl_9	119.281	C_1 – C_2 – Cl_9	119.063	_
C_3 – C_2 – Cl_9	118.623	C_3 – C_2 – Cl_9	117.528	C_3 – C_2 – Cl_9	117.255	_
$C_2 - C_3 - C_4$	119.326	C_2 – C_3 – C_4	119.320	C_2 - C_3 - C_4	119.393	120.0
C_2 – C_3 – H_7	119.740	C_2 – C_3 – H_7	119.746	C_2 - C_3 - H_7	120.123	_
C_4 – C_3 – H_7	119.433	C_4 – C_3 – H_7	119.455	C_4 – C_3 – H_7	119.283	_
$C_3 - C_4 - C_5$	117.130	C_3 – C_4 – C_5	117.069	$C_3 - C_4 - C_5$	111.124	120.0
$C_3-C_4-C_{12}$	120.925	C_3 - C_4 - C_{12}	120.983	$C_3-C_4-C_{12}$	120.930	_
$C_5 - C_4 - C_{12}$	120.945	C_5 – C_4 – C_{12}	120.008	C_5 - C_4 - C_{12}	120.346	_
$C_4 - C_5 - C_6$	119.044	C_4 - C_5 - C_6	119.191	$C_4-C_5-C_6$	119.683	120.0
C_4 - C_5 - Cl_{10}	119.333	C_4 – C_5 – Cl_{10}	119.281	$C_4-C_5-Cl_{10}$	119.063	_
C ₆ -C ₅ -Cl ₁₀	118.623	$C_6-C_5-Cl_{10}$	118.528	$C_6 - C_5 - Cl_{10}$	118.255	_
C_1 - C_6 - C_5	119.326	C_1 - C_6 - C_5	119.390	C_1 - C_6 - C_5	119.393	120.0
C_1 - C_6 - H_8	119.433	C_1 - C_6 - H_8	119.455	$C_1 - C_6 - H_8$	119.283	_
$C_5 - C_6 - H_8$	119.740	$C_5 - C_6 - H_8$	119.746	$C_5 - C_6 - H_8$	120.123	_
C_1 - C_{11} - H_{13}	110.273	C_1 - C_{11} - H_{13}	111.307	C_1 - C_{11} - H_{13}	111.271	_
C_1 - C_{11} - H_{14}	111.271	C_1 - C_{11} - H_{14}	111.304	C_1 - C_{11} - H_{14}	111.270	_
$C_1 - C_{11} - H_{15}$	110.763	C_1 $-C_{11}$ $-H_{15}$	110.744	C_1 - C_{11} - H_{15}	110.706	_
H ₁₃ -C ₁₁ -H ₁₄	109.851	$H_{13}-C_{11}-H_{14}$	109.906	$H_{13}-C_{11}-H_{14}$	109.101	_
H ₁₃ -C ₁₁ -H ₁₄ H ₁₃ -C ₁₁ -H ₁₅	108.258	$H_{13}-C_{11}-H_{14}$ $H_{13}-C_{11}-H_{15}$	108.206	H_{13} - C_{11} - H_{15}	108.165	_
		$H_{13}-C_{11}-H_{15}$ $H_{14}-C_{11}-H_{15}$				
H ₁₄ -C ₁₁ -H ₁₅	108.268		108.215	$H_{14}-C_{11}-H_{15}$	108.175	_
C ₄ -C ₁₂ -H ₁₆	111.271	C_4 - C_{12} - H_{16}	111.304	C_4 - C_{12} - H_{16}	111.270	_
C ₄ -C ₁₂ -H ₁₇	110.763	C_4 - C_{12} - H_{17}	110.744	C_4 - C_{12} - H_{17}	110.706	_
C ₄ -C ₁₂ -H ₁₈	111.273	C_4 - C_{12} - H_{18}	111.307	C_4 - C_{12} - H_{18}	111.271	
H ₁₆ -C ₁₂ -H ₁₇	108.268	H_{16} - C_{12} - H_{17}	108.215	$H_{16}-C_{12}-H_{17}$	108.175	_
H ₁₆ -C ₁₂ -H ₁₈	107.851	H ₁₆ -C ₁₂ -H ₁₈	107.906	H_{16} - C_{12} - H_{18}	107.101	_
H ₁₇ -C ₁₂ -H ₁₈	108.258	H ₁₇ -C ₁₂ -H ₁₈	108.206	H ₁₇ -C ₁₂ -H ₁₈	108.165	
Dihedral angle	Value (°)	Dihedral angle	Value (°)	Dihedral angle	Value (°)	
$C_6 - C_1 - C_2 - C_3$	0.013	C_6 - C_1 - C_2 - C_3	0.025	C_6 - C_1 - C_2 - C_3	0.016	_
0 0 0 0	170.001	C C C C	170.000	C C C C	170.000	
C_6 - C_1 - C_2 - Cl_9	-179.981	C_6 – C_1 – C_2 – Cl_9	-179.990	C_6 – C_1 – C_2 – Cl_9	-179.980 179.999	

Table 1	(Contd.)	١
I able 1	COITEU.	,

Dihedral angle	al angle Value (°)		Value (°)	Dihedral angle	Value (°)	_	
C_{11} – C_1 – C_2 – Cl_9	-0.003	C_{11} – C_{1} – C_{2} – Cl_{9}	-0.008	C_{11} - C_{1} - C_{2} - Cl_{9}	-0.001	_	
C_2 - C_1 - C_6 - C_5	-0.014	C_2 - C_1 - C_6 - C_5	-0.014	C_2 - C_1 - C_6 - C_5	-0.013	_	
C_2 - C_1 - C_6 - H_8	-179.992	C_2 - C_1 - C_6 - H_8	-180.000	C_2 – C_1 – C_6 – H_8	-180.000	_	
C_{11} – C_{1} – C_{6} – C_{5}	-179.993	C_{11} - C_{1} - C_{6} - C_{5}	-180.000	C_{11} – C_{1} – C_{6} – C_{5}	-180.000	_	
C_{11} – C_{1} – C_{6} – H_{8}	0.013	C_{11} – C_{1} – C_{6} – H_{8}	0.011	C_{11} – C_1 – C_6 – H_8	0.014	_	
C_2 - C_1 - C_{11} - H_{13}	-59.519	C_2 - C_1 - C_{11} - H_{13}	-59.577	C_2 - C_1 - C_{11} - H_{13}	-59.687	_	
C_2 - C_1 - C_{11} - H_{14}	59.520	C_2 - C_1 - C_{11} - H_{14}	59.575	C_2 - C_1 - C_{11} - H_{14}	59.665	_	
C_2 - C_1 - C_{11} - H_{15}	-179.993	C_2 - C_1 - C_{11} - H_{15}	-180.000	C_2 - C_1 - C_{11} - H_{15}	-180.010	_	
C_6 - C_1 - C_{11} - H_{13}	120.459	$C_6-C_1-C_{11}-H_{13}$	120.405	$C_6-C_1-C_{11}-H_{13}$	120.295	_	
C_6 - C_1 - C_{11} - H_{14}	-120.502	$C_6-C_1-C_{11}-H_{14}$	-120.440	$C_6-C_1-C_{11}-H_{14}$	-120.350	_	
C_6 - C_1 - C_{11} - H_{15}	-0.016	C_6 - C_1 - C_{11} - H_{15}	-0.014	C_6 - C_1 - C_{11} - H_{15}	-0.023	_	
C_1 - C_2 - C_3 - C_4	0.001	C_1 – C_2 – C_3 – C_4	-0.012	$C_1-C_2-C_3-C_4$	-0.004	_	
C_1 - C_2 - C_3 - H_7	-180.005	C_1 - C_2 - C_3 - H_7	-180.020	C_1 - C_2 - C_3 - H_7	-180.010	_	
$Cl_9-C_2-C_3-C_4$	179.995	$Cl_9-C_2-C_3-C_4$	180.005	$Cl_9-C_2-C_3-C_4$	179.997	_	
Cl ₉ -C ₂ -C ₃ -H ₇	-0.011	$Cl_9-C_2-C_3-H_7$	-0.004	$Cl_9-C_2-C_3-H_7$	-0.013	_	
C_2 - C_3 - C_4 - C_5	-0.014	C_2 - C_3 - C_4 - C_5	-0.014	C_2 - C_3 - C_4 - C_5	-0.013	_	
C_2 - C_3 - C_4 - C_{12}	-179.993	C_2 - C_3 - C_4 - C_{12}	-180.000	C_2 - C_3 - C_4 - C_{12}	-180.000	_	
H_7 - C_3 - C_4 - C_5	179.992	$H_7-C_3-C_4-C_5$	-180.000	$H_7-C_3-C_4-C_5$	-180.000	_	
H_7 - C_3 - C_4 - C_{12}	0.013	$H_7-C_3-C_4-C_{12}$	0.011	H_7 - C_3 - C_4 - C_{12}	0.014	_	
C_3 - C_4 - C_5 - C_6	0.013	C_3 - C_4 - C_5 - C_6	0.025	$C_3-C_4-C_5-C_6$	0.016	_	
C_3 - C_4 - C_5 - Cl_{10}	-179.981	C_3 - C_4 - C_5 - Cl_{10}	-179.990	$C_3-C_4-C_5-Cl_{10}$	-179.980	_	
C_{12} – C_4 – C_5 – C_6	179.992	C_{12} – C_4 – C_5 – C_6	180.009	C_{12} – C_4 – C_5 – C_6	179.999	_	
C_{12} – C_4 – C_5 – Cl_{10}	-0.003	C_{12} – C_4 – C_5 – Cl_{10}	-0.008	C_{12} – C_4 – C_5 – Cl_{10}	-0.001	_	
C_{12} – C_4 – C_5 – H_{16}	-120.502	C_{12} – C_4 – C_5 – H_{16}	-120.440	C_{12} – C_4 – C_5 – H_{16}	-120.350	_	
C_3 - C_4 - C_{12} - H_{17}	-0.016	C_3 - C_4 - C_{12} - H_{17}	-0.014	C_3 - C_4 - C_{12} - H_{17}	-0.023	_	
C_3 - C_4 - C_{12} - H_{18}	120.459	C_3 - C_4 - C_{12} - H_{18}	120.405	C_3 - C_4 - C_{12} - H_{18}	120.295	_	
C_5 - C_4 - C_{12} - H_{16}	59.520	C_5 - C_4 - C_{12} - H_{16}	59.575	C_5 - C_4 - C_{12} - H_{16}	59.665	_	
C_5 - C_4 - C_{12} - H_{17}	-179.993	C_5 - C_4 - C_{12} - H_{17}	-180.000	C_5 - C_4 - C_{12} - H_{17}	-180.010	_	
C_5 - C_4 - C_{12} - H_{18}	-59.519	C_5 - C_4 - C_{12} - H_{18}	-59.577	C_5 - C_4 - C_{12} - H_{18}	-59.687	_	
C_4 - C_5 - C_6 - C_1	0.001	C_4 - C_5 - C_6 - C_1	-0.012	$C_4-C_5-C_6-C_1$	-0.004	_	
C_4 - C_5 - C_6 - H_8	-180.005	C_4 - C_5 - C_6 - H_8	-180.020	C_4 - C_5 - C_6 - H_8	-180.010	_	
Cl ₁₀ -C ₅ -C ₆ -C ₁	179.995	Cl_{10} - C_5 - C_6 - C_1	180.005	Cl_{10} - C_5 - C_6 - C_1	179.997	_	
$\text{Cl}_{10}\text{-C}_5\text{-C}_6\text{-H}_8$	-0.011	Cl_{10} – C_5 – C_6 – H_8	-0.004	Cl_{10} – C_5 – C_6 – H_8	-0.013	_	

substitutions of halogens and methyl groups in *para* position. The calculated bond lengths for C_1 – C_{12} & C_4 – C_{12} of about 1.51 Å, the methyl C–H in the range 1.09–1.10 Å and ring C–H as 1.08 Å are correlated well with the standard values. But, the bond length of C–Cl observed as 1.83 Å is higher than the standard values.

In addition, $C_2-C_3-C_4$, $C_1-C_6-C_5$ (119.3°), $C_2-C_1-C_6 \otimes C_3-C_4-C_5$ (117.1°) bond angles are reduced than that of a benzene ring, but the $C_1-C_2-C_3$ (121.6°) bond angle is enlarged. The dihedral angles obtained are represented in Table 1. From the Table, it can be seen that, the dihedral angels $C_6-C_1-C_2-C_3$, $C_2-C_1-C_6-C_5$ and $C_1-C_2-C_3-C_4$ are supported the planer nature of aromatic ring. The optimization geometry of DCPX showed that, two substituted methyl groups are out-of-plane of the ring. $^{20-22}$ This has been evidenced from the dihedral angels of $C_2-C_1-C_{11}-H_{14}\approx 60^\circ$, $C_6-C_1-C_{11}-H_{13}\approx 120^\circ$ and $C_5-C_4-C_{12}-H_{16}\approx 60^\circ$, $C_3-C_4-C_{12}-H_{18}\approx 120^\circ$. Similarly other two $C_{11}-H_{15}$ and $C_{12}-H_{17}$ are in-planebending of ring which has been known from the following dihedral angles $C_6-C_1-C_{11}-H_{15}\approx 0^\circ$ and $C_3-C_4-C_{12}-H_{17}\approx 0^\circ$.

4.2. Assignments of spectra

Normal coordinate analysis provides a detailed description of vibrational modes. For this purpose, the full set of 62 standard internal coordinates containing 14 redundancies have been calculated and listed in Table S2 of ESI.† From these, a non-redundant set of local symmetry coordinates was constructed by appropriate linear combinations of internal coordinates according to Pulay *et al.*²³ and they are presented in Table S3 of ESI.† DFT force fields calculated theoretically have been transformed as a set of vibrational coordinates and then used for all subsequent calculations. The fundamental modes of DCPX in terms of vibrational assignments associated with theoretical IR, Raman intensities and normal mode descriptions (characterized by PED) are listed in Table 2.

When using computational methods to predict theoretical normal vibrations for relatively complex polyatomic molecules, scaling strategies are used to bring computed wave numbers. The observed FT-IR and FT-Raman spectra of the DCPX were presented in Fig. 2 and 3. The 48 normal modes of DCPX are distributed among the symmetry species as $\Gamma_{\rm 3N-6}=33$ A' (inplane) + 15 A" (out-of-plane). The detailed vibrational assignments of fundamental modes of DCPX have been reported in Table S3.† Root mean square (RMS) values of frequencies were obtained in the study using the following expression,

$$RMS = \sqrt{\frac{1}{n-1} \sum_{i}^{n} \left(U_{i}^{\text{calc}} - U_{i}^{\text{exp}} \right)^{2}}$$
 (2)

The RMS frequency error is being 101 cm⁻¹. Earlier report revealed that other methods than B3LYP such as HF showed more deviation in experimental and calculated values for substituted xylene derivative. Hence, B3LYP method has been adopted in the present study to calculate vibrational frequencies for DCPX according to Govindarajan et al.11 Systematic assignment is produced using the unscaled computed frequencies. Then for comparing the theoretical values, with the calculated frequencies, they were multiplied by a common scale factor of less than one to reduce the overall deviation. Pulay²⁴ recommended a set of transferable coordinates to refine the scale factors. The least square refinement of the scale factors led to a mean deviation of 7.09 cm⁻¹ for DCPX. The least square refinement of results relies on the frequencies observed and on the initial band assignments. The uniform scaling force field does not change the nature of calculated intensity patterns of simulated spectra, because it does not changes the vibrational modes. The selective scaling changes the frequencies in addition to changing the normal modes of vibrations. The changes in relative band intensities were caused by the changes in PED values which are corresponding to vibrational vectors.

4.2.1. C-H vibrations. Regularly the bands in the range 3300-3000 cm⁻¹ are assigned to C-H stretching vibrations of aromatic compounds.25 The C-H vibrations of the DCPX are observed at 3249 cm⁻¹ in FT-IR spectrum and 3239 cm⁻¹ in FT-Raman. The bands at 3245 cm^{-1} in FT-IR and 3235 cm^{-1} in FT-Raman from B3LYP/6-311G** method is in good agreement with observed spectral data. These are intense stretching modes and identified from the PED in Table 2. The C-H in-plane bending vibrations, generally occurs in 1000-1300 cm⁻¹ region.²⁵⁻²⁸ The FT-IR bands observed at 1225 cm⁻¹ are assigned to C-H in-plane-bending vibration, which was arrived in FT-Raman spectra at 1262 cm⁻¹ are assigned to C-H in-planebending vibrations. The bands at 1221 cm⁻¹ in FT-IR and 1257 cm⁻¹ in FT-Raman are in good agreement with observed spectral data. The C-H out-of-plane bending vibrational frequencies are mainly identified by the number of adjacent hydrogen atoms on the ring and not vastly affected by the nature of substitutions.

The C–H out-of-plane bending vibrations which are strongly coupled vibrations usually occurred in the region 1000–750 cm⁻¹.^{28,29} For DCPX, the strong bands in experimental FT-IR spectrum at 881 cm⁻¹ (80%) suggested the existence of aromatic C–H out-of-plane bending vibrations and very strong band in experimental FT-Raman spectrum at 895 cm⁻¹ (84%) confirmed the presence of aromatic C–H out-of-plane bending vibrations which are well correlated (885 cm⁻¹ and 890 cm⁻¹) with calculated vibrational spectra using B3LYP/6-311G** methods. The observation of smaller deviation of calculated vibrational frequencies with experimental vibrational frequencies validated the adoption of the above said functional in the present study. The theoretically computed frequencies are presented in Table 2.

4.2.2. C-C **vibrations.** The ring C-C and ring C=C stretching vibrations (semicircle stretching region) are generally appears in the region $1625-1400 \text{ cm}^{-1}$ and $1380-1280 \text{ cm}^{-1}$, respectively. The ring stretching vibrations are expected within

the region 1620–1390 cm⁻¹. In the present study, the absorption at 1605, 1526, 1401, 1262 cm⁻¹ (FT-Raman), 1447, 1334, 1225 cm⁻¹ (FT-IR) are assigned to C–C stretching vibrations. The FT-IR bands observed at 725 and 530 cm⁻¹ and FT-Raman at 776 and 352 cm⁻¹ are assigned to C–C in-plane and out-of-plane bending vibrations, respectively. These assignments are in quite agreement with the literature and in addition the nature and position of the substitution affected the band in this region.^{30,31} Aromatic ring vibrational modes are pretentious by the substitution in the aromatic ring.

4.2.3. C-Cl vibrations. The vibrations due to C-X (X = F, Cl)etc.) bonds formed between the ring and the halogen atoms, are quite informative since mixing of vibrations are possible due to the lowering of molecular symmetry and the presence of heavy atoms. 31-33 The C-Cl stretching frequency is generally observed in the region 800-500 cm⁻¹ which is depending on the configuration and conformation of the compound. Regarding the above information the following absorption bands in both FT-IR and FT-Raman spectra arose at 726, 479 cm⁻¹ and 775 cm⁻¹ respectively were corresponded to planar C-Cl stretching of DCPX. The planar C-Cl bending modes appeared to be relatively pure. In addition, the C-Cl out-of-plane bending vibrations were appeared at 530, 479 cm⁻¹ and 352, 295, 243 cm⁻¹ in FT-IR and FT-Raman, respectively. There is a very slight deviation in both the in-plane and out-of-plane vibrations of DCPX have been observed. After scaling of computed values for C-Cl in-plane and out-of-plane bending vibrations by B3LYP/6-311G** functional, those slight deviations have also been reduced. The observation further supported the validation of same functional in the present study.30-33

4.2.4. Methyl group vibrations. In DCPX, there are CH₃ groups substituted in first and fourth position aromatic ring. Regarding vibrational frequencies of CH₃, there is a possibility of the symmetrical (CH3 ips) and asymmetrical (CH3 ops), in-plane stretching modes (i.e. in-plane hydrogen stretching mode); deformation of symmetrical (CH3 ss), and asymmetrical (CH3 ips), the in-plane (CH3 ipr), out-of-plane (CH3 opr) rocking and twisting (t CH₃) modes. 11-13,31 Further, the asymmetric stretching (CH₃ ops) and asymmetric deformation (CH₃ opb) of CH₃ group are assumed to be depolarized for A" symmetry species. The asymmetric stretch is generally appeared as higher wave number than the symmetric stretch. The bands at 3160 ips, 3114 ops, 3054 ss cm^{-1} in FT-IR and 3158 ips, 3061 ops cm^{-1} in FT-Raman spectra are assigned to C-H asymmetric stretching vibration. We have observed the symmetrical methyl deformation mode at 1447 cm⁻¹ in FT-IR and at 1401 cm⁻¹ in FT-Raman. The absorptions appeared at 1078, 1067 cm^{-1} in FT-IR and 1081, 1065 cm^{-1} in FT-Raman are rocking vibrations of CH₃, respectively. Further, the out-of-plane bending vibrations of methyl groups are assigned at 165 cm⁻¹ FT-Raman. 11-13 The fundamental vibrations, for all the modes of CH3 group of DCPX are observed in their respective characteristic regions and they are listed in Table 2.

4.3. NMR

The isotropic chemical shifts are normally used to identify the organic reactive sites. It is important to predict exact molecular

Table 2 Vibrational assignments of experimental frequencies of DCPX along with calculated frequencies by B3LYP/6-311G** functional^a

	Symmetry of species C_s	Observed frequency (cm ⁻¹)		Calculated frequency (cm ⁻¹) with B3LYP/6- 311G** force field				(41)	
S. no.		Infrared	Raman	Unscaled	Scaled	$\operatorname{IR}^b A_{\mathrm{i}}$	Raman ^c I _i	PED (%) among type of internal coordinates ^d	
1	A'	3249		3247	3245	14.762	63.175	vCH (99)	
2	A'		3239	3237	3235	14.611	64.334	νCH (99)	
3	\mathbf{A}'	3160		3159	3157	0.004	152.4	CH_3 ips (98)	
4	\mathbf{A}'		3158	3159	3155	4.946	0.094	CH ₃ ips (98)	
5	\mathbf{A}'			3115	3111	16.428	185.539	CH ₃ ops (96)	
6	\mathbf{A}'	3114		3113	3110	20.01	152.897	CH ₃ ops (96)	
7	A'		3061	3060	3056	12.537	73.311	CH_3 ops (96)	
8	A'	3057		3058	3055	9.822	93.552	νCH ₃ (96)	
9	A'			1656	1653	0	42.075	νCC (67), bCH (12), bring (11)	
10	A'		1605	1602	1601	0.008	23.423	ν CC (53), γ CH ₃ (15), bring (8), β CH ₃ (11	
11	\mathbf{A}'	1535		1532	1530	46.449	1.068	γCH_3 (49), βCH_3 (45)	
12	\mathbf{A}'		1526	1524	1521	44.231	0.632	νCC (33), bCH (22), βCH ₃ (17), γCH ₃ (19	
13	\mathbf{A}'			1521	1518	2.961	20.565	βCH ₃ (37), γCH ₃ (49), βCH ₃ (18)	
14	A'	1515		1517	1514	8.508	14.364	βCH ₃ (91), (CH ₃)r (5)	
15	\mathbf{A}'		1510	1509	1506	3.936	17.38	γCH_3 (72), βCH_3 (22)	
16	A'			1451	1448	0.41	22.795	γCH_3 (79), νCC (7), βCH_3 (5)	
17	\mathbf{A}'	1447		1449	1446	0.271	30.536	γCH_3 (79), νCC (7), βCH_3 (5)	
18	\mathbf{A}'		1401	1399	1397	22.045	0.011	νCC (38), γCH ₃ (12), βCH ₃ (15)	
19	\mathbf{A}'	1334		1332	1329	2.69	0.004	νCC (84)	
20	\mathbf{A}'			1295	1294	0.015	0.098	bCH (72), νCC (9)	
21	\mathbf{A}'		1262	1261	1257	0	26.44	γCH (49), νCC (42), CCl (7)	
22	\mathbf{A}'	1225		1223	1221	1.6	0	γCH (37), νCC (42), bring (15)	
23	\mathbf{A}'			1092	1089	123.645	0	bring (46), vCCl (20), vCC (17), bCH (10	
24	A''		1081	1080	1078	5.982	0.113	gCC (60), CH ₃ r (23), gCH (5), βCH ₃ (5)	
25	A''	1078		1075	1070	0.706	1.022	gCC (57), CH ₃ r (24), tring (8)	
26	\mathbf{A}'	1067	1065	1062	1059	1.492	2.605	CH_3r (56), γCH_3 (13), βCH_3 (9), νCC (7)	
27	\mathbf{A}'			1024	1020	26.262	0.204	CH ₃ r (45), γCC (17), γCH ₃ (11), βCH ₃ (8	
28	Α"	954		952	949	0.001	5.648	gCCl (34), vCC (24), bring (23), vCC (16)	
29	A''		895	893	890	19.957	0.001	βCH (84), tring (8)	
30	A''	881		887	885	0.008	3.534	βCH (80), tring (14)	
31	\mathbf{A}'		775	776	774	26.255	0	γCC (57), bring (21), νCCl (13)	
32	A'	726		725	721	0	16.305	bring (58), γCC (19), νCCl (15)	
33	A''			704	702	0	0.525	tring (62), gCCl (20), gCC (15)	
34	A''		605	603	600	0.161	0	tring (35), gCCl (34), gCC (27)	
35	Α"			536	534	24.613	0	gCCl (67), bring (14)	
36	A'	530		528	525	0	5.872	βCC (46), bCC (23), βCCl (38)	
37	A'	479		478	474	0	9.65	bring (52), βCCl (7)	
38	A''			461	458	3.958	0	tring (61), gCCl (19), gCC (16)	
39	A''		352	350	348	0	1.355	βCC (47), βCCl (47)	
40	A'			331	328	0	9.239	bring (45), νCCl (43)	
41	A'		295	294	290	3.36	0	βCCl (41), bCCl (10)	
42	A"			246	244	0	2.35	tring (31), gCCl (26), gCC (20), gCH (12)	
43	A'		243	241	240	0	1.604	bCCl (28), βCC1 (32), bring (9)	
44	A'			225	220	1.339	0	bCCl ₁ (45)	
45	Α"		165	167	164	0.132	0	βCH ₃ (40), gCCl (20), tring (16), gCH (11) gCC (10)	
46	A''		153	151	147	0.001	0.568	tCH ₃ (56), gCC (7)	
47	A''			139	137	1.009	0	tCH ₃ (6), gCC (20), tring (17), tCH ₃ (16), gCCl (9), gCH (8)	
48	A''			90	88	1.352	0	tring (90)	

 $[^]a$ Abbreviations: ν – stretching, b – bending, γ – in-plane bending, β – out-of-plane bending, t – torsion, t – rocking, g – waging, ips – in-plane stretching, ops – out-of-plane stretching. b Relative absorption intensities normalized with highest peak absorption. c Relative Raman intensities calculated by eqn (1) and normalized to 100. d For the notations used see Table S3.

geometries in order to calculate the magnetic properties of molecules. ^{12,34} Hence, complete optimized geometry of DCPX was performed using TMS B3LYP/6-311+G (2d, p) and TMS HF/6-31G (d) GIAO methods in order to get better understanding in

the structural features of the title compound.¹6 The ¹H and ¹³C NMR chemical shift values of the DCPX have been calculated by Gauge Invariant Atomic Orbital (GIAO) method.¹2,35,36 The experimental and theoretical chemical shift values for ¹H and

¹³C NMR of DCPX are given in Table 3. The theoretical and experimental NMR spectra include ¹H & ¹³C for DCPX is given in Fig. S1 (ESI),† where TMS has been used as a reference. The theoretically calculated and experimentally observed chemical shifts values of DCPX in both ¹H & ¹³C NMR are reliably in good agreement with each other. Ring carbons had greater chemical shift values than the aliphatic carbons, due to the influence of the resonance effect of the aromatic ring.³³ Further, the poor electron donating nature (+I effect) of methyl group increased the chemical shift of the ring carbon (C_1 and C_4). The chlorine substituted carbon atoms have greater chemical shift than all other carbons of DCPX. This could be attributed to the electro negativity of chlorine atoms. The signals of the aromatic proton $(H_7 \& H_8)$ were observed at 5.8 ppm. The H atom is the smallest of all atoms and mostly localized on the periphery of molecules. The methyl group protons showed less chemical shift values than the aromatic ring protons. The ring protons involved in the delocalization of π electrons in the benzene ring. Among the methyl protons, of H₁₆ and H₁₄ have higher chemical shift values as they might be involved in the intra molecular hydrogen bonding (H₁₄-Cl and H₁₆-Cl).³⁷⁻⁴¹

4.4. Natural bond orbital (NBO) analysis

Natural bond orbital (NBO) analysis gives the information about nature of hydrogen bonding and interaction among bonds, and also gives information about transfer or conjugative interactions of molecular systems. 25,35,36 In addition, it gives essential information on the interactions of donor and accepter orbitals of molecules which enable the assessment of intra and intermolecular interactions. This is carried out by considering all possible interactions between filled and empty NBOs and calculating their energy importance by second order perturbation theory. For each electron donor (i) and electron acceptor (j), the stabilization energy (E^2) associates with electron delocalization between Lewis (filled) and non Lewis (unfilled) is estimated as:

$$E^{(2)} = -q_{i} \frac{\left(F_{i,j}\right)^{2}}{\varepsilon_{i} - \varepsilon_{i}} \tag{3}$$

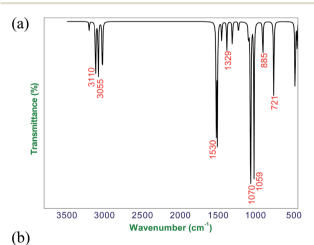
where, q_i , ε_i & ε_j are donor orbital occupancy, diagonal elements, orbital energies, respectively. F(i,j) is the off diagonal NBO Fock matrix element. Larger the stabilization energy $E_j^{(2)}$ value the higher is the interaction between electron filled (donor) and unfilled (acceptor) orbital's *i.e.*, greater the donating ability from donors to acceptor cause the greater extent of conjugation of the whole system. The Lewis structure of occupied orbitals can interact strongly. ^{12,41,42} A filled bonding orbital can act as a donor and an unfilled antibonding can act as an acceptor. Delocalization of electron density between filled Lewis (bond or lone pair) NBO orbital and unfilled non Lewis (antibond or Rydberg) NBO orbital corresponds to a stabilizing donor-acceptor interaction.

NBO analysis has been calculated at the DFT/B3LYP/6-311G** functional in order to clarify the charge transfer or conjugative interaction, the intra compound rehybridization and delocalization of electron density within the molecule. The

second-order perturbation theory of Fock matrix in the NBO of the DCPX showed the presence of strong intramolecular hyperconjugative interactions which are listed in Table 4. In NBO analysis of DCPX, we had observed two types of halogen (Cl₉ & Cl₁₀) antibonding with their respective stabilization energies. They were observed as follows Cl₉ (LP1) undergoes π *C2–C3, σ *C1–C2 interaction with $E^{(2)}$ of 12.45, 11.68 kJ mol⁻¹ where as Cl₁₀ (LP1) σ *C5–C6, π *C4–C5 resulted the stabilization energy $E^{(2)}$ of 11.24, 12.19 kJ mol⁻¹, respectively. This stronger interaction around the ring could enhance the bioactivity of DCPX. The p-character of chlorine lone pair contributes the electron donating ability of DCPX.

4.5. Mulliken population analysis

Mulliken Population Analysis (MPA) of Mulliken atomic charge (MAC) is a significant tool which analyzes the electronic properties of a particular molecule.⁴³ Further, it helps in the determination of atomic charges, dipole moment, molecular polarizability and electronic structure of a molecule. The total atomic charges of DCPX were obtained by Mulliken population analysis with B3LYP/6-311G** functional (Fig. 4). The negative



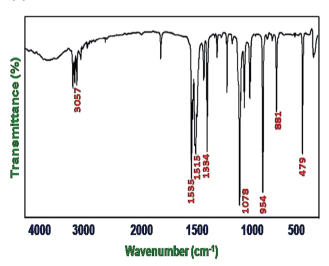


Fig. 2 (a) Calculated (B3LYP/6-311G**) FT-IR spectra of DCPX, (b) observed FT-IR spectra of DCPX.

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values of C₂, C₃, C₅, C₆, C₁₁, C₁₂, Cl₉ and Cl₁₀ in the aromatic ring indicated the redistribution of electron density (Table S4 in ESI†). Due to these strong negative charges C2 and C5 accommodated higher positive charge and become more acidic. The substitution of halogen implied negative values on ring carbons where as the methyl group's substitution kept C4 of aromatic ring as positive.

4.6. Molecular electrostatic potential (MEP)

MEP is an extensive analytical tool for the description of noncovalent interactions particularly hydrogen bonds. MEP analysis enables the identification of high susceptible interactive centre with a proton in the surrounding space of a molecule. 12,40,44 This can be used to predict potential interactions between the molecule and a particular host. To predict the reactive sites of electrophilic and nucleophilic attack for the investigated molecule, MEP at the B3LYP/6-311G** optimized geometry was drawn (Fig. S2 in ESI†). From Fig. S2 (ESI†) negative (red and orange) area of the MEP are relative to the electrophilic reactivity and the positive blue area are related to nucleophilic reactivity. From the MEP map of the DCPX molecule, it is clear that negative potential are on the electronegative halogen atoms (Cl₉ & Cl₁₀), where as the regions having the positive potential are over the hydrogen atoms. Blue color (Fig. S2 of ESI†) indicates the strongest attraction and a red

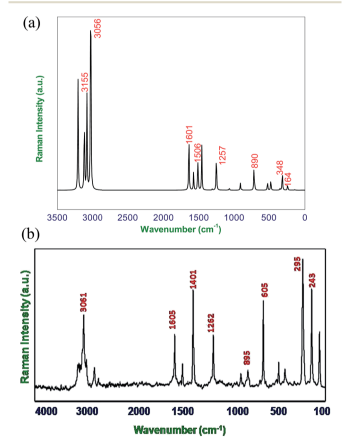


Fig. 3 (a) Calculated (B3LYP/6-311G**) FT-Raman spectra of DCPX, (b) observed FT-Raman spectra of DCPX.

color indicated the strongest repulsion of DCPX towards electrophiles.

Global molecule reactivity descriptors

The structure-activity relationship (SAR) of a particular molecule can well be explained by calculating the global reactivity descriptors using conceptual DFT. Regarding the reactivity of a molecule, the most serious concern is the assumption the direction of reaction and the product formation in various environments. At the time of development of quantum chemical methods, there are several calculation methods have been designed for the calculation of the electronic properties of molecular systems. Among the several concepts derived the one which used to correlate the frontier molecular orbitals with reactivity of molecules designed by Fukui & his co-workers has been utilized extensively to assume the nature of photochemical reaction.45 Meanwhile, Pearson has derived the HSAB concept to explore the reactivity of acids and bases through their hardness and softness. Pearson stated that, there will be an enhanced stabilization when the soft acid and hard acid combine with soft base and hard base, respectively.46,47 HSAB principle is one of the most prominent concepts as it covers almost all the acid-base reactions and it has been become very popular due to the applicability and simplicity.

DFT methods stay better than conventional ab initio method because of economy ratio. The most quantum chemical methods are hybrid DFT based B3LYP functional level which predicts the global and local reactivity descriptors. The value of this innovative molecular reactivity quantity has newly been demonstrated in considerate the toxicity of a range of pollutants in terms of their molecular reactivity and site selectivity. 48,49 The quantum chemical parameters calculated using DFT with B3LYP/6-311G** functional viz., molecular orbital energies (HOMO and LUMO), energy gap (ΔE) , electron affinity (EA), ionization potential (IP), global hardness (η), dipole moment

Table 3 Calculated chemical shifts of carbon atoms and hydrogen atoms of DCPX using TMS B3LYP/6-311G* (2d, p) and TMS HF/6-31G (d) GIAO methods

Atoms	Experimental chemical shift (ppm)	TMS B3LYP/6-311G* (2d, p)	TMS HF/6-31G (d) GIAO
C ₁	138.56	137.526	155.05
C_2	135.12	145.241	162.72
C_3	132.24	128.239	146.42
C_4	131.15	137.526	155.05
C_5	128.24	145.241	162.72
C_6	119.21	128.239	146.42
C_{11}	20.9	12.92	30.46
C_{12}	20.9	12.92	30.46
H_7	7.04	6.069	6.784
H_8	7.02	6.069	6.784
H_{13}	2.29	1.209	1.925
H_{14}	2.24	0.839	1.554
H_{15}	2.12	0.449	1.164
H_{16}	2.22	0.839	1.554
H ₁₇	2.03	0.449	1.164
H_{18}	2.25	1.209	1.925

Table 4 Second order perturbation theory analysis of DCPX using B3LYP/6-311G** functional

	Occupancy	Acceptor (J)	Occupancy	$E^{(2)}$ kcal mol^{-1}	E(i) - E(j) a.u.	F(i,j)
σC_1 – C_2	1.97581	$\pi^*\mathrm{C}_2$ - C_3	0.0248	4.4	1.3	0.068
		σ^*C_3 – H_7	0.0141	2.34	1.14	0.046
$\sigma C_1 - C_{12}$	1.97895	π^*C_2 - C_3	0.0248	2.88	1.18	0.052
πC_1 – C_6	1.96137	π^*C_2 - C_3	0.03546	4.42	1.29	0.067
		σ^*C_2 –Cl ₉	0.03141	4.62	0.85	0.056
		σ^*C_5 -Cl ₁₀	0.03126	4.84	0.85	0.057
σC ₂ -Cl ₉	1.98917	σ^*C_3 - C_4	0.02238	2.27	1.3	0.049
		π^*C_1 - C_6	0.02278	2.7	1.3	0.053
πC_2 – C_3	1.97839	σ^*C_3 – C_4	0.02238	3.1	1.32	0.057
σC_3 – H_7	1.97816	π^*C_2 - C_3	0.0248	0.84	1.09	0.027
σC_3 – C_4	1.96134	π^*C_2 - C_3	0.0248	3.47	1.28	0.06
		σ^*C_2 -Cl ₉	0.03141	4.84	0.85	0.057
		σ^*C_5 -Cl ₁₀	0.03126	4.62	0.85	0.056
$\pi C_4 - C_5$	1.97535	σ^*C_3 – C_4	0.02238	3.7	1.32	0.062
		σ^*C_3 - H_7	0.0141	2.02	1.14	0.043
σC_4 - C_{11}	1.9791	π^*C_2 - C_3	0.0248	2.77	1.18	0.051
$\sigma C_5 - C_6$	1.97841	π^*C_1 - C_6	0.02278	3.14	1.32	0.057
		σ^*C_1 - C_{12}	0.01562	3.11	1.11	0.053
σC_5 - Cl_{10}	1.98913	σ^*C_3 – C_4	0.02238	2.72	1.3	0.053
		π^*C_1 - C_6	0.02278	2.26	1.3	0.049
σC_6 – H_8	1.97804	σ^*C_1 - C_2	0.03546	4.56	1.09	0.063
σC_{12} – H_{14}	1.98275	σ^*C_1 - C_2	0.03546	1.65	1.07	0.038
σC_{12} – H_{15}	1.99086	π^*C_1 -C ₆	0.02278	3.6	1.08	0.056
σC_{11} – H_{16}	1.98226	σ^*C_3 – C_4	0.02238	1.63	1.09	0.038
σC_{11} – H_{17}	1.99186	π^*C_4 - C_5	0.035	3.62	1.07	0.056
Cl ₉ (LP1)	1.96427	π^*C_2 - C_3	0.0248	12.45	0.32	0.062
		σ^*C_{12} - H_{14}	0.01205	0.31	1.74	0.044
		σ^*C_1 - C_2	0.39909	11.68	0.34	0.062
Cl_{10} (LP1)	1.99351	σ^*C_5 - C_6	0.02482	11.24	1.49	0.039
		π^*C_4 - C_5	0.035	12.19	0.32	0.062
		$\sigma^*C_{12}-H_{16}$	0.01526	0.74	1.73	0.041

(μ), global softness (σ), absolute electronegativity (χ) and electrophilicity index (ω) are listed in Table 5.

The molecular energies of HOMO and LUMO are related to the IP and EA, respectively in the framework of Koopmans' theorem.⁴⁸

$$IP = -E_{HOMO} \tag{4}$$

$$EA = -E_{LUMO}$$
 (5)

Using Koopman's theorem for closed shell molecules η, μ and χ can be redefined as

$$\chi = \frac{IP + EA}{2} \tag{6}$$

$$\mu = \frac{E_{\text{HOMO}} + E_{\text{LUMO}}}{2} \tag{7}$$

$$\eta = \frac{\mathrm{IP} - \mathrm{EA}}{2} \tag{8}$$

The σ is defined as the inverse of the η eqn (8).

$$\sigma = \frac{1}{\eta} \tag{9}$$

The electrophilicity index (ω) formula is

$$\omega = \frac{\mu^2}{2\eta} \tag{10}$$

In addition, he described the nucleophilicity (ε) as the multiplicative inverse of electrophilicity ($\varepsilon=1/\omega$).

The chemical hardness and softness of DCPX is clue of the stability of the molecule. Molecules having low energy gap are called as soft molecules which are polarisable, whereas molecules having high energy gap are called as hard molecules which are less polarisable (non-polar). From the Table 5, it is clear that, DCPX having larger energy gap. The values denoted DCPX as hard and less polarisable (non-polar) molecule.

4.8. HOMO and LUMO analysis

Molecules having conjugated electrons are recognized as hyperpolarisabilities and are analyzed in terms of vibrational spectroscopy. In several cases, even in the absence of an inversion center, the intense bands in the Raman spectrum are broad in the IR spectrum and *vice versa*. But the intra-molecular charge transfer from the donor to accepter group through a single-double-bond conjugated path can induce large variations of both the molecular dipole moment and the molecular polarisability, making IR and Raman activity strong at the same time.

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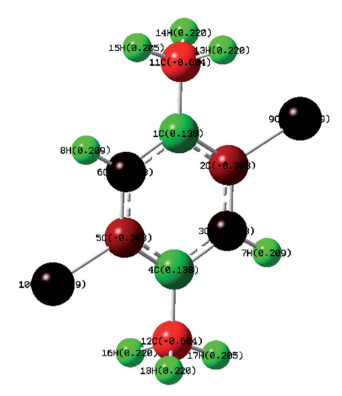


Fig. 4 Calculated Mulliken atomic charges of DCPX.

The above said experimental spectroscopic behavior can well be correlated using DFT calculations in conjugated systems which assume large Raman and infrared intensities for the similar normal modes. It is also observed in the present study that the bands in FT-IR spectrum have their counterparts in the Raman which reveals that the relative intensities in both the spectra are reliable and also explains the result from the movement of electron cloud $\emph{via}~\pi$ conjugated electron donor to acceptor species.

The transition of electrons from ground state to first excited state due to the absorption by means of one electron excitation from the HOMO to LUMO has been known from the wave function analysis. The Gauss Sum program is used to obtain DOS diagram which was drawn by plotting molecular orbital data. The DOS diagram⁵⁰ of DCPX is shown in Fig. 5. It is well known that chemical stability of a molecule is largely affected by the frontier orbitals. The electron donating capability of a molecule can be explained by HOMO–LUMO energy. Thus, it is assumed that the energy gap between the HOMO and LUMO reveals the molecules reactivity.

Fig. 6 shows the energy of HOMO–LUMO energy gap (ΔE) of DCPX. The calculated HOMO–LUMO energy gap of the DCPX is 12.06, 12.23, 10.49, 5.92, 6.07, 5.96 and 12.10, 12.26, 10.60, 5.98, 6.12, 6.03 eV at the HF/SDD, HF/6-311G, HF/6-31G++, B3LYP/6-DD, B3LYP/6-311G++, B3LYP/6-31++G (Gaseous phase) and HF/SDD, HF/6-311G, HF/6-31G++, B3LYP/SDD, B3LYP/6-311G++, B3LYP/6-31++G levels (aqueous phase), respectively. Hence, this value revealed that DCPX has relatively less kinetic stability than other derivatives. Orbitals involved in the electronic transition of (a) HOMO–0 (b) LUMO+0 (c) HOMO–1 (d) LUMO+1 (e) LUMO+2 are represented in Fig. S3 in ESI.† The NLO responses can be understood by examining the energies of frontier molecular orbitals. There is an inverse relationship between hyperpolarizability and HOMO–LUMO.

4.9. First-order hyperpolarizability calculations

Under the applied electric field, first-order hyperpolarizability (a third rank tensor) can be described by a $3 \times 3 \times 3$ matrix. The Kleinman symmetry reduces the components of the 3D matrix to 10 components. 51 The lower tetrahedral format can be used to explain the matrix. It is very clear that, the tetrahedral's lower part is $3 \times 3 \times 3$ matrix. The calculation of NLO properties with high accuracy is challenging and requires consideration of many different issues. Computational techniques are becoming valuable in designing, modeling and screening novel NLO materials. The hyperpolarizability of DCPX is 11.583 imes 10 $^{-30}$ esu (calculated) which is sixty times greater than that of urea $(0.1947 \times 10^{-30} \, \text{esu})$. From the observation, it is clear that DCPX having NLO properties. The dipole moment and hyperpolarizability values are listed in Table S5 in ESI.† Oscillator strength, electronic excitation energies and nature of the respective excited states were calculated by the closed-shell

Table 5 Chemical parameters of DCPX in gaseous phase as well as aqueous phase calculated using various methods

Methods	E_{HOMO} (eV)	$E_{\mathrm{LUMO}}\left(\mathrm{eV}\right)$	I	A	ΔE	η	σ	χ	ω	ε
HF/SDD	-9.35154	2.71436	9.35154	-2.71436	12.06590	6.03295	0.16576	3.31859	0.91274	1.09560
HF/6-311G	-9.40895	2.82837	9.40895	-2.82837	12.23733	6.11866	0.16343	3.29029	0.88467	1.13036
HF/6-31++G	-9.37385	1.11704	9.37385	-1.11704	10.49089	5.24544	0.19064	4.12841	1.62462	0.61553
B3LYP/SDD	-6.77569	-0.84628	6.77569	0.84628	5.92941	2.96470	0.33730	3.81098	2.44942	0.40826
B3LYP/6-311G++	-7.00209	-0.92792	7.00209	0.92792	6.07417	3.03709	0.32926	3.96500	2.58821	0.38637
B3LYP/6-31++G	-6.94440	-0.97690	6.94440	0.97690	5.96750	2.98375	0.33515	3.96065	2.62869	0.38042
DCPX/aqueous pha	ise									
HF/SDD	-9.36705	2.73939	9.36705	-2.73939	12.10644	6.05322	0.16520	3.31383	0.90708	1.10244
HF/6-311G	-9.41004	2.85014	9.41004	-2.85014	12.26019	6.13009	0.16313	3.27995	0.87748	1.13963
HF/6-31++G	-9.37304	1.23405	9.37304	-1.23405	10.60708	5.30354	0.18855	4.06949	1.56130	0.64049
B3LYP/SDD	-6.84834	-0.86669	6.84834	0.86669	5.98165	2.99083	0.33436	3.85752	2.48768	0.40198
B3LYP/6-311G++	-7.04263	-0.91404	7.04263	0.91404	6.12860	3.06430	0.32634	3.97834	2.58251	0.38722
B3LYP/6-31++G	-6.96780	-0.93445	6.96780	0.93445	6.03336	3.01668	0.33149	3.95112	2.58751	0.38647

Paper

3.0 2.5 B3LYP/6-311G** 2.0 1.5 1.0 (6.07417eV) 0.5 0.0 -0.5

Fig. 5 DOS diagram representation of HOMO (left) and LUMO (right) of DCPX.

Energy (eV)

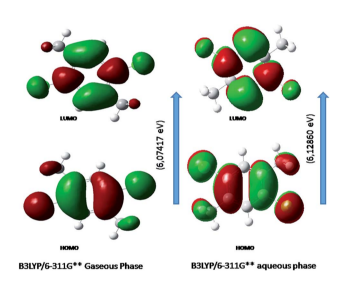


Fig. 6 Representation of the orbitals involved in the electronic transition of DCPX for HOMO and LUMO

singlet calculation method and are summarized in Table S6 of ESI.† Table S6† clearly indicated the higher first static polarizability of DCPX.

5 Conclusions

The symmetry of DCPX has been identified as C_s point group from the optimized structure. In addition, the bond lengths and bond angels of all atoms of DCPX have also been calculated. The difference in bond lengths of various substituents has been due to the inductive and resonance effects in DCPX. The 48 normal modes of DCPX were calculated using DFT coupled with B3LYP/ 6-311+G** functional. Computed vibrational modes include FT-IR and FT-Raman was compared with experimentally observed values and which is in quite agreement with each other. The very least mean deviation of about 7.09 cm⁻¹ for DCPX has been observed after the least square refinement of the scale factors validated the adoption of B3LYP method. The structural features of DCPX have been confirmed from the results of NMR spectral analysis. The occurrence of intra-molecular hydrogen bonding in DCPX has been confirmed from NBO analysis in addition to biological activity. The charge distribution of DCPX is known from Mulliken population analysis. The positive and negative regions followed by nucleophilic and electrophilic reactive centres have been identified from MEP analysis. The SAR of DCPX based on global reactivity descriptors suggested DCPX as hard and non-polar species. The HOMO-LUMO analysis of DCPX revealed the donor-acceptor nature, reactivity and NLO property of DCPX. Further, the NLO property of DCPX has been supported by the observation of larger hyperpolarizability value.

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