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Spectroscopic analysis of 2,5-Disubstituted Pyridine Derivatives: Part III

Krishna Jaiswal, Janardan Prasad Pandey*

M.L.K.P.G. College, Balarampur (U. P.) INDIA 271201

ABSTRACT

Structure and bonding in liquid crystals is important and diverse field in the interface between modern physics and chemistry. Geometrical anisotropy of small molecules with high polarizability may exhibit one or more liquid crystalline phases as well the familiar crystalline and isotropic phases. Structure—property relationship has been a key issue for the study of liquid crystals. Characteristically, it is observed that liquid crystalline compounds have a rigid core, which acts as the mesogenic unit and flexible alkyl chains at the terminal ends. Pyridine derivatives form an important class of molecules exhibiting liquid crystalline behavior. The pyridine derivative has been examined quantum mechanically to understand its physical and spectroscopic details along with various derivatives. The strong dipole-dipole interaction exhibits the crystalline phase and has higher thermal stability. The present work reports a liquid crystal molecules namely 2-[2-(4-butoxyphenyl) ethyl]-5-(4ethylcyclohexyl) pyridine. This molecule undergoes mesogenic phase as Cr 48 Sm 61 N 93 I. Its IR spectra as well as Raman activities is analised. The atomic charges with point dipoles were also computed. The Humo-Lumo gap as representation of band gap, thus computed.

Keywords: 2-[2-(4-butoxyphenyl) ethyl]-5- (4ethylcyclohexyl) pyridine, Liquid Crystals, Mesogen, IR Spectra, Raman Activities

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1. INTRODUCTION

Liquid crystal is an intermediate phase of matter (between isotropic liquid and crystal). these molecules were originally discovered in chemical and biological system. Since long time, liquid Crystals (LCs) have been the subject of theoretical and experimental study. By their role in chemical, biological and elctro-optical system, they have many industrial applications. Liquid crystals are powerful solvent in holding donor-spacer-acceptor systems as well as in maintain the fluid and order property over a wide range of temperatures too. Numerous electronic and optical properties of LC varies with change of substituent. These properties of molecules can be altered by the presence of different substituents. LC molecule have a dipole moment directly perpendicular to the molecular axis.[1-5]

The pyridine derivatives have been examined quantum mechanically to understand the possible reason for varying mesogenic character. The understanding the LC property of these pyridine is of fundamental importance in chemical physics. Raman scattering depends on the polarizability. The Homo-Lumo energy gap is effective for stability of the molecule. Also the electrical transport property determine by the Homo-Lumo gap. [6-7]

Pyridine derivatives form an important class of molecules exhibiting liquid crystalline behavior. The introduction of substituent group in lateral position at terminal and central benzene core decreases thermal stabilities. Theoretical study suggest that the potential around the pyridine molecule in symmetric in nature [8-12].

The present work reports a liquid crystal molecule namely 2-[2-(4-butoxyphenyl) ethyl]-5- (4ethylcyclohexyl) pyridine. These are dependent not only on the electronic properties of individual molecules but also on the properties that are determined in varies physical properties and effect of respective molecule. The vibration states of a molecule are observed theoretically through infrared and Raman spectroscopy.

2. COMPUTATIONAL METHOD

The geometry was optimized using DFT method B3LYP [13,14] using 6-31G** [15, 16] which was found suitable for these type of systems [17] with keeping all atoms free. The analytical frequencies as well as Raman activities were calculated. All calculation were done using Gaussian09 programme suit.[18]

3. RESULTS

The optimized geometry of 2-[2-(4-butoxyphenyl) ethyl]-5- (4ethylcyclohexyl) pyridine molecule is shown in figure 1. The phenyl ring and pyridine ring is connected with ethyl group and phenyl ring and pyridine ring separated with ethyl group by 1.51Å. The alkoxy chain is plannar (06.3°) to phenyl ring and separation is 1.137Å. The cylcohexane is also trans (102.6°) to pyridine ring with separation 1.52Å. The cyclohexane is in boat configuration.

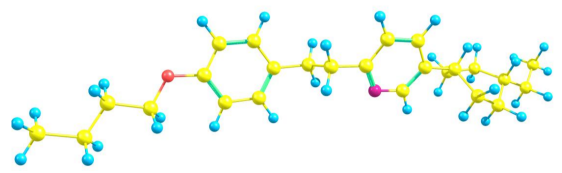


Figure 1: The optimized structure of the 2-[2-(4-butoxyphenyl) ethyl]-5- (4ethylcyclohexyl) pyridine.

Table 1 present the charges as well as multipoles corresponding each atoms of 2-[2-(4-butoxyphenyl) ethyl]-5-(4ethylcyclohexyl) pyridine molecule. Since multipole depends on the coordinates of each atoms therefore coordinates are aslo tabulated here.

Table 1: The charge, coordinates and multipoles corresponding each atoms of 2-[2-(4-butoxyphenyl) ethyl]-5- (4ethylcyclohexyl) pyridine molecule.

Sr.	Coordinates							
Sr. No.	Atom	X	Y	Z	Charge		Multipole (au)	
1	С	0.4226	0.36263	1.04478	-0.19744	0.279309	0.055385	-0.25462
2	С	-0.51499	1.39804	0.99496	0.16719	-0.22913	0.016514	0.126557
3	С	-0.31807	2.40558	0.03659	-0.14186	0.637068	-0.29674	-0.29558
4	С	0.76501	2.37883	-0.8337	-0.56221	0.200796	-0.00718	-0.15813
5	С	1.69399	1.32992	-0.76992	0.328528	-0.14913	-0.0812	0.213946
6	С	1.51927	0.31617	0.17902	-0.54675	-0.05089	-0.07356	0.085252
7	С	-1.7159	1.40904	1.91436	-0.08453	-0.08468	0.142969	-0.20684
8	С	-2.94428	0.68162	1.31272	-0.15874	-0.00641	-0.07488	0.32498
9	С	-4.0996	0.59997	2.28404	0.157068	-0.02495	0.291731	-0.34652
10	N	-3.91082	-0.17242	3.36975	0.050149	-0.73563	0.319643	-0.00672
11	С	-4.89968	-0.26916	4.26078	-0.20143	0.141195	-0.42624	0.609914
12	С	-6.1393	0.38175	4.15606	-0.4536	1.323942	0.704523	-2.26436
13	С	-6.31656	1.18422	3.02589	-0.57029	-0.26815	0.509981	-0.53795
14	С	-5.2957	1.29682	2.08326	-0.40815	-0.18584	-0.00491	0.099593
15	С	-7.22104	0.22416	5.20759	0.357679	2.090386	0.387893	-1.97131
16	С	-7.77046	-1.23065	5.27934	0.786635	-0.97149	0.430553	-0.21776
17	С	-8.41697	-1.55985	6.64403	-0.34704	-0.04862	-0.6158	0.085057
18	С	-9.06337	-0.32393	7.29016	0.933616	0.623042	-0.43485	0.276331

	1			1				
19	С	-7.9703	0.73597	7.59166	-0.67543	-0.21371	1.056851	-0.51459
20	С	-6.7729	0.676	6.61368	1.234102	-0.34693	-1.78745	0.033941
21	С	-9.88471	-0.69168	8.53753	0.220605	-0.17421	0.137853	0.068384
22	С	-10.6342	0.48454	9.17396	0.221989	-0.10159	0.12003	-0.04201
23	О	2.72564	1.39144	-1.66427	-0.72282	-0.5574	0.284516	0.220957
24	С	3.70045	0.35256	-1.65482	1.25442	-0.95325	0.110175	0.803716
25	С	4.71564	0.65357	-2.74916	0.971106	-0.58963	0.553497	-0.00323
26	С	5.81804	-0.40902	-2.83729	0.907287	-0.10931	-0.09546	0.309605
27	С	6.8417	-0.11402	-3.93828	0.492821	0.089659	-0.01966	-0.0818
28	Н	-9.94802	1.25928	9.53111	-0.06807	0.085559	0.069836	0.028026
29	Н	-11.3172	0.95283	8.45533	-0.14973	-0.10592	0.070504	-0.10303
30	Н	-11.2312	0.15399	10.03025	-0.07091	-0.0553	-0.0362	0.091176
31	Н	-10.6083	-1.47143	8.26432	-0.14673	-0.09006	-0.12227	-0.01875
32	Н	-9.21521	-1.14612	9.28209	-0.11393	0.052692	-0.06315	0.093427
33	Н	-9.76317	0.10373	6.55488	-0.07486	-0.03769	0.05631	-0.0685
34	Н	-9.15843	-2.35803	6.51953	-0.13987	-0.11427	-0.09575	-0.02117
35	Н	-7.65803	-1.95707	7.33211	-0.09497	0.107341	-0.04518	0.045706
36	Н	-8.49812	-1.37092	4.47108	-0.45682	-0.19093	-0.08264	-0.2511
37	Н	-6.95843	-1.93927	5.08132	0.111319	0.045769	0.027786	0.025717
38	Н	-8.04876	0.88237	4.91041	0.101745	-0.07422	-0.03081	0.088089
39	Н	-6.28032	1.65279	6.5591	0.131532	-0.01619	0.023053	0.014198
40	Н	-6.0167	-0.02523	6.98843	-0.53786	0.25512	-0.13378	0.123095
41	Н	-8.41562	1.73723	7.56447	-0.17453	-0.05664	0.130678	0.022447
42	Н	-7.60013	0.5945	8.61581	0.241731	-0.01842	-0.03024	-0.05378
43	Н	-4.69548	-0.91051	5.11798	-0.09884	0.075329	-0.08391	0.076752
44	Н	-5.42478	1.91488	1.19937	0.158593	0.006054	-0.00761	0.009795
45	Н	-7.2534	1.7185	2.88275	0.01418	-0.07363	0.020721	0.016301
46	Н	-2.64353	-0.33612	1.03829	0.086179	0.016188	-0.01286	0.025985
47	Н	-3.25779	1.18566	0.39173	0.117927	0.012582	-0.00568	0.015065
48	Н	-1.47217	0.92056	2.86352	0.143881	-0.00626	-0.01017	-0.02222
49	Н	-1.99654	2.44259	2.15216	-0.04519	-0.0043	0.093449	0.02183
50	Н	-1.02468	3.23046	-0.02447	0.317267	0.074135	-0.09624	0.019156
51	Н	0.91766	3.16158	-1.56997	0.191294	-0.03181	-0.0159	0.037051
52	Н	2.22393	-0.50322	0.2576	0.161166	-0.05857	0.017001	0.039116
53	Н	0.30179	-0.43031	1.77941	0.20289	-0.00896	0.045454	-0.02477
54	Н	4.1927	0.30512	-0.67178	-0.41934	0.190561	-0.02574	0.240357
55	Н	3.21737	-0.62016	-1.83187	-0.3677	-0.07415	-0.23889	-0.07257
56	Н	5.15708	1.63954	-2.55793	-0.39383	0.144882	0.27692	0.04478
57	Н	4.18654	0.72772	-3.70734	-0.41487	-0.13689	0.011409	-0.30209
58	Н	5.36371	-1.39312	-3.01553	-0.32607	-0.09955	-0.23948	-0.04877
59	Н	6.33347	-0.48265	-1.87004	-0.37904	0.147197	-0.03192	0.256321
60	Н	7.33777	0.84837	-3.7697	-0.16936	0.084897	0.162009	0.016178
61	Н	6.3624	-0.07035	-4.92268	-0.17029	-0.07664	0.015322	-0.16617
62	Н	7.61681	-0.88591	-3.97893	-0.17981	0.136274	-0.1262	-0.01287
	•							

Various energies components with zero point corrections of 2-[2-(4-butoxyphenyl) ethyl]-5- (4ethylcyclohexyl) pyridine molecule is tabulated in Table 2.

Table 2: Energies Components such as electronic, thermal and Free energies of 2-[2-(4-butoxyphenyl) ethyl]-5- (4ethylcyclohexyl) pyridine molecules.

Energies Components	Hartree
Sum of electronic and zero-point Energies	-1103.21
Sum of electronic and thermal Energies	-1103.19
Sum of electronic and thermal Enthalpies	-1103.19
Sum of electronic and thermal Free Energies	-1103.28

Table 3 presents dipole moment, exact polarizability, approx. and polarizability of 2-[2-(4-butoxyphenyl) ethyl]-5- (4ethylcyclohexyl) pyridine molecules. The band gap (Humo-Lumo) is 5.12 eV. The ionization potential is 5.57 eV and electron affinity is 0.45 eV.

Table 3: Dipole monent, exact polarizability, and approx. polarizability of 2-[2-(4-butoxyphenyl) ethyl]-5-(4ethylcyclohexyl) pyridine molecules.

	 	 3 / 1 3
Dipole Monent		2.7666debye
Exact Polarizability		287.528
Approx Polarizability		390.962

The IR spectra of 2-[2-(4-butoxyphenyl) ethyl]-5- (4ethylcyclohexyl) pyridine molecules molecule is shown in figure 2. From figure 2 it can be visualized that there are several peak and the highest peak (IR intensity) is at 1288.9152 cm⁻¹. This frequency is associated with twisting of phenyl ring attached with alkoxy chain. Second peak is at 1560.8452 cm⁻¹. This frequency is associated with twisting of alkoxy chain. Other peak height is at 3030.0207 cm⁻¹ that is associated with twisting of cyclohexane.

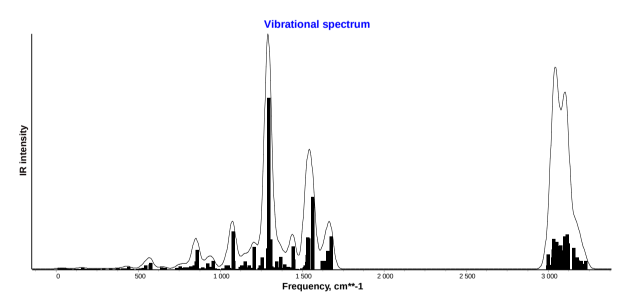


Figure 2: IR Spectra of 2-[2-(4-butoxyphenyl) ethyl]-5- (4ethylcyclohexyl) pyridine molecule.

Figure 3 represent Raman activities of 2-[2-(4-butoxyphenyl) ethyl]-5- (4ethylcyclohexyl) pyridine molecule. There are various peaks and the highest Raman activity is at 1233.2506 cm⁻¹ which are associated with twisting of phenyl ring with alkoxy group. Another peak is at 1652.551 cm⁻¹, which is accosiated with twisting of pyridine ring. Next peak is at 3040.0893 cm⁻¹ that associated with twisting of alkoxy chain tail.

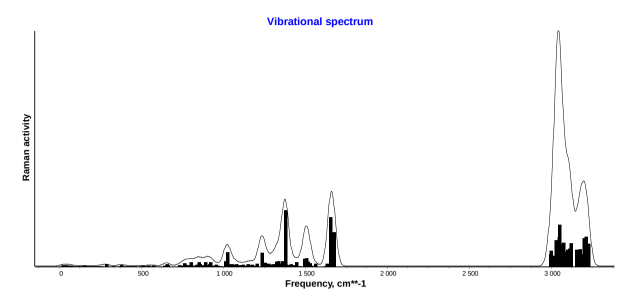


Figure 3: Raman activity of 2-[2-(4-butoxyphenyl) ethyl]-5- (4ethylcyclohexyl) pyridine molecule.

4. CONCLUSION

The electronic structure analysis of on 2-[2-(4-butoxyphenyl) ethyl]-5- (4ethylcyclohexyl) pyridine molecule. The IR peaks and Raman activities peaks were explained.

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