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Mohammed T.Hussien University of Baghdad – college of science – department of physics-Iraq-Baghdad, mohammedtaki97@yahoo.com

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Ab-initio-RHF Methods Calculation to Study the Fundamental Vibrational Frequency of Benzophenone (C₆H₅-CO-C₆H₅)

Mohammed T.Hussien¹

¹ University of Baghdad – college of science – department of physics-Iraq-Baghdad

E-mail: mohammedtaki97@yahoo.com

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Abstract: The electronic distribution and energy level as well as modes of vibration of Benzophenone molecule are studied theortically using ab-initio methods. The geometry optimization of the benzophenone was obtained depending on the Hartree - Fock equation for the restricted system by using STO - 3G basis sets. The modes of vibrationes were calculated under the steady state geometry condition and presented in graphically with frequency, intensity and symmetry for each mode. The molecular orbital calculation and energy level diagram appears that paired electron . The Ionization potential is equal to = 7.4939 eV, while the low electron affinity is equal to = 5.64372 eV. The total energy of benzophenone is equal to = (-15397.47644 eV) where calculated by Ab-initio method compared with different semi-empirical methods, indicate that Ab-initio method is very high accurate method to give the geometry more stable .

Keywords: Ab-initio rhf methods for benzophenone modes .

1. Introduction

Theoretical computation in physics and chemistry using various methods depandant on ab-initio and semi empirical methods are widely used. These methods are very important to the studies of matter and its physical properties. The advancement in computer systems enabled the improvement in the increased accuracy and speed of evaluating theoretical results.

Many searchers used ab-initio and semi empirical methods for example , (Laref,2000) studied the band structure of Germanium crystal using semi empirical methods (Benzair and Aourag , 2003) studied the electronic properties and total energy of Zinc-blende compounds using ab-initio and density functional methods.

In this work, the distribution, energy levels and normal modes of vibration of benzophenone molecule which consist a Carbonal group are studied using ab-initio methods .



The main goal is to classify the benzophenone as an example of non-linear molecule theoretically according to group theory using the linear combination of benzophenone orbitals (LCAO).

2. Methods

In Hartree Fock HF theory, the wave function is represented by a single N-dimentional slater determinant $\phi(x_1, x_2, \dots, x_N)$ made up of N orthonormal spin orbital {Xi(x)}, where x represents both the position r and the spin ω of an electron. Each spin orbital can be have both a spin up α and spin down β part; $\psi(r)$ and $\psi(r)$ respectively (parr and yang 1989).

In restricted Hartree - Fock (RHF) and unrestricted Hartree - Fock (UHF) methods, each spin orbital is either pure α or pure β . In UHF theory, the two sets of molecular orbitals are defined by two sets of

Coefficients $\stackrel{\alpha}{C}_{\mu i}, \stackrel{\beta}{C}_{\mu i}$

For a given nuclear configuration Ri that includes a system of M nuclei and a given set of orthonormal spin orbitals, the electronic energy $E_e(\{ \substack{\alpha \\ \mu i}, \substack{\beta \\ \mu i}, \{ R_i \})$ is (Sherrill,2000):

$$E_{e} = \sum_{i=1}^{N} \langle X_{i} \middle| -\frac{1}{2} \nabla_{I} + \sum_{I=1}^{N} V_{I}(r_{i}, R_{I}) \middle| X_{i} \rangle + \sum_{i=1}^{N} \sum_{j \langle i} \left[\langle X_{i} X_{j} \middle| X_{i} X_{j} \rangle - \langle X_{i} X_{j} \middle| X_{i} X_{j} \rangle \right]$$

$$\tag{1}$$

Where

$$V_I(r_i, R_I) = \frac{Z_I}{|r_i - R_I|}$$
 for the electronic r_i and nuclear R_I

The Born-oppenheimer (Sherrill,2000;klienert,1999) approximation separates the electron and nuclear motions because the nuclear mass is so much larger than that of the electrons, and the nuclei move on a potential energy surface given by :

$$E(\{\stackrel{\alpha}{C}_{\mu i}, \stackrel{\beta}{C}_{\mu i}, \{R_{i}\}) = E_{ele} E(\{\stackrel{\alpha}{C}_{\mu i}, \stackrel{\beta}{C}_{\mu i}, \{R_{i}\}) + \sum_{I,J < 1}^{M} \frac{Z_{I}Z_{J}}{|R_{I}R_{J}|}$$
(2)

Where Z_I , Z_J are the atomic number for I and J atoms respectively .

Π - Ab-initio Methods.

In ab-initio the calculations of electronic structure are based on the HF wave functions. The approximation ab-initio treatments are based on the variation principle which requires an evalution of (szabo, and ostlund 1982).

$$E[\Psi] = \langle \Psi | H | \Psi \rangle / \langle \Psi | \Psi \rangle$$
(3)

$$H=\sum_{i}h(i)+\sum_{i\langle j}\frac{1}{r_{ij}}$$
(4)

Where h(i) represents the single i^{th} electron term which includes the kinetic energy of the electron as well as its energy associated with its interaction with the nucleus . The two electron term r_{ij} denotes the distance between electron i and j.

An application of the variation principle implies that all, integrals for Ψ should factorize into low dimensional cases .

This condition is achieved by building Ψ from the one electron function ϕ i that are called molecular orbitals (Mo), and this leads to the general properties :

$$\psi = \sum_{I} C_{I} \phi_{I} \tag{5}$$

$$\boldsymbol{\phi}_{I} = \begin{bmatrix} \boldsymbol{\phi}_{iI} \dots \boldsymbol{\phi}_{in} \end{bmatrix} \tag{6}$$

$$\left\langle \boldsymbol{\phi}_{i} \middle| \boldsymbol{\phi}_{j} \right\rangle = s_{ij}$$
⁽⁷⁾

3. Results and Discussion

Before starting the calculation, it is necessary to select a geometry that enables the optimization of the compound studied in order to minimize its energy.

In this geometry, the force on the atoms can be calculated by evaluating the gradient of the energy with respect to atomic coordinates analytically.

In quantum mechanics computer programs, such as Mopac 7.21 and hyperchem 6.01, the form of geometry in put called z-matrix . This matrix specifies the positions of an atom(n) by three geometric parameters :

- 1- The bond length r between two atoms r(i,j).
- 2- The bond angle θ at atom j between lines j-i and j-k, $\theta(i,j,k)$.
- 3- The dihedral angle φ between the two planes defined by i-j-k and j-k- \mathbf{k} meeting at the line j-k, φ (i,j,k, \mathbf{k}).

The structure of the benzophenone was optimized at the restricted Hartree-Fock level of theory using the slater-type orbital (STO-3G) basis sets .

Table (1) shows the geometry of Benzophenone molecule as follow .

	• •	
Bond	Bond length(A ⁰)	Bond angle(degree)
C-0	1.21	
C-C	1.52	
O-C-C		122.9

Table 1: Geometry parameters of C₆H₅-CO-C₆H₅.

Using the coordinate system shown in fig (1), one can describe the molecular orbitals of the studied compound in terms of basis orbital derived from ;

- (i) The 2s,2p orbitals of the oxigyen ion .
- (ii) The carbon ion 2s,2p orbitals .
- (iii) The hydrogen ion is orbitals .

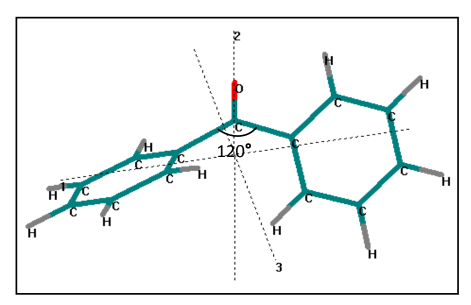


Figure 1: Coordinates system diagram of the Benzophenone C₆H₅-CO-C₆H₅. molecule .

Table (2) shows the net charges and coordinates of the geometry more stable of benzophenone molecule.

Ab-initio calculation of the benzophenone total energy is the sum of the electronic energy plus core-core repulsion and ionization potential as well as dipole moment compare with different semi-empirical methods as presented in table (3).

Atom	Z	Charge		Coordinates(Angstrom	1)	Mass
		(Mulliken)	Х	Y	Z	
0	8	-0.220974	-0.4130539	1.74338527	-2.03575726	15.99900
С	6	0.190394	-0.31069007	0.99759468	-1.06898388	12.01100
С	6	-0.021513	-1.51587369	0.20257209	-0.57271982	12.01100
С	6	-0.021447	1.02391535	0.85043947	-0.34199995	12.01100
С	6	-0.056145	1.88096714	1.94656367	-0.28997973	12.01100
С	6	-0.063444	3.11571284	1.84607265	0.32946562	12.01100
С	6	-0.061943	-1.71874376	-0.06749160	0.77728233	12.01100
С	6	-0.063725	-2.85025254	-0.75178160	1.19149221	12.01100
С	6	-0.057848	-3.78728338	-1.17581619	0.26122335	12.01100
С	6	-0.056099	-2.46976107	-0.21242885	-1.49817130	12.01100
С	6	-0.063413	-3.59626825	-0.90461315	-1.08558521	12.01100
С	6	-0.062001	1.43286833	-0.35673594	0.21685045	12.01100
С	6	-0.057866	3.51224371	0.64124404	0.89061083	12.01100
С	6	-0.063669	2.67201299	-0.46044761	0.82834478	12.01100
Н	1	0.067972	-0.99561071	0.26708158	1.50998550	1.00800
Н	1	0.065506	-3.00209652	-0.95327560	2.24404930	1.00800
Н	1	0.066330	-4.66892835	-1.71372966	0.58645359	1.00800
Н	1	0.066485	-4.32754862	-1.23063650	-1.81421558	1.00800
Н	1	0.073612	-2.31954722	0.01294299	-2.54708331	1.00800
Н	1	0.073550	1.56952409	2.88116766	-0.74026394	1.00800
Н	1	0.066514	3.77067450	2.70682869	0.37256567	1.00800
Н	1	0.066303	4.47841804	0.55905716	1.37247091	1.00800
Н	1	0.065489	2.98383756	-1.40459954	1.25641735	1.00800
Н	1	0.067934	0.78559006	-1.22263168	0.16526170	1.00800

Table 2: Net charge and coordinates of benzophenone molecule.



Quantity	Ab-initio		Semi-empir	ical methods	
	(present work)	MNDO- PM3	MNDO- AM ₁	MINDO/3	MNDO
Total Energy (eV)	-15397.47644	-1986.821	-2121.205	-2105.6	-2124.24
Electronic Energy (eV)	-36473.24409	-11324.4	-11574.23	-11143.27	-11474.5
Core-Core Repulsion (eV)	21075.76765	9337.526	9453.0225	9037.6446	9350.294
Ionization Potential (eV)	7.493904	9.92867	9.79087	8.95864	9.56394
Dipole Moment (Deby)	1.9074	2.78	2.97	3.43	2.8
No. of Filled Levels	48	34	34	34	34
Molecular Weight(amu)	182.22	182.22	182.22	182.22	182.22
Zero Point Energy(eV)	5.9954	5.1587	5.3509	5.2426	5.1593

Table 3: Some physical properties calculated by Ab-initio method of benzophenone molecule, compare with different semi-empirical methods.

The normal modes of vibration of benzophenone nonlinear molecule are calculated which indicate $66 \mod 3$ of vibration , 23 of these modes are stretching and the rest 43 is bending , are presented with frequency, intensity and symmetry for each modes as shown in table (4) .

No	Intensit	v ⁻	(λ)/µm	Types	symmetr	No	Intensit	v ⁻	(λ)/μ	Types	symmet
	у				у		у		m		ry
1	0.472	32.60	306.748	bending	1A	3	0.2845	1205.58	8.2953	bending	34A
2	0.040	55.43	180.407	bending	2A	3	0.1737	1206.97	8.2852	bending	35A
3	0.088	94.60	105.708	bending	3A	3	0.0296	1208.12	8.2773	bending	36A
4	0.849	151.82	65.8674	bending	4A	3	4.1976	1211.25	8.2559	bending	37A
5	2.014	233.05	42.9092	bending	5A	3	1.0979	1270.11	7.8733	bending	38A
6	0.030	238.99	41.8427	bending	6A	3	0.4117	1271.03	7.8676	bending	39A
7	0.041	315.94	31.6515	bending	7A	4	0.1867	1352.1	7.3959	bending	40A
8	4.182	396.94	25.1927	bending	8A	4	4.3062	1368.7	7.3062	bending	41A
9	0.077	476.52	20.9854	bending	9A	4	0.2602	1371.5	7.2912	bending	42A
10	0.015	483.06	20.7013	bending	10A	4	0.9649	1375.35	7.2708	bending	43A
11	0.331	503.86	19.8467	bending	11A	4	0.1371	1378.64	7.2535	Stretchin	44A
12	0.085	514.23	19.4465	bending	12A	4	129.52	1500.64	6.6638	Stretchin	45A
13	0.199	642.12	15.5734	bending	13A	4	1.6277	1546.79	6.465	Stretchin	46A
14	0.501	709.73	14.0898	bending	14A	4	21.978	1547.77	6.4609	Stretchin	47A
15	0.008	713.13	14.0226	bending	15A	4	21.415	1720.5	5.8122	Stretchin	48A
16	32.745	728.68	13.7234	bending	16A	4	22.857	1721.79	5.8079	Stretchin	49A

Table 4: Represents the normal modes of vibration with frequency, intensity, symmetry and types of modes.



179.766776.1712.8837bending17A55.19861784.295.6044Stretch180.354828.3612.0720bending18A51.60741791.595.5816Stretch1918.756835.3511.9710bending19A51.69521912.885.2277Stretch203.780838.6911.9233bending20A52.15411913.15.2271Stretch215.255908.9311.0019bending21A50.73301938.085.1597Stretch	in 51A in 52A
19 18.756 835.35 11.9710 bending 19A 5 1.6952 1912.88 5.2277 Stretch 20 3.780 838.69 11.9233 bending 20A 5 2.1541 1913.1 5.2271 Stretch	in 52A
20 3.780 838.69 11.9233 bending 20A 5 2.1541 1913.1 5.2271 Stretch	
	in 53A
21 5.255 908.93 11.0019 bending 21A 5 0.7330 1938.08 5.1597 Stretch	
	in 54A
22 4.725 940.49 10.6327 bending 22A 5 3.8022 1939.9 5.1549 Stretch	in 55A
23 0.469 1035.1 9.66095 bending 23A 5 30.365 2050.06 4.8779 Stretch	in 56A
24 0.090 1037.3 9.64041 bending 24A 5 0.0805 3708.7 2.6963 Stretch	in 57A
25 65.899 1063.6 9.40132 bending 25A 5 0.1437 3708.8 2.6963 Stretch	in 58A
26 0.282 1123.9 8.89743 bending 26A 5 0.4219 3721.4 2.6872 Stretch	in 59A
27 3.686 1128.1 8.86446 bending 27A 6 0.5859 3723.48 2.6856 Stretch	in 60A
28 1.114 1158.0 8.63498 bending 28A 6 8.9624 3728.78 2.6818 Stretch	in 61A
29 0.027 1160.3 8.61779 bending 29A 6 3.5492 3729.39 2.6814 Stretch	in 62A
30 1.597 1168.2 8.55988 bending 30A 6 12.561 3735.78 2.6768 Stretch	in 63A
31 0.983 1170.8 8.54087 bending 31A 6 13.568 3736.9 2.6760 Stretch	in 64A
32 0.402 1190.9 8.39637 bending 32A 6 4.7428 3744.18 2.6708 Stretch	in 65A
33 0.011 1193.3 8.37956 bending 33A 6 0.9841 3745.3 2.6700 Stretch	in 66A

The vibrational frequencies and its corresponding intensities compare with IR absorption spectrum as shown in fig (2) $^{[8]}$.

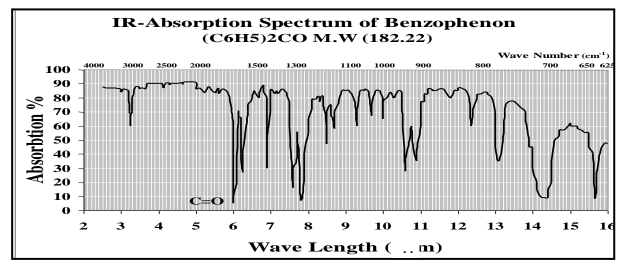


Figure 2: IR absorption spectrum of benzophenone $C_6H_5COC_6H_5$ ^[8]

The total charge density distribution and electrostatic potential of benzophenone molecule in two and three dimension as shown in fig (3) and fig (4) respectively.



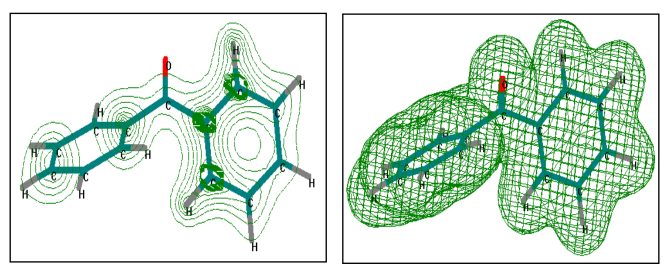


Figure 3: Illustrated total charge distribution for benzophenone molecule in 2D and 3D.

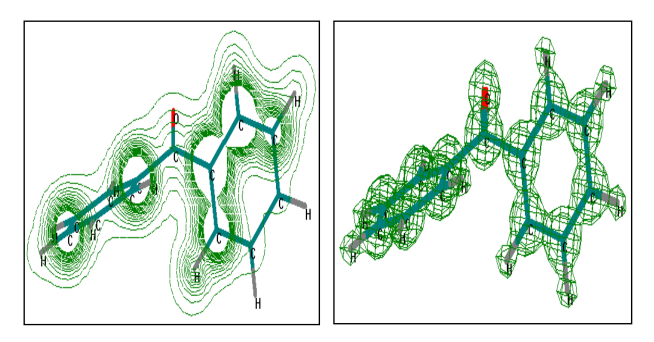


Figure 4: Illustrated electrostatic potential for benzophenone molecule in 2D and 3D.

Fig (5) shows an act final level HOMO and energy value $E_{HOMO} = -7.4939 \text{ eV}$ while symmetry of this level was 48A ,and the first LUMO with energy value $E_{LUMO} = 5.64372 \text{ eV}$ with symmetry 49B. The absolute value of the final level HOMO gives the Ionization potential which is equal to (7.4939) eV, while the first level LUMO represent electron affinity which is equal to (5.64372 eV). They determined the Fermi level as the center of the HOMO and LUMO energies and calculated the gap between them to be (13.1376) eV according to Fischer and Herriksson^[9] is defined as $\Delta E = E_{LUMO} \cdot E_{HOMO}$.



Energy (eV		Symmetry	No.Level
7.2710		53 A ———	53
7.2710		52 A ———	52
7.1270		51 A ———	51
			50
LUMO 5.6437		49 A ———	49
— — — — ———номо -7.4939		48 A	48
-7.8539		47 A	47
-7.8567	11/	46 A	46
-7.9665	11/	45 A ———	45
-8.7877		44 A	44
-11.3066	11/	43 A	43
-11.8433	41,	42 A	42
-12.0406	1	41 A	41
-12.4348	11	40 A	40
-12.4865		39 A ———	39
-12.7329		38 A	38
-13.1819		37 A	37
-14.1298		36 A	36
-14.7623	1/	35 A ———	35
-14.7897	41	34 A ———	34
-14.9691	11/	33 A ———	33
-15.6218	· 1/	32 A ——	32
-15.7358	17	31 A ——	31
-15.7358	14	30 A ———	30
-16.4190	///	29 A ———	29
-17.4530	11,		28
-17.4530		28 A ———	
		27 A —	27
-19.3082	AL.	26 A	26
-20.9014		25 A	25

Figure 5: Schematic diagram for energy levels values for benzophenone shows EHOMO, ELUMO and symmetry .



4. Conclusion

Benzophenone is non-linear molecule has 3N-6 mode of vibration .66 modes are the total number of fundamental modes, 23 of these modes are stretching according to rule (N-1) is the number of stretching , and 43 is the number of bending according to the rule (2N-5) is the number of bending . The Ab-initio calculation of total energy compare with different semi-empirical methods is given too low values, indicate the geometry more stable . Ab-initio method given a data more accuracy, but it taken long time because it take all electrons .The screw (Chair-form) shape of benzophenone molecule is more stable than a boat (Butter-fly) shape.

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