Original Research Article

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Solvent Effects on the Structural, Electronic, Non-Linear Optical

and Thermodynamic Properties of Perylene Based on Density

Functional Theory

6 Abstract:

Perylene (C₂₀H₁₂) is an important member of the polycyclic aromatic hydrocarbons (PAHs) that 7 has a wide applications such as in organic photovoltaic, field effect transistor and bio-sensing. 8 9 Optimized bond lengths and bond angles, HOMO-LUMO energy gap, global chemical indices, total energy, nonlinear optical and thermodynamic properties of Perylene in the gas phase and in 10 solvents (water, chloroform, benzene and acetone) were obtained based on Density Functional 11 Theory with B3LYP/6-311++G(d,p) basis set. All the computations were carried out using 12 Gaussian 03 package. Our results revealed that the solvents have an effect on the optimized 13 parameters. It was observed that the bond lengths increases with an increase in the polarity of the 14 solvents, while the bond angles were found to increase as the polarity of the solvents decreases. 15 The molecule was found to have a higher stability in the gas phase with HOMO-LUMO energy 16 gap of 2.9935eV. The HOMO and HOMO-LUMO energy gap were found to increase with an 17 18 increase in polarity of the solvents. The molecule was found to be harder and less reactive in the gas phase with chemical hardness of 1.4968eV. The maximum value of ionization potential 19 5.3227eV and minimum value of electron affinity 2.2757eV were obtained in water and Gas 20 Phase respectively, as such it is difficult to remove an electron from the molecule in water to 21 22 form an ion and it is also difficult to add an electron to the molecule in Gas Phase. The ground state total energy of the molecule was found to increase with an increase in polarity of the 23 24 solvents. The chemical hardness, chemical softness, electronegativity, chemical potential and electrophilicity index were found to increase with an increase in the dielectric constant of the 25 solvents. In the non-linear optical (NLO) properties calculations, it was observed that Perylene is 26 a neutral molecule. It was found that the specific heat capacity of perylene increases with an 27 increase in the polarity of the solvent while the entropy and the zero-point vibrational energy of 28 the molecule decreases as the polarity of the solvent increases. In the non-linear optical 29 30 properties calculations, it was found that the polarizability ($\langle \alpha \rangle$) of Perylene increases with 31 decrease in the polarity of the solvents and the anisotropic polarizability ($\Delta \alpha$) of Perylene increase with an increase in the polarity of the solvents. In the Natural Bond Orbital (NBO) 32 analysis, high intensive interaction between donor and acceptor electrons of Perylene was 33 34 observed in Chloroform due to large stabilization energy of 4.49Kcal/mol. The result shows that careful selection of the solvents and basis sets can tune the frontier molecular orbitals energy gap 35 and other important optoelectronic properties of Perylene. 36

- **KEYWORDS:** Density Functional Theory, Gaussian 03, HOMO-LUMO, Non-linear Optical
- 38 Properties and Perylene.

1.INTRODUCTION

- With rising demand for sustainably risk free energy, there is no better alternative than organic electronic materials which have proved to be a promising candidates for advanced optoelectronic applications such as in light emitting diodes, photovoltaic[1], organic field effect transistors, organic solar cell and transparent white displays [2, 3]. To fully understand and use these materials, their basic fundamental physical properties must be sufficiently explored. To acquire
- such vital information, we may have to determine their structural, electronic, optical, non-linear
- 47 optical and thermodynamic properties.
- 48 Rylene are series of polycyclic aromatic hydrocarbons (PAHs) with general Chemical formula
- 49 $C_{10n}H_{4n+4}$. Among the family of these compounds, the most intensively investigated is Perylene
- 50 $(C_{20}H_{12})$ and its derivatives such as Perylenebisimides, PTCDA and PDI which due to their
- 51 promising electronic, optical and charge-transport properties are widely used in high-tech
- 52 applications such as organic photovoltaic, organic field effect transistors, bio-labelling, sensors,
- single molecular spectroscopy, super molecular assemblies and opto-electronic devices [4].
- Pleasingly, wide ranges of experimental and theoretical studies have explored the synthesis and
- applications of Perylene and its derivatives. For instance, [5] presented within the framework of
- Density Functional Theory (DFT) a comparative study of the electronic, optical, and transport
- 57 properties of some selected polycyclic aromatic hydrocarbons (Perylene included). Similarly, [6]
- 58 investigated using different exchange correlation functionals the structural, and optoelectronic
- 59 properties of different classes of Perylene; isolated Perylene, diindo[1,2,3-cd:1',2',3'-
- 60 Im]Perylene (DIP) molecule and DIP molecular crystal. [7], studied the effect of aggregation on
- 61 the excited-state electronic structure of Perylene through transient absorption measurements of
- isolated molecules, excimers in solution, monomeric crystal forms ($\beta Perylene$), and dimeric
- crystal forms (α -Perylene). The work of [2], on aggregation of water soluble, dicationic Perylene
- bisimide derivatives using absorption and emission spectroscopies, X-ray and neutron scattering
- 65 techniques as well as electron microscopy, provides evidence for the existence of higher order
- 66 molecular aggregates in solution. [8] investigated theoretically the structure and electronic
- 67 properties of Perylene and Coronene under pressure using density functional theory including

- Vander Waals interactions. The electronic structure of crystalline Perylene was investigated using density functional theory including Vander Waals interaction by [9].
- However, to the best of our knowledge, the effects of solvents on this promising organic material have never been studied before. In the present investigation, computational studies have been carried out to investigate the effects of solvents on the structural, electronic, thermodynamic and non-linear optical properties of Perylene based on density functional theory. The solvents used in this work include Water, Chloroform, Benzene and Acetone with the following dielectric constants; Water ($\varepsilon = 80.37$), Chloroform ($\varepsilon = 4.806$), Benzene ($\varepsilon = 2.284$) and Acetone ($\varepsilon = 37.50$).

2. THEORETICAL BACKGROUND

2.1 Density Functional Theory

Density Functional Theory is a phenomenally successful approach to finding solutions to the fundamental equation that describes the quantum behavior of atoms and molecules [10]. DFT has proved to be highly successful in describing structural and electronic properties in a vast class of materials, ranging from atoms and molecules to simple crystals to complex systems [11]. Density functional theory (DFT) was proposed by Hohenberg and Khon as a method to determine the electronic structure of a system at ground state with a theory stated that all ground state properties for many particle systems are functional of the electron density [12, 13]. In 1965, Khon and Sham (KS) reformulated the problem in a more familiar form and opened the way to practical application of DFT [11]. For a system of non- interacting electrons, the ground state charge density is representable as a sum over one-electron orbitals (KS orbitals) n(r) [11]. That is;

90
$$n(r) = 2\sum_{i} |\Psi_{i}(r)|^{2}$$
 (1)

91 If we assume double occupancy of all states, and the Khon-Sham orbitals are the solution to the Schrodinger equation.

93
$$\left[-\frac{\hbar^2}{2m}\nabla^2 + V_{KS}(r)\right]\Psi_i(r) = \epsilon_i \Psi_i(r)$$
 (2)

94 where,

95
$$V_{KS} = V_{ext} + \int \frac{e^2 n(r) n(r')}{|r - r'|} dr dr' + V_{XC}[n(r)]$$
 (3)

96 is a unique potential having n(r) as its charge density. Thus we have;

97
$$\left(-\frac{\hbar^2}{2m} \nabla^2 + V_H(r) + V_{XC}[n(r)] + V(r) \right) \Psi_i(r) = \epsilon_i \Psi_i(r)$$
 (4)

- where, $V_H(r)$ is the Hartree potential and $V_{XC}[n(r)]$ is the exchange correlation potential.
- 99 A wide variety of different approximations have been developed to take care of the effects of
- 100 electron-electron interactions such as the generalized gradient approximation (GGA), local
- density approximation (LDA) and Hybrid Approximations.
- In the LDA, the exchange correlation energy at a point in space is taken to be that of the
- homogeneous electron gas with local-density \in_{XC} (n). Thus the total exchange correlation
- energy functional is approximated as [14];

$$E_{XC}^{LD} = \int n(r) \in_{XC} (n(r)) dr$$
 (5)

106 From which the potential is obtained as;

$$V_{XC} = \frac{\delta E_{XC}}{\delta n} \tag{6}$$

- where, δE_{XC} and δn are the derivatives of the exchange energy and the electron density
- 109 respectively.
- Whereas, the generalized gradient approximation (GGA) depends on both local density and it's
- 111 gradient, it can be expressed as;

112
$$E_{XC}^{GGA} = \int n(r) \in_{XC} (n|\nabla n|\nabla^2 n) dr$$
 (7)

- where, n(r) is the electron density.
- In different with LDA and GGA, the Hybrid functional is a linear combination Hartree-Fock
- exchanges. It is expressed as [14];

$$E_{XC}^{hybrid} = \alpha E_{XC}^{HF} + E_C \tag{8}$$

where E_{XC}^{HF} is the Hartree-Fock exchange energy and α can be chosen to satisfy particular criteria.

2.2 Global Quantities

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- 119 Global reactivity descriptors such as chemical potential, chemical hardness-softness,
- electronegativity and electrophilicity index are useful quantities in predicting and understanding
- global chemical reactivity trends. The ionization potentials (IP) and electron affinities (EA) of
- the molecule in the gas phase and in solvents are computed using Koopman's Hypothesis,
- through the HOMO and LUMO energy orbitals respectively using the following expressions;

$$IP = -E_{HOMO} \tag{9}$$

$$EA = -E_{LIIMO} \tag{10}$$

- The difference between the highest occupied molecular orbital (HOMO) and lowest unoccupied
- molecular orbital (LUMO) known as energy gap can be obtained from the relation;

$$E_{aan} = E_{LUMO} - E_{HOMO} \approx IP - EA \tag{11}$$

129 Chemical hardness is given by half of the energy band gap [15];

$$\eta = \frac{IP - EA}{2} \tag{12}$$

The softness of a molecule can be obtained by taking the inverse of its chemical hardness [16];

$$S = \frac{1}{\eta} \tag{13}$$

The chemical potential is given by [16];

$$\mu = -\left(\frac{IP + EA}{2}\right) \tag{14}$$

135 The electronegativity is given by [16];

$$\chi = \frac{IP + EA}{2} \tag{15}$$

The electrophilic index is expressed as [16, 17];

$$\omega = \frac{\mu^2}{2\eta} \tag{16}$$

2.3 Non-Linear Optical Properties

- In order to gain an insight on the effects of solvent on the non-linear optical properties (NLO) of
- Perylene, the dipole moment, polarizability, anisotropic polarizability and hyperpolarizability
- were calculated.
- Dipole moment is a property used in describing the polarity of a system. For molecular systems,
- this property can be obtained from [18];

145
$$\mu_{tot} = \left[\mu_x^2 + \mu_y^2 + \mu_z^2\right]^{1/2} \tag{17}$$

- where μ_x , μ_y and μ_z are the components of the dipole moment in x, y and z coordinates.
- 147 Electric dipole polarizability is an important property used in determining the polarizability of a
- molecule or compound. It is a measure of the linear response of an infinitesimal electric field (F)
- and represents second-order variation energy [19];

$$\alpha = -\frac{\partial^2 E}{\partial F_a \partial F_b} \tag{18}$$

where a, and b are coordinates of x, y and z. The mean polarizability is calculated using [16];

152
$$\alpha = \frac{1}{3}(\alpha_{xx} + \alpha_{yy} + \alpha_{zz}) \tag{19}$$

- where the α_{xx} , α_{yy} and α_{zz} quantities are known as principal values of polarizability tensor.
- The anisotropic polarizability is given by [16];

155
$$\Delta \alpha = \left[\frac{(\alpha_{xx} - \alpha_{yy})^2 + (\alpha_{yy} - \alpha_{zz})^2 + (\alpha_{zz} - \alpha_{xx})^2 + 6(\alpha_{xz}^2 + \alpha_{xy}^2 + \alpha_{yz}^2)}{2} \right]^{1/2}$$
 (20)

The mean first hyperpolarizability is defined as [16, 20];

157
$$\beta_{tot} = (\beta_x^2 + \beta_y^2 + \beta_z^2)^{1/2}$$
 (21)

where βx , βy and βz are defined as ;

$$\beta_{x} = \beta_{xxx} + \beta_{xyy} + \beta_{xzz}$$

$$\beta_{y} = \beta_{yyy} + \beta_{xxy} + \beta_{yzz}$$

$$\beta_z = \beta_{zzz} + \beta_{xxz} + \beta_{yyz} \tag{22}$$

- The β_x , β_y and β_z refer to the components of hyperpolarizability along x, y and z components of
- molecular dipole moment.

2.4 Natural Bond Orbital (NBO)

- NBO analysis provides an efficient method for studying intra and intermolecular bonding
- interactions among bonds, and also provides a convenient basis for investigation of charge
- transfer or conjugative interactions in molecular systems [21]. The second-order Fock matrix is
- use to evaluate the donor-acceptor interactions in the NBO basis. For each donor and acceptor,
- the stabilization energy $E^{(2)}$ associated with the electron delocalization between donor and
- acceptor is estimated as [22];

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$$E^{(2)} = -n_{\sigma} \frac{\langle \sigma | F | \sigma \rangle^{2}}{\varepsilon_{\sigma *} - \varepsilon_{\sigma}} = -n_{\sigma} \frac{F_{ij}^{2}}{\Delta E}$$
 (23)

- where $\langle \sigma | F | \sigma \rangle^2$, or F_{ij}^2 is the Fock matrix element between i and j NBO orbitals, ε_{σ^*} and ε_{σ} are
- the energies of $\sigma *$ and σ NBO's and n_{σ} is the population of the donor orbital.

174 3. COMPUTATIONAL METHOD

- 175 The geometry of Perylene was optimized with no symmetry constraint using Becke's three-
- parameter hybrid exchange [23] combined with Lee-Yang-Parr's gradient-corrected correlation
- 177 [24] functional (B3LYP) method with 6-311++G(d,p) basis set. All the parameters were fully
- 178 allowed to relax and each of the calculations converged to an optimized geometry which
- 179 corresponds to a true energy minimum.
- For the study of solvation effects, a Self-Consistent Reaction Field (SCRF) approach based on
- Polarizable Continuum Model (PCM) were employed. The effects of four solvents (water,
- chloroform, benzene and acetone) were investigated by means of the SCRF method based on
- PCM as implemented in the Gaussian 03 [25]. The optimized geometries were then used to

obtain the HOMO-LUMO energy gap, chemical hardness, chemical softness, chemical potential, electronegativity, electrophilcity index, dipole moment, polarizability, anisotropic polarizability, hyperpolarizability, entropy and the specific heat capacity of the investigated molecule at the same level of theory (B3LYP/6-311++G(d,p)). Finally, the NBO calculations [26] were performed using NBO 3.1 program as implemented in the Gaussian 03 package at same level of theory in order to understand the various second-order interactions between the filled orbitals of one subsystem and the vacant orbitals of another subsystem. All calculations were performed within the framework of Density Functional Theory (DFT) as coded in Gaussian 03 package [25].

4. RESULTS AND DISCUSSION

4.1 Optimized Parameters

195 The optimized values of bond lengths and bond angles of the studied molecule were calculated at
196 DFT/B3LYP level using 6-311++G(d,p) basis set in the gas phase and in different solvents
197 (water, chloroform, benzene and acetone). The results are shown in Tables 1 and 2. The distance
198 between the nuclei of two atoms bonded together is termed as bond length while bond angle is
199 the angle between two adjacent bonds of an atom in a molecule [19].

From Table 1, there are little changes in the bond lengths of Perylene when optimized with water, chloroform, benzene and acetone compared with the gas phase. The result shows that the lowest value obtained was 1.0825Å in benzene. However, when compared with results of an isolated Perylene molecule obtained by [6], it was observed that the bond lengths tend to be a little smaller in the present study. It is worth noting that, the smaller the bond length, the higher the bond energy and stronger the bond [27]. Consequently, this has affirmed that the bond lengths of Perylene in the gas phase and in solvents are a little stronger than that of an isolated Perylene molecule. Hence, an enormous amount of energy is required to break these bonds. It was also observed that the bond length increases with an increase in the polarity of the solvents.

Table 1: Selected bond lengths (Å) of the optimized Perylene in the gas phase and in different solvents.

Bond	Gas	Water	Chloroform	Benzene	Acetone	Previous Work
Lengths (Å)	Phase					

R(2,5)	1.4762	1.4773	1.4769	1.4765	1.4772	1.4700 ^a
R(1,4)	1.4339	1.4346	1.4342	1.434	1.4345	1.4360 ^a
R(1,2)	1.4315	1.4324	1.4319	1.4316	1.4323	1.4310 ^a
R(28,32)	1.0858	1.0877	1.0866	1.0858	1.0875	1.0900 ^a
R(14,25)	1.0853	1.0872	1.0862	1.0854	1.087	1.0900 ^a
R(13,24)	1.0827	1.0846	1.0833	1.0825	1.0841	1.0900 ^a

^a(Mohamad *et al.*, 2017)

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Table 2: Selected bond angles (°) of the optimized Perylene in the gas phase and in different solvents.

Bond Angles (°)	Gas Phase	Water	Chloroform	Benzene	Acetone
A(6,3,8)	122.184	122.106	122.130	122.143	122.119
A(3,8,15)	122.001	121.870	121.918	121.944	121.890
A(14,9,18)	120.748	120.919	120.866	120.838	120.900
A(11,20,27)	119.592	119.640	119.626	119.612	119.639
A(20,28,32)	118.965	118.859	118.894	118.905	118.876
A(14,7,16)	117.706	117.733	117.717	117.759	117.679

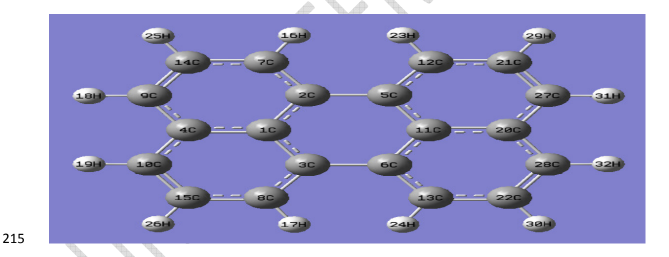


Fig.1 Optimized Structure of Perylene

However, the highest bond length is 1.477Å, which is almost the same in both gas phase and solvents. The structural geometry of the studied molecule that consists of bond lengths and bond angles are found to be in good agreement with the work of [6].

4.2 Frontier Molecular Orbital Energies (FMOEs)

Table 3, presents the highest occupied molecular orbital (HOMO), the lowest unoccupied molecular orbital (LUMO) and HOMO and LUMO energy gaps of Perylene in the gas phase and in solvents calculated at the DFT/B3LYP level in the 6-311++G(d,p) basis set. The values of HOMO, LUMO and their energy gaps reflect the chemical activity of the molecule. The energy gap between HOMO and LUMO determines the kinetic stability, chemical reactivity, optical polarizability and chemical hardness-softness of a molecule [20]. Compounds with large HOMO-LUMO gap value tend to have higher stability [17]. In this work, the order of stability of the molecule is more in the gas phase> water> acetone> chloroform> benzene. Interestingly, the order of stability increases with an increase in polarity of the solvents.

HOMO as an electron donor represents the ability to donate an electron, while LUMO as an electron acceptor represents the ability to accept an electron. The smaller the LUMO and HOMO gaps, the easier it is for the HOMO electron to be excited; the higher the HOMO energies, the easier it is for HOMO to denote electrons; the lower the LUMO energies, the easier it is for the LUMO to accepts electrons [28].

It can be observed from Table 3, that the LUMO energy of Perylene in benzene (-2.2829 eV) is smaller than that in the gas phase and in the rest of the solvents. Hence, the electron transfer from HOMO to LUMO of the molecule in benzene is relatively easier than in the gas phase and in the rest of the solvents. It can also be observed that the HOMO and energy gap increases with increase in polarity of the solvents.

The HOMO-LUMO gap of 2.976 eV obtained in the gas phase is found to be very close to 2.972 eV of an isolated Perylene reported by [6].

Table 3: Calculated HOMO, LUMO and energy gap in (eV) of Perylene in the gas phase and different solvents using B3LYP methods with 6-311++G(d,p) basis set.

Solvents	E_{HOMO}	$E_{LUMO}(eV)$	Gap (eV)	Previous
	(eV)			Work
Gas Phase	-5.2693	-2.2757	2.9935	2.9740^{a}
				2.98^{b}
Water	-5.3227	-2.3332	2.9895	
Chloroform	-5.2878	-2.3005	2.9873	
Benzene	-5.2729	-2.2829	2.9860	

Acetone	-5.3167	-2.3274	2.9893	
^a [6] ^b [5]				

4.3 Total Ground State Energy

Table 4, presents the dielectric constants of the solvents and total energy in atomic mass unit (a.u) of Perylene molecule in the gas phase and in different solvents calculated at the DFT/B3LYP level with 6-311++G(d,p) basis set. The ground state total energy is an important property that describes the physical properties of a molecule. It can be seen from Table 4, that the ground state total energy increases with an increase in dielectric constant of the solvents.

Table 4: Ground state total energies in atomic mass unit (a.u) of the optimized Perylene in the gas phase and different solvents.

Solvents	B3LYP/6-	ε
	311++G(d,p)	
Gas Phase	-769.5611634	
Water	-769.5983746	80.37
Chloroform	-769.5914965	4.806
Benzene	-769.5874674	2.284
Acetone	-769.5954100	37.50

4.4 Ionization Potentials and Electron Affinity

The ionization potential (IP) and electron affinity (EA) measure the tendency of compounds to lose or gain an electron [29]. The IPs and EAs are presented in Table 5. The higher the ionization potential (IP), the more difficult it is to remove an electron to form an ion. The lower the electron affinity (EA), the less easy it is to add an electron.

In Table 5, it can be observed that it is more difficult to remove an electron from water> acetone> chloroform> benzene> gas phase to form an ion. Similarly, it is more difficult to add an electron in terms of their EAs to the molecule in gas phase> benzene>chloroform> acetone > water. It was observed that the ionization potential increases with an increase in the polarity of the solvents while the electron affinity decreases as the polarity of the solvents decreases.

Table 5: Ionization potentials and electron affinities of the optimized Perylene molecule in the gas phase and different solvents.

Solvents	IP (eV)	EA (eV)
Gas Phase	5.2693	2.2757
Water	5.3227	2.3332
Chloroform	5.2878	2.3005
Benzene	5.2729	2.2829
Acetone	5.3167	2.3274

4.5 Global Chemical Indices

The global chemical indices such as chemical hardness, chemical softness, chemical potential, electronegativity and electropilicity index of the molecule in the gas phase and in different solvents were computed and reported in Table 5 using the frontier molecular orbital energy.

Table 6: Global chemical indices of the optimized Perylene in the gas phase and in different solvents.

Solvents	n (eV)	S (eV)	χ(eV)	μ (eV)	ω (eV)
Gas Phase	1.4968	0.6680	3.7725	-3.7725	4.7540
Water	1.4948	0.6690	3.8279	-3.8279	4.9013
Chloroform	1.4937	0.6695	3.7942	-3.7942	4.8188
Benzene	1.4950	0.6689	3.7772	-3.7772	4.7734
Acetone	1.4947	0.6690	3.8221	-3.8221	4.8867

Chemical hardness is proportional to the HOMO-LUMO energy gap. An Increase in the chemical hardness makes the molecule more stable and less reactive. As seen in Table 6, Perylene molecule in the gas phase has slightly the highest value of chemical hardness of 1.4968eV is considered to be harder and more stable than in the rest of the solvents, followed by benzene, water and acetone with chemical hardness of 1.4950eV, 1.4948eV and 1.4947eV

respectively. This indicates that Perylene in chloroform is more stable than in the rest of the solvents.

4.6 Thermodynamic Properties

Table 7, Presents the components and total contribution of the electronic, translational, rotational and vibrational energies to the entropy (S) and heat capacity (Cv) as well as the rotational constants and zero-point vibrational energies (ZPVE) of Perylene in the gas phase and in different solvents.

Table 7: Thermodynamic properties of the optimized Perylene molecule in the gas phase and different solvents.

	Gas Ph	ase	Water		Chlorof	orm	Benzene	2	Acetone	•
Positio	Cv	S	Cv	S	Cv	S	Cv	S	Cv	S
ns	(Kcal/	(Kcal/	(Kcal/	(Kcal/	(Kcal/	(Kcal/	(Kcal/	(Kcal/	(Kcal/	(Kcal/
	Mol)	Mol)	Mol)	Mol)	Mol)	Mol)	Mol)	Mol)	Mol)	Mol)
Electro	0	0	0	0	0	0	0	0	0	0
nic										
Transla	2.981	42.47	2.981	42.47	2.981	42.47	2.981	42.47	2.981	42.47
tional		4		4		4		4		4
Rotatio	2.981	33.24	2.981	33.24	2.981	33.24	2.981	33.24	2.981	33.24
nal		2		6		4		3		6
Vibrati	50.95	37.39	50.96	36.33	50.93	36.74	50.91	37.04	50.96	36.45
onal	1	8	1	1	9	5	3	5	4	4
Total	56.91	113.1	56.92	112.0	56.90	112.4	56.87	112.7	56.92	112.1
	3	14	2	51	1	63	5	62	5	74
Rotation	al 🦪	0.625	0.62372	2	0.62427	1	0.62464	-	0.62388	}
Constant	s	04								
(GHz)		0.330	0.33026)	0.33042		0.33056)	0.33027	'
		6								
•		0.216	0.21593	}	0.21606)	0.21616)	0.21595	
23		23								
ZPVE		158.5	157.371	17	157.801	91	158.123	72	157.487	19
(Kcal/Mo	ol)	5182								

It can be observed in Table 7 that specific heat capacity of perylene is found to increase with an increase in the polarity of the solvents, while the entropy decreases as the dielectric constant

increases. The zero-point vibrational energy (ZPVE) decreases with an increase in the polarity of the solvents.

4.7 Non-Linear Optical Properties

Our investigation also highlighted the effects of solvents on the nonlinear optical properties of the molecule. This is necessary for sufficient understanding of the nonlinear optical response of the molecule. Nonlinear optical (NLO) effect arises from the interactions of electromagnetic fields in various media to produce new fields altered in phase, frequency, amplitude and other propagation characteristics from the incident fields [30].

Hence, Table 8, presents the non-linear optical properties in atomic mass unit (a.u) of Perylene molecule in the gas phase and in solvents. The properties computed and reported are dipole moment (μ_{tot}), polarizability ($\langle \alpha \rangle$), anisotropic polarizability ($\Delta \alpha$) and hyperpolarizability (β_{tot}). **Table 8**: Non-linear optical properties (in Debye) of the optimized Perylene molecule in the gas phase and different solvents.

Solvents	μ_{tot}	$\langle \alpha \rangle$	$\langle \Delta \alpha \rangle$	eta_{tot}
Gas Phase	0.0000	108.2104	24.4360	1.00 E-4
Gas I Hase	0.0000	100.2107	27.7300	1.00 L-4
Water	0.0000	104.3552	31.0839	4.36 E-4
Chloroform	0.0000	105.9012	28.2349	3.00 E-4
Benzene	0.0000	107.0436	26.1246	1.41 E-4
Acetone	0.0000	104.6985	30.4309	2.83 E-4

The dipole moment in a molecule is an important electronic property which results from non-uniform distribution of charges on the various atoms in the molecule [20]. It can be observed in Table 8, that Perylene is a neutral molecule with a dipole moment of $0.000 \, \text{eV}$ in both the gas phase and in the solvents. It can also be observed that the polarizability of Perylene increases as the polarity of the solvents decreases whereas the anisotropic polarizability increases with an increase in the polarity of the solvents. Consequently, increasing or decreasing the polarity of the solvents plays a significant role in determining the values of the non-linear optical properties of Perylene.

4.8 Natural Bond Orbital (NBO) Analysis

Natural Bond Orbital (NBO) analysis provides an efficient method for studying intra-and intermolecular interaction among bonds and also provides a convenient basis for investigating charge transfer or conjugative interactions in molecular systems. Table 9 presents the Natural Bond Orbital analysis of Perylene in the gas phase and in different solvents.

Table 9: Natural Bond Orbital (NBO) of optimized Perylene molecule in the gas phase and in different solvents.

		-(2)		T	T		(2)			
Donor	Acceptor	$E^{(2)}$	E(j)-E(i)	Fji a.u	Donor	Acceptor	$E^{(2)}$	E(j)-E(i)	Fji a.u	
NBO (i)	NBO (j)	Kcal/mol	a.u		NBO (i)	NBO (j)	Kcal/mol	a.u		
		Water			Chloroform					
σC2-C5	σ^* C1-C2	1.40	1.18	0.036	σC2-C5	σ*C1-C3	4.29	1.16	0.052	
σC2-C5	σ*C1-C4	4.03	1.18	0.062	σC2-C5	σ*C1-C4	3.88	1.16	0.060	
σC2-C5	σ*C2-C7	2.81	1.22	0.052	σC2-C5	σ*C2-C5	3.27	1.10	0.054	
	σ*C5-									
σC2-C5	C11	2.33	1.16	0.046	σC2-C5	σ*C2-C7	4.49	1.21	0.066	
	σ*C5-									
σC2-C5	C12	3.74	1.20	0.057	σC2-C5	σ*C3-C8	2.27	1.23	0.047	
	σ*C7-					σ*C4-				
σC2-C5	C14	1.95	1.21	0.044	σC2-C5	C10	2.41	1.18	0.048	
	σ*C11-					σ*C5-				
σC2-C5	C20	2.66	1.22	0.049	σC2-C5	C12	2.30	1.23	0.048	
	σ*C12-		X X			σ*C7-				
σC2-C5	C21	1.87	1.20	0.042	σC2-C5	H16	2.37	1.11	0.046	
σC11-					σC11-					
C20	σ*C2-C5	2.61	1.13	0.049	C20	σ*C2-C5	3.10	1.09	0.052	
σC11-		44			σC11-					
C20	σ *C3-C6	2.58	1.11	0.048	C20	σ*C3-C6	2.91	1.09	0.050	
σC11-	σ*C5-				σC11-	σ*C5-				
C20	C11	4.04	1.17	0.062	C20	C11	3.07	1.16	0.053	
σC11-	σ*C6-				σC11-	σ*C6-				
C20	C11	3.89	1.17	0.060	C20	C11	3.36	1.15	0.056	
σC11-	σ*C20-				σC11-	σ*C20-				
C20	C27	2.69	1.20	0.051	C20	C27	3.73	1.18	0.060	
σC11-	σ*C20-				σC11-	σ*C20-				
C20	28	2.71	1.20	0.051	C20	C28	3.72	1.18	0.060	
σC11-	σ*C27-				σC11-	σ*C27-				
C20	H31	2.35	1.11	0.046	C20	H31	1.81	1.10	0.040	
σC11-	σ*C28-				σC11-	σ*C28-				
C20	H32	2.36	1.11	0.046	C20	H32	1.85	1.10	0.041	
		Benzene			Gas Phase					
Belizerie					233 : 11000					

								, ,	
σC1-C2	σ*C1-C3	4.31	1.16	0.063	σC1-C2	σ*C1-C3	4.30	1.16	0.063
σC1-C2	σ*C1-C4	4.31	1.17	0.063	σC1-C2	σ*C1-C4	4.31	1.17	0.063
σC1-C2	σ*C2-C5	2.49	1.12	0.047	σC1-C2	σ*C2-C5	2.49	1.12	0.047
σC1-C2	σ*C2-C7	3.75	1.25	0.061	σC1-C2	σ*C2-C7	3.75	1.25	0.061
σC1-C2	σ*C3-C8	2.29	1.21	0.047	σC1-C2	σ*C3-C8	2.29	1.21	0.047
	σ*C4-					σ*C4-			
σC1-C2	C10	2.67	1.20	0.051	σC1-C2	C10	2.67	1.20	0.051
	σ*C5-					σ*C5-			
σC1-C2	C12	2.23	1.21	0.047	σC1-C2	C12	2.23	1.21	0.047
	σ*C7-					σ*C7-	*		
σC1-C2	H16	2.42	1.12	0.047	σC1-C2	H16	2.45	1.11	0.047
σC11-	# 62 65	2.07	4.40	0.050	σC11-	# CO CE	2.00		0.050
C20	σ*C2-C5	3.07	1.12	0.052	C20	σ*C2-C5	3.06	1.12	0.052
σC11-	-*C2 C6	2 24	1 11	0.045	σC11- C20	σ*C3-C6	2.24	1 11	0.045
C20 σC11-	σ*C3-C6 σ*C5-	2.24	1.11	0.045	σC11-	σ^*C5-C6 σ^*C5-	2.24	1.11	0.045
C20	C11	4.25	1.16	0.063	C20	C11	4.24	1.16	0.063
σC11-	σ*C6-	4.23	1.10	0.003	σC11-	σ*C6-	4.24	1.10	0.003
C20	C11	3.95	1.16	0.061	C20	C11	3.94	1.16	0.061
σC11-	σ*C20-	0.00		0.002	σC11-	σ*C20-	0.0 .	2.20	0.001
C20	C27	3.83	1.18	0.060	C20	C27	3.86	1.18	0.060
σC11-	σ*C20-				σC11-	σ*C20-			
C20	C28	4.03	1.19	0.062	C20	C28	4.06	1.19	0.062
σC11-	σ*C27-				σC11-	σ*C27-			
C20	H31	1.92	1.10	0.041	C20	H31	1.95	1.10	0.042
σC11-	σ*C28-				σC11-	σ*C28-			
C20	H32	1.80	1.10	0.040	C20	H32	1.83	1.10	0.040
	ı	Acetone							
σC1-C2	σ*C1-C3	4.30	1.16	0.063					
σC1-C2	σ*C1-C4	3.89	1.16	0.060					
σC1-C2	σ*C2-C5	3.27	1.10	0.054					
σC1-C2	σ*C2-C7	4.48	1.21	0.066					
σC1-C2	σ*C3-C8	2.27	1.23	0.047					
	σ*C4-								
σC1-C2	C10	2.24	1.18	0.048					
	σ*C5-								
σC1-C2	C12	2.31	1.23	0.048					
	σ*C7-								
σC1-C2	H16	2.34	1.12	0.046					
σC11-	1.00.07	2.44		0.0=5					
C20	σ*C2-C5	3.11	1.09	0.052					
σC11-	-*C2 CC	2.02	1.00	0.054					
C20	σ*C3-C6	2.92	1.09	0.051					
σC11- C20	σ*C5-	2 07	1 16	U UE3					
CZU	C11	3.07	1.16	0.053					

σC11-	σ*C6-			
C20	C11	3.37	1.15	0.056
σC11-	σ*C20-			
C20	C27	3.71	1.18	0.060
σC11-	σ*C20-			
C20	C28	3.70	1.18	0.059
σC11-	σ*C27-			
C20	H31	1.78	1.10	0.040
σC11-	σ*C28-			
C20	H32	1.38	1.10	0.040

The larger $E^{(2)}$ value, the more intensive is the interaction between electron donors and acceptors. The more donation tendency from electron donors to electron acceptors, the greater is the extent of conjugation of the whole system[22]. It can be seen from Table 9, that larger value of stabilization (4.49 Kcal/mol) energy $E^{(2)}$ of Perylene was obtained in chloroform. Hence, there is high intensive interactions between σ C1-C2 and σ *C2-C7 and has greater conjugation in the molecule. The order of this interaction is more in Chloroform>Acetone>Benzene>Gas Phase>Water. This phenomenon occurs as the polarity decreases between chloroform and benzene.

5. CONCLUSION

To understand the effects of solvents on structural, electronic, thermodynamic and non-linear optical properties of Perylene molecule, we have carried out an extensive computational study of the HOMO, LUMU, HOMO-LUMO energy gap, ionization potential, electron affinity, chemical hardness, chemical softness, chemical potential, electronegativity, electrophilicity index, dipole moment, polarizability, anisotropic polarizability, hyperpolarizability, entropy, heat capacity, rotational constants and zero-point vibrational energy using the B3LYP methods under 6-311++G(d,p) basis set.

In the structural properties calculations, Our findings revealed that the bonds of Perylene tend to be stronger in the gas phase and in solvents compared to that of an isolated Perylene as reported in the literature. It was observed that the bond lengths increases with an increase in the polarity of the solvents, while the bond angles were found to increase as the polarity of the solvents decreases. In the global quantities calculations, it was found that, the electron transfer from HOMO to LUMO was found to be relatively easier in chloroform than in the gas phase an in the

- rest of the solvents. The global quantities, HOMO and HOMO-LUMO energy gaps were found to increase as the polarity of the solvents increases. The ground state energy of Perylene was found to increase with decrease in polarity of the solvents.
- 351 In the Thermodynamic part of our work the specific heat capacity of Perylene increases with an 352 increase in the polarity of the solvents while the entropy and the zero-point vibrational energy decreases as the polarity of the molecule increases. In the non-linear optical properties 353 354 calculations, the polarizability increases with decrease in the dielectric constant of the solvents while the anisotropic polarizability increases as the polarity of the solvents increases. In the NBO 355 analysis, high intensive interaction between donor and acceptor electrons was observed in 356 chloroform due to large value of stabilization energy. The results also show that careful selection 357 358 of basis set and solvents can be utilized to tune the optoelectronic properties of Perylene.

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