

Research Article

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Adsorption of a biological active ethionamide over the Surface of a Fe- porphyrin induced carbon nancone (Fe-PICNC) system Through a Density Functional Theory (DFT)

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1.Introduction

Ethionamide (ET) is a compound with molecular formula of $C_8H_{10}N_2S$ which is a drug used for pulmonary tuberculosis and extra-pulmonary tuberculosis. The ET drug fits to a group of the drugs named antibiotics. A number of scientists recently have interested in the environmental effects of the drugs that gathered in the environment in large amounts and their side effects on living organisms are dangerous which should be important the investigation of this matter [1, 2]. In many studies, more than 70 drugs have been found as the drug contaminants in soil and water of different countries [3, 4]. The Drug accumulated in the environment affect health of human, animal, and plants [5-8]. It is necessary that the accumulated drugs should

ABSTRACT

This study used Density Functional Theory (DFT) calculations to investigate the adsorption behavior of ethionamide (ET) on Fe-porphyrin induced carbon nanocones (Fe-PICNCs). The B3LYP method with the 6-31G* basis set was employed to determine total energies, physical parameters, and electronic analysis. Results indicated that the ET molecule strongly adsorbs onto Fe-PICNCs, with adsorption energies ranging from -30.96 to -15.00 kcal/mol, indicating very strong chemical interactions. The findings suggest that Fe-PICNCs are effective adsorbents for removing unwanted ET drug emissions from the environment. The energy gap (work function) of isolated Fe-PICNCs was calculated as 2.00 eV (3.82 eV). ET adsorption had an insignificant effect on the energy gap (work function) of the Fe-PICNC structures, indicating that they cannot be used as electronic conductivity type or work function type sensors for the ET drug.

be identified by a sensor to find where the drug gathered. Nanoparticles are the materials that can be used as the sensors for the identification of the released drugs [9, 10]. These nanoparticles have received much attention as chemical sensors and other applications as drug delivery, batteries and catalysts [11-13]. Graphene is provided by a variety of methods, such as chemical vapor deposition (CVD) mechanical exfoliation, chemical production, epitaxial growth on silicon carbide (SiC) [14-16].

Porphyrins are important aromatic macrocycle organic structures that play a vital role in biological functions such as respiration, electron transport, and photosynthesis [17-19]. These structures consist of four functionalized pyrroles connected by methine bridges

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(=CH-). Porphyrins and their related compounds have a wide range of applications in medicine, the energy sector, and the chemical industry [20]. Because of their strong light absorption capabilities, porphyrins are being studied for their potential use in non-invasive cancer treatment through photodynamic therapy (PDT), as well as for their potential as anticancer and antioxidant materials [21-23]. Metalloporphyrins are porphyrin cycles that have metal ions inserted into them and have been extensively studied as catalysts in organic oxidation synthesis, of organic compounds, photocatalytic water splitting, and dye-sensitized solar cells [24-27]. They have also been evaluated for use in chemical sensors, such as detecting trace amounts of TNT vapors and volatile organic compounds in human breath [28-33]. Carbon nanomaterials have also attracted significant interest among researchers due to their exceptional properties and high surface area, making them useful for a wide range of applications, including gas adsorption, detection, and catalysis [34-42].

The manuscript explores the potential of Feporphyrine-induced carbon nanocone (Fe-PICNC) as a sensing material for identifying and adsorbing the ET drug, using DFT calculations. The findings of this study have significant implications for the development of an effective adsorbent for the ET drug, as well as for designing a ssionsuitable sensing mechanism.

2.Computational Details

This study employed the DFT method at the B3LYP/6-31G(d) theoretical level, utilizing GAMESS programs for all calculations [43]. The B3LYP method is commonly used in the study of materials based on nanostructures [44-46]. Moreover, a review of the literature suggests that it is a dependable and appropriate density functional for predicting the energetic, electronic, structural, and optical properties of various nanostructures [47-53]. The DFT method is one of the most powerful approaches for quantum calculations.

During the optimization process, desirable molecular structures are optimized to reach the minimum energy, and the energy of the targeted systems is subsequently computed [54] To introduce a defect in the carbon nanocone, a PI ring consisting of 4 nitrogen atoms and 20 carbon atoms was added at the apex of the cone, resembling the structure of PI. The Gaussview software was employed to substitute the PI structure with the defect in the carbon nanocones (PICNC). Optimization calculations were conducted for both the complete Nanocone and the PICNC structure. Finally, the GaussSum program was used to generate DOS plots [55]. The adsorption energy (E_{ads}) is calculated as follows:

 $E_{ads} = E_{(drug/adsorbent)} - [E_{(adsorbent)} + E_{(drugt)}] + E_{BSSE}$ (1)

The adsorption energy of an ET molecule onto a Fe-PICNC molecule is denoted by the term Eads. Specifically, $E_{(drug)}$ represents the energy of the ET molecule, while $E_{(adsorbent)}$ is the total energy of the Fe-PICNC molecule. The total energy of the adsorbed ET molecule on the surface of the Fe-PICNC molecule can be expressed as $E_{(drug/adsorbent)}$. Additionally, the Basis Set Superposition Error (BSSE) is accounted for using the counterpoise method [56], and in this study, we evaluate some interaction indicators that are useful for analysis. In this study, the Natural Bond Orbitals (NBO) charge analysis [57] was conducted at the same level of theory. The HOMO-LUMO energy gap (E_g) is defined as:

 $E_g = E_{LUMO}-E_{HOMO}$ (2) The E_{HOMO} and E_{LUMO} orbitals represent the energy levels of the highest occupied and lowest unoccupied orbitals, respectively. It is typical to assume that the Fermi level (E_F) is positioned near the center of the molecule's energy gap (E_g) at 0 K.

3.Results and Discussions

3.1. Optimizing the Geometry of Pristine Carbon Nanocones (CNC) and Porphyrin-Induced Carbon Nanocones (PICNCs)

In our calculations, we observed a low energy gap value (0.59 eV) between the HOMO and LUMO levels in the carbon nanocone (CNC) structure. However, replacing the porphyrin ring in a CNC resulted in a slight increase in the energy gap value (1.22 eV) (Table 1), indicating a potential for the modified structure to act as a semiconductor. Despite this change, the binding energy and overall stability of the system remained relatively unchanged. After optimization calculations on the substituted porphyrins in the carbon nanocone (PICNC), we found that the resulting hole size is suitable for accommodating different metals.

We anticipate that binding the nitrogen atoms in the cavity with the atoms of the transition metals could effectively prevent clustering of these metals. The results of the optimization calculations, encompassing the energy levels of the HOMO and LUMO orbitals, the energy gap (E_g) between them, Fermi level energy, bipolar moment, and work function, are presented in Table 1.



Fig 1. A schematic formation of porphyrins and Fe-porphyrins-induced in carbon nanocones.

Table 1: Energy of HOMO and LUMO orbitals, energy difference of HOMO and LUMO orbitals (E_g) , Fermi level energy (E_F) , bipolar moment (DM) and working function (Φ) for ethionamide molecule (ET), carbon nanocone (CNC), porphyrins-induced carbon nanocone (PICNC), and Fe-PICNC at the theoretical level of B3LYP/6-31G(d)

Systems	E _{HOMO} (eV)	E _F (eV)	E _{LUMO} (eV)	E _g (eV)	DM(Debye)	Φ
ET	-5.78	-3.46	-1.15	4.63	-	3.60
CNC	-3.98	-3.68	-3.39	0.59	7.79	3.68
PICNC	-5.00	-4.39	-3.78	1.22	10.35	4.39
Fe- PICNC	-4.82	-3.82	-2.82	2.00	7.80	3.82

3.1. Investigating the effects of Fe-porphyrin-induced on the structure of carbon nanocone (Fe-PICNC)

 Fe^{2+} can be located at the center of the porphyrin central cavity depending on its size. To investigate how porphyrin-induced defects affect carbon nanocones, divalent iron ion substituted at the center of porphyrin molecules from the transition elements in carbon nanocones. The team then performed optimization calculations for three models, namely, the carbon nanocone (CNC), the porphyrin ring-induced model (PICNC), and the Fe-porphyrin-induced model in carbon nanocones (Fe-PICNC), as illustrated in Fig 1. These calculations aimed to identify the structural changes resulting from these modifications.Table 1 presents evidence that the HOMO and LUMO orbitals of Fe-PICNC, a type of porphyrin-induced carbon nanocone, undergo alterations when a Fe ion is inserted into the molecule's vacant cavity. Notably, replacing the Fe cation in the porphyrin central cavity leads to a significant increase in Eg as compared to isolated PICNC (as shown in Table 1). These findings imply that the introduction of a Fe cation into PICNC induces a shift in the energy levels between the HOMO and LUMO orbitals, which suggests the emergence of new HOMO and LUMO orbitals with lower energy levels. The TDOS diagram of Fe-PICNC has been shown in Fig 2, too. The absorption of Fe^{2+} causes a shift in the Fermi level. The energy gap of the porphyrin molecule in the carbon nanocone is approximately 1.22 eV, indicating that the compound exhibits semiconductor characteristics. Changes to the Fermi level of a semiconductor due to the absorption

of metal ions can modify the field emission currents. The carbon nanocone being studied has a conjugated structure, resulting in low polarity. However, by creating a cavity and adding nitrogen with high electronegativity, the dipole moment vector in this structure will be oriented towards the nitrogen atoms, resulting in higher polarity (See Table 1). On the other hand, by placing Fe cation with lower electronegativity than nitrogen in the empty cavity, the polarity becomes neutral, and as a result, the dipole moment decreases. As seen from Table 1, placing a Fe ion at the center of the cavity equals 7.80 Debye in comparison; the dipole moment of PICNC equals 10.35 Debye.

Energy (eV)

TDOS of Fe-PICNC

3.2. Adsorption of ET molecule on Fe-PICNC

Fig 2. TDOS diagram of Fe-PICNC structure.

Our study investigated the interaction between the ET molecule and Fe-PICNC. To determine the most favorable adsorption energy, we analyzed how the ET molecule interacts with the Fe cation within Fe-PICNC from external sites.

Fig 3 illustrates the ET molecule interacting with Fe-PICNC, with the ET molecule located at the center of the porphyrin ring from an external site. DFT calculations using the B3LYP method and 6-31G(d) basis set were conducted to investigate the electrostatic properties of the ET molecule and the outer surfaces of Fe-PICNC. We conducted scans of various dihedral angles to identify the most stable adsorption configuration (as shown in Fig 3). Table 2 presents the E_{ads} values computed using Equation (1) for the adsorption of ET onto Fe-PICNC molecules.



Density of state

0 -12 -10 -8



S2



S3

Fig 3. Optimized structures for the adsorption of ET molecules from the outer surface of Fe-PICNC.

Table 2. Absorption energies (E_{ads}) , energy of HOMO and LUMO orbitals, energy difference of HOMO and LUMO orbitals (E_g) , Fermi level energy (E_F) , and dipole moment (DM) and working function for (Fe-PICNC) after adsorption of ET molecule at the theoretical level.

Systems	E _{ads} (kcal/mol)	E _{HOMO} (eV)	E _F (eV)	E _{LUMO} (eV)	Eg (eV)	$\%\Delta E_g$	DM (Debye)	Ф (eV)	%ΔΦ
S1	-30.96	-4.71	-3.71	-2.71	2.005	0.25	4.20	3.71	-2.92
S2	-23.76	-4.76	-3.76	-2.76	2.002	0.1	6.39	3.76	-1.71
S3	-15.00	-4.78	-3.80	-2.82	1.955	-2.25	4.89	3.80	-0.59

Table 2 shows that following optimization, three stable adsorption configurations were observed, where the ET molecule is in contact with Fe-PICNC structure. These configurations are: S1) the S atom of ET molecule is bonded with Fe atom of the external surface of Fe-PICNC structure, S2) the N atom of pyridine ring is bonded with Fe atom of Fe-PICNC structure, and S3) the N atom of amine group is bonded with Fe atom of Fe-PICNC. The adsorption energies (E_{ads}) for each configuration were also determined and are listed in Table 2 as -30.96, -23.76, and -15.00 kcal/mol for S1, S2, and S3 configurations, respectively.

The negative adsorption energy values in Table 2 indicate that the structures are stable due to a very

strong chemisorption interaction between the ET molecule and Fe-PICNC. This suggests that the ET molecule can interact with the Fe atom in Fe-PICNC structures via its S, N (pyridine), or amine groups, resulting in stable configurations (as shown in Fig 3). The S-Fe and N-Fe interaction distances for these configurations are 2.213, 1.895, and 1.986 \Box , respectively (as listed in Table 3). Fe-N bond lengths in Fe-PICNC increase after ET adsorption, as shown in Table 3. Therefore, Fe-PICNC can be used as a suitable adsorbent for removing unwanted ET drug emissions in the environment.

Table 3. Bond length and angle bonds obtained from optimization calculations at the theoretical level B3LYP/6-31G(d) related to Fe-PICNC after ET molecule adsorption process							
Systems	Fe-N1	Fe-N2	Fe-N3	Fe-N4	Fe-ETH		
Fe- PICNC	1.947	1.948	1.947	1.948	-		
S1	2.023	1.978	1.993	1.977	2.213		
S2	2.002	2.003	2.010	2.009	1.895		
S 3	2.003	1.993	1.980	1.982	1.986		







Fig 4. Total density of states (TDOS) and partial density of states (PDOS) of S1, S2, and S3 configurations (The dash line shows Fermi level).

The electronic properties, specifically E_{HOMO} , E_{LUMO} , and E_g , were also investigated in our study. These parameters are important in explaining the behavior of molecules. Table 2 shows that the change in E_g for the configurations resulting from ET adsorption on the outer surface of Fe-PICNC, compared to the respective adsorbents, is insignificant (% ΔE = 0.25, 0.1, and -2.25). The negligible change in E_g is due to the HOMO and LUMO levels of these configurations remaining unchanged compared to their adsorbents. We also analyzed the electronic properties of Fe-PICNC after ET

When it comes to detecting potential, there are two key factors to consider: E_{ads} and E_g . If the Eads falls within a certain range, the adsorption of the ET molecule onto Fe-PICNC can be reversed. However, stronger interactions are not ideal for sensing ET because they result in longer recovery times and make it harder to remove the drug molecule from Fe-PICNC. This relationship can be mathematically expressed as follows: [58, 59]:

 $\tau = v_0^{-1} \exp\left(-E_{ads}/kT\right)$ (3)

The equation given establishes a relationship between recovery time (τ) and attempt frequency (v_0), and the physical parameters of temperature (T) and Boltzmann constant (k). [60]. Equation (3) demonstrates an exponential correlation between E_{ads} and recovery time, indicating that as E_{ads} increases, the recovery time for the sensor also increases. Sensor recovery is a crucial process that can be performed at room adsorption in three configurations, as presented in Table 2 and Fig 4.

Its show an insignificant change in the TDOS of the considered configurations compared to Fe-PICNC. Specifically, we did not observe new LUMO and HOMO levels in these configurations compared to Fe-PICNC. Furthermore, the TDOS and PDOS plots confirm that the valence and conduction levels in S1, S2, and configurations did not shift notably, leading to an insignificant change in the E_g value. According to the PDOS diagrams, HOMO and LUMO of these configurations consist of valence orbitals of Fe atom of Fe-PICNC.

temperature or higher, as suggested by some sources. [61]. The sensing ability of Fe-PICNC is also impacted by another crucial parameter, namely the HOMO-LUMO energy gap (E_g) in the presence of the ET molecule. Equation (4) demonstrates that E_g is directly proportional to the population of conduction electrons (σ), which increases when ET is adsorbed onto Fe-PICNC, leading to a reduction in the HOMO-LUMO energy gap (E_g). In contrast, increasing the value of % ΔE_g also enhances the sensing potential. The relationship between E_g and the electrical conductance of nanoparticles can be mathematically expressed as follows:

 $\sigma = A T^{3/2} \exp(-E_{\sigma}/2kT)$ (4)

The equation includes a constant A (measured in electrons/ $^{m3}K^{3/2}$) and the Boltzmann constant k. Moreover, the outcomes obtained through this method exhibit a clear correlation with the experimental methods described in the scientific literature. [62].