DFT Study About Adsorption of a C\textsubscript{14} Carbon Ring on An Organochlorinated DDE Compound

Yasminne C Esquivel Aguilar, Juan H Pacheco Sánchez*, Genoveva García Rosales

TecNM / Instituto Tecnológico de Toluca, Mexico

*Corresponding author: Juan H Pacheco-Sánchez, TecNM / Technological Institute of Toluca, Av. Tecnológico s/n, Col. Agrícola Bellavista, Metepec, 52149 Mexico.

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Abstract

DFT study about adsorption of one C\textsubscript{14} carbon ring molecule on an organochlorinated DDE (Dichlorodiphenyl dichloroethylene) compound has been developed by taking DDE as a metabolite of DDT (Dichlorodiphenyltrichloroethane) pesticide, which has had negative environmental effects causing diseases in living beings (including human beings), so it is necessary to analyze alternative models allowing to remove this pollutant in order to minimize its effect. Molecular modeling is a computational alternative that will allow geometry optimization of DDE and calculation of its reactivity properties, for which DMol\textsubscript{3} of Materials Studio is used. The latter uses Density Functional Theory (DFT) methodology based on GGA/PW91 functional, spin non restricted; at the level of self-consistent field electron (SFC), with DND basis, and valence atomic basis 6-31G*.

The properties of reactivity such as chemical potential, electronegativity, hardness and chemical softness and Fukui indexes yielded data that will allow molecular interactions between C\textsubscript{14} carbon rings and DDE to help its adsorption and thus minimize the damage caused by this organochlorine compound. Taking C\textsubscript{14} carbon ring as activated carbon structure, and adsorbing it on DDE molecule, it is observed that placing such structures in a certain orientation where HOMO and LUMO of both DDE and carbon complement each other to achieve adsorption, because of the reactivity study in DFT.

Keywords: DFT; Geometry optimization; Activated carbon; DDE

Introduction

The aim in this case is to get a DFT study on the interaction between a C\textsubscript{14} activated carbon ring and an organochlorinated DDE (Dichlorodiphenyl dichloroethylene) compound for its possible adsorption, in order to increase DDE compound. The latter is a metabolite of DDT (Dichlorodiphenyltrichloroethane) pesticide, which has had negative environmental effects causing diseases in living beings, so it is necessary to analyze alternative models allowing to remove this pollutant in order to minimize its effect. Molecular modeling is a computational alternative that will allow geometry optimization of DDE and calculation of its reactivity properties, for which DMol\textsubscript{3} of Materials Studio is used. The latter uses Density Functional Theory (DFT) methodology based on GGA/PW91 functional, spin non restricted; at the level of self-consistent field electron (SFC), with DND basis, and valence atomic basis 6-31G*.

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Figure 1: a) Rings C\textsubscript{14} activated carbon b) DDE

One of the carbon properties is that with atoms of the same element, it is possible to give place to different allotropic forms of car-
bon materials: graphite, graphene, diamond, nanotubes, fullerenes, nanofoam and carbyne type rings by to mention some [7]. They are differentiated by the way in which the carbon atoms are structurally arranged, hence the chemical, physical and mechanical properties of each material arise. For this reason, both theoretical and experimental, a variety of studies have been carried out on how carbon structure geometry impacts in the properties of each material [8]. The adsorption as carbon property, is a process by which the atoms on a solid surface, attract and retain molecules of other compounds. These attraction forces are known as “Van Der Waals” forces (physisorption) [9]. Therefore, given that this is a phenomenon occurring on the surface, the greater the surface area available on a solid, the better adsorbent can be

On the interaction of DDE molecule with C$_{14}$, carbon ring, reactivity properties of each one of them were estimated: chemical potential, chemical hardness and electrophilic index, considering the molecular orbitales HOMO and LUMO for the adsorption analysis by applying geometry optimization [10,11]. In addition, the GAP is also calculated using HOMO-LUMO energies [12]. Geometry optimization is an iterative procedure that tries to locate a minimum of energy by varying lengths and angles up to equilibrium state where the molecular structures are in stable [13]. When atoms and molecules approach each other, intermolecular forces begin to emerge. At very short distances these are repulsion forces; however, as the separation increases, they change to be of attraction. These forces generate a well of asymmetric potential similar to that of Lennard-Jones, which shows the position of the minimum potential energy, indicating equilibrium distance and bond force magnitude of the potential well [14].

To calculate reactivity indexes by means of the formulas proposed by Koopmans [15], the HOMO and LUMO energies of each C$_{14}$ and DDE molecules have been obtained. HOMO (the highest occupied molecular orbital), is the occupied OM of greater energy and the first empty molecular orbital is the LUMO (the lowest unoccupied molecular orbital) which is of lower energy. The reactivity between two molecules increases according to the extent that the HOMO of one of them and the LUMO of the other are closer [16].

In the Density Functional Theory (DFT), the energy can be expressed in terms of electrons number N, and the external potential V, which provides a series of mathematical expressions that form a set of global and local quantities that allow to quantify the reactivity concept [17]. To study reactivity indexes of molecules, the following descriptors are calculated: Chemical potential $\mu$ which measures the tendency of electrons to escape from the electronic cloud, chemical hardness $\eta$ global ownership of the system that measures the resistance imposed by this to the change of its electronic distribution [18]. Similarly, the electronegativity $\chi$ is calculated, a property that measures tendency of electrons to remain in balance system [19].

Fukui indexes are local reactivity indexes giving us information on the atoms of the molecule that have a greater tendency to lose or accept an electron, those who are more likely to suffer a nucleophilicity or electrophilicity attack [20]. These indexes represent the sensitivity of the chemical potential $\mu$ due to the change of electronic density when the number of electrons changes [21]. The electrophilicity $\omega$ is associated with the energy of lowest unoccupied molecular orbital (LUMO) and measures reactivity towards a donor reagent. The nucleophilicity $N$ is associated with the highest energy molecular orbital (HOMO) and measures reactivity towards an acceptor receptor [22]. The electrophilicity $\omega$ scale allowed the classification of organic molecules as strong electrophiles with $\omega>$1.5 eV, moderate electrophiles with $0.8<$ $\omega$ $<$ 1.5 eV and marginal electrophiles with $\omega$ $<0.8$ eV [23].

Electronic density and electrostatic potential are important aspects in the reactivity, as shown on C$_{14}$ and DDE molecules as these are generated by DMol3 software, respectively; becoming in a useful application to predict reactivity of molecules on each point of the space. It allows to visualize the regions of the space towards which particular are attracted either positively or negatively (regions of either positive or negative electrostatic potential, respectively) [24].

**Methodology**

In the analysis of the ground state for electronic structures on model systems, a variational procedure is used; in which the functional of the electronic energy is minimized with respect to the electronic density by means of Density Functional Theory DFT [25] on Dmol3 developed by Delley [26], who used to achieve modeling and geometry optimizations, and the calculation of equilibrium distances and dissociation energy. For all electron calculations, a correlation functional proposed by Perdew-Wang in 1991 [27] using Generalized Gradient Approximations (GGA-PW91 level of theory) has been used for non-restricted spin, self-consistent field (SFC) with DND basis functions equivalent to 6-31G* basis [28].

The reactivity indexes for a system of N-electrons with total energy $E$ and external potential $V(r)$ are defined as the first and second derivative of $E$ regarding $N$ with the following expressions [29]:

$$\mu = (\frac{\partial E}{\partial N})_V(r) = -\chi$$  (1)

$$\eta = (\frac{\partial^2 E}{\partial N^2})_V(r) = (\frac{\partial \mu}{\partial N})_V(r)$$  (2)

The equations 1 and 2, for $\mu$ and $\eta$ respectively, are numerically determined by approximations based on finite differences of three points, as shown in the following expressions 3 and 4 [30], using the energies $E_{LUMO}$ and $E_{HOMO}$ for GAP (ELUMO - EHOMO) calculation [31]. The softness $S$ is the inverse of $\eta$ chemical hardness, and constitutes a useful concept on the prediction of chemical reactivity [32], as shown in the following expressions:

$$\mu = \frac{1}{2}(E_{LUMO} + E_{HOMO})$$  (3)

$$\eta \approx \frac{1}{2}(E_{LUMO} - E_{HOMO})$$  (4)

$$S = \frac{1}{2} \eta$$  (5)

The electrophilicity index [33] is calculated by the expression:
It is modeled a closed structure of 14 carbon atoms, to which a geometry optimization is carried out. This allowed to obtain a uniform ring type of 14 carbon atoms with alternating single- and triple-bonds, and bond length of 1.285 Å, as shown in (Figure 2), also with an approximate radius of 2.874 Å, a binding energy of -3830.310 kcal/mol, approximate perimeter of 18.027±0.023 Å, and an area of 25.949 Å².

\[ \omega = \frac{\mu^2}{2\eta} \]  

**Results**

Table 1: Reactivity parameters the C\(_{14}\) 

| Structure | EHOMO (eV) | ELUMO (eV) | Energía GAP (eV) | \(\mu\) (eV) | \(\eta\) (eV) | s | \(\omega\) |
|-----------|------------|------------|------------------|-------------|-------------|---|--------|
| C14       | -6.196     | -3.437     | 2.759            | -4.816      | 1.379       | 0.362 | 8.409 |

In (Figure 3) it is represented the molecular orbital borders HOMO and LUMO of structure showing blue lobes positive and yellow lobe negative. Energies \(E_{\text{HOMO}}\) and \(E_{\text{LUMO}}\) of each structure, as well as the energy difference between HOMO and LUMO (or energy GAP), are shown in (Table 1). A large GAP energy difference corresponding to stable and low reactive systems (> 5eV) is considered insulating, while with a small difference (< 0.5eV) hopefully the system is not stable and highly reactive, and it is considered as conductor. In this structure the value is 2.759eV, then it can be considered as a semiconductor [27] as in (Table 1) where the whole parameters are shown.

The electrostatic potential maps on (Figure 4) allow to visualize how electrons are distributed in carbyne molecules, the red color indicates regions where electronic density is higher, while blue regions indicate lower electronic population. The intermediate states follow the order of the rainbow from the center up to the external border, according to a greater or lower presence of electrons, and thus the green and yellow color close to the center border of the structure indicate a homogeneous electronic balance. Carbon atoms only possessing the same electronegativity should give an almost homogeneous electronic distribution, and there is no preferential place where electrons be circulating, but our map tells us that the external part of carbyne structure has major electronic density.
Likewise, the HOMO and LUMO energies of DDE are calculated as shown in (Figure 5) indicating the zones of greater reactivity. HOMO energy -4.445 eV and LUMO energy -4.363 eV (Table 2).

With the active zones of each analyzed structure, a reference is given about in which way DDE is accommodated, and the carbon structure can be put in a mode, that may be estimated in a sustained way that type of adsorption exists between the two molecules. (Figure 6a) shows the way in which the carbon structure with a DDE molecule is placed, and (Figure 6b) exhibits the final state of minimum energy, after achieving geometry optimization by means of SCF calculations.

Interactions are also made by placing the DDE inside the carbyne ring as shown in (Figure 7). Geometry optimization is obtained and what observed is that only the carbyne ring is deformed.

Conclusions

Using molecular modeling with DFT methodology in this work, it was observed adsorption capacity of $C_{14}$ activated carbon, which is to depend of the orientation to be placed with respect to DDE molecules. When the HOMO and LUMO of these molecules are overlapped in some region, adsorption is achieved. With the map of the electrostatic potential it could be observed that in the structure of the carbyne, the part having more electronic density is the external part, hence when placing the DDE molecule in the inside part of the ring there was no adsorption phenomenon only a deformed ring. The part with the most electronic DDE density is where the hydrogens are, then we put them outside the ring and DDE is added into the external part of the carbon structure.

For the prediction of reactivity, it is important to know the reactive sites of the molecular system to visualize specific parts where the adsorption phenomenon was given. DFT as a quantum theory based on electronic density, was useful in calculating the...
reactivity indexes to justify why and how the interactions of the molecules $C_{14}$ and $C_{14}HCl$. The chemical potential $\mu$ calculated in each of the molecules shows us the tendency of escaping of the electrons from a system in equilibrium, so that the electrons flow from the regions of high chemical potential towards regions of low potential $\mu(C_{14})=4.816 \text{ eV}$ and $\mu(C_{14}HCl)=3.554 \text{eV}$, which tells us that there is a low and high potential, thus there would be the possibility of a molecule yielding or accepting electrons respectively. With regard to the resistance to modify the electronic distribution of each of the molecules, called $\eta$, the following values were obtained $\eta(C_{14})=1.379 \text{ eV}$ and $\eta(C_{14}HCl)=1.809 \text{eV}$, which indicates that the DDE is more stable than the $C_{14}$, the electrophilicity $\omega$ tells us when the system is subjected to a electrostatic attack where the electrons involved come from the HOMO. The values of $\omega(C_{14})=8.4 \text{ eV}$ and $\omega(C_{14}HCl)=3.464 \text{eV}$ show that they are strong electrophilic, since they are in the range of $\omega>1.5 \text{ eV}$, stressing that $C_{14}$ have the highest value, and it is more likely to attract electrons. The electrophilicity $\omega$ is related with electronic affinity as they both measure the ability to accept electrons, but the electrophilicity $\omega$ measures the energy decrease due to the maximum electron flow between donor and acceptor. These intrinsic properties of the molecules indicate the availability that an adsorption may exist. The adsorption allows a new molecule to be formed, so that the DDE ($C_{14}H_{2}Cl$) grows to $C_{14}H_{4}Cl$ upon adsorption of the $C_{14}$ carbon ring. This growth helps the human beings to eliminate it, since otherwise it can go to be stored in the fatty tissue with possible consequences to some disease.

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