Investigación teórica a nivel semiempírico de Nanotubos de carbono como Nanotubos de ensayo

Theoretical Investigation of Carbon Nanotubes as Nano Test-tube

Dr. M. Al-anber
M.Ai-Anbar@hull.ac.uk

Department of Physics and Mathematics, Fenner Building, room 081, University of Hull, Cottingham Road, Hull, HU6 7RX, UK

Recibido: 7 de febrero de 2016 Aprobado: 5 de julio de 2016

Resumen

Se realizaron cálculos cuánticos (MINDO/3) de las propiedades estructurales de nanotubos de carbono (NTCs) y la adsorción de varios radicales de la glicina y el ácido butírico. Los radicales glicina N-centered y ácido bitírico C1-centered presentan la mayoría de los complejos estables con NTCs. El diámetro y longitud de los NTCs sobre las energías antienlazantes entre estas dos biomoléculas con los NTCs muestran un decrecimiento cuando el diámetro del NTC aumenta mientras que las energías enlazantes se incrementan con el aumento de la longitud del NTC. Los radicales de glicina N-centered se unen preferentemente a los extremos de los NTCs mientras que las moléculas del ácido graso prefieren hacerlo en el medio de la superficie interior del NTC.

Palabras claves: Nanotubos, métodos semiempíricos, interacción, biomoléculas

Abstract

The quantum calculations (MINDO/3) performed on the structural properties of the CNTs upon adsorption of several glycine and butyric acid radicals. The N-centered glycine and the butyric acid (C1-centered) radicals have most stable complexes with CNT. The diameter and length of the CNT on the anti-binding energies between these two biomolecules with the CNTs show a decrease as the CNT diameter increases while the binding energies increase with CNT length increase. The N-centered glycine radicals prefer to bond at the end of the CNTs while the fatty acid molecules prefer that at the middle of the inside surface of the CNT.

Keyword: nanotubes, semi-empirical methods, interaction, biomolecules.
Introduction

The nanotestubes are one the advantages of the nanotechnology that depend on the characteristics and behaviour of nanomaterials, being limited to nanoscale dimensions (1–100 nm). The quantum nature for the nanostructures is appeared due to their atomic and molecular sizes. The carbon nanotubes (CNTs) as sheets of graphite wrapped into a cylindrical form showed a new phenomena in the physics after Iijima discovered it [1-3]. Nanotubes have been used in the new biologic and medicine applications due to the ability for the immobilisation of proteins and enzymes [4,5]. As well, Wong et al. have shown that nanotubes are suited for use as probe tips in applications such as biomolecular probes to the carboxyl groups that are present at the open tip ends and to measure the binding force between single protein-ligand pairs [6]. Furthermore, the nanotubes have potential to penetrate into cells offer the potential for using them for delivery of drugs and antibiotic molecules without toxicity effects [7–10]. Furthermore, the nanomaterials are used in biosensing, antigen recognition and DNA hybridization due to their unique properties and the vaccine delivery biomedical applications [11], and the nanotubes present big technological advances in bioengineering too [12]. However, few theoretical studies of the interaction mechanism between the nanotubes and biomolecules, such as an attempt was made to explore the feasibility of nanoparticulate adsorbents as a drug delivery tool for the administration of erythropoietin to the small intestine [13] else the theory methods have been used to study the ether-bonded carbon nanotube[14], compared with an expectation of the broad applications of nanotubes, which tried to introduce the description of the nature of these interactions. However, Mavrandonakis et al. have studied the interaction of the amino acid with CNTs[15,16]. Chen et al. have shown that boron nitride nanotubes, which are non cytotoxic, can be functionalized for interaction with proteins and cells [17]. Furthermore, the influences of others factors, such as the diameters and lengths of the CNTs, on the interaction were considered too [18], also the investigating the CNTs as delivery for the anti-cancer drugs [19, 20], likewise Hesabi et al. studied the skin anti-cancer drugs with the CNTs [21]. Also, the fatty acid (butyric acid) interaction with the CNTs was studied [22]. What is more, others have studied constant molecules interaction with the magnesium oxide nanotubes [23] and also with the boron nitride nanotubes (BNN) [24]. Moreover, Zeighami et al. introduced the thermodynamic view of the communication between the Tacrine, which is the drug for Alzheimer's disease, with boron nitride nanotubes [25]. Both of these reports studied this interaction on the outer surface of the nanotubes and
paid little or no attention to study this interaction on the inner surface for these nanotubes.

The aim of the present paper is to introduce a model to the ability to consider the carbon nanotubes CNTs as a nanotest tubes, see figure 1, because there are not available dates about this issue yet. Where we try to examine the interaction of the glycine and the butyric acid radicals on the inner surface of the CNTs, which has an armchair type [26]. Then our purpose is to examine this interaction as a function of the CNT's diameters and lengths. Also, we attempt to investigate the effect of changing the positions of these biomolecules-CNT bonds on the interaction energy. Also, we will try to test the ability to link two glycine molecules on the wall of the CNTs surface, so that the first glycine interact on the inner surface of the CNT and the second one on the outer surface.

Figure 1. Some molecules may be able to be inserted into carbon nanotubes as the test tube model.

Computational details

In many cases, the results of the experimental methods are unable to describe accurately small systems of complex chemicals. Theoretical calculation methods can be used to bridge gaps in understanding experimental results. Furthermore, the quantum molecular methods can be used to show properties beyond the scope of current crystallographic methods. Where, the molecular quantum modelsallow us to study optical, magnetic, and
electronic properties that cannot easily measured experimentally. Also, the molecular quantum provides the interaction energies that are not provided by X-ray and NMR (nuclear magnetic resonance) experiments, so that the theoretical methods can be used to investigate further and to predict the physical and chemical nature of hydrogen bonding interactions. To determinate the structural and electronic properties of CNTs decorated with the glycine and butyric radicals, we used MINDO/3 (Modified Intermediate Neglect of Differential Overlap version 3). MINDO/3 is the earliest of the Dewar methods [27, 28]. MINDO/3 provides more accurate geometries and heats of formation than CNDO or INDO and has been used widely. The limitations of the INDO approximation, on which MINDO/3 is based, frequently lead to problems of accuracy when dealing with molecules containing heteroatoms. MINDO/3 is particularly suitable for describing carbocations, including non-classical carbocations, and polynitro organic compounds. In most molecular computations, to perform accurate calculations for a nano-sized system, how can do that without ending in prohibitively large computations such as the DFT methods. The dangling bonds at of the tubes ends were saturated by hydrogen atoms. The resolution of MINDO/3, as implemented in the HyperChem Release 7.52 for Windows Molecular Modeling System programme package (http://www.hyper.com/) was employed for the geometry estimations.

Results and discussion

At beginning it was important to determine the most stable isomers of the glycine and the butyric acid radicals on the inside wall of CNTs. Among these possible isomers are the ones from which one hydrogen atom is abstracted from either the N atom or the C atom (see Fig. 2). So here we adopt three isomers of butyric acid and two isomers from the glycine. Figure 3 shows the linking of these isomers on the inside wall of the CNTs (for a constant length equal to 10.0 ±0.025 Å). Note that in each case; we link the radical–CNT bond in the middle of the inside wall of the CNTs.
Figure 2. The geometry optimized of the isomers of the butyric acid and the glycine radicals.

Figure 3. The linking of the butyric acid and the glycine radicals (isomers) on the inside wall of the CNTs.

Figure 4 shows the binding energies (BE) results for these isomers on the inside wall of the CNTs as a function of the CNTs diameters. The binding energy (BE) of the radicals with the CNTs is according to the equation \( BE = E_{\text{Radical}+\text{CNT}} - (E_{\text{Radical}} + E_{\text{CNT}}) \), where \( E_{\text{Radical}+\text{CNT}} \) is the energy of the complex of the radical with the CNT. However, there is no binding energy (anti-binding) between these biomolecules on the inside wall of CNTs. As the CNTs diameter increases the anti-binding energies decrease, where this behaviour is the same for all these issues, approximately, but there are shifting among these matters. It was found that the C\(^1\)-centered butyric acid and N-centered glycine...
radicals have lower anti-binding energies compare with the other isomers approximately, and this behaviour becomes semi-constant for all issues after the CNTs diameter equal to 10.22 Å.

Figure 4. The binding energies (BE) for the isomers of butyric acid and the isomers of glycine radicals on the inside wall of the CNTs as a function of the CNTs diameters.

The effect of the CNTs diameters reflects the possibility to make control on the ability of these molecules to pass inside the CNTs. We find that reaction with the single tube wall of the CNT, the butyric acid (C1-centered) and glycine (N-centered) radicals show lower anti-binding with CNTs, thus we further study only these two molecules on the inside wall of the CNTs. The second important factor is the interaction of the butyric acid (C1-centered) and the glycine (N-centered) radicals on the inside walls but with different lengths of the CNTs evaluated (for constant diameter equal to 10.22 Å). Also, here we put the radical–CNT bond in the middle of the inside wall of the CNTs. The CNTs lengths during their synthesis is a dynamic property.

The binding energy of the both radicals (N-centered glycine and C1-centered butyric acid) on the inside wall of the CNTs are fluctuated with the lengths, as shown in Fig. 5, so these fluctuating is between the binding and anti-binding as the length of the CNTs increase. However, binding energy between the CNTs with these two biomolecules due to the CNTs lengths increase. There is same behavior for these two biomolecules with CNTs lengths. On another side, the glycine shows more probability to interaction with
the CNTs than the butyric acid radical. Thus, we may conclude that the binding of these two molecules on the inside wall of the CNTs depends on the lengths of CNTs more than their diameters.

Figure 5. The binding energy of the butyric acid (C1-centered) and glycine (N-centered) radicals on the CNT as a function of CNTs length. The binding of the N-centered glycine radicals on the inside wall of the CNTs increases as a function of the N\textsubscript{glycine}–CNT bond position on the internal surface of CNT as this post changes from the middle towards one of the CNT two ends from 1, 2, … 7, as shown in Fig. 6.

Figure 6. The positions of the N-glycine- CNT bond and the C1-centered butyric acid- CNT bond on the inside wall of the CNTs that is expecting.
For this purpose the CNTs that adopt with diameter 11.74 Å and length 17.62 Å. We did same for the C1-centered butyric acid radical. The complexes formed by the glycine radical on the inside of the single tube wall are more stable when the reaction occurs at the ends of the CNT, see Fig. 6. While the binding of the C1-centered butyric acid radicals on inside wall of the CNTs decreases and then increases as a function of the C_{butyric acid}–CNT bond position on the internal surface of CNT, as shown in Fig. 7.

![Graph showing binding energy of radicals on CNTs](image)

**Figure 7.** The binding energy of the butyric acid (C1-centered) and glycine (N-centered) radicals on the inside wall of the CNTs as a function of the position of the radical-CNT bond, from the middle of the CNTs forwards their two ends.

So we expect there is the probability that the radical glycine interaction with the ends of CNTs and may be will not enter inside the CNTs. The complexes that formed by the C1-centered butyric acid radicals on the inside wall of the CNTs are more stable when the reaction occurs at the middle of the CNT. The butyric acid form complex stable than the Glycine molecule inside the CNTs surface. Due to this point, we expect that the butyric acid radicals will collect in the middle of the cavity of the CNTs and may be will not pass inside the CNT quickly. May be the CNTs can be modified to be like a filter to separate the amino acid and the fatty acid molecules. We tried to investigation the ability of interaction two glycine molecules on the wall of the CNTs surface, so that the first molecule interaction on the inside surface of the CNTs and the second one on the outer surface, see Fig. 8.
Figure 8. The expecting positions to link two N-glycine molecules with the CNTs.

For this purpose the CNTs that adopt here with diameter 11.34 Å and length 12.53 Å. There are many expecting to make linking on this issue, so we start with the same atom on the CNTs as in issue 1. The results in Fig. 9 show that the outer glycine has binding on the CNT surface while the inner one without binding.

Figure 9. The binding energy for the outer glycine radical (and the inner glycine radical) on the CNT surface.
Also, when the two molecules become far from each other, the outer one becomes more binding, while the anti-binding for inner one decreases. The inner glycine radical will reduce the binding of the outer glycine that on the CNTs surface, so there is fluctuated in this behaviour may be due to the distances between them. This action becomes constant after position 8, may be attributable to the change the distance between the nitrogen and oxygen atoms of these two glycine molecules. The glycine molecule on the outer surface of the CNTs will decrease the probability to anti-binding the glycine radical on the inside wall of the CNTs, and maybe this issue will not allow the glycine to flow inside the CNTs but without probability to binding on its internal wall.

Conclusions

In summary, we have performed MINDO/3 calculations on the structural properties of the CNTs upon adsorption of several glycines and butyric acid radicals. Among these isomers, the N-centered glycine and the butyric acid (C1-centered) radicals form stable complexes with CNT compare with the other issues. Our results about the interaction between these two biomolecules with the CNTs are summed up by the following:

1- The results of the diameter and length of the CNT on the anti-binding energies between these two biomolecules with the CNTs show a decrease as the CNT diameter increases while the binding energies increase with CNT length increases.

2- The N-centered glycine radicals prefer to bond at the end of the CNTs while the fatty acid molecules prefer that at the middle of the inside surface of the CNT.

3- The interaction two glycine molecules on the wall of the CNTs surface, one molecule on the inside surface of the CNTs and the second one on the outer surface has an influence on the binding of the both molecules.

4- Maybe the CNTs can be modified to be as a filter to separate the amino acid and the fatty acid molecules.

References


