

Theoretical Study Toward Understanding the Electronic Properties of Armchair Hetero Nanotubes

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Abstract

Using the density functional theory ‘DFT’ code ‘SIESTA’ combined with a Green’s function scattering approach ‘GFSA’ code ‘GOLLUM’, we investigate the electronic properties of hetero nanotubes, density of states ‘DOS’, electronic band structure BS, transmission coefficient $T(E)$ and current-voltage curve. We simulated four different hetero structures of nanotubes which are: 1- Hetero nanotube with one graphene ring and one boron nitride ring. 2- Hetero nanotube with two graphene ring and one boron nitride rings. 3- Hetero nanotube with three graphene ring and one boron nitride rings. 4- Hetero nanotube with four graphene ring and one boron nitride rings nanotubes. The results show that there is a clear reduction in energy gap ‘ E_g ’ of hetero nanotubes comparing with the insulator boron nitride ($E_g \approx 4.6$ eV) and also this kind of hetero nanotubes have zero states at Fermi energy comparing with conductor carbon nanotube which has finite states at Fermi energy. This is could be a good method to design a semiconductor with desirable band gap.

Keywords: carbon nanotubes, boron nitride nanotubes, hetero nanotubes, energy gap.

دراسة نظرية نحو فهم الخصائص الإلكترونية للأنابيب النانوية المغايرة التركيب

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الخلاصة

باستخدام نظرية الكثافة "DFT" جنباً إلى جنب مع تقريب دالة كرين لمنطقة التشتت 'GFSA'، تم دراسة الخصائص الإلكترونية للأنابيب النانوية المغايرة التركيب (المركبة من الكربون والبيرون نترأيد)، كثافة الحالات "DOS"، تركيب الحزم الإلكترونية EBS، معامل الانتقال $T(E)$ ومنحنى التيار-الجهد. تم محاكاة أربعة تراكيب مختلفة للأنابيب النانوية المغايرة التركيب وهي: 1- أنبوب مغاير التركيب يتكون من حلقة واحدة من الكرافين حلقة واحدة من البورون نترأيد. 2- أنبوب مغاير التركيب يتكون من حلقتين من الكرافين حلقة واحدة من البورون نترأيد. 3- أنبوب مغاير التركيب يتكون من ثلاث حلقات من الكرافين حلقة واحدة من البورون نترأيد. 4- أنبوب مغاير التركيب يتكون من أربع حلقات من الكرافين حلقة واحدة من البورون نترأيد. النتائج أظهرت أن هناك انخفاضاً واضحاً في فجوة الطاقة E_g للأنابيب المغايرة التركيب بالمقارنة مع الأنابيب المثالية للبورون نترأيد الذي يعتبر عازلاً مثالياً ($E_g \approx 4.6$ eV) وكذلك هذا النوع من الأنابيب المغايرة التركيب لا تحتوي على حالات متوفرة عند وحول طاقة فيرمي عند مقارنتها مع الأنابيب الكربونية النانوية والذي يحتوي على حالات متوفرة محدودة عند طاقة فيرمي.

الكلمات المفتاحية: أنابيب الكربون النانوية، أنابيب البورون نترأيد النانوية، الأنابيب المغايرة التركيب النانوية وفجوة الطاقة.

1. Introduction

The discovery of carbon nanotubes (CNT) in 1991 by S. Iijima [1-3] has reserved enormous intentions and intensive research interest because of their unique electronic properties and potential application in nanodevices [4-6]. Carbon nanotubes are cylindrical or we can describe it as extended-fullerene structures. They exhibit outstanding physical properties that may be

exploited in areas from advanced composites to nanoelectronics [7]. Single walled carbon nanotubes (SWCNTs) exhibit either metallic or semiconducting behaviour depending only on diameter and helicity. This ability to display fundamentally distinct electronic properties of SWCNT was experimentally confirmed by scanning tunnelling microscopy (STM) [4, 5, 8].

Basically the SWCNTs can be considered as a strip cut from a graphene sheet and then rolled up into a tube. The diameter and helicity of a SWCNT are defined by the roll-up vector $\vec{Ch} = n\vec{a}_1 + m\vec{a}_2 \equiv (n, m)$, which connects crystallographically equivalent sites on this sheet as shown in figure 1. The \vec{a}_1 and \vec{a}_2 are the graphene lattice vectors, n and m are integers. Electronic band structure calculations could predict the (n, m) parameters, which determine whether a SWCNT will be a conductor or a semiconductor [4, 7, 9-11].

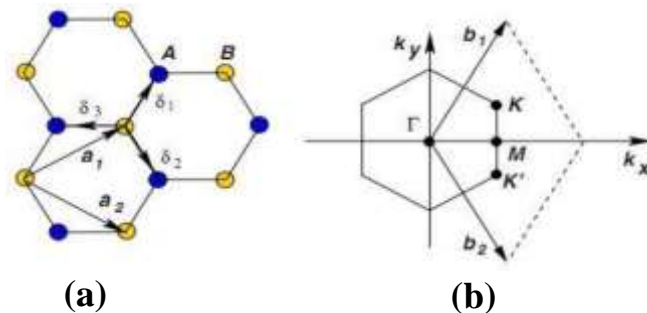


Figure 1: The hexagonal lattice and its Brillouin zone. (a) graphene lattice structure, made out of two interpenetrating triangular lattices which are \vec{a}_1 and \vec{a}_2 are the lattice unit vectors, and $\delta_i, i= 1, 2, 3$ are the nearest-neighbour vectors. (b) represent the corresponding Brillouin zone. The Dirac cones are located at the K and K' points [9].

There are three types of SWCNTs; first armchair carbon nanotubes which are always behaves as conductor, that is depend on n and m parameters ($n=m$), second one is zigzag carbon nanotubes which are exhibit as conductor and semiconductor depends on n parameter ($m=0$) and thirdly is the chiral carbon nanotubes, they are also behave as a conductor or semiconductor depend on m and n parameters ($m \neq n$) [12, 13]. This means

the behaviour of zigzag and chiral carbon nanotubes depends on the diameter of the nanotube [14].

It is well-known in the periodic table, that the III-V material most closely related to C is boron nitride (BN), which is like carbon, both of them have sp^2 - and sp^3 -bonded structures [15]. Boron nitride nanotube has a wide band Energy gap around Fermi energy which is approximately from 5.5 eV to 5.8 eV [15-17], this means all types of bulk boron nitride nanotubes are perfect insulator [15, 16, 18]. Many calculation reported that boron nitride nanotube energy gap is almost independent of tube diameter, chirality, and the number of tube walls [16, 17].

Hetero nanotubes received a huge interest from many scientists and research groups to understand their electronic properties and their possible applications [19-22], these studies reported that boron nitride has a remarkable effect on the energy gap of ideal carbon a nanotube. The importance of the interface between graphene and BN has been investigated [19, 23, 24], for these impurities, a characteristic peak close to the Fermi energy is found, with the peak associated with boron below the Fermi energy and the peak associated with nitrogen above the Fermi energy [19].

2. Theoretical Tools

The calculations are performed by using the SIESTA implementation of DFT [25] to find the optimized geometry. We followed the procedure explained in reference 27 and the *ab-initio* calculation was employed using the local density approximation (LDA) of the exchange and correlation functional with Ceperley-Alder exchange 'CA' parametrization, double zeta polarized (DZP) basis sets of pseudoatomic orbitals, a real-space grid defined with a plane-wave cutoff energy of 300 Ry and a maximum force tolerance of 10 meV/Å. The density of states (DOS) and electronic band structures (EBS) were calculated with $1 \times 1 \times 200$ *k-points*. The Mean-Field Hamiltonian (MFH) obtained from SIESTA was used as an input data to calculate transmission coefficient $T(E)$. to obtain the transmission coefficient $T(E)$, we used the non-equilibrium Green's function code GOLLUM [26] ,

which utilises the DFT-based hamiltonian from SIESTA. For ideal periodic structure, the number of open channels, is equivalent to the transmission coefficients $T(E)$.

3. Electronic Properties of Ideal Nanotubes

In this section, we calculate the electronic properties of ideal carbon and boron nitride nanotubes to make sure that the used theoretical tools (SIESTA and GOLLUM codes) give the correct results. Figure 2 shows the supercell of the ideal carbon and boron nitride nanotubes.

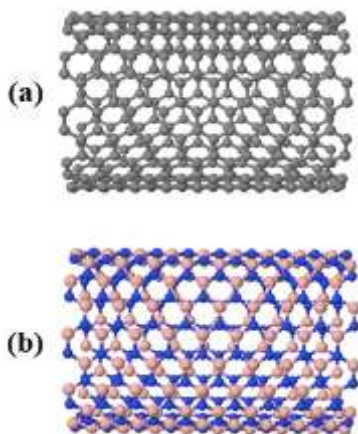


Figure 2: Shows the ideal nanotubes (a) (10,10) carbon nanotube (360 carbon atoms), (b) (10,10) boron nitride nanotube (360 boron and nitrogen atoms).

Figure 2 shows the ideal conductor (10,10) carbon nanotube and ideal insulator (10,10) boron nitride nanotube that we obtained from SIESTA. The obtained results (figure 3) show good agreement with the publications which are reported that all sizes of armchair carbon nanotube are exhibiting a finite states at Fermi energy (conductors) and zero energy gap with two open channels at and around Fermi energy [12, 14]. Also, the boron nitride results confirmed that armchair boron nitride nanotube behaved as a perfect insulator with energy gap of approximately 4.5 eV, our result has a good

agreement with the publications [15-18, 20]. Figure 3 show the results of ideal armchair carbon nanotube and boron nitride.

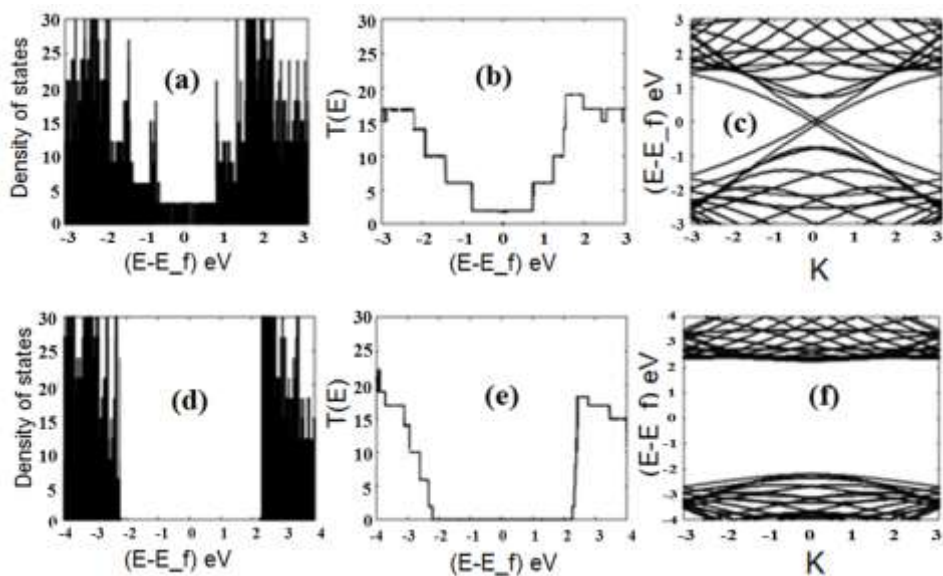


Figure 3: The electronic properties of the perfect carbon and boron nitride nanotubes. Figure2 (a-c) show the density of states (DOS), transmission coefficient $T(E)$ and electronic band structure (EBS)of the ideal (10,10) carbon nanotube respectively. Figure2 (d-f) show the density of states (DOS), transmission coefficient $T(E)$ and electronic band structure (EBS) of the ideal (10,10) boron nitride respectively. All calculations carried out with $1 \times 1 \times 200$ k -points.

4. Electronic Properties of Hetero Nanotubes

Here, we simulated the (10,10) hetero nanotubes with varying the hexagonal carbon rings (from one ring to four rings of carbon in between boron nitride rings) as shown in figure 4. By calculating the mean-field Hamiltonian (MFH) from SIESTA, we calculated the density of states (DOS) and the electronic band structures (EBS). Also, the obtained mean-field Hamiltonian (MFH) used to feed the GOLLUM code to investigate the transmission coefficient $T(E)$. The calculations carried out with $1 \times 1 \times 200$ *k*-points to make sure that the structures shown in figures (3a-3d) are periodic in Z direction and also to obtain the Van Hove singularity.

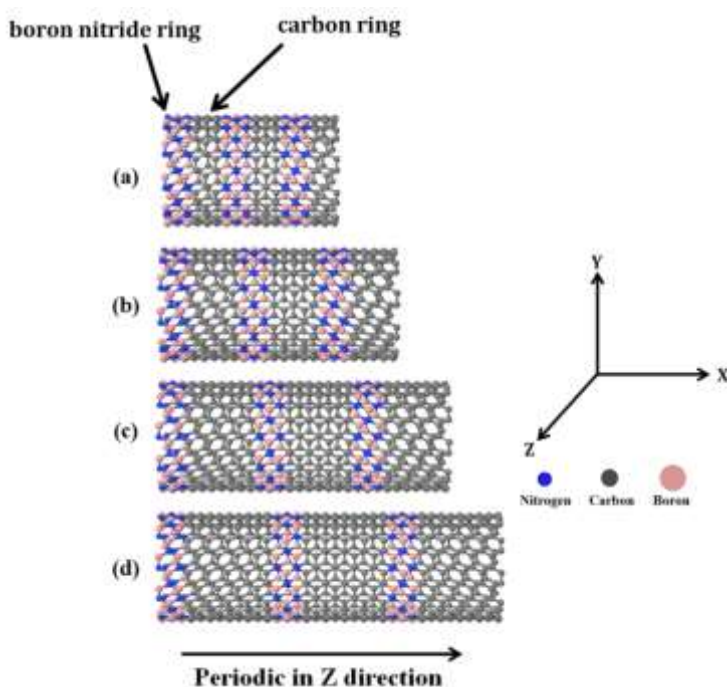


Figure 4: The hetero nanotubes, **(a)** with one boron nitride and one hexagonal carbon ring, the structure contains 360 carbon, boron and nitrogen atoms, **(b)** with one boron nitride and two hexagonal carbon rings (480 carbon, boron and nitrogen atoms), **(c)** with one boron nitride and three hexagonal carbon rings (600 carbon, boron and nitrogen atoms), and **(d)** with one boron nitride and four hexagonal carbon rings, the structure consist of 720 carbon, boron and nitrogen atoms. All structures are periodic in Z direction.

The structures shown in figure 4 obtained using the methodology explained in [27]. This methodology leads to create unique forms of pure sp^2 -bonded carbon and unprecedented heteromolecules, which are formed by sculpting selected shapes from bilayer graphene, heterobilayers, or multilayered materials and allowing the shapes to spontaneously reconstruct [27]. The resulting electronic properties of the structures (figure 4) are shown in figure 5.

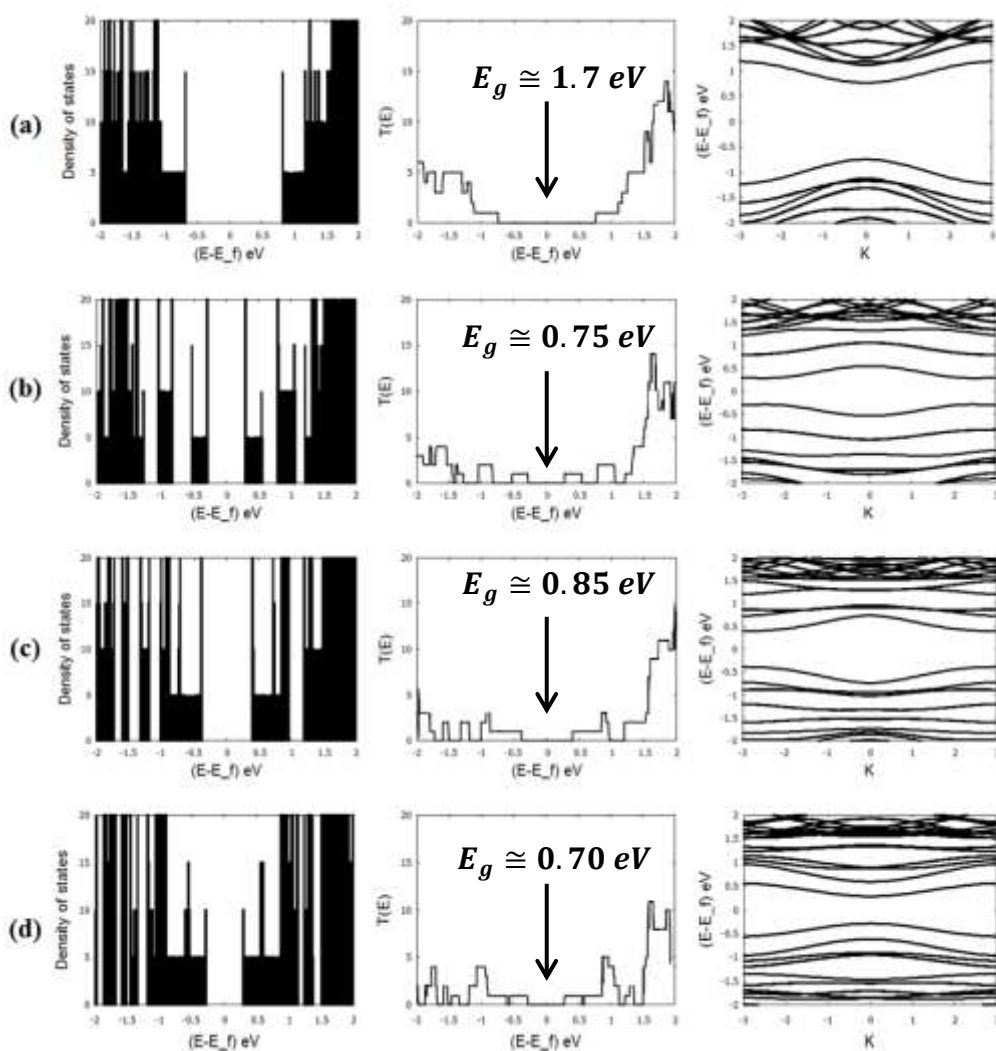


Figure 5: (a) the electronic properties of the structure shown in figure 3a, the density of states, transmission coefficient and band structure. (b) the density of states, transmission coefficient and band structure of the structure shown in figure 3b. (c) the density of states, transmission coefficient and band structure of the structure shown in figure 3c. (d) the density of states, transmission coefficient and band structure of the structure shown in figure 3d. In all figure a, b, c and d, the left column represents the DOS, the middle column represent the T(E) and the right column represents the EBS.

From figure 5 (a-d, the far left figures), it is easy to see that the energy gap changed dramatically. For example, the energy gap in figure 4a is about 1.7 eV which is belong to the structure shown in figure 4a, this means there is a huge increasing in the energy gap comparing with the ideal (10,10) carbon nanotube (conductor) and also there is an enormous reduction in the energy gap comparing with the ideal (10,10) boron nitride nanotube . By having look at the figure 5 (a-d, middle and far right figures), we can see a similar outcome. Overall, the energy gap around Fermi energy for all hetero structures shown in figure 4 are varying from approximately 0.7 eV to 1.7 eV.

5. Current-Voltage features

Current-voltage characteristics are enormously measured experimentally, therefore calculating the current-voltage curve is useful for characterizing the hetero nanotubes shown in figure 4. The current (I), with respect to voltage (V), can be calculated from the transmission probability [19]:

$$I(V) = \frac{2e}{h} \int_{E_f - \frac{eV}{2}}^{E_f + \frac{eV}{2}} T(E, V) dE \quad (1)$$

From equation 1, we can see that the transmission depends on the bias voltage. The maximum current I(V) carried by the hetero nanotubes shown in figure 4 at a finite voltage V, is given by:

$$I(V) = \frac{2e}{h} \int_{E_f - \frac{eV}{2}}^{E_f + \frac{eV}{2}} T(E) dE \quad (2)$$

The resulting I-V plots for the optimized hetero nanotubes are shown in figure 5.

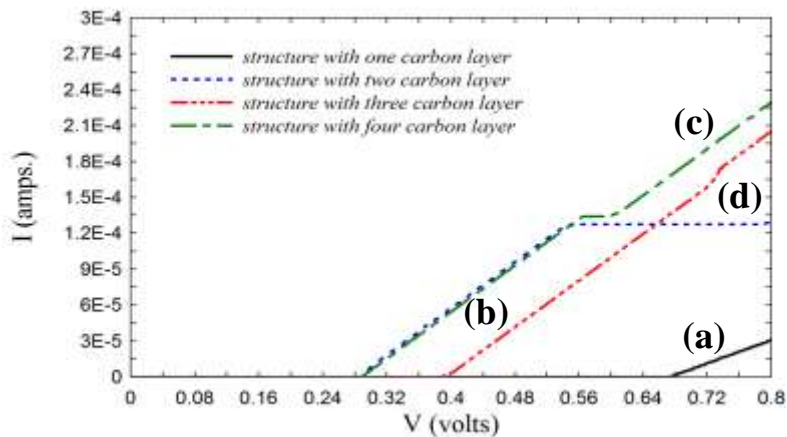


Figure 6: The obtained I(V) curve for (a) the hetero nanotube shown in figure 4a, (b) the hetero nanotube shown in figure 4b, (c) the hetero nanotube shown in figure 4c, and (d) the hetero nanotube presented in figure 4d.

Figure 6 shows clearly that at specific voltage value, for example at 0.8 volt the hetero structure with one hexagonal ring has the lowest current and the hetero structure with four hexagonal rings has the highest current. This result is due to the available states at and around the Fermi energy which means increasing the carbon rings layer will lead to enhance the conductance of the hetero nanotubes.

6. Conclusions

In this work, the energy gap was manipulated of the hetero nanotubes from about 0.7 eV to 1.7 eV by varying the number of graphene rings from one ring to four rings in between the boron nitride rings. These values of the energy gap values are fall in the range of the semiconductors energy gap, for example the energy gap of silicon at room temperature is about 1.12 eV [28]. Boron nitride has non-trivial effect on the energy gap of carbon nanotube. In other word, doping the (10,10) carbon nanotube by boron nitride led to open the energy gap hugely. Also, increasing the graphene rings led to reduce and control the size of the energy gap of the hetero nanotubes. This is could be a good methodology to obtain a semiconductor with desirable band gap.

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