Fe@B₆H₆ Aggregates: From Simple Building Blocks to Graphene Analogue

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Research Article

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Abstract

We suggest the possibility to build graphene analogue with the planar hexacoordinate wheel-type Fe@B$_6$H$_6$ cluster as the building block through studying theoretically the geometry, stability and electron structure of its dimer and trimer as well as the dimerization of the two trimers. Employing the dehydrogenation route to polymerization, we can obtain the hexagonal boron sheet that are partly and uniformly filled by Fe atoms in the center of the holes, achieving uniform chemical doping and a very large hexagonal-hole-density. Thus, we may offer a novel cluster-assembled material for experimental chemists to construct graphene analogue.

1. Introduction

Graphene, the first perfect monatomic two-dimensional carbon crystal, was isolated successfully from graphite in 2004$^1$. Its excellent properties, such as extremely high carrier mobility, high thermal conductivity and high specific surface area, have sparked not only the intensive studies on its synthesis and functionalized applications$^{2,3}$ but also the extensive discoveries towards graphene analogous$^{4,5}$, consisting of compositions other than carbon. Boron clusters have been found theoretically$^6$–$^9$ and experimentally$^{10}$–$^{12}$ to possess a two-dimensional (quasi)planar structure termed boron sheet. However, boron cannot form the stable honeycomb hexagonal-hole framework as graphene because of its electron deficient character. Instead, part of the hexagonal holes need to be filled by boron atoms adopting buckled form so that the boron sheet can maintain its (quasi)planar structure$^{11,12}$. Therefore, the boron atoms filling the holes serve as an important role in boron sheet and obviously distinguish from the ones forming the hole framework. Under the context, can we choose other atom to partially fill the holes and achieve the better results than that from boron atom? If possible, the more crucial question is how we can fill the holes in boron sheet. The challenging and appealing questions arouse our research interest.

The (quasi)planar hexacoordinate wheel-type clusters, such as CB$_6^{2−}$13 and X@B$_6$H$_6$ (X = V, Cr and Mn)$^{14,15}$, give us some inspiration to answer the questions. The existence of these clusters indicates that it is possible to fill the hole with atoms other than boron. And then, we can use these clusters as building blocks for constructing graphene analogue through polymerization. As we have shown in our recent work$^{16}$, the planar wheel-type $D_{6h}$ Fe@B$_6$H$_6$ with good chemical stability is the global minimum isomer and is therefore more attractive candidate for cluster-assembled materials. Here, we studied theoretically the geometry, stability and electron structure of the dimer and trimer of Fe@B$_6$H$_6$ as well as the graphene analogue FeB$_6$. Our calculated results indicate that the structures and charges of the dimer and trimer of Fe@B$_6$H$_6$ are similar to those of the monomer Fe@B$_6$H$_6$. Based on π-electron molecular orbital, Huckel rule and nucleus-independent chemical shifts, the trimer of Fe@B$_6$H$_6$ can be considered as the triphenylene analogue. Moreover, the phonon dispersion indicates that the graphene analogue FeB$_6$ has good dynamical stability. These results indicate that the Fe@B$_6$H$_6$ can be used as the building block to build the graphene analogue FeB$_6$. 

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2. Calculation Method

The structures of the monomer, dimer and trimer of Fe@B₆H₆ were optimized by employing B3LYP¹⁷,¹⁸ method, as implemented in Gaussian 09¹⁹, with 6–31 + G(d) basis sets for all atoms except the Fe, which were described by the Stuttgart–Dresden (SDD) effective core potential²⁰. Using the optimized geometries, the HOMO–LUMO energy gap ($\Delta_{H-L}$), vertical ionization potential (VIP)¹⁶, vertical electron affinity (VEA)¹⁶, natural population analysis (NPA)²¹ and nucleus-independent chemical shifts (NICS)²² were carried out at B3LYP/6-311 + G(d,p) level. The VIP of monomer, dimer and trimer is defined by the energy difference between the cationic $E$(monomer/dimer/trimer)$^+$ and neutral $E$(monomer/dimer/trimer) calculated at the equilibrium neutral geometry, the VEA of monomer, dimer and trimer is the energy difference between the neutral $E$(monomer/dimer/trimer) and the anionic $E$(monomer/dimer/trimer)$^−$ calculated at the equilibrium neutral geometry, as given in the following equations:

\[
VIP = E(\text{monomer/dimer/trimer})^+ - E(\text{monomer/dimer/trimer})
\]

\[
VEA = E(\text{monomer/dimer/trimer}) - E(\text{monomer/dimer/trimer})^-
\]

For graphene analogue FeB₆, its optimized structure is carried out by the DMol³ module implemented in Material Studio 2018 using Perdew-Burke-Ernzerhof (PBE)²³ generalized gradient approximation (GGA) and double-numerical properties plus polarization (DNP). In the convergence tolerance, the energy, force, and displacement were set as $10^{-5}$ Ha, 0.002Ha/Å, and 0.005Å, separately²⁴. A vacuum layer of 20Å is added to avoid the influence of periodic adjacent layers. And Monkhorst-Pack k-mesh of 6×6×1 is adopted in Brillouin zone.

3. Results And Discussions

3.1 Optimized structures and some parameters of the monomer, dimer and trimer of Fe@B₆H₆

Figure 1 shows the optimized geometries of Fe@B₆H₆ and the dimer (Fe@B₆)₂H₁₀ and the trimer (Fe@B₆)₃H₁₂. The bond lengths and the charges of monomer, dimer and trimer are listed in Table 1 and Table 2, respectively. The monomer Fe@B₆H₆ forms the perfect regular hexagon with the planar hexacoordinate Fe atom, possessing the highest $D_{6h}$ symmetry. And the bond lengths and charges agree well with those obtained at BP86/6-311 + G(3df,3pd) level¹⁶. For the dimer (Fe@B₆)₂H₁₀, it has the highest $D_{2d}$ symmetry and the dihedral angle between the two monomer is 90° and the distance between them is 1.660 Å which is in good agreement with the experimental value of B-B bond length (1.691 Å)²⁵, suggesting the interaction between the monomers are very strong. While comparing with the monomer Fe@B₆H₆, the bond lengths and charges the dimer (Fe@B₆)₂H₁₀ do not change significantly in
each monomer except the B atom that links the other monomer. For the dimer \((\text{Fe@B}_6)_2\text{H}_{10}\), all atoms in each monomer are coplanar, which indicates that the character of monomer \(\text{Fe@B}_6\text{H}_6\) is well maintained during the dimerization. The dimer \((\text{Fe@B}_6)_2\text{H}_{10}\) with \(D_{2h}\) symmetry is a transition state for the conversion to the \(D_{2d}\) conformation. For the trimer \((\text{Fe@B}_6)_3\text{H}_{12}\), it has the highest \(C_{2v}\) symmetry and the three monomers are perfectly coplanar. The hole in the trimer is not regular hexagon and it is composed of two different types of B–B distances, which are \(R_{\text{B1-B5}} = 1.671 \, \text{Å}\) and \(R_{\text{B5-B6}} = 1.846 \, \text{Å}\), respectively.

The \(\Delta H_{\text{L-VIP}}\) and \(\Delta H_{\text{L-VEA}}\) of the monomer, dimer and trimer of \(\text{Fe@B}_6\text{H}_6\) are also listed in Table 2. It can be seen that the \(\Delta H_{\text{L-VIP}}\) values of monomer, dimer and trimer are 3.58, 3.22 and 2.80 eV, respectively. Although the \(\Delta H_{\text{L-VIP}}\) value of trimer is the smallest among them, it is larger than the \(\Delta H_{\text{L-VIP}}\) value (2.63 eV) of triphenylene\(^{26}\), indicating the monomer, dimer and trimer of \(\text{Fe@B}_6\text{H}_6\) are chemically stable. The \(\Delta H_{\text{L-VEA}}\) values of monomer, dimer and trimer are 8.06 eV, 7.68 eV and 7.57 eV, respectively, increasing gradually while the \(\Delta H_{\text{L-VEA}}\) values of monomer, dimer and trimer are 1.69 eV, 2.15 eV and 2.53 eV, respectively, decreasing gradually.

3.2 The molecular orbital and aromaticity of the trimer \((\text{Fe@B}_6)_3\text{H}_{12}\)

Since \(\text{Fe@B}_6\text{H}_6\) exhibits the similar \(\pi\) molecule orbitals to benzene\(^{16}\), the trimer \((\text{Fe@B}_6)_3\text{H}_{12}\) may be the triphenylene analogue. In order to confirm our conjecture, the \(\pi\)-electron molecular orbitals (MOs) of \((\text{Fe@B}_6)_3\text{H}_{12}\) and triphenylene are plotted in Fig. 2. It can be seen that the shape of these MOs of \((\text{Fe@B}_6)_3\text{H}_{12}\) and triphenylene are similar. For example, the HOMO-5 of \((\text{Fe@B}_6)_3\text{H}_{12}\) is bond MO which is similar to the bond MO of HOMO-9 in triphenylene. In addition, both \((\text{Fe@B}_6)_3\text{H}_{12}\) and triphenylene have three degenerate MOs (HOMO, HOMO-6 and HOMO-8 in \((\text{Fe@B}_6)_3\text{H}_{12}\) and HOMO, HOMO-2 and HOMO-6 in triphenylene). As a result, their nine \(\pi\)-electron MOs accommodate 18 \(\pi\) electrons that satisfy the \((4n + 2)\) Hückel rule. Thus, the trimer \((\text{Fe@B}_6)_3\text{H}_{12}\) exhibits the aromaticity and can be considered to be the triphenylene analogue.

\(\text{NICS}\) is a simple and efficient criterion to characterize aromatic nature. To better understand the aromaticity, the calculated \(\text{NICS}(d)\) (\(d = 0\) and \(1\) for inside and above the hole, respectively.) of \((\text{Fe@B}_6)_3\text{H}_{12}\) and triphenylene are also shown in Fig. 2. The \(\text{NICS}(0) = -0.53 \, \text{ppm}\) and \(\text{NICS}(1) = -0.28 \, \text{ppm}\) of the hole in the trimer \((\text{Fe@B}_6)_3\text{H}_{12}\) are less negative than the \(\text{NICS}(0) = -1.72 \, \text{ppm}\) and \(\text{NICS}(1) = -5.09 \, \text{ppm}\) of the hole in triphenylene, which indicates that the hole of trimer \((\text{Fe@B}_6)_3\text{H}_{12}\) is less aromatic than that of triphenylene. While the monomer in the trimer \((\text{Fe@B}_6)_3\text{H}_{12}\) has very strong aromatic character since its \(\text{NICS}(1) = -15.2 \, \text{ppm}\) is more negative than \(-9.8 \, \text{ppm}\) for the monomer in triphenylene, which can compensate the aromaticity of the hole in trimer \((\text{Fe@B}_6)_3\text{H}_{12}\).

3.3 The structure and stability of graphene analogue \(\text{FeB}_6\)
Before building the graphene analogue FeB$_6$, we examined the bigger stable aggregates. Two kinds of different dimerization of the trimer are shown in Fig. 3. The six monomers reveal perfect coplanarity in each of them, indicating the trimer possesses very good ability of plane expansion. Thus, assembling the stable trimers (Fe@B$_6$)$_3$H$_{12}$ can provide the possibility of building graphene analogue FeB$_6$ as the triphenylene in graphene$^{27,28}$.

And then, we optimized the graphene analogue FeB$_6$, as shown in Fig. 4. The FeB$_6$ with $P6/mmm$ symmetry is completely planar structure. The boron-ring with Fe atom in the FeB$_6$ has the B-B bond length of 1.860 Å and B-Fe bond length of 1.860 Å, which are slight longer than those (1.824 Å) of Fe@B$_6$H$_6$ monomer. And the bond lengths of two different B-B in the boron-ring without Fe atom are 1.860 Å and 1.661 Å, which are similar to those of boron-ring in Fe@B$_6$H$_6$ trimer. Therefore, the graphene analogue FeB$_6$ preserves the structural features of monomer and trimer of Fe@B$_6$H$_6$.

We also studied the hexagon hole density of the FeB$_6$. The hexagon hole density ($\eta$) is defined as$^{26,29a}$:

$$\eta = \frac{\text{No. of hexagon holes}}{\text{No. of atoms in the original triangular sheet}}$$

According to the formula, the triangular boron sheet has $\eta = 0$, the hexagonal boron sheet $\eta = 1/3$$^{29}$. For the FeB$_6$, it represents a hexagonal hole density of $\eta = 2/7$, which is bigger than those in pure boron $\alpha$ and $\beta$$^{29}$ and very close to the hexagonal boron sheet $\eta = 1/3$.

Besides, we also investigated the dynamical stability of the FeB$_6$. The phonon dispersion is shown in Fig. 5. The unit cell of FeB$_6$ monolayer has seven atoms, suggesting that the phonon band structures should have 21 phonon branches. The highest frequency reaches up to 1204 cm$^{-1}$, and is higher than the highest frequency of 1036 cm$^{-1}$ in BSi$^{30}$ and 924 cm$^{-1}$ in Ti$_2$B$_2$$^{31}$, indicative of robust Fe-B and B-B interactions in FeB$_6$ monolayer. Furthermore, the absence of virtual frequencies at any high-symmetry direction also confirms the dynamic stability of the FeB$_6$.

4. Conclusions

A proposal that is the possibility of building graphene analogue FeB$_6$ with the Fe@B$_6$H$_6$ aggregates is presented theoretically in the present study. Adopting the dehydrogenation route to polymerization, we can not only place a Fe atom into hexagonal hole instead of B atom in boron sheet, achieving the uniform chemical doping, but also obtain a very large hexagonal-hole-density that is very near to 1/3. The graphene analogue FeB$_6$ await experimental verification because it may possess some novel electronic and chemical properties.

Declarations
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Conflicts of interest/Competing interests (include appropriate disclosures)

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Availability of data and material (data transparency)

All data generated or analysed during this study are included in this published article

Code availability (software application or custom code)

We own the copyright of Gaussian09 and Dmol3.

Authors' contributions

Chao Wang: performed the data analyses and wrote the manuscript;

Jianhua Hou: contributed to the conception of the study;

Qian Duan: helped perform the analysis with constructive discussions.

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Tables

Due to technical limitations, table 1,2 is only available as a download in the Supplemental Files section.

Figures
The optimized geometries of D6h monomer Fe@B6H6, D2d dimer (Fe@B6)2H10 and C2v trimer (Fe@B6)3H12.

Figure 1
Figure 2

Valence electron molecular orbitals of trimer (Fe@B6) 3H12 and triphenylene with NICS in ppm.
Figure 3

The optimized geometries of two kinds of different dimerization of the trimer (Fe@B6)3H12.
Figure 4

The optimized geometries of graphene analogue FeB$_6$ with bond distances in angstroms (Å).
Figure 5

Calculated phonon dispersion curves of FeB6 monolayer.

Supplementary Files

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- Table1.jpg
- Table2.jpg