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Graphene/Cu composites: electronic and mechanical properties by first-principles calculation

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Abstract: Graphene characterized with ultrahigh intrinsic strength and excellent electronic properties is an ideal material to reinforce metals without despairing their thermal and electrical properties. Here, the electronic and mechanical properties of graphene intercalated copper (graphene/Cu) composites are investigated using density functional theory calculations. Graphene/Cu systems present an excellent electrical conductivity and increasing Debye temperature from 335K for pure Cu to over 535K in regardless of stacking models. In addition to greatly enhanced Young's modulus (149%), shear modulus (156%) and bulk modulus (108%) compared to copper, the ultimate strength of graphene/Cu composites are enhanced by 174% and 162%, in x and y directions, respectively. The strengthening and toughening effects of graphene in the composites is originated from strain strengthening and load transfer, which is consistent with the experimental results. Based on this calculation, the strengthening mechanism can be understood, which explains many experimental observations and also provides us a guide to improve graphene/metal composites quality.

Keywords: Graphene; first principles; physical properties; mechanical properties; deformation mechanism
1、Introduction

Copper (Cu) is one of the most extensively used material due to its high electrical and thermal conductivities combined with environment-friendly and easy available material with good corrosion resistance. Thanks to these excellent properties, Cu has wide variety of applications, such as in integrated circuit, electrical conductors, radiators, and electrodes and so on [1,2]. As a result of its poor mechanical properties, Cu is not considered as a good material for structural applications, which has greatly limited its application. Recently, scientists and engineers have developed various strategies to get rid of weak mechanical properties without deterioration of original physical properties, such as metal matrix composites, nanolaminated structure, and nano-twinned structure, gradient structure, multi- and bimodal microstructures [3-6]. Among which, introducing graphene reinforcement is an effective method to obtain a Cu matrix composite with stable structural and outstanding functional performances. Recently, graphene has aroused considerable interest around the world due to its superior mechanical properties and stiffness, excellent thermal properties and low weight [7-10]. Several experimental investigations demonstrated that graphene-reinforced metal composites have higher strengths, better electrical conductivity and thermal conductivity in combination with lighter weights, compared with conventional metals and composites [11-14].

In order to meet the constantly increasing demands for structural strength and energy efficiency, graphene reinforced Cu matrix composites were analyzed through some experimental investigations. The results revealed that adding a small amount of
graphene resulted in valuable improvements in the mechanical properties of Cu matrix. For example, according to Wang et al. [13] by means of reducing the graphene oxide fabricated then spark plasma sintering at 600°C for 5 min with an applied pressure of 40 MPa synthesis of graphene-copper composites within the micro-layered grains, which results showed that the graphene not only improved the strength greatly, but also improved the conductivity by adding 2.5% volume of graphene oxide. Jiang et al. [14] used simple slurry based process and powder metallurgy at 30 MPa and 973 K for 5 min to fabricate bulk graphene/Cu composite, yield strength was up to 172 MPa, which was 90% higher than pure Cu with adding 0.3wt% of graphene to matrix. Additionally, Chen et al. [15] through a graphene in-situ grown approach, which involved ball-milling of Cu powders with PMMA as solid carbon source, in-situ growth of graphene on flaky Cu powders and vacuum hot-press sintering fabricated copper composite with 0.95 wt% graphene with a noticeable increase in yield strength of 244 MPa, which was 177% enhancement. As a two-dimension lamellar material, graphene has unique deformation mechanism. Yet, the strengthening mechanism of graphene is confined to second-phase strengthening [16-17]. To optimize strength and toughness, it is highly required to understand the underlying atomic details of strengthening mechanism. Unfortunately, at the atomic scale, our knowledge regarding the interaction mechanism between single layer graphene and copper is still very limited. For example, even the most fundamental question, “what is the deformation mechanism of composites?” is still unclear.

A few atomistic simulations were also carried out to investigate the role of
graphene in improving the mechanical properties of composites. Duan et al and He et al. conducted a series of molecular dynamics simulations to examine the tensile behavior of graphene-embedded copper nanocomposites [18-19]. It was found that the graphene-embedded copper has better Young's modulus, tensile strength and fracture strength in comparison with pure Cu, which attributed to inherent axial stiffness of graphene and considerable interface slip stresses. Using a similar method, shear strength and toughness of graphene copper nanolayered composites were investigated by Liu et al [20]. Their findings indicated that interfacial constraining effect between graphene and copper layer play an important role in improving the shear strength and the toughness of nanolayered composites. Comprehensively it can be known that although the graphene have a remarkable effect on copper matrix composites, the simulation of deformation mechanism of these composites is only limited to the nanoscale. The deformation mechanism of composites in atomic level remains undefined, and the mechanism of interaction between atoms is not very distinct. Here, we use first-principles to simulate the tensile process of composites, hoping for more detailed analyses to reveal the strengthening mechanism of graphene in composites and understand the deformation mechanism of Cu(111)/graphene/Cu(111) (graphene/Cu) composites at the atomic-scales.

This paper is organized as follows. Section 2 deals with the computational methods. Optimized geometric structure models are described in Sec. 3.1. In Sec. 3.2 we carried out the analysis of energies, structure and interface stability models. Section 3.3 contrasts physical properties of composites and pure copper. In Sec. 3.4
we performed tensile simulation and study their mechanical stability. In Sec. 3.5, deformation mechanism of composites is discussed by means of change of magnitude of atomic charge, displacement change of atoms and generalized stacking fault energies curves. Finally, Section 4 concludes with a summary.

2. Computational methods

First-principles calculations are performed based on pseudopotential and plane-wave basis, set within the density functional theory (DFT) using the Cambridge Serial Total Energy Package (CASTEP) software. We used the generalized gradient approximation (GGA) with the Perdew-Burke-Ernzerhof parametrization (GGA-PBE) for exchange-correlation function [21-22]. A plane-wave cut-off kinetic energy was taken as 340 eV. During geometry optimization, the Brillouin zone sampling was carried out with a \( 5 \times 5 \times 1 \) Monkhorst–Pack k-point grid. To obtain the correct calculation results, we employed the DFT-D method proposed by Grimme [23] to account the long-range van der Waals interactions and optimized the coordinates of all atoms in the slab and the lattice parameters (a, b, c, \( \alpha \), \( \beta \) and \( \gamma \)). The models calculations were followed by cell optimization with an energy convergence tolerance of \( 1.0 \times 10^{-5} \) eV/atom, a maximum displacement of 0.001 Å, and maximum force of 0.03 eV/Å.

The results presented in this work are obtained using a \( 2 \times 2 \) supercell for pure copper, having 24 copper atoms and graphene/copper systems with 24 copper atoms and 8 carbon atoms in graphene layer. The composites models are composed by monolayer graphene, and six atomic layers of copper, and the stacking way about
three layers of copper atoms upper graphene is exactly same with copper atoms under graphene. The mismatch between Cu(111)’s surface lattice constant of 2.56 Å and graphene’s lattice constant of 2.46 Å is 3.8%. The experimental results show that 60% of graphene grains incline to rotate by 3° with respect to the substrate Cu(111) [24]. In order to account for the lattice mismatch [25] and to describe graphene as realistically as possible [26], the simulation cell in-plane dimensions chosen the experimental graphene lattice constant, then adapting the lattice constant of copper accordingly.

To validate the accuracy of the current theoretical scheme, the crystal lattice constants of the composites were calculated to be \( a = b = 2.49 \) Å, which is in good agreement with the calculated graphene’s lattice constant of 2.46 Å [25], the C–C bond length was calculated to be 1.435 Å on graphene/copper systems, which is in good agreement with the calculated value of 1.425 Å [9]. The interlayer distances between graphene and Cu(111) are also in good agreement with previous investigation [27,28]. These validate the application of the methodology to study physical and mechanical properties.

3. Result and discussion

3.1 Geometric structure

In the vertical direction of the Cu(111) one distinguishes three nonequivalent Cu planes/layers, we name the copper layers as top, hcp and fcc, as shown in Fig. 1 (top, hcp and fcc layers of coppers expressed in blue, pink and yellow respectively). According to which carbon atoms sit over in copper layer, we test three different commensurate configurations, which could give rise to three possible graphene
physisorbed positions named as top-fcc (model a), top-hcp (model b), and fcc-hcp (model c) configurations [25]. Fig. 1a, b, c and d show the side and top views of the pure Cu and three geometric structures. As shown in Fig. 1b, in top-fcc (model a), where one carbon atom is on top of a copper atom of the top layer, while another nonequivalent carbon is over the fcc Cu layer. For top-hcp (model b), the two nonequivalent carbon atoms are above on the top or hcp Cu layer. Similarly to fcc-hcp (model c) configuration, the two nonequivalent carbon atoms are situated above the Cu atoms of the hcp and fcc sites. Moreover, top view can be seen that yellow, blue copper atoms and graphene make up the interfaces of composites.

After optimizing those models, the distances between graphene and adjacent copper atomic layers are 2.813 Å/2.803 Å, 2.965 Å/2.881 Å, 2.855/2.964 Å for model a, model b and model c as shown in Fig. 1b, c and d, respectively.

![diagram](image)

**Fig. 1** Top and side view of pure Cu, model a (top-fcc), model b (top-hcp) and
model c (fcc-hcp). Blue, pink and yellow (large) spheres indicate the copper atoms, black (small) spheres are the carbon atoms. The copper layers are labeled by top, hcp, and fcc (top, hcp, and fcc layers coppers expressed in blue, pink and yellow respectively).

### 3.2 Energies, structure and interface stability

To examine the structure stabilities of different models, the binding energies ($E_b$) [29] are calculated based on the following formulas:

$$E_b = \frac{1}{m+n} (E_{\text{total}} - mE_{\text{Cu iso}}^{\text{Cu}} - nE_{\text{iso}}^{\text{C}})$$

where $E_{\text{total}}$ is the total energy of the bulk model, $m$ and $n$ represents copper atoms and carbon atoms in graphene respectively. Based on the computation models, we know $m=24$, $n=8$; $E_{\text{Cu iso}}^{\text{Cu}}$ and $E_{\text{iso}}^{\text{C}}$ are energies of the isolated copper atom and carbon atom in graphene.

In the case of composite materials, especially in mechanical performance, the interface plays a decisive role. Interfacial energy ($E_{ib}$) and interfacial separation work ($W_{sep}$) properties between graphene and Cu (111) are calculated as [30]:

$$E_{ib} = E_{GR} + E_{\text{slab}}^{\text{Cu}} - E_{\text{total}}$$

where $E_{ib}$ is the total energy of the graphene/copper interface structure; $E_{GR}$ is total energy of monolayer graphene with 8 carbon atoms. $E_{\text{slab}}^{\text{Cu}}$ is the six-layer reference Cu slab with interface areas.

$$W_{sep} = \frac{(E_{1}^{\text{tot}} + E_{2}^{\text{tot}} - E_{12}^{\text{tot}})}{S}$$

where $E_{i}^{\text{tot}}$ is the total energy of separated slab i; $E_{12}^{\text{tot}}$ is the total energy of whole system, and $S$ is the total interfacial area of the unit cell. As shown in Fig.1, Model b,
c and d all have two interfaces, the upper named $W_{\text{upper}}$, the lower named $W_{\text{lower}}$.

**Table 1** $E_{\text{b}}$ (eV per atom), $E_{\text{ib}}$ (eV) and $W_{\text{sep}}$ (J/m$^2$) of different models

| Construction | $E_{\text{b}}$ | $E_{\text{ib}}$ | $W_{\text{upper}}$ | $W_{\text{lower}}$ |
|--------------|--------------|--------------|-----------------|-----------------|
| Model a      | -2.946       | 10.12        | 6.03            | 6.03            |
| Model b      | -2.944       | 10.04        | 5.99            | 6.03            |
| Model c      | -2.943       | 10.03        | 6.01            | 5.99            |

The calculated values of $E_{\text{b}}$, $E_{\text{ib}}$, $W_{\text{upper}}$ and $W_{\text{lower}}$ of three different configurations are shown in Table 1. Table 1 tells us in three models $E_{\text{b}}$, $E_{\text{ib}}$ and $W_{\text{sep}}$ all have a very small difference. The absolute values of $E_{\text{b}}$ indicates that model a is the most stable configuration with $E_{\text{b}}$ of -2.946 eV per atom; on the contrary, model c is the most unstable structure with $E_{\text{b}}$ of -2.943 eV. The interface energy $E_{\text{ib}}$ is also called the interface formation energy, the greater the value, the more stable the interface. Analysis of interface energy $E_{\text{ib}}$ indicates that interfacial stability follows the order like model a (10.12eV) > model b (10.04eV) > model c (10.03eV). $W_{\text{sep}}$ is closely related to the layer distances and the atomic bonding at the interface [29]. The greater the separation work, the smaller the interval between the layers. The layer distance of model a is the smallest, indicating that the adhesion of the interface is the strongest, whose values of $W_{\text{upper}}$ and $W_{\text{lower}}$ are almost equal, the interlayer of the upper and lower copper atoms while graphene in the middle are nearly same. This change of law also applies to model b and model c. Table 1 indicates that energies, structure and interface stability among model a, model b and model c have a little difference, which result in their electrical conductivity, Debye temperature and tensile properties mentioned below having a tiny discrepancy.

### 3.3 Physical properties
3.3.1 Electrical conductivity

According to Boltzmann transport equation and Einstein relation, electrical conductivity is proportional to density of states [31]. As shown in Fig.2(a), at the Fermi level ($E_F$), the numeric values of total density of states for copper atoms in model a, b and c are 7.18, 7.18 and 7.17 states per unit cell per eV, respectively. However, the numeric value of total density of states of pure copper is 5.82 states per unit cell per eV, as shown in Fig.2(b). Hence one can conclude that the addition of graphene promotes the conductivity of copper matrix [32].

![Fig.2](a) PDOS of atom Cu and C in Cu/graphene composites ; (b)DOS for pure Cu

3.3.2 Debye temperature

Debye temperature is a fundamental parameter for crystals, closely related to many physical properties of bulk materials, such as specific heat, thermal conductivity, melting temperature and so on, also closely bounded up with structural stability and the strength of bonds of the solids. It can be estimated by the average sound velocity ($v_m$) [33].

\[
\Theta_D = \frac{\hbar}{K_B \left( \frac{3n}{4\pi n v_m} \right)^{1/3}}
\]

Where $\hbar$ is the Plank’s constant, $K_B$ is the Boltzmann’s constant, $n$ is the number of
atoms in cell and \( V_a \) is the cell volume. The average sound velocity can be calculated in terms of the longitudinal \( (v_l) \) and transverse \( (v_t) \) elastic wave velocity as \([33, 34]\).

\[
v_m = \left[ \frac{1}{3} \left( \frac{1}{v_l} + \frac{2}{v_t} \right) \right]^{-1/3}
\]

\[
v_l = \left( \frac{3B+4G}{3\rho} \right)^{1/2}
\]

\[
v_t = \left( \frac{G}{\rho} \right)^{1/2}
\]

Where \( \rho \) represents the density of materials; \( v_l \) and \( v_t \) is the longitudinal and transverse sound velocity, which are obtained by the bulk modulus \( (B) \) and shear modulus \( (G) \). Bulk modulus \( (B) \) and shear modulus \( (G) \) are firstly calculated from the single-crystal elastic constants by Voigt-Reuss average scheme \([35, 36, 37]\).

**Table 2** The calculated Debye temperature \( (\theta_D, \text{K}) \), sound velocities \( (\text{in m/s}) \), density \( (\text{g/cm}^3) \), bulk modulus \( (\text{GPa}) \) and shear modulus \( (\text{GPa}) \) of different models

| Phase    | \( \rho \) | B   | G   | \( v_l \) | \( v_t \) | \( V_m \) | \( \theta_D \) |
|----------|------------|-----|-----|-----|-----|-----|-----|
| Pure Cu  | 8.93       | 137.4 | 46.3 | 4723 | 2277 | 2560 | 335\[^{[38]}\] |
| Model a  | 7.88       | 148.7 | 72.0 | 5573 | 3024 | 4181 | 564 |
| Model b  | 7.79       | 133.1 | 65.3 | 5317 | 2896 | 3972 | 535 |
| Model c  | 7.80       | 153.3 | 67.7 | 5587 | 2945 | 4058 | 546 |

The calculated Debye temperatures and sound velocities of graphene/Cu composite materials are listed in Table 2. Generally, the addition of graphene significantly changes the Debye temperature of copper. The numerical values increase from 335K (pure Cu) to 564K (Model a), 535K (Model b) and 546K (Model c). The disparity in the Debye temperature among graphene/copper composites is small, which is consistent with their structural characters above mentioned. Owning to higher Debye temperature, graphene/copper composites are supposed to be more thermodynamically stable and stronger bonding force between atoms compared to pure Cu \([39]\).
3.4 Stretching simulations

Except the excellent electrical conductivity and thermodynamically stable favorable condition for graphene/copper composites, the mechanical stability is crucial to the feasibility of crystal existence. The criteria of mechanical stability could be evaluated by means of stretching simulations.

In our stretching calculations, the most stable graphene/copper structures (model a, b and c in Fig. 1) are taken as the starting configuration, and pure Cu is stretched as well to make a comparison. Then the uniaxial strains are carried out in certain axis direction (x-, y- or z-axes displayed in Fig. 1, where x-, y-axes represent parallel to graphene layer and z-axes represents vertical directions of graphene layer) step by step. On each of the step, all the atoms are allowed full freedom of motion until the system reaches the minimum energy, and every initial structure for next step is based on the pro-optimum structure.

For x and y directions, graphene/Cu systems get obvious strength enhancement, which are separately 174% and 162% enhancement over pure Cu. Stretch along the x axis in Fig. 3 (a), for pure copper, ultimate tensile strength is 16.41 GPa and the failure strain is 44%. The ultimate tensile strength of model a, b and c are 28.48, 28.49 and 28.46 GPa respectively and failure strains are 23%. In Fig. 3 (b) stretch along the y axis, the ultimate tensile strength of pure Cu is 14.27 GPa and the failure strain is 40%. The ultimate tensile strength of model a, b and c are 26.43, 26.66 and 26.78 GPa respectively and failure strains are 23%. When composite materials are stretched parallel to the direction of graphene (in x-, y-axes), the $sp^2$ bonds in
graphene begins to break at 21%, as shown in Fig. 3(a) and 3(b). During tensile deformation, the rupture of $sp^2$ bonds will lead to the rupture of other bonds in another graphene layer and the copper layer, eventually result in the failure of the materials. Stress has a dramatic decrease in the stress–strain curve as the $sp^2$ bonds are broken, which indicates that the tensile strength and fracture strain of the graphene/Cu composites are dominated by the $sp^2$ bonds of the graphene [18, 40].

Stretch along the z axis, about pure Cu, the ultimate tensile strength is 21.8GPa and the failure strain is 32%. The ultimate tensile strength of model a, b and c are 4.55, 4.42 and 4.40 GPa, but materials is failure in a small amount of deformation, as shown in Fig. 3 (c). Finally, bulk modulus (B) and shear modulus (G) obtained for Cu and graphene/Cu systems calculated from the single-crystal elastic constants by Voigt-Reuss average scheme, where the Young’s modulus (E) is estimated by this equation [37]:

$$E = \frac{9GB}{G+3B}$$

Bulk modulus are 137.4(pure Cu), 148.7(model a), 133.1(model b) and 153.3(model c) GPa; shear modulus are 46.3(pure Cu), 72.0(model a), 65.3(model b) and 67.7(model c) GPa; Young’s modulus are 124.9(pure Cu), 186.1(model a), 168.4(model b), 176.9(model c) GPa. Apparently, graphene/Cu composites have better ability of deformation resistance than pure Cu. Above results clearly show that the mechanical properties of graphene/Cu composites are greatly improved and graphene is a good reinforcing agent in the Cu matrix. Compared with pure Cu, the mechanical enhancement in paralleled direction is mainly contributed to graphene layer with an
outstanding mechanical property. In z-axis direction (vertical direction of graphene layer), mechanical properties get decreased after inserting graphene, suggesting that graphene has a poor interfacial interactions with both sides of Cu [24]. In the future, the mechanical properties can be improved by improving the bonding ability of the interface.

3.5 Mechanical properties analysis

The tensile properties of model a, model b and model c in same drawing direction are almost identical, which shows that for graphene/Cu composites, whose stacking ways has a small effect on mechanical properties. Consequently, we select model c as a representative to intensive study in terms of changes of charge, atomic position and generalized stacking fault energies to reveal the strengthening mechanism of graphene in composites.

3.5.1 Change of magnitude of atomic charge during tensile process

Change of magnitude of atomic charge is closely related to the interaction between atoms. Under different tensile strains, magnitude of atomic charge about model c at different layers has been counted, as shown in table 3(charge means total charge of all the atoms in one layer). \(Cu_i, \ i = 1, 2, 3, 4, 5\) and 6, which represents the copper atoms in specified layers (for example, \(Cu_1\) represents the first layer of model c in which layer copper atoms also has a math numeric “1”, as shown in Fig. 4(a); \(Gr\) represents total charge of graphene transfer). For x and y directions, the magnitude of atomic charge in \(Gr\), \(Cu_3\) and \(Cu_4\) are increasing, while the other layers are almost unchanged. In z-axes, the amount of charge in \(Gr\) is decreasing, while in \(Cu_2, Cu_3,\)
Cu₄ and Cu₅ gradually increasing. The above results suggest that in parallel to the direction of graphene (in x-, y-axes), the interaction between graphene and adjacent copper atoms (Cu₃ and Cu₄) became stronger, while in vertical directions of graphene layer (in z-axes), the interaction between interfaces is gradually diminished, and the interaction between the matrix copper atoms is becoming stronger.

### 3.5.2 Images of atomic arrangements at different tensile strains

To better understand the deformation process of graphene/Cu composites, different deformation images of atomic arrangements about model c as shown in Fig. 4. Fig. 4(a) and 4(b) show in x-axes and y-axes, the images of the atomic arrangements at 11%, 17% and 21% tensile strains. Stretching along x-axes, atoms slip along [101̅] direction in a (111) plane; Stretching along y-axes, atoms slip along [110] direction in a (111) plane. As shown in Fig. 4(a) and 4(b), in the atomic glide direction, the copper atoms in Cu₃ and Cu₄ (dark yellow and dark blue) near the graphene has an obvious tendency to close to each other compared with copper atoms in Cu₁ and Cu₆ (light yellow and light blue, which sit over same carbon atoms as Cu₃ and Cu₄). It demonstrates that graphene could effectively prevent cracks initiation and improve the mechanical properties of the materials. These change of laws are the same as model a and model b.
Fig. 3 Strain-stress curves for pure Cu and graphene/Cu systems in x (a), y (b), and z (c) axes directions (top views of graphene in model c at 8%, 15% and 21% tensile strains are shown as in sets, side views of model c under 8% and 19% strains
are also shown as in sets; $\sigma$ and arrow indicate the direction of external stress).

Table 3 Under different tensile strains, atomic charge of model c in different layers

| Phase | Species | Stain(%) | 0  | 5  | 8  | 11 | 13 | 15 | 17 | 19 | 21 |
|-------|---------|----------|----|----|----|----|----|----|----|----|----|
|       |         |          |    |    |    |    |    |    |    |    |    |
| Model c-x | Cu$_1$ | 0.12     | 0.12 | 0.08 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 |
|         | Cu$_2$ | 0.36     | 0.36 | 0.36 | 0.36 | 0.36 | 0.36 | 0.4  | 0.36 | 0.36 | 0.36 |
|         | Cu$_3$ | 0        | 0.04 | 0.08 | 0.36 | 0.36 | 0.36 | 0.4  | 0.4  | 0.4  | 0.4  |
| Model c-y | Gr     | -0.84    | -0.88 | -0.92 | -1.4 | -1.44 | -1.44 | -1.6 | -1.64 | -1.76 | -1.76 |
|         | Cu$_4$ | -0.08    | -0.08 | -0.08 | 0.12 | 0.16 | 0.16 | 0.24 | 0.32 | 0.4  | 0.4  |
|         | Cu$_5$ | 0.36     | 0.36 | 0.36 | 0.32 | 0.32 | 0.32 | 0.32 | 0.32 | 0.32 | 0.36 |
|         | Cu$_6$ | 0.08     | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 |
| Model c-z | Cu$_1$ | 0.12     | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 |
|         | Cu$_2$ | 0.36     | 0.32 | 0.32 | 0.36 | 0.36 | 0.36 | 0.36 | 0.36 | 0.36 | 0.36 |
|         | Cu$_3$ | 0        | 0.04 | 0.12 | 0.4  | 0.4  | 0.4  | 0.36 | 0.4  | 0.4  | 0.4  |
| Model c-z | Gr     | -0.84    | -0.88 | -0.96 | -1.44 | -1.44 | -1.48 | -1.64 | -1.64 | -1.76 | -1.76 |
|         | Cu$_4$ | -0.08    | -0.08 | -0.08 | 0.12 | 0.12 | 0.12 | 0.2  | 0.24 | 0.32 | 0.4  |
|         | Cu$_5$ | 0.36     | 0.36 | 0.36 | 0.32 | 0.32 | 0.32 | 0.32 | 0.32 | 0.32 | 0.36 |
|         | Cu$_6$ | 0.08     | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 |

Fig. 4 atomic configuration images of model c at various strains in x (a), y (b),
3.5.3 Parallel to graphene layer, displacement change of atoms in different layers

In order to further understand the effect of graphene in parallel to graphene layer direction, displacement change of different layers of atoms is discussed, as shown in Table 4. Throughout the study process, we find copper atoms in same layer have the same atomic displacement change, and the change rule of nonequivalent carbon atoms are inconsistent, other carbon atoms consistent with them. We select one copper atom in each layer and two nonequivalent carbon atoms to study their displacement change. Research indicates that the change of atomic displacement in the same layer is exactly the same, yet the variation of displacement in different layers has a big disparity, which indicating that the tensile deformation of the composites are carried out in a layer, that is to say, deformation of composites is achieved by means of the slip deformation between layer to layer.

In table 4, $\Delta X$ and $\Delta Y$ represent the differences between the atomic fractional coordinates of 21% and 0% separately. The displacement change of the copper atoms ($\text{Cu}_3$ and $\text{Cu}_4$) near the graphene is larger. However, the shift/displacement change of copper atoms far away from graphene has a tendency to decrease. Graphene plays a great influence on tensile deformation. In the process of deformation, stress concentration can be delayed effectively because of stress transfer from graphene to matrix through the atomic displacement. Reducing the stress concentration could achieve good plasticity of composites. Meanwhile, strengthening mechanism could
result from deformation hardening from graphene and adjacent copper atoms. It should be noted that the effects of graphene from the present calculation is in good agreement with similar observations in the experiment which state that strengthening effect of graphene/Cu composites is contributed to load transfer [15].

**Table 4** Stretching along x- or y- axis, atomic displacement change in different layers of model c

| Layer | x-axis | y-axis |
|-------|--------|--------|
|       | ΔX     | ΔY     | ΔX    | ΔY     |
| Cu_1  | 0.02   | 0.026  | -0.022| -0.035|
| Cu_2  | -0.012 | -0.006 | 0.011 | -0.001|
| Cu_3  | -0.046 | -0.047 | 0.054 | 0.035 |
| C_1   | 0.108  | 0.137  | 0.124 | 0.09  |
| C_2   | 0.125  | 0.138  | 0.124 | 0.108 |
| Cu_4  | 0.128  | 0.151  | -0.147| -0.143|
| Cu_5  | 0.089  | 0.095  | -0.091| -0.104|
| Cu_6  | 0.054  | 0.056  | -0.053| -0.069|

### 3.5.4 Effect of graphene on stacking fault energy of pure Cu

To validate the aforementioned results, the effect of graphene on stacking fault energy of Cu is investigated. The generalized stacking fault energies (GSFE) could be derived as follows [41]:

$$\gamma = \frac{E_r - E_0}{A}$$

(9)

where $E_r$ is the total energy of the model after the slip of the displacement ($r$) along the Burgers vector (a/6<112>), $E_0$ is the total energy of the model before the slip, and $A$ is the area of the surface. Consequently, the generalized-stacking-fault-energy
(GSFE) curve for pure Cu and model c have been successfully calculated by rigidly shearing at a (1 1 1) plane along a <112> slip direction, as shown in Fig. 5. Unstable stacking fault energy ($\gamma_{us}$) is used to represent the energy barrier of the dislocation emission [42]. Values of $\gamma_{us}$ about pure Cu and model c from the present study are 197mJ/m$^2$ and 193mJ/m$^2$ respectively. The present study of pure Cu is consistent with the $\gamma_{us}$ values (180mJ/m$^2$) from first principles calculation [43, 44]. The addition of graphene in Cu would increase the width of extended dislocations and facilitate the slip of the dislocation [45].

![Fig.5 GSFE curves of pure Cu and model c slipping along (111)/<112> direction, the displacements are in units of their Burgers vectors $b_p$, and a represents the lattice constant (illustrations are models slipping different displacement)](image)

It is of importance to further find out the effects of graphene on the ductility of Cu. Specifically, the ductility parameter (D) could be expressed by the following equation [45, 46].
\[ D = \frac{0.3\gamma_s}{\gamma_{us}} \]  

(10)

Where \( \gamma_{us} \) is the unstable stacking fault energy, \( \gamma_s \) is the surface energy can be calculated as below [47]:

\[ \gamma_s = \frac{E_{slab} - nE_{bulk}}{2A} \]  

(11)

Where \( E_{slab} \) and \( E_{bulk} \) represent for the total energy of the constructed slab model and bulk model, respectively, and \( A \) is the area of the slab model. It should be pointed out that the calculated surface energies of pure Cu and model c are 1.87 and 3.24 J/m². The present study of pure Cu is consistent with the \( \gamma_s \) values (1.83 J/m²) from experiment [48]. Ductility parameter (D) of pure Cu and model c are greater than 1, implying that they are plastic materials. Graphene considered here to show a tendency to decrease both the unstable fault energy and intrinsic fault energy for the current slip system, which makes the dislocation of glide easier and slow the stress concentration. As shown in Fig.6, the copper atoms adjacent to graphene have a large amount of deformation and transfer the stress to the distance. Generally, graphene/copper systems are expected to show superior strength due to strain strengthening meanwhile maintaining good ductility during the load transfer.
Fig. 6 The schematic illustration of the distribution of stress under specific deformation

4. Conclusion

In this work, the structure and interface stability, electrical conductivity, Debye temperature and mechanical properties of graphene/Cu composite materials are systematically investigated by the first principles calculation. Compared with pure Cu, composites undergo excellent electrical conductivity, Debye temperature increase from 335K (pure Cu) to 564K (Model a), 535K (Model b) and 546K (Model c). Young's modulus, shear modulus and bulk modulus are significant improved. The following conclusions could be achieved. Parallel to the graphene, the interaction between graphene and adjacent copper atoms (Cu$_3$ and Cu$_4$) became stronger with the increase of strain and when the strain is too high it broke due to the failure of the sp$^2$ bonds of graphene. Besides, graphene could effectively prevent crack’s initiation and growth due to stress transfer between atoms. Deformation strengthening improves the
strength of composites. Perpendicular to the graphene, the interaction between interfaces is diminished quickly due to weak force between graphene and copper atoms. In practical application, making graphene’s parallel grow in copper to prepare the layered structure through which we could obtain composites with eminent comprehensive properties.

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**Highlights**

- Deformation mechanism of graphene/Cu composite materials was systematically summarized.
- Graphene/Cu systems present an excellent electrical conductivity and increasing Debye temperature compared with pure Cu.
- Compared to copper, the tensile strength of graphene/Cu composites are enhanced by 174% and 162% in parallel to the direction of graphene.
- The strengthening and toughening effects of graphene in the composites is originated from strain strengthening and load transfer.