



Article Interfacial Characteristics of Graphene-Reinforced Iron Composites: A Molecular Dynamics Study

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Abstract: Interface has a significant effect on mechanical properties of graphene reinforced metal composites. Taking graphene nanosheet reinforced iron composite (Gr/Fe) as an example, the interfacial characteristics of Gr/Fe (110), (111), (11 $\overline{2}$), and (001) interfaces have been studied using molecular dynamics (MD) simulations. Two types of interfacial bonding have been examined: physical and chemical bonding. The results show that when the graphene and iron form a physical adsorption (weak-bonded) interface, the interactive energy of the graphene and Fe (110), (111), $(11\overline{2})$, and (001) interface is -1.00 J/m^2 , -0.73 J/m^2 , -0.82 J/m^2 , and -0.81 J/m^2 , respectively. The lengths of the Fe-C bonding are distributed in the range of 2.20–3.00 Å without carbide formation, and no distinct patterns of atomic structure are identified. When the graphene and iron form a chemical (strong-bonded) interface, the corresponding interactive energy is -5.63 J/m^2 , -4.32 J/m^2 , -4.39 J/m², and -4.52 J/m², respectively. The lengths of the Fe-C bonding are mainly distributed in the ranges of 1.80–2.00 Å and 2.30–2.50 Å, which the carbides such as Fe_3C and Fe_7C_3 are formed at the interface. Moiré patterns are observed at different-oriented interfaces, because of the lattice geometrical mismatch between graphene and different-oriented iron crystal structures. The pattern of diamond stripe is at the (110) interface, which is in good accordance with the experiment. Other patterns are the hexagonal pattern at the (111) interface, the wavy stripe pattern at the $(11\overline{2})$ interface, and the chain pattern at the (001) interface. These moiré patterns are formed through the competition and coordination of the three binding sites (Hollow, Bridge, and Top) of graphene with Fe atoms.

Keywords: graphene-reinforced iron matrix composite; interfacial characteristic; molecular dynamics simulation; moiré pattern; crystallographic orientation

1. Introduction

Graphene, as a model of two-dimensional carbon material with excellent mechanical, physical, and chemical properties, is potentially used in new-generation composites. By combining unique electricity [1–3], heat [4], mechanics [5–8], and other properties [9–11], graphene reinforced metal matrix composites such as copper [12–14], aluminum [15–17], nickel [18,19], and iron [20,21] are prosperous in the applications of future aerospace, military, transportation, and energy industries. For instance, compared with unreinforced aluminum matrix, the tensile strength and elongation of nano-graphene reinforced aluminum matrix composites increased from 233 MPa to 287 MPa and 5.5% to 5.8%, respectively [17]. The yield strength of graphene nanosheet reinforced iron matrix composite can be increased up to 1647 MPa, compared with 110 MPa of pure iron matrix [22]. Since graphene has a large contact area with metal matrix, graphene/metal interface, as a transition region connecting the reinforcement phase and matrix, significantly affects the properties of composites.

The interfacial characteristics of graphene/metal are crucial in graphene reinforced metal matrix composites, where the interfacial reaction often occurs, for instance Al-C



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Copyright: © 2022 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (https:// creativecommons.org/licenses/by/ 4.0/). reaction [23] and Fe-C reaction [24]. In graphene/metal composites, the structure and bonding features at the interface are very complex. By utilizing advanced technology, such as X-ray photoelectron spectroscopy (XPS), scanning tunneling microscopy (STM), and lowenergy electron diffraction (LEED), the orderly cyclical atomic structures (termed as moiré patterns) were often observed at the graphene/metal interfaces [25–27]. Jacobberger et al. presented a comprehensive study of the evolution of graphene/Cu interface as a function of the matrix orientation, which provided a new way to adjust the structure and morphology of graphene reasonably [28]. Rasool et al. studied the atomic-scale characterization of graphene grown on Cu (100) surface by STM and graphene was firstly observed over copper steps, corners, and screw dislocations [29]. Through LEED and STM, Vinogradov et al. found that on the graphene/Fe (110) interface showed a new periodic moiré pattern, and the observed topography of the graphene/Fe superstructure was well reproduced by density functional theory (DFT) calculations, resulted from a unique combination of the lattice mismatch and strong interfacial interaction [30]. Based on near-edge X-ray absorption fine structure (NEXAFS) and photoelectron spectroscopy (PES), Preobrajenski et al. found that the degree of corrugation of graphene critically depended on the strength of chemical bonding at the interface, which was rising in the series of Pt (111)-Ir (111)-Rh (111)-Rh (001) [31].

Meanwhile, interfacial characteristics of graphene reinforced metal matrix composite have been studied through simulations. Pang et al. investigated the effect of defects and grain boundaries on the morphology of graphene on different planes of single crystalline copper substrate through molecular dynamics (MD) simulations, which provided a theoretical basis for the manipulation of graphene morphology in engineering applications [32]. Wang et al. used MD simulations to study the dislocation and graphene interaction in graphene nanosheet reinforced iron matrix composite (Gr/Fe) and the results showed that the yield stress of the composite with a strong interfacial bonding could be enhanced up to 200 MPa over that in the weak-bonded Gr/Fe composite [33]. The relationship between dislocation evolution and graphene orientation in the Gr/Fe composite during compression was investigated [34], in which the yield stress was around 10 GPa and three possible interaction mechanisms of transmission, reflection, and gliding were revealed to be dependent on graphene location. Furthermore, the effects of the number of graphene layers, amorphous layer structure, the bond strength of C-C atoms in graphene, and boundary conditions on dislocation-graphene interaction were also studied by MD simulations [35,36].

The Gr/Fe composites were successfully synthesized through the DC electro–plating method, where this process was carried out under low temperature for avoiding the reaction between Fe and C elements to form carbide phases and ensuring the stability of graphene morphology and structures for excellent performance [37]. Moreover, through a controlled carburization process, graphene-Fe₃C-Fe composites can be obtained [24]. In our previous work, adhesion energy and structure of monolayer, bilayer or trilayer graphene nanosheets located on four differently oriented iron surfaces were studied by MD simulations using two types of Fe-C potentials [38]. However, the interfacial characteristics in Gr/Fe composites are still elusive.

In this work, taking graphene nanosheet reinforced iron composite as an example, the interfacial characteristics of Gr/Fe (110), (111), (11 $\overline{2}$), and (001) interfaces are studied using molecular dynamics (MD) simulations, in which two types of interfacial bonding (weak and strong) are considered. The interaction energy of different graphene/iron interfaces is calculated and the corresponding moiré pattern is further analyzed.

2. Molecular Dynamics (MD) Model and Setup

The MD simulations were performed by Large-scale Atomic/Molecular Massively Parallel Simulator (LAMMPS) software [39]. In Figure 1, the simulation cell was about $200 \times 200 \times 200 \text{ Å}^3$, which included a cubic single crystalline α -iron matrix and a graphene nanosheet with an area of $100 \times 100 \text{ Å}^2$, parallel to the (110), (110), (111), (112), and

(001) crystallographic plane of matrix, respectively. The corresponding X, Y and Z axis settings were [110]-[001]-[110], [112]-[111], [110]-[112]-[111], [111]-[110]-[112], and [100]-[010]-[001], respectively, the periodic boundary conditions were applied to the three directions. Among them, the first and second orientations represented two different {110} interfaces, defined as the type-I and type-II {110} interface. The lattice constant of iron and graphene is 2.856 Å and 1.420 Å, respectively.

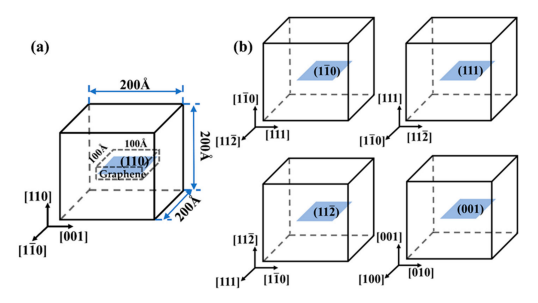


Figure 1. Schematic of molecular dynamic (MD) setting for the graphene nanosheet embedded into iron aligned along different crystallographic plane in the Gr/Fe composites: (**a**) (110) and (**b**) other four plane of $(1\overline{10})$, (111), (11 $\overline{2}$) and (001), respectively.

In the simulation, the C-C interactions were described by Adaptive Intermolecular Reactive Empirical Bond Order (AIREBO) potential [40], the Fe-Fe interactions were modeled by the Embedded-Atom Method (EAM) potential [41]. Considering the complex interactions among Fe-C atoms, two different types of Fe-C potential were selected to describe the chemical (strong-bonded) or physical adsorption (weak-bonded) interfaces of Gr/Fe composites. The results calculated by Lennard-Jones (LJ) potential [42] ($\varepsilon = 0.0168 \text{ eV}$ and $\sigma = 2.221 \text{ Å}$) agreed well with first-principle calculation results, so it was selected to simulate weak/von der Waals force of Fe-C interaction. The EAM potential proposed by Hepburn [41] can reproduce the main structure of cementite and it was selected to describe strong chemical bonding of Fe-C atoms.

The MD simulation process was performed as follows. Firstly, conjugate gradient algorithm [43] was employed to minimize the energy of the system, preliminary static equilibrium of the system was carried out. Graphene and iron matrix were allowed to relax in three directions, and the fluctuation of the stress component in all three directions of the final system was controlled less than 10 MPa. Then, the system was heated to 300 K, the isothermal-isobaric (NPT) ensemble [44] was employed and the system temperature kept at 300 K using Nose-Hoover algorithm [45] running 100 ps. Graphene and iron were initially bound in the thermally activated state. Finally, when the system reached the thermodynamic equilibrium state, the quenching process was used to reduce the temperature of the system to 0 K. After minimizing the whole system energy, the stable interfacial structure of the graphene and iron was obtained. The OVITO software [46] was used to visualize atomic structure (e.g., moiré pattern) at the Gr/Fe interface.

3. Results and Discussion

3.1. Interfacial Characteristics of Gr/Fe Interface

The interactive energy *E*, defined as the energy per unit area to connect graphene nanosheet with iron matrix, is calculated by:

$$E = \sum_{i=1}^{N_{Fe}} \sum_{j=1}^{N_C} \phi_{ij} / A_{Gr}$$
(1)

where ϕ_{ij} is the potential energy between the *i*th Fe atom and the *j*th C atom, N_{Fe} and N_C is the total number of Fe and C atoms in the system, respectively, and A_{Gr} is the contact area between the graphene nanosheet and the iron. The interactive energy (*E*) and average vertical distance ($\overline{D_i}$) of Gr/Fe type-I {110}-(110), type-II {110}-(110), (111), (112), and (001) interfaces are summarized in Table 1. Compared with adhesion energy, the interactive energy excludes the interactions between C-C and Fe-Fe atoms, which could be more directly to measure the interaction between graphene and iron.

Table 1. Interactive energy (*E*) and average vertical distance $(\overline{D_1}, \overline{D_2})$ between the graphene nanosheet and the 1st- or 2nd-closed iron atom layer at the interface, where the graphene nanosheet is aligned on different crystallographic plane of iron matrix, and $\overline{D_0}$ is the initial setting distance between the graphene and iron matrix.

Interface Properties	(110)		_ (110)		(111)		(112)		(001)	
	LJ	EAM	LJ	EAM	LJ	EAM	LJ	EAM	LJ	EAM
$E(J/m^2)$	-1.00	-5.63	-1.00	-5.63	-0.73	-4.32	-0.82	-4.39	-0.81	-4.52
$\overline{D_0}$ (Å)	2.02		2.02		2.06		2.32		2.15	
$\overline{D_1}$ (Å)	2.15	1.73	2.15	1.73	2.05	1.66	2.17	1.75	2.15	1.76
$\overline{D_2}$ (Å)	4.16	3.78	4.16	3.78	3.61	2.36	3.27	2.76	4.16	3.19

Note: LJ and EAM represent the weak and strong bonding of Fe-C, respectively.

The result shows that when graphene and iron form a weak-bonded interface, the interactive energy of the Gr/Fe (110), (110), (111), (112), and (001) interface is -1.00 J/m^2 , -1.00 J/m^2 , -0.73 J/m^2 , -0.82 J/m^2 , and -0.81 J/m^2 , respectively. When graphene and iron form a strong-bonded interface, the corresponded interactive energy is -5.63 J/m^2 , -5.63 J/m^2 , -4.32 J/m^2 , -4.39 J/m^2 , and -4.52 J/m^2 , respectively, which is roughly five times larger than the weak-bonded one. Both the weak- and strong-bonded interfaces show the similar tendencies: the {110} plane is the most stable crystallographic plane among the graphene-iron interfaces, while the {111} one has the lowest stability, which is consistent with the observations in other work [47].

In the weak- and strong-bonded Gr/Fe composites, the average vertical distance (D_1) between the graphene nanosheet and the 1st-closed iron atom layer at the Gr/Fe (111) interface is the shortest one, with the value of 1.66 Å and 2.05 Å, respectively. The $\overline{D_1}$ at other interfaces are very close within the range of 2.15–2.17 Å in the weak-bonded Gr/Fe and 1.73–1.76 Å in the strong-bonded one. Compared to the initial distance ($\overline{D_0}$), there is no significant difference between the $\overline{D_0}$ and $\overline{D_1}$ in the weak-bonded composite. However, the $\overline{D_1}$ in the strong-bonded one changes considerably, indicating that the 1st-closed iron atoms layer are greatly attracted by graphene.

The average vertical distance (D_2) between the graphene nanosheet and the 2nd-closed iron atom layer is 4.16/3.78 Å (weak/strong), 3.61/2.36 Å, 3.27/2.76 Å, and 4.19/3.19 Å at the (110), (111), (112), and (001) interface, respectively. The $\overline{D_2}$ in the weak-bonded composite equals to the $\overline{D_1}$ plus the layer spacing of iron matrix, which means little interaction between graphene and 2nd-layer Fe atoms at the interface. However, the $\overline{D_2}$ in the strong-bonded composite are much shorter, indicating the existence of strong interfacial interaction. The smaller the Fe-Fe interlayer spacing, the stronger the attraction between graphene and Fe atoms. The distributions of the length (*L*) of Fe-C bonding at the Gr/Fe (110)/(110), (111), (112), and (001) interface is provided in Figure 2. It is found that when graphene and iron form a weak-bonded interface, the *L* is distributed in 2.20–3.00 Å without carbide formation. However, when graphene and iron form a strong-bonded interface, the *L* is divided into two ranges of 1.80–2.00 Å and 2.30–2.50 Å, indicating the carbide formation (e.g., Fe₃C and Fe₇C₃) at the interface.

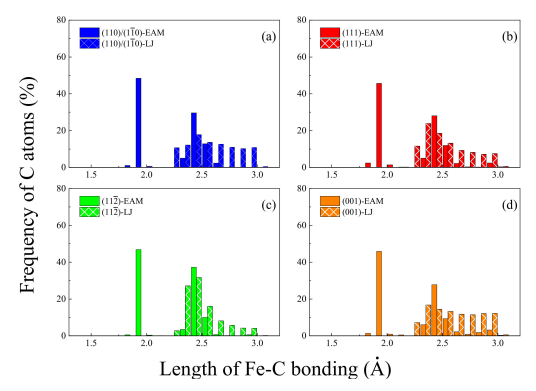
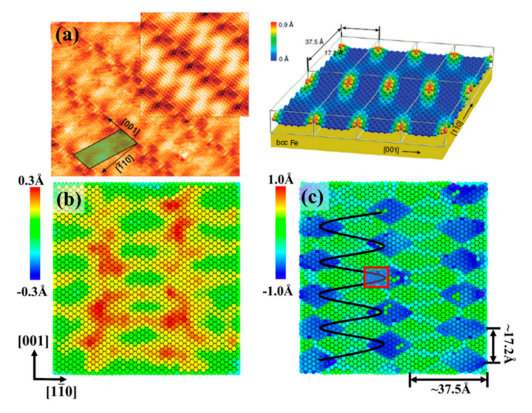


Figure 2. Distribution histograms of the length of Fe-C bonding (*L*) at the Gr/Fe (**a**) (110) or ($\overline{110}$), (**b**) (111), (**c**) ($\overline{112}$), and (**d**) (001) interface, respectively, calculated from MD simulations using LJ and EAM potentials.

3.2. Moiré Pattern of Gr/Fe Typed-I {110} Interface

The Gr/Fe {110} interface, with the lowest interactive energy and the most stable atomic structure, has always been recognized as the easiest crystallographic plane for graphene to grow on. In 2012, Vinogradov et al. made a graphene grown on Fe {110} crystallographic plane at relatively low temperature, and a rhombic moiré pattern was observed periodically along the [001] direction through LEED and STM with an amplitude of 40 Å along the [110] direction [30], as shown in Figure 3a. After enlarging the moiré structure, it could be found that the fluctuation of graphene in the vertical direction were about 0.60 \pm 0.20 Å. Furthermore, the moiré structure at the Gr/Fe (110) interface was found by DFT calculations, as shown in Figure 3b, in which the vertical distance between the graphene layer and the iron surface fluctuated within the range of 2.12–3.03 Å, and the vertical fluctuation of graphene was about 0.90 Å [30].

Due to the limitation of experimental conditions and the spatial scale of DFT calculations, an atomic-scale analysis of the Gr/Fe {110} interface in details is still lacking. Therefore, MD simulation is used to analyze this type of interface. The setup of MD model shows in Figure 1a, in which the armchair and zigzag edge of graphene nanosheet is along [110] and [001] direction, respectively. Figure 3b,c show the atomic structure at the weakand strong-bonded Gr/Fe (110) interface. When graphene and iron form a weak-bonded interface, the (110) interface cannot reveal a clear interfacial topography, the amplitude of Fe atoms at the interface is only 0.2 Å. However, for the strong-bonded interface, the fluctuation range of Fe atoms is two times larger than that of weak-bonded one and a clear



shaped pattern can be observed. Therefore, in the following section, we only focus on moiré patterns in the strong-bonded Gr/Fe composites.

Figure 3. (a) Close-up STM image and positions of Gr/Fe (110) interface calculated by DFT [30], atomic structures from MD simulations using (b) LJ and (c) EAM potentials of Fe-C.

In Figure 3c, a diamond-shaped moiré pattern presented by the displacement of Fe atom is characterized by 35×17 Å² corrugation structure, in which every 18 graphene rings match 19 columns of Fe atoms (-35 Å) in the zigzag direction (i.e., the [110] direction) and 6 graphene rings match 7 Fe–Fe distances (-17 Å) in the armchair direction (i.e., the [001] direction), in consistent with the pattern in Figure 3a. The lattice compatibility of graphene and Fe matrix is the main factor to dominate the periodicity of this pattern.

Figure 4 shows a detailed atomic structure of this moiré pattern. After the system reaches the equilibrium state, Fe atoms are located in three sites: graphene ring center (Hollow-H site), C-C covalent bond center (Bridge-B site), and the site directly above the C atom (Top-T site). After relaxation, the position of C atom is nearly unchanged, while the 1st-layer Fe atom produces a large displacement along the normal direction of the interface, changing from the initial value of 2.02 Å to -1.3 Å (H site) and -2.0 Å (B or T site), and the displacement of 2nd layer Fe atoms is relatively small. To study the stability of different binding site, a single Fe atom is placed in the highly symmetrical H, B, and T site, respectively, and the results show that the stable sequence is H > B > T site, consistent with the DFT results [48–50]. Therefore, the Fe atom in the moiré pattern is mainly located in the H site. However, due to the effect of the lattice mismatch along other orientations of the pattern edge, the Fe atoms can move from the H site to the B or T site.

3.3. Moiré Pattern at Different Gr/Fe Interface

When graphene nanosheet is parallel to different crystallographic plane of the iron, the moiré patterns at the interface shows differently.

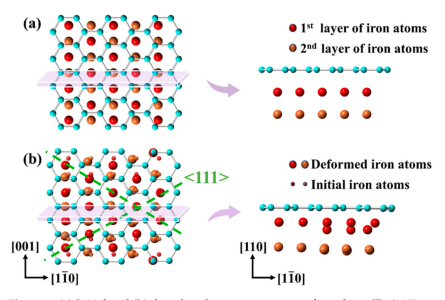


Figure 4. (a) Initial and (b) the relaxed atomic structures of graphene/Fe (110) interface, corresponding to the red region in Figure 3c.

After rotating the graphene nanosheet on the {110} plane of the iron, type II-(110) interface is built with the armchair along the $[11\overline{2}]$ and the zigzag along the [111] direction. In Figure 5a, the moiré pattern at the (110) interface exhibits a linear stripe structure, which is different from the (110) interface. The fringes at the interface extend along the [111] direction with the Gr/Fe lattice mismatch less than 0.5%, where the lattice interval of graphene and Fe is 2.46 Å and 2.47 Å, respectively. Along the [112] direction, the stripe width is about 4.30 Å. The atoms inside the stripe are compressed up to 7% after relaxation and form a clear boundary. In the [112] direction, there is no obvious regularity of the stripe spacing. The Fe atoms are undulating along the vertical direction at the interface with the positions of 1.27–2.04 Å, which is 13–41% larger than that at the (110) interface.

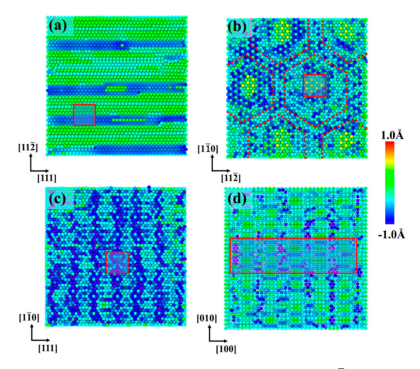


Figure 5. Displacement distribution of graphene/Fe (**a**) $(1\overline{10})$, (**b**) (111), (**c**) $(11\overline{2})$, and (**d**) (001) interface, calculated from MD simulations using EAM potential of Fe-C.

When graphene sheet with the armchair along the $[11\overline{2}]$ and the zigzag along the

[110] direction is embedded into the Fe (111) plane, the moiré structure is identified in Figure 5b. A remarkable highly symmetrical hexagonal structure is observed, and its six sides are along the {111} and (110) directions. There is a transition zone between the hexagons with the width of 7.0 Å. The periodicity of the hexagons is about 59.5 Å, which is similar to the graphene structure in ruthenium (0001), nickel (111), rhodium (111), and copper (111) interfaces.

When the embedded graphene sheet is on the (112) plane, a wavy linear stripe pattern along the [110] direction with an equal spacing of 9.3 Å in the [111] direction is observed, as shown in Figure 5c. In the normal direction of interface, the 1st-layer Fe atoms move toward the graphene and the displacement amplitude is up to 0.87 Å. A chain-like pattern is formed at the Gr/Fe (001) interface in Figure 5d. The Fe atoms fluctuate greatly at the chain junction, compared with those inside.

Moreover, atomic structure characteristics in the red regions of Figure 5 are further analyzed in detail, as shown in Figure 6. The 1st–6th layer Fe atoms are marked with the color of red, orange, purple, blue, pink, and green, respectively.

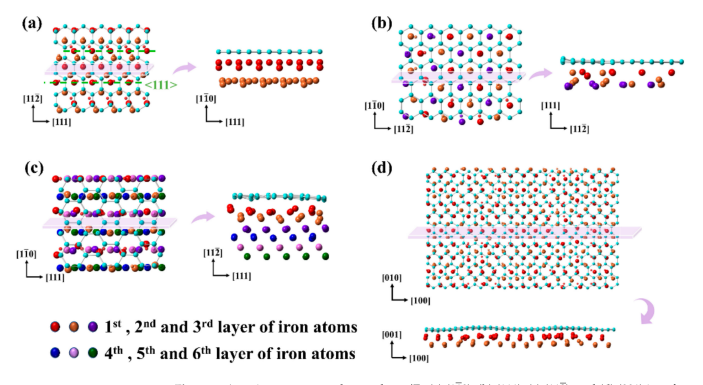


Figure 6. Atomic structures at the graphene/Fe (**a**) $(1\overline{10})$, (**b**) (111), (**c**) $(11\overline{2})$, and (**d**) (001) interfaces, corresponding to the red region in Figure 5a–d, respectively.

Figure 6a shows the atomic structure at the Gr/Fe Type-II (110) interface. The $\overline{D_0}$ is 2.02 Å. After the relaxation, the vertical spacing of the H, B, and T site is 1.27–1.40 Å, 1.75–1.81 Å, and 1.99–2.04 Å, respectively. Therefore, the 1st-closed layer Fe atoms present a wave-shaped undulation with the amplitude of 0.77 Å. The 2nd-closed layer Fe atoms also present relatively less fluctuations in the vertical direction with the corresponding spacing of 3.45–3.50 Å (H site), 3.62–3.71 Å (B site), and 3.95–4.10 Å (T site) and the amplitude of 0.65 Å.

The atomic structure at the Gr/Fe (111) interface shows in Figure 6b. The D_0 is 2.06 A, and the average spacing between the 1st- and 2nd-closed layer Fe atoms is only 0.82 Å. Due to the sparse arrangement of Fe atoms on the {111} plane, the graphene has enough space to form wrinkles. Moreover, owing to a smaller planar spacing between the {111} planes of iron, the graphene has a strong interaction with the three layers of Fe atoms, leading to a displacement fluctuation up to 0.89 Å. After relaxation, the Fe atoms in the central area of

the hexagon are mainly located at the H site, while those at the boundary are located at the B and T sites. The vertical spacing between the graphene and the 1st-, 2nd- and 3rd-closed layer Fe atoms reduces 1.27–1.94 Å, 1.51–2.87 Å, and 2.78–3.64 Å, respectively.

Figure 6c shows the atomic structure at the Gr/Fe (112) interface. The six-layered Fe atoms are periodically arranged along the [111] direction and only the 1st, 2nd and 3rd-closed layer Fe atoms produce obvious displacements. The Fe atoms on the {112} plane are also sparsely arranged with a smaller planar spacing, similar to the case of {111} plane. After relaxation, the graphene sheet fluctuates up to 0.88 Å. The $\overline{D_0}$ changes from the 2.32 Å to the $\overline{D_1}$ of 1.45–2.03 Å with a maximum fluctuation of 0.84 Å. It is also found that the fluctuation of graphene plays a certain role in changing the layer spacing. The vertical spacing between the graphene and 2nd and 3rd-closed layer Fe atoms is 2.04–3.21 Å and 3.78–4.24 Å, respectively, indicating that the vertical displacement of the 2nd layer atoms is significantly greater than the 3rd ones.

Figure 6d shows atomic structure at the Gr/Fe (001) interface. The graphene sheet exhibits a wave-like undulation with the amplitude of 0.55–0.61 Å and the periodic spacing of 8.61 Å. The wrinkles cause the fluctuation of the vertical spacing between the graphene and Fe atoms, leading to the rearrangement of Fe atoms, especially the 1st-layer Fe atoms. It is also found that along the [100] direction, every 3–4 Fe atoms collectively move to the adjacent stable binding position in a row, and finally stabilize at or near the H, B or T site. These atoms are periodically rearranged to form a chain-like pattern. In additional, most of the 2nd-closed layer Fe atoms are only displaced in vertical direction, but due to the wrinkles of the graphene, their bonding positions also changes to a certain extent.

Generally, the proposed approach and the results obtained, especially in terms of considering moiré patterns, can be further applied to a wide class of graphene reinforced metal matrix materials.

4. Conclusions

In this work, interfacial characteristics in graphene reinforced iron matrix composite (Gr/Fe) are systematically investigated using molecular dynamics simulations, associated with the chemical bonding (strong) and physical adsorption (weak) interfaces. The main conclusions are as follows:

(1) When the graphene sheet is embedded in the iron and forms a weak interface, the Gr/Fe (110), (110), (111), (112), and (001) interface interactive energy is -1.00 J/m^2 , -1.00 J/m^2 , -0.73 J/m^2 , -0.82 J/m^2 , and -0.81 J/m^2 , respectively. The average vertical distance between the graphene sheet and the iron interface fluctuates in the range of 2.05–2.15 Å, the bond length of Fe-C atoms is continuously distributed in the range of 2.20–3.00 Å, and the Fe-C atoms tend to be combined by van der Waals force without specific bonding types.

(2) When the graphene sheet is embedded in the iron and forms a strong interface, the interactive energy of Gr/Fe (110), (110), (111), (112) and (001) interfaces are -5.63 J/m^2 , -5.63 J/m^2 , -4.32 J/m^2 , -4.39 J/m^2 , and -4.52 J/m^2 . The average vertical distance between the graphene sheet and the iron interface fluctuates in the range of 1.66–1.76 Å. The bond length of Fe-C atoms is mainly discontinuously distributed in the two intervals of 1.80–2.00 Å and 2.30–2.50 Å. The Fe-C atoms are bonded by chemical bonds, and the carbides such as Fe₃C and Fe₇C₃ are formed at the interface.

(3) When the graphene sheet and the iron form a strong interface, the Gr/Fe (110) interface presents a periodic diamond-shaped moiré structure with size of 35×17 Å², which is consistent with the experimental observation.

(4) Under the strong interface, different Gr/Fe interface shows different moiré pattern: The Gr/Fe ($1\overline{10}$) interface has a linear stripe pattern, the Gr/Fe (111) interface has a highly symmetrical hexagonal pattern, the Gr/Fe ($11\overline{2}$) interface has a wavy linear stripe pattern, and the Gr/Fe (001) interface has a chain-like pattern.

(5) Due to the lattice mismatch between graphene and iron crystal structures, the system can reach the equilibrium state through the competition and coordination of interfacial

Fe atoms at the three stable binding positions of Hollow (H), Bridge (B) and Top(T) sites of the graphene, leading to the formation of moiré pattern at the interface.

Author Contributions: J.J., L.W. and Q.P. conceived and designed the simulations; M.S., L.W. and H.W. performed the simulations; M.S., J.J., S.L., L.W., S.T. and Q.P. analyzed the data; M.S., J.J. and L.W. wrote the paper. All authors have read and agreed to the published version of the manuscript.

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Conflicts of Interest: The authors declare no conflict of interest.

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