Winding a nanotube from black phosphorus nanoribbon onto a CNT at low temperature: A molecular dynamics study

Kun Cai, Lingnan Liu, Jiao Shi, Qing H. Qin

HIGHLIGHTS
• Fabricating a black phosphorene (BP) nanotube from a BP nanoribbon is studied.
• Winding a BP ribbon on a CNT may form a perfect BP nanotube at low temperature.
• At T ≤ 50 K, the BP nanoribbon with perfect length can form a BP nanotube on the CNT.
• A BP ribbon longer than perfect length cannot form a nanotube on the CNT.

GRAPHICAL ABSTRACT

ABSTRACT
Black phosphorus (BP) has become a popular two dimensional semiconducting material. Investigation of a BP nanotube, theoretically and experimentally, is therefore a hot topic attracting many recent researchers. In this paper, a nanotube system is proposed by placing a BP ribbon near a carbon nanotube (CNT) to form a BP nanotube. The system is formed based on the following two aspects. The first is that the CNT has a perfect surface that provides only van der Waals attraction to the BP ribbon. The BP ribbon does not damage the CNT. The second aspect is that a BP ribbon can feature a large curvature without breaks of the phosphorus-phosphorus bonds in it. If the length of the BP ribbon is well chosen to be a ‘perfect length’, the ribbon can form a perfect BP nanotube winding on a CNT at very low temperature. At higher temperature, e.g., ≥50 K, it is difficult to obtain a perfect BP nanotube. With a BP ribbon whose length is greater than the perfect length, it is difficult to create a perfect BP nanotube even at ultralow temperature.

1. Introduction
With the rapid development of nanotechnology, low dimensional nano materials can be obtained easily and their physical properties studied profoundly. Typical low dimensional materials include graphene [1–5], MoS$_2$ [6,7], BN [8,9], and black phosphorus (BP) [10, 11], carbon nanotubes [12–16]. Although the electrical properties of BP have been reported for a few decades [17,18], it has not become popular until recently due to the works of Refs [10,19,20]. In these works, the electric field effect of a field effect transistor made from few-layered BP was studied and excellent transistor performances of such 2D material at room temperature were obtained. There are several allotropic forms of phosphorus. Unlike carbon allotropes that include 3D diamond, 2D graphene, and 1D carbon nanotube (CNT), no nanotube
the axial distance between the two outer tubes. The width of BP ribbon and is identical in all models.

Phosphorus allotropes, the 3sp^3 hybrid orbital based phosphorus-

Fig. 1. Schematic geometric model (in x-y-z coordinate system) of a nano-system consisting of a double walled carbon nanotube and a black phosphorus ribbon. (a) Top view of the system; (b) axial view of the system. \( l_i \) is the length of the inner tube. \( L_0 \) is the axial distance between the two outer tubes. The width of BP ribbon w is \(-2.32\) nm, and is identical in all models. \( l_o \) is the unit length of BP ribbon along the armchair direction, here \(-0.44\) nm. \( l_o \) is the perfect length of BP ribbon along the armchair direction. The internal edges of the two outer tubes are hydrogenated to avoid strong interaction between the CNT and BP ribbon. The atoms on the outer face of BP ribbon are colored blue and those on the inner face are red. The phosphorus atoms on the edge that are the furthest from the inner CNT are yellow and are used to judge the generation of new phosphorus-phosphorus (P–P) bonds. For BP ribbon, the y-direction is the armchair direction and the z-axis is along the zigzag direction.

from a phosphorus allotrope has ever been discovered in laboratory. Among the phosphorus allotropes, BP is the most stable, as each phosphorus atom in a single-layered BP is covalently bonded with three adjacent phosphorus atoms, and all the three valence electrons of each phosphorus atom are occupied. Compared to 2sp^2 or 2sp^3 orbitals in carbon allotropes, the 3sp^3 hybrid orbital based phosphorus-phosphorus (P–P) bond is much weaker, which leads to the lower perfect length of BP ribbon along the armchair direction. Stillinger-Weber potential, with new parameters developed by Jiang [32,33], AIREBO potential [34] is employed to estimate the interaction among carbon and/or hydrogen atoms in a CNT. The interactions among atoms on different components, e.g., inner CNT, outer CNT and BP ribbon, are evaluated through a 12-6 Lennard-Jones potential [35] using the constants listed in Table 2. The interactions between phosphorus and hydrogen atoms are neglected in the present simulations.

It is known that attraction between BP ribbon and inner CNT can change their initial configurations from those shown in Fig. 1. If the interaction between the two components is strong enough, the BP ribbon will curve and cover the inner CNT to reduce the surface energy. To estimate the stable configurations of BP ribbon after a period of attraction by an inner CNT, we perform simulations in the Large-scale Atomic/Molecular Massively Parallel Simulator (LAMMPS [36,37]) package. The whole system is placed in a canonical NVT ensemble with specified temperature T (8 K, 50 K, 100 K, and 150 K) for a period of no more than 1000 ps. The positions in the system are updated every 0.001 ps. In the present study, the temperature is evaluated according to the state

<table>
<thead>
<tr>
<th>Model no.</th>
<th>CNTs</th>
<th>Number of atoms on CNTs and BP ribbon</th>
<th>Perimeter of outer CNT</th>
<th>( l_i ) (nm)</th>
<th>( l_o ) (nm)</th>
<th>( L_0 ) (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>(5.5)/(10,10)</td>
<td>640C + 160C + 40H + 330P</td>
<td>4.26</td>
<td>7.75</td>
<td>5.58</td>
<td>4.75</td>
</tr>
<tr>
<td>2</td>
<td>(6.4)/(11,9)</td>
<td>608C + 160C + 40H + 330P</td>
<td>4.27</td>
<td>7.40</td>
<td>5.67</td>
<td>4.75</td>
</tr>
<tr>
<td>3</td>
<td>(8.0)/(17,0)</td>
<td>608C + 272C + 34H + 330P</td>
<td>4.18</td>
<td>7.95</td>
<td>5.55</td>
<td>4.75</td>
</tr>
<tr>
<td>4</td>
<td>(9.9)/(14,14)</td>
<td>1152C + 224C + 56H + 450P</td>
<td>5.96</td>
<td>7.75</td>
<td>5.58</td>
<td>4.75</td>
</tr>
<tr>
<td>5</td>
<td>(13.4)/(23,2)</td>
<td>1614C + 410C + 50H + 450P</td>
<td>5.92</td>
<td>9.60</td>
<td>5.79</td>
<td>6.49</td>
</tr>
<tr>
<td>6</td>
<td>(15.0)/(24,0)</td>
<td>1140C + 384C + 48H + 450P</td>
<td>5.90</td>
<td>7.95</td>
<td>5.55</td>
<td>6.49</td>
</tr>
<tr>
<td>7</td>
<td>(13.13)/(18,18)</td>
<td>1664C + 288C + 2H + 570P</td>
<td>7.67</td>
<td>7.75</td>
<td>5.58</td>
<td>6.68</td>
</tr>
<tr>
<td>8</td>
<td>(14,12)/(19,17)</td>
<td>1726C + 304C + 72H + 570P</td>
<td>7.67</td>
<td>8.10</td>
<td>5.68</td>
<td>6.68</td>
</tr>
<tr>
<td>9</td>
<td>(22,0)/(31,0)</td>
<td>1672C + 496C + 62H + 570P</td>
<td>7.62</td>
<td>7.95</td>
<td>5.55</td>
<td>6.68</td>
</tr>
</tbody>
</table>

Table 1
Parameters of the nano-system in Fig. 1, and number of carbon (C), phosphorous (P) and hydrogen (H) atoms.
The first factor is the deformation-based elastic potential energy of the BP ribbon. It depends on the radius of curvature of the wound BP ribbon. According to the theory of elasticity, the value of $\Delta P E$ can be calculated according to the equation:

$$\Pi_{\text{Deform}} = \int_{r_1}^{r_2} \frac{E}{2\rho r} \, dr$$

where $E$ is the equivalent Young’s modulus of the BP ribbon along the armchair direction, i.e., $-44$ GPa [22]. $I$ is the inertia moment of the x-z system. The second factor is the deformation-based elastic potential energy of the BP ribbon. The deformation-based elastic potential energy of the BP ribbon leads to an increase rather than a decrease of the potential energy of the system. For that reason, we choose three BP ribbons of different lengths among the three components in system can be expressed as

$$\Pi_{\text{vdW}} = \Pi_{I-O} - \Pi_{\text{I-BP}}$$

where the first part of Eq. (3), i.e., $\Pi_{I-O}$, the interaction between the inner CNT and BP ribbon, is the van der Waals (vdW) interaction energy between CNTs and a BP ribbon. As the width of BP ribbons remains the same ($w = 2.32$ nm), the value of $\Pi_{I-O}$ depends only on the length of BP ribbon. The deformation-based potential energy of the three isolated components from the total potential energy of the system as the distance between two edges of BP ribbon they attract each other. On the other hand, from our previous study [25,26], curvature of BP ribbon along the armchair direction occurs more easily than that along zigzag direction. Hence, the winding direction would be identical to the armchair direction during winding. This prediction is checked via numerical tests. The second item in Eq. (3) is the interaction energy between the outer CNT and the BP ribbon. At the initial stage of the winding process, we place the ribbon at the center of the inner tube, such that the distance between the BP ribbon and the two outer tubes is $>1.5$ nm, i.e., $0.5(L_p-w) > 1.5$ nm. Therefore, the value of $\Pi_{I-O}$ is initially zero. However, if the BP ribbon attracted on the inner tube moves towards an outer tube, the value of $\Pi_{I-O}$ will not be zero. That change may occur soon after the start of winding of the BP ribbon on a chiral CNT. Both $\Pi_{I-O}$ and $\Pi_{I-O}$ lead to a decrease in the $\Delta P E$ of the system. The third part of Eq. (3), i.e., $\Pi_{\text{I-O}}$, represents the interaction between CNTs attributable to the vibration of atoms on CNTs. Because outer CNTs are fixed in simulation, the value of $\Pi_{I-O}$ stabilizes quickly after starting the simulation. The value of $\Pi_{I-O}$, depends on the temperature $T$, i.e., the value of $\Pi_{I-O}$ is higher at higher temperature.

The second factor is the variation of surface potential energy that depends on the interactions among the curved BP ribbon and the inner and outer CNTs. The value of $\Delta P E$ due to surface interactions among the three components in system can be expressed as

$$\Pi_{\text{vdW}} = \Pi_{I-O} - \Pi_{\text{I-BP}}$$

where $N_{\text{unit}}$ is the number of units along the armchair direction of the BP ribbon. The deformation-based elastic potential energy of the BP ribbon leads to an increase rather than a decrease of the potential energy of the system. For that reason, we choose three BP ribbons of different lengths near different CNTs (see Table 1) to reveal the radius/curvature effect on the winding of the BP ribbon.

Table 2

<table>
<thead>
<tr>
<th>Atom $i$</th>
<th>Atom $j$</th>
<th>$\sigma$ (Å)</th>
<th>$\varepsilon$ (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>P</td>
<td>C</td>
<td>3.4225</td>
<td>0.06878</td>
</tr>
<tr>
<td>P</td>
<td>C</td>
<td>4.348</td>
<td>0.01594</td>
</tr>
<tr>
<td>C</td>
<td>C</td>
<td>3.400</td>
<td>0.002844</td>
</tr>
<tr>
<td>C</td>
<td>H</td>
<td>3.025</td>
<td>0.002065</td>
</tr>
<tr>
<td>H</td>
<td>H</td>
<td>2.650</td>
<td>0.001409</td>
</tr>
</tbody>
</table>

During interactions among the components in such a system, the potential energy of the system may change due to variation of configuration. Three major factors influencing the variation of potential energy ($\Delta P E$) of the system are attributed to changes of configuration of the system.

For a bulk atom in BP, the value is $\sim 5.12$ eV, which is estimated by Stillinger-Weber potential [32]. Hence, the generation of every new $P$ bond between two layers with edge atoms (red and yellow) in simulation. $P$ bonds on the curved outer surface of the BP ribbon is very weak, which leads to instability of the BP at high temperature. During the winding of a BP ribbon on a CNT, the $P$-$P$ bonds on the curved outer surface of the BP ribbon are easily broken, leading to an increase in the $\Delta P E$ of the system. This should be avoided if we want to obtain a perfect nanotube from BP ribbon. In Fig. 1b, the phosphorus atoms at the edges along the CNT axis are unsaturated. If there is no broken $P$-$P$ bond, and simultaneously the free edges of BP ribbon along the axis of the CNT join together, new $P$-$P$ bonds could be generated. In this situation, the $\Delta P E$ of the system drops. Mathematically, the chemical reaction energy can be expressed as the total number of new $P$-$P$ bonds, i.e.,

$$\Pi_{\text{React}} = N_{\text{bnd}} \times d \Pi_{P-P}$$

where $N_{\text{bnd}}$ can be considered as the variation of the number of $P$-$P$ bonds in BP ribbon during winding. If there is neither broken bond nor new bond generated, $N_{\text{bnd}} = 0$. As predicted by [39], the potential energy of an edge atom on BP ribbon depends on its position. For the yellow atoms in the BP ribbon shown in Fig. 1, the potential energy of edge atoms is between $[-4.82, -4.76]$ eV (mean value is $-4.79$ eV). For a bulk atom in BP, the value is $-5.12$ eV, which is estimated by Stillinger-Weber potential [32]. Hence, the generation of every new $P$-$P$ bond between two layers with edge atoms (red and yellow) in simulation.
the BP leads to potential energy drop of about twice \(-0.33\) eV, i.e., \(\Delta \Pi p = -0.66\) eV.

From the analysis above, the deformation-based elastic potential energy is formed from both bonded and non-bonded interaction among not only the atoms on BP ribbon but also the neighbor atoms on CNTs. When deformation of BP ribbon needs more energy than the interaction energy between BP and CNTs (Eq. (3)), BP cannot wind on CNT even if the length of BP is high enough. On the other hand, if BP ribbon can be attracted by and tightly upon a CNT, it indicates that the deformation potential energy is far less than the interaction energy among the atoms on both BP ribbon and CNTs. In this case, the variation of the potential energy of the system is able to illustrate the variation process of the configuration of the system.

3. Results and discussion

3.1. Winding of BP ribbon at ultralow temperature, e.g., 8 K

At 8 K, the \(\Delta PE\) histories of the systems with different components are shown in Fig. 2. When the BP ribbon with \(L_p = 4.75\) nm is attracted by axially commensurate double walled CNTs, i.e., (5,5)/(10,10) or (8,0)/(17,0), the \(\Delta PE\) curves are aligned very well (Fig. 2a). A similar phenomenon can also be found at the earlier stage when \(L_p\) is greater (see Fig. 2b, c). During the process, the drop in the potential energy of the system is caused mainly by both vdW interaction and the released energy in generation of new \(P-P\) bonds on the BP ribbon. From the previous theoretical analysis, the quantity of \(\Delta PE\) of the system is actually greater than that given in Table 3 because of the growth of potential energy for deformation of the BP ribbon (\(H_{deform}\) in Eq. (1)) during winding.

<table>
<thead>
<tr>
<th>(L_p) nm</th>
<th>CNTs</th>
<th>Initial PE/eV</th>
<th>(\Delta PE_1 + \Delta PE_2/eV)</th>
<th>WT1/WT2/ps</th>
</tr>
</thead>
<tbody>
<tr>
<td>4.75</td>
<td>(5,5)/(10,10)</td>
<td>-509.99</td>
<td>9.85 + 0</td>
<td>33/33</td>
</tr>
<tr>
<td></td>
<td>(6,4)/(11,9)</td>
<td>-576.47</td>
<td>8.87 + 1.63</td>
<td>30/108</td>
</tr>
<tr>
<td></td>
<td>(8,0)/(17,0)</td>
<td>-659.25</td>
<td>9.84 + 0</td>
<td>32/32</td>
</tr>
<tr>
<td>6.49</td>
<td>(9,9)/(14,14)</td>
<td>-10,326.4</td>
<td>20.50 + 0</td>
<td>43/43</td>
</tr>
<tr>
<td></td>
<td>(13,4)/(23,2)</td>
<td>-13,666.3</td>
<td>20.70 + 0.80</td>
<td>43/88</td>
</tr>
<tr>
<td></td>
<td>(15,0)/(24,0)</td>
<td>-11,436.5</td>
<td>20.40 + 0</td>
<td>44/44</td>
</tr>
<tr>
<td></td>
<td>(13,12)/(18,18)</td>
<td>-14,638.6</td>
<td>30.30 + 0.90</td>
<td>54/267</td>
</tr>
<tr>
<td></td>
<td>(14,12)/(19,17)</td>
<td>-15,290.7</td>
<td>30.50 + 0.70</td>
<td>54/270</td>
</tr>
<tr>
<td></td>
<td>(22,0)/(31,0)</td>
<td>-16,261.6</td>
<td>30.00 + 0</td>
<td>55/55</td>
</tr>
</tbody>
</table>

At 8 K, the \(\Delta PE\) histories of the systems with different components are shown in Fig. 2. When the BP ribbon with \(L_p = 4.75\) nm is attracted by axially commensurate double walled CNTs, i.e., (5,5)/(10,10) or (8,0)/(17,0), the \(\Delta PE\) curves are aligned very well (Fig. 2a). A similar phenomenon can also be found at the earlier stage when \(L_p\) is greater (see Fig. 2b, c). During the process, the drop in the potential energy of the system is caused mainly by both vdW interaction and the released energy in generation of new \(P-P\) bonds on the BP ribbon. From the previous theoretical analysis, the quantity of \(\Delta PE\) of the system is actually greater than that given in Table 3 because of the growth of potential energy for deformation of the BP ribbon (\(H_{deform}\) in Eq. (1)) during winding.

![Fig. 3. Snapshots of the winding process of BP ribbon on different inner CNTs at 8 K.](image-url)
When the BP ribbon is attracted by incommensurate double walled CNTs, e.g., (6,4)/(11,9), the potential energy of the system usually has two jumps, as can be seen from the configurations of the blue curves in Fig. 2. For example, when the BP ribbon winds on a (6,4) tube, the potential energy of the system first drops due to the interaction of BP ribbon and inner CNT ($I_{I-BP}$ in Eq. (3)) during the first 30 ps (Table 3). After a period of running, the second drop in the potential energy of the system occurs. From Table 3 we know that the value of $WT_2$ with respect to the second drop in potential energy of the system is 108 ps. Comparing the configurations of the system from 0 to 108 ps (see group (6,4)/(11,9) in Fig. 3b, and Movie 1), we conclude that the second drop in potential energy of the system is caused by two features occurring simultaneously, namely generation of new P–P bonds (occurs during 100 and 108 ps) ($I_{React}$ in Eq. (4)) and interaction between the BP tube and an outer CNT ($I_{O-BP}$ in Eq. (3)). As $L_p$ is different from 4.75 ns, the second drop in potential energy of the system is no more than 1.0 eV. For example, $\Delta PE_2 = 0.80$ eV when $L_p = 6.49$ nm or $\Delta PE_2 = 0.70$ eV when $L_p = 7.81$ nm (see Table 3). The reason is that the second drop in potential energy of the system is only caused by the attraction between the BP tube and an outer CNT. The fluctuation in the $\Delta PE$ curve in Fig. 2b or 2c is due to frequent collisions between the BP tube and an outer CNT (Movies 2, 3). The reason for the BP ribbon sliding towards an outer CNT is that the potential barrier on the inner chiral CNT shows a helix[40]. The relative rotation between the BP ribbon and the inner CNT determines which of the outer CNTs will become closer to the BP ribbon.

When investigating the winding speeds of the BP ribbon on an inner CNT, we find that the same BP ribbons display the same duration of winding on different CNTs. However, with a chiral inner CNT on which the BP ribbon winds, extra time may be involved with the generation of new P–P bonds once the CNT goes closer to the BP. For example, after 32 ps the BP winds on the inner CNT, whereas at 108 ps the two edges of the BP meet and bond perfectly (see group (6,4)/(11,9) in Fig. 3b).

### 3.2 Winding of BP ribbon at temperature ≥ 50 K

As the temperature increases, the winding process of the BP ribbon on a CNT becomes complex and different from that at 8 K. For example, the $\Delta PE$ of the system shows a quick jump (circled in Fig. 4a, b & c-50 K) at the start of winding. The jump is caused by the sharp variation in the configuration of both the CNT and the BP ribbon after being relaxed in a canonical NVT ensemble at a temperature higher than 8 K. The jump also occurs in other cases at a temperature higher than 50 K. Moreover, the variation of the jump is greater at higher temperature.

Observation of the winding process of the BP ribbon on a CNT at temperatures higher than 8 K shows that fortunately, the BP ribbon can still be perfectly wound on any CNT at 50 K (see the inserts in Fig. 4a, b & c-50 K, Movie 4). However, at a temperature higher than 50 K, e.g., at 100 K, the BP ribbon cannot be perfectly wound on such CNTs as (13,4), (15,0), (13,13), (14,12), and (22,0). That means that it is very difficult to wind a BP ribbon on a CNT with larger radius at the same temperature. As we observe the process of the BP ribbon winding on a CNT with a larger radius, e.g., (15,0) (Movie 5) or (14,12) (Movie 6), we find that the edge of the BP ribbon near the CNT is first attracted onto the CNT. As well, during the winding process, the free edge of the BP also moves close to the CNT (at 20 ps in Fig. 5a, b). However, as the overlap between the BP ribbon and CNT increases, the middle part of the BP ribbon becomes a curve that is opposite to the curving direction of the CNT (at 20 ps in Fig. 5), and simultaneously, the free edge of the BP keeps moving to the CNT due to the inertia effect. Hence, a critical curvature of the middle part of the BP ribbon develops and is followed by the breakage of P–P bonds on the red layer (at 26–27 ps in Fig. 5).
If the length of the BP ribbon is small, e.g., $L_p = 6.49\, \text{nm}$, the free part of the BP ribbon will continue to be attracted onto the CNT after the breakage. The reason is that the inertia of the free part is low, i.e., the number of phosphorus atoms is low, and the difference in relative velocity between the two parts of the broken BP ribbon is not great. Hence, the free part is under strong attraction from the CNT as the rest of the BP ribbon rotates together with the CNT. If the length of the BP ribbon is greater, e.g., $L_p = 7.81\, \text{nm}$, the BP ribbon will break into two parts due to the higher inertia effect, the free part of the BP ribbon will move closer to CNT, and it also will finally be attracted to the CNT. However, the two parts of the BP ribbon cannot be united into a perfect BP tube (at 100 ps in Fig. 5a, b). Besides the effect of the length of the BP ribbon on the winding process, the chirality of the CNT also influences the winding results. From Fig. 5, we also find that the BP ribbon is attracted symmetrically by a zigzag CNT, causing the motion of the atoms at the free edge of the BP ribbon to have the same velocity as the CNT. Hence, the broken P—P bonds are almost on the same column (along the tube axis or $z$-direction) and the free part of the BP ribbon can still be attracted almost symmetrically by the CNT (at 40 ps in Fig. 5a). If the CNT is a chiral one, e.g., (14,12), it provides asymmetric attraction to the BP ribbon. The free part of the BP ribbon rotates while moving close to the chiral CNT. The rotation of the free part of the BP ribbon is caused by the helical potential barrier on the surface of the CNT (at 100 ps in Fig. 5b).

From Fig. 4, we also know that a curve of the $\Delta PE$ of the system with a lower rate of decrease implies that the free part of the BP ribbon is not attracted onto the CNT. Curves with a higher rate of decrease illustrate that the phosphorus atoms are attracted onto the CNT even if the BP ribbon has broken into several parts.

From the inserts in Fig. 4a, b & c at 150 K, we find that the BP ribbon can be successfully wound on the (5,5) CNT only when the system is at 150 K. It is known that the vibration of atoms on the BP ribbon becomes more marked at 150 K than at temperatures below 100 K. Strong vibration of phosphorus atoms results in large deformation of the ribbon. As the P—P covalent bond is far weaker than the C—C bond, it is more...
easily broken under such large deformation. Taking this reason into consideration, to obtain a perfect BP tube from ribbon wound on a CNT, vibration of the free part of the BP should be reduced.

3.3. Can a BP ribbon longer than \( L_p \) be wound successfully?

In our previous discussion, we assumed that the length of a BP ribbon was a special value that led to the perfect nanotube from the ribbon, i.e., Eq. (2) \( L_p = N_{\text{unit}} \times L_0 = -2\pi (r + C/P + 0.5d) \). Can the ribbon still be perfectly wound onto a nanotube at a very low temperature, e.g., \( \leq 50 \) K, if we choose a BP ribbon with a length greater than \( L_p \)? Here we choose a standard BP ribbon with \( L_p = 8.24 \) nm for comparison. First, we add just one more unit cell along the armchair direction, so that the length of the new BP ribbon is \( L_p + L_0 \) and put it near three different CNTs, i.e., CNT (13,13), (14,12), and (22,0).

Fig. 6 illustrates the \( \Delta P_e \) histories of the system during the winding process of the longer BP ribbon at (a) 8 K and (b) 50 K. It can be seen that most of the curves have only one jump before 100 ps. The inserts in Fig. 6 show that the two edges of the BP ribbon along tube axis are not joined, as can be judged easily from the positions of the yellow atoms with respect to the red atoms on the other edge of the BP ribbon. In other words, the free edge (with yellow atoms) overlaps with the other edge rather than attaching to the CNT. It can be concluded that the BP ribbon with length of \( (L_p + L_0) \) can be wound onto the CNT but cannot form a perfect BP nanotube at the temperature of \( T \leq 50 \) K.

On the other hand, we also try to wind a BP ribbon of length \( L_p + 5L_0 \) or \( L_p + 10L_0 \) onto a CNT (13,13). From the snapshots in Fig. 7, it can be seen that the free edge of the BP ribbon has a different configuration in the final state. For example, when the length of the BP ribbon is \( L_p + 5L_0 \), the free edge finally overlaps with the other edge of the ribbon, and the BP ribbon becomes a nanoscroll. However, if the BP ribbon is longer, e.g., with the length of \( L_p + 10L_0 \), the free edge of the BP ribbon may be distant from the inner CNT. The reason is that the blue phosphorus atoms near the broken \( P-P \) bonds attract one another and the atoms lack sufficient kinetic energy to change the configuration further. From these results, we conclude that a BP ribbon with a length longer than \( L_p \) may be wound, but cannot form a perfect BP nanotube even at very low temperature.

4. Conclusions

Using a molecular dynamics simulation approach, we investigated the possibility of forming a nanotube from BP nanoribbon via winding on a CNT. After a series of discussions of the variations of configuration of a BP ribbon in the winding process at very low temperature, the following six conclusions can be drawn. First, during the winding process, the potential energy of the system decreases due to both vdW interaction among components in the system and released energy in the generation of new \( P-P \) bonds on the BP ribbon. A greater decrease of potential energy leads to a more stable configuration of the system. Second, if the two edges of a BP ribbon along the tube axis do not join soon after the BP ribbon is attracted onto the inner tube, the history curve of the \( \Delta P_e \) of the system contains at least two sharp decreases. The first jump is attributed to the interaction between the BP ribbon and the inner CNT. The second jump could be caused by the generation of new \( P-P \) bonds. If the jump occurs after formation of a perfect BP tube, it is mainly due to the BP tube being attracted by an outer CNT. Third, a BP nanotube can be formed on the condition that the BP ribbon has perfect length, i.e., \( L_p = N_{\text{unit}} \times L_0 = -2\pi (r + C/P + 0.5d) \), and the temperature is very low, e.g., \( \leq 50 \) K. Fourth, at 8 K, the formation of a perfect nanotube from the same BP ribbon (with perfect length) on a chiral inner CNT takes longer because extra time is required for the generation of new \( P-P \) bonds after the BP ribbon has been compactly attracted by the CNT. Fifth, at higher temperatures, e.g., \( \geq 50 \) K, even with perfect length, it is difficult to perfectly wind a BP ribbon on a CNT with higher radius. That task becomes possible if the vibration of the free part of the BP ribbon is reduced. Final, if the length of a BP ribbon is greater than the perfect length, the ribbon cannot form a perfect BP nanotube even at very low temperature.

In the present study, defect-free CNTs are considered. If there is an unsaturated carbon atom in the CNT existing nearby the BP nanotube, chemical reaction between the unsaturated carbon atom and a phosphorus atom on the BP nanotube may happen, which will lead to failure of fabrication of an ideal BP nanotube. Fortunately, an ideal CNT can be easily obtained, and the present simulations are reasonable for the guidance of future fabrication of BP nanotubes.

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