Systematic search for low-enthalpy sp^3 carbon allotropes using evolutionary metadynamics

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We present a systematic search for low-energy metastable superhard carbon allotropes by using the recently developed evolutionary metadynamics technique. It is known that cold compression of graphite produces an allotrope at 15–20 GPa. Here we look for all low-enthalpy structures accessible from graphite. Starting from 2H-or 3R-graphite and applying a pressure of 20 GPa, a large variety of intermediate sp^3 carbon allotropes were observed in evolutionary metadynamics simulation. Our calculation not only found all the previous proposed candidates for "superhard graphite," but also predicted two allotropes (X-carbon and Y-carbon) showing unusual types of 5+7 and 4+8 topologies. These superhard carbon allotropes can be classified into five families based on 6 (diamond/lonsdaleite), 5+7 (M- and W-carbon), 5+7 (X-carbon), 4+8 (bct-C₄), and 4+8 (Y-carbon) topologies. This study shows that evolutionary metadynamics is a powerful approach both to find the global minima and systematically search for low-energy metastable phases reachable from given starting materials.

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Introduction. Carbon can adopt a wide range of structures, from superhard/superdense insulating [diamond, lonsdaleite, hypothetical phases hP3, tI12, and tP12 (Ref. 1)] to ultrasoft semimetallic (graphite, fullerenes) and even superconducting (doped diamond² and alkali-doped fullerenes³). The quest for carbon materials with desired properties is of great interest in both fundamental science and advanced technology. One important direction in carbon research is the discovery of carbon allotropes with advanced mechanical and electronic properties.

It is well known that graphite transforms to the thermodynamically stable diamond at high pressures (>12 GPa) and high temperatures (1900–2500 K).⁴ On the contrary, several experiments reported that cold compression of graphite produces a metastable superhard and transparent phase, clearly different from diamond or lonsdaleite, but the exact crystal structure could not be determined.^{5–9} The difficulty to experimentally resolve the crystal structure has stimulated theoretical efforts. 10-20 Several structural models were found using different techniques. The physical properties of these models [M-carbon, 10,11 W-carbon, 12 oC 16 (also called Zcarbon), $\overline{^{13-15}}$ R/P carbon, $\overline{^{16}}$ and bct-C₄ (Refs. 17–19)] have been intensely studied. Simulated x-ray diffraction patterns and band gaps of these models are mostly in good agreement with experimental data, making it even harder to decide which one is the metastable product observed in experiments. On the other hand, it is not guaranteed that there is not even a better solution for this experimental puzzle. Furthermore, it is likely that different metastable phases will be obtained by roomtemperature compression of different polytypes of graphite, or under various nonhydrostatic conditions. This motivates us to do a systematic search for low-energy metastable carbon allotropes.

So far, there are several methods to find the ground state structures of unknown materials. However, none of them are designed to search for metastable states. Our recently proposed evolutionary metadynamics method²¹ can focus on that task. Starting from a reasonable initial crystal structure,

with this technique one can produce efficiently both the ground state and metastable states accessible from that initial structure. In this Rapid Communication, we applied this technique to systematically search for metastable carbon allotropes accessible from graphite. Starting the calculation at 20 GPa from two polytypes of graphite (2H and 3R), we easily found the diamond structure (ground state) and a number of low-energy metastable structures with sp^3 hybridization which could possibly explain "superhard graphite."

Methodology. The idea of metadynamics is to introduce a history-dependent potential term, which fills the minima in the free energy surface so that the system could cross the energy barriers and undergo phase transitions.²² This technique is usually applied as an extension of molecular dynamics (MD) simulation technique.²³ Since it relies on MD simulation to equilibrate the system, it often leads to trapping in metastable states and amorphization rather than a transition to a stable crystal structure. We recently proposed a hybrid method, basically a metadynamics-like method driven not by local MD sampling, but by efficient global optimization moves.^{10,24}

In this approach, 21 we start from one known initial structure at a given external pressure P. Following Martonak $et\ al.$, we used the cell vectors matrix h_{ij} (also representable as a six-dimensional vector h) as a collective variable to distinguish the change of state of the system. 23 For a given system with volume V under external pressure P, the derivative of the free energy G with respect to h is

$$-\frac{\partial G}{\partial h_{ij}} = V[h^{-1}(P-p)]_{ji}.$$
 (1)

At each generation (or metastep), many structures are produced and relaxed at fixed h, and we select the lowest energy structure and compute it internal tensor p. The technique used here generates many structures at each metastep, while in traditional metadynamics, 23 only one structure is produced at each metastep. The cell shape is then updated with a stepping

parameter δh ,

$$h_{im}(t+1) = h_{im}(t) + \frac{\delta h}{|f|V^{1/3}} S_{ijkl} f_{kl} h_{jm}(t), \qquad (2)$$

where S is the elastic compliance tensor corresponding to an elastically isotropic medium with Poisson ratio 0.26, which corresponds to the border between brittle and ductile materials²⁵ and is a good average value to describe both metals and insulators. The driving force $f = -\frac{\partial G}{\partial h}$ in Eq. (2) comes from a history-dependent Gibbs potential G^t where a Gaussian has been added to G(h) at every point h(t') already visited in order to discourage it from being visited again,

$$G(t) = G^{h} + \sum W e^{-\frac{|h - h(t')|^{2}}{2\delta h^{2}}},$$
 (3)

where W is the Gaussian height. Then we compute the vibrational modes for the selected structure according to the dynamical matrix constructed from bond hardness coefficients

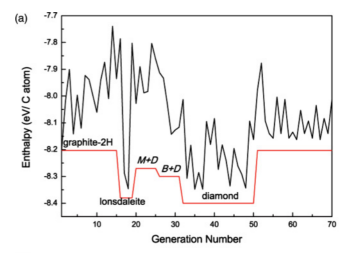
$$D_{\alpha\beta}(a,b) = \sum_{m} \frac{\partial^{2}}{\partial \alpha_{a}^{0} \partial \beta_{b}^{m}} \left(\frac{1}{2} \sum_{i,j,l,n} H_{i,j}^{l,n} \left(r_{i,j}^{l,n} - r_{0i,j}^{l,n} \right)^{2} \right)$$
(4)

where coefficients α , β denote coordinates (x,y,z); coefficients a,b,i,j describe the atoms in the unit cell; coefficients l,m,n describe the unit cell number; and $r_{i,j}^{l,n}$ is the distance between atom i in the unit cell l and atom j in the unit cell n, while $r_{0i,j}^{l,n}$ is the corresponding bond distance. $H_{i,j}^{l,n}$ is bond hardness coefficient computed from bond distances, covalent radii and electronegativities of the atoms. Above that the dynamical matrix corresponds to zero wavevector (extension to non-zero wavevectors is straightforward) and unit masses.

The simulated vibrational modes are used to produce the next generation (typically 20–40 softmutated structures). To perform softmutation, 21,24 we move the atoms along the eigenvector of the softest calculated mode. One structure can be softmutated many times using different nondegenerate modes and displacements. The magnitude of the displacement (d_{max}) along the mode eigenvector is an input parameter: With relatively small d_{max} and displacements represented by a random linear mixture of all mode eigenvectors, the method becomes similar to MD metadynamics in crossing energy barriers and equilibrating the system. With large d_{max} along the softest mode eigenvectors, we obtain the softmutation operator, 24 capable of efficiently finding the global energy minimum.

The next generation of softmutated structures is produced and relaxed in the updated cell. Repeated for a number of generations, this computational scheme leads to a series of structural transitions and is stopped when the maximum number of generations is reached. For technical details and discussion of evolutionary metadynamics, see Ref. 21.

In this work, structure relaxations were done using density functional theory (DFT) within the generalized gradient approximation (GGA)²⁶ in the framework of the all-electron projector augmented wave (PAW)²⁷ method as implemented in the VASP^{28,29} code. We used a plane-wave kinetic energy cutoff of 550 eV for the plane-wave basis set and a Brillouin zone sampling resolution of $2\pi \times 0.08~\text{Å}^{-1}$, which showed excellent convergence of the energy differences, stress tensors and structural parameters.



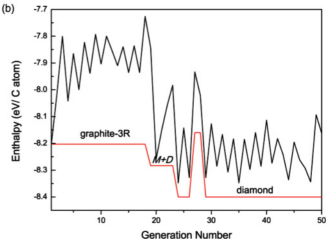


FIG. 1. (Color online) (a) Enthalpy evolution during the compression of graphite-2H at 20 GPa. (b) Enthalpy evolution during the compression of graphite-3R at 20 GPa (black line: enthalpies for best structures with constant cell matrix; red line: enthalpies for best structures after full relaxation).

Results and discussions. In a compression experiment at low temperatures (low enough to preclude transition to the stable state), the product depends on the nature of the starting materials, and on the energy landscape (in particular, energy barriers). To fully investigate the possible candidate materials, we performed simulations starting from two different graphite polytypes (graphite-2H and graphite-3R), which differ in the stacking of graphene layers.

Starting from graphite-2H. We did a preliminary test at 20 GPa starting from the graphite-2H structure, and successfully found diamond as the ground state, and M-carbon and bct-C₄ as metastable states.²¹ In the calculation we set $d_{\text{max}} = 2.5 \text{ Å}$, $W = 4000 \text{ kbar Å}^3$, and $\delta h = 0.6 \text{ Å}$.

Figure 1 shows the enthalpy evolution. Graphene layers [Fig. 2(a)] persisted until the 15th generation. Then, upon sufficient cell deformation, the layers began to buckle, and the planar structure transformed into three-dimensional (3D) networks of sp^3 -hybridized carbon atoms. Lonsdaleite with six-membered rings [Fig. 2(b)] appeared as the best structure in the 16th generation. We observed in the same generation the bct-C₄ structure with 4+8 membered rings [Fig. 2(c)],

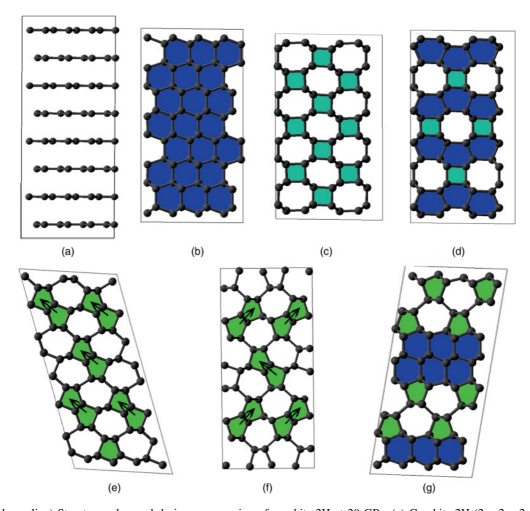


FIG. 2. (Color online) Structures observed during compression of graphite-2H at 20 GPa. (a) Graphite-2H $(2 \times 2 \times 2 \text{ supercell of the calculation model})$; (b) lonsdaleite; (c) bct- C_4 with 4+8 membered rings; (d) *Z*-carbon (belonging to B+D type); (e) *M*-carbon with 5+7 membered rings; (g) M+D type carbon. Polygons are highlighted by different colors (quadrangle: turquoise; pentagon: green; hexagon: blue).

and M- and W-carbon structures containing 5+7 membered rings [Figs. 2(e) and 2(f)] appeared shortly after. Lonsdaleite survived for a few generations until it transformed into a hybrid structure made of alternating layers of M-carbon and diamond [Fig. 2(g), we refer to it as M+D type], followed by the transition to another hybrid structure made of bct-C₄ and diamond [Fig. 2(d), similarly, we refer to it as B+D type]. Diamond was dominant in the following generations. At the 51st generation, the system reverted to graphite.

The power of the evolutionary metadynamics method lies in that it is highly suitable for harvesting low-energy metastable structures in addition to the ground state. Those previously proposed candidate structures for the product of cold compression of graphite, bct- C_4 , M-carbon, and W-carbon are all easily recovered in a single simulation. More interestingly, we also observed many low-energy structures based on 5+7 or 4+8 topology. The B+D type structure [Fig. 2(d)] observed in the simulation is actually the oC16 structure (sometimes called Z-carbon), recently suggested as a candidate for superhard graphite. Since oC16 inherits layers of bct- C_4 and diamond, there is no surprise that its thermodynamic properties are intermediate between these

two structures. The 5+7 class of structures shows a larger diversity. In some of these structures, five-membered rings form pairs, while in others these they are single. The difference of the five-membered ring pairs' orientation leads to two allotropes: M-carbon [Fig. 2(e)] and W-carbon [Fig. 2(f)]. Some structures can be thought of as combinations of layers of the M-carbon and diamond structures [M+D] carbon, as shown in Fig. 2(g)].

Starting from graphite-3R. Starting the calculation at 20 GPa from another polytype, graphite-3R, which contains three layers per lattice period, we again easily found the diamond structure and a number of low-energy metastable structures with sp^3 hybridization. Figure 3 shows the results. Since the initial model has three graphene layers, it could form a large variety of M+D and B+D structures based on 4+6+8 or 5+6+7 topologies. For instance, we observed a B+D structure containing $2\times(4+8)$ layers and 1×6 layer [Fig. 3(c)], or $1\times(4+8)$ layers and 1×6 layer [Figs. 3(e) and M+D structure containing $2\times(5+7)$ layers and a 1×6 layer [Figs. 3(e) and 3(f)]. Most strikingly, we also observed another structure with a 5+7 topology, which is shown in Fig. 3(g). The projections of pentagons and heptagons along

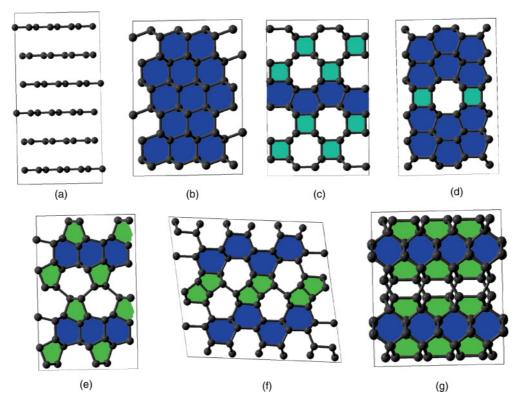


FIG. 3. (Color online) Structures observed during compression of graphite-3R at 20 GPa. (a) Graphite-3R $(2 \times 2 \times 2 \text{ supercell of the calculation model})$; (b) diamond; (c), (d) B+D type structures; (e), (f) M+D type structures; (g) X+D type structures. Polygons are highlighted by different colors (squares: turquoise; pentagons: green; hexagons: blue).

the c axis could not be separated as in M-carbon, but overlap each other. We extracted the 5+7 part from the complex structure, and obtained a different configuration with pure 5 + 7 topology. This crystal structure (which we call X-carbon, and the hybrid structure from X-carbon and diamond is referred to as X + D type) is shown in Fig. 4. It is a monoclinic structure with C2/c symmetry, and contains 32 atoms in the conventional cell. We also found an unexpected 4+8topology in an allotrope that we call Y-carbon with unique 4+8 ring topology from another separate metadynamics run. It is an orthorhombic structure with *Cmca* symmetry, containing 16 atoms in the conventional cell. The simulated x-ray diffraction patterns of all these structures are in good agreement with experimental data (as shown in Fig. 5), suggesting that both X-carbon and Y-carbon could explain the experiments on cold compressed graphite. Although all these structures show a satisfactory agreement with experimental x-ray data, our recent transition path sampling calculations³⁰ suggest M-carbon to be kinetically the likeliest product of cold compression of graphite-2H. Using other polytypes of graphite, or different conditions (nonhydrostatic or dynamical compression), one might produce alternative allotropes found here. Synthesis of these allotropes would be desirable in view of their physical properties.

Properties. From evolutionary metadynamics simulations, five families of sp^3 -hybridized structures made by stacking corrugated graphene layers and having competitive enthalpies were discovered: 6 (diamond and lonsdaleite), two classes of 5+7 topologies (one includes M- and W-carbon and the other

X-carbon), and two classes of 4+8 (one is bct-C₄, and the other Y-carbon). The enthalpies of different carbon phases as a function of pressure are presented in Fig. 6. Apart from the prototypes, we also included hybrid structures (lowest enthalpy B+D, M+D, and X+D carbon—see the crystallographic

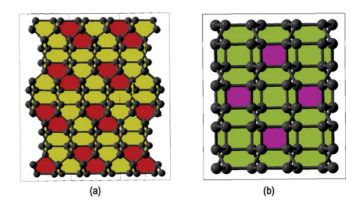


FIG. 4. (Color online) (a) Allotropes with 5+7 topology, X-carbon, space group C2/c, a=5.559 Å, b=7.960 Å, c=4.752 Å, $\beta=114.65^{\circ}$. This structure has five nonequivalent Wyckoff positions: C1 (0.250, 0.083, 0.949), C2 (0.489, 0.809, 0.982), C3 (0.000, 0.200, 0.250), C4 (0.247, 0.913, 0.801), C5 (0.000, 0.816, 0.250). (b) Allotrope with 4+8 topology, Y-carbon, space group Cmca, a=4.364 Å, b=5.057 Å, c=4.374 Å. This structure has one Wyckoff position: C (0.681, 0.635, 0.410). Polygons are highlighted in different colors to show the 5+7 and 4+8 topology.

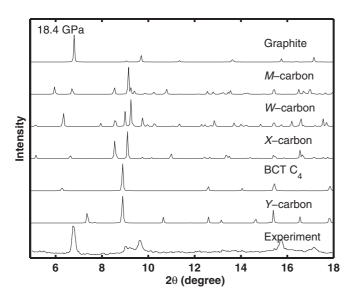


FIG. 5. Comparison of simulated x-ray diffraction patterns of proposed models and graphite with experiment (Ref. 9). The patterns were simulated using Accelrys Materials Studio 4.2 software with an x-ray wavelength of 0.3329 Å.

data in the Supplemental Material³¹). At elevated pressures, all these allotropes become more stable than graphite. For the prototypes, M- and W-carbon are energetically more favorable than bct-C₄ and X-carbon. Lower enthalpies are obtained by combining layers of these structures with layers of diamond. For the models under consideration (up to four graphene layers), B + D [1 × (4 + 8) + 2 × 6 layers, Fig. 3(d)] tends to have the lowest enthalpy, while M + D [2 × (5 + 7) + 2 × 6 layers, Fig. 2(h)] is quite competitive (only 8 meV/atom higher than B + D). X + D [Fig. 3(h)] is 50 meV/atom higher than B + D, indicating that X-carbon has the poorest possibility to interface with diamond.

We also computed the mechanical and optical properties (see the Supplemental Material³¹). Similar to previous theoretical investigations, ^{11–16,18,19} all of these candidate allotropes exhibit high hardnesses³² and bulk moduli, which

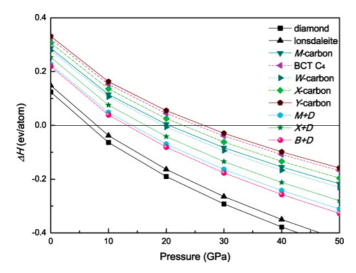


FIG. 6. (Color online) Enthalpies of various carbon structures relative to graphite-2H.

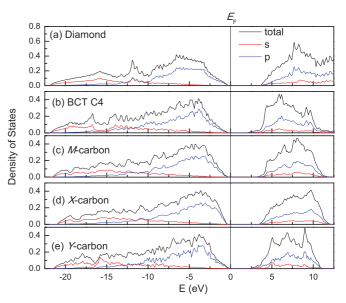


FIG. 7. (Color online) Total and partial densities of states of different carbon allotropes.

are comparable with those of diamond. Figure 7 shows the calculated total and partial electronic densities of states in both systems. It can be clearly seen that 2p states exhibit a larger overlap with 2s states in diamond, which makes diamond the most stable allotrope among sp^3 forms of carbon. The magnitude of overlap determines the order of stability: M-carbon > X-carbon > bct- C_4 > Y-carbon. The DFT band gaps of M-carbon, X-carbon, bct- C_4 , and Y-carbon are 3.6, 3.8, 2.7, and 2.9 eV, and we should bear in mind that DFT always underestimate the band gaps—so the real gaps are larger, and all of these sp^3 allotropes should be transparent, colorless insulators.

Conclusions. In summary, we performed a systematic search for metastable allotropes of carbon that can be synthesized by cold compression of graphite by using the recently developed evolutionary metadynamics technique.²¹ Starting from 2H- or 3R-graphite, at 20 GPa we easily found diamond as the ground state and observed a large variety of low-energy metastable sp^3 carbon allotropes accessible from graphite. Apart from diamond, lonsdaleite, and their polytypes, we summarize four other families of low-enthalpy carbon allotropes which can be obtained by cold compression of graphite: (i) 5+7 topology (M- and W-carbon), (ii) 5+7 topology (X-carbon), (iii) 4+8 topology (bct-C₄), and (iv) 4+8topology (Y-carbon). All of these structures are consistent with experimental data on "superhard graphite," and are predicted to have excellent mechanical properties, but transition path sampling calculations³⁰ unequivocally show M-carbon to be the likeliest product of cold compression of graphite. Yet, the allotropes predicted here could be synthesized with a different experimental protocol and starting from different materials (graphite-3R, turbostratic graphites, fullerenes, or nanotubes, etc.). We find it particularly encouraging that all the previously proposed structures and two topologically interesting and previously not described ones (X- and Y-carbon) were found in a systematic way, using just one calculation per starting material (graphite-2H or graphite-3R). Our work shows that evolutionary metadynamics is a powerful method for efficiently finding not only stable but also low-energy metastable structures.

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