Bottom-Up Synthesis of Heteroatom-Doped Chiral Graphene Nanoribbons

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ABSTRACT: Bottom-up synthesis of graphene nanoribbons (GNRs) has significantly advanced during the past decade, providing various GNR structures with tunable properties. The synthesis of chiral GNRs, however, has been underexplored, and only limited to (3,1)-GNRs. We report herein the surface-assisted synthesis of the first heteroatom-doped chiral (4,1)-GNRs from a rationally designed precursor 6,16-dibromo-9,10,19,20-tetraoxa-9a,19a-diboratetrabenzo[afjo]perylene. The structure of the chiral GNRs has been verified by scanning tunneling microscopy, non-contact atomic force microscopy, and Raman spectroscopy in combination with theoretical modeling. Due to the presence of oxygen-boron-oxygen (OBO) segments on the edges, lateral self-assembly of the resulting GNRs has been observed, realizing well-aligned GNR arrays with different modes of homochiral and heterochiral inter-ribbon assemblies.

Graphene nanoribbons (GNRs), nanometer-wide strips of graphene, have been recognized as the next-generation of semiconductors compared with the non-bandgap graphene. The bottom-up synthesis of GNRs has proven as a powerful tool to fabricate atomically precise graphene nanostructures by synthetic design. Especially, the on-surface synthesis under ultrahigh vacuum (UHV) offers new opportunities to obtain well-defined GNRs with different edges and widths. So far, armchair GNRs (AGNRs) have been extensively investigated, displaying width-dependent bandgaps while the long-pursued zigzag GNRs (ZGNRs) possessing localized edge states have also been achieved in 2016 (Figure 1a,b). On the other hand, chiral GNRs (CGNRs) with a combination of armchair and zigzag edges have been largely underexplored, although they represent an important member of the GNR family (Figure 1c). The edge configuration of CGNRs is described by a translation vector \( (n,m) \) of the graphene lattice, and different chiral indices result in distinct electronic properties. However, only the synthesis of (3,1)-CGNRs has been achieved from 9,9’-bianthracene-based molecular precursors and the investigations of bottom-up synthesized CGNRs have been limited to this structure. Further precise variations of the chiral edges have remained elusive.

Furthermore, incorporating heteroatom dopants into GNRs via the bottom-up approach can finely modulate their band structures and electronic properties, as well as self-assembly behavior. Major achievements on heteroatom-doped GNRs in recent years include chevron-type GNRs with nitrogen, sulfur, or oxygen atoms on the edges, and AGNRs with boron dopants or boron-nitrogen co-dopants inside the backbone as well as sulfur or nitrogen atoms on the edges. Nevertheless, heteroatom doping in other types of GNRs, e.g. ZGNRs and CGNRs, has never been explored. Herein, we report the bottom-up synthesis and characterization of the first heteroatom-doped (4,1)-CGNRs using the rationally designed molecular precursor 6,16-dibromo-9,10,19,20-tetraoxa-9a,19a-diboratetrabenzo[afjo]perylene.

Figure 1. Representation of the three classes of graphene nanoribbons (GNRs). (a) Armchair GNRs. (b) Zigzag GNRs. (c) Chiral GNRs. The edge geometry of chiral GNRs is characterized by the translation vector \( C_h \), defined as \( C_h = n a_1 + m a_2 = (n,m) \), where \( a_1 \) and \( a_2 \) are the unit vectors of the graphene lattice. An exemplary case of chiral (4,1)-GNRs is illustrated here.
formed into iodo groups by treatment with iodine monochloride couplings, forming 1D polymer (ICl) to give 1,4-diiodo-2,5-bis(2'-methoxyphenyl)benzene (compound 7 in Scheme 1). The monomer features stable zigzag edges substituted with OBO units, and the resulting GNRs represent the first case of OBO-doped GNRs.

The synthetic route to obtain OBO-doped (4,1)-CGNR 1 is described in Scheme 1. 1,4-Diiodobenzene (2) was first deprotonated by lithium disopropylamide (LDA) and then reacted with trimethylsilyl chloride (TMSCl) to provide 1,4-diiodo-2,5-bis(trimethylsilyl)benzene (3) in 76% yield. Suzuki coupling of 3 with (2-methoxyphenyl)boronic acid afforded 1,4-bis(5'-bromo-2'-methoxyphenyl)benzene (4) in 90% yield. Subsequently, the TMS groups on 4 were efficiently transformed into iodo groups by treatment with iodine monochloride (ICl) to give 1,4-diiodo-2,5-bis(2'-methoxyphenyl)benzene (5), which was subjected to another Suzuki coupling with (5-bromo-2-methoxyphenyl)boronic acid to produce 1,4-bis(5'-bromo-2'-methoxyphenyl)benzene (6) in 62% yield. Then a tandem demethylation-borylation reaction was applied to 6 with BBBr in o-dichlorobenzene (ODCB) to give monomer 7 in 42% yield. Monomer 7 was characterized by 1H and 13C NMR spectroscopies and high-resolution mass spectrometry.

Scheme 1. Synthetic Route to Chiral (4,1)-GNR 1.

To fabricate OBO-doped (4,1)-CGNR 1, monomer 7 was first sublimed onto an Au(111) substrate held at room temperature under UHV conditions. Constant-current scanning tunneling microscopy (STM) images of the resulting molecular layer reveal intact monomers self-assembled into linear chains (see Figure S1). A closer look allows one to discern R and S enantiomers within the same chain, stabilized mainly by lateral intermolecular O···H interactions with a projected length of 2.7 ± 0.2 Å, forming assembled structures along the perpendicular dimension, were also observed (see Figure S1, where lateral intermolecular lengths were obtained from properly scaled atomistic models of the STM images). Annealing of the substrate at 200 °C induced aryl–aryl couplings, forming 1D polymer 8 after dehalogenation of 7. Importantly, single chirality of monomers, assembled side by side, was preserved within each polymeric chain (Figure 2a). This fact is attributed to steric repulsion between monomers with opposite chirality inhibiting the aryl-aryl coupling between them. A closer view of a single polymeric chain reveals a periodic appearance of bright protrusions (apparent height of 2.4 Å), which result from the highly nonplanar geometry adopted by the polymer due to the steric hindrance between the H atoms in neighboring units (Figure 2b). The periodic distance of these bright protrusions in the STM image is 11.2 Å, in good agreement with the structural model (11.0 Å, Figure 2c), demonstrating the successful polymerization.

Figure 2. STM images of monomer 7 after annealing at 200 °C on Au(111), producing debromination and polymerization. (a) Large-scale STM image of the resulting polymer 8. (b) Magnified view of the single polymeric chain shown in panel a evidencing the presence of periodic bright protrusions. Scanning conditions: $V_b = −1.5$ V, $I = 20$ pA. (c) Structural model of polymer 8. The benzene rings in red correspond to the bright protrusions in panels a and b. Further annealing of the substrate at 400 °C induced minor modifications of the 1D polymeric chains, which were attributed to the onset of the cyclodehydrogenation reaction (see Figure S2). Finally, annealing the sample at 450 °C completed cyclodehydrogenation, providing the fully planarized OBO-doped (4,1)-CGNR 1 with an apparent height of 1.8 Å (Figure 3a,b) and an average length of 61 nm (see the histogram in Figure S3). To confirm the chemical structure of the resulting CGNRs, non-contact atomic force microscopy (nc-AFM) measurements using a CO-functionalized tip were performed. Figure 3c depicts the resulting constant-height frequency-shift image where the periodic aromatic carbon atoms together with the oxygen atoms are clearly unveiled, while the boron atoms appear with a darker contrast (more negative frequency shift) due to a stronger interaction with the gold substrate as previously reported by Kawai et al. in the formation of boron-doped AGNRs. It is worthwhile mentioning that the OBO-doped CGNRs are found to laterally align on the Au(111) substrate. The regular alignment of GNRs arises from inter-ribbon O···H interactions (see Figure S4 for STM images and corresponding atomistic models of homochiral and heterochiral inter-ribbon interactions) and illustrates a particular aspect of heteroatom-doped GNRs, which can be of crucial importance for future applications. To gain more insight into the lateral alignment of OBO-doped (4,1)-CGNRs 1, density functional theory (DFT) calculations have been performed on the self-assembled structures (Figure 3d,e). The interaction energy of two OBO-doped CGNRs of opposite chirality is $−0.30$ eV/unit cell with O···H interaction distances of 2.43 and 2.72 Å, in good agreement with experimental data. In the case of two OBO-doped CGNRs with the same chirality, the interaction energy is $−0.24$ eV/unit cell with O···H interaction distances of 2.28 and 3.49 Å. These computational results suggest that the interaction of ribbons of opposite chirality is stronger than that of ribbons with the same chirality, though no clear experimental preference in the interaction of ribbons with the same/different chirality has been found. In addition, we have computed the interaction energy of the structurally equivalent pristine (4,1)-CGNR to be $−0.10$ eV/unit cell, indicating that the OBO units on the edges endow the GNRs with stronger inter-ribbon interactions.
Figure 3. Formation of OBO-doped (4,1)-CGNR on Au(111) after annealing at 450 ºC. (a) STM overview of (4,1)-CGNR. $V_b = 0.1\ V$, $I = 70\ pA$. (b) High-resolution STM image of a single ribbon with the chemical structure of a segment superimposed on it. $V_b = -1.5\ V$, $I = 50\ pA$. (c) Constant-height frequency-shift nc-AFM image of the ribbon segment reported in the dashed blue rectangle of panel b acquired with a CO-functionalized tip ($z$ offset $-20\ pm$ below STM set point: $-5\ mV$, 10 pA). (d,e) DFT models of inter-ribbon interactions for heterochiral and homochiral (4,1)-CGNRs, respectively.

Next, scanning tunneling spectroscopy (STS) studies were performed on (4,1)-CGNR and its differential conductance ($dI/dV$) spectra were recorded. The conduction band edge and the valence band edge are clearly identified at 1.77 eV and −1.56 eV, respectively, corresponding to a STS-bandgap of 3.33 eV (Figure S5). This is in good agreement with the bandgap of 2.96 eV, calculated within the image charge corrected 18 GW approximation. Additionally, DFT calculations reveal that the OBO-doped (4,1)-CGNR exhibits a larger bandgap (1.50 eV) in vacuum compared with the pristine all-carbon (4,1)-CGNR (0.50 eV). This can be explained by the weak conjugation of the $p_z$ orbitals of the OBO segments with the extended $\pi$-system of the carbon backbone of the GNR. Although weak, this conjugation lowers the bandgap with respect to that of the GNR without OBO units by 0.40 eV (1.90 eV) (see Figure S6 for details).

Finally, we used Raman spectroscopy to further characterize its structure. Raman spectra were measured in ambient conditions with an excitation wavelength of 532 nm (2.33 eV) on Au (111). To assign the measured Raman lines, DFT calculations were performed (Figure 4). The experimental result agrees well with the DFT-simulated spectrum, further confirming the structure of (4,1)-CGNR and indicating that the GNR structure is preserved in ambient conditions. Interestingly, DFT calculations reveal that the typical D-mode vibrations localize on two different regions of (4,1)-CGNR, namely the armchair (A) and zigzag (Z) segments of the ribbon. Therefore, the D signal splits into two components arising from the A and Z regions (1325 cm$^{-1}$ and 1481 cm$^{-1}$, respectively). Similarly, the G band exhibits contributions from the A and Z segments, which both display longitudinal ($G_{A1}$ and $G_{Z1}$) and transversal ($G_{A1}$ and $G_{Z1}$) components (Figure 4, see Figure S7 and S8 for details).

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Notes
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