Ultrathin graphdiyne film on graphene through solution-phase van der Waals epitaxy

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Graphdiyne (GDY) is an ordered two-dimensional (2D) carbon allotrope comprising sp- and sp2-hybridized carbon atoms with high degrees of π-conjugation, which features a natural band gap and superior electric properties. However, the synthesis of one- or few-layer GDY remains challenging because of the free rotation around alkyne-aryl single bonds and the lack of thickness control. We report the facile synthesis of an ultrathin single-crystalline GDY film on graphene through a solution-phase van der Waals epitaxial strategy. The weak molecule-substrate interaction at the heterostructure drastically relaxes the large lattice mismatch between GDY and graphene. It allows the fast in-plane coupling of admolecules and slow out-of-plane growth toward the formation of an incommensurately stacked heterostructure, which is composed of single-layer graphene and few-layer ABC-stacked GDY, as directly observed by electron microscopy and identified from Raman fingerprints. This study provides a general route not only to the bottom-up synthesis of intriguing 2D acetylenic carbon allotropes but also to the device fabrication for the direct measurement of their intrinsic electrical, mechanical, and thermal properties.

INTRODUCTION

The exploitation of low-dimensional carbon allotropes has long been a research topic of particular interest because of their fascinating properties and outstanding performance in diverse fields. The discovery of fullerenes (zero-dimensional (0D)) (1), carbon nanotubes (1D) (2), and graphene (2D) (3) as typical low-dimensional carbon allotropes with similar chemical scaffolds containing sp2-hybridized carbon has reached great success because of their unprecedented electrical, mechanical, and thermal properties. In 1997, Haley et al. (4) proposed a unique 2D carbon allotrope with a totally different chemical scaffold combining both sp- and sp2-hybridized carbon, namely, graphdiyne (GDY). The presence of acetylenic linkages endows GDY with distinct physical and chemical properties from sp2-hybridized carbon allotropes (5). For example, GDY exhibits a predicted natural moderate band gap of 0.44 to 1.47 eV while maintaining a high carrier mobility of 104 to 105 cm2 V−1 s−1 at room temperature (6–13). In addition, the highly π-conjugated structure and the ordered porous topology enable its great potential in applications such as catalysis (14), gas separation (12), and energy-related fields (15, 16).

Encouraged by these intriguing properties of GDY, tremendous efforts have been made to experimentally synthesize such 2D acetylenic carbon allotropes. For example, dry chemistry approaches have been used, but only noncovalent assemblies or oligomers are derived on the metal surface under ultrahigh vacuum (17, 18). In contrast, the wet chemistry approach achieved the scalable synthesis of GDY and further improved the quality of GDY. Li et al. (19) conducted a pioneering work in preparing a uniform GDY film with ~1-μm thickness via Glaser coupling reaction in solution, where copper foil was used as both the catalyst and the substrate. Following this work, several studies have been carried out to further control the growth of GDY. Zhou et al. (20) reported a feasible synthetic route of GDY nanowalls by using a modified Glaser-Hay coupling. Different morphologies of GDY on various substrates were also prepared by using anodic aluminum oxide template (21) and copper envelop strategy (22). Multilayer GDY nanosheets were prepared via a successive alkyne-alkyne homocoupling reaction at the interface between monomer/catalyst solutions (23). More recently, we proposed a template-assisted approach for synthesis of thin-film GDY (3 to 20 nm) by using Hiyama coupling reaction at 65° to 70°C (24). Despite the promising advances in pursuing high-quality GDY, the synthesis of a single-crystalline GDY film thin enough to retain its “2D character” (that is, typically one- to few-layer) is, however, very challenging. The major obstacle lies in the fact that (i) the free rotation around alkyne-aryl single bonds in monomers usually leads to a highly branched or cross-linked framework with less ordering rather than the desired crystalline planar framework of GDY during polymerization; (ii) conventional epitaxial growth requires severe lattice matching conditions between the epilayer and the substrate with dangling bonds due to their strong interaction; and (iii) the step-down diffusion of monomers adsorbed on the epilayers is hindered by the Ehrlich-Schwoebel (ES) barrier (25), which thus leads to the accumulation and nucleation of admolecules on the epilayer and results in a large thickness via layer-by-layer out-of-plane growth.

Van der Waals (vdW) epitaxy, on the other side, greatly alleviates the lattice matching conditions due to the weak vdW-type interaction between the epilayer and the substrate with saturated surface bonds. The weak binding between the epilayer and admolecules largely promotes the in-plane versus out-of-plane growth of the epilayer by effectively diminishing the ES barrier. In addition, by using substrates with a 2D structure like graphene, its atomically flat surface favors the in-plane coupling rather than out-of-plane cross-linking of admolecules that usually adopt a flat-lying adsorption geometry (26). Here, we developed a facile solution-phase vdW epitaxial strategy to confine the GDY growth on graphene by Eglinton reaction using hexaethynylbenzene...
(HEB) as monomers, which can occur at room temperature even with a low monomer concentration. Benefiting from the higher binding energy of HEB monomers on graphene than GDY (1.47 eV versus 0.896 eV, as calculated), the monomers also thermodynamically prefer to adsorb on graphene toward the in-plane Eglinton coupling reaction, especially at low coverage. Together, a much faster in-plane than out-of-plane growth of GDY is expected on graphene through vdW epitaxy. Using this method, we achieved an ultrathin single-crystalline GDY film on graphene, which exhibits excellent electric properties for diverse applications, such as gas sensing. The proposed solution-phase vdW epitaxial strategy here may advance the design of novel 2D acetylenic carbon allotropes with unique structural and physical properties.

RESULTS AND DISCUSSION
Figure 1 (A and B) shows a schematic illustration of the synthetic process of an ultrathin single-crystalline GDY film on graphene. First, single-layered graphene was transferred onto a SiO2/Si substrate as the substrate for the synthesis of GDY. The optical image and Raman spectra are shown in fig. S1. The high-resolution transmission electron microscopy (HRTEM) images show typical contrast for monolayer graphene (fig. S2). The as-prepared graphene substrate was immersed in a CH2Cl2 solution of HEB monomers, where HEB molecules are adsorbed to graphene. Then, a pyridine solution of copper acetate was added dropwise, serving as the catalyst of acetylenic coupling reaction for the synthesis of GDY. To further control the kinetics of the coupling reaction and avoid the side reactions of HEB oxidation, we adopted a low concentration of both the monomer (0.04 mM) and the catalyst (0.7 mM) for the growth of GDY. After reaction under an argon atmosphere at room temperature for 24 hours, the ultrathin GDY film can be synthesized on the graphene surface. Figure 1 (C and D) shows typical optical microscopy (OM) and scanning electron microscopy (SEM) images of GDY/graphene films. The GDY film on the graphene surface is continuous with a lateral dimension of 50 μm. Atomic force microscopy (AFM) analysis (Fig. 1E) exhibits an ultraflat surface with a thickness of only 1.74 nm, inclusive of monolayer graphene.

In addition, GDY was prepared on the exfoliated graphene with different domains for further thickness evaluation. We measured the thickness of these domains before and after growth of GDY, respectively. The typical AFM image is shown in fig. S3 (A and B). The statistical thickness data of as-grown GDY were extracted from 66 independent domains. The data exhibited a narrow bimodal distribution centered at 1.05 ± 0.02 nm (major) and 2.04 ± 0.04 nm (minor), respectively (fig. S3C). Accordingly, most of the as-grown GDY film was determined to have a trilayer thickness.

The as-grown film was separated from the SiO2/Si substrate and transferred onto a copper grid by a poly(methyl methacrylate) (PMMA)–assisted transfer method (fig. S4), which was then inspected by TEM (Fig. 2A). Figure S5 shows that the as-grown film is mainly composed of carbon. Apart from the graphene layer, a second layer with...
dark contrast was observed by TEM. The dual-layer structure was more clearly visualized from the edges, as marked in Fig. 2A. Electron diffraction pattern of such transferred film exhibits two sets of overlapping single-crystal diffractions spots, one of which has smaller d spacings (that is, 0.214 and 0.124 nm) and can be easily indexed by the (1/2100) and (11/2200) reflections of graphene, respectively (Fig. 2B). The thickness of graphene was further confirmed to be single-layer due to the more intensive (1/2100) reflections than (11/2200) reflections. Another set of hexagonally arranged diffraction spots with a larger d spacing of 0.470 nm was observed, which can be indexed by the (11/2200) reflections of GDY ([23](23)). Apparently, the dual-layer heterostructure is composed of incommensurately stacked single-crystalline graphene and GDY, which follows the vdW epitaxy growth mode and allows a nearly complete relaxation of large lattice mismatch at the heterointerface. As-grown GDY is oriented with a ~14° interlayer rotation with respect to the underlying graphene, which has also been theoretically predicted to be an energetically preferred vdW heterostructure, as shown in Fig. 2 (I and J). Notably, the (1100) reflections of as-grown GDY are absent, which contradicts with either monolayer, AA-stacked, or AB-stacked GDY structural models with P6/mmm, P6/mmm, or P63/mmc symmetries, respectively (fig. S6). Among all stacking modes of GDY structures with high symmetry, the observed systematic extinction corresponds to rhombohedral (R)–centering and the ABC stacking mode with an R-3m symmetry, as similarly observed in multilayer GDY ([23](23), [27](27)). This is verified by the fact that theoretically predicted ABC-stacked GDY has slightly lower energy than other stacking modes (fig. S7). We were able to directly observe the as-grown GDY/graphene vdW heterostructure at high resolution by using aberration-corrected and monochromated TEM under low voltage (80 kV) and low dose while maintaining its structural integrity (fig. S8). Despite the fact that the amorphous (PMMA) layer used for transferring the GDY/graphene film cannot be completely removed, hexagonal honeycomb lattice contrast with a nearest neighbor distance of ~0.53 nm for bright dots is unambiguously observed, which could be attributed to the highly crystalline ABC-stacked GDY sheets, as schematically shown in Fig. 2K. This point is further confirmed by the good agreement between the experimental HRTEM image of as-grown GDY and the simulated one using an ABC-stacked GDY model, as shown in Fig. 2 (E and F). The GDY/graphene vdW heterostructure can be more clearly identified from the fast Fourier transform (FFT) of the HRTEM image in Fig. 2C, where two sets of reflections belonging to randomly stacked graphene bilayers, as well as another set of reflections for the GDY film, are discriminated from each other. The FFT contains structural information of GDY transferred up to 2.6 Å, with which the contrast transfer function (CTF)–corrected and lattice-averaged (symmetry-imposed) images resolve two neighboring motifs composed of superimposed acetylenic groups and match the simulated potential map of ABC-stacked GDY very well in Fig. 2 (G and H) ([28](28)). Systematic extinction arising from the R-centering of the ABC-stacked
GDY is also identified in the FFT, and the measured interlayer rotation angle between GDY and one of the graphene layers is ~16°, all of which are consistent with previous observations.

The electronic structure of the as-grown GDY was probed by x-ray photoelectron spectroscopy (XPS), electron energy-loss spectroscopy (EELS), and photoluminescence (PL) spectroscopy, respectively. Figure 3A presents a high-resolution asymmetric C 1s XPS spectrum of the GDY/graphene film. The C 1s peak can be deconvoluted into four subpeaks at 284.6, 285.3, 286.9, and 288.5 eV, corresponding to C–C (sp²), C–C (sp), C=O, and C≡O, respectively (19). According to the GDY structural model, the benzene rings are connected to each other by conjugated diyne linkers. The ratio of the sp- to sp²-hybridized carbon atoms should thus be exactly 2. The experimentally observed ratio is, however, less than 2, owing to the contribution from the underlying graphene fully with sp²-hybridized carbon. The small amounts of C=O species may originate from traces of impurities or defects in the sample. Moreover, the sp²- and sp-hybridized carbon in the as-grown film were also identified at nanoscale using site-specific EELS by showing two distinct π* resonance features located at 285.9 and 287.1 eV, respectively (fig. S9). In addition, the PL properties of as-grown GDY were investigated on the hexagonal boron nitride (h-BN) substrate (Fig. 3C and fig. S10). The PL spectrum exhibits a single emission peak, with a maximum at a photon energy of 1.79 eV, which is close to the predicted band gap of GDY (fig. S11).

Raman scattering contains the fingerprint structural information of GDY, especially for those Raman-active diyne linkers arranged in the specific topology of GDY. Although the typical Raman spectra of GDY were systematically studied by group theory and first-principles calculations (29), experimentally, the reported Raman spectra always show broadened and poorly resolved bands (15, 19, 23, 27), which probably originate from the poor crystallinity and/or highly defective structure and cannot match the predicted spectral features of GDY well. In our experiments, the intrinsic high-resolution Raman fingerprints of GDY were observed by using the graphene substrate via graphene-enhanced Raman scattering (GERS) (30), for the first time. Figure 3B shows seven sharp Raman peaks in the range of 1000 to 3000 cm⁻¹, including 1344.8 cm⁻¹ (defined as G′), 1430.1 cm⁻¹ (defined as G′₁), 1523.9 cm⁻¹ (defined as G), 1586.1 cm⁻¹ (defined as G₂), 2105.9 cm⁻¹ (defined as Y), 2189.2 cm⁻¹ (defined as Y′), and 2696.5 cm⁻¹ (defined as 2D₂). G₂, 2D₂, and 2D bands are assigned to the lattice vibration modes of graphene, as reported (31). The remaining Raman bands are from the GDY structure and in agreement with the predicted Raman fingerprints of GDY (fig. S12) (29). For example, the Y band (centered at 2105.9 cm⁻¹) and Y′ band (centered at 2189.2 cm⁻¹) are attributed to the in-phase and out-of-phase stretching modes of C≡C triple bonds, respectively, as shown in fig. S13. Polarization-dependent Raman spectra of GDY were further measured in XX (red) and XY (blue) polarization configurations to accurately assign the different vibration modes in GDY (see Materials and Methods). As shown in Fig. 3B, G′ and Y′ are assigned to the A₁g mode, of which the intensity significantly decreases in the XY configuration compared with the XX configuration. In contrast, the E₂g modes (G and Y′) are nearly the same in XX (red) and XY (blue) polarization configurations. These results are consistent with the calculated ones using a 2D GDY model (29). Notably, compared with HEB monomers, there is an obvious blue shift of Y′ band in the Raman spectra of GDY (fig. S14), likely caused by the coupling reaction of HEB monomers on graphene. The coupling reaction is further confirmed by observing the intensity changes of Y′ band in the in situ Raman spectroscopy experiments of GDY growth (figs. S15 and S16). In addition, the Raman mapping of the Y′-to-2Dg intensity ratio over the marked area further verified the uniformity of as-grown GDY at the microscopic scale, with a rather uniform contrast shown in Fig. 3 (D to F). The stability of

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**Fig. 3. Spectroscopic characterization of as-grown GDY film.** (A) High-resolution core-level XPS spectrum of C 1s. CPS, counts per second; a.u., arbitrary units. (B) Typical Raman spectrum (black) of GDY grown on graphene transferred to SiO₂/Si substrate and polarization-dependent Raman spectra of GDY measured in XX (red) and XY (blue) polarization configurations by fixing the incident light and the scattered signal’s polarization directions in the parallel-polarized configuration and cross-polarized configuration (excitation at 514.5 nm). (C) PL spectrum of as-grown GDY on h-BN. (D) OM image of as-grown GDY on graphene with a mark indicating the mapping region. (E) Typical Raman spectra randomly collected across the corresponding sample in (D). (F) Integrated intensity ratio Raman maps of Y band (2189.2 cm⁻¹) and 2D band (2696.5 cm⁻¹) over the marked area in (D), confirming the uniformity of the GDY on the graphene surface in macroscopic scales.
as-prepared GDY on graphene was also investigated by Raman spectroscopy. As shown in fig. S17, after being stored in air for 15 days, the Y' to 2Dg intensity ratio decreased from ~0.55 to ~0.47, but it was still kept at a high level, exhibiting an air stability.

To provide mechanistic insights into the solution-phase vdW epitaxial growth of GDY, we investigated the Eglinton oxidative homocoupling of HEB monomers both on graphene and in solution. Specifically, GDY was synthesized under the optimized conditions on SiO2/Si half-covered with single-layer graphene. It is interesting to find that the 2D GDY film would only grow on the surface of graphene rather than that of SiO2/Si, as observed by Raman spectroscopy and SEM, as shown in Fig. 4 (A and C). The products collected in solution show cross-linked and amorphous structures (Fig. 4B), likely originating from the free rotation around alkyne-aryl single bonds during polymerization. All these results indicate that vdW epitaxial type growth is essential to achieve the ultrathin highly crystalline GDY structures. First-principles calculations propose detailed energetics and kinetics of reaction steps for vdW epitaxy (see Materials and Methods). As shown in fig. S18A, the absorbed HEB molecule shows a flat-lying geometry parallel to the graphene surface with a binding energy of 1.47 eV per molecule. We note that the adsorbed HEB monomers have an energetically preferred rotation angle of ~15° with respect to the underlying graphene in fig. S18 (B and C), which surprisingly coincides with the case of GDY on graphene. Both energy barriers and reaction energies of the vdW epitaxial growth strategy are lower than those of the solution growth strategy, as shown in Fig. 4D and fig. S19. For instance, the energy barrier of the rate-determining step (that is, HEB deprotonation) (steps II to III) of graphene-mediated vdW epitaxy is only 0.174 eV, which is significantly lower than that in solution (0.612 eV). The reduced barrier and the reaction energy are possibly caused by the enhanced interaction between the Cu(OAc)2 catalyst and HEB molecules that are coadsorbed onto the graphene substrate and in the vicinity of each other (Fig. 4E). As a result, the consecutive in-plane coupling of HEB admolecules catalyzed by the coadsorbed Cu(OAc)2 species becomes the fast and energetically favored reaction route that leads to the formation of an ultrathin single-crystalline GDY film (Fig. 4C).

Finally, as a proof-of-concept study of the electrical property and applications of GDY, the GDY/graphene film has been fabricated as an NH3 gas sensor first, showing great potential in gas sensing (fig. S20). Furthermore, we replace graphene with h-BN for electric measurement. To evaluate the electrical property of the as-synthesized GDY, we fabricated field-effect transistors with SiO2 as the gate dielectric and conducted the measurement at room temperature. A schematic depiction of the device and I-V curve is shown in figs. S21 and S22. The conductivity is calculated as 3180 S m−1, which is comparable with the reported results (19, 32), and demonstrates that our as-prepared GDY film exhibits p-type semiconducting properties.

In summary, we reported a facile synthetic route to ultrathin single-crystalline GDY through a solution-phase vdW epitaxial strategy. The GDY film has a trilayer structure with an ABC stacking order as directly observed by HRTEM. Here, the high quality of the as-synthesized GDY film and the GERS effect ensure that the predicted Raman fingerprints belonging to a perfectly ordered 2D GDY structure are experimentally observed. This study paves the way for the design and synthesis of 2D acetylenic carbon allotropes with different degrees of sp and sp2 hybridizations by using monomers with diverse structures.

MATERIALS AND METHODS
Preparation of GDY through vdW epitaxy in solution
Chemical vapor deposition–grown graphene was transferred onto a 300-nm SiO2/Si substrate as the substrate for the synthesis of GDY. The

Fig. 4. Proposed mechanism for synthesis of GDY film on graphene through a solution-phase vdW epitaxial strategy. (A) Raman spectra of GDY on different reaction areas (on graphene, in solution, and on SiO2/Si), synthesized under the same conditions. SEM images of as-grown GDY and corresponding selected-area electron diffraction (SAED) pattern in inset, synthesized in solution (B) and on graphene (C). (D) Proposed mechanism for the Eglinton reaction and computed free-energy profiles on graphene (blue) and in solution (red). (E) Optimized geometries from I to VI (see fig. S19 for details; Cu, blue; O, red; C, gray, and H, white).
as-prepared graphene substrate was immersed into the dichloromethane solution of HEB (0.1 mM, 5 ml). Then, a pyridine solution (5 ml) of copper acetate (1.6 mM) and toluene (2 ml) were added to the dichloromethane solution of HEB. Under an inert argon atmosphere at room temperature, the successive catalytic coupling reaction for 24 hours led to the growth of the 2D covalent network, generating an ultrathin GDY film at the graphene surface. Using the same synthetic procedures, GDY can be synthesized on h-BN by using h-BN as the substrate for the synthesis of GDY. The as-prepared GDY on h-BN was used for PL and the electronic measurement. One- to few-layer graphene samples mechanically exfoliated on a 300-nm SiO2/Si substrate were also used to synthesize GDY for the thickness analysis.

**TEM image processing and simulation**

HRTEM images and EELS spectra were taken on an FEI Titan 60–300 electron microscope equipped with a spherical aberration (Cs) corrector in the image-forming (objective) lens and a monochromator, operated under 80 kV. HRTEM simulation was performed by using the multislice method implemented in QSTEM software (33). Simulation conditions used were as follows: 80 kV; Cs = 10 μm; Cc = 1 mm; ΔE = 0.13 eV; Δf = 68 nm; α = 0.2 mrad; focal spread = 5 nm; an objective aperture was used.

**Polarized Raman measurement**

Polarized Raman spectra were measured using JY Horiba HR800 with a 514.5 nm laser. The optical layout is shown in fig. S23. The incident polarization direction was determined by polarizer I, and polarizer II was in front of the detector. The Z(XX)Z and Z(XY)Z geometries, but with the polarization direction of the polarizers. GDY belongs to the D6h point group, and the phonon modes at the Brillouin zone center can be presented as

$$\Gamma_{\text{graphdiyne}} = 2A_{2u} + 5E_{1u} + 3B_{1u} + 3B_{2u} + 3E_{2u} + 3A_{1g} + 3A_{2g} + 3B_{1g} + 3E_{1g} + 6E_{2g}$$

$A_{1g}$, $E_{1g}$, and $E_{2g}$ modes are Raman-active under backscattering geometry, but $E_{1g}$ modes cannot be detected.

The Raman tensors of the $A_{1g}$ and $E_{2g}$ modes can be presented as follows

$$R(A_{1g}) = \begin{pmatrix} a & 0 & 0 \\ 0 & a & 0 \\ 0 & 0 & b \end{pmatrix}$$

$$R(E_{2g}) = \begin{pmatrix} d & 0 & 0 \\ 0 & d & 0 \\ 0 & 0 & d \end{pmatrix}$$

For $A_{1g}$ modes, under the XX configuration, the Raman intensity can be calculated as

$$I_{XX} = \left| \begin{pmatrix} 1 & 0 & 0 \\ 0 & a & 0 \\ 0 & 0 & b \end{pmatrix} \right|^2 = a^2$$

while under the XY configuration, the Raman intensity is zero. For $E_{2g}$ modes, under XX and XY configurations, the Raman intensities are $d^2$. With polarized Raman spectra, the Raman modes can be assigned.

**Free-energy computation during the synthesis of GDY**

First-principles calculations were performed using the density functional theory method implemented in the Vienna Ab-initio Simulation Package code to obtain the energy variation during the synthesis of GDY. The generalized gradient approximation with the Perdew-Burke-Ernzerhof–type exchange-correlation functional and the projector augmented wave method were adopted in all calculations, as well as the Tkatchenko-Scheffler (TS) method for the vdW corrections. Spin-polarized calculations were considered for all models. The plane-wave energy cutoff was fixed at 400 eV. The climbing image nudged elastic band (cNEB) method was exploited to locate the transition states during the process of deprotonation. The convergence of energy and force was set as $10^{-4}$ and $10^{-2}$ eV/Å (0.05 eV/Å for cNEB calculations), respectively.

**SUPPLEMENTARY MATERIALS**

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/4/7/eaat6378/DC1

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