Graphene Oxide Functionalization

Controlled Covalent Functionalization of Thermally Reduced Graphene Oxide To Generate Defined Bifunctional 2D Nanomaterials

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Abstract: A controlled, reproducible, gram-scale method is reported for the covalent functionalization of graphene sheets by a one-pot nitrene [2+1] cycloaddition reaction under mild conditions. The reaction between commercially available 2,4,6-trichloro-1,3,5-triazine and sodium azide with thermally reduced graphene oxide (TRGO) results in defined dichlorotriazine-functionalized sheets. The different reactivities of the chlorine substituents on the functionalized graphene allow stepwise post-modification by manipulating the temperature. This new method provides unique access to defined bifunctional 2D nanomaterials, as exemplified by chiral surfaces and multifunctional hybrid architectures.

Graphene is a two-dimensional (2D) carbon network with unusual properties, including extraordinary mechanical stiffness, strength, and elasticity, outstanding electrical and thermal conductivity, and many others.[1-3] Despite these unique properties, its low solubility, poor reactivity, and the limited accessibility of a well-defined basal plane are major challenges that must be addressed for future applications. An ideal method to overcome these problems is the covalent attachment of functional molecules to its surface.[3-12] The nitrene [2+1] cycloaddition reaction is one of the most useful approaches to conjugate diverse functional groups to the surface of graphene.[13] However, this approach requires high temperatures to functionalize carbon-based nanomaterials efficiently.[14,15] Therefore, controlling the functionality and surface structure of the obtained 2D nanomaterials is challenging.[16] The functionalization of graphene under ambient conditions by perfluoroazides as precursors of the highly reactive nitrenes has been explored to overcome this drawback.[13] The mild reaction conditions and well-defined functionality provide superior efficiency and simplicity compared to common methods for the functionalization of nitrenes.[17,18] In addition to efficiency and simplicity, a universal and affordable functionalization method should also involve readily available starting materials and provide reactive functional groups for flexible post-modification. 2,4,6-Trichloro-1,3,5-triazine is a cheap, commercially available compound and is highly reactive towards nucleophiles such as amines. The reaction between sodium azide and 2,4,6-trichloro-1,3,5-triazine at low temperature results in 2-azido-4,6-dichloro-1,3,5-triazine, which is a precursor to the highly reactive nitrene intermediates.[19] Inspired by perfluoroazides,[20,21] the lifetime of these electron-poor nitrenes could be long enough to conjugate to the π-system of graphene through a [2+1] cycloaddition reaction.

Herein, dichlorotriazine-functionalized TRGO (TRGO-Trz) was successfully synthesized by the reaction of TRGO with 2-azido-4,6-dichloro-1,3,5-triazine (Figures 1 and S1). The nitrogen content of TRGO-Trz was determined by elemental analysis to calculate the degree of functionalization (DF; triazine/carbon atoms, see the Supporting Information, page S12). Since NMP interacts with the surface of graphene strongly,[22] control experiments were performed to subtract any nitrogen content originating from the solvent (Supporting Information, page S7). To increase the DF of TRGO-Trz, it was repeatedly functionalized up to three times by nitrene [2+1] cycloaddition reactions. TRGO functionalized once, twice, or three times are denoted as TRGO-Trz1, TRGO-Trz2, and TRGO-Trz3, respectively (Table S1). Subsequent functionalizations dramatically increase the DF of TRGO-Trz from 1:50 to 1:25 after three runs (Table S2). The DF values of TRGO-Trz1, TRGO-Trz2, and TRGO-Trz3 that were calculated on the basis of mass loss in the thermogravimetric analysis (TGA) diagrams at 450°C are in good agreement with the obtained elemental analysis results (Table S2 and
The correlation between the DF and the number of reactions allows this method to be considered as a controlled covalent functionalization. The binding energy (BE) range between 285 and 289 eV in the highly resolved C1s XPS spectrum is assigned to contributions of the new dichlorotriazine groups containing C@Na and C@Cl bonds (Figures 2a and S5). The stepwise increase in the intensity of these component peaks from TRGO-Trz1 to TRGO-Trz3 supports the hypothesis of an increase in the DF after repetitive runs. The CK-edge NEXAFS spectrum for pristine TRGO surfaces is displayed in Figure 2b (and Figure S3). The spectra arise from transitions from the C1s orbital to various empty, bound final states. At the low-energy side, the NEXAFS spectrum shows a sharp resonance at a photon energy of 285.4 eV (d), which corresponds to a C1s→π* transition.[22,24,25] After functionalization of TRGO, all the TRGO-related resonances are found again in the C K-edge spectrum. New resonances (e–g) related to the dichlorotriazine groups and their covalent bonds to TRGO occur in the “fingerprint” region at photon energies between 286 and 290 eV. The intensities of these resonances increase from TRGO-Trz1 to TRGO-Trz3, thus revealing an improved DF. At the same time, the intensity of the TRGO-related π*-resonance decreases. This can be explained by a consumption of sp2 carbon bonds during the reaction and the formation of other types of carbon bonds, which are represented by the new resonance features (e–g) in the “fingerprint” region.[25]

TRGO-Trz can be used as a platform with a defined DF. Furthermore, it is expected to have a similar chemical reactivity as dichlorotriazine derivatives.[26] Stepwise nucleophilic substitution of the dichlorotriazine groups results in the formation of well-defined bifunctional 2D nanomaterials (Figure 1).

To prove the controlled bifunctionalization, l- and d-cysteine (l-Cys and d-Cys) were conjugated selectively and in a stepwise manner to TRGO-Trz, by tuning the temperature of the nucleophilic reactions (Figure 1). Based on the XPS results and elemental analysis, the ratio of cysteine molecules to TRGO carbon atoms for TRGO-l-Cys and TRGO-l,d-Cys was approximately 1:47 and 1:22, respectively (Figures S6 and S7, and Table S5). This clearly indicates a quantitative substitution of the first and second chlorine atoms of TRGO-Trz, by controlled post-modification (Supporting Information, page S3). Since they are enantiomers, substitution of the first chlorine atom of the dichlorotriazine groups by l-Cys should produce chiral 2D nanomaterials.[27]

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The binding energy (BE) range between 285 and 289 eV in the highly resolved C1s XPS spectrum is assigned to contributions of the new dichlorotriazine groups containing C=N and C=Cl bonds (Figures 2a and S5).[23] The stepwise increase in the intensity of these component peaks from TRGO-Trz1 to TRGO-Trz3 supports the hypothesis of an increase in the DF after repetitive runs. The C K-edge NEXAFS spectrum for pristine TRGO surfaces is displayed in Figure 2b (and Figure S3). The spectra arise from transitions from the C1s orbital to various empty, bound final states. At the low-energy side, the NEXAFS spectrum shows a sharp resonance at a photon energy of 285.4 eV (d), which corresponds to a C1s→π* transition.[22,24,25] After functionalization of TRGO, all the TRGO-related resonances are found again in the C K-edge spectrum. New resonances (e–g) related to the dichlorotriazine groups and their covalent bonds to TRGO occur in the “fingerprint” region at photon energies between 286 and 290 eV. The intensities of these resonances increase from TRGO-Trz1 to TRGO-Trz3, thus revealing an improved DF. At the same time, the intensity of the TRGO-related π*-resonance decreases. This can be explained by a consumption of sp2 carbon bonds during the reaction and the formation of other types of carbon bonds, which are represented by the new resonance features (e–g) in the “fingerprint” region.[25]

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Meanwhile, substitution of the second chlorine atom would racemize the system. Accordingly, the controlled and stepwise modification of TRGO-Trz by L- and D-Cys is also demonstrated by the chiroptical properties of the synthesized 2D stereoisomers. The circular dichroism (CD) analysis shows characteristic peaks of L- and D-Cys at 210–220 nm (Figure 2c) for TRGO-L-Cys and TRGO-D-Cys, thus confirming successful conjugation of those amino acids to the TRGO-Trz under ambient conditions (Figure 2d). However, conjugation of the next enantiomer to those 2D stereoisomers results in a racemized 2D nanomaterial without any special absorption in the CD spectra (Figure 2d).

The utility of our method is further demonstrated by stepwise attachment of differently functionalized macromolecules, that is, azido-polyglycerol (hPG-N₃) and polyglycerol amine (hPG-NH₂) or amino-β-cyclodextrin (β-CD-NH₂) to TRGO-Trz. To minimize the steric hindrance between the macromolecules, we synthesized a TRGO-Trz with a low DF (TRGO-TrzL). The synthesis and characterization of TRGO-TrzL is explained in the Supporting Information (page S9, Tables S2 and S3, and Figure S8). Nucleophilic substitution of the first chlorine atom of TRGO-TrzL by propargylamine at 25°C resulted in a TRGO derivative suitable for subsequent click reactions (Figure 3a). Accordingly, the click reaction between hPG-N₃ and propargylamine-functionalized TRGO led to water-dispersible 2D nanomaterials with low polymer coverage (TRGO-hPGL, Figure 3a). A subsequent nucleophilic reaction between TRGO-hPGL and β-CD-NH₂ or hPG-NH₂ at 60°C afforded 2D nanomaterials with hPG/β-CD dual functionality (TRGO-hPGL-CD) or fully coated by polyglycerol derivatives (TRGO-hPGL₃), respectively (Figures 3a, S14, and S12). The two-step post-modification of TRGO-Trz by macromolecules has been evaluated by different thermal and spectroscopic methods (see Supporting Information Figure 3.

**Figure 3.** a) Stepwise attachment of polyglycerol and cyclodextrin to TRGO-Trz. i) Propargylamine, DMF, Et₃N, 25°C, 24 h; ii) polyglycerol azide 4%, sodium ascorbate, CuSO₄·5H₂O, DMF, 50°C, 24 h; iii) β-CD-NH₂, Et₃N, DMF, 60°C, 12 h; iv) polyglycerol amine 50%, Et₃N, NMP, 60°C, 48 h. b) Highly resolved C1s spectra of TRGO-hPG₃ and TRGO-hPGL-CD (for assignments and full spectra see Figures S8–S11, and Tables S3 and S4). c) UV spectra of TRGO-hPG₃/rhodamine 6G and TRGO-hPGL-CD/rhodamine 6G. d) Host–guest interaction of TRGO-hPG₃/rhodamine 6G and TRGO-hPGL-CD/rhodamine 6G, Milli-Q water, rhodamine 6G, 30 min incubation (see page S28 in the Supporting Information for loading capacity).
After a click reaction, the ratio of the XPS peak areas of C1s components related to polyglycerol and TRGO \((C_{\text{PG}}/C_{\text{TRGO}})\) was 0.46:1, thereby suggesting that TRGO-hPG\(_L\) contains about 31 wt.% polyglycerol and about 69 wt.% TRGO carbon atoms (Figure S11). This result agrees with TGA data showing a 30 wt.% polyglycerol content for TRGO-hPG\(_L\) (Figure S15b). In the XPS spectrum, the peak area ratio of C1s at 286.6 eV increased upon conjugation of β-CD-NH\(_2\) to TRGO-hPG\(_L\). Since this peak is assigned to components with C–O bonds, it proves the successful attachment of cyclodextrin to graphene sheets in the second step of post-modification (Figures 3b and S12, and Table S3).

Furthermore, UV measurements showed that the loading capacity of TRGO-hPG\(_L\)-CD with rhodamine 6G is 1.6 times higher than TRGO-hPG\(_L\) (Figure 3c; Supporting Information page 28). As a consequence of the presence of the conjugated cyclodextrins, TRGO-hPG\(_L\)-CD can deliver suitable guest molecules not only by π-π stacking but also through host–guest interactions (Figure 3d).

In a second example of post-modification, the nucleophilic reaction between TRGO-hPG\(_L\) and hPG-NH\(_2\) results in a positively charged 2D nanomaterial (TRGO-hPG\(_H\)). In the XPS spectrum of TRGO-hPG\(_H\), the ratio of the peak areas of C1s components related to polyglycerol and TRGO \((C_{\text{PG}}/C_{\text{TRGO}})\) increased from 0.46:1 to 1.23:1 (Figure S11, Table S3). This clearly proves efficient conjugation of hPG-NH\(_2\) to the carbon surface. The C K-edge spectrum (Figure S10) shows that the intensity of the TRGO-related π*-resonance \((j)\) decreases with increasing polymer coverage. This is due to the increased thickness of the TRGO with polyglycerol, which attenuates the electron signal originating from the TRGO substrate. The other prominent change in the C K-edge NEXAFS is the occurrence of the resonance \((n)\) at 289.4 eV, which is assigned to C–H* resonances.[29] This is taken as proof of the successful synthesis of TRGO-hPG\(_H\), and TRGO-hPG\(_H\). Scanning force microscopy (SFM) in tapping mode (TM) and quantitative nanomechanical mapping mode (QNM) were used to investigate the TRGO functionalization (Figure S16). SFM height images of the TRGO-hPG\(_H\) show spherical objects on the TRGO surface (Figures 3a and 4a). Both TRGO-hPG\(_L\) and TRGO-hPG\(_H\) have a height of \((11 ± 2)\) nm at 35–40% (ambient) relative humidity (Figures 4b,4c). However, in the case of TRGO-hPG\(_H\), the polyglycerol branches were tightly packed (Figure S18). SFM suggests an integrated polyglycerol layer with a height of \(4–5\) nm on both sides of the functionalized TRGO-Trz, thereby leading to an overall thickness of \(11–12\) nm. Furthermore, sample deformation images obtained from QNM-SFM provides complementary information that demonstrates the attachment of hPG to TRGO-Trz. The softer regions are positions where spherical objects are observed. Regions with deformations of \(1–2.5\) nm under applied forces of \(0.3–1\) nN by the SFM tip are attributed in the case of TRGO-hPG\(_L\) to single or a few polyglycerol molecules (Figures S19a,b). The more deformable objects observed on TRGO-hPG\(_H\) are attributed to the tightly packed polyglycerol branches (Figures S19c,d).

Finally, the ability of TRGO-hPG\(_H\) to function as a multi-valent platform to bind gold nanoparticles was investigated. Although no specific interactions between negatively charged gold nanoparticles (AuNPs) and TRGO-hPG\(_H\) were observed in phosphate buffered saline (PBS buffer, Figure 4d), the immobilized AuNPs (ca. 5 nm) can be detected on the surface of TRGO-hPG\(_H\) in stereoscopic cryo-TEM image pairs (Figures 4e and S21). This is due to the electrostatic interactions between AuNPs and positively charged polyglycerolamine branches that are attached to the surface of TRGO-hPG\(_H\) (Figure S15a).

In conclusion, we have developed a simple and mild method for the controlled covalent functionalization of TRGO by an in situ nitrene cycloaddition reaction using commercially available reagents. The functionalized graphene sheets can be transformed into defined 2D surfaces by stepwise post-modification in high yields. The advantage of this method is the construction of bifunctional carbon nano-materials with controlled ligand densities, which even allows the conjugation of sensitive bio(macro)molecules at room temperature.

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Conflict of interest

The authors declare no conflict of interest.

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