Chemical Modification of Graphene Oxide by Nitrogenation: An X-ray Absorption and Emission Spectroscopy Study

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Nitrogen-doped graphene oxides (GO:Nx) were synthesized by a partial reduction of graphene oxide (GO) using urea [CO(NH₂)₂]. Their electronic/bonding structures were investigated using X-ray absorption near-edge structure (XANES), valence-band photoemission spectroscopy (VB-PES), X-ray emission spectroscopy (XES) and resonant inelastic X-ray scattering (RIXS). During GO:Nx synthesis, different nitrogen-bonding species, such as pyrrolic/graphitic-nitrogen, were formed by replacing oxygen-containing functional groups. At lower N-content (2.7 at%), pyrrolic-N, owing to surface and subsurface diffusion of C, N and NH is deduced from various X-ray spectroscopies. In contrast, at higher N-content (5.0 at%) graphitic nitrogen was formed in which each N-atom trigonally bonds to three distinct sp²-hybridized carbons with substitution of the N-atoms for C atoms in the graphite layer. Upon nitrogen substitution, the total density of state close to Fermi level is increased to raise the valence-band maximum, as revealed by VB-PES spectra, indicating an electron donation from nitrogen, molecular bonding C/N/O coordination or/and lattice structure reorganization in GO:Nx. The well-ordered chemical environments induced by nitrogen dopant are revealed by XANES and RIXS measurements.

The chemical toughness of a single-layered graphene sheet is important for electron transport and electro-catalytic reactions, and especially for interior (crystalline structure) and exterior (nano-composite) modifications of graphene-like structures. The semiconducting properties that support nano-electronics and light-energy harvesting applications require opening of the energy gap of graphene. Chemical doping, substrate induction, and crystallographic orientation can all affect the host state, charge transfer, and changes in the electronic density of states (DOSs). Doping with boron or nitrogen atoms as the electron-acceptors or donors through carbon substitution, respectively, is commonly conducted to control the Fermi level (E_F) and metal-semiconductor transition. Decoration of the graphene structure with hydroxyl and epoxy groups, to form chemically derived graphene oxide, reduces graphene oxide (r-GO) in a controllable oxidation and reduction step, favors large-volume synthesis and chemical solubility in a variety of solvents. In fact, the chemical doping of graphene can lead to the formation of external covalent bonds in contrast to the six-membered structure and prevent the conjugation of π-states. The removal/substitution of carbon atoms, topological defects/vacancies, structural disorder, molecular absorption and symmetry breaking have been demonstrated to result in the formation of localized states and electron/hole states in the electronic structures of GO and r-GO. Chemical doping affects the surface of GO and is also involved in dislocating the lattice of graphene. Therefore, the mechanisms of...
the changing gapless property of graphene from dopant-induced perturbation and defect/vacancy-induced rearrangement need to be understood in order to develop energy storage and environmental applications\(^{26,27}\).

Among the various processes for synthesizing, the Hummers method\(^{18}\) is commonly used to extract individual graphene sheets that comprise several oxidation groups (i.e. C-OH, -COOH, C-O-C and C=O bond). In this process, highly oriented pyrolytic graphite (HOPG) is exfoliated and then dispersed into an aqueous solution. Various methods for controllably de-oxidizing functional groups, such as reduction with hydrazine\(^{19}\), sodium borohydride\(^{20}\), hydrothermal treatment\(^{21}\) and thermal annealing\(^{22}\), are used to inactivate the chemical connection between epoxide/hydroxyl groups in the GO-basal network. However, these processes desorb the covalently bonded oxygen and have two important effects, namely mixing the metallic \(sp^2\) domain and the insulating \(sp^3\) matrix, and forming structural defects/vacancies in the structural configuration\(^{23}\). Instead of intensively breaking oxygen molecules by these reduction processes and relieving unpaired electrons, nitrogen-associated replacement can be used to decompose the oxygen-containing functional groups because nitrogen has a similar atomic radius to oxygen and the chemical hybridization\(^{24}\). Li et al.\(^{25}\) developed a chemical method to obtain N-doped GO through thermal annealing of GO in ammonia at various temperatures; where they found that the content of doped nitrogen achieved by the replacement of oxygen-containing functional groups is proportional to the increase in temperature. The electrical conductivity measurement revealed that the Dirac point is at a negative gate voltage because nitrogen is an \(n\)-type electron dopant\(^{26}\). Since nitrogen and oxygen co-exist in the graphene sheet, along with structural defects, the effects of chemical doping and/or nitrogenation in GO warrant intensive study. Among the various techniques for studying element-specific absorption, the X-ray absorption near-edge structure (XANES) spectroscopy can be used to detect the unoccupied states in the conduction-band\(^{24}\) and X-ray emission spectroscopy (XES) is used to probe the occupied states in the valence-band of the material\(^{27}\). XANES/XES is element-specific and site-specific, making it a powerful tool for elucidating detailed electronic properties of materials. Owing to the momentum conservation between the excited electron in the conduction band and decayed valence-band hole, resonant inelastic X-ray scattering (RIXS) is used to probe the symmetry points along the band structures relying on its doping, reduction and lattice effect\(^{28}\). The graphene-related and GO/r-GO materials have been found to demonstrate novel electronic and magnetic properties resulting from different electronic structure and bonding environment, offering the working mechanism for the future use\(^{21,28-31}\).

In the present study, GO was used as a starting material to elucidate the mechanism of the nitrogen substitution by replacing the oxygen-related bonds, and to verify the modification of electronic/bonding structures and changes in the binding energy of GO:N\(_x\) by the partial reduction of GO with urea using household microwave. This process is simple and differs from several other reported approaches\(^{19,25,26,32,33}\). In this approach, the chemical environments in GO:N\(_x\) are monitored by studying the C, N, and O at various concentrations of nitrogen; the effect on band gap opening (or energy separation) of the activation of nitrogen and oxygen atoms in the graphene-basal plane network is also examined. For this purpose, various forms of spectroscopy, including Raman spectroscopy, X-ray photoelectron spectroscopy (XPS), valence-band photoemission spectroscopy (VB-PES), and combined XANES/XES and RIXS techniques were used.

**Results and Discussion**

Initially, GO is synthesized herein by the Hummers method\(^{18}\), and then mixed with urea to synthesize GO:N\(_x\) in a household microwave at 700 W for various durations. The surface morphologies of GO and GO:N\(_x\) are investigated using transmission electron microscopic (TEM) images, which show the uniform stacking of multi-layered GO/GO:N\(_x\) as presented in Fig. 1(a). The elements in GO/GO:N\(_x\) (at\% of C, O and N) are quantified using TEM-energy dispersive spectroscopy (EDS). The TEM-EDS study revealed that the oxygen content gradually decreases in the order from 32.5 (GO) \(\rightarrow\) 26.2 (GO:N\(_{2.7}\)) \(\rightarrow\) 19.3 at\% (GO:N\(_{5.0}\)) as the carbon (nitrogen) content increases in the order from 67.5 (GO) \(\rightarrow\) 71.1 (2.7) \(\rightarrow\) 75.7 (5.0) at\% in GO and GO:N\(_x\). These quantitative results imply that treatment with nitrogen induces oxygen reduction and carbon purification, facilitating nitrogen substitution in GO:N\(_x\) (x \(=\) 2.7 \& 5.0 at\%). The microstructural properties of GO and GO:N\(_x\) are studied by Raman spectroscopy, as presented in Fig. 1(b). The Raman spectrum of the pristine GO exhibits three characteristic features: the \(D\) band at \(\sim\) 1362 cm\(^{-1}\), the \(G\) band at \(\sim\) 1587 cm\(^{-1}\) and the \(2D\) band at \(\sim\) 2702 cm\(^{-1}\). The \(G\) and \(D\) band yield information about the lattice structure and about graphene-related \(E_{2g}\) phonon and defect-resolved \(A_{1g}\) symmetry, respectively\(^{24}\). The intensities of all features (\(D, G\) and \(2D\)) of GO:N\(_x\) significantly decrease. In GO and GO:N\(_x\), the splitting of \(G\)-peaks in two components (\(G\) at \(\sim\) 1587 and \(D\) at \(\sim\) 1629 cm\(^{-1}\)) is attributed to the mixed signal from the disordered graphene lattice upon nitrogen doping\(^{25}\). The peak at \(\sim\) 2455 cm\(^{-1}\) and a very wide peak at \(\sim\) 2920 cm\(^{-1}\) arise owing to a combination of \((D + D')\) bands, which is seen in defect-activated graphene oxides\(^{26}\). The feature of \(D\) is also defect-activated in an inter-valley double resonance process and its intensity is a convenient measure of the degree of disorder\(^{29}\). The decrease in the intensity of \(D\) (1362 \pm 1 cm\(^{-1}\)) and the presence of a new band \((D')\) at \(\sim\) 1629 (\(\pm\) 1) cm\(^{-1}\) can be attributed to the attachment of nitrogen on the top layer or the inter-layer of GO, compared with the well-ordered lattice of HOPG. Upon nitrogenation, the \(2D\) band of GO:N\(_x\) is reduced and slightly shifted, and the \(I_{2D}/I_G\) ratio changes, as displayed in Fig. 1(b) and (c). The intensity ratio \(I_{2D}/I_G\) decreases, and the \(I_G\) and \(I_{2D}\) ratio increase with an increase in \(N\) at\% doping. The second-order \(D\) peak \((2D)\) at \(\sim\) 2702 cm\(^{-1}\) is associated with the two-phonon mode, in which the phonons have opposite momentum, close to the \(K\) point of the Brillouin zone. Various \(N\) at\% in GO:N\(_x\) causes the shape and intensity of the peak to differ significantly from that of the highly ordered HOPG. The \((I_{2D}/I_G)\) ratios of both GO and GO:N\(_x\) are less than one, which is indicative of the formation of tri- or multi-layered GO/GO:N\(_x\), as discussed in our earlier study\(^{29}\).

Liu et al.\(^{26}\) probed layer-dependent GO using Raman scattering and observed that oxidation increases the frequencies of the \(D, G\) and \(2D\) modes of tri-/multi-layer graphene. Liu et al. concluded that oxidation by \(O_2\) creates strong ‘hole’ doping (\(p\)-type) in the graphene surface. In the present study, GO is converted into GO:N\(_x\) by the removal of oxygen-containing functional groups in a de-oxidation process, so the peaks are shifted downward in frequency close to HOPG. Also, nitrogenation shifts the \(G\)-band from 1587 cm\(^{-1}\) (GO) to 1592 (1589) cm\(^{-1}\).
in GO:N\textsubscript{2.7} (GO:N\textsubscript{5.0}) by changing the charge density\textsuperscript{37}. This observation clearly indicates the doping of nitrogen along with the simultaneous removal of oxygen-containing functional groups and intrinsic n-type conduction.

Figure 2(a–c) show the N, C and O 1\textit{s} core-level XPS of the individual chemical binding environment in GO and GO:N\textsubscript{x}. All measured spectra are plotted as black circles and their fitted curves are in different colors; these are obtained by the peak-fitting routine for a minimum of the difference function (\(\chi^2\)) as for the purely statistical noise. The components of the fitting procut are described by the parameters, Shirley/linear background and individual Voigt-function peak (position constrained within \(\pm0.1\) eV). In Fig. 2(a), the wide asymmetric C 1\textit{s} complex band of GO has two maxima and one minimum features which arise gradually as one strong and two weak features, owing to the formation of GO:N\textsubscript{x}. All C 1\textit{s} core-level XPS spectra are well-fitted by four features using the minimum \(\chi^2\) search\textsuperscript{38,39}. These features are attributed to C = C (plum) at ~284.6 eV, C-OH (green) at ~286.3 eV, C-OOH (blue) at ~288.7 eV and C-N bonds (filled yellow) at ~285.4 eV\textsuperscript{40,41}. The number of oxygen-containing functional groups in the GO:N\textsubscript{x} structure gradually decreases as N-doping increases. This change in the reduction of the oxygen-containing functional groups is observed in the O 1\textit{s} XPS spectra in Fig. 2(b). Figure 2(b) presents a wide asymmetric O 1\textit{s} core-level XPS spectrum complex band of GO, that is comprised of various oxygen-related bonds, and is converted to less intense features in GO:N\textsubscript{x}. The fitting yields three features, which are C = O (shaded plum) at ~531.6 eV, C-OH (green) at ~532.4 eV and C-OOH (blue) at ~533.2 eV\textsuperscript{42}. In Fig. 2(c), the intensity of N 1\textit{s} XPS spectrum increases with the nitrogen concentration. The fitting of N 1\textit{s} spectra yields three features, which are centered at ~398.2, ~399.8, and 401.4 eV, and are assigned the pyridinic N at the edge of six-membered ring (plum), the pyrrolic N structure (yellow), and the quaternary N structure (green), respectively\textsuperscript{25}. The peak at ~401.4 eV becomes more intense as the nitrogen concentration increases\textsuperscript{25}, indicating the formation of a more graphitic structure. Overall, the above results reveal that the intensities of oxygen-containing functional groups in C 1\textit{s}/O 1\textit{s} spectra decrease and the intensity of the N 1\textit{s} XPS spectra increases with the increase in nitrogen content in the GO:N\textsubscript{x}. These further confirmed the removal of oxygen-containing functional groups (C = O and C-OH) and an increase in the numbers of nitrogen-related pyrrolic, pyridinic, quaternary-N groups in the GO:N\textsubscript{x}.

The occupied valence DOSs of GO and GO:N\textsubscript{x}; the total DOSs below valence-band maximum (VBM) or \(E_T\) was obtained by using the VB-PES of GO, GO:N\textsubscript{x} and HOPG for a reference spectra, as presented in Fig. 2(d). In this figure, HOPG has four distinct hybridized states, which are C 2p-\(\pi\) at ~3.0 eV, C 2p \(\sigma + \pi\) at 6.0 eV, C 2p-\(\sigma\) at 8.0 eV and C 2s-\(\sigma\) at 10.0 eV\textsuperscript{43}. In pristine GO, the features at ~12.1, 8.6, and 6.7 eV are attributed to the \(\sigma\) and \(\pi\) bonding of C = O, and the O lone pair\textsuperscript{44} and overlaps C 2p-(\(\sigma + \pi\)) and C 2p-\(\sigma\) aromatic features of HOPG. The prominent nitrogen-associated bands, C = N at ~7.3 eV and the N lone pair at ~5.0 eV, are clearly observed in GO:N\textsubscript{x} (x = 5.0 at\%\textsuperscript{45}). These two features are related to the \(p-\pi\) and \(\pi-\pi\) contributions to the DOSs, respectively, as also observed in our earlier study of a-CN\textsubscript{x} thin films\textsuperscript{46}. The VBM was determined from the extrapolation of the threshold-edge of the VB-PES spectra of HOPG and GO:N\textsubscript{x} to the baseline, shown in the upper inset in Figure 1.
Fig. 2(d). The VBM position as 5.1 eV (GO) → 3.1 eV (GO:Nx2.7) → 2.6 eV (GO:Nx5.0) → 1.8 eV (HOPG), indicating an increase of DOSs at/near VBM or $E_F$ with an increase in nitrogen concentration in GO:Nx. This effect could be due to the formation of the N-lone-pair electrons, π-electrons of C-N bonds and σ-electrons of C-N bonds as nitrogen content increases; however, an open question for VBM shift should be related to the lattice rearrangement and oxygen/nitrogen bonding groups, decreasing work function of GO:Nx. This result is consistent with the electrical conductivity measurements, which reveal that the Dirac point is at a negative gate voltage owing to nitrogen as $n$-type electron doping in nitrogen-doped graphene, confirming the change in chemical composition and bonding modification with the active involvement of nitrogen in the network of oxygen-containing functional groups and the carbon matrix structure.

Figure 3(a) displays the normalized and fitting C K-edge XANES spectra of GO and GO:Nx. De-oxidation and nitrogenation significantly change the spectral features and line-shapes of GO. The spectra of GO and GO:Nx are fitted into four features $C_1$, $C_3$, $C_4$, and $C_6$ and six features $C_1$, $C_3$, $C_4$, $C_5$, and $C_6$, respectively, as presented in Fig. 3(a). These features are assigned to $C_1$ at ~285.5 eV ($C = \pi$ bond)49, $C_2$ at ~286.4 eV ($1s \rightarrow \pi^*$ (e$_{2u}$) transition similar to that of pyridine ($C = N$))46,50,51, $C_3$ at ~287.4 eV ($\pi^* C$-H/C-OH)46,52, $C_4$ at ~288.7 eV ($\pi^* C$-OOH/C = O)46,52, $C_5$ at ~289.5 eV (C-N bond with the carbonyl functionality nearby)46,50,51, and $C_6$ at ~290.1 (likely C-H bond)53. The feature of GO observed at ~285.2 eV ($C_1$) is the $C = \pi$ bonds in the six-membered ring, which is shifted to a higher energy of ~285.5 eV. Its intensity increases upon chemical decoration by nitrogen-related functional groups to form GO:Nx owing to de-oxidation in the presence of urea. During de-oxidation, nitrogen in GO:Nx structure is bonded to carbon ($C_5$: 286.4 eV and $C_5$: 289.5 eV)46,50,51 by the changing the numbers of $\pi^* C$-OH ($C_6$: 287.4 eV), $\pi^* C$-OOH/C = O ($C_3$: 288.7 eV)46,52, and $C$-H bonds ($C_3$: 290.1 eV)53. The intensities of all these features vary upon de-oxidation/nitrogenation, as clearly shown in the intensity bar diagram in the inset of Fig. 3(a), consistent with the above XPS results. The increase in the number of $sp^2$ clusters in the form of pyridine-like N-C and graphite-N-like ($C_2$ and $C_3$) bonds and the resonant energy and charge transfer from the N and O atoms to the $sp^2$ clusters involve de-oxidation, nitrogenation, change in structural-ordered and, therefore, changes in the chemical, electronic structure and bonding coordination of GO:Nx. However, to verify directly the increase in the number of substitution-induced localized electronic states in GO:Nx upon nitrogenation, a detailed analysis of the DOSs in the region of $\pi$ or in the $\pi^*$ gap was conducted using RIXS. This technique is a powerful method for investigating the incident coherent excitation ($h\nu_m$) and emitted-energy dependence that is element specific and allows sensitive probe of the symmetry points of the Brillouin zone. Figure 3(b) displays the C K$_\alpha$ RIXS spectra of GO, GO:Nx and HOPG, obtained by excitation with various $h\nu_m$. All of the emission spectra were obtained within the same period and were normalized to yield an intensity of unity for the maximum inelastic emission peak in each spectrum. According to RIXS theory, the total energy and momentum must be conserved in resonant inelastic scattering in absorption-emission processes. The $h\nu_m$-dependent emission features that are identified from the HOPG results from a transition at a well-orientated crystal momentum. Figure 3(b) shows that the C K$_\alpha$ RIXS spectra of GO, GO:Nx, and HOPG depend strongly on the excitation energy, and this dependence is believed to arise from the non-resonant inelastic scattering, related to the similarity in the...
band structures of GO and GO:N_x. The GO/GO:N_x spectra exhibit different line-shapes and emission features (σ and π) that appear slightly dispersive, particularly within the range 280–285 eV, differing clearly from the HOPG spectrum. The π emission derives from transitions from p_z states that are oriented perpendicular to the graphitic plane. Emission from the σ bands is much more isotropic because these bands are derived partially from the p_x and p_y states that are parallel to the surface. The emitted photon energies (hν_\text{em}) vary with hν_\text{in} from 285.1 to 290.1 eV. The hν_\text{em}-dependent emission features arise from transitions from states with a well-defined crystal momentum^{36,37}. The σ orbital at ~273.4 eV and π orbital at ~284.0 eV of HOPG are shrunken by the k-conserving partial DOSs in the well-defined crystal momentum as a function of the hν_\text{in} energy^{58}. The emission spectra of GO:N_x are insensitive to on-site elastic scattering and become broader in different hν_\text{in}-energies as compared to that of HOPG spectrum, which is characteristic of structural disorder effects as additional oxygen and nitrogen atoms are dispersed over the band structures^{59}. The π-lines in the C K-edge, RIXS spectra of the GO/GO:N_{5.0} change with incident photon energy from 285.5 to 285.1 eV, as presented in right-hand panels of Fig. 3(b), in a very different way from that of HOPG. This is because the excitation energies at the various absorption edges of carbons that are bonded to oxygen and/or oxygen-containing functional groups generate more resonant π-states in GO/GO:N_{5.0} than in HOPG. The DOSs of the pristine GO is higher than that of GO:N_x with a shoulder just above E_f and is observed more prominently at hν_\text{in} = 285.5 eV [inset in Fig. 3(b)]. This DOS is highly sensitive to the electronic DOS close to E_f, as observed by Zhong et al^{60} in the C K-edge emission spectra of single wall nanotubes. This feature of GO at ~283.3/282.6 eV (hν_\text{in} = 285.1/285.5 eV) is not observed in the spectrum of the GO:N_x structure because the localizations of carbon core-hole orbitals changed upon the reconstruction of the nitrogen-doped states^{61}. This spectral change is caused by mixing of the π-σ range upon de-oxidation/nitrogenation and the contribution of the C-OH-like feature, C_3-, in the C K-edge XANES spectra. This revealed that the removal of oxygen and/or oxygen-containing functional groups from the GO lattice and induced nitrogen-related states, changed the DOSs in the π-region and/or in the gap between π and π* states.

Figure 4(a) presents the normalized O K-edge XANES spectra of GO and GO:N_x. All spectra are fitted into four features (O_1–O_4), and their intensities are shown in Fig. 4(a). A sharp 1 s → π* C = O (O_2) resonance at ~530.9 eV and intense 1 s → σ* C-OH resonance at ~538.5 eV (O_3) with two shoulders π* C-O-C/C-OH at ~533.9 eV (O_2) and σ* C-O/C = O ~542.4 eV (O_4) are observed^{52,62}. The intensities of these features decreased upon de-oxidation/nitrogenation, as clearly revealed in the intensity bar diagram in the inset of Fig. 4(a). These results confirmed the removal of different oxygen-containing functional groups (C = O and C-OH) from the GO surface and the inclusion of nitrogen that is bonded with carbon and oxygen (pyrrolic, pyridinic, quaternary-N groups) in GO:N_x as the nitrogen content increases in the GO matrix. To investigate the potential removal of oxygen-containing functional groups, the O K-edge RIXS of GO and GO:N_x with hν_\text{in} of 531.1–539.9 eV are used, as shown in Fig. 4(b). All spectra exhibit an intense feature at ~526.5 eV with a shoulder at ~522.1 eV at various hν_\text{in} energies, to which π → 1 s and σ → 1 s in the valence band contribute. A strong resonant contribution is observed at hν_\text{in} = 534.4 eV (O_3); the intensity of the π at 524.0–525.5 eV and the σ electron at 520.9 eV decreased as nitrogen concentration increased^{63}, indicating the reduction of the C-O/C-OH bond in the valence band. The emission width of the π-derived state shrunk at hν_\text{in} = 539.9 eV owing to a loss of oxygen-containing bonds as
the concentration of N increased. The RIXS profile measured at $h\nu_{\text{in}} = \sim 537.5$ and 539.9 eV reflects insensitive in the $\pi$- and $\sigma$-derived states, which may be a result of the incoherent yield and fluorescent superposition effect.

Figure 5(a) displays the normalized N K-edge XANES spectra of GO:N$_x$, which provide evidence concerning the local electronic states that are bound to the nitrogen sites. The spectra are fitted into five features, whose intensity bars are presented in the inset of Fig. 5(b). In the $\pi^*$ region, a strong feature (N2) at $\sim 400.2$ eV is assigned to the -C≡N triplet in GO:N$_2$. Close to this feature, two shoulders at $\sim 399.2$ eV (N1) and at $\sim 401.5$ eV (N3) are attributed to the pyridinic/pyrrolic N-related group with two carbon neighbors in six/five-membered ring and the nitrogen-induced substitution in graphite and/or N-O bonding.

In the $\sigma^*$-region, the 1$s$→$\sigma^*$ transition at $\sim 406.4$ eV (N4) is associated with the superposition of graphite-like and pyridine-like nitrogen structures and/or photoionization of N 1$s$ electrons, whereas the broad feature at $\sim 415$ eV (N5) indicates the $\sigma^*$ state of pyridinic structure on the surface of GO:N$_x$. The intensity of the N1/N2 feature slightly decreases; whereas those of features N3 and N4 increases as the nitrogen concentration (5 at.%) increases, indicating the decrease of non-graphene-like structure (pyridinic/pyrrolic N and -C≡N triplet) and increase of graphene-like structure (N-substitutional site).

Figure 6 displays C, O and N K-edge XANES and $K_\alpha$ XES spectra of GO and GO:N$_x$, to obtain the band gap [or energy separation between the conduction-band maximum (CBM) and VBM] of GO and GO:N$_x$. Details of the derivation of energy separation in the GO from C and O K-edge XANES and $K_\alpha$ XES spectra measurements can be found elsewhere. At the resonant energy $h\nu_{\text{in}} = 406.4$ eV, the RIXS profiles with the high N-content (5.0 at.%) show more intense features associated with $\pi$-bonding electrons at $\sim 394.5$ eV and $\sigma$-bonding electrons at $\sim 391.7$ eV. Interestingly, the dispersive features between $\sigma$ and $\pi$ orbitals and their line shapes differ greatly from each other. When the resonant ($h\nu_{\text{in}}$) energies are 399.2 and 400.2 eV, the RIXS profiles at low N content (2.7 at.%) include more intense features at the $\pi$-state ($\sim 391.7$ eV) and the $\sigma$-state ($\sim 392.7$ eV) than at a higher nitrogen content (5 at.%), owing to the pyridine (N1) and -C≡N triplet bond (N2) in the GO:N$_x$. During de-oxidation/nitrogenation process, various N-O bonds are expected to be formed in the GO network, and the oxygen/oxygen-containing functional groups in GO:N$_x$ (x = 5.0 at%) are strongly reduced, as revealed by the O K$_\alpha$ RIXS emission spectra in Fig. 6(a). At the resonant energy $h\nu_{\text{in}} = 406.4$ eV, the RIXS profiles with the high N-content (5.0 at.%) show more intense features associated with $\pi$-bonding electrons at $\sim 394.5$ eV than that with the lower N-content (2.7 at.%), indicating the formation of a pyridine-like N-bonding structure, which was also observed by Hellgren & Guo et al. in carbon nitride films and is consistent with the appearance of the N lone-pair in VB-PES, as presented in Fig. 2(d). In the pyridine-like N-bonding structure, nitrogen not only forms a hexagonal ring structure, but is substituted for one carbon atom in the hexagonal ring, accordingly, the percentage of carbon-bonded nitrogen bonds must increase as the nitrogen concentration in the structure increases. Therefore, the atomic fraction of nitrogen in a locally ordered C-N structure differs from that of a disordered structure.
energy separation between CBM and VBM is close to 1.1(±0.1) eV for GO:Nx. This result clearly indicates that both oxygen and nitrogen are responsible for the change in the electronic structures at/near $E_f$ in the GO:Nx.

Although the combination of individual XES spectra (as partial DOS) is equal to VB-PES spectra (as total DOS), the dipole cross-section dependence between X-ray polarization and molecular orientation during absorption and de-excitation process is the key to observe the difference between XES and VB-PES. Since the off-resonant (XES) probing is the integrated effect of whole functional group, the on-resonant (RIXS) probing intensifies the spectral weight of discrete ones with the dipole symmetry selection for electronic structure mapping. It could be related to the molecular orientation of functional groups modulated with the nitrogen concentration in GO:Nx, revealing DOSs at/near VBM or $E_f$ (VB-PES result) increases with the decrease of work function$^{47}$. Souto et al.$^9$ calculated the DOSs of N-substituted graphite clusters and found that as the N concentration in the cluster increases, the $\pi$ and $\sigma$ bonds are pushed to higher binding energies and new features close to $E_f$ are formed.

![Figure 5](image1.png)

**Figure 5.** N 1 s (a) XANES and de-convoluted features, The inset is the intensity bar diagram of N 1 s XANES features of GO:Nx (x = 2.7 and 5.0), obtained by fitting. (b) N Kα RIXS spectra of GO:Nx (x = 2.7 and 5.0).

![Figure 6](image2.png)

**Figure 6.** (a) Normalized C K-edge XANES and Kα XES spectra of GO and GO:Nx and HOPG. (b) Normalized O K-edge XANES and Kα XES spectra of GO and GO:Nx. (c) Normalized N K-edge XANES and Kα XES spectra of GO:Nx (x = 2.7 and 5.0).
because $sp^2$-N in a planar graphite structure contributes two electrons to the $\pi$-system. Souto et al. further argued that, as the N content increases, the system adopts a three-dimensional $sp^3$-structure, in which the N $\pi$-electrons become nonbonding lone-pairs as other electrons occupy the $\pi$ orbitals of C-N bonds and the $\sigma$ orbitals of C-N bonds at cluster boundaries. These results are consistent with this study, as the VBM or $E_f$ in VB-PES spectra changes from 5.1 eV (GO) to 2.6 eV (GO:N$_{1.3}$), indicating that DOSs at/near VBM or $E_f$ increases with nitrogen concentration and binding environment. It further supports the band gap (or energy separation) behavior that was obtained from Fig. 6 and is consistent with the results reported by King et al. for amorphous hexagonal boron nitride films with/without silicon doping.

In summary, different nitrogen-bonded carbon/oxygen-containing functional groups are formed by the partial removal of oxygen-containing functional groups from the GO surface through simultaneous de-oxidation/nitrogenation in the presence of urea. The dependence of the electronic and bonding properties to the N species and its concentration was extensively elucidated. The number of oxygen-containing functional groups (C-OH and C-OOH) gradually decreases from GO upon the chemical reaction of nitrogen atoms, yielding distinct features associated with the aromatic C=C state and various N-bonded carbon states, such as nitrile, pyridine-like, and graphitic-like structures in the GO:N$_x$ matrix. Specific chemical environments of nitrogen, as in a disordered graphene-sheet give rise to multiple states in the valence-band structures owing to the $h_{\nu_{\text{eh}}}$-dependent at C $K_{\alpha}$ RIXS. The N $K_{\alpha}$ RIXS features are quite sensitive to non-resonant $h_{\nu_{\text{eh}}}$ reflecting an ordered structure at nitrogen sites, which highlights the complexity and disorder in GO:N$_x$. The VBM or $E_f$ in the VB-PES spectra changes considerably from 5.1 to 2.6 eV upon de-oxidation/nitrogenation, suggesting the preferred metallic behavior of GO:N$_x$. Resonant and non-resonant emission result provides an alternative means to determine element-specific binding and electronic structure between C, O, and N sites, indicative of work function decreasing and VBM increasing. The advantage of using different X-ray spectroscopic methods could trace the electronic and magnetic binding and electronic structure between C, O, and N sites, indicative of work function decreasing and VBM increasing. This study finds that the electronic and structural properties of GO/GO:N$_x$ can be tailored by de-oxidation/nitrogenation in the presence of urea, rendering GO/GO:N$_x$ attractive for use in the fabrication of junction devices and field effect transistors for semiconducting and optoelectronic devices.

**Methods**

**Preparation of GO and GO:N$_x$** Initially, GO is synthesized by the Hummers method. 50 mL of aqueous solution of GO (0.08 g) was prepared by ultrasonating for 50–60 minutes. 0.08 g of urea was then added and sonicated for another 5 minutes. The mixture was then put in a standard household microwave (2450 MHz, 700 W) and heated for 100, and 400 s at maximum output power. The solution was cooled to room temperature and centrifuged at 3000 rpm for 30 minutes. The clear supernatant obtained after centrifuge was separated and the brown/black precipitate was further washed in deionised water a few times by centrifuging at 3000 rpm for 30 minutes. The final slurry was left to dry in the vacuum oven at 50 °C overnight to result in GO with 2.7 at% nitrogen (GO:N$_{2.7}$) and GO with 5 at% nitrogen (GO:N$_{5.0}$) as confirmed by XPS measurements.

**Characterization.** XANES and XES/RIXS were performed at the undulator of beamline 8.0.1 at the Advanced Light Source (ALS), Lawrence Berkeley National Laboratory. The absorption spectra of C, O, N K-edge are recorded in total electron yield and fluorescence yield modes at 300 K and normalized by the current of the Au grid upstream of the beamline. The energy resolution of the absorption spectra is approximately 0.10 eV for C $K_{\alpha}$-edge, 0.30 eV for O $K_{\alpha}$-edge, 0.25 eV for N $K_{\alpha}$-edge XANES measurements. The XES/RIXS spectra are separately calibrated using the elastic features of HOPG, the O site of TiO$_2$, and the N site of BN. Core-level XPS is accurately calibrated using the elastic features of HOPG, the O site of TiO$_2$, and the N site of BN. Core-level XPS is performed at $E_{\text{bvp}} = 621$ eV with an energy resolution of $\sim 0.08$ eV, and VB-PES is measured at a photon energy of 144 eV, using Beamline-09A1 at the National Synchrotron Radiation Research Center (NSRRC), Taiwan. Raman spectroscopy was conducted using an ISA LabRam system that was equipped with a 632.8 nm He-Ne laser with a spot size of approximately 2–3 mm, yielding a spectral resolution of better than 2 cm$^{-1}$. Due care was taken to minimize heating of sample by using a low laser power of less than 2 mW to minimize desorption and/or oxidation by the laser-induced heating of de-oxidized GO. The surface morphologies of GO and GO:N$_x$ were studied using TEM images. Elementals were quantified using TEM-EDS and XPS study.

**References**

1. Geim, A. K. & Novoselov, K. S. The rise of graphene. *Nat. Mater.* **6**, 183–191 (2007).

2. Yoo, E. J. et al. Enhanced Electrocatalytic Activity of Pt Subnanoclusters on Graphene Nanosheet Surface. *Nano Lett.* **9**, 2255–2259 (2009).

3. Eda, G. et al. Large-area ultrathin films of reduced graphene oxide as a transparent and flexible electronic material. *Nat. Nanotech.* **3**, 270–274 (2008).

4. Yeh, T.-E. et al. Graphite Oxide as a Photocatalyst for Hydrogen Production from Water. *Adv. Funct. Mater.* **20**, 2255–2262 (2010).

5. Li, S.-S. et al. Solution-processable graphene oxide as an efficient hole transport layer in polymer solar cells. *ACS Nano* **4**, 3169–3174 (2010).

6. Wehling, T. O. et al. Molecular Doping of Graphene. *Nano Lett.* **8**, 173–177 (2008).

7. Zhou, S. Y. et al. Substrate-induced bandgap opening in epitaxial graphene. *Nat. Mater.* **6**, 770–775 (2007).

8. Schultz, B. J. et al. Imaging local electronic corrugations and doped regions in graphene. *Nat. Commun.* **2**, 372 (2011).

9. Souto, S. et al. Electronic structure of nitrogen-carbon alloys ($a$-CNx) determined by photoelectron spectroscopy. *Phys. Rev. B* **57**, 2536–2540 (1998).

10. Panchakarla, L. S. et al. Synthesis, Structure, and Properties of Boron- and Nitrogen-Doped Graphene. *Adv. Mater.* **21**, 4726–4730 (2009).

11. Chang, C.-K. et al. Band Gap Engineering of Chemical Vapor Deposited Graphene by in Situ BN Doping. *ACS Nano* **7**, 1333–1341 (2013).

12. Wang, B. Y. et al. Nonlinear bandgap opening behavior of BN co-doped graphene. *Carbon* **107**, 857–864 (2016).

13. Kotakoski, J. et al. From Point Defects in Graphene to Two-Dimensional Amorphous Carbon. *Phys. Rev. Lett.* **106**, 105505 (2011).

14. Pereira, V. M. et al. Disorder Induced Localized States in Graphene. *Phys. Rev. Lett.* **96**, 036801 (2006).
65. Zheng, W. T. et al. Chemical bonding in carbon nitride films studied by X-ray spectroscopies. Diamond Relat. Mater. 10, 1897–1900 (2001).
66. Shimada, H. et al. Nitrogen K-edge X-ray absorption near edge structure (XANES) spectra of purine-containing nucleotides in aqueous solution. J. Chem. Phys. 141, 055102 (2014).
67. Bhattacharyya, S. et al. Structure of nitrogenated amorphous carbon films from NEXAFS. Diamond and Related Materials 11, 8–15 (2002).
68. King, S. W. et al. Valence and conduction band offsets at amorphous hexagonal boron nitride interfaces with silicon network dielectrics. Appl. Phys. Lett. 104, 102901 (2014).

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Author Contributions
C.H.C., S.C.R. and W.F.P. designed the experiments having prior discussion with J.H.G. The GO and GO:Nx samples were synthesized by S.S., A.G. and P.P. All measurements are performed by C.H.C., J.W.C., H.M.T., H.W.S., C.H.C. and H.J.L. The data analysis and manuscript writing are done by C.H.C., S.C.R., D.M. and W.F.P. All authors discussed the results and contributed to finalization of the manuscript.

Additional Information
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