Synthesis, Structure and Electronic Properties of Graphitic Carbon Nitride Films

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Supporting Information

ABSTRACT: Dark-colored shiny flakes of graphitic carbon nitride materials produced by reacting dicyandiamide C2N4H4 in a KBr/LiBr molten salt medium were determined to have a C/N ratio near 1.2:1. The compounds also contained 2.3–2.5 wt % H incorporated within N–H species identified by Fourier transform infrared spectroscopy. One recent study revealed analogous results for thin films produced by a similar synthesis method, while a previous investigation instead reported formation of crystalline gC3N4 flakes with a triazine-based graphitic carbon nitride (TGCN) structure. The structures of the materials produced here were studied using a combination of high resolution transmission electron microscopy, X-ray diffraction, IR and Raman and X-ray photoelectron spectroscopy, along with series of density functional theory (DFT) calculations carried out for a range of model layered structures. The results indicate the graphitic layered gC3N4 materials contain a mixture of sp3-hybridized C–N and C–C bonded structures, with TGCN to graphene-like domains existing within the layers. Paramagnetic centers localized on the C3N rings revealed by electron paramagnetic resonance spectroscopy correspond to potential defect structures within the graphitic layers predicted by DFT calculations. Our results combined with those of previous researchers indicate that a range of graphitic carbon nitride materials could exist with different C/N/H ratios leading to tunable electronic properties for catalysis, semiconducting, spintronics and energy applications, that could be targeted by controlling the synthesis and thin film deposition procedures.

1. INTRODUCTION

Layered carbon-based materials are being developed for applications ranging from optoelectronics, energy storage and conversion to catalysis and photocatalysis.1–18 Graphitic carbon nitride (gC3N4) materials are typically reported to be wide-gap semiconductors, with bandgaps ranging between 2.4 and 2.7 eV. Most C3N3Hx compounds currently under investigation correspond to amorphous to nanocrystalline polymers to layered materials formed by thermal condensation of N-rich precursors such as dicyandiamide (C2N4H4: DCDA), melamine or urea by elimination of NH3. Although the end result of such a process is predicted be a fully-polymerized gC3N4 phase, this is not achieved due to evaporation of CN-rich components at higher temperatures, and the products typically have a limiting composition near C2N3H. However, synthesis of gC3N4 films has been achieved by chemical vapour deposition techniques using single-source precursors.1,2 The nanocrystalline materials produced were characterized in that study by high resolution transmission electron microscopy (HRTEM), electron energy loss spectroscopy, Fourier transform infrared (FTIR) and Rutherford backscattering. Fully condensed gC3N4 could be based on tri-s-triazine (heptazine, C6N7) or s-triazine (C3N3)
Figure 1. Schematic diagrams of different structural motifs for carbon nitride layered structures. (a) PH gC₃N₄ (b) polytriazine gC₃N₄ (c) polytriazine gC₄N₃.

Figure 1a,b) units, with polyheptazine (PH) layered structures predicted to give rise to more thermodynamically stable forms. However, such structures have not been observed experimentally. Instead, the materials prepared by Kouvetakis et al. were found to be based on C₃N₃ rings linked by three-coordinated nitrogen (sp²-bonded NC₃) units, forming graphene-like sheets containing C₆N₆ voids within the layers (Figure 1b). Various gC₃N₄ polymorphs based on different stacking arrangements of both the polytriazine (PT) and PH layer motifs have been studied theoretically. Theoretical predictions of Teter and Hemley indicated that the material was gC₃N₄ with a triazine-based layered structure (TGCN), that then represented the first bulk crystalline example of the nanocrystalline phase found by Kouvetakis et al., and that was included in the ab initio theoretical predictions of Teter and Hemley. It was proposed that the semiconducting TGCN phase could then be developed for a wide range of optoelectronics, catalysis and solar energy harvesting applications.

Other layered phases that have also been described as layered “carbon nitrides” include N-doped graphites and graphenes, that have metallic properties. Recent work has predicted the existence of a new compound gC₆N₉, containing C atoms substituted for sp²-bonded N on sites linking the triazine rings within TGCN layers (Figure 1c). This compound has an unsaturated valency count resulting in interesting half-metallic magnetic properties. Doping such structures with H or Li could suppress the metallicity to provide optically transparent materials with photocatalytic properties.

Ladva et al. recently reported carbon nitride thin films deposited on different supports from vapors obtained by reacting DCDA in a eutectic LiBr/KBr molten salt medium. Their chemical analyses indicated a C/N ratio ~1.2:1, that is similar to the materials produced in our own study, but that is substantially more C-rich than the TGCN compound investigated by Algara-Siller et al. Here we present results of our detailed characterization studies of layered gC₆N₆H₂ nanomaterials produced by condensation reactions from DCDA in a LiBr/KBr molten salt medium.

2. METHODS

2.1. Synthesis. Solid DCDA (C₂N₄H₄; 0.83 g, Sigma-Aldrich) was ground under argon together with a pre-dried mixture of KBr (5.99 g, Alfa Aesar) and LiBr (6.48 g, Aldrich) in proportions corresponding to the eutectic melt composition (Tₑ₅ = 329 °C). The mixture was transferred to a quartz glass ampoule (OD 25 mm, ID 22 mm, length15 cm), evacuated to 10⁻⁵ mbar (Leybold vacuum systems, PT70G compact), then sealed under vacuum. The ampoules were heated to 400 °C at 40 °C/min, maintained for 4 h at this temperature, and then raised to 600 °C for 60 h, using a horizontal tube furnace (Carbolite MTF 12/38/400). Some synthesis experiments were also carried out in a vertical (Deltech) furnace but yields of the dark-colored shiny product to be studied here were found to be substantially lower. After cooling, the reaction vessel contained a solid LiBr/KBr block mixed with C₆N₆H₂ reaction products, along with black shiny flakes coating the upper internal walls. The solid residue was filtered and washed to remove residual salts. Larger flakes up to 1–5 mm in dimension (~70 mg aliquots from each run) were also hand separated from the reaction products. One unexpected result was that X-ray analysis of the powdered run products from the LiBr/KBr salt block did not reveal the presence of any crystalline PTI-LiBr, that has previously been reported to form under similar reaction conditions. However, we did observe formation of this phase within the molten salt medium in experiments with the ampoule mounted vertically in our Deltech furnace. Previously, Wirnhier et al. have noted differences in the nature and morphology of PTI materials produced as a function of the reaction chamber geometry.

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Figure 2. XRD patterns obtained for single flakes of layered carbon nitride (gC3N4) materials (Cu Kα: λ = 1.5418 Å). (a) XRD pattern taken in transmission geometry while rocking the sample by ±5° about the incident X-ray beam axis. (b) XRD pattern taken in reflection mode with the scattering vector aligned normal to the sample surface.

2.2. Bulk Chemical Analyses and X-ray Photoelectron Spectroscopy. Samples of separated flakes and powdered materials from three different synthesis runs were sent to the University of Sheffield, Department of Chemistry CCIAS service for CNH analysis. 5–10 mg aliquots of each sample were catalytically combusted in a Vario Micro cube CHN/S analyser to produce N2, CO2 and H2O that were quantified using thermal conductivity detection. XPS measurements were carried out at UCL with a Thermo Scientific spectrometer using monochromated Al Kα radiation. Samples were mounted on conductive carbon tape and a charge compensating flood-gun was applied to avoid charging effects. XPS experiments were carried out both on as-prepared samples and following Ar+ ion-milling erosion to remove surface contaminant species.

2.3. XRD and HRTEM Studies. XRD patterns for sample flakes on a glass support were first obtained with Cu Kα radiation (λ = 1.5418 Å) using a Philips X-Pert diffractometer in reflection geometry, with the scattering vector aligned normal to the sample surfaces. Additional studies were carried out using individual flakes mounted with their largest dimension normal to the X-ray beam in a Stoe Stadip diffractometer, with data collected while rotating the sample by ±5° about the incident beam axis to introduce orientation averaging. HRTEM studies were carried out for samples prepared by sonication in methanol followed by dropping the resultant suspension onto 300 mesh TEM grids with a holey carbon film. TEM images, selected area diffraction (SAED) patterns and energy dispersive X-ray (EDX) imaging results were obtained using a JEOL JEM-2100 instrument.

2.4. IR, Raman and Electron Paramagnetic Resonance Spectroscopy. FTIR data were obtained by transmission through individual flakes placed on a CaF2 slide using a Bruker FTS-66v microbeam instrument. Samples of different thicknesses were examined. Unpolarized microbead Raman spectra were recorded for individual flakes on CaF2 supports using a Renishaw InVia system with 325 nm (He–Cd), 514.5 nm Ar· or 785 nm diode laser excitation. All spectra presented here were obtained with the incident beam normal to the sample flakes. Additional data obtained at different angles to the incident beam showed no differences in the relative band intensities, and the spectra did not change during rotation about the incident beam axis. Electron paramagnetic resonance (EPR) measurements were performed using a Bruker EMXplus spectrometer at 9.85 GHz (X-band) equipped with a 4122SHQE resonator. Spectra were acquired with a magnetic field sweep of 8 mT, microwave power 0.5 μW, modulation amplitude 0.1 mT and frequency modulation 100 kHz. To minimize instabilities and drift in the applied magnetic field, the electromagnet was left to stabilize for 8 h prior to collection. The applied field was calibrated against the Bruker strong pitch standard (g = 2.0028). A single flake of the carbon nitride material was mounted with its plane parallel to the long axis of an EPR tube, which was rotated in the resonator by means of a laboratory-built goniometer (resolution ±5°). Spectra were collected every 20°, and at every 10° near the extrema. Simulations of the continuous wave EPR spectra were carried out to determine orientation-dependent g-values and linewidths using the Easyspin toolbox running in Matlab. Magnetic susceptibility data shown in the Supporting Information (Figure S7) were acquired for single flakes using a Quantum Design MPMS-7 magnetometer with the field applied perpendicular to the flake surface. Temperature scans were taken while warming from 5 to 300 K in 1000 and 2000 Oe magnetic fields.

2.5. Computational Studies. DFT calculations were carried out for different model structures based on gC3N4 and gC5N3 motifs, implemented using the hybrid exchange-correlation B3LYP functional26,27 within the Crystal14 code.28,29 We used 6–21G** Gaussian basis sets and all atoms were treated at all-electron level. We applied Monkhorst–Pack sampling of 6 and 16 for semiconducting and metallic systems, respectively, with 20 and 130 k-points in the irreducible Brillouin zone. Truncation thresholds were set to (6 6 6 6 12) for the Coulomb and exchange series. SCF convergence was set to 10−7 hartree for structure optimization and to 10−3 hartree for calculation of IR and Raman spectra. The fractional coordinates and cell parameters were fully optimized within P1 space group. Structure optimization was deemed to have converged when the largest gradient component was 1.2 × 10−3 a.u. and the largest estimated nuclear displacement was 1.8 × 10−3 a.u. Vibrational spectra were calculated within the harmonic approximation. IR intensities were evaluated using the localized Wannier function approach and Raman intensities by a coupled perturbed Hartree–Fock–Kohn–Sham method.30 DFT cluster calculations implemented within Gaussian 0931 using B3LYP/ EPR-II were used to predict localized electron spin densities and hyperfine coupling (hfc’s) to interpret the EPR data.26,27

3. RESULTS AND DISCUSSION

3.1. XRD and HRTEM. X-ray reflection data for flakes of our gC3N4N3 sample mounted on a glass support showed a single
broad peak near 27° 2θ (Cu Kα radiation) indicated a layered material with interlayer spacing ∼3.24 Å, comparable to the d₀₀₂ reflection of crystalline graphite (d = 3.28 Å). Similar results were obtained in transmission experiments for a single flake obtained while rocking the sample by ±5° about the beam axis (Figure 2). A similar pattern was observed by Ladva et al.34 for their carbon nitride films. However, Algara-Siller et al. obtained different results for their TGCN samples prepared using a similar synthesis procedure. The background-subtracted XRD patterns reported by these authors showed a main peak at 27.2° 2θ accompanied by a smaller peak near 24.4° 2θ that they assigned to the 101 reflection of the graphitic structure, along with additional weak broad features at ∼50° and 56° 2θ.8 They refined their X-ray data within space group P̅6m2 consistent with AB stacking of the triazine-based gC₃N₄ layers, although HRTEM image analysis of thin specimens of the same samples were interpreted to indicate ABC layer stacking (i.e., space group P₆₃/mcm).8

In their XRD analyses, Algara-Siller et al. assumed a planar model for the gC₃N₄ layers, although both their and our DFT calculations indicated that the equilibrium geometry should be buckled.8 Here we calculated XRD patterns for both AB and ABC stacked models of the TGCN structure, using planar and fully-relaxed buckled layer geometries (Figure 3). We note that the character of the diffraction patterns changes significantly once layer buckling is taken into account. In particular, the relative intensities of the two main peaks in the 25°–30° 2θ range are reversed for the buckled layer solutions that correspond to the most stable TGCN arrangements.

We also obtained powder XRD patterns for the brown/black powdered residue recovered after removing the soluble components from the solidified salt block by washing. We note that reacting DCDA in molten LiBr/KBr is typically expected to result in formation of crystalline PTI-LiBr.12–14,25 That phase was found among the run products from reactions carried out in the vertical furnace geometry, that led to significantly lower yield of the shiny dark colored carbon nitride flakes. However, we found no evidence for the PTI-LiBr phase among the run products from synthesis experiments in the horizontal tube furnace. Instead, the XRD pattern showed a main broad peak at 27.2° 2θ as observed for the larger individual flake samples, along with a shoulder at 28.2° 2θ and weaker sharp reflections at 21°, 24.2°, 32.3° and 27.2° 2θ (Figure 4). We note that similar peaks were observed by Algara-Siller et al. for their ground “TGCN” sample in a synchrotron powder XRD experiment (λ = 0.827127 Å).8 Those peaks do not match the predicted reflections for the TGCN structure. We suggest that they could correspond to an as yet unidentified carbon nitride phase that is formed during the condensation/polymerization reaction from DCDA in the molten salt medium. We note here that several new crystalline phases are now predicted theoretically to occur within the C−N−H system, containing both layered and open-framework structures based on sp² as well as sp³ bonding.9

As shown in previous studies, optical microscopy and SEM images reveal uniformly flat surfaces and terraced edges for the layered gC₃N₄ structures (Figure S1).5,34 HRTEM images for uncleaved thin edge regions of our samples (Figure 5a–d) show crystalline in-plane lattice fringes, and Fourier transformation of the imaged intensity data demonstrate the characteristic 2.6 Å hexagonal lattice spacing typical of graphitic carbon-based nanomaterials.35,36 SAED patterns exhibit multiple spots and the
of separated flakes and powdered material obtained after washing the salt block in water were sent for CNH analysis (Table 1). We note that only \( \sim 80\% \) of the total mass was accounted for in these analyses. Our first response was to potentially assign the missing mass to LiBr/KBr salts that were either incompletely removed by washing, or that were redeposited on the sample surface afterwards. Our EDX analyses carried out during TEM studies did reveal some Br signals scattered randomly thoughout the samples (Figure S2). The XPS survey spectra indicated that \( \sim 1 \) at. % Br was present within the material (Figure 6). We did not detect the presence of K, although its 2p \( \frac{1}{2} \) doublet near 300 eV is notoriously weak, or Li 1s near 55 eV, so that it is not impossible that a very small amount of Br might be incorporated within or attached to the surface of the gC\(_{3}N_{x}H_{y}\) layered sample. Our XPS results also showed that O atoms incorporated at the surface within the bulk might also account for some of the missing mass component. Prior to Ar\(^{+}\) etching \( \sim 15\% \) O was observed at the sample surface. Although the O signal decreased in intensity during ion milling, it was not completely eliminated, with \( \sim 6 \) at. % recorded after this process (Figure 6). That result could indicate a small percentage of O atoms bound within the structure. Algara-Siller et al. reported similar results from their XPS characterization of TGCN, although their chemical analysis obtained using a CHNOS analyzer reported no evidence for oxygen component within the bulk material, and they observed C, N, H values that summed to 98.7 wt %.

Bulk chemical analyses of our samples indicated that 2.3–2.5 wt % H component was present (Table 1). That value is similar to that reported by Ladva et al. for thin film materials prepared using an analogous synthesis approach. Our FTIR spectroscopic results presented below clearly showed that N–H groups were incorporated within the structure. The C/N ratio determined by bulk analysis ranged between 1.21 and 1.24:1 for both thin film and powdered materials prepared in different synthesis experiments (Table 1). This composition is significantly more C-rich than the TGCN material described by Algara-Siller et al., although it agrees with that reported by Ladva et al. for gC\(_{3}N_{x}H_{y}\) films deposited from vapors produced by DCDA reacted in molten LiBr/KBr.\(^{34}\) These results indicate that the structure must contain C–C as well as C–N bonded species within its sp\(^2\)-bonded graphic layers.

Detailed analysis of the C 1s and N 1s XPS signals provides additional information on the local bonding environments and chemical composition of C\(_{3}N_{x}\) materials (Figure 6). The C 1s spectra of crystalline graphite and graphene exhibit an asymmetric peak at 284.5–284.6 eV. Most samples examined

| Table 1. The C\(_{x}N_{y}H_{z}\) Compositions of Carbon Nitride Materials Obtained in Different Synthesis Runs, or Collected from Different Parts of the Reaction Vessel, Determined by Bulk CHN Analysis* |
|-----------------|--------|--------|--------|--------|--------|
| synthesis run 1 (flakes) | C (wt %) | N (wt %) | H (wt %) | total | C/N ratio |
| synthesis run 1 (powder) | 43.03 | 37.09 | 2.40 | 78.5 | 1.23:1 |
| synthesis run 2 (flakes) | 42.54 | 40.68 | 2.38 | 85.6 | 1.22:1 |
| synthesize run 1 (powder) | 42.29 | 40.75 | 2.39 | 85.4 | 1.21:1 |

*The undetermined ~20 wt % could include Br (~7 wt %) and/or O (~9 wt %) components, observed by XPS analysis.
by XPS show a feature near 284.5−285.0 eV that is typically assigned to “adventitious” carbon species at the sample surface, that can include oxygenated components. Graphitic and graphene oxide (GO) samples typically show a strong peak emerging at ∼286.1 eV due to COH and COC species incorporated within the layered materials. N-doped graphene exhibits a main peak at 284.6 eV due to sp2-bonded atoms occurring within the majority C−(C)3 environments,

Figure 6. (a) XPS survey spectra of carbon nitride flake samples before (red) and after (blue) Ar+ ion beam milling to remove surface contaminant species. The first carbon content indicated is for those C atoms that do not occur in “adventitious” component environments, whereas the value in brackets indicates the amount that could be assigned to adventitious carbon, but might also partially correspond to signal from the sample (see text for discussion). (b) C 1s and (c) N 1s spectra showing deconvolution into Gaussian−Lorentzian components to show individual contributions to the overall lineshape.

Figure 7. Experimental and calculated IR and Raman spectra for layered carbon nitride materials. (a) IR absorption spectrum obtained by transmission through a single flake of graphitic carbon nitride (gCxNyHz) material prepared in this study. This is compared with the result of a DFT calculation for a triazine based gCxNyHz layered structure. (b) Raman spectra of a typical gCxNyHz flake obtained with different laser excitation wavelengths (325 nm: black; 514.5 nm: green; 785 nm: blue) compared with DFT calculation for triazine based gC3N4.
while additional peaks emerging near 285.8 (shoulder) and 288.2 eV are coincident with the appearance of a strong N 1s signal at 400.6 eV and a weaker feature at 398.6 eV. A C 1s peak at 285.5 eV was observed to grow as a function of N content in N-doped graphene and was assigned to the emergence of C=N bonding. However, detailed assignment and interpretation of C 1s and N 1s XPS signals for carbon nitride compounds remains an active research problem, while many different authors have either accepted without question, or have modified and added to the suggested assignments, without fully supporting their arguments.

The C 1s data for our gC3N4Hx compound exhibited two clear maxima, along with a shoulder at higher binding energy. These features were best fit using three Gaussian–Lorentzian contributions. The 288.5 eV peak occurs at a similar position to that for sp2-bonded C atoms in amorphous carbon,45 and it is readily assigned to N-C=N=C=N- bonded species that occur within the polymerized triazine rings of the gCN samples. A fully polymerized TGCN structure should give rise to a single C 1s peak near this position, whereas our data indicate three component signals (Figure 6). It would be usual to assign the 285.3 eV component to “adventitious” carbon, and then ignore that contribution in order to obtain a C/N ratio for the gC3N4Hx layer structure. However, that solution would not account for our bulk chemical analyses or the results of Ladva et al., showing that the C/N ratio is greater than expected for the C3N4 stoichiometry. Taking account of those results then indicates that at least part of the 285.3 eV C 1s peak intensity must be due to non-adventitious carbonaceous bonding environments within the layered gC3N4Hx materials. Assignment of the 286.7 eV C 1s component is likewise unclear. This feature was observed in the XPS data of Algarra-Siller et al., where it was suggested to arise from −C≡N (nitrile) functional groups. We observed no evidence for these species by IR spectroscopy (Figure 7a). However, N-doped graphene exhibits a secondary C 1s peak at the same position. We propose that the feature observed in our study could be associated with sp2 C atoms bound to either one or two N atoms within the graphene-like layers of our gC3N4Hx materials.

The N 1s profile is best fit using three Gaussian–Lorentzian components (Figure 6c). The main contribution in the NIs profile at 399.3 eV is readily assigned to sp2-bonded −C≡N=N− environments such as those expected for triazine-based or related C3N4 ring units. A similar peak has been documented for both melanime46 and N-doped graphene.47 The 400.6 eV contribution is assigned to N−(C)3 bridging environments between the rings, as proposed by Algarra-Siller et al., although the intensity ratio between the two features does not correspond to that expected for the TGCN structure. The minor peak at 401.6 eV is typically associated with N−H bonded environments, that have been confirmed to be present in our study by FTIR spectroscopy (Figure 7a).

3.3. Vibrational Spectroscopy and DFT Modelling. Microbeam IR transmission spectra obtained through individual flakes of our gC3N4 materials are dominated by strong in-plane C−N/C=C stretching modes at 1276 and 1590 cm−1 (Figure 7a). A small absorption maximum can be observed between the two main bands at 1420 cm−1, and a weak sharp peak occurs at ~700 cm−1. We note an unusual “dip” in the IR spectra just above the intralayer vibrational modes that resembles the Fano resonance lineshape observed for graphene and related 2D materials, caused by coupling between the IR-active vibrational modes and a broad spectrum of electronic excitations. Our IR spectra also exhibit absorption maxima at 3179 and 3311 cm−1 that resemble the N−H stretching modes observed for crystalline PTI-Br and PTI-Cl compounds (Figure S3). However, we found no evidence for the presence of PTI phases from our XRD or TEM studies, indicating that the N−H stretching vibrations are intrinsic to the gC3N4Hx flakes. Another possibility might be that these modes are related to O−H stretching vibrations, that could be associated with the small proportion of O atoms concentrated at the surface and detected by XPS measurements. However, the N−H absorptions are different from the broad O−H stretching feature observed for GO materials (Figure S3).

We also obtained microbeam Raman scattering spectra for our gC3N4Hx samples using laser excitation wavelengths ranging from the near-IR (785 nm) to visible (514.5 nm) and UV (325 nm) (Figure 7b). The appearance of the spectra depends on the excitation wavelength, as has been observed previously for C-graphite, graphene and gC3N4 samples. It is known that different vibrational modes can exhibit resonant behaviour with various parts of the electronic absorption manifold, and the resulting spectra can probe different spatial extents of vibrational coherence and structural ordering within the samples. In general, UV excitation tends to favour more localized vibrations whereas near-IR Raman studies probe responses over longer length scales, that could include structurally disordered regions. Our UV-Raman data exhibit main peaks near 1700 and 1300 cm−1 that resemble the “G” and “D” modes of graphite and graphene materials, and they can be assigned to C−N and/or C=C stretching vibrations of the sp2-bonded network. As the excitation wavelength moves into the near-IR range the phonon response becomes broadened and loses its intrinsic structure (Figure 7b). Our data indicate wavelength dispersion occurring for the fundamental (1200−1700 cm−1) stretching modes as well as the Raman active overtone in the 2500−3500 cm−1 region. N-doped graphene and graphite samples exhibit IR absorption bands in the 1200−1700 cm−1 region that resemble the G and D band profile observed in the Raman spectra of structurally disordered or nanocrystalline pure carbon materials. Rodil et al. have analyzed the effects giving rise to the IR band profiles obtained from different types of amorphous C3N4 films.

To aid our understanding of the vibrational spectra and structural nature of the layered carbon nitride materials prepared here, we carried out DFT calculations for series of models representing structural motifs that might be present among the gC3N4Hx layered phases (Figures 7a,b; 8a,b). We began with a single sheet of ideal triazine based gC3N4 (TGCN), that was fully relaxed into its buckled conformation. The simulated IR spectrum for this material shown in Figure 7a displays three main peaks in the C−N stretching region between 1200 and 1600 cm−1. The predicted strong in-plane C=N modes agree generally with the dominant IR bands observed for our sample, although the N−H stretching features are absent. We have compared our simulated spectrum with IR data for nanocrystalline H-free gC3N4 reported by Kouvetakis et al. That compound exhibits a single broad band between 1100 and 1700 cm−1, with its main peak at approximately 1300 cm−1, covering the range of our DFT results. Broadening in the experimental spectrum might occur due to finite particle and domain size effects, along with local differences in layer buckling and stacking patterns within the nanoparticles. However, the gC3N4 layered model predicts the occurrence of strong Raman active modes at 1040-1050 cm−1 associated with symmetric
breathing vibrations of the C₃N₃ triazine rings, that are absent from our data (Figure 7b). That observation caused us to examine the nature of the vibrational mode symmetries in more detail, and to construct further models that could account for the observed spectra.

Analysis of the IR-active normal modes of our simulated gC₃N₄ (TGCN) shows that the 1566 cm⁻¹ peak arises from in-phase stretching of two parallel C−N bonds within the same triazine ring, combined with parallel out-of-phase movement of the remaining N atom that connects them together. For convenience, we refer to these two-coordinated N atoms within the triazine ring as N(2), and denote the bridging three-coordinated N atoms that link the triazine units as N(3). The ~1314 cm⁻¹ vibrations involve the same sets of bonds and atoms, but the C−N(2) stretching is out-of-phase. The ~1220 cm⁻¹ peak is a doublet, constituting in- and out-of-phase stretching combinations of the C−N(3) bonds.

We further developed our modelling analysis by assessing how to introduce N−H into our model structure, while removing the high local symmetry giving rise to the 1044 cm⁻¹ Raman active mode (Figure 8a,b). A suitable model could be found for the gC₃N₄ framework by introducing a neutral H atom that forms a bond with a N(2) atom in the triazine ring (Figure 9). We note that this structural model does not imply any "protonation" of the gC₃N₄ layers. Instead, it corresponds to generation of a chemically reduced gC₃N₄,H structure, with the reducing H-atom bonded to the triazine ring and donating an unpaired electron to the system. Such radical species are well documented to occur among various N-containing compounds, including carbon nitrides. Within such a model, the unpaired electron could occur as a polaronic state displaying localized paramagnetism, or might contribute to a conduction band. Our DFT calculations predict that a dispersionless polaronic state constitutes the most stable configuration, ~1 eV below the CB minimum, with the spin density localized in the ring to which the H is bound. We found that these polaronic defects were stable as the H-content was varied, from a single H-atom per 12 triazine rings, to a maximum in which all the triazine rings bear 1 H⁺ atom. A reduced gC₃N₄ material with such a gC₃N₄,H stoichiometry would constitute a system with a very high density of localized polarons (1.78 × 10⁻⁶ e/μm²). A metallic conductive state was predicted to occur at only ~0.26 eV above the ground state, indicating that a temperature-dependent equilibrium between localized paramagnetic and conductive states might be established for gC₃N₄,H doped materials. We describe the results of EPR experiments and magnetic susceptibility studies to investigate the magnetic properties of our synthesized gC₃N₄,H materials below.

Although the reduced gC₃N₄,H model yields a system that could satisfy the constraints of the experimental IR and Raman observations, our analytical measurements now indicate that the prepared materials were significantly more C-rich than the C₃N₄ stoichiometry. To introduce the effects of this observation into our DFT studies, we constructed a layered model with C₄N₃ stoichiometry, with triazine (C₃N₃) rings linked by bridging C(5) rather than N(3) atoms (Figure 1c). Although gC₄N₃ structures have been predicted theoretically to occur, no such material has been demonstrated experimentally. These would have unsaturated valency containing unpaired electrons, and are...
predicted to exhibit interesting and potentially useful “half-metallic” properties. We calculated the IR and Raman spectra for gC$_3$N$_4$ layers, as well as the “reduced” form, gC$_3$N$_4$H. The addition of neutral H to form gC$_3$N$_4$H motifs generates a charge-balanced system with no unpaired electrons. Although the simulated Raman spectrum for gC$_3$N$_4$ displays the characteristic symmetric ring-breathing mode at $\sim$1040 cm$^{-1}$, forming gC$_3$N$_4$H removes this feature as observed previously for our gC$_4$N$_3$H model structure.

We then examined further structures containing triazine (C$_3$N$_3$) rings linked by both N(3) and C(3) atoms, that could provide models for graphene-like sheets with gC$_3$N$_4$ compositions (Figure 8a,b). We also included H in the structures investigated. Our first model represented gC$_3$N$_4$H$\cdots$C$_3$N$_4$, with added H-atoms bonded to ring N(3) atoms on 50% of the rings, and H--N bonds aligned to point towards an N(3) bridging atom across the layer “void”. That stoichiometry corresponds to a valency-saturated system, without unpaired electrons. Analysis of the atomic displacements for the calculated vibrational modes shows that the two Raman active ring-breathing modes have very low intensity, and might not be detected experimentally. Our final model represented a gC$_3$N$_4$H$\cdots$C$_3$N$_4$H layer, in which all the rings have an H atom attached and N--H groups occur in two different environments, pointing across the voids at either bridging N(3) or C(3) atoms. That structure definitely eliminates the occurrence of a $\sim$1040 cm$^{-1}$ Raman mode, while the simulated IR spectrum reveals a double peak in the N--H stretching region at 2880 and 3020 cm$^{-1}$, as is observed experimentally (Figure 7a).

3.4. EPR Spectroscopy, Paramagnetic Defects and Magnetic Properties. We carried out EPR studies on single flakes of our gC$_3$N$_4$H$_x$ material at room temperature to investigate the presence of localized paramagnetic electron species (Figure 10). Spectra were recorded from samples of gC$_3$N$_4$H$_x$ flakes over a full rotation ($360^\circ$) in 20$^\circ$ increments or 10$^\circ$ near the turning points. The spectra at 0$^\circ$ and 180$^\circ$ correspond to the sample oriented parallel ($\pm 5^\circ$) to the applied magnetic field, whereas spectra at 90$^\circ$ and 270$^\circ$ correspond to the field held perpendicular ($\pm 5^\circ$) to the field. The dependence of g-values as a function of the relative orientation ($\pm 5^\circ$) between the sample flakes and the applied magnetic field is also shown. The g values showed a small dependence on orientation, varying between $g = 2.0035$ for parallel and 2.0034 for perpendicular orientation. However, the periodic behaviour observed throughout a full rotation indicates orientation-dependent g-anisotropy. The EPR linewidth shows a more marked dependence on the sample orientation in the applied magnetic field with values of 0.7 and 1.0 mT recorded for parallel versus perpendicular orientations. It is likely that the linewidth is due to unresolved hfc interactions ($A$) between the unpaired electron and the neighbouring magnetic nuclei (N and H). The red line in Figure 10b,c represents a fit according to: $A^2 = A_1^2 \sin^2 \theta + A_2^2 \cos^2 \theta$, where $A_1$ and $A_2$ are the linewidths at 0$^\circ$ and 90$^\circ$ and $\theta$ is the angle between the applied magnetic field and a vector normal to the planar sheets, that is expected to contain the $p_z$ orbital of a localized unpaired electron species, as found in organic radical species containing N-atoms including neutral flavin radicals. We note that the EPR signal saturates readily at room temperature, that is also typical for organic radicals. In contrast to a recent study on a related carbon nitride material, no electron spin echo was observed. This indicates that $T_2^*$ is short in the system studied here that could indicate a high local concentration of spins.

We carried out DFT calculations for a molecular model based on a reduced triazine unit (C$_3$N$_3$H) surrounded by C$_3$N$_4$ groups to study the potential distribution of unpaired spin densities on atoms surrounding the C$_3$N$_4$ ring (Figure 10e, Table S1). The unpaired spin densities were found to be concentrated on the 3 C atoms. However, given that 99% of these are $^{12}$C nuclei, this has no appreciable impact on the observed EPR spectrum.

![Figure 10](image-url)
However, the 3 N atoms (>99% 14N) in the reduced ring also have significant spin densities. These are predicted to have axial Hfc’s with the largest component perpendicular to the nearly planar gC3N4 units, consistent with our observation of the largest EPR linewidths when the plane of the flake is perpendicular to the magnetic field (Figure 10, Table S1). In related organic systems such as triphyrins, tautomerism allows the proton to hop between the three neighbouring N atoms, giving an unusually broad NMR spectrum at room temperature.6 Analogous rapid H* transfer could also occur in the radicals proposed here, reducing the N Hfc’s by approximately a factor of three and potentially contributing to the fast T2.

4. CONCLUSIONS

We prepared layered carbon nitride samples from DCDA in a molten salt (LiBr/KBr) flx environment sealed glass ampoules at 400–600 °C. Dark-colored shiny flakes were either deposited on the walls of the container or formed within the molten salt. Synthesis of analogous materials under similar conditions have been reported previously to yield either crystalline triazine-based graphitic carbon nitride8 (TGCN) gC3N4 or nanocrystalline flakes with much higher C/N ratio (~1.2:1) deposited from the vapor phase.34 Our results agreed more closely with the second study. However, our experiments also indicate that the reaction products are highly dependent on details of the synthesis conditions, including furnace geometry. Our XRD and HRTEM results confirmed the graphitic structure of the flakes. The presence of moiré patterns and multiple diffraction spots in our HRTEM studies indicated overlapping layers with different orientations or domain structures. Bulk chemical analyses showed a composition that was significantly more C-rich than gC3N4, with C/N ratio ~1.22 compared with 0.75 expected for gC3N4. That result corresponds to observations made by Ladva et al.,34 but differs from the results of Algarra-Siller et al.8 for materials prepared using similar techniques. Analysis of the C 1s and N 1s XPS signals indicated that sp2-bonded domains with mixed −C≡C− and −C≡N= bonding is likely to present among the layered compounds. Such materials are expected to have interesting and potentially useful electronic and magnetic properties. The presence of H* defects could lead to localized unpaired electron spins or conducting domains. Taken together, the results from this and previous studies indicate that a range of gC3N4 solid solutions or nanocomposite materials with tunable electronic properties might be produced by scalable thin film production techniques, by adjustment of the synthesis and deposition conditions.

ASSOCIATED CONTENT

# Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpcc.8b07972.

Additional data including IR, Raman spectroscopy, SEM, EDX and optical microscopy images (PDF)

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Notes

The authors declare no competing financial interest.

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