

Fast Exfoliation and Functionalisation of 2D Crystalline Carbon Nitride by Framework Charging

Jingjing Jia, Edward R. White, Adam J. Clancy, Noelia Rubio, Theo Suter, Thomas S. Miller, Paul F. McMillan, Veronika Brázdová, Furio Corà, Chris A. Howard, Robert V. Law, Cecilia Mattevi,* and Milo S. P. Shaffer*

Abstract: 2D layered graphitic carbon nitride nanosheets offer tunable electronic and chemical properties. However, exfoliation and functionalisation of gCN for specific applications remains challenging. We report a scalable one-pot reductive method to produce solutions of single and few layer 2D gCN nanosheets with excellent stability in a high mass yield (35%) from polytriazine imide. High resolution imaging confirms the intact crystalline structure and identifies an AB stacking. The first successful deliberate organic functionalisation of dissolved gCN is illustrated, providing a general route to adjust their properties.

Graphitic carbon nitride (gCN) has triggered tremendous interest due to its 2D structure, analogous to graphene, but with complementary characteristics.^[1] In particular, it offers inherent semiconductivity with tunable band gap and optical absorption,^[2] whilst the different chemical valences of N and C create empty sites within the layers.^[3] Monolayer/few layer carbon nitride nanosheets (FL-CNs) have been isolated as a new family of 2D layered materials, motivated by their unique photocatalytic activity.^[4] Several methods have been adopted to synthesize FL-CNs of various thicknesses/sizes.^[5] Unfortunately, many of these processes damage the structure, altering the properties of interest; they are also time-consuming and provide low yields and dilute suspensions, and most work has focused on the disordered heptazine-based gCNs. Polytriazine imide (PTI) has been previously synthesized and characterized using a number of bottom-up approaches.^[6] PTI is more crystalline than its heptazine-based counterpart, containing genuine planar layers of imide-bridged triazine units,^[7] and its exfoliation into high quality 2D FL-CN crystals is, therefore, attractive. Achieving a non-damaging preparation of 2D few-layered PTI (FL-PTI) in a high yield is still in its infancy, although slow dissolution has recently been reported.^[8] Moreover, while covalent functionalisation is a vital tool in tailoring the

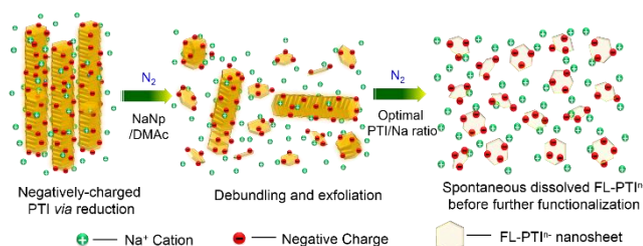
properties of nanomaterials,^[9] to date there has been little direct covalent functionalisation of PTI.

This paper demonstrates a simple, one-pot exfoliation, dissolution, and optional functionalisation of FL-PTI via reduction. Reductive charging has been used previously to dissolve a variety of 2D nanomaterials,^[10] via metal-ammonia solutions and organic charge transfer agents (CTA). The use of sodium naphthalide (NaNp) dissolved in N,N-dimethylacetamide (DMAc) was recently found to be especially effective for the dissolution and functionalisation of single wall nanotubes in a single step.^[14] Here the methodology is adapted to gCNs, specifically PTIs.

Successful exfoliation of PTI was achieved by framework charging process (Fig. 1). Sodium is used as the electron source to form naphthalide ions, which act as a CTA. DMAc is an excellent room temperature solvent for naphthalene/ide and anionic nanocarbons, and can be expected to be a good solvent for FL-PTI.^[11] NaNp/DMAc solution has a characteristic green color which simplifies reaction monitoring. NaNp/DMAc solutions were added into dried PTI powders at controlled stoichiometry. The reduction and exfoliation process was observed by the color change from the initial brown PTI suspension to dark green after addition of NaNp/DMAc, before finally forming a brilliant orange dispersion of FL-PTI (FL-PTIⁿ⁻) within minutes (Fig. S1). After removing the remaining insoluble PTI fragments by centrifugation (5000 g, 30 min, Fig. S2), a homogeneous golden FL-PTIⁿ⁻ solution with a concentration up to 1.2 mg·mL⁻¹ was obtained (Fig. S1c), which was stable under N₂ for >1 year (Fig. S3a). Deposited FL-PTIⁿ⁻ nanosheets display hexagonal geometry with a height of 1-2 nm (Fig. S3), indicating that they comprise only a few PTI layers, based on a 3.52 Å layer thickness.^[6b] The negative charges from the naphthalide are likely to be rapidly transferred to the PTI, due to the high reduction potential of naphthalide (ca. 3.0 eV vs SHE);^[12] accelerated by the small size of the PTI platelets and the intrinsic pores.^[13] The partially dissociated Na⁺ counterions leave a net unscreened negative charge on the PTIⁿ⁻ (Fig. S4a), leading to short range Coulombic repulsions and hence exfoliation of PTI into solvated FL-PTIⁿ⁻ sheets (Fig. 1), analogous to reduced nanocarbons^[10a, 14] and transition metal dichalcogenides.^[10b] Upon air exposure, the FL-PTI reaggregate slowly (~2 months, Fig. S5b); the reduced rate compared to charged SWCNTs,^[14b] likely relates to the lower aspect ratio and localisation/low mobility of the charges on the framework.^[15]

[*] J. Jia, E. R. White, A. J. Clancy, N. Rubio, M. S. P. Shaffer, Dept. Chemistry, Imperial College London, London, SW7 2AZ, UK.
E-mail: m.shaffer@imperial.ac.uk
C. Mattevi
Dept. Materials, Imperial College London, London, SW7 2AZ, UK
E-mail: c.mattevi@imperial.ac.uk
T. Suter, T. S. Miller, P. F. McMillan, V. Brázdová, F. Corà, Dept. Chemistry, University College London, London, WC1H 0AJ, UK.
C. A. Howard
Dept. Physics and Astronomy, University College London, London, WC1E 6BT, UK.

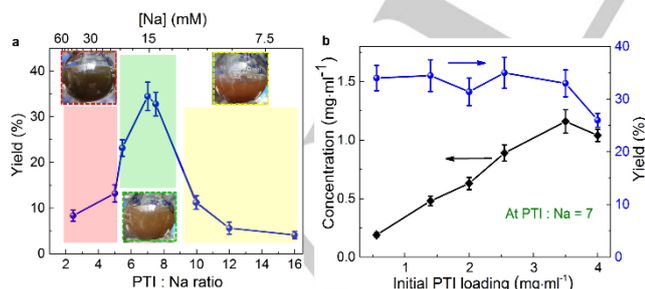
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81 **Figure 1.** Schematic of charging and exfoliation of PTI.

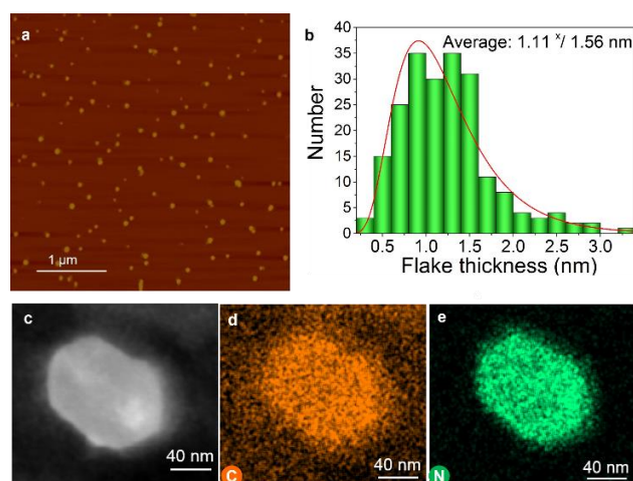
82 The charging ratio (molar [PTI framework atoms]:Na (ESI),
 83 weighted $M_w(\text{PTI}) = 13.14$) and initial PTI loading (mg PTI/
 84 mL of DMAc) are two vital factors affecting the exfoliation,
 85 controlling both the yield (mass fractions of PTI) and
 86 concentration of solubilised FL-PTIⁿ⁻. Increasing the degree
 87 of charging (*i.e.* lower PTI:Na), at a static PTI loading initially
 88 led to an improved yield of FL-PTIⁿ⁻ (4.1 wt% to 34.5 wt%)
 89 due to the enhanced Columbic repulsion (Fig. 2a). However,
 90 further increasing charge (PTI:Na < 7) reduced the yield.
 91 Similar effects have been observed in charged nanocarbon
 92 solutions, attributed to Na⁺ condensation and charge
 93 screening.^[11, 16] The optimum Na concentration for exfoliation
 94 of PTI is 15 mM (*i.e.* 7:1 PTI:Na for 1.4 mg·mL⁻¹, Fig. 2a),
 95 comparable to ~10 mM identified for the exfoliation of Na-
 96 reduced graphite of similar geometry.^[16] At the highest
 97 charge regimes, the charge on the PTI saturates (at PTI:Na
 98 ratio of ~5), as observed by the green tinge of unreacted
 99 NaNp (Fig. 2a). On varying the initial PTI loading (Fig. 2b),
 100 the concentration of dissolved FL-PTIⁿ⁻ scales linearly, giving
 101 a consistent yield between 31–35 wt%, indicating that there
 102 may be an intrinsically soluble portion of the starting material.
 103 The residue, isolated after centrifugation may contain defects
 104 that bind the layers; indeed, qualitatively, the undissolved
 105 residue appears disordered by SEM (Fig. S2c). The
 106 maximum concentration of FL-PTIⁿ⁻ is ~1.2 mg·mL⁻¹ from a
 107 PTI loading of 3.5 mg·mL⁻¹. Further increases in PTI loading
 108 did not increase the concentration, indicating the solution is
 109 saturated (Fig. 2b).
 110



111 **Figure 2.** (a) FL-PTIⁿ⁻ yield versus PTI:Na ratio/[Na], 1.4 mg·mL⁻¹ PTI.
 112 Yellow, green and red rectangles correspond to low, to high [Na]
 113 respectively with inset photographs showing resultant FL-PTIⁿ⁻
 114 dispersions. (b) Effect of initial PTI loading on the concentration/yield of
 115 FL-PTIⁿ⁻ dissolution (7:1 PTI:Na ratio).
 116

117 Aqueous FL-PTI dispersions are desirable for
 118 environmentally benign processing; however, pristine PTI

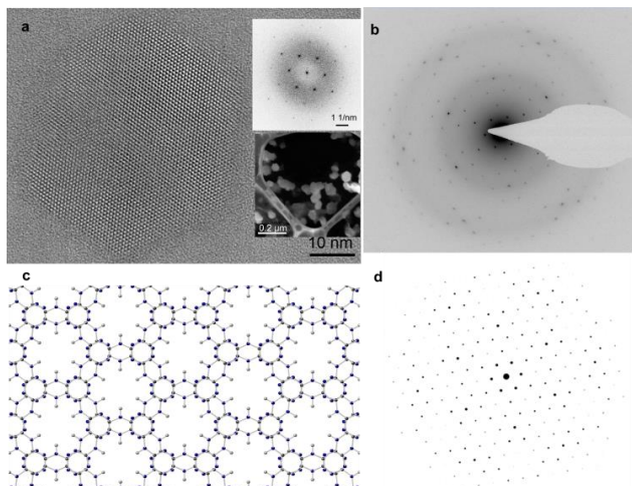
119 are poorly soluble in water due to their strong interlayer
 120 interactions (6 h probe sonication was found to give a
 121 concentration <0.2 mg mL⁻¹). The framework charging
 122 process overcomes the strong interlayer interactions and
 123 accelerates solubilisation in DMAc, allowing the removal of
 124 the intrinsically insoluble fraction of the pristine PTI. The FL-
 125 PTIs can then be recovered from the DMAc by solvent
 126 exchange and the resultant FL-PTI can be transferred into
 127 water by solvent exchange, reaching a saturated
 128 concentration of 3.5 mg·mL⁻¹. Atomic force microscopy
 129 (AFM) confirms an excellent dispersion of FL-PTI in water
 130 (Fig. 3a), with corresponding heights of 0.33–3.2 nm (avg.
 131 1.11 nm, Fig. 3b and S6), suggesting that the nanosheets
 132 mostly comprise ~3 PTI layers, although some monolayers
 133 are present. Representative energy dispersive X-ray
 134 spectroscopy (EDX) maps show uniform dispersion of C and
 135 N throughout the whole hexagonal area of the exfoliated FL-
 136 PTI nanosheet (Fig. 3c–e). High-resolution transmission
 137 electron microscopy (HRTEM) micrographs show intact PTI
 138 crystallites with regular hexagonal geometry and clear facets
 139 (Fig. 4a). Notably, no defect holes or dislocations were
 140 observed, confirming the non-destructive nature of the
 141 framework charging exfoliation, as well as the high quality of
 142 the starting material. The fast Fourier transform (FFT) of the
 143 unfiltered HRTEM image shows a hexagonal lattice,
 144 demonstrating a single crystal exfoliated FL-PTI (Fig. 4a and
 145 further examples in S9).^[17] The minimum reciprocal lattice
 146 vector, G_{min} , is 1.4 nm⁻¹, giving a lattice constant $a = (2/\sqrt{3})G_{min} = 8.5 \text{ \AA}$, consistent with the reported values from Br
 147 intercalated PTI.^[15] Two possible stacking models can be
 148 considered for the FL-PTI: AB stacking with aligned voids
 149 forming c-axis channels, and AC stacking without channels
 150 in two adjacent layers (Fig. S10a). Comparing simulated
 151 electron diffraction patterns of these two models with the
 152 experimental selected area electron diffraction (SAED) data,
 153 the AB stacking structure is the better fit for the FL-PTI
 154 nanosheets (Fig. 4b–d and S10b).
 155



156

157 **Figure 3.** (a) AFM image of FL-PTI nanosheets. (b) PTI thickness
 158 histogram ($n > 200$). Mean value is derived from a lognormal distribution.
 159 (c) STEM image and EDX elemental maps of C (d) and N (e) on a FL-PTI

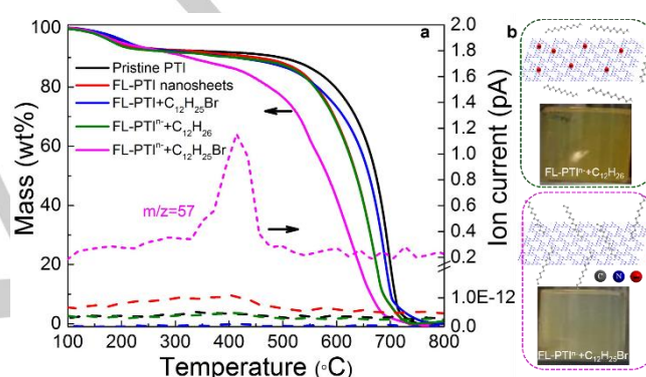
160 nanosheet. The background C signal in (d) is due to the carbon TEM
161 support.



162
163 **Figure 4.** (a) HRTEM image of a FL-PTI nanosheet. Top inset: FFT of the
164 HRTEM image, showing a single crystal hexagonal structure. Bottom
165 inset: STEM image, indicating the layered structure of FL-PTI (b) SAED
166 pattern of a FL-PTI nanosheet. (c) Schematic of the AB stacking in a
167 bilayer. (d) Simulated electron diffraction pattern from the AB crystal
168 structure.

169 The FL-PTIⁿ⁻ synthesised in DMAc solution provides a
170 versatile platform for covalent functionalisation of PTI,
171 comparable to negatively charged graphene and boron
172 nitride nanotube counterparts.^[16, 18] Pristine PTI and FL-PTI
173 nanosheets are thermally stable up to ~700 and 670 °C,
174 respectively (Fig. 5a and Figure S12); the slightly depressed
175 decomposition temperature for FL-PTI reflects its few-
176 layered character. Functionalisation can be performed *via*
177 simple addition of an alkyl halide to the reduced
178 nanomaterial. After functionalisation by reaction with dodecyl
179 bromide, a 10 wt% mass loss can be observed in
180 thermogravimetric analysis/evolved-gas mass spectrometry
181 (TGA-MS), relative to controls; the weight loss correlates
182 with a *m/z* peak at 57, attributed to C₄H₉⁺ from the grafted
183 C₁₂H₂₅ alkyl chain (Fig. 5a). During the alkylation reaction,
184 the solution changes from a clear golden, to a turbid pale
185 yellow appearance; as the depletion of the negative charges
186 progresses, the Coulombic stabilization is lost, leading to
187 agglomeration and increased light scattering (Fig. 5b, S1c
188 and S12). X-ray photoelectron spectroscopy (XPS)
189 measurements confirmed the covalent attachment of alkyl
190 chains to the PTI structure. Core level N 1s spectra can be
191 divided into two components: 398.6 eV corresponding to C-
192 N=C groups and 400.9 eV attributed to secondary and
193 tertiary amines (NH/N-(C)₃) ((Figure S13 and S14).^[19] An
194 increase in the NH/N-(C)₃ peak was observed for the
195 dodecyl-functionalised PTI compared to both FL-PTI and
196 physisorption controls (Table S1). Quantitatively, the XPS
197 data indicate 74 framework atoms per alkyl chain, comparing
198 favourably with the TGA estimate of 101 atoms per chain.
199 Controls of air quenched FL-PTI mixed with C₁₂H₂₅Br and
200 FL-PTIⁿ⁻ with unreactive C₁₂H₂₆ showed similar TGA curves

201 to the unfunctionalised FL-PTI, precluding a contribution
202 from physisorption. The XPS measurements also exclude
203 physisorption since the core level Br 3d was not observed in
204 the grafted products, although it was visible in positive
205 controls at relevant concentrations (Figure S15). Reaction
206 with a shorter alkyl chain was also investigated; when
207 charged FL-PTI reacted with octylbromide a mass loss of 12
208 wt% and the corresponding *m/z* peak at 57 were observed
209 (Figure S16). XPS showed a similar increase in the NH/N-
210 (C)₃ peak as observed for the dodecyl-functionalised PTI
211 (Figure S17 and Table S1). In this case, XPS indicates one
212 alkyl chain every 47 PTI atoms, closely matching the TGA
213 estimate of one chain per 55 framework atoms. Given the
214 uncertainties in these measurements, the agreement is
215 excellent, and provides direct evidence of grafting to the PTI
216 layers.



217
218 **Figure 5.** (a) TGA and TGA-MS (dashed) of pristine PTI and alkylated FL-
219 PTI. Inset shows 100-400 °C TGA region. (b) Photographs of control
220 sample (top) and FL-PTI functionalised with dodecyl bromide (bottom).

221 The observed change in the XPS nitrogen components upon
222 functionalization can be attributed to the attachment of alkyl
223 chains to the nitrogen of the triazine ring. Density functional
224 theory (DFT) calculations suggest that the sodium ion
225 bridges two triazine rings, with the extra electron delocalized
226 over those two rings (Figure S19). The extra electron
227 appears to be primarily located on the nitrogen atoms of the
228 ring, which may therefore be most susceptible to react with
229 the alkyl bromide molecule (Scheme S2)

230 In summary, framework charging provides a new, simple
231 route for exfoliation and functionalisation of PTI nanosheets,
232 *via* NaNp/DMAc reduction. By avoiding damage, the intrinsic
233 properties of the PTI structure can be retained and HR
234 images indicate highly exfoliated hexagonal, crystalline FL-
235 PTI nanosheets, averaging 1.1 nm thick (~3 layers) with AB
236 stacking. The as-prepared FL-PTI solutions had a yield of 35
237 wt%, with excellent stability

238
239 y. Stabilized dispersions of FL-PTIs are useful feedstocks for
240 a wide range of promising multifunctional applications. The
241 small flake size is particularly relevant to potential applications
242 in (electro)catalysis and photochemistry.^[20] The FL-PTIⁿ⁻
243 was successfully functionalised with alkyl chains *via* the

244 framework charge, suggesting a route to a wide range of
245 functionalised species to modulate surface chemistry and
246 functional properties.

247 **Acknowledgements**

248 Financial support within framework of the European Flagship
249 (grant agreement No. 696656–GrapheneCore1). CM
250 acknowledges the award of a Royal Society University Re-
251 search Fellowship by the UK Royal Society and the EPSRC
252 award EP/K033840/1. Work by TS, TS, VB, FC, CAH and
253 PFM at UCL was also supported by EPSRC grant
254 EP/L017091/1.

255 **Conflict of interest**

256 The authors declare no conflict of interest.

257 **References**

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