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Developing extended visible light responsive polymeric carbon nitrides for photocatalytic and photoelectrocatalytic applications†

Sanjit Mondal,^a Gabriel Mark,^a Liel Abisdris,^a Junyi Li,^a Tirza Shmila,^a Jonathan Tzadikov,^a Michael Volokh,^{ID, a} Lidan Xing^{ID, b} and Menny Shalom^{ID, *a}

Polymeric carbon nitride (CN) has emerged as an attractive material for photocatalysis and photoelectronic devices. However, the synthesis of porous CNs with controlled structural and optical properties remains a challenge, and processable CN precursors are still highly sought after for fabricating homogenous CN layers strongly bound to a given substrate. Here, we report a general method to synthesize highly dispersed porous CN materials that show excellent photocatalytic activity for the hydrogen evolution reaction and good performance as photoanodes in photoelectrochemical cells (PEC): first, supramolecular assemblies of melem and melamine in ethylene glycol and water are prepared using a hydrothermal process. These precursors are then calcined to yield a water-dispersible CN photocatalyst that exhibits beneficial charge separation under illumination, extended visible-light response attributed to carbon doping, and a large number of free amine groups that act as preferential sites for a Pt cocatalyst. The optimized CN exhibits state-of-the-art HER rates up to $23.1 \text{ mmol h}^{-1} \text{ g}^{-1}$, with an AQE of 19.2% at 395 nm. This unique synthetic route enables the formation of a homogeneous precursor paste for substrate casting; consequently, the CN photoanode exhibits a low onset potential, a high photocurrent density and good stability after calcination.

Introduction

Polymeric carbon nitride (CN) has emerged as an attractive material for photocatalytic and photoelectronic devices, such as photoelectrochemical cells (PEC), light-emitting diodes (LED), and biosensors.^{1–5} Some of the key factors determining the photoactivity of CN materials are their structural properties

New concepts

Photocatalytic and photoelectronic devices, such as photoelectrochemical cells (PEC), require complex materials that combine a range of structural (including morphology and specific surface area) and optoelectronic properties (such as band gap, electronic band positions, and charge separation under illumination). Polymeric carbon nitride (CN) has emerged as an attractive material for these applications thanks to its low cost its overall desirable and highly tunable properties. Here we introduce a general approach for the rational synthesis of porous CN materials with state-of-the-art photocatalytic activity for the hydrogen evolution reaction (HER) and excellent performance as photoanode materials in photoelectrochemical cells: we integrated strategies of supramolecular chemistry, hydrothermal treatment, and solvent mixtures to prepare a paste composed of monomers that affords CN after calcination. The presence of EG in the condensation process leads to C-doping, high surface area, and a large number of free NH_2 groups in the final CNs; as a result, these materials exhibit a unique electronic structure (including homojunction and C-doping), extended light-harvesting properties, excellent dispersibility in water, and high cocatalyst loading capacity. The as-obtained homogenous paste precursor can be directly used for electrode preparation.

(including morphology and specific surface area) and their optical and electronic properties (such as band gap, electronic band positions, and charge separation under illumination).^{6–11} For heterogeneous photocatalysis, the CN powders must also have good dispersibility in water, which is both the dispersant and the reactant in the hydrogen evolution reaction (HER); an additional pivotal requirement is the availability of free sites favoring the anchoring of co-catalysts.^{12,13} As for photoelectronic devices, the precursors should be amenable to uniform casting and must remain bound to the substrate during the ensuing thermal polymerization. Finding a simple synthetic path toward a material that encompasses all these features is highly desired. Recently, 2,5,8-triamino-tri-s-triazine (melem), the repeating unit in CN, has been used for the synthesis of CNs for its high thermal stability and low sublimation.^{14–16} The reaction of melem with other small monomers allows the CN properties to be adjusted.^{17,18} However, the insolubility of

^a Department of Chemistry and Ilse Katz Institute for Nanoscale Science and Technology, Ben-Gurion University of the Negev, Beer-Sheva 8410501, Israel.

E-mail: mennysh@bgu.ac.il

^b School of Chemistry, South China Normal University, Guangzhou 510006, China

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melem in most solvents hinders the construction of supramolecular structures in high yield.

The design of supramolecular assemblies composed of CN monomers as reactants is a powerful tool for synthesizing tailored CN materials.^{19–22} The supramolecular approach enables good control over the morphology of CN and over its optical, electronic, and catalytic properties.^{23–26} The formation, uniformity, and reproducibility of the self-assembly rely strongly on the monomers having adequate solubility in a given solvent: the solubility of the monomers will direct their interactions, thus allowing a molecular ‘blueprint’ to evolve into the morphology on the macroscale. To date, most studies have been done in water, where monomers have limited solubility.^{18,27,28} The construction of complex structures of several monomers and solvents may pave the path toward novel CN materials. As for photoelectronic devices, the reproducible preparation of a paste with high control over its composition remains challenging.

Here we report a general approach to synthesizing a highly photoactive CN as a photocatalyst for HER and as a photoanode in photoelectrochemical cells. To do so, we construct supramolecular assemblies based on a combination of melem and melamine in a mixture of ethylene glycol (EG) and water. We hypothesized that the hydrogen-bonding EG would favor the self-assembly of the monomers and also allow casting *via* doctor-blading.

To optimize the assembling the monomers and the organization process, we used a hydrothermal technique to improve solubility. Theoretical and experimental data show that EG strongly improves the solubility of the monomers by binding to the melem and melamine units, leading to the formation of a homogeneous precursor paste of carbon-rich monomers.

Upon calcination, a highly porous CN with extended light absorption and high concentration of NH₂ surface groups is obtained. The homogeneous precursor paste may be directly used to fabricate CN films on conductive substrates for use as photoelectrochemical water-splitting photoanodes.

Results and discussion

Nanostructured polymeric carbon nitrides (CNs) were synthesized in a three-step synthesis procedure as shown in Fig. 1 (a detailed synthesis procedure is provided in the ESI[†]).

Synthesis of CN precursors

The first two synthetic steps involve the preparation of the precursors that will be thermally polymerized into the final CN materials, as discussed later. First, melem (mlm) was synthesized by the thermal polymerization of melamine (M) for 12 h at 400 °C in air.²⁹ Supramolecular assemblies of melamine and melem at a 1:1 molar ratio were prepared by dispersing these two monomers in a water/EG mixture, followed by overnight shaking. To improve the solubility of the monomers and optimize the formation of supramolecular assemblies, the dispersion was heated at 160 °C for 20 h in a PTFE-lined autoclave. After that, the final supramolecular assemblies were collected by centrifugation; they are labeled as P_{M+mlm} (control precursors were prepared using only melamine and melem and denoted as P_M and P_{mlm} respectively). These supramolecular assemblies were directly used as precursors for (a) the rational synthesis of a powder photocatalyst or (b) the preparation of a photoelectrode. Density functional theory (DFT) simulations reveal that the interaction energy (E_{int}) in M-EG-mlm

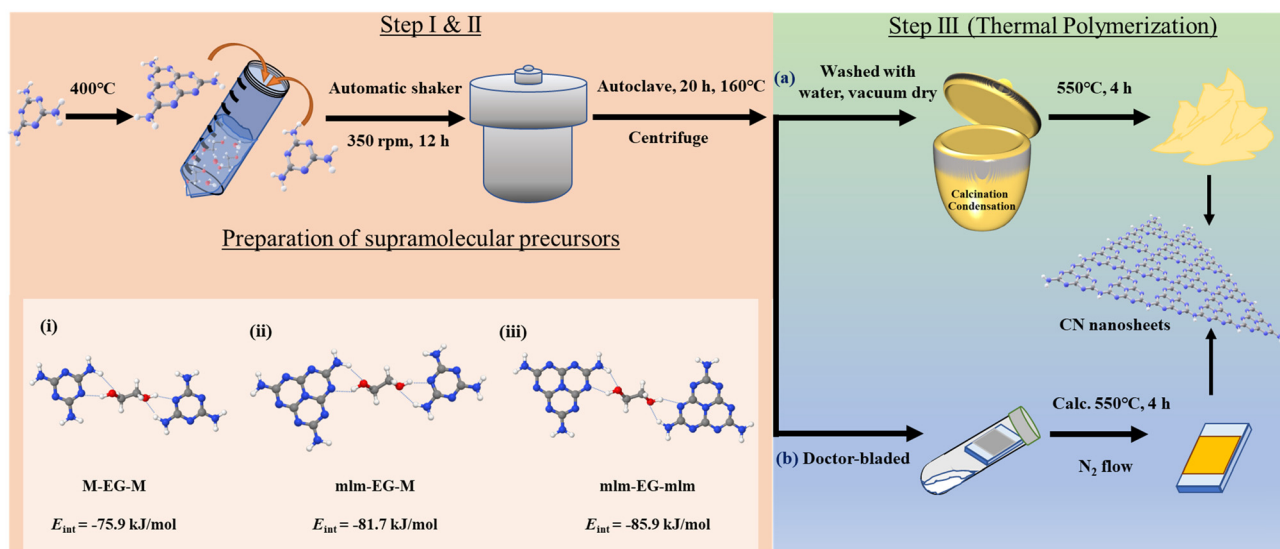


Fig. 1 Procedure for the preparation of CN powder photocatalysts and photoelectrodes *via* doctor-blading. First, melem is synthesized from melamine in one step. Second, a two-step process is used to prepare supramolecular assemblies. Then, they are calcined in a crucible (photocatalyst powder) or deposited on a transparent conductive substrate and calcined (photoelectrode with a CN film). The inset represents the calculated interaction energy values between ethylene glycol (EG) and (i) two melamine molecules (M-EG-M), (ii) one melem (mlm) and one M (mlm-EG-M), and (iii) two mlm molecules (mlm-EG-mlm); all three options can serve as precursors in CN preparation.

structures is more negative than that in M-EG-M structures (-81.7 vs. -75.9 kJ mol^{-1} , respectively), indicating that EG favors the formation of a stable supramolecular structure including both M and mlm (Fig. 1, structure i and ii, Table S1, ESI[†]). Besides, the calculated E_{int} for mlm-EG-mlm is -85.9 kJ mol^{-1} , suggesting that EG easily binds to two melem units (Fig. 1, structure iii). In all three cases, the hydrogen bonding EG facilitates supramolecular assembly of two melem unit and melem-melamine.

Structural characterization of the prepared supramolecular precursors

The powder X-ray diffraction (XRD) pattern of the $\text{P}_{\text{M+mlm}}$ precursor after hydrothermal treatment is shown in Fig. S1 (ESI[†]). Fourier transform infrared spectroscopy (FTIR) inspection of the hydrothermally treated CN precursors (P_M , P_{mlm} , and $\text{P}_{\text{M+mlm}}$) and pristine M and mlm in Fig. 2a show characteristic stretching vibrations related to the breathing mode of triazine units centered at 790 cm^{-1} and vibrations of the heptazine units at $1220\text{--}1610\text{ cm}^{-1}$. Furthermore, the hydrothermally treated samples additional vibrational bands at 2920 cm^{-1} and 2865 cm^{-1} , originating from C-H stretching, and other bands at 1070 cm^{-1} , 1034 cm^{-1} , and 883 cm^{-1} , corresponding to the C-O stretching, C-C-O asymmetric, and C-C-O symmetric stretching modes, respectively, indicating the presence of the EG linker in the final precursor.^{30,31} For the hydrothermally treated samples, an additional weak band is observed at 1682 cm^{-1} , which is attributed to the O-H bending mode.³² In general, the broad bands ($3000\text{--}3600\text{ cm}^{-1}$) are attributed to the stretching modes of primary and secondary amines or their intermolecular H-bonding interactions; the peak intensities and positions ($\sim 3070\text{ cm}^{-1}$) of the hydrothermally treated samples are slightly different from those of pristine melamine and melem precursors, verifying the H-bonding between monomers and EG molecules. Finally, the FTIR spectra of the $\text{P}_{\text{M+mlm}}$ precursors during hydrothermal treatment

(Fig. S2, ESI[†]) show that increasing amounts of EG lead to an increased intensity of the bands at 1070 cm^{-1} and 1034 cm^{-1} , ascribed to the C-O stretching and C-C-O asymmetric modes, respectively, along with a minor shift of their positions. These changes in the FTIR spectra confirm our hypothesis that bonding interactions between EG and the precursors are present during hydrothermal treatment.

Scanning electron microscopy (SEM) images of the precursors indicate that the morphologies of the hydrothermally treated P_M , P_{mlm} , and $\text{P}_{\text{M+mlm}}$ precursors in the presence of EG are entirely different from those of untreated M and mlm (Fig. 2b-d and Fig. S3, ESI[†]). The hydrothermal treatment of $\text{P}_{\text{M+mlm}}$ without EG induces the formation of ordered 1D microstructures as shown in Fig. S4a (ESI[†]). In contrast, the SEM images of $\text{P}_{\text{M+mlm}}$ samples treated with various amounts of EG are presented in Fig. S4b-d (ESI[†]) and show a significant change in their morphology with increasing EG incorporation. The mixed morphology, both 1D microstructure and macroaggregates particles, was observed in the precursors obtained from 16, 32, and 50 v% EG. However, the changes in their morphology were random with the increase in EG concentration.

Structural characterization of the nanostructured CNs

The use of EG in the hydrothermal process has several advantages: (i) the sufficiently high solubility of the CN precursors in EG enables the establishment of highly ordered supramolecular assemblies; (ii) moderate carbon doping shifts the optical and photocatalytic response to longer wavelengths, as shown later; (iii) the prepared precursor paste is readily applicable onto substrates for optoelectronic devices—the precursors, after centrifugation, can be directly used for the preparation of CN photoelectrodes over transparent conductive oxide (TCO) substrates, followed by calcination under N_2 atmosphere (as will be shown later). For the preparation of photocatalyst powders, the slurry remaining after centrifugation was washed once with water and then dried overnight in a vacuum oven at $60\text{ }^\circ\text{C}$. The obtained powder was calcined in a muffle furnace at $550\text{ }^\circ\text{C}$ for 4 h in air. Light brown CN photocatalysts were obtained; their characterization is described in the following subsections. The resulting polymeric carbon nitride materials are denoted as 'CN' with the precursor name in subscript (namely, CN_M , CN_{mlm} , and $\text{CN}_{\text{M+mlm}}$, which are investigated in detail).

The diffraction pattern of $\text{CN}_{\text{M+mlm}}$ (Fig. 3a) exhibits two significant diffraction signals at 13.2° and 27.7° , corresponding to in-plane tri-*s*-triazine units (100) and interlayer lattice (002) planes of melon-based CN, respectively.³³ The diffraction patterns of CN_M and CN_{mlm} similarly exhibit these signature graphitic diffractions. Furthermore, the (002) diffraction plane of $\text{CN}_{\text{M+mlm}}$ is at a higher angle than CN_M and CN_{mlm} , stemming from a decrease in the interlayer-stacking distance in the CN aromatic units. The intensity of the (002) peak is also noticeably diminished relative to those of the CN_M and CN_{mlm} samples, indicating a smaller sheet size of the layers.³⁴

The FTIR spectra of the produced CN samples (Fig. S5, ESI[†]) show the presence of the typical stretching modes of C-N

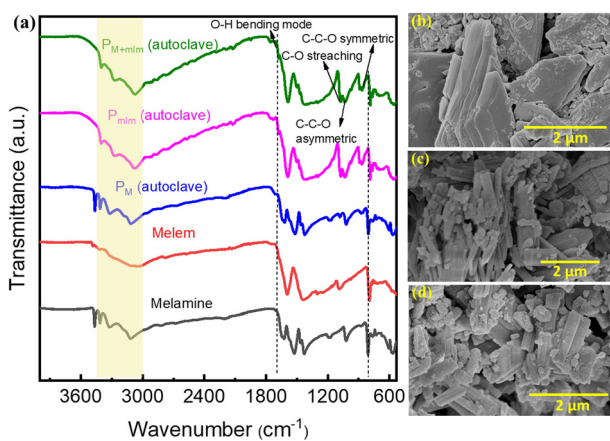


Fig. 2 Precursors characterization. (a) FTIR spectra of the precursors before (melem and melamine) and after (P_M for melamine, P_{mlm} for melem, and $\text{P}_{\text{M+mlm}}$ for the mixture) hydrothermal treatment in an EG/water mixture. SEM images of (b) P_M , (c) P_{mlm} , and (d) $\text{P}_{\text{M+mlm}}$. The spectra in (a) are offset for clarity.

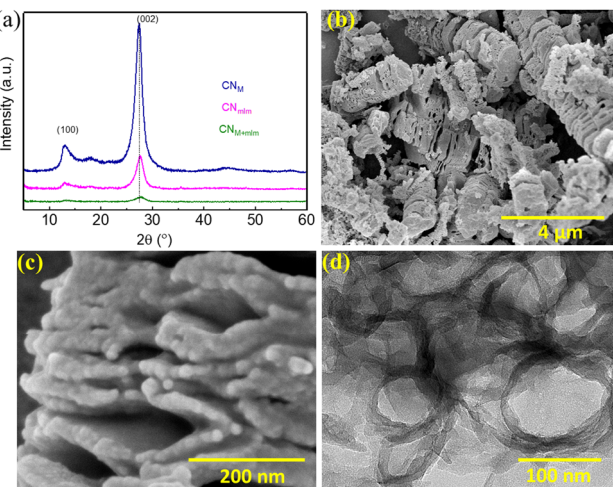


Fig. 3 CN characterization. (a) Powder XRD patterns of CN_M, CN_{mlm}, and CN_{M+mlm}. SEM images of (b) CN_{M+mlm}, and (c) CN_{M+mlm} at higher magnification. (d) TEM image of CN_{M+mlm}.

heterocycles at 1220–1610 cm^{−1} and the breathing mode of the triazine units at 804 cm^{−1}. No noticeable difference is seen between the three spectra, except for a slight change in the broad bands located at 3000–3400 cm^{−1}.^{35,36} No major changes were observed in the FTIR spectra of the resulting CN_{M+mlm} samples when varying the amount of EG during synthesis.

As observed in the SEM images Fig. 3b and c, CN_{M+mlm} retains the both 1D and particle morphology of the precursor (P_{M+mlm}). Moreover, a careful examination reveals that CN_{M+mlm} is composed of thin nanosheets attached to one another (Fig. 3c). We suggest that the condensation process during calcination of the supramolecular assemblies (1D and micro-particles) used as precursors is responsible for this unique morphology. In contrast, CN_M shows the formation of dense solid CN agglomerates several micrometers in size (Fig. S6a, ESI[†]). CN_{mlm} consists of a nanosheet architecture with irregular porous structures (Fig. S6b, ESI[†]).

Fig. S7 (ESI[†]) indicates that the morphologies of the precursors and the final CN samples are significantly affected by the amount of EG in the precursor subjected to hydrothermal treatment. The ordered 1D microstructures of the precursor have completely changed and formed nanosheet architecture with irregular porous structures in the case of CN_{0 v%} EG (Fig. S7a, ESI[†]). However, CN obtained from various amounts of EG (16, 32 v%) shows a porous exfoliated 1D morphology composed of nanosheets attached to one another (Fig. S7b and c, ESI[†]). It can be noted that the CN_{50 v%} EG consists of both porous exfoliated 1D and porous particles (Fig. 7d).

To study the role of EG, we varied the EG content in the preparation of CN_{M+mlm}. Fig. S7 (ESI[†]) indicates that the morphologies of the precursors and the final CN samples are significantly affected by the amount of EG in the precursor subjected to hydrothermal treatment. Specifically, an optimum amount of EG is required to prepare well-textured CN nanostructures. We hypothesize that the amount of EG must be so that it provides sufficient solubility to promote the formation of

well-defined supramolecular assemblies while still acting as a binder (Fig. 1, structures i, ii, and iii) that holds the structure together during condensation. Transmission electron microscopy (TEM) images of CN_{M+mlm} (Fig. 3d and Fig. S8, ESI[†]) disclose a morphology based on porous and thin nanosheets.

The chemical properties of CN_M, CN_{mlm}, and CN_{M+mlm} were examined by X-ray photoelectron spectroscopy (XPS). The survey spectra (Fig. S9, ESI[†]) confirm the presence of C, N, and O; the weak O 1s signal can be ascribed to surface-adsorbed CO₂ or H₂O, suggesting a negligible amount of impurities in the CN samples. The high-resolution C 1s XPS spectra of CN_M, CN_{mlm}, and CN_{M+mlm} are deconvoluted into three peaks (Fig. 4a). The peaks at 284.8, 286.7, and 288.1 eV are ascribed to graphitic carbon (C–C–H), sp²-bonded carbon in the heptazine or triazine rings (N=C–N), and sp² carbon in the aromatic ring attached to an NH₂ group (C–(N)₃), respectively.^{37–39} The high-resolution N 1s spectra (Fig. 4b) are deconvoluted into four peaks centered at 398.6, 399.2, 400.1, and 401.1 eV, which are attributed to pyridinic N, tertiary N (N–(C)₃), primary N (from –NH₂ groups), and quaternary N, respectively. It is worth noting that the apparent peak shifts in the C 1s and N 1s spectra of CN_{M+mlm} with respect to CN_{mlm} and CN_M indicate that the local environment (chemical and physical) is significantly different.^{40–43} The higher binding energy of C 1s and N 1s in the CN_{M+mlm} spectrum can also be described based on the increased C–N bond covalency and the presence of defects. Furthermore, to find out the defects in the CN with various precursors, we have recorded the C 1s XPS (etched for 60 s) for CN_M, CN_{mlm}, and

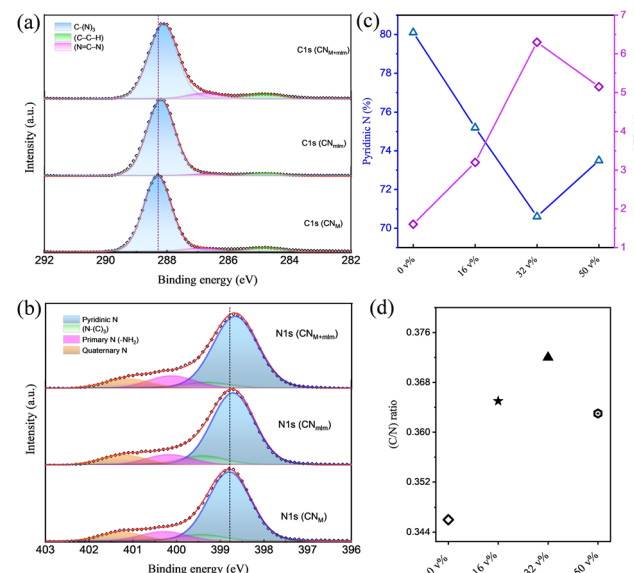


Fig. 4 XPS CN analysis. High-resolution (a) C 1s and (b) N 1s XPS spectra of CN_M, CN_{mlm}, and CN_{M+mlm}. (c) Calculated percentage of pyridinic N and free NH₂ species for various amounts of EG (v%) used in the hydrothermal treatment during CN_{M+mlm} syntheses, calculated from N 1s spectra. (d) Atomic C/N ratio obtained from survey scans of CN_{M+mlm} prepared with varying EG content. The experimental points in (a) and (b) are fitted with a red line, which is deconvoluted into peaks corresponding to the indicated chemical species. The lines connecting the diamonds and triangles in (c) serve as a guide to the eye.

$\text{CN}_{\text{M+mlm}}$ samples (Fig. S10, ESI[†]). After etching, a new peak appeared for all the samples at 285.9 eV, which is attributed to the presence of a C–C bond, originating through the N-vacancy in the tri-*s*-triazine.⁴⁴ In addition, the amount of C–C bonds is higher in the case of $\text{CN}_{\text{M+mlm}}$ compared to CN_{M} and CN_{mlm} , suggesting that the precursor induces C-defects in the final CN structure.

The effect of the EG content in the precursor on the chemical properties of the resulting $\text{CN}_{\text{M+mlm}}$ was also examined using XPS. The C/N ratio was calculated from the survey spectra of samples prepared with various amounts of EG. The C/N ratio increases with increasing EG content until ~ 32 v% as shown in Fig. 4c (and Fig. S11, ESI[†]). However, the C/N ratio was lowered as the EG content was further increased to 50 v%; a decrease attributed to the excessive solubility of melamine in EG, which disfavors the formation of a supramolecular assembly (Fig. S12, ESI[†]).

The high-resolution C 1s and N 1s spectra of $\text{CN}_{\text{M+mlm}}$ samples that were prepared using various EG contents (v% in water) are shown in Fig. S13 (ESI[†]); they exhibit an increase in the amount of free NH_2 and a decrease in the amount of pyridinic N, as the EG content is increased up to 32 v% (Fig. 4d).^{45–47} EG with its carbon chain creates a large distance between two monomer units through H-bonding; therefore, during the calcination process, some of the EG molecules leave and result in free $-\text{NH}_2$ groups rather than polymerization between the two $-\text{NH}_2$ moieties. A table of relative ratios of groups in N 1s XPS spectra for CN with various EG content is shown in Table S2, ESI[†]. Importantly, an increased number of free NH_2 groups provides a larger number of electron-rich centers in CN, which favor the adsorption of metal ions (Pt^{2+}) during co-catalyst photodeposition. This interaction, in turn, facilitates the formation of multiple Pt(s) nucleation sites,⁴⁸ resulting in well-dispersed small Pt nanoparticles, which serve as cocatalysts during photocatalysis. Elemental analysis (EA) of CN_{M} , CN_{mlm} , and $\text{CN}_{\text{M+mlm}}$ (Table S3, ESI[†]) indicates that their carbon content also increases with increasing EG content, probably owing to the incorporation of carbon originating from the EG molecules. Solid state ¹³C NMR spectra of $\text{CN}_{0\text{ v\% EG}}$ and $\text{CN}_{32\text{ v\% EG}}$ samples indicate that the EG contributes trace amount of $-\text{CH}_2$ species to the final CN, as shown in Fig. S14, ESI[†].

Optical properties of CN nanostructures

The UV–vis diffuse reflectance spectra of $\text{CN}_{\text{M+mlm}}$ (Fig. S15, ESI[†]) show a redshift of the absorption band edge and improved light absorption in the visible range, suggesting the presence of defects states below the conduction band in the CN framework.^{49,50} The calculated optical direct band gap of $\text{CN}_{\text{M+mlm}}$, E_g , is 2.82 eV; higher E_g values of 2.93 eV and 3.01 eV were obtained for CN_{M} and CN_{mlm} , respectively (Fig. 5a).

The amount of EG used in the hydrothermal process has a direct effect on the light-absorption properties of $\text{CN}_{\text{M+mlm}}$, as shown in Fig. 5b. A noticeable redshift of the absorption band was observed with increasing EG content, indicating the incorporation of carbon into the CN structure (as confirmed by EA,

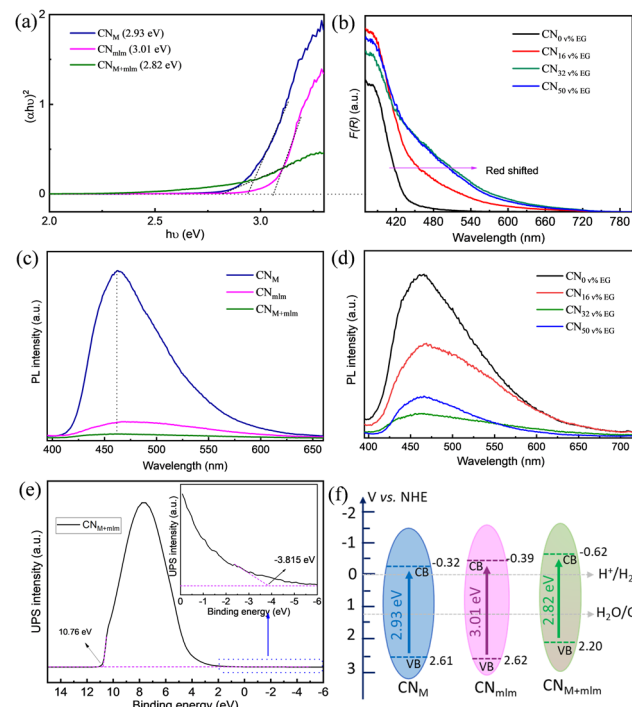


Fig. 5 Optical characterization and band positions estimation. (a) Tauc plot analysis of the CN samples assuming a direct E_g . (b) UV-vis diffuse reflectance spectra of $\text{CN}_{\text{M+mlm}}$ prepared using varying EG content (v%). Photoluminescence emission spectra of (c) CN_{M} , CN_{mlm} , and $\text{CN}_{\text{M+mlm}}$, and (d) $\text{CN}_{\text{M+mlm}}$ prepared using varying EG content in the precursor preparation. (e) UPS valence band spectrum of $\text{CN}_{\text{M+mlm}}$. (f) Schematic representation of the electronic band structure (on the normal hydrogen electrode (NHE) scale) of CN_{M} , CN_{mlm} , and $\text{CN}_{\text{M+mlm}}$, determined using the UPS-measured VB position and the optical E_g determination.

Table S3, ESI[†]). The latter observation suggests that even after its removal, residual amounts of EG remain in the precursor, bound within its structure. The photoluminescence (PL) spectra for the CN_{M} , CN_{mlm} , and $\text{CN}_{\text{M+mlm}}$ samples under an excitation wavelength (λ_{ex}) of 375 nm are shown in Fig. 5c. CN_{M} shows a strong emission band centered at 462 nm, which is attributed to direct-photogenerated electron–hole recombination. Significant quenching was observed for $\text{CN}_{\text{M+mlm}}$ as compared with CN_{M} and CN_{mlm} , possibly stemming from an alternative charge recombination process, which suggests that charge separation was improved by the creation of new defect states.^{51,52} The pronounced decrease in the PL emission of the CN samples with increasing EG amounts (Fig. 5d) is attributed to the higher carbon content. This quenching is maximal for $\text{CN}_{\text{M+mlm}}$ with 32 v% EG, suggesting an optimum EG content with respect to photogenerated charge recombination.

In addition to having an appropriate E_g , a semiconductor photocatalyst must have conduction band (CB) and valence band (VB) positions that match the redox potentials required for the targeted photocatalytic reactions. To study the VB positions of the CN materials, ultraviolet photoelectron spectroscopy (UPS) was recorded (Fig. 5e and Fig. S16, ESI[†]). The VB energy (E_{VB}) positions of CN_{M} , CN_{mlm} , and $\text{CN}_{\text{M+mlm}}$ were calculated to be 2.61, 2.62, and 2.20 V, respectively. A schematic

of E_{VB} and E_{CB} for the CN samples is illustrated in Fig. 5f. The E_{CB} of CN_{M+mlm} is more negative (-0.62 V) than that of CN_M and CN_{mlm} (-0.33 V and -0.38 V, respectively).^{38,53} The higher E_{CB} position results in a higher driving force towards the HER. The E_{VB} is positioned below the water oxidation level (H_2O to O_2), suggesting that CN_{M+mlm} is also suitable for O_2 production.

The specific surface area (S_A) of the CN materials was analyzed by N_2 adsorption–desorption measurements (Fig. S17, ESI†). The introduction of melem into the precursor leads to significant enhancement of the S_A of the final material: the S_A of CN_{mlm} and CN_{M+mlm} are 13.4 and 9.6 times as high as that of CN_M (S_A of 187, 134, and $14\text{ m}^2\text{ g}^{-1}$, respectively). The introduction of melem in the precursor creates a larger cavity in the supramolecular assembly endowing higher specific surface area in the final CN.⁵⁴ Moreover, the S_A of CN_{M+mlm} is also a function of the amount of EG in its precursor (Fig. S18, ESI†).

Photocatalytic hydrogen evolution reaction (HER) performance

The HER activity under white light illumination of the CN_{M+mlm} samples synthesized using various amounts of EG (0, 16, 32, and 50 v%) discloses an enhancement in activity until 32 v% EG, corresponding to the trends in optical absorbance, PL quenching, NH_2 groups content, and S_A (Fig. 6a). The corresponding average hydrogen production rates for these photocatalysts (normalized to the total mass of the catalyst) were 10.8, 15.2, 23.1, and $13.0\text{ mmol g}^{-1}\text{ h}^{-1}$, respectively. Because the HER performance of CN_{M+mlm} (32 v% EG) is the highest among all samples, in line with its material properties, all the photoactivity measurements hereinafter are of CN samples

from P_{M+mlm} precursors containing 32 v% EG. A comparison table of recently reported results on CN-based HER photocatalysis is displayed in Table S4, ESI† showing the excellent performance of CN_{M+mlm} (32 v% EG) over other catalysts.

The HER activities of the CN_M and CN_{mlm} samples were also determined, as displayed in Fig. 6b. The HER rate of CN_{M+mlm} ($23.1\text{ mmol g}^{-1}\text{ h}^{-1}$) is about 1.6 and 17 times as high as those of CN_M ($14.3\text{ mmol g}^{-1}\text{ h}^{-1}$) and CN_{mlm} ($1.34\text{ mmol g}^{-1}\text{ h}^{-1}$), respectively, thanks to its better light adsorption, suitable CB position, and defective nanostructure. The corresponding hydrogen production of the three samples over time (2 h) is shown in Fig. 6c, expressing a linear relationship between the total produced H_2 and time.

To investigate the stability of CN_{M+mlm} , cycling tests, that is repeated tests in the same experimental setup and reaction conditions, were performed. Fig. 6d shows the CN_{M+mlm} activity during five consecutive cycles; it exhibits only a 10% decline after the 5th catalytic cycle, illustrating the durability of the material. Post-test characterization of the CN_{M+mlm} catalyst (after 5 catalytic cycles) was performed using TEM (Fig. S19, ESI†) and shows uniformly dispersed Pt nanoparticles on the CN surface, with an average size of 5 ± 3 nm and narrow size distribution, explaining the excellent photocatalytic performance.

The apparent quantum efficiency (AQE) of CN_{M+mlm} was measured at several wavelengths in the 395–595 nm range as a proxy for photocatalytic activity evaluation (Fig. 6e). The maximum AQE of CN_{M+mlm} , $\sim 19.2\%$, was obtained at 395 nm. Importantly, CN_{M+mlm} also exhibits good AQE in the visible range, even in the 490–595 nm range, suggesting extended

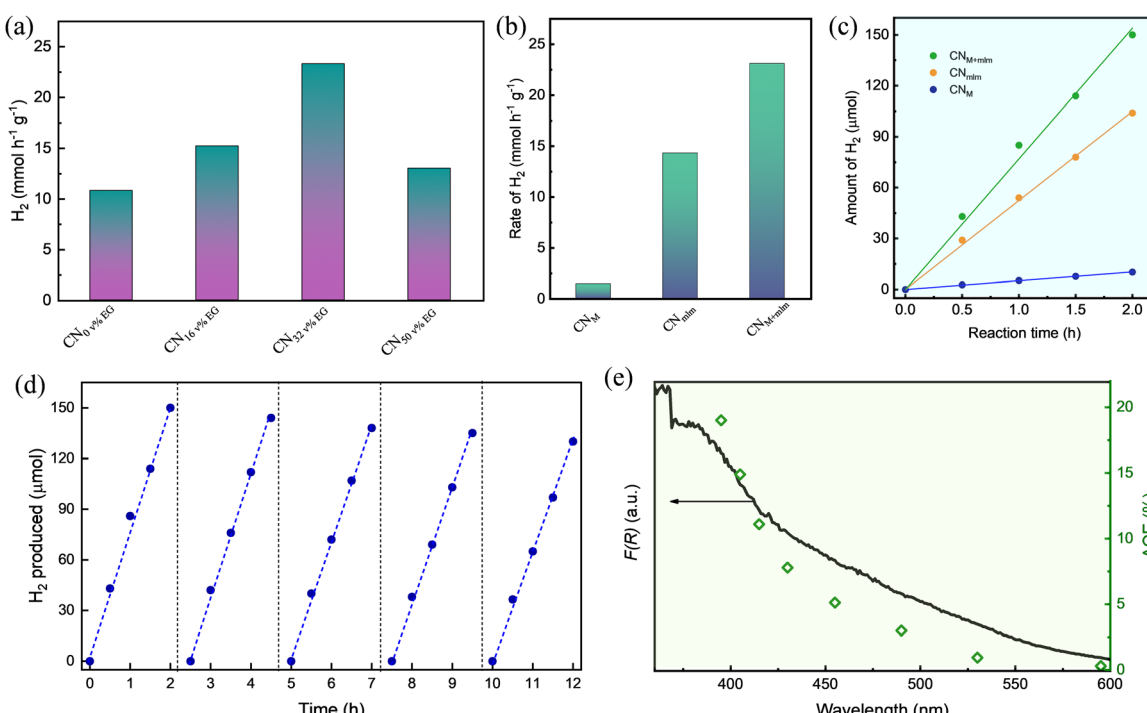


Fig. 6 Photocatalytic characterization. (a) HER rates of CN_{M+mlm} prepared using varying EG content, normalized to catalyst mass. HER performance of CN_M , CN_{mlm} , and CN_{M+mlm} ; (b) photocatalyst mass-normalized HER rate, (c) amount of H_2 vs. time, (d) evolved $H_2(g)$ vs. time in a photocatalytic HER stability test for 5 consecutive cycles, and (e) AQE of CN_{M+mlm} measured at several illumination wavelengths and corresponding absorbance spectrum.

visible light absorption relative to common graphitic carbon nitrides. As shown in Fig. 6e, the photocatalytic performance of $\text{CN}_{\text{M+mlm}}$ coincides well with its absorbance, indicating that the HER performance is directly linked to the formation of electron-hole pairs by photoexcitation.

The higher HER activity of the 32 v% EG $\text{CN}_{\text{M+mlm}}$ may be explained by the combination of several factors: (1) high content of free NH_2 groups (indicated by N1s XPS, Fig. 4b); these groups facilitate the dispersion of the photodeposited Pt nanoparticle cocatalysts and may serve as ligands for nanoparticles, enhancing their stability during the HER. In addition, the free NH_2 groups augment the dispersibility of the CN, as can be seen in Fig. S20 (ESI[†]), allowing better reactivity in water. (2) Formation of a favorable chemical interaction owing to bridging between melem and melamine units in the precursor; upon calcination, favorable defect sites in the CN matrix are formed, as indicated by the observed PL quenching. Furthermore, the EG-mediated C-bridging might result in the formation of a CN homojunction (CN from melem connected C-doped CN from melamine; the homojunction should improve charge separation and catalytic activity. (3) Enhanced light absorbance in the visible light region, corresponding to non-zero AQE at lower energies. (4) High specific surface area.

Photoelectrochemical performance

To show the advantage of the as-prepared precursor containing EG for optoelectronic device fabrication, we prepared photoelectrodes by doctor-blading the obtained slurry directly after the hydrothermal process of the precursors (P_M , P_{mlm} , and $\text{P}_{\text{M+mlm}}$, all using 32 v% EG), on fluorine-doped tin oxide (FTO) coated glass, followed by calcination under N_2 at 550 °C. Note that our illustrated method would not obtain an effective CN film under aerial conditions. Due to the thin layer of precursors formed by the doctor blade on FTO, most of the precursor surface is highly exposed, leading to sublimation and degradation of the monomers during the high-temperature calcination process as shown in Fig. S21, ESI[†]. In this facile procedure, CN photoelectrodes are prepared without using an additional binder, which is typically needed to achieve the required specific viscosity values for the doctor-blade deposition.^{9,10,55} Thus, it eliminates the time-consuming mixing step and improves reproducibility.

The XRD and FTIR spectra of the electrodes (Fig. S22a and b, ESI[†]) prove the formation of a CN layer on the conductive substrate. UV-vis diffuse reflectance spectroscopy (DRS) spectra of the electrodes (Fig. S22c, ESI[†]) confirm an extended visible

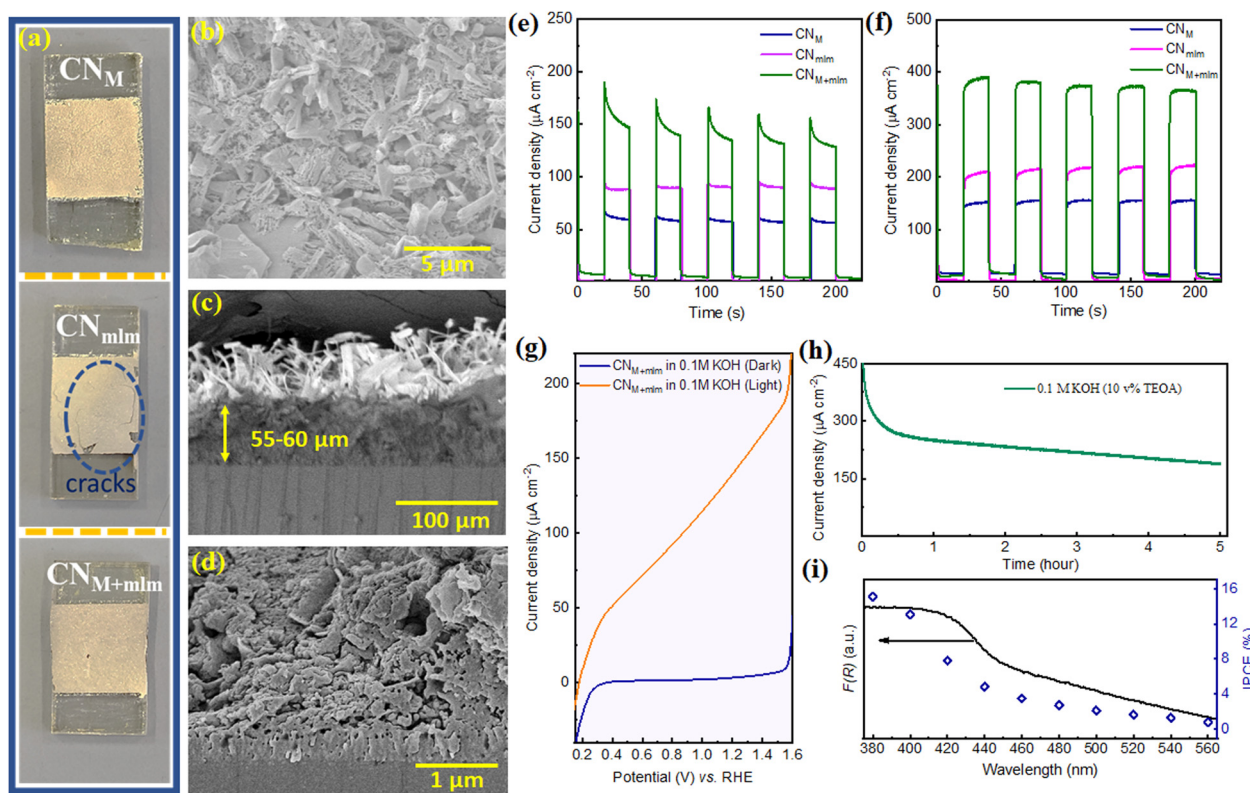


Fig. 7 (a) Digital photographs of CN_M , CN_{mlm} , and $\text{CN}_{\text{M+mlm}}$ photoelectrodes on FTO. (b) SEM image (top-view) of $\text{CN}_{\text{M+mlm}}$. (c and d) Cross-sectional SEM images of $\text{CN}_{\text{M+mlm}}$ electrodes at different magnifications. (e) Chronoamperometry (current densities vs. time) of CN_M , CN_{mlm} , and $\text{CN}_{\text{M+mlm}}$ electrodes in 0.1 M KOH at 1.23 V_{RHE} upon on/off 1 sun illumination. (f) Chronoamperometry of CN_M , CN_{mlm} , and $\text{CN}_{\text{M+mlm}}$ electrodes in 0.1 M KOH aqueous solution containing 10 v% TEOA as a hole scavenger at 1.23 V_{RHE}. (g) Linear sweep voltammetry (LSV) of $\text{CN}_{\text{M+mlm}}$ in 0.1 M KOH in the dark and under 1 sun illumination. (h) Stability measurement (current density under continuous 1 sun illumination) of $\text{CN}_{\text{M+mlm}}$ electrode at 1.23 V_{RHE} in 0.1 M KOH containing 10 v% TEOA during 5 h. (i) Incident photon-to-current conversion efficiency (IPCE) of the $\text{CN}_{\text{M+mlm}}$ electrode at different wavelengths (380–560 nm; using band-pass filters) in 0.1 M KOH containing 10 v% TEOA at 1.23 V_{RHE}.

range absorption for the $\text{CN}_{\text{M+mlm}}$ electrode relative to CN_{M} and CN_{mlm} (Fig. S22c, ESI[†]), in accordance with powder CNs. In addition, the intensity of the PL emission band is lower for $\text{CN}_{\text{M+mlm}}$ films (Fig. S22d, ESI[†]). Fig. 7a shows a photograph of the photoelectrodes obtained after the calcination process. Because the CN_{mlm} film has multiple cracks, hydrothermally treated melem alone is not a suitable precursor to obtain good adhesion between the CN film and the substrate (FTO). We ascribe this incompatibility to partial polymerization, hindering intimate contact with the substrate. In contrast, $\text{CN}_{\text{M+mlm}}$ exhibits a sheet-like structure on FTO, thanks to the fast sublimation of melamine during calcination, which results in the formation of a continuous film (Fig. 7b). A cross-sectional analysis (Fig. 7c and d) of $\text{CN}_{\text{M+mlm}}$ reveals a compact CN film with a thickness of 55–60 μm and a strong attachment to the FTO substrate. SEM images of CN_{M} and CN_{mlm} electrodes are displayed in Fig. S23 (ESI[†]).

The PEC performance of the CN electrodes was evaluated under 1 sun illumination using an aqueous KOH solution (0.1 M) as the electrolyte in a three-electrode system. The photocurrent upon back illumination gives higher photocurrent densities than front illumination (Fig. S24, ESI[†]), owing to the shorter distance that the photogenerated electrons have to travel before reaching the FTO. Hence, the photocurrent measurements were performed using back illumination. The $\text{CN}_{\text{M+mlm}}$ electrode exhibited a photocurrent density ($160 \pm 10 \mu\text{A cm}^{-2}$ at 1.23 V vs. RHE (V_{RHE}))) nearly 2.7 and 1.8 times as high as those of the CN_{M} and CN_{mlm} electrodes (60 ± 5 and $90 \pm 5 \mu\text{A cm}^{-2}$), respectively (Fig. 7e). The improved photocurrent for the $\text{CN}_{\text{M+mlm}}$ electrode indicates that the supramolecular precursor, which contains both melamine and melem, plays a vital role in the formation of a compact CN film over FTO.

When triethanolamine (TEOA, 10 v%) was added to the KOH aqueous solution as a hole acceptor, the photocurrent densities of the CN_{M} , CN_{mlm} , and $\text{CN}_{\text{M+mlm}}$ electrodes were enhanced to 140, 210, and 380 $\mu\text{A cm}^{-2}$, respectively (Fig. 7f). As expected, it indicates that a sluggish hole extraction is the primary limiting factor in a PEC setup. In addition, $\text{CN}_{\text{M+mlm}}$ photoanode photocurrent densities were measured over a wide pH range (0.5 M Na_2SO_4 , pH 6.27; and 0.5 M H_2SO_4 , pH 0.27); values of 80 and 60 $\mu\text{A cm}^{-2}$ were recorded in the neutral and acidic media, respectively (Fig. S25, ESI[†]).

Linear sweep voltammetry (LSV) curves of $\text{CN}_{\text{M+mlm}}$ in Fig. 7g correspond to the typical behavior of a photoanode in a PEC setup: no noticeable change in dark current density was measured until applying a voltage bias of *ca.* 1.55 V, which was sufficient to induce electrocatalytic water splitting, whereas a linear increase in photocurrent density was observed under illumination. The remarkably low onset potential (0.153 V vs. RHE) is complementary evidence of facile charge separation under visible-light illumination of CN.

A stability test for the photoanode was performed under continuous 1 sun illumination for 5 h using similar conditions but in the presence of TEOA as a hole scavenger (pH = 13.1), as shown in Fig. 7h. The current density decreased by nearly 50% after 5 h ($\sim 190 \mu\text{A cm}^{-2}$). To examine the reasons for decreased

stability after 0.5 h, we have recorded an N 1s XPS spectrum of CN electrode after the stability test (Fig. S26, ESI[†]). The deconvoluted N 1s spectrum shows that the primary NH_2 content decreased after operation and a new peak appears at 403 eV, attributed to N–O species. This suggests that the decrease in the current after 0.5 h is due to (partial) oxidation of the NH_2 groups in CN structure. SEM images of the CN electrode after the stability test suggest that the structure of CN remains similar to the unmeasured one (Fig. S27a and b, ESI[†]). We also noticed that the CN is well attached with the FTO even after a reaction of 5 h, as can be seen from the inset of Fig. S27a, ESI[†]. In addition, the cross-section image (Fig. S27c, ESI[†]) indicates that the CN is well attached to the FTO. However, the upper layer thickness of the CN film has been decreased compared to the fresh sample, which could also lead to the decrease stability (Fig. S27c and d, ESI[†]).

The incident photon-to-current conversion efficiency (IPCE) of the $\text{CN}_{\text{M+mlm}}$ photoanode was measured from 380 to 560 nm at 1.23 V_{RHE} (Fig. 7i). The IPCE of the $\text{CN}_{\text{M+mlm}}$ electrode at $\lambda = 380$ nm is calculated to be 15.2%. Notably, the IPCE value at $\lambda = 560$ nm is $\sim 0.8\%$, showing extended activity in the visible range at lower incident photon energies, in line with the measured optical absorption of the films.

With this data in hand, we conclude that the origin of the improved photocurrent and stability of $\text{CN}_{\text{M+mlm}}$ films as photoanodes stems from their good adhesion and intimate contact with the conductive substrate, extended light absorption, better charge separation, low charge recombination, and enhanced electronic conductivity due to carbon doping.

Conclusions

In summary, we introduced a straightforward method to synthesize polymeric carbon nitride materials and electrodes with an enhanced optical response, high surface area, excellent dispersibility in water, and good activity as photocatalysts for the HER and as photoanodes in photoelectrochemical cells. To do so, we employed a hydrothermal route with an aqueous solution of EG to form highly ordered and thermally stable supramolecular assemblies based on the melem–EG–melamine unit as a precursor. The addition of EG enables the formation of highly organized supramolecular structures thanks to the high solubility of melem and melamine. Moreover, EG binds the precursor molecules, leading to the formation of homogeneous precursor paste with carbon-rich monomers. The obtained paste can be cast on a transparent conductive substrate without further treatment, allowing the easy fabrication of CN photoanodes for photoelectrochemical water splitting.

Upon thermal condensation, a highly porous CN with homojunctions and extended light absorption into the visible region is synthesized. The best sample, $\text{CN}_{\text{M+mlm}}$, exhibits excellent, state-of-the-art HER rates up to $23.1 \text{ mmol h}^{-1} \text{ g}^{-1}$ under a white LED irradiation with an AQE of 19.2% at 395 nm. We also developed an original approach for the fabrication of CN photoelectrodes, where no additional binder is used for

doctor-blading, significantly facilitating this process and improving its reproducibility. Compared with the CN_M and CM_{mlm} photoanodes, the constructed $\text{CN}_{M+\text{mlm}}$ photoanodes exhibit a low onset potential of ~ 0.153 V vs. RHE, a high photocurrent density of $160 \pm 10 \mu\text{A cm}^{-2}$ at 1.23 V in 0.1 M KOH ($380 \mu\text{A cm}^{-2}$ in the presence of a hole scavenger) and high IPCE values ranging from 15.2% at $\lambda = 380$ nm to $\sim 0.8\%$ at $\lambda = 560$ nm, and their activity is extended to longer wavelengths.

Author contributions

S. M. performed most of the experiments, analyzed the data, and wrote the initial draft of the manuscript. G. M. helped in photocatalysis studies. L. A. performed XPS characterization. J. L. measured elemental analysis. J. T. has performed TEM analysis. T. S. helped in IPCE measurements. L. X. performed the DFT calculations. M. V. took part in analysis, SEM imaging, and manuscript writing and review. M. S. supervised the study, co-wrote and reviewed the paper, and acquired funding. All the authors discussed the results and reviewed the manuscript.

Conflicts of interest

There are no conflicts to declare.

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