Robust Carbon Nitride-Based Thermoset Coatings for Surface Modification and Photochemistry

Baris Kumru, Jesús Barrio, Jianrui Zhang, Markus Antonietti, Menny Shalom, and Bernhard V. K. J. Schmidt

†Max Planck Institute of Colloids and Interfaces, 14424 Potsdam, Germany
‡Department of Chemistry and Ilse Katz Institute for Nanoscale Science and Technology, Ben-Gurion University of the Negev, 8410501 Beer-Sheva, Israel

Supporting Information

ABSTRACT: Herein, the convenient visible light-induced photografting of hydroxyl ethyl metacrylate onto graphitic carbon nitride (g-CN) is described, leading to well-dispersible g-CN-based precursor polymers that can be injected. Mixing with citric acid as the cross-linker and heating leads to stable thermoset coatings. The process is versatile and easy to perform, leading to g-CN-based coatings. Moreover, the coating can be further functionalized/modified via grafting of other polymer chains, and the resulting structure is useful as photocatalytic surface or as photoelectrode.

KEYWORDS: carbon nitride, dispersion, coating, thermoset, photoactive surface, photoelectrochemistry, photopolymerization

INTRODUCTION

Photoactive nanomaterials have been in the focus of nanoscience in recent years. Among the various photoactive materials, quantum dots, conjugated polymers, or metal oxides have been studied very frequently. In particular, applications such as bioimaging, energy conversion, or photocatalysis are intriguing and drive the research forward. A frequently investigated material is graphitic carbon nitride (g-CN), which is mainly because of its photocatalytic and chemical properties as well as facile synthesis. g-CN has been utilized as the catalyst in applications such as CO₂ conversion, hydrogen evolution, or synthesis of organic molecules, or as promoter for the photoinitiation of polymerizations. Recently, porosity and grain size of g-CN was correlated with hydrogen evolution efficiency, which shows how the material textures affect the utility of g-CN. Moreover, g-CN was doped with metals to gain access to diversified catalysts mechanisms in antibiotic degradation. In electro-oxidation of formic acid or methanol, g-CN was combined with Pd and carbon black to obtain stable and reliable catalysts. One of the major disadvantages of g-CN lies in its low dispersibility in water or organic solvents. The latter limits its maximum concentration and the range of applications. Thus, various approaches have been investigated to tackle the dispersibility issue, for example, surface functionalization or treatment with strong acids, just to name a few. Recently, photo-induced functionalization reaction has been introduced as a versatile tool to enhance dispersibility of g-CN and tailor the surface structure according to specific needs. In addition, photoreactive surfaces have found significant interest recently. In such a way, surface properties can be altered effectively and with spatial control, for example, for polymer grafting, placement of cells, or protein functionalization, or light-emitting diodes (LEDs). g-CN has remarkable photocatalytic properties; thus, the formation of g-CN films and coatings is a topic of significant interest for further exploitation of g-CN in photoelectric devices. An early example of g-CN film formation utilized a sputtering approach that allowed the formation of uniform coatings with thicknesses up to 2 μm. One of the methods that are frequently used is based on vapor deposition, which allows film formation on various substrates such as indium tin oxide, silica, or glass. Such g-CN films can be utilized as actuators reacting to various external triggers. Wang and coworkers investigated the formation of g-CN films and coatings via the formation of a sol. In this work, the sol was formed via oxidation of the g-CN in an acidic environment. Another approach is the direct growth of CN on the surface, for example, via a supramolecular preorganization route or precursor paste formation. Wang and co-workers showed the catalytic activity of g-CN films in water splitting. The films were formed on fluorine-doped tin oxide glass directly, which formed an effective system for photocatalysis. Structured and patterned surfaces were generated via templating methods, for example, soft or hard templating as well as a combination of both, an additional way to patterned surface growth of g-CN inside well-defined porous substrates. Moreover, film...
formation enables the formation of flexible devices if flexible substrates are employed, which is of significant interest for organic electronics. Nevertheless, the formation of g-CN containing polymer coatings and films in a straightforward and easy way is still a matter of research. Hence, film formation can be consigned to a polymer, whereas the photochemical and catalytic properties are provided by the g-CN. Here, we approach these coatings by embedding g-CN into a polyester thermoset, which is a fundamentally different approach compared to literature-known g-CN films. The polymer-based route is largely scalable, reproducible, and affordable. Moreover, the utilization of a polymer matrix allows the formation of various shapes and structures, which is hardly achieved with g-CN itself. Polyesters are common polymer materials that have various applications in everyday life, and they are considered to be robust against CN oxidation. Using colloidal precursors, carbon materials were introduced into polyester thermosets for enhancing mechanical properties, or polyester thermosets were formed from renewable resources. Polyester thermosets combined with inorganic compounds can be indeed considered model systems to obtain hybrid materials with enhanced mechanical and thermal properties.

Herein, the convenient photografting of hydroxyl ethyl methacrylate (HEMA) onto g-CN is described, leading to well-dispersible g-CN precursor colloids [cyanuric acid–melamine (CM)–HEMA] (Scheme 1) that can be processed or injected. The precursors can be mixed with citric acid as a cross-linker and converted into stable coatings via heating. Moreover, the coating can be further functionalized/modified via grafting of other polymer chains, utilizing the photocatalytic surface.

Scheme 1. Schematic Overview of CM–HEMA Precursor Formation and Subsequent Film Formation via Thermal Curing

### EXPERIMENTAL SECTION

**Materials.** Al_{2}O_{3} basic (Sigma-Aldrich), citric acid (ACS grade 99.5%, Sigma-Aldrich), cyanuric acid (98%, Sigma-Aldrich), D_{2}O (99.9%, Sigma-Aldrich), dichloromethane (DCM, anhydrous HPLC grade, Merck), ethylene glycol (EG, 99%, Fluka), hydrochloric acid (1 M solution, Merck), hydroquinone (ReagentPlus, Sigma-Aldrich), melamine (99%, Sigma-Aldrich), potassium hydroxide (Sigma-Aldrich), rhodamine B (RhB, >95% Sigma-Aldrich), and tetrahydrofuran (THF, anhydrous, Sigma-Aldrich) were used as purchased. 2-Hydroxyethyl methacrylate (HEMA, Sigma-Aldrich), N,N-dimethylacrylamide (DMA, 99%, Sigma-Aldrich), and styrene (ReagentPlus, Sigma-Aldrich) were passed through basic alumina column prior to use to remove the inhibitor. For visible light irradiation, 50 W LED chips (Foxpic High Power 50 W LED Chip Bulb Light DIY White 3800LM 6500 K) were connected to a self-made circuit and cooling system. g-CN from the cyanuric acid-melamine complex (noted as CM) was synthesized according to the literature.

**Synthesis of CM-Based Precursor.** CM (200 mg) was mixed with 4.5 g of deionized water and 4.5 g of EG and ultrasonicated at 50% amplitude for 20 min (10 × 2 min portions) to yield a g-CN dispersion. Afterward, 0.8 g of HEMA was added to the dispersion, and the mixture was put between two 50 W LED daylight sources (20 cm apart from each other) and reacted for 3 h under mild stirring (conversion 94% obtained via 1H NMR). The resulting highly viscous and disperse material was later utilized as the coating precursor after the addition of citric acid.

**Synthesis of g-CN Coating.** The g-CN-based precursor (2 g) was mixed with 100 mg of citric acid and stirred until dissolved. Later on, the precursor mixture can be applied to the desired surface and cured at 120 °C for 3 h for complete cross-linking and evaporation of solvents.

**Further Polymer Grafting on CM–HEMA Coatings.** CM–HEMA-coated glass substrate surfaces were covered with monomer (DMA or styrene) and illuminated via visible light for 8 h from the top (illumination applied from 20 cm above the sample). After reaction, the DMA-grafted substrate was washed with 20 mL of distilled water, whereas the styrene-grafted substrate was washed with 20 mL of THF for the removal of unreacted/nongrafted impurities.

**Dye Degradation via CM–HEMA Coatings.** The photoactivity of the CM–HEMA films was tested by the adsorption of an organic dye and its degradation under white light irradiation. For this experiment, glass-coated CM–HEMA films (2 × 5 cm²) were submerged in a solution of RhB (20 mg mL⁻¹) overnight and then exposed to illumination with a 50 W white LED array (Bridgelux BXRA-50CSS0; λ > 410 nm). The degradation of the dye to CO₂ with the CM–HEMA films was confirmed by gas chromatography (GC).
characterization methods. Scanning electron microscopy (SEM) was performed using JSM-7500F (JEOL) equipped with an Oxford Instruments X-MAX 80 mm² detector for g-CN-based coating surface morphology. Fourier transform infrared (FT-IR) spectra were taken on Nicolet iS 5 FT-IR spectrometer. Sessile drop contact angles of water on the coatings were measured at room temperature about 5–10 s after placing the drop on the surface with a DSA 10 video contact angle measuring system G10 (Krüss, Germany), and data evaluation was done with software DSA version 1.80.02. Thermogravimetric analysis (TGA) was performed via TG 209 Libra from Netzsch in a nitrogen atmosphere with a heating rate of 10 K min⁻¹ using an aluminum crucible for samples. Ultrasonication was performed via an ultrasonicator at 50% amplitude (Branson D450). For rheological investigations, Anton Paar MCR 301 rheometer equipped with a cone plate 12 (CP-12) was used. Measurements were performed at a constant angular frequency (10 rad s⁻¹) with changing frequency in the range of 0.1 to 100% with 31 measuring points and 0.02 mm gap. Frequency-dependent measurements were performed at constant strain (0.1%) with changing frequency in the range of 1–100 rad s⁻¹. Viscosity measurements were performed at ambient temperature with changing shear rates between 1 and 40 s⁻¹. The flow curve was obtained via measuring the shear stress and viscosity with changing shear rates from 0 to 100 s⁻¹. Atomic force microscopy (AFM) from Digital Instruments was employed for the investigation of half-coated glass samples. Glass substrates were utilized as the reaction medium that allows to increase the solid content of the reaction via enhanced dispersibility of CM. Hence, HEMA-grafted CM (CM–HEMA) could be obtained in a facile fashion. To characterize the formed material and prove successful grafting, FT-IR and elemental analysis were performed (Figure S2 and Table S1). FT-IR showed the presence of bands that can be assigned to poly(HEMA) (PHEMA), for example, the band around 1750 cm⁻¹ corresponding to the carbonyl stretching vibration (Figure S2). Both CM and CM–HEMA materials present absorption bands corresponding to the triazine breathing of CN around 800 cm⁻¹. Moreover, stretching modes of CN heterocycles can be detected between 1150 and 1600 cm⁻¹, and asymmetric CH stretching arising from HEMA can be observed around 2900 cm⁻¹ in CM–HEMA (Figure S2). XRD of the CM–HEMA precursor shows no crystallinity probably because of the undefined nature of the material before cross-linking (Figure S3). X-ray photoelectron spectroscopy proves the formation of modified g-CN films (Figure S4). C 1s spectra display three different chemical states corresponding to sp³ C–H (284.6 eV), sp² C–O (286.0 eV) and sp C–N (287.7 eV) bonds, which confirm the presence of high amounts of the HEMA precursor. Furthermore, the contribution at higher binding energies (288.7 eV) certifies the presence of C≡N–C units within the films. N 1s spectra show the typical features of CN materials, three different contributions can be observed that belong to C≡N–C coordination (398.7 eV), N–(C)₂ (399.9 eV), and the remaining amino groups (N–H, 401.0 eV).

To gain insights into the polymerization process and to verify the free radical polymerization mechanism, blank reactions were performed. Without light irradiation, no conversion of monomer was observed (Figure S5). After the addition of a polymerization inhibitor, namely hydroquinone, insignificant conversions (below 5%) compared to the inhibitor-free polymerization (conversion above 90%) were observed. The dispersion of CM–HEMA in water/EG medium could be utilized directly as precursor for coatings. Citric acid was added as a cross-linker to obtain stable coatings. An efficient coating process was established via adjustment of the polymerization conditions to yield CM–HEMA precursor with a suitable viscosity for film formation (Figure S6) that features values around 25 Pa s at a shear rate of 1 s⁻¹ and a storage modulus (G″) of 5550 Pa at 1% strain (Figures S7 and S8). As the G′ is higher than the loss modulus (G″), the precursor has viscoelastic behavior. Significant shear thinning was observed leading to G′ values of 68 Pa at 50% strain. Between 32 and 40% strain, the gel-like behavior shifts to a sol-
like behavior ($G'' > G'$) (Figure S8). The precursor was obtained as a yellow viscous dispersion that can be processed and even injected with a syringe as well (Figure S6), utilizing the shear-dependent viscosity of the precursor (Figure S7), for example, at a shear rate of 15 s$^{-1}$, the viscosity drops to 1.3 Pa s. This way, the precursor can be applied to various surfaces in a spatially controlled manner. Viscosity values at a low shear range is needed for applications via dipping, however, significantly decreased viscosity at higher shear rates provides plausible applications via spraying or injection.

To facilitate the cross-linking, a carboxylic acid cross-linker such as citric acid was added to the CM–HEMA dispersion. After its formation, the film could be cured via thermal treatment at 120 °C to form the cross-linked polyester (Scheme S1). Homogenous films were obtained after curing as observed via AFM and SEM (Figure 1). AFM shows a smooth film with homogenous thicknesses around 500 nm when compared to the uncoated glass slide (Figure S9). The AFM results are confirmed by SEM that shows a film with minor surface alteration and nonporous as well as rigid structure (Figures 1 and S10). The thickness of films can be adjusted conveniently via the applied method (spraying or brushing) and the applied cycles. However, a detailed study regarding the thickness and photocatalytic properties is work in progress. The incorporation of CM was observed via FT-IR as well as the naked eye because of the light-yellow color of the films. XRD of the coating is dominated by the amorphous polymeric network and shows only a minor diffraction peak at 28° (Figure S11). Interestingly, films could be formed and cured on various substrates, for example, on glass, copper, poly(styrene) (PS), or wood, which shows the versatility of the presented method (Figures S12 and S13). In addition, the formed coatings can be detached from the substrate, leading to a free-standing film (Figure 1). The cross-linking process can be followed easily via TGA. A weight loss of around 87% can be observed between 50 and 170 °C with the steepest weight loss around 139 °C (Figures S14 and S15). There are several processes happening during heating of the precursor mixture. First of all, solvent, that is, water and EG, evaporates. Second, a condensation reaction between citric acid, CM–HEMA, and most likely EG takes place. The final product consists of a polyester between citric acid, EG, and PHEMA. To quantify the incorporation of EG, gravimetry was utilized. After curing at 120 °C for 24 h, an EG incorporation of 52 wt% was observed (refer to the Supporting Information for details of the calculation). Thus, EG acts as a bridging molecule to obtain an extended cross-linked gel-like structure that enables the photochemical features of the coating (see below). The curing temperature of 120 °C is an intermediate temperature below the temperature of significant evaporation of solvent molecules that allows slow curing to facilitate homogenous coatings. The rather low-curing temperature for the polyester formation can be explained with the organocatalytic property of CN in esterification.\textsuperscript{65,66} In the absence of citric acid, curing yields a flexible film that is soluble in water (Figure S16). As soluble structures are less applicable as coatings, further investigations were focused on cross-linked structures.

The final coating has a thermal stability up to temperatures of around 290 °C (Figure S11), which is sufficient for many applications, especially in photocatalysis, where usually the temperature ranges from ambient to 120 °C. The stability of the CM–HEMA coatings against chemical exposure was investigated as well. For that, films on glass slides were introduced into various chemical environments, namely water, acid (1 M HCl), base (2 M KOH), THF, and DCM. Excluding the base treatment, in all cases, no response of the films was observed (Figure S17) also after treatment with visible light (Figure S18). Treatment with a base led to the complete dissolution of the films after 24 h, which is due to the ester-based linkage of the coatings. As esters can be readily hydrolyzed via basic saponification, the low stability in basic medium was expected.\textsuperscript{57} In comparison, a reference film from
linear PHEMA dissolves in organic solvents easily, which clearly supports the colloidal cross-linking also by the CN-based precursor (Figure S19).

To study whether the formed CM–HEMA films are still photoactive, photopolymerization was probed. Therefore, CM–HEMA films were swollen with a hydrophobic monomer, styrene, and irradiated with visible light. A photoinitiated growth of PS from the coated layer could be observed via various methods. For example, contact-angle measurements show an increase in water contact angle from 42.1° to 79.5° (Figure 2b), which is in accordance with an increase in hydrophobicity of the surface. Moreover, the grafted PS could be located via FT-IR showing both bands from the CM–HEMA precursor as well as PS (Figure 2b), for example, the HEMA carbonyl stretching band around 1750 cm⁻¹ and the aromatic stretching of PS around 1500 cm⁻¹. Notably, the coating could be functionalized with poly(N,N-dimethylacrylamide) (PDMA) via visible light photopolymerization as well. The hydrophilic PDMA leads to a decrease in water contact angle to 19.0° as expected because of its hydrophilic nature (Figure 2b). FT-IR revealed the incorporation of PDMA, for example, both carbonyl stretching band from PDMA and HEMA are visible around 1650 and 1750 cm⁻¹, respectively. Because of the photo-induced process of polymer grafting, patterned surfaces could be prepared as well. For example, a glass slide was coated with CM–HEMA, cured, and one-half was covered. Subsequently, PS was grafted in the uncovered area, followed by covering of the PS-grafted area, and another step of PDMA grafting was performed. In such a way, a surface containing a domain of PS and a domain of PDMA were prepared that have different water contact behavior in two domains (Figure S20).

Functionalization of CM–HEMA films with PS and PDMA does not only prove the photoactivity of the incorporated CM but allows for versatile modification of the coating properties. In such a way, surface properties and selectivity can be tailored according to an envisioned application and combined with a photocatalytic activity in a convenient way.

The incorporated CM does not only act as a photoinitiator but can be utilized for photocatalysis as well. To test the photocatalytic activity of the as-prepared films, we performed the uptake followed by the photodegradation of RhB dye. For that, a CM–HEMA-coated glass slide was immersed overnight in an RhB solution to achieve a complete adsorption of dye within the films. After the uptake, the films obtain the typical pink color of the organic dye (Figure 3), and the photochemical experiment starts by switching on a white LED. The film showed an excellent photocatalytic activity, degrading the dye to CO₂ in 90 min, as confirmed by the analysis of the head space during illumination by GC. Clearly, the color of the dye vanished, and the CM–HEMA film recovered its pristine yellowish color. Furthermore, the coated glass slide can be recycled and utilized in consecutive cycles of dye adsorption and photodegradation maintaining the initial activity. On the downside, the photocatalytic activity decreases compared to pure g-CN films. Nevertheless, this experiment shows one of the key futures of these CM-containing coatings: they are easily applied onto various surfaces and can act as a recyclable photocatalyst, which might be interesting for waste water remediation in the future.

In addition to photocatalysis, CN photoelectrodes have attracted widespread attention lately, and achieving uniform CN layers is one of the main challenges to address in CN-based photoelectrochemical cells. Therefore, photoelectrochemical measurements were performed, employing our coating as photoelectrode. To obtain a photoelectrode, CM–HEMA precursor was deposited on an FTO electrode and thermally cured. The photoresponse was evaluated in a photoelectrochemical cell with an Ag/AgCl reference electrode and a Pt counter electrode employing an acidic electrolyte (0.5 M H₂SO₄, pH = 0.2) and one sun illumination (1.5 A) (Figure S21). Stable photocurrent was observed with values in the range of 0.3–0.7 μA cm⁻² (Figure S22). The value of the observed photocurrent strongly depends on the thickness of the layer as observed via a comparison of illumination from the front or the back. In the case of back illumination, the light has a shorter path through the photocatalytic material, which leads to an approximately doubled photocurrent. The observed photocurrents are significantly lower than those reported in literature. A reason for the low photocurrent might be the low conductivity of the synthesized CM–HEMA coatings, which hinders effective charge transport. Although the obtained photocurrent does not reach literature values, the results are still relevant, considering the otherwise electrically insulating character of the polyester film matrix. An option for the further improvement of photoelectrochemically active layers might be the incorporation of conductive additives or more conductive polymers. As porosity is an important factor for photocatalytic activity, controlling the cross-linking density or utilization of porogens during curing might be an option to enhance the performance of CN-based polyester materials in the future.

**CONCLUSIONS**

In conclusion, a convenient method to obtain CN-containing polymer coatings was presented. The CM–HEMA precursor can be synthesized easily via visible light irradiation, processed, injected, or printed, for example, for spatially controlled deposition. Curing is performed via heat treatment leading to smooth and stable coatings. Furthermore, the coatings can be functionalized with polymers via visible light-induced free-radical polymerization, as shown with PS and PDMA. The coatings slightly swell and are thereby are not only active in polymerization reactions but also in photocatalysis, as shown via dye degradation experiments and photoelectrochemical measurements. Overall, the CM–HEMA can be utilized to
cover surfaces on a large scale in a convenient way and serve as a kind of platform to obtain photocative coatings.

**ASSOCIATED CONTENT**

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsami.8b21670.

Additional analytical data (rheology, viscometry, photographs, and photoelectrochemistry) and procedures (PDF)

**AUTHOR INFORMATION**

**Corresponding Authors**
*E-mail: mennysh@bgu.ac.il (M.S.).
*E-mail: bernhard.schmidt@mpikg.mpg.de (B.V.K.J.S.).

**ORCID**
Jesus Barrio: 0000-0002-4147-2667
Markus Antonietti: 0000-0002-8395-7558
Menny Shalom: 0000-0002-4506-4177
Bernhard V. K. J. Schmidt: 0000-0002-3580-7053

**Author Contributions**
B.K. and J.B. contributed equally. The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

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The authors declare no competing financial interest.

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**REFERENCES**


