Doping Carbon Nitride Quantum Dots into Melamine-Silver Matrix: An Efficient Photocatalyst with Tunable Morphology and Photocatalysis for H₂O₂ Evolution under Visible Light

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Carbon nitride quantum dots (CNQDs) were doped into melamine (MA) and silver (Ag) matrix yielding CNQDs@MA–Ag nanocomposites by the supramolecular self-assembly route. To our surprise, the resulting mesoporous nanocomposites could present different morphologies (i.e., nanowires, nanoflowers, and nanosheets) and especially tunable visible-light photocatalysis performances depending on the percentages of CNQDs used. It was discovered that the CNQDs@MA–Ag ones with the pinecone-like structure (containing 30% CNQDs) could exhibit the strongest visible-light photocatalysis for splitting water to yield H₂O₂ with a production efficiency of 39.82 μmol·g⁻¹·h⁻¹, which is more than ten-fold higher than that of CNQDs alone.

Herein, the nano-scaled Ag could facilitate the increased light harvesting and the separation as well as transmission of electron-hole pairs of CNQDs, resulting in the enhanced visible-light photocatalysis of nanocomposites. More importantly, the pinecone-like ones might additionally enable more exposed active sites of photocatalysis and larger mesoporous specific area of contacting dissolved oxygen in water, leading to the higher photocatalytic efficiency of H₂O₂ generation. This CNQDs@MA-based fabrication strategy may open a new door towards the design of a variety of photocatalysts with different morphologies and tunable photocatalysis performances for the efficient splitting water toward H₂O₂ evolution under sunlight.

Introduction

Hydrogen peroxide (H₂O₂), an environmentally-friendly oxidant only with the final byproduct of water, has been extensively applied in the fields of environmental treatment, chemical industry, and biological process.[1–4] Traditionally, H₂O₂ is produced industrially by two kinds of routes of anthraquinone oxidation[5] and H₂ and O₂ reactions,[6] which may be trapped by some limitations like high energy consumption, tedious operation, and explosion risk. In recent years, many efforts have been alternatively devoted to the photocatalytic production of H₂O₂ due to the advantages of green sources (i.e., water and oxygen) and clean energy input (i.e., sunlight).[7–10] It is generally recognized that the performances of the photocatalytic systems may largely depend on the functional properties of photocatalysts.[11–13] As a result, how to enhance the photocatalytic functions especially the visible-light ones of photocatalysts has been being a hot topic in the current researches.[14–18] For example, Ge et al. utilized Ag nanoparticles to increase the visible-light photocatalytic activity of polymeric g-C₃N₄.[14] Xu and coauthors fabricated ZnFe₂O₄ composites in combination with Ag/AgBr showing improved photocatalysis for organic pollutant degradation.[19] Moreover, the performances of photocatalysts can largely depend on their micro-morphologies and structures.[19–21] For example, Dong’s group discovered that (BiO)ₓCO₃ nanomaterials doped with different levels of thiourea could present different morphological structures and tuning photocatalysis properties.[19] Wang et al. have reported the fabrication of Ag₃PO₄ photocatalysts with morphologies-depending photocatalysis in the degradation of methylene blue.[20] Unfortunately, most of the current fabrication routes for the control of morphologies and structures of photocatalysts can encounter with some formidable limitations like complicated steps, limited photocatalytic activities, and harsh reaction conditions (i.e., high temperature and pressure). In particular, some efforts have been devoted to the applications of surfactants with large amounts for controlling the morphological structures of photocatalysts.[22,23] A challenge regarding the limited photocatalysis of the photocatalysts might be encountered due to their surface active sites might be greatly hindered by the surfactants adsorbed. Hence, exploring a simple, efficient, and green technology for the fabrication of photocatalysts with controlled morphologies and visible-light-driven photocatalysis performances for splitting water to generate H₂O₂ evolution has been a challenging but attractive target to pursue. Besides, recent years have witnessed the rapid
development of various types of carbon materials like nitride quantum dots (CNQDs) or g-C₃N₄ for the extensive applications of photocatalysis, adsorption, and sensing analysis due to their outstanding merits such as chemical stability, metal-free non-toxicity, cost-effectiveness, and easy preparation. For example, Wang et al. have applied g-C₃N₄ for the photocatalytic hydrogen evolution from water under visible light. Zhang’s group have fabricated nitrogen-rich CNQDs with tunable fluorescence emission by using a solid-phase reaction strategy. Nevertheless, there has been no report on the use of carbon materials for tuning the morphology or structure of functional materials including photocatalysts.

Up to date, the supramolecular self-assembly route has been increasingly applications for fabrication of various functional materials. In the previous works, we discovered that melamine (MA), an organic molecule consisting of amino groups and aromatic nitrogen atoms, could conduct the self-assembly with various heavy metal ions like Ag⁺ and Cu²⁺ yielding the mesoporous nanocomposites. In the present study, we initially doped CNQDs of different levels into the MA and silver (Ag) matrix by the supramolecular self-assembly route. To our surprise, the yielded mesoporous CNQDs@MA–Ag nanocomposites could present the CNQD-controlled morphologies (i.e., nanowires, nanoflowers, and nanosheets) and especially the tunable visible-light photocatalysis performances depending on the percentages of CNQDs used. It was found that the nanocomposites with the pinecone-like nanoflower structure could exhibit the largest visible-light photocatalysis in splitting water towards H₂O₂ evolution. The main fabrication and photocatalytic H₂O₂ evolution procedures of CNQDs@MA–Ag nanocomposites are schematically illustrated in Scheme 1 with the pinecone-like ones as example. It is thought that the nano-scaled Ag with the surface plasmon resonance (SPR) effect might increase the light harvesting as well as the separation and transmission of electron-hole pairs of CNQDs, leading to the enhanced visible-light-driven photocatalysis of nanocomposites. To the best of our knowledge, this is the first report on the synthesis of the photocatalysts with CNQD-tuned morphological structures and visible-light-driven photocatalysis performances for the efficient water splitting to generate H₂O₂ under sunlight.

Results and Discussion

Fabrication and characterization of CNQDs@MA–Ag with CNQD-tuned morphologies and photocatalysis performances

It is well established that MA with rich nitrogen elements (66.7 %) can conduct the self-assembly with Ag⁺ ions to yield the mesoporous nanocomposites. In the present work, CNQDs were doped with different percentages into the MA and Ag matrix to yield structure-different CNQDs@MA–Ag nanocomposites by the supramolecular self-assembly route, as schematically illustrated in Scheme 1A. Figure 1 displays the SEM images of the morphological structures of the resulting CNQDs@MA–Ag nanocomposites, with the amplified views of SEM images (insert). It was observed that the yielded products could present the changing morphologies, such as nanowires, nanoflowers, and nanosheets, depending on the CNQD percentage from 0 to 40% (Figure 1A–E).

For example, the CNQDs@MA–Ag with the pinecone-like nanoflower structure could be obtained at the CNQD percentage of 30% (Figure 1D). The results indicate that CNQDs with amino groups should participate the reactions between amino...
groups-containing MA and silver by following the supramolecular self-assembly route \(^{26,35}\) in which CN\(_{QDs}\) might competitively interact with silver to induce the MA–Ag reunion reactions with different degrees depending the percentages of CN\(_{QDs}\) used. As a result, the CN\(_{QDs}@MA\)–Ag nanocomposites would be yielded with changing morphologies typically varying from nanowires (Figure 1A) to nanosheets (Figure 1E) as increasing the CN\(_{QDs}\) percentages up to 40%. Also, the polymeric matrix of nitrogen-rich MA could allow for the controlled silver growth in nanocomposites through the strong Ag–MA chelating reactions, which might additionally aid to increase the photostability of nano-scaled Ag during the photocatalytic reactions. Moreover, Figure 2 manifests the TEM images of the pinecone-like nanocomposites by taking CN\(_{QDs}\) as the control. One can note that CN\(_{QDs}\) could be fabricated uniformly with an average size of about 4.3 nm (Figure 2A). A high crystallinity with a lattice parameter of 0.34 nm could be obtained as revealed in the amplified image (Figure 2C), which is in agreement with the 002 plane of graphitic carbon nitride previously reported.\(^{26}\) Furthermore, Figure 2B manifests the TEM image of an individual nanocomposite of CN\(_{QDs}@MA\)–Ag with the pinecone-like structure, in which numerous nano-scaled Ag could be produced in the coagulated form as disclosed in the amplified view (Figure 2D).

The composition of the pinecone-like nanocomposites was investigated separately by elemental mapping and EDS analysis, with the data shown in Figure 3. As expected, the C, N, and Ag elements could be distributed uniformly throughout the nanocomposites. Besides, the Brunauer-Emmett-Teller (BET) analysis was performed for the pinecone-like CN\(_{QDs}@MA\)–Ag nanocomposites in comparison with MA–Ag nanocomposites without CN\(_{QDs}\) (Figure 4). The textural surface parameters obtained from the isotherm data, i.e., BET surface area, mean pore diameter, and total pore volume, are included in Figure 4B. Accordingly, the pinecone-like CN\(_{QDs}@MA\)–Ag nanocomposites can possess apparently the mesoporous characteristics with the average pore diameter of about 5 nm. Especially, they can display the larger BET surface area than the nanocomposites without CN\(_{QDs}\).

More importantly, the as-prepared nanocomposites with different morphological structures could exhibit the CN\(_{QDs}\)-tunable photocatalysis performances, which would be demonstrated in splitting water to generate H\(_2\)O\(_2\) under sunlight afterwards. Figure 5A describes the UV-vis diffuse reflectance spectra (DRS) of CN\(_{QDs}@MA\)–Ag nanocomposites by taking MA–Ag nanocomposites and CN\(_{QDs}\) for comparison. To our surprise, no significant peak of UV-vis diffuse reflectance spectra could be obtained in the visible region for the CN\(_{QDs}@MA\)–Ag nanocomposites (curve c), in contrast to those for MA–Ag (curve a) and CN\(_{QDs}\) (curve b) exhibiting the large peaks in the
visible region. Moreover, as visually illustrated in the photographs (Figure 5A, insert), the MA–Ag nanocomposites could be produced as the white suspensions. When CN$_{QDs}$ were introduced into the MA–Ag reaction matrix, the resulted CN$_{QDs}$@MA–Ag nanocomposites could be formed as the light yellow precipitates, with the changed morphologies revealed in the SEM images above (Figure 1). Accordingly, the CN$_{QDs}$@MA–Ag nanocomposites could be yielded substantially as new nanocomposites during the CN$_{QDs}$-tuned self-assembly of MA and silver matrix, which might otherwise be formed as the nanowires-shaped products in the absence of CN$_{QDs}$ as aforementioned (Figure 1A). Besides, it should be pointed out that when the CN$_{QDs}$@MA–Ag nanocomposites were applied for the photocatalytic splitting of water for H$_2$O$_2$ evolution under sunlight, the enhanced absorption could be observed for the used nanocomposites showing the broadened UV-vis diffuse emission in the visible region (Figure 5A, insert), due to that nano-scaled Ag might be increasingly formed during the illumination procedure. Importantly, such a broadening of the visible-light response region of nanocomposites would help to further improve their light harvesting or the utilization efficiency of solar energy, as confirmed elsewhere for other kinds of photocatalysts.$^{[36–38]}$

In addition, a comparison of the FTIR spectra was conducted among the CN$_{QDs}$@MA–Ag nanocomposites, CN$_{QDs}$ and MA–Ag nanocomposites (Figure 5B). It is found that all of the samples could possess the peaks ranging from 1200 to 1650 cm$^{-1}$ that may belong to the typical stretching modes of C–N heterocycles. The broad peak around 3300 cm$^{-1}$ (curve a) can be assigned to the stretching vibration of -NH or -NH$_2$ groups of MA there. Moreover, as shown in Figure 5B, the amounts of -NH$_2$ groups could be increased in the CN$_{QDs}$@MA–Ag nanocomposites (curve c), which might aid to further improve the hydrophilicity of the nanocomposites to facilitate the promoted electron transferring of nanocomposites, as described for N-doped carbon nanofibers elsewhere.$^{[19]}$ which might lead to the increased photocatalysis of CN$_{QDs}$@MA–Ag nanocomposites for H$_2$O$_2$ evolution afterwards.

Investigations on the photogenerated carriers behaviors of CN$_{QDs}$@MA-Ag nanocomposites containing different CN$_{QDs}$ percentages

The photogenerated carriers behaviors of CN$_{QDs}$@MA–Ag nanocomposites containing different CN$_{QDs}$ percentages (morphological structures) were explored comparably by monitoring the separation rates and transferring performances of photogenerated electron-hole pairs by using electrochemical impedance spectroscopy (EIS) and transient photocurrent experiments (Figure 6). As can be seen from Figure 6A, the lowest resistance of charge transferring, which can be reflected by the frequency semicircles in the EIS Nyquist diagram, could be obtained for the pinecone-like nanocomposites containing 30% CN$_{QDs}$ showing the superior charge transfer capability. Furthermore, Figure 6B reveals the transient photocurrent response curves of the morphology-different nanocomposites on the typical cycles of visible light irradiation on or off, in which the 0% refers to MA–Ag nanocomposites for the blank control. One can note that with the light on, all of the CN$_{QDs}$@MA–Ag nanocomposites could immediately produce the photocurrents under visible light irradiation, in contrast to the MA–Ag nanocomposites with no significant change in photocurrent. The data demonstrate that CN$_{QDs}$ could endow the CN$_{QDs}$@MA–Ag nanocomposites with the enhanced photo-sensitivity and more efficient spatial charge separation. In particular, the pinecone-like CN$_{QDs}$@MA–Ag (containing 30% CN$_{QDs}$) could display the largest transient photocurrents, promising the strongest capacity of photocatalytic splitting water to yield H$_2$O$_2$ under sunlight.

**Photocatalytic H$_2$O$_2$ generation performances of CN$_{QDs}$@MA-Ag nanocomposites containing different CN$_{QDs}$ percentages**

The photocatalytic performances of CN$_{QDs}$@MA–Ag nanocomposites containing different CN$_{QDs}$ percentages (morphological structures) were comparably evaluated by the H$_2$O$_2$ concentrations produced in the photocatalytic water splitting (Figure 7), in which a standard curve for H$_2$O$_2$ determination was scheduled (Figure 7A). Figure 7B shows the variation of H$_2$O$_2$ concentrations for the tested CN$_{QDs}$@MA–Ag photocatalysts containing different CN$_{QDs}$ percentages, taking CN$_{QDs}$ and
The photoresponse of the CN_{QDs}@[MA–Ag] nanocomposites was studied by using the pinecone-like ones as example, in which four consecutive runs of photocatalytic water splitting reactions were performed under visible light irradiation (Figure 7C). It was observed that the H_{2}O_{2} production could show no significant change during the four runs of illumination experiments. Such a high photoresponse of the photocatalysts may promise for long-term applications due to the polymeric matrix of MA for nano-scaled Ag and CN_{QDs} in nanocomposites as aforementioned. Moreover, the trapping experiments were performed for active species yielded in the photocatalytic water splitting reactions separately by using p-benzoquinone (BQ) and isopropanol (IPA) as the capture agents for \( \cdot \text{O}^- \) and \( \cdot \text{OH} \) free radicals, respectively (Figure 7D). It was witnessed that the photocatalytic reactions could be completely restrained by BQ showing the dramatically decreased H_{2}O_{2} amounts. The results indicate that the generation of H_{2}O_{2} could be resulted from the formation of \( \cdot \text{O}^- \) free radicals during the photocatalytic water splitting reactions. To further confirm such a fact, the \( \cdot \text{OH} \) removal experiments were carried out by using the \( \cdot \text{OH} \) quencher of IPA. It was expected, no obvious decrease in the H_{2}O_{2} production was observed. Therefore, the photocatalysis process of CN_{QDs}@[MA–Ag] nanocomposites in splitting water to produce H_{2}O_{2} might be mainly dominated by the reduction of photo-generated electrons rather than the oxidation of holes of the photocatalysts.

CN_{QDs}-tuned formation process and photocatalytic H_{2}O_{2} generation mechanism of CN_{QDs}@[MA–Ag] nanocomposites

Based on the results above, the CN_{QDs}-tuned self-assembly process and photocatalytic H_{2}O_{2} generation mechanism of CN_{QDs}@[MA–Ag] nanocomposites are schematically illustrated in Scheme 1 using the pinecone-like ones as the model. As shown in Scheme 1A, CN_{QDs} synthesized under the NH_{3} atmosphere should possess rich nitrogen atoms that can interact with silver elements, so does the nitrogen-rich polymer molecule of MA. Once CN_{QDs} were introduced into the MA and Ag matrix, a supramolecular self-assembly process would be triggered to yield the nanocomposites showing the different morphological structures regulated by CN_{QDs}. Moreover, a photocatalytic H_{2}O_{2} generation mechanism is thereby proposed for the developed photocatalysts of CN_{QDs}@[MA–Ag] nanocomposites as illustrated in Scheme 1B. Under the visible light irradiation, the CN_{QDs}@[MA–Ag] nanocomposites should be excited to generate the electrons and holes with efficient spatial separation. As revealed by reactive species-trapping experiments above, here-in, the superoxide (\( \cdot \text{O}_2^- \)) so yielded should play a major role in photocatalytic splitting water to generate H_{2}O_{2}. Accordingly, the dissolved oxygen in water should mainly react with the excited electrons to restore O_{2} into \( \cdot \text{O}_2^- \), which should be further reduced into H_{2}O_{2} with the photocatalytic reactions described as follows:

\[
\text{O}_2 + e^- \rightarrow \cdot \text{O}_2^- + e^- \rightarrow H_2O_2 \quad (1)
\]
\[ \text{OH} + h^+ \rightarrow \text{H}_2\text{O}_2 \]  

Besides, it is thought that new photocatalysis active sites of the synergistic CN\textsubscript{QDs}, and nano-scaled Ag might be produced in the CN\textsubscript{QDs}\textsubscript{@MA–Ag} nanocomposites. In particular, during the photocatalytic \( \text{H}_2\text{O}_2 \) generation, the nanocomposites with the pinecone-like structure may enable more exposed photocatalytic active sites and larger specific area for contacting with the dissolved oxygen in water, thus achieving the higher efficiency of \( \text{H}_2\text{O}_2 \) generation under visible light irradiation.

**Conclusions**

In summary, CN\textsubscript{QDs} were successfully doped into the MA–Ag matrix yielding the mesoporous CN\textsubscript{QDs}\textsubscript{@MA–Ag} nanocomposites for the efficient photocatalytic \( \text{H}_2\text{O}_2 \) evolution under sunlight. It was discovered that the resulting nanocomposites could present the different morphologies changing from one-dimensional nanowires to three-dimensional nanoflowers or nanosheets depending on the percentages of CN\textsubscript{QDs} used. More interestingly, CN\textsubscript{QDs}-tuned photocatalysis activities could be expected for the CN\textsubscript{QDs}\textsubscript{@MA–Ag} nanocomposites in splitting water to produce \( \text{H}_2\text{O}_2 \) under visible light. It is thought that the nano-scaled Ag with the SPR effect might increase the light harvesting as well as the separation and transmission of electron-hole pairs of CN\textsubscript{QDs} could present the different morphologies changing from one-dimensional nanowires to three-dimensional nanoflowers or nanosheets depending on the percentages of CN\textsubscript{QDs} used. More interestingly, CN\textsubscript{QDs}-tuned photocatalysis activities could be expected for the CN\textsubscript{QDs}\textsubscript{@MA–Ag} nanocomposites in splitting water to produce \( \text{H}_2\text{O}_2 \) under visible light. It is thought that the nano-scaled Ag with the SPR effect might increase the light harvesting as well as the separation and transmission of electron-hole pairs of CN\textsubscript{QDs} could present the different morphologies changing from one-dimensional nanowires to three-dimensional nanoflowers or nanosheets depending on the percentages of CN\textsubscript{QDs} used. More interestingly, CN\textsubscript{QDs}-tuned photocatalysis activities could be expected for the CN\textsubscript{QDs}\textsubscript{@MA–Ag} nanocomposites in splitting water to produce \( \text{H}_2\text{O}_2 \) under visible light.

**Synthesis of photocatalyst materials**

**Synthesis of CN\textsubscript{QDs}**

Carbon nitride quantum dots (CN\textsubscript{QDs}) were synthesized according to a modified procedure previously reported.\textsuperscript{[26]} Briefly, 0.101 g urea and 0.081 g sodium citrate were thoroughly ground into the powders in an agate mortar in dry environment to be uniformly mixed. Then, the mixed powders were transferred into a Teflon-lined autoclave to be heated at 180 °C for 1 h. After that, the reaction vessel was stewed overnight in the atmosphere NH\textsubscript{3} discharged from urea. Furthermore, the obtained yellowish products were washed for three times with ethanol, and finally dried in an oven at 60 °C for further usage.

**Synthesis of CN\textsubscript{QDs}\textsubscript{@MA–Ag} nanocomposites**

The CN\textsubscript{QDs}\textsubscript{@MA–Ag} nanocomposites were synthesized by the controlled supramolecular self-assembly procedure using CN\textsubscript{QDs} of different percentages. Briefly, under vigorous stirring, an aliquot of MA (50.4 mg) was dissolved in 10 mL water at 60 °C, followed by addition of CN\textsubscript{QDs} of different amounts (9.37, 21.08, 36.13, and 56.2 mg). After being cooled down to room temperature, 10 mL AgNO\textsubscript{3} (20 mM) was separately introduced into each of the mixtures to be aged for 1 h. Afterwards, the mixtures were centrifuged for about 15 min at 4000 rpm, and then washed with water and alcohol, each for three times. Subsequently, the light yellow products of CN\textsubscript{QDs}\textsubscript{@MA–Ag} nanocomposites were collected to be dried under vacuum and then stored in dark for further usage. In addition, the BET surface areas of CN\textsubscript{QDs}\textsubscript{@MA–Ag} nanocomposites were measured by the nitrogen adsorption-desorption method.
Photocatalytic H$_2$O$_2$ generation experiments

The photocatalytic activities of the functional materials were evaluated under the visible light irradiation, in which a 300 W Xenon lamp with a 420 nm cutoff filter was applied as the visible light source. During each of the photocatalytic experiments, an aliquot of 0.020 g photocatalyst samples was initially dispersed into 10 mL water in a container with cooling water jacket outside. Prior to the visible light irradiation, the photocatalytic reaction proceeded in dark by stirring for 1 h to ensure the adsorption-desorption equilibrium among the photocatalysts, dissolved oxygen, and water. Moreover, the continuous irradiation was carried out to trigger the photocatalytic reactions for H$_2$O$_2$ evolution. An aliquot of 0.50 mL suspensions was taken from the reaction system at the given time intervals of one hour. After the centrifugation for removing the photocatalyst precipitations, the fluorescent measurements of the products of catalytic H$_2$O$_2$-HPAA reactions were performed according to a fluorometric analysis method previously reported.¹⁹

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