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A post-grafting strategy to modify g-C$_3$N$_4$ with aromatic heterocycles for enhanced photocatalytic activity

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A novel and facile post-grafting strategy, instead of conventional copolymerization, via Schiff base chemical reaction between aldehyde and -NH$_2$ groups has been developed to construct aromatic heterocycenter-grafted graphitic carbon nitride (g-C$_3$N$_4$) photocatalysts for the first time. The high-resolution N 1s XPS spectrum and the CO$_2$ TPD analysis confirmed the successful introduction of heterocycles and the reduced structure defects (unreacted -NH$_2$ groups during the copolymerization formation of g-C$_3$N$_4$). The post-grafting of aromatic rings into g-C$_3$N$_4$ network did not disrupt the original framework of g-C$_3$N$_4$, but effectively expanded its π-delocalized system, enlarged its surface area and promoted the separation and transfer of photo-excited charge carriers. As a result, the obtained copolymer composites exhibit significantly enhanced visible-light photocatalytic activity for H$_2$ evolution over the pristine g-C$_3$N$_4$. This strategy is general and can be used to graft numbers of aromatic rings of different molecular structures onto g-C$_3$N$_4$.

1. Introduction

Global energy shortage and environmental pollution stimulates scientists to explore new clean and sustainable energies that may replace fossil fuels. Semiconductor-mediated harvest and conversion of solar energy through water photolysis to produce hydrogen has been considered as a sustainable and prospective solution to alleviate the energy crisis and environmental issues. 1-2 Various semiconductors or their composite materials have been developed as possible photocatalysts in past decades. 3-5 Recently, graphitic carbon nitride (g-C$_3$N$_4$), a metal-free photocatalyst, has attracted intensive interest. In detail, g-C$_3$N$_4$ has a bandgap of ca. 2.7 eV and consequently can absorb light up to 450 nm. 6 its suitable band gap, together with its thermal and chemical stability, is especially suitable for applications in photochemistry and photocatalysis. However, it still suffers from a high recombination probability of photoexcited charge carriers (electron-hole), inefficient utilization of visible-light and thus low photocatalytic conversion efficiency. 7 To overcome this problem and improve its photocatalytic performance, numerous methods have been developed, such as synthesizing mesoporous structures 8, 7 or nanocomposite structures, 10 introducing heteroatoms 11-14 copolymerization, 15 coupling with dyes 16 or other semiconductors 8 etc.

Among the various strategies, copolymerization with organic monomers or constructing polymer-polymer composites can effectively modify the electronic properties of this π-conjugated polymer. 15 Hao et al. 17 adopted a facile one-step thermal condensation to synthesize PAN/g-C$_3$N$_4$ composites and effectively improved the interfacial charge transfer. Zhu et al. 18 fabricated P3HT-g-C$_3$N$_4$ heterojunction by assembling p-type P3HT particles on n-type g-C$_3$N$_4$ nanoparticles via a ball milling method. The photocatalytic activity was enhanced since the addition of P3HT increased the n-conjugated length of g-C$_3$N$_4$ network. Zhang et al. 19 incorporated a little amount of pyrimidine rings into the network of g-C$_3$N$_4$ to substitute one nitrogen atom with carbon atom in the tri-s-triazine ring. Wang et al. 20-21 grafted aromatic motifs into the conjugated g-C$_3$N$_4$ networks via one-pot co-condensation of urea (or dicyandiamide) and organic co-monomers (such as 2,4-diaminoquinazoline, 2-aminothiophene-3-carbonitrile, barbituric acid, dianaminoaleneitrile, triophene etc.). It is conducive to extend the delocalization of π electrons, change the intrinsic semiconductor properties, and consequently, greatly improves the photocatalytic activity of g-C$_3$N$_4$. Our group 22, 23 constructed g-C$_3$N$_4$-based copolymer via nucleophilic substitution/addition reaction between aromatic bromides and urea or schiff base chemical reaction between aromatic aldehydes and urea, respectively. All the as-synthesized copolymers showed remarkably enhanced and stable visible-light photocatalytic hydrogen evolution performance. However, organic monomers copolymerization with active groups of precursors may form interface defects (e.g. -NH groups) and destroy reactive centers, which cause polymer growth terminated and tri-s-triazine network interrupted at some point. The reduced
polymerization degree of g-C_{3}N_{4} shows a decrease the photocatalytic activity.

As is well known that, there are abundant residual amino groups on the g-C_{3}N_{4} frameworks due to the incomplete condensation of the precursors such as urea, cynamamide, or melamine etc.\textsuperscript{21} Guo et al\textsuperscript{22} found that the photocatalytic activity of g-C_{3}N_{4} for hydrogen evolution decreased with the increase of –NH_{2} content. These surface exposed terminal –NH_{2} groups have been regarded as defects, where the charge carriers recombine during the photocatalytic reaction. Lowering the defect content in g-C_{3}N_{4} is therefore conducive to improve its photocatalytic performance. Fortunately, –NH_{2} groups are very reactive and can participate in many types of reactions. It means that we can diminish this kind of defect to a large extent in g-C_{3}N_{4} based on suitable amino chemical reactions. Wang et al\textsuperscript{23} synthesized PNA–g-C_{3}N_{4} through post-treatment via the hydrogen bonding interaction of p-nitrobenzoic acid (PNA) and g-C_{3}N_{4} to improve the photo-degradation property for methyl orange. However, such a weak binding force is unstable and easily broken.

Here in this work, we have successfully developed a series of novel g-C_{3}N_{4}-based composite photocatalysts based on the Schiff base interaction of aldehyde groups and surface exposed terminal amino groups in g-C_{3}N_{4} after the thermal polymerization of urea was complete. In this way, π-delocalized system of g-C_{3}N_{4} has been expanded by the grafted aromatic rings through covalent C=N bond. In the mean time, the amount of the residual amino groups have been largely reduced. As expected, compared with the pristine g-C_{3}N_{4}, the obtained composites show remarkably enhanced photocatalytic H_{2} evolution performance by water-splitting under visible light. According to our knowledge, this is the first report on the covalent post-grafting modification of g-C_{3}N_{4} instead of most reported copolymerization approaches, for improved visible-light photocatalytic hydrogen evolution.

2. Experimental and theoretical methods

2.1 Chemicals and reagents.

Urea (>99%) was purchased from Sinopharm Chemical Reagent Co., Ltd.; terephthalaldehyde (>98%), 2-naphthaldehyde (>98%), phenanthrene-3-carboxaldehyde (>97%), pyrrole-2-carboxaldehyde (>98%), indole-3-carboxaldehyde (>98%), 2-quinoxalinecarboxaldehyde (C_{9}H_{8}NO) (>96%) and benzo[b]thiophene-3-carboxaldehyde (C_{9}H_{8}OS) (>98%) were purchased from TCI Co., Ltd.; 2,5-thiophenedicarboxaldehyde (C_{7}H_{10}O_{2}) (>99%), triethanolamine (>97%) and chloroalatitic acid hexahydrate (>37.5%, Pt basis) were purchased from Sigma–Aldrich Co., LLC. All chemicals were used as received without further purification.

2.2 Sample preparation.

2.2.1 Preparation of g-C_{3}N_{4}. Sample g-C_{3}N_{4} was prepared by thermally decomposing urea at 550 °C in an alumina crucible with a cover for 2 h in static air with a ramp rate of 5 °C min\textsuperscript{-1}. Then, the crucible was cooled to room temperature naturally. The resulting yellow product was collected and ground into fine powder for further use and was denoted as CN.

2.2.2 Post-grafting modification of g-C_{3}N_{4}. Various aromatic aldehydes were used to post-modify g-C_{3}N_{4}. Taking terephthalaldehyde as an example, in a typical procedure, 1 g of g-C_{3}N_{4} and a certain amount of terephthalaldehyde were mixed thoroughly. Then the mixture was put in an alumina crucible with a cover and heated to 250 °C (the boiling point temperature of aromatic aldehydes) for 5 h at a heating rate of 5 °C min\textsuperscript{-1} in air. The products were denoted as CNTE-x, in which x represents the weight percentage of terephthalaldehyde in the mixture (x% = 0.1, 0.5, 1, 5 and 10%).

2.2.3 Post-heating treatment of g-C_{3}N_{4}. As comparison, the obtained g-C_{3}N_{4} (CN) was heated at 250 °C in an alumina crucible with a cover for 5 h in static air with a ramp rate of 5 °C min\textsuperscript{-1}. The resulting product was denoted as CN-250.

2.2.4 Copolymerization modification of g-C_{3}N_{4}. For comparison, aromatic ring copolymerized g-C_{3}N_{4} was synthesized according to our earlier report.\textsuperscript{21} Typically, 20 g urea and 2 mg terephthalaldehyde were mixed thoroughly and then the mixture was put in an alumina crucible with a cover and heated to 550 °C for 2 h at a heating rate of 5 °C min\textsuperscript{-1}. The resulting product was denoted as CNAL-2.

2.3 Characterization.

Elemental analysis (C, H, N) was performed on a Vario EL microanalyzer. X-ray diffraction (XRD) measurements were conducted on a Rigaku D/Max 2200PC diffractometer (Cu Kα radiation). Fourier transformed infrared (FTIR) spectra were recorded with a Nicolet i510 FTIR spectrometer. X-ray photoelectron spectroscopy (XPS) measurement was carried out on a Thermo Scientific Escalab 250 spectrometer with Al Kα radiation as the excitation source. Binding energies for the high resolution spectra were calibrated by setting C 1s to 284.6 eV. Transmission electron microscopy (TEM) images and electron energy loss spectroscopy (EELS) results were obtained with a FEI Magellan 400 instrument. The BET surface areas were measured on a Micromeritics Tristar 3000 system. The UV-visible absorption spectra were recorded in the range of 300-800 nm with a UV-3600 PC Shimadzu spectroscope using BaSO_{4} as a reference. Photoluminescence spectra (PL) of the samples were obtained at room temperature excited by incident light of 380 nm using a fluorescence spectrometer (Shimadzu RF-5301PC). The temperature-programmed desorption of CO\textsubscript{2} (CO\textsubscript{2}-TPD) was carried out on a Micromeritics Chemisorb 2750. The zeta potentials of the samples in water were measured using a Zetasizer nano ZS90 analyzer (Malvern Instruments Ltd., UK).

2.4 Electrochemical Analysis.

Electrochemical measurements were conducted on a CHI 760E electrochemical workstation (Chenhua Instruments, Shanghai, China) with a standard three-electrode cell, which employed an FTO electrode deposited with samples as the working electrode, a platinum sheet as the counter electrode and saturated Ag/AgCl as the reference electrode. A 300 W Xe lamp (PLS-SEX300C, Perfectlight Limited, Beijing) with a 420 nm cut-off filter was used as visible light source. The working electrodes were prepared by electrophoretic deposition in an acetone solution (25 mL) containing sample powder (20 mg) and iodine (40 mg). The FTO
glass was kept in the above solution for 5 min with a 10 V bias under potentiostat control. After being calcined for 2 h in an oven at 150 °C, a homogeneous film was obtained. 50 mL of 0.2 M Na$_2$SO$_4$ was used as electrolyte solution, which represented the best compromise between activity and stability of the composites.

### 2.5 Photocatalytic hydrogen evolution test.

The visible light-induced H$_2$ evolution was carried out in a Pyrex top-irradiation reaction vessel connected to a closed glass gas-circulation system (Lab-Solar-III AG, Perfectlight Limited, Beijing). A 300 W xenon lamp (CEL-HXF300, Ceaulight, Beijing) with a 420 nm cut-off filter was chosen as a visible light source, and the light intensity was 230 mW cm$^{-2}$ (tested by FieldMaxII-TO, Coherent). Photocatalyst (100 mg) was suspended in an aqueous solution (100 mL) containing triethanolamine (10 vol %), and 3 wt% Pt was loaded on the surface of the catalyst by in situ photodeposition using H$_2$PtCl$_6$ as the precursor. The reactant solution was evacuated several times to remove air prior to the irradiation experiment. The temperature of the reaction solution was maintained at 10 °C by a flow of cooling water during the photocatalytic reaction. The evolved gases were analyzed by gas chromatography (GC7900, Techcomp) equipped with a thermal conductive detector (TCD) and a 5 Å molecular sieve column, using nitrogen as the carrier gas.

### 3. Results and discussion

The post-grafting incorporation of aromatic rings into the conjugated system of g-C$_3$N$_4$ based on Schiff base reaction of aromatic aldehydes and the terminal –NH$_2$ groups is depicted in Fig. S1. The g-C$_3$N$_4$ sample was synthesized by the thermal polymerization of urea at elevated temperature. Because of the kinetic problems during the thermo-polymerization process of urea, it is difficult to obtain ideally condensed and completely crystallized g-C$_3$N$_4$ phase, leaving a number of -NH$_2$ groups at the edge of s-triazine ring (Fig. S1A). Then aldehyde groups in terephthalaldehyde can be subjected to shiff base reaction with the marginal –NH$_2$ groups in g-C$_3$N$_4$ to form C=N bonds. Consequently, benzene rings can be grafted onto g-C$_3$N$_4$ network by dint of this chemical post-grafting method, creating surface/interfacial heterojunctions as well as reducing amino defects. Results of elemental analysis are shown in Table 1, as expected, the molar ratio of C to N gradually increases with more benzene rings being incorporated. For example, the C/N molar ratio increases from 0.692 of the pristine g-C$_3$N$_4$ (CN) to 0.717 of CNTE-10, indicating successful grafting of benzene rings into the network of g-C$_3$N$_4$.

The X-ray diffraction (XRD) patterns of CN and CNTE-x samples are showed in Fig. S2a. All the samples exhibit two distinct diffraction peaks, which can be ascribed to the typical diffraction peaks of g-C$_3$N$_4$, indicating that the molecular framework of g-C$_3$N$_4$ has been well retained after the post-grafting modification. The stronger peak at around 27.6° represents the (002) interlayer reflection of the graphitic-like structure with d=0.322 nm. The other small reflection peak at about 12.8° is attributed to the (100) inplane repeated units of tri-s-triazine. Notably, the (002) peak has shifted from 27.68° to 27.58° with the gradual increase of benzene ring content, corresponding to the increase of the interplanar distance from 0.322 nm of CN to 0.323 nm of CNTE-10. This indicates that the post-grafted benzene rings have led to slightly expanded packing of the graphitic-like g-C$_3$N$_4$ layers. This increase in the interplanar distance may improve the accessibility of the active sites on g-C$_3$N$_4$ layers thus enhance the photocatalytic activity. Additionally, no apparent diffraction peaks of terephthalaldehyde can be observed in all the CNTE-x samples, suggesting no aggregation of terephthalaldehyde in the g-C$_3$N$_4$ matrix.
The multiple bands between 1200-1600 cm$^{-1}$ are identified as the aromatic carbon atoms and the sp$^2$ deconvoluted into two peaks centred at 284.6 and 288.0 eV, which was carried out and the corresponding spectra are shown in Fig. 2. XPS spectra reveals noticeable difference between CN and CNTE-1 (Fig. 2). After further analysis of the obtained data, the peak-area ratio of CN and CNTE-1 samples, X-ray photoelectron spectroscopy (XPS) and FT-IR results. As shown in Fig. 2b, the high-resolution N 1s spectra shows four main signals at 398.5 eV, N-C, 399.9 eV, N-H, 401.0 eV, and 404.2 eV, respectively. The main signal shows the occurrence of the sp$^2$ hybridized nitrogen in s-triazine rings (C-N=Cs, 398.5 eV, N-Cs) and tertiary nitrogen groups (N-(C)N, 399.9 eV, N-N). Both of them, together with the aforementioned sp$^3$ hybridized carbon (288.0 eV) constitute the s-heptazine heterocyclic ring units as elementary building blocks of the CN polymers. The CO$_2$ molecules.

In order to further check the chemical nature of the obtained CN and CNTE-1 samples, X-ray photoelectron spectroscopy (XPS) was carried out and the corresponding spectra are shown in Fig. 2. The high-resolution C 1s signals (Fig. 2a) of the two samples can be deconvoluted into two peaks centred at 284.6 and 288.0 eV, which are identified as the aromatic carbon atoms and the sp$^2$ hybridized carbon in the s-triazine ring, respectively. It is worth noting that the peak at 284.6 eV (i.e., the aromatic carbon atoms, 7.6%) of CNTE-1 is much stronger than that of CN (5.6%), suggesting again aromatic rings have been introduced into the g-C$_3$N$_4$ matrix. As shown in Fig. 2b, the high-resolution N 1s spectra shows four peaks centred at about 398.5, 399.9, 401.0 and 404.2 eV, respectively. The main signal shows the occurrence of the sp$^2$ hybridized nitrogen in s-triazine rings (C-N=Cs, 398.5 eV, N-Cs) and tertiary nitrogen groups (N-(C)N, 399.9 eV, N-N). Both of them, together with the aforementioned sp$^3$ hybridized carbon (288.0 eV) constitute the s-heptazine heterocyclic ring units as elementary building blocks of the CN polymers. Another weak peak at 401.0 eV indicates the defective amino groups (C-NH, N-N). The CNTE-1 shows that the specific surface area and pore volume increases firstly and then decreases with the increased amount of grafted aromatics rings. The CNTE-1 samples shows a maximum specific surface area (120.31 m$^2$ g$^{-1}$), which is remarkably higher than that of CN (65.20 m$^2$ g$^{-1}$). The increased BET surface area might be attributed to the exfoliation and tailoring of the bulk layered structure of g-C$_3$N$_4$ during the post-treatment process. However, grafting excessive aromatic rings may lead to over-distortion or curvature of g-C$_3$N$_4$ layers, which decreases the specific surface area.

Due to the incomplete condensation of urea, there are high amounts of residual –NH$_2$ groups in the polymeric networks of g-C$_3$N$_4$, which generally become the recombination sites of photogenerated charge carriers in photocatalytic reactions. These –NH$_2$ groups can also act as Lewis basic sites and adsorb slightly acidic CO$_2$ molecules. The CO$_2$ temperature programmed desorption (TPD) measurement was conducted on CN and CNTE-1. As shown in Fig. 3, CN sample shows a larger peak area of CO$_2$ desorption than CNTE-1, suggesting a decreased content of surface –NH$_2$ groups on CNTE-1 in comparison with CN. We further tested zeta potentials of CN and CNTE-1 samples dispersed in water. Because of the free lone pair electrons on nitrogen atoms, –NH$_2$ groups can act as proton acceptors and acquire positive surface charges. It can be seen from Fig. S4, the zeta potential of CNTE-1 (-41.1 mV) is more negative than that of CN (-22.2 mV). This further proves the significantly decreased content of terminal –NH$_2$ groups after post-grafting aromatic rings. The analyses above support the reaction mechanism depicted in Fig. S1. It can be concluded that the aromatic rings have been successfully grafted onto the g-C$_3$N$_4$ networks.

### Table 1. Bulk contents of C, N, and H by elemental analysis, textural parameters, and band gap of CN and CNTE-x samples

<table>
<thead>
<tr>
<th>Sample</th>
<th>C (%)</th>
<th>N (%)</th>
<th>H (%)</th>
<th>C/N atomic ratio</th>
<th>BET (m$^2$ g$^{-1}$)</th>
<th>Pore volume (cm$^3$ g$^{-1}$)</th>
<th>$E_g$ (eV)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CN</td>
<td>35.08</td>
<td>59.16</td>
<td>1.73</td>
<td>0.692</td>
<td>65.20</td>
<td>0.50</td>
<td>2.65</td>
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<tr>
<td>CNTE-0.1</td>
<td>35.27</td>
<td>59.29</td>
<td>1.83</td>
<td>0.694</td>
<td>66.10</td>
<td>0.52</td>
<td>2.64</td>
</tr>
<tr>
<td>CNTE-0.5</td>
<td>35.45</td>
<td>59.03</td>
<td>1.77</td>
<td>0.701</td>
<td>83.66</td>
<td>0.61</td>
<td>2.64</td>
</tr>
<tr>
<td>CNTE-5</td>
<td>35.80</td>
<td>59.26</td>
<td>1.71</td>
<td>0.705</td>
<td>120.31</td>
<td>0.88</td>
<td>2.62</td>
</tr>
<tr>
<td>CNTE-10</td>
<td>35.78</td>
<td>58.39</td>
<td>1.63</td>
<td>0.710</td>
<td>100.28</td>
<td>0.81</td>
<td>2.62</td>
</tr>
</tbody>
</table>

Fig. S2b shows the FT-IR spectra of CN and CNTE-x samples. All of the samples exhibit several typical absorption bands revealing the characteristic structure of g-C$_3$N$_4$. The intense peak at ca. 810 cm$^{-1}$ is assigned to the out-of-plane deformation vibration modes of the s-triazine ring. The multiple bands between 1200-1600 cm$^{-1}$ correspond to the stretching modes of aromatic C-N heterocycle.$^{12}$ The broad peaks at around 3100-3400 cm$^{-1}$ are assigned to stretching vibration modes of the -NH group and -OH group of the adsorbed H$_2$O from air, which become weaker after post-grafting modification, indicating the decreased content of surface amino groups in CNTE-x. The analyses above support the reaction mechanism depicted in Fig. S1. It can be concluded that the aromatic rings have been successfully grafted onto the g-C$_3$N$_4$ networks.

The morphology of CN, CNTE-1, and CNTE-10 samples were investigated by TEM. As shown in Fig. 4, all samples exhibit typical layered structure. Whereas, after the incorporation of aromatic rings, a curved/distorted structure dominates. The EELS spectra of these samples were collected at the layer edges, where amino groups were located, and the results are summarized in Fig. 4d. Not surprisingly, the C/N atomic ratio increases with the increased incorporation of benzene rings, further indicating the aromatic rings have been hung to the position of surface amino groups. Table 1 show that the specific surface area and pore volume increases firstly and then decreases with the increased amount of grafted aromatics rings. The CNTE-1 samples shows a maximum specific surface area (120.31 m$^2$ g$^{-1}$), which is remarkably higher than that of CN (65.20 m$^2$ g$^{-1}$). The increased BET surface area might be attributed to the exfoliation and tailoring of the bulk layered structure of g-C$_3$N$_4$ during the post-treatment process. However, grafting excessive aromatic rings may lead to over-distortion or curvature of g-C$_3$N$_4$ layers, which decreases the specific surface area.
Fig. 3 CO₂ TPD profiles of CN and CNTE-1.

The optical absorptions of CN and CNTE-x composites with different amounts of aromatic rings grafted were characterized using UV-vis absorption spectra. It can be seen from Fig. 5a that, compared with CN, the incorporation of aromatic rings can effectively improve the light absorption in both UV and visible region. Moreover, CNTE-x sample with increased amounts of aromatic rings displays intensified and red-shifted light absorption. This result may be attributed to the following two aspects. One is the extended electron delocalization in the 2D networks of g-C₃N₄ because of the incorporation of aromatic rings and the other is the activation of n to π* electron transition because of the distortion of the co-polymeric units in g-C₃N₄ networks. The band gap of the obtained samples has been estimated by a classical Tauc approach (Fig. S5) and shown in Table 1. E_g value decreases from 2.65 eV of CN to 2.60 eV of CNTE-10, implying that the electronic structure of g-C₃N₄ has been remarkably modified by the grafting of aromatic rings. And at the same time, the visible light harvesting of the photocatalyst has also been enhanced.

Photoluminescence spectra (PL) were performed to determine the separation/recombination of photogenerated charge carriers. In Fig. 5b, the CNTE-x samples have similar PL spectra to CN, but the PL emission intensity of the CNTE-x samples gradually decreases as the amount of grafted aromatic rings increases. Because grafting aromatics into carbon nitride networks can extend conjugated π-electron system and make the carbon nitride layer more curved/distorted. The distortion of CNTE-x layers increases the intramolecular collision frequency, resulting in the rapid loss of exciton energy and the aggravation of PL quenching. This indicates that the grafting of aromatic rings has effectively improved the separation of photogenerated charge carriers in the modified g-C₃N₄ photocatalysts under visible-light illumination.

Based on the above discussion, it is safe to conclude that the grafted aromatics onto the g-C₃N₄ framework has brought following benefits for photocatalysis: (1) enlarged specific surface area, which exposes more catalytic active sites; (2) red-shift of absorption edge, which improves the visible light utilization; (3) a lower density of amino defects, which decreases the recombination possibility of charge carriers in photocatalysis reactions. The photocatalytic hydrogen evolution rates of all the samples are presented in Fig. 6a. Significantly enhanced hydrogen evolution rate can be found on all of the CNTE-x samples, especially for CNTE-1, which has the highest photocatalytic performance with a H₂ evolution rate of 92.4 μmol h⁻¹, above 4.1 times that of CN (22.2 μmol h⁻¹) among the samples prepared. When the dosage of terephthalaldehyde is higher than 1 wt%, the photocatalytic H₂ evolution will decrease with the further increased addition of terephthalaldehyde. This can be ascribed to the over-aggravated distortion of g-C₃N₄ layers with more aromatics grafted, resulting in decreased surface area, easier dissipation of photo-generated charge carriers into vibrational or thermal energy, and thus gradually decreased photocatalytic activity. In short, a proper amount of aromatic rings is important to optimize the photocatalytic activity of CNTE-x photocatalysts. Fig. S6 shows that the ratio of BET surface area and the ratio of H₂ evolution rate of CNTE-x/CN evolve in a similar way, indicating that large surface area would benefit the enhancement of photocatalytic performance. However, the H₂ evolution rate normalized to the BET surface area...
(H₂ evolution rate ratio/surface area ratio) of CNTE-x (especially CNTE-1) is considerably higher than 1 of CN, suggesting that the surface area can hardly play a decisive role in this photocatalytic system and the other factors also contribute much to the enhanced photocatalytic activity.²⁵,³⁹ It is well known that the photo-reduction ability of the semiconductor is determined by the potential of its conduction band (LUMO).⁴² Based on the Mott-Schottky plots (Fig. S7), the LUMOs of CN and CNTE-1 are estimated to be -1.02 eV and -1.15 eV, respectively. The higher LUMO of CNTE-1 implies the stronger reduction ability of its photo-induced electrons than those in CN. The stability of CNTE-1 has been also demonstrated by repeating the photocatalytic experiments under the same condition for four cycles (16 h) (Fig. 6b). Clearly, there is no significant decrease in the H₂ evolution rate after four consecutive cycles. Furthermore, after photocatalytic reactions, the used CNTE-1 sample was collected by centrifugation. Fig. S8 compares the XRD patterns and FT-IR spectra of CNTE-1 sample before (fresh CNTE-1) and after (used CNTE-1) the photocatalytic reaction. It can be found that, there are no noticeable alternations in the crystal and chemical structure, suggesting the excellent reusability and stability of this photocatalyst.²³

To further explore how the grafting of aromatic rings promotes the photocatalytic performance of g-C₃N₄, several other analyses were performed to characterize the post-heating treatment sample CN-250. As indicated by XRD, FT-IR, UV-vis absorption, PL spectra and N₂ adsorption-desorption (Fig. S9a-e), the crystal structure, morphology, optical absorption/luminescence properties, the surface areas and pore structures have not changed much after post-heat treatment. For comparison, the photocatalytic H₂ evolution on CN-250 was also examined. Only a very slight improvement in photocatalytic activity was obtained, which is far less significant than that enhanced by the post-grafting of aromatic rings (Fig. S9f). This provides powerful evidences that the remarkably enhanced photocatalytic performance of CNTE-x samples has nothing to do with the post-heating treatment.

The electrochemical impedance spectra (EIS) were recorded to investigate the electrochemical properties of the samples. In general, the radius of the arc reflects the reaction rate on the surface of the electrode. As shown in Fig. 7a, the arc radius of the samples follows the order CN < CNAL-2 < CNTE-1, suggesting that post-grafting modification of g-C₃N₄ can more efficiently promote the separation and transfer of photogenerated charge carriers at the material surface/interface than copolymerization.⁴³ Both for post-grafting and copolymerization modified g-C₃N₄, surface/interfacial heterojunctions can be created, so the spatial separation of photogenerated charge carriers can be improved. However, during the copolymerization formation of g-C₃N₄, there are more defects derived from incomplete heptazine coupling, which act to increase the charge carrier recombination rate and thus hinder the efficient utilization of charge carriers. In comparison, post-grafting modification of g-C₃N₄ dissipates terminal –NH₂ defects without deceasing its crystallinity, meanwhile, introduces electron acceptors to enhance charge transfer transition and consequently improves the separation/transfer and surface reactivity of charge carriers. Furthermore, the considerably enhanced photocurrent generated (Fig. 7b) on CNTE-1 also indicates the substantially accelerated charge separation and transfer, well in line with the above results.
To confirm that the hydrogen evolution reaction is indeed driven by light absorption on the catalyst, wavelength dependence of the photocatalytic H₂ evolution on CNTE-1 was examined using different wavelength optical filters, and its wavelength-dependent apparent quantum yield (AQY) is shown in Fig. 8, which is well coincident with its optical absorption spectrum, suggesting that it is the harvested visible photons that dominate the driving energy force of the photocatalytic reaction.44

To illustrate the general feasibility of this post-grafting strategy based on Schiff base chemistry, a series of other aromatic molecules with active aldehyde groups (e.g. 2,5-thiophenedicarboxaldehyde (C₆H₄O₂S), 2-quinoxalinecarboxaldehyde, (C₆H₄HNO) and benzo[b]thiophene-3-carboxaldehyde (C₆H₄O₃S) etc.), instead of terephthalaldehyde, were chosen as precursors to post-graft thiophenedicarboxaldehyde (C₆H₄O₂S) molecules with active aldehyde groups (e.g. 2,5-thiophenedicarboxaldehyde (C₆H₄O₂S), 2-quinoxalinecarboxaldehyde, (C₆H₄HNO) and benzo[b]thiophene-3-carboxaldehyde (C₆H₄O₃S) etc.), instead of terephthalaldehyde, were chosen as precursors to post-graft other kinds of aromatic rings onto g-C₃N₄ at their individual boiling points. As shown in Fig. 9, all the carbon nitride copolymers incorporated with these aromatic rings show enhanced hydrogen evolution activity. Moreover, the considerably decreased arc radius of Nyquist plots and the enhanced photocurrent of these carbon nitride copolymers implies their improved electronic conductivity, and the promoted separation and transfer of the charge carriers between them. The photocatalytic activities of these modified carbon nitride polymers are mainly determined by the monocone of aromatic molecules as well as their optimized dosages.

4. Conclusions
A novel post-grafting strategy, rather than copolymerization, based on Schiff base chemistry has been developed to fabricate novel aromatics-grafted carbon nitrides. The introduction of aromatics enables to modify the intrinsic electronic properties, enhance the visible light response of g-C₃N₄, and meanwhile, reduce amino defects on its surface, which finally promote the separation and transfer of photogenerated charge carriers and the photocatalytic activity enhancement. Hence the as-obtained copolymers show substantially enhanced photocatalytic hydrogen evolution performance compared with that of the pristine g-C₃N₄. Through this facile strategy, large numbers of aromatics can be grafted onto g-C₃N₄ networks. We anticipate that this post-modification strategy will provide novel routes to construct efficient carbon-nitride-based copolymer photocatalysts with reduced structure defects and enhanced photocatalytic activities.

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Notes and references


A novel and facile post-grafting strategy has been developed to construct aromatic hetercycle-grafted graphitic carbon nitride photocatalysts.