Half-metallic carbon nitride nanosheets with microgrid mode resonance structure for efficient photocatalytic hydrogen evolution

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Photocatalytic hydrogen evolution from water has triggered an intensive search for metal-free semiconducting photocatalysts. However, traditional semiconducting materials suffer from limited hydrogen evolution efficiency owing to low intrinsic electron transfer, rapid recombination of photogenerated carriers, and lack of artificial microstructure. Herein, we report a metal-free half-metallic carbon nitride for highly efficient photocatalytic hydrogen evolution. The introduced half-metallic features not only effectively facilitate carrier transfer but also provide more active sites for hydrogen evolution reaction. The nanosheets incorporated into a micro grid mode resonance structure via in situ pyrolysis of ionic liquid, which show further enhanced photoelectronic coupling and entire solar energy exploitation, boosts the hydrogen evolution rate reach up to 1009 μmol g⁻¹ h⁻¹. Our findings propose a strategy for micro-structural regulations of half-metallic carbon nitride material, and meanwhile the fundamentals provide inspirations for the steering of electron transfer and solar energy absorption in electrocatalysis, photoelectrocatalysis, and photovoltaic cells.
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	on the journey to pursue renewable green energy to solve the global energy crisis and environmental pollution, hydrogen is regarded as an ideal energy owing to its environmental friendliness and high energy capacity\textsuperscript{1,2}. It is of great interest to utilize solar energy to acquire hydrogen from water, especially in light of the environmental cost. However, in many applications, the lack of stable, efficient, and inexpensive catalysts for such systems remains a principal problem. Inspired by rapidly developing exploratory study, the economical and abundant metal-free photocatalysts, such as carbon quantum dots\textsuperscript{3,4}, black phosphorus\textsuperscript{5,6} and graphitic carbon nitrides\textsuperscript{7,8,9} have been proposed to replace the metal-containing, especially for noble metal Pt catalyst. Among the above metal-free semiconducting materials, photocatalysts with large number of exposed active sites, low resistance for carrier transfer and wide solar light response have been identified as promising candidates for hydrogen evolution reaction (HER)\textsuperscript{10}. In this context, the introduction of metal-free half-metallic feature\textsuperscript{11-13} (a material with complete spin polarization at Fermi level: the majority is metallic and the minority is semiconducting) into semiconducting nanosheets can meet the above-mentioned requirements. The half-metallicity, ensuing both the maintenance of quick electron transfer and efficient electron-hole separation owing to introduction of triplet and singlet excited states, is conducive to solar exploitation\textsuperscript{14}. More importantly, some new active sites are expected to appear on half-metallic nanostructure surfaces.

Herein, we present a solar \text{H}_2\text{ production system using the half-metallic carbon nitride [hm-C(CN)\textsubscript{3}] nanosheets incorporated into an artificial nanotube array as the photocatalyst. The designed various nanostructures are widely used in solar cells to enhance light absorption\textsuperscript{15-18}. In our system, this artificial nanotube array acts as not only micro grid mode resonance to tune optoelectronic coupling process for enhanced utilization of solar energy but also the template to control the size of carbon nitride nanosheets. Such a use of the half-metallic carbon nitride nanosheets into a photocatalytic system in combination with the micro grid artificial microstructure leads to efficient HER.

## Results

### Preparation and characterization of half-metallic carbon nitride samples

The density functional theory (DFT) prediction was implemented and the calculated results show that the layered carbon nitride [C(CN)\textsubscript{3}] has the expected intrinsic half-metallicity and suitable band gap (2.06–2.33 eV) to harvest more solar energy (Supplementary Fig. 1)\textsuperscript{14}. The incorporation of two-dimensional (2D) hm-C(CN)\textsubscript{3} nanosheets into the artificial nanotube arrays (nanoporous anodic aluminum oxide membrane, AAM) [named as MG@hm-C(CN)\textsubscript{3}] to form micro grid mode resonance structure was achieved via in situ pyrolysis of imidazolium-based ionic liquid, and meanwhile the bulk hm-C(CN)\textsubscript{3} samples were prepared without AAM for characterization and comparison. Here, the pyrolytic condition for hm-C(CN)\textsubscript{3} sample was determined via the scanning thermal gravimetric analysis of the ionic liquid precursor (Supplementary Fig. 2). Detailed description is given in the Methods and Supplementary Fig. 3. Energy dispersive X-ray (EDX) spectra disclose that hm-C(CN)\textsubscript{3} bulk sample and nanosheets derived from the ionic liquid precursor have the same C/N ratios (~ 58 at% for C and ~ 42 at% for N). These values are close to 4/3 which is the expected value for hm-C(CN)\textsubscript{3} sample.

The experimental and calculated X-ray diffraction (XRD) patterns (Fig. 1a) confirm that the strongest diffraction peak located at ~ 25.75° is attributed to the (002) peak of bulk hm-C(CN)\textsubscript{3} and the high angle peak at ~ 43.86° is indexed into (200) peak, respectively\textsuperscript{19}. The photo of the hm-C(CN)\textsubscript{3} powder in the inset displays a black metallic luster, which is favorable to solar absorption\textsuperscript{20}. The bulk hm-C(CN)\textsubscript{3} has a sheet-like shape with a planar size of 5–250 μm (Supplementary Fig. 4a). Atomic force microscopy (AFM) image indicates these sheets are of 0.7 ~ 1.1 μm in thickness (Supplementary Fig. 4b). EDX analysis confirms that these hm-C(CN)\textsubscript{3} sheets are mainly composed of C and N elements (Supplementary Fig. 5). The corresponding element mapping images in Supplementary Fig. 6 show a homogeneous nature of the as-prepared sheets, in which C and N are uniformly distributed in the mapping image of hm-C(CN)\textsubscript{3}. XPS further

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**Fig. 1** Characterization of hm-C(CN)\textsubscript{3}. a XRD patterns of the bulk hm-C(CN)\textsubscript{3} materials (Scale bar, 1 cm). b FTIR spectra of g-C\textsubscript{3}N\textsubscript{4} and bulk hm-C(CN)\textsubscript{3}. c Schematic representation of chemical structure for hm-C(CN)\textsubscript{3} material. d Top panel: applied TEM image (Scale bar, 20 nm) and SAED pattern of R@hm-C(CN)\textsubscript{3} nanosheets. Bottom panel: TEM image of R@hm-C(CN)\textsubscript{3} nanosheets (Scale bar, 200 nm). e HR-TEM image of R@hm-C(CN)\textsubscript{3} nanosheets (Scale bar, 2 nm). Left inset: locally amplified HR-TEM image (Scale bar, 1 nm). Right inset: atomic structural model. f ESR spectra of g-C\textsubscript{3}N\textsubscript{4} and hm-C(CN)\textsubscript{3} nanosheets.
evidences the element composition and the successful preparation of hm-C(CN)$_3$ (Supplementary Fig. 7). To better understand the structure of hm-C(CN)$_3$, the Fourier transform infrared (FTIR) spectra were provided to compare with g-C$_3$N$_4$. As shown in Fig. 1b, the FTIR spectrum of g-C$_3$N$_4$ shows strong vibration bands at 1200–1650 cm$^{-1}$ (marked by black box) and 809.9 cm$^{-1}$, which are attributed to the stretching vibrations of the CN heterocycles and out of plane bending vibration of the aromatic ring, respectively.\cite{9,20,21} However, the FTIR spectrum of hm-C(CN)$_3$ mainly displays the stretching vibrations of C-S(C)$_3$ (669.1 cm$^{-1}$), CN (1000–1500 cm$^{-1}$) (marked by black box) and C=N (1608.3 cm$^{-1}$). Obviously, the difference in vibrational energy is ascribed to their different chemical structures. Based on XRD, XPS analysis and previous report\cite{19}, the chemical structure of hm-C(CN)$_3$ can be inferred as shown in Fig. 1c, which is further confirmed by the solid state $^{13}$C nuclear magnetic resonance (NMR) spectra (Supplementary Fig. 8).

The transmission electron microscope (TEM) images indicate that the obtained hm-C(CN)$_3$ nanosheets after facile removal of AAM using 0.5 M NaOH solution emerge with a planar size of 20–60 nm (bottom panel in Fig. 1d). These nanosheets, without further liquid exfoliation, have much smaller size than that of bulk hm-C(CN)$_3$, demonstrating that the structure-assisted in situ pyrolysis of ionic liquid is a potential strategy to obtain carbon nitride nanosheets. The selected area electron diffraction pattern reveals that faint diffraction rings are assigned to the [002] and [200] with lattice spacing of 0.47 and 0.21 nm (upside in Fig. 1d), indicating the polycrystalline structure of hm-C(CN)$_3$ nanosheets. This is well consistent with the broad diffraction peaks from bulk hm-C(CN)$_3$ in XRD pattern. The high-resolution transmission electron microscope (HR-TEM) image for a typical crystalline region shows the arrangement of atoms (left inset in Fig. 1e), which is according to the theoretical simulation displayed in the right inset of Fig. 1e. It is worthwhile pointing out that some disordered structures may exist in our system, leading to more dangling bonds\cite{19,22}, which can improve optical absorption (Supplementary Fig. 9). However, this disordered structure makes the protons difficult to bond to the catalyst surface, resulting in the slow HER kinetics (Supplementary Fig. 10). Therefore, we point out that the main contribution to the HER is from active sites at crystalline region of hm-C(CN)$_3$ material. As shown by electron spin resonance (ESR) spectra in Fig. 1f, compared with C$_3$N$_4$ materials\cite{19}, the superfluous C element in hm-C(CN)$_3$ introduces larger number of lone pair of electrons out of the aromatic rings (marked by red circle, see atomic model in Supplementary Fig. 7), which can be used as additional active sites. The increased ESR intensity of hm-C(CN)$_3$ corroborates the improved delocalization of electrons and appearance of carbon atoms as adsorption sites.

**Electronic structures of the half-metallic carbon nitride nanosheets.** For highly efficient photocatalytic hydrogen evolution, the relative position of the conduction band minimum (CB) and valence band maximum (VB) and their absolute energies with respect to the reduction and oxidation levels are decisive elements\cite{7}. To determine these parameters of hm-C(CN)$_3$ nanosheets, the work function of the hm-C(CN)$_3$ nanosheets was obtained by scanning Kelvin probe force microscopy, as shown in Fig. 2a, b. The Au line was patterned on the hm-C(CN)$_3$ nanosheets as a reference (Fig. 2a), which has a known work function of 4.9 eV (the valley value in Fig. 2b). The line-scan data (Fig. 2b) show the uniformity of the as-prepared hm-C(CN)$_3$ nanosheets and the average Fermi level ($E_F$) across the middle area is $\sim$ 5.84 eV. The band gap is evaluated as 2.19 eV from the linear potential scans (Supplementary Fig. 11). Taking the ultraviolet photoelectron spectroscopy (UPS) (Supplementary Fig. 12)\cite{4}, band gap and Fermi level into consideration, the CB and VB positions of hm-C(CN)$_3$ nanosheets are evaluated as $-0.47$ and 1.72 V, respectively, as shown by the blue and pink arrows in Fig. 2c. The state-of-the-art hybrid functional (HSE06) calculations show that band gap of hm-C(CN)$_3$ decreases from 2.33 eV for monolayer to 2.06 eV for multilayer (Supplementary Fig. 1), and the band structure of monolayer is shown in Fig. 2c. The hm-C(CN)$_3$ displays a non-isotropic electronic structure
with a direct band gap (blue line: ~ 2.33 eV) at the G point, which is in coincidence with the experimental measurement. Importantly, the spontaneous spin polarization makes spin-down bands (marked by down) obviously across the $E_F$ to display metallic characteristic\(^{14}\), whereas the spin-up bands (marked by up) are obviously lower than $E_F$ and show semiconducting feature. This interesting half-metallic feature is experimentally observed in the valence band spin-resolved photoemission spectra (PES) near $E_F$ at 300 K of the hm-C(CN)\(_3\) sample and confirmed by DFT calculations (Supplementary Figs. 13–18, Supplementary Table 1 and Supplementary Note 7). The spectrum originated from the majority spin is extended to $E_F$ and displays an obvious metallic Fermi cutoff, although that of the minority spin reduces sharply at binding energy of ~ 0.6 eV and the spectral weights disappear around $E_F$, displaying an insulating gap. Furthermore, DFT calculations disclose that the radical C sites, resulting in half-metallic feature, can be easily decorated by hydrogenation than that of defects. (Supplementary Figs. 13–14 and Supplementary Table 1). Simultaneously, the \(^{13}\)C NMR spectra in Supplementary Fig. 15 indicates that the peak 1 originating from radical C sites slightly shifts after hydrogenation treatment, further confirming that hydrogen deactivation occurs at radical C sites and no additional defect structures are introduced. After hydrogenation treatment, the spin splitting between majority and minority in PES spectra disappear completely, which indicates that spin polarization is mainly originated from radical C sites (Supplementary Figs. 16–17 and Supplementary Note 1). The spectral differences between PES spectra and spin-resolved DOSs (densities of states) stem from lattice distortion or deformation in sample preparation, which can be confirmed by broad XRD pattern and DFT calculations (Supplementary Figs. 18–19).

Besides, the spontaneous spin polarization brings about remarkable ferromagnetic hysteresis as shown in Supplementary Fig. 20. Under optical excitation, the photogenerated carriers occupying different spin states can transform each other. The spin-resolved total density of state in Fig. 2d shows that the hm-C(CN)\(_3\) surface is fully filled with metallic spin-down states (see inset), which is more advantageous to carrier transfer. In addition, orbital analysis reveals that the half-metallicity mainly originates from the $p$ orbital of three N atoms at triazine ring\(^{1,14}\). The C atom outside of triazine ring (marked by circle in Supplementary Fig. 7) will inject a hole into this system, which may significantly alter its electronic properties and photocatalytic activity. These results evidence that hm-C(CN)\(_3\) is a potential photocatalyst for hydrogen evolution.

**Design of micro grid mode resonance structure for hydrogen evolution.** The hm-C(CN)\(_3\) nanosheets were combined with micro grid mode resonance structure for better capture of light. Figure 3a schematically depicts the in situ pyrolysis process. The original transparent nanoporous anodic aluminum oxide membrane (AAM), hm-C(CN)\(_3\) represents the sample of hm-C(CN)\(_3\) bulk material, MG@hm-C(CN)\(_3\) represents the sample of hm-C(CN)\(_3\) nanosheets incorporated into AAM, respectively.

Figure 3a schematically depicts the in situ pyrolysis process. The original transparent nanoporous anodic aluminum oxide membrane (AAM), hm-C(CN)\(_3\) represents the sample of hm-C(CN)\(_3\) bulk material, MG@hm-C(CN)\(_3\) represents the sample of hm-C(CN)\(_3\) nanosheets incorporated into AAM, respectively.

![Image](image1.png)

**Fig. 3 Micro grid mode resonance based artificial microstructure.** a Schematic key steps involved in MG@hm-C(CN)\(_3\) synthesis. Inset: digital camera images of a bare nanoporous template (white) sample and a MG@hm-C(CN)\(_3\) sample (black). b Self-assembly of MG@hm-C(CN)\(_3\) nanosheets on nanoporous templates to form artificial microstructure (Scale bar, 150 nm). c Top-view (Scale bar, 1 μm) and d cross-sectional (Scale bar, 200 nm) SEM images of the MG@hm-C(CN)\(_3\)_a. e Experimental absorption spectra measured by an integrated sphere in the visible and near-infrared range. MG represents the sample of nanoporous anodic aluminum oxide membrane (AAM), hm-C(CN)\(_3\) represents the sample of hm-C(CN)\(_3\) bulk material, MG@hm-C(CN)\(_3\) represents the sample of hm-C(CN)\(_3\) nanosheets incorporated into AAM, respectively.
perpendicular to the incident surface is called the S wave (marked by red arrows: $E_{1s}$). Compared with absorption of S waves in Fig. 4c, the P waves have two strong absorption areas in the large incidence angle (Fig. 4d), which indicates that the P wave behavior is more beneficial to light absorption in packed hm-C(CN)$_3$ layer. In this special columnar micro grid structure, the incident natural lights are all expressed as P wave on all the planes perpendicular to the nanotube inner walls (Supplementary Fig. 23 and Supplementary Note 2). Therefore, more energy will be introduced into the hm-C(CN)$_3$ nanosheets via P wave behavior, leading to higher absorption capacity. Besides, hm-C(CN)$_3$ nanosheets are incorporated into AAM to construct cylindrical resonators (void resonators), of which the inner walls are rough. Theoretical calculations indicate that cylindrical resonators with rough inner wall can reduce reflections and increase effective optical length, which makes incident lights travel with coherent superposition, resulting in a linear near-infrared (NIR) absorption (Supplementary Fig. 23).

**Photocatalytic hydrogen evolution performance.** Figure 5a shows the electrical conductivity of the multilayered hm-C(CN)$_3$ nanosheets reaches a high magnitude ($10^6$), which is advantageous to the migration of photogenerated carriers. To clarify the influence of van der Waals forces in multilayered hm-C(CN)$_3$ nanosheets on its high electrical conductivity, we plotted the electronic wave function distribution across multilayered hm-C(CN)$_3$ nanosheets for the CB in Fig. 5b. Owing to weak interlayer coupling effect in this 2D construction, built-in electric field induced by photogenerated carriers restrict electrons to intralayer so as to promote electron transfer, resulting in high electrical conductivity and excellent HER performance. The photocatalytic hydrogen production activities of different samples were detected in a mixed 10% (v/v) triethanolamine aqueous solution under a Xe lamp irradiation with an AM 1.5 G filter (Supplementary Movie 1). As shown in Fig. 5c, the pristine g-C$_3$N$_4$ sheets exhibit a low HER activity because of the rapid recombination of electron–hole pairs though they have similar size to our bulk hm-C(CN)$_3$ sample (~200 μm see Supplementary Fig. 24). The bulk hm-C(CN)$_3$ sheets bring about a enormous improvement in the hydrogen production rate (up to 301 μmol g$^{-1}$ h$^{-1}$) because the excellent optical absorption induces more photogenerated carriers and its half-metallic feature facilitates the rapid separation of photogenerated electron–hole pairs. As expected, MG@hm-C(CN)$_3$ sample shows a much higher hydrogen evolution rate of 1009 μmol g$^{-1}$ h$^{-1}$, ~ 62.8 times higher than that of pristine g-C$_3$N$_4$ sheets. After removal of the nanotubes [named as R@hm-C(CN)$_3$], the photocatalytic activity drops obviously, but still much higher than that of bulk hm-C(CN)$_3$ sheets. This is because the R@hm-C(CN)$_3$ nanosheets of much smaller size (20 ~ 60 nm, Supplementary Fig. 25) have more active sites than bulk hm-C(CN)$_3$ sample although the capture of light reduces without micro grid structure. In order to clarify the contributions from lights of different wavelengths to the hydrogen evolution, photocatalytic H$_2$ evolution and On/Off photocurrent curves were carried out under different irradiation lights (Supplementary Fig. 26 and Supplementary Note 3). Both the hydrogen evolution rates and the photocurrents decrease obviously in the absence of NIR or ultraviolet (UV) lights, evidencing that the entire solar energy can make contributions to the hydrogen evolution. The calculated solar-to-hydrogen conversion efficiencies for different samples are described in the Supplementary Fig. 27 and Supplementary

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**Fig. 4** Simulation of micro grid mode resonance enhanced absorption based polarization optics. a Schematic diagram of the model system: the hexagonal nanotube arrays with deposited hm-C(CN)$_3$ (top) and Cross-section (bottom) of the simulated electric field distribution of MG@hm-C(CN)$_3$. b Ep and Es for the two mutually perpendicular components of Electric vector. Absorption zones of c S-wave and d P wave versus wavelength and incident angle.
Methods, and MG@hm-C(CN)₃ has a highest value of 2.24 (ΔGₘₚ=10⁻⁶) %, which corresponds to an apparent quantum efficiency of 1.03% (see Supplementary Methods for details)²³. Besides the efficiency of hydrogen evolution, the long-term stability is another important parameter to evaluate the photocatalytic performance. As shown in Fig. 5d, after twelve cycles of repeated trials, MG@hm-C(CN)₃ shows negligible change in HER performance, which indicates that MG@hm-C(CN)₃ has the excellent photochemical stability. The amount of evolved hydrogen is proportional to the sunlight irradiation time, indicating of the constant hydrogen production under continuous irradiation. More importantly, for practical application, our designed MG@hm-C(CN)₃ sample can be easily collected without need of an additional recycle technology, which has obvious advantages over conventional photocatalyst nanomaterials.

**Photocatalytic hydrogen evolution mechanism.** To elucidate the HER mechanism, the potential energy profile is calculated to estimate the adsorption of H⁺ on the different sites of hm-C(CN)₃ nanosheets. The results show that the favorite adsorption sites are located in the neighboring region of C or N atom, especially for C1 and C2 positions (Fig. 6a). When these active sites are covered by H⁺ atom, the corresponding free energy for hydrogen adsorption is calculated and shown in Fig. 6b. It is generally known that a material displays a good catalytic activity when the free energy of adsorbed hydrogen tends to thermoneutral, i.e., ΔGᵣ = 0. If hydrogen cannot efficiently adsorb onto catalyst or forms a strong chemical bond, the hydrogen release and proton/electron transfer process also be limited, leading to lower catalytic activity²²-²⁷. DFT calculation indicates that the C2 sites successively introduce nonequilibrium stoichiometry to allow for favorable hydrogen adsorption on this active site as follows: H⁺ + e⁻ + *→H₂, where * denotes a binding site. After spontaneous hydrogen adsorption occurred at C1 and C2 sites (Fig. 6b), the ΔGᵣ at the C2 sites becomes almost thermoneutral, reaching an optimal ΔGᵣ. This makes the release of molecular hydrogen easier via reactions 2 H⁺→H₂↑+ 2e⁻. In actual HER process, H⁺ can unavoidably adsorb onto C4 and C6 sites, but this adsorption cannot lead to hydrogen generation. This is because if the H⁺ adsorbed onto the most active termination C2 site intends to generate molecular hydrogen more efficiently, it has to bond to neighboring H⁺ at C1 site. Thus, the influence of C4 and C6 terminations on HER can be neglected. To further understand the process of hydrogen production, the climbing nudged elastic band method is adopted to study the adsorption, activation, and reaction processes of H⁺ on the hm-C(CN)₃ nanosheets, and the results are shown in Fig. 6c. Generally, the spin triplet state energy (marked by black T line) is smaller than that of spin singlet state (marked by red S line), and triplet-to-singlet conversion happens between S6 and S7. Naturally, the HER process will follow lower energy route (marked by a black arrow). First, a single H atom appears away from site C1 at a distance of 4.903 Å (S1), then spontaneously adsorbs physically onto the site C1 after energy release of ~1.72 eV (S3) before overcoming the negligible barrier at S2. Subsequently, another H atom closes (S4) and adsorbs onto the C2 site, then two metastable H⁺ atom configuration forms with a distance of 1.865 Å (S5). Upon photoexcited perturbation, the two adjacent H⁺ atoms begin orbital hybridization and bridge a transition state at a H−H distance of 1.864 Å (S6). To complete this process, activation energy of 0.58 eV relative to the energy difference between the potential energies of two metastable states is required. With phonon assistance, a molecular hydrogen with diagonal bridge configuration is dissociated from the hm-C(CN)₃ surface. In this process, the degeneracy of its π* orbitals is lifted owing to the hybridization with the C1 site lone pairs, and the singlet state becomes more stable than the triplet (between S6 and S7)²⁸. Therefore, for the generation of H₂ there must be at least one point in configuration space where the triplet and singlet potential energy surfaces cross (S7), and triplet-to-singlet conversion likely takes place nonradiatively. Eventually, the H₂ molecule is released from the hm-C(CN)₃ surface with an energy gain of 1.24 eV (S8). The triplet-to-singlet conversion can be calculated as: \( P_{\text{r}} = 2\left[1 - \text{exp}(-V^2/h\nu[\text{Ft-Fs}])\right]^{29} \), here h, V, Fs, and Ft...
are Planck’s constant, the spin-orbit matrix element, slopes at energy crossing point between triplet and singlet states, respectively. By using $V = 122 \text{ cm}^{-1}$ for describing $H_2$ dynamic process, we find $P_\text{ts} = 0.78$ at 300 K$^{28}$, which confirms that the triplet-to-singlet conversion plays an irreplaceable role in this HER mechanism.

To experimentally confirm the singlet-to-triplet conversion, the streak image and the time-resolved photoluminescence (PL) spectra are shown in Fig. 6d. The transient photoluminescence streak image and the time-resolved photoluminescence (PL) spectra can be resolved into the prompt emissions ($< 50 \text{ ns}$) and the long tail components ($> 50 \text{ ns}$). The prompt emissions mainly come from singlet excited state, and long tail components are originated from the intersystem crossing from singlet-to-triplet excited states. The obvious red-shifts in two time-dependent PL spectra disclose that the triplet excited states have been generated successfully via intersystem crossing process. (See Supplementary Fig. 30, Supplementary Table 2 and Supplementary Note 8)30,31.

In conclusion, we have demonstrated that half-metallic C(CN)$_3$ nanosheets can be facilely synthesized and incorporated into designed micro grid mode resonance systems via in situ pyrolysis of ionic liquid to tune optical transmission for highly efficient photocatalytic hydrogen evolution. The artificial microstructure not only sharply enhances solar exploitation but also obviously improves photocatalyst’s stability and recyclability. More importantly, this microstructure makes it possible to prepare size-controllable carbon nitride materials without liquid exfoliation. In view of the great variety of microstructures and ionic liquids,
other materials of special features are foreseen in the areas of photocatalysis, electrocatalysis, and photovoltaic cells.

Methods
Synthesis of MG@hm-C(CN)3 micro grid resonance structures. All chemicals were of analytical grade and used as received without further purification. First, 0.01 mol 1-butyl-3-methylimidazolium chloride and 0.01 mol potassium triglycinate (toxic by inhalation, in contact with skin and if swallowed) were dissolved in 30 ml deionized water. Then, the mixture was stirred under the nitrogen current for 30 min at room temperature to ensure complete reaction (step 1). Subsequently, the reaction solution was evaporated in 80 °C water bath to remove the solvent. The residual was dissolved in 10 ml absolute ethanol followed by centrifugation to remove the precipitant. The supernatant was evaporated in 80 °C water bath in nitrogen current for 30 min at room temperature to ensure complete reaction (step 2). The pyrolytic temperature for bulk hm-C(CN)3 sample under N2 carrier gas was set to 550 °C. In preparation of MG@hm-C(CN)3 samples, the AAM template was put vertically downstream of the oil precursor, at a distance of 15 cm. Before heating, the system was purged with 517 sccm (standard cubic centimeters per minute) high-purity nitrogen (N2 99.9999%) for 30 min. Then, the pressure was reduced to 7.5 × 10−2 Torr for the duration of the reaction. After that, the furnace was heated, with a heating rate of 5 °C min−1, to a temperature of 650 °C. It was kept at this temperature for 2 h with an N2 flow of 526 sccm (step 3). After the system was cooled down to room temperature, a metalacene product was found deposited onto the AAM. The pyrolytic temperature for bulk hm-C(CN)3 sample under N2 carrier gas was determined via TGA (Supplementary Fig. 2). The TGA profile discloses that the characteristic liquid peaks at 300 °C results in the formation of partially solidified liquid–solid reaction and heating in the range of 450–550 °C gives solid products. Polymerization of [(C6H5)2C]=2 in the temperature range of 300–400 °C may follow a similar dynamic cyclotrimerization reaction to the condensation of aromatic nitriles, cyanamide, and acetylenes, which is accompanied by the decomposition of the corresponding ionic liquids cations. We prepared the bulk hm-C(CN)3, template with the temperature maintained at ~ 500 °C. In preparation of MG@hm-C(CN)3, the AAM template was put vertically downstream of the oil precursor, and the annealing temperature was increased to ~650 °C to optimize the formation of hm-C(CN)3. After considering a series of distances (5, 10, 15, 20 cm) from the reaction precursor, we found that the 15 cm was the optimal separation. The preparation route is schematically displayed in Supplementary Fig. 3.

Characterization. The morphology and microstructure were characterized by field-emission scanning electron microscopy (Hitachi, S-4800), HR-TEM (JEOL-JEOL-2100) equipped with X-ray energy dispersive spectrum, X-ray photoelectron spectroscopy (XPS, PHI5000 VersaProbe), Tapping-mode AFM (Nanoscope IIIA) and half-metallicity in a primitive carbon nitride unit cell. The activation energies of the corresponding ionic liquids cations. We prepared the bulk hm-C(CN)3, with the temperature maintained at ~500 °C. In preparation of MG@hm-C(CN)3, the AAM template was put vertically downstream of the oil precursor, and the annealing temperature was increased to ~650 °C to optimize the formation of hm-C(CN)3.

Photocatalytic hydrogen evolution. The photocatalytic hydrogen evolution activities of different samples were tested by using a photocatalytic activity evaluation system (LabSorb-II ASC), Perfectlight, Beijing). The light source was a 300 W Xe lamp including an AM 1.5 G filter. The Xe lamp was placed 10 cm away from the reactor. The focused intensity (J) on the reactor was adjusted to 100 mW cm−2. Various photocatalysts with the same quality were directly dispersed into 100 ml of the 10% (v/v) triethanolamine mixture solution. Before irradiation, the whole system was sealed and vacuumed by a mechanical pump to eliminate any gas impurities, after the generated hydrogen was extracted every 1 hour with an on-line gas chromatograph (9790 II, Fulig, Zhejiang) equipped with a TCD detector and Ar gas carrier. The generated amount of hydrogen was evaluated according to the fitted standard curve. During the photocatalytic process, the fluid cooling water was used to guarantee the reactant solution at room temperature.

Electric near-field simulation details. The electromagnetic simulation is performed using a Finite-Element Method (FEM). In addition, the refractive index of water is set to be dispersive. All the physical domains are surrounded by perfectly matched layers to absorb all the outgoing waves, thus minimizing any reflection. Plane waves are used as excitation sources in all simulations along the z axis. For better clarification, the intensity of electrical field distribution is visualized in a square scale.

Maxwell’s equations are rigorously solved by three-dimensional finite-element method. The incident, internal, and scattered electromagnetic fields of nanosheet can be described by introducing a series of cubic harmonics. The absorption coefficient of the light wave is related to the polarization, wavelength, and incident angle of the light. According to the Fresnel formula, the relationship between the absorption coefficient of the medium for TE polarization (s wave) and TM polarization (p wave) and the wavelength and incident angle can be calculated separately.

Theoretical calculation. The calculations are performed by using plane-wave basis. All calculations are performed with plane-wave basis. Vienna ab initio simulation package code with generalized gradient approximation. The cutoff energy for expanding Kohn-Sham wave functions is chosen to be 460 eV and the vacuum space of 20 Å is used to avoid the interaction between periodical images, respectively. The Monkhorst-Pack k-points grid is 8 × 8 × 1 and all forces on the free ions is 0.03 eV/Å, which have been tested to be well converged. The state-of-the-art hybrid functional calculations based on the Heyd-Scuseria-Ernzerhof (HSE06) functional have been adopted to calculate the spin polarization and half-metallicity in a primitive carbon nitride unit cell. The activation energies are calculated using the climbing nudged elastic band method. Electromagnetic simulations are obtained by a FEM.

Data availability. The data that support the findings of this study are available from the corresponding authors on request.

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Author contributions
G.Z. performed the experiments and co-wrote this manuscript. Y.S and X.Y.X prepared the samples and proofread the manuscript. J.D performed the theoretical calculation and co-wrote the manuscript. J.C.S plotted the figures. J.L.Z and J.H.G designed the experimental setup. L.Y.L and S.L provided the calculation resources. Y.Y.H co-wrote the manuscript. L.Z.L and X.L.W analyzed the data and wrote the manuscript.

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