Promoting Photocatalytic Hydrogen Evolution Activity of Graphitic Carbon Nitride with Hole-Transfer Agents


Visible light-driven photocatalytic reduction of protons to $\text{H}_2$ is considered a promising way of solar-to-chemical energy conversion. Effective transfer of the photogenerated electrons and holes to the surface of the photocatalyst by minimizing their recombination is essential for achieving a high photocatalytic activity. In general, a sacrificial electron donor is used as a hole scavenger to remove photogenerated holes from the valence band for the continuation of the photocatalytic hydrogen ($\text{H}_2$) evolution process. Here, for the first time, the hole-transfer dynamics from Pt-loaded sol–gel-prepared graphitic carbon nitride (Pt-sg-CN) photocatalyst were investigated using different adsorbed hole acceptors along with a sacrificial agent (ascorbic acid). A significant increment (4.84 times) in $\text{H}_2$ production was achieved by employing phenothiazine (PTZ) as the hole acceptor with continuous $\text{H}_2$ production for 3 days. A detailed charge-transfer dynamic of the photocatalytic process in the presence of the hole acceptors was examined by time-resolved photoluminescence and in situ electron paramagnetic resonance studies.

Conversion of photon energy to chemical energy by reducing water into molecular $\text{H}_2$ has the highest prospective of providing clean and sustainable energy to replace fossil fuels.[1] An efficient photocatalyst with a suitable band structure can harvest solar light to convert it into chemical energy.[2] Among various photocatalysts reported in the literature, the organic semiconductor polymeric graphitic carbon nitride (g-CN) has attained growing interest for the visible-light-induced photocatalytic $\text{H}_2$ evolution reaction (HER) in recent years.[3] The appropriate bandgap, conduction band position, high physicochemical stability, and facile large-scale synthesis make g-CN proficient for the photocatalytic proton reduction.[4] Moreover, the band structure can be engineered by doping with heteroatoms or changing the conjugated structure of the g-CN to avoid better photocatalytic performance.[5] Generally, the photocatalytic activity of g-CN largely depends on the choice of the cocatalyst, pH of the reaction medium, and a sacrificial electron donor.[6] A noble metal cocatalyst like Pt has shown improvements in the photocatalytic HER rate compared to the bare g-CN.[7] The use of non-noble cocatalysts with g-CN to attain efficient photocatalytic $\text{H}_2$ evolution has also been recently demonstrated.[8–10]

Irradiation of g-CN by visible light excites electrons to the conduction band (CB) generating holes in the valence band (VB).[8] The electrons in the CB can reduce protons to $\text{H}_2$ in the presence of a suitable cocatalyst whereas holes in the VB can be extracted to a certain extent by sacrificial electron donors during the photocatalytic $\text{H}_2$ evolution reaction (HER). The recombination of holes and electrons, on the other hand, decreases the photocatalytic activity.[6] For the efficient photocatalytic $\text{H}_2$ evolution reaction, effective transfer of the electrons from the CB to the catalytic site as well as the transfer of the holes from the VB to the sacrificial agent, are equally important.[10] Therefore, the hole extraction from the...
VB by minimizing the recombination of charges is crucial to improve the proton reduction efficiency.

In this regard, a hole acceptor, phenothiazine (PTZ), has been effectively employed to extract the holes from the VB of CdS.\textsuperscript{[11]} Similarly, PTZ-modified CdSe quantum dots were shown to improve hole transfer during photocatalytic HER.\textsuperscript{[12]} Different derivatives of PTZ were also studied for the photocatalytic HER by combining them with a TiO\textsubscript{2} photocatalyst.\textsuperscript{[13]} In addition, PTZ-based molecules were successfully explored as hole-transfer agents to improve the efficiency of solar cells.\textsuperscript{[14]} Two main factors directly control the use of PTZ as an ideal hole-transfer agent during the photocatalytic process: (i) one-electron oxidation of PTZ to form the cationic radical (PTZ\textsuperscript{+}) with high stability and (ii) thermodynamically favorable hole transfer from the VB of g-CN [1.47 V vs. normal hydrogen electrode (NHE)] to the highest occupied molecular orbital (HOMO) of PTZ (0.9 V vs. NHE).\textsuperscript{[15]}

Although Durrant and co-workers reported that the photophysics of g-CN materials are similar to that of inorganic semiconductors,\textsuperscript{[16]} the role of the hole-transport agents (HTAs) on the photocatalytic properties of g-CN or, in a broader context, to photoactive organic polymers remains unexplored. In this work, the role of different HTAs for extracting the holes from the VB of sol–gel-prepared graphitic carbon nitride (sg-CN), when irradiated with visible light, is explored (Figure 1). Three different substrates, PTZ, phenazine (PZN), and phenoxazine (POZ), were employed to understand the role of hole acceptors during photocatalytic H\textsubscript{2} evolution with Pt deposited on sg-CN (Pt-sg-CN) in the presence of ascorbic acid (AA) as the sacrificial electron donor (Scheme 1 and Tables S1–S5 in the Supporting Information). A remarkable increase in the rate of the H\textsubscript{2} production was observed when PTZ and POZ were used as the hole acceptors. The highest photocatalytic activity was obtained when PTZ was added to a mixture of Pt-sg-CN and AA (4.84 times increment in the rate of H\textsubscript{2} production).

Electron paramagnetic resonance (EPR) studies confirmed the effective hole extraction from the VB of sg-CN in the presence of PTZ with a decrease in the recombination of the photogenerated charges. Photoluminescence (PL) spectroscopic studies also revealed the improved charge transfer by minimizing electron–hole recombination in the presence of PTZ. Notably, an optimum amount of the acceptor for efficient hole transfer is observed reaching its maximum when 0.25 wt% of PTZ is used. Moreover, Pt-sg-CN-PTZ 0.25% continuously produced H\textsubscript{2} for 3 days indicating the long-term durability of the photocatalytic system.

Pt was deposited on sg-CN by a microemulsion method following our previously reported procedure (see Experimental in the Supporting Information).\textsuperscript{[17]} First of all, three different organic molecules, PTZ, POZ, and PZN, were loaded (0.25 wt%) on Pt-sg-CN and characterized by powder X-ray diffraction (PXRD; Figure S1). PXRD analyses showed the peaks at 2\texttheta values of approximately 13 and 27° corresponding to the intra-planar heptazine unit distance and the inter-planar stacking arrangement in layered g-CN, respectively.\textsuperscript{[8]} Solid-state cross-polarization magic angle spinning (CP/MAS) \textsuperscript{13}C NMR spectra of the PTZ-loaded Pt-sg-CN exhibited peaks at 154 and 162 ppm for CN\textsubscript{g} and CN(\textsubscript{N})\textsubscript{N}(\textsubscript{H}) moieties of the heptazine based system, whereas no extra peak corresponding to PTZ was detected (Figure S2).\textsuperscript{[8]} UV/Vis spectra revealed an increment in the visible light absorption after loading of the HTAs on Pt-sg-CN (Figure S3) whereas no significant change was detected in the IR spectra (Figure S4). Transmission electron microscopic (TEM) studies displayed 3–4 nm Pt nanoparticles (NPs) supported on sheet-like carbon nitride in PTZ-loaded Pt-sg-CN (Figure S5). Energy-dispersive X-ray spectroscopy (EDX) and inductively coupled plasma atomic emission spectroscopy (ICP-AES) further quantified the presence of Pt (≈2 %, Figure S6 and Table S6). Elemental analyses could not detect any change in the atomic ratio of C/N (≈1.46 for sg-CN) after the loading of the HTA on sg-CN (Table S7).

The oxidation state of Pt in PTZ-Pt-sg-CN was analyzed by X-ray photoelectron spectroscopy (XPS) (Figure S7). The Pt 4f\textsubscript{7/2} and 4f\textsubscript{5/2} peaks showed two pairs located at 71.6/74.9 and 72.5/75.5 eV indicating the presence of Pt\textsuperscript{6} and Pt\textsuperscript{4} species, respectively.\textsuperscript{[18]} The C 1s, N 1s, and O 1s XPS spectra of PTZ-Pt-sg-CN were matched with previously reported literature and discussed in the Supporting Information (Figures S8–S10).\textsuperscript{[8]} EDX mapping showed a homogeneous distribution of the elements C, N, O, and Pt along with S, after the loading of PTZ (Figure S11). The Brunauer–Emmett–Teller (BET) surface area of Pt-sg-CN was determined to be 147.8 m\textsuperscript{2} g\textsuperscript{−1}, and a reduction of the surface area was observed after loading of the HTAs, which points to the
incorporation of PTZ within the pores accompanied by a partial pore blocking (Figure S12, Table S8).

The role of PTZ, POZ, and PZN was investigated as the HTA for the photocatalytic H₂ evolution with Pt-sg-CN in the presence of AA as the sacrificial electron donor under sunlight simulation (AM 1.5 G, 300 W). A significant improvement in the photocatalytic HER was observed when PTZ or POZ were employed as the hole carrier along with AA (Figure 2a). However, no such increment in H₂ production was recorded with PZN. The amount of produced H₂ (mmol) with 2 % PTZ was 2.74 times higher compared to the reaction carried out without any HTA after 20 h of photocatalytic reaction (Figure 2). The difference is more pronounced (4.84 times) when 0.25 % PTZ was used as the HTA in a 20 h experiment (Figure 2a). In the presence of 2 and 0.25 % POZ, the increment was 1.76 and 2.30 times higher compared to the reaction with no hole acceptor. As the sacrificial electron donor was the same (AA), the enhancement in the H₂ production confirmed the effective hole transfer from the VB of Pt-sg-CN to the HOMO of PTZ or POZ under irradiation with the sunlight simulator.

The photocatalytic HER was further optimized by varying the amount of PTZ to determine the best reaction condition (Figure S13). It was observed that the amount of H₂ production was highly dependent on the amount of PTZ, which attained a maximum at 0.25 % and slightly drops to reach a plateau in the presence of an excess amount. This result indicates a limiting amount of hole transfer from the VB of Pt-sg-CN to the HOMO of PTZ, and the excess amount of PTZ leads to a saturation point. It is worth to mention here that the photocatalytic H₂ evolution with PTZ-CdS or PTZ-CdSe was studied without loading of any cocatalyst.[11–12] As a cocatalyst significantly facilitates the electron transfer from the CB of the photocatalyst to protons, without cocatalyst the overall photocatalytic process is affected. Therefore, we chose to investigate the photocatalytic H₂ evolution with a cocatalyst (Pt)-loaded sg-CN and only bare sg-CN in the presence of PTZ and AA. Without Pt loading, the amount of produced H₂ with sg-CN and AA is negligible, and even the addition of PTZ does not improve the H₂ production (Figure S14).

Under the best reaction condition (Pt-sg-CN-PTZ 0.25 % and AA), the rate of photocatalytic H₂ production was 0.05 mmolH₂g₅₉₉₉₋₁ h⁻¹. This rate is higher than the rates obtained with other graphitic carbon nitride-based catalyst systems when Pt is used as the cocatalyst in the presence of sacrificial agent AA (pH = 3; Table S9).[19] Moreover, a continuous H₂ evolution for almost 3 days was recorded with PTZ-Pt-sg-CN under sunlight simulation irradiation that showed the long-term durability of the system. Furthermore, we also compared the activity of Pt-sg-CN-PTZ 0.25 % with respect to the effect of the light source by using 300 W Xe lamp with a 420 nm cut-off filter that indeed displayed significantly enhanced H₂ production (almost 1.5 times) compared to the sunlight simulator and was highly promising compared to the related systems (Figure S15, Table S10). We have carried out electrospray ionization (ESI) MS measurements to confirm that during photocatalysis the adsorbed PTZ on the surface of Pt-sg-CN is responsible for the hole transfer and it does not leach out into the solution (Figure S16). ESI-MS of the solution after photocatalysis did not detect any peak of PTZ and showed signals only related to AA, which confirmed our hypothesis. It is noteworthy that a better photocatalytic activity can be achieved with sg-CN or related materials when triethanolamine (TEOA) is used as a sacrificial agent, rather than AA.[41] However, the photocatalytic activity studies using PTZ as the HTA cannot be carried out in TEOA, as PTZ is oxidized to phenothiazinone in an aqueous solution at pH > 5 (Figure S17).[20] Therefore, we have carried out the photocatalytic hydrogen evolution reactions in AA solution (pH = 2.73) instead of the TEOA solution (1:9 TEOA/H₂O solution: pH = 7.76).

A negligible amount of photocatalytic H₂ evolution was detected from the control experiments of Pt and AA (without the addition of sg-CN). Further, we have carried out the photocatalytic hydrogen evolution with Pt-sg-CN-PTZ 0.25 % in D₂O to confirm the source of protons. GC-MS confirmed D₂ as the major product (87 %) while a small amount of HD (9 %) and H₂ (4 %) was detected because of D⁻ exchange with AA (Figure S18). This result clearly established water as the source of hydrogen and not the sacrificial agent AA. For a comparison purpose, the apparent quantum efficiency (AQE) with Pt-sg-CN-PTZ 0.25 % was determined at two different wavelengths. At 415 and 440 nm, the highest AQE was determined to be 0.32 and 0.25 %, respectively.

The charge-transfer properties of Pt-sg-CN and PTZ-Pt-sg-CN were further examined by transient photocurrent measurements and electrochemical impedance spectroscopic (EIS) analyses (Figures 2b and S19). Nyquist plots confirmed a better charge transport in the presence of PTZ compared to only Pt-sg-CN under visible light irradiation and even in the dark (Figure S19). Improvement in transient photocurrent with Pt-sg-CN was also recorded in the presence of PTZ at a constant potential (at 0.5 V vs. Ag/AgCl, Figure 2b).

The effect of the HTA on charge separation, migration, and recombination was studied in detail by time-resolved photoluminescence (TRPL). Recently, Durrant and co-workers...
reported sub-ps time-scale charge separation in g-CN, while the long-lived (> μs) holes and electrons are unavailable for the next step, to facilitate photocatalytic H₂ production. They further demonstrated that the photoinduced electron accumulation in the CB of carbon nitride accelerates the charge recombination. Similarly, Merschjann et al. disclosed that the electron transport in g-CN is in the perpendicular direction of the parallel sheets of the 2D material.

The emission peak in the PL spectra originates from the separation of photogenerated electrons and holes, and its intensity can reflect the separation efficiency of charge carriers. Consequently, a lower intensity indicates a more effective charge separation and trapping in non-emissive states. In the present case, the Pt-sg-CN exhibited a strong emission peak at 465 nm corresponding to the bandgap charge separation and the PL emission peak intensity of Pt-sg-CN-PTZ 0.25% was found to be lower than that of Pt-sg-CN (Figure S20). This suggested that the electron–hole pair separation is higher in Pt-sg-CN-PTZ 0.25% resulting in higher H₂ production compared to Pt-sg-CN. In contrast, we previously reported a decrease in the recombination of the charges and improved electron transport when the cocatalyst Ni,P was combined with sg-CN.

Photoinduced hole transfer from Pt-sg-CN to HTA was also investigated by TRPL spectroscopy (Figure 3a, Table S11). Photoluminescence lifetime measurements showed that the presence of the HTA reduced the exciton lifetime of sg-CN and the quenching increased from PZN to POZ to PTZ. It should be mentioned here that not only the hole transfer but also electron transfer and energy transfer from sg-CN to the adsorbate can lead to the exciton quenching. In the present study, we did not consider the energy transfer from sg-CN to the adsorbate due to a lack of spectral overlap of sg-CN emission (λ_max = 475 nm) and PTZ absorption (λ_max = 265 nm) (Figure S21). In addition, we were able to detect the radical cation (PTZ⁺) generated by the hole transfer from photoexcited sg-CN (Figure S22). PTZ⁺ showed a strong absorption at 324 nm with a shoulder at 380 nm, well separated from the absorption of PTZ at 265 nm. The above results clearly demonstrate that the HTA promotes the dissociation of the photogenerated electron–hole pairs to free charge carriers in excited sg-CN.

The process of transferring holes from the photocatalyst to water molecules is more complicated than the transfer of electrons to protons. The current understanding of the hole-transfer process has been mostly deduced from the photoluminescence quenching of the exciton in the presence of hole acceptor. Previously, Kamat and co-workers were able to detect the adsorbate cation formed by the hole transfer from CdSe quantum dots to p-phenylenediamine. Similarly, the transfer of holes from CdS to PTZ has been described by photoluminescence spectroscopy.

Further, in situ EPR studies were carried out to understand the charge recombination and transport properties. Pure sg-CN, even in the dark condition, exhibits a narrow isotropic singlet with Lorentzian line shape at a g-value of 2.0046, which results from unpaired electrons trapped at sp² carbon atoms in a typical heptazine unit (Figure S23). The EPR signal intensity increased under visible light irradiation due to enhanced excitation and trapping of electrons in the CB and decreased again after light switch-off due to the recombination of photoexcited electrons with holes (Figure S24). This intensity decreases in the following order: Pt-sg-CN > Pt-sg-CN-PZN > Pt-sg-CN-POZ > Pt-sg-CN-PTZ (Figure 3b, black symbols), due to the transfer of excited electrons from the CB of sg-CN to Pt, where they cannot be detected by EPR. This means, the lower the EPR signal intensity under irradiation, the more effective is electron transfer to the Pt nanoparticles. Comparison of Figure 3b and Figure 2a reveals that the EPR intensity under visible light irradiation is the lowest, that is, electron transfer to Pt is the highest for the most active catalyst Pt-sg-CN-PTZ. This is easy to understand, considering that proton reduction takes place on the Pt cocatalyst and a high concentration of electrons might speed up H₂ formation.

Interestingly, the EPR signals of the catalysts containing HTAs are smaller and broader compared to the signal of pristine Pt-sg-CN (Figure S24). This suggests that some transfer of trapped electrons from sg-CN to the HTA may occur already in the dark. In principle, this should give rise to radical formation. In Figure S24, a hyperfine structure typical for such kinds of radicals is not resolved, yet it could contribute to the line width, which is highest for the most active Pt-sg-CN-PTZ. The significantly stronger dark signal of pristine Pt-sg-CN may be due to the fact that an HTA, which could take up a part of the trapped electrons, is missing.

In conclusion, we demonstrated here the effect of hole acceptor molecules to transfer the photogenerated holes from the VB of Pt-sg-CN to the sacrificial electron acceptor. EPR and PL studies have shown that the transport of the photogenerated holes becomes faster in the presence of the molecules PTZ and POZ resulting in improved electron–hole separation compared to PZN. In the case of PTZ, the most effective hole transport has been observed with a 4.84 times increase in the rate of photocatalytic HER. Moreover, the PTZ-Pt-sg-CN can produce H₂ for 3 days to exhibit long-term
stability. It is observed that the formation of the radical cation (PTZ\textsuperscript{+}) by one-electron oxidation is the key factor for the facile hole transfer. Further, the photogenerated PTZ\textsuperscript{+} can be utilized as the super-photooxidants to carry out photocatalytic organic reactions. Besides, this study will open a new avenue to develop hole transfer agents, which are stable in alkaline medium (especially in TEOA). This will be highly important to further improve the photocatalytic activity of Pt-sg-C\textsubscript{N} combining HTA and using TEOA as the sacrificial electron donor.

### Experimental Section

#### Chemicals
The chemicals were purchased from Sigma Aldrich or Alfa-Aesar and used without further purification.

#### Instruments
Powder XRD was recorded in Bruker AXS D8 advanced diffractometer equipped with a position-sensitive detector (PSD) and curved germanium (111) primary monochromator and the radiation used was CuK\textsubscript{α} (λ = 1.5418 Å).

The solid-state CP/MAS \textsuperscript{13}C\textsubscript{(1H)} NMR spectra were recorded in a Bruker Advance 400 spectrometer. UV/Vis spectroscopic measurements were performed in Lambda 35 UV/Vis spectrometer from Perkin Elmer (USA). IR spectroscopic studies were performed in BIORAD FT5 6000 Fourier-transform (FT) IR spectrometer under attenuated total reflection (ATR) conditions.

XPS studies were performed in a Kratos Ultra X-ray photoelectron spectrometer (Kratos Analytical Ltd., Manchester, UK) with an AlK\textsubscript{α} monochromatic radiation source (1486.7 eV) and 90° take-off angle maintaining at 2×10\textsuperscript{-5} Torr pressure in the analyzing chamber. The high-resolution XPS spectra were recorded with a pass energy of 20 eV and a step with 0.1 eV. The binding energies were calibrated with respect to the C 1s peak energy position at 285.0 eV. Data analyses were performed employing Casa XPS (Casa Software Ltd.) and Vision data processing program (Kratos Analytical Ltd.).

SEM was carried out on a LEO DSM 982 microscope integrated with an EDX spectrometer (EDAX, Apollo XPP). Data handling and analyses were achieved with the software package EDAX. Most abundant elements were selected from the EDX spectra. TEM was accomplished on an FEI Tecnai G2 20 S-TWIN transmission electron microscope (FEI Company, Eindhoven, Netherlands) equipped with a LaB\textsubscript{6} source at 200 kV acceleration voltage. EDX analyses were achieved with an EDAX r-TEM SUTW detector (Si (Li) detector) and the images were recorded with a GATAN MS794 P CCD camera. 5 independent EDX measurements were done to determine the C:N ratio and Pt content on Pt-sg-CN. The SEM and TEM experiments were conducted at the Zentrum für Elektronenmikroskopie (ZELMI) of the TU Berlin.

The elemental (CHN) analyses were performed on a Thermo Scientific LQF orbitrap XL. The analyzed solution was taken from the supernatant after 20 h of photocatalysis and injected directly into the spectrometer.

#### Synthesis of Pt-sg-CN
Carbon nitride, sg-CN was synthesized according to the literature-reported procedure.

Initially, concentrated solutions (5 mg mL\textsuperscript{-1}) of PTZ, PZN, and POZ were prepared by dissolving 25 mg of the hole transfer agent in 5 mL EtOH. The concentrated solution was diluted to prepare the solutions of desired concentration (0.10, 0.25, 0.50, 1.2, and 10% of loading in Pt-sg-CN). 60 mg of Pt-sg-CN was dispersed in 5 mL solution of the HTA in EtOH (Tables S1–S3) to give materials with different amounts of the incorporated HTA. The mixture was sonicated for 2 h, followed by stirring for 1 h at 500 rpm. The solid catalyst was separated from the solution by centrifugation (5000 rpm, 10 min). The solid was washed with acetone and dried overnight at room temperature. The final concentration of the HTA in the solution was determined by UV/Vis spectroscopy. The amount of incorporated HTA was calculated from the difference in the initial concentration of the solution and after the loading on Pt-sg-CN.

#### Photocatalytic hydrogen evolution
Photocatalytic hydrogen evolution reactions were carried out in a 60 mL Teflon reactor fitted with a quartz glass side window. The temperature of the reaction mixture was maintained at 30±1°C by using a temperature controller. 25 mg catalyst, 1.33 g AA, and 38 mL water were kept inside the photoreactor, degassed by purging Ar for 30 min and irradiated with a sunlight simulator (LOT LS0308). This consists of a 300 W Xe lamp with an AM 1.5G filter, which creates a spectrum similar to sunlight. The distance between the lamp and the reactor is always 10 cm. The long-term experiment was also done using the 420 nm cut-off filter. The increase of pressure was measured for 20 h (70 h for the long-term experi-
Photoexcitation of PTZ and generation of radical cation PTZ$^+$

A mixture of 2 mg PTZ and 5 mg AA in 10 mL of water was sonicated for 30 min to prepare a suspension, and the UV/Vis absorption spectrum was recorded. 5 mg of Pt-sg-CN were added to the mixture and sonicated for 10 min with visible light irradiation (400–700 nm, LUMATEC light source, SUPERLITE 400). The UV/Vis absorption spectrum was measured immediately to detect the generated PTZ$^+$. 

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Conflict of Interest

The authors declare no conflict of interest.

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