Charge-transfer regulated visible light driven photocatalytic H₂ production and CO₂ reduction in tetrathiafulvalene based coordination polymer gel

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The much-needed renewable alternatives to fossil fuel can be achieved efficiently and sustainably by converting solar energy to fuels via hydrogen generation from water or CO₂ reduction. Herein, a soft processable metal-organic hybrid material is developed and studied for photocatalytic activity towards H₂ production and CO₂ reduction to CO and CH₄ under visible light as well as direct sunlight irradiation. A tetrapodal low molecular weight gelator (LMWG) is synthesized by integrating tetrathiafulvalene (TTF) and terpyridine (TPY) derivatives through amide linkages and results in TPY-TTF LMWG. The TPY-TTF LMWG acts as a linker, and self-assembly of this gelator molecules with Zn²⁺ ions results in a coordination polymer gel (CPG); Zn-TPY-TTF. The Zn-TPY-TTF CPG shows high photocatalytic activity towards H₂ production (530 μmol g⁻¹ h⁻¹) and CO₂ reduction to CO (438 μmol g⁻¹ h⁻¹, selectivity > 99%) regulated by charge-transfer interactions. Furthermore, in situ stabilization of Pt nanoparticles on CPG (Pt@Zn-TPY-TTF) enhances H₂ evolution (14727 μmol g⁻¹ h⁻¹). Importantly, Pt@Zn-TPY-TTF CPG produces CH₄ (292 μmol g⁻¹ h⁻¹, selectivity > 97%) as CO₂ reduction product instead of CO. The real-time CO₂ reduction reaction is monitored by in situ DRIFT study, and the plausible mechanism is derived computationally.

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Artificial photosynthesis, i.e. the conversion of sunlight into fuels, is a green approach and has the potential to solve the global energy crisis. In recent years, a significant amount of research has been carried out to develop artificial systems for mimicking the sophisticated methodology of nature’s water splitting as well as CO2 reduction. Natural photosynthesis rely on the occurrence of precise sequences of proteins/enzymes to the several elementary steps of inter-component light-absorption, charge separation, and migration. The synthetic assimilation of these parameters precisely in a spatial organization of molecular components is indeed a challenging task. The recent upsurge of converting CO2 into fuels like CH3OH, CH4, or different chemical feedstock has gained widespread attention. The conversion of CO2 to hydrocarbon fuels would mitigate not only the effect of CO2 concentration in the atmosphere but also reduce the dependency on fossil fuel-based economy. However, the photoreduction of CO2 to CH4 has not been efficient. A handful of metal, metal oxide, and chalcogenides can faciliate the facile diffusion of reactants to the active sites and will show efficient electron transfer between different components. These artificial hybrid synthetic systems can mimic the intricate functioning of the natural photosystem and can eventually show impressive H2 evolution from water or CO2 reduction. Extended face-to-face arrays of the donor–acceptor π-chromophoric systems would be an ideal candidate for light harvesting. These systems will allow greater exciton mobility, which, in turn, leads to charge generation and subsequent electron transfer to the catalyst. To this end, tetrahiafulvalene (TTF) moiety is a well-known p-type semiconductor possessing high electron donation capability with excellent photostability and good charge carrier mobility. Further, the integration of a suitable electron acceptor unit to the TTF moiety could result in a system with excellent charge-transfer characteristics. Thus, designing a TTF containing donor–acceptor based LMWG could be an elegant approach for developing new photocatalyst material. Such systems are likely to show low energy charge transfer in the visible range, which will further reduce the bandgap, and corresponding visible-light photocatalyst can be realized. Coordination driven array of donor–acceptor pairs would further improve photocatalytic performances by enhancing charge transfer to the catalytic centre. Thus, the introduction of a suitable metal-binding moiety like terpyridine on TTF containing LMWG could lead to the formation of CPG and provide an opportunity for further improving the photocatalytic activity.

Here, we aim to report an emerging class of materials known as ‘coordination polymer gel’ by integrating ZnII with TTF-based LMWG and explored as a photocatalyst for solar fuel production based on water and CO2 reduction. The intermolecular charge transfer is regulated by the innovative design of LMWG, where TTF derivative is connected with metal-binding terpyridine units (TPY) through a flexible alkyl amide chain. The coordination polymer gel (Zn-TPT-TTF CPG) provides a suitable platform for light harvesting as well as a catalytic center for H2 production from water (530 μmol g−1 h−1) and CO2 reduction to CO (438 μmol g−1 h−1) with 99% selectivity (Fig. 1). Furthermore,  

![Fig. 1 Schematic presentation of visible/sunlight-driven photocatalysis in TTF-based coordination polymer gel (CPG).](image-url)

A TPY-TTF low molecular weight gelator (LMWG) based linker and suitable metal ions, could be an excellent design approach in the realm of photocatalysis. Such hierarchical soft nanofibrous materials can facilitate the facile diffusion of reactants to the active sites and will show efficient electron transfer between different components. These artificial hybrid synthetic systems can mimic the intricate functioning of the natural photosystem and can eventually show impressive H2 evolution from water or CO2 reduction. Extended face-to-face arrays of the donor–acceptor π-chromophoric systems would be an ideal candidate for light harvesting. These systems will allow greater exciton mobility, which, in turn, leads to charge generation and subsequent electron transfer to the catalyst. To this end, tetrahiafulvalene (TTF) moiety is a well-known p-type semiconductor possessing high electron donation capability with excellent photostability and good charge carrier mobility. Further, the integration of a suitable electron acceptor unit to the TTF moiety could result in a system with excellent charge-transfer characteristics. Thus, designing a TTF containing donor–acceptor based LMWG could be an elegant approach for developing new photocatalyst material. Such systems are likely to show low energy charge transfer in the visible range, which will further reduce the bandgap, and corresponding visible-light photocatalyst can be realized. Coordination driven array of donor–acceptor pairs would further improve photocatalytic performances by enhancing charge transfer to the catalytic centre. Thus, the introduction of a suitable metal-binding moiety like terpyridine on TTF containing LMWG could lead to the formation of CPG and provide an opportunity for further improving the photocatalytic activity.
Zn-TPY-TTF CPG with nanoribbon morphology is conjugated with Pt co-catalyst, and the Pt@Zn-TPY-TTF CPG shows many folds enhanced photocatalytic activity towards H₂ production (14727 μmol g⁻¹ h⁻¹). Interestingly, this Pt@Zn-TPY-TTF CPG produces CH₄ (292 μmol g⁻¹ h⁻¹) as a photoproduct with high selectivity (97%) and impressive quantum efficiency. Importantly, both Zn-TPY-TTF CPG and Pt@Zn-TPY-TTF CPG shows the potential to perform sunlight-driven photocatalytic activity under ambient conditions. The in situ DRIFT study and DFT calculations help to elucidate the plausible mechanism of charge transfer regulated CO₂ reduction to CO/CH₄ formation using CPG catalysts under visible light as well as direct sunlight irradiation.

Results
Preparation and characterization of TPY-TTF OG and Zn-TPY-TTF CPG. The TPY-TTF LMWG was synthesized by the amide coupling reaction between 2,2′,6′,2″-terpyridin-4′-yl-propane-1,3-diamine (TPY-NH₂) and 1,3,6,8-tetrakis (benzoic acid) tetrathiaphthalvalene (TTF-COOH) (details are provided in method section and supplementary information (SI), Supplementary Figs. 1–2). The newly synthesized TPY-TTF LMWG was characterized by H and C Nuclear Magnetic Resonance (NMR), Fourier Transform Infrared (FT-IR) Spectroscopy, and Matrix Assisted Laser Desorption/Ionization–Time of Flight (MALDI-TOF) mass spectrometric analysis (Supplementary Figs. 3–6). UV-vis absorption study was performed for a well-characterized TPY-TTF LMWG in methanol (10⁻⁶ M) and showed distinguished absorption bands at 270 nm and 320 nm corresponding to π−π transition for TPY unit and TTF core, respectively (Fig. 2g). Notably, a low energy absorption band appeared at 520 nm that can be ascribed to intramolecular charge-transfer (CT) interaction between TTF core and benzo-amide moiety. The CT property of TPY-TTF has also been supported by time dependant-density functional theoretical (TD-DFT) computation where the highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO) are centered in TTF and benzo-amide groups, respectively (Supplementary Fig. 7).

We have examined the gelation propensity of TPY-TTF LMWG in several solvent compositions (Supplementary Table 1). The purple-coloured opaque gel of TPY-TTF organogel (OG) was obtained in the solvent mixture (methanol (MeOH), dichloromethane (DCM) and water (H₂O) in 2:1:1 ratio) upon heating at 60 °C followed by cooling to room temperature as shown in Fig. 2a and characterization of gel was performed by different techniques (Supplementary Figs. 8–11). The strain-sweep rheology experiments for TPY-TTF OG at 25 °C showed the values of storage modulus (G') and loss modulus (G'') move constantly in the linear viscoelastic (LVE) region, consisting of the larger G' value under less strain range as compared to the G'', indicating the stable viscoelastic nature, which is a characteristic feature of a gel material (Supplementary Fig. 8). TPY-TTF OG was dried under vacuum at 80 °C to prepare the xerogel and studied the properties of the material (Supplementary Figs. 9–11). Morphology of the TPY-TTF OG xerogel was recorded by the Atomic Force Microscopy (AFM) and Field Emission Scanning Electron Microscopy (FE-SEM) that showed micrometer size stacked layer type of morphology (Fig. 2b, c). We have also prepared aerogel using a critical point dryer (CPD) that also showed similar layered type morphology (Supplementary Fig. 11b). Transmission Electron Microscopy (TEM) images have further confirmed such morphologies (Fig. 2d). The distance between the layers in the stacked morphology was found to be 3.4 ± 0.4 nm based on AFM measurement. The high-resolution TEM analysis showed that the lattice fringes at ~3.7 Å (2d: inset) for the layered morphology suggesting the self-assembly in TPY-TTF OG is driven by the intermolecular π−π interactions. This was also supported by the powder X-ray diffraction (PXRD) study of the xerogel that exhibited a peak at 2θ = 24° with a d-spacing of 3.7 Å (Fig. 2f).

Further, we have performed DFT calculations which also support that the self-assembly is driven by intermolecular π−π stacking interactions between TTF––TTF units at a distance of 3.68 Å (Fig. 2e) and TTF––TTF units at a distance of 4.02 Å (Supplementary Fig. 12)⁵⁰. The UV-vis absorption study for TPY-TTF OG xerogel displayed slightly red-shifted absorption as compared to the methanolic solution of TPY-TTF LMWG (Fig. 2g). It showed absorption bands at 300 nm and 330 nm, which can be assigned for π–π transitions of TPY and TTF units of LMWG, respectively. Notably, a broad absorption between 480 and 615 nm was also observed, indicating the existence of charge transfer (CT) in TPY-TTF OG in the xerogel state. The closer analysis of the CT band showed that it consists of two distinguishable adjacent bands with absorption maxima at 510 and 565 nm (Fig. 2g: inset). Further, in-depth analysis through TD-DFT calculations showed CT absorption band at 498 nm and 564 nm, displaying quite fair agreement with the experimental results. The theoretical absorption at 498 nm was consist of both intramolecular (TTF to PhCONH-) and intermolecular (TTF to TPY) CT transitions (Fig. 4a, Supplementary Table 2). Whereas absorbance at 564 nm was attributed to the intramolecular (TTF-PhCONH-) CT transition. Based on the experimental and theoretical observations, the supramolecular assembly of TPY-TTF OG is represented in Fig. 2h, which showed that for both, TTF––TTF as well as as TTF––CTF stackings are feasible. The optical bandgap for TPY-TTF OG, calculated by the Kubelka-Munk plot derived from UV-Vis diffuse reflectance spectrometry, was found to be 2.26 eV (Supplementary Fig. 13).

Next, the presence of four terpyridine units in the TPY-TTF gelator has prompted us to investigate further their metal-binding ability to develop coordination polymer gel (CPG) for widening their applications. To this end, we have chosen Zn⁺ as a metal node for binding with TPY as such self-assembly is well explored due to soft acid-base interaction⁴⁵,⁵¹. We have performed titration of TPY-TTF (8 × 10⁻⁷ M in MeOH) with a methanolic solution of Zn(NO₃)₂·6H₂O (8 × 10⁻⁴ M), and corresponding UV-vis absorption spectra were recorded (Fig. 3b). Notably, the presence of isosbestic point in the Zn⁺ titration suggested the complex formation between Zn⁺ and TPY-TTF LMWG (Supplementary Fig. 14a, b)⁵². The UV-vis titration study illustrated the binding of Zn(NO₃)₂ and TPY-TTF in the ratio of 2:1.⁴⁷ The association constant (Kₐ) of the Zn⁺ ion with TPY-TTF was calculated to be 2.8 × 10⁸⁷ by the Benesi-Hildebrand plot (Supplementary Fig. 14c). Next, Zn(NO₃)₂ and TPY-TTF gelator was taken in a molar ratio of 2:1 in the solvent mixture of MeOH, DCM and H₂O in 2:1:1 ratio, respectively. Heating the reaction mixture to 60 °C followed by cooling to room temperature has afforded a deep purple-coloured coordination polymer gel (Zn-TPY-TTF CPG) (Fig. 3a). Similar to OG, strain-sweep rheology experiments were performed for Zn-TPY-TTF CPG at 25 °C (Supplementary Fig. 15). The value of G' and G'' up to 0.01% was found to be ~10 times larger for the Zn-TPY-TTF CPG than the OG, illustrating higher stability of the former one, which is possibly attained due to coordination of the Zn⁺ ion⁵³. More importantly, in both cases, tan δ = G'/G'' value was found to be lower than one before reaching yield strain point, which is the intrinsic property of the gel phase (Supplementary Fig. 16). Next, the morphology of the Zn-TPY-TTF CPG was evaluated by FE-SEM upon drying the sample at 80 °C under vacuum, which showed the 3D entangled nanofibrous morphology (Fig. 3c). The aerogel of CPG was also prepared by the CPD method, which showed a
Fig. 2 Preparation and characterizations of TPY-TTF OG. a Photograph for organogel formation. Morphological analysis for xerogel. b AFM image. c FE-SEM image. d HR-TEM image (inset: showing lattice fringes). e Interplanar spacing in the optimized structure of TTF—TPY stacked model obtained through DFT calculation. f PXRD pattern for xerogel. g Comparison of absorption spectra of TPY-TTF LMWG in solution (8 × 10^{-6} M) with TPY-TTF OG xerogel state. h Pictorial representation for self-assembly in TPY-TTF OG forming layered sheet-like morphology.

Fig. 3 Preparation and characterizations of Zn-TPY-TTF CPG. a Photograph for Zn-TPY-TTF CPG formation. b UV-visible titration of TPY-TTF LMWG with Zn^{II} ion. c FE-SEM image of Zn-TPY-TTF CPG xerogel. d HR-TEM image of Zn-TPY-TTF CPG xerogel. e Lattice fringes of the selected region. f AFM image of Zn-TPY-TTF CPG xerogel and corresponding, g height profile and h height histogram. i Comparison of absorption spectra of TPY-TTF OG and Zn-TPY-TTF CPG. j Interplanar spacing in the optimized structure of TTF—[Zn(TPY)_{2}]^{2+} stacked model obtained through DFT calculation. k PXRD of Zn-TPY-TTF CPG xerogel. l Schematic representation for self-assembly of Zn-TPY-TTF CPG.
similar morphology suggesting that the fibrous morphology of Zn-TPY-TTF CPG is driven through metal coordination with LMWG (Supplementary Fig. 17). TEM studies revealed that nanoribbons form 3D interconnected fibrous morphology (Fig. 3d). The AFM images of Zn-TPT-TTF CPG revealed the height of nanoribbon is ~7 nm and the diameter is in the range of 40–80 nm (Fig. 3f–h). The elemental mapping of the xerogel exhibited the uniform distribution of ZnII in a 3D network of the CPG (Supplementary Fig. 18). EDAX and elemental analyses also correlated the 2:1 ratio of ZnII: TPT-TTF in the CPG (Supplementary Fig. 19). The high-resolution TEM analysis exhibited ordering in the nanoribbon, and lattice fringes were observed with a distance of 3.6 Å, which could be attributed to the intermolecular π–π stacking between the TTF–TPY units from the [Zn(TPY)]22+ (Fig. 3e), which was also observed in the gel state, justifying the presence of π–π stacking (Fig. 3k, Supplementary Fig. 20). In order to evaluate the nature of stacking, TD-DFT calculations were performed, which showed that the packing of TTF with [Zn(TPY)]22+ on top of each other was stabilized with a distance of 3.56 Å (Fig. 3j, l) which is in good agreement with the experimental observations50,54. On the contrary, stacking through TTF–TTF units on the top of each other was optimized, which revealed the TTF–TTF distance >11 Å due to steric repulsion among the [Zn(TPY)]22+ units attached to TTF core, and therefore, the possibility of TTF–TTF stacking was ruled out (Supplementary Fig. 22a). Further, the UV-vis absorption spectrum of Zn-TPT-TTF CPG in xerogel state was found to be similar to the TPT-TTF OG with an enhanced absorption in the visible region as shown in Fig. 3i. To analyse the reason behind enhanced absorption in the visible range, TD-DFT calculations were performed for the stacked model of Zn-TPT-TTF CPG (Fig. 4b). The result showed that the band

observed at 510 nm was similar to TPT-TTF OG. Whereas the experimental band at 565 nm is mainly attributed to the theoretical band at 553 nm (Fig. 4b, Supplementary Fig. 22b). Importantly, the transition at 553 nm was observed due to the significant contribution of intermolecular CT from TTF core to [Zn(TPY)]22+ unit as shown in Supplementary Table 255. The dominated intermolecular CT transition in Zn-TPT-TTF CPG as compared to TPT-TTF OG is mainly triggered by the planarization of terpyridine ligand, which occurred after the complexation with ZnII ion in CPG56. Additionally, ZnII complexation also increased the electron-accepting tendency of terpyridine ligand from the TTF moiety, which further facilitated the intermolecular CT transition57. The optical bandgap for Zn-TPT-TTF CPG was calculated to be 2.27 eV, which is closer to the bandgap of TPT-TTF OG (Supplementary Fig. 13). Next, as a controlled study, we found energies of the theoretical band at 553 nm was observed due to the transition at 500 nm for Zn-TPY-TTF CPG was calculated to be 2.17 eV, respectively, in the aqueous medium (Fig. 4c), which indicates that excited-state electron transfer is energetically favourable from TTF(PhCONH2)4 core to [Zn(TPY)]22+ centre.

Mott-Schottky analysis was performed for the xerogel of both TPT-TTF OG and Zn-TPT-TTF CPG to evaluate experimental feasibility for water and CO2 reduction (see SI for details). The M-S plots exhibited n-type nature with a positive slope for both TPT-TTF OG and Zn-TPT-TTF CPG (Fig. 5a). The flat band potentials ($V_{fb}$) were found to be ~0.54 V versus RHE (at pH 7) for TPT-TTF OG and Zn-TPT-TTF CPG, respectively. Based on the bandgaps obtained using UV-vis diffuse reflectance spectrometry (Supplementary Fig. 13), the electronic band structures versus RHE at pH 7 could be elucidated and are displayed in Fig. 5b58. Interestingly, the band alignments are shown in Fig. 5b illustrate that both TPT-TTF OG and Zn-TPT-TTF CPG possess suitable band edge positions to perform water and CO2 reduction under visible-light irradiation.
H₂ production activity of OG and CPG under laboratory condition. We have examined the potential of Zn-TPY-TTF CPG for photocatalytic H₂ production from water under visible-light (400–750 nm) irradiation using 300 W xenon lamp as the light source (Supplementary Figs. 23–25). The photocatalytic H₂ production activity of Zn-TPY-TTF CPG was screened with different sacrificial electron donors, and the best activity was found with triethylamine (TEA) as a sacrificial agent (Supplementary Fig. 26). Photocatalytic activity of Zn-TPY-TTF CPG was examined in both gel and xerogel state, and similar H₂ evolution was observed for both under similar conditions (Supplementary Fig. 27a). However, catalytic activities in different conditions were performed using the xerogel because of the ease of handling the catalyst in comparison to the gel state. After optimizing the catalyst loading (Supplementary Fig. 28), 1 mg of Zn-TPY-TTF CPG in xerogel state was dispersed in 38 ml of water for the photocatalytic H₂ production, and 2 ml triethylamine (TEA) was added into it, which acted as a sacrificial electron donor. The photocatalytic activity of Zn-TPY-TTF CPG was monitored by gas chromatography (GC) analysis and showed 10.60 mmol g⁻¹ of H₂ evolution in 20 h (activity = 530 μmol g⁻¹ h⁻¹) upon visible-light irradiation as shown in Fig. 5c. The amount of H₂ evolved was reached saturation in 20 h, and the turnover number (TON) was calculated to be 23.5, as shown in Fig. 5d. The activity is higher than many other transition metal-based photocatalysts (Supplementary Tables 6–8). The apparent quantum efficiency (AQE) for H₂ production using Zn-TPY-TTF CPG catalyst was calculated to be 0.76% at 550 ± 10 nm. Further, the absence of H₂ formation with Zn-TPY-TTF CPG under dark condition (absence of light) confirming light is an essential component for catalysis. Next, we have also performed the recyclability test by recollecting the catalyst followed by reusing the photocatalysis for four additional cycles of 6 h each time (Supplementary Fig. 27b). Interestingly, the amount of H₂ evolution was found to be similar in every cycle (>99%). We also performed ICP analysis for as-synthesized Zn-TPY-TTF CPG sample as well as recovered sample after photocatalysis. In both cases, the amount of Zn content was calculated to be 5.9 (±0.2) wt%. Next, recycled catalyst (Zn-TPY-TTF) was collected at the end of the fourth cycle and analysed by FE-SEM and TEM studies and suggested no significant change in the structure and morphology after the catalytic reaction indicating high stability of the catalyst (Supplementary Fig. 27c, d).

Next, we have examined photocatalytic activity for the OG to compare the importance of morphology, i.e. the spatial arrangement of the chromophore and also the role of metal directed assembly in CPG. Experimental conditions employed for the TPY-TTF OG was similar to the CPG. Interestingly, the H₂ evolution by the TPY-TTF OG upon visible-light irradiation was increased with irradiation time and reached saturation in 22 h (Fig. 5c). The maximum H₂ evolution in 20 h was calculated to be 2 mmol g⁻¹ (activity = 100 μmol g⁻¹ h⁻¹), which is albeit lesser than the CPG but higher than most of the reported metal-free photocatalysts for H₂ evolution59. To understand the significant difference in photocatalytic activities between TPY-TTF OG and Zn-TPY-TTF CPG, photocurrent measurements were performed for both (Fig. 5e) in the presence and absence of light. The photocurrent for Zn-TPY-TTF CPG in the presence of light was found to be double as compared to TPY-TTF OG. This indicates the facile charge separation in Zn-TPY-TTF CPG under light irradiation; therefore, expected to show better photocatalytic activity as compared to TPY-TTF OG. This argument has been further validated by the EIS measurement, where the charge-transfer resistance for Zn-TPY-TTF CPG was observed to be significantly lesser as compared to TPY-TTF OG under both dark and light irradiated conditions (Fig. 5f). This can be attributed to the nanoribbon morphology of Zn-TPY-TTF CPG that provides a continuous charge-transfer pathway (via co-facial intermolecular charge delocalization) for the photogenerated electrons, which ultimately enhances the photocatalytic activity.
We have also performed the photocatalytic study with individual structural units of Zn-TPY-TTF CPG to evaluate the importance of coordination driven assembly of CPG in catalysis. Notably, individual components, such as TEA, TTF(COOH)₄, TPY-NH₂, and [Zn(TPY-NH₂)₂]²⁺ as well as the physical mixture of TTF(COOH)₄ and [Zn(TPY-NH₂)₂]²⁺ were found to be not efficient in catalysing H₂ evolution reaction (Supplementary Fig. 29). Next, the photocatalytic study has also been performed by making a blend of Zn⁹⁺ salt with PTY-TTF (in 2:1 ratio) (details are given in SI). This showed aggregated spherical morphology (sphere diameter 80 ± 10 nm) as confirmed by FE-SEM study (Supplementary Fig. 30), and the corresponding H₂ evolution within 6 h was found to be three times lesser as compared to the CPG (Supplementary Fig. 29). This signifies the impact of nano-structuring in photocatalytic performances. To validate the role of intermolecular CT interaction, we have also synthesized a coordination polymer of Zn⁹⁺ (Zn-CP) with TTF(COOH)₄ and characterized by FE-SEM, EDAX, elemental mapping, TGA, PXRD and UV-vis absorption study (Supplementary Figs. 31–34). Zn-CP showed micron-sized spherical particles with a diameter in the range of 2–3 µm. The Zn-CP showed 0.8 mmol g⁻¹ of H₂ production from water in 12 h which is eight times lesser in activity in comparison to Zn-TPY-TTF CPG photocatalyst under a similar condition (Supplementary Fig. 29). Overall, control experiments have unambiguously indicated that the coordination driven spatial arrangement of donor–acceptor chromophores and corresponding CT interaction have high significance in visible-light photocatalytic performances of CPG (Zn-TPY-TTF).

Preparation and characterizations of Pt@Zn-TPY-TTF CPG. Further, we envisioned that the entangled hierarchical fibrous structure of the coordination polymer gels could easily immobilize co-catalyst like Pt on the surface⁶⁰, which would facilitate the separation of photogenerated charge carriers by decreasing diffusion length and eventually enhance the photocatalytic activity⁶¹. Thus, we have successfully executed in situ generation and stabilization of platinum nanoparticles (Pt NPs) in the self-assembled interconnected network of Zn-TPY-TTF CPG (Pt@Zn-TPY-TTF) (Fig. 6a). High-resolution TEM and FE-SEM analysis have confirmed the stabilization of Pt NPs in self-assembled networks of Pt@Zn-TPY-TTF CPG within the size range of 2–3 nm (Fig. 6b, Supplementary Fig. 36a). Lattice fringes were observed for Pt NPs with the d-spacing value of 0.23 nm, indicating the presence of Pt (111) planes (Fig. 6c). Inductively coupled plasma-optical emission spectroscopy (ICP-OES) measurement indicated the presence of ~2.7 wt% Pt in Pt@Zn-TPY-TTF CPG, which is also supported by EDAX analysis (Supplementary Fig. 36b). The elemental mapping was ensured for the uniform distribution of Pt NPs in the gel matrix (Supplementary Fig. 37). Further, the Mott-Schottky analysis for Pt@Zn-TPY-TTF CPG also revealed that the conduction band edge occurs at ~0.51 V vs RHE at pH 7, which is lesser compared to the Zn-TPY-TTF CPG (~0.54 V) catalyst (Supplementary Fig. 38a).

H₂ production activity of Pt@Zn-TPY-TTF CPG. Next, photocatalytic activity towards water reduction was examined for Pt@Zn-TPY-TTF CPG under a similar condition as employed for CPG as well as OG. Interestingly, Pt@Zn-TPY-TTF CPG has shown remarkably enhanced catalytic activity under visible-light irradiation, and the amount of H₂ was calculated to be 162.42 mmol g⁻¹ in only 11 h (average activity = 14727 µmol g⁻¹ h⁻¹), and the corresponding TON value was found to be 1176.9 (w.r.t. 2.7 wt% of Pt, Supplementary Table 3) as shown in Fig. 6d. We have also investigated photocatalytic H₂ production activity by varying loading amount of Pt NPs, which displayed the highest catalytic activity in the presence of 2.7 wt% of Pt to the CPG (Supplementary Fig. 35). Drastically increased H₂ evolution after Pt NPs stabilization could be ascribed to the efficient charge separation in Pt@Zn-TPY-TTF CPG as Pt centre is a well-known electron acceptor that accumulates a pool of electrons and subsequently exhibits efficient water reduction. Here, the photoexcited electrons from Zn-TPY-TTF CPG using the harvested light energy transferred to Pt NPs surface to reduce water into H₂. Furthermore, the recyclability and reusability of Pt@Zn-TPY-TTF CPG towards photocatalytic hydrogen evolution were evaluated upto four cycles similar to the Zn-TPY-TTF CPG (Supplementary Fig. 39). Photocatalytic activity of the recycled catalyst was found to be retained >95% after 4th cycle. The formation of the Schottky junction in Pt@Zn-TPY-TTF CPG was helpful to separate the photogenerated electron-hole pairs. This argument was further validated by the EIS measurement, where the charge-transfer resistance for Pt@Zn-TPY-TTF CPG was found to be almost half as compared to Zn-TPY-TTF CPG under both dark and visible-light irradiation (Supplementary Fig. 38b). Furthermore, approximately four-folds higher photocurrent was observed for Pt@Zn-TPY-TTF CPG as compared to Zn-TPY-TTF CPG (Supplementary Fig. 38c), which corroborated the facile electron transfer from the Zn-TPY-TTF CPG to the Pt centre. Further, the photoluminescence (PL) spectra of Zn-TPY-TTF CPG showed weak emission with a maximum at 581 nm due to the intermolecular charge-transfer interaction (λex = 510 nm). (Supplementary Fig. 40). PL spectra for Pt@Zn-TPY-TTF CPG upon excitation at 510 nm showed significantly quenched emission compared to Zn-TPY-TTF CPG. Therefore, to gain more insight into the advantages of Pt NPs in the increased charge separation, the time-resolved photoluminescence (TRPL) decay was studied on the Zn-TPY-TTF CPG and Pt@Zn-TPY-TTF CPG, as shown in Supplementary Fig. 41 (Supplementary Table 4). A higher average lifetime Zn-TPY-TTF CPG system (1.95 ns) compared to Pt@Zn-TPY-TTF CPG system (0.22 ns) obtained by the TRPL studies confirms that migration of photoexcited electrons is much faster in Pt@Zn-TPY-TTF CPG compared to Zn-TPY-TTF CPG. The decrease in the lifetime is attributed to the enhanced separation of photogenerated electrons in Pt@Zn-TPY-TTF CPG, which play decisive roles in enhancing the efficiency of photocatalytic processes. To understand the charge-separation dynamics, transient absorption (TA) experiments were performed with Zn-TPY-TTF CPG and Pt@Zn-TPY-TTF CPG. The TA spectra of Zn-TPY-TTF CPG (Supplementary Fig. 42a) showed a broad excited-state absorption (ESA) band covering 450–700 nm region at early time delay. Such a broad ESA band is due to TTF cation and TPY anion radical formed by the electron transfer process by the photoexcitation. TTF cation has an absorption band at ~440 nm with a long tail extending up to 700 nm, and Zn-TPY anion radical complex has a broad absorption band covering the range of 500–600 nm. Thus, the appearance of a broad band at early time delay suggested that the electron transfer from TTF to Zn-TPY is very efficient in Zn-TPY-TTF CPG and takes place within the time resolution of the TA instrument (120 fs). The fitting of the decay trace at 570 nm due to TPY anion radical complex gives an average lifetime of 563 ps (Supplementary Fig. 42b). Pt@Zn-TPY-TTF CPG showed qualitatively similar TA spectra as of Zn-TPY-TTF CPG. However, the transient signal at 570 nm decays significantly faster in Pt@Zn-TPY-TTF CPG (average lifetime is 174 ps, see Supplementary Fig. 42c). Such faster decay kinetics in Pt@Zn-TPY-TTF CPG suggest electron transfer from reduced Zn-TPY to Pt NPs. Besides the TRPL and TA results,
DFT calculations also revealed that in Pt@Zn-TPY-TTF CPG, the electron transfer could take place from TTF to Pt NPs via [Zn(TPY)]2+. We have investigated the possible loading positions of Pt NPs in the CPG theoretically and the details have been given in SI (Supplementary Figs. 43–44). We were able to perform the stabilization energy calculations up to four atoms Pt cluster due to computational limitation (Supplementary Fig. 44) and detailed experimental study related to Pt NPs stabilization will be carried out in future works. Furthermore, AQE for the water reduction to H2 was determined for the Pt@Zn-TPY-TTF CPG upon illuminating with monochromatic light of the wavelength of 400 ± 10 nm, 450 ± 10 nm, 500 ± 10 nm, 550 ± 10 nm, 600 ± 10 nm, 650 ± 10 nm, and 700 ± 10 nm (Supplementary Table 5, Supplementary Fig. 45). Notably, the highest AQE was obtained to be 14.47% at 550 ± 10 nm. These experiments suggested that the photocatalytic activity is mainly driven through intermolecular charge-transfer interaction. The H2 evolution using Pt@Zn-TPY-TTF CPG was examined under both light and dark conditions (Supplementary Fig. 39c). No H2 evolution was detected under dark condition, indicating the importance of light for water reduction. Next, photocatalysis was also performed for Pt@Zn-TPY-TTF and Zn-TPY-TTF CPG without any sacrificial donor (TEA). The aqueous dispersion of Zn-TPY-TTF CPG produced 0.92 mmol g−1 of H2 in 20 h, which is 11 times lesser than with TEA. Similarly, Pt@Zn-TPY-TTF CPG showed 30 times lesser activity without TEA (Supplementary Table 6).

CO2 reduction activity of OG and CPG under laboratory condition. As mentioned above, the theoretical and experimental bandgap alignment of Zn-TPY-TTF CPG and TPY-TTF OG has the potential to reduce CO2 as well. Therefore, we have performed visible-light-driven photocatalytic CO2 reduction with the xerogel of Zn-TPY-TTF CPG and also compared it with TPY-TTF OG. TEA was used as a sacrificial electron donor for CO2 reduction. First, screening of the solvent composition for CO2 reduction has been performed (Supplementary Figs. 47–48, Supplementary Table 10), and the mixture of acetonitrile: water (3:1) have shown the best activity which could be attributed to high solubility of CO2 in acetonitrile, whereas, water acted as a proton source65. Visible-light-driven CO2 reduction by Zn-TPY-TTF CPG yielded 3.51 mmol g−1 of CO in 8 h with >99% selectivity (activity = 438 μmol g−1 h−1) as shown in Fig. 7b. The AQE of CO2 photoreduction for Zn-TPY-TTF CPG at 550 ± 10 nm was calculated to be 0.96%. Such a selective CO production is noteworthy and one of the best results among various reported hybrid photocatalysts systems (Supplementary Table 12). The TON for Zn-TPY-TTF CPG in 8 h was calculated to be 7.8 (Fig. 7b). Recyclability test for the Zn-TPY-TTF CPG photocatalyst was performed for four cycles, similarly as employed for the water reduction. The photocatalytic performances of Zn-TPY-TTF CPG were found to be almost unchanged (Supplementary Fig. 49). Further, the photocatalytic activity of Zn-TPY-TTF CPG was examined upon isotopic labelling with13CO2. This showed the formation of 13CO, which confirms that the produced CO was originated from CO2 (Supplementary Figs. 50–51). Next, visible-light-driven photocatalytic CO2 reduction has also been investigated for TPY-TTF OG under a similar condition as employed for Zn-TPY-TTF CPG. The TPY-TTF OG has displayed 1.12 mmol g−1 CO formation in 11 h with >99% selectivity (activity = 140 μmol g−1 h−1), and the corresponding TON was estimated to be 2.1 as shown in Fig. 7a.

We have performed in situ Diffuse Reflectance Infrared Fourier Transform (DRIFT) spectroscopic study for Zn-TPY-TTF CPG to track the reaction intermediates formed in the course of CO2 reduction to CO (Fig. 8a)38. Two peaks that appeared at 1514 and 1692 cm−1 were assigned for the COOH− and COO− intermediates, respectively, which are the signature intermediates formed during CO2 reduction process (Fig. 8a)66. Peak observed at 1454 cm−1 could be attributed to symmetric stretching of the HCO3− 67. A noteworthy peak at 2074 cm−1 was indicated for the formation of the CO−. Most importantly, the peak intensity
the CO\(^\ddagger\) intermediate was substantially increased with reaction progress, suggesting CO\(^\ddagger\) formation increases with time. Based on the experimental results and in situ DRIFT study, we have computed a plausible mechanism for CO\(_2\) reduction, which is also in good agreement with the earlier report\(^{68}\) (Supplementary Figs. 65–66). The photocatalytic cycle was initiated by the light absorption due to CT interaction in Zn-TPY-TTF CPG, and electron transfer took place from the excited state of TTF\(^*\) to [Zn(2-TPY)\(_2\)]\(^{2+}\) units followed by reductive quenching of TTF\(^*\) by TEA. As a result, the [Zn(2-TPY)\(_2\)]\(^{2+}\) converted to the radical cation species [Zn(2-TPY\(^−\))\(_2\)]\(^{+}\). Notably, the electron spin density distribution plot revealed that the electron was localized at the terpyridine unit of the catalyst that resulted in [Zn(TPY\(^−\))\(_2\)]\(^{+}\) species with the stabilization energy of 0.52 eV. (Supplementary Fig. 65d and Supplementary Table 35). Next, as the reaction progress, the acetonitrile (CH\(_3\)CN), solvent molecule replaced one ligating site of terpyridine to afford [Zn(TPY\(^−\))\(_2\)(\(\eta^2\)-TPY)(CH\(_3\)CN)]\(^+\) which is slightly uphill (\(\Delta G = +0.77 \text{ eV}\)) with a low activation barrier of 0.81 eV. This [Zn(TPY\(^−\))\(_2\)(\(\eta^2\)-TPY) (CH\(_3\)CN)]\(^+\) intermediate subsequently binds with CO\(_2\) molecule with a distance of 2.41 Å to Zn\(^{II}\) ion and yielded the [Zn(TPY)\(_2\)(\(\eta^2\)-TPY)(COO\(^−\))]\(^+\) species (\(\Delta G = +0.46 \text{ eV}\)). The formulation of the intermediate as [Zn(TPY)(\(\eta^2\)-TPY)(COO\(^−\))]\(^+\) rather than [Zn(TPY\(^−\))(\(\eta^2\)-TPY)(COOH)]\(^+\) is supported by the spin density distribution plot (Supplementary Table 38). Notably, the CO\(_2\) molecule acted as a monodentate ligand, and the \(\angle\text{O–C–O}\) angle was found to be 139.36\(^\circ\), keeping in mind that the free CO\(_2\) has \(\angle\text{O–C–O}\) = 139.48\(^\circ\). Notably, the CO\(_2\) complex gets readily protonated and results in the formation of CO\(_2\)⋅H\(_2\)O. As a result, the CO\(_2\) complex favours the binding of CO\(_2\) molecule by increasing electron density around the metal centre, which is a prerequisite for the nucleophilic attack to the CO\(_2\). However, the complex [Zn(TPY)(\(\eta^2\)-TPY)(COOH)]\(^+\) was further reduced to form the singlet species [Zn(TPY\(^−\))(\(\eta^2\)-TPY) (COO\(^−\))]\(^−\) (\(\Delta G = −0.79 \text{ eV}\)) (Supplementary Fig. 65g and Supplementary Table 39). Next, the carboxylate centre of the singlet complex gets readily protonated and results in the formation of [Zn(TPY\(^−\))(\(\eta^2\)-TPY)(COOH)]\(^+\) complex (\(\Delta G = −2.88 \text{ eV}\)) (Supplementary Fig. 65h and Supplementary Table 40). In the next step, subsequent protonation and water elimination
from [Zn(TPY−)(η2-TPY)(COOH)]+ intermediate lead to the formation of [Zn(TPY)(η2-TPY)(CO)]2+ which is found to be a highly downhill process (ΔG = −2.55 eV). (Supplementary Fig. 65i and Supplementary Table 41). In this complex, the CO molecule is loosely attached to the ZnII centre at a distance of 2.66 Å. As a result, the CO molecule can be easily released from the metal centre. Thus, the intermediate [Zn(TPY)(η2-TPY)(CO)]2+ gets readily reduced, leading to the subsequent removal of CO (ΔG = −1.51 eV) which regenerates the active species [Zn(TPY−)(TPY)]+ and re-enters into the catalytic cycle. Specifically, in this photocatalytic CO2 reduction mechanism, the Zn-terpyridine complex involves retention of the bisterpyridine ligand to gain the original coordination environment similar to as-synthesized Zn-TPTY-TTF CPG.

**CO2 reduction activity of Pt@Zn-TPTY-TTF CPG.** Next, the photocatalytic activity of Pt@Zn-TPTY-TTF CPG towards CO2 reduction has also been examined under a similar experimental condition as applied for CPG and OG. Pt@Zn-TPTY-TTF CPG showed excellent CO2 reduction activity, and more interestingly, CH4 was produced rather than CO as obtained for Zn-TPTY-TTF CPG (Fig. 7d). It has already been reported in the literature that the presence of Pt NPs on the surface of semiconductor plays a key role in the formation of CH4.60 The formation of CH4 has reached saturation in 30 h, and the corresponding yield was calculated to be 8.74 mmol g−1 (activity = 292 μmol g−1 h−1) (Fig. 7d). During CO2 reduction, a small amount of H2 evolution was also observed (~ 0.20 mmol g−1 in 30 h), which was more in the initial hours but significantly decreased as the reaction progressed with time. Thus, the selectivity of CO2 reduction to CH4 formation in 30 h was noted to be more than 97% (Fig. 7e). The TON for CH4 was calculated to be 63.4 (w.r.t. Pt loading on Zn-TPTY-TTF CPG; Fig. 7e and Supplementary Table 3) in 30 h, which is better than many of the earlier reports (Supplementary Table 12). The recyclability of the catalyst, Pt@Zn-TPTY-TTF CPG was examined for 6 h up to four cycles, and the amount of CH4 formation in the fourth cycle was found to be >95%, indicating high stability of the catalyst (Supplementary Fig. 56). The ICP-OES analysis of Pt@Zn-TPTY-TTF CPG was also performed after the fourth catalytic cycle, which showed 2.52 wt% of Pt present in the recovered catalyst. The partial decrease in catalytic activity (<5%) could be attributed to the minimal loss of Pt NPs during the recycling process. The AQE of the Pt@Zn-TPTY-TTF CPG towards CO2 reduction was calculated at different wavelengths using monochromatic lights (Supplementary Fig. 54 and Supplementary Table 9). The highest AQE for the CH4 formation was obtained as 0.81% at 550 ± 10 nm, which further confirms that the photocatalytic activity of Pt@Zn-TPTY-TTF is attributed to the intermolecular charge-transfer interactions. The CO2 reduction was also performed under both light and dark conditions using Pt@Zn-TPTY-TTF (Supplementary Fig. 55). The CH4 formation was not increased under dark condition, justifying the importance of light in photocatalysis. Further, photocatalysis was performed with labelled 13CO2 (isotopic labelling) using Pt@Zn-TPTY-TTF (Supplementary Figs. 52–53). This showed the formation of labelled 13CH4 and confirming that CO2 is the actual source for the CH4 formation. Next, we have performed in situ DRIFT experiment for Pt@Zn-TPTY-TTF in order to monitor the real-time progress of the CO2 reduction reaction (Fig. 8b). IR-stretching peaks observed at 1610, 1570, and 1426 cm−1 could be attributed to the intermediates COOH*, COO*, and HCO3−, respectively.58 Weak intensity peak for the CO* at 2062 cm−1 illustrated that the CO* could be readily converted to other multi-electron reduction intermediates. Moreover, the characteristic intermediates for CH4 formation were observed at 1081 cm−1 (CHO−), 1015 and 1120 cm−1 (CH3O)58. Based on the DRIFT study, the plausible mechanism for the CH4 formation was elucidated with the help of free energy calculation of different intermediates using DFT, which suggested that the Pt NPs is likely to be the catalytic centre during the CO2 reduction (Supplementary Figs. 67–70).

**Sunlight-driven H2 production.** The above discussions have clearly shown that visible-light photocatalytic activity and stability of both Zn-TPTY-TTF CPG and Pt@Zn-TPTY-TTF CPG in the xerogel state is indeed impressive. Notably, the amount of H2 evolution using photocatalyst Zn-TPTY-TTF CPG is higher than previously reported non-precious metal-based photocatalyst materials (Supplementary Table 8). Therefore, our next goal was to examine the potential of Zn-TPTY-TTF towards H2 evolution upon direct sunlight irradiation at ambient condition (Supplementary Fig. 24). The experiment was performed with Zn-TPTY-TTF CPG under sunlight from 10:00 am to 4:00 pm for one week from 12th to 17th April 2019 on the rooftop of our institute. The weather condition corresponding to the above-mentioned period can easily be found out on the web (www.timeanddate.com). Interestingly, maximum H2 evolution of 5.14 mmol g−1 in 6 h (activity = 857 μmol g−1 h−1) was observed on 15th April, which is comparable with the amount of H2 obtained under laboratory conditions (Xe-lamp irradiation). The TON for H2 evolution was calculated for the above-mentioned period (Fig. 5g). The highest TON value of 11.9 was obtained on 15th April 2019. Whereas the lowest TON was calculated to be 7.2 on 12th April 2019 because of partially cloudy weather (Supplementary Table 7). Next, similar to Zn-TPTY-TTF CPG, we have also examined sunlight-driven photocatalytic H2 evolution for Pt@Zn-TPTY-TTF CPG in xerogel state (Fig. 6e). The experimental condition for Pt@Zn-TPTY-TTF CPG was similar to the Zn-TPTY-TTF CPG. Nevertheless, experimental timing was different for the Pt@Zn-TPTY-TTF CPG. The experiment with Pt@Zn-TPTY-TTF CPG was performed from 27th to 29th June 2019. The highest H2 evolution was calculated to be 72 mmol g−1 in 6 h (activity = 12,000 μmol g−1 h−1) on 27th June 2019, and the corresponding TON was calculated to be 521.8 (w.r.t. Pt loading), which is indeed noteworthy (Fig. 6e; Supplementary Table 3). Whereas the lowest TON value for H2 evolution was found to be 376.48 (w.r.t. Pt loading) on 28th June due to partially cloudy weather.

**Sunlight-driven CO2 reduction.** Interestingly, the potential of Zn-TPTY-TTF CPG for CO2 reduction has also been examined under direct sunlight between 10:00 am to 4:00 pm for three days from 28th to 30th September 2019. The highest CO formation of 1.79 mmol g−1 was observed in 6 h (activity = 298 μmol g−1 h−1) on 30th September 2019, and the corresponding TON was calculated to be 3.9 (Fig. 7c). Next, sunlight-driven CO2 reduction has been performed with Pt@Zn-TPTY-TTF CPG for three days from 1st to 3rd October 2019 (Fig. 7f). Similar to the laboratory conditions, Pt@Zn-TPTY-TTF CPG upon sunlight irradiation has displayed CH4 formation (Supplementary Table 11). The highest CH4 formation of 0.96 mmol g−1 in 6 h (activity = 160 μmol g−1 h−1) was observed on 1st October 2019, and the corresponding TON value was calculated to be 6.9 (w.r.t. Pt loading; Supplementary Table 3). Whereas the lowest CH4 evolution with the TON value of 6.24 in 6 h (w.r.t. Pt loading) took place on 2nd October 2019. The CO or CH4 formation under direct sunlight is almost linearly related to the laboratory condition, but the obtained amount under ambient condition is quite exciting and promising because of practical application. Furthermore, after performing sunlight-driven photocatalysis with Zn-TPTY-TTF CPG and Pt@Zn-TPTY-TTF CPG catalysts were recovered.
through centrifugation and washed with fresh water 3–4 times. The FE-SEM and TEM analysis, PXRD, FT-IR and UV-vis absorption experiments were performed for the recovered catalysts and found to be similar to the as-synthesized material, indicating that the structural integrity of the material remained intact after photocatalysis (Supplementary Figs. 57–64). Whereas the EDAX analysis ensured the presence of all the elements in similar quantity as obtained for as-synthesized Zn-TPY-TTF and Pt@Zn-TPY-TTF CPG (Supplementary Figs. 57 and 61). This unambiguously demonstrated the high stability of these catalysts during sunlight irradiation.

Discussion

In a nutshell, we have successfully demonstrated a TTF-based soft processable metal-organic hybrid gel as a visible-light photocatalyst for H₂ evolution and CO₂ reduction to carbonaceous fuel such as CO/CH₄. Here, charge-transfer-driven photocatalysis based on coordination polymer gel has been exploited, where earth-abundant metal ions play a crucial role in the spatial organization of donor–acceptor π-chromophores to drive the catalytic activity. Further, we have shown the catalytic activity of the CPG after decorating Pt NPs as co-catalyst. It enhances the rate of H₂ production in 20 folds and dramatically changes the CO₂ reduction product from CO to CH₄. We have also demonstrated efficient catalytic activity of the CPG and Pt decorated CPG under sunlight with high selectivity. The real-time reaction progress of CO₂ reduction was monitored by DRIFT studies, and based on that, a plausible mechanism of CO₂ reduction was elucidated for CPG. The easy processability and structural tunability of LMWG offers a lot of room for designing efficient photocatalyst materials for practical application. Our work will pave the way toward designing coordination driven hybrid soft processable photocatalyst systems for solar energy driven fuel production.

Methods

Reagent. Tetraphiafulvalene (TTF), 1,3-diaminopropane, 4′-chlooro-2,2′,6′,2′-terpyridine, Zinc nitrate (Zn(NO₃)₂·6H₂O), Ethyl-4-bromobenzoate (Br–C₆H₄COOEt), Cesium carbonate (Cs₂CO₃), Palladium acetate, Tri-tert-butylphosphonium trifluoroborate (Pbuno, HBF₄), Thionyl chloride (SOCl₂), Chloro-platinic acid (H₂PtCl₆·6H₂O) were purchased from Sigma–Aldrich chemical Co. Ltd. Spectroscopic grade solvents were used for all spectroscopic experiments without further purification.

Synthesis of TPY-TTF LMWG. TPY(COOH), (634 mg, 1.65 mmol) was dissolved in 50 ml of dry tetrahydrofuran (THF), and thionyl chloride (SOCl₂) (2.4 ml, 33 mmol) was added into it under inert condition. The reaction mixture was refluxed for 2 h at 65 °C. Then the reaction mixture was distilled at 120 °C to remove excess SOCl₂ and yielded a solid precipitate of acid chloride. Next, the solid precipitate was dissolved in 40 ml of dry THF. Now the solution of TPY-NH₂ (2.21 g, 7.26 mmol) in 10 ml of dry THF along with triethylamine (1.25 ml, 9 mmol) was added to the solution of acid chloride dropwise at 0 °C. The reaction mixture was stirred overnight and then heated at 80 °C under vacuum for 8 h. Finally, the Pt nanoparticle stabilized CPG sample (Pt@Zn-TPY-TTF) was thoroughly washed with deionized (DI) water and dried under vacuum at 80 °C for 12 h. More detailed characterization of Pt@Zn-TPY-TTF CPG is provided in Supplementary Information.

Preparation of Zn-TPY-TTF CPG. We have used Zn(NO₃)₂·6H₂O salt to synthesize Zn-TPY-TTF CPG. The formation of Zn-TPY-TTF CPG was confirmed by XRD test.

Preparation of Pt@Zn-TPY-TTF CPG. For in situ platinum nanoparticles (Pt NPs) stabilization, 1 ml of H₂PtCl₆·6H₂O was dispersed in 40 ml of water containing 10 mg of Zn-TPY-TTF CPG. After continuous stirring for 1 h in a closed system, the well-dispersed solution was irradiated using 300 W xenon lamp (Newport) with a 6.0 cm long IR water filter for 2 h. Finally, the Pt nanoparticle stabilized CPG sample (Pt@Zn-TPY-TTF) was thoroughly washed with deionized (DI) water and dried under vacuum at 80 °C for 12 h. More detailed characterization of Pt@Zn-TPY-TTF CPG is provided in Supplementary Information.

Characterization

General. 1H-NMR spectra were recorded on a Bruker AVANCE-400 NMR spectrometer (at 400 MHz) and JEOL-ECZR NMR spectrometer (at 600 MHz) with CDCl₃ as the solvent and all spectra were referenced against TMS. 13C-spectrum was recorded at 150 MHz frequency using a Varian Inova 600 MHz NMR spectrometer. UV-Vis spectra were recorded in a Perkin-Elmer Lambda 900 spectrometer. For the UV-vis absorption studies, TPY-TTF OG and Zn-TPY-TTF CPG and Pt@Zn-TPY-TTF CPG were coated on a quartz plate as a thin film. Time-resolved photoluminescence (TRPL) and photoluminescence (PL) studies were performed on an Edinburgh instrument (FLS 1000). Fourier transform infrared spectra (FT-IR) were recorded by making KBr pellets using Bruker IFS 66 v/S Spectrophotometer in the region 4000–400 cm⁻¹. Thermal stability of the materials was measured using Mettler Toledo TGA 850 instrument in the temperature range of 30–800 °C with a heating rate of 5 °C min⁻¹ in N₂ atmosphere. Powder X-ray diffraction (PXRD) patterns were measured by a Bruker D8 Discover instrument using Cu Ka radiation. Atomic force microscopy (AFM) measurements were carried out with a Nanoscope model Multimode 8 Scanning Probe Microscope to analyze the morphologies of the sample surface. For this analysis, samples were deposited on silicon wafer followed by gold (Au) sputtering for FE-SEM measurements. Transmission Electron Microscopy (TEM) studies were done on JOEL JEM - 3010 with an accelerating voltage of 300 kV. For this study, the xerogels were dispersed in ethanol and then drop-cast onto a small piece of silicon wafer followed by gold (Au) sputtering for FE-SEM measurements.

Rheology. The rheological study was done in Anton Paar Rheometer MCR 302. Rheological measurements were performed in a 25 mm cone-and-plate configuration with a 0.5° cone angle. The rheology experiment was performed using the amplitude sweep method over strain % at a constant frequency (ω = 1 Hz). For each rheology measurement, the gel was prepared in 10 ml glass vial. Next, ~20 mg of gel sample was loaded onto the rheometer plate with the help of a spatula in a single shot to avoid any damage to the loaded sample. Further, data accuracy was ensured by repeating these experiments a minimum of three times.

Critical point drying. Toussain Automods®931 was used for critical-point drying (CPD) of the gel samples. After gel preparation, solvents present in CPG were exchanged with ethanol using a gradient of ethanol/water mixtures (40–100 %). Next, the ethanol exchanged gel samples were then transferred to a stainless-steel cage with wire mesh followed by critically point dried with supercritical CO₂.

Transient absorption. Femtosecond transient absorption (TA) experiments were performed using an amplified Ti:sapphire laser system (800 nm, 60 fs, 1 kHz, 3 mJ) from Amplitude Technologies, France, and a pump-probe set-up (Exciproc) from CDP Corporation, Russia. A fraction of the fundamental laser beam was used to 80 °C under vacuum for 8 h. Selected FT-IR (KBr, cm⁻¹) for xerogel state of TPY-TTF OG: 3392 (b), 3051 (s), 2945 (w), 1649 (s), 1581 (s), 1460 (s), 1362 (w), 1294–1096 (w), 976 (w), 839 (s), 786 (s), 619 (b).
generate 400 nm pump pulse in a 0.2 mm thick BBO crystal. Another small fraction of 800 nm pulse was used to generate broad continuum pulse (350–750 nm) in a rotating calcium fluoride window. The time delay between the pump and probe pulses was maintained by using an optical delay stage in the probe path. To minimize the noise in the transient signal, a part of the probe beam, known as reference pulse, was passed through the unexcited region of the sample and detected simultaneously with the probe pulse using a dual diode array-based spectrometer. The reference pulse was focussed in the same geometry with the focussed pump pulse. To obtained the pump induced changes in the absorbance of the sample (AA), the alternate pump pulses were blocked with the help of a synchronized chopper. Each transient spectra were collected after averaging 1000 Hz, from each sample. Each sample was dispersed in methanol and taken in a rotating cell to avoid their photodecomposition. The instrument response function of the TA set-up was 120 fs.

Photocatalytic experiments
Experiment set-up for photocatalytic water reduction. Photocatalytic H₂ evolution experiments were carried out in an 80 mL self-designed borosilicate glass cell containing a magnetic stir bar sealed with a small septum (picture of photocatalytic cell is given in Supplementary Fig. 23). For photocatalytic experiment, 1 mg catalyst was dispersed in 38 ml water (500 µl) and water (500 µl) was drop cast over the GC electrode and allowed to dry for 3 h under ambient condition. EIS was recorded at −1.2 V vs. Ag/AgCl applied bias from 0.1 Hz to 100 kHz (under the dark condition and visible-light irradiation).

for production. 1.0 mg of TPY-TTF OG and Zn-TPY-TTF CPG and Pt@Zn-TPY-TTF CPG was depicted by the Mott–Schottky (MS) analysis (at 1000 Hz, from −2.0 to +2.0 V) using ITO as a working electrode (WE) in N₂-purged aqueous solution of 0.5 M Na₂SO₄ at pH = 7, Pt as a counter electrode (CE) and Ag/AgCl as a reference electrode (RE). An electrochemical ink was prepared by making a dispersion of a mixture of catalyst (2.0 mg) in the solvent mixture of isopropanol (500 µl), water (500 µl), and Naflon (14 µl). Upon sonication for 20 min, a well-dispersed ink (3.5 µl) was drop cast over the ITO electrode and allowed to dry for 3 h under ambient condition. TON was calculated for the pump induced changes in the probe path. To obtained the pump induced changes in the probe path. To obtained the pump induced changes in the probe path. To obtained the pump induced changes in the probe path. To obtained the pump induced changes in the probe path.

**Lifetime calculations**. Lifetime data for Zn-TPY-TTF CPG and Pt@Zn-TPY-TTF CPG were collected upon exciting at 510 nm. The average lifetime is calculated using the below formula.

\[
\tau_{avg} = \frac{2}{\Delta \lambda} \frac{\tau_1}{\tau_2}
\]

Where, \(\tau_{avg}\) = average lifetime in nano-seconds, \(\Delta \lambda = \) sum of percentage of all the components that exist in the excited state, \(\Sigma \tau_{i} = \) sum of excited-state lifetime of all components. The details are provided in Supplementary Table 4.

**Turnover Number**. The turnover number (TON) is calculated by the below formula.

\[
TON = \frac{\text{Amount of product evolved (µmol)/amount of catalyst (µmol)}}{n}
\]

For hydrogen production. 1.0 mg of TPY-TTF OG is equivalent to 0.545 µmol and the amount of H₂ evolved was found to be 2 µmol from 1 mg of TPY-TTF in 20 h. Therefore, TON was calculated for TPY-TTF OG in 20 h. The unit of Zn-TPY-TTF CPG was calculated based on a binding ratio of TPY-TTF LMWG with Zn(NO₃)₂ in CPG (i.e. 1:2). Therefore, unit formula weight of Zn-TPY-TTF CPG is 2212.94. Therefore, 1 mg Zn-TPY-TTF CPG catalyst contains 0.451 µmol and the corresponding TON was found to be 23.5 in 20 h. Similarly, TON was also calculated for CO formation in presence of Zn-TPY-TTF CPG photocatalyst. The detailed TON calculation for H₂ production and CH₄ formation in presence of Pt@Zn-TPY-TTF CPG are provided in Supplementary Table 3.

**Apparent quantum efficiency.** The apparent quantum efficiency (AQE) is defined by the ratio of the effective electron used for product formation to the total photon flux. The AQE is calculated by using the below formula.

\[
AQE(\%) = \left( \frac{\text{Effective electrons}}{\text{Total photons}} \right) \times 100 = \left( \frac{n \times X \times Y}{S \times Y + S} \right) \times 100
\]

Where, \(n\) is the number of electrons used in the photocatalysis process, \(X\) is the yield of evolved gas from the sample (mol), \(Y\) is the Avogadro’s number (6.022 × 10²³ mol⁻¹), \(S\) is the photon flux, \(T\) is the irradiation time, and \(S\) is the illumination area. The photon flux was calculated at 400 ± 10 nm, 450 ± 10 nm, 500 ± 10 nm, 550 ± 10 nm, 600 ± 10 nm, 650 ± 10 nm, and 700 ± 10 nm by using separate bandpass filters with the help of power meter. The power meter (model: LaserCheck: 0623G19R) used for the experiment was purchased from Coherent. Since the reduction of a water molecule to the H₂ molecule requires two-electron (n = 2). The following calculation is based on data from water photoreduction with Zn-TPY-TTF CPG for 1 h, AQE was calculated as 0.76% at 550 ± 10 nm (the amount of H₂ evolved in 1 h was found to be 0.094 µmol). Meanwhile, the AQE of H₂ for the Pt@Zn-TPY-TTF CPG was also calculated at the various wavelength (400–700 nm), and the maximum was found at 550 ± 10 nm, which was calculated to be 14.47%. The detailed calculation for H₂ production in presence of Pt@Zn-TPY-TTF CPG at different wavelength is provided in Supplementary Table 5.

The reduction of a CO₂ molecule to the CO molecule requires two-electron (n = 2). The following calculation is based on data from CO₂ photoreduction with Zn-TPY-TTF CPG at 550 ± 10 nm, AQE was calculated as 0.96% (the amount of CO evolved was found to be 0.121 µmol in 1 h). Similarly, the reduction of a CO₂ molecule to the CH₄ molecule requires eight-electron (n = 8). Therefore, the AQE for CH₄ formation by Pt@Zn-TPY-TTF CPG was calculated at various wavelengths (400–700 nm) and the maximum efficiency at 550 ± 10 nm as 0.81%. The detailed calculation for CO₂ photoreduction to CH₄ in presence of Pt@Zn-TPY-TTF CPG at different wavelength is provided in Supplementary Table 9.
Data availability
All other data supporting the findings are available in the article as well as the supplementary information files and can be found from the authors on reasonable request.

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**Author contributions**

P.V. and T.K.M. designed research. P.V. performed the experimental works and analysed data. P.V., A.S. and T.K.M. contributed to analysing experimental data and wrote the manuscript. F.A.R. and S.K.P. carried out the computational studies. S.N. performed the transient absorption studies. All the authors discussed the results and commented on the manuscript.

**Competing interests**

The authors declare no competing interests.

**Additional information**

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