Structure-Activity Relationships in Pt-Functionalized Graphitic Carbon Nitride Photocatalysts

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Graphitic carbon nitrides (g-CN$_3$H$_y$s) offer immense potential as inexpensive photocatalysts for solar fuel generation (e.g., H$_2$ from water) owing to their facile synthesis from a range of precursors, ability to absorb visible light, and high surface area [1]. Typically, g-CN$_3$H$_y$s are functionalized with Pt via photodeposition at high loadings (e.g., ~1-5 wt%) but single-atom Pt cocatalyst can also be used for H$_2$ evolution at much lower loadings [2]. Recent experimental and theoretical work has highlighted the importance of native N-H$_2$ defects in facilitating interfacial charge transfer to supported Pt cocatalyst [3]. Nanoscale characterization should reveal more precise structure-function relationships leading to rational design strategies for Pt/g-CN$_3$H$_y$s photocatalysts that maximize energy conversion efficiency and minimize Pt consumption.

A systematic study correlating the H$_2$ evolution rates (HERs), bulk characterization of the g-CN$_3$H$_y$s supports, and active site densities determined from annular dark field scanning transmission electron microscopy (ADF-STEM) was performed to understand photocatalytic activity in Pt/g-CN$_3$H$_y$s at multiple length scales. Three g-CN$_3$H$_y$s demonstrating a range in structural condensation (based on X-ray diffraction) were selected including a commercially-produced “Nicanite” and two urea-derived g-CN$_3$H$_y$s. Nicanite is the most structurally condensed with the smallest interplanar spacing. The urea-based g-CN$_3$H$_y$s were synthesized by calcining urea at 550°C for 30 and 240 minutes to yield U$_{30}$-gCN and U$_{240}$-gCN, respectively, which show more narrowed layer stacking distances with calcination time. HERs were determined from gas chromatography of the photoreactor headspace under visible light irradiation (λ>400 nm) and in 10 vol% aqueous triethanolamine. Pt was loaded onto each g-CN$_3$H$_y$s via photodeposition (PD) from PtCl$_4$ at a loading 1.6 wt%, yielding active sites for H$_2$ evolution.

When each support is loaded with Pt via PD, the HER of U$_{240}$-gCN is >2x higher than that of Nicanite or U$_{30}$-gCN (Figure 1). ADF-STEM was performed to understand photocatalytic activity in Pt/g-CN$_3$H$_y$s at multiple length scales. Three g-CN$_3$H$_y$s demonstrating a range in structural condensation (based on X-ray diffraction) were selected including a commercially-produced “Nicanite” and two urea-derived g-CN$_3$H$_y$s. Nicanite is the most structurally condensed with the smallest interplanar spacing. The urea-based g-CN$_3$H$_y$s were synthesized by calcining urea at 550°C for 30 and 240 minutes to yield U$_{30}$-gCN and U$_{240}$-gCN, respectively, which show more narrowed layer stacking distances with calcination time. HERs were determined from gas chromatography of the photoreactor headspace under visible light irradiation (λ>400 nm) and in 10 vol% aqueous triethanolamine. Pt was loaded onto each g-CN$_3$H$_y$s via photodeposition (PD) from PtCl$_4$ at a loading 1.6 wt%, yielding active sites for H$_2$ evolution.

When each support is loaded with Pt via PD, the HER of U$_{240}$-gCN is >2x higher than that of Nicanite or U$_{30}$-gCN (Figure 1). ADF-STEM imaging of the used Pt/g-CN$_3$H$_y$s (Figure 2a-c, e) reveals that the Pt particle size distribution varies widely with the support structure and U$_{240}$-gCN facilitates the most dispersed cocatalyst species. By normalizing the HER by the total number of active sites (i.e., the number of surface-exposed Pt atoms) estimated from the particle size distributions, different g-CN$_3$H$_y$s’ turn over frequency (TOF) can be compared. The TOF describes the rate of H$_2$ produced per active site and unlike the HER, is not influenced by ensemble effects. This analysis reveals the following trend in TOF: Nicanite (213 h$^{-1}$) > U$_{240}$-gCN (146 h$^{-1}$) > U$_{30}$-gCN (103 h$^{-1}$). Alternatively, predominately single-atom Pt was deposited onto Nicanite following a chemical reduction (CR) approach (Figure 2d-e), achieving a low loading of 0.15 wt% and peak HER of 880 µmol/h/g (Figure 1) [3]. The TOF of this photocatalyst, at 196 h$^{-1}$, is similar to Nicanite prepared with PD at high loading. Ultimately, the stacking distance of the light-harvesting support was found to strongly correlate with the TOF, which we speculate originates from higher mobilities in more structurally condensed g-CN$_3$H$_y$s meaning more photogenerated electrons may reach the surface for performing water reduction [4]. In Nicanite, the low active site densities resulting from both PD and CR appear to limit the overall HER suggesting this factor is critical to achieving high energy conversion efficiencies. Methods for evaluating photocatalytic
activity in complex systems like Pt/g-CN₃H₅s will also be emphasized.

References:
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**Figure 1.** H₂ evolution rates (HER), per mass of photocatalyst, of different Pt/g-CN₃H₅s. Pt was loaded onto the supports either through photodeposition (PD) or chemical reduction (CR).

**Figure 2.** ADF-STEM images of photocatalysts prepared via PD at a loading of 1.6 wt%: (a) Nicanite, (b) U₂₄₀-gCN, and (c) U₃₀-gCN. (d) ADF-STEM image of Nicanite loaded with 0.15 wt% Pt using CR. (e) Pt features size distributions and violin plots of each photocatalyst used to estimate active site densities; inset shows a more detailed histogram of Pt on Nicanite prepared with CR.