Structure–property relationships for covalent triazine-based frameworks: The effect of spacer length on photocatalytic hydrogen evolution from water

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1. Introduction

Until recently, direct photocatalytic hydrogen production using particulate photocatalysts has been dominated by inorganic materials [1–3], but there is now a growing interest in organic and polymeric photocatalysts [4]. Graphitic carbon nitride (g-C3N4) is the most extensively studied material in this class for hydrogen evolution from water [5–10] and it has been reported to drive overall water splitting via a four-electron pathway [11,12]. One of the drawbacks of graphitic carbon nitride is the limited scope for fine-tuning the structure and properties as the synthesis is carried out at temperatures above 520 °C [9]. Materials are typically modified by varying the nitrogen rich starting materials [13,14] or by doping methods; for example, by adding sulfur [15] or potassium hydroxide [16]. This difficulty in tailoring the photophysical properties of g-C3N4 makes it attractive to study polymeric organic photocatalysts, which can be synthesized by coupling a diverse range of monomers at low temperatures, thus allowing fine synthetic control over the primary structure. Poly(p-phenylene)s [17,18], oligo(p-phenylene)s [19] and poly(pyridine-2,5-diyi) [20] show limited catalytic activity for hydrogen evolution from water in the presence of a sacrificial hole-scavenger, mostly under UV light. More recently, conjugated microporous polymer (CMP) photocatalysts synthesized by Suzuki-Miyaura polycondensation of substituted pyrenes and benzenes [21,22] have shown promising photocatalytic activities, and the absorption spectrum can be shifted into the visible region by varying the co-monomer composition. A further study on CMPs [23] explored the influence of substitution patterns and conjugation lengths on the hydrogen evolution rates. CMPs based on 4,8-di(thiophen-2-yl)benzo[1,2-b:4,5-b']dithiophene [24,25] have also shown good activity. Besides these CMPs, linear conjugated copolymers of planar units were shown to have even higher activities for photocatalytic hydrogen evolution in the presence of triethylamine as sacrificial hole-scavenger [26]. Subsequently, poly(phenyleno-benzothiadiazole)s [27] and poly(fluorene-co-benzothiadiazole) [28] were both shown to promote hydrogen evolution from water in the presence of sacrificial hole-scavengers.

Covalent triazine-based frameworks (CTFs) have also been studied for photocatalytic hydrogen evolution from water. CTFs are
similar to g-C3N4 in terms of their high nitrogen content and their synthesis which often, but not always, involves ionothermal approaches in molten salt or molten eutectic salt mixtures with temperatures over 350 °C [29–32]. CTFs for photocatalysis were prepared previously by condensation of 1,3,5-tris(4-formyl-phenyl) triazine and 2,5-diethoxy-terephthalohydrazide to form hydrazones at 120 °C via microwave synthesis [33] and also obtained by a thermal azine formation form 4,4’,4”-terazine-2,4,6-triyl[(benzaldehyde) and hydrazine at 120 °C [34]. Other CTF photocatalysts were synthesized by acid catalyzed, low temperature trimerizations of nitride precursors, such as tetra(4-cyanophenyl) ethylene (25 °C) [35], terephthalonitrile and methylterephthalonitrile (0 °C) [36].

In this study, we prepared a range of structurally related CTFs using low temperature synthesis routes and explored the effect of structure and synthesis conditions on photocatalytic hydrogen evolution under sacrificial conditions.

2. Methods

All reagents were obtained from Sigma-Aldrich or from Alfa Aesar and used as received, except for [1,1’:4’,1”-terphenyl]-4,4”-dicarbonitrile (M3) and [1,1’:4’,1”-terphenyl]-4,4”-dicarbonitrile (M4), which were synthesized according to literature procedures [37,38]. 2,4,6-Tris[4-bromomethyl]-1,3,5-triazine (M5) was synthesized via acid catalyzed trimerization and 2,4,6-tris-[4-(4,4,5,5-tetramethyl-1,3,2-dioxaborolan-2-yl)phenyl]-1,3,5-triazine (M6) was synthesized using a modified Miyaura-boration protocol [21]. Water for the hydrogen evolution experiments was purified using an ELGA LabWater system with a Purelab Option S filtration and ion exchange column (p = 15 MΩ cm) without pH level adjustment. Reactions were carried out under nitrogen atmosphere using standard Schlenk techniques. CHN Analysis was performed on a Thermo EA1112 Flash CHNS-O Analyzer using standard microanalytical procedures. The UV-visible absorption spectra of the polymers were recorded on a Shimadzu UV-2550 UV-Vis spectrometer as powders in the solid state. The ball milled polymer powders were prepared by grinding the polymer samples in a Retsch 400 MM mixer mill at a frequency of 30 Hz for 30 min. The mixer mill was equipped with 10 mL stainless steel grinding jars that contained two stainless steel grinding balls with a diameter of 7 mm. (TD-)DFT calculations were performed using the Turbomole 6.6 code [39], the B3LYP [40,41] density functional and the COSMO solvation model [42], for more details see the supporting information.

2.1. Hydrogen evolution experiments

A flask was charged with the polymer powder (25 mg), water (20.0 mL), triethanolamine (5.0 mL), H2PtCl6 solution (0.02 mL, 8 wt % in H2O) or with the polymer powder (25 mg), water (20.0 mL), triethanolamine (2.5 mL), methanol (2.5 mL), H2PtCl6 solution (0.02 mL, 8 wt % in H2O) and sealed with a septum. The resulting suspension was ultra-sonicated for 10 min until the photocatalyst was well dispersed before degassing by N2 bubbling for 30 min. The reaction mixture was side-illuminated with a 300 W Newport Xe light-source (Model: 6258, Ozone free) for the time specified. The lamp was cooled and IR light cut out by water circulating through a metal jacket with quartz windows. Gas samples were analyzed with a Bruker gas chromatograph.

2.2. General procedure for the acid catalyzed trimerization polymerization

Trifluoromethanesulfonic acid and CHCl3 were added to a flask under nitrogen and the monomer dissolved in chloroform was added with a syringe pump at a rate of 2 ml min⁻¹. The reaction was stirred for 48 h before it was poured onto an aqueous ammonia solution (28.0–30.0% NH3) to neutralize the trifluoromethanesulfonic acid. The suspension was left stirring for 1 h after the addition of the ammonia solution. The resulting solid was filtered, washed successively with dichloromethane, ethanol, water and methanol. The samples were dried under vacuum at 75 °C overnight to obtain the polymer.

2.2.1. CTF-1

Trifluoromethanesulfonic acid (2.83 mL, 32.0 mmol), CHCl3 (5.00 mL) and 1,4-dicyanobenzene (513 mg, 4.40 mmol) in 40 mL CHCl3 were used in the polymerization. After the addition of the monomer, the solution was heated to 50 °C for 1 day, cooled to room temperature and continued with the general procedure. After work-up, the product was obtained as a white powder (328 mg) in 63% yield. Calc. for (C9H2N)n: C, 74.99; H, 3.15; N, 21.86%; Found: C, 57.71; H, 3.72; N, 15.88%.

2.2.2. CTF-2

Trifluoromethanesulfonic acid (2.83 mL, 32.0 mmol), CHCl3 (5.00 mL) and 4,4’-biphenyldicarbonitrile (817 mg, 4.00 mmol) in 40 mL CHCl3 were used in the polymerization. After work-up, the product was obtained as a pale yellow powder (600 mg) in 73% yield. Calc. for (C13H14N)n: C, 82.33; H, 3.95; N, 13.72%; Found: C, 75.88; H, 3.81; N, 12.21%; For the ball milled sample C, 75.62; H, 3.81; N, 12.17%.

2.2.3. CTF-3

Trifluoromethanesulfonic acid (0.40 mL, 4.48 mmol), CHCl3 (0.10 mL) and [1,1’:4’,1”-terphenyl]-4,4”-dicarbonitrile (157 mg, 560 μmol) in 0.56 mL CHCl3 were used in the polymerization. After work-up, the product was obtained as a pale green powder (110 mg) in 70% yield. Calc. for (C18H14N)n: C, 85.69; H, 4.31; N, 9.99%; Found: C, 81.54; H, 4.57; N, 8.90%.

2.2.4. CTF-4

Trifluoromethanesulfonic acid (0.40 mL, 4.48 mmol), CHCl3 (0.10 mL) and [1,1’:4’,1”-terphenyl]-4,4”-dicarbonitrile (200 mg, 560 μmol) in 0.56 mL CHCl3 were used in the polymerization. After work-up, the product was obtained as a pale green powder (164 mg) in 82% yield. Calc. for (C19H12N)n: C, 87.62; H, 4.52; N, 7.86%; Found: C, 82.39; H, 4.47; N, 7.55%.

2.3. General procedure for the Pd(0)-catalyzed Suzuki-Miyaura polycondensation

The flask was charged with the monomers, N,N-dimethylformamide, an aqeous solution of K2CO3 (2.0 M) and degassed by bubbling with N2 for 30 min [Pd[PPh3]4] (1.2 mol %) was added and the solution was degassed by bubbling with N2 for 10 min before heated to 150 °C for 48 h. The mixture was cooled to room temperature and poured into water. The precipitate was collected by filtration and washed with H2O and methanol. Further purification of the polymer was carried out by Soxhlet extraction with cyclohexyl methyl ether for two days and the product was dried under reduced pressure at 125 °C.
After work-up, the product was obtained as a pale yellow powder (440 mg) in 98% yield. Calc. for (C_{10}H_{14}N)n: C, 82.33; H, 3.95; N, 13.72%; Found: C, 69.26; H, 4.84; N, 9.00%.

2.3.2. CTF-3 Suzuki
1,4-Benzene diboronic acid (182 mg, 1.10 mmol), 2,4,6-tris(4-bromophenyl)-1,3,5-triazine (400 mg, 733 μmol) N,N-dimethyformamide (16.5 mL), aqueous K2CO3 (2.0 M, 3.3 mL) and [Pd(PPh3)4] (15 mg, 1.2 mol %) were used in the polymerization. After work-up, the product was obtained as a pale green powder (225 mg) in 73% yield. Calc. for (C_{10}H_{14}N)n: C, 85.69; H, 4.31; N, 6.12%; Found: C, 79.60; H, 4.49; N, 8.97%.

2.3.3. CTF-4 Suzuki
4,4′-Bisphenylidiboronic acid bis(pinacol) ester (71.7 mg, 177 μmol), 2,4,6-tris(4-bromophenyl)-1,3,5-triazine (64.3 mg, 118 μmol) N,N-dimethyformamide (2.65 mL), aqueous K2CO3 (2.0 M, 0.53 mL) and [Pd(PPh3)4] (2.40 mg, 1.2 mol %) were used in the polymerization. After work-up, the product was obtained as a pale green powder (44.9 mg) in 71% yield. Calc. for (C_{10}H_{14}N)n: C, 87.62; H, 4.52; N, 7.86%; Found: C, 67.89; H, 3.89; N, 6.12%.

3. Results and discussion
We present here a series of CTFs and compare the influence of the phenylene spacer length and the synthesis methods on the photocatalytic hydrogen evolution rate. We further compare the measured properties of these materials with theoretical (TD-)DFT predictions.

Two series of CTFs were synthesized (Fig. 1). CTF-1 to CTF-4 were obtained by acid catalysis and CTF-2 Suzuki to CTF-4 Suzuki were made using palladium chemistry. The numerals in these names refer to the length of the para-phenylene spacers between the triazines in the material (i.e., CTF-1 = phenyl, CTF-2 = biphenyl, etc.). CTF-1 to CTF-4 were synthesized via trifluoromethanesulfonic acid (TfOH) catalyzed cyclotrimerization of the corresponding dicyano arenes [43]. The reaction proceeded well at ambient temperature for 4,4′-dicyano-diphenylether (M2), [1′,1′′-terphenyl]-4,4′-dicarbonitrile (M3) and [1′′,1′′′,1′′′′]-quaterphenyl]-4,4′′′-dicarbonitrile (M4). However, in the case of 4,4′-dicyano benzene (M1) [36], it was necessary to increase the reaction temperature to 50 °C.

CTF-2 Suzuki to CTF-4 Suzuki were prepared via palladium(0)-catalyzed Suzuki-Miyaura type polycoupling of 2,4,6-tris(4-bromophenyl)-1,3,5-triazine (M5) and 2,4,6-tris-[4-(4,4,5,5-tetramethyl-1,3,2-dioxaborolan-2-yl)phenyl]-1,3,5-triazine (M6), 1,4-benzenediboronic acid (M7), or 4,4′-diphenyldiboronic acid bis(pinacol) ester (M8) in N,N-dimethyformamide (DMF) at 150 °C in the presence of aqueous K2CO3. For both sets of materials, the polymers were obtained as powders and ball milled prior to analysis and photocatalysis experiments.

All CTFs were insoluble in common organic solvents tested (i.e., acetone, methanol, hexane, dichloromethane, toluene, tetrahydrofuran, and N,N-dimethyformamide). Fourier-transform infrared spectroscopy (FT-IR) shows a significant decrease in intensity for the aryl nitrite band around 2230–2220 cm⁻¹ compared to the monomer for the TfOH catalyzed series (CTF-1 to CTF-4), qualitatively indicating a high degree of polymerization (Fig. S-1, ESI). The FT-IR spectra of the Suzuki-coupled polymers show similar features compared to those prepared by TfOH catalysis, except for the expected absence of a nitrite band. Even though no distinct features in the FT-IR can be assigned to hydrolysis products, partial hydrolysis of unreacted nitrite groups at the end of the polymerization or during work-up in polymers CTF-1, CTF-2, and CTF-4 is conceivable. All polymers are porous to nitrogen at 77 K after ball milling, except CTF-1 and CTF-4. CTF-2 shows the highest apparent Brunauer-Emmett-Teller surface area (SA_BET) of 560 m² g⁻¹, while all others have SA_BET in the range from 200 to 400 m² g⁻¹. Polymers CTF-1 and CTF-2 were reported previously to have similar FT-IR spectra and surface areas [43]. The polymers also show moderate gas uptakes for H₂, CO₂, Kr, and Xe (Table S-1, ESI), with CTF-2 showing the highest uptakes for H₂ (5.35 mmol g⁻¹ at 77 K, 1 bar), CO₂ (1.25 mmol g⁻¹ at 298 K, 1 bar), Kr (0.45 mmol g⁻¹ at 298 K, 0.9 bar), and Xe 1.73 mmol g⁻¹ (298 K, 1 bar). Powder X-ray diffraction patterns were featureless, indicating that both synthesis routes result in amorphous materials without any long-range order (Fig. S-7, ESI). This is probably due to the irreversible nature of the low temperature synthesis routes employed, since dynamic bond formation is believed to be necessary to give crystalline materials [44,45]. By contrast, limited crystallinity in CTFs has been reported for materials synthesized using ionothermal [29,30] and microwave routes [43]. Residual palladium was determined via ICP-OES and found to be 0.27 wt % for CTF-2 Suzuki, 0.07 wt % for CTF-3 Suzuki, and 0.65 wt % for CTF-4 Suzuki (Fig. S-28 and Table S-3, ESI). Energy-dispersive X-ray spectroscopy also confirms that residual palladium is present in the materials synthesized via Suzuki-Miyaura coupling (Fig. S-4, ESI). Thermogravimetric analysis (TGA) shows that the materials exhibit thermal stability up to 245 °C, except for CTF-1 which has a decomposition onset of 265 °C, possibly indicating a lower degree of polycrystallinity. Comparing both series of materials, the materials synthesized via TfOH tri- merization show an earlier decomposition on-set compared to the polymers made by Suzuki coupling. This could be due to lower molecular weight of the polymers, which might also affect the photo-physical properties (infra vide). In contrast to previous reports [36,43], we found that when CTF-1 was synthesized at room temperature, the onset of decomposition was even lower (245 °C; Fig. S-3, ESI) and a photocatalytically inactive material was obtained. The samples were isolated after work-up as macroscopic particles (CTF-1, CTF-2 Suzuki, CTF-3 Suzuki, CTF-4 and CTF-4 Suzuki) or as macroscopic flakes (CTF-2 and CTF-3). After ball milling, SEM images of all materials (Fig. S-30–S-33, ESI) were obtained to estimate the decrease in particle size. The particle size of CTF-1 could be reduced from around 100 μm to a few hundred micrometers. By contrast, the particle size of CTF-2, CTF-3, CTF-2 Suzuki and CTF-3 Suzuki were decreased from several millimeters to smaller particles in the range of a few hundred micrometers. By contrast, the particle size of CTF-4 and CTF-4 Suzuki was not decreased after ball milling, probably due to the softness and lower density of the networks.

UV-visible spectra (Fig. 2) show that the optical gap of the polymer depends on the length of the 1,4-phenylene linker between the triazine cores. Within the series CTF-1–CTF-4, the optical gap decreases from 2.95 eV to 2.48 eV (Table 1). A similar trend is observed for series CTF-2 Suzuki to CTF-4 Suzuki, but these materials fall within a much narrower range (2.93 eV–2.85 eV). As such, the synthesis route seems to have an important impact on the photophysical properties of the materials, possibly due to differences within the material and end-groups. In both series, the material with four-phenylene spacers (CTF-4 and CTF-4 Suzuki) is the most red-shifted and has the smallest optical gap (see Fig. 3).

All materials are weakly fluorescent upon excitation at 280 nm and show two characteristic maxima (Fig. S-5 and Fig. S-6, ESI). The maxima fall between 367 and 374 nm for the first peak and 448 and 476 nm for the second peak. No coherent trends can be found within either series of polymers, and the observed Stokes’ shifts are small, possibly an indication of rigidity.

We tested all materials for photocatalytic hydrogen evolution from water in the presence of triethanolamine (TEOA, 20 vol %) acting as a sacrificial hole-scavenger [32,34,36]. For improved performance, the materials were loaded with 3 wt % platinum as
the co-catalyst by \textit{in situ} photodeposition of H$_2$PtCl$_6$ \cite{24,36,46}. For CTF-2, Pt particles with diameters in the range of 17–33 \(\mu\)m were observed (Fig. S-34, ESI). All materials, with the exception of CTF-4 and CTF-4 Suzuki, were found to produce hydrogen under visible-light irradiation (>420 nm). The hydrogen evolution rates of all as-synthesized CTF samples were low, as the polymer powders were macroscopic flakes that were only poorly dispersible in the aqueous mixture (Fig. S-35, ESI). These issues were solved by ball milling the samples and we obtained hydrogen evolution rates of 35 \(\mu\)mol g\(^{-1}\) h\(^{-1}\) for CTF-1, 296 \(\mu\)mol g\(^{-1}\) h\(^{-1}\) for CTF-2, and 45 \(\mu\)mol g\(^{-1}\) h\(^{-1}\) for CTF-3 under >420 nm irradiation. When the materials were synthesized via Pd(0)-cross coupling reaction, slightly lower photocatalytic activities for the analogous polymers CTF-2 Suzuki (265 \(\mu\)mol g\(^{-1}\) h\(^{-1}\)) and CTF-3 Suzuki (44 \(\mu\)mol g\(^{-1}\) h\(^{-1}\)) were found when tested in the presence of the platinum co-catalyst, although these differences are probably smaller than the reproducibility of these measurements. These CTF materials show lower, but still significant, activity even without added platinum co-catalyst; e.g. for CTF-2 Suzuki, the hydrogen evolution rate drops from 265 \(\mu\)mol g\(^{-1}\) h\(^{-1}\) with platinum co-catalyst to 121 \(\mu\)mol g\(^{-1}\) h\(^{-1}\) without added platinum (Fig. 19, ESI). As for organic solar cells \cite{47–49}, residual palladium is
found as particles in these polymers, which have been suggested to act as co-catalysts in a recent report on photocatalytically active CMPs [24]. Palladium was also reported to be active in conjunction with g-C3N4 [50], albeit with a lower activity than platinum, in line with the trends observed here. In contrast, CTF-2 does not produce hydrogen in absence of the Pt co-catalyst from water/triethanolamine solution (Fig. S-19, ESI). No hydrogen could be detected from suspensions of CTF-2 in water (Fig. S-20, ESI), CTF-2 water platinum particle suspensions (Fig. S-20, ESI), and from water/triethanolamine solutions in the absence of photocatalysts (Fig. S-21, ESI). The apparent quantum yield (AQY) [51] of CTF-2 was calculated to be 1.6% at 420 nm ($\pm$10 nm, FWHM, Fig. 4), which is higher than poly(p-phenylene) (AQY420 nm $\approx$ 0.4%) but lower than poly(dibenzo[b,d]thiophene sulfone-co-phenylene) (AQY420 nm $\approx$ 7.2%) [26]. No photocatalytic activity was found under 600 nm, 550 nm or 500 nm illumination using band-pass filters, indicating that the hydrogen evolution is indeed a photocatalytic process. Furthermore, photocurrents were measured for CTF-1, CTF-2, and CTF-4 under $>420$ nm illumination, these show that the process is indeed photocatalytic (Fig. S-29, ESI).

The photocatalytic stability of CTF-2 was also tested under 1 Sun illumination (using an ABA-certified solar-simulator as the light source) for a total of 38 h. The performance slightly decreased after three hours, but stabilized to a constant evolution rate of 526 $\mu$mol g$^{-1}$ C$^{-1}$ h$^{-1}$. After 38 h, the polymer evolved more hydrogen than present in the material, ruling out decomposition of the polymer as the source of hydrogen.

Calculations were performed using our established computational approach [52,53] based around density functional theory calculations on cluster models of the CTF materials (Fig. S-44, ESI). The results of these calculations are presented in Fig. 5 and suggest that the reaction involves the metal nanoparticles, which are formed in situ during the photocatalytic process.

### Table 1

| Polymer | Optical gap a/eV | $\lambda_{\text{max}}$ b/nm, eV | Predicted $\lambda_{\text{max}}$ b/eV | $\lambda_{\text{em}}$ b/nm | HER >420 nm c/$\mu$mol g$^{-1}$ h$^{-1}$ |
|---------|------------------|-------------------------------|-----------------------------------|-----------------|----------------------------------|
| CTF-1 | 2.95 | 354, 3.50 | 3.85 | 374, 469 | 35 ± 1 |
| CTF-2 | 2.73 | 398, 3.12 | 3.49 | 367, 448 | 296 ± 11 |
| CTF-3 | 2.62 | 404, 3.07 | 3.27 | 372, 468 | 45 ± 1 |
| CTF-4 | 2.48 | 382, 3.25 | 3.17 | 372, 476 | 0 ± 1 |
| CTF-2 Suzuki | 2.93 | 365, 3.40 | $\lambda_{\text{em}}$ | 373, 466 | 265 ± 5 |
| CTF-3 Suzuki | 2.88 | 377, 3.29 | $\lambda_{\text{em}}$ | 370, 452 | 44 ± 2 |
| CTF-4 Suzuki | 2.85 | 387, 3.20 | $\lambda_{\text{em}}$ | 371, 467 | 0 ± 1 |

a The optical gap was calculated from the on-set of absorption in the solid-state UV-vis spectrum.

b $\lambda_{\text{max}}$ was determined from the peak maximum of the most red-shifted peak in the solid state UV-vis spectrum.

c $\lambda_{\text{em}}$ was recorded at $\lambda_{\text{ex}}$ = 280 nm (Fig. S-5 and Fig. S-6, ESI).

d No hydrogen evolved during a typical 5 h run.

e The hydrogen evolution rate (HER) was measured with 25 mg of the ball milled samples in water, 20% triethanolamine and 3 wt % platinum as a co-catalyst.

f See CTF-2 to CTF-4 for predicted values.
that the thermodynamic driving force for proton reduction is similar for all materials (difference between EA; red line, and the proton reduction potential; blue line). However, the predicted driving force for the oxidation of the sacrificial hole-scavenger decreases within the series from CTF-1 to CTF-4. In Fig. 5 the oxidation of triethylenamine to diethylamine and acetaldehyde is shown (difference between IP; blue line, and the triethylenamine oxidation potential; grey line), but similar conclusions can be drawn for any sacrificial hole-scavenger as the oxidation potential of a sacrificial hole-scavenger is independent of the material studied. Results for the excited state EA* and IP* potentials are discussed in more detail in the supporting information.

In comparison to our measurements, this at first appears to be inconsistent with the fact that CTF-2 is the most active material, especially under illumination with a solar simulator. The photocatalytic activity of these materials is, however, the product of several properties, such as the driving force for both half-reactions and the number of photons absorbed, as well as the ease of charge transport [54]. We therefore investigated in more detail the spectral effects on the observed activity for CTF-1 and CTF-2.

Going from >420 nm irradiation to broad spectrum illumination with a Xe lamp and a >295 nm cut-off filter (Fig. S-24, ESI) leads to an increase in the hydrogen evolution rate from 35 μmol g⁻¹ h⁻¹ to 94 μmol g⁻¹ h⁻¹ for CTF-1 and from 296 μmol g⁻¹ h⁻¹ to 374 μmol g⁻¹ h⁻¹ for CTF-2 (Fig. S-11, ESI). This increased activity is caused by absorption of UV-photons, showing that CTF-1 is more active under >295 nm UV-light and has limited absorption of visible light photons. We also compared the activity of CTF-1 and CTF-2 when illuminated with exclusively UV light using a U-340 band-pass filter (260–390 nm, see Fig. S-24, ESI for transmission characteristics). Under these conditions, CTF-1 slightly outperforms CTF-2, evolving 118 μmol g⁻¹ h⁻¹ vs. 111 μmol g⁻¹ h⁻¹, respectively. Considering all these measurements and our calculations, it appears that the experimentally observed maximum in visible light activity in the series for CTF-2 is the result of a trade-off between the total amount of light absorbed and the thermodynamic driving force for the oxidation reaction. Along the series from CTF-1 to CTF-4, the amount of light absorbed increases as the optical gap decreases, while the driving force for the oxidation of a sacrificial hole-scavenger, i.e. triethylenamine or triethylenamine, decreases within the series.

Calculations also predict that CTF-4 should evolve hydrogen in the presence of triethylenamine, which is in contrast to the observed lack of activity in experiment when using triethylenamine as sacrificial hole-scavenger. As in previous studies [21,23,26], we therefore tested the materials in water/triethylenamine/methanol mixtures under >420 nm illumination. The addition of methanol has been found previously to act as a co-solvent, thus avoiding phase separation between water and triethylenamine [23,26], and also to improve surface wettability for hydrophobic polymers. For ball milled CTF-2, the hydrogen evolution rates under visible-light are slightly higher when using water/methanol/triethylenamine mixtures compared to water/triethylenamine mixtures (358 vs. 296 μmol g⁻¹ h⁻¹). For CTF-1 and CTF-3, the hydrogen evolution rates nearly double from 35 to 66 μmol g⁻¹ h⁻¹ and from 45 to 79 μmol g⁻¹ h⁻¹, respectively, with the water/triethylenamine/methanol system. More importantly, this mixture leads to a low hydrogen evolution rate of 24 μmol g⁻¹ h⁻¹ for CTF-4, in line with our prediction that this polymer should be able to evolve hydrogen. The fact that no visual improvement could be observed in polymer dispersibility in water/methanol/triethylenamine mixtures in comparison to water/triethylenamine mixtures suggests that the increased hydrogen evolution activity for CTFs might result from the easier oxidation of triethylenamine compared to triethylenamine.

4. Conclusions

The photocatalytic hydrogen evolution rates of seven CTFs using sacrificial hole-scavengers were studied and the results were compared with the predictions of (TD)-DFT calculations. The photocatalytic activity is not significantly different for materials synthesized using catalyzed trimerization or Suzuki-Miyaura type polycondensation, despite some differences in the optical properties. In both series of polymers, UV-vis spectra show a trend of decreasing optical gaps as the phenylene spacer length increases. CTF-2 showed the highest hydrogen evolution rate under visible light using triethylenamine as sacrificial hole-scavenger and this shows that microporosity does not preclude efficient photocatalytic water reduction. This is noteworthy, since in principle microporosity might reduce charge transport rates in the solid state, which could offset gains in mass transport. Under UV light, the activity of CTF-2 increased slightly, whereas CTF-1 evolved nearly twice the amount of hydrogen in comparison to the irradiation >420 nm. These results suggest that the absorption of visible photons is limited in CTF-1 and therefore decreases its hydrogen evolution rate under these conditions. This observation is supported by calculations predicting CTF-1 to have the largest thermodynamic driving force in the series from CTF-1 to CTF-4. In general, the trend in the activity of the CTF series and the origin of the maximum in hydrogen evolution rate for CTF-2 appears to be a combination of the thermodynamic driving force and the optical gap, highlighting how the activity of a material can be controlled by structural property tuning. Residual palladium in the Suzuki-Miyaura derived materials seems to act as a co-catalyst, however, the photocatalytic activity can be further improved by the addition of platinum. Finally, CTF-2 shows a stable hydrogen evolution rate under simulated solar light for at least 38 h, evolving more hydrogen than present in the material and showcasing the stability of the material.

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