Rational Design of Covalent Cobaloxime–Covalent Organic Framework Hybrids for Enhanced Photocatalytic Hydrogen Evolution

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ABSTRACT: Covalent organic frameworks (COFs) display a unique combination of chemical tunability, structural diversity, high porosity, nanoscale regularity, and thermal stability. Recent efforts are directed at using such frameworks as tunable scaffolds for chemical reactions. In particular, COFs have emerged as viable platforms for mimicking natural photosynthesis. However, there is an indisputable need for efficient, stable, and economical alternatives for the traditional platinum-based cocatalysts for light-driven hydrogen evolution. Here, we present azide-functionalized chloro(pyridine)cobaloxime hydrogen-evolution cocatalysts immobilized on a hydrazone-based COF-42 backbone that show improved and prolonged photocatalytic activity with respect to equivalent physisorbed systems. Advanced solid-state NMR and quantum-chemical methods allow us to elucidate details of the improved photoreactivity and the structural composition of the involved active site. We found that a genuine interaction between the COF backbone and the cobaloxime facilitates recoordination of the cocatalyst during the photoreaction, thereby improving the reactivity and hindering degradation of the catalyst. The excellent stability and prolonged reactivity make the herein reported cobaloxime-tethered COF materials promising hydrogen evolution catalysts for future solar fuel technologies.

INTRODUCTION

Identifying competitive alternatives to fossil-fuel-based energy constitutes one of the main research goals of this decade. Nature-inspired processes, like artificial photosynthesis, guide the way to a green and sustainable solution. Covalent organic frameworks (COFs) have been emerging as new materials in this context. COFs consist of light elements only, and their bottom-up synthesis enables high versatility and tunability on a molecular level, while benefiting from high stability and crystallinity due to covalent bonding in-plane and π−π-stacking out-of-plane. Most reports of COFs as photosensitizers for light-driven hydrogen evolution use platinum as a cocatalyst; hydrogen evolution rates up to 16.3 mmol h⁻¹ g⁻¹ have been reported in this context. Recent studies showed that the precious metal platinum can be replaced by earth-abundant molecular cocatalysts, namely, chloro(pyridine)cobaloxime and related complexes. These cocatalysts are well-known and well-defined, while offering high tunability, which facilitate their incorporation into photocative organic and inorganic systems. Cobaloximes feature low overpotential for the hydrogen evolution reaction and have been used in heterogeneous systems with metal–organic frameworks and carbon nitrides, as well as physisorbed to COFs. A major drawback of molecular proton reduction catalysts physisorbed to photosensitizers is their photodeactivation over time and rate limitations due to diffusion-controlled mechanisms. While previous attempts used molecular cobaloxime catalysts in solution, in this work we report photocatalytic hydrogen evolution with molecular cobaloxime catalysts covalently tethered to the COF backbone, yielding unprecedented insights into the nature of the active site and the COF–cocatalyst interface. By comparison with equivalent unbound, i.e., physisorbed, systems, we show how the modification of the hydrazone-based COF-42 and attachment of functionalized chloro(pyridine)cobaloxime lead to more efficient hydrogen evolution in a water/acetonitrile mixture under visible-light illumination in the presence of a sacrificial electron donor. The structural composition of the photoreaction is verified by computational and experimental methods including advanced high-resolution solid-state NMR techniques.

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Supporting Information

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techniques. These results combine the advantages of fully heterogeneous systems with the tunability of molecular cocatalysts and lead the way toward true single-site COF-based photocatalytic systems with a high level of interfacial control.

RESULTS AND DISCUSSION

In previous studies, COF-42 has been shown to be active in photocatalytic hydrogen evolution reactions with conventional hydrogen evolution cocatalysts such as platinum nanoparticles or molecular chloro(pyridine)cobaloxime. At the same time, this COF is a well-known and versatile platform that is chemically robust due to its hydrazone-linked structure. In this study, we used COF-42 as a platform for covalent postsynthetic modification with cobaloxime complexes. The synthesis of COF-42 by solvothermal acid-catalyzed condensation of 1,3,5-triformylbenzene (TFB) and 2,5-diethoxyterephthalohydrazide (DETH) followed published protocols. In order to provide functional sites for the covalent attachment of the cocatalyst, 10 mol % of DETH was replaced by the propargyl-containing 2,5-bis(prop-2-yn-1-xyloxy)terephthalohydrazide (DPTH) to obtain the propargyl-modified pCOF. The COFs were characterized by FT-IR spectroscopy, sorption analysis, powder X-ray diffraction (PXRD), magic-angle-spinning solid-state NMR (ssNMR), and quantum-chemical calculations.

The successful transformation of the starting materials to pCOF was proven by the lack of a residual aldehyde stretching vibration in its FT-IR spectrum. Characteristic C==O vibrations and signals originating from the hydrazone bonds overlap at 1680 cm⁻¹ [see Figure S13 of the Supporting Information (SI)]. New vibrations emerged at 2250 cm⁻¹ that could be assigned to the propargyl groups, confirming the successful incorporation of DPTH building blocks into the COF backbone. This was further supported by a 1D 13C{1H} ssNMR spectrum, where 13C signals at 79 and 58 ppm can be assigned to the propargyl functional group (Figure 1C). These shifts match the corresponding chemical shift of the liquid-state NMR of the DPTH linker (see the Supporting Information for experimental details) and are also confirmed by quantum-chemical calculations (see Table S3, SI).

PXRD analysis confirmed the crystalline structure of pCOF. The PXRD pattern shows a strong reflection at 3.3° 2θ, followed by smaller ones at 5.9°, 7.0°, and 9.1° and a very broad one at 26° 2θ. The experimental powder pattern was compared to a simulated one (see Figure 1E), and the diffraction peaks were assigned as the 100, 101, 200, 201, and 001 reflections, respectively. The peaks are broadened due to...
small domain sizes in the COF particles, especially in the z direction, where the interlayer interactions are defined by π−π-stacking only. Different possible orientations for the propargyl functionality as well as slightly shifted AA' stacking modes lead to very similar powder patterns; due to broadening of the reflections in the experimental data, the different orientations cannot be distinguished. One of these possible structural models is shown in Figure 1B, featuring an AA stacking mode with an interlayer distance of 3.5 Å, which is typical for structurally similar COFs.10,30,31 Note that in the underlying structural model, one out of six DETH linkers per pore was replaced by DPTH, which results in a functionalization degree of 16.6% instead of the statistically distributed 10% in the experimentally prepared pCOF10.

Pawley refinement of the structure in the idealized AA stacking mode suggests P2/m symmetry. For the modeled structure, the resulting cell parameters are a = 51.09 Å, b = 3.50 Å, c = 29.48 Å, α = γ = 90.00°, and β = 89.94°. Sorption analysis revealed a mesoporous structure with pore size of 2.3 nm and a Brunauer–Emmett–Teller (BET) surface area of 1839 m² g⁻¹, which matches the theoretically expected values of the structural model well (see Figure 1D).

For the covalent attachment of the cobaloxime catalyst to pCOF10, a postsynthetic click-chemistry approach was chosen. The copper(I)-catalyzed Huisgen-type cycloaddition of azines and alkynes is known to be broadly applicable with high yields and a high tolerance for functional groups.32–36 Therefore, the pyridine, which acts as an axial cobaloxime ligand, was functionalized with an azide group to yield the para-functionalized pyridine [Co-1a], which forms the azide-functionalized complex [Co-1a], and likewise, the meta-functionalized analogues [Co-1b] were synthesized, as depicted in Figure 2. Additionally, the equatorially functionalized chelating ligand 2 was synthesized as described in the Supporting Information. It forms the azide-functionalized catalyst [Co-2] by metal complexation as before. Two strategies were tested for the attachment of the cobaloxime complex to pCOF10: (i) metal complexation of azide-functionalized ligands with subsequent COF modification by click-reaction with the azide-functionalized complexes, termed route I, and (ii) COF modification by click-reaction with azide-functionalized ligands with subsequent complexation, termed route II (see the Supporting Information for experimental details). The resulting COF–cobaloxime hybrid samples are labeled as follows with the respective numbering according to Figure 2: [1a]−COF for clicked ligands and [Co-1a]−COF for COF–cobaloxime hybrid samples.

To verify the success of the tethering of the cobaloxime and the unperturbed structural integrity of the covalently modified hybrid COF–cobaloxime systems, we performed the same systematic experimental analysis as for the intact pCOF10. PXRD shows that the crystallinity of the COF is preserved and the stacking mode does not change with respect to pCOF10 (Figure S6, SI). Sorption analysis shows the expected reduction of the surface area according to Table S1 (SI). Pore size distributions for the clicked samples were calculated from Ar sorption isotherms, as shown in Figure S5 (SI). In all samples, the 2.3 nm pore size, as found in pCOF10, is preserved with lower pore volume fraction, while additional smaller pores up to 1.9 nm occur, as seen from optimized pore models (see Figure S19, SI). FT-IR spectra display all expected vibrations of the COF, including propargyl vibrations at ca. 3300 and 2300 cm⁻¹. These vibrations are still visible in ligand-tethered samples, which hints at partial transformation. New triazole peaks are hidden in the region around 3100 cm⁻¹ due to low intensity. The success of the click-reaction was further confirmed by the reduced intensity of the propargyl signals relative to the other signals in the 1D ¹³C{¹H} CP ssNMR spectrum upon addition of the azide compounds. We did not observe any additional signals arising from the clicked compound, which is probably due to signal superposition, especially in the aromatic region, and due to lower signal intensity caused by a low functionalization degree. UV–vis diffuse reflectance spectra show two additional broad absorption bands at 500 and 600 nm for the cobaloxime-containing samples (Figure S15, SI). These bands are due to the electronic transitions of the azide-functionalized cobaloximes. Depending on the reaction conditions (see the Supporting Information for more details), the cobaloxime loading can be adjusted within limits. For all samples, the total cobaloxime amount was determined by ICP analysis, and for [Co-1a]−COF, it was additionally confirmed by fast-MAS ¹H-detected NMR spectra. The values range from 0.47 to 2.4 wt % for route II, while route I resulted in higher cobaloxime amounts between 1.2 and 8.5 wt %. The highest cobaloxime content was found for [Co-1a]−COF, as can be seen in Table S2 (SI). The resulting functionalization degrees ranging from 2.0 to 15% are also listed there. Scanning electron microscopy shows a flower-like morphology for all samples. Elemental mapping showed a uniform distribution of carbon, nitrogen, oxygen, and cobalt in the samples, as can be seen in the Supporting Information.

### SSNMR analysis of the COF–cobaloxime hybrid systems

While powder diffraction analysis provides long-range spatial information, such as approximate interlayer separations, ssNMR provides us with short-range interatomic proximities and hints about the position of the cobaloxime inside the pore. To this end, we performed an in-depth structural analysis of the clicked samples 1a−COF and [Co-1a]−COF using ¹H-

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**Figure 2.** (A) Structure of the azide-functionalized ligands 1a, 1b, and 2 and (B) the azide-functionalized complexes Co-1a, Co-1b, and Co-2. (C) Exemplary postsynthetic COF modification toward [Co-1b]−COF. Synthesis conditions can be found in the Supporting Information.
detected, fast-MAS ssNMR at $\nu_{\text{rot}} = 55.55$ kHz at 700 MHz $^1$H Larmor frequency (16.4 T). The samples based on $[\text{Co-1a}]$ were chosen due to their higher molecular symmetry compared to that of $[\text{Co-1b}]$. Both $1\text{a}$$\rightarrow$COF and $[\text{Co-1a}]$$\rightarrow$COF were studied by 1D and 2D $^1$H and $^{13}$C solid-state NMR techniques. All 2D measurements were $^1$H-detected, which significantly improved the sensitivity of the natural abundance $^{13}$C measurements. In addition to the sensitivity gain, we could exploit the $^1$H chemical shifts as well as the $^1$H$\rightarrow$H correlations as sources of structural information.

Figure 3B compares the 1D $^1$H spectra of $1\text{a}$$\rightarrow$COF (yellow) and $[\text{Co-1a}]$$\rightarrow$COF (blue). The high structural order of these two-dimensional crystalline polymers is reflected in the good resolution of the $^1$H signals; $^1$H line widths vary between 800 and 1300 Hz for $1\text{a}$$\rightarrow$COF and between 1000 and 2000 Hz for $[\text{Co-1a}]$$\rightarrow$COF. In the $^1$H spectra, we could directly observe four ($1\text{a}$$\rightarrow$COF) and five ($[\text{Co-1a}]$$\rightarrow$COF) distinct proton resonances, which correspond to the amide proton (10.9 ppm), aromatic protons overlapping with the olefin proton (7.2 ppm), methylene protons (3.9 ppm), and methyl protons (1.7 ppm). For $[\text{Co-1a}]$$\rightarrow$COF, we also observe a well-separated, downfield-shifted, low-intensity peak that belongs to the pyridine aromatic proton (H13), as well as between the resonances of the strongly hydrogen-bonded oxime proton (19.1 ppm). Note that all $^1$H signals are broader in the spectrum of $[\text{Co-1a}]$$\rightarrow$COF relative to that of $1\text{a}$$\rightarrow$COF, which indicates that the cobaloxime functionalization process disrupted the overall COF crystallinity to some extent. Cobaloxime contains $\text{Co(III)}$, which is, unlike $\text{Co(II)}$, diamagnetic; therefore, the observed line broadening of $[\text{Co-1a}]$$\rightarrow$COF cannot be a consequence of paramagnetic relaxation enhancement. Also, residual $\text{CoCl}_2$ salt is washed out during the sample preparation process. It is more likely that the postsynthetic modification reduced the crystalline domain size and increased the sample’s heterogeneity, leading to a wider range of chemical shifts for each site.

The good $^1$H resolution of the fast-MAS $^1$H spectrum prompted us to measure 2D homonuclear correlation experiments to gain a deeper insight into the intramolecular interaction between the COF backbone and the cobaloxime cocatalyst. We probed the relative $^1$H$\rightarrow$H distances using 2D double quantum$\rightarrow$single quantum (DQ$\rightarrow$SQ) correlation experiments employing the R-symmetry-based R14$_r$ SQ recoupling sequence. The R14$_r$ SQ correlation experiment relies on the generation of double-quantum coherences via homonuclear dipole$\rightarrow$dipole coupling and chemical shift terms in the first-order Hamiltonian. We used a $R = \pi_1$ element as the basic R-symmetry block with a nutation frequency of 97.22 kHz (3.5$\nu_{\text{rot}}$). The homonuclear 2D $^1$H$\rightarrow$H DQ$\rightarrow$SQ recoupling experiment relies on the generation of double-quantum coherences via homonuclear dipole$\rightarrow$dipole coupling to obtain through-space information on nearby protons. Due to the double-quantum filter, the spectrum exhibits cross-peaks only between protons that share direct dipolar interactions with each other, and thus, no relayed magnetization transfer occurs. For protonated organic solid materials, such as the COFs of this study, the observation of a DQ peak is indicative of a proton$\rightarrow$proton proximity that is $\leq 3.5$ Å. The relative signal intensities could well-approximate interatomic distances.

Parts C and D of Figure 3 show the DQ$\rightarrow$SQ correlation spectra of $1\text{a}$$\rightarrow$COF (yellow) and $[\text{Co-1a}]$$\rightarrow$COF (blue). The spectra reveal double-quantum correlations between both distinct and identical environments, appearing at the off-diagonal and diagonal positions, respectively. Diagonal peaks are expected for the signals of the methyl and methylene groups, as well as between the resonances of the chemically equivalent aromatic sites. However, the weak diagonal peak for the NH protons corresponds to an NH$\rightarrow$NH autopeaks, which is indicative of the dipolar interaction between COF layers; the separation of NH protons within one layer is $<7$ Å, while the layer-to-layer distance is 3.5 Å according to powder crystal analysis. The two spectra look almost identical, the only considerable difference being the $^1$H cross-peaks of the oxime $^1$H at 19.1 ppm with resonances at 8.7 and 3.4 ppm. In order to assign these two peaks, and thus uncover the position of cobaloxime inside the pore, we performed a detailed quantum-chemical study (vide infra). On the basis of these studies, we conclude that the resonances at 8.7 and 3.4 ppm belong to the pyridine aromatic proton (H13), as well as to a downfield-shifted methyl proton of a neighboring ethoxy group with which the cobaloxime is in close contact.

Next, we assessed the relative flexibility of the two compounds using 1D $^{13}$C NMR spectroscopy. Three different 1D $^{13}$C MAS spectra of $1\text{a}$$\rightarrow$COF and $[\text{Co-1a}]$$\rightarrow$COF are given in parts A and B of Figure 4, respectively. These spectra include $^{13}$C($^1$H) cross-polarization (CP) MAS and $T_1$-weighted, direct-polarization (DP) $^{13}$C spectra recorded with short (1 s) and long (25 s) recycle delay times. These latter spectra were used to elucidate the relative mobility of certain sites in the COF samples. In the $^{13}$C spectra recorded with
not only on the amplitude of nanosecond time-scale motion but also on the number of directly attached protons: the more protons that are directly bound to a carbon, the faster it relaxes via heteronuclear dipolar relaxation. This is reflected in the relative change of signal intensities among the aromatic carbons. Besides, the methyl resonance relaxes rapidly due to the free rotation around the C–C axis in the ethyl group. The methyl resonance line shape in the DP spectrum of [Co-1a]−COF is markedly distorted, presenting a shoulder at lower resonances. This signal could be assigned to the methyl carbons of the cobaloxime ligand. Otherwise, the signals of the covalently tethered ligand do not show any obvious sign of increased fast time-scale flexibility, neither for [1a]−COF nor for [Co-1a]−COF. In the spectrum of [Co-1a]−COF recorded with $d_1 = 1$ s, the intensified resonances at 128 ppm indicate rather flexible aromatic sites, but due to strong overlaps in this region, we could not identify if this signal belongs to the ligand or to some residual impurities that tend to show up more strongly in $T_1$-weighted experiments. Selective $^{13}$C or $^{15}$N labeling at specified positions of the ligand would help us to quantify the amplitude and time-scale of the ligand motion.

The apparent lack of high-amplitude fast time-scale dynamics of the two COF frameworks was further validated by comparing $^1$H-detected 2D CP-based $^1$H−$^{13}$C correlation spectra with INEPT-based 2D HSQC spectra (Figure 4D, E). High-amplitude nanosecond time-scale motion results in inherent decoupling and thus leads to increased coherent lifetimes in INEPT-based experiments and to decreased transfer efficiencies in CP-based experiments. In the HSQC spectrum of both [1a]−COF and [Co-1a]−COF, we observe only a single methyl peak, indicating that the COF backbone is generally rigid on the nanosecond time-scale.

**COMPUTATIONAL STUDIES**

In order to provide a structural model for the position and the orientation of the covalently tethered cobaloxime cocatalyst inside the pore, we conducted a detailed *in silico* structural investigation of [1a]−COF and [Co-1a]−COF. Atom positions and lattices of the periodic COF structure of [1a]−COF were optimized at the RI-PBE-D3/def2-QZVP40−43 level of theory using an acceleration scheme based on the resolution of the identity (RI) technique and the continuous fast multipole method (CFM)44−46 implemented47,48 in Turbomole ver. V7.1.49 The obtained structure for the [1a]−COF was then used to prepare parameters for molecular dynamics simulations using antechamber.50 Force field minimizations and subsequent dynamics were performed with the NAMD program package51,52 using GAFF parameters53 afterward. NMR chemical shifts were then calculated with the NAMD program package and used them to identify features that are also present in the experimentally obtained ssNMR spectra. Such features include the number of cross-peaks, especially cross-peaks of the oxime proton, their relative intensity ratios, and their peak positions. The most distinctive factor in the simulated $^1$H−$^1$H DQ−SQ spectra is the presence of oxime (H15) cross-peaks with resonances at around 8.7 and 3.4 ppm.
which was used to categorize the simulated spectra. These distinct chemical shifts suggest that the oxime proton is interacting with an aromatic proton (at 8.7 ppm) and with either an upfield-shifted methylene proton or with a downfield-shifted methyl proton (at 3.4 ppm). There are four different aromatic protons in [Co-1a]−COF, H1, H4, H12, and H13, out of which only H4 and H13 can get closer than 3.5 Å to H15.

To decide which resonance leads to the 3.4 ppm cross-peak with H15, we analyzed the shielding effects of the glyoxime group on the nearby ethoxy methyl and methylene protons. The approach of the glyoxime oxygen toward the ethoxy group induces a deshielding effect; consequently, both the methyl and the methylene protons resonate at higher frequencies (see Figure S16 and SI text for more details), this rules out the possibility that the cross-peak at 3.4 ppm would stem from an upfield-shifted methylene proton and leaves only a downfield-shifted methyl proton as a possible interaction partner. Besides, we excluded the possibility that the oxime proton shows a trivial intraligand cross-peak with the glyoxime methyl protons, since (i) the distance between the H15 and H16 protons is >3.5 Å and (ii) the calculated chemical shift is below 2.9 ppm.

Out of the 200 simulated $^1$H−$^1$D Q−SQ spectra, 27 (13) contained two (three) oxime cross-peaks, among which 22 spectra have these peaks in the expected ppm range. By considering the relative peak intensity ratios between the oxime cross-peaks, only 15 spectra have a more intense aromatic−oxime than a methyl−oxime cross-peak. Two such spectra, together with the simulated $^1$H−$^1$C 2D spectra and corresponding structures, are displayed in Figure 5A–F. As counter-examples, Figures 5G, H, and J, K displays the spectra of such structures (Figure S1 and L) where three equally intense peaks (Figure S5H) or no oxime proton cross-peak (Figure S5K) appears in the simulated DQ−SQ spectra. The possibility that, in reality, in a fraction of the [Co-1a]−COF pores the cobaloxime does not interact with the pore wall cannot be
ruled out; nonetheless, our current data suggest that when it
does, it gets in close contact with the nearby ethoxy group. It is
also likely that this genuine interaction stabilizes the complex
and restricts the cocatalyst’s degradation during the photocat-
calytic cycles. Note that, at this stage, both the ssNMR
measurements and the in silico calculations were performed in
a solvent-free environment. Future ssNMR measurements with
added acetonitrile/water mixture accompanied by simulations
in explicit solvent could reveal if the cobaloxime stays attached
to the pore wall or whether it gains more flexibility and drifts
toward the pore center.

To inspect the spatial arrangement inside the pore, we
modeled $[\text{Co-1a}]^{-}\text{COF}$ including one tethered cocatalyst
based on the MD-simulated structures (Figure 6). The

displayed ligand has the same orientation as in Figure 5C.
From the side and front views it is apparent that the ligand
spreads over multiple layers and occupies a substantial portion
of the pore. Due to spacial confinements, our model suggests
that no more than three $[\text{Co-1a}]$ over three layers can fit into
the backbone; i.e., the maximum number of $[\text{Co-1a}]$ per layer
is one. In our case, we have 13 mol % functionalization, which
translates into one $[\text{Co-1a}]$ for every seven layers.

## PHOTOCATALYTIC ACTIVITY

To probe whether there is a possible benefit of covalent
cocatalyst immobilization over simple physisorption, $^{14,15}$ the
COF−cobaloxime hybrid samples were tested for photo-
catalytic activity. In a typical photocatalysis experiment, 5 mg
of COF hybrid was suspended in 10 mL of acetonitrile and
water in a ratio of 4:1 at pH 8 containing 100 μL of
triethanolamine (TEOA) as sacrificial donor. A housed Xe
lamp was used to illuminate the suspension interface with a
nominal beam spectral distribution similar to that of AM1.5G.
The beam intensity before experiments was then adjusted to
100 mW cm$^{-2}$. See the SI for more details. Photocatalytic
hydrogen evolution reaction (HER) rates were quantified in a
continuous flow reactor as previously reported $^{13}$ (Figure 7A).
As a reference system, we compared the hybrid systems to
samples where $[\text{Co-1a}]$ or $[\text{Co-1b}]$ was added to the
suspension and physisorbed to COF-42 with a BET surface
area of 2336 m$^2$ g$^{-1}$ during photocatalysis. The maximum
photonic efficiencies after in situ photoactivation of the samples
ranging from 2 to 8 wt % cobaloxime catalyst according to ICP
results can be found in Figure 7A. In the physisorbed samples,
an increase of the photonic efficiency was found when
increasing the catalyst amount from 2 to 4 wt % with a
maximum efficiency of 0.06% for $[\text{Co-1a}]$ and 0.07% for $[\text{Co-
1b}]$ at 4.0 wt %, while the efficiency is fairly constant at higher
percentages (0.06%–0.08% at 5.0 and 8.0 wt %) for $[\text{Co-1b}]$.
This behavior is expected for the system; as in the low-loading
region, the photocatalytic activity scales linearly with the
cocatalyst amount, while it reaches a maximum in the higher-
loading region where the availability of the cocatalyst is not
limiting anymore.

In the hybrid samples, an activity maximum rather than a
constant behavior is found for each hybrid type. For the para-
functionalized $[\text{Co-1a}]$, the highest photonic efficiency was
found at 4.1 wt %, while for the meta-functionalized $[\text{Co-1b}]$
the maximum was found at 3.2 wt %. As before, a linear
increase of the photonic efficiency in the low-loading regime
was observed. However, a further increase in cobaloxime
loading resulted in lower activity in the immobilized samples.
We attribute this to a predominant pore-clogging effect of the
active sites with increasing functionalization. In general, the
highest photonic efficiency was achieved with $[\text{Co-1a}]^{-}\text{COF}$
at 0.14% followed by $[\text{Co-1b}]^{-}\text{COF}$ at 0.11%. Compared to
the physisorbed samples with the corresponding cobaloxime

![Figure 6. Front and side views of the MD-simulated structural model of $[\text{Co-1a}]^{-}\text{COF}$ showing a possible arrangement of the cocatalyst. The linker and the cobaloxime group are depicted by spheres and their carbon atoms are displayed in orange. Co, Cl, O, N, and H atoms are displayed in pink, lime, red, blue, and white, and C atoms of the backbone are light blue.](image)

![Figure 7. (A) Comparison of photonic efficiencies for hybrid samples and COF-42 with physisorbed $[\text{Co-1a}]$ and $[\text{Co-1b}]$. (B) Comparison of the hydrogen evolution rate of $[\text{Co-1b}]^{-}\text{COF}$ containing 3.2 wt % $[\text{Co-1b}]$ and COF-42 with 4.0 wt % physisorbed $[\text{Co-1b}]$ and coarse-grained model fits of both systems. (C) Projection of the hydrogen evolution of $[\text{Co-1b}]^{-}\text{COF}$ containing 3.2 wt % $[\text{Co-1b}]$ and COF-42 with 4.0 wt % physisorbed $[\text{Co-1b}]$ based on the coarse-grained models.](image)
content, the activity doubles for both systems. Additionally, to emphasize the role of the complex environment of the cobaloxime over the pure presence of Co(II), we performed a measurement where we added CoCl₂ to a suspension of pCOF-42 and triethanolamine in the photocatalysis medium, as well as experiments where one of the components (COF, TEOA) was excluded (see Supporting Information). None of the reference samples showed hydrogen evolution after several hours of irradiation. For the hybrid samples, the close contact between the cobaloxime and the COF pore wall—revealed by representative solid-state NMR and computational studies (vide supra) with [Co-1a]−COF—might facilitate charge transfer to the cobaloxime catalyst from the COF pore wall, as also observed from photoluminescence measurements (see Figures S11 and S12, SI) where [Co-2]−COF shows a significantly lower activity in CH₃CN/H₂O, which is a known effect for cobaloximes that lack equatorial protons. The protonation of the oxime oxygen, which is necessary for the catalytic process, is hindered in those cases.⁵⁸,⁵⁹ The catalytic activity could not be improved by lowering the pH to 4. In this case, different acids (ascorbic acid, acetic acid, and citric acid) were tested that simultaneously served as sacrificial electron donors instead of the amine base TEOA. Even though the stability of [Co-2]−COF is predicted to be higher than that for the other tested cobaloximes, the complex proved not to be appropriate in our case. We compared the best performing [Co-1b]−COF sample (containing 3.2 wt % cobaloxime) to COF-42 physisorbed with [Co-1b]. A sample with the same amount of physisorbed cobaloxime was qualitatively active in photocatalytic hydrogen evolution, but for precise quantification, we increased the catalyst amount to 4.0 wt %. Even though it contained 20% less catalyst, the hybrid sample was 47% more active than the physisorbed one (163 vs 111 μmol h⁻¹ g⁻¹) (see Figure 7A). Additionally, the long-term stability increased significantly. After 20 h, the physisorbed sample shows 52% of its initial activity, while the hybrid sample maintains 80% of its initial activity. To get an estimate of the longevity of the systems, we fitted the hydrogen evolution rates of both samples with a coarse-grained model (Figure 7C) that was established in an earlier study on photocatalysis with COFs and a nickel-based oligomer as cocatalyst.¹⁵ The model resulted in very precise fitting for the physisorbed catalyst because of similarities to the original nickel-based system from where the coarse-grain fitting model was obtained, while the hybrid sample showed a more complex behavior that is not perfectly mapped with this simplified model. On the basis of the coarse-grained fits, we projected the total amount of hydrogen evolved by the samples at full depletion (see Figure 7C). After 780 h, the projection of the physisorbed sample reaches 35 μmol of hydrogen evolved, while the value is 59 μmol for the hybrid sample, which is a gain of 69%. Comparing the estimated turnover numbers (TONs) of both systems, the deviation gets even more obvious. While the TON after 780 h is simulated to be 81 for the physisorbed sample, it increases by 110% to a value of 170 in the hybrid sample. We attribute this activity enhancement to the local confinement in the COF hybrid samples, as supported by MD simulations.

Cobaloximes are known to slowly decompose under photocatalytic conditions. The labile axial pyridine ligand decoordinates in the catalytic cycle due to a square-planar Co(II) transition state. The catalyst in solution can then possibly be reduced, which limits its stability. Due to the confinement between the ligand and catalyst in the COF pores, the recoordination might be enhanced, hence counteracting degradation, which leads to reactivation of the catalyst. Additionally, charge transfer is favored in the case of the spatial proximity of the cocatalyst and the pore wall. Both effects result in higher overall activity as well as longevity.

Interestingly, the activation period for the hybrid samples is significantly longer than for the physisorbed ones. This may be attributed to the time-delayed accessibility of the catalyst in the pores. Both limitations could be addressed via a method that was recently published by Thomas and co-workers,⁶⁰,⁶¹ where silica spheres were used to create an inverse-opal architecture in the COF material. The so created macropores could serve as channels for reagents and products. Also, immobilization of the cocatalyst in a COF with larger pores might have a similar effect.

**CONCLUSION**

In summary, we have developed a platform derived from COF-42 as a support for the immobilization of cobaloxime catalysts. The postsynthetic modification of propargyl-functionalized COF-42 enabled the covalent tethering of three different cobaloximes to form COF−cobaloxime hybrid systems. This tethering significantly enhanced the photocatalytic activity of the system by more than 100% compared to that of the physisorbrates with the corresponding cobaloxime amount. The high crystallinity of our materials allowed for an in-depth solid-state 2D NMR characterization using fast MAS and proton detection. In the 1D ¹H spectrum of [Co-1a]−COF, we could clearly identify the resonance corresponding to the oxime proton on the basis of its highly downfield-shifted resonance. The 2D ¹H−¹H DQ−SQ experiment showed two cross-peaks for the oxime proton, consistent with the incorporation of the cocatalyst into the COF material. MD simulations with subsequent quantum-chemical NMR chemical shift calculations allowed us to locate the position of the tethered ligand inside the pore on the basis of the experimentally observed oxime proton cross-peaks. Our analysis suggests that the cobaloxime in [Co-1a]−COF closely interacts with the pore wall. We surmise that this interaction is responsible both for the improved photocatalytic activity and for the prolonged activity of the hybrid samples with respect to the physisorbed variant. We anticipate that larger pore channels or the addition of dedicated transport pores will further improve the pore accessibility and prevent back-reaction via local confinement of the products, thereby increasing the hydrogen evolution activity of the system even further.

**ASSOCIATED CONTENT**

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.0c02155.

Experimental procedures, COF synthesis, and details of molecular dynamic simulations, quantum-chemical calculations, and additional measurements (PDF)

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Notes

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