Light-active Metal-Organic Frameworks for Photocatalytic Hydrogen Evolution.

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Abstract

The general scope of this thesis lies in the application of MOFs in photocatalysis. MOFs demonstrate inherent properties – such as high porosity, tunable optoelectronic and catalytic properties, which render them promising candidates for photocatalysis. The main research activity relative to this thesis aims at the advancement of MOF-based photocatalytic systems in terms of activity, cost and sustainability. This was achieved by employing different strategies that can lead to the favorable modification of the three major photocatalytic steps of light absorption, charge separation-migration and catalysis, which govern the overall performance of a photocatalytic system.

More specifically, Chapter 2 describes the impact of different co-catalysts on the photocatalytic activity of a MOF-based system, based on the well-known MIL-125-NH2. Variation of the co-catalysts can significantly influence the two major photocatalytic steps of charge separation and the catalytic reaction. All the metal oxide and phosphide co-catalysts investigated are found to significantly improve the activity of MIL-125-NH2, with the system using Ni3P nanoparticles (NPs) exhibiting a high H2 evolution rate (1230 μmol h⁻¹ g⁻¹) and an apparent quantum yield of 6.6% at 450 nm, which is comparable to the state of the art. Comparison of Ni3P with Pt showed that the noble-metal-free Ni3P/MIL-125-NH2 system significantly outperforms Pt/MIL-125-NH2. These results are attributed to the enhanced electronic interactions between MIL-125-NH2 and Ni3P, and prove that earth abundant co-catalysts can challenge the commonly used noble metals.

The low cost and high efficiency of Ni3P/MIL-125-NH2 prompted me to focus on another component used in photocatalytic systems, which is the electron donor. Chapter 3 shows that variation of the electron donors highly influences the activity of Ni3P/MIL-125-NH2, with triethylamine significantly boosting the H2 evolution rate. However, the utilization of electron donors is another factor hindering the industrialization of such systems, since these substances can be toxic and expensive. Inspired by this challenge, we then replaced the electron donor with rhodamine B (RhB) – a simulant organic pollutant – envisioning dual-functional photocatalysis for simultaneous H2 generation and organic pollutant degradation. This research project revealed the first example of a MOF-based dual-functional photocatalytic system able to generate H2 in a high rate and degrade RhB under visible-light.

Chapter 4 describes a strategy for enhancing the photocatalytic performance of MOF-based systems by favourably altering the photocatalytic steps of light harvesting and charge separation. This was achieved by developing MOF/MOF heterojunctions. The combination of MIL-125-NH2 with MIL-167 – a Ti-based MOF with complementary light absorption properties – leads to the formation of a type II heterojunction MIL-167/MIL-125-NH2, with enhanced optoelectronic properties. MIL-
167/MIL-125-NH$_2$ significantly outperforms its single components MIL-167 and MIL-125-NH$_2$, in terms of photocatalytic H$_2$ production. This strategy contributes to the discovery of novel MOF-based photocatalytic systems that can harvest the solar energy and exhibit high catalytic activities in the absence of co-catalysts.

Finally, Chapter 5 shows an alternative role of MOFs in the photocatalysis field, which lies in their utilization as precursors for the synthesis of highly active metal-oxide semiconductor photocatalysts. MIL-125-NH$_2$ was calcined to form pure and mixed anatase-rutile phase TiO$_2$ NPs. The MIL-125-NH$_2$-derived TiO$_2$ with anatase to rutile ratio of approximately 3/1 exhibits higher photocatalytic H$_2$ evolution rates than the commercial P25 Degussa, TiO$_2$ synthesized from precipitated titanium hydroxide and MIL-167-derived TiO$_2$. This is attributed to the templated morphology of MIL-125-NH$_2$-derived TiO$_2$, which forces the anatase and rutile NPs into effective contact with each other, leading to an efficient electron-hole separation.

**Keywords**
RESUME

La portée générale de cette thèse réside dans l'utilisation des MOFs pour des applications photocatalytiques. Les MOFs présentent des propriétés inhérentes — telles qu'une porosité élevée, des propriétés optoélectroniques et catalytiques accordables — qui en font des candidats prometteurs pour la photocatalyse. La principale activité de recherche relative à cette thèse vise à faire progresser les systèmes photocatalytiques basés sur MOFs en termes d'activité, de coût et de durabilité. Cela a été réalisé en employant différentes stratégies qui peuvent conduire à l'amélioration des trois principales étapes photocatalytiques qui régissent les performances globales d'un système photocatalytique : l'absorption de la lumière, la séparation-migration de charge et la catalyse.

Plus précisément, le Chapitre 2 décrit l'impact de différents co-catalyseurs sur l'activité photocatalytique d'un système basé sur le MOF bien connu MIL-125-NH₂. La variation des co-catalyseurs peut influencer de manière significative les deux principales étapes photocatalytiques de séparation des charges et la réaction catalytique. Tous les co-catalyseurs d'oxyde et de phosphure métalliques étudiés améliorent considérablement l'activité du MIL-125-NH₂, le système utilisant des nanoparticules de Ni₂P (NPs) présentant un taux d'évolution élevé de H₂ (1230 μmol h⁻¹ g⁻¹) et un rendement quantique apparent de 6,6% à 450 nm, ce qui est comparable à l'état de l'art. La comparaison du Ni₂P avec le Pt a montré que le système Ni₂P/MIL-125-NH₂ sans métal noble surpasse de manière significative le système Pt/MIL-125-NH₂. Ces résultats sont attribués aux interactions électroniques améliorées entre le MIL-125-NH₂ et le Ni₂P, et démontrent que les co-catalyseurs à base de matériaux abondants peuvent remplacer les métaux nobles couramment utilisés.

Le faible coût et la haute efficacité du système Ni₂P/MIL-125-NH₂ m'ont incité à me concentrer sur un autre composant utilisé dans les systèmes photocatalytiques, qui est le donneur d'électrons. Le Chapitre 3 montre que la variation des donneurs d'électrons influence fortement l'activité du Ni₂P/MIL-125-NH₂, la triéthylamine augmentant considérablement le taux d'évolution de H₂. Cependant, l'utilisation de donneurs d'électrons est un autre facteur entravant l'industrialisation de tels systèmes, car ces substances peuvent être toxiques et coûteuses. Inspirés par ce défi, nous avons ensuite remplacé le donneur d'électrons par de la rhodamine B (RhB) - un polluant organique simulant - envisageant une photocatalyse à double fonction pour la génération simultanée de H₂ et la dégradation des polluants organiques. Ce projet de recherche a révélé le premier exemple d'un système photocatalytique à double fonction basé sur un MOF capable de générer du H₂ à un taux élevé et de dégrader le RhB sous la lumière visible.
Résumé

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Enfin, le Chapitre 5 montre un rôle alternatif des MOF dans le domaine de la photocatalyse, qui réside dans leur utilisation comme précurseurs pour la synthèse de photocatalyseurs semi-conducteurs à oxyde métallique très actifs. Le MIL-125-NH₂ a été calciné pour former des NPs TiO₂ en phase anatase-rutile purs et mélangés. Le TiO₂ dérivé de MIL-125-NH₂ avec un rapport anatase/rutile d'environ 3/1 présente des taux d'évolution photocatalytique de H₂ plus élevés que le matériel commercial Degussa P25, un TiO₂ synthétisé à partir d'hydroxyde de titane précipité et TiO₂ dérivé de MIL-167. Cela est attribué à la morphologie basée sur des modèles de TiO₂ dérivé de MIL-125-NH₂, qui force les NPs anatase et rutile à entrer en contact efficace les uns avec les autres, conduisant à une séparation efficace électron-trou.

Mots-clés

Photocatalyse, Structure Métal-Organique, Réduction de l'Eau, Photocatalyse Bifonctionnelle, Photo-oxydation des Polluants Organiques, Hétérojonctions, Synthèse Médiée par le MOF
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List of Abbreviations and Symbols

° Degree
Å Ångström
•OH Hydroxyl radical
ºC Degree Celsius
BDC Terephthalic acid
CB Conduction Band
CH₃CN Acetonitrile
CO₂ Carbon dioxide
COF Covalent-organic frameworks
DMF Dimethylformamide
e.g. Exempla gratia
e⁻ Electron
EDX Energy Dispersive X-ray Spectroscopy
EPR Electron paramagnetic resonance
EtOH Ethanol
eV Electronvolt
g Gram
h Hour
h⁺ Hole
H⁺ Proton
H₂ Hydrogen
HOCOs Highest occupied crystalline orbitals
HOMO Highest occupied molecular orbital
i.e. Id est
LUÇOs Lowest unoccupied crystalline orbitals
LUMO Lowest unoccupied molecular orbital
M Molar (mol L⁻¹)
MeOH Methanol
mg Milligram
MIL Material from Institute Lavoisier
min Minutes
mL Millilitre
### List of Abbreviations and Symbols

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>MOF</td>
<td>Metal-organic framework</td>
</tr>
<tr>
<td>MW</td>
<td>Molecular weight</td>
</tr>
<tr>
<td>N₂</td>
<td>Nitrogen</td>
</tr>
<tr>
<td>NHE</td>
<td>Normal Hydrogen Electrode</td>
</tr>
<tr>
<td>NPs</td>
<td>Nanoparticles</td>
</tr>
<tr>
<td>NH₂-BDC</td>
<td>2-Aminophthalic acid</td>
</tr>
<tr>
<td>(NH₂)₂-BDC</td>
<td>2,5-Diaminoterephthalic acid</td>
</tr>
<tr>
<td>nm</td>
<td>Nanometer</td>
</tr>
<tr>
<td>O₂</td>
<td>Oxygen</td>
</tr>
<tr>
<td>O₂⁻</td>
<td>Superoxide radicals</td>
</tr>
<tr>
<td>OH-BDC</td>
<td>2-Hydroxyterephthalic acid</td>
</tr>
<tr>
<td>(OH)₂-BDC</td>
<td>2,5-Dihydroxyterephthalic acid</td>
</tr>
<tr>
<td>PL</td>
<td>Photoluminescence</td>
</tr>
<tr>
<td>ppm</td>
<td>Parts per million</td>
</tr>
<tr>
<td>PXRD</td>
<td>Powder X-ray Diffraction</td>
</tr>
<tr>
<td>ref.</td>
<td>Reference</td>
</tr>
<tr>
<td>RhB</td>
<td>Rhodamine B</td>
</tr>
<tr>
<td>SEM</td>
<td>Scanning Electron Microscopy</td>
</tr>
<tr>
<td>t</td>
<td>Time</td>
</tr>
<tr>
<td>T</td>
<td>Temperature</td>
</tr>
<tr>
<td>TEA</td>
<td>Triethylamine</td>
</tr>
<tr>
<td>TEM</td>
<td>Transmission Electron Microscopy</td>
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<tr>
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<td>Thermogravimetric Analysis</td>
</tr>
<tr>
<td>TPL</td>
<td>Transient photoluminescence</td>
</tr>
<tr>
<td>UIO</td>
<td>University of Oslo</td>
</tr>
<tr>
<td>UV</td>
<td>Ultraviolet</td>
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<tr>
<td>VB</td>
<td>Valence band</td>
</tr>
<tr>
<td>Vs</td>
<td>Versus</td>
</tr>
<tr>
<td>ΔG</td>
<td>Gibbs free energy</td>
</tr>
<tr>
<td>λ</td>
<td>Wavelength</td>
</tr>
<tr>
<td>μmol</td>
<td>Micromole</td>
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Note: additional abbreviations for specific compounds are defined within the thesis.
Chapter 1 Introduction

Parts of this chapter are based on published work:

‘Dual-Functional Photocatalysis for Simultaneous Hydrogen Production and Oxidation of Organic Substances’

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S. Kampouri’s contribution: writing the manuscript and construction of figures
1.1 Introduction to Photocatalysis

The images at Figure 1.1 represent humanity’s heritage from the 20th century; the increasingly growing environmental pollution and water scarcity, abused natural resources, climate change and extensive deforestation. These major problems are consequences of our society’s activities and are associated with the industrial rise, the increasing growth of population and an economy relying on the exploitation of fossil fuels. Thus, it is our urgent obligation to develop sustainable strategies to mitigate these environmental problems. Among renewable energy technologies, a promising approach lies in the use of energy from light to excite a material that can enable different redox reactions. This broad field is known as photocatalysis. Each potential redox reaction that can occur represents an individual photocatalytic field. More specifically, the major photocatalytic fields that have been extensively studied include water splitting (reduction: H2 production, oxidation: O2 production), carbon dioxide (CO2) reduction, synthesis of molecules, oxidation of organic pollutants, metal reduction and disinfection through the inactivation of pathogenic microorganisms, such as viruses, bacteria and protozoa (Figure 1.2).

Photocatalytic water splitting into H2 and O2 constitutes a field of significant importance, since H2 is regarded as an ideal replacement of fossil fuels, allowing for an energy cycle free of greenhouse gasses. The fundamental principles in this field will be thoroughly described in chapter 1.3. Another major field in photocatalysis includes the light-driven oxidation of organic substances, which can be used to either achieve full degradation of hazardous organic contaminants or to synthesize useful...
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organic molecules. The increasing amounts of wastewater and high toxicity of several organic substances in streams have prompted extensive research in this field. Photocatalytic reduction of metal ions is another efficient approach to alleviate water pollution induced by inorganic pollutants and is based on the fact that the toxicity of a given metal is highly correlated with its oxidation state. This photocatalytic field serves to transform such metals to a more environmentally benign form (e.g. Cr\textsuperscript{VI} to Cr\textsuperscript{III}) or to reduce them to their elemental form and remove it for safe disposal. Last, considering that CO\textsubscript{2} represents the most common greenhouse gas contributing to global warming, the field of CO\textsubscript{2} photocatalytic reduction into useful chemicals (e.g. methane, methanol, formaldehyde, formic acid, and others) is regarded as a promising technology to avoid further increase in the concentration of CO\textsubscript{2} in the atmosphere. In principle, this photocatalytic process involves the reduction of CO\textsubscript{2} along with the oxidation of water, which leads to the production of solar fuels.

1.2 Fundamental Principles of Photocatalysis

One of the milestones in the photocatalysis field constitutes the exploration of titanium dioxide (TiO\textsubscript{2}) for light-induced water splitting, reported in 1972 by Fujishima and Honda. Since then, immense research efforts in photocatalysis has focused on TiO\textsubscript{2} and other semiconducting materials. In contrast to conductors, where the valence and conduction bands (VB and CB, respectively) overlap, a given semiconducting material is characterized by a VB and CB that are separated by a band gap (Figure 1.3A). In this case, the band gap is significantly smaller than that of insulating materials (within the energy range of UV, visible or infrared light) and thus, when sufficient light energy is employed,
electron-hole pairs can be generated. The main requirement for this process to occur is that the photons’ energy should be equal to or higher than the band gap energy of a given semiconductor. Subsequently, the photogenerated charge carriers will be transferred to the surface of the photocatalyst where redox reactions can be initiated. Conclusively, the major steps in every photocatalytic reaction involve i. the light harvesting by the photocatalyst ii. the charge separation and transportation to the surface of the photocatalyst, followed by iii. the redox reactions (Figure 1.3B). The overall equilibrium of the kinetics and thermodynamics of these three distinct steps governs the photocatalytic performance of a light-active material.

**Figure 1.3:** (A) Band gap diagram of conductors, semiconductors and insulators. (B) Schematic illustration of major photocatalytic steps. Step 1: Light absorption, Step 2: Generation of electron-hole pairs and their migration to the surface of the photocatalyst and Step 3: Surface redox reactions.

### 1.2.1 First Major Photocatalytic Step: Light harvesting

Regarding the first step of light absorption, it is highly important not only for the material to exhibit enhanced light absorption (high value of absorbance coefficient), but also to harvest light with specific energy. The ultimate goal in the development of new technologies is to utilize energy from renewable sources such as the sun, wind and wave power. Thus, in the field of photocatalysis it is of crucial importance to harvest the abundant solar energy. The power of the solar energy incident on the Earth’s surface is around 1000 W m\(^{-2}\). Should all this energy be exploited, it could completely cover the global energy demand.\(^{10}\) The solar spectra at sea level is comprised of around 43% visible and 53% infrared light. UV light, on the other hand, accounts for only 4% of the solar radiation (Figure 1.4). While it is known that the most stable semiconductors are characterized by wide band gaps, materials with narrow band gaps, which are capable of absorbing light in the visible or infrared region, are highly desirable.

Numerous research studies have been devoted to the development of strategies in order to enhance or extend the absorption of materials (photocatalysts) in the visible region of the solar spectrum.
Approaches to achieve this include the utilization of dyes as photosensitizers,\textsuperscript{11} doping a given semiconductor with metal or non-metal ions,\textsuperscript{12} utilizing plasmonic nanoparticles for surface plasmon resonance,\textsuperscript{13} or coupling with narrow band gap semiconductors for the formation of heterojunctions (Figure 1.5).\textsuperscript{14}

![Soral Irradiance Spectra](image)

**Figure 1.4:** Soral irradiance spectra (G-173). Blue, gray and black lines represent the extraterrestrial, the global total on 37° sun-facing tilted surface and the direct normal spectral irradiance, respectively. The ultraviolet (UV), visible (Vis) and infrared (IR) regions are highlighted. Data taken from the American Society for Testing and Materials (ASTM).\textsuperscript{15}

### 1.2.2 Second Major Photocatalytic Step: Charge Separation and Migration

As mentioned above, the second step in a photocatalytic reaction involves the separation and migration of the photogenerated charge carriers, which is of significant importance. A photocatalyst with a large visible-light absorption does not guarantee photocatalytic efficiency. The excited state lifetime of the photocatalyst should be long enough for the photogenerated charge carriers to be transported to the surface of the material and, subsequently, to initiate a chemical reaction. High electron-hole recombination rates within the photocatalysts can explain why many photocatalytic reactions demonstrate low efficiencies. Improvement of this second step has attracted intense research interest, with numerous successful strategies being reported, including the design of high crystalline...
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photocatalysts with less grain boundaries or other defects, the formation of heterojunctions (Figure 1.5), or incorporation of co-catalysts to inhibit the undesired electron-hole recombination, by withdrawing the generated charges. Moreover, modification of the optical and electronic properties of a photocatalyst by following the aforementioned approaches could also result in enhanced charge separation efficiency, in addition to improving the first step of light harvesting.

Figure 1.5: Spatial separation of the photogenerated charge carriers within a type II heterojunction with a staggered gap.

1.2.3 Third Major Photocatalytic Step: Catalytic Redox Reaction

The third step of the photocatalytic reaction has been rather neglected in terms of scientific attention, as compared to the previous two steps. This final step can be significantly promoted by inclusion of catalytic active sites, such as the co-catalysts. Co-catalysts are frequently incorporated in photocatalytic systems, since they play a key role in inhibiting the undesired electron hole recombination and reducing the activation energy of a given reaction. An appropriate co-catalyst should exhibit thermodynamic compatibility with the photocatalyst, high intrinsic catalytic activity for a given reaction, and affinity with the reactants. For example, in the case of the H₂ evolution reaction, the free energy of H₂ adsorption, ΔG_H₂, is highly correlated with the catalytic activity of a co-catalyst and thus is considered as a descriptor for evaluating co-catalysts for this reaction. In principle, such co-catalysts should be characterized by a ΔG_H₂ value close to zero, exhibiting adequate interactions with H_ads but avoiding H_ads being too strongly bound to the co-catalyst (Figure 1.6). There have been few recent reviews summarizing the advances in improving this final step through the incorporation of catalytic active sites.
1.3 Photocatalytic Hydrogen Evolution

The continuously growing energy consumption, along with severe environmental problems caused by the utilization of fossil fuels, have induced immense research efforts directed toward the development of an alternative energy “path”, free of pollutants or greenhouse gases, such as N₂O and CO₂. H₂ is considered a clean fuel and a sustainable energy carrier, since it can be produced from natural sources (e.g. water, biomass), it stores high mass-specific energy density, and it only produces water as result of its combustion. Currently, most of the industrially produced H₂ is derived from fuel-based processes; more specifically steam methane reforming and coal gasification. These H₂ production methods however, contribute to the global warming, since CO₂ is also generated as a by-product in high quantities. Interestingly, 5% of the industrially produced H₂ is generated through water electrolysis, which can be a sustainable process when the electricity used is generated by renewable resources (e.g. solar, wind, hydropower). That being said, the high cost of this technique has hindered its broad industrialization.

As mentioned above, H₂ can also be directly generated through photochemical water splitting. Over the last few decades, photocatalytic H₂ evolution has been widely studied, and, solar-driven photocatalysis is regarded as an ideal method to sustainably produce H₂. Pure water splitting involves two half reactions of H₂ and O₂ generation. For this process to occur, it is necessary that the highest level of a material’s VB is more positive than the water oxidation level (1.23 eV versus NHE, normal hydrogen electrode), while the lowest level of the CB should be more negative than the H₂ evolution.

Figure 1.6: Calculated free energy diagram for H₂ evolution at a potential U = 0 relative to the standard H₂ electrode at pH = 0. Reprinted with permission from Ref 20. Copyright 2005 American Chemical Society.
potential (0 eV vs NHE, Figure 1.7). Therefore, the minimum band gap for a suitable water splitting photocatalyst should be 1.23 eV. It should be mentioned that the water oxidation half reaction involves the transfer of four electrons and is more energetically demanding than the water reduction process (minimum potential of 1.23 V per electron transfer).\textsuperscript{27} In addition, since the Gibbs energy for overall water splitting is positive (\( \Delta G > 0 \)), it is challenging to avoid the back reaction.\textsuperscript{28} Consequently, the oxidation half-reaction represents the bottleneck of overall water splitting. Thus, sacrificial electron donors are usually employed, obstructing the \( \text{O}_2 \) evolution reaction, to promote the \( \text{H}_2 \) generation.\textsuperscript{29}

\[ \begin{align*}
\text{H}_2^+ + \text{e}^- & \rightarrow \text{H}_\text{ads} \\
\text{H}_2^+ + \text{H}_2 & \rightarrow \text{H}_2
\end{align*} \]

\[ \begin{align*}
\text{H}_2^+ + \text{H}_2 & \rightarrow \text{H}_2
\end{align*} \]

\[ \begin{align*}
\text{H}_2^+ + \text{H}_2 & \rightarrow \text{H}_2
\end{align*} \]

**Figure 1.7:** Energy diagrams of photocatalytic water splitting relative to NHE, at pH = 0, with the \( \text{H}_2 \) evolution and \( \text{O}_2 \) evolution reactions redox potentials.

On the other hand, the \( \text{H}_2 \) evolution half reaction involves two electrons and can be expressed through three possible steps. The first step is well established and known as the Volmer step, whereby in acidic media a proton reacts with an electron to form atomic hydrogen adsorbed on the surface of the photocatalyst (\( \text{H}^+ + \text{e}^- \rightarrow \text{H}_\text{ads} \)) or in an alkaline environment, a water molecule reacts with an electron to form atomic hydrogen and also hydroxide (\( \text{H}_2\text{O} + \text{e}^- \rightarrow \text{H}_\text{ads} + \text{OH}^- \)). The second part of this reaction can involve the Heyrovsky step as well as the Tafel step. According to the Heyrovsky step, the adsorbed atomic hydrogen will react with an electron and either a proton to form \( \text{H}_2 \) in acidic media (\( \text{H}_\text{ads} + \text{H}^+ + \text{e}^- \rightarrow \text{H}_2 \)), or with a water molecule to form \( \text{H}_2 \) and hydroxide anions in alkaline media (\( \text{H}_\text{ads} + \text{H}_2\text{O} + \text{e}^- \rightarrow \text{H}_2 + \text{OH}^- \)). The Tafel step is the same in both acidic and alkaline environments and involves the combination of two adsorbed atomic hydrogens to form \( \text{H}_2 \) (\( \text{H}_\text{ads} + \text{H}_\text{ads} \rightarrow \text{H}_2 \)).

In addition to the photocatalyst, the following auxiliary components can assist a photocatalytic \( \text{H}_2 \) evolution system (Figure 1.8):
i) Co-catalysts are employed in most cases to promote the photocatalytic activity. After the photoexcitation of the photocatalyst, the presence of both electrons and holes in the material can result in the recombination of these charge carriers. This phenomenon is highly correlated with the excited state lifetime of the photocatalyst and is the key reason why the efficiencies of such reactions can be low. The co-catalysts can significantly increase the H\textsubscript{2} evolution reaction yield, since they serve a two-fold purpose, inhibiting the undesired recombination of the photo-generated charge carriers and reducing the activation energy for the H\textsubscript{2} evolution reaction.\textsuperscript{21a} There are innumerable examples of noble metal-based (e.g. Pt, Pd, Au),\textsuperscript{30} transition-metal-based (e.g. Co, Ni, MoS\textsubscript{2}, Ni\textsubscript{2}P, CoP, CuO),\textsuperscript{31} nanocarbon-based (e.g. graphene, carbon nanotubes)\textsuperscript{32} and molecular-based (e.g. cobaloximes, Mo\textsubscript{3}S\textsubscript{13}\textsuperscript{2-})\textsuperscript{33} co-catalysts. Among them, Pt is considered the paradigm of efficient co-catalysts for H\textsubscript{2} production due to its low overpotential for this reaction and the suitable Fermi level for withdrawing the photoexcited electrons.\textsuperscript{34}

ii) Photosensitizers can also be utilized to extend or enhance the light absorption of the photocatalyst in the visible region (44\% of the solar energy), through electron transfer from their antibonding to the CB of the photocatalyst. Typical examples of photosensitizers are organic dyes, such as eosin Y,\textsuperscript{35} [Ru(bpy)\textsubscript{3}]\textsuperscript{2+},\textsuperscript{36} or porphyrins.\textsuperscript{37}

iii) Reagents such as electron acceptors (e.g. methyl viologen cation, MV\textsuperscript{2+})\textsuperscript{38} or sacrificial electron donors (triethylamine, triethanolamine, ethanol, methanol) can be used to either mediate the transfer of the photogenerated electrons or balance the H\textsubscript{2} evolution half reaction, respectively. In particular, electron donors are regularly used since they can remarkably enhance the photocatalytic performance.

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**Figure 1.8**: Schematic illustration of the possible constituents of a photocatalytic H\textsubscript{2} evolution system.
1.4 Photocatalytic Oxidation of Organic Compounds

Although industrial development has engendered countless advancements in technology, amenities in lifestyle and rise in the economy, it has also inflicted extended deterioration of the environment. One of the most critical environmental problems faced over the last century is water pollution, mainly caused by human activities. Textile or food processing industries, pharmaceutical companies, agricultural activities, and power plants are some of the main sources of hazardous contaminants. Wastewater contains a variety of organic pollutants, such as organic dyes, phenols, fertilizers and hydrocarbons, which can cause incontrovertible environmental problems. A large fraction of these contaminants is mutagenic, carcinogenic, or toxic and their presence poses significant risks for public health. In addition, the vast majority of these organic substances can damage aquatic ecosystems and inhibit the exchange of \( \text{O}_2 \) in water. Moreover, severe water scarcity is projected to impact a quarter of the world’s population within the next few years. Thus, increasing generation of wastewater along with the water shortage projected in the near future has provoked the urgent development of methodologies that can offer effective decontamination of water so as to enable an efficient recycling of wastewater.

So far, conventional water treatment methods (e.g. coagulation, sedimentation, filtration, adsorption) have been characterized by high operating costs, as well as the fact that they can generate other undesired hazardous by-products, or they cannot fully degrade the targeted organic pollutants. Another option to decompose these organic substances from water is through advanced oxidation processes (AOPs), which are based on the generation of strongly oxidizing radical species (e.g. \( \cdot \text{OH} \)) that can efficiently and non-selectively destruct even the most resistant organic species. The AOPs include a range of techniques, such as wet, electrochemical or super critical water oxidation, ozonation, hydrogen peroxide-based methods, and photolysis. In particular, heterogeneous photocatalytic oxidation is a promising approach that can lead to complete degradation of a wide range of hazardous organic substances into easily biodegradable compounds or less toxic molecules, by utilizing light.

It is widely known that \( i. \) the holes (\( h^+ \)), \( ii. \) the hydroxyl radicals (\( \cdot \text{OH} \)) and \( iii. \) the superoxide radicals (\( \text{O}_2^- \)) are the three main active species involved in photocatalytic oxidation processes (Figure 1.9). As mentioned above, upon irradiation of a photocatalyst, generation of excited high-energy states of electron and hole pairs occurs. The hole in the VB is a powerful oxidizing agent that can react either directly with organic molecules and successfully oxidize them to less hazardous products such as \( \text{CO}_2 \) and \( \text{H}_2\text{O} \), or it can react with water to generate highly reactive hydroxyl radicals (\( \cdot \text{OH} \)) (equation 1).
Moreover, the promoted electron in the CB (e\textsuperscript{−}) can initiate oxygen ionosorption by reacting with O\textsubscript{2} to form an anion superoxide radical (O\textsubscript{2}\textsuperscript{•−}) (equation 2). The latter can further react with hydrogen cations (H\textsuperscript{+}) and be protonated to the hydroperoxyl radical (HOO\textsuperscript{•}), which acts as an e\textsuperscript{−} scavenger, being reduced to HOO\textsuperscript{−}, elongating the lifetime of the excited state and subsequently being protonated to hydrogen peroxide (H\textsubscript{2}O\textsubscript{2}).\textsuperscript{4a}

\[
h^+ + \text{H}_2\text{O} \rightarrow \cdot\text{OH} + \text{H}^+ \quad \text{(equation 1)}
\]

\[
e^- + \text{O}_2 \rightarrow \text{O}_2\textsuperscript{•−} \quad \text{(equation 2)}
\]

Figure 1.9: Schematic representation of the photoexcitation of a photocatalyst in the presence of organic pollutants. The mechanism for the generation of anion superoxide and hydroxyl radicals, as well as that for the direct organic pollutant’s oxidation is also illustrated.

Titanium dioxide (TiO\textsubscript{2}) has been considered a very promising photocatalyst for the mineralization of wastewater as well, and has thus been extensively studied.\textsuperscript{48} However, its incapability of operating under visible light restricts its utilization in practical applications. Therefore, immense research endeavours have focused on extending its absorption into the visible light region.\textsuperscript{49}

In contrast with photocatalytic H\textsubscript{2} evolution systems, photocatalytic organic degradation systems are usually less complicated as they usually consist of less components. In the majority of examples reported in the literature, only a photocatalyst is used to harvest the light. It should be mentioned that the photocatalytic oxidation of organic substances is commonly performed under aerated conditions, where O\textsubscript{2} acts as an electron scavenger, capturing the photogenerated electrons and being transformed
to superoxide radicals, which can promote the degradation process. Furthermore, additional electron scavengers are often included, such as H$_2$O$_2$, persulfate (PS), and peroxymonosulfate (PMS), to further suppress the electron-hole pair recombination, thus enhancing the photodegradation efficiency.\textsuperscript{50} In contrast to the photocatalytic H$_2$ evolution field, co-catalysts are not as frequently used for this application.\textsuperscript{47,51}

In the field of photocatalytic degradation of organic substances, parameters such as the pH of the solution and the initial concentration of the pollutant can strongly influence the photocatalytic efficiency.\textsuperscript{52} In particular, variations in the pH can have a profound impact on the photocatalytic degradation rate, since it is a critical parameter for the adsorption of a targeted compound on the surface of the photocatalyst. This is of paramount importance, especially when the photocatalyst is amphoteric (e.g. TiO$_2$) and consequently, the charge of its surface is pH-dependent.\textsuperscript{53} Therefore, it is crucial that these parameters are varied and optimized to obtain legitimate and comparable results.

1.5 Dual-Functional Photocatalysis

Dual-functional photocatalysis constitutes a hybrid field where different photocatalytic fields are combined for a two-fold purpose. In particular, the photocatalytic generation of H$_2$ coupled with the simultaneous photocatalytic oxidation of organic substances has attracted radically increasing research interest over the last few years.\textsuperscript{54} This field can be divided in two subcategories depending on the nature and target of the oxidation process. Both of these categories involve the photocatalytic generation of H$_2$ from water (as the main source) coupled with either the degradation of organic pollutants or the transformation of organic substances into value added products (Figure 1.10).

The utilization of organic substrates for H$_2$ evolution through photocatalysis was introduced in the late ‘70s and early ‘80s,\textsuperscript{55} when the conversion of carbohydrates to H$_2$ was explored, commonly known as photoreforming.\textsuperscript{56} Similar to dual-functional photocatalysis, photoreforming focuses on the production of H$_2$ by utilizing organic substrates. However, in photoreforming, the sacrificial organic molecules to be oxidized are not necessarily hazardous substances and can be useful carbohydrates, such as methane and ethanol.\textsuperscript{57} In addition, in the field of dual-purpose photocatalysis, the H$_2$ should be mainly derived from the water reduction reaction. For example, earlier this year, Zhang \textit{et al.} reported the photocatalytic dehydrogenation of benzyl alcohol for the simultaneous H$_2$ and benzaldehyde production using a Ni-decorated Zn$_{0.5}$Cd$_{0.5}$S photocatalytic system.\textsuperscript{58} However, in this work, the authors attributed the H$_2$ generation to its extraction from benzyl alcohol and thus, it cannot be considered a paradigm of dual-functional photocatalytic activity. Consequently, although these two
fields can overlap, only a fraction of studies in the photoreforming field can be considered dual-functional photocatalysis, which is when either the targeted organic molecules are pollutants or when the aim of the oxidation is the synthesis of value-added products (e.g., aldehydes, furoic acid or imine).

Figure 1.10: Schematic representation of the mechanism for dual-functional photocatalysis for H$_2$ generation coupled with oxidation of organic pollutants to either achieve full degradation of hazardous organic contaminants or to synthesize useful organic molecules.

Even though H$_2$ evolution and organic oxidation have been studied separately in photocatalytic systems,$^{59}$ their integration in a single process is relatively new. The basic principles of the individual photocatalytic fields are combined, generating new mechanistic aspects and criteria for the selection of appropriate dual-functional photocatalysts. Desired characteristics of a dual-purpose photocatalytic system include those applied in general photocatalysis, comprising low cost, extended solar light absorption, prolonged excited state lifetime and high electron-hole separation efficiency. In addition, a dual-purpose photocatalyst for H$_2$ production coupled with organic oxidation should have VB and CB with suitable energy and be capable of oxidizing the organic molecules with holes as the dominant active species and with the electrons being transferred to water molecules or protons. Since O$_2$ is a good electron scavenger, almost every study in the field is carried out under anoxic conditions.


1.6 Metal-Organic Frameworks for Photocatalysis

Metal-organic frameworks (MOFs) represent an interesting type of porous, crystalline materials, that can be self-assembled through the coordination of organic linkers with metal ions/clusters (Figure 1.11). A wide variety of MOF structures has been reported since the term ‘metal-organic-frameworks’ was first conceived by Yaghi in 1995. Owing to their modular nature, large porosity and high surface area (up to 7000 m$^2$/g), MOFs are considered promising candidates for a variety of applications, such as gas storage and separation, magnetic materials, chemical sensing, catalysis, and biomedicine.

![Figure 1.11: Schematic representation of MOFs’ constituents.](image)

The understanding and elucidation of the main optical and vibrational properties of MOF-5 represent one of the milestones in the field, paving the way for MOFs’ implementation in light-related applications. In fact, MOFs have burst onto the light-driven catalysis field only a decade ago, and have since been considered promising candidates for photocatalytic applications, including H$_2$ evolution, organic pollutants degradation, CO$_2$ and metal ions reduction. On the basis of a strict definition, MOFs cannot be regarded as semiconductors, which is the traditional type of materials for light-driven catalytic applications. However, they can exhibit a semiconductor-like behavior upon irradiation, deriving from their optically active ligands and their electronic interactions with the metal nodes. Upon illumination of a light-responsive MOF, an electron will be excited from the highest occupied crystalline orbitals (HOCOs) to the lowest unoccupied crystalline orbitals (LUCOs), analogous to the VB and CB in the band theory of solids, or the bonding and antibonding orbitals in the molecular orbital theory.

The main factor triggering MOFs’ exploration in photocatalytic applications lies in the limitations of the traditional materials (photocatalysts) that have been predominantly employed in this technology. Especially for solar-driven photocatalysis, the bottleneck of such technologies is related to the
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shortcomings of the traditional semiconductor photocatalysts. More specifically, typical wide band-gap semiconductors (i.e., TiO$_2$) suffer from limited sunlight harvesting, while the ones characterized by narrow band-gap (i.e., a-Fe$_2$O$_3$, BiVO$_4$, CdS) demonstrate high charge carrier recombination, poor surface kinetics or instability.$^{67-68}$

MOFs are endowed with extremely high structural versatility, which allows a level of tunability of the optical, electronic and catalytic properties.$^{69}$ By carefully selecting light-absorbing linkers and catalytically active metals with comparable energy of their frontier orbitals, one can design numerous MOFs that would be excellent candidates for photocatalytic applications. Therefore, the tremendous scientific attention received by MOFs in the general catalysis field can be anticipated, considering that one can target and tune the desired features of a MOF through careful choice of ligand and metal. In other words, MOFs offer the exciting possibility to combine the advantages of homogeneous catalysts – such as high tunability – with those of heterogeneous catalysts; robustness and insolvency.

1.6.1 MOFs stability under water splitting conditions

Water stability is a crucial property for many materials to be industrially applicable, since water is abundant in the preparation, storage, transportation and application processes. Especially when it comes to water splitting, it is indispensable for the MOFs to demonstrate stability under aqueous conditions, as well as being resistant to acidic or basic conditions (e.g. water reduction in the presence of sacrificial donors).$^{70}$ The robustness of a given MOF toward water depends on the strength of the bonds between the metal and the oxygen or nitrogen atoms from the ligand comprising it. Unfortunately, due to the lability of ligand-metal bonds, a considerable amount of reported MOFs are sensitive to water content, which results in their gradual decomposition when the environment contains moisture. $^{71}$

Water stable MOFs by definition are classified as those that do not exhibit structural breakdown under exposure to water content. In order to determine if a MOF structure remains stable in a water stability test, we can compare some of their typical characteristics such as crystallinity (powder X-ray diffraction) or structural porosity (gas adsorption capacity) between the pristine and post-exposure to water samples. During the last few years, plenty water-stable MOFs has been reported, mainly due to the improved understanding toward MOF structural stability in water. These water stable MOFs could be categorized into three groups: $i.$ metal carboxylate frameworks consisting of high-valence metal ions (e.g. Ti$^{4+}$, Fe$^{3+}$, Cr$^{3+}$, and Zr$^{4+}$), such as the Ti-based MIL-125, and the Cr-based MIL-101$^{73}$ and the Zr-based UIO-66, $^{74} ii.$ metal azolate frameworks containing nitrogen-donor ligands, such as the MAF
(metal azolate frameworks) series of materials developed by Chen’s group,\textsuperscript{75} iii. MOFs functionalized by hydrophobic pore surfaces or with blocked metal ions, such as CALF-25.\textsuperscript{76} These MOFs are likely to retain their structure under the exhaustive water splitting conditions.

### 1.6.2 MOFs’ Optical Response

One of the key advantages of MOFs is their tunable optical properties deriving from the rational design through careful choice of the ligand and metal comprising it. The overall optical behavior of a MOF is governed by the optical properties of the metals and ligands composing it, and their interactions. Similar energy of the ligands’ and metals’ frontier orbitals and spatial overlap between them can give rise to charge transfer between those two constituents and a smaller optical gap of the MOF comprising them, allowing for light absorption in the visible and infrared, which are the dominant regions of the solar spectrum. Charge transfer between those two components can occur either from the ligand’s orbitals to those of the metal (Ligand to Metal Charge Transfer, LMCT) or vice versa (Metal to Ligand Charge Transfer, MLCT). On the other hand, limited or no energetic and spatial overlap of the linkers’ and metals’ orbitals results in the light harvesting properties of the MOF being similar to those of the individual free ligand and metal comprising it.\textsuperscript{77} The presence or absence of such electronic interactions between the ligand and metal of a given MOF can be explored through UV-Vis absorption spectroscopy, while electron paramagnetic resonance (EPR) could be a useful tool to shed light into the charge transfer mechanism (LMCH or MLCT), through observation of the potential change in the oxidation state of the metal.

As mentioned before, MOFs have the ability to behave as semiconductors, with their electrons being excited from their VB (or HOCOs) to their CB (or LUCOs) upon exposure to light with appropriate energy. However, a great number of MOFs exhibit light absorption only in the UV-region, while undoubtedly one of the main challenges in photocatalysis lies in the ability of the photocatalyst to exploit sunlight. Several approaches have been investigated in order to tackle this issue, including the utilization of light absorbing organic and metal-organic chromophores as building units for the synthesis of MOFs,\textsuperscript{78} doping different photosensitizers into the cavities of MOFs,\textsuperscript{79} or linker functionalization with electron donating functional groups, such as -NH\textsubscript{2} or -OH.\textsuperscript{80}

### 1.6.3 MOF-based Photocatalytic Systems for Water reduction

Similar to the traditional semiconductor photocatalysis field, MOF-based photocatalytic system for the H\textsubscript{2} evolution reaction can consist of several components in addition to the MOF, which is the main
component and typically acts as an antenna harvesting light. So far, electron donors are always used in such systems, since to our knowledge, currently there are no reports of MOF photocatalysts splitting pure water into H\textsubscript{2} and O\textsubscript{2}, and these substances are used to avoid the water oxidation reaction. Recently, Ziao et al. reported a Cd-based MOF photocatalyst, Cd-TPAPy – comprised of 1,3,6,8-tetrakis(p-benzoic acid)pyrene (H\textsubscript{4}TBAPy) as the ligand – that is capable of catalyzing both the water reduction and oxidation reaction with the assistance of Pt or CoPi as co-catalysts.\textsuperscript{81} However, the authors did not investigate the photocatalytic activity of Cd-TPAPy for both the water reduction and oxidation reactions simultaneously, but explored these two reactions separately, using a sacrificial electron donor (triethanolamine, TEOA) and an electron scavenger (AgNO\textsubscript{3}) for the H\textsubscript{2} and O\textsubscript{2} production, respectively (Figure 1.12).

**Figure 1.12: Schematic illustration of the proposed mechanism for photocatalytic H\textsubscript{2} and O\textsubscript{2} evolution over Cd-TBAPy. Reprinted with permission from Ref. 81 Copyright © 2018 John Wiley and Sons.**

In addition to electron donors, another type of sacrificial reagent that could be used is a redox shuttle, mediating the electron transfer from the photocatalyst to the co-catalyst. However, their utilization is far less frequent compared to electron donors. In this context, Fateeva et al. designed and investigated a MOF-based on porphyrin linkers and aluminum (known as AlPMOF), using ethylenediaminetetraacetic acid (EDTA) as the electron donor, Pt nanoparticles as the co-catalyst, with and without methyl viologen (MV) as the electron mediator.\textsuperscript{82} AlPMOF represents one of the very early examples of MOF photocatalysts and demonstrates visible-light absorption similar to its porphyrin ligand. Although this system exhibited adequate photocatalytic activity, the actual rate of hydrogen production was lower when MV was present, which was attributed to diffusion limitations of MV in the pores of the MOF.
Dyes can also be incorporated in MOF-based photocatalytic systems for water reduction, acting as photosensitizers. Similar to the dye sensitized solar cells in which a ruthenium-based dye is typically adsorbed on the surface of TiO$_2$, in MOF photocatalytic systems, upon illumination the dye absorbs light – ideally in the region where the MOF is incapable of absorbing. Then, the photoexcited electron is transferred from the antibonding orbitals of the dye to the LUCOs/CB of the MOF from which it can be involved in the reduction reaction process. Interestingly, in MOF-based systems, the dye molecules can be adsorbed not only on the exterior surface of the MOF but also within its pores, increasing the amount of dye molecules able to electronically interact with the MOF. Dyes are often used in such systems, especially in the cases where the MOF absorbs light only in the UV region. A typical example of UV-active MOF photocatalysts is the Zr-based UIO-66, synthesized by Cavka et al. at the University of Oslo. Considering that UIO-66 represents one of the most stable photocatalysts for this type of materials, there are several photocatalytic studies in the literature showing that UIO-66 can be assisted by dyes (e.g., Rhodamine B and Erythrosin B) for the absorption of light in the visible region. More specifically, He et al. explored the photocatalytic activity of UIO-66 with Pt as the co-catalyst, TEOA as the electron donor, with and without Rhodamine B as the photosensitizer. Under visible light irradiation – in which UIO-66 is mainly inactive – the Pt/UIO-66 photocatalytic system demonstrated 30-fold greater activity when Rhodamine B was used (116 versus 3.9 μmol g$^{-1}$ h$^{-1}$ with and without Rhodamine B, respectively). The proposed mechanism is illustrated at Figure 1.13 and involves electron transfer first, from the antibonding orbitals of Rhodamine B to the LUCOs/CB (denoted as LUMO at Figure 1.13) of UIO-66 and subsequently electron migration to the Pt NPs, where protons are reduced to H$_2$.

![Figure 1.13: Schematic representation of proposed mechanism for photocatalytic H$_2$ generation over Rhodamine B (RhB) sensitized Pt/UiO-66 under visible-light. Reproduced from Ref. 84 with permission from The Royal Society of Chemistry.](image)
Another important component frequently used in MOF-based photocatalytic systems is the co-catalyst, in order to inhibit the undesired electron-hole recombination and offer catalytic active sites for the reaction to occur. So far, the vast majority of MOF photocatalysts reported are combined with expensive noble metal nanoparticles, such Pt.\textsuperscript{85} Due to their high porosity, MOFs offer the possibility to incorporate the co-catalysts within their cavities. In this context, Xiao \textit{et al}. investigated the impact of the co-catalyst’s location relative to the MOF photocatalyst, on the overall activity of the system.\textsuperscript{85a} More specifically, they supported Pt nanoparticles either inside the porous assemblies of UIO-66-NH$_2$ or on its surface (Figure 1.14). The group showed that encapsulating the Pt co-catalysts within the pores of UIO-66-NH$_2$ results in higher photocatalytic efficiency of 257.38 (versus 50.26 µmol g$^{-1}$ h$^{-1}$ for UIO-66-NH$_2$ with Pt nanoparticles simply deposited on the surface). This enhancement of the photocatalytic activity was attributed to the shorter electron-transport paths between the co-catalyst and the photocatalysts, further promoting the charge separation efficiency.

![Figure 1.14: Schematic representation of the synthesis procedure of UiO-66-NH$_2$ with encapsulated Pt NPs (Pt@UiO-66-NH$_2$) and of Pt NPs deposited on the exterior surface of the MOF (Pt/UiO-66-NH$_2$), with the mechanism for photocatalytic H$_2$ production over Pt@UiO-66-NH$_2$ being highlighted. Reprinted with permission from Ref. 85a Copyright © 2016 John Wiley and Sons.](image)

Summarizing, these extra components typically incorporated in MOF-based photocatalytic systems for H$_2$ evolution, are significantly increasing the overall cost and complexity of such systems. Furthermore, some of these components – such as dyes and electron donors – can be toxic and hazardous (e.g., Rhodamine B, triethylamine, TEOA and methanol). To overcome these disadvantages, several challenging issues need to be tackled. First, MOFs that exhibit strong and extensive light absorption must be developed to avoid the use of dyes as photosensitizers. Second,
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abundant photocatalysts that exhibit high electron-hole separation efficiencies or noble-metal-free cocatalysts are highly desired materials and, hence, research efforts should focus on this aspect. Finally, the incorporation of highly toxic or expensive sacrificial electron donors can be avoided by either developing photocatalysts capable of splitting water into H₂ and O₂ or able to perform dual-functional photocatalysis, for simultaneous production of H₂ and oxidation of targeted organic compounds (e.g., degradation of organic contaminants or synthesis of useful chemicals).

1.7 Objectives of this Thesis

In line with the scientific endeavors to optimize and extend the range of materials used for photocatalytic applications, the main objective of this thesis lies in the optimization of MOF-based photocatalytic systems. In an ideal case, the photocatalyst should demonstrate extended solar light absorption, high electron-hole separation and migration efficiency, and high catalytic activity. In fact, these desired features are correlated to the three major photocatalytic steps (as explained in Chapter 1.2). It is thus the main focus of this thesis to improve the photocatalytic activity of MOF-based photocatalytic systems, by exploring and promoting these three major steps of i. light harvesting, ii. charge separation and transport and iii. catalytic redox reactions. More specifically, Chapter 2 highlights how using abundant co-catalysts can modify the last two photocatalytic steps of charge separation and catalytic reaction, and thus significantly improve the photocatalytic activity and lower the cost of MOF-based systems. In Chapter 3, first, the influence of different electron donors on the activity of the abundant MOF-based system described in the previous chapter is investigated. The second part of Chapter 3 shows that the overall cost and sustainability of such systems can be further improved by replacing the electron donors with organic contaminants, performing dual-functional photocatalysis for concurrent H₂ production and pollutants degradation. Chapter 4 describes a strategy to promote the first and second photocatalytic steps of light absorption and charge separation, which lies in the design of MOF/MOF heterojunctions. This strategy exploits the advantages of MOFs as photocatalysts in their maximum potential (e.g. high tunability of the optoelectronic and catalytic properties), and simultaneously tackle their common challenges (e.g. high charge recombination rates and susceptibility to photocorrosion). Finally, Chapter 5 focuses on an alternative use of MOFs in the photocatalysis field, where the MOF is used as a precursor for the synthesis of more active semiconductor photocatalysts with interesting morphology.
Chapter 2 Variation of Abundant Co-catalysts in a Metal-Organic Framework Photocatalytic System

Parts of this chapter are based on published works:

‘Photocatalytic Hydrogen Generation from a Visible-Light Responsive Metal-organic framework system: the impact of nickel phosphide nanoparticles’

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and

‘Concurrent Photocatalytic Hydrogen Generation and Dye Degradation Using MIL-125-NH₂ under Visible Light Irradiation’

S. Kampouri, T. N. Nguyen, M. Spodaryk, R. G. Palgrave, A. Züttel, B. Smit, K. C. Stylianou,
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S. Kampouri’s contribution: Conceiving the project’s idea, synthesis of materials, characterization (powder X-ray diffraction, UV-Vis absorption and photoluminescence spectroscopy, nitrogen sorption isotherms, Fourier-transform infrared measurements), designing and performing experiments (photocatalytic experiments, quantum yield calculations), writing text and constructing figures.
2.1 Introduction

Several MOF-based photocatalytic systems have been tested for visible-light driven H₂ production; most of them are based on reductive metal ions, such as Ti⁴⁺ and Zr⁴⁺ and visible-light-active ligands that can harvest solar light and convert it into H₂. Recently, Cu¹ or Cu²⁺-based MOFs were found to be good candidates for the photocatalytic water reduction reaction under UV/vis or visible irradiation. Despite the great promise of MOFs toward water splitting, the majority of these MOF-based systems demonstrate very low apparent quantum yields and still utilize expensive noble-metal-based co-catalysts, such as Pt nanoparticles (NPs).

The broad utilization of Pt NPs as co-catalysts – especially with MOF-based systems – lies in their low overpotential for the H₂ evolution reaction, their suitable Fermi level for withdrawing photoexcited electrons and their facile preparation. However, their incorporation does not guarantee the best photocatalytic performance, while with dwindling supplies of precious metals it is crucial to eliminate their utilization.

Although very few examples have been reported in the literature, the synergy between MOFs and co-catalysts is proven to be a key factor for the photocatalytic performance of the system, as different hydrogen generation rates and quantum yields are observed when the same MOF is used. Key factors for achieving optimal synergy include energetic alignment and kinetic compatibility between the MOF photocatalyst and the co-catalyst, so as to ensure high electron transfer rates. Consequently, there is a need to investigate the activity of earth-abundant co-catalysts – as alternatives to noble metals – and explore their interactions with a given MOF. It is worth noting that since the number of potential earth-abundant metal oxides or phosphides is essentially limitless, it should be possible to boost the H₂ generation performance of a MOF-based photocatalytic system by simply choosing the appropriate co-catalyst.

This chapter describes the investigation of different transition metal-based co-catalysts with MIL-125-NH₂. MIL-125 is the first titanium-based MOF reported, consisting of a titanium-oxo cluster and terephthalate (BDC) as the organic linker. Replacing this linker with 2-aminoterphthalate (NH₂-BDC) results in the same structure (MIL-125-NH₂), but with visible light absorption as opposed to the UV offered by MIL-125, making it a promising photocatalyst. Variation of the co-catalysts alters the second and third major photocatalytic steps of charge separation and catalytic reaction, which can significantly promote the overall activity of the MIL-125-NH₂-based photocatalytic system. Indeed, all the co-catalysts (Ni₂P, NiO, CoP, Co₃O₄, Fe₂O₃, CuO and Pt nanoparticles) investigated were found to significantly improve the activity of MIL-125-NH₂, with the Ni₂P/MIL-125-NH₂ system demonstrating the highest catalytic activity.
2.2 Synthesis and Characterization

MIL-125-NH$_2$ was synthesised based on a reported procedure (see section S2.1, Appendix)$^{94}$ and subsequently characterised by means of powder X-ray diffraction (PXRD), nitrogen adsorption-desorption isotherms, scanning electron microscopy (SEM) and UV-vis absorption spectroscopy. The PXRD pattern of the bulk MIL-125-NH$_2$ is in great agreement with the pattern derived from the single-crystal structure and the nitrogen isotherms revealed the microporous nature of the MIL-125-NH$_2$, with a BET surface area of ~1200 m$^2$ g$^{-1}$ (Figure 2.1A, B). The MIL-125-NH$_2$ crystals display a circular-disc-type morphology with a mean particle size of around 800 nm, as illustrated by the SEM images (Figure 2.1C). The UV-vis absorbance spectrum of the free ligand NH$_2$-H$_2$BDC and the Kubelka-Munk representation of the UV-vis diffuse reflectance spectrum of MIL-125-NH$_2$ (Figure 2.1D), indicate that the latter exhibits a red shift. This red shift leads to a strong absorption in the 330-500 nm range and is ascribed to the ligand-to-metal charge transfer (LMCT).$^{86}$

**Figure 2.1:** (A) PXRD patterns, (B) $N_2$ adsorption-desorption isotherm and (C) SEM image of the as-synthesized MIL-125-NH$_2$. (D) UV-Vis absorbance of free ligand H$_2$BDC-NH$_2$ and Kubelka-Munk representation of diffuse reflectance of MIL-125-NH$_2$. 

− 23 −
Variation of Co-catalysts

Stavroula A. Kampouri

A range of transition metal phosphide and oxide co-catalysts (Ni<sub>2</sub>P, NiO, CoP, Co<sub>3</sub>O<sub>4</sub>, Fe<sub>2</sub>O<sub>3</sub> and CuO) was selected, based on their availability and low cost compared to noble-metal nanoparticles, which are commonly used in photocatalytic water splitting. The selected co-catalysts were synthesised based on a MOF-derived technique (Figure 2.2). When this synthetic method is used, the self-sacrificing MOF acts as a structure directing template allowing for a more rational preparation of NPs. Specifically, the use of MOFs as precursors can lead to the generation of nano-sized materials, with higher specific surface areas, thermal stability and catalytic activity. For example, Yu et al. developed an efficient electrocatalyst through an easy and scalable one-pot thermal treatment of bimetallic zeolitic imidazolate frameworks (ZIFs). For more information about the synthetic protocols employed see section S2.1 in the Appendix.

![Figure 2.2: Schematic illustration of the synthetic procedure employed for the generation of transition metal oxide and phosphide NPs.](image)

The formation, phase purity and morphology of the MOF-derived NPs were explored through PXRD, nitrogen sorption isotherms and transmission electron microscopy (TEM). The PXRD patterns of the as-synthesised NiO, Ni<sub>2</sub>P, Co<sub>3</sub>O<sub>4</sub>, CoP, Fe<sub>2</sub>O<sub>3</sub> and CuO NPs are in good agreement with the simulated and previously reported patterns (Figure 2.3). The TEM images revealed the morphology and size of these NPs; the NiO, Ni<sub>2</sub>P, Co<sub>3</sub>O<sub>4</sub>, and CuO NPs illustrate a spherical shape with a mean size of 10 - 20 nm, whereas Fe<sub>2</sub>O<sub>3</sub> NPs demonstrate a size of 200 nm (Figure 2.4). The dimensions of the rod-shaped CoP NPs are approximately 20 nm x 6 nm, which is in agreement with previous reports of MOF-derived CoP NPs.

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Figure 2.3: PXRD patterns of MOF-derived NiO, Fe$_2$O$_3$, Co$_3$O$_4$, CoP, Ni$_2$P and CuO NPs.

Figure 2.4: High-resolution TEM images of: (A) NiO, (B) Fe$_2$O$_3$, (C) Co$_3$O$_4$, (D) CoP, (E) Ni$_2$P and (F) CuO NPs.
2.3 Photocatalytic Activity

In order to assess the photocatalytic performance of the different co-catalyst/MIL-125-NH$_2$ systems, the crystalline powder of MIL-125-NH$_2$ was physically mixed with varying amounts of the co-catalyst NPs. Such simple incorporation of the co-catalyst (physical mixture) is advantageous, considering that complicated steps are prevented (e.g. photodeposition$^{90a, 101}$ or encapsulation$^{102}$) and the loaded amounts of the co-catalysts are highly controllable and reproducible. The ratio between the co-catalyst and the MIL-125-NH$_2$ amount is crucial, since there is a volcano-type trend between the added amount of a given co-catalyst and the photocatalytic activity of a system. In principle, the addition of the co-catalyst increases the production of H$_2$, since these NPs play a key role in facilitating the electron-hole separation within the MIL-125-NH$_2$ (Figure S.2). However, further increase in the amount of the co-catalyst decreases the photocatalytic performance of the system. This is attributed to the fact that an excess amount of the co-catalyst leads to restricted penetration of the incident light in the photocatalytic solution, and can hinder the contact between MIL-125-NH$_2$ and the electron donor.$^{21a}$

The photocatalytic solution used for these experiments includes acetonitrile (CH$_3$CN), triethylamine (TEA) as the electron donor and water in a volumetric ratio of 79: 16: 5 v/v/v, respectively. The optimal amounts of each co-catalyst (i.e., the amount that produced the highest H$_2$ evolution rate) varies within the range of 7.9 - 10.2 wt.% (Figure S.2). Interestingly, as illustrated in Figure 2.5, the optimised Ni$_2$P/MIL-125-NH$_2$ photocatalytic system displays the highest hydrogen evolution rate of 1230 μmol h$^{-1}$ g$^{-1}$ (over 8 h), followed by NiO/MIL-125-NH$_2$, which exhibits a rate of 1084 μmol h$^{-1}$ g$^{-1}$. The performances of the Co$_3$O$_4$/MIL-125-NH$_2$ and CoP/MIL-125-NH$_2$ systems are comparable. The optimised Fe$_2$O$_3$/MIL-125-NH$_2$ system exhibits a H$_2$ evolution rate of 435 μmol h$^{-1}$ g$^{-1}$, followed by the CuO/MIL-125-NH$_2$ system which demonstrates the lowest rate of 139 μmol h$^{-1}$ g$^{-1}$. It is worth mentioning that when only the co-catalysts were subjected to photocatalytic test, only traces of H$_2$ were detected (Table S.1). Likewise, without any co-catalysts, MIL-125-NH$_2$ exhibits a very low H$_2$ evolution rate of 51.2 μmol h$^{-1}$ g$^{-1}$. The significantly inferior performance of solely the MIL-125-NH$_2$ highlights that the incorporation of the co-catalysts in the system remarkably promotes the H$_2$ evolution rate.

Inspired by these results, we wanted to evaluate the efficiency of the best performing Ni$_2$P/MIL-125-NH$_2$ using a metric based on the incident radiation. For that, we calculated the apparent quantum yield (AQY) at 450 nm through Ferrioxalate actinometry (see section S2.3, Appendix).$^{103}$ Ni$_2$P/MIL-125-NH$_2$ demonstrates an AQY value of 6.6 %, which is among the highest reported efficiencies for MOFs$^{69b, 90b, 104}$ and comparable to leading photocatalysts (Table S.2).$^{105}$
Figure 2.5: Comparison of H₂ evolution rates (with respect to MIL-125-NH₂) of 9.2 wt.% Ni₂P/MIL-125-NH₂, 8.6 wt.% NiO/MIL-125-NH₂, 7.9 wt.% Co₃O₄/MIL-125-NH₂, 8.8 wt.% CoP/MIL-125-NH₂, 10.2 wt.% Fe₂O₃/MIL-125-NH₂ and 7.9 wt.% CuO/MIL-125-NH₂ under visible light irradiation for 8 hours.

After the photocatalytic test the stability of MIL-125-NH₂ and co-catalysts was confirmed by means of PXRD, X-ray photoelectron spectroscopy (XPS), SEM, Energy-dispersive X-ray spectroscopy (EDX) and N₂ sorption isotherms (for more information see section S2.4 in the Appendix). Recycling experiments for three consecutive cycles of 12 hours were performed on both Ni₂P/MIL-125-NH₂ and NiO/MIL-125-NH₂ systems. The results showed that the photocatalytic activity of both systems is stable throughout the cycles (Figure 2.6), while PXRD experiments confirm that their crystallinity is retained (Figure S.8).

Figure 2.6: Recycling photocatalytic performance of (A) Ni₂P/MIL-125-NH₂ and (B) NiO/MIL-125-NH₂ for 3 consecutive cycles.
2.4 Mechanism

In order to obtain insights into the charge separation efficiency and the synergy between the MIL-125-NH$_2$ and the different co-catalysts, photoluminescence (PL) and transient PL experiments for each co-catalyst/MIL-125-NH$_2$ system were performed (Figure 2.7 and Section S2.5, Appendix). In principle, the addition of the co-catalyst can quench the PL emission of the photocatalyst, since they withdraw the photoexcited electrons and thus prevent the undesired electron-hole recombination. In preparation for the PL experiments, MIL-125-NH$_2$ was mixed with different amounts of co-catalyst NPs and the mixtures were subsequently suspended in the photocatalytic solution. To assess how efficiently each co-catalyst inhibits the electron-hole recombination, the emission quenching was quantified by integrating the PL curves of the MIL-125-NH$_2$ and the samples with the optimal amount of each co-catalyst, PL$_{co-cats}$/PL$_{MOF}$ (Table 1). The charge separation efficiency follows the order: Ni$_2$P > CoP ≈ Co$_3$O$_4$ > Fe$_2$O$_3$ > CuO > NiO. The PL results are consistent with the photocatalytic performance (H$_2$ generation rates) of each system, apart from the NiO/MIL-125-NH$_2$ system. As can be seen in Figure 2.7, the NiO NPs are less efficient at attracting the photogenerated electrons, yet this system exhibits one of the highest photocatalytic H$_2$ generation rates.

![Figure 2.7: PL spectra of the suspensions of MIL-125-NH$_2$ with different amounts of: (A) Ni$_2$P, (B) CoP, (C) Co$_3$O$_4$, (D) Fe$_2$O$_3$, (E) CuO and (F) NiO. The excitation wavelength was 420 nm.](image-url)
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Table 1: Photoluminescent emission quench (PL_co-cat/PL_MOF) and current density for different co-catalysts.

<table>
<thead>
<tr>
<th>Co-catalyst</th>
<th>Ni2P</th>
<th>CoP</th>
<th>Co3O4</th>
<th>Fe2O3</th>
<th>CuO</th>
<th>NiO</th>
</tr>
</thead>
<tbody>
<tr>
<td>PL_co-cat. / PL_MOF, %</td>
<td>0.7</td>
<td>2.6</td>
<td>2.5</td>
<td>6</td>
<td>17.2</td>
<td>28.8</td>
</tr>
<tr>
<td>Current Density, mA cm(^{-2}) at -0.9 V versus Ag/Ag(^+)</td>
<td>-0.07</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>-0.22</td>
</tr>
</tbody>
</table>

In order to gain a better understanding of the discrepancy between the PL results and the photocatalytic H\(_2\) generation rate of the NiO/MIL-125-NH\(_2\) system, electrochemical measurements were performed (cyclic voltammetry, CV and linear scan voltammetry, LSV). The intrinsic activity of NiO in catalysing the H\(_2\) evolution reaction was determined and compared with that of Ni\(_2\)P NPs. The intrinsic catalytic activities of the co-catalysts were compared in terms of current density. To ensure that the conditions of the electrochemical experiments are comparable to those applied in photocatalysis, a solution containing acetonitrile and water (90:10) was used, with 0.1 M tetrabutylammonium perchlorate (TBAP), to ensure the availability of conductive ions in the electrolyte. It was established that the current density (intrinsic catalytic activity) of NiO NPs is notably higher than that of the Ni\(_2\)P NPs (i.e., -0.22 and -0.07 mA cm\(^{-2}\) for NiO and Ni\(_2\)P, respectively at -0.9 V versus Ag/Ag\(^+\)), compensating for the inferior performance of the former in withdrawing electrons (Table 1 and Figure S.11). Hence, the good photocatalytic activity of the NiO/MIL-125-NH\(_2\) system with respect to H\(_2\) generation can be attributed to the superior intrinsic activity of NiO in catalysing the reduction of protons/H\(_2\)O to H\(_2\). For more information about the electrochemical measurements see section S2.6 in the Appendix.

2.5 Comparison of Ni\(_2\)P to Pt

In order to compare the performance of Ni\(_2\)P/MIL-125-NH\(_2\) with that of Pt/MIL-125-NH\(_2\), we synthesized “naked” Pt NPs\(^{106}\) with a size of ~19 nm (for more information, see section S2.7 in the Appendix). We then mixed them with MIL-125-NH\(_2\) and investigated the performance of this system under the same conditions that were used for Ni\(_2\)P/MIL-125-NH\(_2\). The amount of Pt NPs was varied and the optimum Pt/MIL-125-NH\(_2\) system produces H\(_2\) with a maximum evolution rate of 646 μmol g\(^{-1}\) h\(^{-1}\), which is almost half of that demonstrated by the noble-metal-free Ni\(_2\)P/MIL-125-NH\(_2\) system (Figure 2.8).
Variation of Co-catalysts

The effectiveness of charge separation with the Pt/MIL-125-NH₂ system was studied by PL emission spectroscopy. Figure 2.9A compares the PL spectra of MIL-125-NH₂ with various amounts of Pt NPs and the corresponding amounts of Ni₂P NPs. It is apparent that the overall PL quenching with the Ni₂P/MIL-125-NH₂ system is more drastic than that of Pt/MIL-125-NH₂, while the optimized amount of Pt NPs (2 wt.%, in terms of photocatalytic activity) does not induce complete quenching of the PL emission, which is the case for the optimized 9.2 wt.% Ni₂P/MIL-125-NH₂ system. In order to gain further insights into the charge separation efficiency, photochemical chronoamperometry measurements were carried out (Figure 2.9B) and showed that upon illumination both Ni₂P/MIL-125-NH₂ and Pt/MIL-125-NH₂ exhibit comparable initial photocurrents at the early stage, but the photocurrent in the presence of Pt NPs decays faster over time. These results are in agreement with those obtained through PL spectroscopy and suggest that the charge recombination in Pt/MIL-125-NH₂ is more pronounced and faster than in the Ni₂P/MIL-125-NH₂ system.

Figure 2.8: Comparison of the photocatalytic activity of MIL-125-NH₂ with optimized amounts of Ni₂P (green) and Pt (gray) NPs as co-catalysts.

Figure 2.9: (A) Comparison of the PL spectra of MIL-125-NH₂ with and without various amounts of Pt NPs (solid lines) and the corresponding amounts of Ni₂P NPs (dotted lines). Excitation at 420 nm. (B) Photocurrent responses of MIL-125-NH₂ with the optimized amounts of Ni₂P (9.2 wt.%, green) and Pt (2.05 wt.%, black).
2.6 Summary and Conclusions

There are relatively few examples of noble metal-free MIL-125-NH$_2$-based photocatalytic systems explored toward the H$_2$ evolution reaction. The majority of the investigated systems in this study outperform other systems based on MIL-125-NH$_2$ with noble metal co-catalysts – such as Pt and Au NPs,$^{69b, 90a, 107}$ as well as with abundant co-catalysts (e.g., Co-dioxime-diimine).$^{69b}$ More detailed comparison between the best performing Ni$_2$P/MIL-125-NH$_2$ with other MOF-based systems in the literature can be found in section S2.9 in the appendix.

From this study it is apparent that the photocatalytic performance of a system is not only influenced by the individual performance of the MOF (in terms of light absorption and charge separation within the structure) or the co-catalyst (in terms of intrinsic catalytic activity), but is also highly determined by the synergy between these two components. The last step of H$_2$ formation and desorption is associated solely with the intrinsic catalytic activity of the co-catalyst. However, despite the importance of high intrinsic activity, the last step cannot take place if there is not sufficient electron transfer from the MOF to the co-catalyst.

The literature contains many examples of visible light-active MOFs whose photocatalytic activity could be substantially improved by carefully selecting an appropriate co-catalyst.$^{86, 90, 108}$ The criteria for the co-catalyst selection should relate mainly to its synergy with the photocatalyst, rather than to its intrinsic catalytic activity. The target of this strategy is to achieve efficient interactions between the two components, ensuring efficient migration of electrons from the MOF to the co-catalyst. Transition metal phosphides and oxides have emerged as promising co-catalysts for the H$_2$ evolution reaction and their use in MOF-based photocatalytic systems can lead to performances similar to those achieved by traditional semiconductors.
Chapter 3 Variation of Electron-Donors and Dual-Functional Photocatalysis

Parts of this chapter are based on published work:

‘Concurrent Photocatalytic Hydrogen Generation and Dye Degradation Using MIL-125-NH₂ under Visible Light Irradiation’

S. Kampouri, T. N. Nguyen, M. Spodaryk, R. G. Palgrave, A. Züttel, B. Smit, K. C. Stylianou,

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S. Kampouri’s contribution: Conceiving the project’s idea, synthesis of materials, characterization (powder X-ray diffraction, UV-Vis absorbance spectroscopy), designing and performing experiments (photocatalytic experiments, trapping experiments of active species), writing text and constructing figures.
3.1 Introduction

Similar to semiconductor photocatalysis, in the field of MOFs, sacrificial agents are frequently employed. In particular sacrificial electron donors are always employed with MOF-based photocatalytic systems for the H$_2$ evolution reaction. As previously mentioned, electron donors are used to scavenge the photogenerated holes, preventing the thermodynamically more demanding O$_2$ production reaction, thus boosting the H$_2$ generation.$^{109}$ In principle, an efficient electron donor should exhibit thermodynamic adequacy with the photocatalyst, irreversible transformation into inert molecules and fast kinetics of oxidation.$^{29b}$ Depending on individual photocatalytic systems, other potentially desirable characteristics for electron donors include stability across a certain pH range, diffusion into the pores of the photocatalyst, and being used in high concentrations without causing any degradation or collapse to the other components present in the system (e.g. photocatalyst, co-catalyst).

Currently, many of the substances used as electron donors are toxic and hazardous (e.g., TEA, TEOA, methanol) or an energy resource themselves (e.g. methanol and ethanol) and their introduction increases the cost of the photocatalytic system.$^{27,110}$ Optimizing the activity and abundancy of the MIL-125-NH$_2$-based photocatalytic system in the previous chapter prompted us to focus on another important factor inhibiting the real-life application of such systems, which is the utilization of electron donors. In this chapter, first the photocatalytic performance of the Ni$_2$P/MIL-125-NH$_2$ system with different typical examples of electron donors is explored. Then, in an attempt to address the limitations related to the utilization of electron donors, an organic pollutant (Rhodamine B) is used as a replacement of typical electron donors. Considering that the active species for the H$_2$ evolution reaction (e$^-$) can be complementary to those for the organic pollutant’s degradation reaction (h$^+$), our proof-of-concept approach envisions dual-functional photocatalysis for the simultaneous production of hydrogen and pollutant degradation using the Ni$_2$P/MIL-125-NH$_2$ system, under visible light radiation. It should be mentioned here that to date, there are relatively few reports of systems combining the photocatalytic hydrogen evolution with the degradation of organic compounds, with the UV-active semiconductors (e.g. TiO$_2$) being the most studied materials.$^{111}$
3.2 Variation of Electron-Donors

In this section, I describe the photocatalytic performance of the Ni$_2$P/MIL-125-NH$_2$ system with different sacrificial reagents as electron donors. In addition to TEA, four other commonly used electron-donors were investigated; L-ascorbic acid (H$_2$A), methanol (MeOH), ethanol (EtOH), and TEOA. The amount of water was kept constant at 5% v/v in all examined photocatalytic solutions. For more information about the different photocatalytic solutions explored see section S3.1 in the Appendix.

As shown at Figure 3.1A, varying the electron donors has a profound impact on the H$_2$ production rate. Using TEA and TEOA promotes the H$_2$ generation, however the H$_2$ evolution rate in the case of TEOA is ~6 times lower than that when TEA is employed (215 vs. 1230 μmol h$^{-1}$ g$^{-1}$). When EtOH or MeOH is used, the Ni$_2$P/MIL-125-NH$_2$ system exhibits remarkably inferior activity, with H$_2$ evolution rates of only 38 and 26 μmol h$^{-1}$ g$^{-1}$ for EtOH and MeOH, respectively. When L-ascorbic acid is introduced as the electron donor, no H$_2$ evolution is observed. Both Ni$_2$P and MIL-125-NH$_2$ retain their stability after all these photocatalytic tests (Figure 3.1B). The inactivity of L-ascorbic acid can be associated with the fact that the photocatalytic solution contains significant amount of acetonitrile, which is a much less polar solvent than water and therefore can inhibit the dissociation of H$_2$A into HA$^-$ and the subsequent oxidation of HA$^-$ to HA$^•$ as in aqueous media. EtOH and MeOH are weaker bases than TEA and TEOA, which may explain why they are less effective electron donors.

![Figure 3.1](image-url)
Generally, in every photocatalytic reaction the electron transfer processes should be faster than the charge recombination. Furthermore, since the same number of electrons and holes are to be used, the rate of the oxidation reaction of the electron donors should be sufficiently high so as not to slow down the water reduction process. It appears that the oxidation of EtOH and MeOH is not adequate within the Ni$_2$P/MIL-125-NH$_2$ system, leading to the inferior performance of these photocatalytic systems. Finally, the higher H$_2$ evolution rate of the photocatalytic system with TEA compared to that with TEOA can be attributed to the difference in the concentrations used (TEA: 7.2 M, TEOA: 0.01 M, Table S.3). This effect can also be observed from the electrochemical measurements (CV) of TEA and TEOA at these concentrations. More specifically, when scanned toward positive potential, the current density of the anodic peak with TEA (onset at 1.4 V vs. Ag/Ag$^+$) is an order of magnitude higher than that of TEOA (and the other electron donors). In principle, the current density should be proportional to the rate of oxidation and the concentration of the electron donors. This indicates that even if the oxidation of TEA and TEOA occurs at comparable rates, the ability to use a much higher concentration of TEA with the Ni$_2$P/MIL-125-NH$_2$ system is important for the efficiency of the electron donation process. It is worth noting that we attempted to increase the concentration of TEOA in the photocatalytic system, but this led to the partial or complete degradation of MIL-125-NH$_2$. For more information about the electrochemical experiments, see section S3.1.1 in the Appendix.

### 3.3 Dual-Functional Photocatalytic Activity

As the above results demonstrate, using electron donors can favourably influence the performance of a photocatalytic system toward the H$_2$ evolution reaction. Nevertheless, incorporating electron donors into a photocatalytic system can induce toxicity (in the case of TEA, TEOA, and MeOH) and increase the cost of the system. In an attempt to address these limitations and provide a proof-of-concept for an alternative approach, we explored a system in which the electron donor was replaced by a typical organic pollutant, envisioning H$_2$ generation coupled with water remediation.

As mentioned previously, the main active species required for pollutants oxidation can be complementary to those needed for the photocatalytic H$_2$ evolution reaction. However, the challenge of combining these two reactions lies in the fact that the photocatalytic H$_2$ production is unlikely to occur in aerated conditions, since O$_2$ acts as an electron scavenger, whereas the photocatalytic oxidation of organic pollutants is significantly promoted in the presence of O$_2$. Hence, the photocatalyst should be capable of degrading the organic pollutant with the holes being the dominant active species and the electrons being transferred to protons (H$^+$) or water molecules to generate H$_2$. 

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Variation of Electron-Donors and Dual-Functional Photocatalysis

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This idea of dual-functional photocatalysis was explored by using the optimized 9.2 wt.% Ni$_2$P/MIL-125-NH$_2$ system and Rhodamine B (RhB) as a representative of organic pollutants in wastewater. RhB is a hazardous dye widely used in the cosmetic, textile, paint and plastic industry, and its photocatalytic degradation has been extensively studied with MOFs. RhB is toxic and its presence in water can result in teratogenic and carcinogenic effects on public health.

The RhB concentration was varied in the CH$_3$CN-based solution with water, in order to identify the favourable conditions for H$_2$ production. As illustrated in Figure 3.2A, when the dye concentration initially increases, the H$_2$ evolution rate is promoted, indicating that more RhB molecules are available to interact with the photoexcited holes. Further increase in RhB concentration causes a decrease in the amount of H$_2$ generated, since a portion of the incident light is inhibited from approaching the MIL-125-NH$_2$. The H$_2$ evolution rate reaches a maximum value of 335 μmol h$^{-1}$ g$^{-1}$ at a RhB concentration of 1.2 ppm in the photocatalytic solution (0.02 mg in 17 mL). The H$_2$ evolution rate is lower when RhB is used as an electron donor than when TEA is used. This is in accordance with the basicity of these substances and the fact that the TEA concentration used is significantly higher than that of RhB. In addition, the smaller size of the TEA molecules could allow them for more efficient diffusion through the porous assemblies of MIL-125-NH$_2$, and thus provide more effective electron donation. The overall advantage of using RhB as an electron donor relies on the significant benefit of dual-functional photocatalysis.

Figure 3.2: (A) Photocatalytic H$_2$ evolution rates of Ni$_2$P/MIL-125-NH$_2$ in 17 mL of photocatalytic solutions with different concentrations of RhB, under visible light irradiation for 8 h. Inset: photograph showing the difference in the color of the photocatalytic solution before and after the photocatalytic test, with initial RhB concentration of 1.2 ppm. (B) UV-Vis spectra of reference samples and supernatants after the photocatalytic RhB decolorization process.
Variation of Electron-Donors and Dual-Functional Photocatalysis

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After the photocatalytic test, the RhB concentration was monitored for the three best performing solutions (initial RhB concentration of 0.6, 1.2 and 1.8 ppm, Figure 3.2B). For this purpose, the UV-Vis absorbance spectra of the supernatants after the photocatalytic test were collected. With an initial RhB concentration of 0.6 ppm, the dye is completely degraded and with concentration of 1.2-1.8 ppm, the dye concentration substantially decreases. Control experiments were performed to confirm that RhB was photodegraded and not absorbed by MIL-125-NH2. The Ni2P/MIL-125-NH2 system was immersed in 1.2 ppm RhB solution and kept in the dark for 24 h. The supernatant was then collected and subjected to UV-vis absorbance spectroscopy. Interestingly, there is no notable difference between the RhB concentrations before and after this test (Figure 3.3A). Moreover, when 1.2 ppm RhB solution was irradiated in the absence of MIL-125-NH2, there was no change in the dye concentration, confirming that no self-photolysis of RhB occurs (Figure 3.3B). These results indicate that RhB is indeed involved in the photocatalytic process for H2 evolution.

Figure 3.3: Control experiments: (A) UV-vis spectra of 1.2 ppm RhB solution (reference), the supernatant of the same solution combined with Ni2P/MIL-125-NH2 after 24 h in the dark and after the photocatalytic test. (B) UV-Vis spectra of 1.2 ppm RhB solution (reference) and same solution after 3 h irradiation with visible light.

To provide further confirmation of the role of RhB and elucidate the photocatalytic mechanism, trapping experiments of active species were carried out using the optimal RhB concentration (1.2 ppm, 2.5 μM) and 0.01 M tert-Butanol as •OH radical scavenger or 0.01 M TEOA as hole scavenger (more information in section S3.2, Appendix). As displayed in Figure 3.4, the addition of TEOA and tert-Butanol leads to 11.2 % and 18.7 % decrease in the RhB concentration, respectively. This confirms that both the holes and •OH radicals are involved in this process, with the former being the more dominant species. These results further prove that RhB acts as a sacrificial electron donor, scavenging the photoexcited holes and promoting the H2 generation, which leads to its decomposition, as
Variation of Electron-Donors and Dual-Functional Photocatalysis

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demonstrated by UV-Vis spectroscopy. After all the photocatalytic experiments, the stability of the samples was verified by means of PXRD (Figures S18 and S19, Appendix). For more information about the control experiments see section S3.2 in the Appendix.

Figure 3.4: The impact of the incorporation of TEOA as hole scavenger and tert-butanol as •OH radical scavenger on the RhB degradation rate after 3 h photocatalytic test.

3.4 Summary and Conclusions

In summary, variation of the electron donors highlights their significant influence on the overall photocatalytic performance toward the hydrogen evolution reaction. When TEA is used, the Ni$_2$P/MIL-125-NH$_2$ demonstrates significantly higher H$_2$ evolution rates, which can be associated with the higher basicity of TEA, its compatibility with MIL-125-NH$_2$ (in terms of MIL-125-NH$_2$ being stable in high concentrations of TEA) and the fast kinetics of the TEA oxidation process.

Drawing on these insights, we then integrated the photocatalytic H$_2$ evolution and degradation of the organic dye RhB in a single process using the Ni$_2$P/MIL-125-NH$_2$ system under deaerated conditions. This dual-functional photocatalytic system generates H$_2$ at a relatively high rate and the organic dye – acting as an electron-donor – is degraded under visible-light irradiation. Although the dual-functional photocatalysis field is in its infancy, there are several scientific studies focused on the simultaneous photocatalytic H$_2$ production and organic pollutants degradation.$^1$ However, the majority of these studies are based on wide-band-gap semiconductors (i.e., TiO$_2$), restricting the profitable irradiation to only 4% of the solar spectrum that constitutes the UV region.$^{116}$ For example, Cho et al. and Kim et al. recently investigated the concept of dual-functional photocatalysis for H$_2$ production with simultaneous degradation of 4-chlorophenol and urea or urine.$^{111}$ These photocatalytic systems however are based on UV-active semiconductors such as SrTiO$_3$ and TiO$_2$, and noble metals NPs (e.g.
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Rh, Pt) as co-catalysts. In our work, the Ni₂P/MIL-125-NH₂ photocatalytic system operates under visible light irradiation – a dominant region of the solar spectrum – and is free of noble metals. To the best of our knowledge this is the first example of a dual-functional MOF-based photocatalytic system able to simultaneously produce H₂ and degrade organic pollutants under visible light irradiation. This study showcases the great potential of MOFs in photocatalytic applications and paves the way for the generation of new visible light-active and water-stable MOF-photocatalysts that can accomplish simultaneous water remediation as well as H₂ generation.
Chapter 4 Design of MOF/MOF Heterojunctions

Parts of this chapter are based on submitted manuscript:

‘Development of a MOF/MOF Heterojunction for Enhanced Photocatalytic Performance’

S. Kampouri, F. M. Ebrahim, M. Fumanal, P. A. Schouwink, B. Smit,* Christopher P. Ireland* and K. C. Stylianou*

S. Kampouri’s contribution: Conceiving the project’s idea, synthesis of materials, characterization (powder X-ray diffraction, UV-Vis absorption, steady-state and transient photoluminescence spectroscopy, nitrogen sorption isotherms, scanning electron microscopy), designing and performing experiments (photocatalytic experiments, quantum yield calculations), writing text and constructing figures.
4.1 Introduction

As mentioned above, only a minor fraction of photons in sunlight corresponds to the UV region; the vast majority being in the visible and infrared regions. Considering the importance of exploiting the abundant solar energy, light-active materials with narrow band gap (< 2.95 eV), capable of absorbing a large part of the spectrum that composes sunlight, are highly desired. Besides light harvesting (first major photocatalytic step), the separation of the photogenerated charge carriers and their efficient migration (second major photocatalytic step) are additional important factors that define the overall photocatalytic activity of a system. This is the main reason why the majority of photocatalytic systems utilize metal-based nanoparticles or molecular complexes as co-catalysts that can inhibit the undesired electron-hole recombination. In semiconductor photocatalysis, an alternative strategy to optimize the optical, electronic and catalytic properties is to combine different materials in a form of a heterojunction. By carefully matching materials with different band gaps, absorption of a wider region of light can be achieved. In addition, depending on the position of the conduction and valence band of the materials that are combined, the photogenerated electrons can be spatially separated from the holes, which can decrease the probability of the undesired charge carrier recombination (e.g. type II heterojunction of staggered gap, Figure 4.1).

![Figure 4.1](image_url)

**Figure 4.1:** Schematic representation of three different types of heterojunctions: (A) type I, with straddling gap (B) type II with staggered gap and (C) type III with broken gap.

Similar to semiconductor photocatalysis, also in the field of MOF photocatalysis, one of the main challenges is to develop a system that has both a narrow band gap as well as sufficient charge separation efficiency. Inspired by semiconductor systems, in this chapter, we explored the strategy of combining different MOFs to form a MOF/MOF heterojunction for photocatalysis. Heterojunctions with MOFs recently started to be explored, with the majority of studies focusing on combining MOFs with either semiconductors or COFs (covalent-organic frameworks). Recently, synthetic protocols for the design of MOF/MOF core-shell/satellite architectures have been reported.
However, the synthesis of MOF-on-MOF structures does not guarantee the formation of a heterojunction, with the latter involving electronic interactions that alter the optoelectronic properties of the hybrid materials. In the ideal case, fully optimized MOF/MOF heterojunctions should demonstrate extended solar-light harvesting properties, high charge separation efficiency, enhanced catalytic activity and high porosity (allowing for facile diffusion of the photocatalytic solution). Understanding and elucidating the impact of the altered optoelectronic properties of MOF/MOF heterojunctions on the photocatalytic activity – although neglected – is of key importance for the strategic design of highly active MOF-based photocatalysts. Development of such a photocatalytic system would eliminate the necessity of additional expensive components, such as co-catalysts and/or dyes as photosensitizers.

This chapter describes the implementation of this strategy to improve the activity of the well-studied MIL-125-NH$_2$. To select an optimum component to form a heterojunction, we need a MOF with complementary light harvesting properties to those of MIL-125-NH$_2$. As MIL-125-NH$_2$ absorbs light early in the visible (< 500 nm), we need a MOF with more extended visible-light absorption. A promising candidate is MIL-167, also a stable Ti-based MOF, which shows immense light absorption in the extended visible (up to ~700 nm). By combining these two MOFs, we envisioned enhanced photocatalytic activity driven by light energy harvested from almost the whole visible region.

### 4.2 Synthesis and Characterization

The synthetic protocols of the individual MOF components MIL-125-NH$_2$ and MIL-167 are based on reported procedures and are described in sections S2.1 and S4.1 in the Appendix. For the synthesis of the MIL-167/MIL-125-NH$_2$ materials, we developed a strategy in which we use the MIL-167 crystals as additional nucleation centers for the growth of MIL-125-NH$_2$ (Figure 4.2). As anticipated based on the thermogravimetric analysis (TGA) profile (Figure S.22, Appendix), MIL-167 retains its structural integrity throughout the heating process, thus its amount was varied resulting in the synthesis of several MIL-167/MIL-125-NH$_2$ hybrid materials, with different ratios between their MOF components (Figure S23, Appendix). SEM images show that the large crystals of MIL-167 (~ 60 μm) are homogeneously covered by the significantly smaller crystals of MIL-125-NH$_2$ (~ 300 nm, Figure 4.3). More SEM images as well as more details about the characterization of the samples can be found in Section S4.2, in the Appendix.
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Figure 4.2: Schematic representation of the synthetic protocol of the MIL-167/MIL-125-NH₂ heterojunction. MIL-125-NH₂ is synthesized in the presence of MIL-167 crystals.

Figure 4.3: SEM images of (A) MIL-167, (B) MIL-125-NH₂, (C) and (D) MIL-167/MIL-125-NH₂, with different magnification.

4.3 Photocatalytic Activity

The photocatalytic activities of three different MIL-167/MIL-125-NH₂ materials with 3, 8, and 13 wt.% of MIL-167 were investigated toward the H₂ evolution reaction. For this purpose, the samples were irradiated under visible light (≥420 nm), with TEA as the electron donor and without the assistance of any co-catalysts (see Section S3 for more details). As shown in Figure 4.4, the MIL-
167/MIL-125-NH₂ materials significantly outperform their constituent components MIL-167 and MIL-125-NH₂. In fact, MIL-167 produces only traces of H₂, while the best-performing 8 wt.% MIL-167/MIL-125-NH₂ outperforms MIL-125-NH₂ by a factor of almost ten. As a comparison, the photocatalytic activities of the analogous physical mixtures of MIL-125-NH₂ and MIL-167 were investigated. The physical mixtures (with the same MIL-167:MIL-125-NH₂ ratios) exhibit inferior photocatalytic activities, ranging from 30 to 50 μmol h⁻¹ g⁻¹, attributed to the activity of solely the MIL-125-NH₂ component. After the photocatalytic experiments, the stability of the samples was verified by means of PXRD and N₂ sorption isotherms (Figure S26). Furthermore, recycling experiments revealed that the MIL-167/MIL-125-NH₂ heterojunctions consistently produce hydrogen for more than 36 h (Figure S27), while no significant oxidation of the MOFs’ linkers was observed after the cycles (Table S5).

**4.4 Mechanism**

To understand this performance increase, we carried out a combined experimental and computational study to explore the optoelectronic properties of these materials. One of the aims of creating a heterojunction is to extend the visible-light absorption of MIL-125-NH₂. **Figure 4.5A** demonstrates that indeed the MIL-167/MIL-125-NH₂ materials exhibit an extended light absorption in the region from 500 to 750 nm. As with the photocatalytic experiments we use only visible-light (λ ≥ 420 nm), this enhancement has a significant effect on the number of photons absorbed. To determine the formation of a heterojunction and its type, we carried out PL spectroscopy by suspending the different
samples in acetonitrile. Figure 4.5B compares the PL emission of the heterojunction materials with that of MIL-125-NH$_2$, MIL-167 and their analogous physical mixtures. In the MIL-125-NH$_2$ emission spectrum, we observe three dominant peaks at around 470, 490 and 550 nm. In this region, MIL-167 weakly emits and shows no clear emission peaks. The MIL-167/MIL-125-NH$_2$ materials, exhibit more similar PL emission to that of MIL-125-NH$_2$. However, with the MIL-167/MIL-125-NH$_2$ materials, the weaker emission peak of MIL-125-NH$_2$ at 470 nm is significantly intensified. The magnitude of this intensification depends on the amount of MIL-167 in the samples. This increase in intensity indicates that within MIL-167/MIL-125-NH$_2$ there are now more electrons in the conduction band (or LUCOs) of MIL-125-NH$_2$ when they recombine with the hole, they intensify the emission peak at 470 nm. As these electrons can only come from the CB of MIL-167, its CB should be higher in energy than that of MIL-125-NH$_2$ and the MOFs should exhibit good electronic contact.

![UV-vis absorption spectra](image1)

![PL emission spectra](image2)

**Figure 4.5:** (A) UV-vis absorption spectra and (B) PL emission of MIL-125-NH$_2$ (yellow), MIL-167 (red), MIL-167/MIL-125-NH$_2$ heterojunctions (solid lines, mustard: 3 wt.%, orange: 8 wt.% and dark orange 13 wt.%) and corresponding physical mixtures (dotted lines, mustard: 3 wt.%, orange: 8 wt.% and dark orange: 13 wt.%). The excitation wavelength was 420 nm.

However, this is not the case for the physical mixtures of MIL-167 and MIL-125-NH$_2$, which demonstrate similar emission to that of MIL-125-NH$_2$. The absence of the intensified MIL-125-NH$_2$ emission peaks with the physical mixtures, manifests the lack of interactions between the two MOF crystals when they are physically mixed. This can be correlated with their different morphology compared to the heterojunction materials, as the MIL-125-NH$_2$ crystals are randomly aggregated on small regions of the MIL-167 crystals (Figure 4.6). These results demonstrate that the synthetic method of growing MIL-125-NH$_2$ in the presence of MIL-167 results in the formation of a heterojunction.
Figure 4.6: SEM images comparing the morphology of (A) the 8 wt.% MIL-167/MIL-125-NH$_2$ heterojunction sample with that of (B) its analogous physical mixture of MIL-167 and MIL-125-NH$_2$.

To further confirm these results and determine the relative positions of the valence bands (or HOCOs) of MIL-167 and MIL-125-NH$_2$, density functional theory (DFT) calculations were used (for details, see section S4.4.1, Appendix). Figure 4.7A illustrates the positions of the VBs and CBs of both MOFs, with respect to the vacuum level. The calculations reveal that the CB of MIL-167 is 0.6 eV above the CB of MIL-125-NH$_2$, which is in nice agreement with the PL results. As the VB of MIL-167 is above that of MIL-125-NH$_2$, the MIL-167/MIL-125-NH$_2$ materials demonstrate the characteristics of a type II heterojunction; staggered gap (Figure 4.7B). In this heterojunction, MIL-167 is the photosensitizer (harvesting light in the extended visible region where MIL-125-NH$_2$ is incapable) and MIL-125-NH$_2$ is the (photo)catalyst.

Figure 4.7: (A) Computed electronic band alignment relative to the vacuum level of MIL-167 and MIL-125-NH$_2$. (B) Schematic representation of a type II heterojunction with MIL-167 and MIL-125-NH$_2$. 

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To further prove the photosensitization effect by the MIL-167 counterpart, we performed a new photocatalytic test, irradiating the MIL-167/MIL-125-NH₂ heterojunction with wavelength higher than 515 nm. Although MIL-125-NH₂ slightly absorbs in this region (λ ≥ 515nm), the MIL-167/MIL-125-NH₂ heterojunction is photocatalytically active, producing significantly higher amounts of hydrogen compared to that of MIL-125-NH₂ (197 versus 5 μmol g⁻¹, for MIL-167/MIL-125-NH₂ and MIL-125-NH₂, respectively, after 8 h irradiation). These results further highlight the substantial improvement of the MIL-125-NH₂ photocatalyst by combining it with MIL-167, and the extended light absorption properties of the MIL-167/MIL-125-NH₂ heterojunctions. Furthermore, the apparent quantum yields (AQY) of the MIL-167/MIL-125-NH₂ heterojunction were calculated at 450 and 500 nm (see section S4.4.2, Appendix). The AQYs were 2.5 % and 0.7 %, at 450 and 500 nm, respectively, which are high values for this type of materials and comparable to semiconductor photocatalysts (Table S6). These results are in line with the light absorption properties, as both MOF counterparts of the heterojunction absorb photons at 450 nm. However, at 500 nm, it is mainly the MIL-167 counterpart that absorbs photons, thus the AQY at this wavelength is lower, but further confirms the photosensitization effect.

Having a type II heterojunction is necessary but not sufficient for enhanced photocatalytic activity. It is also important to ensure that the long lifetime (3 - 5 μs) of the electrons in MIL-125-NH₂ is not influenced negatively by the formation of the heterojunction. Sufficiently long lifetimes are required for an electron to be able to be involved in other processes (i.e. electron transfer within the heterojunction or to protons/water molecules) before recombining with the hole. Transient PL spectroscopy experiments reveal that the lifetime associated with the main PL emission peaks of MIL-125-NH₂, displayed in Figure 4.5B, is not influenced by the formation of the heterojunctions (for details see section S.4.4.3, Appendix).

The photocatalytic performance shows an optimum activity with the 8 wt.% MIL-167/MIL-125-NH₂ sample (Figure 4.4). This volcano type trend between the photocatalytic performance and the amount of MIL-167 in the heterojunctions can be associated with different competing mechanisms taking place. MIL-125-NH₂ is photosensitized further in the visible region by MIL-167 (Figure 4.5A). However, the bulky large crystals of MIL-167 can compete with MIL-125-NH₂ in the absorption of photons with higher energy (corresponding to less than 500 nm) and in excess amounts they could also act as recombination centers. In the case of 8 wt.% MIL-167/MIL-125-NH₂, optimum interactions between the two MOFs occur, thus resulting in superior photocatalytic activity. This is further supported by the PL data, where the emission peak of MIL-125-NH₂ at ~ 470 nm is the most intensified.
for the 8 wt.% MIL-167/MIL-125-NH₂, compared to the other heterojunction materials, revealing a higher density of electrons being transferred from MIL-167 to MIL-125-NH₂.

### 4.5 Heterojunction with UIO-66-NH₂ and MIL-125-NH₂

With the aim of successfully applying the developed methodology with other materials, we synthesized UIO-66-NH₂/MIL-125-NH₂ heterojunctions, following a similar synthetic and characterization protocol (see section S4.5, Appendix). DFT calculations show that the energetic positions of the UIO-66-NH₂ CB and VB are higher than those of MIL-125-NH₂, which is also in agreement with the literature (Figure 4.8A).⁴²³ We can thus expect a type II heterojunction, but now with comparable band gaps of the individual MOF components. This is not ideal as none of the MOF components can photosensitize the other. Indeed, the UV-vis spectra of the different samples demonstrate that increasing the amount of UIO-66-NH₂ in the UIO-66-NH₂/MIL-125-NH₂ materials, results in an increasing small blue-shift (from around 500 nm for MIL-125-NH₂ to a maximum of 450 nm for UIO-66-NH₂), attributed to the slightly smaller visible-light absorption of UIO-66-NH₂ (Figure 4.8B).

![Figure 4.8](image_url)

**Figure 4.8:** (A) Computed electronic band alignment relative to the vacuum level of UIO-66-NH₂ and MIL-125-NH₂. (B) UV-vis absorption spectra of MIL-125-NH₂ (yellow), UIO-66-NH₂ (black) and UIO-66-NH₂/MIL-125-NH₂ materials (dark-light gray).

As expected, the photocatalytic experiments shown in Figure 4.9A, illustrate that there is no enhancement in the photocatalytic activity. On the contrary, increasing the amount of UIO-66-NH₂ has a negative impact on the H₂ evolution, which attains a minimum value of 0.8 μmol h⁻¹ g⁻¹, when only UIO-66-NH₂ is irradiated. It is noteworthy that a heterojunction is still formed in this system, as manifested by the PL experiments. The PL spectra of the UIO-66-NH₂/MIL-125-NH₂ materials follow
a similar trend to that of the MIL-167/MIL-125-NH$_2$ heterojunctions; the weak emission peak of MIL-125-NH$_2$ located at around 470 nm, and generally the entire emission is significantly intensified with increasing amounts of UIO-66-NH$_2$ in the heterojunction materials (Figure 4.9B). These results indicate that, upon irradiation, photoexcited electrons are transferred from the CB of UIO-66-NH$_2$ to that of MIL-125-NH$_2$, enhancing the intensity of the MIL-125-NH$_2$ emission. However, due to their comparable light absorption, photons with appropriate energy reaching the heterojunction can now either excite an electron within MIL-125-NH$_2$ or UIO-66-NH$_2$. This competition is not beneficial for photocatalysis, as it increases the number of possible charge recombination pathways.

Figure 4.9: (A) Photocatalytic performance and (B) PL spectra of MIL-125-NH$_2$ (yellow), UIO-66-NH$_2$ (black) and UIO-66-NH$_2$/MIL-125-NH$_2$ heterojunctions (dark-light gray).

4.6 Summary and Conclusions

Summarizing, this chapter describes a strategy to improve the photocatalytic activity of MOF-based systems, through the construction of MOF/MOF heterojunctions. As a proof-of-concept, MIL-125-NH$_2$ was synthesized in the presence of either MIL-167 or UIO-66-NH$_2$ crystals. The MIL-167/MIL-125-NH$_2$ heterojunction materials exhibit significantly greater photocatalytic performance, which is attributed to the charge spatial separation and the extended visible-light absorption. Combined experimental and computational efforts demonstrate that MIL-167/MIL-125-NH$_2$ is a type II heterojunction with a staggered gap, where MIL-167 acts as the photosensitizer and MIL-125-NH$_2$ withdraws the electrons and acts as the photo(catalyst). The inferior activity of another type II heterojunction with UIO-66-NH$_2$ and MIL-125-NH$_2$ (UIO-66-NH$_2$/MIL-125-NH$_2$) highlights the importance of selecting appropriate MOFs for this strategy, exhibiting complementary light absorption.
properties and suitable band alignment. Future efforts should focus on further elucidating the mechanism of this novel type of heterojunctions and exploring the impact of the morphological characteristics on the photocatalytic activity.
Chapter 5 MOF-derived Titanium Dioxide for Photocatalysis; The Impact of the Templated Morphology

Parts of this chapter are based on published work:

‘Mixed-Phase MOF-Derived Titanium Dioxide for Photocatalytic Hydrogen Evolution: The Impact of the Templated Morphology’

S. Kampouri, C. P. Ireland, B. Valizadeh, E. Oveisi, P. A. Schouwink, M. Mensi, K. C. Stylianou,

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S. Kampouri contributed in conceiving the project’s idea, the synthesis of materials, characterization (powder X-ray diffraction, UV-Vis absorption, photoluminescence spectroscopy, nitrogen sorption isotherms, scanning electron microscopy), designing and performing experiments (photocatalytic experiments), writing text and constructing figures.
5.1 Introduction

Titanium dioxide (TiO$_2$) was the first material investigated for the photocatalytic splitting of water, and despite its very limited solar light absorption (UV-active), it is still considered a promising photocatalyst, due to its chemical and thermal stability, non-toxicity, high reactivity and low cost. Although tremendous research efforts have focused on the use of TiO$_2$ nanoparticles in a variety of photocatalytic applications (water splitting, water remediation, CO$_2$ reduction), the impact of the shape and morphology of TiO$_2$ NPs on their activity, as well as the interplay between the two most common phases of TiO$_2$ (rutile and anatase) remain questionable.

The low temperature TiO$_2$ anatase phase is generally recognized as a better catalyst toward the H$_2$ evolution reaction. This is attributed to its high surface reactivity, meaning that a higher concentration of active sites is present compared to the TiO$_2$ rutile phase. Nevertheless, the more thermodynamically stable TiO$_2$ rutile phase has a smaller band gap implying that a higher proportion of early visible light can be absorbed, thus careful engineering of TiO$_2$ rutile phase particles can lead to enhanced photocatalytic performance. On the other hand, materials based on mixed TiO$_2$ rutile and anatase phases have demonstrated superior photocatalytic performance in comparison with the single phase TiO$_2$. P25 Degussa, a mixed phase commercial TiO$_2$ is widely used as a standard material for photocatalysis and was studied by Bickley in 1990 to ascertain its success compared to a single phase TiO$_2$. The conclusion was that the excellent electronic contact between the two phases allowed enhanced electron-hole separation.

Subsequent literature has both agreed and disputed these findings and specifically the morphology of the phases’ interface and the interactions between them. However, ample evidence has been provided through both experimental and computational approaches to prove that there is indeed a synergy between the anatase and rutile phases, allowing more efficient electron-hole separation. A general understanding has therefore been developed that TiO$_2$ materials with the anatase and rutile phases in a 3:1 ratio can outperform comparably formed single phase TiO$_2$. The different structural arrangement of TiO$_2$ anatase and rutile phases affords distinct phase electronic structures. As conclusively demonstrated by Scanlon et al., close electronic contact between the two phases allows electrons on rutile TiO$_2$ to irreversibly be transferred to the complementary TiO$_2$ anatase phase, with this synergy promoting the electron-hole separation efficiency.

In order to engineer the electronic contact between the two TiO$_2$ phases, several synthetic techniques have been utilized; approaches of physically mixing anatase and rutile TiO$_2$ have been attempted, but mixing does not allow the two phases to sufficiently interact with each other and achieve this
MOF-derived TiO$_2$ for Photocatalysis

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synergy.\textsuperscript{132} Synthetic efforts on mixed phase TiO$_2$ have employed the sol-gel route, hydro/solvo-thermal method, electrodeposition, direct oxidation method or chemical vapor deposition.\textsuperscript{133} Many of these techniques, however, lack of control, are impractical or may not provide highly active and stable TiO$_2$ NPs.

While MOFs themselves hold great potential for catalytic applications, limitations related to their thermal and chemical stability under harsh conditions (e.g., in highly acidic/basic media or high temperatures, $> 400 \, ^\circ C$) have recently triggered their utilization as precursors for the preparation of very stable metal oxides with high catalytic activities.\textsuperscript{95,97,99,134} More specifically, the MOF-mediated synthesis involves the use of MOFs as crystalline self-sacrificing templates, in order to tailor the shape, composition, surface area and stability of the resulting materials.\textsuperscript{135} Calcination of MOFs in air leads to the partial or complete burning of the organic ligands and formation of highly active metal-oxide NPs, while carbonization at high temperatures under inert conditions can result in a carbon matrix encapsulating the metal phase (NPs).\textsuperscript{136}

Among the Ti$^{IV}$-based MOFs, MIL-125-NH$_2$ was recently utilized as a precursor for the synthesis of TiO$_2$ decorated with either palladium or gold NPs for photocatalytic applications, such as CO$_2$ reduction and hydrogen evolution.\textsuperscript{137} The employment of expensive noble metal NPs as co-catalysts was necessary to enhance the otherwise inadequate photocatalytic performance of TiO$_2$, attributed to high electron-hole recombination rates. Although these studies provided important insights, there is still a need to better understand the impact of templated MOF-derived TiO$_2$ rutile and anatase phases, and the effect of the morphology on the synergy between the two phases in photocatalysis.

In this chapter, the photocatalytic performance of TiO$_2$ prepared using the MIL-125-NH$_2$ as a starting material is primarily investigated. By calcining MIL-125-NH$_2$ over a range of temperatures, the ratio of the anatase and rutile phases can be controlled and optimized for the photocatalytic H$_2$ evolution reaction. For comparison, TiO$_2$ synthesized from precipitated amorphous titanium hydroxide and MIL-167-derived TiO$_2$ was prepared. Interestingly, the photocatalytic activity of the best-performing mixed-phase MIL-125-NH$_2$-derived TiO$_2$ is higher than all the other TiO$_2$ samples, including the commercially available P25 Degussa. Morphological studies reveal the unique nature of the templated MIL-125-NH$_2$-derived TiO$_2$ with nanosized anatase and rutile NPs forced into effective contact with each other, leading to an efficient electron-hole separation. This work highlights the role of effective interactions of TiO$_2$ anatase and rutile phases in a template and offers an intriguing outline for the development of efficient photocatalysts.
5.2 Synthesis and Characterization

Prior to the MOF-mediated synthesis of TiO$_2$, TGA of MIL-125-NH$_2$ was carried out. Figure 5.1A shows the TGA profile of MIL-125-NH$_2$. Two major weight losses are observed from 30 to ~200 °C, owed to the removal of guest molecules from the pores of MIL-125-NH$_2$, and from 360 to 600 °C related to the structural collapse of MIL-125-NH$_2$. The final product obtained after heating MIL-125-NH$_2$ at 600 °C corresponds to the remaining metal oxide, TiO$_2$, as confirmed by PXRD.

Acquiring a clear perception regarding the thermally-induced structural collapse of the MIL-125-NH$_2$, allowed us to continue with the synthesis of MOF-derived TiO$_2$ at selected temperatures (400 - 800 °C) under air (see section S5.1, Appendix). As can be seen in Figure 5.1B, calcination at different temperatures yields TiO$_2$ with differing ratios between the TiO$_2$ anatase and rutile phases. The anatase to rutile transformation is reconstructive, meaning that it involves the breaking of chemical bonds. The transformation temperature itself can vary depending on the starting material particle size, shape and volume and experimental factors such as atmosphere, heating rate and sample container. Reported temperatures for the TiO$_2$ anatase to rutile transformation vary from 400 to 1200°C.$^{138}$ Presently, heating the MIL-125-NH$_2$ crystals up to 400 °C leads to the formation of pure TiO$_2$ anatase phase. Increasing the temperature to 500°C induces the formation of the rutile phase (giving rise to a mixed phase TiO$_2$), which gradually becomes the dominant phase at 800 °C. Here, the anatase to rutile phase transformation is at the lower end of the range of temperatures reported for this transformation.$^{139}$

![Figure 5.1: (A) TGA curve of MIL-125-NH$_2$ obtained under airflow with a heating rate of 5 °C min$^{-1}$. (B) PXRD patterns of the as-synthesized MIL-125-NH$_2$-derived TiO$_2$.](image-url)
Further insights into the specific crystallization process can be gained from a variable temperature (VT) PXRD experiment, where MIL-125-NH₂ enclosed in a quartz capillary was heated from 30 to 800 °C (Figure 5.2A). Although direct comparison of ex-situ and in-situ experiments is not applicable (due to the crystallization process being rather complex and related to heating rate, dwell time and volume of the heating vessel), the in-situ VT PXRD experiment offers an insight into the TiO₂ anatase and rutile phase formation. Upon heating, the PXRD patterns show that the Bragg reflections of MIL-125-NH₂ are eliminated due to decomposition, leading to an amorphous powder. Upon further heating, initial formation of the TiO₂ anatase phase is observed as a broad hump at 25.2 ° 2θ, at furnace temperature of 560 °C. The peak width (Full width half maximum: FWHM 1.73 2θ) manifests a lack of crystallinity which can be indicative of defects and a small domain size (Figure 5.2B). This poor crystallinity may be related to the removal of the organic ligand which disrupts the crystallization process at the initial stages of TiO₂ formation. Interestingly, further heating to 660°C reveals the formation of a more crystalline TiO₂ rutile phase with a FWHM of 0.31 2θ. The integral intensity of the TiO₂ anatase Bragg peak remains constant during initial rutile formation, suggesting that the initial formation results from the same matrix as anatase. Both phases co-exist before the TiO₂ rutile phase becomes the dominant phase at 785 °C. The TiO₂ anatase and rutile phase composition, and the average domain size were quantified by Rietveld refinement from the as-prepared TiO₂ samples (Table 2). It is found that the domain size for both TiO₂ anatase and rutile phases is comparable at the same temperature, and increases with temperature.

![Figure 5.2: (A) Variable temperature PXRD patterns of MIL-125-NH₂ heated up to 810 °C and (B) Full Width Half Maximum (FWHM) values of the TiO₂ anatase and rutile peaks derived from the variable temperature PXRD patterns of MIL-125-NH₂.](image-url)
To serve as a reference, different pure and mixed phase TiO$_2$ were also obtained by calcining (400 - 800 °C) precipitated amorphous titanium hydroxide (TiH$_4$O$_4$) as a starting material (see section S5.1, Appendix). The TGA profile of the TiH$_4$O$_4$ shows a weight loss up to 450 °C, related to the removal of hydroxide, coordinated water and/or organic residues derived from the titanium isopropoxide (Figure 5.3A). Upon calcination, the TiO$_2$ anatase to rutile phase transformation starts at 700 °C and above this temperature, the TiO$_2$ rutile phase becomes dominant (Figure 1d). VT PXRD experiments collected on TiH$_4$O$_4$, using the same protocol used with MIL-125-NH$_2$ revealed that the formation of TiO$_2$ anatase phase begins at a significantly lower temperature than the MIL-125-NH$_2$-derived TiO$_2$, with peaks of anatase identified as low as 180°C (Figure 5.3B, C). Interestingly, the qualitative trend of domain-sizes of anatase and rutile is inversed between MIL-125-NH$_2$ and TiH$_4$O$_4$-derived samples, with initial rutile formation from TiH$_4$O$_4$ showing a much larger FWHM than from MIL-125-NH$_2$ (Figure 5.3D). This suggests that nucleation and growth proceeds through different mechanisms, which is likely related to the topological differences between a pre-existing framework/matrix of Ti-O polyhedral in TiH$_4$O$_4$ as opposed to discrete Ti-O clusters separated by the organic ligands in MIL-125-NH$_2$. Furthermore, the ex-situ determined domain size of the TiO$_2$ anatase phase is significantly greater than that of the MIL-125-NH$_2$-derived material, with the FWHM only 0.73 2θ, implying a much higher degree of crystallinity. This improved crystallinity delays the formation of the rutile phase to 710 °C.  

### Table 2: PXRD-derived data revealing the size and composition of TiO$_2$ synthesized using MIL-125-NH$_2$ and TiH$_4$O$_4$ as precursors.

<table>
<thead>
<tr>
<th>Synthesis Temp. (°C)</th>
<th>MIL-125-NH$_2$-derived TiO$_2$</th>
<th>TiH$_4$O$_4$-derived TiO$_2$</th>
</tr>
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<tbody>
<tr>
<td>400</td>
<td>100</td>
<td>400</td>
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</tbody>
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Anatase (w.t. %)  
Rutile (w.t. %)  
Size Anatase (nm)  
Size Rutile (nm)  

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Figure 5.3: (A) TGA curve of precipitated TiH$_4$O$_4$ obtained under airflow with a heating rate of 5 °C min$^{-1}$. (B) PXRD patterns of TiH$_4$O$_4$-derived TiO$_2$. (C) Variable temperature PXRD patterns of TiH$_4$O$_4$ heated up to 810 °C. (D) FWHM values of the TiO$_2$ anatase and rutile peaks derived from the variable temperature PXRD patterns of amorphous titanium hydroxide (TiH$_4$O$_4$).

Figure 5.4 displays the Kubelka-Munk representation of the UV-vis diffuse reflectance spectra of TiO$_2$ synthesized from MIL-125-NH$_2$ and TiH$_4$O$_4$ as precursors. The spectra of both families are in agreement with the literature, with the presence of rutile – which has a smaller band gap – causing a red shift in the light absorption edge compared to phase pure anatase TiO$_2$. In the case of the MIL-125-NH$_2$-derived TiO$_2$ synthesized through calcination at 400 °C, there is an extended light absorbance in the visible region, which is attributed to the residual carbon from the MIL-125-NH$_2$ covering the surface of TiO$_2$ and is in agreement with the TGA data (Figure 5.1A).
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Figure 5.4: Kubelka-Munk representation of the UV-Vis diffuse Reflectance of (A) MIL-125-NH$_2$-derived and (B) TiH$_4$O$_4$-derived TiO$_2$.

The morphologies of MIL-125-NH$_2$- and TiH$_4$O$_4$-derived TiO$_2$ were investigated through SEM and TEM. As can be seen in Figure 5.5, the well-defined crystal shape of MIL-125-NH$_2$ is retained during heating. It is found that the MIL-125-NH$_2$-derived TiO$_2$ exhibits a cuboid shape with round corners and its size is significantly reduced compared to the MIL-125-NH$_2$ crystals (0.5 - 2 μm, Figure 5.5A). The size reduction is associated with the high calcination temperature and subsequently to the removal of the guest-molecules and organic ligand in MIL-125-NH$_2$. TEM images show that the crystal-templates consisted of nanosized TiO$_2$ NPs (Figure 5.6). This is further confirmed by dark-field TEM, where the incident electron beam is tilted so that only the reflections of specific diffracting crystallographic planes contribute to image formation (Figures S32 and S33). Depending on the diffraction angle, TiO$_2$ NPs of specific crystallographic orientation are illuminated in the dark-field image, confirming the multi-particle nature of the crystal-template. The co-existence of both TiO$_2$ rutile and anatase phases produced at 600 °C was additionally proven by selected area electron diffraction (SAED) pattern measurements (Figure 3c). The observed diffraction pattern is comparable with Bikley’s seminal work on P25, with the many nanosized TiO$_2$ particles diffracting to give Debye-Scherrer rings.$^{130a}$ These rings can be indexed to both TiO$_2$ anatase and rutile phases, confirming the multiphase nature of this material (Figure 5.6C).
Figure 5.5: SEM images of (A) MIL-125-NH$_2$ and MIL-125-NH$_2$-derived TiO$_2$ samples synthesized at (B) 400 °C, (C) 500 °C, (D) 600 °C and (E) 700 °C for 1 h. The images are of the same magnification except that of MIL-125-NH$_2$, which is of 5 times less magnification in order to display the morphology of the larger MOF crystals.

In contrast to the MIL-125-NH$_2$-derived TiO$_2$, the morphology of TiH$_4$O$_4$-derived TiO$_2$ resembles disordered arrangement of bulk aggregates (Figure 5.7), typical of precipitated and co-precipitated metal oxide NPs and reminiscent of P25 TiO$_2$ NPs. Further morphological characterization can be found in section S5.2.2 in the Appendix. The differing morphology of the MIL-125-NH$_2$- and TiH$_4$O$_4$-derived TiO$_2$ offer up an intriguing question of how this could affect the synergy between the TiO$_2$ anatase and rutile phases.
Figure 5.7: (A) SEM, (B) bright-field TEM and (C) dark-field TEM images of the TiH₄O₄-derived TiO₂ sample synthesized at 500 °C.

5.3 Photocatalytic Activity

After characterization, the performance of the MIL-125-NH₂-derived TiO₂ NPs was investigated toward the photocatalytic H₂ evolution reaction, with MeOH used as the sacrificial electron donor (20 v/v%) in the absence of co-catalyst, under UV-Vis light radiation. As shown in Figure 5.8A, the ratio between the TiO₂ anatase and rutile phases was found to have a profound impact on the photocatalytic performance. More specifically, increasing the rutile phases induces a volcano type trend with respect to the H₂ evolution rate. The sample with a composition of 66 w.t.% TiO₂ anatase and 34 w.t.% TiO₂ rutile (synthesized through calcination of MIL-125-NH₂ at 600°C) exhibits the highest H₂ evolution rate of 1394 μmol h⁻¹ g⁻¹. Materials with higher amounts of TiO₂ rutile phase exhibit inferior photocatalytic performance and the H₂ evolution rate for the TiO₂ with composition of 11 w.t.% TiO₂ anatase, 89 w.t.% TiO₂ rutile is 372 μmol h⁻¹ g⁻¹. The same trend between the activity and the ratio between the TiO₂ anatase and rutile phases is observed for the TiH₄O₄-derived TiO₂, with the 92 w.t% anatase and 8 w.t.% rutile TiO₂ exhibiting the best photocatalytic performance, with a H₂ evolution rate of 259 μmol h⁻¹ g⁻¹ (Figure 5.8A).
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Figure 5.8: Comparison of H$_2$ evolution rates of (A) MIL-125-NH$_2$-derived TiO$_2$ samples synthesized at different temperatures and B) TiH$_4$O$_4$-derived TiO$_2$ samples synthesized at 600°C (anatase), 700°C (anatase-rutile) and 800°C (anatase-rutile).

The higher photoactivity of the mixed-phase TiO$_2$ (66 w.t.% TiO$_2$ anatase and 34 w.t.% TiO$_2$ rutile) as compared to phase pure TiO$_2$ anatase or rutile phase is thought to be owed to the synergistic effect between the two polymorphs, allowing for a more efficient charge carrier separation, provided that the two phases are in good electronic contact. The photogenerated electrons flow from the TiO$_2$ rutile to the TiO$_2$ anatase phase, which finally catalyzes the H$_2$ evolution reaction.$^{131}$ Recycling experiments were carried out for the best performing TiO$_2$ for three consecutive cycles. The photostability and the morphology of the MIL-125-NH$_2$-derived TiO$_2$ are fully preserved, with consistent H$_2$ rates produced over 3 cycles (Figure 5.9). The SEM images of TiO$_2$ after photocatalysis demonstrate that the templated morphology of the material is conserved, and PXRD shows no loss of crystallinity with the same TiO$_2$ anatase-rutile phase composition (Figure S.36).

Figure 5.9: (A) Recycling photocatalytic experiments and (B) SEM image of MIL-125-NH$_2$-derived TiO$_2$ (with optimum TiO$_2$ anatase to rutile ratio of 66:34) after recycling experiments.
5.4 Comparison of MIL-125-NH2-derived TiO2 to other TiO2 Samples

Intrigued by these results, the photocatalytic activities of two additional TiO2 samples consisting of both the TiO2 anatase and rutile phases were explored; the commercial P25 Degussa TiO2 (with anatase-rutile phases in a composition of approximately 70:30) and another MOF-derived TiO2 using MIL-167 as the precursor. Figure 5.10A illustrates that the MIL-167-derived TiO2 NPs were successfully synthesized through calcination of the precursor at 600 °C for 1 h in air (for more information, see section S1.2 in the Appendix). Under these conditions, a mixed phase TiO2 with composition of TiO2 75 w.t.% anatase and 25 w.t.% rutile was prepared (Table S9). SEM images (Figure 5.10B, C) show that the morphology of MIL-167-derived TiO2 is based upon irregularly shaped aggregates with a large size of 5-20 μm. Despite the use of a crystalline MOF as precursor, the morphology of MIL-167-derived TiO2 is more comparable with that of the TiH4O4-derived TiO2 rather than that of the MIL-125-NH2 derived TiO2. TEM images indicate that the size of the MIL-167-derived TiO2 NPs is 30-40 nm, significantly larger than the TiO2 NPs derived from MIL-125-NH2 (Figure 5.11A).

![Figure 5.10: (A) PXRD patterns of the MIL-167- and MIL-125-NH2-derived TiO2, synthesized through calcination at 600°C. SEM images of MIL-167-derived TiO2 at (B) smaller and (C) larger magnification.](image)

![Figure 5.11: (A) TEM image and (B) selected area electron diffraction pattern of MIL-167-derived TiO2.](image)
The photocatalytic performance of the different mixed phase TiO₂ is shown in Figure 5.12A. The best MIL-125-NH₂-derived TiO₂ outperforms all the other TiO₂ samples, including the commercially available P25 Degussa. This is despite the relatively higher BET surface area of P25 Degussa compared to the other TiO₂ samples (Figure 5.13A). More specifically, the P25 Degussa TiO₂ exhibited a H₂ evolution rate of 624 μmol h⁻¹ g⁻¹, followed by the MIL-167-derived sample with a rate of 274 μmol h⁻¹ g⁻¹, which is comparable to that achieved by the TiH₄O₄-derived TiO₂ (259 μmol h⁻¹ g⁻¹). For the best performing MIL-125-NH₂-derived TiO₂ and P25 Degussa, the dependence of the H₂ evolution rate on the concentration of photocatalyst in solution was further explored and found that the MIL-125-NH₂-derived TiO₂ reaches a higher plateau than that of the P25 Degussa TiO₂ (Figure S38). This implies that the required amount of MIL-125-NH₂-derived TiO₂ photocatalyst is lower, indicative of a well dispersed sample in solution. The Kubelka-Munk representation of the UV-vis diffuse reflectance spectra of the examined samples is illustrated in Figure 5.13A. The indirect band gap for each of the materials was extrapolated from the linear section of the Tauc plot for an indirect bandgap. It can be seen that the indirect band gap is similar for all samples; 2.96 eV for MIL-125-NH₂-derived and MIL-167-derived TiO₂, 2.98 eV for TiH₄O₄-derived and 3.08 eV for P25 Degussa.

The superior photocatalytic performance of the MIL-125-NH₂-derived TiO₂ NPs over the other TiO₂ samples can be attributed to the morphology differences observed in SEM and TEM images (Figure 5.12B). The fact that the precursors in MOF-mediated syntheses are themselves well-defined crystals allows the resulting templated TiO₂ NPs to retain an exceptional close contact throughout the calcination and transformation process. These findings highlight that the TiO₂ anatase to rutile phase ratio is not the only predominant factor determining the photocatalytic performance, since excellent electronic contact between the two TiO₂ phases is a prerequisite for successful electron-hole separation and thus high photocatalytic activity.

Qualitatively, this is supported by an observation from the photocatalysis experiments, where the MIL-125-NH₂-derived TiO₂ suspension retained its white color throughout the photocatalytic test, in contrast to TiO₄H₄-, MIL-167-derived TiO₂ and P25, which obtained a blue hue (Inset at Figure 5.12A). The blue color is attributed to the presence of Ti³⁺ in the TiO₂, indicative of inefficient electron transport from the TiO₂ NPs to the water molecules or protons, in contrast to MIL-125-NH₂-derived TiO₂, where no Ti³⁺ was observed, indicating superior electron transport and water reduction.
Figure 5.12: (A) Comparison of photocatalytic activity and (B) SEM images of the MIL-125-NH₂-derived, MIL-167-derived, TiH₄O₄-derived TiO₂ and P25 Degussa. Inset: Photograph of the light-blue-colored P25 Degussa (left) and the MIL-125-NH₂-derived TiO₂ (right), after the photocatalytic test.

Figure 5.13: (A) N₂ sorption isotherms for the best performing MIL-125-NH₂-derived, TiH₄O₄-derived, MIL-167-derived and P25 Degussa TiO₂ and corresponding BET surface areas. (B) Kubelka-Munk representation of UV-Vis diffuse reflectance of MIL-167-derived, MIL-125-NH₂-derived, TiH₄O₄-derived TiO₂ and P25 Degussa. Inset: band gap values for each of the materials, extrapolated from the linear section of the Tauc plot for an indirect bandgap.

To further study this, PL experiments were performed to follow the irradiative recombination of the photo-excited charge carriers in the different TiO₂ samples and thus to gain further insights into the charge separation efficiency (Figure S39). In principle, the electron-hole recombination can occur through different pathways, including irradiative recombination through trap states, in addition to the irradiative band gap emission, which is less observed with TiO₂ – being an indirect band gap.
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For comparison reasons, the irradiative recombination of photogenerated charge carriers was quantified by integrating the PL emission curves of the different TiO$_2$ samples (Table 3). The PL emission follows the order: MIL-167-derived TiO$_2$ > TiH$_4$O$_4$-derived TiO$_2$ > P25 Degussa > MIL-125-NH$_2$-derived TiO$_2$. The PL results exhibit a comparable trend with the photocatalytic results. The MIL-125-NH$_2$-derived TiO$_2$ (best-performing photocatalyst) demonstrates the lowest PL emission, followed by the P25, which is the second best-performing photocatalyst.

Table 3: Integration of the PL emission curves of MIL-167-derived, TiH$_4$O$_4$-derived, MIL-125-NH$_2$-derived TiO$_2$ and P25 Degussa.

<table>
<thead>
<tr>
<th>MIL-167-derived TiO$_2$ (a.u. 10$^7$)</th>
<th>TiH$_4$O$_4$-derived TiO$_2$ (a.u. 10$^7$)</th>
<th>P25 Degussa (a.u. 10$^7$)</th>
<th>MIL-125-derived TiO$_2$ (a.u. 10$^7$)</th>
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5.5 Summary and Conclusions

In summary, this chapter describes a synthetic method in which MIL-125-NH$_2$ crystals are used as a precursor for the preparation of TiO$_2$ NPs composed of pure or mixed rutile and anatase phases. The crystals of MIL-125-NH$_2$ act as a template for the synthesis of TiO$_2$ and the templated mixed-phase TiO$_2$ (with anatase to rutile composition of 66:34) produces a H$_2$ rate of 1394 μmol h$^{-1}$ g$^{-1}$, without the assistance of any co-catalysts. The optimum MIL-125-NH$_2$-derived TiO$_2$ notably outperforms the commercial P25 Degussa, the conventionally synthesized TiH$_4$O$_4$- and the MIL-167-derived TiO$_2$. Two major aspects explain the high photocatalytic activity of the MIL-125-NH$_2$-derived TiO$_2$: i. the unique morphology that facilitates the constrained contact of NPs between the TiO$_2$ anatase and rutile phases and ii. the distribution of the TiO$_2$ anatase and rutile phases within the MIL-125-NH$_2$-derived template. These factors combine to result in an enhanced synergistic effect, which significantly suppresses the undesired electron-hole recombination. Our study highlights that high charge separation efficiency can be morphologically-induced within TiO$_2$ by using crystalline starting materials.
Conclusions and Outlook

Chapter 6 Conclusions and Outlook

In line with the scientific endeavors to optimize and extend the range of materials used for photocatalytic applications, over the past few years, the relatively new application of MOFs in photocatalysis has received considerably increasing scientific attention. When this field was in its infancy, most of the scientific studies demonstrated activities far from the state-of-the-art or lacked advanced characterization techniques for the evaluation of the MOF photocatalysts. However, since then, MOFs in photocatalysis have come a long way. Recent studies show high activities and involve sophisticated optoelectronic characterization techniques. In addition, the scientific focus in this field has been slowly shifted toward MOFs’ adaptation for ‘real-world’ applications, in terms of lower-cost and potential scale-up.

In line with these intense scientific efforts, the research activity described in this thesis contributed in the general advancement of the MOFs photocatalysis field, by favorably altering their properties associated with light absorption, charge separation and catalysis, which constitute the three major photocatalytic steps. Gaining insights into how established strategies for traditional photocatalysts can impact MOFs’ properties gave rise to noble-metal-free MOF-based systems with significantly improved photocatalytic activities. These scientific efforts led to diminishing the gap between academic research and industry, highlighting the great potential of MOFs as photocatalysts and bringing this field a step closer to viability as a real-life application. However, there is still a lot of room for improvement in order for this type of photocatalytic systems to become industrialized.

In general, MOFs’ synthetic protocols can involve the utilization of environmentally hazardous or expensive solvents, as well as result in low yields. These factors complicate the commercialization of MOFs. Therefore, optimizing the MOF’s synthetic protocols in terms of sustainability, cost and yields of products is crucial for the advancement of the general field. In particular for MOFs in photocatalysis, there is also a need for novel materials with high intrinsic properties (optoelectronic and catalytic properties) that can be further optimized through the aforementioned strategies described in this thesis. Furthermore, considering the fact that water stability is a crucial property – especially for photocatalytic applications – future work should be directed toward improving the stability of MOFs. This way the utilization of organic solvents in the photocatalytic solution could be avoided, and thus photocatalytic hydrogen production could be achieved in pure water. Last but not least, the form (suspensions) and size of the materials used in photocatalysis (e.g. nanoparticles) can be challenging for scale-up and applicability. To tackle these last challenging issues, research efforts should be devoted to incorporating these materials within a device (i.e., PEC device).
Appendix

S1 Experimental

S1.1 Materials and General Methods

2-amino benzenone-1,4-dicarboxylic acid (NH$_2$-H$_2$BDC, 99%), 2,5-dihydroxyterephthalic acid (H$_2$DHBDC, 98%), titanium isopropoxide (Ti(OCH(CH$_3$)$_2$)$_4$, 97%), nickel nitrate hexahydrate (Ni(NO$_3$)$_2$·6H$_2$O, 99.999% trace metal basis), cobalt nitrate hexahydrate (Co(NO$_3$)$_2$·6H$_2$O, 99.999% trace metal basis), copper nitrate trihydrate (Cu(NO$_3$)$_2$·3H$_2$O, ≥ 99%), iron nitrate nonahydrate (Fe(NO$_3$)$_3$·9H$_2$O, ≥ 99.95% trace metals basis), chloroplatinic acid hexahydrate (H$_2$PtCl$_6$·6H$_2$O, ACS reagent, ≥ 37.50% Pt basis), Zirconium(IV) Chloride (≥ 99.9% trace metals basis), anhydrous N,N-dimethylformamide (DMF; (CH$_3$)$_2$NCHO, 99.8%), anhydrous methanol (CH$_3$OH, 99.8%), N,N-diethylformamide (DEF, 99%), ethanol (EtOH, reagent grade), acetonitrile (CH$_3$CN, ≥ 99.5), sodium hypophosphite (NaH$_2$PO$_2$, ≥ 99%), isopropanol ((CH$_3$)$_2$CHOH, ≥ 98%), triethylamine (TEA, ≥ 99%), triethanolamine (TEOA, reagent grade, 98%), tetrabutylammonium perchlorate (TBAP, ≥ 99.0%, for electrochemical analysis), Rhodamine B (analytical standard) and tert-Butanol (≥ 99.0%, ACS reagent) were purchased from Sigma Aldrich. Benzene-1,3,5-tricarboxylic acid (C$_6$H$_3$(COOH)$_3$, H$_3$BTC, > 98.0%) was purchased from TCI. N,N-dimethylformamide (DMF, ≥ 99.5%) and Methanol (MeOH, ≥ 99.9%) were purchased from Carl Roth and Fisher scientific, respectively.

S1.2 Characterization

**Powder X-ray Diffraction** (PXRD) data were typically collected on a Bruker D8 Advance diffractometer at ambient temperature using monochromated Cu Kα radiation (λ = 1.5418 Å), with a 2θ step size of 0.02° and a 20 range of 2 to 70°. The Al$_2$O$_3$ NIST standard SRM1976b was used for size analysis. Reported phase composition and domain sizes were determined by Rietveld refinement using Topas, the latter are based on integral breadths in an isotropic size approximation.

**Nitrogen adsorption-desorption isotherms** were collected using a BEL-SORP mini (BEL Japan, Inc.) at 77 K. Prior to data collection, the samples were degassed at 423 K for 12 h. The BET surface areas were estimated from the amount of N$_2$ adsorbed at 77 K using the BET (Brunauer-Emmett-Teller) equilibrium equation.

**High-resolution transmission electron microscopy** (HR-TEM) images of the co-catalysts were collected on FEI Tecnai G2 Spirit Twin instrument, equipped with a 80-120 kV LaB6 gun.
Scanning electron microscopy (SEM) images and energy dispersive X-ray maps (EDX) were collected on FEI Teneo instrument, equipped with an energy dispersive X-ray detector (XFlash Silicon drift detector). Prior imaging, the samples were deposited on a carbon tape and were coated with a ~7 nm thick Iridium layer. TEM images were collected on a FEI Tecnai Osiris instrument.

UV-Vis absorbance and diffuse reflectance spectra were obtained with a PerkinElmer UV-Vis Spectrometer. For the UV-Vis absorbance, 1 cm optical path length quartz cuvettes were used. The diffuse reflectance was collected by depositing the powders within quartz slides and the Kubelka-Munk function was applied to the raw data in order to eliminate scattering.

Steadt-State Photoluminescence (PL) spectra were measured with a Fluorescence Spectrometer LS 55 (PerkinElmer). For the MOF-based photocatalytic systems, the excitation wavelength was 420 nm and the experiments were performed using either suspensions of 17 mg of MIL-125-NH$_2$ with different amounts of each co-catalyst in 2.5 mL of the photocatalytic solution (co-catalysts/MIL-125-NH$_2$ project) or 3.5 mg of the MOFs in 3.5 mL of either acetonitrile or the photocatalytic solution (MOF/MOF heterojunctions project). For the TiO$_2$-based photocatalytic systems, the excitation wavelength was 340 nm and the experiments were performed on suspensions of 0.45 mg of TiO$_2$ in 3 mL of the photocatalytic solution (20 v/v % MeOH and water). In all cases, the PL spectra were acquired using chemical-resistant, 1 cm optical path length, quartz cuvettes.

Time-Resolved photoluminescence performed with a LP980 laser flash spectrometer (Edinburgh Instruments). The samples were excited by 5 ns laser pulses at $\lambda = 420$ nm (nanosecond tunable Ekspla NT340 laser with 5 Hz repetition rate). In a typical experiment, 0.35 mg of the investigated materials was dispersed in 3.5 mL of acetonitrile. The suspension was sonicated for ~30 min and then placed in a 1 cm path-length quartz cuvette.

X-ray photoelectron spectroscopy (XPS) was carried out using a Thermo Fisher Scientific Theta Probe spectrometer, with a base pressure of $5 \times 10^{-10}$ mbar and utilizing monochromated Al K alpha radiation (1486.6 eV). The analysis area was defined by the X-ray spot size which was 400 mm. Photoelectron kinetic energy was measured using a hemispherical analyser operated in constant analyser energy (CAE) mode with a pass energy of 50 eV for high resolution spectra and 200 eV for survey spectra. The electron lens axis is 50° from the sample normal, and photoelectrons are accepted over an angular range of 60°. A 2D position sensitive detector was used. A dual beam (electron and Ar ion) charge compensation device was used to prevent charging of the sample surface.

Fourier-transform infrared spectroscopy (FTIR) measurements were carried out with the Spectrum Two FTIR spectrometer (PerkinElmer).
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**Electrochemical experiments** were conducted in a standard three-electrode electrochemical cell, using Autolab 302N (Metrohm). Pt wire was used as counter electrode and the reference electrode was Ag/Ag⁺ with 0.01M AgNO₃ and 0.1 M TB-AP in acetonitrile (ALS Japan). The amount of electrolyte used in a typical experiment was approximately 50 mL.

**Inductively coupled plasma optical emission spectrometry** (ICP-OES) measurements were performed using a NexIon 350 (Perkin Elmer) spectrometer.

**Thermogravimetric analyses** (TGA) was carried out with a standard TG-DTA analyzer under air atmosphere, at a heating rate of 5 °C/min for all measurements.

**Variable Temperature (VT) Powder X-ray Diffraction** data was collected on a Bruker D8 Advance diffractometer equipped with a LynxeyeXE detector using non-monomerized Cu Kα radiation. The sample was packed in a capillary and placed in a furnace, where it was heated at 30 °C min⁻¹ to set temperatures at 25 °C intervals. Data was collected in the 20 range 22 to 31°, with a 0.02° step size.

**CHN elemental analysis** was performed on a Thermo Scientific Flash 2000 Organic Elemental Analyzer.

### S1.3 Photocatalysis

The photocatalytic experiments with all the MOF-based systems were carried out using a 25 mL Pyrex glass reactor, under continuously visible light irradiation using a 300 W Xe lamp equipped with a UV cut-off filter (λ ≥ 420 nm, **Figure S.1**). In a typical experiment, 17.0 mg of the MOF photocatalysts were mixed with different amounts of each co-catalyst and were suspended in 17.0 ml of a photocatalytic solution consisting of acetonitrile, deionized water and a given electron donor. For the experiments with the different co-catalyst/MIL-125-NH₂ systems and the MOF/MOF heterojunctions, the electron donor used was TEA (Conditions: 2.8 mL TEA in 0.8 mL H₂O and 13.4 mL CH₃CN). Subsequently, the suspensions were purged with nitrogen for 20 min under gentle stirring (to remove dissolved oxygen) and sonicated for 20 min. The amount of hydrogen produced was determined by abstracting 200 μL of the gaseous product from the head space of the reactor and analyzing it by means of gas chromatography (PerkinElmer Clarus 480 GC, equipped with a thermal conductivity detector and a molecular sieve 5 A column, with N₂ as the carrier gas). When TiO₂ was explored as photocatalyst, similar procedure was followed, apart from the facts that 10 mg of the photocatalysts were suspended in a photocatalytic solution comprised of water and MeOH (20 v/v%) as the electron donor, while the samples were irradiated with UV-Vis light (no cut-off filter).
Figure S.1: Schematic illustration of the photocatalytic set-up used.
S2 Supporting Information for Chapter 2: Variation of Co-catalysts

S2.1 Synthetic Procedures

Synthesis of MIL-125-NH\(_2\)

The MOF was synthesized by following a reported procedure.\(^9\) 0.286 g of NH\(_2\)-H\(_2\)BDC was dissolved in a mixture of 4.0 mL anhydrous DMF and 1.0 mL anhydrous MeOH. 0.286 mL of titanium isopropoxide was added in the mixture, which was sonicated for 30 min and then heated up to 120 °C for 72h. The obtained product was washed several times with DMF and MeOH.

Prior to photocatalytic testing, the MIL-125-NH\(_2\) was subjected to a washing protocol, in order to remove uncoordinated free ligand and exchange the solvents within the pores of the MOF. The washing procedure involved the immersion of 500 mg of the as-synthesized MIL-125-NH\(_2\) in 25 mL of DMF for 12 h and after centrifugation, the product was twice submerged in 25 mL of MeOH for 12 h. Subsequently, the sample was dried at 70 °C and then suspended in 0.5 L of the photocatalytic solvent mixture of acetonitrile (CH\(_3\)CN), TEA and deionized water. The suspension was stirred overnight under nitrogen atmosphere and finally washed with 100 mL of acetone.

Synthesis of co-catalysts

Ni\(_2\)P nanoparticles (NPs) were synthesized based on a reported procedure.\(^{145}\) First, the Ni-BTC MOF was synthesized by mixing 0.255 g of Ni(NO\(_3\))\(_2\)·6H\(_2\)O and 0.102 g H\(_3\)BTC in 14.0 mL in MeOH; the suspension was then heated for 24 h at 150 °C in a Teflon liner inserted in a stainless steel autoclave. The obtained solid was washed several times with MeOH and then dried under vacuum at 60 °C. In a typical preparation of Ni\(_2\)P NPs, 0.10 g of the as-prepared Ni-BTC and 0.30 g of NaH\(_2\)PO\(_2\) were mixed together, loaded into a ceramic crucible, and heated to 275 °C for 2 h in a furnace. After cooling to room temperature, the product was washed with water and EtOH and dried at 75 °C for 12 h.

CoP NPs were synthesized following similar procedure to that of Ni\(_2\)P, except that Co(NO\(_3\))\(_2\)·6H\(_2\)O (0.255 mg) was used instead of Ni(NO\(_3\))\(_2\)·6H\(_2\)O, for the preparation of the Co-BTC powder.

NiO and Co\(_3\)O\(_4\) NPs were prepared by loading 0.1 g of the as-synthesized Ni-BTC and Co-BTC powders, respectively into a ceramic crucible and heated up to 375 °C for 2 h in a furnace. After cooling to room temperature, the product was washed with water and EtOH and dried at 75 °C for 12 h.

CuO NPs were prepared similar to NiO and Co\(_3\)O\(_4\), except that Cu(NO\(_3\))\(_2\)·3H\(_2\)O was used as the metal source for the synthesis of the Cu-BTC powder.
FeO3 NPs were synthesized similar to NiO and Co3O4 NPs, however in this case the precursor Fe-BTC powder was prepared by dissolving 0.51 g of Fe(NO3)3·9H2O and 0.102 g H3BTC in MeOH.

Pt NPs were synthesized based on a reported procedure. More specifically, 119.5 μL of H2PtCl6·6H2O was added in 16 mL aqueous solution with 0.2 M acetic acid concentration. The solution was sonicated for 5 min and then put into a Teflon-lined autoclave and heated up to 180 °C for 2 h. After cooling to room temperature, the product was washed with EtOH and dried at 60 °C for 12 h.

### S2.2 Photocatalytic Experiments

![Graphs showing photocatalytic H2 evolution rates of MIL-125-NH2 (17.0 mg) against different adding amounts of NiO, Co3O4, CoP, Fe2O3, and CuO NPs under visible light irradiation for 8 h.]

**Figure S.2:** Photocatalytic H2 evolution rates of MIL-125-NH2 (17.0 mg) against different adding amounts of (a) NiO, (b) Co3O4, (c) CoP, (d) Fe2O3 and (e) CuO NPs under visible light irradiation for 8 h.

**Table S.1:** Photocatalytic experiments carried out under irradiation of each co-catalyst for 14 h.

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S2.3 Calculation of Apparent Quantum Yield

Moles of photons emitted by the xenon lamp

The moles of photons emitted by the xenon lamp at 400 nm was calculated by means of Ferrioxalate actinometry:\textsuperscript{146} Stock solutions were prepared using Millipore water:

**0.40 M Iron (III) stock solution (1)** was prepared by dissolving 4.025 g iron(III) nitrate nonahydrate (Fe(NO$_3$)$_3$·9H$_2$O) in 20.0 mL of water. This was acidified with 1.375 mL sulfuric acid (H$_2$SO$_4$), and filled up to 25 mL with additional water.

**1.20 M Di-potassium oxalate stock solution (2)** was prepared by adding 4.975 g di-potassium oxalate monohydrate (K$_2$C$_2$O$_4$) to 25.0 mL of water.

**Phenanthroline/buffer solution (3)** was prepared by dissolving 0.025 g anhydrous 1,10-phenanthroline (C$_{12}$H$_8$N$_2$) and 5.625 g sodium acetate trihydrate (NaCH$_3$COO·3H$_2$O) in 20.0 mL of water. 1.22 mL sulfuric acid (H$_2$SO$_4$) was added and the solution was filled up to 25.0 mL with additional water.

The reaction solution was prepared by adding 1.7 mL of solution (1) and 1.7 mL of solution (2) with 30.6 mL of water in the dark. After stirring, 17. mL of the solution was transferred into a reactor (reactor A) which was irradiated by a 300 W Xe lamp equipped with a monochromatic filter (450 nm) for 46 min, while the remaining 17 mL was kept in the dark (reactor B). 0.5 mL of sample from reactor (A) was extracted, then added in 1.0 mL of solution (3) and made up to 5.0 mL with water. 0.5 mL of sample from reactor B was treated the same. UV-Vis absorbance (400-600 nm) was recorded for both samples, with the difference in absorbance between the two recorded at 510 nm. The moles of Fe$^{2+}$ were calculated by using the formula:

$$n_{Fe(II)} = \frac{V_1 V_3 \Delta A_{10^3 V_2 L \epsilon}}{V_2}$$

Where:

$V_1$: irradiation volume, $V_2$: sample taken, $V_3$: final volume, $L$: optical path-length, $\Delta A$: absorbance difference at 510 nm, $\epsilon$: absorbance coefficient.

From this, the number of photons per min was calculated using the formula:

$$\frac{n_{photons}}{\text{min}} = \frac{n_{Fe(II)}}{\Phi_{\lambda} t F}$$

$\Phi_{\lambda}$: Quantum yield for iron production at wavelength $\lambda$, $t$: time, $F$: mean fraction of light absorbed

Calculation: $n_{Fe(II)} = \frac{V_1 V_3 \Delta A_{10^3 V_2 L \epsilon}}{V_2}$

Where:
**Appendix Variation of Co-catalysts**

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- \( V_1 \): irradiation volume (17.0 mL), \( V_2 \): sample taken (0.5 mL), \( V_3 \): final volume (5.0 mL), \( L \): optical pathlength (1.0 cm), \( \Delta A \): absorbance difference at 510 nm: \( \Delta A_1 = 0.870758 \), \( \Delta A_2 = 0.892268 \), \( \varepsilon \): molar extinction coefficient (11100 L mol\(^{-1}\) cm\(^{-1}\))

Moles of Iron(II):

1\(^{st} \) measurement \( n_{Fe(II)} = 1.36654 \times 10^{-5} \)

2\(^{nd} \) measurement \( n_{Fe(II)} = 1.33359 \times 10^{-5} \)

\[
\frac{n_{\text{photons}}}{\text{min}} = \frac{n_{Fe(II)}}{\Phi \times t \cdot F}
\]

\( \Phi \): Quantum yield for iron production = 1.11 at ~450 nm

\( t \): time = 46 min

\( F \): mean fraction of light absorbed = 1

\[
\frac{n_{\text{photons}}}{\text{min}} = 5.1713 \times 10^{-7}
\]

**Moles of hydrogen generated under 400 nm and 450 nm radiation**

The amount of hydrogen generated under 400 nm radiation was carried out using the same reaction mixture and set-up as the photocatalytic experiments with the exception of using 450 nm band pass filter with the xenon lamp. Amount of H\(_2\) for Ni\(_2\)P/MIL-125-NH\(_2\): \( 2.0937 \times 10^{-6} \) mol in 240 min (4 h).

**Apparent quantum yield (AQY) determination**

The apparent quantum yield was calculated by using the following formula: \( \text{AQY(\%)} = \frac{2 \frac{N_{H_2}}{\text{min}}}{N_{\text{photons}}/\text{min}} \times 100 \)

Where \( \frac{N_{H_2}}{\text{min}} \) = number of hydrogen molecules evolved per minute and \( \frac{N_{\text{photons}}}{\text{min}} \) = number of incident photons per minute.

**Calculation:**

- Ni\(_2\)P/MIL-125-NH\(_2\):

  AQY = 6.6 \% at 400 nm
Appendix Variation of Co-catalysts

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S2.4 Stability

Figure S.3: PXRD patterns of the as-synthesized MIL-125-NH₂ and the different co-catalysts, and the co-catalyst/MIL-125-NH₂ systems after the photocatalytic test.

Figure S.4: XPS data on the CuO/MIL-125-NH₂ (left) and CoP/MIL-125-NH₂ system (right) after photocatalysis, confirming that the co-catalysts are stable.
Appendix Variation of Co-catalysts

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Figure S.5: (a) SEM images after photocatalytic test and corresponding EDX elemental analysis for (a) Ni$_2$P/MIL-125-NH$_2$, (b) NiO/MIL-125-NH$_2$, (c) CoP/MIL-125-NH$_2$, (d) Co$_3$O$_4$/MIL-125-NH$_2$, (e) Fe$_2$O$_3$/MIL-125-NH$_2$ and (f) CuO/MIL-125-NH$_2$. Scale bar: 1 μm.
Figure S.6: $N_2$ adsorption-desorption isotherms of Ni$_2$P/MIL-125-NH$_2$ (green) (a) before (with a calculated BET surface area of 1175 m$^2$ g$^{-1}$) and (b) after the photocatalytic experiments (with a calculated BET surface area of 1176 m$^2$ g$^{-1}$). $N_2$ adsorption-desorption isotherms of NiO/MIL-125-NH$_2$ (orange), (c) before (with a calculated BET surface area of 1288 m$^2$ g$^{-1}$) and (d) after the photocatalytic experiments (with a calculated BET surface area of 1233 m$^2$ g$^{-1}$).

Figure S.3 shows that the MIL-125-NH$_2$, NiO, Co$_3$O$_4$, Fe$_2$O$_3$ and Ni$_2$P NPs retained their crystallinity after the photocatalytic water reduction reaction. XPS measurements were carried out for the CuO/MIL-125-NH$_2$ and CoP/MIL-125-NH$_2$ systems, since the characteristic Bragg reflections of CuO and CoP NPs were not apparent in the PXRD patterns due to their low concentration and crystallinity compared to the amount of MIL-125-NH$_2$ used (Figure S.4). The Cu and Co 2p peaks were characteristic of Cu(II) and Co(III) respectively, and showed no significant change after the photocatalytic test. The SEM mapping images of each co-catalyst/MIL-125-NH$_2$ system confirmed that all the compositional elements of each system are present after the photocatalytic test (Figure S.5). Furthermore, the $N_2$ adsorption-desorption isotherms collected on the best-performing Ni$_2$P/MIL-125-NH$_2$ and NiO/MIL-125-NH$_2$ systems before and after the photocatalytic experiments showed...
Appendix Variation of Co-catalysts

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comparable BET surface areas, demonstrating the retention of the porous structure of MIL125-NH$_2$ (Figure S.6).

**Figure S.7:** FTIR spectra of Ni$_2$P/MIL-125-NH$_2$ before (red) and after (blue) photocatalytic test.

**Figure S.8:** PXRD patterns of (a) Ni$_2$P/MIL-125-NH$_2$ and (b) NiO/MIL-125-NH$_2$. 
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S2.5 Photoluminescence spectroscopy

Figure S.9: PL spectra of the MIL-125-NH$_2$ and the suspensions of MIL-125-NH$_2$ with the optimized amounts of: NiO (orange), CuO (blue), Fe$_2$O$_3$ (red), Co$_3$O$_4$ (violet) CoP (navy blue) and Ni$_2$P (green, optimized for photocatalytic hydrogen evolution). The co-catalyst-induced quenching of the PL follows the descending order of Ni$_2$P > CoP ≈ Co$_3$O$_4$ > Fe$_2$O$_3$ > CuO > NiO. The excitation wavelength was 420 nm.

Figure S.10: (a) Time Resolved PL decay curves and (b) lifetime of MIL-125-NH$_2$ and MIL-125-NH$_2$ with different co-catalysts. The excitation wavelength was 420 nm. Since the lifetimes of the co-catalyst/MIL-125-NH$_2$ systems were not highly different from each other, drawing further conclusions based on these results might be inconclusive.
S2.6 Electrochemical experiments with co-catalysts

The intrinsic catalytic activity of Ni$_2$P and NiO in acetonitrile and water (90:10), with 0.1M tetrabuthylammonium perchlorate (TBAP, ensuring the availability of conductive ions) have been explored. This investigation was conducted by means of cyclic voltammetry (CV) at 50 mV s$^{-1}$ scan rate and by linear scan voltammetry (LSV) at 1 mV s$^{-1}$ in a standard 3-electrode electrochemical cell. The suspension containing the co-catalysts was prepared by mixing 10 mg of each co-catalyst with 1 mL solution of EtOH and water (2:1) and 20 μL of Nafion (5 wt.%). The suspension was deposited as a thin film on a polished glassy carbon electrode with diameter of 3 mm (S= 0.07065 cm$^2$), which was used as a working electrode. The loading of each co-catalyst was 0.7 mg cm$^{-2}$. Pt wire was used as counter electrode, while Ag/Ag$^+$ with 0.01 M AgNO$_3$ and 0.1 M TBAP in acetonitrile (ALS Japan) was used as reference electrode.

Prior the LSV measurement, the CV of the electrode with the co-catalysts was obtained in a non-aqueous electrolyte with 0.1 M TBAP. Subsequently, the electrode was transferred to the cell with water-containing electrolyte and the CV at 50 mV s$^{-1}$ was measured again in order to define the hydrogen evolution reaction (HER) region (-0.7 V to -1.2 V). This potential window was selected to perform further investigation of the intrinsic catalytic activity of the co-catalysts. In order to compare the intrinsic catalytic activity of Ni$_2$P and NiO, the current densities were collected under similar conditions to those applied for the photocatalytic test.

It was established that the current density (intrinsic catalytic activity toward HER) of the examined Ni$_2$P is lower than that of the NiO co-catalyst. For example, at -0.9 V versus Ag/Ag$^+$, the current density of the Ni$_2$P and NiO co-catalysts is -0.07 and -0.22 mA cm$^{-2}$, respectively.

Figure S.11: Cathodic polarization curves of the electrodes with Ni$_2$P and NiO NPs obtained by linear scan voltammetry.
S2.7 Pt NPs as co-catalysts with MIL-125-NH$_2$

**Figure S.12**: (a) PXRD patterns of the simulated (blue) and as-synthesized (red) Pt NPs. (b) N$_2$ adsorption-desorption isotherm of Ni$_2$P NPs.

**Figure S.13**: (a) TEM image and (b) electron diffraction pattern of Pt NPs. The size of Pt NPs is estimated to be around 20-30 nm.
Figure S.14: $H_2$ evolution rate of 17.0 mg MIL-125-NH$_2$ against different adding amounts of Pt NPs.

Figure S.15: PXRD patterns of MIL-125-NH$_2$ simulated (dark gray), Pt NPs as-synthesized (light gray) and 2 ($\pm 0.4$) wt.% Pt/MIL-125-NH$_2$ after photocatalytic test.
S2.8 **Comparison of Ni₂P/MIL-125-NH₂ with other studies in the literature**

**Table S.2:** *Comparison of Ni₂P/MIL-125-NH₂ with other MOF-based hydrogen generation photocatalytic systems.*

<table>
<thead>
<tr>
<th>Photocatalyst</th>
<th>Co-catalyst</th>
<th>Light Source ( \lambda ) (nm)</th>
<th>( \text{H}_2 ) Evolution Rate with respect to the MOF</th>
</tr>
</thead>
<tbody>
<tr>
<td>MIL-125-NH₂ (Ti)</td>
<td>Pt NPs photodeposited</td>
<td>( \geq 420 )</td>
<td>*333 ( \mu \text{mol h}^{-1} \text{g}^{-1} )</td>
</tr>
<tr>
<td>Uio-66-NH₂ (Zr/Ti)</td>
<td>Pt NPs</td>
<td>( \geq 420 )</td>
<td>3.5 mmol mol(^{-1}) in 5 h</td>
</tr>
<tr>
<td>Ti-MOF-Ru(tpy)(_2)</td>
<td>Pt NPs</td>
<td>( \geq 420 )</td>
<td>*100-200 ( \mu \text{mol h}^{-1} \text{g}^{-1} )</td>
</tr>
<tr>
<td><strong>AQY: 0.2 % at 500 nm</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MIL-125-NH₂ (Ti)</td>
<td>Pt NPs photodeposited</td>
<td>( \geq 420 )</td>
<td>*525 ( \mu \text{mol h}^{-1} \text{g}^{-1} )</td>
</tr>
<tr>
<td>Al-PMOF</td>
<td>Pt NPs</td>
<td>( \geq 420 )</td>
<td>200 ( \mu \text{mol g}^{-1} \text{h}^{-1} )</td>
</tr>
</tbody>
</table>

Figure S.16: (a) SEM image of 2 (± 0.4) wt.\% Pt/MIL-125-NH₂ after photocatalytic testing, and corresponding EDX maps of (b) Titanium, (c) Pt and (d) both Ti and Pt.
### Appendix Variation of Co-catalysts

<table>
<thead>
<tr>
<th>Material Description</th>
<th>Pt-loading</th>
<th>Turnover Rate</th>
<th>Additional Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Uio-66 (Zr)</td>
<td>2.87 wt.% Pt</td>
<td>≥ 380</td>
<td>257.38 μmol h⁻¹ g⁻¹</td>
</tr>
<tr>
<td>MIL-125(Ti)</td>
<td>Pt</td>
<td>≥ 320</td>
<td>*155 μmol h⁻¹ g⁻¹</td>
</tr>
<tr>
<td>MIL-100 (Fe)</td>
<td>0.8 wt.% Pt photodeposited</td>
<td>≥ 420</td>
<td>109 μmol h⁻¹ g⁻¹</td>
</tr>
<tr>
<td>MOF-253 (Al)</td>
<td>Pt complex</td>
<td>≥ 420</td>
<td>100 μmol h⁻¹</td>
</tr>
<tr>
<td>[Cu(enMe)₂][PNb₁₂O₆(VO₆)] (OH)₃ 6H₂O</td>
<td>0.75% Pt</td>
<td>Hg lamp</td>
<td>43.86 μmol g⁻¹ h⁻¹</td>
</tr>
<tr>
<td>[Gd₆(abtc)(OH)₂]₂(OH)₂ 2H₂O</td>
<td>1.5 w.t.% Ag</td>
<td>Xe lamp</td>
<td>* 212 μmol h⁻¹ g⁻¹</td>
</tr>
<tr>
<td>Zr₆(μ₃-O)₄(μ₃-OH)₄(bpdc)₅.₉₄ (L₁)₀.₆</td>
<td>Pt NPs photodeposited</td>
<td>≥ 420</td>
<td>3400 turnovers after 48hrs</td>
</tr>
<tr>
<td>Uio-66 (Zr)</td>
<td>Pt NPs in sol</td>
<td>Xe doped mercury</td>
<td>2.4 ml in 3hrs</td>
</tr>
<tr>
<td>Uio-66-NH₂ (Zr)</td>
<td>Pt NPs in sol</td>
<td>Xe doped mercury</td>
<td>2.8 ml in 3hrs</td>
</tr>
<tr>
<td>Uio-66 (Zr) + Rhodamine B</td>
<td>Pt NPs on MOF</td>
<td>≥ 420</td>
<td>116 μmol h⁻¹ g⁻¹</td>
</tr>
<tr>
<td>Uio-66 (Zr) + Erythrosin B</td>
<td>0.5 w.t.% Pt (photo-deposited)</td>
<td>≥ 420</td>
<td>*444 μmol h⁻¹ g⁻¹</td>
</tr>
<tr>
<td>Cu(RSH)(H₂O)₆ + Eosin Y (1D-polymer)</td>
<td>N/A</td>
<td>≥ 420</td>
<td>7880 μmol g⁻¹ h⁻¹</td>
</tr>
<tr>
<td>[Fe-Fe]@ZrPF</td>
<td>[FeFe] based complex</td>
<td>≥ 420</td>
<td>3.5 μmol h⁻¹ in 120 min</td>
</tr>
<tr>
<td>Uio-66 Fe-Fe + Ru(bpy)₃</td>
<td>Fe-Fe</td>
<td>Blue LED 470 nm</td>
<td>* 280 μmol h⁻¹ g⁻¹</td>
</tr>
<tr>
<td>POM@Uio-67</td>
<td>POM</td>
<td>≥ 400</td>
<td>699 μmol h⁻¹ g⁻¹ wrt POM</td>
</tr>
<tr>
<td>MIL-125-NH₂ (Ti)</td>
<td>Co(III)-oxime</td>
<td>≥ 408</td>
<td>*~637 μmol h⁻¹ g⁻¹</td>
</tr>
<tr>
<td>MIL-125-NH₂ (Ti)</td>
<td>Nickel(II) Species</td>
<td>≤ 360</td>
<td>*6693 μmol h⁻¹ g⁻¹</td>
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<tr>
<td>Cu-1-bpy</td>
<td>UV-light Irradiation</td>
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<td>7090 μmol h⁻¹ g⁻¹</td>
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</table>
### Appendix Variation of Co-catalysts

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Co-catalyst</th>
<th>λ range (nm)</th>
<th>Rate (μmol h⁻¹ g⁻¹)</th>
<th>AQY (%)</th>
<th>Source</th>
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</thead>
<tbody>
<tr>
<td>MIL-125-NH₂</td>
<td>Ni₃P NPs</td>
<td>320-780</td>
<td>*32</td>
<td>6.6</td>
<td>This work ^86</td>
</tr>
<tr>
<td>MIL-125-NH₂</td>
<td>Ni₃P NPs</td>
<td>320-780</td>
<td>1230</td>
<td>6.6</td>
<td>This work ^86</td>
</tr>
</tbody>
</table>

* = calculated μmol h⁻¹ g⁻¹

Unless otherwise stated, H₂ rate in respect of MOF

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[^86]: [Link to reference]

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S3 Supporting Information for Chapter 3: Variation of Electron-Donors and Dual-Functional Photocatalysis

S3.1 Variation of electron donors

Table S.3: Concentration of each electron-donor within the photocatalytic solution (with acetonitrile and water). The selection of these specific concentrations was based on other scientific studies with MOF-based photocatalytic systems in the literature. The amount of water was kept constant at ~5 v/v% in all the examined photocatalytic solutions.

<table>
<thead>
<tr>
<th></th>
<th>TEA (v/v%)</th>
<th>TEOA (v/v%)</th>
<th>MeOH (M)</th>
<th>EtOH (M)</th>
<th>L-ascorbic acid (M)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Concentration</td>
<td>16</td>
<td>-</td>
<td>36.3</td>
<td>36.3</td>
<td>-</td>
</tr>
<tr>
<td>(v/v%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Concentration</td>
<td>7.2</td>
<td>0.01</td>
<td>24.7</td>
<td>17.1</td>
<td>0.1</td>
</tr>
<tr>
<td>(M)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

S3.1.1 Electrochemical experiments

The oxidation/reduction behavior of five sacrificial agents: EtOH, MeOH, TEA, TEOA and L-ascorbic acid in a non-aqueous solution of acetonitrile with 0.1 M tetrabuthylammonium perchlorate (TBAP) has been explored. This investigation was conducted by means of cyclic voltammetry at different scan rates in a standard three-electrode electrochemical cell, using the Bipotentiostat 2325 (ALS Japan). Pt wires were used as working and counter electrodes and Ag/Ag+ with 0.01 M AgNO3 and 0.1M TBAP in acetonitrile (ALS Japan) was used as reference electrode. The amount of electrolyte used in a typical experiment was approximately 55 mL. Prior to the investigation of the electrolyte with sacrificial reagents, the blank electrolyte (0.1 M TBAP in acetonitrile) was tested in order to define any possible influence from it. Two oxidation peaks are observed with current densities of less than 0.1 mA cm⁻² at -0.57 V and 0.625 V (Figure S17a).

The electrolyte with TEA shows an order of magnitude higher current density at the oxidation peak (~40 mA cm⁻² at scan rates of 50 and 20 mV s⁻¹, ~12 mA cm⁻² at scan rate of 5 mV s⁻¹), than the other compounds in electrolyte. However, due to higher currents the two-step oxidation observed in the case of TEA, transformed into one wide peak with maximum at 1.4 V (Figure S17b). The electrolyte with TEOA shows two oxidation peaks at 0.797 V and 1.073 V (scan rate: 50 mV s⁻¹), which can be associated with the two following steps: 1) positively charged aminyl radical is formed (at 0.797 V),
which 2) directly reacts with the TEOA and forms iminium species (at 1.073 V). The range of the obtained current densities is between 1.2 to 2 mA cm$^{-2}$ (Figure S17c).

The electrolytes with EtOH (Figure S17d) and MeOH (Figure S17e) show one small oxidation peak at: 1.88 V and 0.963 V (scan rate: 50 mV s$^{-1}$), respectively. The current density of the oxidation peaks is 0.8 and 1.61 mA cm$^{-2}$, for EtOH and MeOH, respectively. According to Han et. Al., there are two kinds of protons in the ethanol molecule. One is bonded to the oxygen atom and the other bonded to the carbon atom. If the proton bonded to the oxygen atom is firstly oxidized, then the first reaction step is: $\text{CH}_3\text{CH}_2\text{OH} \rightarrow [\text{CH}_3\text{CH}_2\text{O}]_{\text{ad}} + \text{H}^+ + \text{e}^-$

If the proton bonded to the carbon atom is firstly oxidized, then the first reaction step should be: $\text{CH}_3\text{CH}_2\text{OH} \rightarrow [\text{CH}_3\text{CHOH}]_{\text{ad}} + \text{H}^+ + \text{e}^-$

It cannot be concluded which one of the two reaction pathways is right and thus the observed peak for EtOH corresponds to one of the above-mentioned reactions. Similar mechanism is assumed for MeOH.

L-ascorbic acid exhibits 2 oxidation and 2 reduction peaks at all scan rates (Figure S17f). While the small peak could be the reducible artefact from acetonitrile, the highest peak at 1.1 V corresponds to the oxidation of ascorbic acid. It has a corresponding reduction peak at -0.8 V ($\text{C}_6\text{H}_8\text{O}_6 \rightarrow \text{C}_6\text{H}_6\text{O}_6 + 2\text{H}^+ + 2\text{e}^-$). The maximum current density of 1.8 mA cm$^{-2}$ was observed at a scan rate of 100 mV s$^{-1}$
S3.2 Photocatalytic Hydrogen evolution coupled with Rhodamine B degradation

Figure S.18: (a) Photocatalytic H\textsubscript{2} production of Ni\textsubscript{2}P/MIL-125-NH\textsubscript{2} in aqueous acetonitrile solution with variant concentrations of Rhodamine B (RhB), (b) PXRD patterns of the as-synthesized MIL-125-NH\textsubscript{2} and Ni\textsubscript{2}P, and the Ni\textsubscript{2}P/MIL-125-NH\textsubscript{2} after the photocatalytic test in 17 mL solution with 0.02 mg of RhB (1.2 ppm).

Figure S.19: (a) Trapping experiments of active species: UV-Vis spectra of 1.2 ppm RhB solution (blank) and supernatants after photocatalytic test of the same solution with 0.01 M TEOA (e\textsuperscript{−} scavenger), 0.01 M tert-butanol (•OH radical scavenger) and no scavenger, and (b) corresponding PXRD patterns. The duration of the photocatalytic tests was 3 h.
Figure S.20: Control experiment: $H_2$ evolution of Ni$_2$P/MIL-125-NH$_2$ in an aqueous acetonitrile solution, without any RhB or other electron donors, after 3 h visible light irradiation (solid red line). This result was compared with the photocatalytic performance of Ni$_2$P/MIL-125-NH$_2$ in aqueous acetonitrile solution with variant concentration of RhB (dashed lines).

Figure S.21: Calibration curve of RhB in acetonitrile solution with water (5 v/v% $H_2O$). Absorbance peak at 543 nm.
S4 Supporting Information for Chapter 4: Construction of MOF/MOF Heterojunctions

S4.1 Synthetic Procedures

Synthesis of MIL-167

MIL-167 was synthesized based on reported procedures,\textsuperscript{121} as follows:
The linker H$_4$DOBDC (1.585 g) was dissolved in 2.2 mL DEF and 2.2 mL MeOH. After stirring of
the mixture for few minutes, 0.592 mL of titanium isopropoxide was added. Then, the suspension was
placed in a Teflon liner and were heated up to 180 °C for 24h. The obtained dark red powder was first
washed several times with DEF then it was stirred overnight in MeOH and the next day it was washed
several times with MeOH. Finally, the obtained powder was dried at 60 °C for ~5 h.

Synthesis of MIL-167/MIL-125-NH$_2$

The MIL-167/MIL-125-NH$_2$ samples were synthesized following similar procedure to that of MIL-125-NH$_2$, except that various amounts of MIL-167 (5-110 mg) were introduced in the MIL-125-NH$_2$-precursor solution, prior to heating.

S4.2 Characterization

![TGA curves](image_url)

*Figure S.22: TGA curve of a) MIL-125-NH$_2$ and b) MIL-167 obtained under airflow with a heating rate of 5 °C min$^{-1}$.\textsuperscript{120}.*
Appendix Design of MOF/MOF Heterojunctions
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Figure S.23: PXRD patterns of a) the as-synthesized (light colors) and simulated (dark colors) MIL-125-NH$_2$ (yellow) and MIL-167 (red), b) MIL-125-NH$_2$ synthesized in the presence of increasing amounts of MIL-167 (5 - 100 mg, increasing toward the top of the figure) and c) MIL-125-NH$_2$ synthesized in the presence of 20, 40 and 60 mg of MIL-167.

Table S.4: X-ray diffraction-derived data (measurements in 0.4 mm borosilicate glass capillaries).

<table>
<thead>
<tr>
<th>Synthesis of MIL-125-NH$_2$ in:</th>
<th>MIL-125-NH$_2$ composition (wt.%)</th>
<th>MIL-167 composition (wt.%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>20 mg MIL-167</td>
<td>97</td>
<td>3</td>
</tr>
<tr>
<td>40 mg MIL-167</td>
<td>92</td>
<td>8</td>
</tr>
<tr>
<td>60 mg MIL-167</td>
<td>87</td>
<td>13</td>
</tr>
</tbody>
</table>

Figure S.24: SEM image of MIL-167/MIL-125-NH$_2$ heterojunction sample. In this case, prior to microscopy the samples were added in a MeOH solution, sonicated and then drop-casted on the SEM holder, which was allowed to dry.
Appendix Design of MOF/MOF Heterojunctions

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Figure S.25: SEM images of a), b): 3 wt.% MIL-167/MIL-125-NH$_2$, c), d): 8 wt.% MIL-167/MIL-125-NH$_2$ and d), e) 13 wt.% MIL-167/MIL-125-NH$_2$, with different magnification. The samples were prepared by spreading a small amount of the MIL-167/MIL-125-NH$_2$ dry powder on a carbon type SEM holder and then blowing it with N$_2$ to remove excess material.

MIL-125-NH$_2$ crystals demonstrate a cylindrical disc type of morphology with an approximate diameter of 400 nm. On the other hand, MIL-167 crystals are spheres of significant greater size, with their diameter approximated at 30-60 μm. The notable difference between the morphology of these two MOFs, allows the visual distinguishing of MIL-167 and MIL-125-NH$_2$ within the heterojunction samples (Figures S24 and S25). It is thus apparent that the bulky spherical crystals of MIL-167 are covered by the significantly smaller crystals of MIL-125-NH$_2$, with the latter being also spread around MIL-167 at Figure S24.
S4.3 Stability

Figure S.26: a) PXRD patterns of 8 wt.% MIL-167/MIL-125-NH$_2$ before and after photocatalysis, and b) N$_2$ sorption isotherms of the as-synthesized MIL-125-NH$_2$ and MIL-167, and 8 wt.% MIL-167/MIL-125-NH$_2$ after photocatalysis. The BET (Brunauer-Emmett-Teller) surface area of the 8 wt.% MIL-167/MIL-125-NH$_2$ is smaller than that of the as-synthesized MIL-125-NH$_2$ (1230 versus 1560 m$^2$ g$^{-1}$), which is associated with the presence of the non-porous MIL-167, contributing to the overall mass of the sample.

Figure S.27: Photocatalytic recycling experiments of MIL-167/MIL-125-NH$_2$ for three consecutive cycles of 12 h each. MIL-167/MIL-125-NH$_2$ consistently produces hydrogen for more than 36 h.
Table S.5: Inductively coupled plasma mass spectrometry (ICPMS) results. Concentration of Ti ions in the MIL-167/MIL-125-NH₂ suspensions before and in the supernatant after 36 hours irradiation. A negligible concentration of Ti ions (0.2 % of the initial concentration) is present in the supernatant after the recycling experiments, showing that there is no significant oxidation of the MOFs’ linkers.

<table>
<thead>
<tr>
<th>Ti ions Concentration (ppm)</th>
<th>Supernatant after 36 hours photocatalysis</th>
</tr>
</thead>
<tbody>
<tr>
<td>(1 mg mL⁻¹) MIL-167/MIL-125-NH₂ suspension before photocatalysis</td>
<td>42.5</td>
</tr>
<tr>
<td></td>
<td>0.09</td>
</tr>
</tbody>
</table>

S4.4 Mechanism

S4.4.1 Density Functional Theory Calculations

DFT calculations were performed using the CP2K computational chemistry package. The atomic coordinates and cell parameters were optimized using the PBE functional with D3 dispersion correction. The Goedecker-Teter-Hutter pseudopotentials were used with a cut-off of 450 Ry and DZVP-MOLOPT basis set for H, C, O and N atoms and TZVP-MOLOPT basis set for Ti atoms. The initial structure of MIL-125-NH₂ was built from the experimental one by means of adding -NH₂ groups. In the case of MIL-167, the solvent and counterions were removed and 32 -[HN(CH₃)₃]+ cations were added in order to neutralize the unit cell. After ground state relaxation of MIL-125-NH₂ and MIL-167 crystals at PBE level, ground state calculations were performed with PBE0 functional. Auxiliary Density Matrix Method (ADMM) was used in order to reduce the computational cost, pFIT3 was used for H, C, O and N atoms and cFIT11 for Ti atoms. Following the methodology previously reported by Walsh and coworkers, the electrostatic potential average in the pore of the material was taken as analogous to the vacuum level and the PBE0 Kohn-Sham energy levels were aligned relative to this potential:

\[ \varepsilon_{KS}^{\text{absolute}} = \varepsilon_{KS} - \Phi_{av} \]

Then, the highest occupied eigenvalue was taken as the ionization potential (IP), and the electron affinity (EA) from the difference between the IP and the computed bang gap \( E_g \):

\[ EA = IP - E_g \]
S4.4.2 Calculation of Apparent Quantum Yield

Moles of photons emitted by the xenon lamp

For the AQY calculations a similar procedure to that described in section S.24 was followed. First the three following stock solutions were prepared (as described in section S.24): 0.40 M iron (III) stock solution (1), 1.20 M di-potassium oxalate stock solution (2) and phenanthroline/buffer solution (3). Then, the reaction solution was prepared by adding 1.70 mL of solution (1) and 1.70 mL of solution (2) with 30.6 mL of water in the dark. After stirring, 17.0 mL of the solution was transferred into a reactor (reactor A) which was irradiated by a 300 W Xe lamp equipped with a monochromatic filter (450 or 500 nm) for 25 min, while the remaining 17 mL was kept in the dark (reactor B). 0.1 mL of sample from reactor A was extracted, and added 0.2 mL of solution (3), and made up to 5.0 mL with water. 0.1 mL of sample from reactor B was treated the same. UV-Vis absorbance (400-600 nm) was recorded for both samples, with the difference in absorbance between the two recorded at 510 nm. The moles of Fe$^{2+}$ were calculated by using the formula: 

$$n_{Fe(II)} = \frac{V_1 V_3 \Delta A}{10^3 V_2 L \varepsilon}$$

as follows:

$V_1$: irradiation volume (17.0 mL),
$V_2$: sample taken (0.1mL),
$V_3$: final volume (5.0 mL),
$L$: optical pathlength (1.0 cm),
$\Delta A$: absorbance difference at 510 nm:

- At 450 nm irradiation: $\Delta A_1 = 0.75837$, $\Delta A_2 = 0.75531$
- At 500 nm irradiation: $\Delta A_1 = 0.76362$, $\Delta A_2 = 0.72065$

$\varepsilon$: molar extinction coefficient (11100 L mol$^{-1}$ cm$^{-1}$)

- **Moles of iron(II) at 450 nm irradiation:**
  1$^{st}$ measurement $n_{Fe(II)} = 5.8074 \times 10^{-5}$
  2$^{nd}$ measurement $n_{Fe(II)} = 5.7839 \times 10^{-5}$

- **A Moles of iron(II) at 500 nm irradiation:**
  1$^{st}$ measurement $n_{Fe(II)} = 5.8475 \times 10^{-5}$
  2$^{nd}$ measurement $n_{Fe(II)} = 5.5185 \times 10^{-5}$

From this, the number of photons per min was calculated using the formula:

$$\frac{n_{photons}}{min} = \frac{n_{Fe(II)}}{\Phi_\lambda t F}$$

as follows:

$\Phi_2$: Quantum yield for iron production at wavelength $\lambda$ (1.11 at ~450 nm and 0.86 at ~500 nm)
Appendix Design of MOF/MOF Heterojunctions

Stavroula A. Kampouri

\( t \): time (25 min)

\( F \): mean fraction of light absorbed = 1

- At 450 nm: \( \frac{N_{\text{photons}}}{\text{min}} = 2.08851 \times 10^{-6} \)
- At 500 nm: \( \frac{N_{\text{photons}}}{\text{min}} = 2.64326 \times 10^{-6} \)

Moles of hydrogen generated under 450 nm and 500 nm radiation

The amount of hydrogen generated by MIL-167/MIL-125-NH₂ under 450 or 500 nm radiation was carried out using the same reaction mixture and set-up as the photocatalytic experiments (Figure S6) with the exception of using a 450 nm or 500 nm band pass filter with the xenon lamp.

- Amount of H₂ at 450 nm: 6.34382 \times 10^{-6} mol in 240 min (4 h)
- Amount of H₂ at 500 nm: 4.66284 \times 10^{-6} mol in 480 min (8 h)

Apparent quantum yield determination

The apparent quantum yield (AQY) was calculated by using the following formula:

\[
\text{AQY} \, (\%) = \frac{2 \times N_{\text{H}_2}}{N_{\text{photons}}} \times 100, \\
\text{where } \frac{N_{\text{H}_2}}{\text{min}} = \text{number of hydrogen molecules evolved per minute and } \frac{N_{\text{photons}}}{\text{min}} = \text{number of incident photons per minute.}
\]

Thus, \( \text{AQY} = 2.53 \% \) and \( 0.74 \% \) at 450 and 500 nm, respectively.

Table S.6: Comparison of the AQY values at wavelengths corresponding to the visible range (\( \lambda \geq 420 \text{ nm} \)) of different photocatalytic systems in the literature.

<table>
<thead>
<tr>
<th>MOF-based System</th>
<th>AQY</th>
<th>Wavelength</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pt/UiO-66 (Zr)</td>
<td>0.25 %</td>
<td>420 nm</td>
<td>112</td>
</tr>
<tr>
<td>Pt/CdS/UiO-66 (Zr)</td>
<td>1.2 %</td>
<td>420 nm</td>
<td>112</td>
</tr>
<tr>
<td>Pt/Zr6[μ3-O]6[μ1-OH]4(bpdc)3.94(L1)0.06</td>
<td>5.6 \times 10^{-4} %</td>
<td>440 nm</td>
<td>85c</td>
</tr>
<tr>
<td>Pt/MIL-125-NH₂ (Ti)</td>
<td>1.3 %</td>
<td>450 nm</td>
<td>107b</td>
</tr>
<tr>
<td><strong>This Work</strong></td>
<td>2.5 %</td>
<td>450 nm</td>
<td></td>
</tr>
<tr>
<td>Ni₂P/MIL-125-NH₂</td>
<td>6.6 %</td>
<td>450 nm</td>
<td>86</td>
</tr>
<tr>
<td>Pt/MIL-125-NH₂ (Ti)</td>
<td>0.3 %</td>
<td>500 nm</td>
<td>107b</td>
</tr>
<tr>
<td>Pt/Ti-MOF-Ru(tpy)₂</td>
<td>0.2 %</td>
<td>500 nm</td>
<td>884</td>
</tr>
<tr>
<td><strong>This Work</strong></td>
<td>0.7 %</td>
<td>500 nm</td>
<td></td>
</tr>
<tr>
<td>Pt/Al-PMOF</td>
<td>0.1 %</td>
<td></td>
<td>82</td>
</tr>
</tbody>
</table>

**Other Systems**

<table>
<thead>
<tr>
<th></th>
<th>AQY</th>
<th>Wavelength</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ni₂P/TiO₂</td>
<td>0.9%</td>
<td>400 nm</td>
<td>86</td>
</tr>
<tr>
<td>MoS₃/CdS</td>
<td>7.3 %</td>
<td>420 nm</td>
<td>166</td>
</tr>
<tr>
<td>Ni/CdS</td>
<td>11.2%</td>
<td>450 nm</td>
<td>167</td>
</tr>
<tr>
<td>Ni/CdS</td>
<td>5.9 %</td>
<td>520 nm</td>
<td>167</td>
</tr>
</tbody>
</table>
**Appendix Design of MOF/MOF Heterojunctions**

Stavroula A. Kampouri

### S4.4.3 Transient Photoluminescence Spectroscopy

The fluorescence lifetimes of our samples were collected through time-resolved PL, performed with a LP980 laser flash spectrometer (Edinburgh Instruments). The samples were excited by 5 ns laser pulses at $\lambda = 420$ nm (nanosecond tunable Ekspla NT340 laser with 5 Hz repetition rate) and the lifetimes were recorded at different wavelengths (460, 490 and 550nm) based on the PL emission peaks of the samples (Figure 3B). In a typical experiment, 0.35 mg of MIL-125-NH$_2$, MIL-167 or MIL-167/MIL-125-NH$_2$ was dispersed in 3.5 mL of acetonitrile. The suspension was sonicated for ~30 min and then placed in a 1 cm path-length quartz cuvette. The transient PL spectroscopy data were fitted to a monoexponential decay to determine the lifetimes.

**Table S.7: Comparison of the lifetimes of MIL-125-NH$_2$, 8 wt.% MIL-167/MIL-125-NH$_2$ and MIL-167, recorded at different wavelengths based on the PL emission peaks of the samples.**

<table>
<thead>
<tr>
<th></th>
<th>460 nm (μs)</th>
<th>490 nm (μs)</th>
<th>550 nm (μs)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MIL-125-NH$_2$</td>
<td>5.364 ± 0.009</td>
<td>4.650 ± 0.013</td>
<td>3.113 ± 0.017</td>
</tr>
<tr>
<td>8 wt.% MIL-167/MIL-125-NH$_2$</td>
<td>5.273 ± 0.011</td>
<td>3.676 ± 0.028</td>
<td>3.259 ± 0.017</td>
</tr>
<tr>
<td>MIL-167</td>
<td>-</td>
<td>0.721 ± 0.018</td>
<td>0.532 ± 0.007</td>
</tr>
</tbody>
</table>
Appendix Design of MOF/MOF Heterojunctions

Stavroula A. Kampouri

S4.5 UIO-66-NH$_2$/MIL-125-NH$_2$ Heterojunctions

S4.5.1 Synthetic Protocol

Synthesis of UIO-66-NH$_2$

UIO-66-NH$_2$ was synthesized by slightly modifying a reported procedure, as follows:

First, two stock solutions were synthesized. The first one by dissolving 0.240 g of the linker NH$_2$-H$_2$BDC in 20 mL DMF and the second one by dissolving 0.320 g of zirconium (IV) chloride in 20 mL DMF. After sonicating the mixtures for sufficient time (in order for the linker or metal salt to be dissolved in the solvent), 2 mL of each solution was added in 10 mL vials and were heated up to 120 °C for 48 h. The obtained light-yellow solids were first washed several times with DMF, then stirred overnight in MeOH and the next day they were washed several times with MeOH. Finally, the obtained powder was dried at 60 °C for ~5 h.

Synthesis of UIO-66-NH$_2$/MIL-125-NH$_2$

The UIO-66-NH$_2$/MIL-125-NH$_2$ samples were synthesized following similar procedure to that of the MIL-167/MIL-125-NH$_2$ except that 10, 20, 40 or 60 mg of UIO-66-NH$_2$ were introduced in the MIL-125-NH$_2$-precursor solution (instead of MIL-167), prior to heating.

S4.5.2 Powder X-ray Diffraction

Figure S.28: PXRD patterns of a) the as-synthesized (light colors) and simulated (dark colors) MIL-125-NH$_2$ (yellow) and UIO-66-NH$_2$ (gray), and b) MIL-125-NH$_2$ synthesized in the presence of 10, 20, 40 and 60 mg of UIO-66-NH$_2$. 

– 98 –
Table S. 8: X-ray diffraction-derived data (measurements in 0.4 mm borosilicate glass capillaries).

<table>
<thead>
<tr>
<th>Synthesis of MIL-125-NH₂ in:</th>
<th>MIL-125-NH₂ composition (wt.%)</th>
<th>Uio-66-NH₂ composition (wt.%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>10 mg UIO-66-NH₂</td>
<td>84</td>
<td>16</td>
</tr>
<tr>
<td>20 mg UIO-66-NH₂</td>
<td>72</td>
<td>28</td>
</tr>
<tr>
<td>40 mg UIO-66-NH₂</td>
<td>54</td>
<td>46</td>
</tr>
<tr>
<td>60 mg UIO-66-NH₂</td>
<td>42</td>
<td>58</td>
</tr>
</tbody>
</table>

S4.5.3 Scanning Electron Microscopy

![SEM images](image_url)

Figure S.29: a) SEM image of UIO-66-NH₂ and b) 58 wt.% UIO-66-NH₂/MIL-125-NH₂.

S4.5.4 Stability

![PXRD patterns](image_url)

Figure S.30: PXRD patterns of UIO-66-NH₂, 16 wt.%, 28 wt.% and 46 wt.% UIO-66-NH₂/MIL-125-NH₂ after photocatalysis.
Supporting Information for Chapter 5: MOF-derived TiO₂ for Photocatalysis

S5.1 Synthetic Procedures

Synthesis of MIL-125-NH₂-derived TiO₂

In a typical preparation, 200 mg of the as-synthesized MIL-125-NH₂ powder was placed in a ceramic crucible covered with foil and calcined for 1 h at either 400 °C, 500 °C, 600 °C, 700 °C or 800 °C. The obtained product was washed several times with EtOH and water, and then dried at 75 °C for 12 h.

Synthesis of precipitated TiH₄O₄-derived TiO₂

The TiH₄O₄-derived TiO₂ was synthesized following a similar procedure to a reported one.⁶¹ 5 mL of titanium isopropoxide were mixed with 15 mL isopropanol. The solution was slowly added in 250 mL millipore water, under vigorous stirring. The precipitated titanium hydroxide (TiH₄O₄) solid was filtered with 400 mL of distilled water and then dried for 3 h at 120 °C. The obtained powder was put in a ceramic crucible covered with foil and calcined for 1 h at either 400 °C, 500 °C, 600 °C, 700 °C or 800 °C. The resulting white powders were washed several times with EtOH and water, and then dried at 75 °C for 12 h.

Synthesis of MIL-167-derived TiO₂

MIL-167-derived TiO₂ was synthesized following similar procedure to that of MIL-125-NH₂-derived TiO₂, except that 200 mg of the MIL-167 was used instead of MIL-125-NH₂ and the sample was heated up to 600 °C for 1 h.

S5.2 Characterization

S5.2.1 X-ray Photoelectron Spectroscopy

X-ray photoelectron spectroscopy (XPS) was performed, to analyze the surface of the mixed phase material. XPS revealed a titanium 2p peak at a binding energy of 458.5 eV. Each TiO₂ rutile and anatase phase has a distinct binding energy separated by 0.4 eV, allowing us to quantify the composition of both phases on the surface of TiO₂.⁶³ From the fitting, the TiO₂ anatase and rutile phase composition of the MIL-125-NH₂-derived TiO₂ at 600 °C is 15:85 (Figure S31a) indicating a higher rutile content on the surface compared to the bulk (66:34). This higher surface rutile content is also the case with the TiH₄O₄-derived TiO₂ (Figure S31b). This analysis is consistent with the thermal
heating process involved, whereby the surface of the particle is where the initial transformation between anatase and rutile takes place.

Figure S.31: XPS showing the Ti 2p binding energy of a) the best performing MIL-125-NH$_2$-derived TiO$_2$ sample (synthesized at 600 °C) and b) the best performing TiH$_4$O$_4$ sample (synthesized at 700 °C). Included is the fitting for the TiO$_2$ anatase and rutile phase, determined using 2 Voigt functions; insert is the relative area of the fits related to each line.
S5.2.2 Transmission and Scanning Electron Microscopy

Figure S.32: a) Bright-field and b) dark-field TEM images of MIL-125-NH$_2$-derived TiO$_2$ synthesized through calcination at 400°C for 1 h. c) Selected Area Electron Diffraction (SAED) pattern. Dark-field TEM image was acquired from the reflections shown by a circle on the SAED pattern.

Figure S.33: a) Bright-field and b), c), d) dark-field TEM images of MIL-125-NH$_2$-derived TiO$_2$ synthesized through calcination at 600°C for 1 h. e) SAED pattern. For the acquisition of the dark-field images, the incident electrons beam is tilted so that only reflections of specific diffracting crystallographic planes contribute to image formation. Depending on the angle of diffraction, specific nano-sized particles are illuminated, manifesting the multi-particle nature of the template. Dark-field TEM images correspond to the diffracting planes shown by a green circle in the SAED pattern.

Figure S.34: SEM image of the TiH$_4$O$_4$-derived TiO$_2$ samples synthesized at a) 500 °C, b) 600 °C and c) 700 °C for 1 h. The images are of the same magnification.
Figure S.35: a) Bright-field TEM image and b) SAED pattern of TiH₄O₄-derived TiO₂ synthesized through calcination at 500°C for 1 h.

S5.2.1 PXRD after Recycling Experiments

Figure S.36: PXRD pattern of MIL-125-NH₂-derived TiO₂ (with optimum TiO₂ anatase to rutile ratio of 66:34) before and after recycling experiments.
S5.2.3 TGA of MIL-167-derived TiO₂

![TGA curve of MIL-167](image)

**Figure S.37:** TGA curve of MIL-167 obtained under airflow with a heating rate of 5 °C min⁻¹.

**Table S. 9:** X-ray diffraction-derived data.

<table>
<thead>
<tr>
<th></th>
<th>Anatase (w.t. %)</th>
<th>Rutile (w.t. %)</th>
<th>Size Anatase (nm)</th>
<th>Size Rutile (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MIL-167-derived TiO₂</td>
<td>74.5(6)</td>
<td>25.5(8)</td>
<td>22</td>
<td>33</td>
</tr>
<tr>
<td>P25 Degussa TiO₂</td>
<td>88.3(3)</td>
<td>11.7(8)</td>
<td>20</td>
<td>34</td>
</tr>
</tbody>
</table>
S.5.3 Comparison of MIL-125-NH$_2$-derived TiO$_2$ to other TiO$_2$ samples

S5.3.1 Dependence of the Photocatalytic Activity on the Concentration of Photocatalyst

![Graph showing the dependence of H$_2$ production on the concentration of photocatalyst in solution.](image)

**Figure S.38:** Dependence of the H$_2$ evolution on the concentration of photocatalyst in solution, for the best-performing MIL-125-NH$_2$-derived and P25 Degussa TiO$_2$.

S5.3.2 Photoluminescence Spectroscopy

![Graph showing photoluminescence spectra.](image)

**Figure S.39:** PL spectra of the MIL-167-derived, TiH$_4$O$_4$-derived, MIL-125-NH$_2$-derived TiO$_2$ and P25 Degussa. The excitation wavelength was at 340 nm.
### S5.3.3 Elemental Analysis

**Table S.10:** Elemental analysis for the best performing MIL-125-NH$_2$-derived TiO$_2$ sample (synthesized at 600 °C). A small amount of carbon, traces of hydrogen and no nitrogen were observed.

<table>
<thead>
<tr>
<th>Nitrogen (%)</th>
<th>Carbon (%)</th>
<th>Hydrogen (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MIL-125-NH$_2$-derived TiO$_2$</td>
<td>0</td>
<td>1.154</td>
</tr>
</tbody>
</table>
References


References


References

- e) Fu, Y.; Yang, H.;
under visible light irradiation. Nanoparticles on UiO-112 surface can produce hydrogen with the degradation of organic pollutants using TiO2 photocatalyst modified with dual Rh/SrTiO3.

Simultaneous H2 evolution and mineralization of organic compounds enabled by a C111 framework.

M. A.; Durrant, J. R.; Cooper, A. I.; Tang, J., Current understanding and challenges of solar hydrogen generation.


MOF-derived nanostructured cobalt phosphide assemblies for efficient hydrogen evolution reaction. RSC Advances 2015, 5, 90265-90271.


Synthesis of Pt@NH2-MIL-125(Ti) as a photocathode material for photoelectrochemical hydrogen production. RSC Adv. 2013, 3, 19820-19823.


References


References


Curriculum Vitae

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Education
Strategic design and characterization of Metal-Organic Framework-based systems with enhanced light absorption properties and charge separation efficiencies. Their application involves the photocatalytic hydrogen evolution reaction and dual-functional photocatalysis for Simultaneous organic pollutant degradation and hydrogen evolution.
Laboratory of Molecular Simulation (LSMO),
Supervisors: Prof. B. Smit and Dr. K. Stylianou.

Sep. 2010 - 2015 Diploma in Environmental Engineering at Technical University of Crete (TUC), Greece.
Thesis: ‘Catalytic Decomposition of Nitrous Oxide (N₂O) on Ir-based and Metal Phosphide Catalysts’ (7 months).
Synthesis and characterization of metal phosphide and Iridium-based catalysts supported on mixed metal-oxides, and their investigation for the catalytic decomposition of nitrous oxide.
Laboratory of Physical and Chemical Processes,
Supervisor: Prof. I.V. Yentekakis.

Employment History
Sept. - Nov. 2015 Internship within the Laboratory of Physical and Chemical Processes, under the supervision of Profs. I. V. Yentekakis and P. Panagiotopoulou at the Technical University of Crete (TUC), in Greece. Project related to the synthesis of different mixed-metal oxides as supports for Iridium catalysts.

May - Jun. 2016 Internship within the Water and Air Quality Laboratory, under the supervision of Prof. A. Stasinakis at the University of the Aegean, in Greece. Project related to the investigation of duckweed Lemna minor as a wastewater treatment system.
Teaching Experience

Spring Semesters 2017 - 2019

Teaching assistant, Chemical engineering practical work for 1st and 2nd year bachelor students, *TP: Mass Transfer Phenomena*, EPFL.

Nov. - Dec. 2019

Supervision of visiting student Matthew Garvin, from the Heriot-Watt University (United Kingdom). *Project related to the design and characterization of metal-organic framework/semiconductor heterojunctions for photocatalysis*, EPFL.

Jul. - Aug. 2019

Supervision of visiting bachelor student Aylin Kinik, from the Istanbul Technical University (Turkey). *Project related to the synthesis of metal-organic framework-derived TiO$_2$ for photocatalysis*, EPFL.

Nov. 2018 - Jul. 2019

Supervision of visiting student Farrokh Mohammadnezhad, from the Razi University (Iran). *Project related to the investigation of the impact of linker doping on the optoelectronic properties and photocatalytic activity of Metal-Organic Frameworks*, EPFL.

Winter Semester 2018

Teaching Assistant, Advanced practical work of physics for 3rd year bachelor students, *TP: Internal photoemission - Schottky barrier*, EPFL.

Winter Semester 2018

Teaching Assistant, Advanced practical work of physics for 3rd year bachelor students, *TP: Optical Spectroscopy*, EPFL.

Prizes and Awards

- **Chemistry Travel Award**, Platform Chemistry of Swiss Academy of Sciences SCNAT and Swiss Chemical Society (SCS), 2019 (Sion, Switzerland).
- **Best Poster Prize** at 12th Int. Symposium Hydrogen & Energy 2018 (Lausanne, Switzerland).
- **Best Poster Prize** at EuroMOF conference 2017 (Delft, Netherlands).
- Award from **Mathematical Olympiad of Greece** E.M.E. Thales 2007 (Mytilene, Greece).

Personal Skills

Languages:

- **Greek** *ILR Level 5* Native or bilingual proficiency.
- **English** *ILR Level 5* Native or bilingual proficiency.
- **French** *ILR Level 2* Limited working proficiency (*Diplôme d’Etudes en langue Françaises, DELF B1*).

Job-related skills:

- **Software**: Excellent competence in Microsoft Office, ChemDraw, Diamond, CCDC software package (e.g., Mercury) Origin.

Outreach activity: **Scientastic Festival EPFL Valais** (2017-2019). Swiss annual festival to communicate science and technology to young people and the general public. Tasks included presenting my research projects in simple, straightforward terms and organizing the LSMO laboratory space for the event.

**Contributions to International Conferences**

1. MRS Fall 2019, Material Research Society Fall Meeting and Exhibit (Boston, U.S.A.).

2. AAAFM 2019, American Association for Advances in Functional Materials (Los Angeles, U.S.A.).


4. NAGC 2018, 8th North America-Greece-Cyprus Workshop on Paramagnetic Materials (Sparta, Greece).

5. Latsis symposium: 12th Int. Symposium Hydrogen & Energy 2018 (Lausanne, Switzerland)


**Published/Submitted Work**


Publications Under Preparation