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Variation of Pore Metrics in Metal-Organic Frameworks for Enhanced Storage and Catalytic Applications

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Variation of Pore Metrics in Metal-Organic Frameworks for Enhanced Storage and Catalytic Applications

A dissertation submitted in partial satisfaction
of the requirements for the degree
Doctor of Philosophy in Chemistry

by

Jonathan Ward Brown

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Variation of Pore Metrics in Metal-Organic Frameworks for Enhanced Storage and Catalytic Applications

by

Jonathan Ward Brown

Doctor of Philosophy in Chemistry
University of California, Los Angeles, 2015
Professor Louis-Serge Bouchard, Chair

With the principles of reticular chemistry, metal-organic frameworks (MOFs) with enhanced storage and catalytic capabilities have been prepared. This dissertation presents the synthesis of azo-IRMOF-74-III for controllable release of cargo molecules as well as the catalytic testing of MOF-525-Mn and an isoreticular series of MOFs based on the MOF-5 framework. The variation of pore metrics in these frameworks show the versatility of reticular chemistry and their application in catalytic reactions.

First, an azobenzene containing MOF-74 analogue was prepared by incorporating the azobenezne moiety into the organic linker used to prepare MOF-74. Cargo molecules were trapped inside the structure based on size constraints and controllably released via modulation of UV irradiation to induce a trans to cis isomerization on the organic linker. The variation in the linker length shows that specific cargo molecules can be trapped by changing the size of the one-dimensional pore in reticular frameworks.

In addition to changing pore size, density of catalytic sites inside a pore was investigated through the isoreticular synthesis of a series of MOFs, modeled after
MOF-5 but containing various amounts of 2-aminoterephthalic acid. These amine groups were post-synthetically modified to contain catalytically active Pd sites and heterogeneous Heck coupling was performed. A inverse relationship was observed between the amount of framework metalation and reaction yield and selectivity. This is attributed to pore blocking from post-synthetic modification at higher metal loadings.

Finally, a manganese porphyrin containing MOF, MOF-525-Mn, was synthesized and its activity tested through the epoxidation of alkenes using molecular oxygen. High catalytic activity was observed for all alkenes and this is the first instance of MOF epoxidation using molecular oxygen, a ubiquitous industrial oxidant. Another MOF containing the same catalytically active porphyrin but in a different topology, MOF-545-Mn, was also tested for catalytic activity to determine the effect of MOF topology on heterogeneous catalysis.

The variation of pore metrics is demonstrated through each chapter and show how powerful and useful reticular chemistry is through variation of structural elements for specific uses. While catalytic MOFs are in their infancy, the future is bright for catalytically active MOFs to come to the forefront of chemical industry.
The dissertation of Jonathan Ward Brown is approved.

Xiangfeng Duan

Paula L. Diaconescu

Dante Simonetti

Louis-Serge Bouchard, Committee Chair

University of California, Los Angeles

2015
For my parents, Nelson and Debbie Brown
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Vita

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Publications


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CHAPTER 1

Introduction

Metal-organic frameworks (MOFs) are made by joining metal-oxide clusters with organic linker molecules to produce extended, crystalline structures [1–4]. These frameworks are structurally defined due to the rigidity of organic linkers and defined geometries of the metal oxide units. MOFs exhibit several hallmark properties including high porosity as well as chemical and thermal stability [5]. Framework integrity remains intact even after all guests have been removed from the pores and the material has been heated to 450$^\circ$ C [6, 7]. Due to this high framework integrity, MOFs often have very high permanent porosity [8,9]. In addition to these hallmark properties, the ability to vary functional groups on the organic links combined with the choice of metal-oxide secondary building units (SBUs) gives MOFs a large degree of flexibility in terms of topology, pore size and geometry. Due to their large surface area and variable pore environment, MOFs have been receiving much attention for applications such as heterogeneous catalysis [10, 11], gas storage/separation and purification [12, 13], high-sensitivity sensors [14,15], porous electrodes [16,17] and magnets [18–21], size-, shape- and enantio-selective chromatography [22–24], and drug storage/delivery [25,26].

Reticular chemistry provides the blueprint for building MOFs. This means that we can predict the structural outcome of a chemical reaction. Reticular chemistry is defined as, “Logical construction of chemical structures that are held together by strong bonds” [1]. This enables us to predetermine the topology and structure and, utilizing rational design, synthesize MOFs with a desired topology,
periodicity and functionality for a desired structural outcome. One of the most ubiquitous MOFs, MOF-5 \([\text{Zn}_4\text{O}(\text{C}_8\text{H}_4\text{O}_4)_3]\) [27] is constructed by connecting \(\text{Zn}_4\text{O}(\text{CO}_2)_6\) with the linear molecule 1,4-benzenedicarboxylic acid as the linker bridging two metal oxide units to form a primitive cubic structure, Figure 1.1. Utilizing the principles of reticular chemistry, this MOF can be expanded or altered to include different functionalities [28]. We know that reactions of carboxylic acids and selected transition metals under specific conditions form octahedral \((\text{M}_4\text{O}(\text{CO}_2)_6)\), square planar \((\text{M}_2\text{(CO}_2)_4)\), or triangular \((\text{M}_3\text{(CO}_2)_6)\) SBUs [29] (Figure 1.1B), and these provide points of extension. The directionality (linear, trigonal, etc.) of the linker that will connect the points of extension helps determine the overall framework topology and allows us to predict the overall structure. Thus, utilizing a \(\text{Zn}_4\text{O}(\text{CO}_2)_6\) cluster that has six points of extension, and connecting those points with 1,4-benzenedicarboxylate links, a cubic topology is obtained where an octahedral \(\text{Zn}_4\text{O}(\text{CO}_2)_6\) cluster is connected by six 1,4-benzenedicarboxylate links that can be shared by six different \(\text{Zn}_4\text{O}(\text{CO}_2)_6\) clusters, thereby forming an extended simple cubic network.

Varying the link type or altering functional groups on the link can drastically change the pore metrics of the structure without disturbing the underlying topology. The multivariate (MTV) principle showed that different organic linkers of similar size and geometry can be used in the synthesis of one MOF, allowing for the fine tuning within MOF structures. Furthermore, the MTV principle has allowed for links incompatible with MOF synthesis to be incorporated in smaller quantities into a MOF framework [30]. Some links were unable to form crystalline structures via solvothermal synthesis either due to steric constraints or unfavorable electronic properties, yet the MTV principle circumvented these problems by mixing multiple types of linkers in a single pot and producing MOFs with several functionalities in the final framework [31]. This opened up new possibilities for frameworks in terms of both functionalities inherent to the MOF and its result-
Figure 1.1: A) Synthesis of prototypical MOF-5. B) Reaction of carboxylate linker and metal salt yields different SBUs and by extension different topologies and properties, such as introduction of hydrophobic or hydrophilic links, Lewis acid and base pairs, and alteration of pore geometry through steric hindrance [11, 32–35].

Another tool for MOF chemists, postsynthetic modification (PSM), allows for introduction of functionalities into a MOF following solvothermal synthesis [36]. PSM works by first forming a framework then performing chemical reactions on that structure to insert a desired functional group or catalytically active site [37–40]. This allows one to take an unreactive framework with specific properties or desired topology and modifies it to take on new and interesting properties such as improved gas sorption, catalytic activity, and more robust physical properties [41].

Both the MTV principle and PSM have been used to introduce new function-
alities or catalytically active sites into a framework, giving the MOF field two important tools to expand the use of MOFs in heterogeneous catalysis. These tools allow for the modification of pore environment and geometry. My research described herein has used both as a way of designing higher order materials that have desirable properties such as enhanced catalytic activity and selectivity or the ability to trap and controllably release molecules of interest.

The objective of this dissertation is to explore catalytic applications of MOFs by altering pore metrics such as geometry and environment. Some of the most common catalytic reactions that have been seen in MOFs are hydrogenation, oxidation, and epoxidations [42–45]. While MOF catalysis is relatively new, the field is rapidly expanding as more and more catalytically active structures are synthesized [46, 47]. One major advantage of using a MOF in heterogeneous catalysis is the ability to vary the organic link in the framework. This opens up many possibilities for lining the pore cavity with catalytic active species and also allows us to control the density of active sites. Both PSM and the MTV principle have been crucial in the development of MOFs as catalysts as many MOFs were viewed as both unreactive and unsuitable for catalysis, either due to instability of the framework or lack of catalytically active sites. While the field of MOF catalysis is growing, many papers report catalytic yields that are unoptimized. My research has sought to fill this gap by optimizing reaction yield and selectivity for a number of different heterogeneous catalyst. Another interesting development in MOF catalysis is the rise of chemically and thermally stable SBUs that are rapidly expanding the types of reactions available for MOFs [48–50]. The synthesis of MOFs utilizing a Zr (the UiO-66 series), Cr or Fe (MIL-101 series), or Al (MIL-53 series) backbone have helped in terms increase the number of frameworks which are chemically or thermally stable enough to withstand rigorous catalytic conditions [51–53].

One of the first goals in my research was to determine whether a molecule
Figure 1.2: Synthetic representation of idealized azo-IRMOF-74-III. The yellow balls represent largest free volume in idealized azo-IRMOF-74-III when azobenzene functionalities are entirely in cis conformation.

could be trapped within a MOF pore without being chemically bonded to the framework. In chapter 2 this idea is explored by the synthesis of a MOF-74 derivative that contains azobenzene units (Figure 1.2).

By choosing a framework with one-dimensional pore structure, we can use the azobenzene functionality, projecting toward the middle of the pore to capture and controllably release cargo molecules based on size exclusion. While the framework itself is not catalytically active, this MOF shows proof of principle that molecules of interest can be controllably captured based on size selection, introduced into the pores of a structure and then controllably released. This idea could be easily extended to trapping molecules based on properties other than simply size constraints and allows for greater control over pore geometry and environment in known MOFs.
Another major component of catalytically active MOFs is synthesis of the catalytic sites themselves. While many MOFs are capable of having active sites introduced into the framework, there have been relatively few examples where functional catalysts are embedded into known frameworks. Chapter 3 discusses the synthesis and optimization of a catalytically active MOF, which contains catalytically active Pd sites. Using both the MTV principle and PSM to optimize the density of catalytically active sites, we show that metalation and porosity of a framework (Figure 1.3) play an integral role in finding optimal catalytic conversion for Heck coupling to synthesize resveratrol trimethyl ether, a pharmaceutically relevant precursor.

Surprisingly, the highest density of catalytic sites in our framework does not show the highest catalytic conversion. In fact, during our Heck coupling experiments we found that the lowest amount of catalytic loading provided both the highest catalytic yield and selectivity. We have seen that by varying the percentages of available functionality within a structure and by extension the number...
and density of catalytic sites, we can test many different frameworks for catalytic activity and find optimal metal loading and activity for a given framework. This seemingly counterintuitive finding that fewer catalytic sites give higher catalytic conversion brings up interesting questions in regards to framework design as well as the density of catalytically active sites and provides valuable background into future research in rational design of new MOFs for catalytic applications.

In chapter 4, the idea of exploring new methods of catalysis is shown by way of a Zr-MOF containing a manganese metalloporphyrin for the epoxidation of alkenes (Figure 1.4).

There have been other published accounts of MOFs that catalyze epoxidations but none that use molecular oxygen as the oxygen source [24, 45, 54]. Molecular oxygen is regularly used in industry due to it being inexpensive, less corrosive and easier to handle on a large scale than liquid peroxides [55]. The epoxidation of alkenes is particularly attractive to industrial chemical companies as propylene
and ethylene oxide are produced in millions of tons per year and are precursors for widely used chemicals [56,57]. While this study examines the epoxidation of styrene instead of propylene or ethylene epoxidation, the use of molecular oxygen makes it an interesting beginning to the search for an industrially relevant MOF catalyst. In addition to seeing high yields and good selectivity for our heterogeneous system, we also elucidated the reaction rates and activation energies associated with this particular reaction, being the first group to show this information in reference to MOF catalysis.

Finally in chapter 5, future direction for this work is explored and discussed. The framework topology in catalysis and reaction optimization plays a major role in MOF catalysis yet there have been very few studies [58] exploring this aspect of reaction engineering. In addition, epoxidation of long chain alkenes is of major interest to chemical industry and exploration of more open structures could lend itself to high yield and selectivity for industrially relevant molecules.

Catalytically active MOFs are still in their infancy, especially with respect to optimization of reactor and catalytic efficiency. Increasing control over pore metrics holds the key to synthesis of new and more efficient catalytic frameworks, expanding both the types of reactions available for MOF catalysis and the catalytic yield of existing structures. The MTV principle and PSM are indispensible tools in the design of new, higher order MOFs and subsequent control over pore metrics. As is demonstrated here, there are many factors that can affect the catalytic ability of MOFs such as framework design and selection of topology, chemical and thermal stability of MOFs and the density of active sites within the framework. Though MOF catalysis is a nascent field, its future is bright and the discovery of new frameworks with interesting catalytic applications is simply a matter of time.
CHAPTER 2

Photophysical pore control and trapping of cargo in azo-IRMOF-74-III

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MOFs are extended crystalline structures containing metal oxide units bridged by organic linkers. They are highly porous and typically exhibit large surface areas. A wide variety of organic linkers have been incorporated into MOFs, highlighting the isoreticular principle whereby the organic linkers can be changed, yet analogous structural types can be synthesized. [48, 59–61] Variation of the functional groups present in MOFs have led to applications in chemical sensing, [62,63] gas separation, [64] gas storage, [65,66] catalysis, [23] and drug delivery. [67] Despite the large library of MOFs reported in the literature, to date the storage and release of guest molecules has relied [68] solely on the uncontrolled diffusion of cargo into and out of the extended structures of the MOFs. Although many MOFs have been synthesized, none have demonstrated on-command release of stored guest molecules using a controllable external stimulus.

One such functionality, which demonstrated promise for the controlled release of cargo from the pores of a MOF, is the azobenzene unit. Azobenzene provides a source of reversible and photo-controllable cis/trans isomerizations [69–71]. This conformational switching results in significant changes in the length of the molecule (9 Å in the trans form to 5.5 Å in the cis form) [72] and, as a result,
this photoswitchable system has been incorporated into a variety of rigid materials including films and nano-structures. [73–86] Furthermore, the integration of azobenzene units has been employed recently in the one-dimensional (1-D) pores of mesoporous silica nanoparticles where it was demonstrated [87, 88] that the on-command release of cargo molecules from the pores is possible. Photoactive linkers have been incorporated into MOFs and shown interesting adsorption and storage properties that have been directly linked to the photoactivity of those frameworks [89–93]. Recently two MOFs were synthesized [94, 95] containing azobenzene units, however, the nature of the pores in these MOFs produced steric restrictions that limited photoswitching applications. We chose to incorporate the photoswitch into a MOF-74 based structure, to combat these steric limitations. Due to a unique binding motif inherent to MOF-74, the inorganic clusters and organic struts form linear stacks producing large, non-interpenetrated 1-D hexagonal pores. This unique architecture will force the azobenzene units directly into the channels where the isomerization of each unit will have a direct impact on the aperture size of a single 1-D channel. Therefore, the photodynamic switching of azobenzene combined with the 1-D structure of this MOF has the potential to allow for controlled cargo delivery from a MOF.

Here, we report the synthesis and photoisomerization studies of an azobenzene-derivatized MOF, azo-IRMOF-74-III, which contains 1-D pores. In this particular MOF, the size and shape of the apertures are controlled by the conformational changes in the azobenzenes, which can be reversibly switched from trans to cis or cis to trans using UV or visible irradiation, respectively. When all of the azobenzene units are in the trans conformation the pore apertures are 8.3 Å in diameter, but upon switching to the cis conformation the size of the aperture is increased significantly (10.3 Å).

This photodynamic MOF exhibits improved photoswitching over previously synthesized azobenzene-derivatized structures because the azobenzenes are evenly
Figure 2.1: Synthesis of Azobenzene-Functionalized MOF Struts

... spaced within the crystalline framework. The photophysical properties of the MOF have been investigated by $^1$H NMR and UV-Vis spectroscopy, and information obtained from these techniques has been used to choose a wavelength of light capable of isomerizing the azobenzene units back and forth between their two possible conformations. Spectroscopic monitoring of the release of a fluorescent guest molecule reveals that the dye remains contained in the MOF until irradiation at an appropriate wavelength. This successful containment and on-command release demonstrates the utility of self-contained photo-active switches in highly porous MOFs.
2.1 Experimental Detail

The synthesis of the azobenzene-functionalized linkers involves (Figure 2.1) a palladium-catalyzed Suzuki-Miyaura cross-coupling reaction between 2,5-dibromo-nitrobenzene and the phenylboronic ester 1 in a degassed 1,4-dioxane/H$_2$O mixture employing PdCl$_2$(dppf) as the catalyst and CsF as the base to yield the nitro derivative 2. Treatment of 2 with Raney Ni and H$_2$ gas in EtOAc at 50°C, serves to facilitate both the removal of the benzyl protecting groups and reduce the nitro group to produce the aniline derivative 3 in nearly quantitative yield. Subjecting 3 to excess of nitrosobenzene in acetic acid at 80°C yielded the azobenzene 4, which could be converted to the dicarboxylic acid 5, by saponification of the methyl esters.

Azo-IRMOF-74-III was synthesized by combining Mg(NO$_3$)$_2$·6H$_2$O (80 mg, 0.315 mmol) and the azobenzene-functionalized linker (42.5 mg, 0.09 mmol) in a solution of N,N-dimethylformamide (DMF) (7.5 mL), EtOH (0.5 mL) and H$_2$O (0.5 mL) for 24 h at 120°C. After 24 h, the red crystals were collected by filtration.

Powder X-ray diffraction was used for structural determination as the microcrystalline material produced from the MOF synthesis did not diffract sufficiently for single crystal elucidation. Since a higher level of characterization is achieved by producing a solvent and guest free framework, the crystals were sequentially washed with DMF (3 times, 10 mL) and methanol (3 times, 10 mL) and then evacuated on a supercritical CO$_2$ dryer over the course of 12 hours. Following this activation process, thermal gravimetric analysis (TGA) was used to confirm that all of the solvent had been removed from the pores of the framework; i.e., there was no significant weight loss up to 275°C (Figure 3.12).

Azo-IRMOF-74-III is an isoreticular expansion of MOF-74, in which helical MgOC rods are constructed from hexacoordinate Mg(II) centers. The rods are connected via azobenzene functionalized terphenylene unit 5 to produce one-
Figure 2.2: Viewing idealized azo-IRMOF-74-III down the c-axis displays one-dimensional pores and azobenzene functional groups projecting into the pores. The yellow and orange balls represent pore aperture in idealized azo-IRMOF-74-III, when the azobenzene functional groups are in trans and cis conformation, respectively.

One-dimensional pores, which have a diameter of 19.5 Å. This structure was simulated using the Forcite module of Materials Studio and was confirmed with the Rietveld method. Because the one-dimensional pores in azo-IRMOF-74-III are lined with azobenzene groups that project toward the middle of the pore, the size and shape of the pore aperture directly depend on the cis or trans configuration. With all azobenzene functionalities in the trans configuration the idealized pore aperture is 8.3 Å but if all functionalities are cis, the aperture is larger 10.3 Å (Figure 2.2).

The PXRD pattern for azo-IRMOF-74-III was indexed on a trigonal unit cell and refined using the Rietveld method in the Reflex module of Materials Studio, yielding cell parameters $a = 46.771(26)$ Å, $b = 46.771(26)$ Å and $c = 6.869(83)$ Å (residuals: $R_p = 12.08 \%$, $R_{wp} = 17.68 \%$) (Figure 2.3).

Azo-IRMOF-74-III was further characterized by cross-polarization magic-angle
Figure 2.3: Indexed experimental (red) and refined (black) PXRD patterns of azo-IRMOF-74-III after Rietveld refinement. The difference plot is indicated in green. Blue ticks indicate the positions of Bragg reflections.

spinning (CP/MAS) $^{13}$C NMR spectra (Figure 2.13). From experiments on the activated material, the expected resonances at 174 ppm for the carbonyl carbon atoms and 166 and 114 ppm for the aromatic carbon atoms of incorporated 5 were found. No free 5 was observed in the MOF spectra, confirmed by a lack of peaks at 176, 160, and 108 ppm.

To confirm the porosity of azo-IRMOF-74-III, $N_2$ isotherms at 77 K were measured on activated samples. From the Brunauer-Emmet-Teller (BET) analysis a surface area of 2410 m$^2$/g was calculated (Figure 2.4). To date, this is the highest recorded surface area of a MOF containing photoswitchable units.
Figure 2.4: Nitrogen isotherm of azo-IRMOF-74-III carried out on activated sample at 77 K. Adsorption and desorption branches are represented by solid and open circles respectively. A surface area of 2410 m$^2$/g was calculated from Brunauer-Emmett-Teller (BET) analysis.

2.2 Results and Discussion

The azobenzene functionalized linker used to synthesize azo-IRMOF-74-III exhibits the expected isomerization when exposed to UV and visible light. While previous studies of azobenzene and its derivatives have examined the photophysical properties in solution and various molecular environments, incorporation to a new platform warrants investigation of its photochromic properties. The UV-vis absorption experiments discussed in this section will thus demonstrate the performance of the azobenzene moiety within the rigid confines of a MOF.

The UV-vis absorption spectrum of non-irradiated 5 exhibits two prominent
Figure 2.5: Pre- (black trace) and post- (cyan trace) irradiation UV-vis absorption spectra of 5 (a) and cis (red trace) and trans (blue trace) spectra calculated from Equation 7.1 using relative cis and trans peak integrations from $^1$H NMR spectra (b). Spectra for (a) (20 M in DMSO) were irradiated with 15 mW of light from a 377 nm diode laser for 30 minutes.

peaks, close in energy, in the near UV (Figure 2.5 A). Irradiation of 5 with UV light (which typically induces a trans-to-cis isomerization for azobenzene) results in dramatic changes in the absorption spectrum; i.e. the emergence of a new absorption band in the visible region. Although the photoisomerization process is evident, transitions due to the terphenyl moiety overshadow those of the azoben-
zene functional group in both the pre- and post-irradiation absorption spectra. In addition, the fact that the absorption spectra of the photostationary states are not necessarily purely those of the trans or cis isomers, respectively, further complicates analysis. Correlation of absorption spectra with $^1$H-NMR data elucidated spectral assignments and will be discussed in more detail below.

Relative concentration ratios of the isomers were determined with $^1$H NMR and combined with UV-vis spectroscopy to calculate pure cis and trans spectra. Examination of the NMR data for a pre-irradiated solution of azobenzene in DMSO (5 mM for NMR experiments, but diluted to 20 M for absorption spectra) stored in the dark at room temperature revealed the absence of the cis isomer. In contrast, the NMR spectrum of the photostationary state achieved after 30 minutes of irradiation with near-UV light (377 nm, 15 mW) contained peaks from both the cis and trans isomers. Integration of the cis and trans peaks in the irradiated solution yielded a trans/cis ratio of 0.43 (70% of trans-azobenzenes had converted to the cis configuration). Pre- and post-irradiation absorption data were correlated with the relative isomer concentrations to calculate a pure cis absorption spectrum according to the following equation:

$$
\epsilon_c = \frac{1}{r_2 - 1/r_1} \left[ A_2 \left( \frac{1}{r_2} + 1 \right) - A_1 \left( \frac{1}{r_1} + 1 \right) \right]
$$

where $\epsilon_c$ is the calculated cis extinction coefficient (M$^{-1}$cm$^{-1}$), $c$ is the concentration of 5 (M), $l$ is the path-length of the cuvette (cm), $A_1$ and $A_2$ are the pre- and post-irradiation absorbances, respectively, and $r_1$ and $r_2$ are the pre- and post-irradiation trans/cis isomer ratios, respectively. The resulting pure cis and trans spectra (Figure 2.5 B) reveal a clearer representation of the transitions of 5. While each of the calculated cis and trans spectra contain appreciable absorption in the UV region due to the inherent absorption from the ter-phenylene structure, the trans isomer absorbs more than the cis isomer in the UV and the cis isomer absorbs more than the trans in the visible region as expected.
Figure 2.6: Comparison of absorption spectra of 5 (solid trace) with the absorption spectrum of azo-IRMOF-74-III (dotted trace). The transition seen at 315 nm shifts to 337 nm upon coordination of the linker with the magnesium atoms is attributed to a $\pi-\pi^*$ transition involving the C=O functionalities on the outer rings.

The UV and visible absorption peaks of the azo-benzene moiety are assigned to $\pi-\pi^*$ and $n-\pi^*$ electronic transitions, respectively. Both cis and trans calculated spectra contain a peak near 300 nm with a shoulder at 315 nm, which are attributed to the common ter-phenylene moiety. The higher energy transition near 300 nm is assigned to a $\pi-\pi^*$ transition that does not shift when the linker is coordinated to the metal atoms in the MOF. However, the shoulder at 315 nm, which is assigned to a $\pi-\pi^*$ transition involving the C=O functionalities on the outer rings, shifts to 337 nm upon coordination within the MOF structure (Figure 2.6). In the calculated spectrum of the trans isomer, a shoulder near 365 nm is attributed to an azobenzene $\pi-\pi^*$ transition. A weaker $n-\pi^*$ azobenzene transition is found at 438 nm in the calculated cis spectrum.

To characterize the photoproperties of linker 5 used in this MOF, absorption spectra were obtained as a function of irradiation time with UV and blue light.
Figure 2.7: Change in absorbance of the azobenzene linker with (a) 377 nm irradiation followed by (b) 457 nm irradiation. Each trace represents 15 seconds of irradiation. The first trace is plotted in black for clarity.

A 50 M solution of 5 was exposed to 15 mW of 377 nm excitation with continuous stirring; its absorption spectra upon irradiation are plotted in Figure 2.7. Irradiation of 5 in the near UV (377 nm, Figure 2.7a) leads to excitation of the azobenzene $\pi-\pi^*$ transition seen in the calculated trans spectrum.

Since the cis isomer has no appreciable absorbance in the near-UV, the probability of trans to cis isomerization is higher than cis to trans isomerization and thus the population of the cis species increases. Additionally, competing absorp-
tion from the terphenylene portion of the linker is relatively low at 377 nm, so the incident photons are able to reach the azobenzene more efficiently. To convert the molecule back to the trans species, the azobenzene linkers were irradiated in the visible region (457 nm, Figure 2.7b) where the absorbance of the cis isomer is higher than the absorbance of the trans isomer. The absorption spectra show an increase in the n-π* band at 438 nm and a decrease in the 300 nm peaks upon near-UV irradiation and the opposite trend with visible irradiation, indicating reversible isomer population changes without decomposition of the linker.

The cis and trans isomers of the linker have an isosbestic point at 402 nm that does not shift when coordinated to magnesium (Figure 2.16). Excitation wavelengths in this region can be utilized for stimulating repetitive large amplitude motions. These large amplitude motions have been used to activate the release of cargo from porous substances, [75,96] and cis-trans isomerization expels cargo in a similar way to the MOFs in this study.

To test the ability of azo-IRMOF-74-III to store and release cargo, a luminescent dye (propidium iodide, Sigma) was loaded into the hexagonal channels so that the release due to photoisomerization could be observed spectroscopically. This dye was specifically chosen because its size, 8Å×11Å×16Å without counterion, matches well with the pore aperture of azo-IRMOF-74-III, between 8.3 and 10.3Å depending on conformation of azobenzene linker (Figures 2.9 and 2.10). When comparing pore diameter to the dye, it can be seen that the dye cannot easily diffuse into the MOF unless the azobenzene functionalities are in the cis conformation (Figure 2.11). To load the azo-MOFs, 100 mg of particles were stirred in a 1 mM ethanolic solution of propidium iodide for 3 days under exposure to room lights. The loading process is driven by a concentration gradient and the photoisomerization afforded by room lights. UV-vis spectra of a propidium iodide solution acquired before and after loading demonstrate a loading capacity of 0.4 wt% (Figure 2.15). While minimizing light exposure the particles were washed
Figure 2.8: (a) Typical release of propidium dye from azo-IRMOF-74-III as a function of time, (b) detailed view of the baseline and start of laser irradiation (marked by the sharp line), and (c) an on/off trial, where the excitation beam was temporarily removed mid-release. The dye continues to diffuse away from the MOF, but the rate of release is drastically reduced.

thoroughly with ethanol to remove surface adsorbed dye. Subsequently the sample was placed in the corner of a 2 cm × 1 cm fluorimetric cuvette. A stir bar was placed in the opposite corner to facilitate mixing of the solution, and the cuvette was filled with spectroscopic-grade ethanol. The solution in the cuvette was stirred carefully to avoid disturbing the azo-IRMOF-74-III particles yet facilitate the diffusion of the dye. A probe beam (30 mW, 568 nm) was used to excite the propidium dye in the ethanol solution above the particles, and its fluorescence intensity was integrated and plotted as a function of time (Figure 2.8a).

As seen in Figure 2.8b, there is no increase in propidium iodide intensity prior to laser activation of the cis/trans isomerization motions in the azo-IRMOF-74-
III particles. This indicates that the azobenzene-functionalized MOF acts as an excellent container, and that no dye escapes from the MOF channels until an external light source of the proper wavelength is added.

After a baseline was established with the probe beam, a 50 mW pump beam at 408 nm (wavelength near the isosbestic point) activated the cis-trans wagging motions and expelled the dye from the MOF channels. A clear release was observed which began to level off after 40 hours (see Figure 2.8a). This release, which took much longer than in azobenzene-derivatized MCM-41 nanoparticles, can possibly be attributed to the fact that the MOF structure’s pores are longer and narrower [97]. Additionally, the physical interactions between the guest molecules and the host metal-organic framework may be vastly different than those seen in MCM-41. To demonstrate the necessity of the pump beam near the isosbestic point of the azobenzene moiety, a 647 nm laser was employed as a pump beam in a subsequent experiment (Figure 2.14). No release was observed until the 647 nm laser was replaced with one near the isosbestic point. The release seen in Figure 2.8a can also be controlled by increasing or decreasing the incident power or removing the beam entirely; blockage of the 408 nm pump beam caused the release to slow down (see Figure 2.8c), but upon reintroduction of the beam the original release rate resumed. The sharp increases and decreases seen in the fluorescence spectra of Figure 2.8 when the laser is turned on and off, respectively, are due to effects from laser scattering. When the laser impinges upon the MOF particles, the uneven surfaces of these particles scatter the incoming laser light and increase the overall intensity counts in the spectra. When the laser is turned off, the source of scattering is removed, and an equivalent drop is seen in the spectra.

The slope of the release does not immediately go to zero, most likely due to

\(^1\)point where the total absorbance of a sample does not change during a chemical reaction or a physical change of the sample
the following factors: (1) pores have larger apertures in the cis orientation (see Figure 2.2), allowing for some residual diffusion of cargo molecules, (2) a delay caused by diffusion of cargo from the MOF particles to the laser detection area is expected, and (3) thermal back-conversion processes continue to generate a small amount of motion in the dark. Most of these effects are short-lived, however, and discontinuation of laser irradiation will eventually reduce the rate of dye release to a negligible level. Since continuous laser irradiation is needed for efficient expulsion of the dye into solution, the pattern and duration of release can be adjusted as necessary. Azo-IRMOF-74-III, therefore, is useful in applications where longer release or stop-start release is desired [98–100].

2.3 Conclusions

An azobenzene-derivatized MOF has been synthesized and studied spectroscopically. Azobenzene trans and cis isomerizations of the linker molecule have been observed via NMR and absorption spectroscopy to ascertain the relative concentrations and the absorption spectra of the cis and trans species as well as the most appropriate wavelength choices for irradiation.

Irradiation at 408 nm, near the linker’s isosbestic point, causes both cis-trans and trans-cis excitations. The large amplitude motions due to these wagging motions led to an increase in dye mobility, expelling cargo from the structure on demand. The storage and release capabilities of the system could be studied with the aid of fluorescent propidium iodide, and the dye release rate was diminished when irradiation was removed. No release was observed prior to irradiation, indicating that the dye molecules are well contained.

The photo-driven azo-MOF structure, the focal point of this paper, is an improvement over recently made photoswitchable frameworks in that this structure contains only one-dimensional pores. The one-dimensional pores permit more effi-
cient storage and release of the dye, and the microporous particles allow filtration in a new size domain. Here, we have shown that the azo-MOF structure is capable of storing dye molecules and releasing them on command via a photoisomerization process. This new MOF creates unprecedented possibilities for the use of light in the realm of robust dynamics.

2.4 Supporting Information

2.4.1 Materials and general procedures

All reagents unless otherwise stated were obtained from commercial sources (Cambridge isotope laboratories, Sigma Aldrich, TCI) and were used without further purification. Yields reported were unoptimized.

2.4.2 Synthesis of azo-IRMOF-74-III

IRMOF-74-III azo \([\text{Mg}_2(\text{C}_{26}\text{H}_{16}\text{O}_6\text{N}_2])\] was made with \(\text{Mg(NO}_3)_2\cdot6\text{H}_2\text{O}\) (80 mg, 0.315 mmol) and the azobenzene-functionalized linker (42.5 mg, 0.09 mmol) in a solution of N,N-dimethylformamide (DMF) (7.5 mL), EtOH (0.5 mL) and H\(_2\)O (0.5 mL) for 24 h at 120\(^\circ\)C. After 24 h, the red crystals were collected by filtration. The crystals were washed with DMF (3×10 mL) over a thirty-minute period. The DMF was then replaced with MeOH (6×30 mL) over a two-day period. Finally, the volatile MeOH was removed by using a supercritical CO\(_2\) dryer. The crystals were then heated at 250\(^\circ\)C under vacuum (30 mtorr) for 12 hrs.

2.4.3 Structure resolution of azo-IRMOF-74-III from Powder X-Ray Diffractometry (PXRD)

Powder X-ray data were collected using a Bruker D8-Discover \(\theta\)-2\(\theta\) diffractometer in reflectance Bragg-Brentano geometry employing Ni filtered Cu K\(\alpha\) line focused
radiation at 1600 W (40 kV, 40 mA) power and equipped with a Vantec Line detector. Radiation was focused using parallel focusing Gobel mirrors. The system was also outfitted with an anti-scattering shield that prevents incident diffuse radiation from hitting the detector, preventing the normally large background at $2\theta < 3$. Samples were mounted on zero background sample holders by dropping powders from a wide-blade spatula and then leveling the sample with a razor blade. To determine the structure of IRMOF-74-III azo, crystal structure modeling was carried out, followed by Rietveld refinement.

2.4.4 Crystal structure modeling

The model for azo-IRMOF-74-III, including cell parameters and atomic positions, was generated using Materials Studio chemical structure-modeling software (Materials studio version 5.0, 2009, Accelrys Software Inc.) employing the Crystal Building module. The structure was generated using the relevant space group, the cell parameters were used as obtained, and atom positions were generated from the refined data of extended MOF-74. The constructed structure was minimized with the Forcite module using the Geometry Optimization routine with the Universal force field (UFF). The obtained model was used as input for the Rietveld refinement. Positions of atoms in the respective unit cells are listed as fractional coordinates in Table 2.1.
Table 2.1: Fractional atomic coordinates for azo-IRMOF-74-III.

Space Group = R 3, a = b = 46.771(46)Å, c = 6.869(83)Å

Volume = 15028 Å³, α = β = 90, γ = 120

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2.4.5 Comparison of Propidium Iodide to azo-IRMOF-74-III aperture

While it may be possible for a dye which is oriented perfectly perpendicular to the pore channels to escape in dark conditions, other orientations may cause blockage and dye diffusion is limited.

Figure 2.9: Propidium Iodide size without counterion in three dimensions.

Figure 2.10: Space filling model of non-idealized azo-IRMOF-74-III.

Additionally, interactions with the pore wall and collisions with solvent and counter ions hinder motion through the pore. With photoswitching of the azo-
Figure 2.11: Dye diffusion with different orientation through MOF structure.
species from trans to cis, mobility is enhanced.

### 2.4.6 Thermal gravimetric analysis of azo-IRMOF-74-III

![Thermal gravimetric analysis of azo-IRMOF-74-III](image)

Figure 2.12: Thermal gravimetric analysis of azo-IRMOF-74-III while heating sample from 20-800°C.

### 2.4.7 Solid-state $^{13}$C CP/MAS nuclear magnetic resonance spectroscopy.

High resolution solid-state nuclear magnetic resonance (NMR) spectra were measured at ambient pressure on a Bruker DSX-300 spectrometer using a standard Bruker magic angle-spinning (MAS) probe with 4 mm (outside diameter) zirconia rotors. The magic angle was adjusted by maximizing the number and amplitudes of the signals of the rotational echoes observed in the $^{79}$Br MAS FID signal from KBr.

Cross-polarization with MAS (CP/MAS) was used to acquire $^{13}$C data at 75.47 MHz. The $^1$H and $^{13}$C ninety-degree pulse widths were both 4 ms. The CP contact time varied from 1.5 to 5 ms. High power two-pulse phase modulation (TPPM)
$^1$H decoupling was applied during data acquisition. The decoupling frequency corresponded to 72 kHz. The MAS sample-spinning rate was 10 kHz. Recycle delays between scans varied between 3 and 10 s, depending upon the compound as determined by observing no apparent loss in the $^{13}$C signal from one scan to the next. The $^{13}$C chemical shifts are given relative to tetramethylsilane as zero ppm, calibrated using the methylene carbon signal of adamantane assigned to 37.77 ppm as the secondary reference.

Figure 2.13: $^{13}$C NMR of azobenzene link and azo-IRMOF-74-III in the solid state.
2.4.8 Fluorescence monitoring of wavelength specific release experiment

The fluorescence of the solvent above a 100 mg sample of azo-IRMOF-74-III loaded with propidium iodide was monitored with a 448 nm laser in a setup identical to the other release studies within this paper. After a baseline was established, a 647 nm laser was focused on the sample of particles; no release was observed. Subsequently, a 403 nm laser was focused on the sample and an immediate increase in the fluorescence signal was observed.

Figure 2.14: Time-dependent concentration of propidium iodide released from azo-IRMOF-74-III under sequential exposure to 647 nm and 403 nm lasers.
2.4.9 UV-vis determination of loading efficiency of propidium iodide into azo-IRMOF-74-III

Figure 2.15: UV-vis spectra of a propidium iodide solution before and after loading a 100 mg sample of azo-IRMOF-74-III for three days. The 6% difference in concentration corresponds to the amount of dye uptaken by the particles.
2.4.10 UV-vis spectra of an ethanol solution of the azo-linker and magnesium chloride

Figure 2.16: Absorption spectra of a 0.2 mM azo-linker and 0.8 mM MgCl$_2$ solution in ethanol before and after exposure to 377 nm light for 15 s intervals. An isosbestic point at 403 nm is observed.
CHAPTER 3

Heterogeneous Heck Coupling in Multivariate Metal-Organic Frameworks for Enhanced Selectivity

Multivariate metal-organic frameworks (termed MTV-MOFs) are porous crystalline solids in which the pores are lined with multiple organic functional groups. [30, 101–104] This report describes heterogeneous Heck coupling reactions in a series of MTV-MOFs where postsynthetic modification is used to adjust and embed specific amounts of catalytically active Pd(II) ions while avoiding pore blockage. [41, 105–107] MOFs as heterogeneous catalysts have been studied because of their high porosity and their ability to incorporate different catalytic units. [43,44,46,108] A distinct advantage of the MTV platform [47,109] is the ability to control the density of the active sites and the porosity of a catalytic MOF. Varying the percentage of functionalized links, in this case amino-functionalized benzene dicarboxylate (BDC-NH₂) in the original synthesis of the MOF provides a means of tuning the amount of metal catalyst lining the pores and optimization of its activity.

Heck coupling reactions to form aromatic carbon-carbon bonds are one of the most important and useful ways to prepare complex organic molecules. [110] Due to its outstanding stereoselectivity, Heck coupling is of particular interest in the pharmaceutical chemistry [111] and has been demonstrated in both homogeneous and heterogeneous systems previously, including other MOFS. [112,113] To high-
light the advantage of combining the MTV approach with postmodification, we investigated the Heck coupling reaction for the synthesis of resveratrol trimethyl ether, a pharmaceutically relevant precursor. Resveratrol is suspected to play a role in the cardioprotective effects of red wine and is of great interest for possible use in cancer treatment. [114] It is found that MOFs with higher porosity and lower density of metal binding sites achieve higher catalytic turnover and selectivity than MOFs with maximal density of metal binding sites.

The MOF-5 structure with 1,4-benzenedicarboxylate (BDC) was assembled by varying amounts of BDC-NH$_2$ to yield five MOFs, Zn$_4$O(BDC-NH$_2$)$_n$(BDC)$_{(3-n)}$ (n = 3, 2.4, 1.8, 1.2, 0.9, 0.75, 0.6, 0.3 and 0.15). [27, 28] Each compound was postsynthetically modified by reaction with salicylic aldehyde to give the imine derivative and metalated with Pd to give the palladium functionalized framework (Figure 3.1). The structural integrity of the framework was confirmed by a powder X-ray diffraction pattern (PXRD), which was coincident to that of simulated structures. The presence of the imine moiety was confirmed by a resonance at 255 ppm in the $^{15}$N cross-polarization magic angle spinning nuclear magnetic resonance (CP/MAS NMR) spectrum. The MOF was digested in DCl/D$_2$O and analyzed with $^1$H solution-state NMR to ascertain the yield of the imine reaction (See SI).

After fully characterizing our frameworks, the catalytic activity of our MOFs in heterogeneous Heck coupling reaction to form resveratrol trimethyl ether was investigated (Figure 3.2(a)). While Heck coupling has been demonstrated in heterogeneous systems previously, concerns of recyclability, [42,115] selectivity, [116] and catalyst leaching [117] continue to plague current systems and led us to investigate the effect of catalyst loading and porosity on yield and selectivity. Our reaction conditions were selected based on a previously published work involving homogenous palladium catalysts. [118] The Heck coupling was performed by loading 1-bromo-3,5-methoxybenzene (0.434 g, 2 mmol), 4-vinylanisole (0.401
Figure 3.1: Schematic representation of post-synthetic modification and metalation of Zn$_4$O(BDC-NH$_2$)$_3$, X and Y are Cl and CH$_3$CN, respectively.

mL, 3 mmol), triethylamine (0.42 mL, 3 mmol), tetra-n-butylammonium bromide [TBAB] (0.32 mg, 1 mmol), MOF (20 mg), dodecane (1.0 mmol, inert internal standard) and DMA (6 mL) into a Schlenk flask, then heating to 120°C for 12 hours. The reaction mixture was monitored using gas chromatography-mass spectrometry (GC-MS).

The reaction conditions for catalysis were optimized by varying the reaction temperature and monitoring the conversion rate. Temperatures in excess of 120°C did not improve the conversion rate. In addition, each GC-MS spectrum was analysed for both isomers of resveratrol trimethyl ether to determine the trans to cis ratio among the product mixture (Figure 3.2C). Selectively obtaining the trans isomer of resveratrol is of great interest as it has been shown to have greater pharmaceutical activity and cardioprotective activity than the cis isomer. [119]

Figure 3.2B shows the catalytic activity of nine catalysts as well as unmetalated MOF-5 [Zn$_4$O(BDC)$_3$] and their respective surface areas. Interestingly, we see peak catalytic activity for Zn$_4$O(BDC-NH$_2$)$_{0.6}$(BDC)$_{2.4}$-Pd, which has lower metal loading than many other structures but still maintains high surface area. In addition, our MOFs show high selectivity for the trans isomer of resveratrol
trimethyl ether, especially at low metal loading (Figure 3.2C). These results suggest that catalytic conversion varied according to two factors: metalation and porosity. Increased porosity allows for more space in the pore for the aryl halide to perform oxidative addition at the Pd center and for possible cis to trans isomerization at the active site, increasing reaction yield and selectivity in agreement with published reaction mechanisms. [120] Increased BDC-NH₂ loading results in more palladium sites potentially available for catalysis; yet, at higher loadings, these results show that catalytic activity rapidly declines. Peak reaction yield occurs at lower metal loading, but a minimum threshold exists in our frameworks for number of catalytically active sites needed to maintain maximum conversion, in this case 20% BDC-NH₂. The combination of high reactivity and selectivity coupled with exceptional recyclability seen in Figure 3.3 make these MOFs an excellent alternative to current state of the art heterogeneous catalysts such as zeolites [121] and mesoporous silica. [122] However, a direct comparison between our catalyst and these systems cannot be made due to differing reaction substrates and conditions.

We tested two of our synthesized catalysts [Zn₄O(BDC-NH₂)₀.₆(BDC)₂.₄-Pd and Zn₄O(BDC-NH₂)₃-Pd] for catalytic recyclability and possible metal leaching. Our MOFs maintained high crystallinity for up to 10 catalytic cycles (Figure 3.3) as well as high yield and selectivity. In addition, after filtering off the MOF, ICP-AES was performed on the reaction mixture to confirm that no Pd leaching occurred. The heterogeneous nature of the Heck coupling was confirmed by filtration. After the coupling reaction was completed, the MOF was filtered and fresh 1-bromo-3,5-methoxybenzene and 4-vinylanisole were added to the filtrate. Using the filtrate, the reaction was run again under analogous conditions to the initial experiment and no catalytic conversion was observed. In addition, no catalytic conversion was observed if an unmetalated MOF was used in the initial reaction mixture.
Figure 3.2: MOF performance as catalyst in a Heck coupling reaction. (A) Reaction of 4-vinylanisole and 1-bromo-3,5-methoxybenzene catalysed by MOF to form resveratrol trimethyl ether (B) Catalytic conversion as a function of increasing metalation. Indicated specific surface areas (m$^2$/g) were calculated from Brunauer-Emmett-Teller (BET) analysis of isotherms, as shown in the SI section. (C) Selectivity of Heck coupling for each MOF. As metalation decreases, we see higher selectivity for the trans isomer until a plateau is reached below $n = 0.6$ (trans : cis product ratio approximately 106 : 1).

To ensure that no palladium nanoparticles or aggregates were formed during the post-synthetic modification, we performed selective poisoning of our catalyst through use of Hg(0) and PPh$_3$. Hg(0) has been shown to selectively poison metal-particle catalysts by amalgamating the metal and PPh$_3$ selectively poisons molecular single-site catalysts. [123] After adding Hg(0) to our reaction mixture,
Figure 3.3: PXRD analysis for Zn₄O(BDC-NH₂)₃-Pd indicate crystallinity is maintained after multiple catalytic cycles. PXRD spectra of calculated Zn₄O(BDC-NH₂)₃ (black), activated Zn₄O(BDC-NH₂)₃-Pd (blue), Zn₄O(BDC-NH₂)₃-Pd after one catalytic cycle (red), after five catalytic cycles (green), and after ten catalytic cycles (purple). Dark blue ticks indicate the positions of Bragg reflections.

we see no change in reaction yield or selectively, indicating that we have no metal aggregates within our frameworks (see SI). In a separate trial, we added PPh₃ to our reaction mixture and observed an immediate drop-off in catalytic activity (see SI). The results of these tests indicate that we have a single atom Pd catalyst as opposed to Pd nanoparticles or aggregates within the pores.

In conclusion, we have developed a new heterogeneous Heck coupling catalyst and successfully demonstrated its activity by synthesizing resveratrol trimethyl ether. While direct comparisons to other heterogeneous catalysts are difficult due to different reaction substrates and conditions, our catalyst shows an excellent
combination of high activity, selectivity and recyclability. The observed decreased
catalytic activity with increased BDC-NH₂ loading is attributed to the blocking of
access to the palladium sites as more links are reacted with imine functionalities
and Pd, leading to a decrease in overall porosity of the MOFs. The interplay of
metalation and porosity implies that the catalytic activity of the MOFs can be
adjusted to maximize catalytic efficiency.

3.1 Supporting Information

3.1.1 Materials and general procedures

All reagents, unless otherwise stated, were obtained from commercial sources (Alfa
Aesar, Cambridge isotope laboratories, Sigma Aldrich, TCI) and were used with-
out further purification. Reported MOF crystallization yields were unoptimized.
All experiments other than ICP-AES (performed at University of Southern Cali-
ifornia) were performed at the University of California, Los Angeles, Department
of Chemistry and Biochemistry.

3.1.2 Synthesis of all Zn₄O(BDC-NH₂)ₙ(BDC)ₙ₋₃ (ₙ = 3, 2.4, 1.8,
1.2, 0.9, 0.75, 0.6, 0.3 and 0.15) samples

3.1.2.1 Zn₄O(C₈H₄O₄)₂.₈₅(C₈NH₅O₄)₀.₁₅

A mixture of H₂BDC-NH₂ (9 mg, 0.05 mmol), H₂BDC (157 mg, 0.95 mmol)
and zinc nitrate hexahydrate (890 mg, 3.0 mmol) was added to DMF (50 mL)
and sonicated until the components were fully dissolved. The static solution was
heated at 85°C for 24 h to form single crystals. The crystals were collected and
washed with DMF (3 times, 30 mL) over a 3-h period.
3.1.2.2 $\text{Zn}_4\text{O(C}_8\text{H}_4\text{O}_4)_{2.7}\text{(C}_8\text{NH}_5\text{O}_4)_{0.3}$

A mixture of $\text{H}_2\text{BDC-NH}_2$ (18 mg, 0.10 mmol), $\text{H}_2\text{BDC}$ (149 mg, 0.90 mmol) and zinc nitrate hexahydrate (890 mg, 3.0 mmol) was added to DMF (50 mL) and sonicated until the components were fully dissolved. The static solution was heated at 85°C for 24 h to form single crystals. The crystals were collected and washed with DMF (3 times, 30 mL) over a 3-h period.

3.1.2.3 $\text{Zn}_4\text{O(C}_8\text{H}_4\text{O}_4)_{2.4}\text{(C}_8\text{NH}_5\text{O}_4)_{0.6}$

A mixture of $\text{H}_2\text{BDC-NH}_2$ (63 mg, 0.34 mmol), $\text{H}_2\text{BDC}$ (108 mg, 0.65 mmol) and zinc nitrate hexahydrate (890 mg, 3.0 mmol) was added to DMF (50 mL) and sonicated until the components were fully dissolved. The static solution was heated at 85°C for 24 h to form single crystals. The crystals were collected and washed with DMF (3 times, 30 mL) over a 3-h period.

3.1.2.4 $\text{Zn}_4\text{O(C}_8\text{H}_4\text{O}_4)_{2.25}\text{(C}_8\text{NH}_5\text{O}_4)_{0.75}$

A mixture of $\text{H}_2\text{BDC-NH}_2$ (45 mg, 0.25 mmol), $\text{H}_2\text{BDC}$ (124 mg, 0.75 mmol) and zinc nitrate hexahydrate (890 mg, 3.0 mmol) was added to DMF (50 mL) and sonicated until the components were fully dissolved. The static solution was heated at 85°C for 24 h to form single crystals. The crystals were collected and washed with DMF (3 times, 30 mL) over a 3-h period.

3.1.2.5 $\text{Zn}_4\text{O(C}_8\text{H}_4\text{O}_4)_{2.1}\text{(C}_8\text{NH}_5\text{O}_4)_{0.9}$

A mixture of $\text{H}_2\text{BDC-NH}_2$ (116 mg, 0.70 mmol), $\text{H}_2\text{BDC}$ (116 mg, 0.70 mmol) and zinc nitrate hexahydrate (890 mg, 3.0 mmol) was added to DMF (50 mL) and sonicated until the components were fully dissolved. The static solution was heated at 85°C for 24 h to form single crystals. The crystals were collected and
washed with DMF (3 times, 30 mL) over a 3-h period.

3.1.2.6 \( \text{Zn}_4 \text{O} (\text{C}_8 \text{H}_4 \text{O}_4)_{1.8} (\text{C}_8 \text{NH}_5 \text{O}_4)_{1.2} \)

A mixture of \( \text{H}_2 \text{BDC-NH}_2 \) (120 mg, 0.66 mmol), \( \text{H}_2 \text{BDC} \) (100 mg, 0.60 mmol) and zinc nitrate hexahydrate (890 mg, 3.0 mmol) was added to DMF (50 mL) and sonicated until the components were fully dissolved. The static solution was heated at 85°C for 24 h to form single crystals. The crystals were collected and washed with DMF (3 times, 30 mL) over a 3-h period.

3.1.2.7 \( \text{Zn}_4 \text{O} (\text{C}_8 \text{H}_4 \text{O}_4)_{1.2} (\text{C}_8 \text{NH}_5 \text{O}_4)_{1.8} \)

\( \text{H}_2 \text{BDC-NH}_2 \) (136 mg, 0.750 mmol), \( \text{H}_2 \text{BDC} \) (42 mg, 0.25 mmol) and zinc nitrate hexahydrate (890 mg, 3.0 mmol) was added to DMF (50 mL) and sonicated until the components were fully dissolved. The static solution was heated at 85°C for 24 h to form single crystals. The crystals were collected and washed with DMF (3 times, 30 mL) over a 3-h period.

3.1.2.8 \( \text{Zn}_4 \text{O} (\text{C}_8 \text{H}_4 \text{O}_4)_{0.6} (\text{C}_8 \text{NH}_5 \text{O}_4)_{2.4} \)

A mixture of \( \text{H}_2 \text{BDC-NH}_2 \) (163 mg, 0.89 mmol), \( \text{H}_2 \text{BDC} \) (17 mg, 0.10 mmol) and zinc nitrate hexahydrate (890 mg, 3.0 mmol) was added to DMF (50 mL) and the mixture was sonicated until the components were fully dissolved. The static solution was heated at 85°C for 24 h to form single crystals. The crystals were collected and washed with DMF (3 times, 30 mL) over a 3-h period.

3.1.2.9 \( \text{Zn}_4 \text{O} (\text{BDC-NH}_2)_3 \)

The synthesis of IRMOF-3 was analogous to those reported in the literature. [28] H\(_2\)BDC-NH\(_2\) (180 mg, 0.99 mmol) and zinc nitrate hexahydrate (890 mg, 3 mmol)
were added to DMF (50 mL) and the mixture was sonicated until the components were fully dissolved. The static solution was heated at 85°C for 24 h to form single crystals. The crystals were collected and washed with DMF (3 times, 10 mL) repeatedly over a 3-h period. PXRD was used to confirm the phase purity of the as-synthesized samples.

3.1.3 Metalation of Zn₄O(BDC-NH₂)ₙ(BDC)ₙ₋₃ (n = 3, 2.4, 1.8, 1.2, 0.9, 0.75, 0.6, 0.3 and 0.15)

Utilizing a two-step postmodification reaction, imine condensation followed by metalation, all samples were metalated using identical synthetic conditions. [106] For example, a freshly solvent-exchanged sample of Zn₄O(BDC-NH₂)ₙ(BDC)ₙ₋₃ was reacted with salicylic aldehyde in toluene to form an imine bond in the framework. Before metalation could be achieved, the framework was washed to remove excess salicylic acid. After solvent exchange with CH₂Cl₂ was completed, PdCl₂(CH₃CN)₂ in CH₂Cl₂ was added, and allowed to stand at room temperature. After 48 h the sample was exchanged with CH₂Cl₂ to remove excess PdCl₂(CH₃CN)₂. The framework was then exposed to a vacuum (30 mTorr) at 85°C to remove all of the solvent from the pores.

3.1.3.1 Imine formation in Zn₄O(BDC-NH₂)₃

DMF-exchanged Zn₄O(BDC-NH₂)₃ (100 mg) was solvent exchanged with dry toluene (5 20 mL) over a 2-h period. Salicylic aldehyde (1.0 mL, 7.1 mmol) was added to Zn₄O(BDC-NH₂)₃ in toluene and the reaction was allowed to stand at 50°C for 5 days. After 5 days, the sample was exchanged with dry toluene (5 times, 20 mL) to remove unreacted salicylic aldehyde, yielding Zn₄O(BDC-NH₂)₃-Imine.
3.1.3.2 Metalation of Zn$_4$O(BDC-NH$_2$)$_3$

Toluene exchanged Zn$_4$O(BDC-NH$_2$)$_3$-Imine (100 mg) was solvent exchanged with CH$_2$Cl$_2$ (1 times, 20 mL) over a 2-h period. PdCl$_2$(CH$_3$CN)$_2$ (50 mg, 0.19 mmol) was dissolved in CH$_2$Cl$_2$ (40 mL) to which the Zn$_4$O(BDC-NH$_2$)$_3$-Imine was added in a minimum amount of solvent. The reaction was allowed to stand at room temperature for 2 days, at which point it was washed with CH$_2$Cl$_2$ (3 times, 20 mL) over a 2-h period, followed by CH$_2$Cl$_2$ (3 times, 20 mL) over a 3-day period. The solvents were removed from the pores of the framework by vacuum pumping (at 30 mTorr) for 12 h at 85°C to yield Zn$_4$O(BDC-NH$_2$)$_3$-Pd.

3.1.3.3 Imine formation in Zn$_4$O(BDC-NH$_2$)$_n$(BDC)$_{(3-n)}$ (n = 2.4, 1.8, 1.2, 0.9, 0.75, 0.6, 0.3 and 0.15)

DMF exchanged Zn$_4$O(BDC-NH$_2$)$_n$(BDC)$_{(3-n)}$ (100 mg) was exchanged with toluene (5 times, 20 mL) over a 2-h period. Salicylic aldehyde (1.0 mL, 7.1 mmol) was added to Zn$_4$O(BDC-NH$_2$)$_n$(BDC)$_{(3-n)}$ in toluene and the reaction was allowed to stand at 50°C for 5 days. After 5 days, the sample was exchanged with fresh toluene (5 times, 20 mL) to remove unreacted salicylic aldehyde, yielding Zn$_4$O(BDC-NH$_2$)$_n$(BDC)$_{(3-n)}$-Imine.

3.1.3.4 Metalation of Zn$_4$O(BDC-NH$_2$)$_n$(BDC)$_{(3-n)}$-Imine

Toluene-exchanged Zn$_4$O(BDC-NH$_2$)$_n$(BDC)$_{(3-n)}$-Imine was solvent exchanged with CH$_2$Cl$_2$ (20 mL) over a 2-h period. PdCl$_2$(CH$_3$CN)$_2$ (50 mg, 0.19 mmol) was dissolved in CH$_2$Cl$_2$ (20 mL) to which the Zn$_4$O(BDC-NH$_2$)$_n$(BDC)$_{(3-n)}$-Imine was added in a minimum amount of solvent. The reaction was allowed to stand at room temperature for 2 days, at which point it was washed with CH$_2$Cl$_2$ (3 times, 20 mL) over a 2-h period, followed by CH$_2$Cl$_2$ (3 times, 20 mL) over a 3-day period. The solvents were removed from the pores of the framework by vacuum pumping (at 30 mTorr) for 12 h at 85°C to yield Zn$_4$O(BDC-NH$_2$)$_3$-Pd.
(30 mTorr) for 12 h at 85°C to yield Zn₄O(BDC-NH₂)ₙ(BDC)ₙ₋ₚd.

### 3.1.4 Powder X-ray diffraction data

The structural integrity of the unfunctionalized [Zn₄O(BDC-NH₂)ₙ(BDC)ₙ], the imine functionalized framework [Zn₄O(BDC-NH₂)ₙ(BDC)ₙ-Imine], and the Pd functionalized framework [Zn₄O(BDC-NH₂)ₙ(BDC)ₙ-Pd] (n = 3, 2.4, 1.8, 1.2, 0.9, 0.75, 0.6, 0.3 and 0.15) was confirmed by PXRD.

![Figure 3.4: PXRD patterns of unfunctionalized (A, D), imine functionalized (B, E), and metalated frameworks (C, F).](image-url)

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3.1.5 Gas adsorption at 77 K

Upon activation of \( Zn_4O(BDC-NH_2)_n(BDC)_{(3-n)} \) \((n = 3, 2.4, 1.8, 1.2, \text{ and } 0.6)\), the \( N_2 \) isotherms of each material were measured at 77 K. Isotherms at 77 K indicated that metalation had decreased the porosity of \( Zn_4O(BDC-NH_2)_3-Pd \), with the BET surface area of 210 m\(^2\)/g, analogous to other studies of metalation in \( Zn_4O(BDC-NH_2)_3 \). [105] However, \( N_2 \) isotherms at 77 K confirmed other \( Zn_4O(BDC-NH_2)_n(BDC)_{(3-n)} \) samples maintained high porosity upon metalation; the accuracy is 10 m\(^2\)/g for all samples. The maintenance of porosity displays the advantage of the application of the MTV-process in the metalation of MOFs with pores smaller than 1 nm. It is known that MOF samples after postsynthetic modification reactions sometimes show a significant hysteresis loop, [105] and this is most likely due to intercrystalline voids in the sample related to lowered crystallinity of the sample after the postsynthetic modification reaction. [124]

Low-pressure \( N_2 \) adsorption experiments (below 760 Torr) were performed on a Quantachrome NOVA 4200e automatic volumetric gas adsorption analyzer. Ultra-high purity grade \( N_2 \) (99.999% purity) and a liquid nitrogen bath (77 K) were used. For the estimation of surface areas, the BET method was applied using the adsorption branches of the \( N_2 \) isotherms assuming a \( N_2 \) cross-sectional area of 16.2 Å\(^2\)/molecule.
Figure 3.5: Nitrogen isotherms of all Zn\textsubscript{4}O(BDC-NH\textsubscript{2})\textsubscript{n}(BDC)\textsubscript{3-n}-Pd samples.
3.1.6 $^{15}$N and $^1$H solution NMR spectra and ICP-AES analysis

$^{15}$N CP/MAS solid state NMR provides no information about the yield of reactions; therefore, we employed digestion NMR and ICP-MS to quantify the yield of postmodification and metalation techniques that have been used to quantify the yield of postmodification reactions in MOFs. [125] [126] Samples of metlated MOFs were digested in DCl/DMSO-d$_6$, all visible peaks in the spectra were labeled with corresponding hydrogen atoms (letters A-F wherever possible) and the percentage amino-BDC and functionalization for each framework is shown in Figure 3.6. To determine the loading of the frameworks with palladium inductively coupled plasma atomic emission spectroscopy (ICP-AES) was performed, and Zn$_4$O:Pd ratios of 1:0.4, 1:0.7, 1:1.0, 1:1.1, and 1:1.7 for n = 0.6, 1.2, 1.8, 2.4, and 3.0, respectively, could be determined. All ICP-AES measurements have a standard deviation of 10%. MOF samples were digested in DMSO-d$_6$/D$_2$O/DCl.

Liquid-state NMR spectra were performed on a Bruker AX 400 spectrometer.

<table>
<thead>
<tr>
<th>Framework</th>
<th>Expected amino-BDC content (%)</th>
<th>Actual amino-BDC content (%)</th>
<th>Imine formation (% of amino-BDC functionalized)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zn$<em>2$O(BDC-NH$<em>2$)$</em>{0.6}$(BDC)$</em>{2.4}$</td>
<td>5</td>
<td>5.08</td>
<td>90</td>
</tr>
<tr>
<td>Zn$<em>2$O(BDC-NH$<em>2$)$</em>{1.2}$(BDC)$</em>{1.8}$</td>
<td>10</td>
<td>9.97</td>
<td>90</td>
</tr>
<tr>
<td>Zn$<em>2$O(BDC-NH$<em>2$)$</em>{1.8}$(BDC)$</em>{1.2}$</td>
<td>20</td>
<td>20.17</td>
<td>80</td>
</tr>
<tr>
<td>Zn$<em>2$O(BDC-NH$<em>2$)$</em>{2.4}$(BDC)$</em>{0.6}$</td>
<td>25</td>
<td>24.95</td>
<td>80</td>
</tr>
<tr>
<td>Zn$<em>2$O(BDC-NH$<em>2$)$</em>{2}$(BDC)$</em>{2}$</td>
<td>30</td>
<td>30.20</td>
<td>85</td>
</tr>
<tr>
<td>Zn$<em>2$O(BDC-NH$<em>2$)$</em>{2.5}$(BDC)$</em>{1.5}$</td>
<td>40</td>
<td>40.10</td>
<td>80</td>
</tr>
<tr>
<td>Zn$<em>2$O(BDC-NH$<em>2$)$</em>{2.9}$(BDC)$</em>{1.1}$</td>
<td>60</td>
<td>59.06</td>
<td>90</td>
</tr>
<tr>
<td>Zn$<em>2$O(BDC-NH$<em>2$)$</em>{3.4}$(BDC)$</em>{0.6}$</td>
<td>80</td>
<td>79.69</td>
<td>90</td>
</tr>
<tr>
<td>Zn$_2$O(BDC-NH$<em>2$)$</em>{3}$</td>
<td>100</td>
<td>100</td>
<td>70</td>
</tr>
</tbody>
</table>

Figure 3.6: Percentage and degree of functionalization of amino-BDC in MTV–MOFs
Figure 3.7: Digestion NMR spectrum of Zn$_4$O(BDC-NH$_2$)$_{0.15}$(BDC)$_{2.85}$.

Figure 3.8: Digestion NMR spectrum of Zn$_4$O(BDC-NH$_2$)$_{0.3}$(BDC)$_{2.7}$.
Figure 3.9: Digestion NMR spectrum of Zn₄O(BDC-NH₂)₀.₆(BDC)₂.₄.

Figure 3.10: Digestion NMR spectrum of Zn₄O(BDC-NH₂)₀.₇₅(BDC)₂.₂₅.
Figure 3.11: Digestion NMR spectrum of Zn$_4$O(BDC-NH$_2$)$_{0.9}$(BDC)$_{2.1}$.

Figure 3.12: Digestion NMR spectrum of Zn$_4$O(BDC-NH$_2$)$_{1.2}$(BDC)$_{1.8}$.
Figure 3.13: Digestion NMR spectrum of $\text{Zn}_4\text{O} (\text{BDC-NH}_2)_{1.8} (\text{BDC})_{1.2}$.

Figure 3.14: Digestion NMR spectrum of $\text{Zn}_4\text{O} (\text{BDC-NH}_2)_{2.4} (\text{BDC})_{0.6}$. 

Figure 3.15: Digestion NMR spectrum of Zn₄O(BDC-NH₂)₃.
Figure 3.16: Digestion NMR spectrum of Zn$_4$O(BDC-NH$_2$)$_{0.15}$(BDC)$_{2.85}$-Pd.

Figure 3.17: Digestion NMR spectrum of Zn$_4$O(BDC-NH$_2$)$_{0.3}$(BDC)$_{2.7}$-Pd.
Figure 3.18: Digestion NMR spectrum of Zn$_4$O(BDC-NH$_2$)$_{0.6}$BDC$_{2.4}$-Pd.

Figure 3.19: Digestion NMR spectrum of Zn$_4$O(BDC-NH$_2$)$_{0.75}$BDC$_{2.25}$-Pd.
Figure 3.20: Digestion NMR spectrum of Zn₄O(BDC-NH₂)₀.₉(BDC)₂.₁-Pd.

Figure 3.21: Digestion NMR spectrum of Zn₄O(BDC-NH₂)₁.₂(BDC)₁.₈-Pd.
Figure 3.22: Digestion NMR spectrum of Zn$_4$O(BDC-NH$_2$)$_{1.8}$(BDC)$_{1.2}$-Pd.

Figure 3.23: Digestion NMR spectrum of Zn$_4$O(BDC-NH$_2$)$_{2.4}$(BDC)$_{0.6}$-Pd.
Figure 3.24: Digestion NMR spectrum of Zn₄O(BDC-NH₂)₃-Pd.
3.1.7 Solid-state $^{15}$N CP/MAS NMR spectroscopy

The palladium coordination was determined using $^{15}$N CP/MAS solid-state NMR spectroscopy, which characterizes the imine condensation and the metal binding site, with the use of $^{15}$N labeled BDC-NH$_2$. Zn$_4$O(BDC-NH$_2$)$_{1.2}$(BDC)$_{1.8}$ was synthesized using labeled BDC-NH$_2$, $^{15}$N CP/MAS solid-state NMR clearly showed an amine resonance at 65 ppm. [127] Upon postmodification with salicylic acid an imine resonance is observed at 255 ppm in Zn$_4$O(BDC-NH$_2$)$_{1.2}$(BDC)$_{1.8}$-Imine. In addition, the amine resonance is still observed at 68 ppm indicating the reaction does not proceed to completion. The final step of the postmodification in Zn$_4$O(BDC-NH$_2$)$_{1.2}$(BDC)$_{1.8}$, metalation, was further characterized by $^{15}$N NMR, which contained a broad resonance at 10 ppm, the Pd-$^{15}$N bond of the imine complex. This resonance was confirmed by comparison to a model compound Pd(C$_{13}$H$_{10}$NO)$_2$, which also showed two resonances, at 5 and 13 ppm (Figure 3.25).

![Figure 3.25: $^{15}$N solution spectrum of a model compound Pd(C$_{13}$H$_{10}$NO)$_2$ used to confirm the metal binding in the solid state NMR experiments.](image)

High resolution solid-state NMR spectra were recorded at ambient pressure on a Bruker DSX-300 spectrometer using a standard Bruker MAS probe with 4 mm (outside diameter) zirconia rotors. The magic angle was found by maximizing the number and amplitudes of the signals of the rotational echoes observed in the $^{79}$Br
MAS free induction signal from KBr.

$^{15}$N CP MAS for the isotopically labeled materials were measured at 30.42 MHz. The $^1$H and $^{15}$N ninety-degree pulse widths were both 4 s. The CP contact time varied from 1.5 to 5 ms. The direct excitation NMR was measured at recycle delays of up to 45 seconds to allow for complete relaxation. The $^{15}$N chemical shifts are given relative to liquid N$_2$ at zero ppm, calibrated using the nitrogen signal of glycine assigned to 36.2 ppm as secondary reference.

Figure 3.26: $^{15}$N CP/MAS NMR spectrum of Zn$_4$O(BDC-NH$_2$)$_{1.2}$(BDC)$_{1.8}$.

The peak between 100 and 200 is most likely the result of protonation of the amine. This downfield shift of approximately 100 ppm from the free amine has been observed by our group [128] and other groups. [129]
Figure 3.27: $^{15}$N CP/MAS NMR spectrum of Zn$_4$O(BDC-NH$_2$)$_{1.2}$(BDC)$_{1.8}$-Imine.

Figure 3.28: $^{15}$N CP/MAS NMR spectrum of Zn$_4$O(BDC-NH$_2$)$_{1.2}$(BDC)$_{1.8}$-Pd.
3.1.8 Optical microscopy

Optical microscopy of Zn$_4$O(BDC-NH$_2$)$_{1.2}$(BDC)$_{1.8}$, Zn$_4$O(BDC-NH$_2$)$_{1.2}$(BDC)$_{1.8}$-Imine, Zn$_4$O(BDC-NH$_2$)$_{1.2}$(BDC)$_{1.8}$-Pd was carried out using a Leica optical microscope. The as-synthesized samples were dispersed onto a glass plate for imaging.

Figure 3.29: Optical microscopy of Zn$_4$O(BDC-NH$_2$)$_{1.2}$(BDC)$_{1.8}$, Zn$_4$O(BDC-NH$_2$)$_{1.2}$(BDC)$_{1.8}$-Imine, Zn$_4$O(BDC-NH$_2$)$_{1.2}$(BDC)$_{1.8}$-Pd, from left to right.

3.1.9 Catalytic Testing: Experimental Setup

In a 25 mL Schlenk flask, 1-bromo-3,5-methoxybenzene (0.434 g, 2 mmol), 4-vinylanisole (0.401 mL, 3 mmol), triethylamine (0.42 mL, 3 mmol), tetra-n-butylammonium bromide (0.32 mg, 1 mmol), MOF (20 mg), dodecane (1.0 mmol, inert internal standard) were added to DMA (6 mL). After rapidly stirring for 5 minutes, the reaction was heated to 120°C for 12 hours and was monitored via GC-MS.

Experiments with a palladium homogeneous catalyst used the following procedure:

In a 25 mL Schlenk flask, 1-bromo-3,5-methoxybenzene (0.434 g, 2 mmol), 4-vinylanisole (0.401 mL, 3 mmol), triethylamine (0.42 mL, 3 mmol), tetra-n-butylammonium bromide (0.32 mg, 1 mmol), (MeCN)$_2$Pd(Cl)$_2$ (10 mol%), dodecane (1.0 mmol, inert internal standard) were added to DMA (6 mL). After
rapidly stirring for 5 minutes, the reaction was heated to 120°C for 12 hours and was monitored via GC-MS.

3.1.10 Determination of Catalytic Activity: GC-MS

**Instrument description.** GC-MS measurements were carried out using an Agilent Model 7683 Autosampler, 6890 Gas Chromatograph, and 5975 Inert MassSelective Detector in the Electron Impact (EI) mode. EI energy was set to 70 eV. Data collection was controlled using MSD Enhanced Chemstation software (Agilent). Separation was carried out on an Agilent HP5-MS column with dimensions 30 m×250 m×0.25 m. Ultra High Purity Grade He (Airgas) was used as carrier gas with the flow set to 0.8 mL/min in constant flow mode.

**Data collection parameters.** 1 L of sample dissolved in dichloromethane (Fisher) was injected using a split ratio of 20:1, with the inlet temperature set to 280°C. The initial oven temperature was set to 45°C for 1 minute followed by a 30°C/min ramp to a final temperature of 300°C which was maintained for 3 min. A 4.5 min solvent delay was used. The MSD was set to scan the 40 — 1050 m/z range.

**Data analysis.** Mass Spectrometric data was analyzed using the MSD Enhanced Chemstation software. Product spectra were identified by comparison of the measured fragmentation patterns to those found in the NIST 08 Mass Spectral Library.
Figure 3.30: GC-MS of pure reactants and product for Heck coupling to establish retention times.

To determine the retention times of the isomers of resveratrol trimethyl ether, a sample of trans-resveratrol trimethyl ether was placed under U. V. irradiation (294 nm for 60 mins). GC-MS experiments of the resveratrol trimethyl ether was performed both before and after irradiation and shown in Figures 3.31 and 3.32, respectively.
Figure 3.31: GC-MS of resveratrol trimethyl ether before U.V. irradiation (294 nm for 60 min).
Figure 3.32: GC-MS of resveratrol trimethyl ether after U.V. irradiation.
Figure 3.33: GC-MS of Heck coupling reaction to synthesize resveratrol trimethyl ether.
3.1.11 Catalyst Results and Recycling Experiments

The yield and selectivity of all MOFs and a homogeneous palladium catalyst are shown in Figure 3.34. We can see that some MOFs show higher yield and selectivity than the homogeneous catalyst.

<table>
<thead>
<tr>
<th>Catalyst used</th>
<th>Yield (%)</th>
<th>Trans to cis ratio in product</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zn₄O(BDC-NH₂)₀.₆(BDC)₂.₄</td>
<td>99.3 ± 0.02</td>
<td>105.9 :1</td>
</tr>
<tr>
<td>Zn₄O(BDC-NH₂)₁.₂(BDC)₁.₈</td>
<td>89.7 ± 1.7</td>
<td>81.9 :1</td>
</tr>
<tr>
<td>Zn₄O(BDC-NH₂)₁.₈(BDC)₁.₂</td>
<td>75.5 ± 1.4</td>
<td>64.8 :1</td>
</tr>
<tr>
<td>Zn₄O(BDC-NH₂)₂.₄(BDC)₀.₆</td>
<td>33.4 ± 3.8</td>
<td>53.9 :1</td>
</tr>
<tr>
<td>Zn₄O(BDC-NH₂)₃</td>
<td>15.5 ± 1.6</td>
<td>46.6 :1</td>
</tr>
<tr>
<td>(MeCN)₂Pd(Cl)₂</td>
<td>92.9 ± 0.9</td>
<td>58.5 :1</td>
</tr>
</tbody>
</table>

Figure 3.34: Yield and selectivity for MOF and homogeneous catalysts.

Two MOFs [Zn₄O(BDC-NH₂)₀.₆(BDC)₂.₄-Pd and Zn₄O(BDC-NH₂)₃-Pd] were tested for recyclability to ensure that our frameworks maintain high yield after repeated catalytic cycles. As seen in Figure 3.35, high catalytic activity and selectivity is maintained for 10 cycles.
<table>
<thead>
<tr>
<th>Framework used</th>
<th>Average Yield for 10 cycles (%)</th>
<th>Average trans to cis ratio in product</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zn$<em>4$O(BDC-NH$<em>2$)$</em>{0.6}$(BDC)$</em>{2.4}$-Pd</td>
<td>99.3 ± 0.002</td>
<td>105.8 ± 6.3</td>
</tr>
<tr>
<td>Zn$_3$O(BDC-NH$_2$)$_3$-Pd</td>
<td>15.5 ± 1.6</td>
<td>46.5 ± 2.5</td>
</tr>
</tbody>
</table>

Figure 3.35: Average yield and trans to cis ratio of product for two MOFs after 10 catalytic cycles.
3.1.12 Catalyst control and poisoning results

Control studies were performed to test that our metalated MOFs are the catalytically active species. Heck coupling reactions were run without any MOF present and with unmetalated MOFs to ensure that our frameworks without Pd are not catalytically active. As seen in Figure 3.36, no catalytic conversion was observed if MOF is omitted or an unmetalated MOF is used. Reaction conditions, other than omitted species, are analogous to those of other catalytic runs, namely mixing MOF (20 mg), DMA (6 mL), 1-bromo-3,5-methoxybenzene (0.434 g, 2 mmol), 4-vinylanisole (0.401 mL, 3 mmol), triethylamine (0.42 mL, 3 mmol), tetra-n-butylammonium bromide (0.32 mg, 1 mmol), and dodecane (1.0 mmol, inert internal standard). This mixture was stirred rapidly for 5 minutes before heating to 120°C and monitoring the reaction via GC-MS.

Catalyst poisoning experiments were performed to test that our metalated MOFs have a single site Pd catalyst and that Pd nanoparticles or aggregates have not formed during catalyst synthesis. Heck coupling reactions were run using Zn$_4$O(BDC-NH$_2$)$_{0.6}$(BDC)$_{2.4}$-Pd with the addition of either Hg(0) or PPh$_3$. As seen in Figure 3.37, no change in catalytic conversion was observed if Hg(0) is added to our reaction mixture. Figure 3.38 shows immediate drop-off of catalytic activity followed by cessation of all catalytic activity.

Reaction conditions are analogous to those of other catalytic runs, namely mixing MOF (20 mg), DMA (6 mL), 1-bromo-3,5-methoxybenzene (0.434 g, 2 mmol), 4-vinylanisole (0.401 mL, 3 mmol), triethylamine (0.42 mL, 3 mmol), tetra-n-butylammonium bromide (0.32 mg, 1 mmol), and dodecane (1.0 mmol, inert internal standard). For Hg(0) poisoning experiments, 0.25 mL Hg(0) was added to the reaction mixture and for PPh$_3$ poisoning experiments, 80 mg of PPh$_3$ was added to the reaction mixture. These mixtures was stirred rapidly for 5 minutes before heating to 120 °C and monitoring the reaction via GC-MS.
<table>
<thead>
<tr>
<th>Framework used</th>
<th>Yield (%)</th>
<th>Yield (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zn₄O(BDC-NH₂)₀.₆(BDC)₁.₄</td>
<td>0.0</td>
<td>0.0</td>
</tr>
<tr>
<td>Zn₄O(BDC-NH₂)₁.₂(BDC)₁.₈</td>
<td>0.0</td>
<td>0.0</td>
</tr>
<tr>
<td>Zn₄O(BDC-NH₂)₁.₈(BDC)₁.₂</td>
<td>0.0</td>
<td>0.0</td>
</tr>
<tr>
<td>Zn₄O(BDC-NH₂)₂.₄(BDC)₀.₆</td>
<td>0.0</td>
<td>0.0</td>
</tr>
<tr>
<td>Zn₄O(BDC-NH₂)₃</td>
<td>0.0</td>
<td>0.0</td>
</tr>
</tbody>
</table>

Figure 3.36: Reaction yields of Heck coupling after omission of MOF or using unmetalated MOF. All measurements have an uncertainty of 1%.
Figure 3.37: Reaction progress shows no change after Hg(0) is added to reaction mixture.
Figure 3.38: Reaction progress shows loss of catalytic activity after PPh$_3$ is added to reaction mixture.
3.1.13 ICP-AES of reaction product

Samples of the reaction product were submitted to USC to determine palladium leaching after the Heck coupling reaction had occurred. ICP-AES (standard deviation = 2.8%) indicated leaching was within the experimental error of the experiment and we conclude that no leeching occurred during the Heck coupling reaction.

3.1.14 X-ray photoelectron spectroscopy (XPS)

X-ray photoelectron spectroscopy (XPS) was employed to investigate the oxidation state of the Pd in Zn$_4$O(BDC-NH$_2$)$_{1.2}$(BDC)$_{1.8}$-Pd and Zn$_4$O(BDC-NH$_2$)$_{3}$-Pd. In both samples only photoelectrons corresponding to Pd$^{2+}$ are observed prior to catalysis, as expected for a square planar complex (Figure 3.33). However, after catalytic tests, XPS reveals both MOF samples contain Pd$^{2+}$ and Pd$^0$, indicating that during the reaction, Pd$^{2+}$ is reduced (Figure 3.39). This reduction is also observed in a color change from purple to black in each MOF sample. XPS was performed using a Kratos AXIS Ultra DLD (acquired under NSF CRIF:MU award number 0840531) with the charge neutralizer on. The finely powdered samples were supported using non-conductive double-sided tape. Spectra were analyzed using CasaXPS, fitting the peak areas to a Gaussian/Lorentzian (GL(30)) product form lineshape ($m = p/100$, in which $p = 30$) (Figure 3.39(inset)). Pd 3d spectra were fitted and plotted using the GL(30) lineshape.
Figure 3.39: XPS Spectra of Zn\(_4\)O(BDC-NH\(_2\))\(_3\)-Pd and Zn\(_4\)O(BDC-NH\(_2\))\(_{1.2}\)(BDC)\(_{1.8}\)-Pd.
CHAPTER 4

Epoxidation of Alkenes with Molecular Oxygen Catalyzed by a Manganese Porphyrin-Based Metal-Organic Framework

Here, MOFs are extended crystalline structures comprising metal clusters joined by organic links. [101,130] Due to the tunability of the reticular framework, MOFs may be designed and synthesized for specific catalytic reactions. Heterogenous MOF catalysis [46,131] was demonstrated in a number of reactions including hydrogenation, [132] oxidation, [133] and Knoevenagel condensations. [43] Recently, MOFs with metalloporphyrin linkers have been synthesized and their catalytic applications explored. [134–136] Industrially-viable catalytic MOFs are still unknown due to framework degradation upon exposure to harsh conditions. [137–139]

Epoxidation reactions are important to the chemical industry as oxides of small alkenes are produced in million ton quantities each year and are used as starting points for the production of common chemicals such as polyether polyol and ethylene glycol. [57, 140] The catalysis of epoxidation reactions at mild conditions using molecular oxygen would be an important development. This study demonstrates the epoxidation of several alkenes with molecular oxygen by a manganese porphyrin containing MOF (Figure 4.1).

Molecular oxygen, commonly used in industrial catalysis, is an ideal oxidant for epoxidations due to its low cost and environmentally friendly nature. [55] Earlier studies involving homogenous epoxidation of alkenes using manganese porphyrin
MOF-525-Mn (5.0 mg) CH$_2$Cl$_2$ (10 mL), isobutyraldehyde (10 mmol), O$_2$ (30 sccm), 2-4 hours, room temperature

Free space in the pore is illustrated by a yellow sphere that contacts the van der Waals radii of the framework atoms.

Figure 4.1: Epoxidation of styrene using MOF-525-Mn and molecular oxygen.

show high reactivity when using molecular oxygen as the oxidant. [141] While epoxidations in MOFs using molecular oxygen have been reported, [142] thus far, only low conversion rates and leeching of embedded catalysts has been shown. This article details a zirconium MOF with manganese porphyrin active site which uses molecular oxygen for enhanced catalytic conversion rates and decreased leeching of catalytic species during epoxidation.

Heterogeneous catalysts eliminate the need for steps to separate product from catalyst. The immobilization of homogeneous catalysts has been a long-standing problem in catalysis and MOFs could be useful platforms to this end. We sought to incorporate this high catalytic activity into an extended framework by selecting the MOF-525 structure for its potentially high density of catalytically active sites and the high thermal and chemical stability of the zirconium secondary building unit. [51] While other catalytically-active porphyrin-containing MOFs have been reported, [44, 143, 144] this is the first instance where a porphyrin MOF uses...
molecular oxygen to catalyze epoxidation of alkenes.

The new MOF, termed MOF-525-Mn \([\text{Zr}_6\text{O}_{4}(\text{OH})_4(\text{MgC}_{48}\text{H}_{24}\text{O}_8\text{N}_4\text{Cl})_3]\), is analogous to a previously reported MOF \([145]\) where zirconyl chloride octahydrate (12.5 mg, 0.037 mmol) and Mn(III) meso-Tetra (4-carboxyphenyl) porphine chloride (TCPP-Mn) (2.5 mg, 0.0028 mmol) was added to N,N-dimethylformamide (DMF, 10 mL) in a 20 mL scintillation vial and sonicated for 30 minutes. After sonication, acetic acid (2.5 mL) was added to the vial and heated to 65°C for 72 hours to form MOF-525-Mn. The structure of MOF-525-Mn was confirmed by coincidence of the observed PXRD with the simulated pattern (see SI) and has a pore size of 19 Å. Full manganese metal loading in the framework was confirmed by UV-Vis spectroscopy and ICP-AES. \(\text{N}_2\) adsorption isotherm measurements at 77 K revealed a BET surface area of 1850 \(\text{m}^2/\text{g}\).

Reaction conditions were selected based on a previously published work involving homogenous manganese metalloporphyrins, \([146]\) with the exception that in our heterogeneous system the MOF catalyst could easily be separated from the reaction solution at the end of the catalytic run using a simple filtration step. The epoxidation reactions were performed by bubbling \(\text{O}_2\) (30 sccm) through a mixture of methylene chloride (10 mL), isobutyaldehyde (0.913 mL, 10 mmol), alkene (2 mmol), MOF (5 mg) and heptane (1.0 mmol, inert internal standard) in a reaction vessel for an extended period of time. The reaction mixture was monitored using GC-MS. The results for catalysis of different alkenes by both MOF-525-Mn and the free TCPP-Mn linker are summarized in Figure 4.2, which lists the main reaction products for each alkene reactant and corresponding reaction yields and conversion rates. Note that the size of all alkenes and sacrificial aldehyde are small enough to fit inside the MOF pore (19 Å in diameter), allowing full access to all active sites in the framework. Some yields and conversion rates for the smaller alkenes are close to maximal for MOF-525-Mn. Ethylene and propylene were not investigated in this study due to the gas-phase nature of these
reactions, requiring an entirely different reaction setup than the one used here.

<table>
<thead>
<tr>
<th>Reactant</th>
<th>Product</th>
<th>Time (h)</th>
<th>Conversion (%)</th>
<th>MOF Yield (%)</th>
<th>Yield free linker (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>2.5</td>
<td>99</td>
<td>82.8 ± 0.2</td>
<td>97.2 ± 0.1</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2.0</td>
<td>99</td>
<td>96.9 ± 0.1</td>
<td>97.1 ± 0.2</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3.0</td>
<td>99</td>
<td>96.8 ± 0.3</td>
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<tr>
<td></td>
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<td>4.0</td>
<td>74</td>
<td>70.0 ± 0.1</td>
<td>84.5 ± 0.2</td>
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<td></td>
<td></td>
<td>4.0</td>
<td>75</td>
<td>67.9 ± 0.2</td>
<td>90.3 ± 0.3</td>
</tr>
</tbody>
</table>

Figure 4.2: Epoxidation of alkenes catalyzed by MOF-525-Mn and free TCPP-Mn with molecular oxygen and isobutyraldehyde.

In the case of styrene epoxidation, GC-MS analysis revealed a series of reaction products, with the main reaction product being styrene oxide (see Figure 4.9). Due to the large number of side products, we optimized the styrene reaction yield by varying reactor temperature. While this optimization technique could be applied to the other alkene reactions, we decided to focus on styrene and therefore analyzed the reaction kinetics to determine the main parameters of the epoxidation reaction pathway (Figure 4.3). Industrial applications of styrene epoxidation include epoxy resins and production of chemicals such as phenethyl alcohol and styrene glycol. [147]

The reactor was modeled as a constant-volume batch reactor. Rapid stirring of the reaction mixture served to ensure homogenous mixing of all components in the reactor, and the concentration of dissolved oxygen in the reaction mixture was assumed constant due to the large and continuous flux of oxygen gas into
Figure 4.3: Reaction pathway for the epoxidation of styrene to styrene oxide including side products. Relative yields of each species measured at equilibrium are shown as percentages.
the reaction vessel. The reaction pathway is assigned the pseudo-rate constants \( (k'_1,k'_2,k'_3) \). The latter are assumed to follow Arrhenius behavior:

\[
k'_i(T) = A_i(T)e^{-E_{a,i}/RT}
\]

where \( E_{a,i} \) is the activation energy for the \( i \)-th pathway, \( R \) is the ideal gas constant, \( T \) is the temperature and \( A_i(T) \) is the pre-exponential prefactor. The pseudo-rate constant for styrene epoxidation, \( k'_1 \), is related to the inherent rate constant, \( k_1 \), according to \( k'_1 = k_1 [O_2] \), where \([O_2]\) is the concentration of oxygen dissolved in the solvent.

The apparent rate constants and reaction orders were computed over the temperature range 293-329 K by fitting the decline in styrene concentration over time (see SI section for details of the procedure). The pseudo-rate constants at different temperatures were used to derive the activation energy of the reaction from a fit to 4.1. The temperature dependence of the reaction pseudo-rate constants for MOF-525-Mn was calculated from experimental data. A linearized Arrhenius plot showing the fit over the temperature range 293-329 K is found in Figure 4.4. The styrene epoxidation reaction was found to be first order with respect to styrene, and the activation energy for the epoxidation of styrene was found to be \( 42.6 \pm 5.7 \) kJ/mol. This activation energy agrees with calculated theoretical values within experimental error. [148] (We are unaware of any experimentally measured values for \( E_a \) reported in the literature at time of writing.) The reaction order for this heterogeneous catalyst is in agreement with published results from homogeneous manganese porphyrin systems. [149–151] This result suggests that the manganese porphyrin heterogeneous catalyst likely operates according to a mechanism similar to its homogeneous counterpart.

The heterogeneous nature of the epoxidation reaction was confirmed by filtration. After the initial reaction of styrene, the MOF was filtered and fresh isobu-
tyraldehyde and styrene were added to the filtrate. Using the filtrate, the reaction was run again under analogous conditions to the initial experiment and no catalytic conversion was observed. In addition, no catalytic conversion was observed if isobutyraldehyde, MOF-525-Mn, or metalated porphyrin were not present in the initial reaction mixture.

MOF-525-Mn showed minimal deactivation and maintained its structural stability and crystallinity at the end of the catalytic cycle, as evidenced by structural analysis with PXRD (Figure 4.5). Leeching of manganese and porphyrin from MOF-525-Mn was investigated by performing ICP-AES and elemental analysis on the product solution after filtration. We found no evidence of leeching within experimental error of these measurement techniques. XPS measurements indicated an oxidation state of Mn(III) within the framework and no change in oxidation state was observed after catalysis (Figure 4.5). To test catalytic activity after multiple runs, several cycles of epoxidations were run back-to-back using the
Figure 4.5: Catalytic recycling of MOF-525-Mn (a) PXRD of MOF-525-Mn after multiple catalytic runs indicates crystallinity is maintained after 6 catalytic cycles (b) XPS spectra for Mn 3s of TCPP-Mn linker and MOF-525-Mn. The data has been fit to the sum of two Gaussian, splitting is indicative of Mn(III) and does not change after catalysis (c) Product yield after multiple catalytic runs by MOF-525-Mn. High activity is maintained for all alkenes for 6 catalytic cycles.

The same MOF sample, filtering it between each run while monitoring the reaction via GC-MS. High catalytic activity was maintained for all alkenes after 6 catalytic cycles (Figure 4.5).
The newly developed heterogeneous catalyst, MOF-525-Mn, based on manganese porphyrin, catalyzes epoxidation reactions of alkenes using molecular oxygen. The catalyst exhibits high yields, excellent recyclability and chemical stability with respect to different alkenes tested. The ability to use molecular oxygen in heterogeneous epoxidation reactions may lead to improved strategies for industrially viable processes.
4.1 Supporting Information

4.1.1 Materials and general procedures

All reagents unless otherwise stated were obtained from commercial sources (Frontier Scientific, Cambridge Isotope laboratories, Sigma Aldrich, TCI) and were used without further purification. Yields reported were unoptimized. All synthesis and reaction experiments were performed at the University of California, Los Angeles, Department of Chemistry and Biochemistry. Elemental analysis was performed at UC Davis Stable Isotope Facility and inductively coupled plasma atomic emission spectroscopy (ICP-AES) was performed at USC.

4.1.2 Synthetic procedure for MOF-525-Mn

4.1.2.1 \([\text{Zr}_6\text{O}_4(\text{OH})_4(\text{MgC}_{48}\text{H}_{24}\text{O}_8\text{N}_4\text{Cl})_3]\)

Zirconyl chloride octahydrate (12.5 mg, 0.037 mmol) was added to DMF (10 mL) and sonicated for 30 minutes. After sonication, Mn(III) meso-Tetra (4-carboxyphenyl) porphine chloride (TCPP-Mn) (2.5 mg, 0.0028 mmol) was added to the solution. After 10 minutes of further sonication, acetic acid (2.5 mL) was added to the solution. The solution was placed into a 20 mL scintillation vial and heated to 65°C for 72 hours. The microcrystalline powder was filtered and washed with DMF (5 x 10 mL) over a three-hour period. The DMF was replaced with acetone (5 x 30 mL) over a five-day period. Finally, the acetone was removed by heating by 120°C under vacuum (40 mTorr) for 48 hours.

4.1.3 Powder X-ray diffraction data

The structural integrity of activated MOF-525-Mn was confirmed by PXRD, which showed peak position analogous to MOF-525 (Figure 4.6). The PXRD patterns for synthesized and activated MOF-525-Mn are shown in Figure 4.6.
4.1.4 Gas adsorption at 77 K

Low-pressure gas adsorption isotherms of MOF-525-Mn were measured volumetrically and the BET surface area was found to be 1850 m$^2$/g. The N$_2$ isotherms of MOF-525-Mn were measured using an NOVA surface area analyzer (Quantachrome Instruments). A liquid nitrogen sample bath (77 K) was used for N$_2$ measurements and the N$_2$ gas used was UHP grade. For measurement of the surface areas, the BET method was applied using the adsorption branches of the N$_2$ isotherms assuming a N$_2$ cross-sectional area of 16.2 Å$^2$/molecule.

4.1.5 ICP-AES results for metalation

To ensure the manganese metal content of the frameworks, ICP-AES was carried out, which revealed a Zr:Mn ratio of 1:0.5, confirming the porphyrin is fully metalated inside the framework.
4.1.6 UV-Vis spectroscopy

UV-Vis spectrophotometry measurements were performed on a Shimadzu UV1800 using 1 cm Hellma quartz optical cells and used to confirm the presence of manganese in MOF-525-Mn. A basic solution (1M NaOH) was used to dissolve MOF-525-Mn and the TCPP-Mn link for UV-Vis analysis. Literature values of porphyrins containing manganese show peaks at 468, 562, 596 nm. [152] These peaks are seen in both the TCPP-Mn link and MOF-525-Mn (Figure 4.8)
Figure 4.8: UV-Vis spectra of TCPP-Mn link and MOF-525-Mn show absorbance peaks at 468, 562, 596 nm, confirming the presence of manganese.
4.1.7 X-ray photoelectron spectroscopy (XPS)

XPS was employed to investigate the oxidation state of the Mn in MOF-525-Mn and TCPP-Mn. Both pre and post-catalysis MOF-525-Mn exhibit the same Mn 3s spectrum with a splitting of 5.49 eV, corresponding to Mn(III). The TCPP-Mn linker exhibited a Mn 3s splitting of 5.54 eV which also corresponds to Mn(III). XPS was performed using a Kratos AXIS Ultra DLD (acquired under NSF CRIF:MU award number 0840531) with the charge neutralizer on. The finely powdered samples were supported using non-conductive double-sided tape. Spectra were analyzed using MATLAB, fitting the peak areas to a sum of two Gaussians.

4.1.8 Catalytic Testing: Experimental Setup

In a 25 mL Schlenk flask, isobutyraldehyde (0.913 mL, 10 mmol), alkene (2 mmol), MOF-525-Mn (5 mg) and heptane (1.0 mmol, inert internal standard) were added to methylene chloride (10 mL). After rapidly stirring for 5 minutes, bubbling O₂ (30 sccm) was introduced and the reaction was monitored via gas chromatography-mass spectroscopy (GC-MS).

4.1.9 Determination of Catalytic Activity: GC-MS

Instrument description. GC-MS measurements were carried out using an Agilent Model 7683 Autosampler, 6890 Gas Chromatograph, and 5975 Inert Mass Selective Detector in the Electron Impact (EI) mode. EI energy was set to 70 eV. Data collection was controlled using MSD Enhanced Chemstation software (Agilent). Separation was carried out on an Agilent HP5-MS column with dimensions 30 m x 250 m x 0.25 m. Ultra High Purity Grade He (Airgas) was used as carrier gas with the flow set to 0.8 mL/min in constant flow mode.

Data collection parameters. 1 L of sample dissolved in dichloromethane
(Fisher) was injected using a split ratio of 20:1, with the inlet temperature set to 280°C. The initial oven temperature was set to 45°C for 1 minute followed by a 30°C/min ramp to a final temperature of 300°C which was maintained for 3 min. A 3.2 min solvent delay was used. The MSD was set to scan the 40 – 1050 m/z range.

**Data analysis.** Mass Spectrometric data was analyzed using the MSD Enhanced Chemstation software. Product spectra were identified by comparison of the measured fragmentation patterns to those found in the NIST 08 Mass Spectral Library.
Figure 4.9: GC-MS spectrum of styrene epoxidation reaction.
Figure 4.10: GC-MS spectrum of cyclohexene epoxidation reaction.
Figure 4.11: GC-MS spectrum of cyclooctene epoxidation reaction.
Figure 4.12: GC-MS spectrum of trans-stilbene epoxidation reaction.
Control studies were performed to test that MOF-525-Mn is the catalytically active species and that sacrificial aldehyde, isobutyraldehyde, is necessary. Epoxidation reactions were also run with unmetalated MOF-525 to ensure that unmetalated MOF-525 is not catalytically active. As seen in Figure 4.14, no catalytic conversion was observed if MOF-525-Mn or isobutyraldehyde are omitted or if un-
metalated MOF-525 is used. Reaction conditions, other than omitted species, are analogous to those of other catalytic runs, namely mixing MOF (5 mg), methylene chloride (10 mL), isobutyraldehyde (0.913 mL, 10 mmol), alkene (2 mmol). This mixture was stirred rapidly for 5 minutes before adding bubbling O₂ (30 sccm) and monitoring the reaction via GC-MS.

### 4.1.11 Elemental analysis and ICP-AES of reaction product

Samples of the reaction product were submitted to UC Davis Stable Isotope Facility and USC to determine porphyrin and manganese leeching, respectively, after the epoxidation reaction had occurred. Both elemental analysis (standard deviation = 0.3%) and ICP-AES (standard deviation = 2.8%) indicated leeching was within the experimental error of the experiment and we conclude that no leeching occurred during the epoxidation reaction.
4.1.12 Reaction Rate Kinetics and Fitting of GC-MS data in MATLAB

The reaction vessel is modeled as a constant volume batch reactor due to the absence of liquid streams entering or exiting the vessel. Though oxygen is bubbled constantly through the liquid mixture, this was found to cause minimal volume change from \( \text{O}_2 \) induced evaporation. Under these assumptions, the general mass balance for styrene (denoted with subscript A) in the reaction vessel is

\[
\frac{dC_A}{dt} = \dot{r}_A
\]  

(4.2)

where \( \dot{r}_A \) is the rate of generation of styrene. Since the styrene participates in three reactions (conversion into styrene oxide, benzaldehyde, and benzeneacetaldehyde, see Figure 4.3 in the main text) the rate term has three contributions

\[
\dot{r}_A = -k_1C_A^{\alpha}[\text{O}_2]^\beta - k_2[C_A]^\gamma - k_3[C_A]^\delta
\]  

(4.3)

where the rate constants \( k_1, k_2, \) and \( k_3 \) apply to each of the three major reactions that the styrene takes part in, and \( \alpha, \beta, \gamma, \) and \( \delta \) are the reaction orders for each reaction.

In this study, the dissolved oxygen concentration [\( \text{O}_2 \)] is approximately constant across all trials due to the large excess of gaseous oxygen used, and is therefore incorporated into \( k_1 \) (relabeled \( k'_1 \)) to form the pseudo rate equation:

\[
\dot{r}_A = -k'_1C_A^{\alpha} - k_2[C_A]^\gamma - k_3[C_A]^\delta
\]  

(4.4)

Equation 4.4 was used in equation 4.2 in order to fit rate parameters to the concentration data collected for styrene. Time derivatives were computed by fitting a polynomial equation to \( C_A \) as a function of time and differentiating the resulting expression. All reactions were found to be first order with respect to
Figure 4.15: Temperature dependence of the pseudo kinetic rate constant $k_1'(T)$ of MOF-525-Mn.

styrrene by fitting experimentally measured time-course concentrations of styrene and styrene oxide with Equation 4.4. Using a multiple start-point solver for constrained gradient-based optimization (Global Optimization Toolbox, MATLAB), the rate constants ($k_1', k_2, k_3$) and reaction orders ($\alpha, \gamma, \delta$) that minimized the residuals between the proposed rate model and the empirically observed rate of styrene consumption were identified. The globally optimal parameters for the data fit indicated that exponential values ($\alpha, \gamma, \delta$) were equal to one (first order kinetics). The pseudo rate constants $k_1'$ calculated by the solver at different temperatures are the values reported in the main text. An Arrhenius fit to $k_1'$ values at different temperatures was used to report the activation energy in the main text. The temperature dependent plot of $k_1'$ is shown in Figure 4.15.
CHAPTER 5

Conclusions and Future Directions

The research demonstrated in this dissertation explores several important aspects of MOFs and their use in catalysis. The azobenzene containing MOF-74 shown in Chapter 2 is the first instance of dynamic pores controlling release of cargo molecules, Figure 5.1A. This dynamic pore environment may play a major role in future MOF catalysis where molecules of interest need to controllably captured and released in catalytic systems. In addition to dynamic pore control, MOFs must be able to withstand typically harsh chemical environments often used in industrial catalysis. These necessary qualities have been demonstrated by use of a manganese porphyrin-containing framework in the epoxidation of alkenes, Figure 5.1B. In addition, this is the first instance of using molecular oxygen, the most commonly used industrial oxidant, in epoxidation by MOFs. Finally, viable MOF catalysis benefits from inherent advantages of using MOFs, namely the improvement of catalytic efficiency through varying the density of active sites and ability to introduce new catalytically active sites in frameworks. In Chapter 4, the use of the MTV principle and postsynthetic modification in frameworks show that we have the ability to enhance catalytic activity and selectivity while limiting the amount of catalyst needed, Figure 5.1C.
One area of interest for our group is utilizing a MOF to immobilize a heterogeneous catalyst, specifically for use in NMR signal amplification by reversible exchange (NMR-SABRE). Currently, SABRE is performed using a homogeneous iridium N-Heterocyclic Carbene (NHC) catalyst yielding signal enhancements up to 8100 fold. [153] The SABRE catalyst achieves this enhancement factor by transferring the polarization from para-hydrogen to a molecule of interest without any chemical modification, unlike para-hydrogen induced polarization. [154, 155] If we were
able to successfully immobilize a SABRE catalyst in a water stable MOF, this heterogeneous system would be a significant advancement in the hyperpolarization field and a step forward for catalytically active MOFs.

While there are several approaches to immobilizing catalysts inside MOFs, including cationic exchange [156], host-guest “locking” [157] and incorporating the catalyst as the organic linker [158], we decided to use a well known PSM method, formation of an imine bond, [36] to attach a modified version of the Crabtree catalyst [159] to IRMOF-74-III.

Our method is described in the Figure 5.2, but despite varying our PSM conditions, we were unable to preserve crystallinity after PSM of IRMOF-74-III. This may have been due to the fact that we were attempting to attach an active hydrogenation catalyst to our MOF. In addition, attempted hydrogenation of propylene using the amorphous material after PSM showed no catalytic activity. Catalyst and pore size may have also played a part in the unsuccessful PSM. While we specifically selected IRMOF-74-III for its large pore size and had a relatively small counter-ion with our Crabtree catalyst, perhaps the solvated catalyst was too large to properly diffuse into the pore and rearrange itself and attach to the framework. To remedy this, we would need to expand our linkers to create a larger pore cavity, as has been seen in the IRMOF-74 system, or select a different MOF that has a larger pore cavity such as MIL-101. [52]

Though we were ultimately unsuccessful in attaching a SABRE catalyst to a MOF for heterogeneous hyperpolarization, there are still several avenues that could be explored for a successful heterogeneous SABRE catalyst. One has already been discussed above and would involve expanding the MOF pore to accomodate the large homogeneous catalyst but this still involves attempting to attach an active catalyst via chemical reaction. Two other methods that may be more viable are the host-guest “locking” and cationic exchange demonstrated in other MOF systems for incorporation of homogeneous catalysts. The advantages of both are
that a homogeneous catalyst can be introduced without requiring a covalent bond to be formed between catalyst and framework. This should mitigate the loss of crystallinity that was seen in our systems and still gives us room to play with the steric effects from side groups on the homogeneous catalyst. While changing these side groups could ultimately effect the catalytic activity observed from our SABRE catalyst, this is another factor to take into consideration while attempting to immobilize a homogeneous catalyst.

Another possibility for heterogeneous SABRE catalysis is to use the organic linker or MOF SBU itself as a catalyst. Recently iridium catalysts have been introduced in zirconium based MOFs [160, 161] but there has yet to be a MOF where the SBU itself it made from iridium clusters. While this may be unfeasible
due to the high activity of iridium centers, a MOF with an iridium based SBU would be ideal for this type of SABRE catalysis. In addition to using MOFs, a metal-organic polyhedra (MOP) [162] could be used with catalyst linkers as seen in Figure 5.3. While this is a feasible way to incorporate NHC iridium catalysts into a crystalline structure, the catalyst centers would all be pointing into the middle of the pore, possibly decreasing catalytic activity due to pore space constraints. Since the size of Ir complexes do play an appreciable role in SABRE activity, the idea of cation exchange to incorporate catalysts into a MOF is especially interesting. While the size of both the MOF and catalyst would have to be tuned for proper catalyst encapsulation into the superstructure, this could be accomplished either by modifying the side groups on the catalyst or modifying the linker length to accommodate a catalyst of specific size.
5.2 The Effect of Topologically Distinct MOFs on Alkene Epoxidation Yield

While MOF catalysis is still in its infancy, there are many future directions available for new research. One of the most intriguing aspects of MOFs is the wide array of topologies available for catalytically active MOFs. Changes in topology can effect a wide range of properties including pore size, pore aperture, density of functionalities and environment of the functionality. While these parameters have been explored in terms of gas separation and storage, [163] there is a scarcity of studies involving the effect of topology in heterogeneous MOF catalysis. Nature often finely tunes the environment of catalytic sites in enzymes using many of the same parameters highlighted above. Therefore, by using the same catalyst in different topologic environments, we can explore the possibility of catalytic optimization via different catalytic activities as well as different product selectivity for different topologies, which will likely deviate from homogeneous systems.

We have already begun preliminary work on catalytic activity optimization via change in topology through the epoxidation of alkenes in MOF-545-Mn, MOF-525-Mn and PCN-224-Mn, where the effect of different topologies is being investigated in alkene epoxidation. This work builds upon the work in Chapter 4 on the epoxidation of alkenes using a manganese porphyrin based MOF, MOF-525-Mn [Zr$_6$O$_4$(OH)$_4$(MgC$_{48}$H$_{24}$O$_8$N$_4$Cl)$_3$]. While this project looks simply at optimization of catalytic yield as a function of topology, different frameworks can also influence selectivity of products. This has initially been seen in the oligomerization of ethylene based on the change in pore apertures in MOF-74 analogues. [58] The selectivity of linear versus branched alkene chains could be influenced largely by the topology of the heterogeneous catalysis. These two projects are both extensions of the idea that framework topology research can influence the catalytic optimization for industrially relevant reactions and are interesting new directions.
forward for the research I have completed during my time at UCLA.

The work on alkene epoxidation presented in Chapter 4 will lead to the investigation of the epoxidation of longer alkene chains. With larger substrates, there may be difficulty accessing all active sites in MOF-525-Mn due to its cubic topology and small pore aperture (19 Å diameter, space group Pm3m). An excellent alternative would use the hexagonal topology of MOF-545 that has a large one-dimensional pore (35 Å diameter, space group P6/mmm) or to use PCN-224 which has channel type pores (19 Å diameter, space group Im-3m). We expect different catalytic activity from the three frameworks simply based on steric concerns and size exclusion of longer alkene chains from the MOF-525 structure. We would expect that as alkene chain length increases, the activity of the MOF-545 and PCN-224 structures would increase relative to the MOF-525 structure due to greater number of accessible catalytically active sites. To test this, we can set up a system such as the one seen in Figure 5.4, where we monitor catalytic conversion over time via GC-MS or NMR.

Current work shows that there are some differences in catalytic activity be-
### Reactant Product MOF-525-Mn MOF-545-Mn

<table>
<thead>
<tr>
<th>Reactant</th>
<th>Product</th>
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<th>MOF-545-Mn Yield (%)</th>
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<td>54.9</td>
</tr>
</tbody>
</table>

Figure 5.5: Epoxidation of alkenes using molecular oxygen in MOF-525-Mn and MOF-545-Mn.

Between the two frameworks but more work needs to be done optimizing the MOF-545-Mn reaction parameters. The yields shown in Figure 5.5 have been optimized for MOF-525-Mn; however, only preliminary data at room temperature has been collected for the MOF-545-Mn reactions. We are currently working on optimizing the MOF-545-Mn reactions as well as synthesizing and testing the activity of PCN-224. Initial attempts at synthesizing PCN-224 indicates that the framework cannot be made by starting with a metallated TCPP link. Instead an unmetallated TCPP linker must be used in the initial framework synthesis, then the framework post-synthetically modified to include the Mn metal center on the porphyrin link.
Figure 5.6: PXRD of simulated and activated A) MOF-525-Mn B) PCN-224-Mn and C) MOF-545-Mn.

We have confirmed our porphyrin MOF structures by PXRD as seen in Figure 5.6. Surface area and XPS measurements still need to be taken for MOF-545-Mn and PCN-224-Mn.

We expect MOF-545-Mn and PCN-224 to show enhanced catalytic yield as alkene size increases due to the larger space in the pore cavity of MOF-545-Mn. There may be some tradeoff in this increased free space as the isobutyraldehyde co-catalyst also needs to interact at the active site of the manganese porphyrin and there may be some optimal density of co-catalyst to active site that needs to be achieved. Discussion of optimal density of catalyst to co-catalyst within a given pore volume is beyond the scope of this thesis but these are simple experiments that can be quickly carried out on these systems to find optimal catalytic yield for epoxidations of various alkenes.
5.3 Bimetallic Porphyrin-based MOFs for Propylene Epoxidation

In addition to small alkene epoxidation using known structures, preliminary work on bimetallic MOF systems for propylene epoxidation was explored through collaboration with the Senkan group at UCLA. Our idea was to insert Cu and Ru porphyrin sites into the MOF-525 framework as the Senkan group has observed propylene epoxidation from mixing Cu and Ru nanoparticles. [164] We tried two approaches for incorporation of two metalated porphyrins. First, mixing premetalated Cu porphyrin linkers in specific ratios (Cu to unmetalated ratios of 90:10, 80:20, 60:40, and 50:50) with unmetalated linkers, synthesizing the framework, and then inserting the Ru into the unmetalated porphyrins. Second, using premetalated Cu and Ru porphyrins and then mixing the links in the initial synthesis (using the same ratios as the previous method). We observed crystalline MOF-525 with incorporation of both metals into the final structures using the second method, but observed no appreciable conversion of propylene into propylene oxide when our MOF samples were used in the Senkan group catalytic set-up. This may have been due to several factors including the two catalytic centers not being in close proximity to each other, a lack of appropriate reaction conditions for this new system, and an inability to saturate the MOF with NaCl which helped achieve higher yields in different systems. [165] Additionally, while the oxidation states of the Cu and Ru linkers (II and IV respectively) should match those in the system used by the Senkan group, these may have been altered by solvated ions in our MOF system. We did not run XPS analysis to confirm the oxidation states in our system, but that should be a major component for this project going forward to confirm the identity of each metal center within the MOF.

Another major factor in this epoxidation system is whether a cocatalyst is necessary for epoxide formation. In our other epoxidation project, the sacrificial
Figure 5.7: Schematic representation of 2D layered Cu-porphyrin MOF joined by axial Ru-porphyrins to make a bimetallic 3D MOF.

aldehyde isobutyraldehyde is a integral part of the catalytic system. For this system, and more specifically for propylene epoxidation, it remains to be seen if a cocatalyst is necessary to observe propylene oxide formation. The Senkan group observed that the introduction of NaCl into the space between nanoparticles facilitates higher reaction yield of propylene oxide but we are unsure whether the porous structure of a MOF needs the same NaCl introduction and whether a cocatalyst will remain inside the pores during catalysis. We attempted to saturate the pores with NaCl by soaking our bimetallic MOF with an aqueous NaCl solution but no propylene oxide was observed after attempted catalysis. While I believe the NaCl is necessary for higher propylene oxide formation, the distance between Cu and Ru centers seems to be of greater importance for this reaction and should be investigated first before varying cocatalysts.

Finally, it may be prudent to validate this bimetallic system in a gas phase epoxidation using a different alkene before attempting propylene epoxidation. At this point, we are unable to assess whether this bimetallic MOF system is even catalytically active due to the lack of observed epoxide products from our pro-
plyene reaction but proving that our system can form epoxides of longer chain alkenes, such as 1-hexene, 1-pentene or butene, at elevated temperatures would allow for both proof of concept and optimization of reaction parameters. This validation and optimization of our system is vital before pursuing the epoxidation of the more thermodynamically stable propylene.

One of the major problems working with bimetallic systems in MOF-525 is the idea of mixing different linkers that have similar symmetry and connectivity. The ability to use different linkers in the same framework is one of the major advantages of MOFs but specific placement of distinct linkers, relative to each other, requires more thought. A simple way of positioning each catalytic site in a well defined orientation is to first build a 2D framework using one type of linker, then the connect separate layers using a second linker with different connectivity. Similar systems have been seen before in several MOF systems [166] but have yet to be applied to bimetallic porphyrin systems. Our general overview can be seen in Figure 5.7, where we synthesize a 2D layer of Cu-porphyrins and then add Ru-porphyrins to the axial sites on the copper paddlewheel SBU to form a 3D MOF. The different connectivity of the two porphyrins allows for control of the distance and orientation between the two catalytic centers and should give greater pore control within the framework.

5.4 Effect of MOF Topology on Oligomerization of Propylene

While density of catalytic sites can play a role in catalytic yield, it can also influence selectivity of products from catalytic reactions. One recent example of this selectivity was seen in the degree of dimer branching after the oligomerization of propylene over a nickel MOF catalyst. [58] While this system used the same topology in each case, the pore size was varied and this led to different degrees of
branching in each case. An alternative method is to totally change the topology of the MOF catalyst from one dimensional pores to three dimensional pores of various apertures and environments and to study the selectivity of product formation resulting from each topology. [167]

A general outline of our experimental set-up can be seen in Figure 5.8 where we synthesize three MOFs of different topology, test their catalytic activity and determine their selectivity using NMR and GC-MS. Nickel analogues of MOF-253, [168] MOF-199 [169] and MOF-74 [58] were selected because they are well known MOFs and can be reliably reproduced in high yield and quantity. In addition, each of these three MOFs has known analogues where pore diameter can be varied by changing the length of the organic link while still maintaining the same topology and catalytically active site of the overall MOF structure.

With these three structures we also can examine the differences between catalysts that are part of the metal SBU of the MOF and catalysts that have been added to the framework through PSM. [36] MOF-253 has a square planar nickel
site [168] on the organic link while the active sites in MOFs 199 [170] and 74 [171] are part of the backbone of the MOF. This illustrates yet another instance where catalytic site density can be varied amongst different MOFs, especially those using PSM to add a catalytically active site to a framework.

The idea of bimetallic MOFs for propylene epoxidation and varying topology for oligomerization are just a two examples of future projects but the field as a whole has many avenues for future research. Some important topics still unsolved are the development of industrially viable MOF catalysts, [172] biomimetic catalysts [134] and the mixing of frameworks and catalysts for cascade and tandem reactions . [173, 174] While MOFs hold several advantages over commonly used industrial supports, zeolites and silicas, there has yet to be an industrially viable MOF mainly due to their lack of chemical and thermal stability. The development of zirconium frameworks [51] has led to new testing in harsher chemical environments and development of new catalysts with Zr backbones, with industrially viability the key focus, should be given high priority in the MOF catalyst community.

In addition to industrially viable MOFs, the development of biomimetic frameworks should also be of great interest for future study. While MOF catalysis is still relatively new, the development of highly selective catalysts is of key importance for future MOF chemists. With the ability to position catalysts in precise locations inside frameworks, inspiration should be taken for enzymes to mimic active sites that are highly selective and show molecular preference during reactions. MOFs have already demonstrated selective gas uptake [64] and there is no reason to believe that similar molecular selectivity cannot be shown for heterogeneous catalysis. With the ability to alter the chemical environment, steric, and size of the MOF pore; the ability to synthesize complex structures with lock and key or induced fit specificity [175] is an exciting idea for future MOF catalysts.
5.5 Cascade Catalysis in MOF Systems

Finally, one major topic that should be developed is the idea of tandem or cascade catalysis [176] utilizing MOFs. This has already been seen in homogeneous catalysis [177] but the development of solid-state catalysts should be of great importance due to their inherent advantages such as lack of necessary filtration and easy recyclability. [178]

The biggest challenge when discussing heterogeneous cascade catalysis is whether to have multiple catalysts in one framework or to sequester each catalyst into its own framework and then physically mix the frameworks. The development of heterogeneity in MOFs is a recent advancement [179] and there are still questions to be answered about the simplest way to obtain heterogeneity while maintaining crystalline order. In my experience, the easiest way to incorporate multiple catalysts is to keep them sequestered in their own framework. This makes the development of each catalyst simpler by having to crystallize out a structure with only one catalyst and keeps each framework relatively uncomplicated. While a single MOF containing multiple catalysts is a very interesting idea, the crystallization step becomes increasingly difficult as the pore environment becomes more complex. In addition, attempting to synthesize a MOF with catalytically active species is difficult due to the suppression of catalytic active site during solvothermal synthesis of the framework. This is one reason why PSM of frameworks has been such an important development for MOF catalysis. I have found it easier to add a catalytically active site after the MOF has already been synthesized. While PSM adds an additional step in the overall MOF synthesis, it allows MOF chemists to ensure that framework integrity is maintained throughout all necessary synthetic steps. One difficulty with PSM is the reaction yield of the PSM step, as it is often not quantitative as was demonstrated in Chapter 4 with our MTV MOF series. As pore space increases the yield of PSM also increases but
this would be something to study in further depth if PSM were to be used on an industrial scale. While it is often not necessary to have all available functionalities modified during the PSM step, inhomogeneity within a MOF structure could cause difficulties when discussing pore and, by extension, catalyst environment. While PSM is a very useful tool for inserting new functionalities and catalysts into a MOF, more work needs to be done to determine the relationship between pore environment and the efficiency of PSM reactions.

While the field of catalytically active MOFs is growing with each passing year, there is still much work to be done, specifically in testing the influence of MOF topology on catalytic yield and selectivity of different reactions. The exploration of topological effects on the yield in alkene epoxidation and the selectivity of dimer branching in propylene oligomerization could open up new avenues and tools for catalytic optimization in MOFs. In addition, the development of specific placement of multiple catalysts, in our case Cu and Ru porphyrins, within a single framework should be of highest priority as it would aid the development of an industrially viable MOF for propylene epoxidation. Selection of MOF topology to optimize specific catalytic reactions could soon be the next tool to join the MTV principle and PSM as commonly used techniques in applying the flexibility of reticular frameworks to industrially relevant projects.
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