Crystals versus Electrochromic Films: Pathway-Dependent Coordination Networks

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Supporting Information Placeholder

ABSTRACT: We demonstrate the formation of metal-organic frameworks (MOFs) and thin-film coatings that have entirely different compositions and structures, distinct by cation exchange, despite using the same set of starting materials. The reaction of an iron polypyridyl complex with a copper salt by diffusion of one solution into another resulted in iron-to-copper exchange, concurrent ligand rearrangement and the formation of MOFs. This observation shows that polypyridyl complexes can be used as expendable precursors for the growth of MOFs. In contrast, alternative depositions of the iron polypyridyl complex with a copper salt by automated spin-coating on conductive metal-oxides resulted in the formation of electrochromic coatings, and the structure and redox properties of the iron complex were retained. The possibility to form such different networks by “in solution” versus “on surface” coordination chemistry from the same set of molecular building blocks broadens the synthetic space to design functional materials.

Coordination chemistry has been used to control the shape, size, and topology of supramolecular structures and to limit the possibilities to produce mixtures of multiple sets of structures. Early examples of self-assembled architectures are compounds made from cryptands and crown ethers and were studied by Pederson, Lehn, and Cram in the 1960s. In the following years, highly complex helicates were introduced and are composed of oligobipyridine strands coordinated to copper cations. The coordination chemistry of carboxylic acids and late transition metals has been extensively used for the formation of metal-organic frameworks (MOFs). This strong metal-ligand interaction resulted in highly robust and porous materials. Metal-pyridine coordination chemistry has also been used for the generation of self-assembled structures in solution and on surface. Although the interaction between metals and pyridine is weaker than with carboxylic acids, stable and functional materials can be isolated. Examples include cages, MOFs, and thin films. The control of material properties by external stimuli (e.g., light, voltage) has resulted in diverse functionalities, including memory elements, biomedical applications, and electrochromism.

Structures and functionalities of supramolecular assemblies formed in solution or on surfaces are difficult to predict. Assemblies formed from the same starting materials can have the same or different molecular arrangements and functions. Interfacial chemistry has been used to generate assemblies that cannot be formed in solution otherwise. Monolayer chemistry has been used control chirality and morphology of crystals, and network interpenetration of MOFs. In general, the development of defined supramolecular structures with desirable properties occurs with retention of the structural integrity of the molecular building blocks, but the assemblies can have different molecular arrangements and appearances. Synthetic routes that involve changes in the molecular structures prior to the assembly of the components are rare and examples of pathway dependence on such processes are unknown to the best of our knowledge.

We show here the formation of two assemblies having strikingly different molecular compositions, although the same starting materials were used (Schemes 1, 2). Reacting a structurally well-defined iron polypyridyl complex with copper nitrate by diffusion of one solution into another resulted in an exchange of the metal cations followed by the formation of MOFs. The initial disassembly of the iron complex is followed by formation of a coordination polymer consisting of the polypyridyl ligand and the copper salt. In contrast, alternative spin-coating of the same iron polypyridyl complex with copper nitrate on fluorinated tin-oxide (FTO) resulted in electrochromic coatings.

![Scheme 1. Iron polypyridyl complex](image)
The electrochromic activity originates from the iron polypyridyl complex. The free pyridine moieties of the polypyridyl ligands are coordinated to the copper cations forming a dense network of iron complexes. This network stabilizes the iron complexes as cation exchange was not observed, even not after prolonged exposure to a solution containing an excess of copper salt.

Crystals were obtained by slow diffusion of solutions into another at room temperature. We used a thin tube (\( \phi = 5 \) mm) containing three layers with the top and bottom layers with solutions of complex 1 or \( \text{Cu(NO}_3\text{)}_2 \). The layer in the center was a co-solvent. During the reaction the color of the solution changed from purple to colorless. These coordination organic networks (SolCONs) were isolated after 20 days by centrifugation and washed with ACN. Two different solvent combinations were used and resulted in the same crystallographic structures and morphologies, but of slightly different dimensions.

**Scheme 2.** Formation of divergent coordination networks in solution (SolCON) versus on surface (SurCON)a

a Water molecules and anions are omitted for clarity. The diffusion experiment shown is for SolCON-A. For SolCON-B, the solutions with iron complex 1 and the copper salt are the top and bottom layers, respectively.

SolCON-A was formed by the reaction of complex 1 (DCM:MeOH, 1:1 v/v) with \( \text{Cu(NO}_3\text{)}_2 \cdot 3\text{H}_2\text{O} \) in ACN in a molar ratio of 1:2. DCM:MeOH:ACN (0.5:0.5:3 v/v/v) was used as a co-solvent in the center. SolCON-B was formed by using ACN as solvent for complex 1 and DMF for \( \text{Cu(NO}_3\text{)}_2 \cdot 3\text{H}_2\text{O} \). ACN:DMF (1:1 v/v) was used as a co-solvent in the center (Figure 1). Scanning electron microscopy (SEM) analysis revealed the formation of crystals that have the appearance of a parallelepiped (Figure 1, Chart 1). Although these crystals were uniformly-shaped, their dimensions vary. For SolCON-A the size distribution was 2.1 ± 0.9 µm (~50 crystals) and for SolCON-B sizes were between 1-4 µm (~50 crystals). In addition, larger crystals of 38-70 µm (~10 crystals) were also observed for SolCON-B. The different solvent combinations did not affect the overall crystal morphology as is sometimes observed.35

The single-crystal X-ray analysis of SolCON-B showed the formation of metal-organic framework based on copper cations and the ligand of complex 1 (Figure 1, Chart 2). The formation of the framework involved ligand transfer from complex 1 to the copper salt. The 3D framework is formed by mono- and bidentate binding of copper centers to the pyridine moieties of the ligand. The ligands are coordinated in square-pyramidal fashion around the copper centers. Two nitrate counter anions are present in the asymmetric unit and hydrogen bonding is observed between the oxygen atom of NO\(_3\) and hydrogen atoms of the polypyridyl ligand. The nanobeam electron diffraction (NBED) patterns of SolCON-A consist of sharp spots that match with the according zone-axis patterns calculated from the refined structure of SolCON-B, demonstrating that these MOFs have very similar crystallographic structures (Figure 1, Chart 3, Figure S1).

Reacting a solution of [Fe(bpy)\(_3\)](PF\(_6\)) in ACN with excess of Cu(NO\(_3\))\(_2\) (40 equiv.) resulted within 60 h in the disappearance of the typical red color associated with the iron complex.36 This observation indicates that the vinylpyridyl moieties of complex 1 are not essential for the cation exchange. To demonstrate the differences between bulk crystallization versus on surface chemistry, a thin film (SurCON) was prepared by Layer-by-Layer deposition of solutions containing complex 1 and Cu(NO\(_3\))\(_2\) (Figure 2). Using this approach, complex 1 retains its structure and electrochromic properties. SurCON was assembled on fluorine doped tin-oxide (FTO) on glass (2 cm × 2 cm) using an automated spin coating, and solutions of Cu(NO\(_3\))\(_2\)·3H\(_2\)O (4.0 mM, ACN) and complex 1 (0.6 mM, DCM:MeOH, 1:1 v/v). This deposition sequence was repeated 18 times to obtain SurCON. The SurCON was coated with a thin layer of platinum, and was milled using a focused ion beam (FIB) (Figures 2, Chart 1). The transmission electron microscopy (TEM) images of a cross section of SurCON shows a homogeneous film having a thickness of ~178 nm. EDS mapping clearly indicates the uniform distribution of both the iron and copper cations.

This deposition sequence was repeated 18 times to obtain SurCON. UV-vis spectra recorded for different numbers of deposition cycles showed the broad metal-to-ligand charge transfer (MLCT) bands related to complex 1 at \( \lambda_{\text{max}} = 458 \) nm and \( \lambda_{\text{max}} = 596 \) nm (Figures 2, Chart 2A). An intense π-π* transition band of the ligand was also present at \( \lambda_{\text{max}} = 332 \) nm. Plotting the absorption intensity (\( \lambda_{\text{max}} = 596 \) nm) versus the number of deposition cycles indicated linear growth with retention of complex 1.
X-ray photoelectron spectroscopy (XPS) data of the SurCON confirmed the presence of iron complex 1, and copper cations as cross-linkers (Figures 2, Chart 2B). Two characteristic bands for Fe$^{2+}$ are present at 708 eV (2p$_{1/2}$) and 720 eV (2p$_{3/2}$). The ratio N$_{pyr}$/Fe of 11.9 is in excellent agreement with the expected ratio for complex 1 (N$_{pyr}$/Fe = 12). The bands for Cu$^{2+}$ are observed at 935 eV (2p$_{3/2}$) and 955 eV (2p$_{1/2}$) and the satellite bands at 941-945 eV and 962-965 eV. The observed ratio of Cu/Fe (~2.7) indicates the formation of a fully network (Cu/Fe = 3), where the copper centers are bound by two pyridine groups.

Electrochemical measurements unambiguously confirmed the presence of the electrochromic complex 1 (Figure 2, Chart 3). Cyclic voltammograms (CVs) showed reversible one-electron redox processes as expected for the Fe$^{2+/3+}$ couple with a half-wave potential ($E_{1/2}$) of 1.1 V and a peak-to-peak separation of 310 mV at a scan rate of 100 mV/s. The color of the SurCON changed from gray (at 0.4 V) to colorless (at 1.8V) upon oxidation of Fe$^{2+}$ to Fe$^{3+}$. This reversible process could be monitored using spectroelectrochemical (SEC) measurements (Figure 5a). The changes of the oxidation states were accompanied by variations in the absorption intensities of the MLCT bands. The time required to reach 90% of the maximum transmittance ($\Delta T \sim 40\%$) was ~2.1 s. The switching stability was indicated by 250 redox cycles with >80% retention of the initial $\Delta T$. The coloration efficiency (CEs) was 1.48 cm$^2$/C. The SurCON is densely packed as indicated by the molecular density of ~1.1 x 10$^{-6}$ molecule/cm$^3$ for a charge density (Q) of 1.77 mC/cm$^2$. Exponential and linear dependencies of the anodic and cathodic peak currents on the scan rate and square root of these scan rates, respectively, were observed indicating a redox process controlled by diffusion. The calculated diffusion coefficients ($D_h \sim 3.37 \times 10^{-9}$ cm$^2$ s$^{-1}$ (oxidation) and $\sim 3.64 \times 10^{-9}$ cm$^2$ s$^{-1}$ (reduction) are similar and derived from the Randles-Sevcik equation. The SurCON is remarkably stable as no cation exchange was observable by UV-vis spectroscopy and electrochemical measurements. Immersion of a SurCON is an ACN solution containing Cu(NO$_2$)$_2$·3H$_2$O (4.0 mM, ACN) for three days did not result in ligand transfer (Figure 5c). Clearly, the formation of a network containing both the copper salt and complex 1 enhances its stability.

In conclusion, the reactions demonstrated here are two examples of coordination-based polymerization processes: metal-ligand exchange followed by crystallization and on-surface deposition. The composition of the assemblies is controlled by the applied method. We showed that iron polypyridyl complexes can be used as sacrificial precursors for the formation of MOFs by slow diffusion of solutions. The on-surface polymerization is much faster, this prevents the metal-ligand exchange. However, reacting palladium salts with iron polypyridyl complexes in solution (diffusion or by fast mixing) did not result in metal-ligand exchange, as was observed here.
Figure 2. Characterization and electrochromic properties of SurCON. Chart 1: (A) Scanning transmission electron microscopy (STEM) image showing a cross section of a SurCON sample. (B, C) EDS elemental map showing the distribution of iron and copper metals. Scale bar: 100 nm. 

Chart 2: (A) Ex-situ absorption spectra recorded during the film formation. FTO/glass was used for the baseline (black). Inset: Absorbance intensity of the metal-to-ligand charge transfer (MLCT) band (λ_{max} = 596 nm) versus the number of deposition cycles. (B) X-ray photoelectron spectroscopy (XPS) spectra. 

Chart 3: (A) Photographs of the colored (0.4 V, Fe^{2+}) and bleached (1.8 V, Fe^{3+}) states using an electrolyte solution of 0.1 M TBA PF_6 in ACN. Additional details are shown in Figure S2. (B) Cyclic voltammograms (CVs) recorded at a scan rate of 100 mV/s. (C) Absorption spectra showing the reduced and oxidized states. FTO/glass was used for the baseline (black).

for complex 1 and a copper salt (Figure S4). A similar on-surface polymerization was observed by us and others with palladium salts for the formation of electrochromic coatings.16,17,29,34,35 The use of copper rather than palladium salts to form electrochromic coatings is advantageous due to lower toxicity and cost. Others have reported the formation of related coordination structures based on terpyridine iron complexes and copper salts in solution and on surface.29,39,36,37 No exchange of the iron and copper cations have been reported. The previous finding with palladium chemistry and the abovementioned reports by Constable, Housecroft, Gupta, Mondal, Zharnikov highlight that our example of divergent coordination chemistry is rare and can offer new opportunities in the molecular engineering of functional materials. 

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MOF

Cu for Fe

Cu salt

Cu + Fe

Fe complex

n

FTO

Electrochromic film