Superior Charge Storage and Power Density of a Conducting Polymer-Modified Covalent Organic Framework

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Supporting Information

ABSTRACT: The low conductivity of two-dimensional covalent organic frameworks (2D COFs), and most related coordination polymers, limits their applicability in optoelectronic and electrical energy storage (EES) devices. Although some networks exhibit promising conductivity, these examples generally lack structural versatility, one of the most attractive features of framework materials design. Here we enhance the electrical conductivity of a redox-active 2D COF film by electropolymerizing 3,4-ethylenedioxythiophene (EDOT) within its pores. The resulting poly(3,4-ethylenedioxythiophene) (PEDOT)-infiltrated COF films exhibit dramatically improved electrochemical responses, including quantitative access to their redox-active groups, even for 1 μm-thick COF films that otherwise provide poor electrochemical performance. PEDOT-modified COF films can accommodate high charging rates (10−1600 C) without compromising performance and exhibit both a 10-fold higher current response relative to unmodified films and stable capacitances for at least 10 000 cycles. This work represents the first time that electroactive COFs or crystalline framework materials have shown volumetric energy and power densities comparable with other porous carbon-based electrodes, thereby demonstrating the promise of redox-active COFs for EES devices.

INTRODUCTION

Pseudocapacitors can combine the high energy density of batteries and superior power densities of double-layer capacitors by storing electricity through both the non-faradaic formation of an electrochemical double layer (EDL) and reversible faradaic processes of surface bound/immobilized species. Nanoporous electrodes, often carbon-based materials, feature high specific surface areas that maximize electrochemical double-layer formation. Although redox-active groups have been covalently bonded or adsorbed to these electrodes, their performance is often compromised by charge transfer and counterion transport limitations. In addition, their poorly defined structures complicate characterization and rational improvement. Two-dimensional covalent organic frameworks (2D COFs) address these limitations by predictably and deliberately organizing redox-active groups into insoluble, high-surface area polymer networks with uniform micropores. However, the modest conductivity of existing 2D COFs has limited devices to thin films of the active material (50–250 nm) grown on Au or carbon nanotube electrodes that only operate at slow charge/discharge rates, limiting high power performance. Here we address this challenge by electropolymerizing EDOT into the pores of redox-active 2D COF films. The resulting poly(3,4-ethylenedioxythiophene) (PEDOT)-modified COF films (Figure 1) exhibit quantitative electrochemical accessibility of their redox-active groups and enable the use of at least 1 μm thick films that can sustain fast charging rates (up to 1600 C) without compromising performance. This improved performance relative to as-synthesized COF films corresponds to a 30-fold increase in volumetric energy density and a 12-fold increase in volumetric power density, two important metrics for evaluating thin film capacitors.

The electropolymerization of PEDOT within a nanoporous COF template represents a means to organize the conductive polymer at the nanometer length scale, which is an increasingly important capability for electrochromic, drug delivery, electrocatalytic, and EES devices, among other applications. Templated electropolymerizations form conducting polymers, such as polypyrrole, polythiophene, or polyaniline, as nanowire arrays, which respond rapidly in electrochemical devices largely because of short counterion diffusion lengths. In these examples, the porous templates are removed. Here we observe a synergistic effect in which the pore-confined PEDOT effectively wires the redox-active groups of the COF to the electrode, enabling the use of thicker films and dramatically faster charging rates.

RESULTS AND DISCUSSION

PEDOT was electropolymerized within a 2,6-diaminoanthraquinone-2,4,6-triformylphloroglucinol (DAAQ-TFP) thin-film working electrode via cyclic voltammetry in a 100 mM CH3CN solution of EDOT containing 100 mM (n-Bu)4NClO4 (TBAP)
as the supporting electrolyte. DAAQ−TFP thin films were prepared by adding a DMF solution of TFP over 1 h to a DMF solution of DAAQ containing a gold substrate.20 Similar to our previous report, we isolate polycrystalline thin films whose thickness was controlled by varying the initial monomer concentration. For example, an initial DAAQ concentration ([DAAQ]0) of 22 mM yielded DAAQ−TFP thin films of 810 ± 224 nm thickness, as determined by atomic force microscopy (AFM, Figures S1−S4). The chemical composition of the COF films was assessed using Fourier transform infrared (FT-IR) and X-ray photoelectron spectroscopy (XPS). Their morphology was characterized by scanning electron microscopy (SEM, Figures S5 and S6), and their crystallinity was measured using grazing incidence X-ray diffraction (GIXD, Figures S7−S9). These measurements are consistent with those of thinner films reported previously20,29,30 and indicate the formation of crystalline DAAQ-TFP films whose layered crystallites are preferentially oriented parallel to the substrate (Figures S4−S8). The amount of PEDOT incorporated into the 2D COF film was characterized by two independent measurements: the current passed during each electropolymerization cycle (−0.5−1.0 V vs Ag/AgClO4, 20 mV s−1 scan rate) and the mass deposited onto a COF-modified electrode, as detected using an electrochemical quartz crystal microbalance (EQCM, Figure 2A,B). In the first cycle, 15 μg of PEDOT and electrolyte were deposited per cm² of the working electrode’s geometric area, which increased to 30 μg cm⁻² for the second and subsequent cycles. Because of viscoelastic losses over the course of the experiment, the Sauerbrey equation overestimates these masses (see Supporting Information for further discussion, Figures S10 and S11). During the reductive sweep of each electropolymerization cycle, there is a mass loss corresponding to removal of perchlorate counterions (dedoping) from the film to maintain electroneutrality.31 The voltammetric response exhibited an increase in current with each scan, which is characteristic of PEDOT electropolymerization (Figure 2B), and the redox processes occurred at voltages consistent with those expected for EDOT oxidation.32−35 FT-IR spectra of the films, acquired before and after the electropolymerization, suggest the formation of PEDOT and that the DAAQ−TFP’s chemical linkages are retained. After two electropolymerization cycles, the FT-IR spectrum of the film was almost identical to that of the as-synthesized DAAQ−TFP film. Both spectra exhibited peaks at 1250 cm⁻¹, 1560 cm⁻¹, and 1615 cm⁻¹, corresponding to intact β-ketoenamine C=N, C=C, and C=O stretches, respectively. After nine electropolymerization cycles, spectral features resembling bulk PEDOT emerge, though absorbances associated with the β-ketoenamine remain, albeit with lower relative intensities (Figure 2C). Grazing incidence X-ray diffraction (GIXD) experiments, performed at the Cornell High Energy Synchrotron Source (CHESS), indicate that the as-synthesized DAAQ−TFP films are
crystalline, with a peak at 0.23 Å\(^{-1}\) that corresponds to the (100) reflection of the hexagonal lattice. This peak was also observed in films that were subjected to two electropolymerization cycles, indicating that the periodicity of the DAAQ–TFP thin film is retained under the electropolymerization conditions. After nine cycles, this peak is not observed, which is not unexpected after depositing amorphous PEDOT within the ordered DAAQ–TFP COF (Figures S7–S9). For example, Russell et al. reported that the scattered intensity diminishes when a porous host is filled with a polymer of similar electron density.\(^{39}\) The surface area of the films before and after PEDOT polymerization, as measured using Kr adsorption (Figure S12), was also consistent with the deposition of the polymer into the pores of the DAAQ–TFP COF. Unmodified DAAQ–TFP films exhibited Brunauer–Emmett–Teller surface areas (\(S_{\text{BET}}\)) of 73 cm\(^2\) per cm\(^3\) of substrate. This value decreased to 40 cm\(^2\) per cm\(^3\) after two electropolymerization cycles, and films subjected to nine electropolymerization cycles appeared nearly nonporous (\(S_{\text{BET}} = 6\) cm\(^2\) cm\(^{-3}\)). An X-ray photoelectron spectroscopy (XPS) depth profile of the elemental composition of a PEDOT-modified DAAQ–TFP film subjected to nine electropolymerization cycles further indicates the presence of PEDOT throughout the 2D COF film. XPS spectra were recorded after a series of exposures to an Ar ion beam that etches both the PEDOT and the COF. Prior to etching, an intense S\(_{2p}\) signal (164.9 eV) is observed, along with no signal above baseline in the N\(_{1s}\) region (399.2 eV), which we attribute to a thin PEDOT overgrowth layer on top of the COF film. After the first etching cycle, N\(_{1s}\) and S\(_{2p}\) signals are both observed in five subsequent consecutive spectra (Figure 2D), and, after prolonged etching, the entire film is removed. These combined observations indicate that PEDOT electropolymerization occurs in the pores of the COF film and that the conductive polymer effectively infiltrates the COF structure.

The PEDOT-modified DAAQ–TFP composite films exhibited dramatically enhanced current responses in cyclic voltammetry (CV) experiments when compared to an unmodified DAAQ–TFP film (0.5 M H\(_2\)SO\(_4\) 20 mV s\(^{-1}\) scan rate, Figure 3A). While both CVs exhibit reversible electrochemistry consistent with electron transfer between the anthraquinones and the working electrode, the PEDOT-modified DAAQ–TFP films exhibited more than an order of magnitude increased current. These responses correspond to a faradaic charge storage of only 0.230 mC, corresponding to only 3% of the available anthraquinones for a representative unmodified, 1 μm-thick DAAQ–TFP film (see Supporting Information for a sample calculation of electrochemically addressed anthraquinones, Figure S13). This value increased to 9.3 mC after the same DAAQ–TFP sample was modified with PEDOT, corresponding to a 40-fold increase in accessible charge. In contrast to the unmodified COF films, PEDOT-modified films retain their well-defined redox responses and charge densities at scan rates up to 500 mV s\(^{-1}\) (Figure 3B). In contrast, the already small percentage of accessible anthraquinones (3%) measured at 20 mV s\(^{-1}\) decreased to less than 1% at sweep rates above 100 mV s\(^{-1}\) (Figure 3B).

Having established the dramatic enhancement in performance associated with modifying COF thin films with PEDOT, we determined the optimal electropolymerization conditions (nine cycles, 20 mV s\(^{-1}\); see Supporting Information for details Figures S14–S16) for maximizing the charge stored and accessed at high scan rates (100–500 mV s\(^{-1}\)) while maintaining electrolyte access to the framework. While additional electropolymerization cycles increased the capacitance associated with additional PEDOT coverage, the faradaic contribution remained constant at a value corresponding to a quantitative access to the quinone sites (Figure S14). These experiments indicate that the electropolymerization of PEDOT in a redox-active COF film provides dramatically enhanced charge storage capacitance and rate. These effects most likely arise from intimately mixing PEDOT in the COF for improved conductivity of the PEDOT-modified DAAQ–TFP film, which was assessed using electrochemical impedance spectroscopy (EIS, Figures S17 and S18, Table S1). We used a two constant-
phase-element (2CPE) model, in series with the solution resistance \((R_s)\), to model the EIS data. The \(R_s\) was more than an order of magnitude lower for the PEDOT-modified DAAQ–TFP film (\(R_s = 20 \Omega\)) than the unmodified DAAQ–TFP film (\(R_s = 430 \Omega\)), demonstrating an increased conductivity when PEDOT is incorporated into the films.

Given the superior electrochemical performance of the PEDOT-modified DAAQ–TFP films, their capacitance was evaluated through galvanostatic charge–discharge experiments performed at various charging rates \((C)\), where \(nC\) corresponds to charging or discharging of the film over \(1/n\) hours (Figures S19–S24, Table S2). PEDOT-modified DAAQ–TFP electrodes consistently showed higher capacitances than those lacking PEDOT and retained more than 80% of their capacitance when charged or discharged at 10 and 100 C, corresponding to charge times of 360 and 36 s, respectively. The PEDOT-modified DAAQ–TFP film even retains 50% of its maximum capacitance \((350 \text{ F cm}^{-2})\) at the extremely high charging rate of 1600 C, corresponding to a charging time of only 2.25 s (Figure 4A). In contrast, the unmodified DAAQ–TFP films show only moderate capacitances at 10 C \((20 \text{ F cm}^{-2})\), which decreased further at higher charge/discharge rates. The PEDOT-modified DAAQ–TFP films also showed outstanding stability over 10 000 charge–discharge cycles (Figure 4B). The capacitance of the film was measured at 10 C for three cycles, then at 100 C for 10 000 cycles, and finally three more cycles at 10 C. No decrease in capacitance was observed under these conditions.

To further probe the stability, we conducted a potentiostatic experiment where we held the composite film at a reducing potential \((-0.3 \text{ V vs Ag/AgCl})\) for 15 h, after which no degradation in the CV response was observed (Figure S25). We examined the possibility for slow internal electron transfer between the PEDOT in its oxidized, conducting form and the reduced anthraquinone moieties by performing a scan rate dependence experiment at slow cycling rates (0.5, 1, 5, 10, 20, 50, 100, 300, 500 mV s\(^{-1}\)). Since the integrated charge for anthraquinone reduction is greater than that of the integrated oxidative wave, we hypothesize that a portion of the reduced anthraquinone moieties transfer electrons and reduce the oxidized PEDOT (effectively a “self-discharge” mechanism). However, the Coulombic efficiency becomes quantitative at scan rates above 10 mV s\(^{-1}\), suggesting that process is not occurring at the operational C rates and is not detrimental to the composite electrode performance (see Figure S26 and accompanying discussion).

To determine the relative contributions of the faradaic and nonfaradaic processes at both fast and slow scan rates, GCDC experiments were performed between 0.35–0.6 V vs Ag/AgCl\(_o\) where the DAAQ moieties are redox-inactive, and compared them to an experiment performed over the full \(-0.3–0.6 \text{ V vs Ag/AgCl}_o\) range (Figure S22, Table S2). This comparison indicates that both the DAAQ moieties and the PEDOT contribute nearly equally to the capacitance at all tested charging rates, demonstrating the synergistic effect of combining the two materials. We also examined the performance of a PEDOT composite of a non-redox-active COF based
on 1,4-diaminobenzene (DAB), DAB–TFP COF, which showed capacitances comparable to PEDOT contribution of the PEDOT-modified DAAQ–TFP composite (Figure S23). Furthermore, Au electrodes modified only with electropolymerized PEDOT exhibited a similar double-layer capacitance as the PEDOT/COF hybrid, but lacked the enhanced charge storage associated with the DAAQ redox couple, as determined from the lack of voltage plateau in the potential/capacity plots (Figures S20 and S21).

As described by Ruoff and co-workers, performance metrics are more appropriately evaluated using a two electrode configuration; therefore, we probed the PEDOT-modified DAAQ–TFP films using a high surface area carbon counter and quasi-reference, where the size and mass of the carbon counter was significantly greater than that of the active film. Under these conditions (20 mV s⁻¹, 0.5 M H₂SO₄), the films exhibit reversible oxidation and reduction waves (Figure 4C) that are associated with pseudocapacitive features in a two-electrode configuration. GCDC profiles exhibited well-defined voltage plateaus consistent with previous experiments (Figure 4D) and capacitances comparable to those observed in a three-electrode system at charging rates up to 800 C. At still faster charging rates, device performance was limited by the counter electrode, not the COF film (Figure S27).

As a proof-of-principle for integrating larger amounts of the COF into devices, we fabricated coin cell devices, which were characterized and used to power a light emitting diode (LED, Figure 4E). Since the quantity of the PEDOT-modified COF obtained from electropolymerization is on the order of micrograms, we turned to insoluble polycrystalline DAAQ–TFP COF powder and a Fe(Clo₄)₂ chemical polymerization of EDOT within the pores to afford bulk PEDOT-modified DAAQ–TFP COF (see Supporting Information for fabrication procedures, Figures S28–S32 and Table S3). The crystallinity and chemical composition of these materials were confirmed using X-ray powder diffraction and IR spectroscopy (Figures S29 and S30). A 4 mg portion of PEDOT-modified DAAQ–TFP COF powder (1:1 PEDOT/COF by mass as active electrode and activated carbon counter electrode) was integrated into coin cells. When two of these cells were connected in series, they successfully powered a green LED for 30 s (see Supplemental Video). Electrochemical performance testing of the cells show that they exhibit well-defined redox waves associated with the reduction and oxidation of the anthraquinone moieties in both the CV and GCDC responses (Figures S31 and S32) with a capacitance of 197 F g⁻¹ based on the active composite electrode or 30 F g⁻¹ when the mass of the active and counter electrodes are considered. Notably, these cells are not optimized, and the gravimetric capacitances are expected to increase if the minimal mass of counter electrode and PEDOT to elicit the same electrochemical performance are identified. However, they demonstrate a means to fabricate working charge storage devices from conducting polymer-modified COFs, even as methods to access thicker films continue to emerge.

### CONCLUSION

Prior to this work, 2D COFs had shown promise for EES devices only when used instead of few-layer sheets or very thin films (~50 nm) because of their limited conductivity. Although some coordination polymers circumvent this issue through elegant linkage chemistries, such approaches limit the scope of accessible frameworks. The facile electropolymerization of EDOT within the pores of comparably thick COF films (~1 μm) enhances the framework’s conductivity to provide complete electrochemical addressability of redox-active groups within the COF, even at very high scan rates. The complete electrochemical accessibility and fast charging rates of the COF/PEDOT composites show significant improvements in volumetric energy and power densities relative to unmodified COF films, as well as outstanding stability to cycling. These findings justify follow-up efforts to access and probe even thicker COF films, further optimize the COF and conducting polymer structures, and explore other applications that will benefit from the enhanced electrical conductivity of the hybrid films. Positive developments in these directions will continue to demonstrate the promise of designed, structurally precise organic materials for EES devices.

### METHODS

#### Synthesis of DAAQ–TFP Films.

DAAQ (17 mg, 0.071 mmol) in N,N-dimethylformamide was added to a glass vial. A gold electrode (2.5 cm × 1.3 cm) was submerged in the solution, and a septum was used to seal the vial. The solution was placed on a hot plate preheated to 90 °C. Subsequently, TFP (10 mg, 0.048 mmol) was added over the course of 1 h via syringe from a 10 mg mL⁻¹ solution in DMF. During the course of the addition, the reaction mixture was gently swirled. After the addition, the reaction was allowed to proceed at 90 °C for an additional 3 h. The total reaction time was 4 h (including TFP addition), and the final volume was 4.2 mL after TFP addition. After the reaction was complete, the film which covered electrode was removed, rinsed three times with DMF and twice with acetone, and then dried in air.

#### Electropolymerization of 3,4-Ethylenedioxythiophene (EDOT).

Electropolymerization was carried out in a three electrode configuration with a DAAQ–TFP COF film on gold (prepared as described above) as the working electrode, a Ag/AgClO₄ reference electrode, and either a coiled Pt wire or high surface area carbon counter electrode. A controlled area (0.64 cm²) surface cell was used for electrochemistry experiments. A 0.1 M solution of EDOT was prepared in 0.1 M TBAP, and nine electrolypolymerization cycles between ~0.5 and 1.1 V vs Ag/AgClO₄ at 20 mV s⁻¹ were carried out. After electropolymerization, the PEDOT-modified DAAQ–TFP on gold was rinsed 3X with acetonitrile and 2X with acetone before electrochemical testing.

#### Height Analysis of Films.

AFM was used to determine film thicknesses both before and after electropolymerization of EDOT. Film heights were obtained from averaging step edges at three locations on the film (see Supporting Information for further details).

#### Electrochemical Testing of PEDOT-Modified DAAQ–TFP Films.

After electropolymerization, the prepared films were rinsed as described above and submerged in 0.5 M H₂SO₄. For initial testing, cyclic voltammetry with scan rate dependence or galvanostatic charge/discharge experiments were performed in a three electrode configuration. Capacitances were obtained from the discharge curves by multiplying the applied current by the time of discharge and normalizing by the examined voltage window and geometric film volume. Film volumes were obtained from the electrochemical cell area and the experimentally determined film thicknesses as described above. Sample calculations can be found in the Supporting Information. Thin films were also tested in a two-electrode thin
film setup where films were countered to a high surface area carbon after shorting the reference and counter electrode together. Sample calculations for volumetric capacitance and electrochemically accessible anthraquinones can be found in the Supporting Information.

Sample calculations of volumetric capacitances can be found in the Supporting Information.

**ASSOCIATED CONTENT**

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acscentsci.6b00220.

Experimental details, sample calculations, and additional electrochemical characterization (PDF)

Supplemental video (AVI)

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**Notes**

The authors declare no competing financial interest.

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