Covalently Linked Polyoxometalate–Polypyrrole Hybrids: Electropolymer Materials with Dual-Mode Enhanced Capacitive Energy Storage

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ABSTRACT: Lindqvist-type polyoxometalates (POMs) derivatized with pyrrole (Py) via an aryl-imido linkage [MoO$_3$O$_5$NPhPy]$^{2-}$ (1) and [Mo$_5$O$_{11}$NPhCCPhPy]$^{2-}$ (2) undergo coelectropolymerization with pyrrole, producing the first electropolymer films with covalently attached POM “molecular metal oxides”. X-ray photoelectron spectroscopy (XPS) and energy-dispersive X-ray (EDX) elemental analyses indicate that the loadings of POM achieved are far higher than in conventional, noncovalent inclusion films, and covalent attachment prevents loss of POM on initial reduction cycles. Cyclic voltammetry, electrochemical impedance spectroscopy, and galvanostatic charge–discharge measurements together indicate that the POMs enhance the specific capacitance (up to 5×) and decrease the charge-transfer resistance of the films by both modifying the behavior of the polypyrrole (PPy) film and introducing a substantial additional faradaic contribution through the POM redox processes. Increasing the length of the POM–PPy linker improves both capacitance and stability, with PPy-2 retaining 95% of its initial capacitance over 1200 cycles.

INTRODUCTION

Polyoxometalates, a vast class of anionic, molecular metal oxides,[1,2] offer a wealth of catalytic,[3–5] magnetic,[6,7] and electro-optical[8–10] properties suitable for exploitation in materials. Among the most important is their ability to act as stable, multielectron acceptors (up to 24 electrons in the solid state)[11] with structures and properties that can be tailored for specific demands at the molecular level, e.g., by connection of organic moieties[12–15] or inclusion of almost any heterometal.[16] Coupled with their facility for rapid electron transfer, this makes them an excellent base for electrochemical energy storage in batteries and supercapacitors,[17–20] as demonstrated by the use of [PMo$_7$V$_2$O$_{40}$]$^{3-}$ and [PW$_7$V$_2$O$_{40}$]$^{3-}$ as cathode materials for lithium-ion batteries.[20] However, the direct use of POMs in both these and other devices is restricted by high solubility and low conductivity as they are typically salts with negligible conductivity.[21] Interfacing of POMs with devices can be achieved by electrically “wiring” the POM to conductive substrates, such as conductive polymers (CPs). These are an attractive choice as their electronic and optical properties, including charge storage, can complement those of the POM and there is a growing interest in composite materials that introduce other functional components (metal oxides, metal nanoparticles) to CPs, resulting in enhanced, emergent properties.[24–26] Polypyrrole (PPy) is the most widely studied CP due to its combination of facile synthesis and device construction by electropolymerization, with high electrical conductivity, high energy storage capacity, high stability, and biocompatibility.[27] POMs have been incorporated into PPy by noncovalent inclusion in the cationic PPy matrix, producing materials with enhanced properties for energy storage[28–30] as well as catalysis and sensing applications.[31–34] However, POM loadings are restricted by the level of positive charge on the PPy and the anions can be vulnerable to leaching, particularly upon redox cycling. Loading and stability could be increased by covalent linkage, and the increased control over spatial relationships and electronic coupling provided by covalent materials is expected to provide enhanced and novel properties. In supercapacitors specifically, performance may be enhanced by covalently wiring charge storage components together at the molecular level so as to maximize the electron-transfer activity within the material and to the external circuit. Yet, there are relatively few examples of POMs covalently connected to polymers[12,25–37] and, to our knowledge, none involving PPy or other common CPs (e.g., polyaniline, polythiophene) that can be grown by electropolymerization.

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Thus, motivated by the general desirability of a covalent route to POM–PPy hybrids and recent reports indicating the enhanced capacitive energy storage of PPy noncovalently doped with Lindqvist (M₆O₁₉⁻²) polyoxometalate anions, we report the synthesis of PPy covalently linked to arylimido–Lindqvist anions [Mo₆O₁₈NAr]²⁻ through coelectropolymerization of pyrrole with POM functionalized monomers (Scheme 1). Study of the resulting films by DC and AC electrochemical techniques, electron microscopy, X-ray photoelectron spectroscopy (XPS), and energy-dispersive X-ray analysis (EDX) indicates stable film formation and bulk morphological studies). By taking advantage of the combination of FIB with an SEM, it was possible to have high-resolution images for the surface of the polymer hybrid. All Electrochemical Experiments. Electrochemical experiments were performed using high-resolution SEM at the Leeds EPSRC National Nanoscience Facility (LENNF). Sample preparation involved the use of FEI Nova200 dual-beam SEM/FIB fitted with a Kleindiek micromanipulator and operated at 30 keV (5 nA), technique for cross-section TEM (for both film-thickness determination and bulk morphological studies). By taking advantage of the combination of FIB with an SEM, it was possible to have high-resolution images for the surface of the polymer films.

EXPERIMENTAL SECTION

All Chemicals. Chemicals were purchased from Sigma-Aldrich (Gillingham, UK). Acetonitrile solvent was dried by reflux over CaH₂ and collected by distillation. Tetraethylammonium tetrafluoroborate (Bu₄NBF₄) was prepared following a literature method and used for all experiments as the electrolyte. Monomers 1 ([NBu₄]₂[Mo₆O₁₈NC₆H₄CC₆H₄NC₆H₄]) and 2 ([NBu₄]₂[Mo₆O₁₈NC₆H₄CC₆H₄NC₆H₄]) were synthesized by procedures previously developed and published by our group. All Electrochemical Experiments. Electrochemical experiments were performed under nitrogen using a Schlenk line or glovebox and performed using an Autolab PGSTAT302N potentiostat/galvanostat (Metrohm) with a three-electrode system. The glassy carbon (GC) working electrode was polished using polycrystalline diamond suspensions 3 and 1 μm sequentially and then alumina 0.3 μm. The counter electrode was Pt, and Ag or Ag/AgCl was used as a reference electrode, where the ferrocene (I/III) couple (Fc/Fc⁺) is at 0.43 V vs Ag/AgCl. Cyclic voltammetry experiments were performed using a three-compartment cell. Electrodeposition experiments were performed by cyclic voltammetry at the potential range from −2 to 1.8 V, using glassy carbon (GC), fluorine tin oxide (FTO) glass, and Pt electrodes immersed in dry acetonitrile with 0.1 M Bu₄NBF₄ electrolyte, 7 × 10⁻⁴ M POM derivative monomer 1 or 2, and 3 × 10⁻⁴ M pyrrole.

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Electrochemical Impedance Spectroscopy (EIS) Measurements. Electrochemical impedance spectroscopy (EIS) measurements were carried out using 0.9 to 1.5 V potentials at current range 1 mA with a frequency range from 10 to 10⁶ Hz, to evaluate the charge-transfer resistance and electrolyte diffusion of copolymer films with 10 deposition cycles on GC electrodes. The electrolyte was dry acetonitrile with 0.1 M Bu₄NBF₄ and the reference electrode was Ag wire.

Galvanostatic Charge–Discharge Experiments. Galvanostatic charge–discharge experiments were performed on GC electrodes coated with 10-deposition-cycle films of PPy, PPy-1, and PPy-2, using a potential range of 2 to −2 V against an Ag wire pseudo reference electrode. The electrolyte was 0.1 M Bu₄NBF₄ in dry acetonitrile.

EQCM Measurements. EQCM measurements used a carbon-coated 6 MHz quartz AT-cut crystal electrode (ø = 6.0 mm) immersed in dry acetonitrile with 0.1 M Bu₄NBF₄ electrolyte, 7 × 10⁻⁴ M POM derivative monomer 1 or 2, and 3 × 10⁻⁴ M pyrrole.

Scanning Electron Microscopy (SEM) and Transmission Electron Microscopy (TEM) Measurements. SEM and TEM measurements were performed using high-resolution TEM at the Leeds EPSRC National Nanoscience Facility (LENNF). Sample preparation involved the use of FEI Nova200 dual-beam SEM/FIB fitted with a Kleindiek micromanipulator and operated at 30 keV (5 nA), technique for cross-section TEM (for both film-thickness determination and bulk morphological studies). By taking advantage of the combination of FIB with an SEM, it was possible to have high-resolution images for the surface of the polymer films. SEM images and EDX analysis were also measured at UEA using a Philips CM200 FEGTEM fitted with a Gatan SC200 Orius CCD camera and Oxford Instruments 80 mm2 EDX SDD running AZtec software (Scheme 2).

RESULTS AND DISCUSSION

Cyclic Voltammetry of Monomers and Their Electropolymerization into Films. Electropolymerization of pyrrole

Scheme 1. Organoimidoo-Hexamolybdate Anion-Based Monomers 1 and 2

![Scheme 1](image1)

Scheme 2. Electropolymerization of Monomer 1 to Form PPy-1a

![Scheme 2](image2)

“aThe polypyrrole is shown in its reduced state with the monomer 1/pyrrole ratio approximately consistent with EDX, XPS, and electrochemical measurements. PPy-2 is synthesized by an identical procedure with monomer 2.
POM derivatives is important, as a change in the oxidation capacitive region. The positive shift in this oxidation for the deposited at the electrode and the onset of its conductive, corresponds to oxidation of the polypyrrole backbone. 

PPy-1 copolymers derivatives electrochemical process centered at ca. 0.5 V for the POM and S2. Film formation is indicated by growth of an monomer and the negatively charged \([\text{Mo}_6\text{O}_{18}\text{NAr}]\) possibly a weak interaction between the radical cation of the monomer. Thus, we found that the derivative monomers currents with no subsequent oxidations resulted in poor and broad peak (see Figure 1a) but no sign of electropolymerization: process peaking at +1.4 V associated with the pyrrole units oxidatively to up to +1.8 V reveals an irreversible oxidative Ag/AgCl for both (Figure S1). Sweeping these monomers sterically demanding phenyl ring and the \{\text{Mo}_6\} cluster, and lack of polymerization, due to steric hindrance (presence of the N-substituted pyrroles often show a negative shift in potential of ca. 180 mV introduced by the electron-donating organoimido group, to around −0.5 V vs Ag/AgCl for both (Figure S1). Sweeping these monomers oxidatively to up to +1.8 V reveals an irreversible oxidative process peaking at +1.4 V associated with the pyrrole units (see Figure 1a) but no sign of electropolymerization: subsequent oxidations resulted in poor and broad peak currents with no film deposited onto the electrode surface. This is not surprising as N-substituted pyrroles often show a lack of polymerization, due to steric hindrance (presence of the sterically demanding phenyl ring and the \{\text{Mo}_6\} cluster), and possibly a weak interaction between the radical cation of the monomer and the negatively charged \([\text{Mo}_6\text{O}_{18}\text{NAr}]^{2-}\) cluster.

Copolymerization has been shown to be one solution for such problems, where a polymerizable monomer such as pyrrole (see Figure 1b) is mixed with the desired monomer. Thus, we found that the derivative monomers would coelectropolymerize with underivatized pyrrole at up to 70 mol % of Monomers 1 and 2 to form corresponding copolymers PPy-1 and PPy-2, respectively; see Figures 1c,d and S2. Film formation is indicated by growth of an electrochemical process centered at ca. 0.5 V for the POM derivatives PPy-1 and PPy-2 and ca. 0.2 V for PPy, which corresponds to oxidation of the polypyrrole backbone deposited at the electrode and the onset of its conductive, capacitive region. The positive shift in this oxidation for the POM derivatives is important, as a change in the oxidation potential of the film implies polymer chains that are chemically different to PPy, i.e., possessing a distribution of derivatized and underivatized pyrrole units. This is further supported by a larger positive shift (ca. +0.4 V) in the peak monomer oxidation potential observed in the synthesis of PPy-1 and PPy-2 vs PPy, as they imply the involvement of monomers 1 and 2 in the polymerization reactions. It is also consistent with the literature, where functionalization of pyrrole, especially at the N-position, may introduce electronic effects that shift the oxidation potential of the monomer.

### ELECTROCHEMICAL QUARTZ CRYSTAL MICROBALANCE STUDY

Electrochemical quartz crystal spectroscopy (EQCM) is a powerful technique to investigate changes in mass by the change in frequency during the deposition of the polymeric film. It enables measurement of the mass of the deposited film according to the Sauerbrey equation:

\[
\Delta f = -C_f \frac{\Delta m}{A}
\]

where \(\Delta f\) is the change in frequency (Hz), \(C_f\) is the sensitivity factor of the crystal (0.0815 Hz ng\(^{-1}\) cm\(^2\)), \(\Delta m\) is the mass change per unit (ng), and \(A\) is the area of the substrate (cm\(^2\)). EQCM was recorded with the swept potential during formation of PPy, PPy-1, and PPy-2 in acetonitrile with 0.1 M Bu\(_4\)NBF\(_4\) (Figure 2). In all cases, a frequency decrease is observed with each cycle in the potential range between −0.2 and 1.6 V, indicating increased mass due to film deposition. Interestingly, it can be seen that while PPy-1 (2.08 × 10\(^{-6}\) g average per cycle on 0.07 cm\(^2\)) deposits more mass than PPy (1.04 × 10\(^{-6}\) g average per cycle), consistent with the high formula weight of monomer 1, PPy-2 shows less deposited mass (0.86 × 10\(^{-6}\) g average per cycle) compared to either of the other materials. This is consistent with the increased steric bulk provided by the longer \(\pi\)-bridge in monomer 2 impeding...
the polymerization and deposition process, leading to slower and thus more ordered polymer chain growth.

### ELECTRON MICROSCOPY

The surface morphology of the resulting electrogenerated films on FTO glass electrodes, at different film thicknesses, were analyzed by scanning electron microscopy (SEM) and high-resolution transmission electron microscopy (TEM) with cross-section analysis. The morphology of PPy-1 and PPy-2 copolymers is uneven; however, other N-substituted pyrroles have been found to show a more even morphology. This implies a higher surface area and thus potentially higher performance for these POM-derivatized materials due to improved charge transport between the film and the electrolyte. Figure 3 shows the SEM images of PPy-2 of 10-cycle film thickness. In the high-magnification image, the amorphous nature with densely packed surfaces based on spherical particles can be clearly observed, with no phase-separated domains. TEM images show clearly that the topography of the films is relatively similar to that of the FTO substrate (Figure 4). Film thicknesses were between 60 and 120 nm depending on the number of deposition cycles, and elemental mapping shows that Mo is present through the entire thickness of the film, confirming that there is no phase separation.

### REFLECTANCE FOURIER TRANSFORM IR CHARACTERIZATION

The PPy-1 copolymer was studied by reflectance Fourier transform infrared (FTIR) spectroscopy to identify the surface functionalities. Fresh films of the copolymer were deposited on a Pt working electrode, washed with fresh MeCN, and allowed to air-dry with no marking of cracking or peeling. Spectra were measured against the uncoated polished Pt electrode, and the spectra were compared with those for the monomers and PPy-
only films. The FTIR spectrum of monomer 1 (see Figure 5) showed the characteristic Mo = O stretch at 940 and an imido stretch Mo≡N at 947 cm$^{-1}$.

When comparing monomer 1 with its corresponding PPy-1 copolymer (Figure 5, black trace), the latter showed a shift in the position and a combination of these functional groups (Mo=O and Mo≡N) between 950 and 916 cm$^{-1}$. Figure 6 presents the reflectance FTIR spectra of PPy-1 (black trace) and PPy only (red trace), showing a clear difference from PPy in the presence of the BF$_4^-$ peak at 1070 cm$^{-1}$ in both films, which is consistent with the literature. However, the strength of the BF$_4^-$ peak relative to the other vibrations is far weaker in PPy-1, showing that the presence of the negatively charged POM reduces or eliminates the need to include anions to balance any positive charge on the PPy backbone. Indeed, at higher frequencies (Figure S3), the C–H stretching band for PPy-1 is far stronger than for PPy, consistent with substantial inclusion of NBu$_4^+$ cations in the film to compensate for the presence of the self-doping POM monomers. Inclusion of NBu$_4^+$ implies that the PPy-1 polymer carries a negative charge and, combined with the different potentials needed for electropolymerization (vide supra) and the different behavior to POM noncovalent inclusion films upon redox cycling (vide infra), shows that the POM monomers must be covalently connected as part of the backbone.

**X-RAY PHOTOELECTRON (XPS) AND ENERGY-DISPERSIVE (EDX) X-RAY SPECTROSCOPIC**

XPS was carried out on the FTO electrode coated with copolymer films in the polypyrrole reduced (neutral) state. Figure 7 shows an XPS spectrum representative of the Mo 3d core level for PPy-1 (blue) and PPy-2 copolymers (red) of 20-deposition-cycle film thickness. The data were calibrated with the carbon signal at 288 eV.

The FTIR spectrum of monomer 1 using the ATR technique showing the characteristic Mo = O stretch at 940 and an imido stretch Mo≡N at 947 cm$^{-1}$.

Figure 5. FTIR spectrum of monomer 1 using the ATR technique showing the characteristic Mo = O stretch at 940 and an imido stretch Mo≡N at 947 cm$^{-1}$.

![Figure 6. Reflectance FTIR spectrum of a Pt disc electrode coated with PPy only (red trace) and the PPy-1 copolymer (black trace).](image)

![Figure 7. XPS spectrum of the Mo 3d core levels for PPy-1 (blue) and PPy-2 copolymers (red) of 20-deposition-cycle film thickness. The data were calibrated with the carbon signal at 288 eV.](image)
Table 1. XPS Surface Elemental Analysis

| material, thickness | elemental abundance [atomic %] |
|--------------------|-------------------------------|
|                    | C 1s | F 1s | Mo 3d | N 1s | Na 1s | O 1s | S 2p |
| PPy-1, 20 cycles   | 69.16 | 6.96 | 5.23 | 0.34 | 17.25 | 1.05 |
| PPy-2, 10 cycles   | 75.76 | 4.70 | 4.63 | 14.00 | 0.92 |
| PPy-2, 20 cycles   | 75.49 | 5.00 | 5.16 | 0.01 | 14.17 | 0.17 |
| PPy-2, 30 cycles   | 72.34 | 5.90 | 4.19 | 0.00 | 17.27 | 0.31 |
| PPy, 20 cycles     | 73.36 | 0.38 | 0.22 | 8.93 | 0.25 | 14.44 | 2.43 |

Conditions: electrolyte 0.1 M NBu4BF4 in MeCN, scan rate 100 mV/s.

Consecutive cyclic voltammograms of a PPy-1 film showing minimal changes from redox cycles 1 to 15. (d) Plot of current peak I vs scan rate (ν) for reduction of the PPy-1 film. Conditions: electrolyte 0.1 M NBu4BF4 in MeCN, scan rate 100 mV/s, working electrode 0.07 cm² GC, Ag/AgCl as the reference electrode.

Figure 8. (a) CV showing the \[\{\text{Mo}_6\}\] redox wave in PPy-1. (b) CVs of a \(\{\text{Mo}_6\}\)-PPy inclusion film showing the loss of POM after the first redox cycle. (c) Consecutive cyclic voltammograms of a PPy-1 film showing minimal changes from redox cycles 1 to 15. (d) Plot of current peak I vs scan rate (ν) for reduction of the PPy-1 film. Conditions: electrolyte 0.1 M NBu4BF4 in MeCN, scan rate 100 mV/s, working electrode 0.07 cm² GC, Ag/AgCl as the reference electrode.

Figure 9. CVs of PPy-1 (black trace) and PPy-2 (red trace) of 10-cycle film thickness on the FTO electrode in 0.1 M NBu4BF4/McCN at 25 °C vs Ag/AgCl at scan rate 250 mV/s.

visible, consistent with the high level of POM functionalization determined by XPS and EDX. Integrating the oxidation wave of the PPy (from −0.2 to 0.6 V) and the reduction wave of \(\{\text{Mo}_6\}\) from (−0.2 to −0.8 V) indicates that around 20–30% of pyrrole monomers are derivatized with a POM. This is slightly higher than suggested by XPS but represents a good agreement given the considerable uncertainties involved in integrating a cyclic voltammogram—such as choice of baseline and the contribution of double-layer capacitance.

Electrochemical impedance spectroscopy (EIS) was performed in the conductive, capacitive voltage region of the polypyrroles to obtain comprehensive information about the properties of capacitors based on the PPy component, and results are shown in Figure 10. Nyquist plots include three parts, a semicircle in the high-frequency region corresponding to the charge-transfer resistance, a transition region possibly associated with Warburg impedance (W) within the film, and the straight-line (C) part at low frequency corresponding to the ion (electrolyte) charging during the charge/discharge process. The semicircles of PPy-1 and PPy-2 in the high-frequency region are much smaller than for the PPy electrode, indicating a much lower apparent charge-transfer resistance \(R_{ct}\). As the POMs cannot be participating in faradic reactions at the potential used for the measurement (+0.6 V), the likely explanation for this is a more open structure with a higher surface area for charge transfer between the film and electrolyte.\(^{2,5}\) In the low-frequency domains, both PPy-1 and PPy-2 show similar behavior.

The Bode and phase angle plots (Figure 10b/c) show that at low frequencies (10 Hz), the phase angle of PPy-2 is 53°,

...from 0 to −0.75 V on films of 10 deposition cycles. These show that on the CV time scale the resulting film of the PPy-1 copolymer can undergo around 15 redox cycles (see Figure 8c) with only minimal changes, indicating the high stability of these films, due to the covalent linkage between the Lindquist derivative and pyrrole units. This is clearly evident when comparing PPy-1 with an analogue made by the inclusion of \([\text{Mo}_6\text{O}_{19}]^{2-}\) in underderivatized PPy; the latter showed a large decrease in current intensity and shift to a less negative potential when moving from redox cycle 1 to 2; see Figure 8b. Similar results indicating loss of POM were obtained for an analogue made by the inclusion of the arylimido-derivatized \([\text{Mo}_6\text{O}_{19}\text{NArI}]^{2-}\) cluster in PPy (Figure S6 in the SI), showing that the improved stability of PPy-1 and PPy-2 is not simply a result of the bulky, appended organic group helping entrap the POM in the PPy matrix.

Full-range cyclic voltammograms of PPy-1 and PPy-2 are shown in Figure 9. Both POM and PPy processes are clearly...
The device has the most ideal capacitive performance. At higher frequencies, the phase angles of PPY-2 and PPY-1 fall to 19 and 22°, respectively. This is (in part) due to Warburg impedance and arises from the porous structure of the film. The difference from PPY may be linked to inclusion of NBu₄⁺ cations, rather than BF₄⁻ anions, in the film: NBu₄⁺ is large, and upon oxidation, there may be processes ejecting it from the film, as well as taking in the smaller BF₄⁻ anions. Figure 10d shows the electrochemical circuit fit where Rₛ is the solution resistance, R₁ and C₁ are the resistance and capacitance of the double-layer capacitor, respectively, and C₂ is the capacitance of the pseudocapacitor, resulting from the film (Table 2).

Table 2. Electrode Behavior (PPY, PPY-1, and PPY-2) Calculated from the Electrochemical Circuit Fit of the Experimental EIS Data

| sample  | Rₛ [Ω] | R₁ [Ω] | C₁ [F] | C₂ [F] | Yₑ [mS s^0.5] |
|---------|--------|--------|--------|--------|--------------|
| PPY     | 316    | 1280   | 2.06 × 10⁻⁹ | 8.85 × 10⁻⁶ | 68.6         |
| PPY-1   | 282    | 308    | 3.55 × 10⁻⁹ | 41.6 × 10⁻⁶ | 659          |
| PPY-2   | 278    | 375    | 3.68 × 10⁻⁹ | 17.9 × 10⁻⁶ | 486          |

“Real part of Yₑ where the Warburg impedance Zₑ = 1/[jωR₁]^{1/2}.

CVs showing clear polyoxometalate peaks for PPY-1 and PPY-2 indicate that their energy storage mechanism is not purely pseudocapacitive—they have some battery-like character. Accordingly, below we also provide capacities in mAh for these two materials.

Compared to underivatized polypyrrole (PPy) measured under the same conditions, films with covalently connected {Mo₆} anions show a significant increase in the specific capacitance of the polypyrrole—to 22 F g⁻¹ for PPY-1 and to 34 F g⁻¹ for PPY-2; see Table 3. This corresponds to a specific capacitance increase of ca. 2× vs PPY for PPY-1 and 3× for PPY-2, roughly in line with that seen for noncovalent inclusion of [Mo₆O₁₉]²⁻ in PPY (2X).38 However, the large loading of polyoxometalate in these films means that it also contributes a significant pseudocapacitance. This was assessed in the same way as the PPy contribution, revealing that the POM contributes around 50% of the capacitance of the polypyrrole film, to give a total of 33.5 F g⁻¹ for PPY-1 and 52 F g⁻¹ for PPY-2 and thus an enhancement vs PPY of 3–5X. Considering the battery-like character of these systems, this translates to capacities of ca. 14 and 22 mAh g⁻¹, respectively. The overall POM contribution to charge storage is approximately consistent with the ratio of POMs to pyrroline units estimated by XPS, considering that each POM stores one electron on reduction, and each pyrrole 0.33 positive charges on oxidation. Overall, the results show that the modification of PPy with hexamolybdate {Mo₆} anions via covalent linkage gives access to composites with enhanced capacitive charge storage that can be ascribed both to steric bulk, which improves the structure

### SPECIFIC CAPACITANCE AND CHARGE–DISCHARGE STABILITY

The specific charge (C_{charge}) and discharge (C_{discharge}) capacitances associated with the polypyrrole and {Mo₆} anions over a potential range of 0.7 to −0.8 V were calculated by integrating the current density of the CV curve in the oxidation wave for polypyrrole and in the reduction wave for the {Mo₆} anions, respectively (Figure 9). This energy storage property has been calculated, and tabulated as capacitance (F g⁻¹), because most of the storage occurs in the polypyrrole, which is considered to be a pseudocapacitor. There is, however, considerable current discussion of systems that lie at the boundary between batteries and pseudocapacitors,19,54–56 and
(surface area/porosity) of the film, and to a direct faradic contribution from the POM electron acceptor.

Galvanostatic charge–discharge (GCD) measurements (Figure 11) were also used to evaluate the specific capacitance and electrochemical stability of PPy, PPy-1, and PPy-2 over 1200 cycles. The specific capacitance was calculated according to the following equation

$$\text{Cs} = \frac{I \Delta t}{A \Delta E}$$

where $I$ is the maximum constant current (A), $t$ is time (s), $E$ is the potential (V) during the charge–discharge process, and $A$ is the area of the electrode. The GCD curves are almost symmetrical, indicating good capacitive behavior. Notably, the curves for PPy-1 and PPy-2 extend to negative voltages, consistent with involvement of the POM in the process, and the time per charge/discharge cycle is shorter than for PPy, at around 21 vs 36 s, indicating better charge transport in the POM-derivatized films. The specific capacitances (Table 4) follow the same trend as those obtained by cyclic voltammetry, with PPy-1 and PPy-2 showing excellent agreement with the total POM + polypyrrole capacitance calculated from the CV. Calculation of capacity produces values of ca. 11 mAh g$^{-1}$ for both materials, the value for PPy-2 dropping substantially compared to that assessed by CV because of the smaller voltage range observed in the GCD curves.

Interestingly, PPy-2 shows better multicycle stability compared with PPy-1. Figure 12 shows the capacitance retention of both copolymers where PPy-1 exhibits 81% and PPy-2 shows 95% capacitance retention over 1200 cycles. Thus, while the multicycle stability of PPy-1 is comparable to that shown for noncovalent inclusion of $\text{Mo}_6\text{O}_{19}$$^{2-}$, PPy-2 demonstrates the potential of covalent linkage to improve the stability of POM–polymer composite systems. The improved performance of PPy-2 compared to PPy-1 in both stability and specific capacitance is consistent with the longer spacer between pyrrole-N and POM forcing larger ion channels, increasing the active surface area, and reducing disruption of the film by ion migration.

### CONCLUSIONS

Polypyrrole supercapacitors with covalently linked Lindqvist polyoxometalates were successfully prepared by electropolymerization. These films are the first covalent POM–PPy hybrids, and as the first demonstration of electropolymerization of a POM-derivatized monomer, they open new possibilities for construction of POM hybrid materials. The covalent linkage gives access to far higher loadings of POM than obtained in noncovalent inclusion films, prevents loss of POM in early reduction cycles, and increasing its length increases both the specific capacitance and stability of the film.

| sample | specific capacitance $[\text{F g}^{-1}]$ |
|--------|----------------------------------------|
| PPy    | 19                                     |
| PPy-1  | 35                                     |
| PPy-2  | 54                                     |

![Figure 11. Galvanostatic charge–discharge curves for 10-cycle PPy, PPy-1, and PPy-2 films on GC electrodes. Electrolyte: 0.1 M NBu$_4$BF$_4$/MeCN at 25 °C vs the Ag wire reference.](image1)

![Figure 12. Capacitance retention % of (a) PPy-1 and (b) PPy-2 of 10-cycle deposited film thickness on GC electrodes for the first 1200 cycles. Electrolyte: 0.1 M NBu$_4$BF$_4$/MeCN at 25 °C vs the Ag wire reference.](image2)
to cycling, such that 95% capacitance retention over 1200 cycles is achieved for the longer diphenylacetylene spacer. Crucially, the POM enhances specific capacitance both by modifying the polypyrrole itself, with reduced charge-transfer resistance implying that a more open structure results from the steric bulk of the POMs, and by a direct faradic contribution through the POM redox process. Future work will focus on optimizing the loading of the POM for the best specific capacitance and expand the approach to POMs capable of storing multiple electrons.

■ ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.macromol.0c02354.

Additional cyclic voltammograms of monomers and electropolymer films, photographs of coated electrodes, and additional FTIR data and EDX spectra (PDF)

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The authors declare no competing financial interest.

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■ ABBREVIATIONS USED

PPy, polypyrrole; POM, polyoxometalate; TEM, transmission electron microscopy; SEM, scanning electron microscopy; CP, conducting polymer; GC, glassy carbon; EIS, electrochemical impedance spectroscopy; GCD, galvanostatic charge-discharge; EQCM, electrochemical quartz crystal microbalance; FTIR, Fourier transform infrared; XPS, X-ray photoelectron spectroscopy; EDX, energy-dispersive X-ray spectroscopy; FTO, fluorine tin oxide

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