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Electron transport kinetics for viologen-containing polypeptides with varying side group linker spacing[†]

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Studies investigating the influence of the length of linkers between redox-active moieties and peptidebased polymer backbones were conducted to advance fundamental knowledge toward the design and development of sustainably-sourced, recyclable, and degradable materials for energy applications. In this work, precursor polypeptides were synthesized through the ring-opening polymerizations of Ncarboxyanhydrides decorated with varying lengths of alkylchloride side chain groups, followed by postpolymerization installation of the viologen moieties. Electrochemical interrogation of the viologen-based polypeptides provided estimates of the electron transfer rate constants, both heterogeneous (k^0) and electron self-exchange (k_{ex}) , the apparent diffusion coefficient (D_{ap}) , and their device-based energy storage performance. For the first redox couple (viologen dication state to viologen radical-cation state), it was found that the rate of electron transfer among the pendant groups in all viologen-based polypeptides, kex, was not significantly impacted by linker length. In contrast, for the second redox couple (viologen radical-cation state to the neutral viologen), k_{ex} varied with linker length and was fastest during reduction from the viologen radical-cation state to the neutral viologen. Most interestingly, a linear relationship was identified between $log(k^0)$ and $log(k_{ex})$ with a slope of 1.85, indicating that electron transport in the viologen-based polypeptides followed most closely to Marcus-Hush theory with diffusion limitations or Laviron-Andrieux-Savéant (LAS) theory. Finally, the polypeptides were studied in lithium metal half cells to determine the relationship between k_{ex} and energy storage performance. The viologen-based polypeptide with the moderate length linker exhibited the highest capacity and lowest degree of swelling, but only moderate k_{ex} , demonstrating that the device performance was primarily influenced electrode swelling. Taken together, the viologen-polypeptide backbone dictated the mechanism of electron transfer, whereas the linker length could be used to alter the rate of electron transfer (k_{ex}). Balancing the rate of electron transfer (k_{ex}) and degree of swelling will be a major challenge to identify polymers for high performance energy storage devices.

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Introduction

With the increase in demand for lithium-ion batteries and the strained supply of strategic elements, alternative active materials are needed for battery electrodes. One such material class of interest is redox-active polymers (RAPs), which include both conjugated polymers and non-conjugated polymers with redox-

active pendant groups.1-5 Extensive work has been done to investigate structure-property relationships for conjugated polymers in the solid-state to reveal linkages among the conduction mechanism, regiochemistry, and crystallinity.6-9 However, there has been significantly less work done for nonconjugated RAPs, due to their amorphous nature for which "crystal-engineering" is precluded. For cathodic RAPs with pendant redox-active groups, such as 2,2,6,6-tetramethylpiperidin1oxyl (TEMPO) containing polymers, there have been several studies investigating the electron transport mechanism.¹⁰⁻¹² However, there has been less work on anodic pendant RAPs,13,14 and most studies used flexible or semiflexible backbones. Previously, our group reported redox-active polypeptides in degradable solid-state and redox flow batteries,^{15,16} for which the polypeptide backbones displayed

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solution-state α -helical chain conformations, but the electron transfer mechanism was not examined.

Prior work shows that local ordering and mobility of redoxactive groups influence electron transport. Akhoury et al. studied non-crosslinked and crosslinked ferrocene-based grafts on carbon particles.¹⁷ The authors found that bounded diffusion and electron hopping influenced the rate of electron transfer for the non-crosslinked grafts, whereas only electron hopping contributed in the crosslinked grafts, due to reduced mobility.17 Similarly, Whiting et al. studied poly(triphenylamine acrylate) brushes and spin-coated films and found that the ordered brushes led to increased rates of electron transfer as compared to the more disordered spin-coated films.18 In addition to the influence of the helical polypeptide backbone on electron transfer among redox-active pendant groups, electron tunneling through the helical backbone could also influence electron transport. Arikuma et al. demonstrated that for short ferrocene-terminated helical polypeptides (less than 16 repeat units), electron tunneling contributed to the overall rate of electron transfer.¹⁹ Deng et al. reported the synthesis of viologen-containing helical polyacetylenes; however, effects of the helical backbone and pendant group spacing on the electron transport mechanism were not studied.20

Three prominent theories consider the diffusion of polymerbound redox active moieties: Dahms–Ruff (D–R) theory,^{21,22} Laviron–Andrieux–Savéant (LAS) theory,^{23,24} and the diffusion cooperative model (Fig. 1).¹⁰ D–R theory describes the electron transfer of freely-diffusive redox-active groups.^{21,22} In this model, charge transport is limited by the diffusion of the molecules, where electron transfer occurs only as rapidly as molecules contact each other. Explicitly stated, this model is applied in systems where the physical diffusion (D_{phys}) of the redox-active moieties is larger than the diffusion of electron transfer (D_e). Experimental studies have shown that D–R theory accurately describes electron transfer in metal complexes,²⁵ viologen-polymer and dimer solutions,^{14,26} ferrocene polymer solutions,²⁶ and viologen-based ionic liquids.²⁷ As the physical diffusion $(D_{\rm phys})$ of the redox-active moieties is further restricted and the redox-active moieties are immobilized $(D_{\rm phys} \approx 0)$, $D_{\rm e}$ becomes larger than $D_{\rm phys}$, and LAS theory is applicable.^{23,24} Previously, LAS theory has been used to describe electron transfer of TEMPO-containing polyether thin films,²⁸ ferrocenecontaining polyacrylamide gels,²⁹ crosslinked TEMPOcontaining polynorbornene,³⁰ ferrocene-based metal–organic frameworks,³¹ and viologen and TEMPO nanoparticles.³²

Recently, Burgess et al. and Sato et al. reported deviation from both D-R and LAS theory for viologen-based polymeric solutions and TEMPO-based polymer thin films, respectively.^{10,14} To account for this deviation, Sato et al. proposed the diffusion cooperative model, which considers the effects of Brownian motion of the polymer.¹⁰ In this model, the diffusion of electron transfer has contributions from both $D_{\rm e}$ and $D_{\rm phys}$. The D_{phys} considered here is significantly reduced compared to the free diffusion of the D-R theory but greater than the lack of $D_{\rm phys}$ assumed in LAS theory. This intermediate $D_{\rm phys}$ model has also been applied to an immobilized TEMPO-containing polyamine,33 TEMPO-substituted poly(ethylene sulfide),34 and a variety of nitroxide-containing polymers with varying backbone and linker groups.10 However, much of the work described in those reports involved physically-restricted crosslinked polymers or polymers with a flexible backbone.

Here, we report the synthesis of viologen-based polypeptides with varying spacing between the backbone and viologen pendant group. The influence of backbone structure is also considered through comparison of the helical polypeptide backbone with a RAP having an aliphatic backbone. The polymers were processed to produce thin-film coatings, and the electron transfer rate constants were determined from chronoamperometry and cyclic voltammetry. The determined parameters included the apparent diffusion coefficient (D_{ap}), the rate constant for electron self-exchange between pendant viologen groups (k_{ex}), and the rate constant for heterogenous charge transfer (k^0). The physical diffusion coefficients (D_{phys}) for the polymers in solution and the viologen pendant group

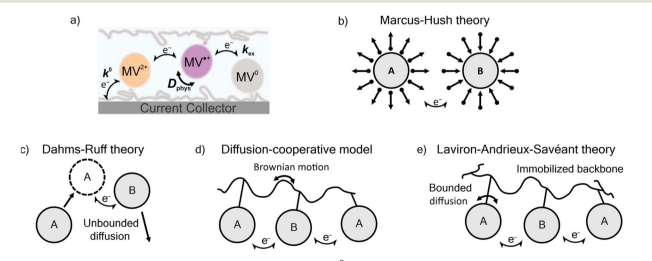


Fig. 1 (a) Schematic of rate constant for heterogenous electron transfer (k^0) or electron transfer between viologens and the current collector, rate constant for homogenous electron transfer (k_{ex}) or electron transfer/exchange between viologens, and the coefficient for diffusion of the viologens (D_{phys}). (b) A schematic of Marcus–Hush theory, as well as (c–e) representation of diffusion in three descriptions of electron-transfer.

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relative to the backbone were determined using dynamic light scattering (DLS) and computational methods, respectively. The relationship between D_e and D_{phys} was analyzed to determine the predominate mechanism of electron transport in the system. Finally, we evaluated the performance of the viologenbased polypeptides in half-cell batteries to relate the electron transfer kinetics to battery performance. Overall, these results present the dominant theory and redox couple for charge transport in viologen-functionalized polypeptides.

Experimental

Materials

Lithium chloride (LiCl, powder, \geq 99.98% trace metals basis), and γ -butyrolactone (GBL, \geq 99%) were used as received from Sigma-Aldrich. Dichloromethane (DCM, \geq 99.5% stabilized) was used as received from VWR. Ultrapure water (MQ water) was collected from a Milli-Q® integral water purification system (18 M Ω cm). Methanol was used as received from VWR. Lithium metal was purchased from Alfa-Aesar. Indium tin oxide (ITO)-coated glass (CUV5B) was purchased from Delta and cleaned by subsequent sonication in soap water, water, acetone, and isopropyl alcohol, followed by drying with N₂ and ozone plasma treatment for 10 min. Au-coated QSense quartz crystal microbalance sensors with a Ti adhesion layer (QSX 338) were purchased from Biolin Scientific and cleaned with 5:1:1 MQ H₂O: ammonium hydroxide : hydrogen peroxide at 75 °C for 5 min before use. Materials for synthesis can be found in the ESI.†

Chemical characterization

Attenuated total reflectance-Fourier transform infrared (ATR-FTIR) spectra were recorded for powder samples on an IR Prestige 21 system (Shimadzu Corp.). The data were analyzed using IRsolution v. 1.40 software. Circular dichroism (CD) spectra of the viologen polypeptide solutions in a quartz cell were recorded, between 180 and 280 nm, using a wavelength step of 1.0 nm, and had a path length of 1.0 cm, on a Chirascan CD spectrometer from Applied Photophysics, Ltd (Leatherhead, UK) equipped with a 150 watt xenon arc lamp. The viologen polypeptides and control viologen polymer were dissolved in H2O (HPLC grade) to generate solutions for analysis. The resulting CD spectra were analyzed using Pro-Data v. 5 software. ¹H and ¹³C nuclear magnetic resonance (NMR) spectra were recorded on a Bruker AVANCE NEO 400 or Bruker AVANCE III 500 spectrometer. Thermogravimetric analysis (TGA) was conducted using a Mettler-Toledo TGA2/1100/464. The measurements were performed at a heating rate of 10 °C min⁻¹ across a temperature range of 25 °C to 500 °C under a nitrogen atmosphere. Data were analyzed using Mettler-Toledo STAR^e v. 17.00 software to determine the onset degradation temperature (T_d) for all the polymers. For the final viologen chloride polymers, the glass transition temperatures (T_{α}) were determined from the inflection point of the second heating cycle measured using modulated differential scanning calorimetry (MDSC) on a Q200 DSC (TA Instruments) with a heat-cool-heat cycle. The samples were ramped from -40 °C to 200 °C at a rate of 5 °C min⁻¹ with an amplitude of 1.272 °C for a period of 60 s with nitrogen purge at 50 mL min⁻¹. For all other polymers, the glass transition temperatures (T_g) were determined from the inflection point of the second heating cycle measured using DSC on a Mettler-Toledo DSC3/700/1190 (Mettler-Toledo, Inc., Columbus, OH) under a nitrogen atmosphere. The measurements were performed with heating and cooling rates of 10 °C min⁻¹ over three heating and cooling cycles. Data analysis was conducted using Mettler-Toledo STAR^e v. 17.00 software. Finally, dynamic light scattering (DLS) data were collected, using the same aqueous solutions from the CD measurements, on a Malvern Zetasizer Nano ZS and analyzed with Zetasizer v7.13 to obtain the hydrodynamic radius values. Using the Stokes–Einstein equation, eqn (1), the solution state polymer diffusion coefficient (D_{DLS}) can be calculated from the measured hydrodynamic radius (r_h) in DLS.

$$D_{\rm DLS} = \frac{k_{\rm B}T}{6\pi\eta r_{\rm h}} \tag{1}$$

where $k_{\rm B}$ is Boltzmann's constant, *T* is the temperature (in K), and η is the viscosity of the medium (H₂O).

Synthesis of viologen polypeptides with varying alkyl spacer lengths

The viologen polypeptides were synthesized following the method of Nguyen *et al.*¹⁶ For the C3 and C9 viologen polypeptides, the first synthetic step was altered slightly by replacing 6-chloro-1-hexanol. Instead, 3-chloro-1-propanol and 9-chloro-1-nonanol were used to produce the C3 and C9 viologen polypeptides, respectively. Details of the synthetic steps can be found in the ESI.[†]

Synthesis of the control viologen polymer

Poly(vinylbenzyl methyl viologen) was synthesized following a modified method as reported by Nagarjuna *et al.*³⁵ First, methyl viologen chloride (MV-Cl) was synthesized following the procedure reported by Nguyen *et al.* followed by ion exchange using Dowex® A2 ion exchange resin.¹⁶ Poly(vinylbenzyl chloride) (PVBC) was synthesized *via* reversible additionfragmentation chain-transfer (RAFT) polymerization following a modified method as reported by Moraes *et al.* and Nagarjuna *et al.*; however, 4-cyano-4-(((dodecylthio)carbonothioyl)thio) pentanoic acid was used as the RAFT chain transfer agent.^{35,36} Details of the synthetic steps can be found in the ESI.[†]

Determination of thickness swelling ratio

Quartz crystal microbalance with dissipation (QCM-D) (QSense E1, Biolin Scientific) was used to determine the thickness swelling ratio of the viologen polymer films as a ratio of their dry film thickness (h_{dry}) and swollen film thickness ($h_{swollen}$). All QCM-D measurements were conducted in an Ar-filled glovebox. Baseline measurements were conducted on uncoated gold-coated AT-cut quartz crystal sensor in Ar and in 0.5 M LiCl in GBL. After measuring the baseline, the sensor was washed with MQ water and dried, followed by drop casting of a viologen polymer onto the sensor surface. The drop casting solution was 8 mM of repeat unit for each polypeptide in 1:1 (v:v)

methanol: MQ water. 20 μ L of the prepared solution was dropcast on a gold-coated AT-cut quartz crystal sensor (coating area $\approx 1.13 \text{ cm}^2$). The change in frequency and dissipation was determined for the polymer-coated sensor in Ar, followed by exposure to 0.5 M LiCl in GBL. The films were allowed to equilibrate for 16 h in 0.5 M LiCl in GBL and the swollen thickness was taken from the last two minutes of data. The change in frequency for each step was recorded and converted to thickness using the Sauerbrey equation.³⁷ The thickness swelling ratio was calculated as $h_{\text{swollen}}/h_{\text{dry}}$. The above protocol was repeated for each viologen polymer on new QCM-D sensors.

Experimental determination of rate constants

Thin films of each viologen polypeptide were prepared by dissolving 8 mM of repeat unit for each polypeptide in 1:1 (v:v) methanol: MO water, drop-casting 80 µL of the prepared solution onto ITO glass (coating area ≈ 1.3 cm²), and allowing to dry at ambient conditions for 24 h, followed by drying under vacuum at r.t. for 24 h. After drying, the mass of each film was measured to determine the mass loading and the thickness was determined using profilometry (KLA Alpha-Step D-500). Finally, the fabricated electrodes were utilized in a three-electrode cell (coated ITO glass working electrode with lithium metal reference and counter electrodes) with 10 mL of 0.5 M LiCl in GBL as the electrolyte. All electrochemical measurements were performed using a potentiostat (Solartron Electrochemical Interface 1287). Chronoamperometry (CA) was performed in two steps (1) from 3 V to 2.2 V vs. Li/Li^+ and (2) from 2.2 V to 1.6 V vs. Li/Li⁺ to determine the properties for both redox couples separately and cyclic voltammetry (CV) was performed from 3.2 V to 1.5 V vs. Li/Li⁺ at 10, 25, 50, and 100 mV s⁻¹.

The CA data were further analyzed to estimate the apparent diffusion coefficient (D_{ap}) by fitting using the Cottrell equation (eqn (2)) assuming semi-infinite diffusion:¹⁰

$$i = \frac{nFAC_0^*}{\sqrt{\pi t}}\sqrt{D_{\rm ap}} = \tau \frac{nFAC_0^*}{\sqrt{\pi}}\sqrt{D_{\rm ap}}$$
(2)

where *i* is the current output from CA (in mA), *n* is the number of electrons transferred per repeat unit, *A* is the electrode area (in cm²), and C_0^* is the total concentration of redox sites (or repeat units in mol cm⁻³). Following the method of Paulse *et al.*, the slope was determined from the line tangent to the CA data and the origin (0,0).³⁸ If $D_{ap} \ge D_{phys}$, the rate constant for electron self-exchange, k_{ex} was found using the Dahms–Ruff equation (eqn (3)):

$$k_{\rm ex} = \frac{6(D_{\rm ap} - D_{\rm phys})}{C_0^* \delta^2} \tag{3}$$

where δ is the average site distance, which is determined from MD simulations as discussed below. However, if $D_{\text{phys}} < D_{\text{ap}}$ and $D_{\text{phys}} \approx 0$, k_{ex} was calculated using the Laviron–Andrieux–Savéant equation (eqn (4)):

$$k_{\rm ex} = \frac{6D_{\rm ap}}{C_0^* \delta^2} \tag{4}$$

From the peak separation of the CV data at 10 mV s⁻¹, the heterogeneous rate constant, k^0 , was calculated using the simplified method of Nicholson, eqn (5). The simplified form assumed the diffusion coefficient of the oxidized and reduced species are equal to D_{ap} :

$$\psi = \frac{k^0}{\left(\pi D_{\rm ap} f \upsilon\right)^{1/2}} \tag{5}$$

where ψ is a dimensionless rate parameter from the method of Nicholson, ν is the scan rate of CV (in V s⁻¹), and D_{ap} is the apparent diffusion coefficient from CA (in cm² s⁻¹), respectively. f = F/RT, where *F* is Faraday constant (or 96 585C mol⁻¹ e⁻¹), *R* is the gas constant (or 8.3124 J mol⁻¹ K⁻¹) and *T* is the temperature (in K).

Evaluation of the capacity performance

The fabricated films were also used as working electrodes in a half cell with a lithium metal reference and filter paper soaked in 0.5 M LiCl in GBL as the separator and electrolyte, respectively. Galvanostatic charge–discharge (GCD) was performed on the half cells at different C-rates (1C is the current required to completely charge the battery in 1 h). GCD was performed for 5 cycles at 0.2, 0.4, 0.6, 0.8, 1.0, and 1.2C followed by 10 cycles at 0.2C.

Molecular dynamics simulations

Atomistic MD simulations were performed using LAMMPS software package using OPLS-2005 force field extracted by the FFLD Server toolkit from Schrodinger utilities.39,40 A real space cutoff distance of 10 Å was used for non-bonded interactions including truncated Lennard-Jones potential and coulombic interactions. Long-range coulombic interactions were treated by using the particle-particle particle-mesh solver with 10⁻⁴ accuracy. Initial configurations of an oligomer (up to 26 monomers), 7600 GBL molecules as the solvent, and the corresponding number of chloride counterions for each state of charge within a 100 Å per side cubic box were packed using Packmol package.⁴¹ The LAMMPS input files were generated using Moltemplate tool.42 At the beginning of the simulation, energy minimization was performed to relax the system. The atomic velocities were initialized with Gaussian-distributed velocity associated with 300 K. After initialization, the system was maintained at 300 K for 50 ps under NVT ensemble followed by 50 ps simulation under NPT ensemble as preequilibration steps. Subsequently, the system was heated up linearly to 1000 K for 0.5 ns, maintained at 1000 K for 0.5 ns, and cooled down linearly from 1000 K to 300 K for 2 ns to search for a lower energy configuration under NVT ensemble throughout the process. The temperature was kept at 300 K and pressure at 1 bar under NPT ensemble for 2.5 ns as the production run. A Nosé-Hoover barostat or thermostat was applied to control the pressure or temperature of the system. The timestep was set to 0.5 fs initially using a standard velocity-Verlet integrator during the pre-equilibration steps and then set to 1.0 fs during the initial heating of the system. Each type of oligomer was simulated five times under randomized initial

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conditions. During the production run, the physical diffusion coefficient of viologen groups relative to the backbone (D_{phys}) were approximated by the regression slope of mean squared displacement of viologen groups relative to the center of mass of the whole oligomer divided by six. For the simulations in the solvated state (7600 GBL molecules and a single polypeptide oligomer) the repeat unit concentration (C_0^*) was *ca.* 44.9 mM, whereas the in the solvated state (with 1 : 1 atom ratio of GBL to six polypeptide oligomers), C_0^* was *ca.* 3.85. M. The diffusion coefficients of the redox groups for the two different redox processes (MV^{2+}/MV^{*+} and MV^{*+}/MV^{0}) were estimated by taking the average values of diffusion coefficients found at each state, Table S1.†

Electronic coupling and reorganization energy calculations

The geometry of methyl viologen was optimized at B3LYP/Def2-SV(P) level of theory with Gaussian 16 (revision B.01).43 To estimate the electronic coupling between these states of charge, the obtained geometry was further stacked as a dimer at a fixed plane-to-plane distance (height) at 3.5 Å then displaced from 0 to 7 Å along the long axis (displacement). The displacement was also fixed at 0 and 1 Å separately then shifted from 3 to 7 Å along the height. Dimer snapshots were extracted from MD trajectories after equilibration with the nitrogen atom linking with the carbon linker capped with a methyl group. Electronic coupling calculations were evaluated upon these dimer structures using ALMO (MSDFT2)44 implemented in Q-Chem 5.3 at PBE0/6-31+G(d) level of theory.⁴⁵ This method was shown to be robust on electronic coupling calculations across different functional, and we conducted additional tests with the B3LYP and ω B97X-D functionals using same basis sets.

Reorganization energy calculations were conducted using Gaussian 16 at the LC-wHPBE/Def2-SV(P) level of theory. The calculation of the reorganization energy is one of the most sensitive parameters in modeling the kinetics of these systems, since the quantity is part of an exponential term for electron hopping rate calculation. Thus, we choose LC-ωHPBE which includes range-separated and dispersion-corrected method to capture the subtle changes that occur in the different charge states of the viologen molecules. A generic solvent was chosen with self-defined static and optical dielectric constants (41.78 and square of refractive index, 1.437 respectively) selected to correspond to the experimental results of GBL for the polarizable continuum model. The reorganization energy, λ can be expressed as the sum of inner and outer reorganization energy, λ_{in} and λ_{out} separately. With Nelsen's four-point method, λ_{in} can be estimated as following eqn (6):46

$$\lambda_{\rm in} = \left(E_{\rm neu}^{\rm redox} + E_{\rm redox}^{\rm neu} \right) - \left(E_{\rm neu}^{\rm neu} + E_{\rm redox}^{\rm redox} \right) \tag{6}$$

where *E* represents the total energy of the molecule under certain optimized geometry (subscript) and redox state (superscript). λ_{ou} can be further estimated from the solvent parameters, as originally given by Marcus following eqn (7):^{47,48}

$$\lambda_{\text{out}} = (\Delta e)^2 \left(\frac{1}{2r_{\text{D}}} + \frac{1}{2r_{\text{A}}} - \frac{1}{R_{\text{DA}}} \right) \left(\frac{1}{\epsilon_{\text{op}}} - \frac{1}{\epsilon_{\text{s}}} \right)$$
(7)

where Δe is the amount charge being transferred, $r_{\rm D}$, $r_{\rm A}$ are the donor and acceptor radii, respectively, $R_{\rm DA}$ is the donor/acceptor separation, and $\epsilon_{\rm op}$, $\epsilon_{\rm s}$ are the optical and static dielectric constant for GBL. The donor and acceptor radii, being same for the methyl viologen dimers, were estimated from the UFF radii given in Gaussian 16. $R_{\rm DA}$ was estimated by assuming it equals $r_{\rm D} + r_{\rm A}$, where $r_{\rm D} = r_{\rm A} = 4.08$ Å, at the closest approach.

Results and discussion

Synthesis and chemical characterization of viologen polypeptides and control viologen polymer

The three viologen polypeptides were synthesized through the post-polymerization modification of iodoalkyl-containing polypeptide precursors having C3, C6 or C9 alkyl chain lengths. The spacing between the polypeptide backbone and the viologen redox-active group was determined by this incorporation of alkyl chains of varying lengths within the precursors (Fig. 2a) to systematically study side chain effects. In addition to these polypeptides, poly(vinylbenzyl methyl viologen) (PV) was synthesized as a control (Fig. 2b) using post-polymerization modification of a chloro-containing poly(vinylbenzyl chloride) (PVBC) precursor. PV was selected as the control because it bears a hydrocarbon (non-peptide) backbone and because it has been extensively studied in the literature.^{35,49–51} By this design, the effect of having a peptide and non-peptide backbone can be compared.

ATR-FTIR spectroscopy, NMR spectroscopy and elemental analysis were used to confirm the chemical structures during the synthetic steps for the studied polymers (Fig. S1-S14 and Table S1[†]). The secondary structure of the viologen polymers was confirmed by ATR-FTIR and CD spectroscopies. Absorbance values around 1650 cm^{-1} (amide I) and 1547 cm^{-1} (amide II) in the FTIR spectrum (Fig. S15†), along with bands at 208 nm and 222 nm in the CD spectrum (Fig. S16[†]), suggested α-helical conformations for C6 and C9 polypeptides. In contrast, PV and C3 did not exhibit helical structures, as evident by ATR-FTIR or CD spectroscopies in either organic or aqueous solutions.52,53 From ¹H-NMR spectroscopy measurements and calculations, the molar mass for the precursor (prior to viologen installation) of C3 was 10.4 kDa, C6 was 12.4 kDa,16 C9 was 14.6 kDa, and PV was 9.8 kDa. From SEC, the number average molar mass (M_n) of the precursors were 10.6 kDa, 15.6 kDa,¹⁶ 12.4 kDa, and 11.0 kDa for C3, C6, C9, and PV, respectively (Fig. S17[†]). The

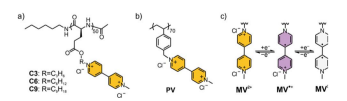


Fig. 2 The chemical structure of (a) viologen-based polypeptides with alkyl spacer lengths of three (C3), six (C6), or nine (C9) carbons. (b) The chemical structure of the control viologen (PV), bearing a non-peptide backbone. (c) The redox reaction of methyl viologen consisting of the dication (MV^{2+}), radical cation (MV^{++}), and the neutral (MV^{0}) states.

dispersities (*D*) for the three polypeptides were *ca.* 1.1, whereas *D* of PV was 1.25. All the viologen-functionalized polymers did not exhibit glass transitions or crystallization phenomena within the studied temperature range (Fig. S18†). From DLS, the hydrodynamic radii, $r_{\rm h}$, were determined to be 75 ± 3 nm, 51 ± 2 nm, 61 ± 3 nm, and 14 ± 1 nm for PV, C3, C6, and C9, respectively (Fig. S19†).

These relatively large $r_{\rm h}$ values with low molar masses, suggest that each polymer sample formed aggregates in the aqueous solutions used for DLS measurements. Given that the peptides each have the same degree of polymerization, it can be inferred that both C3 and C6 formed multimolecular aggregates of greater numbers of chains than did the C9 analog. Finally, the swelling ratio of each polymer film in 0.5 M LiCl in γ butyrolactone (GBL, the electrolyte used for electrochemical tests) was quantified using quartz crystal microbalance with dissipation monitoring (QCM-D), Fig. S20–S22.† The swelling ratios were 1.80, 1.22, 1.21, and 1.25 for PV, C3, C6, and C9, respectively (Table S2†). Additionally, no mass loss was observed for any of the polymers during the swelling period, suggesting that PV exhibited the greatest extent of swelling but no dissolution.

To understand how the side chain length affects the nature of the polypeptides' redox response, cyclic voltammetry (CV) was conducted using thin films on ITO glass (mass loadings: 0.25 to 0.40 mg cm⁻² and thicknesses: 200 to 275 nm), Fig. 3. For the CVs obtained at 10 mV s⁻¹, the three polypeptides exhibited two redox couples at similar potentials ($E_{1/2} = 2.12$ and 2.54 V vs. Li/Li⁺), consistent with that of PV and previous literature.³⁵

For C3 and C6, the MV^{2+}/MV^{*+} redox couple was more reversible with a peak separation (ΔE_p) of *ca*. 0.25 V as compared

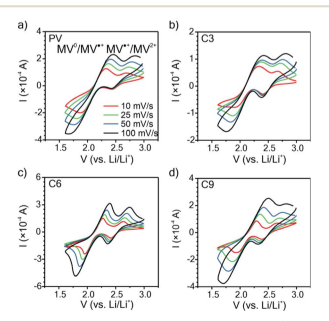


Fig. 3 Cyclic voltammograms at 10, 25, 50, and 100 mV s⁻¹for (a) PV, (b) C3, (c) C6, and (d) C9 thin films on ITO-coated glass as working electrodes in a three-electrode beaker cell configuration. The supporting electrolyte was 0.5 M LiCl in GBL. Lithium metal was used as counter and reference electrodes.

to 0.42 V *ca.* for the MV^{+/}/MV⁰ redox couple. C9 exhibited the same $\Delta E_{\rm p}$ of 0.32 V for both redox couples. PV exhibited a $\Delta E_{\rm p}$ consistent with C3 and C6. These results generally follow past literature for PV, for which MV⁺⁺/MV⁰ is less reversible than MV²⁺/MV^{+,54} All polymers exhibited diffusion-limited redox reactions, as indicated by the linear fit of peak current *vs.* square-root of scan rate (Fig. S23†).

Experimental determination of diffusion within the viologenbased polymer thin films

To elucidate the apparent diffusion coefficient (D_{ap}) for the polymer films, Cottrell plots from chronoamperometry (CA) were examined for oxidation and reduction and decoupled for MV^{2+}/MV^{*+} and MV^{*+}/MV^{0} , Fig. 4, S24 and S25.† In general, PV and all three polypeptides exhibited linear CA responses with similar slopes for the reduction step of MV^{2+}/MV^{*+} and the oxidation step of MV^{0}/MV^{*+} (Fig. 4a–c and S25a†), which is consistent with previous literature^{13,14} and is indicative of similar rates of electron transfer. In contrast, the slopes for the reduction step of MV^{*+}/MV^{0} differed and followed the trend of

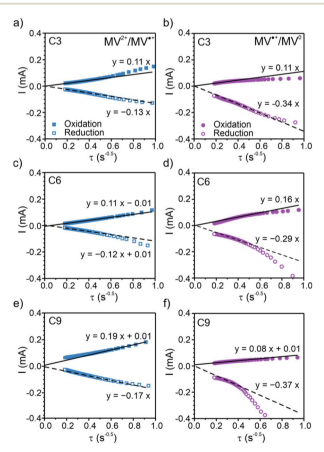


Fig. 4 Chronoamperometry (CA) Cottrell plots for the two redox couples of the three polypeptides. (a), (c), and (e) CA response for the MV^{2+}/MV^{+} redox couple. (b), (d), and (f) CA response for the MV^{+}/MV^{0} redox couple. Viologen polypeptides coated on ITO glass were used as the working electrodes in a three-electrode beaker cell configuration. The supporting electrolyte was 0.5 M LiCl in GBL. Lithium metal was used as counter and reference electrodes. The Cottrell plots for PV are found in Fig. S25.†

C6 < C3 < C9 \approx PV, which would suggest PV or C9 would exhibit the highest electron transfer rates.

The slope of the tangent lines from the Cottrell plots were used to estimate the apparent diffusion coefficient (D_{ap}) using eqn (2). The C_0^* value of each polymer repeat unit was determined to be 2.03 M for PV, 2.92 M for C3, 2.65 M for C6, and 2.42 M for C9, assuming a polymer film density of 1.2 g cm⁻³. Despite changing the alkyl spacer length, the site-to-site distance (δ) of the pendant viologen groups in all three polypeptides remained similar at 4 Å (estimated using MD simulations), which is consistent with previous reports for PV.^{13,14} The D_{ap} values were similar for all the polymers investigated and on the order of 10^{-13} cm² s⁻¹ (Fig. S26†), which is smaller by *ca*. 7 orders of magnitude than previous solution state reports (10^{-6} – 10^{-7} cm² s⁻¹).^{55–57} The low diffusion coefficients are hypothesized to be due to the polymer chains being more immobilized in the thin film state than in solution.

To evaluate the mechanism of electron transfer, the physical diffusion coefficient of the system must be compared to $D_{\rm ap}$. $D_{\rm ap}$ is the sum of the physical and electron diffusion coefficients. Two different methods were used to approximate physical diffusion of the viologen-based polymers within the thin films. The first method determined the solution-state diffusion coefficient of the polymer using DLS ($D_{\rm DLS}$) and the second used MD simulations to obtain the physical diffusion of the viologen pendant groups ($D_{\rm phys}$). DLS is one experimental method to measure the diffusion coefficient of a polymer chain, but it does not directly measure the diffusion of the redox-active pendant group, yet past literature assumes their equivalency.¹⁰ Therefore, MD trajectories were analyzed to provide a computational method to determine the redox-active group diffusion coefficient.

Initial MD simulations identified that the behavior of polypeptide oligomers converged at approximately 20 repeat units. Therefore, to minimize computational costs without sacrificing accuracy, oligomers with 26 repeat units were simulated (instead of the 50 repeat units studied experimentally). With these criteria, the diffusion coefficients of the redox-active methyl viologen groups (D_{phys}) were extracted from the trajectories of MD simulations of the oligomers in solvated states, taking GBL as the solvent. Table 1 shows the diffusion coefficients (D_{phys}) of the redox groups for the two different redox processes $(MV^{2+}/MV^{+} and MV^{+}/MV^{0})$ calculated using MD simulations. For comparison, the DLS-measured $r_{\rm h}$ of the polypeptides in the MV^{2+} form and eqn (1) were used to experimentally estimate the diffusion coefficient of the polymer in solution $(D_{\rm DLS})$.

Table 1 Experimentally determined D_{DLS} and calculated D_{phys} constant values (×10⁻⁷ cm² s⁻¹) from MD simulations for both redox couples of PV, C3, C6, and C9

	$D_{ m DLS}$	D _{phys}		
	MV^{2+}		MV^{*+}	MV ⁰
PV	0.83 ± 0.10	N/A	N/A	N/A
C3	1.2 ± 0.25	2.2 ± 0.9	3.0 ± 0.9	11.9 ± 3.1
C6	1.0 ± 0.12	1.3 ± 0.6	8.9 ± 3.0	11.1 ± 2.7
C9	$\textbf{4.6} \pm \textbf{0.74}$	4.0 ± 1.8	$\textbf{9.2}\pm\textbf{3.3}$	11.9 ± 3.5

The obtained D_{DLS} and D_{phys} values were on the order of 10^{-7} cm² s⁻¹ (Table 1), consistent with prior D_{ap} values for solution state viologen molecules.⁵⁵⁻⁵⁷ Notably, both approximations are solution-state, but the electrochemical testing conditions are solid-state, which may result in overestimates of D_{DLS} and D_{phys} . Previous studies on charged polymers and protein hydrogels suggest that the solution state diffusion coefficient is approximately 10 orders of magnitude larger than thin film diffusion (*ca.* 10^{-12} to 10^{-19} cm² s⁻¹).^{58,59} Using this approximation, the diffusion of the viologen polypeptides would be *ca.* 10^{-17} cm² s⁻¹. This value was smaller than the D_{ap} (10^{-13} cm² s⁻¹) suggesting Laviron–Andrieux–Savéant (LAS) theory is the appropriate mechanism of electron transport in the viologen polypeptides.²⁸

Experimental determination of rate constants k_{ex} and k^0

To elucidate how the side chain lengths affect the rate constants of electron transfer of the polypeptides, thin films were investigated experimentally using CA and CV. Specifically, D_{ap} from CA was used to calculate the rate constant for electron selfexchange (k_{ex}) and the peak separation from CV was used to calculate the heterogeneous rate constant (k^0) using the method of Nicholson.

Using eqn (4), the homogeneous electron transfer rate constant k_{ex} was determined and the values are summarized in Table S3[†] and Fig. 5. For all the polymers studied (independent of linker group), k_{ex} for the redox couple MV⁺/MV⁰ was greater than that for the redox couple of MV⁺/MV²⁺, except for C9 which exhibited a higher k_{ex} for oxidation of MV^{*+}/MV^{2+} versus MV⁰/MV^{*+}. This trend was also observed by Dalton et al. and Hatozaki et al. for an electrolyte-swollen viologen film, where k_{ex} for MV^{*+}/MV⁰ was larger than MV²⁺/MV^{*+}.^{13,60,61} The greatest variability in k_{ex} for the polymers was for oxidation of MV^0/MV^{*+} , where PV exhibited the largest value $(1.9 \times 10^3 \text{ M}^{-1} \text{ s}^{-1})$ and C9 the smallest (1.5 imes 10² M⁻¹ s⁻¹). The larger $k_{\rm ex}$ value for PV could be due to the increased mobility of the polymer's random coil backbone (compared to the stiffer polypeptide backbones) leading to enhanced dimerization and a higher rate of electron transfer. Further discussion of trends found in the k_{ex} and k^0 data can be found in the ESI (Fig. S27[†]).

One additional consideration for k_{ex} and the mechanism of electron transfer in the viologen polypeptides is the possibility of electron tunneling through the helical backbone of C6 or C9. According to LAS theory, k_{ex} should be equal to the activationlimited rate constant (k_{act}) determined from Marcus-Hush theory (eqn (S2), Fig. S28 and 29).† However, if electron tunneling occurs it would manifest in an enhanced experimentally determined k_{ex} comparable to k_{act} . Here, k_{act} values were estimated using the electronic coupling (H_{AB}) and reorganization energy (λ) determined from DFT calculations, which are described in detail in the ESI.[†] k_{act} was *ca.* 10⁹ M⁻¹ s⁻¹, which is significantly larger than the experimentally determined k_{ex} values for C6 and C9 (the helical polypeptides) and previously studied TEMPO-based monomer value (ca. 10^8 M^{-1} s⁻¹).¹⁰ This finding suggests that there is no significant electron tunneling in the viologen polypeptide thin films, which is

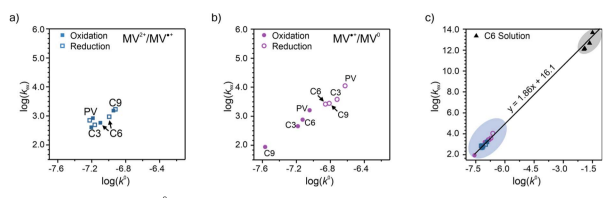


Fig. 5 Summary of the $log(k_{ex})$ and $log(k^0)$ for PV, C3, C6, and C9. The results are presented as separate experimental values of oxidation and reduction for the (a) MV^{2+}/MV^{+} and (b) MV^{++}/MV^{0} redox couples. (c) Plot showing all the $log(k_{ex})$ vs. $log(k^0)$ for experimental data for both redox couples. Panels (a) and (b) show the enlarged region containing the CV/CA-determined rate constants for the viologen polypeptides studied in this work. In panel (c), the data points in the blue circle are from polymer thin-films and the data points in the black circle are from solution state values for a 0.2 mM solution of C6 dissolved in 0.5 M NaCl in H₂O (raw CA and CV data can be found in Fig. S33†).

consistent with previous results for alanine and aminoisobutyric acid α -helical polypeptides with 48 repeat units (*ca.* $1.4 \times 10^{-13} \text{ M}^{-1} \text{ s}^{-1}$).¹⁹

Finally, the previously discussed $\Delta E_{\rm p}$ values from CV at 10 mV s⁻¹ were used to determine the heterogeneous rate constant (k^0) following eqn (5). The calculated k^0 values are summarized in Fig. 5a and b and are classified by redox couple, with the values for MV⁺⁺/MV⁰ being slightly smaller than MV²⁺/MV⁺⁺. The k^0 values are plotted along with the $k_{\rm ex}$ values, for which combined values lying in the upper right quadrant indicate faster overall kinetics. For the MV²⁺/MV⁺⁺ redox couple in Fig. 5a, no strong trend was observed among the polymers tested. For the MV⁺⁺/MV⁰ redox couple in Fig. 5b, PV demonstrated faster kinetics. Further discussion of trends found in the $k_{\rm ex}$ and k^0 data can be found in the ESI.[†]

Given the determined k_{ex} and k^0 values, the electron transport mechanism was further confirmed through evaluation of the slope of the relationship between $\log(k_{ex})$ and $\log(k^0)$ in Fig. 5c. Remarkably, the plot resulted in a linear trend even for polymers of different linker lengths, including the control, which suggests that these viologens transfer charge by a similar mechanism. The slope was determined to be 1.85 and was compared to the slopes from Marcus–Hush theory and the diffusion cooperative model. First, the relationship from Marcus *et al.* is presented in eqn (8), similar to the relation found in LAS theory:⁴⁷

$$\left(\frac{k_{\rm ex}}{A_{\rm ex}}\right)^{\frac{1}{2}} = \frac{k^0}{A_{\rm el}} \tag{8}$$

where A_{ex} and A_{el} are the pre-exponential factors for selfexchange and the electrode reaction. From this linear relationship, if electron transfer follows Marcus–Hush theory, it is expected that the slope for $\log(k_{\text{ex}})$ vs. $\log(k^0)$ is 2. From the diffusion cooperative model, a relationship between k_{ex} and k^0 is found when substituting for D_{phys} , eqn (9):¹⁰

$$\frac{k_{\text{ex,app}}}{16\pi a N_{\text{a}}} = \frac{k^0 2L}{3\kappa_{\text{el}} e^{\left(-\frac{\lambda}{4k_{\text{b}}T}\right)}}$$
(9)

When reorganized, the linear relationship is $\log(k_{ex,app}) = \log(k^0) + \log(b)$, where $b = 2L16\pi aN_a/3\kappa_{el}\exp(-\lambda/4k_bT)$. For the diffusion cooperative model, the expected slope from the $\log(k_{ex}) vs. \log(k^0)$ is 1. Given the slope analysis of 1.85, we conclude that the studied viologen polymers do not follow the diffusion cooperative model. Instead, the slope of 1.85 is closer to that of Marcus–Hush theory with diffusion limitations, which supports transport *via* LAS theory (Fig. 5).

Impact of transfer kinetics on electrode performance

An important consideration for the future design of RAPs is the relationship between electron transfer rates and the charge-discharge performance. To determine the capacity of the viologen polypeptides as thin films, galvanostatic charge-discharge (GCD) was utilized at a variety of C-rates in a half cell with lithium metal, where 1C is the current required to charge the battery in 1 hour. The theoretical capacities for the studied polymers were 149.2 mA h g⁻¹ for PV, 130.3 mA h g⁻¹ for C3, 118.3 mA h g⁻¹ for C6, and 108.0 mA h g⁻¹ for C9. For the half cells, all the polymers studied exhibited two peaks in the CV at 2.2–2.5 V and 1.7–2.1 V vs. Li/Li⁺ (Fig. S30†), which was similar to those observed in the three-electrode beaker cells (Fig. 3). In addition to CV, two characteristic plateaus at 2.15 and 2.55 V vs. Li/Li⁺ were observed in the GCD curves at 0.2C associated with the two redox couples (Fig. S31†).

Fig. 6a summarizes the discharge capacity at varying Crates for the viologen polypeptides and control viologen polymer. All the polymers exhibited capacities significantly lower than theoretical (*e.g.*, C6 exhibited the maximum at *ca.* 40% of theoretical), due to the lack of conductive additives and insulating backbones. These additives were not added to the thin film electrodes so as to isolate the response of the polymer alone. Additionally, all the polymers exhibited capacity fade and low coulombic efficiencies, likely due to dissolution of the polymer into the electrolyte (Fig. S32†) during cycling. For all the polymers, as the C-rate increased there was a decrease in capacity and the two plateaus of the galvanostatic charge–discharge curves (Fig. S31†) became less pronounced.

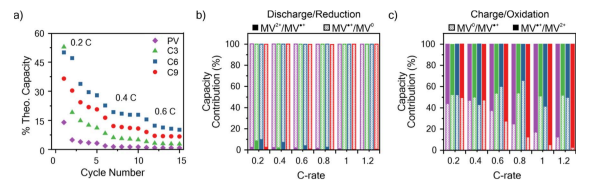


Fig. 6 (a) Discharge capacity summary for the polymers studied at varying C-rates. (b) The percent of total discharge (reduction) capacity from the MV^{2+}/MV^{+} and MV^{++}/MV^{0} redox couples at varying C-rates. (c) The percent of total charge (oxidation) capacity from the MV^{0}/MV^{+} and MV^{++}/MV^{2+} redox couples at varying C-rates. The data is taken from the first galvanostatic charge–discharge cycle at each C-rate in Fig. S31.† The colors corresponding to each polymer in panel (a) are applicable to panels (b) and (c). The working electrodes were tested in a half cell against lithium metal. The working electrode was polymer thin film on ITO-coated glass and 0.5 M LiCl in GBL on filter paper as the supporting electrolyte and separator, respectively.

At 0.2C, C6 exhibited the highest capacity of 1.28×10^{-2} mA h cm⁻², and PV exhibited the lowest (0.45 $\times 10^{-2}$ mA h cm⁻²). At C-rates smaller than 1C, C6 showed the best performance followed by C9, C3, and PV. It is surprising that C6 should show the best capacity, especially because its kinetics were slower than that of PV. However, capacity can also be influenced by other factors, such as swelling and ion diffusion. For example, we have shown previously that some swelling of the electrode is needed, but too much swelling can cut off interchain electron transfer pathways.^{62,63} Here PV, which exhibited the worst capacity at 1C, swelled the most.

The respective contributions of the MV^{+}/MV^{0} and $MV^{2+}/$ MV^{*+} redox couples were considered by identifying the inflection point in the second derivative of the charge-discharge curves (Fig. 6b, c and S31⁺). In discharge (reduction), MV⁺⁺/ MV⁰⁺ contributes almost entirely to the observed discharge capacity for all the polymers. However, in charge (oxidation), MV^{*+}/MV⁰⁺ contributes to only about half of the observed charge capacity for C-rates less than 1C for all polymers; above 1C, C9 and PV exhibited decreased contributions from MV⁺⁺/MV⁰⁺, whereas for C3 and C6 the contribution remains approximately 50% (Fig. 6c). These observed trends in capacity contribution for MV^{*+}/MV^{0+} mirror similar trends to those observed for k_{ex} . Specifically, k_{ex} for reduction of MV^{+/}/MV⁰ > reduction of MV^{2+/} MV^{*+} , meaning in reduction the contribution of MV^{*+}/MV^{0} is much higher than MV^{2+}/MV' . However, upon charging k_{ex} for oxidation of MV^{*+}/MV²⁺ is only slightly larger than oxidation of MV⁰/MV^{*+}, so the two redox couples contributed equally to the charge capacity.

Conclusions

The successful synthesis of viologen-containing polypeptides with varying linker lengths between the redox-active viologens and the polypeptide backbone was demonstrated and confirmed using a variety of characterization methods. Only the longer alkyl spacer lengths of C6 and C9 allowed for the polypeptide backbone to adopt an alpha-helical conformation. The

polymers were cast into thin films to study the electron transfer kinetics using cyclic voltammetry (CV) and chronoamperometry (CA). From CV studies of C3, C6, and PV, it was found that MV^{+/}/ MV^0 was less reversible than MV^{2+}/MV^{*+} , whereas for C9 the two redox couples exhibited the same reversibility. Additional investigation using Cottrell analysis of the CA data led to determination of D_{ap} and k_{ex} . k_{ex} followed the pattern of reduction of MV^{*+}/MV^{0} > reduction of MV^{2+}/MV^{*+} and oxidation of MV^{0}/MV^{*+} > oxidation of MV^{*+}/MV^{2+} , except of C9 which exhibited oxidation of MV^{+}/MV^{2+} > oxidation of MV^{0}/MV^{+} . The largest difference in k_{ex} for the studied polymers was for oxidation of MV^0/MV^{+} , where PV exhibited the highest k_{ex} and C9 exhibited the smallest. Upon analysis of the linear relationship between $log(k_{ex})$ and $log(k^0)$, we conclude redox mechanism follows Marcus-Hush theory of electron transfer with diffusion limitations, supporting electron transport by LAS. Finally, galvanostatic-charge discharge of lithium metal half cells was used to evaluate the capacity performance. C6 showed highest capacity at C-rates less than 1C.

However, additional studies are needed to confirm structure-property relationships for anodic RAPs. In varying the length of the linker group, we accessed both coil (PV, C3) and helix (C6, C9) conformations, but neither conformation appears to have strong trends in affecting the kinetics or capacities. It is possibly a product of using the viologen moiety, which can exhibit dimerization, stiffening the polymer through the redox units instead of the backbone. Further, swelling may influence capacity by influencing intermolecular electron transfer. Although swelling is needed for improving the access of ions into the electrode, too much swelling (as with PV) can lead to poor capacity. Future work should consider utilizing several different redox-active moieties on helical backbones to further resolve backbone effects. Also, swelling should be closely monitored. Last, composite electrodes with conductive additives will be needed to create practical organic battery electrodes with higher thicknesses. Besides these future works, the polypeptide RAPs remain interesting due to their on-command degradation and bio-inspired structure.16

Data availability

The data supporting this article have been included in the main text and as part of the ESI.[†]

Author contributions

A. D. E., T. P. N., J. L. L., and K. L. W. developed the study. T. P. N. and S.-G. L. synthesized and structurally characterized the viologen polypeptides. A. D. E., S.-G. L., and K.-H. M. K. synthesized and structurally characterized the control viologen polymers. A. D. E. performed the electrochemical characterization of the materials. A. D. E. and S.-G. L. conducted thermal characterization. C.-H. L. and D. P. T. conducted all DFT calculations and MD simulations. The manuscript was written by A. D. E., with help from S.-G. L., C.-H. L., T. P. N., K.-H. M. K., D. P. T, J. L. L. and K. L. W.

Conflicts of interest

There are no conflicts to declare.

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