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Parallel Variation of Mass Transport and Heterogeneous and Homogeneous Electron Transfer Rates in Hybrid Redox Polyether Molten Salts

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Metal complexes can be prepared as highly viscous (semisolid), room temperature molten salts by combining them with oligomeric polyether substituents. The fluidity and transport properties of these hybrid redox polyether melts can be systematically manipulated by changing the oligomeric chain lengths and by adding unattached oligomers as plasticizers. This paper describes the voltammetrically measured transport properties of several Co(II) polypyridine (2,2'-bipyridine, phenanthroline) melts. The properties evaluated are the physical self-diffusion coefficient (D_{PHYS}) of the cationic complex in its melt, the diffusivity of its counterion (D_{CION}), the heterogeneous electron-transfer rate constant (k_{HET}) of the Co(III/II) oxidation at the electrode surface, and the rate constant (k_{EX}) for homogeneous electron self-exchange between Co(II) and Co(I) in the mixed valent layer next to the electrode. These dynamics parameters change in parallel manners, over a large ($>10^3$) range of values, when the melt fluidity is changed by plasticizers or temperature. While k_{HET} and k_{EX} both change systematically with D_{PHYS} , they change on a more nearly proportional basis with D_{CION} . The latter relationship is interpreted as a kind of solvent dynamics control in which both the homogeneous Co(II/I) and heterogeneous Co(III/II) reaction rates are controlled by the ionic atmosphere relaxation time constant, namely, the time constant of redistribution of counterions following an electron-transfer step that has produced a nonequilibrium charge distribution. D_{CION} provides a measure of the ion atmosphere relaxation rate.

Introduction

When polyether oligomers are covalently bonded to normally crystalline redox active compounds, or are bonded to their counterions, the result is a highly viscous, amorphous, room-temperature molten salt or molecular melt.¹ Such redox polyether hybrid materials, in their neat state, can exhibit large viscosities (as large as 10^6 cP at 25 °C)^{1b} and are model semisolids. The melts consist of hard cores of redox moieties surrounded by soft, deformable polyether shells.^{1c} Their viscosities are large because the redox moiety contributes little to the material's free volume, which increases and viscosity decreases as the molar volume fraction of the redox polyether hybrid material that is polyether oligomer increases.^{1a} The number and length of the oligomer chains determine the latter and greatly affect both mass and charge transport.^{1c,2} Added, unattached polyether oligomer also enhances transport properties, as does sorption from a liquid CO₂ phase;³ this is called diffusion plasticization, or just plasticization.⁴

This report describes mass transport and Co(III/II) and Co(II/I) electron-transfer dynamics in a series of molten salts of Co polypyridine metal complexes (structures in Figure 1). Melt **I** is a Co phenanthroline (phen) complex, in which a MePEG₃₅₀ oligomer (methyl-terminated polyethylene glycol, average MW = 350) has been covalently bonded to its sulfonate counterion (MePEG₃₅₀SO₃⁻). This [Co(phen)₃](MePEG₃₅₀-SO₃)₂ material (**I**) is plasticized in two ways, by adding unattached, neutral MePEG₃₅₀ oligomer and by adding an ammonium salt of the MePEG₃₅₀ tailed counterion, (NH₄)(MePEG₃₅₀SO₃). Melt **II_n** in Figure 1 is a bipyridine complex in which the bpy ligands

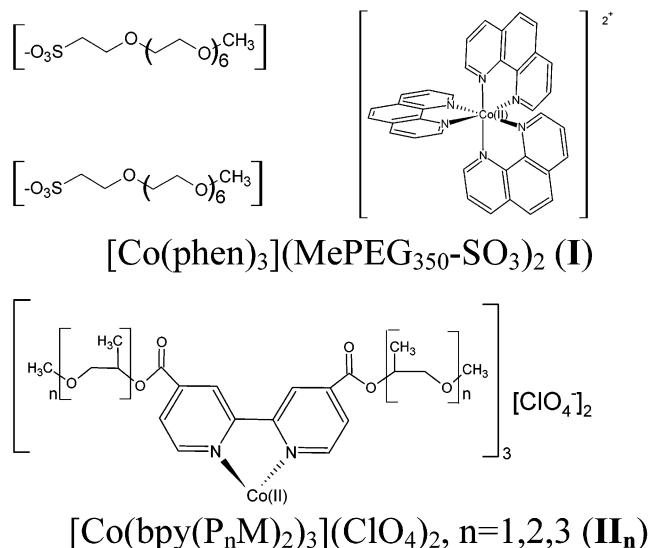


Figure 1. Structure of cobalt trisphenanthroline counterion tailed melt ([Co(phen)₃](MePEG₃₅₀SO₃)₂) (**I**) and of cobalt trisbipyridine perchlorate melt ([Co(bpy(P_nM)₂)₃](ClO₄)₂) (**II_n**) where $n = 2$ or 3.

bear attached (P_nM) methyl-terminated polypropylene oxide oligomers having either 2 or 3 repeat units.^{2b} These perchlorate complexes ([Co(bpy(P_nM)₂)₃](ClO₄)₂) can exhibit exceptionally low fluidity owing to the small number of propylene oxide repeat units; in practice, the fluidity of a given melt **II_n** also varies widely as a result of minor variations in the stoichiometry of the prepared complex (analytically established as having either excess P₃M or P₂M ligand or excess Co ions) and by adventitious traces of adsorbed moisture. The **II_n** melts do not crystallize owing to the diastereomeric-mixture character of the

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propylene oxide chains; their ethylene oxide analogues slowly crystallize over weeks of standing.

In this study, we describe the electron-transfer rate constants for homogeneous Co(II/I) electron self-exchanges (k_{EX}) and for heterogeneous Co(III/II) electron transfers (k_{HET}), and the self-diffusion coefficients of the Co complexes (D_{PHYS}) and of their counterions (D_{CION}), in the neat and plasticized melts. Co polypyridine complexes are convenient for this purpose because, as first demonstrated⁵ by Buttry and Anson, the Co(II) physical diffusion rate (D_{PHYS}) can be measured voltammetrically from oxidation currents of the Co(II) complex. Similarly, the Co(II/I) electron self-exchange rates can be measured voltammetrically from the enhanced apparent diffusivity (D_{APP}) upon reducing the Co(II) complex. The Co(II) apparent diffusivity is enhanced by the rapid electron hopping within the Co(II/I) mixed valent diffusion layer generated around the electrode; the contribution of hopping, or “electron diffusion” (D_{E}) to D_{APP} can be estimated using the cubic lattice model⁶

$$D_{\text{APP}} = D_{\text{PHYS}} + D_{\text{E}} = D_{\text{PHYS}} + k_{\text{EX}} \delta^2 C / 6 \quad (1)$$

where k_{EX} is the homogeneous Co(II/I) electron self-exchange rate constant, δ the equilibrium center–center distance between complexes,⁷ and C the total concentration of cobalt complex sites in the melt.

The Co(III/II) electron self-exchange is quite slow,^{8a} so the D_{APP} of Co(II) oxidation measures solely D_{PHYS} . The Co(III/II) reaction rate (k_{HET}) is measured as a heterogeneous reaction since, in the voltammetry, the Co(III/II) electrode reaction is electrochemically quasireversible. The counterion transport rate (D_{CION}) is measured from a combination of D_{PHYS} and the melt ionic conductivity, using the Nernst–Einstein relation. The D_{CION} results are supported by direct voltammetry of iodide as a model counterion. By varying the added plasticizers and the melt temperature, a roughly 10^3 range of the above dynamics constants was achieved in melts of **I** and **II_n**.

The motivation of the above measurements is to understand why electron-transfer rates in semisolids such as **I** and **II_n**, and in a number of other redox polyether hybrids, are generally slower and exhibit larger thermal activation energy barriers^{1b,c,d} than predicted by Marcus theory⁹ or observed experimentally for analogous reactions in dilute, low-viscosity fluid solutions. We reported previously that, over a huge (10^{11} -fold) dynamics range, (a) the Co(III/II) k_{HET} varied inversely with the longitudinal relaxation time in a series of Debye solvents^{10,11} and b) in a series of polyether solutions and redox hybrid melts k_{HET} varied systematically with D_{PHYS} of the Co complex.^{1e} Others¹² have also seen correlations (albeit over more narrow ranges) between k_{HET} and physical diffusion rates of metal complexes. These correlations have been interpreted as control of the electron transfer rate by the dynamics of solvent repolarization upon charge transfer. Solvent repolarization rates are presumed to analogously dictate mass transport dynamics. In the case of diffusants such as **I** and **II_n**, the “solvent” is the attached shell of polyether oligomer chains.

Analogous correlations have also been observed for homogeneous Co(II/I) k_{EX} rate constants. Recent measurements^{3b} in Co complex melts such as **I**, plasticized by sorption from a liquid CO₂ bath, revealed a strong correlation between k_{EX} and D_{PHYS} of the complex, suggesting that the Co(II/I) reaction was dominated by polyether dipole repolarization kinetics.¹³ That study also, however, revealed an even stronger (proportional) correlation between k_{EX} and the diffusivity (D_{CION}) of the melt counterion. The possibility of a different kind of solvent-based rate control of electron transfer was raised and termed ion

atmosphere relaxation. In the present study, we set out to assess the relation of both k_{EX} and k_{HET} to both D_{PHYS} and D_{CION} , in several redox polyether hybrid melts. The observations for both Co(III/II) and Co(II/I) reactions show that their rates vary in a 1:1 proportionality with D_{CION} , consistent with an ion atmosphere relaxation control.

Experimental Section

Synthesis of (Na)(MePEG₃₅₀SO₃). This was prepared by a previous¹⁴ procedure, modified as described in Supporting Information.

Synthesis of [Co(phen)₃]Cl₂. CoCl₂·6H₂O dissolved in ethanol was added dropwise to an excess of sublimed phenanthroline in ethanol while stirring; the phenanthroline solution changed from clear to brown. After the solution was stirred thoroughly, it was concentrated through rotary evaporation, heated, then cooled to 5 °C. Crystals were collected and rinsed with cold ethanol. Purity was checked with UV–vis spectroscopy.

Synthesis of [Co(phen)₃](MePEG₃₅₀SO₃)₂. A published literature procedure was used.^{3b} Anal. Calcd C, 54.3; H, 6.00; N, 5.76; S, 4.40; Co, 4.04; Na, 0; Cl, 0. Found (average of 5 determs.): C, 53.2(±2); H, 6.38(±0.2); N, 5.71(±0.7); S, 4.45(±0.3); Co, 4.22(±0.7); Na, 0.296(±0.6); Cl, 0.293(±0.2).

Synthesis of (NH₄)(MePEG₃₅₀SO₃). An aqueous solution of ~300 mg (Na)(MePEG₃₅₀–SO₃) (0.7 mmol) was passed through a cation exchange column in the H⁺ form (pretreated with 4 M HCl), the eluent dripping into 17.2 mL (0.0687 mol) of ammonium hydroxide chilled in an ice bath. The resulting solution, still in the ice bath, was stirred for 30 min, and then for 30 min at room temperature. Water and excess ammonia were removed by rotary evaporation and with a vacuum pump on a Schlenk line. The product was dissolved in methylene chloride, filtered through 0.2-μm Nalgene PTFE syringe filters, and dried under vacuum.

Plasticization of [Co(phen)₃](MePEG₃₅₀SO₃)₂. Weighed amounts of MePEG₃₅₀ and (NH₄)(MePEG₃₅₀SO₃) plasticizers were added as described in Supporting Information. The quantities of plasticizer added are stated as moles plasticizer/ moles Co.

Synthesis of [Co(P_nM)₃](ClO₄)₂ ($n = 2$ or 3), **II_n.** The **II_n** propylene oxide oligomer-tailed complexes were prepared as previously described.² In the preparation of [Co(P₃M)₃](ClO₄)₂ (**II₃**), a stoichiometric excess of the tailed bpy ligand was mixed with a Co(II) salt. The subsequent cleanup steps were only partly successful in removing the excess ligand, and the several preparations contained varied, residual amounts of tailed ligand (analyzed by nmr), which acts as plasticizer.² In one case (Table 3), a larger amount of tailed bpy ligand was deliberately added after cleanup.

Electrochemical Measurements. Potential step chronoamperometry of the Co(III/II) and Co(II/I) electrochemical reactions (for D_{PHYS} and D_{APP} values) and cyclic voltammetry of the Co(III/II) reaction (for k_{HET}) were conducted on neat films of the redox polyether hybrids resting on microelectrode assemblies, as before.¹ For measurements of [Co(phen)₃](MePEG₃₅₀SO₃)₂ (**I**), the melt rested on an epoxy disk with exposed electrode wire tips, Pt (radius = 15 μm) microdisk working electrode, Pt counter electrode, and Ag quasireference electrode, as done before.¹ See Supporting Information for further details. The working electrode in studies of [Co(P_nM)₃](ClO₄)₂ (**II_n**) melts were either Pt microdisks (radius = 5, 13, 50, or 95 μm) as above or Pt microbands,^{2,15} fashioned in one case by exposing the edge of a 2.7-μm Pt foil (area 3.1×10^{-3}

TABLE 1: Physical Dynamics and Electron-Transfer Results for [Co(phen)₃](MePEG₃₅₀SO₃)₂ with Added MePEG₃₅₀ Plasticizer

moles MePEG ₃₅₀ /Co complex	0.65:1	2:1	3:1	6:1 ^a
concentration (M) ^b	0.74	0.57	0.47	0.33
δ (Å) ^c	13.1	14.3	15.2	17.1
D_{PHYS}^d (25 °C) (cm ² /s)	5.3×10^{-12}	1.3×10^{-10}	4.3×10^{-10}	4.4×10^{-9}
D_{CION}^e (25 °C) (cm ² /s)	na ^m	1.9×10^{-9n}	3.3×10^{-9}	2.0×10^{-8}
D_{APP}^f (25 °C) (cm ² /s)	na	3.4×10^{-9}	7.9×10^{-9}	4.5×10^{-8}
D_{E}^g (25 °C) Co ^{III} (cm ² /s)	na	3.3×10^{-9}	7.5×10^{-9}	4.1×10^{-8}
$E_{\text{A,PHYS}}^h$ (kJ/mol)	na	60	54	43
$E_{\text{A,ION}}^i$ (kJ/mol)	na	46	39	32
$E_{\text{A,EX}}^j$ (kJ/mol)	na	47	41	30
k_{EX}^k (25 °C) (M ⁻¹ s ⁻¹)	na	1.7×10^6	4.1×10^6	2.5×10^7
k_{EX}^l (M ⁻¹ s ⁻¹)	na	3.4×10^{14}	5.0×10^{13}	4.4×10^{12}
k_{HET}^f (25 °C) (cm/s)	1.0×10^{-7}	1.1×10^{-6}	1.5×10^{-6}	na

^a Data presented for 6:1 mol MePEG₃₅₀ is at 30 °C. D_{PHYS} is calculated from microdisk equation.¹⁹ ^b Concentrations are calculated from density measurements: $\rho = 1.24$ g/mL for 0.65:1 MePEG₃₅₀; 1.23 g/mL for 2:1 MePEG₃₅₀; 1.18 g/mL for 3:1 MePEG₃₅₀; 1.18 g/mL for 6:1 MePEG₃₅₀. ^c δ is redox center–center distance calculated from the density assuming cubic packing. ^d From Co(III/II) reaction, Cottrell slope chronoamperometry.¹⁸ Each value is an average of 2 or 3 trials. ^e Calculated from eq 2. Each value is an average of 2 or 3 trials. ^f From microdisk equation.¹⁹ ^g Calculated via eq 1. Each value is an average of 2 or 3 trials. ^h From slopes of activation plots in Figure 4. ⁱ From slopes of activation plots in Supporting Information. ^j Calculated via eq 1. ^k Intercepts of activation plots of k_{EX} (Figure 4). ^l Average values given: 5 trials for 3:1 mol, 10 trials for 2:1 mol, 1 trial for 0.65:1 mol. ^m Data not available. ⁿ Values for 2:1 mol calculated assuming 1:1 ion pairing between cobalt cation and sulfonate anion (ref 21).

cm²) and in another being a lithographically defined¹⁶ microband (“LDM”, 10 μ m wide, $A = 2.0 \times 10^{-4}$ cm²). The microband electrodes were flanked on both sides by microband reference

electrodes, 2–3 μ m away, an arrangement making these electrodes exceptionally robust against uncompensated resistance effects.² Films of the cobalt complex melts (ca. 1 mm thick) were cast onto the microelectrode platforms and thoroughly dried under vacuum (ca. 1×10^{-3} Torr) in a temperature-controlled cell enclosure at 70 °C for at least 12 h. Films of the melts were equilibrated at each temperature for at least 1 h prior to measurements. The temperature of the cell enclosure was controlled as described in Supporting Information.

Cyclic voltammetry and chronoamperometry were performed using a locally built low-current potentiostat.¹⁷ Experimental control was exercised with a PC interfaced with a Keithley DAS-HRES 16-Bit A/D board, using locally written software. Ionic conductivities of the melts were measured using a Solartron Model SI 1260 impedance/gain phase analyzer, SI 1287 electrochemical interface combination. Impedance measurements from 1 MHz to 1 Hz were performed at 0 V dc bias and 10–50 mV ac amplitude. Conductivity was calculated as the product of the geometric cell constant (22.5 cm⁻¹ for microdisk measurements of melt **I** and 0.0848 cm⁻¹ for LDM measurements of melts **II_n**) with resistance taken from the low-frequency real-axis intercept of the complex impedance semicircle.

Results and Discussion

Mass Transport and Electron-Transfer Rate Measurements. The effect of adding MePEG₃₅₀ as plasticizer on the microdisk cyclic voltammetry of melt **I**, [Co(phen)₃](MePEG₃₅₀SO₃)₂, at 70 °C is shown in Figure 2. The added plasticizer causes increases in currents for both Co(II) oxidation (at positive potentials) and Co(II) reduction (at negative potentials). The general waveshape reflects a mixed radial–

TABLE 2: Physical Dynamics and Electron-Transfer Results for [Co(phen)₃](MePEG₃₅₀SO₃)₂ with Added (MePEG₃₅₀SO₃)(NH₄) Plasticizer

moles plasticizer/Co	0	1:1	1.5:1	2:1 ^f	3:1 ^f	4:1 ^f	5:1
conc. (M) ^a	0.74	0.68	0.59	0.58	0.50	0.42	0.34
density (g/mL)	na ^e	1.30	1.26	na	na	na	1.24
δ (Å) ^b	na	13.5	14.1	na	na	na	17.0
D_{PHYS}^c (25 °C) (cm ² /s)	7.1×10^{-13}	5.5×10^{-12}	7.1×10^{-12}	1.0×10^{-11}	2.3×10^{-11}	4.8×10^{-11}	1.6×10^{-11}
k_{HET}^d (25 °C) (cm/s)	3.6×10^{-9}	1.1×10^{-7}	1.1×10^{-7}	1.7×10^{-7}	3.0×10^{-7}	3.2×10^{-7}	1.3×10^{-7}

^a Concentrations are calculated from density measurements. Calculated as (moles of melt + moles of plasticizer)/(liters of melt + liters of plasticizer). ^b δ is redox center–center distance taken from the molecular volume assuming packed cubes. ^c From Co(III/II) reaction, Cottrell slope chronoamperometry.¹⁸ Values are averages from 2–4 trials except for 0 mol that is a single trial value. ^d From Nicholson-Shain analysis.²² Each value is an average from 2 to 4 trials except for the 1.5 and 4 trials which are only one. See Supporting Information for full scan rate dependent data. ^e Data not available. ^f Concentrations of 2:1, 3:1, and 4:1 (MePEG₃₅₀SO₃)(NH₄) were extrapolated from a plot of concentration vs moles (MePEG₃₅₀SO₃)(NH₄):Co because insufficient material was available for density measurements of those samples.

TABLE 3: Physical Dynamics and Electron-Transfer Results for Co(bpy(P_nM)₂)(ClO₄)₂

melt	density g/cm ³	conc M ^a	δ Å ^b	mol % plast. ^c	trial no.	D_{PHYS} 25 °C cm ² /s	D_{CION} 25 °C cm ² /s	k_{HET} 25 °C cm/s	$E_{\text{A,PHYS}}$ kJ/mol	$E_{\text{A,CION}}$ kJ/mol	$E_{\text{A,HET}}$ kJ/mol	k_{HET}^d cm/s			
Co(P ₃ M) ₃	1.21	0.57	14.3	<1	1	4.9×10^{-16}	na ^d	2.0×10^{-10}	na	na	na	na			
					5.1	1	3.2×10^{-15}	na	2.5×10^{-9}	na	na	na	na		
				36	5	1	2	2.0×10^{-14}	na	3.8×10^{-9}	na	na	na	na	na
							3	6.4×10^{-14}	na	3.1×10^{-9}	na	na	na	na	
							1	6.1×10^{-12}	na	5.1×10^{-8}	na	na	na		
							2	1.6×10^{-14}	4.3×10^{-12}	2.4×10^{-9}	108 (± 2)	66 (± 2)	69 (± 1)	$3.4 (\pm 0.1) \times 10^3$	
							3	7.1×10^{-15}	4.3×10^{-12}	1.6×10^{-8}	130 (± 4)	64 (± 3)	$3.4 (\pm 0.1) \times 10^3$		
							4	1.5×10^{-14}	4.3×10^{-12}	1.3×10^{-9}	99 (± 16)	72 (± 12)	$9.2 (\pm 4.3) \times 10^3$		
							5	na	na	na	104 (± 4)	65 (± 4)	$2.0 (\pm 0.4) \times 10^3$		
							6	2.3×10^{-15}	4.4×10^{-12}	6.7×10^{-9}	76 (± 13)	e	e		
Co(P ₂ M) ₃	1.26	0.71	13.3	1	1	2.0×10^{-17}	1.2×10^{-13}	2.7×10^{-11}	135 (± 7)	133 (± 2)	e	e			
					8	1	3.2×10^{-15}	na	1.1×10^{-10}	e	na	66 (± 9)	$4.4 (\pm 3.9) \times 10^1$		

^a Concentrations are calculated from density measurements. ^b δ is redox center–center distance taken from the molecular volume assuming packed cubes, using the given values for density. ^c The plasticizer used is 2,2'-bipyridyl-4,4'-bis(tripropylene glycol monomethyl ether carboxylate) (P₃M) for all Co(P₃M)₃ samples except for the 5% sample, which is plasticized with Co(ClO₄)₂(H₂O)₆. The plasticizer used for the Co(P₂M)₃ samples is 2,2'-bipyridyl-4,4'-bis(dipropylene glycol monomethyl ether carboxylate) (P₂M). ^d Data not available. ^e Outlier results not reported.

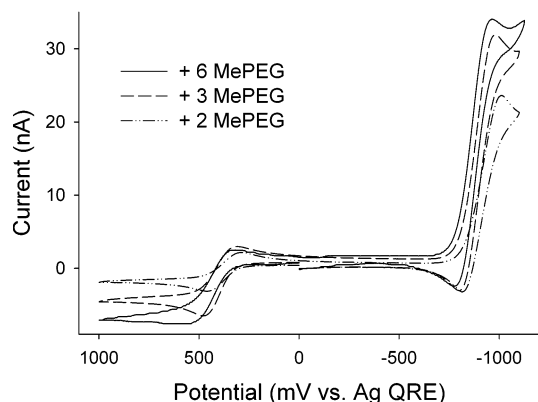


Figure 2. Cyclic voltammetry at 5 mV/s and 70 °C of $[\text{Co}(\text{phen})_3]\text{-(MePEG}_{350}\text{SO}_3)_2$ melt at a 15- μm radius Pt microdisk electrode with 6:1, 3:1, and 2:1 mol plasticizer/moles Co complex. k_{EX} is 2.5×10^7 , 4.1×10^6 , and $1.7 \times 10^6 \text{ M}^{-1} \text{ s}^{-1}$, and D_{PHYS} is 4.4×10^{-9} , 4.3×10^{-10} , and $1.3 \times 10^{-10} \text{ cm}^2/\text{s}$ respectively for +6, +3, and +2 mol added plasticizer.

linear diffusion geometry and quasireversibility for the Co(III/II) reaction. Figure 3 shows further examples of Co(III/II) voltammetry, for **I** plasticized by different amounts of $(\text{NH}_4)(\text{MePEG}_{350}\text{SO}_3)$ (Figure 3A) and for **II**₂, $[\text{Co}(\text{P}_2\text{M})_3](\text{ClO}_4)_2$, at various temperatures (Figure 3B). (The Co(II/I) electron-transfer reaction was not part of the experiments on **II**_n nor was it studied when **I** was plasticized with $(\text{NH}_4)(\text{MePEG}_{350}\text{SO}_3)$, owing to excessive background currents caused by this plasticizer at negative potentials.)

Physical diffusion coefficients (D_{PHYS}) of the Co(II) complexes in the melts were measured with potential step chronoamperometry, stepping from the double layer region in the middle of the Figure 2 voltammogram to the diffusion-controlled

plateau of the Co(III/II) wave. Usually, currents could be measured at times sufficiently short to maintain a linear diffusion geometry, and D_{PHYS} was obtained from the slopes of Cottrell plots (current vs $t^{-1/2}$).¹⁸ Results for D_{PHYS} values at 25 °C in melt **I** are given in Table 1 for MePEG_{350} plasticizer and in Supporting Information for other temperatures and in Table 2 for melts containing the $(\text{NH}_4)(\text{MePEG}_{350}\text{SO}_3)$ plasticizer. Table 3 presents D_{PHYS} values at 25 °C for **II**₂ and **II**₃ melts with various amounts of plasticizer and in Supporting Information for other temperatures (the sole data point obtained for **II**₁ is included in Supporting Information).

Apparent diffusion coefficients (D_{APP}) for the reduction of Co(II) to Co(I) in **I**, plasticized with MePEG_{350} , were obtained chronoamperometrically by stepping the electrode potential to the diffusion-controlled plateau of the Co(II/I) wave. Charge transport in this case is faster because of Co(II/I) electron hopping, and after ca. 10^3 s , the currents are steady state and controlled by radial diffusion. Diffusion coefficients (D_{APP}) obtained from such currents with the microdisk equation¹⁹ are found in Table 1. Combining D_{APP} with D_{PHYS} (eq 1) gives D_{E} and electron self-exchange rate constant k_{EX} values, also given in Table 1.

The diffusion coefficients of the ClO_4^- and $\text{MePEG}_{350}\text{SO}_3^-$ counterions were evaluated indirectly, based on measured ionic conductivities (σ_{ION}) and the D_{PHYS} of the Co complexes, using the Nernst–Einstein relation²⁰

$$\sigma_{\text{ION}} = F^2/RT[z_{\text{Co}}^2 D_{\text{PHYS}} C_{\text{Co}} + z_{\text{ClON}}^2 D_{\text{ClON}} C_{\text{ClON}}] \quad (2)$$

where z , D , and C are charge, diffusion coefficient, and concentration of the indicated species, respectively. Table 1 gives the resulting D_{ClON} values for the $\text{MePEG}_{350}\text{SO}_3^-$ counterion at 25 °C in the melt **I**; Table 3 gives 25 °C D_{ClON} values

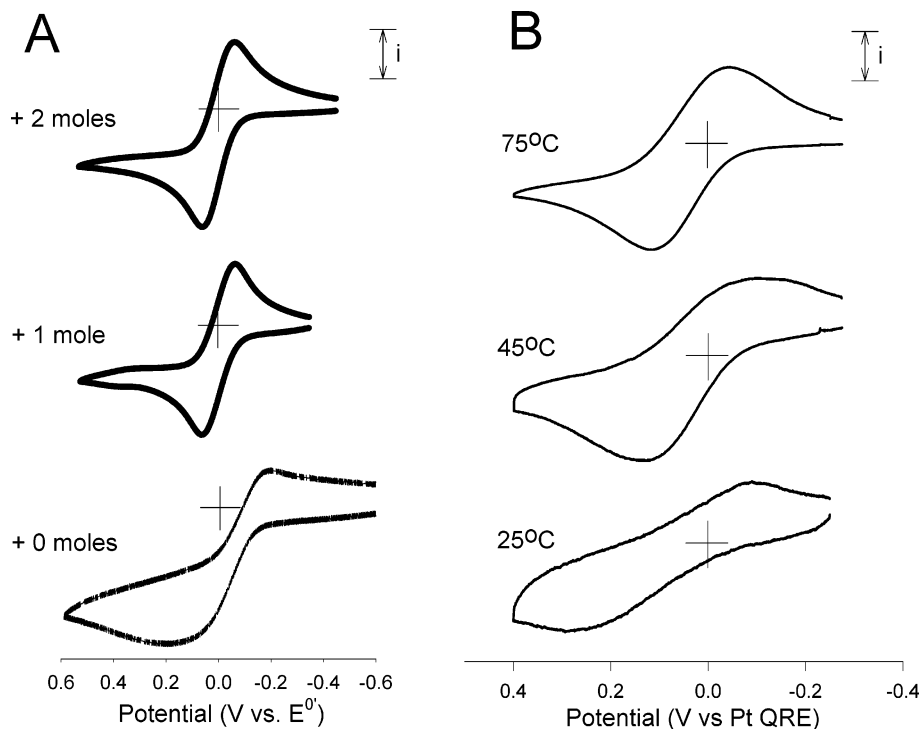


Figure 3. (A) Cyclic voltammetry at 25 °C and at 0.2 mV/s of $[\text{Co}(\text{phen})_3](\text{MePEG}_{350}\text{SO}_3)_2$ at a 15- μm radius Pt microdisk electrode. Added +2, +1, and 0 mol of $(\text{MePEG}_{350}\text{SO}_3)(\text{NH}_4)$ plasticizer (relative to Co) are indicated on plots, for which the current bars are 24, 18, and 3 pA, respectively. $k_{\text{HET}} = 3.6 \times 10^{-9}$, 1.1×10^{-7} , and $1.7 \times 10^{-7} \text{ cm/s}$, and $D_{\text{PHYS}} = 7.1 \times 10^{-13}$, 5.5×10^{-12} , and $1.0 \times 10^{-11} \text{ cm}^2/\text{s}$, respectively. (B) Cyclic voltammetry at indicated temperatures of $[\text{Co}(\text{bpy}(\text{P}_2\text{M})_2)_3](\text{ClO}_4)_2$ at a $2.0 \times 10^{-4} \text{ cm}^2$ area microband (LDM). Current bars are 200, 8, and 2 pA, potential scan rates are 5, 1, and 0.5 mV/s, $k_{\text{HET}} = 5.3 \times 10^{-8}$, 1.6×10^{-9} , and $2.7 \times 10^{-11} \text{ cm/s}$, and $D_{\text{PHYS}} = 5.8 \times 10^{-14}$, 8.2×10^{-16} , and $2.0 \times 10^{-17} \text{ cm}^2/\text{s}$ for the 75, 45, and 25 °C trials, respectively. See Supporting Information for scan rate dependent data.

TABLE 4: Counterion Transport Measured in [Ru(bpy(CO₂MePEG₃₅₀)₂)₃][ClO₄]₂ Containing 30 mol % of [Ru(bpy(CO₂MePEG₃₅₀)₂)₃][I]₂ Using Ionic Conductivity and Iodide Oxidation Voltammetry

<i>T</i> °C	<i>D</i> _{IODIDE} × 10 ⁹ cm ² /s ^a	<i>σ</i> _{ION} × 10 ⁶ mhos ⁻¹ cm ⁻¹ b	<i>D</i> _{ClON} × 10 ⁹ cm ² /s ^c
45.0	1.3	4.7	1.5
50.0	1.9	6.1	2.0
55.0	2.9	7.7	2.6
60.0	3.9	9.7	3.3
65.0	5.2	12	4.2

^a Diffusion measurements were based on chronoamperometry. ^b From AC impedance measurement. ^c Calculated from eq 2 assuming that *D*_{PHYS} of the Ru complex is negligible.

for the ClO₄⁻ counterion in the melt **II_n**. Supporting Information presents *D*_{ClON} results at other temperatures. As discussed previously,^{1d} any ion-pairing interactions present would depress the apparent *D*_{ClON} values, so they represent lower limits. The effect of ion pairing should not be large; assuming complete formation of a 1:1 ion pair for example increases the calculated *D*_{ClON} by only a factor of ~3-fold.²¹ Migration effects were also minor.²¹

To verify the indirect *D*_{ClON} measurements, we conducted a set of direct voltammetric determinations of the diffusion coefficient of another small counterion, iodide, in a closely related Ru complex melt, [Ru(bpy(MePEG₃₅₀)₂)₃](X)₂ where the MePEG₃₅₀ oligomer tail was attached to bipyridine ligands and X⁻ is a mixture of ClO₄⁻ and (30% mol percent) I⁻. Iodide is oxidized to tri-iodide in the voltammetry. The average *D*_{ClON} was also determined from ionic conductivity of the melts, using eq 2. Table 4 shows that the directly measured *D*_{IODIDE} and the indirectly measured *D*_{ClON} agree very well. The experiments leading to Table 4 and analogous measurements will be described more fully in another publication.²²

The Co(III/II) heterogeneous electron-transfer rate constant (*k*_{HET}) was measured from reduction–oxidation separations (ΔE_{PEAK}) between peak currents in cyclic voltammograms as in Figures 2 and 3, using the classical Nicholson approach.²³ ΔE_{PEAK} decreases with increased amounts of plasticizer, reflecting an increase in the reaction rate. The measured *k*_{HET} values are generally independent of the potential sweep rate used to obtain the voltammogram (see illustrative data in Supporting Information). The validity of *k*_{HET} measurements and the avoidance of IR_{UNC} effects in the Co complex melts was thoroughly examined in a recent study.^{2b} *k*_{HET} results for the Co(III/II) reaction in melt **I** with MePEG₃₅₀ and (NH₄)(MePEG₃₅₀-SO₃) plasticizers are given in Tables 1 and 2, respectively, and in Table 3 and Supporting Information for melt **II_n** with various amounts of plasticizer and at other temperatures, respectively.

The temperature dependencies of *k*_{EX}, *k*_{HET}, *D*_{PHYS}, and *D*_{ClON} in melts **I** and **II_n** are shown in Figure 4 as activation plots; the activation barrier energies are found in Tables 1–3. Thermal activation prefactors (*k*_{EX}⁰ and *k*_{HET}⁰) derived from the Arrhenius plot intercepts are given in Tables 1 and 3. Complete tables of the temperature-dependent data are found in Supporting Information.

While the activation barrier energies (*E*_{A,EX}) for the Co(II/I) reaction in Table 1 are enthalpies of activation, the reaction entropies and entropies of activation are zero since the reactions are symmetrical (isotopic) self-exchanges.^{8,9b} The *E*_{A,EX} barriers are thus equated with activation free energies (ΔG^*), and the Co(II/I) reaction rate temperature dependence is expressed by⁹

$$k_{\text{EX}} = K_{\text{p}}\kappa\nu_n \exp[-\Delta G^*/RT] \quad (3)$$

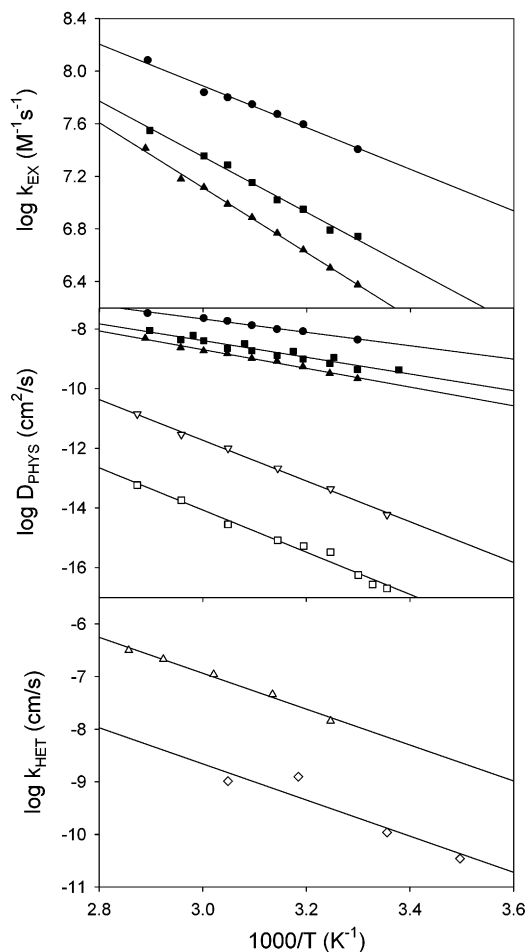


Figure 4. Activation plots of *k*_{EX} (M⁻¹ s⁻¹, upper), *D*_{PHYS} (cm² s⁻¹, middle), and *k*_{HET} (cm/s, lower) with different amounts of plasticizer added: 6:1 mol MePEG₃₅₀ (●), 3:1 mol MePEG₃₅₀ (■), 2:1 mol MePEG₃₅₀ (▲), relative to moles Co complex, 5% Co(ClO₄)₂(H₂O)₆ representative trials (△, ▽), 1% P₂M (□), 8% P₂M (◇). Filled symbols represent [Co(phen)₃](MePEG₃₅₀SO₃)₂. Open triangles represent [Co(bpy(P₂M)₂)₃](ClO₄)₂. Open squares and diamonds represent [Co(bpy(P₂M)₂)₃](ClO₄)₂. See Tables 1 and 3 for activation energy barrier and intercept data and Supporting Information for complete tables of temperature-dependent data.

where the activation enthalpy $E_{\text{A,EX}} \approx \Delta G^*$ ($= \lambda/4$ where λ is reorganization barrier energy), K_{p} is the donor–acceptor precursor complex formation constant, κ the electronic transmission coefficient, ν_n the nuclear frequency factor, k_{B} is the Boltzman constant, and $k_{\text{EX}}^0 = K_{\text{p}}\kappa\nu_n$.

The changes in the rate constants with additional MePEG₃₅₀ can arise from either changes in the activation barrier energies or in the pre-exponential term, according to eq 3. Comparing the results in Table 1 for 3 and 6 mol of added MePEG₃₅₀ plasticizer (relative to moles Co complex), the activation barrier decreases for the latter, with a contravening decrease in the pre-exponential term ($k_{\text{EX}}^0 = K_{\text{p}}\kappa\nu_n$), leaving the Co(II/I) *k*_{EX} a net 6-fold change. The reaction seems to become less adiabatic as plasticizer is added, but since adiabatic electron-transfer reactions (highly coupled reactant/product potential energy surfaces) have⁹ $\kappa\nu \approx 10^{12}$ – 10^{13} s⁻¹, most of the experiments in Table 1 would appear to fall into the adiabatic category.

Correlations between Electron Transfer and Diffusion Rates. Examination of the results in Tables 1–3 shows that both Co(II/I) and Co(III/II) electron-transfer rate constants, *k*_{EX} and *k*_{HET}, increase systematically as the mass transport dynamics

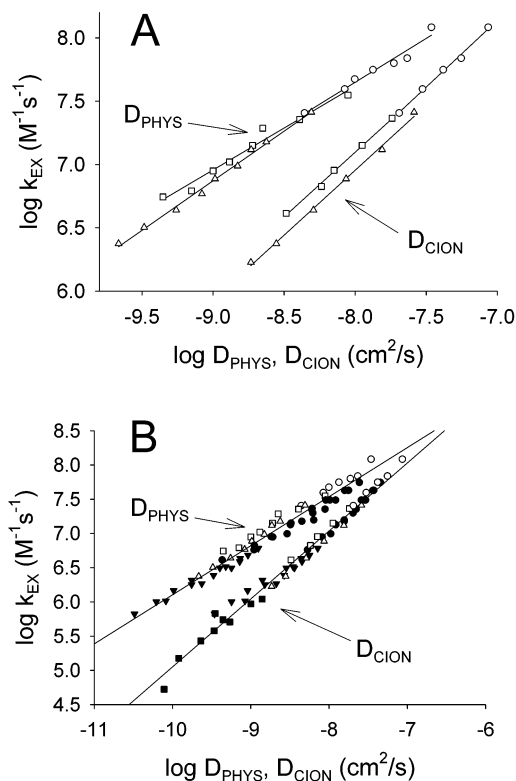


Figure 5. (A) Linear log–log relationship between k_{EX} ($M^{-1} s^{-1}$) for the Co(II/I) electron-transfer reaction vs. D_{PHYS} (cm^2/s) and D_{CION} (cm^2/s) in $[Co(phen)_3](MePEG_{350}SO_3)_2$ plasticized with MePEG₃₅₀ (6:1 mol (O), 3:1 mol (□), 2:1 mol (Δ), relative to moles Co complex) at a range of temperatures (25–70 °C). Slopes for the D_{PHYS} and D_{CION} correlations are 0.7, 0.7, and 0.8 and 1.1, 1.0, and 1.0, for 6:1, 3:1, and 2:1 mol plasticizer relative to Co, respectively. (B) Panel A data (same symbol set) combined with previously published data for $[Co(phen)_3](MePEG_{350}SO_3)_2$ ^{2b} (●) and $[Co(bpy(CO_2MePEG_{350})_2)_3](ClO_4)_2$ ^{3a} (▼), both plasticized with liquid CO₂ (over a range of temperatures) and $[Co(bpy(CO_2MePEG-350)_2)_3](ClO_4)_2 + xLiClO_4$ ^{1d} (■) ($x = 0–1.31$). The open panel B data symbols are the same as for panel A. Slope for the D_{PHYS} correlation is 0.7, and slope for the D_{CION} correlation is 1.0. See Figure S2 in Supporting Information for a colored version of this figure.

parameters increase. These correlations are shown in Figures 5 and 6, respectively.

Figure 5A displays the new results of this study for melt **I** as log–log plots of the *homogeneous* Co(II/I) constant k_{EX} ($M^{-1} s^{-1}$) vs. the Co complex self-diffusion constant D_{PHYS} (cm^2/s) and the MePEG₃₅₀SO₃[−] counterion diffusion constant D_{CION} (cm^2/s), both over a series of temperatures (25–70 °C). The data sets for different levels of plasticization of **I** with MePEG₃₅₀ (6:1 (O), 3:1 (□), and 2:1 (Δ) mol plasticizer/Co) are substantially consistent with one another and give slopes for the D_{PHYS} correlations of 0.7, 0.7, and 0.8 and, for the D_{CION} correlation, slopes of 1.1, 1.0, and 1.0, respectively. These new results are consistent with and extend earlier data on the Co(II/I) electron-transfer reaction rate in melt **I** and in another Co complex melt where the plasticization was accomplished in different ways. Figure 5B presents the new (open symbols) and previous data together, again taken over a series of temperatures, where melt **I**^{2b} (●) and the melt $[Co(bpy(CO_2MePEG_{350})_2)_3](ClO_4)_2$ ^{3a} (▼) had been plasticized by contact of the melt film by liquid CO₂, and the melt $[Co(bpy(CO_2MePEG-350)_2)_3](ClO_4)_2$ contained dissolved LiClO₄^{1d} (■) (mole ratio = ~0–1.31). In the latter case, added LiClO₄ “deplasticizes” the melt through polyether chain cross linking by Li⁺ cation/polyether coordination, resulting in depressed transport and electron-transfer rates. The

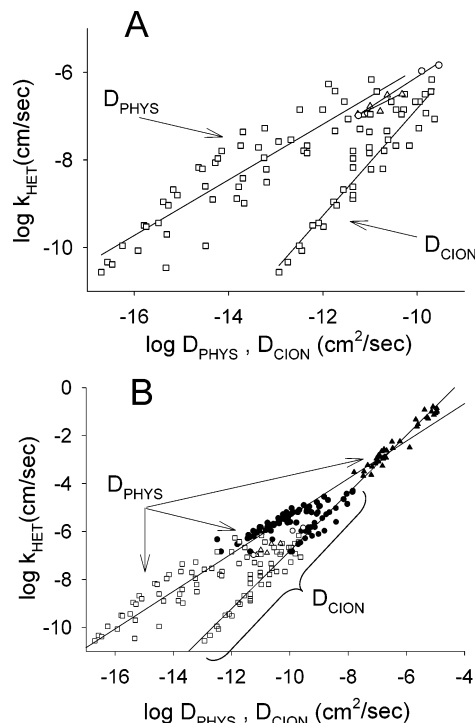


Figure 6. (A) Linear relationship between k_{HET} (cm/s) for the Co(III/II) electron-transfer reaction and D_{PHYS} (cm^2/s) and D_{CION} (cm^2/s) in $[Co(phen)_3](MePEG_{350}SO_3)_2$ plasticized with $(MePEG_{350}SO_3)(NH_4)$ (Δ) and MePEG₃₅₀ (O) and $[Co(bpy(P_nM)_2)_3](ClO_4)_2$ ($n = 2, 3$) plasticized with P_nM ($n = 2$ and 3) or $Co(ClO_4)_2(H_2O)_6$ (□) over a range of temperatures. Slopes are 0.6 and 1.2 for the $[Co(bpy(P_nM)_2)_3](ClO_4)_2$ D_{PHYS} and D_{CION} correlations, respectively. Slopes for the D_{PHYS} correlation of $[Co(phen)_3](MePEG_{350}SO_3)_2$ plasticized with MePEG₃₅₀ and $(MePEG_{350}SO_3)(NH_4)$ are 0.7 and 0.5, respectively. (B) Panel A data combined with previously studied Co(bpy)₃ melts^{1e} (●) and previously studied solutions of Co(bpy)₃¹⁰ (▲). Panel B open data symbols are the same as for panel A. Slopes are 0.8 and 1.2 for the combined D_{PHYS} and combined D_{CION} correlations, respectively. See Figure S3 in Supporting Information for a colored version of this figure.

average slope of the D_{PHYS} correlation in Figure 5B is 0.7, that for the D_{CION} correlation is 1.0. See Figure S2 in Supporting Information for a colored version of this figure.

Figure 6 shows an analogous comparison for the *heterogeneous* Co(III/II) electron-transfer rate constants. Figure 6A plots the new results in this report (over a series of temperatures) for log k_{HET} (cm/s) vs log D_{PHYS} (cm^2/s) and vs D_{CION} (cm^2/s) in melt **I** plasticized with $(NH_4)(MePEG_{350}SO_3)$ (Δ) and with MePEG₃₅₀ (O) and in melts **II**₂ and **II**₃ plasticized as described in the Experimental Section (□). There is substantial scatter but also clear systematic trends in the data for melts **II**₂ and **II**₃, which extend over a large range of values. The slopes of the correlation plots for **II**₂ and **II**₃ are 0.6 and 1.2 for D_{PHYS} and D_{CION} , respectively. Figure 6B combines these new results (open symbols) with previous data on the Co(III/II) reaction, studied in melts of PEG-tailed $[Co(bpy)_3](ClO_4)_2$ complexes^{1e} (●) and solutions of $[Co(bpy)_3](ClO_4)_2$ ¹⁰ (▲) in various solvents including Debye solvents. As in Figure 5B, there is a great consistency of the combined data, for which the log–log slopes are 0.8 and 1.2 for D_{PHYS} and D_{CION} , respectively. The D_{CION} data are more consistent with D_{PHYS} results in Debye solvents¹⁰ (upper right data in Figure 6B, where k_{HET} was also directly correlated with known τ_L values). See Figure S3 in Supporting Information for a colored version of this figure.

Models for Correlations between Electron-Transfer and Mass-Transport Rates. We have recently discussed and compared³ two models that rationalize correlations between

electron-transfer rate constants and physical transport rates such as those in Figures 5 and 6. These are (a) solvent dynamics in the context of the rate of solvent dipole reorganization^{10,11,12,24,25} and (b) a newer model that is a different version of solvent dynamics, termed ion atmosphere relaxation.³ These models will be briefly outlined next.

Solvent dynamics electron-transfer rate control in the context of dipolar reorganization time constants refers to adiabatic reactions where the barrier crossing frequency ν_n is influenced by coupling between the solvent and transition-state motions:²⁴

$$\nu_n = \tau_L^{-1} \left[\frac{\Delta G_{OS}^*}{4\pi RT} \right]^{1/2} \quad (4)$$

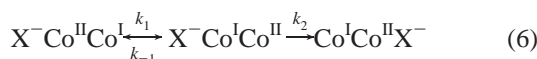
where ΔG_{OS}^* is the outer-sphere reorganizational barrier energy and τ_L is the longitudinal solvent relaxation time or time constant for solvent dipolar reorganization. The value of τ_L can be connected to diffusion constants by^{12,24}

$$\tau_L^{-1} = \left(\frac{\epsilon_s}{\epsilon_{op}} \right) \frac{3r_H D}{2\alpha^3} \quad (5)$$

where ϵ_{op} and ϵ_s are the optical and static dielectric constants, respectively, and α and r_H are molecular and hydrodynamic radii, respectively. Equation 5 provides a relation between τ_L of a solvent (and thus electron-transfer rate) and the physical diffusivity of a solute diffusing in it. A number of reports have described correlations between heterogeneous^{1e,10,11,12,24,25} and homogeneous³ electron-transfer rate constants and τ_L and/or the diffusion coefficient of the reactant (i.e., D_{PHYS}), as predicted by eqs 4 and 5.

The D_{PHYS} and D_{CION} plots in Figures 5 and 6 are consistent with eq 5. The picture drawn is that, in the metal complex melts, the solvent dipoles are considered to be those of the polyether chains, and the repolarization dynamics are associated with the chain segmental motions, as are the rates of physical diffusion of both the metal complexes and their counterions. The slopes of the former, log–log correlation between D_{PHYS} and k_{EX} and k_{HET} , are less than unity, but it has been seen before,^{24b,25} and rationalized,²⁶ that electron-transfer kinetics can vary with $\tau_L^{-\theta}$, where θ is between 0 and 1. Thus, one could interpret the results of Figures 4 and 5 as reflecting some kind of solvent dynamics control in which the dipolar reorganizational frequency and segmental motions of the polyether chains concurrently influence the electron-transfer and mass-transport rates.

The ion atmosphere relaxation model is a newer³ picture of how mass transport and electron-transfer rates could vary in parallel manners. In the metal complex melts, when an electron transfer occurs, the associated relocation of the cationic charge must be accompanied by a redistribution of its charge-compensating counterions. On the basis of the “electron-transfer-first” case discussed²⁷ in the context of ion pairing effects on electron transfers, one can write for the Co(II/I) reaction



where following electron transfer at rate constant k_1 , the counterion redistribution occurs at rate constant k_2 . The counterion mass transport step is in competition with the back electron transfer (rate constant k_{-1}). The net electron-transfer rate constant is given by

$$\frac{1}{k_{EX}} = \frac{1}{k_1} + \frac{k_{-1}}{k_2 k_1} \quad (7)$$

The counterion redistribution can be modeled²⁷ as a diffusion process in which an ion diffuses (D) over a distance “ a ” to relieve the Coulombic imbalance created by the electron transfer

$$k_2 = D \left(\frac{\pi}{2a} \right)^2 \quad (8)$$

Thus, if the right-hand term in eq 7 is dominant, the experimental electron-transfer rate constant k_{EX} becomes proportional to the diffusivity D of the ion atmosphere. In principle, k_2 refers to the rate of all ionic motions, including those of any added indifferent electrolyte and of the metal complex cation itself. In Tables 1 and 3, no electrolyte is added and $D_{CION} > D_{PHYS}$. Thus, the dominant ion atmosphere transport in the Co complex melts is expected to be of the ClO_4^- and $MePEG_{350}SO_3^-$ counterions, which are measured by D_{CION} . The ion atmosphere relaxation model, when the right-hand term of eq 7 is dominant, predicts that the net rate of electron transfers, whether the electron transfers are heterogeneous (k_{HET} for Co(III/II)) or homogeneous (k_{EX} for Co(II/I)), should be directly proportional to the counterion diffusivity D_{CION} . This is indeed what is found in Figures 5 and 6, making the ion atmosphere model a plausible alternative to the solvent dipolar reorganization dynamics model discussed above.

On what basis can one gauge which model is the more appropriate explanation? The following points are relevant: (a) The correlation of k_{EX} and k_{HET} with D_{CION} is more closely 1:1 than with D_{PHYS} . (b) In the ion atmosphere relaxation model, the net electron transfer rate is not actually governed by the electron-transfer step. Therefore the electron diffusion coefficient by which k_{EX} is measured for the Co(II/I) reaction should equal that of the counterion, and the activation barrier energy for the electron transfer should be that of ionic conductivity (from which D_{CION} is obtained). The data in Table 1 and in a previous study using CO_2 plasticization³ show that these parameters are indeed quite close to one another. (c) Finally, the time scales for the Co(III/II) and Co(II/I) electron transfers are rather different. In the context of the solvent dipole reorganization model, this is in conflict with the supposition that the solvent reorganizational time constants are of the same magnitude as the barrier-crossing frequency. The polyether melt would have to exhibit a very wide dispersion of repolarization time constants to simultaneously accommodate the results in Figures 5 and 6, which span many orders of magnitude of time scale. There is no conflict with the ion atmosphere model, on the other hand, since the experimental electron transfer rate is actually of the counterion. (d) Finally, in regard to the behavior of the heterogeneous rate constant k_{HET} , a classical Frumkin effect is an unlikely explanation since the cobalt melt is a concentrated material and overall concentration gradients of the metal complex are difficult to imagine. Likewise, while the interfacial potential gradient could be distorted by the size difference between the cobalt complex and its counterion, it is difficult to conceive how this could systematically change with melt fluidity.

Our overall conclusion is that the ion atmosphere relaxation model is a better representation of the electron-transfer dynamics in the Co complex melt.

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Supporting Information Available: Supplementary information on synthesis, electrodes, temperature-controlled cell, temperature dependence of D_{PHYS} , D_{CION} , σ , k_{EX} , and k_{HET} , and

scan-rate dependence of k_{HET} is available free of charge via the Internet at <http://pubs.acs.org>.

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- (21) (a) When considering the ratio of $D_{\text{CION}}/D_{\text{E}}$, electronic migration is not significant except for the +2 mol MePEG₃₅₀ sample. To correct for this a 1:1 ion pairing relationship between cobalt complex and counterion was assumed in this sample to calculate the reported D_{CION} . In addition, a calculation using Saveant's theory^{21b} showed that electronic migration enhances k_{EX} by a minor ca. 1.1-fold factor in all the samples. Ionic migration was also calculated to be negligible because the transference numbers t_{CION} of the counterions were only modestly less than unity.^{21c,d} (b) Andrieux, C. P.; Saveant, J. M. *J. Phys. Chem.* **1988**, *92*, 6761. (c) $t_{\text{CION}} = z_{\text{CION}}^2 D_{\text{CION}} C_{\text{CION}} / \sum (z_i^2 D_i C_i)$. (d) Bard, A. J.; Faulkner, L. R. *Electrochemical Methods: Fundamental and Applications*; John Wiley & Sons: New York, 1980; pp 66, 123.
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- (23) (a) The characteristic parameter, ψ , relates the ΔE_{PEAK} values to heterogeneous electron-transfer rate constants (k_{HET}) through the equation,^{23b,c} $\Psi = (k_{\text{HET}}(D_o/D_r)^{\alpha/2})/[D_o \pi \nu (nF/RT)]^{1/2}$, where D_o and D_r are the self-diffusion coefficients of the Co(III) and Co(II) complexes respectively,^{23d} α is the transfer coefficient (assumed to be 0.5), and ν is potential sweep rate. Digital simulation (Digisim 2.1)^{23e} of a cyclic voltammogram with a simulated ΔE_{PEAK} will correspond to a simulated k_{HET} that can be used to calculate ψ values. In this way, a standard curve of ψ vs ΔE_{PEAK} can be created and used to determine an experimental k_{HET} when ΔE_{PEAK} is measured. (b) Nicholson, R. S.; Shain, I. *Anal. Chem.* **1964**, *36*, 706. (c) Nicholson, R. S. *Anal. Chem.* **1965**, *37*, 1351. (d) D_o and D_r have been shown to be equal according to ref 2b. (e) Bioanalytical Systems, Lafayette, IN.
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- (26) The fraction θ will be unity if the reaction adiabaticity is strong and there is a small inner-sphere contribution to the activation energy barrier. The dependence of the reaction on solvent is weakened when either of these requirements is not met, causing θ to decrease in value.^{24b, 25}
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