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PREPARATION AND DISCHARGE CHARACTERISTICS OF SOLID REDOX POLYMERIZATION ELECTRODES EMPLOYING DISULFIDE POLYMERS AND COPOLYMERS.

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ABSTRACT

Recent work in our laboratory on polymeric organodisulfides has shown these materials to perform well as positive electrodes in solid-state batteries. The polymeric materials have been named solid redox polymerization electrodes (SRPE's) due to the reversible polymerization/depolymerization reaction that occurs on charge/discharge of the electrode. The cell reaction for SRPE-based cells can be described for a simple case as, $2nM + (SRS)_n = nM_2SRS$, where M is an alkali metal (Li,Na,K) and R is an organic group. In the broader sense SRPE's can have more than two S groups per monomer R unit, and are reversible to other monovalent and divalent metals. In the fully charged state SRPE's consist of polydisulfide polymers and are depolymerized on discharge by scission of sulfur-sulfur bonds, leading to the formation of dithiolate salts in the fully discharged cell. SRPE's are easy to synthesize, are air stable, and should be very inexpensive in bulk quantities. Depending on the redox potential of the polydisulfide and reaction conditions, many disulfides can be copolymerized by oxidizing a mixture of dithiols, x HSRSH + y HSR'SH + $(x+y) I_2 = (SRS)_x(SR'S)_y + 2(x+y) HI$, allowing modification of the physical and/or redox properties of the SRPE's. A series of simple aliphatic dithiols including (HSCH₂CH₂SH), (HSCH₂CH₂CH₂OCH₂CH₂SH), and (HSCH₂CH₂SCH₂CH₂SH) have been oxidized to polydisulfides and mixtures of the dithiols have been copolymerized. All of the resulting polymers and copolymers were evaluated in solid-state lithium cells, with some of the new materials demonstrating high levels of performance. The utilization of available capacity in the positive electrode was observed to be independent of electrode thickness for a number of SRPE's at loading levels up to 45% by weight. At 90°C, relatively thick positive electrodes based on (SCH₂CH₂S)_n have been discharged to surface capacities of over 20 coulombs/cm² at a current density of 0.5 mA/cm². The discharge profiles for most of the aliphatic polydisulfides are exceedingly flat at slightly over 2 volts versus lithium. Although other polydisulfides such as those derived from the dithiazoles exhibit higher cell voltages, the low equivalent weight of materials such as (SCH₂CH₂S)_n [46 g/equiv] and the low cost of such polymers indicates a potential for commercial application.

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INTRODUCTION

Recent work in this laboratory has focussed on the electrochemical properties and charge/discharge characteristics of the organodisulfide/dithiolate redox couple^[1-3]. There is potential for developing practical devices based on this chemistry; polymeric organodisulfides have been shown to perform well as positive electrodes in solid-state batteries^[4,5]. The poly(organodisulfides) have been termed "solid redox polymerization electrodes (SRPE's)" due to the reversible polymerization/depolymerization that occurs on charge/discharge in a reaction cell.

The general redox process for SRPE-based cells is described by (1):

$$xn M + (SRS)_n = n M_x SRS$$
(1)

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As a large number of disulfides are readily prepared^[6] and the choice of metal not restricted by topological effects (as in an intercalation process), the number of realizable reactions conforming to (1) is very large indeed. Initial studies have concentrated on relatively simple poly(organodisulfides) and M = Li and Na. More generally, SRPE's may include more complex poly(organodisulfides), such as copolymers containing disulfide linkages and/or polymers containing more than two disulfides per organic moiety. In addition, SRPE's may be reversible to a number of monovalent or higher valent metals.

The physical and electrochemical properties of the poly(organodisulfide) are of primary concern in the theoretical capacity and energy densities, performance, and fabrication of these cells. The remarkable influence of the organic moiety on cell potential and discharge characteristics has already been described^[4]. Poly(organodisulfides) are typically easy to prepare, air stable and inexpensive (all very desirable properties for commercial applications), and as noted above, numerous. The systematic examination of new SRPE's may therefore optimize other useful properties such as (1) low equivalent weight (for high energy and capacity densities), (2) rapid electrode kinetics at moderate temperatures, (3) high ionic diffusivities (to minimize polarization effects), and (4) processability.

In the present study, a number of organodisulfide polymers and copolymers were examined as SRPE's in lithium cells. Certain desirable traits described above were simple to incorporate. For instance, poly(ethylene disulfide), (SCH₂CH₂S)_n, (X8) has a low equivalent weight (46 g/mole), and therefore has a high theoretical energy density (see Table 1). Poly(ethyl ether disulfide), (SCH₂CH₂OCH₂CH₂S)_n has advantageous physical properties, being an amorphous, nearly colorless, soluble polymer^[7]. Thiokol (Morton Thiokol, Inc.) is an inexpensive commercial product, and was also examined in SRPE's. While the detailed chemical composition is unavailable, Thiokol is mainly composed of poly(ethyl ether disulfide) and structurally related disulfides. Copolymers were also prepared to obtain properties such as increased hardness via cross-linking.

EXPERIMENTAL

2,5-dimercapto-1,3,4-thiadiazole (Aceto Corp.), 2,4-Dithiopyramidine (Sigma), 2-mercaptoethyl ether (Aldrich, tech. grade), 2-mercaptoethyl sulfide (Aldrich, 95%), trithiocyanuric acid (Aldrich, 95%), 1,2 ethanedithiol (Aldrich, 99%), thiokol (Morton Thiokol, Inc.), K₃Fe(CN)₆ (Aldrich, 99+%), LiOH• H₂O (Aldrich, 99.9+%), I₂ (Baker, sublimed), and CH₃CN (Aldrich, HPLC grade) were used as supplied. Poly(ethylene) oxide (avg mw = 5×10^6 , Aldrich), C black (Shawinigan acetylene black, Chevron) and LiSO₃CF₃ were dried under vacuum at 50°C for several days and stored under an inert atmosphere. Lithium foil of approximately 5 mil thickness (Lithco Co, battery grade) was stored under helium and cut into disks (18 mm diameter) prior to use.

The preparation of disulfide polymers and copolymers involved the formation of a soluble lithium salt of the dithiol and subsequent oxidation of the dithiolate to precipitate the disulfide polymer. The general preparative method is illustrated by the following reactions:

$$HSRSH + 2LiOH (aq) = Li_2SRS (aq) + 2 H_2O$$
(2a)

This two-step method is quite simple and usually preferable as some of the dithiols are themselves somewhat difficult to handle even in a well-ventilated hood (stench). In contrast, the aqueous dithiolate solutions have little or no odor and, in the case of copolymerizations, insure a homogeneous phase prior to oxidation. Notably, for the case of nitrogen containing compounds such as the dimercaptodithiazoles, the starting mercaptans have no discernable smell at all. All reactions proceeded rapidly under ambient conditions in aqueous solution. Copolymers were prepared by a similar method using an appropriate combination of dithiols in (2a). The resultant polymers or copolymers were washed copiously with deionized water and a suitable organic solvent and then dried *in vacuo* at 50°C.

Positive electrodes were prepared from a mixture of the disulfide polymer (30 or 45 wt %), carbon black (7 - 10 wt %), carbon dispersant (2 - 3 wt %), and remainder poly(ethylene) oxide. A viscous slurry, typically 2 - 3 wt % total solids, was obtained in CH₃CN by vigorous stirring for several days followed in some cases by use of a high-speed homogenizer (Tekmar). The homogeneity of the dispersion was found to correlate strongly with the performance of the films produced. Films of 20 - 80 μ m thickness were cast and air-dried on a clean Teflon surface, cut into disks (area = 2.0 cm²), and dried *in vacuo* for several days at 50°C. PEO₈LiSO₃CF₃ electrolyte films (30-50 μ m thickness) were prepared by a similar method.

Test cells (see Fig. 1) were assembled under an inert atmosphere and discharged at 90 - 95°C. The Li electrode capacity in all cells exceeded that of the positive electrode by more than 400% (by stacking Li foils if necessary) and therefore did not limit cell performance. Discharge and cycling characteristics were examined with a PAR 173 or 371 potentiostat/galvanostat utilizing software developed in this laboratory^[7]. Cell impedances (typically $\approx 20 - 30 \Omega \text{ cm}^2$) were determined *in situ* by a pulse technique described elsewhere^[8]. Large cell impedances (> 75 $\Omega \text{ cm}^2$) occasionally resulted from inhomogeneous cathode films or poor contact between cell components (high interfacial impedance), and these cells generally showed poor discharge behavior. Galvanostatic cell discharge was performed at 1 mA (0.5 mA/cm²) with cell OCV's monitored periodically.

RESULTS

Structures, equivalent weights, and OCV's are listed in Table 2. The X1 polymer was obtained as a free-flowing off-white to yellow powder. The X7, X8, and X9 polymers were white powders, and X0 was a colorless or faint yellow, amorphous, highly viscous, polymer.

Copolymers were prepared by the oxidation of an appropriate combination of dithiolates in aqueous solution. In some syntheses, inhomogeneous products were obtained which clearly involved a mixture of two polymeric phases. For example, the attempted copolymerization of 50% X0 with 50% X5 produced a white powder along with a tacky, translucent polymer. In other reactions, homogeneous products with physical properties dissimilar to either of the pure polymers resulted. These were therefore taken to be true copolymers. The addition of 5% by weight X5 to X0 resulted in a single-phase polymer which was harder and considerably easier to handle than pure X0 polymer. A 50% X0 / 50% X1 combination resulted in a yellow-white powder similar in appearance to the pure X1 polymer. Products obtained were not directly identified as block, regular, or random copolymers (Fig. 2).

Discharge profiles are given in Figures 3 - 5. Charge efficiency was based on the theoretical capacities derived by weighing the cathode films, and varied from 45% (X7) to more than 80% (X1). Theoretical capacities are shown for each curve. Figure 3 gives discharge profiles for the X7, X8, X9, and Thiokol polymers. Figure 4 shows the effect of stacking cathode films in Li/PEO₈LiTf/X8 cells. Figure 5 shows discharge profiles for the X0 and X1 polymers and copolymers with 50% X0 / 50% X1, and 95% X0 / 5% X5 (X5 = trithiocyanuric acid).

DISCUSSION

The effect of the organic moiety of the poly(organodisulfide), R in (SRS)_n, on the discharge voltage in Li/Li⁺ electrolyte/disulfide cells has been noted previously^[4,5], and further illustrated by these data. Poly(organodisulfides) containing -N = CR'-S-S- linkages (X1,X7) have discharge plateaus near 2.6 - 2.7 V, those with $-CR'_2-CR''_2-S-S$ - linkages (X0,X8,X9) discharge at 2.0 V. The latter class of polymers often give initial OCV's which are some 0.5 V higher, but these potentials fall rapidly upon discharging the cell.

The X8 polymer shows a flat discharge plateau with discharge efficiencies typically above 60% for either 30% or 45% loadings. As seen in Figure 4, the stacking of X8 cathode films does not impair cell performance, and > 20 C/cm² was drawn at 0.5 mA/cm² when 3 cathode films were stacked. Initial studies on deep charge/discharge cycling of X8-based cells do not show the high cycle life and reversibility observed in X1-based cells; the X8-based cells generally lost \approx 10% discharge capacity per cycle. The cycling results may be significantly improved after further testing and modification these cells.

Table 2 gives theoretical coulometric and gravimetric film capacity and energy densities for cathode films containing X0, X1 and X8. The values derived illustrate that the very low equivalent weight of the X8 polymer (42 g/mole) compensates for its lower discharge voltage (2 V vs 3 V for X1), resulting in similar energy densities for X1 and X8- based cathode films. Due to the greater equivalent weight of X0, these polymer films have lower theoretical energy densities. Copolymers produced from 95% X0 monomer and 5% X5 monomer (trithiocyanuric acid) are easier to handle than the X0 polymer, and the discharge

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characteristics were nearly identical (see Figure 5). Cathode films containing the X1 / X0 copolymer have a discharge profile like that of X1-based films, but exhibit a more gradual decrease in potential at the end of the discharge. Thiokol-based cathode films (see Figure 3) performed almost as well as the polymers prepared by us, and clearly show the multiple discharge plateaus characteristic of a mixture of disulfides or block copolymer. The facile modification of discharge profiles and physical properties by employing copolymers or a combination of disulfides will likely be a desirable feature of SRPE-based cells. Research is presently being pursued in two major areas; (1) the investigation of the cycling characteristics and stability of these cells, and (2) the fabrication of ambient temperature cells by incorporating amorphous polymer binders and electrolytes.

ACKNOWLEDGEMENTS

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Table I

Table II

Figure 1. Experimental cell for evaluating SRPE's in solid-state lithium batteries.

Figure 2. Various types of copolymers that might be formed by oxidation of mixtures of dithiolates (-SRS-) having different organic R groups.

Figure 3. Discharge characteristics for X7, X8, X9, and Thiokol polymers (table I) in solid-state lithium cells.

Figure 4. The effect of stacking cathode films in the Li/PEO₈LiTf/X8 battery.

Figure 5. Discharge profiles for X0 and X1 polymers, and copolymers composed of 50% X0 / 50% X1, and 95% X0 / 5% X5 (where X5 = trithiocyanuric acid).

Table I

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active	loading	film density	volumetric capacity density	gravimetric capacity density	volumetric ¹ energy density	gravimetric ¹ energy density
material	wt. pct.	g/cm -	Ah/cm ⁻	Ah/g	Wh/cm	wn/g
Xo	30	1.27	0.15	0.12	0.30	0.24
XU	45	1.27	0.23	0.18	0.45	0.35
X1	30	1.45	0.16	0.11	0.47	0.33
	45	1.54	0.25	0.16	0.75	0.49
Vo	30	1.39	0.24	0.17	0.48	0.35
	45	1.45	0.38	0.26	0.76	0.52
Li	100	0.53	2.0	3.8		

¹Energy densities calculated using nominal voltages

vs Li (X0 = X8 = 2.0 V, X1 = 3.0 V).

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Table II

formula (notation)	name	eq. wt. (g/mole)	OCV *
(x_1)	2,5-dimercapto- 1,3,4-thiadiazole	74.1	3.0
(X7)	2,4-dithiopyrimidine	59.2	2.5 - 2.8
– sch₂ch₂och₂ch₂s -)- п (X0)	2-mercaptoethyl ether	68.1	2.5
- SCH ₂ CH ₂ SCH ₂ CH ₂ S	2-mercaptoethyl sulfide	76.2	2.3
	1,2-ethanedithiol	46.1	2.3 - 2.8

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* OCV measured in Li/PEO₈LiTf/polymer composite

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-(-SRS) + SR'S + (SR'S) + (S

block copolymer

SRS-SR'S-SRS-SR'S-SRS-SR'S \rightarrow

regular copolymer

random copolymer

Figure 2. Various types of copolymers that might be formed by oxidation of mixtures of dithiolates (-SRS-) having different organic R groups.

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Figure 3. Discharge characteristics for X7, X8, X9, and Thiokol polymers (table I) in solid-state lithium cells.

Discharge curves for X8 multilayer cathodes







Figure 5. Discharge profiles for X0 and X1 polymers, and copolymers composed of 50% X0 / 50% X1, and 95% X0 / 5% X5 (where X5 = trithiocyanuric acid).



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