Assessment of Lithium-based Battery Electrolytes Developed under the NASA PERS Program

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Recently, NASA formally completed the Polymer Energy Rechargeable System (PERS) Program, which was established in 2000 in collaboration with the Air Force Research Laboratory (AFRL) to support the development of polymer-based, lithium-based cell chemistries and battery technologies to address the next generation of aerospace applications and mission needs. The goal of this program was to ultimately develop an advanced, space-qualified battery technology, which embodied a solid polymer electrolyte (SPE) and complementary components, with improved performance characteristics that would address future aerospace battery requirements.

Programmatically, the PERS initiative exploited both interagency collaborations to address common technology and engineering issues and the active participation of academia and private industry. The initial program phases focused on R&D activities to address the critical technical issues and challenges at the cell level. A variety of cell and polymeric electrolyte concepts were pursued as part of the development efforts undertaken at numerous governmental, industrial and academic laboratories. Numerous candidate electrolyte materials were developed, synthesized and optimized for evaluation. Utilizing the component screening facility and the “standardized” test procedures developed at the NASA Glenn Research Center, electrochemical screening and performance evaluations of promising candidate materials were completed.

This overview summarizes test results for a variety of candidate electrolyte materials that were developed under the PERS Program. Electrolyte properties are contrasted and compared to the original project goals, and the strengths and weaknesses of the electrolyte chemistries are discussed. Limited cycling data for full-cells using lithium metal and vanadium oxide electrodes are also presented. Based on measured electrolyte properties, the projected performance characteristics and temperature limitations of batteries utilizing the advanced electrolytes and components have been estimated. Limitations for the achievement of practical performance levels are also discussed, as well as needs for future research and development.
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Program Background

• DoD/NASA collaborative effort was formed in 1997 to enhance Li-ion liquid battery and cell performance.
• Lithium-based chemistries with solid-state and/or polymer-based components were envisioned to be enabling next-generation technologies.
• In FY 2000 NASA and the Air Force Research Laboratory expanded collaborative efforts to support development of polymer-based technologies.
• Advancement of next-generation technologies formed the basis of the Polymer Energy Rechargeable System (PERS) Program.
  – Participants selected in FY 2001 via a NASA Research Announcement (NRA)
Objective

Establish a world-class technology capability and U.S. leadership in polymer-based battery technology for aerospace applications.

Anticipated benefits of PERS

• simplified, lightweight designs
• intrinsic safety
• increased specific energy, energy density, life
• favorable manufacturing costs
### PERS Program Participants

#### PERS PROGRAM SELECTED CONTRACTOR AND GRANTEES

<table>
<thead>
<tr>
<th>Organization</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Covalent Associates</td>
<td>Advanced Lithium Polymer Electrolyte for Li-Ion Batteries</td>
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<tr>
<td>Eagle-Picher Technologies, LLC, Joplin, MO</td>
<td>Novel Host Material for Ceramic Electrolytes</td>
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<tr>
<td>EIC</td>
<td>Lithium Based Polymer Electrolyte Battery</td>
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<tr>
<td>Indiana University</td>
<td>Nanostructured Single-Ion SPE for Lithium Batteries</td>
</tr>
<tr>
<td>InvenTek</td>
<td>Development of Solvent-Free High Performance Li-Polymer</td>
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<tr>
<td>Lawrence Berkeley National Laboratory (LBNL)</td>
<td>Advanced Polyelectrolyte Separators for Rechargeable Lithium Batteries</td>
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<tr>
<td>LITHCHEM International</td>
<td>New Stable, More Conductive Solid Polymer Electrolyte</td>
</tr>
<tr>
<td>Lithium Power Technologies</td>
<td>Polymer Electrolytes for Solid-State Lithium Batteries</td>
</tr>
<tr>
<td>Lockheed Martin / Comsat</td>
<td>Lithium Based Polymer Battery Development</td>
</tr>
<tr>
<td>Max Power / Temple University</td>
<td>An Integrated Approach to Develop a High Energy Density Long Cycle Life Lithium Based Polymer Battery</td>
</tr>
<tr>
<td>Naval Air Warfare Center Weapons Division (NAWCWD)</td>
<td>New Solid Polymer Membranes for Rechargeable Lithium Batteries</td>
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<tr>
<td>National Center for Microgravity Research</td>
<td>Thermal Modeling of Lithium Based SPE Battery in Microgravity Applications</td>
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<tr>
<td>Northwestern University</td>
<td>Development of Highly Conductive Polyelectrolytes for Lithium Batteries</td>
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<tr>
<td>Physical Sciences, Inc.</td>
<td>High Performance Li-Ion Polymer Electrolyte with Pendant Anion Receptors</td>
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<tr>
<td>Texas Engineering Experimental Station (TAMU)</td>
<td>Advanced Li-Ion Polymer Batteries for Aerospace Applications</td>
</tr>
<tr>
<td>University of Minnesota</td>
<td>New Polymer Electrolyte Cell Systems</td>
</tr>
<tr>
<td>University of Utah</td>
<td>A Joint Simulation and Experimental Study of Nanocomposite Polymer Electrolytes</td>
</tr>
<tr>
<td>Yardney Technical Products</td>
<td>Development of a Lithium-Based Polymer Energy Rechargeable System for Future Space Applications</td>
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<tbody>
<tr>
<td>NASA Glenn Research Center (GRC)/University of Akron</td>
<td>Novel Molecular Architectures for Improved Solid Polymer Electrolytes for Lithium Polymer Batteries</td>
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<tr>
<td>Air Force Research Laboratory (AFRL) – WPAFB</td>
<td>Ionically-Conducting Channel Polymer Electrolytes</td>
</tr>
<tr>
<td>Jet Propulsion Laboratory (JPL)</td>
<td>Solid Polymer Electrolyte Development</td>
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Goals for Electrolyte Development

GRC Testing Facility was expanded to screen and compare candidate PERS electrolytes in terms of critical properties:

- High ionic conductivity (~10^{-3} S/cm)
- Li^{+} transference number approaching 1
- High salt diffusion coefficient
- Electrochemical Stability Window 0 to >4 V
- High Electronic Resistivity (>10^{12} ohm·cm)
- Favorable Interfacial Properties
- Stable transport properties
- Thermal and dimensional stability
- Mechanical toughness
Ionic Conductivity

- Gels approach PERS goal at room temperature.
- Dry SPEs approach goal at >80°C.
- Ionomers have lower conductivity due to immobilized anion.

Over 330 polymer electrolyte samples submitted (most based on PEO)
The best dry SPE (optimized for conductivity) = $5 \cdot 10^{-5}$ S/cm at 25°C
Ionomers not provided in sufficient quantity for supplemental testing
Transport Properties for Dry SPE

- Li$^+$ transference number ($t_+^0$) and salt diffusion coefficient ($D_s$)
- Influence salt concentration gradients under polarization.
- Limiting current density, power production, electrode utilization and even life are affected.

<table>
<thead>
<tr>
<th></th>
<th>dry SPE at 80°C</th>
<th>Liquid electrolyte at 23°C</th>
</tr>
</thead>
<tbody>
<tr>
<td>$t_+^0$</td>
<td>0.2</td>
<td>0.2-0.4</td>
</tr>
<tr>
<td>$D_s$, cm$^2$/s</td>
<td>10$^{-8}$</td>
<td>10$^{-6}$</td>
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</table>

Salt diffusion coefficient is low in SPE, even at 80°C
Electrochemical Stability Window

- measurements at 80°C shown for dry SPE optimized for conductivity.
- reasonable plating/stripping efficiency for Li metal.
- small oxidation current observed at +3.3 V
- salt and polymer makeup could be factors

anticipated compatibility with 3-Volt class electrodes
Full Cell Testing with Vanadium Oxide

- Vanadium oxide explored at GRC as positive electrode material
- 200 mAh/g (practical capacity for $V_6O_{13}$).
- 3-Volt, fits stability window
- Preliminary cycle-testing in coin cells with non-optimized electrode formulation.

Preliminary GRC cells built using un-modified PEO as well as rod-coil SPE
Full Cell Impedance

- Cell data at 70°C
- Compare SPE cell with control cell using liquid electrolyte (Merck LP30)
- Higher interfacial impedance with SPE

Even if separator thickness was reduced, interfacial impedance would still be high.
Full Cell Data

- Achieved 200 mAh/g in limited cycling at 80°C, 80 μA/cm².
- Variability is due to non-optimized nature of the electrodes.

Preliminary lab cells achieved “practical” capacity in limited cycling.
Battery Energy Estimation

With limitations of dry SPEs:

- Cell operates at elevated temperature (80°C)
- 3V cell potential (1 volt less than Li-ion)
- Battery design to accommodate insulation & heaters (adds weight and volume)
- Parasitic power loss to drive heaters

... will a heated battery based on dry SPE have an energy advantage over Li-ion?
## Comparison of 30 Ah Prismatic Cells

<table>
<thead>
<tr>
<th>Chemistry</th>
<th>Li-ion</th>
<th>PERS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cell energy, Wh</td>
<td>108 (at 3.6 V&lt;sub&gt;avg&lt;/sub&gt;)</td>
<td>72 (at 2.4 V&lt;sub&gt;avg&lt;/sub&gt;)</td>
</tr>
<tr>
<td>components</td>
<td>grams</td>
<td>wt. fraction</td>
</tr>
<tr>
<td>electrochemical materials</td>
<td>633.1</td>
<td>75.1%</td>
</tr>
<tr>
<td>enclosure materials</td>
<td>201.6</td>
<td>23.9%</td>
</tr>
<tr>
<td>terminals</td>
<td>1.8</td>
<td>0.2%</td>
</tr>
<tr>
<td>auxiliary header materials</td>
<td>6</td>
<td>0.7%</td>
</tr>
<tr>
<td>total weight</td>
<td>842.5</td>
<td></td>
</tr>
</tbody>
</table>

Cell specific energy, Wh/kg
- Li-ion: 128
- PERS: 188

Cell dimensions, cm.
- Overall height: 16.71
- Width: 8.89
- Thickness: 2.27
- Total volume, cm<sup>3</sup>: 337.3

Cell energy density, Wh/dm<sup>3</sup>
- Li-ion: 321
- PERS: 324

Lower cell voltage is offset by low component weight. Specific energy is 1.5-times greater at the cell level.
Thermal Model of PERS Battery

Assumptions:
• Uniform cell temperature
• Neglect generation (polarization/entropy)
• Natural convection
• Conduction to platform
• 23°C ambient temperature

Boundary Conditions

Significant heat loss through electrical conductors. Insulated buss extension is essential, to limit heat flow but adds weight.
Comparison of 28-Volt Batteries

<table>
<thead>
<tr>
<th>Electrochemistry</th>
<th>Li-ion</th>
<th>PERS</th>
<th>PERS</th>
</tr>
</thead>
<tbody>
<tr>
<td>Allowance for heat loss</td>
<td>none</td>
<td>none</td>
<td>heat loss at 80°C</td>
</tr>
<tr>
<td>no. of cells</td>
<td>8</td>
<td>12</td>
<td>12</td>
</tr>
<tr>
<td>Specific energy, Wh/kg</td>
<td>113</td>
<td>167</td>
<td>146</td>
</tr>
<tr>
<td>Energy density, Wh/dm³</td>
<td>269</td>
<td>291</td>
<td>178</td>
</tr>
<tr>
<td>net capacity, Ah</td>
<td>30</td>
<td>30</td>
<td>27.6*</td>
</tr>
</tbody>
</table>

Even with allowance for parasitic heat loss, PERS battery has significantly greater specific energy.

Insulation reduces energy density.

* 9% of battery power used to run guard-heaters
Cell Electrode Area and Parasitic Weight

Electrode area is a critical cell-design parameter.
• current density
• active material layer thickness
• relative weight of current collectors

12,000 cm² electrode area was selected for the hypothetical 30-Ah cell design:
  3-mil thick positive electrode active layer
  For C/8 rate (3.75 A) current density is 312 μA/cm²
  30% of the electrochemical material is current collector

For rates greater than C/8, current density or electrode area must be increased.

What if electrode area is increased by a factor of 3?
Electrode-Area Effect in 30-Ah Cell

With 3-fold increase in electrode area, half of the cell component weight is occupied by metal substrate.

Assumes “conventional” foil thicknesses 0.8-mil Al & 0.6-mil Cu
Summary

• Over 20 PERS participants helped increase the body of knowledge for novel SPE approaches and produced materials with new characteristics and physical properties.

• Breakthrough-levels of electrochemical performance have yet to be accomplished - work continues.

• A number of SPEs show promise as viable electrolytes at elevated temperature.

• Specific energy advantages are possible for batteries with thermal management systems.

• Cycle-life, utilization, long-term stability, etc. need to be demonstrated in future work with optimized electrodes.
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Acknowledgments

The authors gratefully acknowledge the support provided by NASA and other collaborating governmental agencies.

All of the formal NASA PERS Program participants.

The efforts of the participants resulted in a wealth of new and fundamental knowledge into the public domain and scientific literature, which, hopefully, will stimulate more new ideas and approaches to address the challenging criteria for the future viability of the technology.