A Fundamental Investigation on the Structural Dynamics of Model Siloxane Networks under Extreme Pressure

Stephen J. Harley (PI), James P. Lewicki

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RESULTS FROM PRIOR LDRD SUPPORT

The PI, Stephen Harley, has not previously been PI of an LDRD; however, Stephen provided analytical support for the multifunctional materials focus area of the Chemistry and Materials Science Directorate ER-LDRD (12-ERD-046) “Predicting Weapon headspace gas atmosphere.” The focus of this element was to develop a code capable of predicting weapon headspace atmosphere in support of aging assessment. Part of this research involved developing experimental and computational methodologies to assess the aging characteristics of these materials. Stephen aids in both the experimental design to obtain relevant parameters and the computational code generation. This LDRD is ongoing and is in year 2, there are 3 publications currently being written and 1 has been accepted:

**PROJECT DESCRIPTION**

**INTRODUCTION**

Everyone is familiar with the elastic properties of a butyl rubber racket ball, bouncing freely when thrown against a wall. People are similarly aware that when submerged in liquid N\textsubscript{2} the same ball becomes brittle and hard, shattering when thrown against a wall. This is a result of butyl rubber undergoing a phase transition as it crossed a critical temperature, T\textsubscript{g}. While the thermal dependence of these phase transitions has been extensively explored, other state variables such as pressure have largely been ignored. However, this gap in our knowledge base does not indicate that pressure has no influence on the morphological state of polymers; rather, it’s simply been overlooked. Mechanical testing is the dominant method to characterize polymers up to approximately 1 MPa\[1\] and diamond anvil experiments span 1 GPa onward\[2\]. There is a clear gap in our knowledge base where pressure induced changes have not been explored. Phase transitions are not benign subtle variations in a polymer’s properties; rather they induce drastic changes in the materials mechanical properties. Consequently, pressure induced crystallization could result in a polymeric part going out of spec. Additionally network failure or unknown degradative mechanisms could arise. At this point, we are looking at an entirely unexplored area of materials behavior.

Functional components based on Poly(siloxane) elastomers are ubiquitous in their use in the fields of engineering, aerospace, defense and the biomedical industries. And as such, this diverse class of material often find application in extreme environments of temperature and/or pressure. While the effects of temperature on polysiloxanes has been well studied and their behavior is well understood, it is startling to note that we know practically nothing about this class of materials behavior or eventual fate at extreme pressures. Consequently no predictive estimate of the lifetime of failure of a polysiloxane material can be offered in the MPa-GPa region, where this critical knowledge gap lies.

![Figure 1: Schematic representation of our gap in testing capabilities for polymeric materials.](image-url)
Nuclear Magnetic Resonance Spectroscopy (NMR) and Broadband Dielectrics Spectroscopy (BDS) are sensitive techniques to assess both the long and short range motions in polymeric materials.[3-9] Incorporation of high pressure probes to these spectroscopies has already been established in the solution state by the PI.[10-13] In addition, a preliminary experiment on a model PDMS network did yield a sudden change in relaxation that could indicate a phase transition, see Fig. 2. Further investigations were not pursued due to the lack of time available to the team. We have the materials, expertise, and equipment to start a rigorous scientific study, funding is sought to cover the time needed for this effort.

**PROJECT PLAN**

**Objective 1: Hardware Development**

Currently we have two functional high pressure probes both adapted for NMR analysis. The first is a large variable temperature probe having dimensions similar to typical NMR probes. The probe circuit coil will need to be tuned to the desired frequency (400 MHz) and sized according to the samples geometry. The PI does not envision any problems with this as the PI has over 8 years of NMR hardware experience. This system has a maximum working pressure of 350 MPa. J. Lewicki (an expert in the field of applied polymer dielectric spectrometry) will support the modification of this probe to accommodate compact, inter-digit electrodes and the circuits required to interface it with a broadband dielectric spectrometer, allowing dielectric reloxmetry data of materials of study to be acquired under pressure and in real time.

The second probe is a much more compact variant that allows for hydrostatic pressures up to 2 GPa. It has been pressure tested with an academic collaboration at UC Davis and NMR spectra for $^{29}$Si solution speciation has been taken. As with the system above, circuit modifications to achieve a 400 MHz resonance does not present a challenge. The team has significant experience.
with microcoil NMR and adaptation to dielectrics will be similar to the challenges mentioned above. A safety note will need to be written to take spectra here under our existing IWS.

Figure 4: High pressure probe capable of reaching 2GPa.

Objective 1 Key Deliverables:
Year 1: Adapt hardware to interface with our spectrometers
Year 2: Continue with maintenance and implement changes as needed.

Objective 2: Demonstrate ability to monitor phase changes

The co-PI has significant experience in synthetically producing model PDMS networks and has optimized the formulation chemistry. We will use these networks as the focus of our investigation. The benefit of these networks is a control in the physical chemistry. This is to say we can understand any bulk morphological changes as a direct result of the PDMS undergoing a phase transition thereby decoupling the effects of fillers typically encountered in commercial formulations. This allows us to understand possible phase transitions from a fundamental scientific perspective.

Figure 5: Example of a model PDMS network where the cure chemistry has been optimized.

These model networks spanning both above and below the entanglement molecular weight of PDMS will be cast into appropriate sample containers and inserted into the high pressure vessels. Pressure will be varied systematically from approximately 100 kPa to 2 GPa and evidence for phase transitions will be monitored with the aforementioned spectroscopies. In dielectric spectroscopy a sudden shift in the α and β relaxation frequencies would be indicative of a phase
transition. In NMR spectroscopy the relaxation properties of the PDMS will be investigated with a combination of classic inversion recovery and CPMG pulse sequences as well as more advanced multiple quantum experiments. It is well established that the relaxation properties of the material are directly related to the motional dynamics of the material.[14] Therefore any drastic stiffening or softening of the material (indicative of a phase change) would be clearly visible in these spectroscopies.

The one possible source for failure in these experiments is the lack of any phase change in the span of pressures investigated. However, we have preliminary evidence (Fig. 2) that indeed there will be.

**Objective 2 Key Deliverables:**
- Year 1: Test ideal network with NMR
- Year 2: Test ideal network with Dielectrics and compare

**Objective 3:** Interpret data towards broader implications in other materials

Model networks provide an excellent foundation for our understanding of pressure related morphological changes. However, these model materials not have the necessary mechanical properties for commercial implementation. Fillers are often added to achieve desired mechanical properties. As part of phase two of this project model networks will be synthetically produced will varying levels of filler loading. Then these materials will be subjected to the procedures outlined in objective 2 and shifts in phase changes will be sought.

This data will directly aid in the tailoring of materials to be more resistant to pressure changes.

**Objective 3 Key Deliverables:**
- Year 1: N/A
- Year 2: Test more complex networks

**MANAGEMENT PLAN**

Principal Investigator Stephen Harley has extensive experience leading small teams in high pressure research will have direct responsibility for ensuring that goals, deliverables and milestones are met throughout the course of the project. S. Harley will also be technical lead in the use, modification of the high pressure equipment and will be scientific lead on NMR spectroscopy. James Lewicki will be responsible for the synthesis the model networks and lead the dielectric spectroscopy effort. Evaluation of all data will be done as a group, communication with the team will occur primarily through regularly scheduled project meetings to discuss project status and results and facilitate the integration of the computational and experimental aspects of the project. We will obtain valuable peer review through publication of results in scientific journals and presentations at conferences, well as other national scientific venues.

We anticipate that the initial results obtained with internal LDRD investment will serve as a foundation upon which we will transition to sponsor funds to expand to a much larger
investigation of materials pertinent to the mission of the lab. This capability will be broadly applicable to any solid or liquid and will provide increased understanding of how materials behave in this little understood pressure region. With the fundamental understanding of the chemistry that governs pressure induced phase transitions, a better selection of materials to mitigate unwanted morphological changes can be made.

**DISSEMINATION**
This work will be published in peer review journals and presented at professional conferences.

**SUMMARY**
Poly(dimethylsiloxane) elastomers are ubiquitous in their application as key components in a range of (often extreme) environments and as such, they often experience pressures in a region where we do not understand their physical response and therefore can offer no predictive estimate of lifetime of failure. We are proposing a fundamental scientific investigation of the pressure induced morphological changes in model PDMS networks. Both NMR and BDS spectroscopies will be used to interrogate pressure induced changes and meaningful phase relations will be sought. The addition of filler to the networks will be used as an attempt to engineer the network to be more resistant to pressure changes. We have the equipment, model networks and expertise to start this project immediately; funding is sought as an investment in our time allocation.

**REFERENCES CITED**


**BIOGRAPHICAL SKETCHES**
Stephen Harley, Ph.D.
Energetic Materials Center
Lawrence Livermore National Laboratory
925.422.5115
harley2@llnl.gov

(a) Professional Preparation

University of California - Davis  Chemical Engineering  B.S., 2006
University of California - Davis  Analytical Chemistry  Ph.D., 2009
Lawrence Livermore National Lab  CSD  2011 (current)

(b) Appointments

March 2013 – Current: Staff Researcher, Lawrence Livermore National Lab, Chemical Sciences Division, Energetic Materials Center.


Feb. 2010 – April 2011: Post-Doctoral Researcher, University of California - Davis, Chemistry Department, Casey Group. Designed high pressure NMR hardware, studied aqueous actinide ligand exchange and characterized new water splitting catalyst using a broad spectrum of analytical techniques.


(c) Awards

Office of Naval Research Award: 2006.


(d) Publications
Related to proposed project:


Other significant publications:


**James P. Lewicki**  
Forensics and Assessments Support  
Lawrence Livermore National Laboratory  
423-1115  
Lewicki1@llnl.gov

**(a) Professional Preparation**

A list of the individual's undergraduate and graduate education and postdoctoral training as indicated below:

University of Strathclyde, Glasgow  
Chemistry  
Msci, 2004

University of Strathclyde, Glasgow  
Physical Chemistry  
Ph.D, 2008

University of Strathclyde, Glasgow  
Polymer Chemistry  
2008-2009

**(b) Appointments**

Dec 2011 to Present: Full time scientific research staff member, Lawrence Livermore National Laboratory, Chemical Sciences Division, Forensics and Assessments Support Group


**(d) Publications**

i) Related to the proposed project


ii) Other significant publications


