Novel High Temperature and Radiation Resistant Infrared Glasses and Optical Fibers for Sensing in Advanced Small Modular Reactors

Reactor Concepts
Research Development and Demonstration (RCRD&D)

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1. Overview

1.1 Abstract

One binary and three series of ternary non-oxide pure sulfide glasses compositions were investigated with the goal of synthesizing new glasses that exhibit high glass transition ($T_g$) and crystallization ($T_c$) temperatures, infrared transparency, and reliable glass formability. The binary glass series consisted of Ge$_2$S$_2$ and La$_2$S$_3$ and the three glass series in the x(nBaS + mLa$_2$S$_3$) + (1-2x)GeS$_2$ ternary system have BaS:La$_2$S$_3$ modifier ratios of 1:1, 1:2, and 2:1 with . With these glasses, new insights were realized as to how ionic glasses form and how glass modifiers affect both structure and glass formability. All synthesized compositions were characterized by Infrared (IR) and Raman spectroscopies and differential thermal analysis (DTA) to better understand the fundamental structure, optical, and thermal characteristics of the glasses.

After a range of these glasses were synthesized, optimal compositions were formed into glass disks and subjected to gamma irradiation. Glass disks were characterized both before and after irradiation by microscope imaging, measuring the refractive index, density, and UV-VIS-IR transmission spectra. The final total dose the samples were subjected to was ~2.5 MGy. Ternary samples showed a less than 0.4% change in density and refractive index and minimal change in transmission window. The glasses also resisted cracking as seen in microscope images. Overall, many glass compositions were developed that possess operating temperatures above 500 °C, where conventional chalcogenide glasses such as As$_2$S$_3$ and have $T_g$s from ~200-300 °C, and these glasses have a greater than $T_c–T_g$ values larger than 100 °C and this shows that these glasses have good thermal stability of $T_g$ such that they can be fabricated into optical fibers and as such can be considered candidates for high temperature infrared fiber optics. Initial fiber fabrication efforts showed that selected glasses could be drawn but larger samples would be needed for further development and optimization.

1.2 Research Motivation

Advanced Small Modular Reactor (AdvSMR) development is being considered as an “energy-bridge” to providing the US with a sustainable, economically viable, and carbon-neutral energy supply. Future AdvSMR deployment requires new sensing and monitoring instrumentation, control, and human-machine interface (ICHMI) architectures that accommodate the extreme temperatures, pressures, corrosive environments, and
radiation fluxes of these new systems. Optical fiber-based sensing approaches have been identified by the DoE as suitable for measuring key AdvSMR process parameters. However, high temperature and radiation flux deployment of current silica-based optical fiber is limited by the fiber jacket properties and susceptibility of dopants to radiation-induced darkening. Overcoming the current optical fiber limitations would enable in-vessel optical access and optical-based ICHMI systems.

Non-oxide glasses conventionally employed for IR fiber-based sensors are resistant to radiation-induced darkening, but their relatively weak chemical bonding limits their working temperature range and thermal stability. This research focused on a family of non-oxide glass compositions that were built upon ionic bonding schemes. Due to their very different bonding, ionic non-oxide compounds can have significantly different properties than their covalent analogues including high melting points and significant radiation hardness.

2. Selection of the Glasses

Due to their widely different chemical bonding, ionic non-oxide glass compositions possess dramatically different properties than their covalent analogues, including melting points of the ionic chalcogenides that they are formed from that can be in excess of 3,000°C and hence they can have very high glass transition temperatures (Tgs). Further, their ionic character gives rise to high electronic binding energies which can foster radiation hardness. Specifically, rare-earth chalcogenides, such as La$_2$S$_3$, and alkaline earth chalcogenides, such as BaS, have melting points of ~ 2,000°C and are strongly glass forming with other refractory sulfides, such as Ga$_2$S$_3$, which produce refractory, fully ionic, and IR transparent glasses.

This project has explored some of the more promising of these less studied, but none-the-less important ionically-bonded non-oxide glasses that possess high thermal stability and radiation hardness. We have developed compositions based upon chemical structure models that develop charge compensated networks, such as mixing high melting point divalent chalcogenides with a high melting point trivalent chalcogenides to produce an overall in situ tetrahedral network (ITN) that lends itself to increased glass formation. The composition MS + $G'$$_2$S$_3$, where M is a divalent metal, such as Ba, and $G'$ is a trivalent metal, such as La$_2$S$_3$ would be typical of this compositional model. Specific examples could be BaS + La$_2$S$_3$ + GeS$_2$.

3. Synthesis of glasses in the xBaS + yLa$_2$S$_3$ + (1-x-y)GeS$_2$ glass system

The handling of sulfide materials necessitates high purity nitrogen and argon glovebox environments where the O$_2$ and H$_2$O levels are consistently below 5 ppm. As high purity germanium sulfide is expensive and not widely available, it was synthesized in the lab. Sixty-gram batches of stoichiometric amounts of germanium and sulfur
powders (germanium and sulfur from Arcos Organics 99.999 % pure) were sealed in a silica ampoule under vacuum.

The prepared ampoule containing the Ge and S powders was placed into a metal chamber wrapped in nichrome wire to secure the chamber lid. The secured chamber was placed inside a tube furnace angled at 5 degrees above horizontal and connected to a motor to rotate the chamber at 10 rpm throughout the synthesis. The furnace was heated at 1 °C/min to 900 °C and held for 8 hours. After melting, the furnace was allowed to cool to room temperature naturally. The sample was removed from the ampoule and subsequently ball milled in a glovebox to a micron sized powder for use as a batch material.

In order to synthesize binary and ternary glasses, 5 gram batches of lab-synthesized GeS₂, and BaS (Sigma Aldrich 99.9 % pure) and/or La₂S₃ (Strem Chemicals 99.9 % pure) were mixed and placed into a carbon coated silica ampoule with a 10 mm inner diameter and 2 mm wall thickness. The prepared ampoule was evacuated down to 60 mTorr and then sealed. Sample batches were heated in a tube furnace at 1 °C/min to 1,150 °C and held at temperature for 12 hours. Samples were homogenized by rocking the furnace from vertical to 10-15 degrees below horizontal for one hour. After mixing, the samples were left vertical for at least one hour so that the glass would pool at the bottom of the ampoule. After the melting process, ampoules were quenched to room temperature for typical glass samples, and annealed for disk samples. A ternary plot showing glass formation in the glass system is shown in Fig. 1.
4. Glass Formability of and Phase Separation in Glasses in the BaS + La$_2$S$_3$ + GeS$_2$ System

All of the compositions in each these four series formed glassy materials except compositions that melt above 1150 °C, which is the upper temperature limit for the silica ampoules. DTA scans showed glass transition and crystallization temperatures. In addition, the broadness of the infrared and Raman peaks indicated a variety of bond angles, indicating an amorphous material was synthesized. However, upon further investigation with an optical microscope, it was found that at low total modifier content (typically 5% < BaS + La$_2$S$_3$ < 30%), phase separation occurs. Figures 2, 3, and 4 provides optical micrographs of select glass compositions in each series showing the progression of microstructure with increasing modifier content.

In each series, the microstructure of the phase separated compositions shown in Figs. 2, 3, and 4 progresses in a similar fashion where initially large droplets dominate the structure. As the modifier content (BaS + La$_2$S$_3$) increases, the average diameter of the droplet phase reduces and eventually the two phases combine to give a single phase homogenous melt. Fortunately, the glasses with the highest Tgs are these single phase glasses that have the highest modifier content and can be considered further for fiber optic applications. Table 1 lists where single phase and two phase glasses occur for each series.
Figure 2. Micrographs showing large droplet and matrix phases in the 5 + 5 + 90, 5 + 10 + 85, and 10 + 5 + 85 compositions.

Figure 3. Micrographs showing droplet and matrix phases in the 15 + 15 + 70, 10 + 20 + 70, and 20 + 10 + 70 compositions.

Figure 4. Micrographs showing single phases glasses in the 20 + 20 + 60, 17.5 + 35 + 47.5, and 40 + 20 + 40 compositions.
Table 1. Compositional ranges for homogenous glass formation and phase separation for all glasses prepared in the series xBaS + yLa$_2$S$_3$ + (1-x-y)GeS$_2$.

<table>
<thead>
<tr>
<th>Composition</th>
<th>Single Phase Glasses</th>
<th>Two Phase Glasses</th>
</tr>
</thead>
<tbody>
<tr>
<td>x = y</td>
<td>20 + 20 + 60</td>
<td>5 + 5 + 90</td>
</tr>
<tr>
<td>2x = y (La$_2$S$_3$ rich)</td>
<td>30 + 15 + 55</td>
<td>10 + 5 + 85</td>
</tr>
<tr>
<td>x = 2y (BaS rich)</td>
<td>12.5 + 25 + 62.5</td>
<td>5 + 10 + 85</td>
</tr>
</tbody>
</table>

5. Glass Transition and Crystallization Temperatures of Glasses in the BaS + La$_2$S$_3$ + GeS$_2$ System

In order to better understand the chemical and structural role played by the glass modifiers (BaS + La$_2$S$_3$) with respect to thermal stability and to measure the maximum usage temperature of glasses in these systems, differential thermal analysis (DTA) was employed. The maximum usage temperature of a glass is typically 10-50 °C below the $T_g$ of the glass, however, many factors influence this such as glass composition, environment and pressure. To measure the onset $T_g$ and $T_c$ values, see Fig. 5 below, and to determine the thermal stability of glasses, $\Delta T$, where $\Delta T = T_c - T_g$, two different types of DTA scans were performed on each sample. First, a survey scan was run to relatively high temperatures that was expected to be well above both the $T_g$ and the $T_c$ of the glass composition. This full temperature survey scan allowed a rough determination of $T_g$ and a determination of the $T_c$ of the glass to be measured. After these values were, determined, a $T_g$ scan was made such that the glass was heated up to $\sim$ 40 °C of the $T_g$ but below the $T_c$ of the glass. The glass was then cycled three times between about 200°C below the $T_g$ to about 40 °C above the $T_g$ of the glass to give three scans of the same reproducible thermal history of the glass. In this way, all of the $T_g$s of all of the different glasses would have the same thermal history and as such the composition dependence of the $T_g$ would only reflect the effect of the composition and not any differences due to different thermal histories.

20 mg samples were sealed hermetically under vacuum inside 4 mm inner diameter and 5 to 8 mm length thin walled silica ampoules. The hermtical sealing of the glass
powder protected them from oxidation during the high temperature DTA thermal scans. As described above, the ampoules were loaded into the DTA and heated ~40 °C above T_g of the glass at 20 °C/min. Three scans was performed in order to impart the same thermal history to each composition and to determine the reproducibility of the T_g values. A final survey scan was run to crystalize the glass in order to obtain T_c values. Figure 5 depicts how the onset of T_g and T_c are calculated for a typical DTA curve. Tangents are taken from the flat portion of the curve and the inflection point in the curve and where the two tangent lines cross is considered to be the T_g and T_c values, respectively.

Figure 5. Example construction for calculating the onset of the glass transition (T_g) and crystallization (T_c) temperatures.

Figures 6, 7, and 8 show the dependence of T_g, T_c and ΔT on the modifier content (x) for the xBaS = xLa_2S_3 (equal molar amounts), xBaS = 2xLa_2S_3 (lanthanum sulfide rich), and 2xBaS = xLa_2S_3 (barium sulfide rich) series, respectively. It is important to note that the three glasses with the highest modifier content in each series are single phase homogeneous glasses, as described in Table 1. GeS_2 glass is also a single phase homogeneous glass. All of the T_g and T_c values measured in this work are given in Table 2 below.

In the case of the phase separated glasses, intermediate levels of modifier content, only the first T_g and T_c values are reported because these are the lower of the two sets of values. For the series with equal parts BaS and La_2S_3, the T_c values remain relatively constant while the T_g values increase for single phase glasses with increasing modifier content. This results in a reduction in thermal stability with increasing modifier content. Table 2 shows that the ΔT values for nearly all of the glasses examined are well above
100°C and as such indicate that the thermal stability of all these particular glassy compositions is sufficient enough for fiber pulling. The highest $T_g$ in this series is the 25-25-50 composition which has a $T_g$ of 570 °C, but this comes at the cost of thermal stability where it has a $\Delta T$ of 107 °C.

Figure 6. Compositional dependence of $T_g$, $T_c$ and $\Delta T$ in the $x = y$ series of the $x$BaS + $y$La$_2$S$_3$ + (1-x-y)GeS$_2$ system.
Figure 7. Compositional dependence of $T_g$, $T_c$ and $\Delta T$ in the $x$BaS = $2x$La$_2$S$_3$ series.

Figure 7. Compositional dependence of $T_g$, $T_c$ and $\Delta T$ in the $x$BaS = $2x$La$_2$S$_3$ series xBaS + yLa$_2$S$_3$ + (1-x-y) GeS$_2$ system.
Figure 8. Compositional dependence of $T_g$, $T_c$ and $\Delta T$ in the $2xBaS = xLa_2S_3$ series $xBaS + yLa_2S_3 + (1-x-y) GeS_2$ system.

For the $La_2S_3$ rich series, $2x = y$, a similar trend is seen in the $T_g$ and $T_c$ values. The high modifier, single phase glasses again have a fairly constant $T_c$ value whereas the $T_g$ values increase with increasing modifier content. This happens with greater effect than in the $BaS = La_2S_3$ (1:1 modifier ratio) series. The glass with the highest $T_g$ is the $17.5BaS + 35La_2S_3 + 47.5GeS_2$ composition with a $T_g$ of 607 °C, $T_c$ of 735 °C, and a $\Delta T$ of 128K. The $T_g$ of this composition is about 30K higher than that of the $25BaS + 25La_2S_3 + 50GeS_2$, and has a larger thermal stability by 21K making it a more optimal glass for high temperature infrared optics.

It was found that glasses in the $BaS$ rich series, $x = 2y$, possessed lower $T_g$ values than did glasses in the other two series, but that they have large $T_c$ values, which leads to the largest $\Delta T$ values for any glass series. The glass with the highest $\Delta T$ is the $35BaS + 17.5La_2S_3 + 47.5GeS_2$ composition, which has a $T_g$ value of 525 °C, a $T_c$ value
of 772 °C, and a ΔT of 247K. This series would be best choice for applications where a large thermal stability is important.

Table 2. Summary of the optimal glass forming compositions for each series in the xBaS + yLa$_2$S$_3$ + (1-x-y)GeS$_2$ glass system.

<table>
<thead>
<tr>
<th>xBaS + yLa$_2$S$_3$ + (1-x-y)GeS$_2$</th>
<th>$T_g$ (±5 °C)</th>
<th>$T_c$ (±5 °C)</th>
<th>ΔT (±10 °C)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>BaS = La$_2$S$_3$ series</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>20 + 20 + 60</td>
<td>554</td>
<td>668</td>
<td>114</td>
</tr>
<tr>
<td>25 + 25 + 50</td>
<td>570</td>
<td>677</td>
<td>107</td>
</tr>
<tr>
<td><strong>La$_2$S$_3$ rich series</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>12.5 + 25 + 62.5</td>
<td>589</td>
<td>742</td>
<td>153</td>
</tr>
<tr>
<td>15 + 30 + 55</td>
<td>599</td>
<td>726</td>
<td>127</td>
</tr>
<tr>
<td>17.5 + 35 + 47.5</td>
<td>607</td>
<td>735</td>
<td>128</td>
</tr>
<tr>
<td><strong>BaS rich series</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>30 + 15 + 55</td>
<td>518</td>
<td>759</td>
<td>241</td>
</tr>
<tr>
<td>35 + 17.5 + 47.5</td>
<td>525</td>
<td>772</td>
<td>247</td>
</tr>
<tr>
<td>40 + 20 + 40</td>
<td>531</td>
<td>695</td>
<td>164</td>
</tr>
</tbody>
</table>

In summary, Table 2 provides $T_g$, $T_c$ and ΔT values for all single phase glass compositions in each series. In applications where the highest $T_g$ value is important, compositions rich in La$_2$S$_3$ are the best choice as they tend to yield the highest $T_g$ values. On the other hand, if large ΔT values are desired, then compositions high in BaS should be chosen. While the series with equal lanthanum and barium sulfides have no distinct advantage in terms of $T_g$, $T_c$ or ΔT, the 20BaS + 20La$_2$S$_3$ + 60GeS$_2$ composition does provide a relatively large $T_g$ and great glass formability as glass disks can be readily produced.

6. **Structural Analysis in the BaS + La$_2$S$_3$ + GeS$_2$ Glass System**
6.1 Raman Spectroscopy

Raman spectroscopy was utilized in order to obtain a better understanding of the local structure of the glass network. Typical structures found in GeS$_2$ glass networks are shown in Fig. 9. Figures 10, 11, and 12 give the Raman spectra for all glasses synthesized in the BaS = La$_2$S$_3$, 2xBaS = La$_2$S$_3$ and xBaS = 2xLa$_2$S$_3$ series, respectively. A ten 10-second sample spectra were collected with a 5x objective at 100% power using a 488 nm laser on a Renishaw inVia Raman spectrometer. Spectra were also taken of low modifier content phase separated glasses with a high magnification 20x and 50x objective to determine the Raman spectra of the individual phase separated regions. Representative comparison spectra of the matrix and droplet phases (see Fig. 13) are given in Fig. 13. All of the Raman spectra of all the different glasses in all of the three different series glasses in this ternary system have a similar Raman spectra envelop. This arises from the common GeS$_4$ tetrahedron that is the main building block of the network structure of all of these different glasses. All of the spectra were normalized such that the highest intensity band has the same intensity for each composition in each spectra set.
Figure 9. Fundamental Ge
 tetrahedral units in germanium sulfide glasses.
Figure 10. Compositional dependence of the Raman spectra of $x = y$ glasses and two binary glasses in the $x\text{BaS} + y\text{La}_2\text{S}_3 + (1-x-y)\text{GeS}_2$ system.

Figure 11. Compositional dependence of the Raman spectra of glasses in the $2x\text{BaS} + x\text{La}_2\text{S}_3 + (1-3x)\text{GeS}_2$ series.
The spectra for pure glassy GeS$_2$ glass is well established in the literature and our Raman spectra here, Fig. 10, for example, is in good agreement with those studies [1,2]. The most intense peak located at 342 cm$^{-1}$ is associated with $A_1$ symmetric stretching of the Ge$^4$ corner sharing tetrahedral units, or a tetrahedral unit that bonds to four other Ge tetrahedra by four bridging sulfur (BS) ions [2,3]. The vibrational mode centered at 372 cm$^{-1}$ is assigned to the $A^1_2$ companion to the $A_1$ mode, which is also associated with corner sharing Ge$^4$ tetrahedra [2]. The germanium tetrahedral vibrational mode centered at 435 cm$^{-1}$ is due to edge sharing tetrahedra where two germanium tetrahedra share two of the same sulfur atoms compared to only sharing one when connected at the corners [1].

The Raman spectra of the lowest total modifier content glass in each series, as well as the 0 + 7.5 + 92.5 binary glass, have spectra similar to the GeS$_2$ glass. However, at 260 cm$^{-1}$ a small peak associated with vibrations of S$_3$Ge-GeS$_3$ unit is detected [4]. This structural unit is due to a small deficiency of sulfur, which leads to a Ge-Ge homopolar bonds between Ge atoms. While these Raman spectra look similar, there are slight...
variations in relative peak intensity. The peak centered at 372 cm\(^{-1}\) in the Raman spectra of 5 + 10 + 85 glass has a higher intensity than that in the Raman spectra of the 10 + 5 + 85 glass. This is likely the result of more Ge\(^0\) short range order (SRO) structural units, where as shown above in Fig. 9, above, there are zero BSs and 4 non-bridging sulfurs (NBS), units in the matrix phase in the 5 + 10 + 85 composition glass. In this case, it is expected that La\(_2\)S\(_3\) forms three times as many NBSs compared to BaS. Another possible explanation is that it could be an artifact of phase sampling. For example, using the larger field of view lower magnification 5x objective of the Raman microscope that averages over a larger surface area of the sample, estimated to be \((200 \mu m)^2\), moving to different locations on the sample does yield slightly different Raman spectra as slight difference in sampled phases occurs. This makes assigning percentages of structural units to phase separated glasses impossible with these experiments.

At the highest modifier content where maximum fractions of NBSs are formed, all Ge tetrahedra should be in the Ge\(^0\) SRO unit and the glass should have excess sulfur that cannot be associated with Ge\(^0\) tetrahedral units. In these spectra, there is a shift in the most intense vibrational mode from 342 to 372 cm\(^{-1}\). Since the \(A_1\) mode decreases with increasing modifier concentration so should the \(A_1\) companion mode thus indicating a new structural feature has formed at 372 cm\(^{-1}\). It is well established that the mode at 372 cm\(^{-1}\) is typically due to the stretching of Ge\(^0\) unit [2,5]. Also, in these spectra, an additional peak centered at 475 cm\(^{-1}\) is observed. This has been associated with the free sulfurs in the structure forming sulfur chains with Ba\(^{+2}\) cations in the glass [6]. These polysulfide chains could explain the glass formation at the extreme ends of modifier content as the chains span through the liquid and prevent crystallization of the anionic species. The last peak centered at 212 cm\(^{-1}\) is associated with La-S bonds. However, it is difficult to assign a structural unit as the Raman cross section of La is small and addition features may not be observable. Similar lanthanum peaks have been shown in other works [4].

As reported by others [2,5], there are distinct changes in the Raman spectra of glasses when the composition changes from predominantly Ge\(^4\) (0% total modifier, BaS + La\(_2\)S\(_3\)) to Ge\(^0\) (67% total modifier). In those works, the most intense peak in the Raman spectra gradually changed from indicating a fully bridging tetrahedral Ge\(^4\) structure to a completely molecular anionic Ge\(^0\) structure, as expected from the composition. At 0% (BaS + La\(_2\)S\(_3\)), there are no anionic NBS structures and the Ge tetrahedral are comprised solely of Ge\(^4\) units. At 2BaS + GeS\(_2\) (67%BaS + 33%GeS\(_2\)), this gives rise to the SRO structure (Ba\(^{+2}\))\(_2\)GeS\(_4\)^{4+} and this is the Ge\(^0\) SRO unit. Likewise, at 2/3La\(_2\)S\(_3\) + GeS\(_2\) (40%La\(_2\)S\(_3\) + 60%GeS\(_2\)) this gives rise to the SRO structure (La\(^{+3}\))\(_{4/3}\)GeS\(_4\)^{4+} which is also the Ge\(^0\) SRO unit. As seen in the Raman spectra
for each of these series, the glasses do not exhibit a majority of Ge$^3$ Ge$^2$ or Ge$^1$ units, and, rather it appears to be a proportional mix of the two end spectra. We hypothesize that, because these glasses phase separate out a modifier rich phase, the germanium tetrahedral units in the matrix phase are converted into Ge$^0$ units, while the droplet phase consist predominantly of Ge$^4$ units. This behavior can be explained as each one mole% addition of BaS and La$_2$S$_3$ adds eight additional moles of NBS to the glass network where each barium cation adds two and each lanthanum cation adds three NBS. This gives rise to the rapid shift from fully bridging Ge$^4$ tetrahedral units to the molecular anionic Ge$^0$ units in the matrix phase for phase separated glasses. At high modifier contents, the phases combine forming predominantly Ge$^0$ units with a small fraction of Ge$^1$ units that vibrate at around 400 cm$^{-1}$. This hypothesis is supported as Raman spectra from each phase were collected with the much smaller field of view 20x and 50x objective for phase separated samples and the comparison between the two phases in the 5 + 5 + 90 composition are shown in Fig 13. The droplet phase spectrum is consistent with the GeS$_2$ spectra, while the matrix phase is more consistent with more heavily modified glass of Ge$^0$ structure. This behavior is consistent across all phase separated compositions.

Typically, as a glass becomes more depolymerized the $T_g$ decreases. However, an increase in the $T_g$ is observed here. This suggests that an ionic bonding scheme centered perhaps on the GeS$_4^{4-}$ molecular anion or barium polysulfide chains may be bridging the heavily modified glassy materials together.
6.2 Infrared Spectroscopy

a. Dilute concentration for Structural Studies

Infrared spectra of the glass are shown in Figs. 14, 15 and 16 for \( x \)BaS = \( y \)La\(_2\)S\(_3\), 2xBaS = \( x \)La\(_2\)S\(_3\) and \( x \)BaS = 2xLa\(_2\)S\(_3\) series, respectively. Sample pellets were produced by milling together 7.5 mg glass and 200 mg CsI and pressing the resultant powder into a pellet die. The linewidth of the peaks suggests that all compositions are in the glassy state as glasses have a larger distribution of bond angles compared to crystals giving rise to significantly broader peaks for glassy compared to polycrystalline samples. The IR spectra average mm distances of the pressed pellet samples and as a result the identification of different phase separated covalent Ge\(^4\) regions from the more ionic Ge\(^0\) phases is not possible. As both phases are present in the CsI pellet, their absorbances are combined and are roughly proportional to the concentration of the phase in the pellet as their molar absorptivities are likely to be similar.
The IR spectra of GeS$_2$ has been well documented in the literature and our spectra reported in Figs. 14, 15, and 16 are very consistent with those reported in the literature, see for example Ref. [3]. The most intense peak in the IR spectra for the base GeS$_2$ glass is located at 376 cm$^{-1}$ followed by near equally intense shoulders at 335 and 430 cm$^{-1}$. These peaks are associated with different vibrational modes associated with Ge-S vibrations in GeS$_{4/2}$ tetrahedron. For the heavily modified ternary glasses, these peaks have more pronounced overlap such that it appears to be one continuous peak making deconvolution more difficult. At highest modifier contents this band shifts to 397 cm$^{-1}$ with the transformation to Ge$^0$ units [2,7,8]. The next peak centered at 240 cm$^{-1}$ is associated vibrations of La-S bonds [4], and the intensity of this peak grows as the concentration of lanthanum sulfide increases as expected with increasing La$_2$S$_3$ content. Overall, the progression of germanium tetrahedra is very similar to the trend found in the Raman spectra seen in the previous section. The one major exception is that the La-S bonds are less pronounced in Raman spectra as lanthanum has a small Raman cross section.

The additional peaks are attributed to vibrational modes associated with oxygen contamination in the glass and are typically observed to some extent. The peak centered at 825 cm$^{-1}$ is associated with Ge-O vibrational modes [2,9] while the peak centered at 675 cm$^{-1}$ is assigned to La-O vibrations. It is worth noting that the only
sample to contain the 825 cm\(^{-1}\) peak is the pure germanium sulfide glass. The other samples do not contain this peak but may instead exhibit the peak at 675 cm\(^{-1}\) associated with La-O vibrations. This suggests that lanthanum has a higher affinity for oxygen than germanium does in this system. Lastly, no spectral features past \(~125\) cm\(^{-1}\) can be resolved as the cesium iodide vibrates at that wavenumber eliminating transmission through the composite sample.

The most intense peak in the low modifier compositions is centered at 343 cm\(^{-1}\). This peak is associated with the symmetric stretching, \(A_1\), vibrational mode of Ge\(^4\) tetrahedra where each germanium tetrahedron is connect to four other germanium tetrahedra at the corners. The second most intense peak in low modified glasses is located at 370 cm\(^{-1}\). This peak is due to vibrations of \(A_c^1\) companion mode to the \(A_1\) Ge\(^4\) stretching. The last peak seen in these spectra is located at 435 cm\(^{-1}\). This peak is due to edge sharing tetrahedron where two tetrahedra share two of the same sulfur ions, instead of sharing one like corner sharing tetrahedra. With increasing modifier concentration the peak at 343 cm\(^{-1}\) begins to decrease in intensity and with the must its companion mode at 370 cm\(^{-1}\). However, despite the concentration of Ge\(^4\) units decreasing with increasing modifier concentration, the peak at 370 cm\(^{-1}\) is growing. This is due to an increasing number of Ge\(^0\) units being formed by modifier additions, which overlaps with the \(A_c^1\) companion mode of the Ge\(^4\) tetrahedral stretching. The main difference between the three series’ spectra is the fact that the intensity Ge\(^0\) peak in the \(x\)BaS = \(2x\)La\(_2\)S\(_3\) compositions grows quicker than that of the BaS = La\(_2\)S\(_3\), and \(2x\)BaS = \(x\)La\(_2\)S\(_3\) series. This can be explained as lanthanum sulfide adds 3 times more non-bridging sulfurs than barium sulfide does. This means Ge\(^0\) unit formation is much slower in those two series resulting in lower intensity at 370 cm\(^{-1}\) when comparing analogues.

b. Thick polished disc studies for transmission range measurements

Figure 17 shows the combined infrared spectra from \(~2\) to 14 \(\mu\)m for GeS\(_2\) and 20BaS + 20La\(_2\)S\(_3\) + 60GeS\(_2\) polished glass disks glass samples. Sample spectra were normalized and then averaged to obtain finalized transmission spectra for each composition. These measurements are important to insure that the glasses transmit well into to IR range so that they can serve as appropriate IR optical fiber sensors.

The GeS\(_2\) glass transmits light from \(~0.5\) to \(~10.7\) \(\mu\)m with a region centered at 7.5 \(\mu\)m where transmission drops to near 0\%. This absorbance is due to Ge-O vibrational modes that arise from Ge-O impurities in the glass. There are also peaks centered at about 3 and 4 \(\mu\)m that are due to –OH and –SH vibrations respectively, and all impurity peaks have been reported by others. The 20-20-60 composition transmits over a similar region however, it has much lower absorbance at 7.5 \(\mu\)m and the peaks at 3 and 4 \(\mu\)m are slightly shifted to higher wavelength and are more broad. This broadening is likely
spectral broadening due to multiple cations present forming a variety of X-OH bonds, where X = Ba, La, and Ge, slightly altering vibration wavelength. The long wavelength cutoff for these samples are at 10.7 µm for the GeS2 samples and 10.2 µm for the 20-20-60 samples. This is due to a combination of the materials strong Ge-S and to a lesser extent Ge-O vibrations. In ternary samples La-S vibrations also absorb heavily in this region.

Figure 17. Infrared absorption spectra through ~ 2mm thick polish lenses of base GeS2 and modified 20BaS + 20La2S3 + 60GeS2 glasses. IR transmission extends down to ~ 10 µm and shows that these glasses would be appropriate for use as IR transmission fiber optics.

The ternary 20-20-60 glass exhibits at least 50% of their maximum transmission from about 0.5 to about 9 µm with a slight interruption from –OH bonds between 2.8-3.5 µm. The GeS2 glass exhibits more impurity absorptions than the ternary glasses. The GeS2 glass transmits from about 0.6 to about 7.2 µm with impurity bands in the region from 3.8- to 4.1 um also due to -OH vibrations and from 7.2-9.7 µm with an interruption from 7.2-8 µm due to Ge-O vibrations. Lastly, that the most intense absorber transitions
from Ge-O vibrations in the germanium sulfide glasses to –OH vibrations in the 20-20-60 compositions.

As with many sulfide glasses, there exists some trace amount of water and oxygen contamination. With purification technique such as sulfur distillation and sulfurization of batch materials by passing H2S gas over them, the purity of these glasses could be improved.58,59 While these techniques were not available for this study, the improved purity would allow for better transmission across the infrared light spectrum and open up new opportunities in the chemical sensing field for these materials. If all contaminants could be removed from the samples the transmission window would span from approximately 0.5 to 10.5 µm.

Even with these contaminate, in the glasses many different chemical species can be detected in the infrared spectrum, and by removing them even more can be detected. Some of the important chemical species that can be detected with these materials are vibrations associated with methane, sulfur dioxide and ammonia from 1300-1700 cm-1 and carbon monoxide and carbon dioxide from 2,100-2,400 cm-1. Many of these chemical species could be especially important for combustion reactions, but can be found in many other processes as well.

While the fingerprint region of the infrared spectrum, from 1400-600 wavenumbers, will be blocked by the intense sulfur bonds in sulfides glasses, there is still a large region of the infrared spectrum that can be utilized for chemical sensing. Many important chemical species are found in the region 4000-1400 cm-1 such as carbon, nitrogen, and hydrogen species. This is a tradeoff as many telluride and selenide compositions can detect chemicals in the molecular fingerprint region, but they cannot operate at elevated temperature. Luckily, most chemical species have multiple characteristic vibrational modes that occur at different locations in the infrared spectrum. In contrast to selenides and tellurides, oxides have significantly higher glass transition, crystallization and melting temperatures. With this comes a reduction in infrared transmission. Pure silica has one of the highest Tgs for traditional glasses at 1200 °C, but only transmits to about 4 µm depending on impurities. While changing glass formers and modifiers can influence these two properties, typically infrared transparency can only be extended to approximately 7 µm as shown in 60 TeO2 – xBi2O3 – (30–x)B2O3 – 10ZnO glasses.60 Again, with these strong IR transmitting oxides they have Tgs below 400 °C for compositions tested exhibiting the strong trade off of these two characteristics.
7. Glass Disk and Rod Formability

Many attempts were made to produce sample disks for irradiation and glass rods for fiber pulling. A series of experiments changing ampoule coating, crucible melting, and processing parameters was conducted in the hopes of defining a process for the consistent formation of glass disks and rods. The most successful approach to disk formation was changing the processing parameters. The annealing temperature, dwell time, and ramp down rate were varied in a series of experiments to come up with the ideal processing parameters of melting at 1150 °C for 12 hours then air quenching for 10-20 seconds followed with annealing at 555 °C for 45 mins and ramping down to room temperature at 0.6 °C/min. With dedication, glass disks made of GeS$_2$ and 20BaS – 20La$_2$S$_3$ – 60GeS$_2$ could be formed with relative ease. This was not the case with preform rods. As glass disks were cut from small rods with length of 3-5 cm, less thermal stresses would form. As more material was added to form a larger rod preform, length greater than 10 cm, the thermal stress on the preform rod was great enough to cause fracture. Forming fiber preforms was the most difficult portion of this project and further work on utilizing a variety of containment and coatings would promote the formation of preforms.

That said, a pure GeS$_2$ sample was drawn into prototype fiber in order to validate the potential for these systems to be fiberized. More specifically, a rod of 10 mm approximate diameter and about 14 mm of drawable length was heated to a temperature of 680°C in a custom soft-glass furnace specially designed for chalcogenide and fluoride glass fibers. Figure 18 provides an image of the resultant fiber and preform remnants.

![Figure 18. Optical image of GeS$_2$ fiber drawn and remnants of the preform.](image-url)
While fiber was drawn, the small preform size was insufficient to yield longer lengths of higher quality fiber. However, the pure GeS₂ did draw into fiber and represents a validation of the fiberization capability of these refractory chalcogenide glasses.

8. Gamma Irradiation and Characterization of Glass Disks

Glass disks were irradiated using a 3000 Curie Co-60 collimated gamma irradiator and a 1300 Curie Co-60 gamma Bunker at PNNL. Ten glass samples in the form of 10 mm diameter by 2 mm thick discs were irradiated. The samples include four 20BaS + 20La₂S₃ + 60GeS₂ samples, and six GeS₂ samples. In order to prevent oxidation, the discs were sealed under inert argon in a ½” diameter silica ampoule prior to irradiation. The ampoule was also pumped down to 100 mTorr to prevent over pressure due to potential material vaporization. The chamber was heated to 300 °C using a coiled heater during the irradiation to simulate a high temperature reactor environment. Three irradiation runs were conducted where each run gave 256, 450 and 1805 kGy Air-Kerma dose respectively, for a total Air-Kerma dose of ~2.5 MGy.

![Figure 19. Dose dependence of density in GeS₂ disk samples.](image1)

![Figure 20. Dose dependence of density in 20BaS + 20La₂S₃ + 60GeS₂ disk samples.](image2)
Figures 19 and 20 show the dose dependence of density for GeS$_2$ and 20BaS + 20La$_2$S$_3$ + 60GeS$_2$ samples, respectively. For each composition, the average density is reported at each dosage. Both sample sets show similar trends where there is a slight decrease in density at the lowest dosage, followed by a slow, steady increase thereafter. The ternary samples showed less density change than the GeS$_2$ samples, which may suggest they are more radiation hard than the base GeS$_2$ glass samples. The density never reaches the initial value for either sets of samples with the received dosages, but with increasing dosage, there is a minimal change in density, which shows promise and suggest these samples can withstand MGy doses of radiation.

Figures 21 and 22 show the dose dependence of the refractive index at 633 nm for GeS$_2$ and 20BaS + 20La$_2$S$_3$ + 60GeS$_2$ samples respectively. For each composition, the average refractive index is reported at each dosage. In both samples, the difference in refractive index at different dosages is very small at 0.0029 for GeS$_2$ samples and 0.004 in ternary samples. With such a small change in refractive index from radiation, it is possible to engineer a core and cladding glass such that internal reflection can be achieved and maintained in a radiation environment as slight changes in refractive index, on the order of 0.05 - 0.1, can be produced with 1-5 mol% change in
composition. This change due to chemistry is much larger than the change arising from irradiation.

Figures 22 and 23 show the transmission spectra over a wavelength range from 0 to 14 µm for the pure GeS₂ and the 20BaS-20La₂S₃-60GeS₂ composition glass disks, respectively, at different gamma irradiation dosages. Curves were shifted up incrementally 20, 40, and 60% with increasing dosage for clarity, and only infrared transmission was collected at 256 kGy dose. For both samples there is little change in the shape of the spectra.

The germanium sulfide samples transmit infrared light from 0.5-10.7 µm with a region centered at 7.5 µm where transmission drops to near 0%. This absorbance is due to Ge-O vibrational modes. There are also peaks centered at about 3 and 4 µm that are due to –OH and –SH vibrations respectively, and all impurity peaks have been reported by others [10]. The 20 + 20 + 60 composition transmits over a similar region however, it has much lower absorbance at 7.5 µm and the peaks at 3 and 4 µm are slightly shifted to higher wavelength and are broadened. This broadening is likely due to multiple cations present forming a variety of X-OH bonds, where X= Ba, La, and Ge, slightly altering vibration wavelength. The long wavelength cutoff for these samples are at 10.7 µm for the GeS₂ samples and 10.2 µm for the 20 - 20 - 60 samples. This is due to a combination of the materials strong Ge-S and to a lesser extent Ge-O vibrations. In ternary samples, La-S vibrations also absorb heavily in this region.
As a result, the ternary BaS + La₂S₃ + GeS₂ glasses transmit from ~0.5 to ~9 μm. The glasses prepared here were not highly purified as is the case with many non-oxide glasses and as a result, our glasses exhibit modest IR absorption in the spectra region of 2.8-3.5 μm arising from contaminating –OH estimated. Using a typical molar extinction coefficient of ~ 50 l/mole-cm and a glass sample thickness of 0.2cm, we estimated the –OH contamination to be in the range of about 100 to 1,000 ppm. Finally, the most intense contamination absorption bands are at about 10 μm and arise from Ge-O vibrations in the glass. A similar estimation based upon the IR absorption suggests that the ≡Ge-O-Ge≡ contamination in the glass is in the range of about 1,000 ppm. These contaminations can be removed by purifying the glass using known methods and as such should not present a huge barrier to these glasses being used as IR fiber optics.

As with many sulfide glasses, there exists trace water and oxygen contamination. With purification techniques such as sulfur distillation and sulfurization of batch materials by passing H₂S gas over them, the purity of these glasses could be improved [11,12]. Many studies have shown that this purification eliminates a significant portion of contaminates, but some oxygen contamination may persist when melting in SiO₂ ampoules that do not have a robust enough coating to prevent contact of the melt and silica. While these techniques were not available for this project, the improved purity would allow for better transmission across the infrared light spectrum and open up new opportunities in the chemical sensing field for these materials. If all contaminants could be removed from the samples the transmission window would span from approximately 0.5 to 10.5 μm.

Even with these contaminant ions many different chemical species can be detected in the infrared spectrum, and by eliminating them even more species can be detected. Some of the important chemical species that can be detected with these materials are vibrations associated with water and hydrogen fluoride from 2.4 to 2.8 μm, carbon monoxide and carbon dioxide from 4.2 to 4.4 μm, and methane from 7.3–8.1 μm. Many of these chemical species are contaminants, corrosion by-products, or reaction by-products that can potentially exist in the AdvSMR headspaces [17]. Furthermore, gamma radiation up to an accumulated dose of 2.5 MGy at a temperature of 300 °C didn’t affect the transmission spectra of these glasses. This further establishes the potential of using these materials in AdvSMR applications.

9. Summary

In summary, a judicious set of ionic non-oxide glasses in the xBaS + yLa₂S₃ + (1-x-y)GeS₂ system were selected and studied under this program. The optimal composition along with the GeS₂ base glass were synthesized and characterized at elevated
temperature under high radiation flux. The ternary glass demonstrated a glass transition temperature higher than 500°C, over 100°C thermal stability, a broad 0.5 to 10.5 um transmission window and radiation hardness up to 2.5MGy gamma irradiation at a temperature of 300°C. Challenges arose in available material purification technique for completely eliminating absorptions from the –OH bonds. Also long preform rods for pulling fibers were difficult to produce without cracking. Future studies are suggested to focus on overcoming barriers in preform rods formation and drawing optical fibers using optimized compositions. Radiation studies may be performed at higher radiation doses and temperatures with a planned improved gamma source at the PNNL, allowing 6000 Curie (1.7 Mrad/hour) vs. 1300 Curie (380 Krad/hour) in the current study.

Publications and Presentations resulting from this program


Papers in Progress
9. Josh Roth, Steve W. Martin, Hong Amy Qiao, Norm Anheier, Mark Murphy, Jacob Peterson, “Structural Analysis and Glass Formability Studies in the xBaS + yLa$_2$S$_3$ + (1-x-y)GeS$_2$ Glass System,”

10. Josh Roth, Steve W. Martin, Hong Amy Qiao, Norm Anheier, Mark Murphy, Jacob Peterson, “Gamma Irradiation of Novel Sulfide Based Glasses at Elevated Temperature.”

References


5 Yao, W., Berg, K. & Martin, S. Structure and properties of glasses in the MI + M$_2$S + (0.1Ga$_2$S$_3$ + 0.9GeS$_2$), M = Li, Na, K and Cs, system. *Journal of Non-Crystalline Solids* **354**, 2045-2053 (2008).


