Nonlinear properties of chalcogenide glass fibers

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Chalcogenide glasses have demonstrated high third-order Kerr ($\chi^{(3)}$) nonlinearities up to 1000x higher than silica glass which make them attractive for applications such as nonlinear switching, optical regeneration, Raman amplification, parametric amplification, and supercontinuum generation. Poling of chalcogenide glasses to induce an effective second order ($\chi^{(2)}$) nonlinearity has also been demonstrated and opens the possibility for the use of poled glass waveguides for applications such as frequency conversion or electro-optic modulation. Stimulated Brillouin scattering (SBS) has also been investigated in As$_2$S$_3$ and As$_2$Se$_3$ single-mode fibers. The threshold intensity for the stimulated Brillouin scattering process was measured and used to estimate the Brillouin gain coefficient. Preliminary results indicate record high values for the figure of merit and theoretical gain, compared to silica, which bodes well for slow-light based applications in chalcogenide fibers.

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1. Introduction

Chalcogenide glasses are based on the chalcogen elements S, Se and Te with the addition of other elements such as Ge, As and Sb to form of stable glasses [1]. Due to their large IR transparency, fibers fabricated from these glasses are ideal for transmission of high power IR light. Several applications of chalcogenide fibers for IR transmission have been documented [2]. Also of interest is the high nonlinearity of these glass compositions. The high $\chi^{(3)}$ nonlinearities of chalcogenide glasses make them excellent candidates for applications such as all nonlinear switching, optical regeneration, Raman amplification, parametric amplifiers and supercontinuum generation. We have extensively studied the nonlinearities of the As-S-Se based system. High strength low loss fibers can be drawn from this system. In this paper, we will report on the nonlinear properties of this system.

2. Glass preparation

Chalcogenide glasses are melted directly in quartz ampoules using chemicals purified via distillation/sublimation [3]. Typical melt temperatures range from 600°C to 900°C, depending upon composition. The liquids are quenched and the glass rods annealed at temperatures around the appropriate softening temperatures. The optical fibers are obtained by the double crucible (DC) process [4]. The DC process enables adjustments to be made in the core/clad diameter ratio during fiber drawing by independent pressure control above each melt. Therefore both multimode and single mode fibers can be drawn with relatively few processing steps.

3. Fiber properties

Fig. 1 compares the losses routinely obtained for a couple of chalcogenide glasses along with the lowest ("champion") losses reported in the literature [5,6]. Depending upon composition, the sulfide, selenide and telluride based fibers transmit between about 0.8-7 µm, 1-10 µm, and 2-12 µm, respectively. Therefore, the practical applications dictate the type of fiber to be used. The As-S fibers have received the most attention to-date in our laboratory and so the loss routinely achieved is about 0.1-0.2 dB/m in fiber lengths of about 500 meters. Losses for As-Se fibers typically range from 0.5 to 1 dB/m in the near IR around 1.5 µm.

![Fig. 1. Transmission loss spectra of (a) lowest loss sulfide fiber, (b) typical sulfide fiber, (c) lowest loss telluride fiber, and (d) typical telluride fiber.](image-url)
4. Nonlinear properties

It is well established that the values of $\chi^{(3)}$ for chalcogenide glasses are about two orders of magnitude larger than silica [7,8]. More recently, glasses have been reported with non-linearities approaching 1000 times silica [9,10]. These large non-linearities would allow small compact low power devices for telecommunications. The subpicosecond response of these non-linearities is ideal for high data rate telecommunication devices.

For efficient nonlinear switches utilizing the optical Kerr effect, the nonlinearity must be high and the nonlinear absorption must be low. A figure of merit FOM $= n_2/(\beta \lambda)$ can be defined as a useful metric to determine optimum compositions, where $n_2$ is the nonlinear index and $\beta$ is the nonlinear absorption. For isotropic medium, one and two photon resonant processes dominate the third-order susceptibility. For frequencies approximately half of the material resonance, two photon processes resonantly enhance the nonlinear index $n_2$. Normally, however, the two photon resonance enhancement is accompanied by two photon absorption which competes with the nonlinear index $n_2$. In the case of amorphous materials such as chalcogenide glass, an exponential Urbach tail exists and its absorption edge extends below the half gap. This edge leads to two photon absorption (TPA) below the half gap and thus $n_2$ may increase faster than TPA absorption in this region. Consequently, we expect to find that the best performance in terms of nonlinear index strength vs. TPA (FOM) will occur just below the gap. Fig. 2 shows the bandgap of the As-S-Se system vs. Se concentration.

Spectrally resolved two beam coupling measurements of As-S-Se system have been formed to determine the magnitude of the nonlinear index $n_2$ and the two photon absorption coefficient. Details of these measurements can be found in [10]. Fig. 3 shows the results of these measurements. Values for As-S were found to be $\sim 220$ times silica at 1.55 $\mu$m and increased with Se substitution of S to a value of $\sim 930$ times silica for As-Se. Likewise, two photon absorption also increases with increasing Se content. We can use this data to calculate the FOM for the As-Se system (Fig. 4). As expected. The glasses with the larger FOM for operation at 1550 nm occurs for $E_g/h\nu$ at $\sim 0.45$ which is the As-Se composition. [11]

![Fig. 3. $n_2$ and TPA absorption of As-S-Se glass system.](image)

Here, the bandgap is defined at the point of $10^3$ cm$^{-1}$ absorption. In the graph, Se content of 0 at. % corresponds to pure As$_{60}$S$_{40}$ while Se content of 60 at. % corresponds to pure As$_{40}$Se$_{60}$. We note that the bandgap of glass system increases with Se content. For operation at 1.55 $\mu$m (0.8 eV), we would expect an optimum composition of As$_{50}$Se$_{50}$ where $E_g/h\nu \sim 0.45$. We will see that this is borne out by experimental data.

![Fig. 2. Bandgap of As-S-Se glass system. Bandgap is defined at the point of $10^3$ cm$^{-1}$ absorption.](image)

5. Raman amplification

Fig. 5 shows the normalized Raman spectra of As$_{60}$S$_{40}$, As$_{40}$Se$_{60}$, and silica. As$_{40}$Se$_{60}$ glass has a much narrower Raman line ($\sim 60$ cm$^{-1}$) than silica glass ($\sim 250$ cm$^{-1}$). In addition, the Raman shift for As$_{40}$Se$_{60}$ glass is much smaller ($\sim 240$ cm$^{-1}$) than the Raman shift of silica glass ($\sim 440$ cm$^{-1}$) due to the heavier atoms present in the chalcogenide glass. Previous studies have looked at stimulated Raman scattering in As$_{40}$S$_{60}$ glass, a very similar glass system to As$_{40}$Se$_{60}$ [12]. These studies found the Raman gain coefficient of As$_{40}$S$_{60}$ to be almost two orders of magnitude higher than that of silica. It was also
found that this enhancement in the Raman gain roughly corresponded to the enhancement in the nonlinear index, \( n_2 \). Consequently, one might expect to see an even larger Raman gain coefficient in As<sub>2</sub>Se<sub>6</sub> since the selenide glass has shown an even larger nonlinearity and also a narrower Raman spectrum.

\[ G_d = \exp \left( \frac{g_R P_r L_{eff}}{A_{eff}} \right) \]  

(1)

Where \( g_R \) is the Raman gain coefficient, \( P_r \) is the pump power, \( A_{eff} \) is the fiber effective area and \( L_{eff} \) is the fiber effective length. The fiber effective length is given by

\[ L_{eff} = \frac{1}{\alpha} \left( 1 - e^{-\alpha \ell} \right) \approx \frac{1}{\alpha} \]  

(2)

Where \( \alpha \) is the fiber loss. For long lengths, \( L_{eff} \) is approx \( 1/\alpha \). From these equations, we can see that the gain is proportional to \( \exp (-g_R / \alpha) \) for long fiber lengths. Thus, we can use the value \( g_R / \alpha \) as a rough FOM for Raman amplification. Table 1 compares the performance of an As-Se Raman fiber laser or amplifier operating at 4 \( \mu \)m to a silica Raman fiber laser or amplifier operating in the telecommunications band at 1.5 \( \mu \)m. Here, the Raman gain coefficient of As-Se, \( g_R \), which is measured to be 780x silica at 1.5 \( \mu \)m is extrapolated to it value in the mid-IR since the Raman gain coefficient scales inversely with wavelength. \( \alpha \) is the fiber loss. For silica, a loss of 0.2 to 0.3 dB/km is typical of telecommunication grade fiber. For As-Se, two losses are given. The loss of 200 dB/km is typical of “champion losses” achieved at NRL for As-Se fiber while the loss of 0.1 dB/km is theoretical loss for As-Se fiber [14]. For the loss of 200 dB/km, we see that \( g_R / \alpha \) for an As-Se fiber Raman amplifier operating at 4 \( \mu \)m is about 0.38 compared to 1.1 for a silica fiber Raman amplifier. For the theoretical loss of 0.1 dB/km, we see that \( g_R / \alpha \) for As-Se fiber operating at 4 \( \mu \)m is 860 times that of silica fiber operating at 1.5-\( \mu \)m.

<table>
<thead>
<tr>
<th>( \lambda ) (( \mu )m)</th>
<th>( g_R ) (cm/W)</th>
<th>Loss (dB/km)</th>
<th>( \alpha ) (cm(^{-1}))</th>
<th>( g_R / \alpha ) (10(^{-6})W(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Silica Fiber</td>
<td>1.5 x 10(^{-12})</td>
<td>0.2 to 0.3</td>
<td>(-6 \times 10^{-7})</td>
<td>1.1</td>
</tr>
<tr>
<td>As-Se Fiber</td>
<td>4 x 10(^{-10})</td>
<td>200(^{11})</td>
<td>5 x 10(^{4})</td>
<td>0.38</td>
</tr>
</tbody>
</table>

Table 1. Figure of merit for Raman amplification in As-Se fiber at 4-\( \mu \)m compared Raman amplification in silica fiber at 1.5-\( \mu \)m. The loss value of 200 dB/km (a) for As-Se is typical of a “champion” loss value. The loss value of 0.1 dB/km (b) is theoretical loss from [14].

Stimulated Raman scattering (SRS) has also been observed in the IR. Fig. 7 shows the SRS in a 1 m length of As-Se fiber under CW CO laser pumping at \( \sim 5.4 \mu \)m. The SRS is seen at \( \sim 6.1 \mu \)m. Raman laser operating in the wavelength range of from 6.1 to 6.4 \( \mu \)m would have applications in laser surgery. These wavelengths correspond to amide II bands in tissues and studies have shown that ablation of soft tissue is possible at these wavelengths with minimal collateral damage, thus accelerating healing [15]. Modeling of a Raman laser operating at 6.45 \( \mu \)m under CO laser pumping at 5.59 \( \mu \)m has shown high slope efficiencies and moderate threshold power operation is possible [16].
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6. Supercontinuum generation

Supercontinuum generation has been demonstrated for between 2 to 3 µm in small core sulfide and selenide fibers as well as photonic crystal selenide fibers [17]. The 1 meter length of fibers were pumped with a Ti:sapphire laser at a wavelength of 2.5 µm using 100 fs pulses and 100 pJ/pulse. The outputs from the fibers are shown in figure 5. The sulfide and selenide fibers were 7 µm core diameter, while the PCF fiber had a 10 µm core diameter.

Broader supercontinuum can be realized by pumping near to the zero dispersion wavelength or in the anomalous dispersions region. Currently, however, pumping is limited by the tuning range of the OPA-800C to ~2500 nm wavelength. Tailoring the dispersion of the As-Se PCF fiber to shift the dispersion wavelength closer to the near IR to enable broader bandwidth supercontinuum is feasible. Work on chalcogenide PCF fiber has shown that that the minimum dispersion wavelength can be shifted over very broad ranges [17].

7. Poling of chalcogenide glass

Isotropic materials such as glasses lack a center of inversion symmetry and thus have no second order nonlinear susceptibility ($\chi^{(2)}$) they should not exhibit second harmonic generation (SHG) [18]. However, undoped and Pr-doped GaLaS glasses have exhibited SHG [19] through optical pumping. This SHG may be due to crystallization or the effect of frozen-in electric fields. The latter arises from the relationship $\chi^{(2)} = E_{fc}\chi^{(3)}$, where $E_{fc}$ is the frozen-in electric field [18]. Electric poling has been successfully used to produce SHG in silica based fiber systems [20]. It is not unreasonable to expect similar results in chalcogenide fibers.

Since $\chi^{(3)}$ is about 2 to 3 orders of magnitude larger in chalcogenides compared with silica, we expect larger SHG efficiencies in electrically poled chalcogenide glasses. However, the question arises as to whether the electric fields can be frozen-in for chalcogenide glasses. We have observed second harmonic generation at 780 nm using electrically poled arsenic sulfide glass when pumping a 1 mm thick arsenic sulfide glass disk at 1560 nm. The sample was electrically poled at 100°C for 5 hours under nitrogen gas atmosphere. At the present time the magnitude appears comparable to silica glass but the mechanism is unknown.

Fig. 7. SRS signal observed at 6.1 µm under ~5.4 µm CO laser pumping.

Fig. 8. Supercontinuum generation in small core chalcogenide fibers. The insert shows the cross-sectional view of the selenide PCF fiber.

Fig. 9. Second harmonic generation in poled As-S glass. Glass was pumped at 1.56 µm. Shown is 780 nm SHG signal.
8. Brillouin scattering

In order to determine the Brillouin gain coefficient, we measured the threshold power of the stimulated Brillouin scattering (SBS) process using the experimental setup detailed below in Fig. 10. The threshold power is easily determined by monitoring the spectrum of the reflected light using a high-resolution optical spectrum analyzer (OSA) as sampled by the circulator which is the interface between the pump delivery system (DFB laser source plus Er amplifier, EDFA) and the chalcogenide fiber. The fiber was coated with liquid gallium on 10-cm lengths on each end to eliminate the radiation leaking into the cladding. The fiber ends were not anti-reflection coated and hence cavity effects were significant due to the high refractive index of the fiber. The losses in the fiber, and the coupling optics (4% for the focusing lens, 14% for the collimating objective), along with the Fresnel loss at the fiber ends (17.7% for As$_2$S$_3$ and 22.6% for As$_2$Se$_3$) are all taken into account in throughput measurements used to estimate the coupling efficiency, and hence the amount of pump launched into the core. We estimate 45% coupling efficiency in the As$_2$S$_3$ case, and 37% in the As$_2$Se$_3$ case. In the future, the coupling efficiency can be optimized and hence the SBS threshold power can be reduced, which is a desired trend from a system design perspective.

The spectral changes of the backward wave propagating through the chalcogenide fiber, as sampled by the circulator, are shown in Fig. 11 for the As$_2$S$_3$ fiber, and in Fig. 12 for the As$_2$Se$_3$ fiber. The cavity effects reduced the accuracy with which we were able to determine the threshold as indicated in the captions. Nevertheless, the threshold is easily identified by the significant jump in the peak of the Brillouin-shifted signal monitored on the OSA. Additionally, we observed clamping of the pump output power as, once the threshold is reached, most of the pump power is transferred to the Stokes wave [21].

**Fig. 10.** Experimental setup used for SBS threshold measurements.

**Fig. 11.** Typical spectra of the reflected light sampled by the circulator for different launched pump powers into the As$_2$S$_3$ fiber core. Fiber length was 10.0 m. Estimated SBS threshold: $(27 \pm 3)$ mW. Tick labels shown only on one plot for clarity.

**Fig. 12.** Typical spectra of the reflected light sampled by the circulator for different launched pump powers into the As$_2$Se$_3$ fiber core. Fiber length was 5.0 m. Estimated SBS threshold: $(127 \pm 7)$ mW. Tick labels shown only on one plot for clarity.

The numerical aperture (NA) of a fiber, essentially the contrast in index between the core and the clad, is an important parameter. It determines the mode-field diameter and hence the effective area of the fundamental mode, with direct implications on the threshold power estimation for stimulated Brillouin scattering. It also determines the number of modes supported by the fiber at a given wavelength, $\lambda$. The $V$-number for a step-index fiber is a function of NA as given in Eq. 2, where $d$ is the core diameter:
Using Eqs. 4-6, the parameters from Table 2, and the fiber lengths and pump threshold values indicated in Fig. 3 and Fig. 4, we determined the Brillouin coefficient to be \((3.9 \pm 0.4) \times 10^{-9}\) m.W\(^{-1}\) for the As\(_2\)S\(_3\) and \((6.75 \pm 0.35) \times 10^{-9}\) m.W\(^{-1}\) for As\(_3\)S\(_3\). The value for the As\(_2\)Se\(_3\) is close to the only other previously published result for this composition [22]. The value for the As\(_2\)S\(_3\) fiber, although lower than the one for As\(_3\)Se\(_3\), is still two orders of magnitude higher than that for fused silica \((-4.4 \times 10^{-11}\) m.W\(^{-1}\)) [22,24].

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1 m and a standard pump power of 1 mW. Then, the theoretical gain is given by Eq. 9:

$$G_{th} [\text{dB}] = 4.34 g_b k \times 1 \text{mW} \times L_{\text{eff}} |_{L=1 \text{m}}$$

(10)

We used this last, fairly simple expression to compare the most representative fibers considered so far: silica [25,26], high-nonlinearity bismuth fiber [27,28], $\text{As}_2\text{Se}_3$ fiber [22], along with the results reported here. The comparison is provided in Table 3, with all the data reported for experiments without polarization control ($k=0.5$). We also include the FOM as defined above for completion. One can easily notice the significant increase in the theoretical gain (or FOM) for the $\text{As}_2\text{S}_3$ fiber due to its smaller core, lower loss and slightly reduced refractive index.

Table 3. Comparison of figure of merit for slow-light based applications at 1.56 µm.

<table>
<thead>
<tr>
<th>Fiber</th>
<th>$n$</th>
<th>$A_{\text{eff}}$ [m$^2$]</th>
<th>Loss [dB.m$^{-1}$]</th>
<th>$L$ [m]</th>
<th>$\text{Left}$ [m]</th>
<th>$g_0$ [m.W$^{-1}$]</th>
<th>$G_{th}$ [dB]</th>
<th>FOM [dB.W$^{-1}$.m$^{-1}$]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Silica</td>
<td>1.47</td>
<td>2.22</td>
<td>0.01</td>
<td>2.0</td>
<td>2.0</td>
<td>4.40x10$^{-11}$</td>
<td>0.076</td>
<td>1</td>
</tr>
<tr>
<td>$\text{Bi-HNL}$ [27]</td>
<td>2.81</td>
<td>2.81</td>
<td>0.84</td>
<td>5.0</td>
<td>3.23</td>
<td>6.10x10$^{-9}$</td>
<td>1.084</td>
<td>77</td>
</tr>
<tr>
<td>$\text{As}_2\text{Se}_3$ [22]</td>
<td>2.81</td>
<td>0.91</td>
<td>0.90</td>
<td>5.0</td>
<td>3.1</td>
<td>6.75x10$^{-9}$</td>
<td>0.719</td>
<td>51</td>
</tr>
<tr>
<td>$\text{As}_2\text{S}_3$</td>
<td>2.45</td>
<td>0.90</td>
<td>0.57</td>
<td>5.6</td>
<td>3.90x10$^{-9}$</td>
<td>3.398</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

9. Conclusions

The large nonlinearities and fast response of the nonlinearity of the As-S-Se system make fibers drawn from these glasses well suited for optical switches, optical regenerators for high speed telecommunication systems. Use of these materials will allow compact devices cm’s in length with optical powers <1W peak power (1 ps in 1 ps pulses). The large Raman gain of the As-S-Se fibers coupled with the large IR transparency make these well suited for compact Raman amplifiers for telecommunications as well as fiber lasers and amplifiers in the mid-IR. These high nonlinearities also allow efficient supercontinuum generation which is useful for broadband sources in the near and mid-IR. Finally these materials can be poled to induce an effective $\chi^{(2)}$ opening up the potential of waveguide parametric amplifiers.

The stimulated Brillouin scattering process was studied in $\text{As}_2\text{S}_3$ and $\text{As}_2\text{Se}_3$ single mode fibers. Values of the Brillouin gain coefficient were measured to be $(3.9 \pm 0.4) \times 10^{-9}$ m.W$^{-1}$ and $(6.75 \pm 0.35) \times 10^{-9}$ m.W$^{-1}$, respectively. An analysis of the figure of merit for slow-light based applications indicates that the smaller core $\text{As}_2\text{S}_3$ fiber performs best due to the lower loss, reduced core size and slightly lower refractive index. The configuration using the small-core $\text{As}_2\text{S}_3$ fiber yields a figure of merit which is about 140 times larger, or a theoretical gain about 45 times larger, than the best silica-based configurations reported to date.

The continued improvement of chalcogenide materials will make such devices feasible in the near term.

References

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