Femtosecond laser induced two-photon absorption in Au-ion embedded glasses

Rabia Ahmad1, M. Shahid Rafique1,2, Ammar Ahmed3, Ali Ajami4, Pavla Nekvindova5, Blanka Svecova5, Shazia Bashir6 and Saman Iqbal1

1Department of Physics, University of Engineering & Technology, 54000 G. T. Road, Lahore, Pakistan; 2Institute of Applied Physics, Vienna University of Technology, Wiedner Hauptstrasse 8-10 Wien-1040, Austria; 3Amido, 43 Worship St, London EC2A 2DW, UK; 4Faculty of Physics, Semman University, 35131-19111 Semnan, Iran; 5Department of Inorganic Chemistry, Faculty of Chemical Technology, Institute of Chemical Technology, Technicka 5, 166 28 Prague, Czech Republic and 6Centre for Advanced Studies in Physics Government College University, Lahore, Pakistan

Abstract

Two-photon absorption (TPA) of Au-ion irradiated glasses in the femtosecond regime has been analyzed by an open-aperture \( Z \) scan technique. Three types of glasses, namely GIL49, BK7, and Glass B were irradiated by using 1700 keV Au\(^+\) ion beams. Samples were post-annealed at 600° C for 5 h. Penetration depth and distribution of Au\(^+\) ions having 1700 keV energy within glass substrates were estimated by transport of ions in matter (TRIM) simulations. Detailed calculations with full-damage cascades were performed for each sample, taking into account the chemical composition of glass substrates. TRIM results reveal that there is no significant change in ion range, straggling, and ion distribution with the change in the substrate composition. However, \( Z \) scan results showed a difference in TPA coefficients for all three glasses. Extent of crosslinking within each of irradiated sample, owing to its chemical composition, may have affected their TPA coefficients.

Introduction

Nonlinear optics has become an essential element of modern technologies invoking extensive research in the field (Armani et al., 2018; Carvalho et al., 2018; Khalbe et al., 2018; Xu et al., 2018). Optical nonlinearity arises from intense light-induced modifications in optical properties of a material. Response of the material therefore, depends nonlinearly on the strength of applied field. Experimental demonstration of this phenomenon turned out to be possible after the advent of first laser in 1960 when intense light, capable of producing such nonlinearity became a reality. Vast range of potential applications has inspired researchers to further investigate and analyze the outcomes in this field of research. Optical limiting, optical data storage, nonlinear spectroscopy, all optical switching are few of such applications (Sutherland 2003; Hainfeld et al., 2004; Huang et al., 2006; Kim et al., 2010; Rumi and Perry, 2010; Razvi et al., 2015; Abed et al., 2016).

Several physical mechanisms are originated from an intense light–matter interaction (Fang et al., 2010; Shaheen and Fryer 2012; Tritic et al., 2013) which result in nonlinear behavior of the medium (Kanapathipillai, 2006; Kumar and Verma, 2011). Two-photon absorption (TPA) is one of these processes in which two photons are simultaneously absorbed by the medium. In this transition from the ground state to an excited state, an intermediate virtual (not real) state is involved. Nonexistence of a real intermediate state requires a simultaneous absorption of both photons. Nonlinear absorption (NLA) is proportional to square of incident intensity if both photons have the same frequency (belong to a single beam) (Sutherland, 2003; Rumi and Perry, 2010). This allows spatial control of excitation within the medium in three dimensions (Rumi and Perry, 2010). TPA applications include, micro-structuring by TPA polymerization, 3D optical data storage, two-photon photodynamic therapy (Ajami et al., 2015).

Ability of a material to allow nonlinearity upon intense light irradiation is suggested by its properties. Measurement of NLA coefficient of a material is essential for realization of TPA applications. Measuring technique which has been employed in this research work is the single beam, open-aperture \( Z \) scan technique (Ajami et al., 2017). \( Z \) scan technique was first introduced by Sheikh-Bahae in 1989 for measurement of nonlinear refraction (NLR) and NLA (Sheik-Bahae et al., 1989, 1990). In open-aperture \( Z \) scan technique, a nonlinear medium is scanned along the direction of propagation of laser beam through the focus of the beam. Transmittance is recorded as a function of sample position. Scan starts far away from focus where irradiance is low and absorption is constant. At focus however, sudden increase in absorption (positive nonlinearity) occurs which is shown as a valley in the transmittance curve. \( Z \) scan trace can be analyzed to find the type of nonlinearity by employing a theoretical fit. Significance of this technique is its simplicity, cost effectiveness, and sensitivity (Menard et al., 2007; Quarterman et al., 2015; Razvi et al., 2015; Samuel et al., 2015).
This research work focuses on analyzing TPA in the femtosecond regime, in Au-ion irradiated glasses namely, GIL49, BK7, and Glass B. Introduction of metallic ions enhances nonlinear properties in transparent media (Stepanov, 2011).

**Experimental**

Three types of glasses namely, GIL49, BK7, and Glass B were used in the current work. All three glasses are of different chemical compositions which have been presented in Table 1. Chemical composition of the glasses differs in terms of concentration variation of monovalent modifiers which affects the extent of crosslinking in the matrix.

Samples were cleaned and then implanted with 1700 keV Au ions by using a Tandetron 4130 MC accelerator at the Nuclear Physics Institute in Prague, Czech Republic. Ion fluence for all samples was \( \sim 10^{16} \text{ ions/cm}^2 \). Samples were post annealed at 600°C for 5 h. Purpose of annealing is to initiate reformation and crosslinking within the glasses.

A well-established and accredited open-aperture Z scan technique is used to measure TPA coefficients. A femto-second laser (FEMTOPOWER Compact PRO) has been utilized in the Z scan setup. Pulse duration is measured by using an autocorrelator from "Femto Lasers." The laser carries average power maxima at 800 mW, a repetition rate of 1 kHz, and 25 fs pulse duration with a Gaussian profile. The wavelength spectrum is centered at 800 nm. A digital power meter is used to measure laser power before each scan. The scans were performed at 30 nJ pulse energy. Laser beam waist radius and Rayleigh range (\( Z_R = k w_0^2 / 2 \)) were found to be 11 μm and 0.28 nm, respectively.

Figure 1 shows the experimental scheme of the setup. Laser beam was split in two by using a beam splitter; one part was taken as the reference beam. Other part, with a beam diameter of 15 mm, was focused on the sample by a plano-convex lens with a focal length of 175 mm. Samples were placed on a linear translational stage (Newport) and were scanned for a distance of ±3 mm from the focal point.

Reference and transmitted beams were detected using Si diodes (Photodiode 1 and 2 in Fig. 1). A neutral density filter was used to adjust absolute intensity at the diodes. A low-pass filter blocked all unwanted emissions (at Photodiode 2). Data were recorded using a Picoscope from "Picotech." Data acquisition and sample movement were performed by utilizing National Instruments’ Labview software. This software allowed averaging over several laser shots at each position. Resultant data comprised of normalized transmittance at known sample positions.

We have developed C# based software to perform fitting of experimental data for open-aperture Z scans. This Z scan software generates the theoretical curves by considering experimental specifications for each sample. The software can further be modified to include other nonlinear phenomena such as NLR (for closed aperture Z scan), multiphoton absorption, etc.

**Results and discussion**

Ion irradiation onto a material initiates a process of energy transfers from ion to the substrate which is strongly influenced by

<table>
<thead>
<tr>
<th>Type of glass</th>
<th>SiO(_2)</th>
<th>Na(_2)O</th>
<th>Al(_2)O(_3)</th>
<th>CaO</th>
<th>MgO</th>
<th>K(_2)O</th>
<th>B(_2)O(_3)</th>
<th>BaO</th>
</tr>
</thead>
<tbody>
<tr>
<td>GIL49</td>
<td>63.2</td>
<td>24.4</td>
<td>1.1</td>
<td>5.6</td>
<td>5.3</td>
<td>0.5</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>BK7</td>
<td>68.3</td>
<td>8.8</td>
<td>–</td>
<td>0.1</td>
<td>–</td>
<td>8.1</td>
<td>12.1</td>
<td>2.5</td>
</tr>
<tr>
<td>Glass B</td>
<td>88</td>
<td>8.7</td>
<td>3.3</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

![Fig. 1. Experimental scheme of Z scan setup.](https://www.cambridge.org/core).
Fig. 2. 3D ion distribution in Au-ion embedded (a) GIL49, (b) BK7, and (c) Glass B obtained by TRIM simulations.
energy of ions and structural density of the medium. Energetic ions collide with target atoms (nuclear stopping) or electrons (electronic stopping). Ion energy transferred to the target by these two processes, results in rapid local temperature rise followed by rapid cooling (Ahmad et al., 2014). Complete stopping of ions involves several other phenomena such as initiation of collision cascades, ionizations of target atoms, evolution of thermal spike defects generation, damage accumulation, melting, shrinkage due to rapid cooling, sputtering at the surface, and structural alterations of the target (replaced and displaced atoms) (Correa et al., 2012). Stopping and range of ions in matter (SRIM)/Transport of ions in matter (TRIM) simulations have been employed to establish ion range and depth profile within the medium.

Ion penetration within the substrates is expected after irradiation employing a high-energy ion accelerator. Penetration depth and distribution of Au+ ions having 1700 keV energy within glass substrates were estimated by TRIM simulations. Detailed calculations with full-damage cascades were performed for each sample, taking into account the chemical composition of glass substrates. Figure 2 shows the Au+ ion distributions within (a) GIL49, (b) BK7, and (c) Glass B. Each simulation was performed for 500 ions for a detailed analysis. Ion distributions show the subsurface implantation of metallic ions. Ion straggling is also evident from ion distributions in each sample.

Ions penetrating in a medium cover a specific distance depending upon ion density, medium density, and ion energy. This distance is referred to as stopping range or ion penetration depth. Ion energy (1700 keV in our case) will cause electronic stopping which does not cause deflection. Near stopping range, however, they lose enough energy to start nuclear stopping. Ion trajectory will then involve straggling within the medium. Hence the angular distribution affects the ion penetration depth within medium until complete stopping. Nuclear stopping not only causes deflection in ion path but also initiates target damaging phenomena. These include phonons generation, ionizations, vacancies, and replacement collisions. Values for all these parameters are not reported here as an obvious difference was not observed for all three glasses.

Table 2 shows ion ranges and straggling values acquired from TRIM simulations for each sample. It also accounts for total phonons and total target damage. There is not a big difference in these parameters for all three glasses. It can be concluded from TRIM results that ion penetration parameters are almost the same for the glasses used in this work.

UV-VIS spectra of all three glasses before and after ion irradiation and annealing did not show one-photon absorption at or around 800 nm. Furthermore, as irradiated samples did not show a recognizable nonlinear response either. Samples were then annealed at 600°C for 5 h, as it was expected to enhance NLA due to reformation and crosslinking within the samples.

TPA measurements of Au-ion irradiated glasses, post annealing, were performed by employing the open-aperture Z scan technique. Normalized transmittance as a function of sample position was recorded. These experimental data were then fitted using Eq. (1) (Ajami et al., 2010):

\[ T(z) = \sum_{n=0}^{\infty} \frac{(-q_0)^n}{(n + 1)^{3/2}(1 + x^2)^y} \]  

where

\[ x = z/z_R, \quad q_0 = \beta L_{eff} I_0, \quad L_{eff} = (1 - \exp(-\alpha_e L))/\alpha_e \]

Here, \( I_0 \) is the maximum on-axis intensity, \( Z \) is the sample position, \( Z_R \) is the Rayleigh length, \( L_{eff} \) is the effective sample length, \( \alpha_e \) is linear-absorption coefficient, and \( \beta \) is the TPA coefficient.

We have developed C# based software which calculates all these parameters to generate \( T(z) \) values corresponding to each position \( z \). An output excel file is generated containing the position values (experimental) and the normalized transmittance values (calculated). These output files were then used to plot the theoretical curves along with the experimental traces (shown in Fig. 2). Plots of theoretical curves along with the experimental traces for each sample are shown in Figure 3. The best fit data were utilized to extract the TPA coefficient (\( \beta \)).

The \( Z \) scan traces shown in Figure 3 (a) Au ions in GIL49, (b) Au ions in BK7, and (c) Au ions in Glass B are typical TPA curves and they have been perfectly fitted using Eq. (1). This equation represents pure TPA. Modeling of this equation assumes negligible excited state absorption, three photon and higher order absorption. A perfect fit confirms the occurrence of TPA. At intensity maxima, there will be maximum absorption and as we move away from focus, it rapidly reduces on both sides of the scan. Au ions embedded BK7 have lowest normalized transmittance value of 0.928 at intensity maxima (\( Z = 0 \)). In cases of GIL49 and Glass B, lowest normalized transmittance values are 0.96119 and 0.97885, respectively. TPA coefficients extracted using \( Z \) scan software for these samples are as follows: for Au ions in GIL49 it is 7.72 cm/GW, Au ions in BK7 has the largest value of 16.3 cm/GW, and Au ions in Glass B it is 5.25 cm/GW.

Figure 4 is a chart showing ion ranges and TPA coefficients for three glasses. Minor change in ion ranges can be observed. Largest TPA absorption coefficient was found in BK7.

In the femtosecond regime, nonlinear optical absorption is enabled by TPA-induced surface plasmon resonance (SPR). SPR are the collective electromagnetic modes. Photon energy is absorbed by electrons inducing thermally nonequilibrium distribution. Electron–electron interactions then lead to thermalization of electronic distribution in a time frame of a few hundred femtoseconds. These heated electrons then transfer their energy to lattice by means of electron–phonon interaction in the time frame of a few picoseconds. Energy thermalization continues by phonon–phonon interactions. Thermal expansion of surrounding medium is then released causing complete recovery of the system (Link and El-Sayed, 2000; Stepanov, 2011; Almeida et al., 2015). Annealing the samples initiates a reformation and crosslinking of the glass matrix.

Chemical composition of the host glass is an important parameter in crosslinking of embedded ions. Glass transition temperature (\( T_g \)) for these glasses are in the following order: BK7 < GIL49 < Glass B (Svecova et al., 2017). \( T_g \) for BK7 (560°C) is

<table>
<thead>
<tr>
<th>Type of glass</th>
<th>Ion range (nm)</th>
<th>Straggling (nm)</th>
<th>Total phonons (keV/ion)</th>
<th>Total target damage (keV/ion)</th>
</tr>
</thead>
<tbody>
<tr>
<td>GIL49</td>
<td>375.7</td>
<td>68.5</td>
<td>576.2</td>
<td>43.92</td>
</tr>
<tr>
<td>BK7</td>
<td>393.8</td>
<td>72.2</td>
<td>577.2</td>
<td>43.29</td>
</tr>
<tr>
<td>Glass B</td>
<td>405.8</td>
<td>72.1</td>
<td>569.6</td>
<td>43.23</td>
</tr>
</tbody>
</table>

Table 2.

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lower than annealing temperature. It can be expected to go through melting and rearrangement with highest crosslinking and new bond formation. This can be the reason for the highest TPA coefficient in BK7 as nonlinear properties of SPR are known to be enhanced by increasing the filling factor of embedded medium.

Conclusion

TPA coefficients were extracted for Au embedded glasses from Z-scan traces. Highest TPA coefficient (16.3 cm/GW) corresponds to Au-embedded BK7. There is no significant change in ion range, straggling, and total target damage with the change in the substrate composition. Difference in TPA coefficient is only caused by extent of crosslinking (attributed to post annealing) which strongly depends on chemical composition and structure of the glass substrate.

References


