1	Fabrication and experimental evaluation of microstructured ⁶ Li silicate fiber arrays for 'igh spatial
2	resolution neutron imaging
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15	Abstract
16	This work presents the fabrication and exp. imental evaluation of instrumentation designed to enable
17	higher spatial resolution neutron r . dio _{b.} and y for those performing research at neutron scattering facilities.
18	Herein, we describe a proof-of-conce, array of microstructured silicate fibers with 6Li doped cores that
19	shows progress towards a d sign for \$\mu m\$ resolution neutron radiography. The multicore fiber was fabricated
20	by drawing stacked unit tements of Guardian Glass (Nucsafe Inc., Oak Ridge, TN, USA), a ⁶ Li scintillating
21	core glass, and a s ⁱⁿ ate c. uding glass. These structured fibers function as an array of sub-10- μm
22	waveguides for scin illatio light. Measurements have shown a significantly increased integrated charge
23	distribution in respons to neutrons, and the spatial resolution of the radiographs is described by edge
24	response at the spread functions of $48 \pm 4 \mu m$ and $59 \pm 8 \mu m$, respectively.
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29	1. Introduction
20	
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31	The properties of novel X-ray opaque materials, like those used it energy storage systems, precision
32	manufacturing technologies, aerospace components, and metalic active manufacturing, are often
33	described using simulation tools lacking experimentally grounded mouels, and are based on first-principle
34	calculations and theoretical assumptions alone. Thus, ongoing .pater.al characterization relies on neutron
35	scattering instrumentation to verify performance predict and and structure to future models. Despite the
36	successes that neutron scattering science facilities achieved in recent years, even higher-impact
37	research into microstructure evolution, thermody, wic, and mechanical properties of advanced materials
38	is sometimes limited by the current spatial resembles of neutron sensing instrumentation.
39	
40	Recent improvements in neutron sens. 'o instrumentation have enabled a few modern neutron scattering
41	facilities to reach the current state-of the- art spatial resolution of approximately 10-25 μ m [1] [2]. However,
42	progress towards µm resolution emains a challenging goal for the neutron imaging community for two
43	primary reasons.
44	
45	Firstly, thermal neut, in faxes at even the most novel user facilities are much lower than similar X-ray
46	imaging sources. For exant, le, the raw power of several tens of W/cm^2 is regularly reported at advanced
47	X-ray sources 'e Auvanced Photon Source (APS) at Argonne National Laboratory. This power
48	corresponds to rang of fluxes that conservatively start several orders of magnitude greater than the
49	estimated no 'e' thermal neutron flux of approximately $1.2 \times 10^9 n/s/cm^2$ at the ODIN neutron imaging
50	beamline planned at the European Spallation Source (ESS) [3].

Secondly, the efficiency of thermal neutron sensitive scintillators is in competition with the spatial resolution. Thick scintillators (mm scale) have higher efficiencies and, therefore, $\tan r$ roduce high contrast radiographs in less time than thin scintillators (μ m scale). However, a thick scintillator will have poor spatial resolution, as the position of captured neutrons is smeared throughout the scintillating volume. Conversely, thin scintillating screens and films bypass spatial resolution smealing by reducing the thickness of scintillating medium at the cost of efficiency. Still, the spatial resolution of the charged particles emitted from neutron captures and the succeeding isotropic emission of scintillation light from neutron converters such as ${}^{6}\text{Li}(n, {}^{3}H)$, ${}^{10}\text{B}(n, \alpha)$, and ${}^{157}\text{Gd}(n, \gamma)$. Thus, there is a clear need to marry the benefits of high efficiency and spatial resolution while a ${}^{10}\text{C}$ overcoming the neutron capture position uncertainty that results from charged particle track var ${}^{10}\text{C}$ cc. Some of our prior work based upon Monte Carlo simulation suggests that tracking these charges basis could allow one to overcome this limiting uncertainty [2] [4].

Not long after the seminal work on reutron so intillating glasses [5] was published, scintillating fiber optic glass faceplates were being researched in 1983 [6]. Three years later, the first results of a neutron scintillating fiber (SCIFI) tracker were reported [7]. In 1994, the first neutron rack ograph, of a pierced sheet of Gd, taken using a SCIFI array was published with resolvable feature on the order of several 100s of µm [8]. Since that time, SCIFIs have been researched for remotoradian and dosimetry and neutron sensitive fusion applications [9] [10] [11]. More recently, interest surrounding coupling optical fiber tapers to existing scintillator based neutron imaging setups has groon with lovel 11 µm resolution results [1]. However, in the last 24 years, it appears that no work has been to build upon the original concept of the SCIFI tracker.

We propose the use of microstructured scintillating optical fiber arrays, capable of heavy charged particle

(HCP) tracking via waveguiding scintillation light following neutron capture, for high resolution neutron
imaging. Specifically, we utilize the well-known capture reaction for Li-glass 6 Li (n, α) , e_1 . 4 ting alpha (α)
and triton (${}^{3}H$) particles back-to-back that ionize primary and secondary e extra ns, exciting a Ce^{3+}
activator, which in turn emits scintillation light at 395-432 nm in the near-UV/v. ble wavelength range.
This light is transported through the neutron sensitive microstructured v. veguides and is observed by a
photodetector, see Figure 1. Provided that the collected and converted light h. s a sufficient signal to noise
ratio (SNR), one should be able to observe the tracks of these particing and precisely estimate the locations
of neutron capture reactions [4].

In our first generation proof-of-concept array, we attempted a use air capillaries in the cladding layer to maximize the refractive index difference between the Li-dop of glass core and the cladding. While the lead oxide cladding glass was mechanically compatible word, the Li core glass, the air capillaries tended to collapse, so we moved to an all-solid glass design. Word the all-solid design, we have made several attempts to better match the thermal fiber pulling properties with the optical properties to achieve the best active scintillating volume and refractive index difference insofar as possible. This work describes the fabrication and characterization of this next generation, proof-of-concept neutron SCIFI tracker.

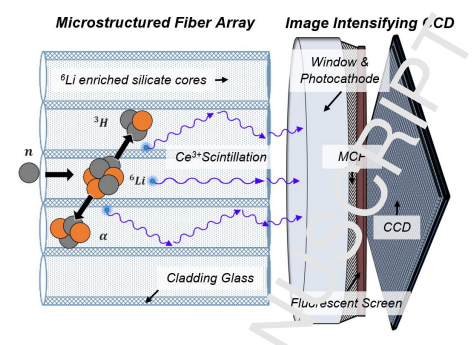


Figure 1. Enriched lithium-6 glass fibers that are doped w .. a bsorb thermal neutrons, emit scintillation light, and act as optical waveguides to channel the scintillation light for imaging. They are surrounded by a cladding of lower refractive index. While an image intensifying CCD is our selected choice for a photosensor in this proof-of-concept study, it is possible that a different photosensor nazy be more appropriate in order to scale to a larger area.

2. Fabrication

To fabricate the neutron SC. $^{\circ}$ I, a multicore design was used. A rod of $^{\circ}$ Li-loaded glass with properties similar to GS20 [5] (fab. $^{\circ}$ Ced $^{\circ}$ Y Nucsafe Inc., Oak Ridge, TN, USA) was inserted in an extruded optical glass cladding tube. Canes $^{\circ}$ f Li-glass cores, embedded in the optical cladding glass, were drawn and then stacked into an $^{\circ}$ Li-glass cores, embedded in the optical cladding glass, were drawn and then stacked into an $^{\circ}$ Li-glass cores, embedded in the optical cladding glass, were drawn and then stacked into an $^{\circ}$ Li-glass cores, embedded in the optical cladding glass, were drawn and then stacked into an $^{\circ}$ Li-glass cores, embedded in the optical cladding glass, were drawn and then stacked into an $^{\circ}$ Li-glass cores, embedded in the optical cladding glass, were drawn and then stacked into an $^{\circ}$ Li-glass cores, embedded in the optical cladding glass, were drawn and then stacked into an $^{\circ}$ Li-glass cores, embedded in the optical cladding glass, were drawn and then stacked into an $^{\circ}$ Li-glass cores, embedded in the optical cladding glass, were drawn and then stacked into an $^{\circ}$ Li-glass cores, embedded in the optical cladding glass, were drawn and then stacked into an $^{\circ}$ Li-glass cores, embedded in the optical cladding glass, were drawn and then stacked into an $^{\circ}$ Li-glass cores, embedded in the optical cladding glass, were drawn and then stacked into an $^{\circ}$ Li-glass cores, embedded in the optical cladding glass, were drawn and then stacked into an $^{\circ}$ Li-glass cores, embedded in the optical cladding glass, were drawn and then stacked into an $^{\circ}$ Li-glass cores, embedded in the optical cladding glass, were drawn and then stacked into an $^{\circ}$ Li-glass cores, embedded in the optical cladding glass, were drawn and then stacked into a single plant glass cores, embedded in the optical cladding glass, were drawn and then stacked into a single plant glass cores, embedded in the optical cladding glass, were drawn and then stacked into a single plant glass

viscosities must be matched for fiber drawing. Previous trials of ours have led to a better understanding of difficulties associated with matching both optical and thermal properties of glass fiber while maintaining the chemical stability required for scintillation. An early attempt to create a square multicore neutron SCIFI, described in detail in [12], yielded an irregular array of cores that had a refractive heav almost equal to the chosen cladding. Although drawing was possible, issues related to glass devitridation during fiber pulling significantly decreased the fiber guiding properties, undermining the relation resolution. Alternatively, the N-FK5 SCHOTT glass makes a more attractive cladding glass with an n_{co} equal to 1.5 and a seemingly compatible transition temperature, (T_g) . However, our initial causes of Li-glass and N-FK5 crystallized during drawing due to an incompatible codrawing temperature for the glass viscosities.

The fabrication of our current multicore fiber began with the extrusion of an N-KF9 cladding tube. A cylindrical glass billet with a 29 mm diameter and 35 n. 1 height and a stainless steel die was used during the extrusion process. A 4 kN force was applied to the die and the glass billet while the glass was held in a furnace at an onset temperature of 650°C (true glass emperature of 620°C). Some metallic inclusions were observed inside the extruded glass. Next the Li- class rod (11.7 mm diameter) was inserted into a 100 mm long section of the extruded N-KF9 tabe, and crawn using the preform drawing method. To determine the optimal drawing temperature, the Li- cored silicate glass, T_g , was experimentally established with differential scanning calorimetry (Netz. th DSC STA 449 F1 JUPITER). The measurement was carried out with a heating rate of 5°C in ap to 1300°C in sealed Pt/Rh pans using \sim 30 mg of fine grain sample, providing a value of $T_g = 47$ $to \pm 2$ °C. So, the fiber drawing took place at a furnace onset temperature of 800°C (730 \pm 10°C glass tem; crature) under a flow of N2 at 2 l/min. The preform was fed into the furnace at a speed of 1 mm/min, ... the fiber was drawn at a speed of 5.4 m/min under a tension of 15 g. About 500 m of fiber were drawn rom the preform, resulting in a fiber with an outer diameter and a core diameter of 185 \pm 4 \pm 4 \pm 7 m, and \pm 700 \pm 4 \pm 4 \pm 7 m, respectively.

The fiber was then cut into 2,700 separate 120-mm-long pieces and stacked as a heragonal array. This array was then thermally consolidated into a single unit at 640°C and the N-KF9 tube (1.° ±0.2 mm OD/ 11.4 ±0.2 mm ID), used to jacket the Li-glass cores, was extruded under the sene conditions previously described for the outer cladding tube. The resulting preform was then drawn into a single multicore fiber 910 ±10 μ m in diameter with individual cores possessing 7–10 μ m diameters, see Figure 2. The hexagonal circle packing geometry allows for the highest fill fraction (active scint dating volume) of circles (cores) to remaining space (cladding), $\frac{\pi\sqrt{3}}{6} \approx 90.7\%$, provided the pitch of cross is equal to the diameter of the cores. Given a conservative estimate of the average cladding spacing where cores of ~2 μ m, we estimate that the active scintillating volume of our hexagonal multicore substantial scincillating volume, while also allowing for sub-10- μ m individual core dimensions.

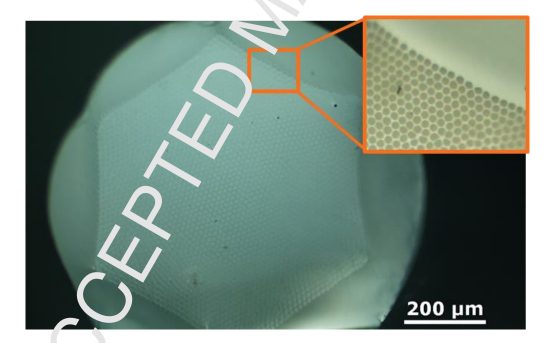


Figure 2. Photographer cross section of a cleaved end of a single multicore SCIFI (hexagonal) inside of its outer jacket (round)

3. Experimental Evaluation

Following fiber drawing, it is necessary to establish that our neutron SCIFI still so intillates as expected. Thus, the radioluminescence emission spectrum of an unpolished Li-glass rod, a sed for the core glass, was measured for an *On Side* (diameter) and an *On End* (length) case. The diameter of the unpolished Li glass rod was 10 mm with a length of 150 mm. A standard emission peak of Ce³⁺ so intillation light near to the 395 nm peak emission was observed for the *On Side* case, see Figure 2. In the *On End* case, the emission peak contracted 15 nm in the near-UV region. Here, the self-absorption effects of the overlapping Ce³⁺ and Ce⁴⁺ emission and absorption bands are observed when the schriften on light is transported further for the *On End* case. In Figure 3, these unpolished rod results ar compand to the arrays of multicore SCIFIs, with and without the outer jacket removed; refer back to Figure 2 for an image of a single multicore fiber possessing an outer jacket. The SCIFI arrays were firm which with a 5 x 5 mm surface. The SCIFI arrays were measured on end, as a faceplate. After the science of the aforementioned heat processing, the SCIFI

emission behaves as expected for Ce³⁺ scintillation. The SCIFI had a slightly broadene¹ emission, 3-5 nm, compared to the *On Side* case for the Li-glass rod.

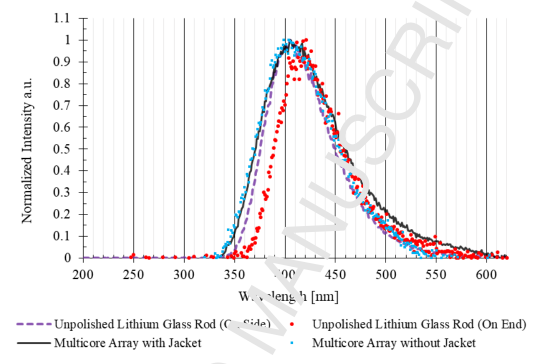


Figure 3. Radioluminescence spectra charact ristic c° Ce³⁺ activator emission for the unpolished ⁶Li glass rod (unprocessed) and the multicore arrays (d awn 11, 24 bers) with and without outer jackets.

Having verified that the scin allace in mechanism is behaving as expected, it is essential to characterize any transport of the Ce activator from the core to the cladding via thermal diffusion to ensure that scintillation light is being produced within the core glass. So, Energy Dispersive X-ray Spectrometry (EDS) was used to examine the accinic concentrations of dopants in cane cross-sections with a Zeiss EVO MA15 Scanning Electron. Aicroscope. The canes of the Li-glass cores cladded with N-KF9 were embedded in epoxy, and EC 3 line soans were acquired across the ends of the canes. The results of the EDS line scans are shown in Figure 4, where Mg inside of the core glass can be seen as the boundary between the core and cladding. No corease, below standard deviation, for the Ce concentration in Li-glass cores at the near-cladding position was observed. EDS point scans targeting 4 randomly selected Li-glass cores, at 3 positions

each, correctly detected a Ce concentration of 2.4 ± 0.2 wt.%. No Ce was detected when point scans targeted the cladding with a lower detectable limit set to 0.1 wt.%. So, the intensity of the Ce concentration in the cladding and surrounding epoxy is assumed to be background. Thus, Ce was well bound within the core glass during fiber drawing. Again, the fiber was drawn at a $730 \pm 10^{\circ}$ C glass temporature, 260° C above T_g , with any given segment of fiber experiencing approximately 10 min of heaving. Previous work of ours has shown that the mobility of Ce in Li-glass, heated at temperatures between $500-600^{\circ}$ C for 5 hr with a subsequent 24 hr anneal, resulted in a few μm of Ce diffusion [13]. Thus, the quasi-immobilization of Ce for the short heating duration agrees with our experimental data

If a significant quantity of the ⁶Li absorber is in the SCIFI clac. ing, then neutron captures in the cladding could decrease overall light output, and potentially create ± 5. im position smearing. So, it useful to know how the ⁶Li absorber is diffusing within the multic re. However, EDS systems are fundamentally insensitive to atomic numbers < 5 due to the absorptic of low energy X-rays within the detection window. X-Ray Photoelectron Spectroscopy could be used to learch for Li in the cladding, but it would not provide an accurate concentration depth profile. Such technique would require micron scale accuracy when removing core from cladding, and while reobing the depth of the cladding. Instead, the mobility of another alkali metal, Na, is particularly in eresting hecause of its chemical similarity, comparable ionic radius, and mobility to the Li absorber in the core glass. The diffusion of Na within Li enriched silicate, ⁷Li/⁶Li = 0.04526, and Na self-diffusion in the core glass. The diffusion of Na within Li enriched silicate, ⁷Li/⁶Li = 1.04526, and Na self-diffusion in the core glass indicates that the activation energy for Li diffusion remains comparable to Na, specificany inside of aluminosilicate glasses; where mixed alkali effects will not immobilize Li or Na [1517, 7]. The self-diffusion of Na within the cladding glass and its diffusion of 30 ±4 μm into the core can be clearly seen in Figure 4. Although the EDS line scan intensities do not represent

relative concentrations between elements, the Li absorber is very likely diffusing in a similar manner to Na from the core into the cladding.

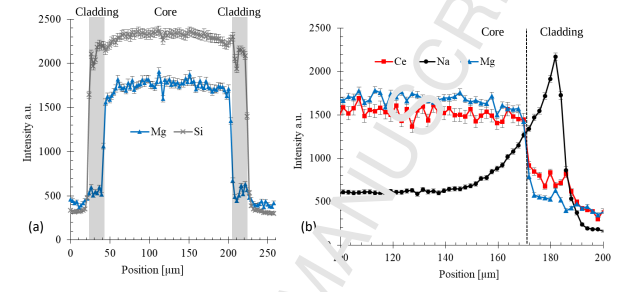


Figure 4. (a) EDS line scan across the entire cane where the core (Mg) and core/cladding (Si) components can be seen, and (b) spectra of the core-cladding interfere for Mg, Ce, and Na.

Preliminary studies on gamma/n tro distrimination performance of the Li-glass within the previous square multicore SCIFI describe in [12], were conducted at the CG-1D neutron imaging beamline at the High Flux Isotope Reactor ('ar.'?) at Oak Ridge National Laboratory (ORNL) [18]. The response of the previous square multicore SCn in array to background radiation (cold neutron flux $< 1 \times 10^1 \ n/s/cm^2$ at target with neutron shutter pen) was observed with an R9779 PMT. The scintillation light was separable from electronic noise or Cherenkov light generated in the PMT window, both in magnitude (integrated charge) and fall in the previous are characteristic of scintillation caused by the Ce³⁺ dopant. While scintillation as the array exhibited a higher light output response to cold neutrons than the

gamma background radiation present in the beamline. Due to the low effective active scintillating volume
of the previous square packing used (~27%), the majority of energy deposited in the multicore from the
charged particles was not producing scintillation light. The current hexagonal design at ows for nearly triple
the amount of energy to be deposited into the scintillating core glass.

Radiographs were taken with the current hexagonal multicore fibers using the Swiss Spallation Neutron Source at the Paul Scherrer Institut (PSI). Images were acquired with the Paul Scherrer Institut (PSI). Images were acquired with the Policy at the Policy Policy (NM) at the Pulse OverLap Diffractometer (POLDI) beamline. The use of the New at the Policy Deamline was uniquely desired due to the requirements of the optical system needed for the experiment. Specifically, the NM possesses the magnification and numeric aperture to enable a reminal pixel size of acquired images of 1.3 μm, and a true spatial resolution of about 5 μm [19]. Additionally, the optics have μm-level repeatable positioning, and are sensitive to near UV Ce scintillar and with other beamlines, a thermal/cold neutron spectrum from 1.1–5 Å, and a flux at the sample position of 6 × 106 n/s/cm² [20].

A diffused, blue light source was used to critically focus onto the ends of hand-cleaved and polished single multicore fibers. Al tape v as the vield to enclose the scintillator. Open Beam (OB) images were taken to focus the optics to the scint. Into light. A quantity of 20 images with 300 s exposures were acquired of the OB. A gade iniv n-based Siemens Star (SS) [21] was then positioned in front of the active area of the scintillator and in aged. Eighty (80) images of the SS with 300 s exposure were acquired. Due to decreased thermal meutron dux, 2 OB and 11 SS images were removed prior to post processing. The remaining SS image with outliers removed, were divided by the OB background, and summed. The resultant image was transformed with a bilinear clockwise rotation of 90° for analysis. The resolvable spokes of the SS spokes possessing approximately 50 µm spatial features. Referencing the 10% to 90% contrast transition for the Edge Response Function (ERF) of the spoke in the highlighted

region, a $48 \pm 4 \,\mu m$ resolution was found. Fitting a Gaussian to the Line Spread Function (LSF) of the same region, a FWHM measurement yields $59 \pm 8 \,\mu m$ spatial resolution.

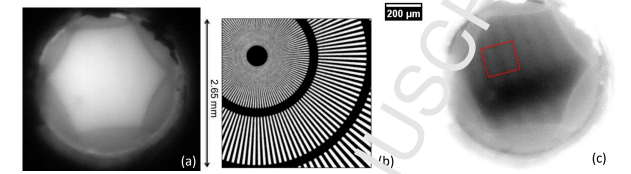


Figure 5. (a) OB neutron radiograph taken with a 2 mm thick must ore neutron SCIFI, (b) visual test object reference, PSI's Gd SS taken with the NM in [18], and (c) neutron and graph of the SS acquired after aforementioned post processing, with the region of interest highlighted.

4. Conclusions and Fut re Work

Li-glass cores with 7-10 μ m diame. Thave been drawn into 500 m of an all-solid glass composite, multicore fiber at a 70% pacting traction. The Li-glass remains active and scintillates as expected following drawing. The Li absorber shows some diffusion into the cladding, but the Ce activator is largely bound during the fabrication process. Hand-cleaved and polished single fibers of lengths 2-5 mm have resolved features on the order of 10, of μ m while utilizing the Neutron Microscope. Ideal light collection in these same fibers should allow for \leq 20 μ m resolution. To increase spatial resolution, the core glass must be drawn to span, wizes which should be possible with the current fabrication process. Image contrast

improvement requires the use of a lower refractive index cladding material. Moreover, fabrication without an outer cladding jacket will allow for easy array assembly.

Near term work is planned to evaluate the radiographic spatial resolution of a. array of 100 multicore neutron SCIFIs with a 5 x 5mm field-of-view. This is a comparably thinne design (1mm thick), with outer jacket removed, and it has a well-polished surface. Since a larger field of-vie v was desired for evaluation of imaging performance, the multicore fiber outer jacket was ground way, and a 10×10 array was stacked together. Assembly was finished by securing the array with structural glass plate siding and polishing the surfaces, see Figure 6.

Meanwhile, a novel phosphate glass cladding, possessing refractive index of $n_c < 1.53$, is being used for another neutron SCIFI fabrication. This cladding is expected to allow for more than twice the current amount of scintillation light to remain internally because. We plan to draw the multicore fiber using the phosphate cladding with a goal of 2 μ m diameter E glass cores while remaining at a packing fraction \geq 70%.



Figure 6. Microsco_P \sim pnotos of the (a) cross section of a microstructured multicore fiber with scale, (b) stacked multicore fiber with outer cladding removed with 8 \pm 1 μ m core diameters, and (c) polished faceplate surface of a multicore SCIFI array with 0.5 x 0.5 mm pixels

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