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Abstract. A nonresonant, fiber-optic raster scanning endomicroscope was developed using a quarter-tubular piezoelectric (PZT) actuator. A fiber lever mechanism was utilized to enhance the small actuation range of the tubular PZT actuator and to increase its field-of-view. Finite element method simulation of the endoscopic probe was conducted for various conditions to maximize its scanning range. After fabricating the probe using a double clad fiber, we obtained two-photon fluorescence images using raster beam scanning of the fiber. The outer diameter of the probe was 3.5 mm and its rigid distal length was 30 mm including a high numerical aperture gradient index lens. These features are sufficient for input into the instrumental channel of a commercial colonoscope or gastroscope to obtain high resolution images in vivo.

Keywords: two-photon microscopy; endomicroscopy; fiber scanning; raster scanning; tubular piezoelectric actuator; endoscopic imaging.

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1 Introduction

A fiber-scanning endomicroscope is one of the powerful tools used for visualizing gastrointestinal and colonic images in vivo at a cellular level. Endomicroscopes were originally developed using a fiber bundle and a gradient-index lens objective. One of the main advantages of the fiber bundle type is that its beam scanning unit is located outside of the endoscopic probe, so the size of the probe can be easily reduced. However, its image quality is quite degraded by the pixelation resulting from gaps between individual optical fibers within the fiber bundles, even if a frequency filter eliminates the honeycomb pattern of the fiber bundle.

Beam scanning can also be conducted using a microelectromechanical systems (MEMS) mirror inside the endoscopic probe instead of a fiber bundle/outer scanning unit. Endoscopic probes that use a MEMS mirror were mainly developed for the side-view type. Due to the bulky volume of the MEMS mirror including electronic parts, it is also hard to assemble into a diameter smaller than about 5 mm.

Fiber-scanning endomicroscopes have been studied in several ways to reduce outer size. Some miniature microscopes have been developed using different light guides between illumination and detection parts. On the other hand, a double clad fiber has also been used to guide illumination/detection light with the fiber and to develop a compact fiber-optic two-photon endomicroscope. In respect to the beam scanning method using a piezoelectric (PZT) actuator, it can be classified into two types: those using a tubular PZT actuator and those using a bimorph (or trimorph) PZT actuator. Most of the studies with the tubular PZT actuator have used spiral or lissajous pattern scanning methods, around the fiber’s resonant modes. These scanning methods have significant nonuniformity between the inner and outer parts of the scanning area. Also, the actuation range of the tubular PZT actuator is too short (tens of micrometers) to provide a suitable field-of-view within a specimen. To obtain a scanning range of hundreds of micrometers, scanning of the fiber near the resonant mode of the fiber is essential. A bimorph PZT actuator can utilize a raster type scanning method. However, one direction of that scanning should be conducted in a nonresonant mode, and this increases the rigid length of the endoscopic probe, because it requires the bimorph PZT to have quite a long length. Also, Li and Fu’s group has studied a resonant fiber-optic PZT scanner to achieve a raster scanning pattern. These are scanned at a resonant mode so the scanning speed cannot be freely changed. Sawinski and Denk’s group has developed a nonresonant raster scanning endomicroscope using a piezolever fiber scanner composed of four trimorph PZT actuators. These studies have advantages to control imaging field-of-view and speed easily, but its outer diameter still could not become smaller than 5 mm due to the size of the four trimorph PZT actuators.

In this paper, a fiber-optic raster scanning endomicroscope was developed using a tubular PZT actuator. By using a nonresonant fiber scanning type, its scanning speed and scanning area can be easily controlled by adjusting the applied voltages of the quarter tubular PZT actuator. Controlling the scanning speed and area is a necessity. Some commercial scanning microscopes have utilized a set of two galvano mirrors or a set of resonant/galvano mirrors for this purpose. In particular, the...
scanning speed, pixel dwell time, and scanning area should be adjusted when the detection period is quite long. This is the case, for example, with a fluorescence lifetime imaging microscope, or when partial scanning is needed, such as fluorescence recovery after photobleaching and fluorescence loss in photobleaching.

2 Methods

The system configuration of the experimental device is shown in Fig.1. A collimated beam from a femto-second pulse laser with a center wavelength of 780 nm (FemtoFiber pro NIR, Toptica, Munich, Germany) was chirped negatively by a customized prism pair and focused into a core of double-clad fiber (SMM900, Fibercore, Southampton, United Kingdom) with 50 cm of length. To compensate for the pulse dispersion caused by propagation of the fiber core, the pulse width was measured by an intensity autocorrelator (Mini, APE, Berlin, Germany) after the laser traveled inside the core. The auto correlation function was measured to be 2115 fs, which corresponds to a pulse width of 1058 fs as shown in Fig.2(a). Therefore, the entire group delay dispersion of our system was calculated to be 34,800 fs² since the initial pulse width was 100 fs. Negative chirp was applied with a customized prism pair which has a short length and high light efficiency. Chirping with one or more gratings was excluded in this study to maximize light efficiency. The prism had an apex angle of 54.85 deg and S-NPH3 (Ohara, Branchburg, New Jersey) was chosen as the glass material due to its superior high refractive index and dispersion characteristics ($n_d = 1.959$, $v_d = 17.47$). The distance between the two prisms was selected to be 51.5 cm using an equation derived from the optical path. The light efficiency of the negative chirping unit was measured to be 79.2%.

Inside the endoscopic probe, one side of a tubular PZT (Customized, PI, Lederhose, Germany) was attached to a PZT holder located inside the probe housing. A hinge made from stainless film was attached to the other side of the tubular PZT. The fiber penetrates through a hole in the hinge. The small actuation range of the tubular PZT ($\pm 23 \mu m$ at $\pm 200$ V) was amplified with a lever mechanism. A commercial gradient-index (GRIN) objective lens (GT-M0-080-018-810, GRINTECH, Jena, Germany) with 0.8 NA was used to get high efficiency from the two-photon effect on a focal spot. Also, the thickness of the hinge affected the deflection range. In our simulation results, when the hinge was thickened, the deflection range was dramatically reduced. Therefore, a stainless film having 10 $\mu m$ of thickness was chosen as the hinge material to provide sufficient range and be robust against tearing. All other mentioned dimensions, $L_1$ and $L_2$, were decided after finite element method (FEM) simulation of the probe model.

3 Results

3.1 Simulation Results

The results of FEM simulation are shown in Fig.3. Figure 3(a) shows a model of the simulation. The fiber length, $L_2$, was fixed at 21 mm, as limited by the length of the tubular PZT (20 mm). Depending on $L_1$, the lateral displacement of the distal end of the fiber varied from $\pm 240.3$ to $\pm 329.4 \mu m$ as shown in Fig. 3(b). Also, contact conditions between the fiber and the hinge affect the results. However, adjusting the tolerance between the fiber and the hole of the hinge was practically very difficult. Instead of making a sliding joint, the minute space inside the hole was filled and fixed by elastic epoxy (EP001, Cemedine, Tokyo, Japan) to permit a relative and flexible movement. To verify the strength of the hinge, maximum stresses were also calculated by FEM simulation as shown in Fig. 3(c). Maximum stresses (Max. principal stress) applied on the fiber varied from 81.0 to 140.4 MPa, which was...
much lower than the rupture stress of the fiber (2350 MPa). Maximum stresses (Max. von Mises stress) applied on the hinge varied from 92.5 to 110.5 MPa, which was lower than the yield strength of the hinge material (Stainless steel, ~200 MPa). The first-resonant frequency of the fiber according to $L_1$ is shown in Fig. 3(d). The simulation result indicates that the length of $L_1$ is almost unaffected by the resonant frequency of the fiber, which ranges from 157.5 to 158.6 Hz. After considering all of these factors, $L_1$ was determined to be 1.2 mm, to achieve the maximum lateral displacement. After fixing the $L_1$ as 1.2 mm, we considered the effect caused by the thickness of the epoxy bonded between the hole of the hinge and the fiber. FEM simulation was conducted and is shown in Fig. 3(e). The larger the thickness of the epoxy, the smaller the lateral displacement of the fiber tip becomes. This is reasonable since the change of epoxy thickness is equivalent to the change of $L_1$. Also, the overall stiffness of the lever structure will increase as thicker epoxy bonding is used. The endoscopic probe was assembled into an aluminum housing produced by a wire cutting process with an outer diameter of 3.5 mm and a wall thickness of 0.35 mm. Soldering of the tubular PZT with an outer diameter of 2.2 mm was achieved using silver paste. A gap between the electric wire and the PZT surface was filled with silver paste (P-100, Elcoat, Tokyo, Japan) and pressed thin. To reduce the overall size of the probe, most of the components were assembled using a high strength epoxy (KEIN 60 min), except for the gap between the hole of the hinge and the fiber (where an elastic epoxy was used). To align the center of the fiber, three-axis manual stages were utilized for precise assembly procedures. The fiber scanning range of the slow axis was about 518 $\mu$m. The final scanning range at the focal plane of the GRIN lens was measured to be 107.9 $\mu$m, due to the magnification of the GRIN lens (4.8×).

### 3.2 Experimental Results

#### 3.2.1 Performance evaluations

After assembling the endoscopic probe, the characteristic of the fiber tip depending on driving frequencies was measured. Sinusoidal input voltages with an amplitude of ±50 V were driven to the fast axis of the PZT actuator. Although the driving frequency changes from 32 to 160 Hz, the displacements of the fiber tip were measured by a charge-coupled device (WAT-202D, WATEC, Kawasaki, Japan). As the driving frequency of the fast axis increased, the displacement of the fiber tip gradually increased as shown in Fig. 4. The first-resonant frequency of the system can be predicted above 160 Hz from Fig. 4, which is somewhat greater than the simulation results presumably due to assembly error. It was not measured until the exact first-resonant frequency due to the worry about the collision between the fiber tip and the interior of the tubular PZT actuator (inner diameter of 1 mm). The results can be utilized to adjust the field of view of the system when the imaging speed has to be changed.

A beam splitter (BSN11, Thorlabs, Newton, New Jersey), a doublet lens (W49-356, Edmund Optics, Barrington, New Jersey), a 30-$\mu$m diameter pinhole, and a photomultiplier tube (H6779, Hamamatsu Photonics, Hamamatsu, Japan) were additionally inserted within the detection light path to obtain reflection confocal images for evaluating the scanning performances as shown in Fig. 5. The reflection image of a distortion grid target (R1L3S5P, Thorlabs) was obtained while the raster scanning pattern was achieved. As a result of the nonlinearity of the sinusoidal
input signal and inherent characteristics of the PZT actuator, image distortion exists and it becomes larger at the periphery of the scanning area as shown in Fig. 5(a). Image reconstruction was conducted to compensate the image distortion by reorganizing the image data using MATLAB software (MathWorks, Natick, Massachusetts) as shown in Fig. 5(b). The same algorithm was applied to the image of a Siemens star target (R1L3S5P, Thorlabs) to show the repeatability and usefulness of the applied algorithm using the circular-symmetric target as shown in Fig. 5(c).

3.2.2 Images

We acquired images of a test target and a biological specimen, as shown in Fig. 7. The slow axis scanned with a saw-tooth pattern instead of the sinusoidal pattern. The image acquisition time was 1.6 s with a pixel size of 256 × 256. A reflection image of a 1951 USAF target (R1L1S1P, Thorlabs) was obtained using the fiber raster scanning method in Fig. 7(a). The smallest pattern of the target (group 7, elements 5 and 6) was magnified two times by the partial scanning of the fiber in Fig. 7(b). The amplitudes of applied voltages to the PZT actuator were ±50 V (fast axis) and ±162.5 V (slow axis) in the case of Fig. 7(a), and ±26 V (fast axis) and ±100 V (slow axis) in the case of Fig. 7(b), respectively. The amplitude does not scale down linearly due to the characteristic of the PZT actuator, but the relationship between the displacement and the amplitude is sufficiently repeatable to obtain different sized images by making a look-up table.

To obtain two-photon fluorescence images of the biological specimen, we utilized a mouse kidney section stained with Alexa Fluor 488 wheat germ agglutinin (F-24630, Invitrogen, Thousand Oaks, California, USA). Images were acquired using the fiber-optic raster scanning two-photon endomicroscope.
Carlsbad, California), a green-fluorescent lectin which binds to convoluted tubules. The fluorescence signal from the specimen was reflected at a dichroic mirror and acquired by a photomultiplier tube (H7422P-40, Hamamatsu). As a result, the convoluted tubules of the kidney are well differentiated in the magnified image of (c), obtained by partial scanning. The lumen of the convoluted tubule is marked with yellow arrows. Scale bar is 20 μm.

4 Conclusions

In summary, we developed a nonresonant fiber-scanning endomicroscope with a tubular PZT, and demonstrated two-photon imaging of a mouse kidney. To optimize the design parameters of the endoscopic probe, we performed FEM simulations. The fiber scanning pattern was observed and demonstrated to verify a flexible field of view with orthogonal and uniform shape. Finally, we obtained reflection and two-photon images using the developed endomicroscope using a raster beam scanning of the fiber instead of a spiral or lissajous scan. For further work, a customized double clad fiber and a customized miniature lens will increase the device performance by increasing its scanning speed and light efficiency. This miniaturized endomicroscope with highly flexible control of the scanning field-of-view and speed, will allow various preclinical studies for investigating gastrointestinal diseases.

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References


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