Developing compact multiphoton systems using femtosecond fiber lasers

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Abstract. We implement a fiber-delivered compact femtosecond fiber laser at 1030-nm wavelength in multiphoton imaging. The laser pulse duration is 150 fs, the average power is 200 mW, and the repetition rate is 40 MHz. The laser measures 200 × 160 × 45 mm in size and its output is delivered through a photonic bandgap fiber. Intrinsic second-harmonic generation signal is excited from rat tail tendon and human skin samples. Two-photon excited fluorescence signal is obtained from human skin tissues stained with exogenous fluorophore. Our results show that femtosecond fiber lasers at 1030-nm wavelength have significant potential in developing compact, all-fiber-based, portable multiphoton systems and endoscopes.

Keywords: multiphoton microscopy; two-photon excited fluorescence; second-harmonic generation; femtosecond fiber lasers.

Finding new ultrashort laser sources that are low cost, compact, and portable is expected to enhance the impact of MPM in research and particularly in clinical studies. Semiconductor lasers have been recently demonstrated for MPM imaging.11–13 By frequency doubling a 1540-nm gain-switched semiconductor laser, Yokoyama et al. have developed a pulsed source with 5-ps pulse duration at 770-nm wavelength, and applied the source for MPM imaging of actin filaments in PtK2 cells.11 Using a gain-switched semiconductor laser, Taira et al. developed a 980-nm wavelength source with 3.5-ps pulse duration and obtained MPM images of mouse kidney tissues.13 The pulsewidth of those semiconductor lasers is limited to several picoseconds, which cannot excite MPM signals as efficiently as femtosecond pulses.

Recently, femtosecond fiber lasers have emerged as a promising compact light source for MPM applications.14–16 Millard et al. used a 1560-nm wavelength femtosecond fiber laser for third-harmonic generation (THG).14 Unruh et al. applied a frequency doubled 1560-nm wavelength femtosecond fiber laser for two-photon fluorescence correlation spectroscopy (FCS).15 The 1560-nm femtosecond fiber lasers and their frequency doubled versions were still output into free space. Therefore, they lacked the capability of direct fiber delivery of femtosecond pulses from the laser sources.

Femtosecond fiber lasers at 1030-nm fundamental wavelength have not been reported in MPM applications. There are several advantages of using the 1030-nm wavelength for MPM. Exploring longer wavelength excitation for MPM can potentially reduce tissue scattering and increase imaging depth in turbid tissues. The 1030-nm wavelength band can be used to excite a large selection of fluorescent proteins and dyes that have been originally developed for confocal and fluorescence imaging.16–18 Squirrell et al. showed that two-photon imaging using 1047-nm excitation wavelength maintained embryo viability over the long term.17

In this work, we introduce a compact short-pulsed, high-power, fiber-delivered femtosecond fiber laser at 1030-nm wavelength for MPM applications. The main advantage of using the fiber laser is the compactness of the laser source and fiber delivery of femtosecond pulses, which makes it an ideal source for developing compact and portable MPM systems. Such advantages are demonstrated by coupling the fiber laser to an MPM microscope system to provide excitation wavelength at 1030 nm, and implementing the fiber laser in an all-fiber connected multiphoton spectroscopy system.

The femtosecond fiber laser (PolarOnyx, Sunnyvale, California) is composed of an oscillator that generates seed femtosecond pulses, and an amplifier that boosts the power to 200 mW. In the oscillator, a high concentration Yb-doped PCF is used as the gain medium. A 980-nm pump laser pumps Yb ions for amplification of intracavity pulses. Passive mode locking is achieved through nonlinear polarization evolution in the slightly birefringent fiber laser cavity.11–18 Polarization filtering is through a fiber-based inline polarization isolator and polarization controllers inside the laser cavity. Amplification of the seed pulse is achieved by using a short piece of high concentration double cladding Yd-doped fiber pumped with another 980-nm pump laser. After the amplifier, a piece of PBF is used to compress the pulsewidth. The PBF

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can provide $\sim 100 \text{ ps/nm/km}$ anomalous group velocity dispersion (GVD) to compress positively chirped pulses from the amplifier. The output fiber is terminated with an FC/APC fiber connector to deliver femtosecond pulses.

Figures 1(a) and 1(b) show the spectrum and autocorrelation trace of the laser output. The pulse duration is 150 fs measured at the end of the output fiber, and the spectral bandwidth is 20 nm, which brings the time-bandwidth product of the laser to be $\sim 0.8$. The pulse is not transform limited, and the residue chirp is mainly due to higher order dispersion from the fibers. The laser output power is 200 mW and pulse repetition rate is $\sim 40$ MHz. The fiber laser has a highly compact size of $200 \times 160 \times 45$ mm, including the oscillator, amplifier, and pump lasers.

Due to its compact size, the fiber laser can be conveniently coupled to any MPM microscope systems to provide excitation wavelengths at 1030 nm. In our experiment, the output of the fiber laser is collimated and coupled into an MPM scanning microscope. Details about the MPM microscope can be found in Ref. 19. Figures 2(a) and 2(b) show the MPM images excited with the 1030-nm wavelength fiber laser. Figure 2(a) shows the backward SHG imaging of rat tail tendon where the collagen fiber bundles are observed. Figure 2(b) shows the TPEF imaging of the cross section (side view) of ex vivo human skin. The skin sample is stained with yellow-orange vital nuclear dye, and the cellular structure of the skin is captured by the TPEF imaging. In Fig. 2(b), the TPEF signal is mainly from the epidermis (outer) layer of the skin, which is cellular. The dermis (inner) layer shows little TPEF signal because it is mainly composed of an extracellular matrix. In this experiment, no dispersion compensation is applied to simplify the system design. Therefore, to minimize pulse broadening of the excitation light, a $10 \times (0.3 \text{ NA})$ objective is used that has less dispersion than a high NA objective. The excitation power at the sample location is $\sim 20$ mW. The images have 256 $\times$ 256 pixels, and the acquisition time is $\sim 6$ s/frame. Higher image resolution can be achieved by using a high NA objective and implementing dispersion precompensation to maintain short pulse at the sample location.

To take full advantage of the fiber delivery of femtosecond pulses, the fiber laser is implemented in an all-fiber connected multiphoton spectroscopy system. Figure 3(a) shows the schematic of the experimental setup. Femtosecond pulses from the fiber laser are delivered to the tissue site by PBF. At the end of the PBF, the laser power is 200 mW and the pulsewidth reaches the minimum of 150 fs. Thus, MPM signal can be excited from the proximate area of the end of the excitation fiber. Signal collection is through a fiber bundle positioned next to the excitation fiber. The other side of the fiber bundle is coupled to a spectrograph (SpectraPro-150, Acton Research, Princeton Instruments, Trenton, New Jersey) with a grating of 300 grooves/mm blazed at 500-nm wavelength, followed by a cooled charge-coupled device (CCD) camera (NT2/CCD-512-EBFT, Princeton Instruments, Trenton, New Jersey). Figure 3(b) shows the spectrum of intrinsic MPM signal excited from ex vivo human skin without staining. A 5-s integration time is used. As shown by the solid line, a strong peak is observed at 515 nm, which corresponds to the SHG signal excited by the 1030-nm wavelength. The strong SHG signal is excited from the dermis layer, composed mostly of collagen fibers. From the epidermis layer, very low SHG signal is observed, as shown by the dotted line. Due to the long excitation wavelength, very low autofluorescence is excited from the skin sample.

Figure 4 shows the comparison of MPM imaging of human skin excited with 1030- and 800-nm wavelengths. The images are obtained using a Zeiss LSM510, where the laser source (Chameleon, Coherent, Santa Clara, California) can be tuned to 1030 and 800 nm, respectively. A $40 \times$ water NA=0.8 objective is used. The laser power before the objective is $\sim 15$ mW in both cases. For the SHG channel, bandpass filters of 500 to 530 nm and 390 to 465 nm are used for 1030- and 800-nm excitation, respectively. The TPEF channel is filtered with a 535 to 590-nm bandpass filter. Both images are collected in a backward mode. In Fig. 4(a), bright SHG
but very low TPEF signal is observed when 1030-nm excitation is used. In Fig. 4(b), both SHG and TPEF signals are observed from the same sample when 800 nm excitation is used. In Fig. 4(b), the TPEF signal shows in yellow, because the 390 to 465-nm filter also collects some TPEF signal in the SHG channel. In our study, only a gradual decrease in backward SHG intensity is observed when the excitation moves to longer wavelengths. From 800 to 1030 nm, a ~50% drop in the SHG intensity is observed, which can be compensated by increasing the laser power by 140%. Therefore, SHG imaging can be implemented using a wide range of laser wavelengths. Figure 4(a) also shows that the autofluorescence background is largely reduced when excitation moves to 1030 nm. This property can be used to provide highly specific fluorescence imaging with no autofluorescence background when used together with exogenous fluorophores.

In conclusion, we have implemented a compact, all-fiber-based femtosecond fiber laser in multiphoton imaging and spectroscopy at the wavelength band of 1030 nm. Optical pulses of 150-fs pulse duration, 200-mW average power, and 40-MHz repetition rate are generated by an all-fiber-based laser system and delivered by a photonic bandgap fiber. The fiber laser is incorporated with a scanning microscopy system to perform multiphoton optical imaging. Our results show that the femtosecond fiber laser can be used to excite intrinsic SHG signals from tissues, and TPEF imaging can be applied in combination with fluorescence staining. An all-fiber-connected multiphoton spectroscopy system is developed to demonstrate the potential of femtosecond fiber lasers in developing all-fiber-connected portable MPM systems. Overall, femtosecond fiber lasers at 1030-nm wavelength appear to have excellent potential in applications where all-fiber-connected systems are needed, such as multiphoton endoscopes, where the fiber laser can be naturally integrated with a fiber optic probe.

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References

Fig. 4 Comparison of MPM imaging of human skin using 1030- and 800-nm wavelength excitation. (a) Human skin excited by 1030-nm wavelength, (b) The same sample excited by 800-nm wavelength, SHG and TPEF are color coded in green and red, and the overlapping areas are in yellow. Scale bar is 50 μm.