

Laser Combiner Enables Scanning Fluorescence Endoscopy

A laser-scanning fiber endoscope uses a plug-and-play laser combiner to enable high-contrast images suitable for detecting early-stage precancerous lesions. BY DAN CALLEN AND MATTHIAS SCHULZE, COHERENT INC.

etection of early cancerous lesions using molecular imaging can improve disease diagnosis and increase survival rate – this is well established. For example, fluorescently labeled antibodies can be combined with laser excitation in various types of imaging endoscopes to deliver images highlighting precancerous tissue. This can enable real-time in vivo screening by allowing the endoscopist to identify areas for testing and/ or excision. However, the presence of strong autofluorescence (AF) can seriously limit image contrast and make this identification difficult, if not impossible, in some instances.

Figure 1. Above: Where's Waldo? Without some type of autofluorescence mitigation, the high autofluorescence background makes it difficult to locate and determine the extent of any fluorescently tagged tissue containing high-grade dysplasia when imaging a model esophagus with a scanning fluorescence endoscope and a single (488 nm) laser excitation wavelength.⁸

Now, a team of researchers at the University of Washington has successfully demonstrated that using multiple laser wavelengths simultaneously can eliminate most of the AF background problem in studies targeting high-grade dysplasia (HGD) cells in human esophageal tissue. A key enabling component in their scanning fiber endoscope (SFE) is a completely new type of plug-and-play fiber combiner that allows up to eight laser wavelengths to be efficiently deployed in a stable manner in just a few minutes, rather than the many hours normally required from a highly skilled optical technician.

The problem of autofluorescence

Professor Dr. Eric Seibel, his students and staff at the University of Washington's Human Photonics Laboratory have been investigating the use of laser scanning fluorescence images for real-time tissue screening for some time. They recently developed a unique wide-field multispectral fluorescence endoscopy imaging system targeted at early-stage cancer detection in the esophagus.^{1,2} In one modality, they demonstrated that two laser-excitation wavelengths can be utilized for a clever background reduction method: mitigating inherent tissue AF background, as well as spurious specular reflections.³

"In this particular study, we have used fluorescein-labeled antibodies to target cell membrane proteins such as EGFR, ERBB2 and C-Met in a molecular imaging diagnostic scheme that is specific for high-grade dysplasia cells in esophageal tissue associated with Barrett's esophagus," said Chenying Yang, a Ph.D. candidate in Seibel's research group. "Without mitigation of the strong background autofluorescence, the limited contrast typically makes it difficult, and often impossible, to obtain usable tissue-specific data from the fluorescently labeled images (Figure 1). And looking forward to possible future clinical use, where it is not possible to quantify the relative fluorescence intensity of targeted biomarkers, then the endoscopist simply would not be able to perform red-flagging and biopsies of targeted tissue.

"The issue of autofluorescence contrast reduction is a commonly encountered problem in fluorescence-based tissue imaging and certainly not unique to our particular targeting system.⁴ And so we wanted to develop a robust method of subtracting the strong autofluorescence out of the raw images to deliver high-contrast images where the targeted tissue can be clearly observed and delineated."

Background subtraction

The researchers successfully investigated a solution using two different laser wavelengths: 445 and 488 nm. The 488-nm wavelength was chosen to match the absorption maximum of the target fluorophore (fluorescein), but they knew from previous studies that it would also excite broadband autofluorescence (Figure 1). The 445-nm wavelength was chosen to be far enough away from the fluorescein absorption band, but close enough to 488 nm to generate only autofluorescence that presumably has a spectral profile similar to the AF excited by the 488-nm laser. They confirmed this assumption in a separate study by dispersing autofluorescence excited at these two wavelengths using tissue samples that had not been treated with fluorescein.

They also showed that any part of the observed tissue field that produced strong spurious specular reflections would be similarly reflective at both 445 and 488 nm.³ This led them to reason that subtracting an appropriately scaled 445-nm excited AF image from the 488-nm excited image should eliminate both the AF background and the specular reflectance background, yielding an image containing

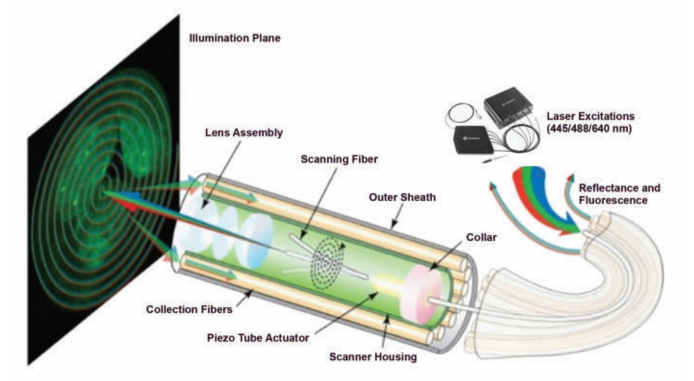


Figure 2. Schematic illustration showing the key components of a scanning fluorescence endoscope developed and tested on esophagus tissue phantoms in studies at the University of Washington.^{3,8}

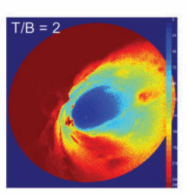
Real-Time 2-NBDG Molecular Imaging with Autofluorescence Mitigation on a Lumenlike Tissue Phantom



Scanning Fiber Endoscope Raw Image

Intensity Color Map





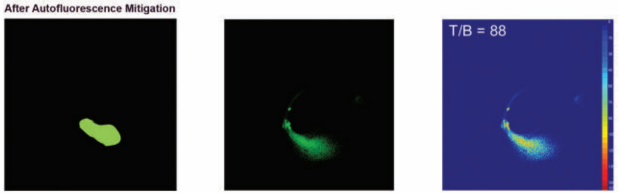


Figure 3. This typical data shows the effectiveness of this dual-wavelength approach, eliminating nearly all of the autofluorescence and specular background to yield a final image clearly delineating the target high-grade dysplasia tissue. In this example, the target background/ratio (T/B) is increased by almost two orders of magnitude (from 2 to 88).⁸

only fluorescence from the fluoresceintagged HGD cells excited by the 488-nm laser.

Figure 2 is a schematic illustration of the SFE that the UW team developed. Light beams from three Coherent OBIS FP lasers at 445, 488 and 640 nm are combined in an OBIS Galaxy plug-andplay beam combiner. The combiner's single-mode output is then connected to the single-mode illumination-delivery fiber in the SFE. At the distal end of the endoscope, a piezo tube actuator deflects a scanning fiber across the field of view.

Why three wavelengths? The 640-nm channel will be used for an additional level of real-time biomarker quantification, as the red reflectance imaging channel provides a means for adjusting the biomarker's fluorescence pixel values for the distance from the endoscope.¹ Thus, a three-channel imaging endoscope can more accurately guide the doctor to taking a biopsy of the most dangerous tissue, while also providing anatomical information when fluorescence is absent.

In operation, the combined multilaser beam is scanned in a spiral pattern by a piezoelectric transducer and focused onto the target surface by a lens assembly. Fluorescence and reflectance are then collected by a concentric ring of highnumerical-aperture optical fibers, which surround the single-mode beam-delivery fiber and lens assembly. This multispectral SFE system has wide-field (80°), high-resolution (50 µm) and real-time (30 Hz) fluorescence-reflectance endoscopic imaging capabilities. An ultrathin 2.1-mm-outer-diameter scope is utilized as part of the multispectral SFE system for high-sensitivity (nanomolar) fluorescence detection.5,6

The collected light passes through a notch filter to block any scattered light from the 488-nm laser and through a longpass filter with a nominal 458-nm cutoff to block any scattered light from the 445-nm laser beam. Dichroic beamsplitters then divide the filtered light into two spectral components: a blue-detection region (AF only) that is passed through a 470-nm (\pm 14 nm) bandpass filter before reaching a blue-optimized photomultiplier tube, and a green-detection region (target and AF) that is passed through a 531-nm (\pm 23 nm) bandpass filter before being detected by a green-optimized photomultiplier tube. The researchers created an algorithm to multiply the blue (AF only) image by a calibration factor and then subtract this, pixel by pixel, from the green (target plus AF) image.⁴

The researchers then demonstrated the efficacy of this novel AF-mitigation algorithm on a tissue phantom grown to simulate an esophagus with regions of HGD cells.⁷ Specifically, digested bovine collagen was reconstituted as a gel to form a support matrix, which was seeded with an HGD Barrett's esophagus cell line. This was then molded into a tubular



Figure 4. The OBIS Galaxy optical bus uses eight wavelength-dedicated plug-and-play input sockets to combine multiple lasers in a single output fiber. Courtesy of Coherent.

3-D structure to roughly simulate the shape of a real esophagus.

Figure 3 shows typical image data from the study, together with a graphical illustration to simplify the interpretation of the images.⁸ "These data show that our approach worked even better than we had hoped," Yang said. "The intensity ratio between the target and background (T/B) was increased by nearly two orders of magnitude. This dramatic increase moves us another step closer to a real-time and quantitative clinical tool that could increase survival rates for esophageal cancer through very early detection and intervention."

Multiple lasers in a single fiber

As with many other multiwavelength applications in bioinstrumentation (e.g., cytometry and microscopy), combining multiple lasers into a single optical fiber was one of the biggest practical challenges in successfully building this multiwavelength SFE. There are two aspects to the well-known challenge of delivering multiple lasers via a single fiber.

First is the basic question of coupling a laser into a single-mode fiber. Many instruments, including this endoscope, must use polarization-preserving singlemode fiber, which has a core diameter of only 3.5 μ m. Efficient coupling of a laser into such a small fiber core necessitates optimization in up to 6° of adjustment (X, Y, Z and three rotational angles) at submicron precision. It also can take several hours, even for an experienced technician. Moreover, these adjustments can drift over time because of thermal or mechanical effects, which is why permanently aligned pigtailed modules such as the OBIS FP series from Coherent have been an important advance for instrument builders.

In instruments with multiple lasers, the situation is much more complex –

accordingly, combining multiple lasers into a single fiber has traditionally been one of most challenging optomechanical tasks in biophotonics. Typically, this has been done in a serial manner using macroscopic, collimated beams: adding a second laser, then a third and so on. This necessitates the cost and complexity of dichroic beamsplitters, polarizers and wave plates at each successive laser wavelength. Then the collinear beams have to be carefully coupled into the output delivery fiber. Moreover, the optical characteristics of the dichroic coated optics are very angle dependent and can result in significant losses. Plus, the numerical aperture (NA) of conventional fiber varies with wavelength. The end result is a typical maximum optical throughput of around 30 percent per laser. This optical hurdle is in complete contrast to the computers and smart devices that make up the electronic part of biophotonics, where USB and Firewire plug-and-play functionality have long since made the soldering iron a relic in most labs.



Fiber combining

A completely different type of beamcombining module – the OBIS Galaxy – brings the same plug-and-play functionality and flexibility to fiber-coupled lasers that USB and other bus technologies brought to computers and smart electronics (Figure 4). It also preserves singlemode output and linear beam polarization, requires no mechanical adjustments whatsoever and is completely passive (i.e., does not use any electronics). And, like USB, this patented module is designed to function as an open-access architecture using standard available fiber connectors (FC/UFC), so that it can be used with any fiber-coupled laser that meets the requisite performance levels in the areas of beam quality, and wavelength fidelity and stability.

As can be seen in Figure 4, the eight wavelength-dedicated laser input sockets use the latest FC/UFC connectors that feature maximum optical registration precision. Each input is then collimated as a macroscopic beam using a rigidly mounted lens before all these collimated beams are integrated in a single monolithic glass element. The light is then coupled into the output fiber that terminates in an FC/APC connector to enable simple connection into an experiment or instrument. The laser inputs span the visible spectrum and include legacy wavelengths such as 488, 514 and 532 nm, as well as the relatively new yellow wavelengths (552 and 561 nm) enabled by optically pumped semiconductor laser technology. And because the combiner uses a refractive optic strategy to combine up to eight different laser inputs, as well as a proprietary output fiber whose NA is independent of wavelength, the performance of this device is both efficient and stable; throughput is up to 75 percent for each laser.

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