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Abstract. We develop a coherent anti-Stokes Raman scattering (CARS) microscopy system equipped with a tunable picosecond laser for high-speed wavelength scanning. An acousto-optic tunable filter (AOTF) is integrated in the laser cavity to enable wavelength scanning by varying the radio frequency waves applied to the AOTF crystal. An end mirror attached on a piezoelectric actuator and a pair of parallel plates driven by galvanometer motors are also introduced into the cavity to compensate for changes in the cavity length during wavelength scanning to allow synchronization with another picosecond laser. We demonstrate fast spectral imaging of 3T3-L1 adipocytes every 5 cm⁻¹ in the Raman spectral region around 2850 cm⁻¹ with an image acquisition time of 120 ms. We also demonstrate fast switching of Raman shifts between 2100 and 2850 cm⁻¹, corresponding to CD₆ symmetric stretching and CH₃ symmetric stretching vibrations, respectively. The fast-switching CARS images reveal different locations of recrystallized deuterated and nondeuterated stearic acid.

Keywords: CARS microscopy; picosecond mode-locked laser; high-speed wavelength scanning; AOTF; fast spectral imaging; adipocytes; deuterated/non-deuterated stearic acid.

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1 Introduction

Raman microscopy is a powerful tool for biochemical observation and analysis that enables nonskinning, noninvasive imaging with high selectivity of specific chemical structures and properties by directly detecting their molecular vibrations. Some biological studies demand observation times on the order of several seconds to minutes, such as observation of lipid droplet dynamics (e.g., exchange and transfer of lipid droplet contents) and fatty acid uptake into lipid droplets for energy storage, and fatty acid oxidation for providing bioenergy in muscle and heart tissue. However, biological processes may occur faster, and therefore, an imaging system capable of acquiring images at video rates or faster would be advantageous. Unfortunately, conventional spontaneous Raman microscopy requires long acquisition times due to the extremely small Raman scattering cross-section.

Coherent Raman scattering (CRS) microscopy, particularly coherent anti-Stokes Raman scattering (CARS) and stimulated Raman scattering (SRS), offers a solution for more-reliable, label-free, fast imaging in biochemical and biomedical applications. CRS microscopy employs two laser sources with slightly different frequencies (ω_pump, ω_Stokes). If the frequency difference between the two lasers coincides with a specific molecular vibration (Ω), CRS images can be obtained due to new emission at ω_СARS = 2ω_pump - ω_Stokes (CARS)[12-14] or energy transfer from ω_pump to ω_Stokes (SRS).[12-14] Though CRS microscopy provides fast imaging with high spatial resolution, it is capable of detecting only a specific wavelength or Raman shift. A further challenge to enhance the potential of CRS microscopy for biological applications is to realize fast spectral imaging in order to instantly detect changes in a specific molecular structure and to simultaneously show multiple molecular structures.

A tunable laser source is required for providing a Stokes beam for exciting various molecular vibrations so that specific vibrational signatures in Raman spectra can be obtained. Therefore, the tunable laser should have high spectral resolution yet wide-range tunability. To observe the dynamics of biochemical processes, the laser should also have a fast tuning ability while providing sufficient, stable power. Moreover, good synchronization between the pump and Stokes beams should be maintained during the wavelength tuning. A picosecond mode-locked laser has been used as a narrowband tunable laser source to achieve high spectral resolution.[12,15-17] In biological applications, Raman spectra are usually observed in fingerprint and CH vibrational regions, typically around 1000 and 3000 cm⁻¹, respectively.[10,11,14] Further enhancement can be achieved by extending the observation range to the silent region at 1800 to 2800 cm⁻¹ where some Raman-tag molecules generate signals without any interference from biological specimens.[18-20]

The most common light sources for CRS microscopy are based on optical parametric oscillators (OPOs).[13,21-27] OPOs have a sufficiently wide tunable range to cover the entire Raman spectral region. However, OPOs require at least several tens of seconds to cover such a wide tuning range. Taking advantage of maintenance-free and compact designs, recent CRS microscopy systems employ all-fiber lasers[28] or a combination of a fiber laser and a picosecond mode-locked laser.[29] Because of material nonlinearity, the length of the fiber should be taken...
into account, and this limits the spectral width and tunable range of the laser pulses.\textsuperscript{28}

In this study, we developed a CARS microscopy system capable of fast acquisition and high-speed spectral scanning. The system employed a picosecond mode-locked laser equipped with an acousto-optic tunable filter (AOTF) inside the laser cavity to change the laser wavelength. We modified the laser to compensate for fluctuations in the repetition frequency of the output laser pulses by installing a moveable end mirror of the laser cavity attached on a piezoelectric actuator and a pair of controllable parallel plates on galvanometer motors. During synchronization, the AOTF laser was tunable from 800 to 940 nm with a typical pulse duration of 10 ps and a resynchronization duration of 20 ms/wavelength change.\textsuperscript{30} An average laser power >300 mW was high enough for obtaining CARS images with multifocus scanning.\textsuperscript{30} For demonstration purposes, fast spectral images of lipid droplets in 3T3-L1 adipocytes were obtained every 5 cm\(^{-1}\) within a Raman spectral region around 2850 cm\(^{-1}\) at 120 ms/image. We also obtained fast-switching CARS images of a mixture of deuterated and nondeuterated stearic acid at 520 ms/image. For that purpose, we rapidly switched the wavelength of the AOTF laser between 844 and 900 nm, whereas another laser wavelength remained at 709 nm. Those wavelengths correspond to Raman shifts of 2100 cm\(^{-1}\) (CD\(_2\) stretching in deuterated stearic acid) and 2845 cm\(^{-1}\) (CH\(_2\) stretching in nondeuterated stearic acid).

2 Development of Fast Spectral Imaging System

2.1 Picosecond Laser with High-Speed Wavelength Scanning System

We found that the development of an ideal tunable laser source for realization of fast spectral CRS microscopy still remains a challenge. We decided to construct a CARS system with two picosecond mode-locked Ti:sapphire lasers, one of which was a high-speed tunable laser (Mega Opto Co. Ltd., Saitama, Japan). The high-speed tunable laser was equipped with an AOTF inside its cavity for tuning the wavelength electrically.\textsuperscript{31} The AOTF was a piezoelectric device made from a birefringent crystal that alters the diffracted wavelength of incident light by applying suitable radio frequency waves without the need for any mechanical parts.\textsuperscript{32} Therefore, the laser could be rapidly tuned from 800 to 940 nm with a typical pulse duration of 10 ps within 1 ms/wavelength change. An average laser power >300 mW was high enough for obtaining CARS images with multifocus scanning.\textsuperscript{30}

Figure 1 shows a schematic diagram of the modified AOTF laser for synchronization with another picosecond mode-locked laser. A prism pair was provided to passively reduce the dispersion of the laser cavity. An end mirror attached on a piezoelectric actuator and a pair of parallel plates driven by galvanometer motors were provided to actively compensate for changes in the repetition frequency.

The repetition frequency of a mode-locked laser depends on the cavity length. During the wavelength scanning of the AOTF laser from 800 to 940 nm, the repetition frequency varied greatly, by a value of 44 kHz. This mainly occurred due to the refractive index dispersion of the AOTF crystal, even though a prism pair (SF-10) was placed inside the laser cavity. The relationship between the repetition frequency, \(f\), and the laser cavity length, \(L\), is given by \(f = c/2L\), where \(c\) is the speed of light. A large change in the repetition frequency leads to difficulty in pulse synchronization; thus, compensation is required to maintain the repetition frequency.

We attached the end mirror of the laser cavity on a piezoelectric actuator (Model 840.10, 15 µm, Physik Instrumente GmbH & Co.KG, Karlsruhe, Germany) to compensate for small changes in the repetition frequency and to synchronize both lasers. Unfortunately, its controllable range of 300 Hz was not enough for compensation of the 44-kHz repetition frequency change. To achieve larger compensation, we placed a pair of parallel plates (BK7, 12 × 4.3 × 10 mm\(^3\)) mounted on galvanometer motors (LSA-20B-30, Harmonic Drive Systems, Tokyo, Japan) inside the laser cavity. The parallel plates were rotated in opposite directions by the same angular increment, which was rapidly and precisely controlled by the galvanometer motors. This configuration allowed repetition frequency compensation of 65 kHz, which was sufficient for our purposes.

2.2 Synchronization System for Picosecond Mode-Locked Lasers

Both laser pulses in CARS microscopy must spatio-temporally overlap. To achieve spatial overlapping, one can adjust the beam pathways in the optical setup. The degree of temporal overlapping is determined by timing jitter during synchronization of both laser pulses. Reduction of timing jitter is very important for improving the signal-to-noise ratio of the CARS image. We employed two types of timing-jitter detection systems to

![Modified acousto-optic tunable filter (AOTF) laser with a moveable end mirror attached on a piezoelectric actuator and a pair of parallel plates on galvanometer motors.](https://www.spiedigitallibrary.org/journals/Journal-of-Biomedical-Optics)
synchronize the lasers: an electronic phase detector for coarse control of synchronization and a balanced cross-correlator for fine control (Fig. 2).33,34

The output from the electronic phase detector represented picosecond-order temporal differences between the two laser pulses.33 This signal was fed back to the galvanometer motors via a digital signal processor (DSP) controller (C6713DSK, Texas Instruments Inc., Texas) to rotate the pair of parallel plates to compensate for variations in the repetition frequency caused by wavelength scanning and to synchronize both the lasers. The electronic phase detector output depended on the time difference and amplitudes of the two input pulses. Therefore, we introduced an autogain amplifier (AD8368, Analog Devices Inc., Boston) after each photodetector (Fig. 3) to keep the amplitudes stable. The autogain amplifier generated a 224 mV rms electrical output signal for any photodetector signal with an amplitude >63 mV rms.

The output from the balanced cross-correlator obtained by optical detection was used for fine control of synchronization because it could detect the femtosecond-order temporal differences between the two laser pulses.34 Since the stability of the balanced cross-correlator output highly depended on the balance of cross-correlation signals detected by two-photon detectors, it was necessary to conserve equal separation of the pump and Stokes pulses to both detectors. As shown in Fig. 4, a polarizing beam splitter (PBS052, Thorlabs Inc., New Jersey) together with two achromatic half-wave plates (AHWP05M-980, Thorlabs Inc., New Jersey) enabled us to easily control and maintain the balance of the transmitted and reflected intensities of the incident pulses so that the balanced cross-correlator could minimize the timing jitter significantly. The output from the balanced cross-correlator was fed back to the piezoelectric actuator via an analog proportional-integral-differential (PID) controller (T-PID01Z, Turtle Industry Co. Ltd., Ibaraki, Japan) and to the galvanometer motors.

2.3 Fast Spectral CARS Imaging System

We developed a fast spectral CARS microscopy system by adopting a multifocus CARS configuration with a microlens array scanner and used the AOTF laser described above as one of the laser sources in the multifocus CARS imaging system (Fig. 5). Details of the multifocus microscopy system have been described in our previous article.35 The system covers the

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Fig. 2 Synchronization system: timing jitter detection system, proportional-integral-differential (PID) and digital signal processor (DSP) for feedback control to galvanometer motors and piezoelectric actuator controllers in the AOTF laser.

![Fig. 2 diagram](https://www.spiedigitallibrary.org/journals/Journal-of-Biomedical-Optics própria)

Fig. 3 Electronic phase detector with autogain amplifier. PD1, PD2 = photodetectors; L1, L2 = lenses.

![Fig. 3 diagram](https://www.spiedigitallibrary.org/journals/Journal-of-Biomedical-Optics própria)

Fig. 4 Balanced cross-correlator with polarizing beam splitter and half-wave plates. PBS = polarizing beam splitter; HWP1, HWP2 = half-wave plates; TPD1, TPD2 = two-photodetectors; OL1, OL2 = objective lenses; MP1, MP2 = dichroic mirrors with highly reflective mirrors.

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Raman spectral range from 400 to 2260 cm$^{-1}$ with a pump beam at 775 nm and from 1600 to 3470 cm$^{-1}$ with a pump beam at 709 nm.

Personal computers (PC 1 and PC 2) were mainly used for controlling the AOTF laser operation and image acquisition, respectively. Before starting spectral imaging, the AOTF laser pulses and the picosecond mode-locked laser pulses should be synchronized. This synchronization was conducted by employing a phase-locked loop mechanism using the electronic phase detector and the balanced cross-correlator as parts of the pulse synchronization system. It was also necessary to store proper amplitude and frequency values of the RF wave in the AOTF driver for determining the wavelength change of the AOTF laser. During the wavelength scanning, PC 1 sent trigger pulses to the AOTF driver to start the wavelength changes. Wavelength changes were performed rapidly within 1 ms or less, and the pulse synchronization system attempted to recover the synchronization within 20 ms. Then, the DSP sent a trigger signal to an electron multiplying CCD (EM-CCD) camera (Ixon, Andor Technology Plc, Belfast, United Kingdom) for image acquisition, and the images were stored in PC 2. The total time required to complete imaging at every wavelength change should include both the image acquisition and the resynchronization process. The wavelength change, the resynchronization, and the image acquisition processes were repeated until the scanning process was completed.

3 Application of Fast Spectral CARS Imaging

Lipids play an important role in living cells for protein sorting and transport, construction of cellular compartments, storage of biofuels, and other biological functions. The observation of lipid metabolism in living cells is challenging since it still remains unclear due to the complex metabolic pathways involved as well as a wide diversity of structures and functions. Observation of lipids using Raman-based microscopy techniques can be easily performed since lipids contain abundant CH bonds, which have a strong Raman band at a specific region around 2850 cm$^{-1}$. Thus, imaging of lipids in living cells with Raman-based microscopy can be achieved by detecting intracellular lipid accumulation in lipid droplets.

3.1 Fast Spectral CARS Imaging of Lipid Droplets in 3T3-L1 Adipose Cells

We demonstrated the performance of our CARS microscopy system for imaging at rates faster than video rates (~33 ms/frame). We tuned the Raman vibration to 2850 cm$^{-1}$ for imaging lipid droplets in 3T3-L1 adipocytes and acquired CARS images at 10 ms/frame. The sample stage was moved during imaging. CARS images observed at some time points are shown in Fig. 6. Movie 1 presents a recording of a 100-fps imaging process, showing timestamps.
Spectral CARS images of 3T3-L1 adipocytes were observed by scanning the wavelength of the AOTF laser every 0.5 nm from 875.0 to 905.0 nm, which corresponded to a Raman shift of 2663.21 to 3049.41 cm\(^{-1}\). For a 100-ms exposure time, the imaging process requires 120 ms/image to cover the wavelength change, laser resynchronization after the wavelength change, and image acquisition. To produce final images, we removed the contributions of dark noise and the fundamental lights by subtraction. We also considered intensity fluctuations of the AOTF laser during wavelength scanning and took this into account by division. Figure 7 shows spectral CARS images of 3T3-L1 adipocytes at (a) 2663.21 cm\(^{-1}\), (b) 2845.31 cm\(^{-1}\), and (c) 2896.26 cm\(^{-1}\), which were assigned to off-resonant vibration, CH\(_2\) symmetric stretching vibration, and CH\(_3\) stretching vibration, respectively. Movie 2 presents a recording of spectral CARS images, showing the Raman shift. A CARS spectral profile in Fig. 7(d) was reconstructed by averaging the intensity in a 5×5-pixel area of interest in Movie 2 [indicated by white arrow in Figs. 7(a)−7(c)]. The reconstructed CARS spectral profile included both resonant and nonresonant contributions and it agrees with the results in previous reports.\(^{25,41,42}\) However, lipid abundantly contained CH bonds and they were highly accumulated in the lipid droplets. It led to the generation of large CARS signal that was relatively higher than the nonresonant background.

**3.2 CARS Imaging by Fast-Switching Between CD and CH Stretching Bands**

Intracellular lipids can be easily detected using CARS microscopy because they contain abundant CH bonds, which have a strong Raman vibration around 2850 cm\(^{-1}\). Lipids cover a large group of biochemical structures with various functions and complex metabolism profiles. Therefore, it is important to distinguish lipid subspecies. One potential solution for distinguishing them is to use a deuterated lipid that consists of CD bonds instead of CH bonds on its carbon chain. A deuterated lipid has a Raman vibration that is significantly downshifted to the silent region around 2100 cm\(^{-1}\) where no interference from biological specimens occurs.\(^{31,43-47}\)

A stable, rapid switching process between two or more different wavelengths is needed for fast imaging to obtain the CARS images of different structures at various Raman shifts. We observed a sample of colocalized deuterated and nondeuterated stearic acids for a rapid switching experiment using our fast spectral CARS microscopy system. Each fatty acid was separately prepared as a stock solution of 40 mM in methanol. First, 5 \(\mu\)L of the deuterated stearic acid solution was dropped onto a glass-based dish (12 mm, AGC Techno Glass Co. Ltd., Shizuoka, Japan). The solvent easily evaporated in the room temperature and the deuterated stearic acid crystals were formed. Then, 5 \(\mu\)L of the nondeuterated stearic acid solution was added. With such small amount of solution, the solvent of the added solution was inadequate to dissolve the deuterated stearic acid crystals because it quickly evaporated. Therefore, we were able to obtain the colocalized deuterated and nondeuterated stearic acids crystals. This sample enabled us to detect sufficiently high Raman signals of CH\(_2\) stretching and CD\(_2\) stretching at 2850 and 2100 cm\(^{-1}\), respectively [Figs. 8(a) and 8(b)]. For that purpose, the AOTF laser was rapidly switched between 833 and 888 nm, and CARS images...
were acquired within 520 ms/image. The merged CARS image indicated the locations where the deuterated and nondeuterated stearic acids were independently and colocatedly recrystallized [Fig. 8(c)]. Hence, the CD bond serves as a powerful chemical tag in CARS microscopy. For instance, in the detection of lipid uptake metabolism by cells, the exogenous lipid can be prepared using its deuterated isotope. The exogenous lipid should be easily detectable because naturally no deuterated isotope presents inside the cells. Thus, it is possible to observe lipid dynamics and metabolism in cells.

4 Conclusion

We succeeded in compensating for changes in the repetition frequency of an AOTF laser by using a pair of parallel plates mounted on galvanometer motors and a moveable end mirror of the cavity attached on a piezoelectric actuator. We employed the AOTF laser as a Stokes pulse source and constructed a fast spectral CARS microscopy system. Fast imaging was achieved by use of a microlens array, and good spectral imaging was demonstrated by using the AOTF laser. To improve the performance of spectral CARS microscopy, we minimized the timing jitter of both laser pulses and optimized the synchronization.

The tunable wavelength range of the AOTF laser was from 800 to 940 nm, which is assigned to Raman shift regions of 400 to 2260 cm\(^{-1}\) and 1600 to 3470 cm\(^{-1}\) with the pump laser set at 775 and 709 nm, respectively. Molecular vibrations in the fingerprint region are highly sensitive to distinguish various molecular species. Therefore, it enables a molecularly sensitive detection for biological applications using CARS microscopy.

Detection of CARS signals in the fingerprint region is still very challenging because the detected signals are typically weak and may possibly be overwhelmed by the nonresonant background contribution. Further experiments and careful observations in the fingerprint region are required to reveal the capability of our developed system. Fast spectral imaging of the Raman shift of the CH stretching vibration was demonstrated by observing lipid droplets in adipose cells. Our system was capable of fast switching and imaging between two significantly different wavelengths corresponding to CD\(_2\) stretching and CH\(_2\) stretching vibrations that belonged to deuterated and nondeuterated stearic acid, respectively. Similar to a four-wave mixing process, we also can employ the AOTF laser to realize fast spectral SRS microscopy.

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