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Abstract. The primary goal of this study is to demonstrate that stimulated Raman scattering (SRS) as a new imaging modality can be integrated into a femtosecond (fs) nonlinear optical (NLO) microscope system. The fs sources of high pulse peak power are routinely used in multimodal nonlinear microscopy to enable efficient excitation of multiple NLO signals. However, with fs excitations, the SRS imaging of subcellular lipid and vesicular structures encounters significant interference from proteins due to poor spectral resolution and a lack of chemical specificity, respectively. We developed a unique NLO microscope of fs excitation that enables rapid acquisition of SRS and multiple two-photon excited fluorescence (TPEF) signals. In the *in vivo* imaging of transgenic *C. elegans* animals, we discovered that by cross-filtering false positive lipid signals based on the TPEF signals from tryptophan-bearing endogenous proteins and lysosome-related organelles, the imaging system produced highly accurate assignment of SRS signals to lipid. Furthermore, we demonstrated that the multimodal NLO microscope system could sequentially image lipid structure/content and organelles, such as mitochondria, lysosomes, and the endoplasmic reticulum, which are intrinsically linked to lipid metabolism.

Keywords: lipid droplets; fluorescence lifetime; stimulated Raman scattering; lysosome; mitochondria; endoplasmic reticulum.

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Lipid droplets (LDs) are highly conserved organelles where fat storage and metabolism are regulated. Quantitative measurement of LDs’ distribution and content is critical for understanding how cellular fat storage is regulated under physiological and pathological conditions, including obesity and related metabolic diseases, such as type 2 diabetes, hypertension, and cardiovascular diseases. Coherent Raman scattering microscopy has been demonstrated as a powerful label-free method for quantitative imaging of subcellular lipid and vesicular structures based on the vibration of carbon-hydrogen (C=H) bonds. Specifically, stimulated Raman scattering (SRS) microscopy provides a straightforward way to quantitatively interpret the optical signal because it is free from nonresonant background and nonlinear concentration dependence. A typical picosecond (ps) SRS takes advantage of the narrow bandwidth (<10 cm⁻¹) to achieve high spectral resolution of specific molecules. More recently, a chirped or spectrally tailored broadband femtosecond (fs) laser, essentially equivalent to a ps laser, has also been demonstrated to achieve spectral imaging and detect multiple components. However, ps sources, with their relatively low peak power, are less suitable than fs sources for multimodal nonlinear optical (NLO) imaging. This is because the efficiency of the two-photon excitation process is inversely proportional to the excitation pulse duration. The ps sources are less efficient in exciting the commonly used NLO signals, such as two-photon excited fluorescence (TPEF) and second harmonic generation.

Despite the increasing popularity of SRS for label-free and quantitative lipid imaging, systems based on fs excitation suffer from the detection of false positive signals from protein interference due to poor spectral resolution of the fs sources. This confounds the use of fs SRS as an optical method for accurate measurements of lipids in living biological systems though it is a more desirable excitation source for multimodal NLO imaging. In this study, we developed a unique fs NLO microscope system for characterizing LDs at the subcellular level in *C. elegans*. The system can sequentially excite and probe the SRS signals from the stretching vibrations for lipids/proteins and the TPEF signals of tryptophan and fluorescent proteins, respectively. The cross-filtering of label-free signals from the multimodal system allowed precise identification of LDs. The accuracy of LD identification is further validated by colocalization of the LD marker short-chain dehydrogenase (DHS-3): green fluorescent protein (GFP).

Using the single-copy transgene technology, we generated a transgenic *C. elegans* strain that expressed a DHS-3::GFP fusion protein in the intestine at physiological levels. DHS-3 was originally identified from proteomic studies as one of the most abundant LD associated proteins in *C. elegans*. All *C. elegans* strains were cultured on nematode growth media plates seeded with OP50 Escherichia coli at 20°C. Worms were paralyzed in 0.2 mM levamisole prior to mounting on microscope slides immediately before SRS and TPEF imaging. We first conducted time- and spectral-resolved measurements of emissions from GFP and autofluorescent entities in lysosome-related organelles (LROs). The two-photon microscope was described in the previous study. Briefly, the standard TPEF microscope was equipped with a spectrograph and a multichannel time-correlated single photon counting (TCSPC) module (PML-16-C-0 and SPC-150, Becker & Hickl GmbH). The detection system provided the capability of time- and spectral-resolved measurements at each pixel of a TPEF image from 400 to 600 nm. Throughout this study, we used one-day-old adult *C. elegans*

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for imaging. As shown in Figs. 1(a) and 1(c), the long-lifetime GFP of C. elegans peaks at ∼510 nm, while the short-lifetime autofluorescence covers a wide range from 450 to 600 nm. The lifetime of GFP and autofluorescence calculated from a dual exponential decay model was 2610 and 970 ps, respectively, indicating that the two fluorescence signals can be well separated in the time domain. In Fig. 1(b), the TPEF signals of each pixel in the wavelength region from 490 to 540 nm, dominated by GFP, were displayed in green, whereas the signals in the regions from 400 to 490 nm and 540 to 600 nm, dominated by the autofluorescence of LROs, were presented in red. In Fig. 1(d), we used a simple nonfitting method based on the ratio of the fluorescence signals in the peak region and the tail region shown in Fig. 1(e) to differentiate the GFP signal and the autofluorescence. As can be seen, both time- and spectral-resolved imaging methods yielded identically accurate separation of GFP signals from LROs. In the following study, genuine signals derived from GFP labeled LDs will serve as the reference for assigning the subcellular origins of SRS signals. It should be emphasized that the GFP signals, localized exclusively on the LD surface, can only be used to identify LDs, but not detect the concentration of lipid in LDs.

It has always been a challenge to distinguish between lipid and protein because their Raman spectra overlap in the CH2 and CH3 bonds region. Though spectral SRS imaging based on the tuning of narrowband ps excitation and the multivariate spectral analysis has enabled accurate measurements of different chemicals, these techniques cannot be directly adopted to a multimodal NLO microscope system based on broadband fs excitations that cover both CH2 and CH3 bonds. However, we previously demonstrated that the TPEF of tryptophan, an essential amino acid serving as one of the building blocks in many proteins, conveys the information on protein content. The TPEF of tryptophan can be efficiently excited by short-wavelength fs source and the signal can be potentially used as a filter to exclude the interfering protein-rich structures subsequent to fs SRS imaging of LDs.

The schematic of the multimodal NLO microscope is shown in Fig. 2. Briefly, an fs Ti:sapphire laser with 80 MHz repetition rate (Chameleon Ultra II, Coherent, Inc.) was tuned at 830 nm. Twenty percent of its output was used as the pump beam for SRS and the excitation for TPEF signals of GFP and LROs, respectively. The rest of the output was used to pump an optical parametric oscillator (OPO) (Chameleon OPO, Coherent, Inc.) to generate 1100 nm as the Stokes beam for SRS. The frequency difference between 830 nm (pump) and 1100 nm (Stokes) matches the center of the CH3 bond at 2950 cm\(^{-1}\) of lipids and proteins. The spectral widths of pump and Stokes beams are 8 and 13 nm, respectively, significantly broader than the spectral separation between lipid and protein. We found that spectral imaging could not resolve the lipids and proteins by tuning the laser wavelength as in ps SRS systems. A pair of prisms introduced precompensation of dispersion to the Stokes beam before passing through a 10.7 MHz acousto-optic modulator (AOMO 3080-122, Crystal Technology). When exciting the TPEF signal of tryptophan, a mirror was inserted to reflect the Stokes beam to a beta barium borate crystal and generate an fs short-wavelength excitation. The excitation beams were combined by dichroic mirrors (DM1: 900DCXR and DM2: 710DCXR, Chroma) and scanned laterally by a pair of galvo mirrors (6210H, Cambridge Technology). A water immersion objective (UAPO40XW3/340, 1.15 NA, Olympus) was driven by an actuator (Z625B, Thorlabs) to achieve the axial scanning. The backward-scattered TPEF signals of GFP, LROs, and tryptophan were recorded by a single channel hybrid detector and TCSPC module (HPM-100-40 and SPC-150, Becker & Hickl GmbH). Two sets of dichroic mirrors and bandpass filters (DM3/4 and Fs1: FF665-Di02, Semrock and ET510/30m, Chroma; FF510-Di01 and FF01-357/44, Semrock) were interchangeable by driving a pair of motorized filter flippers (MFF101/M, Thorlabs) for the measurements of TPEF signals excited at 830 and 550 nm sequentially. At 830 nm excitation, the TPEF signals of GFP and LROs were
recorded simultaneously and separated in the time-domain as described previously based on their lifetime difference. Signals derived from tryptophan containing proteins were obtained at 550 nm excitation. In the forward detection, both of the pump and Stokes beams were collected by a condenser (U-AAC, Achromat/aplanat condenser, NA 1.4, Olympus). The pump beam was further selected by another pair of filters (Fs2: 64335 shortpass at 900 nm, Edmund and FF01-794/160, Semrock) and recorded by a photodiode (FDS 100, Thorlabs). The SRS signal was demodulated by a high-frequency lock-in amplifier (SR844, Stanford Research Systems). The excitation powers of 550, 830, and 1100 nm beams at samples were 6.9, and 10.6 mW, respectively. The image acquisition time was 4 s for each of the three images.

To improve the veracity of LDs’ detection using fs SRS, we examined one-day-old adult transgenic C. elegans that expressed the lipid droplet marker DHS-3::GFP. The representative raw SRS image of aliphatic C–H bonds, raw TPEF image of tryptophan, and color-coded TPEF lifetime image of GFP/LROs acquired at the same site are shown in Figs. 3(a), 3(b), and 3(c), respectively. All raw images were sequentially acquired from a small area of 30 μm × 30 μm (256 pixels × 256 pixels) to ensure high-quality detection of LDs with minimal dwell time. The fs SRS provides information of both lipids and proteins, while the tryptophan fluorescence is derived exclusively from proteins. Therefore, we used the tryptophan fluorescence as a label-free filter to decompose the SRS signals into lipid and protein components. Finally, the filtering accuracy was validated by the GFP signals, which were localized exclusively on the LD surface. The processed images after the decomposition of protein-rich vesicles and LDs are shown in Figs. 3(d) and 3(e), respectively. A standard MATLAB®-based blob detection code was used to identify the vesicular structures in the figures. The false positive signals arising from protein-rich vesicles were subtracted in Fig. 3(f). The TPEF image of GFP labeled LDs is shown in Fig. 3(g), in which the signal of LROs with a short lifetime was coded in red and excluded from the GFP labeled LDs. Figure 3(h) is the merged image of Figs. 3(c)–3(e). As can be seen, the overwhelming majority of LD-like structures in the SRS image overlap with the GFP labeled LDs after filtering of tryptophan TPEF signals. More detailed results based on measurements of 20 SRS and TPEF images from five animals are displayed in Fig. 3(i). Statistically, we found that the colocalization of raw SRS with LD associated GFP signals was only 72 ± 20%. However, after applying the tryptophan fluorescence filtering, the colocalization was improved to 99 ± 1%. This result demonstrates that the false positive features in fs SRS can be effectively eliminated by the cross-filtering technique. It has been reported that the protein droplets largely coincide with LROs and other lysosomal compartments for the breakdown of unwanted polypeptide and lipid species and for protein degradation. Therefore, lysosomes should have a relatively high level of tryptophan and SRS signals.

To demonstrate the multimodal imaging capability, we extended the use of our integrated fs SRS and two-photon
fluorescence microscope system for sequential imaging of subcellular lipid structures and lipid metabolism related organelles based on SRS and TPEF signals, respectively. We chose transgenic *C. elegans* animals in which mitochondria, lysosomes, or the endoplasmic reticulum (ER) was labeled with fluorescent proteins for the study. This was because physical contacts between lipid droplets and the ER and mitochondria are important for the storage and utilization of cellular fat content, respectively.

In addition, mobilized fatty acids may in part be channeled to the LRO, which links aging and metabolic activity. The representative results are shown in Fig. 8. The tryptophan TPEF images were used to identify the false positive structures in the SRS images. The cross-filtered LD signals were then merged with the TPEF images of subcellular organelles labeled with fluorescent proteins. Our results clearly demonstrate that the multimodal SRS and TPEF microscope can not only image the LDs with the critical information on their fat content, but also reveal their engagement with different subcellular organelles. It should be emphasized that with a Ti:sapphire laser of 140 fs pulse duration used in this study, the excitation power for the TPEF imaging of fluorescent protein labeled organelles was always <10 mW. The excitation efficiency is about a factor of 40 higher than the sources of 5 to 7 ps pulse duration that were commonly used for ps SRS imaging. Therefore, the fs system may cause significantly less photodamage to biological samples.

In summary, our fs NLO microscope system is capable of highly specific assignment of LDs by cross-filtering signals from multiple imaging modalities. We showed that false positive signals of SRS are derived from a partially overlapping set of structures which are rich in protein entities. We demonstrated that the multimodal imaging system can sequentially produce quantitative SRS image of lipid structures and the TPEF images of important organelles related to lipid metabolism. Finally, it should be pointed out that the cross-filtering technique may eliminate genuine LDs whose size falls under the diffraction limit. This is because tryptophan signals from proteins on the LD surface will not be optically separated from SRS signals that originate from the lipid core of LDs. Such a concurrence of tryptophan and SRS signals will cause the misassignment of small LDs. In addition, the technique cannot identify protein droplets that do not contain tryptophan.

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