Synchronization-free all-solid-state laser system for stimulated Raman scattering microscopy

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We introduce an extremely simple and highly stable system for stimulated Raman scattering (SRS) microscopy. An 8-W, 450-fs Yb:KGW bulk oscillator with 41 MHz repetition rate pumps an optical parametric amplifier, which is seeded by a cw tunable external cavity diode laser. The output radiation is frequency doubled in a long PPLN crystal and generates 1.5-ps long narrow-band pump pulses that are tunable between 760 and 820 nm with >50 mW average power. Part of the oscillator output is sent through an etalon and creates Stokes pulses with 100 mW average power and 1.7 ps duration. We demonstrate SRS microscopy at a 30-μs pixel dwell time with high chemical contrast, signal-to-noise ratio in excess of 45 and no need for balanced detection, thanks to the favorable noise properties of the bulk solid-state system. Cw seeding intrinsically ensures low spectral drift. We discuss its application to chemical contrast microscopy of freshly prepared plant tissue sections at different vibrational bands.

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INTRODUCTION

Coherent Raman scattering (CRS) is a powerful nonlinear microscopy technique for label-free, three-dimensional imaging of tissues and cells, exploiting the characteristic vibrational response of molecules as an intrinsic contrast mechanism¹. CRS enables real-time, non-destructive chemical identification of compounds at the sub-micrometer scale without addition of any external marker², whose presence could perturb the molecules under study or even inhibit the evolution of dynamical processes, such as drug transport or cell differentiation. CRS microscopy has found many applications in biomedical optics³–⁴; in particular, given the strong Raman response of long-chain hydrocarbons, it is the ideal method for visualizing lipids and studying their metabolism⁵, as well as for differentiating lipids, DNA, and proteins within the cell⁶. In addition CRS promises to be a powerful tool for rapidly discriminate between tumors and healthy tissue, paving the way to ‘virtual histopathology’, both ex vivo and intraoperatively⁷–⁸.

CRS techniques require two synchronized laser pulses at frequencies ωₚ (pump frequency) and ωₛ (Stokes frequency) to drive a vibrational coherence within the ensemble of molecules inside the laser focus. When the difference between pump and Stokes frequencies matches a specific vibrational frequency Ω, i.e., ωₚ − ωₛ = Ω, then all the molecules in the focal volume vibrate in phase, enhancing the Raman response by many orders of magnitude with respect to the incoherent, spontaneous Raman process. The two most widely employed CRS techniques are coherent anti-Stokes Raman scattering (CARS)⁹ and stimulated Raman scattering (SRS)¹⁰,¹¹. In the CARS process the vibrational coherence is read by a further interaction with the pump beam, generating radiation at the anti-Stokes frequency ωₛ = ωₚ + Ω; in SRS coherent interaction with the sample induces stimulated emission from the virtual state of the sample to the investigated vibrational state, resulting in a Stokes-field amplification (stimulated Raman gain, SRG) and in a simultaneous pump-field attenuation (stimulated Raman loss, SRL). CARS can be more simply implemented, since the nonlinear signal is at a frequency ωₛ differing from those of pump and Stokes, and can thus be easily singled out by spectral filtering. SRS is instrumentally more demanding, since it requires the measurement of a tiny differential signal (the SRG or SRL) on top of a large and potentially noisy background (the Stokes or pump intensity). On the other hand, SRS has two important advantages: (i) it is free from non-resonant background, generated by four-wave-mixing¹², which in CARS adds to and often obscures the resonant signal of interest; (ii) it provides a signal that is linear with the density of oscillators, thus allowing quantitative evaluation of the concentrations of the molecular species¹³. In broadband or spectroscopic CARS the non-resonant background acts as a phase-locked local oscillator heterodyning the resonant signal, which can be recovered by numerical processing¹⁴; this procedure is, however, not possible in the single-frequency mode, which allows the highest acquisition speeds up to the video rate¹⁵,¹⁶. For these reasons SRS is currently the technique of choice for a broad spectrum of applications.

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The main stumbling block which has so far prevented widespread adoption of CRS microscopy and, in particular, SRS in the biological and medical communities is the complication and cost of the experimental apparatus. Raman transitions exhibit linewidths of the order of tens of cm\(^{-1}\) in the condensed phase: To efficiently excite them with the highest peak power, while still preserving frequency resolution and molecular selectivity, picosecond pump/Stokes pulses are thus needed. In its single-frequency version, CRS microscopy requires two synchronized trains of narrowband, picosecond pulses (pump and Stokes) with tunable frequency difference, high repetition rate (\(\approx 100\) MHz) and output power \(\approx 100\) mW per branch, which is required to compensate for losses in the optical chain of the microscope. The complexity of the excitation laser is one reason why, after its early demonstration in 1982\textsuperscript{17}, the development of CRS microscopy has stopped for nearly two decades. Following its revival, initial CRS implementations were based on two electronically synchronized picosecond Ti:sapphire oscillators\textsuperscript{18}, while the current ‘gold standard’ is represented by an optical parametric oscillator (OPO) synchronously pumped by a picosecond Nd:YVO\(_4\) oscillator\textsuperscript{19}. Such systems are complex, expensive, and they all critically require the synchronization between two independent laser sources, which must be maintained over time. Drastically simplified excitation architectures, with lower cost and smaller footprint, are thus greatly in demand, as they would lower the technological entrance barriers to CRS microscopy.

In recent years, considerable efforts have been devoted to the development of fiber-format systems for CRS microscopy, as they are intrinsically compact and turn-key. Several architectures, based on Er\textsuperscript{3+}:YAG or Yb-doped fibers combined with nonlinear spectral broadening and frequency conversion, have been demonstrated\textsuperscript{20–29}. While these systems have been successfully applied to CARS imaging\textsuperscript{26–29}, the CRS modality has been much harder and in some cases impossible\textsuperscript{28} to be implemented. This is due to the fact that fiber lasers intrinsically suffer from excess high-frequency noise with respect to their bulk counterparts, because of the much greater length of the active medium which leads to higher amplified spontaneous emission levels. This extra noise pedestal, which can amount to as much as 30 dB, prevents sensitive detection of the weak nonlinear SRS signal, sitting on top of the strong and fluctuating Stokes (pump) signal, unless balanced detection schemes\textsuperscript{22,23,30} with sophisticated noise-cancelling electronics\textsuperscript{23} or complex optical layouts\textsuperscript{30} are used. Such schemes become particularly challenging to be implemented when measuring biological samples, with large variations of transmission/scattering over the scanned area.

Recently we introduced a novel compact and low-cost architecture for CRS microscopy based on a bulk all-solid-state Yb:KGW oscillator followed by an optical parametric amplifier (OPA), employing periodically poled lithium niobate (PPLN)\textsuperscript{31}. OPAs are significantly simpler than OPOs, as they work without a feedback cavity. However, as they require high nonlinear gain levels they are typically driven by amplified laser systems. In our case the high peak power of the Yb oscillator, combined with the large nonlinear coefficient of PPLN, allowed its use to directly drive the OPA at tens of MHz repetition rate, while preserving the excellent noise figure of the bulk solid-state laser. However, a critical aspect of an OPA is the generation of the seed light, which typically requires a nonlinear optical process starting from the pump. Our previous implementation still used a complex optical layout with several OPA stages, and self-phase-modulation in a tapered fiber was required to generate a tunable OPA seed. Here we demonstrate a drastic simplification of the system, in which the OPA is seeded by a tunable continuous wave external-cavity diode laser (ECDL). This configuration completely removes path-length synchronization issues, greatly simplifying the optical setup and enhancing its robustness and insensitivity to environmental perturbations. Exploiting the very low noise of solid-state bulk oscillators in the MHz regime, we demonstrate high speed, low-noise SRS imaging without balanced detection.

**MATERIALS AND METHODS**

Our SRS system as shown in Figure 1 is based on a home-built mode-locked Yb:KGW solid-state oscillator, which provides up to 8 W of output power (with 6.2 W available for the experiment) at 1032 nm with 450-fs pulse width at a repetition rate of 41 MHz\textsuperscript{32}. A 1.5-W fraction of the main laser output is used to pump a single-stage OPA based on a 10-mm-long MgO-doped PPLN crystal, equipped with 9 different poling periods (from \(\Lambda = 27.91\) \(\mu\)m to \(\Lambda = 31.59\) \(\mu\)m). The OPA is seeded by a home-built cw ECDL (gain chip: SAL:1590-100, Sacher Lasertechnik), employing a grating in the Littrow configuration, and generating 30 mW average power over the 1500–1650 nm tuning range with single longitudinal mode operation. Its fibered output provides after collimation a nearly diffraction-limited beam that is mode-matched and collinearly combined with the Yb:KGW output and focused to a 50-\(\mu\)m waist in the PPLN crystal. Cw seeding dramatically simplifies the OPA operation, as there is no need for...
pump-seed synchronization\textsuperscript{3}. Considering the ECDL output power, the pump pulse duration and the repetition rate, there are approximately \(10^7\) seed photons per pulse; this level is sufficient to completely suppress parametric superfluorescence in the OPA, greatly improving the energy stability, removing timing jitter\textsuperscript{24}, and minimizing spectral drift. Further, it enables precise adjustment of the OPA center wavelength by rotating the ECDL grating and changing poling period and crystal temperature accordingly. In the future, a fan-out design will allow for simpler, faster and gapless spectral tuning. Ultimately, using commercially available ECDLs and fast linear translation stages in combination with fan-out crystals, a tuning speed over the entire CH-region below 1 s can be realized. Thanks to the high nonlinear coefficient of PPLN, the output power remains above the 300 mW level across the whole tuning range of the ECDL (Figure 2, lower panel). The spectral bandwidth amounts to 10 nm, dictated by the linewidth of the Yb oscillator, which corresponds to a pulse duration of \(\approx 350\) fs. The Raman pump pulses for the SRS process are obtained by frequency doubling the OPA output in a second 10-mm-long PPLN crystal and exploiting second-harmonic generation (SHG) spectral compression\textsuperscript{35} to reduce the bandwidth to \(\approx 11\) cm\(^{-1}\). A tuning range between 760 and 820 nm, as limited by the available poling periods of the SHG crystal, is obtained with power levels > 50 mW (Figure 2, upper panel). The corresponding pump-Stokes frequency detuning ranges from 2500 to 3460 cm\(^{-1}\), thus fully covering the C-H stretching band that is typically used for the imaging of lipids. By replacing the ECDL and the PPLN crystals, the OPA tuning range can be easily extended to 1770–1870 nm, thus allowing the 1000–1600 cm\(^{-1}\) fingerprint region to be covered after SHG. The fixed frequency 1032-nm Stokes pulses are created from the residual oscillator power by spectral filtering with an etalon with a free spectral range of 85 cm\(^{-1}\) and a finesse of 12, resulting in pulses with \(\approx 8\) cm\(^{-1}\) bandwidth. Considering the 28% transmission of the etalon, the 4.5 W of residual oscillator power would result in 1.26 W maximum Stokes power. However, since typically < 100 mW will be needed, the remaining oscillator power is available for other nonlinear optical microscopy modalities or even to pump additional OPAs.

Pump and Stokes pulses are synchronized by a delay line, collinearly combined by a dichroic beam-splitter and sent to a home-built laser-scanning SRS microscope, working in transmission mode and using a NA = 0.7 focusing objective (Nikon 60x). The transmitted light is collected with a NA = 1.4 achromatic condenser. We chose to detect the SRG of the Stokes beam, which benefits from the extremely good noise properties of the all-solid-state Yb:KGW oscillator. An acousto-optic modulator is used to modulate the pump pulses at 500 kHz, while the Stokes intensity is detected by a single-channel amplified silicon photodiode, which is connected to a high-frequency lock-in amplifier. A pair of galvanometric mirrors is used for the fast spatial scanning of the samples.

**RESULTS AND DISCUSSION**

We first validated the system by measuring SRS spectra of various solvents. Figure 3 shows the SRG spectrum of an acetone solution, as measured by scanning the pump wavelength from 777 to 802 nm; the spectrum and in particular the position of the characteristic peak at 2923 cm\(^{-1}\) is in very good agreement with the spontaneous Raman spectrum, displayed in the inset of Figure 3a, as expected for SRS, which directly measures the imaginary part of the third-order nonlinear susceptibility without lineshape distortions\textsuperscript{10,11}. The stability of the SRS source is monitored in a continuous 24 h long time measurement of the SRG signal at the acetone peak at 2923 cm\(^{-1}\). The system was free-running without any active control or feedback during the whole time. The average SRG signal drift is lower than 3% over this period, with a peak-to-peak difference of 13%, as shown in Figure 3b. Note that the SRG signal depends on multiple parameters, such as pump and Stokes power, wavelength detuning and beam pointing fluctuation; its very low long-term drift without active stabilization is a clear indication of the intrinsic stability of our design. Due to the absence of synchronization issues, the system can provide this performance even after a cold start of the Yb oscillator and the ECDL.

**Figure 2** Power-spectra of external-cavity diode laser seeded OPA (lower panel). Corresponding SHG from PPLN (in green in upper panel) serving as tunable Raman pump. The upper panel also depicts the spectrum of the Stokes beam, filtered by an etalon.

**Figure 3** (a) SRG spectra of acetone solution compared to the spontaneous Raman spectrum shown in inset. (b) Stability of the SRG signal over 24 h at the acetone resonance of 2923 cm\(^{-1}\). The signal is a 5-minute running average, sampled with 1 Hz.
As a first test of the imaging capabilities of our SRS setup, we used a composite sample of poly-methyl methacrylate (PMMA) and polystyrene (PS) beads with a 6-μm diameter, and probed it at frequency detunings of 2953 and 3066 cm\(^{-1}\). These images are acquired with 30 μs pixel dwell time with 16 mW pump (modulated at 500 kHz) and 2 mW Stokes powers respectively, at the sample. The focusing microscope objective (Nikon 60x, 0.70 NA in focus) has ~79% and 44% transparencies for pump and the Stokes wavelengths, respectively. The inset graph (c) shows the measured dependence of the signal-to-noise ratio on integration time for acquisition of PS beads under the same experimental conditions.

Figure 4 (a) and (b): Stimulated Raman gain images (54×54 μm, 150×150 pixels) of a blend of 6-μm PS and PMMA beads taken at their respective Raman resonances of 3066 and 2953 cm\(^{-1}\). These images are acquired with 30 μs pixel dwell time with 16 mW pump (modulated at 500 kHz) and 2 mW Stokes powers respectively, at the sample. The focusing microscope objective (Nikon 60x, 0.70 NA in focus) has ~79% and 44% transparencies for pump and the Stokes wavelengths, respectively. The inset graph (c) shows the measured dependence of the signal-to-noise ratio on integration time for acquisition of PS beads under the same experimental conditions.

Figure 5 (a–c) SRG images (80×80 μm, 400×400 pixels) of cells in an cross-sectional area of Abutilon pictum leaf petiole, containing sclerenchyma (upper left area) and parenchyma cells, acquired at 300 μs integration time at pump-Stokes frequency detunings of 2890, 2945, and 3245 cm\(^{-1}\), respectively. (d) Raman spectra, available in the literature, of key compounds present in the plant cells: polysaccharides including cellulose and pectin, polyphenols (lignin), and water. The intensity in the 2800–3000 cm\(^{-1}\) region of the Raman spectrum originates mainly from C–H stretching vibrations, while the intensity in the 3050–3600 cm\(^{-1}\) region mainly stems from O–H stretching vibrations. (e) Overlay image of all three SRG signals. Exemplary sites of intercellular spaces are indicated by white arrowheads, pits, or plasmodesmata by orange arrowheads. Refer to text for details.

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To demonstrate the performance of the system on a biological sample, we imaged a tissue domain of a leaf petiole prepared from an *Abutilon pictum* plant. Figure 5 shows SRG images of an 80 × 80 μm cross-sectional area of this stem, containing thick-walled sclerenchyma cells (small lumina; top left) and large parenchyma cells (right), at three different pump-Stokes frequency detunings, namely 2890 cm⁻¹ (Figure 5a), 2945 cm⁻¹ (Figure 5b), and 3245 cm⁻¹ (Figure 5c). In Figure 5a and b, the plant cell walls are clearly visible due to the SRG signal originating from the C–H stretching vibrations of the molecules of their major constituents, namely the macromolecular polysaccharides cellulose, xyloglucans, and pectins, as well as xylans and polyphenolic polymers (lignins)³⁶, whose Raman resonances (see spontaneous Raman spectra in Figure 5d) correspond to the adopted frequency detunings. On the other hand, in Figure 5c the SRG signal is dominated by the O–H stretching vibrations of the water molecules, which are homogeneously distributed within the lumina of the cells. In-between adjacent cells, intercellular spaces formed by separation of contiguous walls at junctions of three or more cells can be seen (as indicated exemplarily by white arrowheads in 5e). Pit canals traversing the cell walls, with plasmodesmata as connections between the living protoplasts of adjoining cells, are discernible (exemplified by orange arrowheads). The overlay image 5e in a pseudo-color scheme, using the same color assignments for identifi-

CONCLUSIONS

In this paper we present a novel compact, all-solid-state system for SRS microscopy, based on a mode-locked bulk Yb:KGW oscillator driving an ECDL-seeded OPA. This is a passive, single-pass system, which avoids third-order nonlinearities and synchronization issues, since no delay control is required for the tunable Raman pump generation. Hence, it is not only simpler to set up and operate, with respect to existing solutions based on bulk lasers, but also robust with respect to thermal or environmental changes and well suitable for long term operation. At the same time, our approach avoids the excess high-frequency noise intrinsic in fiber-based laser systems, allowing high-quality SRS imaging without complex balanced detection schemes. In addition, with respect to other solutions for SRS microscopy, our system offers the following important advantages:

i The bandwidths of both pump and Stokes pulses can be easily adjusted to match the linewidth of the vibration of interest by selecting the etalon design and the length of the SHG crystal used for spectral compression³⁵, we chose bandwidths of 10–20 cm⁻¹, corresponding to a pulse duration of 1–2 ps, which is the optimum compromise between peak power and spectral resolution for lipid imaging³⁷.

ii The system also produces high peak power 450-fs pulses, which can be spectrally broadened in a nonlinear fiber and compressed to sub-100-fs duration; these pulses are available for other nonlinear microscopy modalities, such as second and third harmonic imaging or two-photon excited fluorescence.

We believe that our system will significantly reduce the technical entrance barriers to SRS microscopy for label-free chemical contrast imaging, bringing it closer to real-world applications in biomedical and biological research and in diagnostics.

**CONFLICT OF INTEREST**

The authors declare no conflict of interest.

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1 Min W, Freudiger CW, Lu SJ, Xie XS. Coherent nonlinear optical imaging: beyond fluorescence microscopy. *Annu Rev Phys Chem* 2011; 62: 507–530.

2 Camp CH, Cicerone MT. Chemically sensitive bioimaging with coherent Raman scattering. *Nat Photon* 2015; 9: 295–305.

3 Pezacki JP, Blake JA, Danielson DC, Kennedy OC, Lyn RK et al. Chemical contrast for imaging living systems: molecular vibrations drive CARS microscopy. *Nat Chem Biol* 2011; 7: 137–145.

4 Yu Y, Ramachandran PV, Wang MC. Shedding new light on lipid functions with CARS and SRS microscopy. *Biochim Biophys Acta* 2014; 1841: 1120–1129.

5 Zumbusch A, Langbein W, Borri P. Nonlinear vibrational microscopy applied to lipid biology. *Prog Lipid Res* 2013; 52: 615–632.

6 Lu FK, Basu S, Igras V, Hoang MP, Ji MB et al. Label-free DNA imaging in vivo with stimulated Raman scattering microscopy. *Proc Natl Acad Sci USA* 2015; 112: 11624–11629.

7 Ji MB, Orringer DA, Freudiger CW, Ramkissoon S, Liu XH et al. Rapid, label-free detection of brain tumors with stimulated Raman scattering microscopy. *Sci Transl Med* 2013; 5: 201ra119.

8 Ji MB, Lewis S, Camelio-Piragua S, Ramkissoon SH, Snuderl M et al. Detection of human brain tumor infiltration with quantitative stimulated Raman scattering microscopy. *Sci Transl Med* 2015; 7: 309ra163.

9 Zumbusch A, Holton GR, Xie XS. Three-dimensional vibrational imaging by coherent anti-Stokes Raman scattering. *Phys Rev Lett* 1999; 82: 4142–4145.

10 Freudiger CW, Min W, Saar BG, Lu SJ, Holton GR et al. Label-free biomedical imaging with high sensitivity by stimulated Raman scattering microscopy. *Science* 2008; 322: 1387–1386.

11 Nandakumar P, Kovalev A, Volkmer A. Vibrational imaging based on stimulated Raman scattering microscopy. *New J Phys* 2009; 11: 033026.

12 Evans CL, Xie XS. Coherent anti-Stokes Raman scattering microscopy: chemical imaging for biology and medicine. *Annu Rev Anal Chem* 2008; 1: 883–909.

13 Fu D, Zhou J, Zhu WS, Manley PW, Wang YK et al. Imaging the intracellular distribution of tyrosine kinase inhibitors in living cells with quantitative hyperspectral stimulated Raman scattering. *Nat Chem* 2014; 6: 614–622.

14 Camp CH Jr, Lee YJ, Heddleston JM, Hartlhorn CM, Walker ARH et al. High-speed coherent Raman fingerprint imaging of biological tissues. *Nat Photon* 2014; 8: 627–634.

15 Evans CL, Potma EO, Puori/haag M, Cote D, Lin CP et al. Chemical imaging of tissue in vivo with video-rate coherent anti-Stokes Raman scattering microscopy. *Proc Natl Acad Sci USA* 2005; 102: 16807–16812.

16 Saar BG, Freudiger CW, Reichman J, Stanley CM, Holton GR et al. Video-rate molecular imaging in vivo with stimulated Raman scattering. *Science* 2010; 330: 1368–1370.

17 Duncan MD, Reintjes J, Manuccia TJ. Scanning coherent anti-Stokes Raman microscope. *Opt Lett* 1982; 7: 350–352.

18 Potma EO, Jones DJ, Cheng JX, Xie XS, Ye J. High-sensitivity coherent anti-Stokes Raman scattering microscopy with two tightly synchronized picosecond lasers. *Opt Lett* 2002; 27: 1168–1170.

19 Ganikhanov F, Carrasco S, Xie XS, Katz M, Seitz W et al. Broadly tunable dual-wavelength light source for coherent anti-Stokes Raman scattering microscopy. *Opt Lett* 2006; 31: 1292–1294.

20 Krauss G, Hanke T, Seli A, Trudelien D, Leitenstorfer A et al. Compact coherent anti-Stokes Raman scattering microscope based on a picosecond two-color Er: fiber laser system. *Opt Lett* 2009; 34: 2847–2849.

21 Marangoni M, Gambetta A, Marzoni C, Kumar V, Ramponi R et al. Fiber-format CARS spectroscopy by spectral compression of femtosecond pulses from a single laser oscillator. *Opt Lett* 2009; 34: 3262–3264.

22 Gambetta A, Kumar V, Grancini G, Polli D, Ramponi R et al. Fiber-format stimulated-Raman-scattering microscopy from a single laser oscillator. *Opt Lett* 2010; 35: 226–228.

23 Freudiger CW, Yang WL, Holton GR, Peghambarian N, Xie XS et al. Stimulated Raman scattering microscopy with a robust fibre laser source. *Nat Photon* 2014; 8: 153–159.

24 Coluccelli N, Kumar V, Cassiniero M, Galzerano G, Marangoni M et al. Er:Tm: fiber laser system for coherent Raman microscopy. *Opt Lett* 2014; 39: 3090–3093.

25 Baumgartl M, Chemnitz M, Jauregui C, Meyer T, Dietzek B et al. All-fiber laser source for CARS microscopy based on fiber optical parametric frequency conversion. *Opt Express* 2012; 20: 4484–4493.

doi:10.1038/lsa.2016.149

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26 Gottschall T, Baumgartl M, Sagnier A, Rothhardt J, Jauregui C et al. Fiber-based source for multiplex-CARS microscopy based on degenerate four-wave mixing. Opt Express 2012; 20: 12004–12013.

27 Chemnitz M, Baumgartl M, Meyer T, Jauregui C, Dietzek B et al. Widely tuneable fiber optical parametric amplifier for coherent anti-Stokes Raman scattering microscopy. Opt Express 2012; 20: 26583–26595.

28 Lefrancois S, Fu D, Holtom GR, Kong LJ, Wadsworth WJ et al. Fiber four-wave mixing source for coherent anti-Stokes Raman scattering microscopy. Opt Lett 2012; 37: 1652–1654.

29 Lamb ES, Lefrancois S, Ji MB, Wadsworth WJ, Xie XS et al. Fiber optical parametric oscillator for coherent anti-Stokes Raman scattering microscopy. Opt Lett 2013; 38: 4154–4157.

30 Nose K, Ozeki Y, Kishi T, Suminura K, Nishizawa N et al. Sensitivity enhancement of fiber-laser-based stimulated Raman scattering microscopy by collinear balanced detection technique. Opt Express 2012; 20: 13958–13965.

31 Steinle T, Kumar V, Steinmann A, Marangoni M, Cerullo G et al. Compact, low-noise, all-solid-state laser system for stimulated Raman scattering microscopy. Opt Lett 2015; 40: 593–596.

32 Steinmann A, Metzger B, Hegenbarth R, Giessen H. Compact 7.4W femtosecond oscillator for white-light generation and nonlinear microscopy. Proceedings of the Conference on Lasers and Electro-Optics 2011; paper CThA45. Baltimore, Maryland, USA: CLEO, 2011.

33 Steinle T, Kedenburg S, Steinmann A, Giessen H. Combining cw-seeding with highly nonlinear fibers in a broadly tunable femtosecond optical parametric amplifier at 42 MHz. Opt Lett 2014; 39: 4851–4854.

34 Marzoni C, Cirmi G, Brida D, De Silvestri S, Cerullo G. Optical-parametric-generation process driven by femtosecond pulses: Timing and carrier-envelope phase properties. Phys Rev A 2009; 79: 033818.

35 Marangoni M, Brida D, Quintavalle M, Cirmi G, Pigozzo FM et al. Narrow-bandwidth picosecond pulses by spectral compression of femtosecond pulses in a second-order nonlinear crystal. Opt Express 2007; 15: 8884–8891.

36 Evert RF. Esau’s Plant Anatomy: Meristems, Cells, and Tissues of the Plant Body - Their Structure, Function, and Development. Hoboken, New Jersey: John Wiley & Sons, Inc., 3rd ed. 2006.

37 Rimke I, Hehl G, Beutler M, Volz P, Volkmer A et al. Tunable dual-wavelength two-picosecond light source for coherent Raman scattering microscopy. In: Periasamy A, So PTC, König Keditors. Proceedings of SPIE Multiphoton Microscopy in the Biomedical Sciences XIV. San Francisco, CA, USA: SPIE, 2014; 8948: 894816.

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