Development of novel Optics and Analysis Tools for enhancing Biomedical Imaging by Coherent Raman Scattering

PhD Thesis
to obtain the degree of
Doktor der Naturwissenschaften (Dr. rer. nat.)

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on
June 24, 2019
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1. Introduction

Biomedical research has benefitted from the continuous technical developments and improvements that were made to many optical microscopy techniques over the last decades. New, highly relevant topics were addressed in order to understand the origins and early development of human diseases, such as cancer-related growth of cells [Ji et al. 2015], acquired immunodeficiency syndromes [Hübner et al. 2009], age-related affection of the musculoskeletal system [Croft et al. 2019], brain- [Ji et al. 2018] or cardiovascular-related diseases [Huang et al. 2019], to name just a few examples. A specific enabling requirement for targeting these questions was the application of fluorescent dyes for the specific labeling of biological structures of interest in a sample [Taylor and Wang 1980]. The introduction of immunofluorescence staining [Pawley 2006] and genetic modification by gene transfection techniques [Kim and Eberwine 2010] for the active expression of fluorescent dyes in living samples allowed e.g. virus tracking [Chojnacki and Eggeling 2018] or identification of tumor cells and affected tumor tissues [Yang et al. 2017a].

Also the spatial and temporal resolution of the applied microscopy techniques were enhanced, especially by the introduction of novel superresolution techniques, which are able to resolve structures of interest beyond the diffraction limit of light. This task was tackled by exploiting either the stochastic blinking of fluorescent dyes in a multi-image acquisition series and final post-processing [Heilemann et al. 2008], by active engineering of the point spread function of the excitation volume [Klar and Hell 1999] or by applying a structured illumination pattern in a multi-image acquisition scheme followed by post-processing steps [Gustafsson, M. G. L. 2000, Gustafsson, M. G. L. et al. 2000].

All fluorescence-based microscopy techniques share the intrinsic drawback of the need for exogenous fluorescent dyes, which results in laborious preparation protocols, detrimental effects to the sample by the presence of the potentially toxic dyes [Trewin et al. 2018] or influencing time-dependent processes, such as diffusion experiments or lifetime tracking in living cell samples based on the size and weight of the dye. All this affects the response of the sample also due to the necessarily high laser intensities in the visual part of the optical spectrum, resulting in increased phototoxicity [Wäldchen et al. 2015]. Also there are samples of interest, where no sufficient labeling method could be found, yet, e.g. due to the presence of rigid cell walls [Bogen et al. 2013b] or insufficient fluorescent labeling [Cheng and Xie 2015].

An alternative solution to address these biomedical research questions can be found utilizing Raman-based imaging techniques such as label-free stimulated Raman scattering.
coherent anti-Stokes Raman scattering (CARS) [Terhune et al. [1965], Druet, S. A. J. and Taran, J. P. E. [1981]], which offer high chemical specificity and rapid imaging speeds due to their enhanced signal strength by exploiting the coherent nature of the signal. This leads to signals, which are several orders of magnitude stronger than those obtained by spontaneous Raman scattering while they do not suffer from bleaching effects as their fluorescent counterparts. Coherent Raman scattering (CRS) techniques are actively probing the molecular resonance of a large number of the same molecular groups by coherent laser wavelengths, which will perform a pump-probe experiment resulting in the strongly enhanced CRS-signals. Nevertheless, the molecular sensitivity of these techniques is still limited, which hinders the extensive replacement of the established fluorescent microscopy techniques with CRS-microscopy also due to the mostly diffraction limited resolution in relevant biomedical applications [Cheng and Xie [2015]]. This is also the case due to an enhanced level of complexity of the microscope setup as well as to the required cost-intensive and bulky laser systems, which also constrain the clinical use and which is part of ongoing research [König et al. [2011], Saar et al. [2011]].

To address these obstacles a state-of-the-art CRS microscope setup is developed and presented in this PhD-thesis, which utilizes novel optical and analytical tools to further enhance the reproducibility and chemical sensitivity of the CRS techniques. It covers a multimodal acquisition capability of simultaneous CARS and SRS imaging in combination with second harmonic generation (SHG) and two-photon excited fluorescence microscopy enabling high imaging speeds and volumetric scanning of the sample. In addition to that a hyperspectral scanning mode is implemented by computer-controlled wavelengths tuning, which demonstrates the enhanced sample information by sequentially addressing different molecular resonances by CARS [Fussell et al. [2013]]. This technique is further analyzed in terms of reproducibility as well as on specific properties of the laser system used for the experiments and a new level of accuracy is presented to the research community [Pilger et al. [2018]]. In collaboration with several partners the successful application of the newly constructed setup to a range of biomedical problems was investigated and is presented. To further extend the range of applications of CARS microscopy the concept of structured illumination is explored in this scanning-based microscopy technique. The special appeal of this approach is the highly cost-effective way of boosting the resolution using mostly standard components of state-of-the-art two-photon scanning and CRS-microscopes.

For paving the way of CRS microscopy to clinical application a collaborative effort with the University of Hong Kong, with the group of Prof. K. K. Y. Wong, is presented by exploiting fiber-laser technology to construct a novel, low noise all fiber-based two
color CRS laser system, which can replace the current solid-state laser setups offering similar performance. The introduced all-fiber-based laser source is characterized in its performance, which is superior to other published solutions in terms of noise level. This enables rapid, high quality CARS and SRS imaging without the need for additional noise cancellation schemes such as balanced detection devices. The fiber laser is then utilized on several biomedical samples to demonstrate the simultaneous SRS and CARS imaging capability, while also offering SHG and two-photon excited fluorescence imaging by minor changes to the laser setup. This turns it into a cost-efficient, environmentally insensitive, stable, compact and versatile tool, which can be proposed for clinical applications. This domain has additional relevance due to ongoing research on endoscopic applications using CRS-contrast, which would be a valuable match to the fiber laser source [Saar et al. 2011, Lukic et al. 2017].

In the following thesis a short theoretical introduction is provided on coherent Raman scattering as well as on the microscopic image formation with emphasis on resolution enhancement by structured illumination techniques (section 2). Following this part the CRS setup is described introducing novel aspects, which separates it from other recent realizations [3]. The results obtained with this system are presented in section 4: first, CARS and SHG imaging contrast is utilized for the investigation of the osteogenic differentiation of human stem cells, which offer promising treatment for the cure of injured bone structures [4.1]. The results achieved by CARS and SHG imaging are combined with spontaneous Raman spectroscopy and then compared to established methods in this field, such as fluorescence staining and rt-PCR (reverse transcription polymerase chain reaction). Then the successful application of volumetric CARS imaging to the oleaginous microalgae Monoraphidium neglectum is demonstrated in vivo for investigating the lipid content under nitrogen starvation conditions [4.2]. Here, an intensity modulation approach is utilized in combination with lock-in amplification for the suppression of undesired two-photon excited fluorescence of the chloroplast, which enabled a quantitative determination of the lipid content and, finally, the optimal harvesting time point.

Furthermore, the hyperspectral CARS imaging modality is introduced in [4.3]: the hyperspectral control software is compared in its performance to the conventional non-controlled mode and the influence of the optical parametric oscillator as a tunable wavelength source is determined concerning its influence on the laser pulse length in the experiments [4.3.4]. The hyperspectral image acquisition is contrasted to static CARS imaging for determining the fitness parameter of the soil worm Caenorhabditis elegans under exposition to a copper treatment [4.4.1] as well as for visualizing the differentiation process of primary human alveolar epithelial cells [4.4.4].
Subsequently, the spatial resolution enhancement based on structured illumination applied to scanning-based microscopy setups is explored in two-photon excitation mode of several fluorescent samples as well as on polystyrene bead samples with CARS contrast. Two reconstruction approaches are compared to each other for producing the final image, whose resolution and contrast is strongly enhanced.

After that the setup of the novel two-color all-fiber-based laser source is described and the source is characterized in its optical parameters with strong emphasis on the noise performance indicating superior performance as a source in SRS imaging (4.6). Then data of a range of relevant biomedical samples is presented in simultaneous acquired SRS and CARS contrast demonstrating the performance of the setup. The SHG and two-photon imaging capability is showcased by providing additional data sets proving the laser source to be a versatile multimodal imaging tool.

Finally, the thesis is summarized and future steps are indicated in section 5.
2. Theoretical background

In the following thesis coherent Raman scattering (CRS) is utilized for imaging purposes on biomedical samples. This theoretical introduction describes the two mechanisms of spontaneous Raman scattering and the coherent counterpart in sections (2.1) and (2.2). Then the two coherent effects of coherent anti-Stokes Raman scattering (CARS) (section 2.2.2) and stimulated Raman scattering (SRS) (section 2.2.4) are introduced due to their physical dependencies on the occurring light fields as well as to the different detection modes. Finally, the image formation in a microscope as well as the concept of resolution is introduced (2.3.1), while the aspect of resolution enhancement by structured illumination is discussed in (2.3).

2.1. Introduction to Raman scattering

Single atoms form a molecular bond reaching a lower energetic state due to the overlap of their specific electronic orbitals in the bond state. A molecular bond exhibits three energetic regimes corresponding to three different mechanisms in which energy can be deposited: the excitation of electronic states has the highest amount of energy deposition, which can be performed with excitation wavelengths in the range of ultraviolet to near infra-red light (corresponding to several $\text{eV}$). Also vibrational states can be excited along the binding axis analogue to the harmonic oscillator, while the binding strength of the bond determines the potential of the oscillation. Typical energies are $1\text{eV}$ to $\text{meV}$. The rotation of the binding components along the bond axis exhibits the lowest energy regime ranging in several $\text{meV}$ [Haken and Wolf [2006, pp. 180]].

Electromagnetic fields are able to affect the electrons of a molecule leading to polarization effects. Here, the charges of the molecular bond are separated from each other, which is possible due to the nearly instantaneous reaction of the electrons to an applied perturbation such as a light field. The nuclei are not reacting on the same time scale but several magnitudes slower. By applying the Born-Oppenheimer assumption [Born and Oppenheimer [1927]] stating that the electrons are moving in an unaltered potential of the nuclei, the reaction of the electrons on the perturbation can be described. A change of the potential due to an oscillation of the nuclei leads to an instantaneous following of the electrons [Haken and Wolf [2006, pp. 209]]. The polarization induced by an incident light field leads to a non-equilibrium position of the electrons in the potential of the nuclei. This induces a dipole moment $\mu(t)$, which can be stated with only one electron in one dimension as (derivation following Cheng and Xie [2013, pp. 5]):
\[ \mu(t) = -e \cdot r(t) \]  \hspace{1cm} (1)

\( e \)  charge of the electron
\( r(t) \)  displacement from the equilibrium position

The magnitude of the displacement \( r(t) \) is defined by the coupling strength of electrons and nuclei. By summing up all \( N \) electric dipoles of one molecule or unit volume the macroscopic induced polarization \( P(t) \) is found to be:

\[ P(t) = N \cdot \mu(t) \]  \hspace{1cm} (2)

Raman scattering describes the inelastic scattering of incident photons at a molecule, observed for the first time by C. V. Raman in 1928 [Raman and Krishnan [1928]]. The participating light field couples to a molecular bond and leads to an energy transfer. The visible or near infra-red light field \( E(t) \) drives the oscillation of the electrons, which compose the molecular bond. The oscillatory motion contains information of the motion of the nuclei due to the adiabatic electronic potential, which is determined by the nuclear coordinates. Therefore, the motion of the involved nuclei will affect the motion of the electrons through the binding of both by the nuclear force. Describing the influence of the nuclear motion on the electronic motion in a quantitative manner far of an electronic resonance of the system, the electric dipole moment \( \mu(t) \) is weighted with the polarizability \( \alpha(t) \) [following Cheng and Xie [2013, pp. 10]]:

\[ \mu(t) = \alpha(t) \cdot E(t) \]  \hspace{1cm} (3)

By assuming no nuclear modes or nonlinearities, the polarizability can be approximated as a constant \( \alpha_0 \).

In the presence of nuclear modes the polarizability can be described in terms of the nuclear coordinate \( Q \) and can be approximated by a Taylor series [Placzek [1934]]:

\[ \alpha(t) = \alpha_0 + \left( \frac{\delta \alpha}{\delta Q} \right)_0 \cdot Q(t) + ... \]  \hspace{1cm} (4)

The coupling strength of the nuclear and electronic coordinates is considered to be the first-order correction to the polarizability having the magnitude of \( \delta \alpha/\delta Q \). Assuming the nuclear motion to behave as a classical harmonic oscillator leads to:
\[ Q(t) = 2Q_0 \cdot \cos(\omega_\nu t + \phi) = Q_0 [e^{i\omega_\nu t + i\phi} + e^{-i\omega_\nu t - i\phi}] \] (5)

- \(Q_0\) amplitude of the nuclear motion
- \(\omega_\nu\) nuclear resonance frequency
- \(\phi\) phase of the nuclear mode vibration

The incoming light field can be described as
\[ E(t) = A \cdot e^{-i\omega_1 t} + A^* \cdot e^{i\omega_1 t} \] leading to

the expression of the dipole moment as:
\[ \mu(t) = \alpha_0 A e^{-i\omega_1 t} + A \cdot \left( \frac{\delta \alpha}{\delta Q} \right)_0 Q_0 \left\{ e^{-i(\omega_1 - \omega_\nu) t + i\phi} + e^{-i(\omega_1 + \omega_\nu) t - i\phi} \right\} + c.c. \] (6)

The oscillation of the dipole moment occurs at several frequencies and leads to the radiation of photons, whose wavelengths is corresponding to the oscillation frequency of the dipole moment. The elastic scattering (also named as Rayleigh scattering [Strutt 1871, Rayleigh 1881, 1899]) of the incident photons is represented in eq. (6) by the first term, the dipole moment oscillates with the frequency of the initial light field.

The following term represents the inelastic scattering (the Raman shift). The so-called Stokes-shift is given by \(\omega_S = \omega_1 - \omega_\nu\), where the energy of the radiated photons is lower than the one of the incident photons. Thus, the emitted light is red-shifted in comparison to the incident wavelength. Consequently, the anti-Stokes shift occurs as \(\omega_{aS} = \omega_1 + \omega_\nu\) resulting in blue-shifted, higher energy photons (see fig. 1).

The Raman term is directly proportional to \(\delta \alpha/\delta Q\), which states the change of the polarizability along the nuclear mode due to the incident light field. This change is significantly dependent on the symmetry of the nuclear mode in the molecule, which can be exploited for gathering of information about the molecular bond by Raman spectroscopy.
Incident photons (with the frequency $\omega_1$, left hand side) are scattered at a molecular bond formed by two atoms. Rayleigh scattering leads to radiated light, which is unaltered in its frequency ($\omega_1$). Here, the dipole moment of the chemical bond oscillates with the same frequency as the light field. The frequency of the Stokes-shifted light is decreased with $\omega_1 - \omega_\nu$, energy is transferred from the light field to the molecular bond. The anti-Stokes shift occurs in analogue way at the frequency of $\omega_1 + \omega_\nu$, leading to a blue-shifted radiated light. Adapted from Cheng and Xie [2013, p. 12].

The amplitude of the emitted Stokes-shifted field radiating along $r$ in the far-field is given by Cheng and Xie [2013, p. 12]:

$$E(\omega_S) = \frac{\omega_S^2}{4\pi\epsilon_0c^2} |\mu(\omega_S)| \frac{e^{ikr}}{r} \cdot \sin(\theta)$$  \hspace{1cm} (7)

$k$\hspace{1cm} wave vector of the radiated field
$c$\hspace{1cm} speed of light
$\theta$\hspace{1cm} angle relative to the dipole axis
$r$\hspace{1cm} distance from the dipole location to the observation point
$|\mu(\omega_S)|$\hspace{1cm} amplitude of the dipole oscillation at $\omega_S$

Using the energy flux along $r$ calculated as the time-averaged Poynting flux $S(\omega_S) = \frac{\omega}{2} |E(\omega_S)|$ the total intensity $I(\omega_S)$ emitted by a single dipole can be calculated by integrating over the unit sphere using eq. (6) as:

$$I(\omega_S) = \frac{\omega_S^4}{12\pi\epsilon_0c^3} Q_0^2 |A|^2 \left| \frac{\delta\alpha}{\delta Q} \right|^2$$  \hspace{1cm} (8)

The inelastically scattered light has a $\omega_S^4$ intensity dependency (eq. (8)). The initial intensity of the incoming light scales with $I_0 = |A|^2$ and the polarizability has the factor of $|\delta\alpha/\delta Q|^2$. These are major influences utilizing Raman spectroscopy to analyze the chemical properties of a sample of interest leading to the choice of integration times as well as the choice of the intensity and the wavelength of the excitation light field.
The frequency shift of the scattered light offers information about the chemical bond, independent of the initial frequency. Rayleigh scattering leads to coherently radiated light while Raman scattered light is incoherent due to the uncorrelated vibrations of the different molecules. Each molecule vibrates with its own independent phase $\phi$ [Cheng and Xie [2013, p. 13]]. The intensity of the emitted Raman photons is several magnitudes lower than that of the incident light field. In combination with the small shift of the Stokes and anti-Stokes lines a very narrow line width of the incident photons is required, which is generated by laser sources in combination with monochromators [Haken and Wolf [2006, pp. 239]]. The used spectrometer needs a high resolution in order to separate single Raman peaks from each other while the excitation wavelength is adapted to the detection range of the setup. Fig. 2 is showcasing a typical Raman spectrum of a methanol sample, while two characteristic peaks can be observed in the spectrum.

![Figure 2 – Raman spectrum of pure methanol](image)

This figure shows a normalized Raman spectrum obtained of pure methanol utilizing a cw-excitation wavelength of $785\text{nm}$ with a focal power of $20\text{mW}$. A $60\times 1, 2\text{NA}$ air objective lens focused the light into the sample. The spectrum is measured with an Acton SpectraPro 2300i spectrometer using a CCD camera for the detection (Andor, Newton 920-P-BR-DD). Two typical peaks are marked in the spectrum depicting two molecular resonances of the methanol: the symmetric and anti-symmetric $C - H$ stretching mode at $2830\text{cm}^{-1}$ and $2940\text{cm}^{-1}$, respectively.
2.2. Coherent Raman scattering

Utilizing a classical instead of the full quantum mechanical description of coherent Raman scattering (CRS) is effectual for imaging purposes. In general the spontaneous Raman scattering is a rare event exhibiting a low strength of the Stokes and anti-Stokes signals. The difference of CRS to spontaneous Raman scattering is the actively driven nuclear oscillation of a specific molecular bond. This leads to the coherent excitation of a larger number of the same molecular group and finally to an oscillation of these molecules with one phase. Applying the two step model leads to: first, the electronic states of a molecular bond are driven by two incoming light fields, which vary in their energy. The resulting oscillation of the molecular electron cloud induces an effective force exciting the vibrational states of the bond, which finally results in a spatially coherent modulation of the refractive index of the material. The interaction of the material with a third incident light field, which experiences the modulation, is the second step in the model.

Any photons scattered at the bonds are in phase producing a coherent signal. The finally detected signal is enhanced by several orders of magnitude compared to spontaneous Raman scattering due to the constructive and destructive interference of the signal. The Raman shift is observed according to the frequency of the oscillation. Sidebands occur in the spectrum of the third light field, which can be clearly separated from the initial frequency [Cheng and Xie [2013, p. 14]].

2.2.1. Driven Raman mode

Similar to 2.1 the vibrational motion in the molecule can be described for the coherent Raman process (in a classical way) as a damped harmonic oscillator. The resonance frequency of the system is described by \( \omega_\nu \) along the internuclear axis \( Q \). In contrast to spontaneous Raman scattering the molecular vibration is driven by two incident light fields \( E_1 \) and \( E_2 \), assumed to be a plane wave (following [Cheng and Xie [2013, pp. 14]]):

\[
E_j(t) = A_j \ e^{i \omega_j t} + A_j^* \ e^{-i \omega_j t} \quad j = 1, 2 \quad (9)
\]

The fields apply a force on the nuclei (assuming their frequencies \( \omega_1 \) and \( \omega_2 \) to be much higher than the resonance frequency \( \omega_\nu \)):

\[
F(t) = \left( \frac{\delta \alpha}{\delta Q} \right)_0 \left[ A_1 A_2^* \cdot e^{-i \Omega t} + c.c. \right] \quad (10)
\]

With \( \Omega = \omega_1 - \omega_2 \) being the difference frequency, which occurs using high intensities for the creation of non-linear effects in the electron motion. The nuclear displacement
can be described by the solution of the dampened harmonic oscillator:

\[
\frac{d^2 Q(t)}{dt^2} + 2\gamma \frac{dQ(t)}{dt} + \omega^2 Q(t) = \frac{F(t)}{m}
\]  

(11)

\(\gamma\) damping constant
\(m\) reduced mass of the nuclear oscillator
\(\omega_{\nu}\) resonance frequency of the harmonic nuclear mode

The nuclear displacement over time can be stated from eq. (11) as:

\[Q(t) = Q(\Omega) \cdot e^{-i\Omega t} + \text{c.c.}\]  

(12)

The amplitude at the frequency \(\Omega\) is found to be:

\[Q(\omega_{\nu}) = \frac{1}{m} \left( \frac{\delta\alpha}{\delta Q} \right)_0 \frac{A_1 A_2^*}{\omega_{\nu}^2 - \Omega^2 - 2i\Omega\gamma}\]  

(13)

Eq. (13) shows that the joint action of the incident fields drives the addressed nuclear motion at \(\Omega\). The amplitude of the vibrational mode is depending on the amplitudes of the incoming fields \(A_1\) and \(A_2\) as well as on the coupling strength of the nuclear motion to the polarizability along the nuclear mode due to the incident light field \(\left( \frac{\delta\alpha}{\delta Q} \right)_0\). It also scales on the difference of the driving frequency \(\Omega\) and the resonance frequency \(\omega_{\nu}\) of the oscillator. Matching both frequencies leads to the maximal amplitude (fig. 3).
In **a)** two incoming light fields with the frequency $\omega_1$ and $\omega_2$ are coherently driving the nuclear bond with the difference frequency $\Omega = \omega_1 - \omega_2$. **(b)** shows the interaction of the third light field with the driven molecular bond. Due to the presence of the molecular vibration the refractive index of the material is fluctuating leading to two sidebands, which are shifted in their frequencies compared to the fundamental $\omega_3$ with the frequency of the molecular vibration $\omega_{\nu}$. If $\Omega$ matches the resonance of the molecular bond $\omega_{\nu}$ the amplitudes of the sidebands are at their maximum. Adapted from [Cheng and Xie 2013, p. 15].

### 2.2.2. Coherent anti-Stokes Raman scattering

Coherent anti-Stokes Raman scattering (CARS) is a parametric four wave mixing process (FWM), which can be exploited for a label-free imaging contrast. An incoming pump photon $\omega_p$ excites a molecular bond to a higher virtual energy state. The incoming Stokes photon $\omega_S$ de-excites the bond coherently to a certain vibrational mode, matching the molecular vibration of the bond with the frequency $\omega_{\nu}$. It depends on the molecular bond and is specific to the investigated material (following [Cheng and Xie 2013, pp. 26]):

$$\omega_{\nu} = \omega_p - \omega_S$$

A second pump photon $\omega_p$ excites the driven molecular bond to a higher virtual energy state, comparing this energy to the initial one the following relation can be stated ($|E_{\text{virtual}1}| < |E_{\text{virtual}2}|$). The relaxation of this unstable state creates the coherent blue-shifted anti-Stokes signal $\omega_{aS}$, de-exciting the bond to the initial ground state (see fig. 4):

---

1. Meaning an unstable state, which lasts only a short period of time $t \sim \hbar / \delta E$.
2. In most implementations is has the same frequency as the first pump photon while originating from the same laser source.
\[ \omega_{aS} = (\omega_p - \omega_S) + \omega_p \] (15)

Due to the geometry of the participating scatterers the CARS signal occurs in forward and epi-direction. Comparing CARS to spontaneous Raman scattering the signal strength is many orders of magnitude higher, still depending on the concentration of the addressed molecular group.

\[ P = P^{(1)} + P^{(3)} = \epsilon_0 \chi^{(1)} E + \epsilon_0 \chi^{(3)} E \cdot E \cdot E \] (16)

The quantitative description of the signal generation in the CARS process on the macroscopic scale is possible using the Maxwell equations, which describes the spatial properties of the electric and magnetic fields. The wave equation describes the generation and propagation of the participating waves. It predicts the amplitude and phase of the non-linear electric field \( E \), having the non-linear polarization \( P \) as the source term. Using eq. (16) it is found to be:

\[ \nabla^2 E - \frac{n^2}{c^2} \frac{\partial^2}{\partial t^2} E = \frac{1}{\epsilon_0 c^2} \frac{\partial^2}{\partial t^2} P^{(3)} \] (17)

By the plane wave assumption with a constant amplitude and phase propagating in
forward direction the incoming fields of the FWM process can be described as [Cheng and Xie 2013, p. 50]:

$$E_j(z, t) = \frac{1}{2} \left( A_j(z) e^{i(k_j z - \omega_j t)} + A_j^*(z) e^{-i(k_j z - \omega_j t)} \right) \hat{x} \quad j = 1, 2, 3 \quad (18)$$

\(\hat{x}\) polarization direction of the plane waves
\(\omega_j\) angular frequency
\(k_j = \omega_j n_j / c\) scalar wave vector
\(A_j\) amplitude of the incident wave
\(n_j\) refractive index

By introducing the non-linear susceptibility \(\chi^{(3)}(\Omega)\) with \(\Omega = \omega_1 - \omega_2\) it is possible to state the third-order polarization of the non-linear Raman active material \(P^{(3)}\):

$$\chi^{(3)}_e = \frac{G_\nu}{\omega_\nu - \Omega - i \gamma_\nu} \quad (19)$$

$$P^{(3)}(z, t) = \frac{1}{8} \epsilon_0 \chi^{(3)}_e(\Omega) A_1(z) A_2^*(z) A_3(z) e^{i[k_1 z - \omega_4 t]} \hat{x} + c.c. \quad (20)$$

\(G_\nu\) spectral amplitude
\(\gamma_\nu\) damping constant related to the dephasing time of the vibrational level at frequency \(\omega_\nu\)

The induced polarization is the source of the coherent radiation, which oscillates with the angular frequency \(\omega_4 = \omega_1 - \omega_2 + \omega_3\). Substituting the \(x\)-component of the forward propagating plane wave \(E_4(z, t)\) with the wave vector \(k_4\) along the \(\hat{z}\) and \(-\hat{z}\) directions into the wave equation [Druet, S. A. J. and Taran, J. P. E. 1981], the slowly varying envelope approximation (SVEA) can be performed, which assumes that the change of the amplitude \(A_4\) of the wave is incrementally small on a length scale in the regime of the optical wavelength [Potma et al. 2000], leading to:

$$E_4(z, t) = \frac{1}{2} A_4(z) e^{i(k_4 z - \omega_4 t)} \hat{x} + c.c. \quad (21)$$

Starting with the left hand side of eq. (17) the following term is found:

$$\nabla^2 E_4(z, t) - \frac{n_4^2}{c^2} \frac{\partial^2}{\partial t^2} E_4(z, t) = \frac{1}{2} \left[ \left( \frac{\partial^2}{\partial t^2} + 2 i k_4 \frac{\partial}{\partial z} \right) A_4(z) \right] e^{i(k_4 z - \omega_4 t)} + c.c. \approx i k_4 \frac{\partial A_4(z)}{\partial z} e^{i(k_4 z - \omega_4 t)} + c.c. \quad (22)$$
Inserting the non-linear polarization $P^{(3)}$ (eq. [6]) into the wave equation for the field with the amplitude $A_4$ leads to:

$$\frac{\partial A_4(z)}{\partial z} = i \frac{\omega_4}{8n_4c} \chi^{(3)}_e(\Omega) A_1(z) A_2^*(z) A_3(z) e^{i\Delta k z}$$

(23)

$\Delta k = k_1 - k_2 + k_3 - k_4$ describes the wave vector mismatch along the $z$ axis between the effective wave vector of the polarization field $(k_1 - k_2 + k_3)$ and the wave vector of the radiating field ($k_4$). Assuming that the incident field amplitudes are mostly unaltered during the FWM process the amplitudes $A_j$ are considered as constants during the signal generation. The integration of the differential equation (23) in the range of $z = 0$ to $z = L$, covering the length of the material, leads to the signal amplitude $A_4$ [Druet, S. A. J. and Taran, J. P. E. [1981]]:

$$A_4(L) = i \frac{\omega_4}{8n_4c} \chi^{(3)}_e(\Omega) A_1 A_2^* A_3 L \text{sinc} \left( \frac{\Delta k L}{2} \right) e^{i\Delta k L/2}$$

(24)

With $\text{sinc}(x) = \sin(x)/x$. Here, the strong influence on the amplitude of the coherent radiation by the wave vector mismatch becomes apparent. The maximal amplitude is observed when no phase mismatch between the induced non-linear polarization and the radiating field is present. In media this can not be the case, because of the inevitable dispersion of the medium, which always introduces a wave vector mismatch. Having a non-zero $\Delta k$ the radiating field and the induced polarization are dephasing and, therefore, limiting the effective coherence to a length scale of $L_c = 2\pi/|\Delta k|$. This length determines an effective thickness in the sample material, where the signal amplitude can be generated.

Transferring this description of all CRS-signals to the CARS process the third-order non-linear polarization is found to be $\chi^{(3)}_e(\Omega) = \chi^{(3)}(\omega_{aS}; \omega_p, -\omega_S, \omega_p)$. The pump photons have the electric field of $E_1 = E_3$ at the frequency $\omega_p$ and the Stokes photons $E_2$ at $\omega_S$ ($\omega_S < \omega_p$). In the dual-color CARS process the pump beam participates two times the Stokes beam one time, generating the field $E_{aS}$, which is oscillating with the anti-Stokes frequency $\omega_{aS} = 2\omega_p - \omega_S$. The scalar wave vector of the field is $k_{aS} = 2k_p - k_S$ (fig. 3).
On the left the wave vector diagram is depicting the forward propagating CARS process with the generated wave vector of $k_{aS}$. On the right the epi-direction is illustrated. The pump beam wave vector ($k_p$) participates two times in the FWM process generating together with the Stokes beam ($k_s$) the. Adapted form [Evans and Xie [2008]].

Utilizing eq. (24) the intensity of the detected CARS signal can be found by deriving the squared amplitude [Cheng and Xie [2013, pp. 53]]:

$$I(\omega_{aS}) = |A_{aS}(L)|^2 \propto |\chi_e^{(3)}(\Omega)|^2 I_p^2 I_s L^2 \text{sinc}^2\left(\frac{\Delta k L}{2}\right)$$

Eq. (25) shows the dependencies of the CARS signal on the parameters set in the experiment. The CARS intensity scales quadratic to the pump beam intensity and linear to the Stokes beam intensity. During the imaging experiment the pump beam is set with a higher intensity than the Stokes beam for maximizing the finally detected CARS signal.

The wave vector mismatch $\Delta k$ is weighted with the squared sinc-function. In the experimental realization this means that the interacting beams have to be aligned very exactly to minimize a possible mismatch. The divergence as well as the beam size have to be matched to generate a perfect overlap in the focal plane. This process is also affected by the focusing objective lens, which itself introduces chromatic aberrations. All optical lens failures have to be compensated as good as possible by introducing additional beam divergence. Any mismatch introduced by the dispersion by the medium of the sample is considered as small compared to the coherence length of the signals.
in the microscope. The squared amplitude of the non-linear susceptibility $\chi_e^{(3)}$ itself is proportional to the density of the molecules, which participate in the signal generation process. Hence, the CARS signal depends quadratically to the number of molecules in the interacting focal volume and high concentrations of a molecule in the sample will induce high CARS signal strength.

The non-linear susceptibility contains a vibrationally resonant $\chi_r^{(3)}$ and a non-resonant part $\chi_{nr}^{(3)}$ and can be written as:

$$\chi_e^{(3)} = \chi_r^{(3)} + \chi_{nr}^{(3)}$$  \hspace{1cm} (26)

Now the squared susceptibility can be found to be (following [Evans and Xie [2008]]):

$$\left|\chi_e^{(3)} (\Omega)\right|^2 = \left|\chi_r^{(3)} (\Omega) + \chi_{nr}^{(3)}\right|^2 = \left|\chi_r^{(3)} (\Omega)\right|^2 + \left|\chi_{nr}^{(3)}\right|^2 + 2\chi_{nr}^{(3)} Re[\chi_r^{(3)} (\Omega)]$$  \hspace{1cm} (27)

Here, an intrinsic drawback of the CARS signal as contrast mechanism becomes apparent: the first term is resonant to the addressed molecular vibration and delivers the desired chemically sensitive signal. The second term represents the non-resonant background contribution, which obscures the desired resonant signals especially in the case of lower molecular concentrations. The last term refers to the mixing part of resonant and non-resonant contribution. The sensitivity of CARS is limited by the intensity of this non-resonant background.

The reason for the presence of a non-resonant background can be found by considering all possible responses of the system taking into consideration all the optical frequencies described by the third-order susceptibility. Because of the parametric nature of CARS in the $\chi^{(3)}$ process 24 quantum pathways are possible, while combining the incoming light fields in order to detect the CARS signal. 17 of this pathways do not exhibit a Raman coherence and are not fulfilling the Raman resonance condition. They are summed up to $\chi_{nr}$. A semi-classical explanation of this behavior is the possibility of undergoing some system pathways, which contribute to the dipole radiation at $\omega_{aS}$, but having no propagators at the resonance frequency of the molecular bond. In addition, there are so-called electronic resonances, meaning quantum pathways, which are possible if $\omega_2$ or $\omega_{aS}$ is resonant to electronic states of the material or, in case of a populated vibrational state, if $\omega_2$ has a resonance to the electronic states. In the case that no electronic resonance is present the contribution to the CARS signal of the non-resonant pathways is less than the vibrationally resonant part. In case of an electronic resonance the non-resonant signals are significantly higher than the resonant counterpart [Cheng and Xie [2013, pp. 27]].

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Referring to the driven Raman mode.
The non-linear susceptibility is strongly dependent on the difference frequency $\Omega = \omega_1 - \omega_2$ near the molecular resonance. A simulation for the spectral dependence of each part and the overall CARS signal can be found in fig. 6. The detected CARS signal is generated by the interference of all three parts. The mixing term of the resonant and the non-resonant part of the non-linear susceptibility stated in eq. (27) has a dispersive line shape, which can be observed in the CARS signal as a redshift. When comparing CARS spectra to spontaneous Raman spectra this redshift has to be considered, especially identifying the optimal combination of excitation wavelengths in the CARS process [Evans and Xie [2008]].

Figure 6 – Spectral dependence of $\chi^{(3)}$

a) shows the three components of the CARS signal referring to the non-linear susceptibility, while the molecular resonance $\Omega_R = \omega_1 - \omega_2$ is scanned. The constant non-resonant background $|\chi^{(3)}_{nr}|^2$ is depicted by a dotted line, the purely resonant term $|\chi^{(3)}_r|^2$ in solid line and the mixing term of both $2\chi^{(3)}_{nr} \text{Re}\{\chi^{(3)}_r\}$ in dashed line, respectively. b) shows the resulting total CARS signal, whose peak is red-shifted and which has a dispersive line shape arising from $2\chi^{(3)}_{nr} \text{Re}\{\chi^{(3)}_r\}$. The dotted line represents the non-resonant background, the solid line the combination of all signal components from (a). Adapted from [Evans and Xie [2008]].

2.2.3. Detection modes

When the CARS process is exploited as contrast mechanism for imaging two possible detection pathways can be used: the forward and epi-detection. In forward detection the CARS signal is generated in the focal plane and then propagates through the sample. Finally, it is collected by a condenser lens, filtered and detected. The epi-detection describes all back scattered CARS signals, which are collected by the same objective lens, which focused the excitation beams into the sample. The detection is performed similar to the forward direction.

One major difference of both detection modes is the strength and the origin of the signal, referring to the participating sample structures in the CARS process. This re-
results in differences of information content of the investigated sample. As mentioned before the wave vector mismatch induced by the dispersion of the media is small for the forward propagating CARS signal and has no significant effect on the detected signal [Potma et al. [2000], Cheng and Xie [2013, p. 54]]. In epi-CARS a significant wave vector mismatch is introduced, which can be approximated as $|\Delta k| = 2|k_{as}|$ leading to a coherence length of $L_c \approx 0.24 \mu m$ in water. This mismatch affects the signal generation when the investigated structures are smaller than the coherence length in their axial dimensions. If the structures are smaller than the pump wavelength the wave vector mismatch condition is satisfied in both propagation directions. With an increasing thickness $D$ of the structure the forward CARS signal scales to $\sim D^2$ until it becomes saturated, when $D$ exceeds the longitudinal dimension of the focal excitation volume. Depending on the size and the shape of the scatterers as well as on the optical parameters of the setup, forward CARS overwhelms the epi-signal due to the constructive interference in the forward direction (and the destructive interference in the epi-direction respectively) (see fig. 7).

The epi-CARS signal oscillates as a function of the sample thickness with a periodicity of $\lambda_{as} \cdot 2n \quad n = 1, 2, 3, ...$ and has an intensity which is proportional to [Volkmer et al. [2001]]:

$$I_{as}^{epi} \propto \lambda_{as}^2 \cdot sin^2\left(\frac{2\pi n D}{\lambda_{as}}\right) \quad n = 1, 2, 3...$$  (28)

---

\[4\] Using the planar wave approximation.
The graph shows a simulation of the forward and epi-CARS intensities for tightly focused excitation beams as a function of the diameter $D$ of a spherical scatterer. The following parameters are used: pump and Stokes wavelengths are at $\omega_p = 800\,\text{nm}$ and $\omega_S = 917\,\text{nm}$, respectively. The index of refraction of the medium is 1.52 and the NA of the lens is 1.4. The epi-CARS signal oscillates, the forward signal saturates, when the focal volume is filled with material. Note, that the forward CARS signal is rescaled by a factor of 0.05. From [Volkmer et al. 2001].

When the focal excitation volume is small compared to the thickness of the sample linear scattering events of the forward CARS signal can occur, especially in tissues. Then the collected signal in epi-direction is significantly enhanced due to the loss in the forward direction. Monte-Carlo simulations based on typical tissue scattering parameters in skin samples indicate a loss of the forward CARS signal of $40 - 45\%$, enhancing the epi-CARS signal respectively [Saar et al. 2010]. Utilizing a high NA objective lens 10\% of the entire epi-CARS signal can be collected due to the limited cone angle of the lens. Multiple scattered, diffusive light is in most cases beyond the limit of the objective (see fig. 8).

Assuming structures in the $\sim \lambda_p/2$ regime a variation of $\chi_1^{(3)}$ is induced, which leads to an intrinsic backward radiation added to the overall CARS signal in epi-direction [Cheng and Xie 2013 pp. 70]].
Figure 8 – Back-scattering of forward directed CARS signals in tissues

In tissues a portion of the forward-propagating CARS signal is back-scattered and contributes to the epi-CARS signal (adapted from [Cheng and Xie [2013, p. 71]]).

The geometric shape, size and position relative to the focal volume of the single scatterers play also a role for the ratio of forward and epi-CARS signals due to non-linear interference effects. The spectral properties of $\chi^{(3)}_e$ introduce a phase shift of $\pi$ to the resonant signal, while probing the vibrational resonance from lower wave numbers to higher ones [Cheng et al. [2002]].

In the case of scatterers, which are small enough to be considered as single dipole emitters, an axial phase shift along the excitation beam direction introduces a wave vector mismatch, known as the Gouy phase shift. This effect is responsible for destructive and constructive interference detected in the far-field due to the position of the scatterers in the focal volume. If the focus is shifted through a sample an object located in the lower half of the focal volume will get a phase shift of $\pi$ compared to the case, when it is in the upper half. This can lead to artifacts in the CARS images due to the interference effects with the non-resonant background, which has to be considered during the data interpretation [Popov et al. 2011, Cheng and Xie 2013, pp. 72]].
2.2.4. Stimulated Raman scattering

Stimulated Raman scattering (SRS) belongs to the group of coherent Raman scattering processes [Woodbury E. J. and Ng W. K. 1962] and is used as contrast mechanism in biomedical imaging [Ploetz et al. 2007, Freudiger et al. 2008, Ozeki et al. 2009]. The addressed molecular vibration is driven similar to the CARS-technique, whereas several differences in the signal detection as well as an additional intensity modulation of the Stokes beam are necessary. SRS is a dissipative energy transfer process in which energy is transferred between two incident light fields, the pump beam, with the frequency $\omega_p$, and the Stokes beam with $\omega_S$, respectively ($\omega_S < \omega_p$). Both light fields drive the molecular vibration at the resonance frequency $\Omega$ with the difference frequency $\Delta \omega = \omega_p - \omega_S$. Thereby, the molecular transition rate is enhanced because of the stimulated excitation of molecular transitions [Cheng and Xie 2013, pp. 100]. The electronic states of the molecules are excited into a virtual state by the pump beam and de-excited by the Stokes field, transferring the molecular population from the ground state into a vibrationally excited state (see fig. 9). To satisfy the law of energy conservation a pump photon is absorbed by the molecule and a low-energetic Stokes photon is generated. This transfer of the number of photons in the two beams can be detected in the intensities of the pump beam $I_p$ and the Stokes beam $I_S$ after the interaction with the sample. The resulting loss in intensity SRL (stimulated Raman loss) $\Delta I_p$ and the gain SRG (stimulated Raman gain) $\Delta I_S$ are scaling to [Cheng and Xie 2013, p. 100]:

$$\Delta I_S \propto N \cdot \sigma_{\text{Raman}} \cdot I_p \cdot I_S$$

$$\Delta I_p \propto -N \cdot \sigma_{\text{Raman}} \cdot I_p \cdot I_S$$

$N$ number of molecules in the probe volume
$\sigma_{\text{Raman}}$ molecular Raman scattering cross section

By detecting the intensity of the pump or the Stokes beam with a suitable detector such as a large area fast silicon photo diode it is possible to measure the SRG of SRL as a function of the focus position. Because of the proportionality of the number of addressed molecules to the intensity of the detected beam $\Delta I \propto N$, thereby to the concentration of the target species, the obtained signal holds the spatial information of the chemical properties of the investigated sample. While the focus is raster scanned over the sample a two dimensional map of the chemical properties of the sample is acquired.
Figure 9 – Principle of stimulated Raman scattering

(a) shows the energy diagram of SRS. Here, the pump and Stokes beam coherently excite the molecular bond, transferring the molecules into the first excited vibrational state by passing through a virtual state. The selection of the addressed molecular vibration is achieved by tuning the pump and Stokes wavelengths to the molecular resonance frequency $\omega$, with $\Delta \omega = \omega_p - \omega_S$. (b) illustrates the energy transfer process in SRS: the gain and loss of the two participating light fields is obtained. Due to the conservation of energy the number of photons is increased after the interaction with the sample in the Stokes field, the number of pump photons decreases, respectively. Adapted from [Cheng and Xie 2013, p. 101].

With the plane wave approximation described in chapter 2.2.2 (especially eq. (24)) the amplitude for the forward propagating stimulated Raman wave is found to be [Cheng and Xie 2013, pp. 54]:

$$A_{2}^{stim} = i \frac{\omega^2}{8n_2c} \{\chi^{(3)}(\Omega)\} \ast A_1^* A_2 A_1 L$$  \hspace{1cm} (31)

Due to the coherent nature of the stimulated Raman wave and the incident waves these contributions interfere at the detector (in the case of forward detection). The detected SRG intensity with frequency $\omega_S$ can be written as:

$$I(\omega_S) = \left| A_2(L) + A_2^{stim}(L) \right|^2$$

$$= I_S + \left| \chi^{(3)} \right|^2 \cdot I_p \cdot I_S \cdot L^2 + 2 Im \{\chi^{(3)}(\Omega)\} \cdot I_p \cdot I_S \cdot L$$  \hspace{1cm} (33)

The first term of eq. (33) refers to the intensity of the incident Stokes beam, assuming that it is unattained during the non-linear process. The second term is the intrinsic SRS term, which is depending on the material properties as well as on the incident light fields similar to CARS. The last term is the interference term of the incident Stokes field and the stimulated Raman field at the detector. The induced amplitude $A_2^{stim}$
has an intrinsic $\pi/2$ phase shift relative to the incident $A_2$ driving field, which follows from the plane wave approximation. If the sample contains no addressed molecular content $\chi^{(3)}_e$ is purely real and the interference term disappears. Close to a resonance \{\chi^{(3)}_e\} \neq 0 the interference term is prominent. For the investigation of samples the interference term has to be isolated from the remaining terms of eq. (33). Assuming that the amplitude of the incident Stokes beam is bigger by several orders of magnitude compared to the stimulated Raman field ($A_2 \gg A_2^{stim}$) the forward SRG and SRL can be stated as:

$$\Delta I(\omega_S) \propto 2Im\left\{\chi^{(3)}_e(\Omega)\right\} \cdot I_p \cdot I_S \cdot L$$

(34)

$$\Delta I(\omega_p) \propto -2Im\left\{\chi^{(3)}_e(\Omega)\right\} \cdot I_p \cdot I_S \cdot L$$

(35)

In contrast to CARS the detected field in SRS (either SRG or SRL) is the same as the corresponding incident field, ensuring the phase matching condition automatically [Boyd [2003, pp. 451]]. The intensity gain or loss of the transmitted excitation beam is utilized as the measurand to gather information of the sample. In the parametric CARS process the signal is detected at a new optical frequency. Resulting of this difference the detected signal $\Delta I$ has to be separated from the incident beam during the SRS experiment. The rate of change in amplitude compared to the incident beam intensity ($\Delta I/I$) is in the range of $< 10^{-4}$ in both SRL and SRG [Freudiger et al. [2008]]. To separate the desired signal $\Delta I$ from the overall detected signal, a modulation of one of the incident beams with the frequency $f$ is performed to detect the signal achieving the necessary sensitivity. Avoiding the laser noise, which is scaling to $1/f$, the modulation frequency is set to be in the range of $1MHz$ to $20MHz$ utilizing electro- or acousto-optic modulation devices. Higher frequencies enable shorter integration times for rapid live cell imaging [Freudiger et al. [2008], Cheng and Xie [2013, pp. 103]].

The desired SRS signal $\Delta I$ is modulated with the known frequency $f$ and can be extracted from the larger laser intensity (e.g. $I_p$), which has its characteristic arbitrary fluctuations resulting from the laser source. To perform this operation a lock-in amplifier is used as a very narrow bandpass filter at the frequency $f$ to extract the small SRS signal from the overall signal. The modulated beam has to be the complementary beam, which is filtered out behind the sample and is not detected. In case of SRG the pump beam has to be modulated, in SRL the Stokes beam respectively.

Eq. (34) and (35) state that the detected signal is depending only on the imaginary part of $\chi^{(3)}_e$ meaning that in contrast to CARS there is no non-resonant background present. Here, a higher molecular sensitivity compared to CARS can be observed (for
example 50µM investigating a retinol solution using 1s integration time and 40mW on each beam in the focus [Cheng and Xie [2013, pp. 108]].

The interpretation of data obtained by SRS for determining the molecular concentration in a sample is easier compared to CARS due to the linear concentration dependence in SRS. Spectra obtained with SRS are identical to spontaneous Raman spectra because of the absence of the dispersive line shape, which is found during hyperspectral scanning in the distribution of the CARS signal intensity. This enables a direct and easy comparison of SRS to spontaneous Raman spectra.

Using SRS for imaging the signal is the convolution of the object with the point spread function of the optical system. SRS images can be interpreted straightforward while CARS images have to be checked for coherent image artifacts, especially when images contain small structures compared to the pump and Stokes wavelengths [Cheng and Xie [2013, pp. 100]].
2.3. **Concept of structured illumination for resolution enhancement**

The main goal in the development of optical microscopy was to resolve structures and gain information of unknown samples, whose amount of details could not be revealed with state of the art setups. For comparing the capabilities of different approaches and systems the term resolution was introduced. The resolution is the ability of an optical imaging system to resolve two point-like objects in close proximity. Classic examples for a definition of resolution were given by Abbe ([Abbe 1873](https://www.jstor.org/stable/273870), [Hecht 2002](https://www.jstor.org/stable/273870)), Rayleigh ([Rayleigh 1879](https://www.jstor.org/stable/273870)) and C. M. Sparrow ([Sparrow 1916](https://www.jstor.org/stable/273870)), stating that the resolution \( d \) is limited by the diffraction limit determined by the wavelength \( \lambda \) of the radiation used in the experiment and the utilized optics (in the case of Abbe the following equation was stated having \( n : \text{index of refraction} \) and \( \alpha : \text{the half of the acceptance angle} \)).

\[
d = \frac{\lambda}{2n \cdot \sin \alpha}
\]

(36)

In recent efforts the diffraction limit could be circumvented by various so-called “super-resolution” techniques using fluorescent dyes to label specific parts of the sample. Here, the **stimulated emission depletion (STED) microscopy** (theoretically predicted by [Hell and Wichmann 1994](https://www.jstor.org/stable/273870) and practically demonstrated by [Donnert et al. 2006](https://www.jstor.org/stable/273870)), the **photo-activated localization microscopy (PALM)** (theoretically predicted by [Betzig 1995](https://www.jstor.org/stable/273870), demonstrated by [Betzig et al. 2006](https://www.jstor.org/stable/273870) and [Hess et al. 2006](https://www.jstor.org/stable/273870)), the **stochastic optical reconstruction microscopy (STORM)** ([Rust et al. 2006](https://www.jstor.org/stable/273870) and direct STORM [Heilemann et al. 2005](https://www.jstor.org/stable/273870)) as well as the **structured illumination microscopy (SIM)** (shown independently by [Heintzmann and Cremer 1999](https://www.jstor.org/stable/273870), [Frohm et al. 2000](https://www.jstor.org/stable/273870) and [Gustafsson, M. G. L. 2000](https://www.jstor.org/stable/273870)) are the main state of the art super-resolution techniques applied to biological samples. Several extensions and modifications of the mentioned techniques are scope of the ongoing research for the adaption to sample specific problems or higher resolution enhancements [Sahl et al. 2017](https://www.jstor.org/stable/273870).

For these super-resolution microscopy techniques the classical definition of resolution, e.g. by Abbe, can not be applied straight forward. Specifying a numerical value for the resolution at a selected area in an image can not represent the overall obtained resolution of a technique as well as it may not be comparable to other techniques due to the selection of the sample or the utilized fluorescent dyes. A more convenient method is valuing the resolution of an image by estimating supported frequency bandwidth, taking into consideration that the spatial resolution is inversely related to the spatial frequency bandwidth [Nieuwenhuizen et al. 2013](https://www.jstor.org/stable/273870) and following [Mertz 2010](https://www.jstor.org/stable/273870, pp. 371)].
In the following section a brief description of the image formation in a microscope is provided, highlighting the aspects to circumvent the diffraction limit \([2.3.1]\). Structured illumination for resolution enhancement with linear dependence on the excitation power is discussed in \([2.3.2]\). Also the non-linear structured illumination approach is introduced illustrating the resulting effects in the Fourier transformed frequency space \([2.3.3]\). This technique is especially important for this thesis due to the non-linear properties of the utilized striped illumination in CARS and two-photon excited fluorescence microscopy presented in the results part of the thesis.

### 2.3.1. Microscopic image formation

A microscope is an optical setup with a magnification \(M\) to image a sample of interest (e.g. a point-shaped object), whose point size can not be resolved by using solely a classic detector (e.g. the human eye or a camera). The microscope performs an optical imaging process which results in the magnification of the point size allowing the detector to resolve the structure (fig. 10).

This process can be described in the frequency domain by performing a Fourier transformation. The object has frequency components \(I_0(X,Y)\) which will be transferred via the microscope to the detector, which itself has a limited frequency support. The whole imaging process has an overall limited range of supported frequencies, which are summed up with the modulation transfer function (MTF), which defines the ratio of detected image modulation to the object modulation over the whole frequency band. The cut-off-frequency describes the highest frequency which is supported by the optical system and is a useful value for estimating the resolution in modern microscopy.

The optical transfer function (OTF) \(T(k_X, k_Y)\) describes how the frequency components of the object are transformed by the optical system to the final image. It is a spatial frequency-dependent, complex quantity having the MTF as modulus and whose phase is described by the phase transfer function (PTF) \([\text{Hecht}] [2002\text{, pp. 552}]\).

The most applied transfer function is finally the point spread function (PSF) \(H(x, y)\) determining how a point-shaped object is imaged by the microscope. The point size will be broadened due to the imperfectness of the optical components used in the microscope. Mathematically the object will be convolved with the response of the microscope resulting in the detected image \(I_i\), given as [derivation follows \([\text{Hecht}] [2002\text{, pp. 554}]\)]:

\[
I_i(Y, X) = I_0(x, y) \otimes H(x, y)
\]  

\((37)\)

The convolution theorem states that the frequency spectrum of the image irradiance distribution is the product of the frequency spectrum of the object irradiance distribu-
tion and the Fourier transform of the spread function (see fig. [11]). Transforming the image (eq. (37)) into the spatial frequency domain by a Fourier transform using the convolution theorem leads to:

\[ \mathcal{F}\{I_i(X,Y)\} = \mathcal{F}\{I_0(x,y)\} \cdot \mathcal{F}\{H(x,y)\} \]  

(38)

Now the non-normalized OTF can be defined as \( \Theta(k_X,k_Y) = \mathcal{F}\{H(x,y)\} \), because the multiplication with \( \mathcal{F}\{H(x,y)\} \) produces the alteration in the frequency spectrum of the object, converting it into the image spectrum. Dividing \( \Theta(k_X) \) by the zero spatial frequency value \( \Theta(0) = \int_{-\infty}^{+\infty} H(x) \, dx \) the normalized spread function can be described as:

\[ H_n(x) = \frac{H(x)}{\int_{-\infty}^{+\infty} H(x) \, dx} \]  

(39)

Resulting in the normalized OTF:

\[ T(k_X) = \frac{\mathcal{F}\{H(x)\}}{\int_{-\infty}^{+\infty} H(x) \, dx} = \mathcal{F}\{H_n(x)\} \]  

(40)

For the description of the achievable resolution of an imaging system the concept of spatial bandwidth \( \Delta \kappa_\perp \) and the spatial resolution \( \Delta \rho \) is introduced [following Mertz [2010, pp. 36 and pp. 62]]. \( \Delta \kappa_\perp \) describes the amount of lateral wave vectors supported by the OTF. A Gaussian fit over the OTF of the optical system with the determination of the full width at half maximum (FWHM) can be used as a value for the resolution in microscope setups.

The relation \( \Delta \rho = \frac{1}{\Delta \kappa_\perp} \) describes the spatial resolution. The full angular width of the entrance aperture \( \Delta \theta_0 \) of the optical system limits the supported frequencies. For finding the classical resolution in a wide-field imaging system one can state (with \( \kappa = \kappa_\perp + \kappa_z \)):

\[ \Delta \rho_0 = \frac{1}{\kappa \Delta \theta_0} \]  

(41)

The relationship between resolution and bandwidth in the object plane (\( \rho_0 \) and \( \Delta \kappa_{\perp 0} \)) and the image plane (\( \rho_1 \) and \( \Delta \kappa_{\perp 1} \)) lead to the magnification \( M \):

\[ |M| = \frac{\Delta \rho_1}{\Delta \rho_0} = \frac{\Delta \kappa_{\perp 0}}{\Delta \kappa_{\perp 1}} \]  

(42)

Taking the angular width of the aperture pupil of the imaging system into account given by (\( \theta_0 \) denoting the half-angular width seen from the object plane \( \frac{1}{2} \cdot \Delta \theta_0 \) and \( n \) the index of refraction):
\[ NA_0 = n \cdot \sin(\theta_0) \] (43)

In the paraxial limit \((NA_0 \rightarrow n\theta_0)\) eq. (43) becomes:

\[ NA_0 = \frac{n\Delta\kappa_{\perp,0}}{2\kappa} = \frac{\lambda\Delta\kappa_{\perp,0}}{2} \] (44)

This results in the well-known equation for the diffraction-limited resolution by Abbe [Abbe 1873]:

\[ \Delta\rho_0 = \frac{\lambda}{2NA_0} \] (45)

The introduced methods above (STED, (d-)STORM, SIM), commonly described as super-resolution microscopy techniques, refer to the diffraction-limited resolution, meaning that the achieved resolution is below this intrinsic limit. All methods share the fact that the convolution of OTF and the spectral information of the sample can be deconvolved. Therefore, the PSF has to be measured or estimated for a computational deconvolution step, which can be performed by several user-friendly open-source programs.

In addition to that it is possible to actively improve the performance of the optical system by engineering the PSF. Here, the combination of two or more excitation foci can be exploited to reduce the size of the excitation light volume (e.g. STED [Donnert et al. 2006]). The desired ideal PSF is the delta function, which can image the spatial distribution of the sample without any introduced point spreading.

Utilizing a structured illumination pattern for the imaging of a sample is another method to broaden the accessible frequency support of a microscope. Chapter 2.3.2 explains the principle of SIM microscopy, whose technique is applied to CARS-microscopy as a novel approach to enhance the resolution in this label-free microscopy technique.
Figure 10 – Broadening of a point-shaped object by the PSF
The PSF of the microscope broadens the apparent object size of the sample. The magnification $M$ of the optical system is the ratio of object size to image size. $\Delta \theta_0$ denotes the full angular width of the aperture pupil as seen by the object plane. Adapted from [Mertz, 2010, p. 37]].

Figure 11 – Relation of PSF and OTF in the image formation of an optical system
The spatial information of the object $I_0$ (red) is convolved with the PSF of the optical system resulting in the image $I_i$ (real space, fawn, top row). The Fourier transform (FT, bottom) leads to the frequency domain: the finally detected frequency content is limited by the OTF support of the optical system. Adapted from [Mertz, 2010, p. 39]].
2.3.2. Linear structured illumination

Structured illumination microscopy improves the spatial resolution of optical microscopy beyond the diffraction limit by utilizing striped illumination patterns \cite{Gustafsson2000, Gustafsson2000}. In many implementations a sinusoidal stripe pattern is generated by two or three beam laser light interference effects in the focal plane along a specific angle, described by a wave vector $k_{ill}$ and a specific phase $\varphi$. This pattern illuminates a stained sample (in fluorescence-based microscopy) leading to a striped excitation of the specimen. High signal intensities of the dyes are obtained in the regions of the peaks of the illumination pattern, whereas regions with low excitation energies (valleys of the pattern) will exhibit little to no contribution to the image and appear dark \cite{Schüttpelz2014}.

The principle of the method can be described referring to frequency mixing in radio wave transmission. A carrier frequency is mixed with a sample frequency to transmit the signal across the pass-band of the optical microscope. The advantage of the structured illumination is a resolution enhancement (up to all three dimensions) in the case, that the frequency of the pattern is increased to the point where it approaches the cut-off frequency. The user-selected frequency of the illumination pattern $k_{ill}$ is mixed with the unknown frequency of the sample. Consequently, the Moire effect in two dimensions leads to additional contrast in the image (see fig. 12). This method is exploited in fluorescence-based SIM microscopy for enhancing the resolution by acquiring a series of images, which is then reconstructed via specifically designed software \cite{Gustafsson2005, Gustafsson2008}.
The Moiré effect describes the overlapping of two 2D-striped patterns rotated by an angle to each other. The mixing frequency $\omega_M$ (orange) of the overlapped frequencies ($\omega_S$ green and blue $\omega_i$) can be observed [Schüttpelz et al. [2014]].

Considering pure frequency mixing, the addition of two cosine functions with frequencies $\omega_1$ and $\omega_2$ generates a beat mode carrying two new frequencies, the sum and the difference frequency, which can be described as [Schüttpelz et al. [2014]]:

$$\cos(\omega_1 t) + \cos(\omega_2 t) = 2 \cdot \cos \left( \frac{\omega_1 t + \omega_2 t}{2} \right) \cdot \cos \left( \frac{\omega_1 t - \omega_2 t}{2} \right)$$

In the spatial frequency space, having now the transformed $k_1$ and $k_2$, the additional frequency components are:

$$\cos(k_1 x) + \cos(k_2 x) = 2 \cdot \cos \left( \frac{k_1 x + k_2 x}{2} \right) \cdot \cos \left( \frac{k_1 x - k_2 x}{2} \right)$$

This mechanism is called heterodyning and can be exploited to expand the detectable range of frequencies, when one frequency is known. The second, unknown higher (and, therefore, undetectable) frequency can be obtained by measuring the reduced frequency of the envelope of the beat pattern. Hence, the limited pass-band can be enlarged up to twice of its original value. Expanding the pass-band any further by increasing the frequency of the beat pattern is not possible because the beat pattern itself is not supported any more and can not be detected.

Analyzing an arbitrary (e.g. widefield) image in the two-dimensional Fourier space its spatial frequencies appear as a circular pattern. Low frequencies are located near the center of the image, high frequencies at the border of the circle, having longer $k$ vec-
tors. The OTF support of the optical system can be observed as the radius of the circle depicting the cut-off-frequency introduced above.

Applying a sinusoidal pattern on a homogeneous sample two delta peaks appear in the Fourier transform, if the frequency of the illumination pattern is lower than the OTF support. Connecting these two delta peaks by an axis indicates the pattern orientation, while the distance from the peaks to the center of the image describe the frequency of the applied pattern.

For generating a resolution enhancement as high as possible the frequency of the illumination pattern is chosen to be near the edge of the OTF support. The undetectable spatial frequencies of the sample beyond the OTF support will be mixing with the known pattern frequency. By an unmixing step the unknown frequency can be calculated in a post-processing step utilizing the known pattern frequency. The final accessible frequency range will be broadened allowing to detect spatial sample features beyond the classic OTF support only limited by the need of detecting the mixing frequency.

Shown in 2.3.1 the image $I$ is the convolution of the fluorescence emission $F(\vec{r})$ and the PSF $H(\vec{r})$, meaning $I = F \otimes H$, while $\vec{r}$ denotes the lateral coordinates $(x, y)$. The varying illumination pattern $P(\vec{r})$ as well as the spatial distribution of the fluorescent dyes $S(\vec{r})$ are determining $F(\vec{r})$, $F = P \cdot S$. Assuming a sinusoidal stripe pattern along a specific angle $\vartheta_{rot}$, with the frequency $k_{ill}$ and the phase $\varphi$ the pattern can be described as [derivation follows Schüttpelz et al. [2014]]:

$$P(\vec{r}) = \frac{I_0}{2} \left[ 1 - \cos(k_{ill} \cdot \vec{r} + \varphi) \right]$$

Having $I_0$ as the initial illumination intensity. The measured image $I(\vec{r})$ can now be stated as:

$$I(\vec{r}) = (S(\vec{r}) \cdot P(\vec{r})) \otimes H(\vec{r})$$

Performing a Fourier transform leads to the term in reciprocal space, turning the deconvolution of the PSF and the fluorescence emission into a simple product:

$$\tilde{I}(\vec{k}) = \tilde{H}(\vec{k}) \left[ \tilde{P}(\vec{k}) \otimes \tilde{S}(\vec{k}) \right]$$

The sinusoidal illumination pattern occurs in the frequency domain as a series of delta functions $\delta$, which can be used to simplify the equation to:

$$\tilde{P}(\vec{k}) = \frac{I_0}{2} \left[ \delta(\vec{k}) - \frac{1}{2} \delta(\vec{k} - k_{ill})e^{-i\varphi} - \frac{1}{2} \delta(\vec{k} + k_{ill})e^{i\varphi} \right]$$

Finally the observed image in the frequency domain can be stated as the convolution
of the pattern with the fluorophore distribution:

\[
\tilde{I}(\mathbf{k}) = \frac{I_0}{2} \tilde{H}(\mathbf{k}) \left[ \tilde{S}(\mathbf{k}) - \frac{1}{2} \tilde{S}(\mathbf{k} - \mathbf{k}_{\text{ill}}) e^{-i\varphi} - \frac{1}{2} \tilde{S}(\mathbf{k} + \mathbf{k}_{\text{ill}}) e^{i\varphi} \right]
\]

(52)

In the following post-processing step for reconstructing the raw data into the final image, which has enhanced resolution, the parameters of the system’s PSF as well as the properties of the illumination pattern have to be identified beforehand by a parameter estimation. Then the superresolved distribution of the fluorophores in the sample can be calculated. The PSF can be measured experimentally by imaging a sub-diffraction-limited structure such as e.g. a fluorescent quantum dot or a single fluorescent bead (Ø ≤ 100 nm), which is background corrected and rotationally averaged. The OTF can be stated afterwards by calculating the 2D Fourier transform of the PSF.

Typical SIM raw data contains a stack of single images described by eq. (52). For a constant pattern angle ϑ_{rot} = const. the phase ϕ of the pattern is shifted by translating the interference pattern across the sample for a number of images N. (N ≥ 3 for 2D SIM and N ≥ 5 for 3D SIM). The phase shifts are spaced evenly over 2π (∆ϕ = \frac{2\pi}{N}) (see fig. [13]).

In the experiment M separate values of ϑ_{rot} are set in order to cover the focal plane, which are evenly spread across the image space (∆ϑ_{rot} = \frac{2\pi}{M}), with M ≤ N. The data set of M x N images is background-subtracted and intensity-corrected compensating for bleaching effects during the acquisition as well as source fluctuations. Then the data is Fourier transformed into reciprocal space. Here, a pairwise cross-correlation is utilized to identify the values of the frequency shift vector, the phase shift and the modulation depth for each image, while by linear regression the initially estimated values are optimized.

The information of the images of each angle are combined, each phase triplet is separated into three components: the central band identical to the conventional widefield image, and two side bands representing the new spatial information. The side bands are translated by a vector \mathbf{k}_0 in reciprocal space restoring the information to the original position.

Without a final noise filtering step the obtained image would suffer due to additional noise of the high spatial frequency region. The exclusion of undesired noise is performed typically by a Wiener filter or a Richardson-Lucy deconvolution step (referring to FairSIM [Müller et al. [2016]]). In addition a suitable cutoff frequency has to be identified to exclude contributions, which only provide noise and no further sample information. This task is performed by an apodization function applied to the side band information. It is also evening out overlapping frequency components ensuring an even intensity distribution in the reconstructed image. Then the final image is obtained by
transforming back into real space. In the best case a twofold improved spatial resolution is found in the reconstructed image (compared to standard widefield imaging) [Gustafsson, M. G. L. 2000]. For the reconstruction purpose of SIM data sets open source software packages are published recently, which utilize the image processing platform FIJI [Schindelin et al. 2012, Rueden et al. 2017] in the case of FairSIM [Müller et al. 2016] or MATLAB in the case of MAP-SIM [Křížek et al. 2015, Lukeš et al. 2014].

For the generation of the illumination pattern in the focal plane typically the interference of two (or three in 3D SIM) laser beams is utilized, which are focused onto the back focal plane of the objective lens and finally generating two counter propagating evanescent waves in the focal plane utilizing a diffraction grating. Is is rotated and laterally shifted by piezoelectric transducers, but also limits the imaging speed. More recent attempts replace the diffraction grating by active optical elements, such as e.g. spatial light modulators [Hirvonen et al. 2009, Rego and Shao 2015] or galvanometric mirrors [Mandula et al. 2012] allowing rapid changes in the phase and angle of the pattern. The image detection is performed by CCD or sCMOS cameras providing high sensitivity as well as high acquisition speed for live-cell imaging [Kner et al. 2009, Wu and Shroff 2018, Heintzmann and Huser 2017].

Alternative implementations are utilizing micro-mirror arrays for the generation of point-shaped, multi-focal illumination patterns [York et al. 2012], which are swept over the sample for an improved imaging speed in 2D SIM. Here, the optical excitation and detection pathway can be folded and manipulated by a galvanometric mirror, which affects both optical paths. This way parts of the reconstruction process are performed on a direct, optical way for an enhanced imaging speed [York et al. 2013].
A sinusoidal structured pattern with wave vector $k_0$ is exploited for generating the illumination of the sample. It leads to Moire fringes at $k_{Moire} = k - k_0$ (when $k$ is the sample’s frequency content) (a). Side bands at the mixing frequencies with the vectors $k + k_0$ and $k - k_0$ appear and provide additional spatial information of classically unobservable frequency regions. The microscope can observe only frequencies which are located within the OTF support, indicated with the heavy circle (b). The observed data includes the classic widefield information of the OTF support itself as well as the additional information of the side bands, which is shifted by the vector $k_0$ (indicated with the light circles in b). This information has to be decoded and restored to its original position. By rotating the pattern with three or more angles the additional frequency content can be spread evenly over the frequency space, generating a new circle of observable sample information, which has up to twice the diameter of the initial OTF support (c) [adapted from Gustafsson, M. G. L. [2000]].

### 2.3.3. Non-linear structured illumination

The maximum resolution enhancement in a SIM experiment is limited to a factor of two while using a sinusoidal pattern with a linear excitation of the sample. Introducing a non-linearity into the imaging process the theoretical predicted improvement in resolution improvement is unlimited (Heintzmann et al. [2002]). In fluorescent-based microscopy a saturated structured illumination can generate this non-linear feature [Gustafsson, M. G. L., [2005]]. The intensity of the excitation pattern is increased to the point, where the fluorescent dye is saturated and any stronger illumination intensity does not produce a higher fluorescent signal. This is due to the rather long lifetimes of the excited states of the fluorophores. The mechanism limits the number of emitted photons per time per unit. Now the sinusoidal illumination generates flat tops at the peak intensities of the fluorescence response of the specimen. The non-linear pattern combined with an increased number of phase shifts and angles of rotation leads to an optical spatial resolution of $\sim 50\text{nm}$ in recent implementations [Rego et al. [2012]]. The higher intensities of the excitation power compared to linear SIM lead to increased bleaching effects, which are a limiting factor and disadvantages of this method. In addition the penetration depth of the generated pattern in thick, densely labeled fluorescent samples is still limited, due to scattering effects at the sample ma-
terial, which potentially lead to disturbed illumination patterns.

For increasing the accessible sample thickness the combination of line scanning with structured illumination microscopy (LS-SIM) is performed [Mandula et al. [2012]] allowing the suppression of out-of-focus light by line scanning, while still the resolution is enhanced by SIM. To introduce the non-linearity the length of $k_0$ is chosen to be small compared to the linear Gustafsson-SIM method. Now the non-linearity in the data sets becomes apparent analyzing the Fourier transform, while the length of multiples of $k_0$ is in the OTF support of the optical system. The additional peaks at integer multiples of the initial $k_0$ vector of the illumination pattern appear in the frequency space laterally shifted along the $k_0$ direction (see fig. 14 and fig. 15).

The reconstruction of the data needs exact knowledge of the position and length of the $k$ vectors to shift the additional sample information to its origin in the final image, but also holds great potential in terms of final obtained resolution enhancement. Especially the MAP-SIM approach is capable of reconstructing non-linear excitation patterns [Lukeš et al. [2014]], which are also apparent in the line scanning-based SI-CARS approach introduced in 4.5.5. The non-linearity in the galvomirror-based scanning of specific patterns in a sample arises from the non-sinusoidal intensity distribution of the excitation pattern. Here, the OTF function (mostly determined by the objective lens) is smoothing the rather rectangular on-off-like illumination pattern, but this effect is also dependent on the fineness of the pattern. The finer the pattern the more sinusoidal it gets due to the reduced frequency support of the OTF in that particular frequency region.
In (a) the Fourier transform is depicted for a linear SIM image, two additional frequency components occur in the spectra, shifted from the origin with $\pm k_0$. In the case of non-linear SIM the illumination pattern contains spatial frequencies which are integer multiples of the initial illumination pattern, shifted with $k_0$. In (b) a single angle of rotation is shown, for full resolution enhancement in all spatial directions it is shifted in smaller fractions of $2\pi$. (c) depicts the necessity of small angular shifts: the overlap of the frequency content in higher orders of $k$ can be accomplished by reducing the step size of the rotation angle [from Rego et al. 2012].

**Figure 15** – Fourier transform of SI-CARS image indicating second order frequency components

(a) shows a CARS image with an applied very coarse structured illumination pattern utilizing galvanometric scanning mirrors. The sample was a polystyrene bead sample ($\Theta = 1, 1 \mu m$) embedded into a DMSO solution while probing the $2913 cm^{-1} C - H$ symmetric stretching mode of the DMSO solution generating an anti-contrast. More details are provided in 4.5.5. The scale bar depicts $5\mu m$. Fourier transform (b) of the single raw frame acquired by FIJI [Schindelin et al. 2012, Rueden et al. 2017]. The arrows point at the frequency peaks introduced by the illumination pattern while the green circle depicts the estimated cutoff frequency. Note that the Fourier transform was rescaled for visualization purposes.
3. Materials and Methods

In the course of this thesis a state of the art CRS-setup was developed\textsuperscript{5}. It has a variety of imaging modalities and different work modes. In order to show its capabilities the following chapter will first focus on the general instrumentation and setup, which is used in almost all shown different applications in the thesis. In order to report about the successful construction, characterization and application of the CRS-microscopic setup a selection of data is shown, originating from collaborative work where CRS-imaging was beneficial for actual research in different biomedical fields such as the study of stem cell differentiation (section 4.1), the lipid storage of algae for bio-fuels (4.2), the study of lipid storage in C. elegans under the influence of copper treatment (4.4.1) as well as the differentiation of primary human alveolar epithelial cells (4.4.4). Further work, which is addressing new insight into CRS-imaging and novel technical developments is presented: a study and characterization of the OPO pulse length while performing a wavelengths tuning for hyperspectral (HS) CARS and SRS imaging causing possible spectral distortions in the final data as well as presenting a controlled OPO tuning mode to perform HS-CARS with stable pulse length and, therefore, reproducible and undistorted data by a custom developed Matlab-based OPO tuning control program (section 4.3). In addition a novel imaging modality utilizing structured illumination for CARS (SI-CARS) is presented for enhancing the resolution of conventional CARS imaging beyond the diffraction limit (section 4.5). Therefore, two-photon excited fluorescence data is compared to the SI-CARS data in order to prove the applicability of structured illumination in combination with a coherent contrast mechanism. Due to the need for compact, stable, user-friendly and low cost CRS-laser sources in clinical applications a collaboration with the University of Hong Kong was forged in order to apply all fiber-based techniques for the construction of a novel laser source providing CARS- as well as SRS-imaging without balanced detection, SHG and two-photon excited fluorescence imaging in one approach (section 4.6). Furthermore, a fiber-based short pulse laser source is introduced providing fs-laser pulses with up to 500\textit{mW} intensity, which is applied for SHG and two-photon excited fluorescence imaging. This source is a preliminary results of the project and was proposed as a stand-alone laser source interesting for users of two-photon excited fluorescence and SHG imaging.

\textsuperscript{5}In collaboration with Mr. H. Hachmeister.
3.1. Experimental setup

The following section describes the CRS-microscope, which can be categorized in three main parts (fig. 19): The laser source providing a fundamental IR-wavelength for the Stokes beam (see 3.1.1) as well as a frequency doubled wavelength pumping two OPOs, which deliver tunable pump wavelengths for the CRS process (see 3.1.2). It follows the spatio-temporal overlap of the three excitation beams as well as the beam diameter and divergence adaption in combination with intensity and pulse length diagnostics. Finally, a home-built scanning microscope is utilized for the sample analysis enabling a maximum degree of freedom in the detection and manipulation modes as well as superior performance in terms of stability over long time scales.

3.1.1. Solid-state laser system

A solid-state picosecond laser (HighQ, picoTRAIN IC-1064-15000 / 532-10000 ps Nd:VAN) is utilized for the generation of the Stokes beam as well as to pump an double OPO system in the CRS-setup. It is located on an actively damped optical table (TMC) in order to avoid vibrations in the laser system as well as in the microscope. The laser source consists of an oscillator (fig. 16 blue box A), an amplification stage (red box B) and a final frequency doubling unit (green box C). The oscillator has a wavelength of 1064 nm and can deliver up to 10.8 W power (see fig. 16). A Nd:VAN optical crystal is placed in the folded laser cavity, where the population inversion is created by optical pumping using a laser diode array, whose light is focused into the crystal by a cylindrical lens (HighQ [2012]). The mode locking is performed passively with a SESAM, which has a gold metalized back side soldered to a heat sink mounted in a piezo driven mirror mount. It is used for remote cavity alignment through the mirror mount performed by the user. The laser cavity itself is terminated on one side by the SESAM mirror and on the other by the metalized end facet of the Nd:VAN crystal, which directs a definite portion of the IR-beam to the amplifier stage passing an isolator. This additional amplifier stage utilizes a diode pumped amplifier crystal and delivers an output power of 16.1 W. By a motorized ¼-wave plate and a thin-film polarizer a user selected portion of the IR-beam is frequency doubled to 532 nm through second harmonic generation by a non-linear crystal. For a high efficiency of the SHG process a temperature control is applied to the crystal, which enables a maximum output power of 10.4 W (100% 532 nm). For meeting the need of pumping two OPOs at the same time the laser has three outputs, one for the fundamental 1064 nm beam as

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6In case of DR-CARS.
7Semiconductor saturable absorber mirror
well as two outputs for 532nm. Again a motorized attenuator manages the energy ratio between the two outputs of the 532nm beam. In normal operation 10% of the infra-red power are used as the Stokes beam in the CARS experiments, the remaining portion is frequency doubled and leaves the laser at the two remaining outputs for pumping the OPOs. The polarization of the IR-beam as well as of the 532nm beam leaving the laser at exit 2 is horizontally polarized whereas the 532nm beam of exit 1 has a vertical polarization. In order to meet the requirements for the OPO, which needs vertically polarized light at its entrance window, a wave-plate is place into the beam path of exit 2 in order to turn its polarization into vertical. The 1064nm laser wavelength has a pulse length of 7, 4ps, the 532nm frequency doubled portion has 6, 2ps respectively. The configuration of one master oscillator as fundamental laser source is beneficial for the performed CRS-experiments in terms of temporal and spatial coherence of all beams due to the absence of any electronic synchronization mechanism among two separate excitation sources, which are reported as former configurations in the literature [Ganikhanov et al. 2006, Cheng and Xie 2013, pp. 83].

The application of a pulsed laser system with a high pulse peak energy leads to strongly enhanced CRS signals generated in the sample. Due to the resulting moderate mean powers in biological specimens low thermal heating and non-critical photonic stress is observed. A laser system with picosecond instead of femtosecond pulse duration meets the aim of addressing a single Raman mode for a high signal-to-background ratio in CRS as well as a high spectral resolution for hyperspectral imaging. Here, the line width of the laser excitation matches well to the line width of the probed vibrational mode generating higher CRS signal intensities. Also the undesired non-resonant background in the CARS process is rather low in the regime of pulse length applied in this setup [see also Zumbusch et al. 1999, Cheng and Xie 2013, pp. 80-82].
The photograph provides an overview of the three main components of the solid-state laser system (the access covers are open for maintenance): The blue box (a) marks the master oscillator cavity with the pump diode array, Nd:VAN crystal and the SESAM mirror for passive mode locking. In the red box (b) the amplification stage can be seen. Here, the 1064nm beam is focused again into a pumped optical crystal and can be amplified to a maximal output power of $P_{1064\text{nm}} = 16,1 \text{W}$. In the green box (c) the second harmonic generation (SHG) through a temperature controlled crystal with the following division by polarization dependent optical beam splitters into the different output ports takes place [HighQ [2013]].

### 3.1.2. Optical parametric oscillator

An optical parametric oscillator (OPO) is an array of optical components inside an optical cavity used for the conversion of high energetic photons into lower energetic photons with two defined energy levels, whereas the ratio of both generated photons is tunable in a system specific range (see fig. [17]). The energy-shifted photons are named signal and idler, the shorter wavelength is defined as the signal [Cheng and Xie [2013], Ganikhanov et al. [2006]]. The conversion is performed by focusing the light into a non-linear crystal exploiting the optical parametric generation (OPG), which is a $\chi^{(2)}$-process. Here, the law of energy conservation ($h\omega_{\text{pump}} = h\omega_{\text{signal}} + h\omega_{\text{idler}}$) as well as the phase-matching condition ($\vec{k}_{\text{pump}} = \vec{k}_{\text{signal}} + \vec{k}_{\text{idler}}$) has to be satisfied, while in this configuration the collinear non critical type I phase matching interaction is exploited in the LBO crystal [Weiner [2009] pp. 245]].
The optical parametric conversion performed in a non-linear crystal is shown. Incoming pump photons $\omega_{\text{pump}}$ (green) are converted into the lower energetic signal $\omega_{\text{signal}}$ (orange) and idler $\omega_{\text{idler}}$ (red) wavelength (right hand side). For illustration purposes the directions of the generated beams are not congruent.

In the CRS-setup two commercial synchronously pumped OPOs (APE, Levante Emerald) generate tunable wavelengths used as the pump photons in the CARS and SRS process (fig. 18). Two OPOs are chosen to further extend the range of probed wavenumbers by granting more freedom of combining different wavelengths in the CRS process as well as to enable DR-CARS experiments. The solid state laser system pumps each OPO’s LBO crystal (lithium triborate) with $3.5 - 4.0 \, \text{W}$ of $532 \, \text{nm}$ laser light. For a high efficiency the pump beam is focused by a lens into the crystal, which is placed in a folded 8-mirror cavity. The cavity length of $3.75 \, \text{m}$ is matching the repetition rate of the laser source ($\sim 80 \, \text{MHz}$). This way every $12.5 \, \text{ns}$ a pump pulse enters the cavity and generates the signal and idler photons inside the crystal. All photons travel in the cavity and meet with the next pump pulse in the crystal generating a high instantaneous intensity for a high efficiency of the OPG-process. The cavity length must be tuned precisely to every desired signal wavelength as well as to the repetition rate of the incoming pulses, which is performed by a piezo-actuated delay stage (mounting mirror M2). The smallest increment the stage can travel is $85 \, \text{nm}$. The phase matching condition according to the desired signal wavelength is performed via temperature control of the crystal. Due to the type I phase matching condition the incoming vertically polarized $532 \, \text{nm}$ pump photons generate the horizontally polarized signal and idler photons exiting the OPO. Their polarization is then matching the polarization of the $1064 \, \text{nm}$ laser beam in the final CRS experiment, which was initially horizontally polarized \cite{APE, Powers, pp. 94}. The crystal is placed on a resistor heated mount and can be heated in the range of $105 ^\circ \text{C}$ to $160 ^\circ \text{C}$ in $0,1 ^\circ \text{C}$ steps using a thermocouple for control. A look-up table allows the choice of the correct temperature. The natural bandwidth generated by the LBO crystal is in the range of $\approx 15 \, \text{nm}$. For the desired smaller bandwidth and final tunable wavelengths selection a simplified
Lyot filter scheme is applied in the OPO’s cavity. It consists of a birefringent crystal performing a wavelength-dependent polarization according to its rotation position, which is electronically actuated by a rotation mount. Finally, a polarizer selects the supported wavelength by its polarization sensitive transmission. The Lyot filter further suppresses undesired wavelengths with each pass through the filter layout, resulting in a spectral width of approximately $\Delta \lambda = 0.3 \text{nm}$ (full width at half maximum).

The signal and idler photons are finally split with a short-pass filter into two exits. In the current setup only the signal photons are used for the CRS-experiments. A clean-up filter (AHF, FF01-855/210BP) placed inside the signal beam path eliminates the remaining pump photons and residual idler wavelengths of each OPO. A built-in spectrometer and two photo-diodes measure the incoming pump beam intensity as well as the outgoing signal intensity allowing the adjustment and control of the OPO.

Both commercial OPOs are identical in their performance and can be operated simultaneously. Their tuning range is 680 – 990 nm for the signal wavelength whereas the available range in the CRS experiment is more limited due to the dichroic mirrors installed for the spatial overlap (see fig. 19). The resulting available wavelength range is 950 – 835 nm for OPO I and 830 – 680 nm for OPO II. Utilizing the 1064 nm beam as fixed Stokes pulse leads to an addressable wave number range of $\sim 1100 \text{cm}^{-1}$ to $\sim 2578 \text{cm}^{-1}$ (OPO I) and $\sim 2650 \text{cm}^{-1}$ to $\sim 5753 \text{cm}^{-1}$ (OPO II). If a broader spectral range is needed for CRS-experiments the dichroic mirrors can be replaced by 50/50 beam splitters sacrificing 50 percent of the incoming power at each passed splitter.

In general the OPOs have a good long term stability of the optical cavity so that a readjustment is needed every two to three months.
The 532nm pump beam (green) enters the OPO on the upper left hand side. It is focused into the LBO crystal (C1) by a 75mm lens (L1) and generates the signal and idler photons. The folded cavity beam path is shown in blue. A tunable birefringent crystal (BRF) and a polarizer (PL) form a simplified Lyot filter for defining the signal wavelength as well as the bandwidth. A small portion of the generated photons is directed into the spectrometer (SP) for wavelengths monitoring. Signal and idler beam exit the OPO on two separate ports, which contain Brewster windows (not shown).


3.1.3. Spatio-temporal overlap with laser diagnostics

Due to three different sources of laser light it is necessary to overlap all beams in spatial and temporal properties in the specimen. The temporal overlap is adjusted by two delay stages which introduce additional optical path length to each laser source. The Stokes beam is directed onto the first delay stage, which is constructed by a retro-reflector (Thorlabs, PS973M-B) mounted on a linear translation stage (own construction by workshop University Bielefeld) (see fig. 19). Delay stage 1 covers the temporal overlap of the 1064nm beam and OPO I. The second delay stage is home-built constructed by two dielectric mirrors (Thorlabs, BB1-EO3) facing each other in a 45° arrangement (using for mirror mounting Radiant Dyes, MNI-23025) built on a linear translation stage (Owis, LT60-50). The light path of OPO II is directed onto this stage in order to adjust it to the already matched temporal overlapping beams of Stokes and OPO 1. The need for the use of a mirror based delay stage instead of the retro-reflector approach arises from the change of wavelength during OPO tuning in the pump beam path. This avoids a rotation of polarization in the case of passing a retro-reflector with different wavelengths. Here, we give up the benefit of utilizing a retro-reflector in terms of alignment speed and simplicity but gain an undisturbed polarization of the pump beam in the experiments over the whole range of pump wavelengths.
The spatial overlapping of the 1064nm beam with the two pump beams of the OPOs is performed with a dichroic mirror 1 (DM1, AHF, F73-951 beam splitter). It is transmitting the 1064nm beam and reflecting any shorter wavelengths than 950nm, so that the signal wavelengths of both OPOs are reflected to the microscope. The overlap of the two signal wavelengths of the two OPOs is performed in a similar way at dichroic mirror 2 (DM2, AHF, F43-831), which is reflecting any wavelength shorter than 831nm. Half-wave plates (Thorlabs, AHWP05M-980) in combination with polarizing beam splitter cubes (Thorlabs, PBS252) are utilized for variable intensity attenuation by rotating the polarization of the light of each source with the half-wave plate. This way the desired intensity for the different samples can be set with an accuracy of $\sim 0.5mW$.

In order to adjust the beam diameter and control the beam divergence a magnifying telescope (focal length $f_{T1} = 45mm$ and $f_{T2} = 80mm$, all Thorlabs, AC254-XX-B) is implemented for each OPO (T1 and T2). Here, the second lens is mounted on a linear translation stage for precise tuning of the divergence. Finally all three beams enter the laser scanning microscope with a beam diameter of 4,3mm (FWHM) overlapping in time and space. For further analysis of the interacting pump and Stokes beams we implemented an automated intensity measurement setup consisting of three self constructed fast photodiodes (Luna Optoelectronics, PDB-C613-2), which are read out by an analog-to-digital (A/D) converter (National Instruments, PCIe-6363). A pellicle beam splitter (Edmund Optics, 8R/92T 25.4mm) reflects $\sim 2\%$ of each laser beam onto the photodiode. Here, great care was taken for choosing a wavelength unaffected way of beam separation. A Matlab-based custom written program was developed for recording the actual intensity of each participating laser source linking the intensity values of the excitation beams to the measured signal intensity of the specimen. In a post processing step a correction for intensity fluctuations up to 100kHz as well as normalization between different images can be performed (see also 4.3).
The laser system (light blue) generates the 1064nm beam (depicted in dark red) as well as pumping two OPOs (dark blue), which generate the pump beams for the CRS experiments (in light red). In the middle the spatial overlap is performed with two dichroic mirrors (DM1 and DM2). Two delay stages are utilized to add optical path to Stokes and pump beam 2 in order to perform the temporal overlap of all pulsed laser lines. For intensity corrections as well as normalization in the CRS data three photodiodes (PD) detect the actual intensity of all laser sources by using a small portion of each beam separated by pellicle beam splitters. In a similar way an autocorrelator (Autocor.) is in use for recording the actual pulse length of the laser light. An AOM performs an intensity modulation of the Stokes laser for SRS as well as modulated CARS experiments.

3.1.4. Acousto-optic modulator

An acousto-optic modulator (AOM) (Pegasus Optik, MT 200-A0.4-1064) performs an amplitude modulation of the 1064nm Stokes beam. In this way we enable background free SRS experiments, frequency modulated CARS experiments (in order to suppress e.g. auto-fluorescence of samples) as well as it is used for fast intensity switching for the generation of a structured illumination pattern for SI-CARS. An AOM consists of an optical crystal (here TeO$_2$), which is coupled to a piezo-element. The piezo-driver (AA Opto-Electronics, A200 B4-34) applies a high radio frequency (RF) to the piezo-element (200MHz), which acts as a carrier frequency and creates an acoustical standing wave inside the crystal. The density variations lead to a local change of the refractive index in the material generating an optical grating. Incoming light is diffracted at the grating in different orders, well known from Bragg-diffraction [Pegasus Optik 2007]. For precise alignment of the crystal to the light path it is mounted on a pivotable xyz-stage, enabling a high diffraction efficiency into the first order. All other orders are blocked exploiting the $\sim 1,3^\circ$ offset of order +1 to the zero order. The AOM’s crystal is positioned into the focus of a telescope, which is consisting of the
biconvex lenses $L_1$, $f_{L_1} = 300\, mm$ and $L_2$, $f_{L_2} = 400\, mm$ (see also fig. [fig]). This way parallel light with the desired beam diameter is generated behind the AOM as well as the small entrance aperture of the device is matched, which allows fast rise times. For the final amplitude modulation of the Stokes beam the RF-carrier frequency is externally amplitude modulated by a signal-frequency (setting an amplitude of $0 - 5\, V$) applied to the driver with a waveform generator (e.g. *Hameg*, HMF 2550). This leads to a diffraction efficiency into the first order, which is directly proportional to the signal amplitude of the waveform generator. According to the Nyquist-criteria the maximal theoretical frequency of the light modulation is $100\, MH\dot{z}$, meaning that for the successful transformation of the desired frequency onto the beam the carrier frequency has to be doubled at least. The optical properties of the AOM’s crystal further limit the maximal modulation frequency to a value of $60\, MH\dot{z}$.

The focus size of the telescope was chosen to $\sim 98, 5\, \mu m$ (according to the grid spacing of $21\, \mu m$) ensuring a fast rise time for the experiment as well as to avoid disturbing diffraction effects in the outgoing beam [Pegasus Optik [2007]]. This results in a final restriction of the frequency to maximal $\sim 20\, MH\dot{z}$, which is sufficient for all conducted experiments. Due to the crystal orientation in combination with the mounting mechanics a $\frac{\lambda}{2}$-wave-plate turns the polarization of the light matching the polarization direction of the AOM. Finally the transmitted light is turned back by a second halfwave-plate. In the current installation a transmission efficiency of $76\%$ is reached, while having a modulation depth (at $20\, MH\dot{z}$ modulation frequency) of $\sim 75\%$.

### 3.1.5. Setup for a customized CRS-microscope

For CRS, SHG as well as two-photon excited fluorescence imaging a home-built scanning-microscope was developed. The decision for a customized solution instead of a commercial microscopy body was motivated by the need of a higher grade of flexibility in terms of detection pathways as well as more room for additional optical elements. For being competitive to commercial microscopes a lightproof body housing shields the inner part but also allows fast changes of sample or optical elements inside. Therefore, a frame of rectangular construction rails (*Thorlabs*, XE25L600/M) in combination with black hardboard (*Thorlabs*, TB4) was set up adding sealed cable feed through as well as blackened aluminum sheets as doors, which are removable due to a magnetic holding system. This way any disturbing room light was suppressed.

The scanning of the excitation beams over the sample in a line per line manner is performed by a set of $xy$-galvanometric laser scanning mirrors (*Cambridge Technology, Galvanometer Optical Scanner*, Model 6215H). Each mirror of the pair varies the deflection angle of the beam in one direction, according to a control voltage ($0 - 10\, V$) applied
at the power module of the scanning unit. The control voltage is set by a multifunctional scanning program (ScanImage, Janelia Farm) [Pologruto et al. 2003], which operates an analog-to-digital (A/D) converter (National Instruments, PCI-6110S multifunction device connected to a breakout box BNC-2110). The incoming beams are further magnified by a scanning telescope after the scanning mirrors (Thorlabs, AC508-XXX-B-ML, with \(f_{SL1} = 150\text{mm}\) and \(f_{SL2} = 200\text{mm}\), magnification of 1.33) in order to fill the back focal plane of the objective (Ø \(\approx 8\text{mm}\)) enabling a high resolution through high focus quality without cutting parts of the beam during the scan process. According to the chosen integration times\(^8\) the maximum imaging speed reached with the galvo scanners was 8, 9Hz frame rate.

A non-moving illumination of the back focal plane of the objective lens during the scanning process is important for a even and uniform illumination of the final image over the entire field of view. Especially in SRS experiments intensity losses due to beam cutting can lead to imaging artifacts. Therefore, the distances between the scanning mirrors, the telescope and the objectives’ back focal plane are chosen in a special ratio (see fig. 20) [Fällman and Axner 1997]:

\[
d_1 = f_{SL1} \cdot \frac{f_{SL2} \cdot d_2 - d_3 \cdot (d_2 - f_{SL2})}{f_{SL2} \cdot (d_2 - f_{SL1}) - d_3 \cdot (d_2 - f_{SL1} - f_{SL2})}
\] (53)

Using \(d_2 = f_{SL1} + f_{SL2}\) due to the system’s afocallity eq. (53) is further simplified to:

\[
d_1 = f_{SL1} \cdot \frac{f_{SL1} \cdot f_{SL2} + f_{SL2}^2 - d_3 \cdot f_{SL1}}{f_{SL2}}
\] (54)

\[
d_1 = \frac{f_{SL1}}{f_{SL2}} \cdot \left( f_{SL1} + f_{SL2} - d_3 \frac{f_{SL1}}{f_{SL2}} \right)
\] (55)

---

\(^8\)128x128px and 0,5ms per line
The scanning telescope composed of two lenses having a focal length of $f_{SL1} = 150\text{mm}$ and $f_{SL2} = 200\text{mm}$ (both 51mm in diameter) is magnifying the raster scanned laser beams by a factor of 1.33, while it ensures a non-moving beam position at the back focal plane of the objective lens. The distances of galvo scanning unit, telescope and back-focal plane have to be set carefully according to the equation\[55\]. In red the radial line is shown, in blue the parallel beam, which should be congruent with the radial line at the back focal plane for all scanning angles. The parallelism of the scanning telescope was checked in an external experiment with a parallel laser beam using a long imaging distance.

For focusing the excitation beams into the sample an apochromatic water immersion objective (Olympus, UPLSAPO, $60\times$, NA 1.2) is used (see fig. 21). The focus is set with a linear translation stage containing a differential micrometer screw (resolution in manual translation of $\sim 0.5\mu\text{m}$ with a smallest possible incremental move of $\sim 50\text{nm}$). The transmission ratio of objective lens, scanning telescope and scanning mirrors is found to be $\frac{1}{6}$ for the 1064nm Stokes beam and $\frac{1}{3}$ for the pump beams (measured at the wavelength of 816nm).

The sample is fixed on a motorized xy-stage (Märzhäuser, MT mot 100 $\times$ 100 MR) which can be addressed by the scanning control program via RS-232 for moving the sample in an image series enabling a bigger field of view. A closed-loop piezo controlled z-stage (Märzhäuser, S/N 222009) offers z-positioning of the sample in 1nm step size and is also connected to the imaging software. Hence, it is possible to perform a 3D-analysis of a sample in an automated and fast manner. A dichroic mirror (BS1) (Semrock, FF757-Di01-25 $\times$ 36) placed between the second lens of the scanning telescope ($SL2$) and the objective lens is used to detect epi-CARS signals or bright light from the sample. A flip mirror mount switches between epi-CARS imaging via a photomultiplier module (PMT) (Hamamatsu Photonics, H 9656-20) (connected to ScanImage) or a CMOS camera (IDS Imaging, µ-eye SE) for bright-field imaging, while the sample is illuminated from above by a halogen or LED lamp. Here, an additional lens $L_{I4}$ with a focal length of $f_{L_{I4}} = 120\text{mm}$ focuses the desired signals onto the camera chip or, when the mirror is flipped into the beam path, onto the PMT´s detector area. In forward detection a condenser lens (Olympus, U-AAC, NA = 1.4) is used to collect the CRS signals in standard operation. It has a high transmission for CARS signal.
wavelengths as well as for the pump beam. Therefore, it is supporting the SRS experiments with sufficient intensity of pump light on the SRS detection photodiode. After the condenser lens a multichannel imaging system is constructed through a telescope \( f_{I1} = 300\,\text{mm} \) and \( f_{I2} = 180\,\text{mm} \) and the beam splitter BS2 (AHF, T750SPXRXT) in order to separate the pump beam signal for SRS imaging from CARS, SHG and fluorescence signals, which are blue-shifted. Here, two planes are introduced in the microscope separating the wavelengths also by two lightproof optical breadboards into two branches offering space for several detectors and optical elements (fig. 22). A PMT detector array composed of three PMTs placed in a cage system offers three simultaneous acquisition channels for CARS, SHG and fluorescence signals. It is possible to exchange the optical filters and beam splitters (BS3 and BS4) very rapidly according to the desired wavelength separation. Due to the use of an additional telescope after the condenser lens the detected signals are non-moving on the detector area granting an even illuminated image. In the second area the pump beam is detected with an additional lens \( L_{I3} \) \( f_{L_{I3}} = 50\,\text{mm} \) on a fast silicon photodiode in combination with an electronic filter set and a lock-in amplification scheme (see section 3.1.6). The fundamental Stokes beam is filtered out by a 950\,nm short-pass filter after the lens \( L_{I1} \) in order to avoid this wavelength in any detection area. In total the microscope has four forward detection channels (including the SRS channel) and one epi-detection channel, which can be utilized all simultaneously. The typical CARS wavelengths (in the case of the \( CH_2 \) resonance it is \( \lambda_{CARS} = 660\,\text{nm} \)) are filtered by two 775\,nm short pass filters (AHF, F75-775) to eliminate the excitation wavelengths and by a 514\,nm long pass filter (Semrock, Razor Edge LP02-514 RU-25) to suppress second harmonic signals. The remaining signal is focused onto customized fast PMTs (Hamamatsu Photonics, H 9656-20 and H 9656-20 MOD trans-impedance 20k\,\Omega, having 0-5MHz bandwidth), which are connected to the A/D converter using the PMTs current-voltage-converter. The focal position of the excitation beams in the sample and the corresponding detected signal are combined by ScanImage to the final image. The Matlab-based software provides different scanning speeds and binning factors for the user as well as zooming into full field of view by a specified magnification ratio while selecting regions of interest in the sample.

Due to the arrangement of the focusing objective lens and the condenser lens in a \( 4\pi \) configuration it is necessary to move the condenser lens in xyz-direction with high precision without any mechanical instabilities especially when replacing the point detectors (PMT or photodiodes) with a camera (see also section 3.2). Therefore, a customized condenser lens xyz-mount proved to be necessary (fig. 22). It was constructed based on a 60\,mm cage system utilizing a modified xy-tanslation stage offering \( \pm 1\,\text{mm} \) range
(Thorlabs, CXY2) in combination with a self developed z-translation stage composed of two pairs of linear bearings (Schneeberger, RN 3-075-DR-SQ) for high stability and minimal thermal drifts. Here, a range of 25 mm was realized enabling easy replacement of the sample as well as the option to replace the condenser lens with an objective lens. A differential micrometer screw again offers high precision while setting the z-position.

**Figure 21 – Detection modes and optical path**

The scanning microscope is set up in a $4\pi$ configuration of focusing objective lens (Obj.) and condenser lens (Cond.) for the analysis of the sample, which is placed in between on a motorized 3D-stage. In epi-detection a PMT for CARS or a CMOS camera for bright field imaging are set up. In forward detection two lightproof areas for SRS (upper area) and blue-shifted signals, e.g CARS, SHG or two-photon excited fluorescence are offering space for multiple detectors. In the CARS area an array of three PMTs are used to detect up to three different wavelengths at the same time. The photosensitive area of all detectors are illuminated using a diminutive telescope arrangement in order to avoid imaging artifacts due to vignetting effects during the scanning process. All optical signals can be split by beam splitters (BS2 to BS4) and are finally filtered in front of the detectors by optical filter sets.
For a high flexibility of all desired detectors the microscope has three detection areas. One for epi-detection using the optical table and two forward areas constructed with optical breadboards offering space for CARS and SRS detectors. The focusing objective lens is mounted on a z-linear stage, the condenser lens is placed in a home-built XYZ-holder. It offers high stability and low tilt by using two pairs of linear bearings for the z-translation. The implemented 60mm cage system holds a modified xy-stage, which was optimized by setting in two bronze bushings. The condenser lens is screwed inside by an thread adapter, which offers additional freedom in terms of working distance of the installed collecting optics. The shown inner part of the microscope has an additional lightproof housing with several removable covers.

### 3.1.6. SRS modulation and detection scheme

In the case of SRS imaging the 1064nm Stokes beam is intensity modulated (in general with $5 - 20 MHz$) by the AOM before it is focused into the sample (see [3.1.4]). For the highest sensitivity of the setup the modulation imprinted on the Stokes beam has a fixed phase relation to the $80 MHz$ repetition rate of the laser system and an approach to gain tunability of this phase between both frequencies is realized through a custom built frequency divider and phase shifter. The electronic seed signal of the laser system is amplified and fed into an array of home-made oscillators with phase delay loops consisting of Schmidt triggers and comparator circuits. Here, the frequency is divided to 20, 10 and 5MHz while achieving a tunability of phase from $0 - 2\pi$ regarding to the $80MHz$ repetition frequency. Then a commercial function generator (such as *Hameg*, HMF 2550 or *HP*, 8116A) is externally triggered by this signal (case of 5 and 10MHz) and then delivers the final frequency with an amplitude of $0 - 5V$ to the AOM driver.
module. At 20MHz no additional frequency generator is needed, due to the included local sine generation in the phase shifter unit. In the interaction with the sample photons are transferred from the pump to the Stokes beam according to the current intensity of the Stokes beam (see 2.2.4). The rate of change in amplitude compared to the pump beam intensity ($\Delta I_{SRL}/I$) is in the range of $\ll 10^{-4}$, while the possible sensitivity achieved with current setups is $10^{-7}$ to $10^{-8}$ utilizing lock-in techniques (Freudiger et al. [2008]). The pump beam is transmitted to a large area photo diode (e.g. Hamamatsu Photonics, FDS 1010 or Luna Optoelectronics, PDB-C613-2) using a 950nm short pass filter (AHF, F37-950) to suppress the 1064nm beam and a 787nm long pass filter (AHF, F76-787/LP) (in case of probing the $CH_2$ resonance) to block out any remaining blue-shifted wavelength. The electronic signal of the photo diode is carefully filtered by a band-stop filter for the attenuation of the 80MHz native laser frequency as well as a $\pi$ band-pass filter, all constructed of inductive and capacitive elements (see fig. 23). The desired signal is coupled into a lock-in amplifier (APE, S/N S04757 or Zürich Instruments HF2LI) to perform a demodulation of the SRS signal with the reference signal, which is the same as the initial modulation-frequency of the 1064nm beam, provided as a separate TTL trigger signal by the frequency generator. Tunable settings at the lock-in amplifier are the preamplification of the incoming signal, the integration time and the phase of the demodulator, which is set automatically in the case of a digital lock-in amplifier (e.g. Zürich Instruments, HF2LI). These parameters determine the final speed of the imaging and are strongly dependent on the signal strength. Finally, the demodulated signal is amplified with a variable gain factor selected by the user outputting it to the A/D converter, where it is transformed into an image by ScanImage. Due to the very low signal to noise level of the desired electronic SRS signal generated by the photodiode any additional noise coupling into the signal path, e.g. from other electronic equipment, has to be avoided by careful electromagnetic shielding. Also the proper connection of the earth ground to all hardware in a star-like manner (the starting point was set on the optical table) without ground loops and the separation into case ground and signal ground proved to be crucial to a successful SRS experimental result (see fig. 24).
In order to explore the optimization of the silicon photodiode detecting the pump beam for SRS imaging the shown housing was developed placing the electronic filters in close proximity to the photodiode (e.g. Luna Optoelectronics, PDB-C613-2). Here, the BIAS voltage delivered by a battery pack is low-pass filtered in order to avoid the flow of the signal into that branch as well as to cancel out noise collected form the connection cable (not shown). The desired modulated signal itself is passing a band-stop filter suppressing the $80\, MHz$ signal of the laser repetition rate as well as a $\sim 22\, MHz$ low-pass filter suppressing system noise as well as signal reflexions or higher harmonics. All characteristics were tested and fine-tuned using the frequency scanning mode of the digital lock-in amplifier (see inset). Here, the transmission of the filter was tested and optimized in order to transmit the desired signal with $20\, MHz$ with low losses as well as to suppress the laser repetition rate and any other noise. Case ground and signal ground are separated for avoiding ground loops due to the mounting. For connecting the photodiode module to the lock-in amplifier a low loss, special shielded cable was used (SBB, Aircell 7). The connection from the battery pack delivering the BIAS voltage is a shielded twisted pair cable (LEMO, AWG28/19).
Figure 24 – Modulation and detection scheme of SRS imaging

The photodiode is provided by a low-pass filtered BIAS voltage enabling a fast rise time for the detection of the pump beam. The signal voltage is generated by the load resistor ($R_L = 50\Omega$) and then filtered by a band-stop filter and a low-pass filter, all inside the shielded photodiode module. After that the lock-in detector preamplifies the signal and demodulates it with the reference signal, which is also used for the amplitude modulation of the Stokes beam via the AOM. The imaginary as well as the real part of the demodulated signal are summed and amplified in order to be finally detected by the A/D converter card, which is operated by the imaging computer. The reference signal itself is created of the seed signal of the laser system, which has an $80\, MHz$ repetition rate. A frequency divider is utilized in combination with a phase shifting device to generate a phase shiftable $20\, MHz$ signal, which is addressed to the AOM as well as to the lock-in amplifier. Please note that the lock-in amplification scheme is simplified and thus valid for digital as well as analog systems. Also the pulsed laser characteristic is not included in the drawing.

3.2. Structured illumination for resolution enhancement

In the course of the thesis the application of structured illumination for the resolution enhancement in CARS microscopy (SI-CARS) was explored especially in the forward detection, whereas the first application of this idea was performed in the epi-detection as the task of the master’s thesis [Pilger 2014]. Here, it became apparent that the epi-direction offers too low CARS signal intensity for the detection via an sCMOS camera resulting in a beneficial resolution enhancement. Nevertheless, the application of structured illumination proved itself as a promising tool for the resolution enhancement beyond the diffraction limit. The appealing point is that in comparison to other possible approaches in a wide field arrangement the focal intensities of Stokes and pump beam are still high as well as in comparison to also focused approaches with beam profile shaping, a potential spatio-temporal mismatch of both beams is not occurring, granting the same high signal intensity as without the approach for resolution enhancement.
In addition to the SI-CARS experiments two-photon excited fluorescence is explored using the Stokes beam of the CRS-setup as excitation source. These fluorescent data sets, which are the result of incoherent image formation, were also generated to back up the prospect of resolution enhancement beyond the diffraction limit in coherent microscopy. For the implementation of SI-CARS a sCMOS camera (Hamamatsu Photonics, OrcaFlash4.0) was utilized as detector instead of the PMTs. It has a detector area of 2048 × 2048 pixels, while each pixel has the size of 6.5 × 6.5 µm. The camera’s quantum efficiency is ∼ 70% for the typical CARS signal wavelengths (e.g. 660 nm probing the CH$_2$ resonance) [Hamamatsu Photonics [2012]]. For a low noise performance the sCMOS chip is cooled by a Peltier element in combination with an external water cooling system. Using optimal conditions the readout noise is 1,3 electrons$^{10}$ having a dark current of 0.15 electrons/pixel [Hamamatsu Photonics [2012]]. In order to match the camera’s optimal working condition in terms of signal intensity per pixel as well as the optimal magnification ratio of sample size to image size the detection optics for PMT had to be changed. In order to keep the effort low for switching between the standard PMT detection and the detection by the camera a flip mirror mount (Radiant Dyes, RD-KLS-1) was inserted in the optical path. It allows to address the camera without changing the entire beam path (see fig. 25). Only the first lens behind the collecting objective lens had to be replaced entirely ($L_{11}$ to $L_{15}$). The condenser lens itself had to be replaced with an oil objective lens (Olympus, UPlanApoN 60×, NA 1.42), which has an infinity correction and, therefore, the better imaging performance compared to the condenser lens. The optical detection path for the camera had to be adapted as well. It is placed inside the CARS detection area of the microscope and consists of a telescope, comprised of two lenses with $f_{15} = 180$ mm and $f_{16} = 100$ mm (focal points set congruent, magnification of 0.56), which demagnifies the image first. Then a tube lens $f_{17} = 300$ mm in front of the camera is forming the final image on the camera chip. In combination with the magnification ratio of the objective lens (theoretical focal length is 3 mm) the total magnification ratio of these optics is 56, which leads to a imaging ratio of 116 nm per pixel on the camera chip.

$^{9}$Water cooling, water temperature of 20$^\circ$C.

$^{10}$Calculation: ratio of a full well capacity and the readout noise.
For the structured illumination approach the condenser lens was changed to an oil immersion objective lens with high NA and 60× magnification. A new lens array comprised of $L_{I5}$, $L_{I6}$ and $L_{I7}$ is utilized to form the final image on the sCMOS camera with a pixel size of 116nm/px. It is folded by three additional mirrors (not shown) in order to stay on the available space. A suitable filter set cancels out the pump and Stokes beam, while additional bandpass filters are set in to isolate the signal wavelength of interest. A flip mirror mount is implemented to enable rapid changes between PMT and camera detection, which is possible due to the large free space area built into the microscope. Due to the clear separation of all involved wavelengths and the sound lightproof housing of the detection area low noise due to undesired stray light was present on the camera.
3.2.1. Generation of a striped illumination pattern

In the standard operation of the microscope during e.g. a CARS experiment both pump and Stokes beam are raster-scanned over the sample generating a blue-shifted signal of the sample at every position. The technique to perform a striped illumination of the sample with galvanometric scanning mirrors is based on the rapid intensity switching of one of the beams (here in particular the Stokes beam) by an acousto- or electro-optical device according to the position of the focus in the sample. Due to the already used line per line scanning mode of the microscope the Stokes beam was blanked out by the AOM at certain times, generating a CARS signal only at every third scanned line. The fineness of the scanned pattern as well as the angle orientation were set by the scanning control program. The pump beam was present on the sample at every position but did not affect the experiments in any negative way. For the precise and reproducible synchronization of the position of the scanning mirrors and the modulation of the Stokes beam a trigger scheme was developed.

As the fundamental timing signal the “frame clock trigger” was exported as a 5V TTL signal provided by ScanImage via the second timer channel of the A/D converter card. This signal is sent to the external trigger input of the sCMOS camera after a conversion is performed by a fast level converter to a 3.3V LVCMOS trigger signal matching the camera’s hardware requirements.

For the synchronization of the Stokes beam modulation the 5V TTL trigger signal is fed into a four channel digital delay / pulse generator (Stanford Research Systems, Inc., Mod. DG535), which is also used for the phase generation of the pattern. It provides four variable delay settings which are directed to the trigger input of a waveform generator (Hameg HMF 2550) by a external threefold toggle switch. A periodic rectangular waveform is generated by the waveform generator in the “burst” mode, while its timing is synchronized to the scanning process as well as its duration is matched to the chosen imaging parameters. Finally, the waveform is transferred to the Stokes beam intensity via the AOM (see fig. 26).

By switching the output channel of the delay generator a preset additional delay time is added to the starting point of the waveform generator. In additional measurements the delay times were determined leading to a response of the system, where three phases of the pattern were generated according to the selected channel of the delay generator. Summing up this scheme the waveform generator is triggered after the desired number of lines are scanned blocking the Stokes beam and then starts with the periodic waveform pattern releasing the Stokes beam onto the sample. For the calculation of the cycle duration of the waveform a multiple of the “true ms/line” value of the scanning program is used, which is the sum of the illumination time of one line and the moving
time of the scanning mirrors moving to the starting point of the next line. To specify the numbers of repetitions of one cycle of the waveform, the rate

$$\#\text{cycles} = \frac{\#\text{lines/frame}}{3} - 1 - \frac{1}{3}$$  \hspace{1cm} (56)$$

was identified as proper value for the chosen pattern spacing. The first term refers to one period covering three lines. Due to undesired movement of the scanning mirrors at the end of the frame one cycle was subtracted as well as the first scanned line, which is the result of electronic run-times. The “rotate” setting (defined as angle of rotation $\vartheta_{\text{rot}}$) of ScanImage allows the rotation of the pattern with 0.1° accuracy. In the final image series tree angles as well as three phases per angle are utilized granting a resolution enhancement in the x- and y-direction of the field of view.
The reference signal for the camera trigger as well as the AOM-based pattern generation is a 5V TTL signal provided by the A/D converter card, which is created after a frame in the imaging process is completed (left upper half). Before the camera’s external trigger is addressed a fast level converter transfers the 5V TTL signal to 3,3V CMOS logic matching the camera input (left side). The frame clock signal is also directed to a digital-delay-generator, which provides tree timing delays of the original signal selectable by a switch. These are used for the phase setting of the illumination pattern by feeding them into the external trigger input of a waveform generator (right hand side up to low). The waveform generator creates a rectangular periodic waveform, whose starting point is synchronized to the position of the scanning mirror. By adding a delay the starting point is shifted to the next line of the scanning process, which should be illuminated. The periodicity as well as the duty cycles and amplitude of the waveform are set according to the chosen pattern fineness, scanning speed and desired illumination intensity set in ScanImage. Finally, the waveform is directed to the AOM driver module and operates the AOM rapidly blocking and unblocking the Stokes beam.
4. Results

In the following chapter the capabilities of the microscopy setup will be demonstrated by a range of selected applications on different biomedical questions, which were investigated with several collaborators who performed the preparation of the samples as well as subsidiary experiments in order to support the information provided by the CRS-experiments. Here, also hyperspectral CARS data will be shown and discussed. In the course of the thesis the influence of pulse length variations of the OPO while wavelengths tuning during hyperspectral data acquisition was investigated and a customized Matlab-based control software was developed and implemented into the scanning software, which grants undistorted spectra. Following, the results of structured illumination applied on CARS microscopy for resolution improvement are presented and compared to data of two-photon excited fluorescence. Finally, a novel all fiber based CRS laser source is introduced, which was developed in close collaboration with the research group of Prof. Kenneth K. Y. Wong at the University of Hong Kong. The source will be characterized and data on several biological specimen with CARS as well as SRS and multiphoton applications such as SHG and two-photon excited fluorescence as contrast mechanism are provided.

4.1. Investigation of osteogenic differentiation in human stem cells

4.1.1. Exploring label-free CARS microscopy as suitable tool for visualizing the successful differentiation of ITSCs in clinical research

This collaborative work was performed jointly with the research group of Prof. Christian Kaltschmidt and Prof. Barbara Kaltschmidt, faculty of biology, Bielefeld University, in the course of the bachelor thesis of Mr. Arne Hofemeier [Hofemeier [2014]]. For the cure of injured bone structures in human patients, e.g. after an accident, the use of stem cells is a new promising treatment [Meza-Zepeda et al. [2008]], which delivers new bone material to the damaged structure enabling faster recovery and minimally invasive cure [Greiner, Johannes F. W. et al. [2014], Hauser et al. [2012], Greiner et al. [2011], Seebach et al. [2010]]. In this study the osteogenic differentiation of human neural crest-derived inferior turbinate stem cells (ITSCs) was investigated with CARS and SHG microscopy [Hofemeier et al. [2016]], while spontaneous Raman scattering was utilized for supporting the CARS data as well as Alizarin Red-Staining for fluorescent light microscopy in combination with rt-PCR (reverse transcription polymerase chain reaction) in order to confirm the results. Here, the ability of CARS and SHG microscopy as label-free and, therefore, non-contaminating tool was compared to clas-
tical methods of detection such as fluorescent labeling. There is high interest for a confirmation of the successful differentiation of the stem cells without a contamination or destruction of the specimen by the analysis method, because the very same sample could be used in the therapeutic treatment of the human patient assuring the desired effect of the stem cells.

4.1.2. Design of the experiments: CARS and SHG imaging of ITSCs and control group

The ITSCs are routinely isolated during minimally-invasive surgery and were expanded in vitro. The differentiation process resulting in the production of bone material was induced by applying a chemically-defined directed differentiation medium (containing dexamethasone, $\beta$-glycerophosphate and phosphate) or topological cues of a nanoporous titanium surface on the cells. The specimen were handled in custom-built 1-fold and 4-fold dwell holders, which enabled a sealed condition of the specimen as well as the use of quartz cover slides for spontaneous Raman spectroscopy. Here, different time points were selected for analysis and the cells of a dwell were fixed with a para-formaldehyde (PFA) solution for that specific day. The starting point of the experiments was defined as “Day 0”, while four dwells were initially triggered for the differentiation with medium containing differentiation supplements. Additional four dwells called “FCS control” were kept as a control system, which were treated with standard fetal calf serum (Sigma Aldrich, FCS) in the course of the experiment. The selected time points for analysis were chosen to be 0, 7, 14 and 21 days of differentiation. First bright-field microscopy images were acquired for each sample on a separate spontaneous micro Raman spectroscopy setup [Oberländer 2016] and interesting structures were selected. Areas were biomineralization takes place can be identified due to the star-like shape of the surrounding cells, which express the calcium hydroxyapatite in combination with collagen. At these points spontaneous Raman spectra were acquired for the comparison to the following CARS and SHG imaging, while an Alizarin Red staining was performed in the end on the same sample dwell. For CARS and SHG imaging in total three different channels were acquired sequentially to collect information of the specimen utilizing the 60× water objective lens focusing into the sample with a total focal power of less than 30mW for CARS. The generated signals were collected by a 40× air objective lens (Olympus, LUMPlanFLN, NA = 0.8), while all different signal contributions were isolated with narrow-band optical filter sets (950$SP$, 775$SP$, 785$SP$, 514$LP$ for all channels in common) in order to solely detect

\[\text{Setup made by Dr. Elina Oberlander in the scope of her PhD-thesis, data acquisition by Mrs. Lena Nolte during her master thesis.}\]
The desired wavelengths. The $CH_2$ stretching resonance at $2845 cm^{-1}$ is predominantly connected to lipids and was probed by the 1064 $nm$ Stokes beam and a 816, 7 $nm$ pump beam. This channel is utilized for the visualization of the ITSC cell structure, while low signal levels were assigned to the confluent cell layer and the unspecific non-resonant background, thus shown via thresholding in a separate color (see fig. 27, deep red). Higher signal levels originating from lipid accumulations, such as lipid droplets, are shown in cyan. This CARS signal occurring at 660 $nm$ was additionally filtered by a 660/40 band-pass filter. In addition to that the $PO_4^{3-}$ symmetric stretching mode at 959 $cm^{-1}$ was probed for the identification of calcium hydroxyapatite depositions (magenta). Here, both OPOs were utilized as laser sources directing 796 $nm$ as pump and 860, 6 $nm$ as Stokes beam into the sample while isolating the CARS signal at 740, 4 $nm$ by a tunable band-pass filter at 740/20 $nm$. In the last step SHG imaging (green) was performed with the 1064 $nm$ laser beam generating the frequency doubled signal at 532 $nm$ of the collagen secretion by applying 25 $mW$ focal power. This procedure was repeated also on the control samples in order to prove that the response of the ITSCs is depending solely on the differentiation medium (fig. 28). For backing up the claim of successfully imaging the calcium hydroxyapatite deposits via CARS an additional control experiment was performed on a human skull bone of the mastoid cortex (see fig. 29). In order to prove the applicability of SHG microscopy detecting collagen a second additional experiment was performed on a mouse tail specimen (fig. 30), which is an established sample of oriented collagen fibers. Due to the infrared wavelengths regime and rather long pulse length in the picosecond regime of the 1064 $nm$ laser source no disturbing two-photon excited autofluorescence signal was detected. For the comparison of all different signal channels the data were normalized with respect to the intensities used as excitation laser beams, signal integration time and PMT amplification factor. The images with highest contrast (day 21) was normalized to a value of 1000 and serves as overall reference for all channels and days. In addition to that small pixel shifts due to the subsequent imaging of the channels were corrected by pixel-wise manual shifting until the non-resonant signal contents present in all channels were matched via a custom-written Matlab program, which determines the best correlation between the images.
4.1.3. CARS and SHG data are compared to spontaneous Raman spectra and Alizarin Red staining

Comparing the results of spontaneous Raman spectroscopy as well as images by Alizarin Red staining to the CARS and SHG imaging, we find a Raman peak of the calcium hydroxyapatite after 14 and 21 days of differentiation (fig. 31), while no distinct spectra were present in the control cell samples. This is in compliance with the data acquired via the fluorescent staining, here characteristic calcium deposits could be identified on days 14 and 21 (fig. 32). CARS and SHG imaging were in good accordance with the prior results. Here, after 14 days of differentiation calcium hydroxyapatite deposits were identified probing the $959\, cm^{-1}$ resonance co-localized with SHG signal indicating the presence of collagen type I. At day 21 the clear increase of both signals were detected, while in an additional 3D scan of the sample at day 21 (fig. 33) also the volumetric structure of the calcium hydroxyapatite deposits and collagen was investigated. For this purpose a z-scan at 26 different positions was performed via the piezo stage using a $1\, \mu m$ step size, while at each z-position all three different CARS and SHG channels were acquired consecutively. The mineral deposits were observed to be localized external and superior to the cell layer surrounded by a collagen matrix formed of the collagen secretion of the ITSCs.

4.1.4. CARS and SHG imaging leads to successful visualization of the ITSCs differentiation also applicable to clinical research

In conclusion of this work the label-free and non-invasive CARS and SHG imaging modality is a notable alternative to the established methods of fluorescent staining in order to detect the successful differentiation of the ITSCs showing the similar sensitivity as the destructive counterparts while adding beneficial structural information of the sample by providing imaging data in three dimensions. Future clinical application is also in reach taking new endoscopic approaches into account [Lukic et al. 2017, Lombardini et al. 2018], which could possibly enable investigations even in vivo. Therefore, the presented results indicate that the diagnosis of ossification-related diseases investigating bone mineralization density distributions of patients can be performed without the need of biopsies in the future.
Figure 27 – CARS and SHG imaging of the ITSCs showing the osteogenic differentiation

After the induced osteogenic differentiation process of the ITSCs at day 0 only the presence of the ITSCs is visible while probing the $2845\, cm^{-1}$ $CH_2$ stretching resonance for the visualization of the cellular borders (red) and lipid droplets (cyan) inside the cells via thresholding. Also after 7 days of differentiation no signals were recorded probing the $PO_4^{3-}$ stretching mode at $959\, cm^{-1}$ for calcium hydroxyapatite depositions (magenta) and SHG signals (green) indicating the presence of collagen type I secretions. At day 14 calcium hydroxyapatite deposits and also co-localized collagen secretions (arrows) were recorded, while at day 21 the signals increased in both channels and show the ongoing differentiation process with a higher amount of the deposits outside the cells compared to day 14. On the right hand side all channels are overlapped, showing the position of cells and secretions. The scale bar indicates $20\, \mu m$. Published in [Hofemeier et al. [2016]].
Figure 28 – CARS and SHG imaging of the ITSCs under control conditions showing no deposition of calcium hydroxyapatite or collagen fibers

The control samples, which were treated with standard fetal calf serum (Sigma Aldrich, FCS) instead of medium containing differentiation supplements, can also be observed via CARS probing the $CH_2$ resonance (red and cyan). In contrast to the triggered cells they show no signs of calcium hydroxyapatite depositions probing the $PO_4^{3-}$ mode via CARS (magenta) and no traces of collagen due to the absence of any SHG signals (green) during the ongoing experiment. The scale bar indicates 20µm. Published in [Hofemeier et al. 2016].
Figure 29 – Applying CARS and SHG imaging on a human skull bone for the visualization of calcium hydroxyapatite and collagen fibers

A piece of human mastoid cortex was imaged for investigating, if CARS microscopy can be utilized to show the calcium hydroxyapatite probing the 959 cm\(^{-1}\) resonance (magenta) applying 20 mW focal power for the 860 nm pump and 6 mW focal power for the Stokes beam, respectively. The SHG signal (green) is utilized for identifying collagen in the bone structure by detecting the frequency doubled signal of the 1064 nm laser beam (42 mW focal power were applied). By overlaying both channels we see the combined structure of the bone material, collagen structures fit into the frame formed by the calcium apatite. The imaged area of the specimen was an edge of the bone touching the cover slide. The scale bar indicates 25 µm. Published in [Hofemeier et al. 2016].

Figure 30 – SHG imaging of a mouse tail fiber sample showing the presence of collagen

For the visualization of the mouse tail fiber the CARS signal at 1662 cm\(^{-1}\) (red) was probed in addition to the SHG signal (green) at 532 nm utilizing 50 mW focal power of the 1064 nm laser beam, which shows the collagen with a typical fibrillar structure. The CARS signal is rather unspecific compared to the SHG signal. Note that for this CARS measurement two 842 SP filters were replacing the 775 SP and 785 SP filters. The scale bar indicates 30 µm. Published in [Hofemeier et al. 2016].
The differentiated as well as the control cells were imaged via bright-field microscopy on the spontaneous Raman spectroscopy setup identifying potential areas of calcium hydroxyapatite secretions. This was possible due to the characteristic star-shaped arrangement of the cells to each other at areas of deposits. In the center of the arrangement often the secretion can be found, here the spontaneous Raman spectra were acquired. a) shows the comparison of differentiated (purple) and FCS control (green) ITSCs at day 0, where no distinct peaks could be observed. The same behavior was found in b) at day 7. At day 14 (c) a peak at 959 cm$^{-1}$ corresponding to the $PO_4^{3-}$ stretching mode was found in the differentiated cell sample, while no distinct spectra could be recorded for the FCS control sample. This result was detected with increased intensity of the 959 cm$^{-1}$ peak at day 21 (d), while again the control sample showed no traces of peaks at all. The Raman spectra were acquired using 785 nm continuous wave laser light not exceeding 25 mW of focal power utilizing a 60x water objective lens. Adapted from Hofemeier et al. [2016].
In the final step a fixation by a 4% PFA solution was performed on all samples. Then the Alizarin Red staining was applied to the samples using 1% of Alizarin Red (Waldeck) in ddH$_2$O in order to stain specifically the calcium hydroxyapatite. The samples were imaged with a combined fluorescence and bright-field microscope (PeqLab, AMG EVOS xl microscope). The above shown images are the combination of both imaging modalities, here showing the differentiated ITSCs from day 0 to day 21. Similar to the results of spontaneous Raman spectroscopy as well as CARS and SHG at day 14 first signals of the Alizarin Red staining are observed (white arrow) indicating the presence of calcium hydroxyapatite. At day 21 a higher signal strength in the fluorescence channel was recorded originating from larger calcium hydroxyapatite secretions. The FCS control samples exhibit no fluorescent signals (data not shown), which is in good accordance to the results of all applied techniques. The scale bar shows 100µm. Adapted from [Hofemeier et al. 2016].

In a 3D scan performed on the ITSC sample after 21 days of differentiation a large crystal of calcium hydroxyapatite surrounded by the collagen matrix was imaged using a step size of the piezo stage of 1µm covering 26µm in z-direction. Each channel (lipids: cyan, calcium apatite: magenta, collagen: green) was recorded sequentially and then the z-position was altered. The imaging software FIJI was used for the visualization of the data, enabling different angular views. We observe that the mineral deposits are localized externally above the cell layer and are surrounded by the collagen matrix. This sample was stained with Alizarin Red when the large crystal was observed. Hence, a portion of the SHG signal can also arise from the staining, but the localization of both CARS and SHG channels are in good agreement. The scale bar depicts 25µm. Published in [Hofemeier et al. 2016].
4.2. In vivo analysis of intracellular lipid droplets in *Monoraphidium neglectum*

4.2.1. Identification and characterization of lipid rich microalgae as non-food feedstock in bio-diesel production

Facing the worldwide need for energy in respect to the turning away from fossil fuels raises an increased interest in ways to produce renewable energy sources. In the background of the worldwide climate change caused by the human induced emission of global warming gases using bio-material for the power generation in bio-gas plants or the biomass to fuel conversion holds great potential for a positive influence on the carbon footprint [Rupprecht [2009], Medipally et al. [2015]]. On the other hand the direct use of food-related bio-material or the indirect extrusion of acreage is raising new problems in the worldwide nutrition while the world population is increasing [Schenk et al. [2008]]. Hence, a non-food feedstock as a sustainable source for bio-material generated on non-agrarian used areas of the world are of high interest facing the future production of regenerative energy via biomass conversion. In this field one promising candidate meeting these requirements is the harvesting of oleaginous photosynthetic microalgae [Sheehan et al. [1998]], which can be grown in oceanic tanks where they produce lipids by the conversion of nutrients in the course of the photosynthesis [Mata et al. [2010]]. The final algal lipid yield and fatty acid content of the algae at the time of harvesting are important characteristics for the choice of the best type of microalga, which influence the economic feasibility in the production of e.g. bio-diesel utilizing the lipid content [Hu et al. [2008]]. Here, especially the ratio of neutral lipids such as triacylglycerols (TAGs), which are the main components of lipid droplets (LDs) in the cell, versus polar lipids such as membrane glycerolipids, mainly located in the cellular membrane, are a key characteristic of the successful application of microalgae in the fuel production. The TAGs are the desired form of lipids for the generation of bio-diesel and can be most easily extracted in the subsequent processing from the algae when they are accumulated in LDs.

4.2.2. Established methods for analyzing the lipid content of microalgae are laborious and partly not feasible due to rigid cell walls

The formation of the lipid droplets is dependent on the different environmental conditions of the microalgae, which affect their metabolism [Bogen et al. [2013a]]. Here, a common method to trigger the formation of LDs in microalgae is an increased illumination intensity as well as nitrogen starvation conditions. The TAGs content is routinely quantified by solvent extraction in combination with fractionation and gravi-
metrical analysis. Also HPLC (high performance liquid chromatography) or GC-MS (gas chromatography - mass spectrometry) are established tools for the quantification of the fatty acid composition of the algal lipids, but require a not negligible amount of biomass in combination with extensive work protocols and high time consumption. Faster alternatives are specific fluorescent staining, e.g. Nile Red, labeling the lipids of interest for fluorescent detection in combination with flow cytometry or fluorescent activated cell sorting. However, the major drawback of this approach are massive obstacles in the successful staining of the intracellular lipids due to the rigid cell wall of many microalgal species, which act as an impenetrable barrier for the fluorescent dyes.

4.2.3. Raman-based techniques speed up the analysis process of M. neglectum

Label-free alternatives such as Raman micro-spectroscopy can be utilized for the investigation of the relative degree of saturation of the lipid content. For imaging purposes CRS-microscopy is a powerful alternative to the established methods, especially when fluorescent staining is not working. Recently a proof-of-principle study using CARS microscopy was conducted on the microalga Coccomyxa subellipsoidea C169 [He et al. 2012], where the CARS signal of interest had a significant overlap with the two-photon excited fluorescence of the chlorophyll hindering the analysis of the recorded data. In the diatom Phaeodactylum tricornutum the undesired two-photon excited fluorescence was separated from the CARS signal via time gated detection [Cavonius et al. 2015], due to the fact that the fluorescence signal is delayed a few nanoseconds compared to the instantaneous Raman and CARS signals.

In the following collaboratively obtained results with the research group of Prof. Dr. Olaf Kruse (Bielefeld University) are presented [Jaeger et al. 2016] concerning the analysis of the intracellular lipid droplets in the oleaginous microalga Monoraphidium neglectum utilizing CARS microscopy, also as a part of the PhD-thesis of Dr. D. Jäger [Jaeger 2017]. In prior work the microalga M. neglectum was identified as a promising candidate for bio-fuel production but also in having a rigid cell wall, excluding fluorescent staining [Bogen et al. 2013b]. Therefore, the label-free CARS microscopy was the selected solution circumventing the obstacles of fluorescence-based microscopy or laborious and time consuming studies via gravimetical analysis. For further quantitative determination of the degree of saturation spontaneous Raman microscopy [Oberländer 2016][12] was utilized on selected lipid droplets inside the algae.

4.2.4. Design of experiments: modulation approach excludes two-photon excited fluorescence enabling CARS imaging of lipid droplets

The study evaluates the potential of CARS microscopy for imaging the LD formation in M. neglectum under nitrogen starvation conditions, while probing the $CH_2$ stretching mode at $2845\text{cm}^{-1}$ using $11,5\text{mW}$ focal power of the $1064\text{nm}$ beam as Stokes and $23,5\text{mW}$ focal power of the $816,7\text{nm}$ wavelength as pump beam. The feedstock of cells selected for the experiments were held under nutrient replete conditions and at the start of the study they were diluted to a low cell density ($\sim 4 \cdot 10^6 \text{cells ml}^{-1}$) and divided into three reservoirs to generate three different replicates. In a first step a strong two-photon excited autofluorescence signal, which was also present in the CARS signal occurring at $660\text{nm}$, was detected over a large range of wavelengths. In order to suppress this undesired fluorescent background contribution originating from chlorophyll inside the chloroplast of the alga, the following filter set was applied: $950\text{SP}$, $775\text{SP}$ (two times), $785\text{SP}$ for the suppression of the pump and Stokes laser wavelengths as well as a $514\text{LP}$, $660/40\text{BP}$ and a tunable band-pass filter ($Semrock$, $TBP697/13$, $\sim 13\text{nm}$ transmission range) set to $660\text{nm}$ transmission for the isolation of the CARS signal. Still, the fluorescence signal was present in the CARS images, while the main source for the fluorescence excitation wavelength was identified to be the shorter, higher energetic pump wavelength at $816,7\text{nm}$ (see fig. 34 a) Stokes and b) pump beam).

Due to the rather low background signal originating from the Stokes beam an intensity modulation scheme was utilized in order to divide the signals of the CARS channel into the modulated CARS signal contribution and the static fluorescence signal via lock-in demodulation (similar to [Garbacik et al. 2013]). For this purpose the Stokes beam was intensity modulated with a frequency generator ($HAMEG$) and the AOM at $3\text{MHz}$ using the specifically modified high speed PMT ($Hamamatsu$, H 9656-20MOD transimpedance $20k\Omega$, $0 - 5\text{MHz}$ bandwidth$^{13}$) for the CARS signal detection. Thus the fluorescence background was mostly eliminated from the CARS channel (fig. 34 b, d) by the demodulation using the lock-in detector ($Zurich Instruments$, HF2LI), which served as narrow band electronic filter at $3\text{MHz}$ using an integration time constant of $20\mu s$. The signal strength was sufficient for a typical pixel dwell time of $31,25\mu s$ recording a CARS image (see fig. 34 f). Nevertheless, an additional image with solely the Stokes beam present in the sample was acquired sequentially and subtracted from the corresponding CARS image for excluding any fluorescence contribution originating from the Stokes beam for the quantitative analysis. In order to turn the undesired fluorescence signal into an advantage a second fluorescence channel was set up utilizing

$^{13}$The measuring resistor is changed by the manufacturer to $20k\Omega$ instead of $1M\Omega$ sacrificing sensitivity while gaining a higher bandwidth.
an 600\textit{nm} long-pass filter (\textit{Semrock}) in front of the band-pass and long-pass filter sets, which was detecting the chloroplasts distribution via a second PMT (see fig. 35).
Figure 34 – Modulation approach for eliminating undesired two-photon excited fluorescence signals investigating M. neglectum

The left column (a, c, e) show the CARS channel probing 2845 cm\(^{-1}\) while no modulation of the Stokes beam is performed. In a) solely the Stokes beam (1064 nm) is directed onto the sample, in c) the pump beam (816, 7 nm) respectively. An undesired two-photon excited fluorescence signal is visible in both channels, while its intensity is most prominent in c. Hence, the corresponding CARS signal (e) is disturbed by the fluorescence signal and has become unspecific excluding a quantitative analysis of the lipid droplets. The modulation of the Stokes beam at 3 MHz in combination with the demodulation of the CARS signal by a lock-in amplifier (right column, b, d, f) shows significant less fluorescence signal while applying only Stokes (b) or pump beam (d). The resulting demodulated CARS signal (f) is specific to the lipid distribution again. For the exclusion of any artifacts due to the rather low fluorescence residuals of the Stokes beam an additional image (only Stokes beam in the sample) was acquired and subtracted form the corresponding CARS image. The data was recorded after 1 day of nitrogen starvation. Note that figures (a - d) as well as (e - f) have separate look-up tables, the scale bar depicts 5 \(\mu m\). Published in [Jaeger et al. 2016].
Figure 35 – Two channel detection of CARS signal and two-photon excited fluorescence
For the separation of both channels into: 1) CARS signal (red) probing the 2845 cm$^{-1}$ lipid resonance and 2) the two-photon excited autofluorescence signal (green) of the chlorophyll an additional 600LP optical filter was utilized for the detection of the fluorescent signals on a second PMT. Both channels were recorded sequentially. For the visualization of the lipid droplets and the chloroplast a z-scan was performed having a step size of 500 nm covering up to 12.5 µm range. The two-photon excitation was performed by the pump beam at 816 nm applying 23 mW focal power. A three-fold zoom was used for these images, the scale bar depicts 10 µm. a) and b) show a single 2D frame of days 2 and 3 respectively, while c) shows a full 3D scan of day 3 using the visualization tool volume viewer of FIJI. A Gaussian blur was applied on the data to smooth the 3D view. The chloroplast is visible as a large spherical component surrounded by the lipid droplets.
4.2.5. Comparing the results concerning the lipid content acquired by volumetric CARS imaging to established methods

The experiments are conducted over a time course of 8 days beginning the nitrogen starvation at day 0. On days 0, 1, 2, 4 and 8 CARS microscopy was conducted on the samples, while several microalgae were visible in one acquired image using a zoom-factor of $3 \times$ for having a field of view of $48,5\mu m$ (fig. 36). For the comparison of data acquired by CARS microscopy to conventional methods the M. neglectum samples were investigated by cell counting, dry weight measuring (fig. 37a) and gravimetrical neutral lipid determination (fig. 37b). The dry weight increases constantly over time while the cell density doubles after 4 days of starvation and remains constant afterwards. The lipid content of the biomass is separated by gravimetrical analysis into total-, polar- and neutral- lipids, where the polar lipid content decreased continuously at the first two days and is observed as constant afterwards. The neutral lipid content increases constantly over the whole time course and reaches up to 30% of the dry weight at day 8.

Then the quantitative analysis of the cellular volume as well as the volume of the lipid droplets via CARS microscopy was explored in two different approaches: a high number of replicates was imaged by CARS imaging using arbitrary z-positions at random locations in the specimen for the counting of LDs in the cells (fig. 38a). The number of analyzed cells per day were $n = 90$. This leads to a measure of equal or greater number of lipid droplets per cell than the number detected in the specific section. We observed an continuously increasing number of LDs, after 8 days of nitrogen starvation the mean number of LDs per cell section increased from 0 at day 0 to 15 at day 8. Also the diameter of the observed LDs is approximated from the data via assuming the LDs as circular shapes and measuring the diameter with FIJI [Schindelin et al. 2012, Rueden et al. 2017] for plotting the data in respect to the days of nitrogen starvation (fig. 38b). Here, the mean of the diameter is slightly increasing with time while the mean variation is increasing (number of investigated lipid droplets $n = 0, 468, 969, 1255, 1363$ for day 0, 1, 2, 4, 8). In addition the cell area (elliptic shapes were assumed, fig. 38c) was estimated and via the LDs diameter the corresponding LDs areas were calculated for finally plotting the relative lipid droplets area by dividing the LDs area by the total area of the cells (fig. 38d). A five-fold increase over the time of starvation is observed while the mean cell area remained constant. Comparing the results of CARS microscopy in terms of relative LD area to the gravimetrical determined neutral lipid contents shows a strong correlation (fig. 38e, $R^2 = 0, 9981$) and indicates good agreement of both methods.

An alternative and more intuitive way of analyzing the lipid content and cellular vol-
ume of M. neglectum would be directly measuring the volume of the specimen instead of the area of the LDs and cells via single, arbitrarily selected z-positions. Therefore, 3D imaging was performed using the piezo-stage in the z-direction applying a step size of 500 nm on a separate set of cells originating from the same batch of M. neglectum. Then the acquired data were analyzed assuming the cells volume as an ellipsoid with $Vol_{Cell} = \frac{4}{3} \pi \left( \frac{d_x}{2} \right) \left( \frac{d_y}{2} \right) \left( \frac{d_z}{2} \right)$ and the volume of the lipid droplets as a sphere $Vol_{LD} = \frac{4}{3} \pi \left( \frac{d_{LD}}{2} \right)^3$, see also fig. 39 a). Here, the LD volume (fig. 39 b) as well as the relative LD per cell volume (fig. 39 c) are plotted over time and show the similar tendency as the data obtained analyzing the area instead of the volume. Finally, this results again in a strong correlation to the gravimetrical neutral lipid analysis (fig. 39 d, $R^2 = 0.98$), but is not perfectly overlapping to the area analysis method (compare to $R^2 = 0.9981$). The reason for the slight mismatch of both methods very likely originates from the difference in the number of cell replicates, which also shows the drawback of the 3D method: a higher time consumption compared to area analysis results in the rather low number of replicates (investigated number of LDs $n = 0, 46, 121, 106, 205$ at days 0, 1, 2, 4, 8, respectively). For a statistical more solid comparison of both methods in terms of the biological conclusion a far greater number of replicates would be required in the 3D approach, which was not further pursued in the course of the study, due to the good results of the 2D method matching the gravimetrical neutral lipid analysis and being very efficient in time consumption.
Figure 36 – Frequency modulated CARS imaging reveals lipid droplet formation in M. neglectum under nitrogen starvation

M. neglectum was imaged via frequency modulated CARS microscopy over 8 days of nitrogen starvation conditions (a - e). f) shows the merged signals of CARS signal (magenta) probing $2845 \text{ cm}^{-1}$ and two-photon excited fluorescence of the cup-shaped, central chloroplast (green). Increasing lipid droplet formation can be clearly observed and further analyzed by z-scanning. g) shows typical data acquired at day 8 at selected z-positions indicating the intracellular lipid droplet distribution. Note that the green circle depicts the cellular border to guide the eye. The scale bar indicates 5$\mu$m. Published in [Jaeger et al. 2016].
Figure 37 – Cell growth parameters and gravimetric analysis of M. neglectum during nitrogen starvation

(a) shows the cell density (blue rectangles) determined by a cell counter (Beckmann Coulter, Coulter counter) and the dry biomass weight (red circles, 25ml of the cell culture were washed, dried and weighed) over eight days of nitrogen starvation conditions. The error bars represent the standard error of the mean. In (b) the lipid content of the biomass is shown over 8 days of nitrogen starvation investigated by Folch extraction [Folch et al. 1957] of the lipid content and then separated by gravimetric analysis into polar- (depicted in green, PLF), neutral- (yellow, NLF) and total lipids (gray, TLF). The significance was tested by a two-sided t-Test (n.s.: not significant, ***: p < 0.001). Published in [Jaeger et al. 2016].
Figure 38 – 2D CARS imaging of M. neglectum for investigating cell growth and lipid distribution under nitrogen starvation conditions

(a) shows the number of LDs per cell section at random selected z-planes over time \((n = 90\) cells per day) in a box-whisker plot. In (b) the diameter of the LDs is measured via FIJI and averaged, the sample size was \(n = 0, 468, 969, 1255, 1363\) for days 0, 1, 2, 4, 8. The cell area \(A\) was also measured applying \(A = \pi \cdot \frac{d_x}{2} \cdot \frac{d_y}{2}\) where \(d_x\) and \(d_y\) depict the long and short diameter of the cell, respectively.

(c) shows the mean values of the cell area corresponding to days 1, 2, 4 and 8. In (d) the relative LDs area per total cell area is plotted in respect to the days of nitrogen starvation as indicator of the TAG content. (a - d) box whisker plots: mean value indicated by the thick black lines, the gray boxes show the interval between the first and third quartile while the two whiskers mark the respective 1.5 fold interquartile ranges (open circles: outliers). The significance of changes is indicated in regard to the value of the previous time point by a two-sided Wilcoxon Rank Sum Test (n.s.: not significant, *: \(p < 0.05\), **: \(p < 0.01\), ***: \(p < 0.001\)). In (e) the correlation of the relative LD area to the gravimetric neutral lipid content is shown (error bars represent the standard error of the mean, determination coefficient \(R^2 = 0.9981\)). Adapted from [Jaeger et al., 2016].
Figure 39 – Volumetric CARS imaging of M. neglectum for monitoring cell growth and cellular lipid content

a) illustrates the analysis strategy applied in the volumetric CARS data acquired by z-scanning with a step size of 500 nm. The cellular volume $V_{\text{cell}}$ was estimated as ellipsoid, the LDs volume $V_{\text{LD}}$ as circular sphere utilizing several z-slices of the scanned range for the determination of the values of each cell. This analysis leads to b) plotting the LDs volume during the 8 days of nitrogen starvation and c) the relative LDs volume, respectively (cellular volume not shown). (b and d) box whisker plots: mean value indicated by the thick black lines, the gray boxes show the interval between the first and third quartile while the two whiskers mark the respective 1,5 fold interquartile ranges (open circles: outliers). The significance of changes is indicated in regard to the value of the previous time point by a two-sided Wilcoxon Rank Sum Test (n.s.: not significant, *:$p < 0.05$, **:$p < 0.01$, ***:$p < 0.001$).

d) shows the correlation of the calculated relative LDs volume in respect to the cellular neutral lipid content determined by gravimetric analysis. The data correlates with a determination coefficient of $R^2 = 0.98$, the error bars indicate the standard error of the mean. Note that the number of investigated LDs ($n = 0, 46, 121, 106, 205$ at days $0, 1, 2, 4, 8$, respectively) is much less than in the 2D approach, previously shown in fig. [38]. Hence, the statistical significance and biological conclusion is less sound and these 3D results should mainly depict an alternative methodical approach. Published in [Jaeger et al. 2016].
4.2.6. Studying the degree of unsaturation of fatty acids in lipid droplets of M. neglectum by Raman micro-spectroscopy

Raman micro-spectroscopy was applied to investigate the degree of unsaturation of fatty acids in the lipid droplets utilizing the label-free, non-invasive manner of the technique. Therefore, LDs were identified and analyzed by spontaneous Raman scattering for comparing the ratio of two selected peaks, namely $1660\text{ cm}^{-1}$ ($C\equiv C$ double bond) indicating unsaturation as well as the $1442\text{ cm}^{-1}$ peak ($C-H_2$ vibration) as a measure of the overall lipid content. The analysis showed the ratio as a relative degree of unsaturation to be unchanged between day 4 and 8, which is in good agreement with results using GC-MS and fatty acid methyl ester derivatization. Further time points were not investigated, but the method proved as versatile tool and can be further explored in the future. The corresponding data sets will not be included into this thesis but can be found in the published manuscript [Jaeger et al. [2016]].

4.2.7. Successful CARS imaging of lipid contents in M. neglectum under nitrogen starvation conditions indicates optimal harvesting time point enabling comparison to other types of microalgae

Summing up the results obtained on the oleaginous microalga M. neglectum it was possible to utilize CARS microscopy for imaging a statistical significant number of replicates for the investigation of the relative lipid droplet content during a nitrogen starvation over eight days. Due to an intensity modulation of the Stokes beam in combination with lock-in demodulation it was possible to circumvent undesired two-photon excited fluorescence of the chloroplast, without the need of time-gated detection. Comparing the results obtained by CARS microscopy to classic approaches such as solvent extraction and gravimetric determination good agreement of the techniques is observed for the direct quantitative calculation of the TAG content. Also an optimal time point for the harvesting of the microalga M. neglectum could be identified to be 4 days after the start of the nitrogen starvation conditions. In addition to that the mean size of the LDs observed in M. neglectum was now accessible and could be related to other, already more characterized microalgae such as C. reinhardtii and P. tricornutum.
4.3. Developing a hyperspectral CARS imaging modality

4.3.1. Exploring hyperspectral CARS for enhancing the information content of samples by their spectral response

CARS microscopy is a versatile analysis tool for the investigation and visualization of one or more chemical components of a specimen, especially when the chemical components of interest are identified prior to the measurement, e.g. by spontaneous Raman spectroscopy [Huser and Chan [2015], Terjung et al. [2013]]. The received information of the images can consist of several probed CARS resonances or also SHG and fluorescence signal via wavelengths separation and modulation techniques as shown in the previous chapter. An extension of this methodical range in CARS microscopy gathering additional information of the specimen is the hyperspectral tuning of Stokes or pump beam, which are together addressing the molecular resonance [Fussell et al. [2013], Cheng and Xie [2015], Huang et al. [2018]]. In the typical technical implementation using ps-laser setups the beam delivered by the OPO is chosen for spectral tuning [Kong et al. [2013]], meaning the pump beam in the presented setup. An advantage of this method is that there is no need for prior knowledge of the chemical content of the sample and that the portions of the detected CARS signal originating from the overlay of the probed resonances can be further analyzed and interpreted by computational segmentation. Here, different methods such as principal component analysis (PCA) [Schie and Huser [2013]], non negative matrix factorization (NNMF) and other multivariate analysis tools [Miljković et al. [2010], Davis et al. [2011]] can be utilized to identify areas of similar spectral response of the sample and illustrate these with a color look-up table. The benefit of this step is an enhanced information content of the sample, which could not be achieved by probing single resonances via CARS. A drawback is the increased recording time needed of a full hyperspectral scan, e.g probing a typical wavelength range in the lipid resonance branch ($2700 – 3100cm^{-1}$) applying typical imaging conditions elongates the recording time from $\sim 20 – 60$ seconds for one image at a fixed resonance to up to 35 minutes for the entire hyperspectral scan. The scanning time is composed of the recording time of the image at the current wavelength and the time needed for the wavelengths tuning in the OPO system, which is dominated mainly by the rather slow heat-up or cool-down of the BBO crystal temperature determining the wavelengths working range.
4.3.2. Improving the stability of the microscopy setup enables high fidelity HS-CARS data sets

Due to the long recording time high demands are set to the microscopy setup in terms of focal stability. If the sample should move during the hyperspectral scan the gathered information is inconsistent and can not be used for further analysis. Therefore, great effort was taken to optimize all parameters of the setup, which are affecting the sample drift. Here, typical values of sample drift were less than 450 nm over 75 min in all dimensions. For achieving these values the condenser lens 3D-translation mount was optimized by a custom made mechanism and also the technique of dipping the condenser lens into the aqueous medium of the specimen without an additional cover slip proved to be advantageous. In typical measurements a water immersion objective lens focused Stokes and pump beam into the sample. The best focal stability over long terms was achieved by applying Immersol W 2010 (Carl Zeiss Jena GmbH) as immersion media instead of the standard distilled water due to the lack of evaporation and thus additional forces on the specimen.

4.3.3. Customized software controls entire HS-CARS scan process ensuring undistorted spectra

For the analysis of the hyperspectral data sets it is necessary to record the applied laser power of the OPO during the wavelengths tuning due to the varying response and efficiency of the LBO crystal and the OPO cavity. Addressing this need we choose to expand the task of intensity recording also to the Stokes beam for further improvement of the image quality of all recorded data. Hence, in the course of a bachelor thesis [Anselmetti [2016]] the intensity recording of Stokes and both OPOs signal exits was performed by home-embedded Si photodiodes (Luna Optoelectronics, PDB-C6132) and an additional A/D converter card (National Instruments, PCIe-6363) for the fast recording of the photodiode signals. For the time correlation of recorded laser intensities to the image data a custom-written Matlab program was created, which was capable to correct laser intensity fluctuations in a post-processing step up to 100 kHz, also in imaging data with fixed molecular resonance.

For the application to hyperspectral data this program was extended and further improved by implementing an auto-correlation functionality matching photodiode signals and CARS data in a more accurate manner as well as the development of control software for the hyperspectral scanning in the course of two master theses [Greife [2017], [Weiß [2017]]. For the tuning of the OPO’s wavelength a Matlab-based control program was designed (HyperSpectrum, see fig. 40a), which controls the OPO parameters via
the RS-232 interface of the OPO controller as well as it operates the microscope imaging software. Here, the LBO crystal temperature $t$ was precisely characterized and is set according to the desired pump wavelength for each image of the hyperspectral stack. The piezo stage of the OPO cavity determining the cavity length $L_{Cav}$ is addressed and optimized according to each selected wavelength. Finally, the simplified Lyot filter inside the OPO cavity is adjusted for the selection of the desired signal wavelength, which is monitored by the inbuilt spectrometer of the OPO. In general all described OPO parameters are optimized for the highest possible signal output intensity of the OPO according to the selected wavelength (then described as $L_{Cav}^{opt}$ and $t_{opt}$), which is an important factor for the reproducibility of the hyperspectral data set due to possible pulse length variation of the pump pulses (which will be further explained later). When the output wavelength of the OPO has reached the required parameters the control program starts the image acquisition and awaits its end for continuing in the wavelengths tuning process with a user selectable step size between the consecutive wavelengths.

In a post-processing step the hyperspectral image stack is then intensity corrected and each image is normalized according to the applied laser intensity at the specific pump wavelength with the Matlab routine (fig. 40b). For high precision of this post-processing step it proved necessary to characterize the wavelength-dependent transmission / reflectivity of the pellicle beam splitters used for recording a small portion of the pump and Stokes beam on the photodiodes. Here, a transmission curve was recorded and stored in the analysis program, which corrected the photodiode data by a wavelength-dependent weight factor.
**Figure 40** – Block diagram of the control program for hyperspectral CARS image acquisition

In **a)** the block diagram of the custom-written Matlab routine HyperSpectrum shows the parameters of the OPO, which are acquired and optimized during the wavelengths tuning process in a hyperspectral image acquisition. The temperature tuning curve of the LBO crystal was carefully characterized and stored in the routine. The cavity length of the OPO $L_{Cav}$ is adapted to every target wavelength, while the tuning of the OPO’s generated wavelength is performed by the Lyot filter and monitored via the inbuilt spectrometer. All parameters are optimized with HyperSpectrum in order to operate the OPO at its working point for the maximum intensity output at the specific wavelength. This leads to a constant pulse length and reproducible data acquisition.

In **b)** showcases the post-processing by the Matlab routine HS-CARS, which utilizes the data of the applied laser intensities of pump and Stokes beam recorded by the photodiodes for the correction of stripes in the CARS images due to laser jitter up to 100kHz. In the second step the normalization between all single images of the hyperspectral data stack is performed, which is necessary due to variation of output power of the OPO according to the selected wavelength. For the normalization step the wavelength-dependent transmission of the pellicle beam splitters is measured and implemented into the program as additional weight factor. Published in [Pilger et al. 2018].
4.3.4. Pulse length variation induces spectral distortions in hyperspectral CRS-microscopy

During the development of the hyperspectral CARS scanning modality as well as in additional experiments\(^{14}\) it became apparent, that the pulse length of the laser pulses generated by the OPO is depending on the mode of operation and several tuning parameters. Probing e.g. a molecular resonance while switching back and forth between the pump beams of identical wavelength provided by both OPOs with an acousto-optic tunable filter in a switching rate of \(100kHz\) different CARS signal intensities were recorded. The effect of pulse length variation is most significant during a wavelengths tuning of the OPO e.g in a hyperspectral CARS experiment. Even by applying the same focal power or by measuring the power and recalculating the intensity change in the final images as a correction, the pulse length has a significant influence to the final CARS intensity. As seen in eq. \(25\) the intensity of the pump beam \(I_p\) contributes being squared to the final CARS intensity \(I(\omega_{as}) \propto I_p^2\). The pump beam intensity is composed of the constant duty cycle of the laser system (here \(80MHz\)) and the pulse peak energy \(E_{peak}\), which is defined by the pulse length of the applied laser pulses. The significance of this influence on HS-CARS experiments is illustrated in fig. 41, here a U2OS cell sample is scanned probing a range of \(2738 - 2947 cm^{-1}\) in a step size of \(4,7 cm^{-1}\) while operating the OPO during the wavelengths tuning in two modes: a static OPO setting with optimized OPO parameters on one output wavelength (upper left side in fig. 41) as well as with our customized OPO control software described in chapter 4.3 (see also fig. 40), referred to as dynamic optimization. The dynamic optimization utilizes the prior analysis of the OPO and performs an active optimization of two parameters: the cavity length \(L_{Cav}\) as well as the temperature of the LBO-crystal \(t\) assuring a constant pulse length of the pump beam (depicted on the upper right side of fig. 41). In this study [Pilger et al. 2018] the pulse length of the pump beam was recorded simultaneously during the imaging by directing a small portion of the pump beam into an autocorrelator (APE, pulseCheck 50) and assigning its value to the images (see also fig. 19). This way the microscopy setup was independent and undisturbed of the pulse length measurements and the time of a HS-CARS scan was unaltered. The result of the pulse length measurement indicated, that the static optimization of the OPO parameters on one wavelength leads to a variation in pulse length, whereas the dynamic optimization shows unaltered pulse length. Performing the HS-CARS measurement on a U2OS cell and comparing the spectral response in a selected area (here the nucleus) of both OPO modes shows a strong distortion of the spectrum; the

\(^{14}As a part of the master thesis of Mr. Alex Weiss [Weiß 2017].\)
CARS spectrum using constant pulse length (fig. 41, blue solid line) has an offset to the data using a varying pulse length (green dashed line).

Figure 41 – HS-CARS imaging scheme: influence of the OPO operation mode on spectra due to pulse length variation

On the upper left the HS-CARS image stack of a U2OS cell sample is depicted using a static OPO optimization at 819.6 nm pump wavelength. During the spectral scan the pulse length is not constant resulting in a distorted CARS spectrum depicted in green dashed line. On the upper right the dynamic optimization mode of the OPO is shown utilizing the home-built Matlab program. All OPO parameters are optimized during the HS-CARS acquisition, the pulse length is kept constant resulting in the undistorted spectrum depicted in the solid blue line. Both spectra are acquired of a ROI at the cellular nucleus, and were averaged of 3 repeated HS-CARS scans for each spectrum. Published in [Pilger et al., 2018].
4.3.5. Identifying the OPO parameters influencing the pulse length during wavelengths scanning

In order to investigate the behavior of the OPO and to identify all involved parameters in the OPO tuning, which can affect the pulse length of the output beam of the OPO, additional experiments were performed. The aim of this study was to inform the community performing CRS-microscopy on the OPO’s pulse length variation as well as to offer the customized, Matlab-based software for OPO control to avoid disturbed HS-CARS data. Up to the authors knowledge the pulse length of the OPO was considered in general as determined by the pump laser and, therefore, as constant and not a factor, which had to be further regarded or controlled in hyperspectral experiments. This is also the case for hyperspectral SRS due to the similar dependence on the pulse length of the pump beam. For the determination of the range of pulse lengths, which can occur with the present ps-OPO system, a pure DMSO (dimethyl sulfoxide) sample was prepared between two cover slides separated by thin Teflon ring probing the $C–H$ symmetric stretching mode at $2913 \text{ cm}^{-1}$ by using a constant wavelengths combination of $1064 \text{ nm}$ as Stokes and $812, 3 \text{ nm}$ as pump beam. A constant focal power was applied of $12 \text{ mW}$ Stokes and $23 \text{ mW}$ pump beam correcting the OPO’s power variations due to the settings by rotating the halfwave-plate in front of the polarizing beam splitter cube. The OPO’s cavity length remained constant during the experiment. The OPO’s LBO crystal temperature was then tuned over a range of $3, 5^\circ \text{C}$ (see fig. 42a) while measuring the pulse length as well as the CARS signal by acquiring images ($256 \times 256 \text{ px}$ using $15, 6 \mu \text{s}$ pixel dwell time). For the visualization of the CARS signal intensity all pixels were averaged and normalized on the lowest value coinciding with the longest pulse length (fig. 42b). The pulse length varied in a range of $3, 6 - 5, 4 \text{ ps}$, while the highest possible output power of the OPO at that specific wavelength was observed at the longest pulse length defining the optimal crystal’s temperature $t_{\text{opt}}$. The corresponding normalized CARS intensity showed a linear behavior indicated by the linear regression as a guide to the eye. A signal increase with a factor of up to $1, 51$ was observed in the CARS intensity. The behavior of the OPO was not entirely decoupled from setting a temperature below (red crosses) or above (blue circles) $t_{\text{opt}}$, indicated by the different gradients of the linear regressions. The linear dependence can be explained as an overlay of two competing effects: shortening the pulse length of the pump beam scales quadratic to the CARS intensity as already mentioned leading to a quadratic increase in CARS intensity. On the other hand the temporal overlap of the unaltered Stokes beam with the varying pump beam is decreasing with shorter pulse lengths of the pump beam, which can be understood as an “area matching” in temporal space of both pump and Stokes pulses. The combination of both counteract-
ing effects leads to the observed linear increase in CARS intensity. Then the cavity length $L_{\text{Cav}}$ of the OPO was tuned in the order of $\sim 20\mu m$ holding the LBO crystal temperature constant probing again the DMSO sample (fig. 43). The highest output power of the OPO was observed with the longest pulse length defining the optimal set point for the cavity length setting to $L_{\text{Cav}}^{\text{opt}}$. A change in pulse length in the range of $5,1 - 3,2\,ps$ was detected leading to a change in CARS intensity, which was non-linear (fig. 43a). For investigating the unexpected non-linear behavior of the CARS intensity change two additional experiments were conducted: First, the repetition rate of the pump beam generated by the OPO was measured by detecting the laser pulses with a fast Si-photodiode (Thorlabs, DET10A) and a spectrum analyzer (Tektronix, MDO 4054 B-6, utilizing the 6GHz spectrum analyzer function). The repetition rate was constant while the cavity length was changed in a range of $17,5 - 25,5\mu m$ (data not shown). Second, the temporal overlap of Stokes and pump beam was investigated by feeding the monitor signal of the laser as reference signal of the Stokes beam into a real-time oscilloscope (Tektronix, MSO 71254C, 100GS/s 12,5GHz). The pump beam was detected at a fixed position in the optical beam path with the fast Si-photodiode and directed into the second channel of the oscilloscope utilizing a differential SMA probe (Tektronix, P7313SMA). Both signals were compared by the 50% rising edge amplitude and by applying the delay measurement function of the oscilloscope. Then at least 35450 triggered events (caused by the single laser pulses) were detected and compared at two settings of the OPO cavity, elongating the cavity by $\sim 25,5\mu m$ lead to a change of temporal offset to the reference of $1,57 \pm 0,52\,ps$ by subtracting both data sets of each other. Hence, tuning the cavity length in a larger scale can lead to a temporal mismatch of Stokes and pump beam pulses in CARS experiments. This result lead to a repeat experiment changing the cavity length while probing a DMSO sample with CARS, but now the temporal overlap of both beams was manually adjusted to the highest CARS signal possible at one cavity length by tuning the delay stage behind the OPO beam exit (compare fig. 19). Now the expected linear dependence of pulse length alteration and normalized CARS signal was observed (fig. 43b).
Figure 42 – OPO crystal temperature influences CARS intensity due to pulse length variation imaging a DMSO solution

(a) depicts the dependence of the pulse length due to the OPO’s crystal temperature $t$ in a range of 3, 5°C: red crosses indicate $t$ below the optimal temperature $t_{\text{opt}}$, blue circles above $t_{\text{opt}}$, respectively. $t_{\text{opt}}$ coincides with the maximum output power of the OPO’s signal beam as well as with the longest pulse length ($\sim 5,4$ ps). The corresponding normalized CARS intensity probing the CH symmetric stretching mode at 2913 cm$^{-1}$ with a static pump wavelength of 812,3 nm is shown in (b): the pulse length varies due to the change of the crystal temperature in a range of 5, 4 – 3, 6 ps leading to an increase of the CARS intensity of up to 1, 51. The data is normalized to the lowest CARS signal intensity, a linear regression is added as a guide to the eye. The OPO cavity length was kept constant. Published in [Pilger et al., 2018].
Figure 43 – Detuning of OPO cavity length leads to pulse length variation and temporal mismatch resulting in CARS intensity changes

In a) the cavity length of the OPO was de-tuned from the optimal value $L_{\text{Cav}}^{\text{opt}}$ in a range of $\sim$ 20µm having a constant crystal temperature while a DMSO sample was investigated at $2913\text{cm}^{-1}$. The pulse length varies from $5,1 - 3,2\text{ps}$, while a difference in normalized CARS intensity occurs. The CARS values can be separated into two groups: $L_{\text{Cav}}$ below (red crosses) or above $L_{\text{Cav}}^{\text{opt}}$ (blue circles). No linear behavior of CARS signal to pulse length was observed, which proved to be due to temporal mismatch of the pump to the Stokes laser pulses yielded by the change in cavity length. In b) the temporal overlap was manually optimized by a delay stage behind the OPO signal exit resulting in a linear behavior. The observed effect of the pulse length variation showed an increase of up to 71% of the normalized CARS intensity at a pulse length of $3,2\text{ps}$ compared to the longest pulse length of $5,1\text{ps}$ having the optimal cavity length $L_{\text{Cav}}^{\text{opt}}$. Published in [Pilger et al. 2018].

4.3.6. Influence of pulse length induced distortions on HS-CARS spectra shown in methanol

For estimating the influence of the OPO induced pulse length variation in HS-CARS data sets a pure methanol sample was prepared in the same way as the DMSO sample between two thin cover slides. Then a HS-CARS scan was performed by probing $2800 - 2955\text{cm}^{-1}$ (corresponding pump wavelength was $809 - 818\text{nm}$) in two modes of operation: first, by utilizing the home-written Matlab program, and second, by using a fixed OPO setting optimized for one pump wavelength. Then the second mode was applied recording two spectra at two OPO optimization wavelengths ($819, 8\text{nm}$ and $815, 1\text{nm}$) for comparison to the dynamic mode and to indicate the effects on the spectra itself when changing the static optimization wavelengths, which is usually not a criterion mentioned in current hyperspectral CRS work but very likely affects the reproducibility (fig. 44 a). In the scanned wavenumber range two prominent molecular resonances of methanol are visible: the symmetric $C - H$ stretching mode at $2830\text{cm}^{-1}$ as well as the anti-symmetric $C - H$ stretching mode at $2940\text{cm}^{-1}$. The data was
normalized on 2823 cm$^{-1}$ for comparison reasons. The pulse length was simultaneously acquired for the entire wavelengths tuning range of 8 nm and plotted in fig. 44 showing a constant value for the dynamic optimization ($\sim 5$ ps) while the static optimization induces a shift in the pulse length in the range of 5, 0 – 3, 6 ps (blue curve, OPO optimization at 819, 8 nm) and 5, 0 – 4, 0 ps (red curve, OPO optimization at 815, 1 nm), respectively. The longest pulse length is recorded at the wavelength of the initial OPO optimization. Due to the pulse length variation the hyperspectral CARS intensity is affected, the black curve (constant pulse length) serves as reference at the 2940 cm$^{-1}$ peak showing a difference to the static OPO optimization at 819, 8 nm (blue curve) of 41%, to the OPO optimization at 815, 1 nm (red curve) of 16%, respectively. Comparing the obtained spectra to a spontaneous Raman spectrum acquired of the methanol sample with a 785 nm cw-laser system embedded in a home-built Raman micro-spectroscopy setup shows a good overlap of the molecular resonances of the HS-CARS spectrum to the spontaneous Raman spectrum (fig. 45). This indicates that the HS-CARS setup has a sound performance over the scanned wavelengths range as well as a good wavelengths calibration in the range of the probed resonances.

**Figure 44** – OPO operation modes causing intensity distortions on HS-CARS spectra in methanol due to pulse length variations

HS-CARS spectra probing the range of 2800 – 2955 cm$^{-1}$ in pure methanol covering two molecular resonances at 2830 cm$^{-1}$ and 2940 cm$^{-1}$. a) displays OPO operated by the dynamic optimization mode having a constant pulse length (black solid line), static OPO optimization at 819, 8 nm (2800 cm$^{-1}$) (blue dashed line) and at 815, 1 nm (2870 cm$^{-1}$), respectively. A normalization of all spectra is performed at 2823 cm$^{-1}$ for easy comparison. A distortion of the red and blue spectra compared to the reference becomes apparent at the 2940 cm$^{-1}$ methanol resonance. In b) the corresponding pulse lengths are depicted showing an unaltered value utilizing the dynamic optimization (black solid line), whereas the static OPO settings (red and blue dashed lines) show longest pulse length (5 ps) at the wavelengths of the optimization, but a decrease during the wavelengths scan of up to 3, 7 ps, leading to the observed distortion in the HS-CARS spectrum. Published in Pilger et al. [2018].
Figure 45 – Comparison of HS-CARS to spontaneous Raman spectrum acquired on pure methanol
The normalized CARS intensity during a hyperspectral scan (2800 – 2955 cm\(^{-1}\)) (red line) of pure methanol is compared to a spontaneous Raman spectrum (2700 – 3100 cm\(^{-1}\)) (black line) acquired by a home-built spectroscopy setup utilizing a 785 nm cw-laser system. Two resonances are observed: the symmetric and anti-symmetric C–H stretching mode at 2830 cm\(^{-1}\) and 2940 cm\(^{-1}\), respectively. Both spectra are normalized to 2823 cm\(^{-1}\). HS-CARS shows good accordance to spontaneous Raman data; the spectral peaks are at the same Raman shift indicating a successful spectral calibration of the HS-CARS setup. The single acquired data points are linked by solid lines guiding the eye, the pump wavelength depicted in red is valid only for the CARS spectrum in combination with the unaltered 1064 nm Stokes beam.

4.3.7. Investigating the effect of non-controlled OPO wavelengths tuning on HS-CARS spectra in U2OS cell sample

For the investigation of the spectral distortions due to the pulse length variation on more relevant biological samples a U2OS cell sample fixed with 4% para-form aldehyde solution\(^{15}\) was imaged. A hyperspectral CARS scan was performed in the range of 2738 – 2947 cm\(^{-1}\) with a spectral width of 4.7 cm\(^{-1}\) resulting in a data set of 47 images. Again the two previously discussed modes of OPO wavelengths tuning were performed while the HS-CARS scan was repeated for each mode 3 times in alternating order excluding any additional effects on only one mode due to temporal drifts. The duration of each HS-CARS scan was 27 min resulting in 162 min for all 6 scans. For

\(^{15}\)The cell preparation was performed by Dr. Mathias Simonis.
the visualization of the data one set was chosen to be analyzed by a PCA analysis [Bonnier and Byrne 2012], the first 3 loadings were selected for the false-color coding depicted in fig. 46 a. The corresponding pulse lengths are shown in fig. 46 b, the dynamic optimization has a constant pulse length of 4.9ps (solid blue line) and the static OPO optimization at 819,6nm (dashed green line) exhibits a variation from 4.8ps at 819,6nm to 3.5ps at 810,0nm. Two regions of interest are selected for further analysis, an area of a prominent lipid droplet (ROI A, yellow dot) and a larger area of the cellular nucleus (ROI B). In these areas the pixel values are averaged and the 3 repeated experiments are utilized to generate the mean spectrum of each OPO mode depicted in fig. 46 c (ROI A) and d (ROI B). The standard deviation is represented as the error bars of each spectral data point, indicating two different conclusions: first, in general the standard deviation is low for both methods indicating a high reproducibility due to a stable microscopy setup over time (maximal error $\delta_{max} < 10\%$ of the corresponding data point for ROI A static mode, mean error $\bar{\delta} < 4.2\%$; for ROI B $\delta_{max} < 4.5\%$, mean error $\bar{\delta} < 2.4\%$, respectively). Second, comparing dynamic and static OPO mode to each other we observe a lower error value of both ROIs using the dynamic optimization mode, (ROI A dynamic optimization: $\delta_{max} < 9.8\%, \bar{\delta} < 3.9\%$, for ROI B mean error $\delta_{max} < 2.8\%, \bar{\delta} < 1.5\%$, respectively). This indicates a even higher reproducibility of HS-CARS scans utilizing the home-written Matlab program.

The spectral distortions due to the pulse length variation become apparent comparing the blue solid lines to the green dashed lines in figs 46 c, d: for a better comparison the undistorted spectra were utilized as reference and the relative change in CARS intensity was calculated by dividing the distorted spectra by the undistorted (fig. 47 a for ROI A, b for ROI B, respectively). Here, a pulse length induced increase of CARS intensity with a factor of up to 1.52 in ROI A and of up to 1.49 in ROI B was found (both at 2940cm$^{-1}$).

Summing up these findings the use of a controlled OPO tuning mode is crucial for undistorted HS-CARS data acquisition due to OPO induced pulse length variations during wavelengths tuning. The observed effects using a static OPO control mode can vary spectral components in a range of up to a factor of $\sim 1.5$. A custom-written OPO control software proved to be one possible solution excluding the OPO as a source for spectral distortions and can be of help to other research groups, who are active in the field. The experiments were performed with CARS as a contrast mechanism but are also valid for hyperspectral SRS spectra due to the similar dependence to the pulse length. During the experiments the pulse length was permanently recorded to show the contrast to a static OPO tuning mode. This is not necessary using the dynamic OPO control mode in daily HS-CARS data acquisition due to the coincidence of a repro-
ducible and constant pulse length of the OPO generated pump beam with the highest possible output power of the OPO at a specific wavelength. This relation was shown investigating the OPO behavior at pure samples, where the cavity length as well as the LBO crystal temperature were identified as main influence on the OPO tuning. These findings are exploited for the presented dynamic optimization mode via a home-written Matlab-based control program. Hence, also research groups without an autocorrelator are able to perform undistorted HS-CARS scanning utilizing our findings for their OPO control mode, if they use their own program, or adapt our proposed program for their setups. In addition to that the general performance of the microscopy setup in terms of precision of spectral calibration (comparison to spontaneous Raman spectroscopy) as well as in mechanical and focal stability (U2OS cell repeat experiments) were quantitatively evaluated and the reproducibility proved to be sound and solid.
Figure 46 – OPO operation modes influence pulse length visualized in HS-CARS imaging of a U2OS cell

a) shows a PCA analysis of an HS-CARS image stack of 47 images acquired in a spectral range of 2738 – 2947 cm\(^{-1}\) with a step size of \(\sim 4.7\) cm\(^{-1}\) of a fixed U2OS sample; three characteristic PCA loadings are chosen for a false-color coding (right box). Two ROIs are selected for extracting the spectral response of the CARS intensity: ROI A contains a lipid droplet (yellow dot), ROI B covers a part of the cellular nucleus. During the repeated hyperspectral wavelengths scan two modes for OPO operation are realized sequentially: dynamic optimization mode and static optimization of the OPO at 819.6 nm. b) depicts the pulse length of the different scanning modes during hyperspectral tuning, a constant value of \(\sim 4.9\) ps is achieved for the dynamic optimization (blue solid line), the static OPO setting leads to a change in pulse length from 4.8 – 3.5 ps (green dashed line). This leads to a distortion of the HS-CARS spectra shown for ROI A (lipid droplet) in c) and for ROI B (nucleus) in d): Normalizing the CARS intensity on one spectral resonance (2840 cm\(^{-1}\) for c, 2780 cm\(^{-1}\) for d) a clear difference between both modes can be observed in the spectra at the molecular resonance at 2920 cm\(^{-1}\). For each OPO mode the hyperspectral scan was repeated three times in alternating order, the error bars indicate the standard deviation of the different scans. Published in Pilger et al. [2018].
Figure 47 – Relative difference in CARS signal during hyperspectral scanning due to OPO induced pulse length variation

(a) shows the relative variation of the normalized CARS signal of ROI A (fig. 46, lipid droplet) by dividing the curve with varying pulse length (fig. 46, green dashed line) by the spectrum having constant pulse length (fig. 46, blue solid line). An enhancement in CARS signal of up to 1.52 is observed at 2940 cm$^{-1}$ due to the OPO induced pulse length variation. (b) shows the comparison for ROI B (nucleus area) indicating a change in relative CARS intensity of up to 1.49 at 2940 cm$^{-1}$. Published in [Pilger et al. 2018].
4.4. Applying HS-CARS imaging to biological research questions

In the following section the successful application of HS-CARS answering specific biological questions will be demonstrated emphasizing the comparison of the hyperspectral imaging in one image plane to data acquired at a fixed resonance imaging the entire volume of the sample by z-scanning.

4.4.1. Analyzing the influence of copper on the fitness parameters of Caenorhabditis elegans

In a collaboration with the research group of Prof. Walter Traunspurger, Bielefeld University, Caenorhabditis elegans were analyzed during a life history trait experiment under the influence of copper concerning their fitness parameters [Fueser et al. 2018]. Here, the lipid storage is imaged by probing the 2845 cm$^{-1}$ resonance via CARS microscopy as an indicator of energy for cell survival, which is closely connected to the influence of toxic stress on the organism [den Hartigh et al. 2010, Yen et al. 2010]. Changes in the lipid distribution due to aging are already reported to influence the lifespan of C. elegans [Hou and Taubert 2012] and the correlation of fat storage, oxidative stress and lifespan is found [Yen et al. 2010], showcasing the close relation of life history traits during the life cycle experiment and the lipid analysis utilizing CARS microscopy. The label-free manner of the image acquisition is especially beneficial to this problem, due to the already found misleading interpretations of the amount of lipid storage by utilizing fluorescent staining for microscopic imaging [Yen et al. 2010, Yi et al. 2014] as well as a different behavior of C. elegans in the food uptake, metabolic rates and distribution properties [Hellerer et al. 2007] and also the change in molecular physical properties [Cheng and Xie 2015] due to the fluorescent dyes.

In the present study, the life history trait was performed over 42 and 43 days on a control group (24 nematodes) as well as a group treated (24 nematodes) with low (EC5) and moderate (EC20) concentrations of copper based on the experimental design of [Muschiol et al. 2009].

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16 Sample preparation and additional experiments were performed by Mr. H. Füser during his master thesis [Fueser 2016] in the research group of Prof. Traunspurger.

17 Not investigated by (HS)-CARS.
4.4.2. Volumetric CARS imaging reveals changes of the lipid storage in C. elegans

For the investigation of the lipid storage utilizing CARS 4 adult nematodes of the control as well as from the EC20 group were fixed in a 4% para-form aldehyde solution after their natural death. Due to the absence of lipid droplets in the pharynx region of the C. elegans the vulva and the tail region were selected for the imaging (fig. 48 a). The $C-H_{2}$ stretching mode at 2845 cm$^{-1}$ was probed by 816, 7 nm as pump and 1064 nm Stokes beam applying in total less than 30 mW of focal power avoiding any damage to the sample having a pixel dwell time of 25, 6 µs. The detected CARS signal at 660 nm was isolated by the filter set already described in 4.2. For each region a volumetric z-scan was performed covering the entire diameter of the specimen of 13 – 32 µm with a step size of 500 nm (27 to 65 slices). The study contains 187 images of the control and 135 images of the EC20 group. The analysis of the CARS data was performed by applying a background subtraction and creating a composition of the z-stack into one single two-dimensional projection utilizing the FIJI plug-in VolumeJ 1.8 [Abramoff et al. 2004] for the final quantification of the lipid content for vulva (fig. 48 b) and tail region (fig. 48 c), respectively. After adaptive thresholding (range of 110-255 gray values) the images were converted into a binary form for the final identification of lipid areas of the nematodes by the FIJI tool particle analyzer [Schindelin et al. 2012, Rueden et al. 2017]. This way the total area of lipids per nematode was calculated by summing up all voxels identified as lipids (fig. 49 a) and also the mean area per detected lipid increment was calculated (fig. 49 b). Dividing the total area of lipids by the nematodes’ total area [after Hellerer et al. 2007] the relative lipid area fraction was calculated (fig. 49 c). By sorting the identified average areas of lipids into area classes of 6 intervals (2 – 10, 10 – 20, 20 – 30, 30 – 40, 40 – 50, > 50 µm$^2$) the distribution of the occurrences was studied (fig. 49 d). The data indicates a strong influence by the copper treatment on the lipid distribution of the nematodes; the lipid-filled areas as well as the body fraction, which is used for lipid storage, are reduced in adult nematodes of the EC20 group in comparison to the control group showing high significance. Considering the distribution of lipid area size the EC20 group shows the absence of larger lipid increments (> 30 µm$^2$), instead the number of small lipid areas (≤ 10 µm$^2$) was increased. This findings indicated, that CARS microscopy is a powerful tool to study the physiological mechanisms in C. elegans exposed to copper as a toxic substance. The life history trait experiment provides data on hatching time, lifespan, fertility and reproduction of the nematodes, which also proved to be strongly affected by the copper exposition. The detailed findings and conclusions of these experiments will not be included into this thesis.
Figure 48 – 3D CARS imaging studying the lipid distribution in C. elegans during life history trait

(a) shows a bright-field image of an adult C. elegans depicting the areas selected for CARS microscopy
(red boxes, vulva and tail region, respectively). In (b) 3D projections of CARS probing the 2845 cm\(^{-1}\)
resonance are presented imaging the vulva region of nematodes of the control group (left column) and
the EC20 treatment (right column, respectively). The corresponding data of the tail region is shown
in (c) highlighting lipid-rich structures such as tissue or lipid droplets of seven adults. Each image
shows another replicate. The scale bar indicates 30 \(\mu m\). Published in [Fueser et al. 2018].
Figure 49 – Quantitative analysis of 3D CARS data determining lipid size and content of control and EC20 treatment

After summing the 3D-CARS data into two-dimensional arrays the area containing lipids is identified via thresholding; a) compares the total lipid area of control group and EC20 treatment, b) the average area of lipids, respectively. In c) the relative lipid area fraction is calculated by dividing the total area of lipids by the area of the nematode. By sorting the findings of the total lipid area into 6 size classes the relative size frequency was determined (d). Testing for significance: ***: \( p < 0.001 \), **: \( p < 0.01 \), n.s.: \( p < 0.05 \), \( n = 7 \), median depicted by solid line, mean by dashed line, respectively. Error bars represent standard deviation. Published in [Fueser et al., 2018].
4.4.3. Hyperspectral CARS scanning assessing false positive identification of lipid contents

Following, the volumetric CARS data will be contrasted to hyperspectral CARS imaging in one focal plane covering the wavenumber range of $2664 - 3117 \text{cm}^{-1}$ in terms of information content and applicability to the biological question. The specimen was an adult nematode of the control group fixed in 4% para-formaldehyde after the natural death. The overall focal power was set below $54 mW$ while scanning the pump beam from $799 - 829 nm$ in $0,3 nm$ step size applying a pixel dwell time of $13 \mu s$. During the hyperspectral scan 101 images were acquired and finally analyzed with a principal component analysis (PCA, compare Schie and Huser [2013]) in order to separate the different spectral components of the response of the specimen (fig. 50 a). The first 4 loadings of the PCA showed substantial information content and were selected for the illustration of the nematode, where the principal component 3 indicates the lipid response, PC4 showed predominantly the protein content (fig. 50 b). The lipid content in the body of the nematode can be clearly identified due to the characteristic spectral response, which leads to a clear separation to other parts such as the digestive system or the germ cells. Here, the full spectral response of all components of the nematode enables this clear separation, which is more difficult and reliable probing solely the $2845 \text{cm}^{-1}$ resonance due to unspecific signal contributions by the non-resonant background of CARS microscopy. Here, a smaller mistake in false positive identification of lipid droplets can be expected. On the contrary, the extended time consumption of the hyperspectral CARS imaging ($\sim 30 \text{mins} / z$-plane) in comparison to standard CARS imaging ($\sim 15 \text{sec} / z$-plane) hinders a high number of replicates. Even one full volumetric analysis of one nematode assuming an average number of 35 z-slices results in a measuring time of $17,5 \text{hours}$ in the case of hyperspectral CARS. In the conclusion a combination of both techniques is a promising strategy, where the hyperspectral data can evaluate the quality of the volumetric standard CARS images in terms of false positives at lipid droplet identification as well as to grant deeper insight into other parts of the nematode possibly affected by the copper treatment. Thus, future research on this specimen should likely combine both methods granting high data quality as well as high throughput.
An adult nematode was investigated by a hyperspectral CARS scan from 2664 – 3117 cm\(^{-1}\) wavenumbers by tuning the OPO via a customized Matlab program. **a**) shows the image of a PCA analysis performed on the image stack of the 101 images, the first 4 loadings were selected as important contributions for the image and were color-coded for the final image. ROI 1 shows the digestive tract, ROI 2 shows a randomly selected lipid droplet. A clear separation is achieved between desired lipid contribution and other compartments of the C. elegans, especially the non-resonant background, inevitable in CARS imaging. The scale bar indicates 20 µm. **b**) shows the corresponding scores of the PCA’s first 4 loadings according to the Raman resonance probed by the combination of the fixed Stokes beam at 1064 nm and the tuned pump beam. PC3 (blue) indicates the lipid contribution (at \(\sim 2845\) cm\(^{-1}\)), PC4 (black) has a peak at 2940 cm\(^{-1}\), which corresponds to the protein contributions.
4.4.4. Visualizing the differentiation of primary human alveolar epithelial cells by label-free imaging techniques

As conclusion of the research on C. elegans combining standard CARS imaging with hyperspectral scanning this scheme of analysis is adapted to an other specimen of interest. In a collaboration with the research group of Prof. Maike Windbergs, Goethe University Frankfurt a. M., the differentiation of primary human alveolar epithelial cells was investigated with several imaging modalities such as fluorescence microscopy, spontaneous Raman microscopy as well as CARS and hyperspectral CARS microscopy [Vukosavljevic et al. 2019]. The novelty of the study is the combination of spontaneous Raman microscopy and both CARS techniques as label-free imaging modalities for the comparison to classic fluorescence based microscopy. The differentiation of alveolar type II (ATII) cells located in the peripheral human lung into type I (ATI) cells is of interest because of the generation of the air-blood barrier. ATI cells are forming a cellular barrier to the respiratory tract while the ATII cells are producing, accumulating and secreting the alveolar surfactant, which is lining the cells in the deep lung area. In addition to that the ATII cells are responsible for replacing ATI cells via differentiation into type I, which is accompanied by morphological as well as molecular modifications in terms of cellular lipid content and expression of specific marker proteins studied already with confocal laser scanning microscopy, rt-PCR, flow cytometry and immunoblotting [Fuchs et al. 2003, Campbell et al. 1999]. The morphological changes during the differentiation have been visualized in the past by scanning electron microscopy as well as transmission electron microscopy [Cheek et al. 1989, Fuchs et al. 2003] showing the flattening and wide spreading of the cells. The spreading of up to 90% of the alveolar surface area allows the in vivo gas exchange of the blood. Summing up the prior work on the differentiation of ATII into ATI cell type the analysis methods required fixation and disruption of the cells [Pygall et al. 2007]. Therefore, spontaneous Raman scattering was applied in imaging mode on this problem for gaining insight into the Raman shift of the cells during differentiation accessing their molecular changes [Kann et al. 2015], which was performed with a commercial confocal Raman microscope (WITec, alpha 300R+) [18]. For backing up the conclusions of this study fluorescence microscopy (Zeiss, Zeiss LSM710) as a well-established standard was applied to ensure observing the differentiation process by utilizing Laurdan staining for the visualization of the lipid content. The data of spontaneous Raman microscopy as well as fluorescence microscopy will not be shown in this thesis.

[18] Spontaneous Raman and fluorescence microscopy were performed by Dr. B. Vukosavljevic as part of Vukosavljevic 2018.
4.4.5. **Volumetric CARS imaging enables z-sectioning and shows differentiation process over time**

The combination of volumetric CARS microscopy showing the differentiation process probing the $2845\,cm^{-1}$ resonance for the visualization of the lipid content during a time period of 7 days as well as hyperspectral CARS imaging at selected regions of interest (ROIs) of samples at days 1, 2, 3 is centered in this paragraph. The volumetric CARS microscopy was performed by scanning the cell samples with a step size of $300\,nm$ in z-direction resulting in an image stack of 30 to 40 images per replicate and day covering the entire cellular volume (fig. [51]). The focal power of the Stokes and pump beam were set to be below $30\,mW$, isolating the CARS signal with the previously described optical filter set combination. By applying a twofold zoom the field of view was $75\,\mu m \times 75\,\mu m$ while acquiring $512 \times 512\,px$ with a pixel dwell time of $31.25\,\mu s$. This results in an acquisition time of $6\,mins$ for one 3D-CARS scan per replicate. For the visualization of the data FIJI- Volume Viewer [Schindelin et al. [2012], Rueden et al. [2017]] was utilized to generate a conversion summing up all z-slices into one top view. At the beginning of the experiments we observed cell diameters of $\sim 10\,\mu m$ at day 1. Over time cell growth started and more lipid vesicles could be found and the cells were grouped together after day 3. Days 5 and 7 show strongly enlarged, converging cell structures, which exceed even the field of view. These findings were also in good accordance to the other methods applied on the specimen such as spontaneous Raman microscopy as well as fluorescence microscopy (see also [Vukosavljevic et al. [2019]]). In order to back up the claim of applying CARS microscopy to image the differentiation process of the ATI type cells into ATI type repeat experiments of days 1 to 5 were performed on more replicates, while additional data of day 4 is presented fitting in the overall findings of the differentiation process (fig. [52]). In the case of rather low CARS signals it proved useful to add dotted green lines marking the cellular border as well as blue dotted lines marking the cell nucleus as guide to the eye. For showing a sufficient z-sectioning capability of the setup, which is capable to differentiate between the nucleus and the lipid filled cellular volume, selected 3D data is provided showcasing single consecutive z-scans of cells of day 1 (fig. [53]) and day 3 (fig. [54]), respectively. In fig. [55] the 3D-CARS data of day 2 is presented in a $90^\circ$ rotated side view illustrating the z-extent of the ATI cells. Here, also the z-sectioning capability can be evaluated due to the clear display of the cellular border. However this type of presentation of the 3D-CARS data is limited to a strong signal-to-background ratio occurring at high lipid contents in the cells, which were observed predominantly at the first 3 days of differentiation.
Figure 51 – Comparison of representative CARS images illustrating the differentiation process of ATII into ATI cells over 7 days

The differentiation process from ATII cells into ATI imaged by volumetric CARS imaging; 3D projections are utilized for the illustration of the lipid distribution and the extent of the cells, while probing the $2845\text{ cm}^{-1}$ $CH_2$ stretching resonance. The projection made with FIJI Volume Viewer sums the volumetric data of every z-scan for a top view. The cellular extent at day 1 is rather small, the shape is round. Over time the cells flatten out and the number of lipid droplets is decreasing until at day 7 the field of view is filled completely by the ATI cell. Published in [Vukosavljevic et al., 2019].
Figure 52 – Repeat experiments during the differentiation process of ATII cells applying CARS imaging at different time points

For backing up the claims on the differentiation process of the cell samples several replicates were imaged at the selected time points. Here, additional data on day 4 is included confirming the claim of the differentiation from ATII into ATI type. At day 5 the lipid content is already strongly reduced resulting in a rather weak CARS signal, therefore, the green dashed line is included for marking the cellular border and to guide the eye. Published in Vukosavljevic et al. [2019].
Figure 53 – Representative CARS z-scan on ATII cells at day 1
Volumetric imaging is performed for visualizing the lipid content of the cell samples by applying a step size of 300nm in z-direction. Here, a sample of day 1 is shown scanning a z-depth of 5,8µm. Due to the low lipid content of that particular time point the green dotted line is added marking the cellular border as a guide to the eye, the blue dotted line marks the nucleus, respectively. The cell appears to be round and has a diameter of \(19\mu m\). Published in [Vukosavljevic et al. [2019]].

Figure 54 – Representative CARS z-scan on the differentiation of ATII cells into ATI at day 3
A representative sample after 3 days of differentiation is z-scanned in the range of 21µm with a step size of 300nm, while in the image a step size of 1,2µm is selected for presentation. The cellular area is larger compared to day 1 and the cell’s shape is not round anymore, but has flattened out. The green dashed line depicts the cellular border guiding the eye, the blue line depicts the nucleus, respectively. Published in [Vukosavljevic et al. [2019]].
An other way of presenting the volumetric CARS data of the cell samples is presented above: the FIJI plug-in Volume Viewer is utilized for enabling a 90° rotation of the viewing plane gaining insight into the z-expansion of the cells during the differentiation process. This method is only applicable having higher signal levels due to a high lipid content. The corresponding top view image can be found in fig. 51 d2. Published in [Vukosavljevic et al. [2019]].

4.4.6. Non-negative matrix factorization applied on HS-CARS scans reveals cellular compartments

To confirm the claims made due to the volumetric CARS imaging of distinguishing the lipid rich cellular volume from the nucleus hyperspectral CARS imaging was performed in one focal plane covering a range of 2700 – 3050 cm⁻¹ by tuning the pump beam from 826.5 – 803 nm with a step size of 4.7 cm⁻¹ (0, 3 nm), and applying a focal power of less than 30 mW. The pixel dwell time was adapted to the signal to background due to the current lipid content of the particular day of differentiation as well as to the selected zoom level (2× or 3×) ranging from 6,4 – 12.8 μs while the total scanning time per replicate was for days 1 and 3 37 mins and for day 2 52 mins. On the data sets the non-negative matrix factorization (NNMF) as a multivariate statistical analysis tool was performed by a Matlab-based custom written program¹⁹ based on the Matlab Statistics tool box [Matlab [2012]], which performed the differentiation of the different spectral components of the specimen, such as lipid rich vesicles (2845 cm⁻¹) and cellular proteins (2930 cm⁻¹) as well as the non-resonant background (fig. 56).

¹⁹The development of the NNMF analysis program was predominantly performed by Mr. Henning Hachmeister, also in supervision to the NNMF analysis related work in the master thesis of Mr. P. Greife.
For this purpose a ROI was selected, where an offset subtraction and a binning in a $4 \times 4$ pixel frame was performed reducing the arbitrary signal fluctuations, which can hinder the successful application of the NNMF algorithm by the non-relevant changes in intensity due to noise. In the ROI 4 image subsections were selected showing a characteristic spectral response, which were utilized as initial values inputted into the NNMF analysis. The NNMF analysis separates the original input data $A$, as the original matrix, into the two non-negative matrices $W$ and $H$ with the definite rank number $k$ representing the unique vibrational signatures, which were as a priori knowledge inserted into the algorithm by selecting the subsets in the input data. This step also grants the reproducibility of the performed analysis. The NNMF works in an iterative manner in order to find $W$ and $H$, while minimizing the root-mean-squared residual between $A$ and $W \cdot H$ [Berry et al. 2007]. Applied on the presented images $W$ and $H$ are representing $k$ times a two-dimensional image and the corresponding vibrational response, respectively. Here, each two-dimensional image section of $W$ is having a corresponding intensity distribution of the vibrational response imprinted on $H$. For the final visualization $W$ is false-color coded and illustrated by FIJI. For the link to the spectral response indicated by the false-color coding the eigenvectors found by the NNMF for the entire image are shown next to the images. The successful separation of lipid-rich vesicles (depicted in magenta) in the cellular volume and the cell’s nucleus (blue) can be observed, while the background is identified and depicted in black color. For the investigation on the signal-to-background level of the provided NNMF analysis the eigenvectors of the specific channels of interest are presented in a small sub-ROI, which show a characteristic area of the specimen such as e.g. the nucleus showcased in the data set of day 2 (fig. 57). The clear peak structure in the spectral behavior of the eigenvector indicates a sufficient signal-to-noise ratio of the input data. Comparing the eigenvector spectra of lipids (fig. 57a) to proteins indicating the nucleus (fig. 57c) a distinct spectral discriminability becomes apparent. The presented hyperspectral data showed good accordance to the volumetric CARS data in terms of nucleus identification and also fitted well to the spontaneous Raman data (not shown here). In general the study proved that the differentiation of ATII into ATI type primary human alveolar epithelial cells can be visualized with label-free techniques allowing to study morphological and biochemical changes as well as the storage and secretion of the lung surfactant without the labeling influences.

The benefits of the NNMF analysis utilized in this study over the prior presented PCA analysis are the always positive values of the NNMF eigenvectors as well as the input of prior defined spectral information, which enables an easier interpretation of the results. Furthermore, the NNMF analysis is less affected by undesired influences.
of signal gradients, which can occur due to the rolling ball effect in CARS microscopy caused by the spatial mismatch of Stokes and pump beam at the outer regions of the image or due to a tilted sample plane in relation to the focal plane, which often occurs at imaging tissue sections prepared by a microcryotome on thin glass slides.

Summing up the results on the hyperspectral CARS imaging it was possible to apply these technique to relevant biomedical problems, while the volumetric CARS imaging at one particular molecular resonance enabling fast imaging for a high number of replicates in combination to the hyperspectral scanning proved to be a powerful tool. Here, both methods could play on their advantages, while their intrinsic drawbacks were balanced, especially the rather slow acquisition speed and lacking volumetric information of HS-CARS.
Figure 56 – Hyperspectral CARS imaging of ATII differentiation process utilizing NNMF for decoding spectral information

For the hyperspectral CARS imaging a range of $2700 - 3050\,cm^{-1}$ was scanned with a step size of $4.7\,cm^{-1}$ applying a total focal power of less than $30\,mW$ over time (days 1 to 3 - rows d1 to d3). A non-negative matrix factorization (NNMF) analysis was performed on each image stack producing three individual channels which correspond to the lipid rich environment (magenta), proteins (blue) indicating the nucleus and the background (black) used in the overlay images on the left. The scale bar indicates $1\,\mu m$. On the right the NNMF eigenvectors are shown, which are associated to the different spectral components averaged over the entire image. Increasing lipid droplet formation is observed on days 1 to 2. At day 3 the cellular volume starts to increase, while the size and number of the lipid droplets is continuously growing. Published in [Vukosavljevic et al. 2019].
Figure 57 – Investigating the signal-to-background behavior at selected ROIs utilizing NNMF analysis on hyperspectral CARS data at day 2

The data set of fig. 56 day 2 (d2 - false-color encoded image in the middle) is utilized to showcase the ability of the NNMF analysis to distinguish between the different presented channels by the generation of eigenvectors with sufficient signal-to-background ratio. Therefore, three characteristic regions of interest (ROIs) are selected showing the predominant spectral eigenvector value for the ROI. In a) the ROI is chosen to be from a lipid rich cellular area (depicted in magenta), the eigenvector shows a strong peak at the $2845\,cm^{-1}$ $CH_2$ resonance. A ROI of the area outside of the cell is selected in b), the eigenvector shows an even, slightly fluctuating spectral behavior representing the non-resonant background. c) is selected at the nucleus region of the cell (blue color coding), the eigenvector has a peak at the protein resonance at $\sim 2930\,cm^{-1}$. Note that the data points are connected as a guide to the eye. The scale bar indicates 1µm. Published in [Vukosavljevic et al. 2019].
4.5. Resolution enhancement utilizing structured illumination in scanning-based non-linear microscopy

Recently notable efforts were performed in biomedical research in order to circumvent the diffraction limit in super-resolution fluorescence microscopy by various methods such as STED (stochastic emission depletion) [Hell and Wichmann 1994, Klar and Hell 1999], d-STORM (direct stochastical optical reconstruction microscopy) [Fölling et al. 2008, Wolter et al. 2010, Heilemann et al. 2008, Rust et al. 2006] as well as SIM (structured illumination microscopy) [Lukosz and Marchand 1963, Gustafsson et al. 2008, Gustafsson, M. G. L. 2000, 2005] to name just some major applications in this field. All have the need of a sufficient fluorescence label in common enabling the various methods a resolution of up to \(\sim 20-50\,\text{nm}\) in stochastic-based and \(\sim 180-200\,\text{nm}\) in linear structured illumination-based microscopy [Wu and Shroff 2018, Schermelleh et al. 2019, Sahl et al. 2017]. In the following paragraph the combination of an enhanced resolution and a label-free contrast mechanism by CARS microscopy is explored trying to combine the benefits of the super-resolution fluorescence-based techniques without the need for fluorescent labels, which are laborious, toxic and depending on the specimen can not be applied at all due to rigid cell walls or undesired autofluorescence of cellular compartments. Approaches for a resolution enhancement in CRS-microscopy were demonstrated utilizing the saturation of the CARS signals or the beam shaping of the participating pump or Stokes beam [Betin et al. 1975, Duncan et al. 1981, Tolles et al. 1977, Yonemaru et al. 2015, Kim et al. 2012]. More recently the saturation of the SRS signal was demonstrated by applying very high focal powers and, therefore, limiting the range of samples and increasing the phototoxicity [Gong et al. 2019, Gong and Wang 2014].

For the realization of enhancing the resolution in the presented CRS-setup the combination of structured illumination with CARS as a heterodyne detected contrast mechanism was chosen. Here, two strategies are pursued in order to show the resolution enhancement: first, a scanning mirror-based striped illumination pattern generation for two-photon excited fluorescence is introduced showing significant enhancement in contrast and resolution. The selected specimen are fluorescent bead samples, stained murine tissue sections as well as auto-fluorescent Convallaria majalis sections imaged on the TriM Scope II in a collaborative effort with LaVision Biotec, Germany. The purpose of these measurements is showing the easy implementation of the scanning mirror-based structured illumination due to the use of almost all standard components in state of the art two-photon microscopes by just adding a sCMOS camera for the final detection of the images. Second, this concept is transferred to the presented CARS...
microscopy setup: utilizing the experience in epi-CARS detection [Pilger 2014] concerning the signal strength in combination with the use of a sCMOS camera instead of a PMT the final setup contains a $4\pi$ detection in order to maximize the signal by using forward CARS for the camera-based detection. Here, the mechanical stability (see 3.2) as well as the optimal setting of the pattern parameters turned out to be crucial. The presented data on pure samples such as polystyrene bead samples or DMSO solutions can be seen as the first successful realization of structured illumination in combination with CARS for resolution enhancement and contrast improvement.

4.5.1. Generating striped illumination pattern by addressing the galvanometric scanning mirrors

Realizing the striped illumination patterns by using the galvanometric scanning mirrors was explored at first on the CRS-microscope by synchronizing the AOM for rapid intensity switching due to the current position of the mirror (see also 3.2.1). Applying the patterns to pure dye solutions the accuracy and timing of the pattern was checked and first data were acquired on bead samples by using the 1064nm laser source for two-photon excitation. Here, the pattern fineness and the setting of suitable values for pattern line spacing and rotation angles were tested and evaluated for best modulation depth and signal-to-noise ratio in the raw data. The finer the pattern the lower the modulation depth of the illumination pattern will get due to scattering effects in the sample, which will illuminate also the non-illuminated areas to a certain extent. For best results a high fluorescence signal strength was required, which was realized by transferring the experiment to a setup capable of femto-second laser excitation of the fluorophores. Therefore, a industrial collaborative project was initiated with the company LaVision Biotec, which is developing the two-photon scanning microscope TriM Scope II. This system was specifically modified as platform for the presented two-photon excited fluorescence data sets in this paragraph. The excitation source was a $\sim 140\text{fs}$ pulsed laser system (Coherent, Chameleon Ultra I), with $690 - 1040\text{nm}$ tunable wavelengths range in combination with a dispersion compensation stage realized through a prism compressor. The generation of the striped illumination pattern at the TriM Scope II setup was realized by a specifically adapted software program written in Phython, which allowed to change the line spacing, angle of rotation as well as the phase settings of the excitation pattern. In addition to that the program addressed an inbuilt EOM in order to block the excitation beam when the scanning mirrors were illuminating an undesired position in the sample. A $60\times$ oil immersion

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20 The work was supported by Dr. H. Spiecker, Dr. M. Schütte, Mr. M. Ruoff, LaVision Biotec.

21 Developed together with Mr. M. Ahlering, LaVision Biotec.
objective lens (Olympus, APON, NA = 1.42) focused the excitation laser beam into the sample. For detecting the images a home-built tube lens holder was utilized, which also mounted the sCMOS camera (Andor, Neo 5.5) on top of the microscope body. A 700nm SP beam splitter separated the excitation lasers at 780nm from the epi-detected fluorescence signals. Additional bandpass filters were used in front of the camera to avoid undesired signals or to separate the fluorescence signal of the staining of interest from others (if so the filter is mentioned in the figure caption). The tube lens was a $f = 300\, \text{mm}$ apochromatic $\Theta = 50\, \text{mm}$ lens focusing the fluorescence signal onto the camera chip, which had a pixel size of 6.5$\mu\text{m}$. In combination with the objective lens and tube lens pairing this lead to a pixel size in the image of 66nm. In this approach a illumination pattern with three phases and three angles of rotation (separated by 60°) was used.

### 4.5.2. Enhancing the resolution and contrast by striped illumination of fluorescent bead samples in two-photon excitation

For determining the best pattern spacing a series of images were acquired on fluorescent bead samples (beads $\Theta = 200\, \text{nm}$, ThermoFisher, TetraSpeck microspheres) by increasing the pattern fineness. The bead samples were created by diluting the stock solution by a factor of 1 : 5 with distilled water and letting a drop of 60$\mu\text{l}$ dry on a cover slide, while the sample was placed overhead. By this preparation a single monolayer with the desired density of the fluorescent beads was created. For evaluating the best combination of modulation depth of the excitation pattern as well as a high fineness of the pattern for a high resolution enhancement the *FIJI* plugin FairSIM [Müller et al. [2016]](fig. 58) was utilized in the reconstruction process. In the FairSIM plugin the Richardson-Lucy filtering algorithm on both input and output data of the reconstruction was applied setting 15 iterations. The apodization cut off was set to 1.2. The raw data set containing 9 images was averaged and used as the widefield image for the comparison to the reconstructed image. Fig. 59 shows a zoom into a ROI of a group of fluorescent beads. Here, the filtered widefield image is added to the comparison due to the enhancement of contrast by the necessary filtering steps in the reconstruction process. Hence, the filtered widefield image uses the averaged data of the widefield image but then performs the same filtering steps as the final reconstruction. In this way the comparison is more significant and one can make sure that observed improvements are arising from the striped illumination and not only due to filtering of the images, which is similar to a deconvolution step. In the presented reconstruction the contrast is clearly enhanced due to a larger range of intensity values compared to widefield and filtered widefield image, which can be seen by performing a line plot (fig. 60). Also a
group of beads presented in the ROI can not be resolved in the widefield images. Only the reconstruction shows a gap between single beads, also visible in the line plot by a clear drop of intensity (fig. 60 pink arrow).

**Figure 58** – Structured illumination applied on a $200\text{nm}$ TetraSpeck fluorescence bead sample using two-photon excitation

A mono layer of fluorescent beads $\varnothing = 200\text{nm}$ was illuminated with a striped illumination pattern using two-photon excitation by a $780\text{nm}$ femtosecond laser source having $\sim 140\text{fs}$ pulse length. Left hand side shows the average of all nine acquired raw images forming the widefield image, right shows the reconstruction using FairSIM applying a Richardson-Lucy filtering on the final image. The red box is selected as ROI for a zoom in (see fig. 59). The scale bar indicates 10$\mu$m.

**Figure 59** – Comparison of widefield images to the reconstruction in a region of interest of a $200\text{nm}$ bead sample

The selected ROI shows a constellation of several $200\text{nm}$ beads grouped together in a circle. Left hand side shows the widefield image, middle shows the filtered widefield, which corresponds to a deconvolved image, and right hand side shows the final reconstructed image by FairSIM. The red line indicates the location of a line plot performed for comparing all three images (fig. 60). The scale bar indicates 2$\mu$m.
Figure 60 – Line plot of the 200\,nm bead sample: comparing widefield images to the reconstruction shows enhanced resolution and contrast

The line plot compares the normalized intensity values of widefield (red line), filtered widefield (green) and reconstruction (blue) of the 200\,nm TetraSpeck bead sample using a striped illumination pattern and two-photon excitation (compare red line in fig. 59). Widefield and filtered widefield are unable to resolve the gap in a group of the fluorescent beads, whereas the reconstructed image can separate between the single beads and shows a clear decrease of intensity (pink arrow). The increased contrast of the reconstruction becomes apparent due to the larger range of intensity values (0, 0.06 – 1\,a.u.; reconstruction) in comparison to widefield and filtered widefield (0, 2 – 1\,a.u.; filtered widefield).

4.5.3. Exploiting autofluorescence signals of Convallaria majalis for combining z-scanning with striped illumination pattern via MAP-SIM

In order to explore the performance of the presented striped illumination pattern generation in two-photon microscopy on a larger field of view as well as in a 3D scanning experiment the auto-fluorescent sample Convallaria majalis is investigated. Changing to a 25× air objective lens (Nikon, NA = 1.1; WD = 2\,mm) in combination with the $f = 300\,mm$ tube lens lead to a field of view of $\sim 360\,\mu m$ having a magnification factor of 41, 6 and a camera pixel size of 1\,px=$156,3\,nm$. A z-scan over 8\,\mu m with a step size of 0,5\,\mu m was sequentially performed by the built-in z-scanner of the TriM Scope II setup, based on a micro-motor drive. Nine images containing the tree angles of rotation with each three phase settings of the illumination pattern were recorded for each z-plane. Each z-plane data set was reconstructed by the Matlab-based reconstruction toolbox MAP-SIM\textsuperscript{22} [Lukeš et al. 2014, Krříek et al. 2015], which has the ability to utilize even higher harmonics of the applied illumination pattern. By setting a relative

\textsuperscript{22}In collaboration with Mr. J. Pospisil.
coarse illumination pattern and by the intrinsic character of galvanometric scanning-based illumination patterns to be not sinusoidal higher harmonic contributions of the illumination pattern are observable in the Fourier spectra of the raw data sets. The FairSIM reconstruction algorithm is developed predominantly to operate on illumination patterns, whose frequencies are chosen to be at the edge of the detectable OTF for highest resolution enhancement. In the reverse conclusion a higher harmonic of the illumination pattern is arising artifacts in the final reconstructed image, because its frequency content is not separated from the first order contribution and is not shifted to the correct position in the image. This behavior of the reconstruction algorithm is reducing the possible pattern settings in the presented approach to be consisting only of one detectable frequency of the illumination pattern. In the MAP-SIM algorithm second or higher harmonics of the illumination pattern are utilized for the reconstruction process and contribute to the final image.

Hence, the MAP-SIM approach is applied to the Convallaria majalis sample, which showed a coarse pattern fineness. For each z-plane the reconstruction was performed and also an averaged widefield image is generated for comparison reasons. All acquired z-planes were then false color coded by the hyperstack function in FIJI (fig. 61), the color bar indicates the z-position. The contrast of the reconstruction as well as the z-sectioning capability of the reconstructed image stack is enhanced. Two ROIs are selected for a zoom image, indicated by the red boxes. ROI 1 shows an area with a thicker structure of the cell walls of the Convallaria majalis sample (fig. 62), while ROI 2 (fig. 63) shows a thinner cell wall structure. Both ROIs show a clearer z-sectioning capability as well as a clear enhancement in contrast in the reconstructed image. Due to the lower magnification and coarser structure of the sample no resolution enhancement beyond the diffraction limit can be expected, but nevertheless the striped illumination approach in combination with the MAP-SIM reconstruction improves the image quality and the data set was reconstructable in all recorded z-planes. Combining the scanning mirror-based striped illumination with MAP-SIM as reconstruction tool can be further optimized by finding optimal parameters for illumination angles and phase step size and choice of increments in order to fully exploit the second or higher harmonics of the striped illumination pattern in the reconstruction. Both reconstruction algorithms introduced in this paragraph are compared in their performances on the Ø = 200 nm fluorescent bead sample (compare fig. 58). MAP-SIM as well as FairSIM are optimized in its performance on this data set and the final images are compared to the averaged widefield image (fig. 64). Both reconstructions show the resolution improvement due to the applied structured illumination pattern in the two-photon excitation. The MAP-SIM approach has a better background removal whereas FairSIM shows at
the location of the beads a slightly increased performance in terms of resolution. A line plot indicated by the green line in fig. 64 shows that the dip between single beads is more prominent using FairSIM as well as the diameter of the left hand side bead is slightly smaller (fig. 65).

Scattering in the sample material is identified as major factor which still limits the detectable fineness of the illumination pattern. Applying a very fine illumination pattern on a fluorescent bead sample with a bead diameter of Ø = 1µm shows this effect very prominent at the border of the illuminated area (fig. 66). Two effects are visible in this image: first a corona at each bead arising from out of focus light as well as the illumination of beads, which are outside of the excitation area (red arrows). The red line indicates the border between the area with excitation by the laser (left hand side) and the non illuminated area (right hand side). Beads in the non illuminated area are also visible due to scattering of laser light of the excitation or of the generated fluorescence light into the non illuminated area. This effect is also present in the area between single illuminated lines of the applied striped pattern, which is reducing the modulation depth of the pattern leading to the need of coarser patterns in the recorded raw images. Here, MAP-SIM can circumvent this challenge due to more freedom in the choice of pattern fineness.

Figure 61 – MAP-SIM reconstruction performed on a z-scan image series of a Convallaria majalis sample
The two-photon excited autofluorescence of a Convallaria majalis sample was utilized for applying the structured illumination in addition to a z-scan over 8µm with a step size of 0.5µm. In each z-plane nine images were recorded shifting the phase and angle of the illumination pattern. Each image stack per z-plane was reconstructed by the MAP-SIM toolbox and the results of all z-planes were false color coded according to the z-position by FIJI using the hyperstack functionality. Two regions of interest (ROI 1 and 2, red boxes) were selected for a more detailed comparison of the above shown averaged widefield images and the reconstruction (see figs 62 and 63). Note, that for this data set a 25x objective lens (Nikon, NA = 1,1 ; WD = 2mm) was utilized. The scale bar shows 40µm.
Figure 62 – ROI 1 of Convallaria majalis sample comparing widefield to MAP-SIM reconstruction. This ROI compares the averaged widefield (left hand side) and reconstructed image (right hand side) showing several linked cell walls of the sample. The contrast and z-sectioning is enhanced by the MAP-SIM approach, additional features are revealed in the reconstructed image. The color code bar (box) indicates the position during the z-scanning according to the false color in the image. The scale bar shows 10µm.

Figure 63 – ROI 2 of Convallaria majalis sample comparing widefield to MAP-SIM reconstruction. The second ROI compares the averaged widefield (left hand side) and reconstructed image (right hand side) of a single cell. The contrast and z-sectioning is enhanced performing the MAP-SIM reconstruction. The color code bar (box) indicates the position during the z-scanning according to the false color in the image. The scale bar indicates 10µm.
Figure 64 – Comparing the performance of FairSIM and MAP-SIM on fluorescent bead samples. Both reconstruction algorithms are tested on the $\theta = 200\,\text{nm}$ fluorescent bead data set showing a clear increase in resolution due to the ability to resolve a bead cluster, whereas in the averaged widefield image no single bead structures are visible. In FairSIM the same parameters are used as in fig. 58. The green line indicates the location of a comparing line plot. The background suppression is slightly better in the MAP-SIM reconstruction. The scale bar (white) indicates $500\,\text{nm}$.

Figure 65 – Line plot comparing reconstruction algorithms on a fluorescent bead sample. Comparing both reconstruction algorithms (FairSIM red line, MAP-SIM blue line) to the averaged widefield image data (black line) a clear dip between single beads is apparent due to the applied structured illumination pattern in two-photon excitation. FairSIM has a slightly better performance compared to the MAP-SIM reconstruction due to the higher decrease of values in the gap between the beads as well as the slightly smaller diameter of the left hand side bead.
Figure 66 – Fluorescent 1µm bead sample imaged by fine illumination pattern investigating scattering effects

A sparse monolayer of 1µm fluorescent beads was illuminated with a very fine illumination pattern (single lines of the pattern can not be seen in the presented raw image). The selected cutout of the full image shows the border between illuminated (left hand side) and non-illuminated area (right side), indicated by the red diagonal line. This raw image shows two effects: first, a corona around each bead as well as, second, a fluorescent signal of the non-illuminated area showing the outline of additional beads (arrows). This is an example of the limitations of the striped illumination approach due to scattering events in the sample. The scale bar shows 2.5µm.

4.5.4. Investigating the potential of a striped two-photon illumination in tissue utilizing a stained murine kidney section

In the next step a murine kidney tissue section (ThermoFisher, FluoCells prepared Slide #3) stained with three fluorescent dyes (Alexa Fluor 488 WGA, Alexa Fluor 568 and DAPI) was investigated utilizing a laser wavelength of 800nm for the two-photon excitation of the Alexa Fluor 568 dye, which stained the filamentous actin prevalent in the glomeruli and the brush border. A 595/40BP bandpass filter is set in front of the camera isolating the desired signal at ~ 603nm. Again the developed striped illumination scheme is applied to this sample while the reconstruction is performed in MAP-SIM for fig. 67 as well as in FairSIM using similar parameters as in the fluorescent bead sample for a cutout of the data set (fig. 69). Averaged widefield and reconstructed image are compared in fig. 67 showing an enhanced contrast and resolution. A line plot (fig. 68), whose location is indicated by the red line in fig. 67, is confirming the superior performance of the reconstruction over the widefield image. The range of intensity values is larger in the reconstruction indicating a better contrast, edges in
the reconstructed image are steeper showing the improvements in resolution (apparent
in fig. 68). A ROI processed now with FairSIM (fig. 69) compares the performance
of reconstruction to widefield in an overlay image, confirming that both reconstruction
algorithms work well on the generated data sets.

Figure 67 – Striped two-photon excited fluorescence imaging of a stained murine kidney tissue
section comparing widefield to reconstruction

A wavelength of 800 nm was utilized for the two-photon excitation of the Alexa Fluor 568 dye, which
stains the filamentous actin prevalent in the glomeruli and the brush border in the tissue section
[Wiederschain [2011]]. An additional 595/40 BP bandpass filter isolated the detected signal in front
of the camera. Generating averaged widefield (left hand side) and reconstructed image (right hand
side) by the MAP-SIM algorithm allows the comparison: the reconstruction shows enhanced contrast
and resolution. The red line marks the location of a line plot for comparing both images (see fig. 68).
The scale bar indicates 25 µm.
The line plot compares the normalized intensity values of the widefield (red line) to the reconstructed image (blue line) at the ROI indicated by the red line in fig. 67. The reconstruction made with MAP-SIM shows an enhanced contrast due to the larger range of intensity values as well as an enhanced resolution indicated at the steep edges of the filamentous actin prevalent (pink arrows).

A single cell of fig. 67 is selected for comparing the widefield to the reconstructed image utilizing the FairSIM algorithm with similar parameters determined on the bead samples. The reconstruction shows also an enhanced contrast and resolution confirming that the reconstruction process of the generated data sets is working well with both described algorithms. The scale bar indicates 5µm.
4.5.5. SI-CARS: investigating the potential of resolution enhancement by striped illumination patterns

For the generation of the striped illumination patterns utilizing CARS as contrast mechanism the Stokes beam was rapidly switched in its intensity according to the position of the galvanometric scanning mirrors (see 3.2.1). The parameters of the pattern such as spacing of the illuminated lines, angle of rotation and pixel dwell time were set by the scanning control program ScanImage. Finding the best setting of the pattern fineness was performed on pure samples such as butter, DMSO or olive oil. Here, the modulation depth of the detected illumination pattern was compared and balanced to the noise level in the raw images due to the camera exposure time as well as to the desired high fineness of the pattern, which finally generates the resolution enhancement in the reconstructed image. An example of a raw image with a coarse illumination pattern easy visible as bright stripes is presented in fig. 70. The $2913 \text{cm}^{-1}$ C–H symmetric stretching resonance was probed in pure DMSO, which was applied on a dried monolayer of polystyrene beads with a diameter of $Ø = 1,1 \mu m$. The strong CARS signal of the DMSO generated an anti-contrast of the polystyrene beads, which was useful to evaluate the CARS signal strength on a sCMOS camera as well as to adjust the optics generating the final image on the camera chip. The red arrow indicates the scan direction of a single illuminated line.

For the reconstruction of a SI-CARS data stack nine images were acquired consisting of three different angles of the illumination pattern separated by $\sim 60^\circ$. For each angle of the pattern the phase is shifted in three steps in order to fully illuminate the sample area. An example of typical raw data is showcased in fig. 71. A monolayer of polystyrene beads with a diameter of $Ø = 2 \mu m$ was prepared by letting a drop of the diluted stock solution dry out overhead on a cover slip. In the SI-CARS experiment the $3070 \text{cm}^{-1}$ resonance of the polystyrene was probed generating the CARS signal with a wavelength of $643,6\text{nm}$, which was filtered by a stack of optical short-pass and bandpass filters and finally detected on the sCMOS camera. The reconstruction was performed by FairSIM generating again the averaged widefield and the filtered widefield images for the comparison to the reconstructed image (fig. 72). Due to a still rather coarse illumination line spacing the parameters in the reconstruction process were adapted to the applied illumination pattern. The apodization factor was set to 1,2 while a final Wiener filter of 0,056 delivered the best results on this data set. The red box indicates the position of a line plot showing the normalized intensity values of all three presented images (fig. 73). A better contrast as well as a resolution enhancement becomes apparent studying the gap between two beads due to the larger range of intensity values occurring in the reconstruction as well as a decrease of intensity at the
gap structure between two beads (indicated by the pink arrow in fig. 73). The power spectra of the reconstruction process as well as for the final images can be directly extracted from the FairSIM software (fig. 74). The spectra of the raw data for each angle of the illumination pattern show a clear peak of the frequency of the applied pattern, which is utilized for finding the correct length and angle of the $k_0$ parameter fitting routine (fig. 74 red circles and green arrow). Comparing widefield to the reconstructed image in Fourier space clearly shows additional spatial frequencies in the reconstruction, which becomes apparent in fig. 74 right hand side. The green circle in the Fourier space of the reconstruction indicates the maximum spatial frequencies of the widefield image, all values outside of this circle depict the additional spatial frequencies due to the applied structured illumination.

Figure 70 – 1,1$\mu m$ polystyrene beads in DMSO solution illuminated with coarse CARS illumination pattern
A coarse illumination pattern was generated probing the 2913$cm^{-1}$ $C – H$ symmetric stretching mode of the DMSO solution in which 1,1$\mu m$ polystyrene beads were embedded. In this way an anti contrast of the beads was generated. The red arrow indicates the scan direction. The scale bar indicates 5$\mu m$. 
Figure 71 – Raw data set using striped CARS illumination pattern for imaging \(2 \mu m\) polystyrene beads

A suitable fineness of the CARS illumination pattern was chosen probing the \(3070 \text{cm}^{-1}\) polystyrene resonance. The raw image set consists of three angles separated by \(60^\circ\) rotation, while for each angle the illumination pattern is shifted in its phase by three increments in order to illuminate the entire sample space. The red arrows indicate the scan direction for each angle. The scale bars show \(2 \mu m\).

Figure 72 – SI-CARS: comparison of widefield to reconstruction in \(2 \mu m\) polystyrene beads

The SI-CARS raw data set (fig. 71) probing the \(3070 \text{cm}^{-1}\) polystyrene resonance is processed by the FairSIM plugin in FIJI using a Wiener-filter of \(0.056\) setting a apodization factor of \(1.2\). Left shows the widefield image in comparison to the filtered widefield (middle) and the reconstruction (right hand side). The red box shows the location of a line plot (fig. 73). The contrast and resolution of the reconstructed image is enhanced in comparison to both widefield and filtered widefield. The scale bar indicates \(2 \mu m\).
Figure 73 – SI-CARS: line plot of ROI comparing reconstruction to widefield image
Comparing widefield (green) and filtered widefield (red) to the reconstructed image (blue) an enhanced contrast due to a higher range of intensity values as well as a higher drop in intensity at the area between the single beads can be observed (pink arrow and dashed vertical line). The intensity values were normalized for better comparison.
Figure 74 – Reconstruction parameter fitting and band separation in Fourier space using Fair-SIM

In the upper row the single power spectra in the Fourier space are shown for each angle of the illumination pattern. The inner area of the yellow circle indicates the setting of the excluded frequencies searching for the $k_0$ value in the parameter fitting routine in the FairSIM plugin. The red circles show the localized intensity maxima for each angle of the illumination pattern determining the correct parameter fitting of $k_0$ (drawn in green) for the reconstruction step. In the lower row the Fourier spectra of the averaged widefield image (left hand side) is contrasted to the reconstruction (right hand side). The enhanced area in the Fourier space of the reconstruction is due to the additional information content generated by the striped illumination pattern according to each angle of the illumination pattern. The green circle added into the reconstruction indicates the maximum frequencies of the widefield image for comparison.
4.5.6. Structured illumination utilizing galvanometric scanning mirrors proves to be a versatile tool for two-photon fluorescence and CARS microscopy

In summary, the performance of the newly developed structured illumination approach by utilizing the galvanometric scanning mirrors showed significant improvement in contrast and resolution in two-photon excited fluorescence microscopy. The presented data set acquired on fluorescent bead samples as well as on relevant tissue sections and biological applications such as a Convallaria majalis sample were successfully reconstructed. This concept has the appealing point of utilizing most of the standard components of a state-of-the-art two-photon scanning microscope by adding a sCMOS camera to the setup. This way a cost-efficient improvement of existing setups can be easily performed using free reconstruction software such as FairSIM as well as the MAP-SIM toolbox for the final image. This concept is also applicable to CARS microscopy avoiding high excitation intensities as well as problems with the spatial overlap of pump and Stokes beam by avoiding the use of pattern generation by light interference. First successful data acquisition was performed on a polystyrene bead sample showing improvement in resolution and contrast. In future steps, doubly-resonant CARS can be utilized as contrast mechanism for boosting the signal intensity [Weeks and Huser [2010]]. Thus a better modulation depth of the applied illumination patterns is expected and also finer patterns can be applied resulting in even higher resolution enhancement.
4.6. Developing a self-synchronized two-color fiber laser for fast, high contrast CRS-microscopy

CRS-microscopy is a versatile tool for biomedical research addressing questions closely related to clinical research because of the fast, non-invasive and chemical sensitive imaging modality [Yun and Kwok [2017]]. At the same time no fluorescent labels are needed in this technique while the excitation laser powers are moderate and are mostly not affecting the sample of interest [Zumbusch et al. [1999], Freudiger et al. [2008], Nandakumar et al. [2009], Ozeki et al. [2009]]. Hence, the preparation of the sample is minimal compared to fluorescence-based techniques and also in-vivo experiments can be performed even at human patients in the operating room [Evans et al. [2007], Ji et al. 2013 2015, Lu et al. 2016, Orringer et al. [2017]]. Combining CRS-imaging with endoscopic approaches by utilizing optical fibers can allow live examinations in the human patients [Lukic et al. 2017, Lombardini et al. [2018], Latka et al. 2013], which was performed utilizing CARS as contrast mechanism. Ongoing activities are trying to expand this modality also to SRS.

A major drawback of current state-of-the-art CRS-microscopy setups are the traditional solid-state lasers, typically Ti:sapphire or Nd:Van systems, mostly combined with optical parametric oscillators used for the required tunable laser wavelengths in the CRS imaging process. These laser systems have a large footprint, are cost-intensive, sensitive to influences of the environment such as vibrations, temperature changes, dust and humidity, they require large vibration-isolated optical tables as well as maintenance in terms of retuning the optical paths for granting the spatio-temporal overlap and thus can not be easily included into a medical, sterile operating room.

Recent efforts have been performed to exploit fiber laser technology for replacing the traditional laser systems in order to overcome their shortcomings mentioned above: fiber-Ti:sapphire hybrid sources [Ozeki et al. [2010], Wang et al. [2010]], all-fiber lasers based on the following techniques: active synchronization [Nose et al. 2012, Karpf et al. 2015], parametric wavelength conversion [Zhai et al. 2011, Baumgartl et al. 2012a, Lefrancois et al. [2012]], soliton self-frequency shift [Andresen et al. 2007, Pegoraro et al. [2009], Krauss et al. 2009, Xie et al. 2014, Crisafi et al. 2018], and supercontinuum (SC) light generation [Gambetta et al. 2010, Selm et al. 2010, Freudiger et al. 2014, Chen et al. 2016, Tu et al. 2016, Orringer et al. 2017]. Fiber-lasers are robust and the components can be densely packed into sealed containment while they also offer the desired wavelengths tunability addressing the molecular resonance of interest. The recent progress in fiber laser technology allows to include most optical components into the fiber-based optical path [Sobon et al. 2015, Wei et al.].
which reduces the footprint of the system as well as the need of free-space optics and their alignment as well as it improves the overall stability of the optical system. Nevertheless, previous implementations of the fiber laser technology had shown first promising results but still lacked the full performance of the traditional solid-state laser systems especially due to low power levels, a narrow wavelengths tuning range and high laser intensity noise. These shortcomings resulted in poor image quality, low scanning speeds and the need of special noise suppression schemes especially in the case of SRS. Here, the balanced detection is utilized to overcome the shortcomings of laser systems without high quality noise performance [Nose et al. [2012]], but the scheme relies on the delicate adjustment of balanced receivers subtracting the undesired laser noise, which results in obstacles of the systems in terms of continuous performance while wavelengths tuning over a wide range as well as it increases the complexity of the CRS-microscope and the overall costs.

In a collaborative project with the research group of Prof. Dr. K. K. Y. Wong, the University of Hong Kong, China a novel high-power self-synchronized two-color pulsed fiber laser system was developed [Kong et al. [2019]] overcoming the described drawbacks of the fiber laser technology in recent implementations [Kong et al. [2019]]. The presented picosecond fiber laser provides high intensity stability (50dB better performance than competitive systems), low timing jitter (24.3fs), large wavelengths tuning range resulting in an addressable wavenumber range of 2700 – 3550cm⁻¹, low average power fluctuation (< 0.5%), low pulse width variation during wavelengths tuning (< 1.8%) and a fiber-based intensity modulation scheme for SRS imaging with a excellent modulation depth of > 99%. Due to the excellent noise performance and high intensity modulation depth the performed SRS microscopy shows high contrast and high chemical sensitivity without the need of an additional noise cancellation device such as a balanced detection scheme, while the imaging speed is still comparable to other fiber-based as well as solid-state laser systems.

The performance of the fiber laser system is demonstrated in two steps: first, both wavelengths branches are analyzed in terms of spectral and temporal performance by utilizing a real-time oscilloscope, a spectrum analyzer and power measuring devices. Second, the performance is demonstrated by simultaneous imaging with CARS and SRS contrast on several biomedical samples such as living human cell samples, murine arterial tissue sections, murine brain slices as well as pure substances for comparison to spontaneous Raman scattering spectra. In addition to that the introduced fiber-based laser system provides a femtosecond pulse length branch which extends the range of application also to second harmonic generation as well as to two-photon excited fluores-

[Equal contribution with Mrs. C. Kong and Mr. H. Hachmeister.]
cence imaging providing full multimodal imaging ability. This application is showcased on a mouse tail sample (SHG imaging) as well as on a stained murine kidney section and a stained murine brain section visualizing the pyramidal neurons (two-photon excited fluorescence). This fs-pulsed laser setup is also proposed as a stand-alone system with optimized pulse length, which is a powerful and cost-efficient tool for users of two-photon scanning systems [Kong et al. 2017].

4.6.1. Utilizing highly doped ytterbium and erbium non-linear fibers for the low noise, high power two-color laser in CRS-microscopy

Stokes beam generation in the master oscillator branch

The presented two-color pulsed laser system for CRS-microscopy can be separated in two major branches due to the use as Stokes and pump pulses for the final experiment. The master oscillator consists of a passively mode-locked femtosecond laser cavity at 1000nm wavelength realized by pumping a 25cm long piece of ytterbium-doped (Yb) non-linear fiber (Thorlabs, YB1200-4/125) via a 976nm Cw-laser diode (II-VI, LC96Z400; Pump) (fig. 75, left upper corner) [Xu et al. 2011, Spiegelberg et al. 2004]. A fiber-based optically integrated module (OIM), which provides polarization-sensitive isolation, signal extraction and pump/signal multiplexing, closes the cavity and is combined with a drop-in polarization controller (General Photonics, PolaRite; PC). It allows the passive mode-locking by nonlinear polarization rotation (NPR) granting the successful and reliable self-starting of the source with stable fs-pulses having the all-normal-dispersion regime [Chong et al. 2006]. The wavelengths tuning is performed by placing a tunable filter (Semrock, LL01-1064-12,5; F1) inside the cavity, which is used as bandpass filter (∼8nm) and can be tuned by manual rotation (range 50nm) allowing the adjustment of the laser wavelength from 1010 − 1060nm (fig. 81 a). For rapid automatic wavelengths tuning it can be easily replaced by a fiber-coupled tunable filter in later steps of the setup. Also a system trigger is generated by a fast photodiode, which detects the filtered out light. The master oscillator creates ∼20mW power at the systems fundamental repetition rate of 80MHz (fig. 78 a), which is the same as in conventional solid-state laser systems obtaining the advantages of a high repetition rate such as allowing the fast modulation of the laser intensity (typically at 10 − 20MHz) for SRS imaging with high scanning speeds. The generated fs-laser pulse intensity is extracted through the OIM and then split up in two branches with equal intensity by a 50 : 50 splitting fiber optic coupler (OC) feeding each the following Stokes and pump beam generation.

For the Stokes beam generation a fiber-based chirped pulse amplification (FCPA) scheme is applied [Strickland and Mourou 1985]: first, linear chirp is applied to the
fs-pulses by a single mode fiber (Corning, HI60, 50m length; SMF) for reducing the peak power of the pulses. Then a core-pumped single-cladding Yb-doped fiber amplifier (YDFA) is creating $\sim 20\, \text{mW}$ of output power [Paschotta et al. 1997]. It is comprised of a highly-doped Yb fiber (Thorlabs, YB1200-4/125; 25cm length) a 980/1030nm wavelength-division multiplexing coupler (Hapht; WDM) in combination with a fiber-coupled laser diode (again II-VI, LC96Z400) at 976nm for the Cw pump light generation. Then a fiber-based intensity modulator (iXblue, NIR-MX-LN-10; FIM) is placed behind the preamplifier stage and can be used for the fast frequency modulation at 20MHz used in SRS imaging. The modulation is synchronized with the 80MHz fundamental repetition rate of the master oscillator by processing the system trigger signal. The fiber modulator provides a modulation depth of > 99% (fig. 78b), which is an excellent value compared to free-space devices such as e.g. a resonant EOM (electro-optic modulator) with $\sim 80 - 95\%$ modulation depth (compared e.g. to APE, EOM 1030). Second, a high power double-cladding YDFA (DC-YDFA) (similar to Wei et al. 2016) is utilized to boost the Stokes beam to the final power of $> 1\, \text{W}$, which is then coupled out by a fiber collimator and directed to the grating compressor stage (black dotted box, right hand side of fig. 75). This last amplifier stage is composed of a double-cladding Yb fiber (iXblue, IXF-2CF-Yb-O-10-130-NH-V2.0, 1.5m length), which is pumped by a multimode fiber-coupled pump laser diode (QPhotonics, QSP-975-10) with a maximum power of 10W at 975nm wavelength (to generate a maximal Stokes beam power of $\sim 250\, \text{mW}$ in the experiment usually < 4W pump power are used) [Yong Wang and Hong Po 2003, Zenteno 1993, Snitzer et al. 1988, Jeong et al. 2004]. The undesired residual pump power after the non-linear double clad Yb fiber is extracted from the following beam path by removing the outer coating of the fiber and applying UV-curable optical adhesive (Thorlabs, NOA63), which removes the residual pump light due to the higher refractive index of 1.56.

**Grating pulse compressor is optimizing the pulse length of the Stokes beam**

The compressor stage is set up to generate the desired pulse length of the final Stokes beam of $\sim 3,2\, \text{ps}$ (see also fig. 78c) by two high transmission grating pairs (Lightsmyth LSFSG-1000-13212-94) in combination with a telescope (lenses L1 and L2, Thorlabs, AC508-75-B). The telescope has the purpose to change the compression ratio by translation relative to the gratings as well as to allow spectral filtering by a translation slit placed in the focal plane of L1 (spectral bandwidth is 1nm during wavelengths tuning) [Weiner 2009, pp. 412]. By rotating the polarization of the incoming and exiting light by half- and quarter-wave plates in combination with a polarizing beam splitter cube (PBS) the coupling in and out of the grating compressor
is performed with low intensity losses.

**Slaved second laser cavity generates fundamental pump beam light at 1.5µm wavelengths**

For creating the pump beam laser light the second half of the output of the master oscillator cavity at 1µm is injected into a second laser cavity exploiting the coherent wavelength generation process (CWG) utilizing a 1030/1580nm WDM. This second laser cavity is working at 1578nm using the principle of passive self-synchronization for synchronizing the repetition rate to the 1µm laser pulses [Rusu et al. 2004]. The cavity is comprised of a fiber-based OIM and as the gain medium 50cm of a single-cladding erbium-doped fiber (Thorlabs, ER80-8/125; Er), which is pumped by a fiber-based Cw-laser diode at 976nm. The polarization of the laser light inside the cavity is controlled by the second drop-in polarization controller. A narrow optical bandpass filter (BAAR, bandwidth < 1,0nm; F2) placed between two fiber couplers is determining the central wavelength. The cavity length is optimized to match the repetition rate of the master cavity as good as possible by the use of a high precision translation stage mounting one fiber coupler. In this state the 1.5µm cavity can produce a self-sustained pulse train [Haus et al. 1994, Krausz et al. 1990, Tamura et al. 1993], which is not synchronized to the master cavity.

**Synchronization of both laser cavities by exploiting cross-phase modulation and group velocity dispersion effects**

The synchronization of both laser cavities is performed by the passive self-synchronization, which can coherently generate two-color laser pulses. It relies in the interaction of cross-phase modulation (XPM) and group velocity dispersion (GVD), which can occur due to the use of active and passive single mode fibers in the 1.5µm cavity [Rusu et al. 2004, Yoshitomi et al. 2006, Hsiang et al. 2009]. The XPM effect describes the nonlinear phase shift of an electromagnetic field, which occurs when high intensity optical fields of different wavelengths, direction or state of polarization meet in nonlinear optical fibers. The nonlinearity refers to the intensity dependence of the refractive index in the fiber material. XPM allows both wavelengths in the cavity the coupling to each other without an energy transfer between them, leading to an asymmetric spectral broadening of the center wavelength of the laser pulses in the case that both pulses are co-propagating in the fiber ([Agrawal 2012, pp. 13]).

In the described setup the synchronization will start at the first collision of both pulses co-propagating in the fiber of the 1.5µm cavity (see fig. 76) e.g. at the position A to B. The XPM effect initiates a frequency shift of the 1.5µm pulses depending on the
timing of the pulses to each other (the 1,0µm pulses can be considered as temporal reference), leading to a redshift in the case of a faster timing, or a blueshift for a slower timing. This shift is exploited due to having anomalous dispersion at 1,5µm in the cavity by the used silica fiber (Corning, SMF-28e; normal dispersion at 1,0µm) for connecting e.g. the OIM (compare also Agrawal [2012, p. 8]), which then slows down or speeds up the circulating pulse according to the wavelength shift of the XPM effect. By carefully choosing the dispersion coefficient of the fiber both effects lead to a stable range of operation due to the intrinsic mismatch of the repetition rates of both cavities due to the optical path length. Therefore, both cavities are matched in their repetition rate as good as possible to each other in the prior calibration process. Finally, the described method grants an excellent synchronization of both laser branches in this setup without any electronic devices, resulting in the low timing jitter of ~ 24,3 fs between the cavities determined by optical cross-correlation (see fig. 78 e).

For finding the desired material parameters numerical simulations of the XPM-induced wavelengths shifts were performed and finally the self-synchronization was simulated including dispersion and XPM-induced effects (fig. 77 Fisher and Bischof [1973]). a) and b) show the timing evolution of the 1,5µm pulses assuming the repetition rate of the slaved cavity is slightly lower than of the master cavity, without external injected pulses (77 a) the temporal mismatch is continuously increasing, no synchronization takes place. With injected pulses at 1µm less than 50 round trips (~ 625 ns) are enough to perform the synchronization and stabilize the slaved cavity at zero temporal offset (77 b). c) and d) show the configuration, when a higher repetition rate in the slaved cavity is assumed. The gradient of the temporal mismatch is reversed, but the situation with and without the injection of the external 1µm pulses is similar, also resulting in a stable synchronized operation when the 1µm pulses are present in the slaved cavity.

**Pump beam amplification and frequency doubling generate the final pump beam wavelengths**

The 1,5µm laser light is extracted of the cavity by the OIM and then amplified by a heavily-doped double-cladding erbium/ytterbium-doped fiber amplifier (CorActive, DCF-EY-10/128; DC-EDFA), which is pumped by a multimode fiber-coupled pump laser diode (similar to the DC-YDFA) at 975nm. The amplified laser light is coupled out by a FC, which is mounted on a linear translation stage for tuning the temporal overlap to the Stokes beam (delay line, DL). After optimizing the polarization state of the 1,5µm beam by half- and quarter-wave plates it is focused (by lens

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This work was performed by Dr. X. Wei.
L3, $f_{L3} = 75mm$, Thorlabs, AC508-075-C) into a temperature controlled periodically poled lithium niobate crystal (Covesion, MSHG1550-0.5, PPLN) for performing the frequency doubling to 789.6 nm [Taverner et al. 1998]. The bandwidth of the beam is determined to 0.35 nm with a typical power of $\sim 160mW$, which is utilized as the pump beam in the CRS-experiments (fig. 81 d) after performing the collimation and beam diameter matching by an additional lens (Thorlabs, AC254-125-B, $f_{L4} = 125mm$; L4). Changing the central wavelength of the 1.5 µm cavity in combination with temperature tuning of the PPLN crystal an additional wavelengths tuning range can be achieved also in this laser branch extending the addressable range of molecular resonances (fig. 81 c). Stokes and pump beam are spatially overlapped by the already described methods similar to the solid-state laser system (see 3.1.3) and can now be utilized for CRS-microscopy.
Figure 75 – Schematic diagram of the tunable two-color all fiber-based CRS-laser

The passively mode-locked laser cavity (left upper corner) generates the 1µm fs-pulse train with 80MHz repetition rate, which is then split into two branches. One half is amplified by two amplifier stages to generate the Stokes beam (green color), which is compressed by a grating compressor (right middle box) to ps-pulse length. The other half is utilized to synchronize the second slaved laser cavity at 1.5µm (violet color), which is amplified and then frequency doubled by a PPLN crystal to the final pump wavelength (red). Both Stokes and pump beam are spatio-temporally overlapped and then coupled into the scanning CRS-microscope. The right upper box shows a sketch of the self-synchronization utilizing cross-phase modulation and anomalous dispersion. See main text for detailed explanation. Abbreviations: OC - fiber optic coupler, DM - dichroic mirror, DL - delay line, DC-EDFA - double-cladding erbium-doped fiber amplifier, DC-YDFA - double-cladding ytterbium-doped fiber amplifier, Er - erbium doped fiber, F - filter, FC - fiber collimator, FIM - fiber coupled intensity modulator, G - grating, GVD - group velocity dispersion, L - lens, M - mirror, N_{rt} - round-trip number, OIM - fiber-based optically integrated module, PBS - polarizing beam splitter, PC - polarization controller, PD - photodiode, PPLN - periodically poled lithium niobate crystal, SMF - single-mode fiber, YDFA - ytterbium-doped fiber amplifier, \( \lambda/2 \) - half-wave plate, \( \lambda/4 \) - quarter-wave plate. Part of Kong et al. [2019].
Figure 76 – Sketch of the passive self-synchronization mechanism utilized in the fiber laser setup

Half of the 1.0µm laser light (green) serving as master oscillator is coupled into the 1.5µm pump beam cavity (violet), whose repetition rate is already optimized to the value of the master oscillator in the self-sustained pulse train mode but has still a temporal offset ($\Delta \tau_{\text{initial}}$). In the optical path between points A and B a nonlinear fiber enables the interplay between both laser pulses via cross-phase modulation (XPM) inducing a shift of the center wavelength of the 1.5µm pulse train. Due to the anomalous dispersion in the cavity for the 1.5µm wavelength a slowing-down or speeding-up of the pulses is achieved by group velocity dispersion (GVD) according to the frequency shifts induced by XPM. On the right hand side the synchronization process is depicted showing the timing of both pulses to each other while the round-trip number $N_{rt}$ is evolving from left to right. Due to the combined XPM/GVD effect the timing offset of both pulses is decreasing by each round-trip by $\Delta \tau_{\text{xpm}}$ up to the point of synchronization after $n$ round trips. Part of Kong et al. [2019].
Figure 77 – Numerical simulations of the passive self-synchronization mechanism of the 1.5µm laser cavity
Timing evolution of the 1.5µm pulses in the cavity a) without injected 1.0µm pulses and b) with injected pulses, while the repetition rate of the 1.5µm cavity is assumed to be slightly lower than the master cavity in the simulation. The round-trip number is depicted over the temporal mismatch between both pulses. c) and d) show the same situation but having a slightly higher repetition rate of the 1.5µm cavity, respectively. Both simulated situations show a synchronization by the injected 1.0µm pulses after < 50 round trips (temporal mismatch = 0 ps). The simulation assumes an initial timing delay of 3 ps and a round trip time mismatch of 50 fs. Part of [Kong et al. [2019]].

4.6.2. Characterizing the temporal and spectral properties of the all fiber-based CRS-laser source

For the characterization of the constructed fiber laser setup additional experiments were conducted in order to demonstrate the performance also for comparison to other competitive setups. The constant and synchronous repetition rate of both lasers is determined to ~ 80 MHz, 12.5 ns pulse-to-pulse timing by sequentially visualizing the pulse train of both laser wavelengths by a fast photodiode (HP, 11982 A, 10 GHz bandwidth) and a 20 GHz real-time oscilloscope (LeCroy, SDA-820Zi-B) (fig. 78 a). A long term measurement of the pulse train over 500 µs proves a stable operation without any intensity fluctuations or Q-switching events (fig. 79), which also indicates the environmental insensitive operation of the system.

For demonstrating the capabilities of the fiber-based intensity modulator in the Stokes beam path a 20 MHz sinusoidal modulation was applied and again the pulse train was measured showing a modulation depth of > 99% (fig. 78 b), which is an excellent value compared to state-of-the-art resonant EOM solutions in free-space optics (having 80 – 95% modulation depth, compared to e.g. APE, EOM 1030). For comparison a
sinusoidal function is plotted as an envelope to the depicted pulse train of the Stokes beam as a guide to the eye. The pulse length of both laser wavelengths is also characterized by utilizing two autocorrelators (Stokes: Femtochrome Research, FR-103 MN; pump beam: MesaPhotonics, FROGscan) due to the different wavelengths. The Stokes beam has a pulse length of 3, 2\(\text{ps}\) and 2, 7\(\text{ps}\) for the pump beam, respectively. Due to the already described challenges of the pulse length dependence during wavelengths tuning in hyperspectral CRS-experiments utilizing OPO-based laser systems (compare 4.3), the pulse length of the Stokes beam was also monitored during wavelengths tuning (1015–1040\(\text{nm}\)) showing a variation of < 1, 8\% indicating further advantages of the all fiber-based system (green line in 78 c). The power stability performance is determined by detection via the photodiode and the real-time oscilloscope during a time period of 100\(\text{mins}\) for finally calculating the the root-mean-square value to be better than 0, 5\% for the Stokes (setting 550\(\text{mW}\) power) and 0, 1\% for the pump beam (at 100\(\text{mW}\) power), respectively (fig. 78 d). For testing the performance of the synchronization between pulse trains at these two wavelengths, their cross-correlation curve was measured through sum frequency generation (SFG) in a KTP (potassium titanyl phosphate) optical crystal. During the measurement, the temporal delay between both pulse trains was changed using the optical delay line in the optical path of one pulse train. The raw SFG signal is recorded over 10\(\text{sec}\) by an oscilloscope at the peak (gray curve, full temporal overlap) and 50\% (black curve, 2\(\text{ps}\) delay) SFG intensity in (fig 78 e). The timing jitter between these two pulse trains is estimated by calculating the intensity fluctuation of the SFG signal and the rising/falling slope of the cross-correlation trace, both at half maximum, and then divided the former by the latter, yielding a timing jitter of 24, 3\(\text{fs}\) (0, 8\% of the pulse length).

For comparing the noise performance of both laser branches (fig. 78 f, green and red curve) to other widely recognized laser systems the relative intensity noise (RIN) is measured by a fast photodiode (HP, 11982 A, 10\(\text{GHz}\) bandwidth) and an optical spectrum analyzer (Agilent, E4440 A, 26, 5\(\text{GHz}\) bandwidth) and is compared to the values of a standard solid-state fs laser (Spectra-Physics, MaiTai, black solid curve) and a typical super continuum-based (SC, blue and purple curve) fiber laser (similar to [Freudiger et al. 2014]) (fig. 78 f). At the Stokes beam modulation frequency of 20\(\text{MHz}\) for SRS imaging (black dotted box) the introduced fiber-laser system has in both branches similar performance (~147\(\text{dBc/Hz}\) pump beam, ~148\(\text{dBc/Hz}\) Stokes beam) being a unique selling point of the new design offering ~ 50\(\text{dB}\) better performance compared to other recent fiber-based laser systems. These systems suffer from the degraded noise performance due to nonlinear conversion processes of at least one laser wavelength [Dudley et al. 2006], also apparent in (78 f, blue curve). Here,
the use of two laser cavities in the two-color laser system in combination with their synchronization by exploiting the XPM and GVD effects lead to the better performance. The solid-state laser system (black curve) performs close to the shot noise limit (black dotted curve), in this case at $-164 dBc/Hz$, being the edge in terms of achievable noise performance. Further investigations were also performed concerning the frequency spectra at the fundamental repetition rate of the non-modulated Stokes laser (fig. 80 a) and by applying a modulation of the Stokes beam via the fiber-based intensity modulator at $\sim 20 MHz$ (fig. 80 b) as well as of the repetition rate of the pump laser (80 c). The signal-to-noise performance of all acquired RF-spectra show a low laser noise (SNR at $80 MHz$: Stokes 74 db, pump 73 db, respectively), the desired modulation of $20 MHz$ in the Stokes beam has a SNR of 67 dB without any additional frequency contents, which is beneficial for the lock-in amplification process.

The spectral properties of the fiber-based laser system are depicted in fig. 81 by coarse wavelengths tuning via rotation of the intracavity filter in the Stokes beam cavity (F1, bandwidth of $\sim 8 nm$) the accessible wavelengths regime of $1005 - 1065 nm$ is measured in (81 a) by the frequency analyzer. The final Stokes spectrum for CRS imaging is generated by the grating compressor (depicted in fig. 81 b) and has a bandwidth of $\sim 1,0 nm$ offering a continuous wavelengths spectrum from $1010 - 1060 nm$. Also the fundamental central wavelength of the pump beam branch can be tuned from $1540 - 1590 nm$ by rotating filter F2 having a bandwidth of $< 1,0 nm$ (fig. 81 c). The frequency doubling performed by the temperature controlled PPLN has a typical bandwidth of $0,35 nm$ and is showcased at $1578 nm$ doubled to $789 nm$ in fig. 81 d, the transform-limited Gaussian pulse width is 2,6 ps.

The laser setup is capable of high-speed SRS imaging with high chemical sensitivity, which is appealing due to the lack of any non-resonant background contribution in contrast to CARS imaging. For proving the ability to access the Raman resonances by SRS without any shifts of the Raman peaks e.g. due to the dispersive line shapes in CARS, two pure samples are analyzed by wavelengths tuning of the Stokes beam (fig. 82). The Raman resonances of pure methanol and DMSO are acquired in the range of $2800 - 3050 cm^{-1}$ by SRS as well as with spontaneous Raman scattering (SR) with a home-built spontaneous micro-Raman scattering setup utilizing a 785nm Cw diode laser (Innovative Photonic Solutions) in combination with a fiber-coupled grating spectrometer (Princeton Instruments, Acton 2300i) with camera detection (Andor, Newton DU 920p BR-DD). Comparing the SRS to the SR spectra show excellent peak overlap and the same line shapes, whereas in the case of methanol the spectral background between both Raman resonances in the probed wavenumber range is even better in SRS.
Figure 78 – Temporal characterization of the Stokes and pump laser pulses

(a) shows the pulse train acquired with fast photodiodes and a 20GHz real-time oscilloscope at 1017nm (Stokes) and 789nm (pump beam), respectively. (b) depicts the real-time pulse train of the Stokes beam while applying a 20MHz sinusoidal intensity modulation via the fiber-based modulator. The modulation depth shows > 99%. As a guide to the eye a sinusoidal function is plotted as an envelope to the modulated pulses. By performing an autocorrelation of both laser wavelengths the pulse length was determined to 3.2 ps for the Stokes (green) and 2.7 ps for the pump beam (red) in (c). Also the pulse length stability during wavelengths tuning was measured from 1015 – 1040 nm showing < 1.8% variation (green line plot for the Stokes beam). The power stability was proven to be better than 0.5% for Stokes (at ~ 550 mW output power) and 0.1% for the pump beam (at ~ 100 mW output power) acquired over 100 min calculating the RMS value (d). By performing a sum-frequency generation (SFG) of both beams in a KTP crystal and an optical cross correlation by a spectrum analyzer while tuning the delay stage of the pump beam the timing jitter between both laser lines was analyzed (e). The maximum SFG signal (gray line) as well as the half of the signal (black line) are showcased. By fitting a Gaussian function to all measured SFG intensities while tuning the delay between Stokes and pump beam the slope of the Gaussian fit was extracted to determine the jitter to be ~ 24.3 fs. In f) the noise performance of the described home-built laser system (green and red lines) is compared to typical competing laser systems such as a super continuum fiber laser (SC) at 1.0 μm (Stokes, blue line) and 1.5 μm (as source for the pump beam, purple line), a fs-pulsed MaiTai laser (black solid line) as well as the shot noise limit as reference (black dashed line). The typical modulation frequency for SRS imaging is indicated by the black dotted box. Part of Kong et al. [2019].
Figure 79 – Pulse train characterization of both laser wavelengths over long time period
The pulse train of Stokes (green) and pump beam (red) are analyzed by detecting the laser wavelength with fast photodiodes on a 20GHz real-time oscilloscope over a time period of 500µs. The normalized intensity shows very low changes. Part of [Kong et al. 2019].

Figure 80 – Radio frequency (RF) spectra for characterizing the two-color pulsed fiber laser
(a) shows the RF spectrum of the non-modulated Stokes beam centered at the fundamental repetition rate of the laser system at ~ 80MHz showing the signal-to-noise characteristic of 74dB. Applying the frequency modulation at 20MHz with the fiber-based modulator for SRS imaging (b) leads to a detected modulation performance of 67db. In (c) the frequency doubled pump beam is analyzed showing a SNR of 73db at 80MHz. Part of [Kong et al. 2019].
Figure 81 – Spectral characterization of the two-color fiber laser source

(a) shows the accessible coarse tuning wavelengths of the 1,0µm master cavity by rotating the intracavity filter F1 (8nm bandwidth). In (b) the slit-based filter scheme of the grating compressor is utilized for a fine tuning from 1010 – 1060nm. (c) depicts the tuning range of the fundamental pump beam from 1540 – 1590nm by rotating the intracavity filter F2 (<1nm bandwidth). (d) showcases the final pump beam wavelength generated by the PPLN via frequency doubling with a central wavelength at 789nm having a bandwidth of 0.35nm. All spectra were acquired by a spectrum analyzer and are normalized. Adapted from [Kong et al. 2019].

Figure 82 – Comparing SRS and spontaneous Raman spectra on pure samples

By wavelengths tuning of the Stokes beam the molecular resonance of two pure samples (DMSO - red line, methanol - blue line) is acquired from 2800 – 3050cm⁻¹ and compared to spontaneous Raman spectra (SR, green DMSO and violet methanol, respectively). All intensities were normalized and vertically shifted for easy comparison. The SRS spectra were acquired by scanning 5× zoomed images and averaging the intensity values of the pixels. The SR spectra were acquired by the home-built spontaneous Raman setup utilizing a 785nm Cw-laser. Adapted from [Kong et al. 2019].
4.6.3. All fiber-based two-color laser system enables simultaneous SRS and CARS imaging on living human cells

For exploring the performance of the fiber laser system living osteosarcoma (U2OS) cells are investigated by probing the 2845 cm$^{-1}$ $CH_2$ stretching mode in order to highlight the lipid distribution and the cellular border (fig. 83 a CARS and b SRS). The cells were cultivated in DMEM media (ThermoFisher) containing 10% fetal bovine serum (ThermoFisher) on a standard petri dish stored in a humidified incubator at 37°C, 5% $O_2$ and 5% $CO_2$. Shortly before the imaging process the medium was replaced by phosphate buffered saline (Sigma Aldrich, PBS). The focal pump beam intensity was 33mW the modulated Stokes beam 74mW, respectively. The SRS images were acquired by setting a intensity modulation frequency of 20MHz of the Stokes beam (see also 3.1.6). The pump beam and the CARS signal were separated by a dichroic mirror (Semrock, LP02-785RU) and the CARS signal is further filtered (FF01-655/40-25). The pump beam is filtered by carefully suppressing the Stokes beam (Semrock, FF01-855/210-25, F01-950SP), detected on the fast SI-photodiode module and finally demodulated by a fast lock-in amplifier (LIA) (APE Berlin, SRS Detection Set). The integration time constant of the LIA was 2$\mu$s enabling a pixel dwell time of the laser scanner of 12,5$\mu$s for both contrast methods. No balanced detection scheme was needed for the SRS imaging. The CARS wavelengths were separated by the dichroic beam splitter and detected on the PMT. No stripes can be observed in the simultaneously acquired CARS images due to the slower response time of the PMT, which masks the 20MHz Stokes intensity modulation. For easy comparison all presented CARS images are processed with the look-up table (LUT) “cyan hot”, the SRS images are processed with the LUT “red hot” by the image software FIJI [Schindelin et al. 2012, Rueden et al. 2017]. The intensity of each signal is color-coded in the LUT and optimized for best contrast for each image and can differ from image to image. Therefore, a color bar is depicted beside each image to keep the intensity distributions transparent. Single prominent lipid droplets in the cells are highlighted by arrows, the cellular border of a selected cells is indicated by a white dashed circle.

In the next step living primary myoblast cells (PMD) were imaged during their differentiation process by applying 25mW pump and 47mW modulated Stokes focal power setting a pixel dwell time of 51,2$\mu$s combined with a LIA time constant of 20$\mu$s (fig. 83 c CARS and d SRS). The PMD cells were cultivated in DMEM containing 5% horse serum (ThermoFisher) and prepared on petri dishes coated with an extracellular matrix (Sigma Aldrich) at 1 : 500. The cells contain less lipid contents and, therefore,
exhibited less signal at the 2845 cm$^{-1}$ Raman resonance in combination to the lower laser intensity applied to this sample the pixel dwell time had to be increased compared to the U2OS cells. Nevertheless, single lipid droplets (showcased by the arrows) as well as the cellular border (white dashed circle) are clearly discriminable. Also the advantages of SRS due to the lack of the non-resonant background contribution is apparent in the images: the nucleus area of the cell indicated by the white dashed circle in fig. 83c shows an inner structure, which is not visible in the SRS channel. Here, the background-free SRS responds only to the aliphatic CH$_2$ groups, whereas the CARS channel generates also unspecific signals, which lowers the chemical specificity.

Figure 83 – Comparing SRS and CARS imaging performance on living human cells
In a) and b) simultaneous acquired CARS and SRS images probing the 2845 cm$^{-1}$ resonance in living osteosarcoma cells (U2OS) are presented. The focal power was set to be 33 mW pump and 74 mW Stokes beam (20 MHz modulation), respectively. The pixel dwell time was 12.5 µs while setting the time constant of the lock-in amplifier (LIA) to 2 µs. c) and d) show living primary myoblast cells (PMD) in their differentiation process by again probing the 2845 cm$^{-1}$ resonance. The focal power of the pump beam was 25 mW and 47 mW for the Stokes beam, respectively. The pixel dwell time is 51.2 µs in combination with a LIA time constant of 20 µs. The arrows indicate selected lipid droplets in the cellular area. Part of [Kong et al. 2019].
4.6.4. Investigating the performance of the all fiber-based laser on tissue samples utilizing simultaneous SRS and CARS contrast

A tissue section of a 2-month-old mouse, which suffered of severe combined immunodeficiency, was imaged by the dual channel approach using CARS and SRS as described before. The tissue was embedded in paraffin and cut into 50µm thick sections by a cryo microtome and transferred to a thin cover slide. An overview of the investigated slide is created by a bright-light setup with 4× magnification (fig. 84 a) showing the superior Vena cava (border indicated by the violet dotted lines). For the CRS experiments two ROIs were selected (red and green boxes) and a larger field of view is generated by moving the sample and then stitching the overlapping sequentially acquired images (fig. 84 b CARS and c SRS). The focal powers were set to 33mW pump and 74mW Stokes beam with 20MHz intensity modulation frequency. The pixel dwell time was set to 6, 4µs and the lock-in amplifier time constant was 2µs. In the images the border between Tunica media and Tunica intima can be observed, a violet dotted line is inserted to guide the eye. The yellow inset in fig. 84 b shows a zoom image of red blood cells forming a thrombus (yellow dotted circle), which was adherent to the vascular wall. The CRS images of ROI 2 are presenting a lipid rich area in the Tunica media (fig. 84 d and e, CARS and SRS, respectively). The green line indicates the location of a line scan comparing both imaging modalities to each other (fig. 84 f). The normalized intensity profiles show again the superior performance of SRS to CARS due to the small features visible in the SRS channel (indicated by red arrows), where in the CARS channel the non-resonant background masks them leading to a diminished chemical sensitivity.

In a further experiment a murine brain tissue sample was investigated in the same manner. The sample was extracted of the mouse, cut by a cryo microtome into 50µm thick slices and was finally fixed by using a 4% para-form aldehyde solution. Two regions of interest are selected for the dual channel imaging via SRS and CARS investigating the neuron network as well as the distribution of the granule cells in the hippocampus (fig. 85). During the imaging the focal power of the pump beam was set to 33mW the Stokes beam to 67mW by using a pixel dwell time of 12, 8µs and a LIA time constant of 2µs. The image contrast and sensitivity is very satisfactory and will be exploited to perform large area scans in future projects [Ruiz-Perera et al. 2018, Klenke et al. 2013], enabling the identification of the individual brain sections and following the transitions into each other, especially the formation of the granule cells.

26The samples are provided by Prof. B. Kaltschmidt and Prof. C. Kaltschmidt.
**Figure 84** – CRS-imaging of murine superior Vena cava tissue cryosection

*a*) shows a bright-field overview image of the analyzed cryosection. For further analysis with simultaneous SRS and CARS imaging two ROIs are selected (red and green boxes). The violet dotted line indicates the location of the superior Vena cava.  

*b*) shows the CARS and *c*) the SRS images of ROI 1, which were stitched showing a large field-of-view. They allow the discrimination of Tunica media and Tunica intima. The border between them is indicated by the added pink dotted line. The yellow inset shows a zoom image visualizing a thrombus persisting of red blood cells adhering to the vascular wall.  

*d*) and *e*) show the stitched CARS and SRS images of ROI 2 detailing the Tunica media with lipid rich structures. The green line in *e*) depicts the location of a comparing line plot, which is presented in *f*). Here, small lipid rich structures are revealed by SRS, due to the higher chemical contrast (red arrows) by the advantage of having no non-resonant background in SRS. For all images the focal pump power was set to 33$mW$ and 74$mW$ for the Stokes beam, having a pixel dwell time of 6,4$\mu s$. The LIA time constant is set to be 2$\mu s$ in combination with the applied 20$MHz$ modulation frequency in the Stokes branch. Part of [Kong et al.] [2019].
Murine brain sections are investigated by simultaneous CARS and SRS imaging at two different locations on a 50\(\mu\)m thick cryo-section (a CARS and b SRS of ROI 1 as well as c and d of ROI 2, respectively). The focal power was set to 33\(mW\) and 67\(mW\) for the pump and Stokes beam, respectively. The pixel dwell time was 12, 8\(\mu\)s, while having a LIA time constant of 2\(\mu\)s in combination with the applied 20\(MHz\) modulation frequency in the Stokes branch. The scale bar depicts 20\(\mu\)m.
4.6.5. Fs-laser pulses of Stokes cavity are utilized for SHG and two-photon excited fluorescence imaging broadening the range of application

The multimodal imaging capability of the introduced laser system is demonstrated by generating $\sim 200\text{fs}$ pulsed laser light by removing the filtering slit in the grating compressor stage of the Stokes beam generation, which then provides a spectral bandwidth of $\sim 8\text{nm}$ with a central frequency of $\sim 1025\text{nm}$. An unstained mouse tail sample was harvested from a 3-month old C57BL/6 mouse and selected for demonstrating SHG generation at $\sim 512\text{nm}$ (fig. 86a) by applying $\sim 20\text{mW}$ focal power and controlling the polarization of the fs-laser beam by a combination of a half- and a quarter-wave plate (one state of polarization is shown in the image). The collagen in the tendon tissue of the sample generates the detected SHG signal, while the surrounding muscle tissue remains without signal contributions. Two-photon excited fluorescence imaging was performed using 20$mW$ focal power on a murine kidney section (ThermoFisher, FluoCells prepared slide #3), which is stained with three individual fluorescent dyes. Two optical bandpass filters (centered at 519nm and 600nm) are separating two fluorescent stains in the experiments for sequential imaging: Alexa Fluor 568 phalloidin is detected labeling the filamentous actin cytoskeleton prevalent in the glomeruli and brush border (fig. 86b) as well as Alexa Fluor 488 wheat germ agglutinin labeling elements of the glomeruli and the convoluted tubules (fig. 86c) (interpretation based on [Wiederschain [2011]]).

For demonstrating the z-sectioning capabilities of the system a z-scan over 145$\mu\text{m}$ range with a step size of 5$\mu\text{m}$ is acquired of a 200$\mu\text{m}$ thick mouse brain tissue prepared between two thin cover slides. The mouse is expressing the yellow fluorescent protein (YFP) in layer-V pyramidal neurons (Thy1-YFP H-line) [Lai et al. [2012]], which was excited by $\sim 50\text{mW}$ focal power, indicating the neuron distribution in the tissue. Fig. 86d shows a 3D montage of the data stack generated by FIJI volume viewer as well as in fig. 86e a selected single xy-slice of the stack showcasing the neurons and, thereby, the good z-sectioning of the two-photon excitation.

The application of the Stokes-laser branch as a versatile and low-cost source for SHG and two-photon excited fluorescence was also proposed in a stand-alone version [Kong et al. [2017]]. In general also 3-photon excited fluorescence [Wang et al. 2018, Horton et al. 2013] can be performed by this laser source utilizing the fundamental pump beam cavity laser light (1578nm), but due to the lack of sufficient optics (e.g. objective lens corrected for $\sim 1500 - 1700\text{nm}$, mirrors and telescope lenses in sufficient
anti-reflective coating in this range) this modality has not been pursued so far.

Figure 86 – SHG imaging of mouse tail sample and two-photon excited fluorescence imaging of kidney and brain tissue sections

The fs-pulsed Stokes beam (bandwidth $\sim 8\text{nm}$, $\sim 200\text{fs}$ pulse length) can be utilized for e.g. SHG imaging (a) of a mouse tail sample by applying $\sim 20\text{mW}$ focal power. The polarization of the laser was controlled by a stack of half- and quater-wave plate. Two-photon excitation is demonstrated with the same source and focal power on a mouse kidney section (ThermoFisher, FluoCells prepared slide #3) visualizing two different fluorescent stains of the sample sequentially by using a suitable optical filter set (b, Alexa Fluor 568 phalloidin labeling the filamentous actin cytoskeleton prevalent in glomeruli and the brush border; c, Alexa Fluor 488 wheat germ agglutinin labeling elements of the glomeruli and the convoluted tubules [Wiederschain [2011]]. In d) and e) the two-photon excitation of a $200\mu m$ thick mouse brain tissue section prepared between two cover slides is demonstrated in 3D imaging modality. The mouse was expressing the yellow fluorescent protein (YFP) in layer-V pyramidal neurons (Thy1-YFP H-line) [Lai et al. [2012]]. The z-stage was utilized for scanning a range of $145\mu m$ with $5\mu m$ step size. d) shows a 3D montage (performed via FIJI volume viewer) indicating the distribution of the neurons in the tissue. e) showcases a single frame of the stack, the focal power was set to $\sim 50\text{mW}$. Part of [Kong et al. [2019]].
4.6.6. Extending the range of accessible wavenumbers with a third laser cavity enabling the visualization of alkyne tagged samples in the future

The introduced all fiber-based CRS laser source has been characterized and compared to state-of-the-art fiber-based and traditional solid-state laser systems determining the unique selling point of this laser system in having a superior noise performance, which allows the rapid SRS imaging without the need of a balanced detection scheme for laser noise cancellation. The abilities were demonstrated by imaging relevant living human cells as well as different tissue sections by a dual SRS and CARS contrast in simultaneous rapid acquisition manner. For proving the versatility of the setup also SHG and two-photon excited fluorescence imaging are showcased on several samples combined with z-scanning. In table 1 the demonstrated properties of the two-color all fiber-based laser system are compared to other laser systems using fiber-laser technology for generating one or both laser wavelengths for CRS microscopy known from literature. Here, the novelty of the introduced approach becomes apparent. The next steps in the development of the all fiber-based laser system are inserting an AOTF (acousto-optic tunable filter) into the Stokes beam cavity for enabling rapid automated wavelengths tuning in order to allow a hyperspectral image scan similar to the previously described modality based on the solid-state laser system (compare 4.4). Also the use of all fiber-based electronically controlled Fabry-Pérot wavelengths filters inside the laser cavities can replace the manual wavelengths tuning and enable tuning speeds of up to 100 kHz [Huber et al. 2006].

In addition to that the range of accessible Raman shifts will be further extended by adding a third oscillator cavity as alternative second pump beam for CRS-microscopy. Especially samples with alkyne tags such as e.g. EdU (ethynyl deoxyuridine) are of particular interest due to their Raman resonance in the range of $\sim 2100 \text{cm}^{-1}$, which is located in the “Raman-silent region” ($1800 - 2600 \text{cm}^{-1}$) [Zhao et al. 2017]. Here, no Raman resonances of the typical chemical compounds inside biological cells are active. Therefore, also very small concentrations of alkyne-tagged molecules are accessible by CRS-microscopy. Several studies have been performed utilizing the method of introducing alkyne-tagged molecules into the cells metabolism and finally tracing their distribution by using the label-free CRS-imaging [Shi et al. 2018, Chen et al. 2014, Wei et al. 2014a]. Still, up to now the use of fiber-based laser sources is hindered in the range of the desired Raman shifts due to the lack of sufficient material combinations for the fiber doping and the connected pumping schemes. In the ongoing project new gain materials will be explored combined with the use of additional fiber-based SESAM mirrors (semi-saturable absorber mirrors) in the laser cavity [Qi et al. 2016] for improving the mode locking ability of the laser cavity in the case of more
problematic gain media. Promising candidates are e.g. neodymium-doped (∼ 900nm), bismuth-doped (1100 – 1500nm) and thulium-doped (1700 – 2000nm) fibers. When a sufficient combination is identified a third laser oscillator can be added for generating a second pump laser wavelength, which then can be directed into an additional branch for the frequency doubling to the desired final pump beam wavelength. This way a dual contrast of e.g. the lipid distribution in a cell by probing the 2845cm⁻¹ CH₂ resonance and the alkyne tagged molecules, e.g. 2120cm⁻¹ in the case of EdU, can be exploited for studying the distribution of chemicals during the cells live metabolism. For optimizing the robustness of the source while at the same time minimizing the footprint of the setup, the frequency doubling can also be performed by using fiber-integrated PPLN modules, which would allow a sealed one box system ideal for clinical application [Runcorn et al. [2015]].
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<td>Aquatic plant, mouse brain, blood cells</td>
<td>Mouse skin, human surgical specimens</td>
<td>living U2OS cells, living PMD cells, mouse superior Vena cava tissue, mouse brain tissue</td>
</tr>
</tbody>
</table>

**Table 1** – Comparison of CRS laser sources

HFL: hybrid fiber laser [Ozeki et al. [2010], Wang et al. [2010, 2013], Ozeki et al. [2012], ASFL: actively synchronized fiber laser [Karpf et al. [2015], Nose et al. [2012], PWCFL: parametric wavelength conversion fiber laser Zhai et al. [2011], Baumgartl et al. [2012a], Lefrancois et al. [2012], Yang et al. [2018], [Baumgartl et al. [2012b], Chemnitz et al. [2012], Lamb et al. [2013], Gottschall et al. [2014]], SSFSFL: soliton self-frequency shift fiber laser [Andersen et al. [2007], Pegoraro et al. [2009], Krauss et al. [2009], Xie et al. [2014], Crisafi et al. [2018]], SCFL: super continuum fiber laser Gambetta et al. [2010], [Sehn et al. [2010], Freundiger et al. [2014], Chen et al. [2016], Tu et al. [2016], [Orringer et al. [2017], Riek et al. [2016], Coluccelli et al. [2014]], *: referring to solid-state lasers. Red color highlights drawbacks compared to the introduced home-built fiber-based laser system (shown in blue).
5. Summary and outlook

This PhD thesis describes the development of a coherent Raman scattering microscope and its use for multimodal imaging of several biomedically relevant samples. This instrument enables the simultaneous acquisition of images with SRS and CARS contrast in addition to second harmonic generation and two-photon excited fluorescence imaging in forward and epi-detection. The sample can be moved in all three spatial dimensions allowing volumetric imaging and large fields of view by automated sample movement and image stitching [Hofemeier et al. [2016]]. By utilizing rapid intensity modulation in combination with lock-in amplification undesired fluorescence signals can be successfully excluded from the desired CARS contrast, which was demonstrated with different samples for quantitative analysis [Jaeger et al. [2016]]. The setup can address a wide range of molecular resonances due to the use of two optical parametric oscillators, which also allow hyperspectral scanning by automated wavelength tuning of the pump beam in the CRS experiment. Here, it was possible to introduce a substantially improved level of reproducibility due to the optimization of the mechanical stability of the setup by customizing optomechanical components and by developing a specifically adapted control software, which was shared with other research groups [Pilger et al. [2018]].

This hyperspectral scanning capability was successfully applied to a range of biomedical samples. The combination of rapid volumetric imaging addressing a single molecular resonance in addition to hyperspectral CARS at selected sample positions was found to be a useful match to answer specific research questions that could not be analyzed by other means [Vukosavljevic et al. [2019], Fueser et al. [2018]]. For exploring the possibility of enhancing the spatial resolution of CARS microscopy in a gentle and bio-compatible way the concept of structured illumination was introduced in the setup. This was realized by rapid intensity switching and the precise control of the scanning mirrors in order to generate a striped illumination pattern in the sample. By simply adding a modern sCMOS camera for detection most of the components are already part of modern CRS or two-photon scanning microscopes, which offers enhanced resolution by low additional costs. This technique was successfully demonstrated with fluorescent samples by two-photon excitation as well as on polystyrene bead samples with CARS contrast.

For the support of CRS microscopy in clinical applications a novel low noise two-color all-fiber-based laser source was developed in a collaboration with the University of Hong Kong [Kong et al. [2019]]. This source provides very low relative intensity noise in combination with superior laser power compared to other, fiber-based sources, which makes
it a perfect tool for CRS microscopy, especially for SRS. Here, several biomedical samples were investigated by simultaneously acquiring SRS and CARS signals showcasing the superior performance of the laser source without the need for balanced detection in the SRS detection path. The laser source is characterized in additional experiments and its performance is compared to other fiber laser setups reported in the literature. It also offers multimodal imaging capability through minor changes in the setup allowing SHG and two-photon excited fluorescence imaging by acting as femtosecond pulsed laser source. This solution is also proposed as a stand-alone setup, making it a cost-effective source for two-photon scanning microscopes [Kong et al. [2017]].

In the future, the CRS microscope will be further enhanced through the development and implementation of an additional laser oscillator into the fiber laser setup, which extends the range of addressable wavenumbers to the Raman-silent region (1800 – 2600 cm$^{-1}$), where in typical biomedical samples no natural molecular resonances are present. This can be exploited for imaging alkyne-tagged or deuterated samples, which offer a unique molecular signature and, thereby, lead to an enhanced sensitivity [Shi et al. [2018], Wei et al. [2014a], Chen et al. [2014]].

Also, this technique can be applied in time-lapsed experiments to the tracking of alkyne-tagged virus particles or circulating tumor cells, which would address a topic of high interest in biomedical research [Hübner et al. [2009]]. The SI-CARS approach can be further explored to allow for the imaging of virus particles, the size of which is well below the typical spatial resolution of a CRS microscope. Here, the low non-resonant background in the Raman-silent region will lead to an optimized signal to noise performance, which is beneficial for the SI-CARS application. Ultimately, the use of doubly-resonant CARS, which is actively probing two molecular resonances at the same time leading to a strongly enhanced signal strength [Weeks and Huser [2010]], can be utilized at specifically designed molecules. This will enable investigating the metabolic incorporation of these molecules into living cells. With these tools it will then be possible to follow the distribution of the molecule of interest over time, unlimited by bleaching effects and without the side-effects of fluorescence labeling.
6. List of publications


- *Compact high-power fs ytterbium fiber laser at 1010nm for biomedical applications*; Kong†, C., Pilger†, C., Hachmeister†, H., Wei, X., Cheung, T., Lai, C., Huser, T., Tsia, K. K., Wong, K. K. Y., Biomedical Optics Express: 8(11), 4921-4932, (2017) {†: equal contribution}


A. Statutory declaration

I hereby declare that I completed the submitted thesis with no other help and literature than the one known to the reviewers. Sentences or parts of sentences quoted literally are marked as quotations, identification of other references with regards to the statement and scope of this work is quoted.

Bielefeld, 24. 06. 2019

Christian Pilger
B. Acknowledgments

I would like to thank Prof. Thomas Huser for the chance to produce this thesis and for his mentoring and continuous interest in all things related to the CARS lab and the work we did there. Thomas, I really enjoyed the amount of freedom you allowed me during the realization of all tasks in the lab and the great and steady input of new ideas, which seemed to come in a steady natural flow to you. A big additional “Thank you!” for the great and fruitful idea of forming our collaboration with the University of Hong Kong and the financial support you provided for this work, which also allowed me to experience a new fascinating city and find new friends.

I also want to thank Prof. Anselmetti, Prof. Hütten and Prof. Borghini for joining the examination committee of this thesis. Furthermore, I thank Dr. Gerd Wiebusch for the support with his experience during the work in the lab, providing good council when there was a problem to be solved. Also I want to express my gratitude for proofreading this thesis, which is always a laborious task.

Special thanks go to Henning Hachmeister: we started the study of physics together a long time ago and shared a great part of the last 10 years solving problems together while acting as our sparring partners. I really enjoyed our time in the lab, building our own setup and you keeping the spirits up during long measurement sessions, while we always had joint interests besides the work. Thank you so much!

Really big thanks go to Prof. Kenneth Wong and his research group at the HKU for providing the chance for the fruitful collaboration. I enjoyed the heartfelt hospitality you always provided me during the visits in Hong Kong very much.

In addition to that I would like to thank Sherry Kong and Dr. Xiaoming Wei for the great time we had together during the entire collaboration applying fiber lasers to CRS microscopy, which took a longer time than initially expected. I want to thank both of you for the friendship we formed and also for introducing me to the “very local” places in Hong Kong, which a European tourist would never find on his own.

I’m thanking Alex Weiß and Paul Greife for the great collaboration during their master theses, it was a pleasure to work with you.

I thank Prof. Olaf Kruse and Dr. Daniel Jäger for the well planned and structured collaboration on the microalgae-related work. Daniel, I really appreciated our joint time in the lab.

I also want to thank Prof. Maike Windbergs and Dr. Branko Vukosavljevic for the collaboration on the human lung cell differentiation: Branko, we had a tough long week
of measurements and an even tougher way until the paper was accepted, but finally it worked out for all of us.

I want to thank Prof. Barbara and Prof. Christian Kaltschmidt for the collaboration on multiple topics and the great interest they showed during our collaboration in the CRS-technique, always being generous with their time to come in our lab for having a look at things and also for providing so many various samples.

Many thanks go to Prof. Walter Traunspurger and to Hendrik Füser for the collaboration on C. elegans, which was a very interesting task and worked well when our laser was not making trouble.

Dr. Marcel Müller I want to thank for his the support on topics related to structured illumination and FairSIM.

I also thank Jakub Pospíšil for the collaboration concerning MAP-SIM and all the nice discussions on related topics.

Additional thanks go to Dr. Mathias Simonis for always being generous providing cell samples.

I’m thanking all collaborators from LaVision Biotec for the exciting time during the collaboration; it was a great experience to get insight into industrial-related research. Your support was very generous and friendly!

To all my office mates I say thanks a lot for the fruitful atmosphere and the nice conversations about not work-related topics.

My thanks also go to all members of the research group Biomolecular Photonics for the good working atmosphere and the team spirit, which I enjoyed very much!

I thank Dr. Mark Schüttpelz for his support concerning MatLab and also for nice conversations on brewing techniques.

Furthermore, I’m thanking Reinhild Pätzmann for all her labors concerning the administration matters such as contracts, travel expenses or even custom matters. You always supported the scientific work, many thanks for that.

Also big thanks go to all members of the mechanical and electronic workshop of the faculty of physics, University of Bielefeld, especially to Karl-Heinz Eichner, Wolfgang Gronemeyer, Uwe Walkenhorst, Herbert Bergmeier and Dieter Gollay; without them many technical nice and home-made solutions and devices utilized in this thesis would have not been realized or at least not with that high quality and love to the detail.

Special thanks go to my family, who supported me with their love, encouragement and their interest in my studies! An additional “Thank you!” I want to direct to my sister Diana for proofreading of this thesis while being not an expert on the topic but instead on the language. This fact turned out to be beneficial for finding errors but
also laborious sizing “our physicist” terms.
Finally, my biggest heartfelt thanks go to my wife Nadine Pilger for keeping me grounded all the time with her love and endurance!
References


G. Donnert, J. Keller, R. Medda, M. A. Andrei, S. O. Rizzoli, R. Luhrmann, R. Jahn, C. Eggeling, and S. W. Hell. Macromolecular-scale resolution in biological fluores-


C. W. Freudiger, W. Min, B. G. Saar, S. Lu, G. R. Holton, C. He, J. C. Tsai, J. X. Kang, and X. S. Xie. Label-Free Biomedical Imaging with High Sensitivity


Andrew Fussell, Erik Garbacik, Herman Offerhaus, Peter Kleinebudde, and Clare Strachan. In situ dissolution analysis using coherent anti-Stokes Raman scattering (CARS) and hyperspectral CARS microscopy. *European Journal of Pharm-


Hamamatsu Photonics. *OrcaFlash 4.0 vers. 1.1, C11440-22C. Instruction manual*, 2012.


Stefan Hauser, Darius Widera, Firas Qunneis, Janine Müller, Christin Zander, Johannes Greiner, Christina Strauss, Patrick Lüningschröer, Peter Heimann, Hartmut


Wei-Wei Hsiang, Chia-Hao Chang, Chien-Po Cheng, and Yinchieh Lai. Passive synchronization between a self-similar pulse and a bound-soliton bunch in a two-color


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Keisuke Nose, Yasuyuki Ozeki, Tatsuya Kishi, Kazuhiko Sumimura, Norihiko Nishizawa, Kiichi Fukui, Yasuo Kanematsu, and Kazuyoshi Itoh. Sensitivity enhancement of fiber-laser-based stimulated Raman scattering microscopy by collinear


Lingyan Shi, Yihui Shen, and Wei Min. Invited Article: Visualizing protein synthesis


Wenlong Yang, Ang Li, Yuanzhen Suo, Fa-Ke Lu, and X. Sunney Xie. Simultaneous two-color stimulated Raman scattering microscopy by adding a fiber amplifier to a 2


Dai Yoshitomi, Yohei Kobaishi, Masayuki Kakehata, Hideyuki Takada, Kenji Torizuka, Taketo Onuma, Hideki Yokoi, Takuro Sekiguchi, and Shinki Nakamura. Ultralow-jitter passive timing stabilization of a mode-locked Er-doped fiber laser by


