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Spectroscopic infrared near-field microscopy and x-ray reflectivity studies of order and clustering in lipid membranes
Label-free spectroscopic detection of membrane potential using stimulated Raman scattering

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Hyperspectral stimulated Raman scattering microscopy is deployed to measure single-membrane vibrational spectrum as a function of membrane potential. Using erythrocyte ghost as a model, quantitative correlation between transmembrane potential and Raman spectral profile was found. Specifically, the ratio between the area under Raman band at \(\sim 2930\) cm\(^{-1}\) and that at \(\sim 2850\) cm\(^{-1}\) increased by \(\sim 2.6\) times when the potential across the erythrocyte ghost membrane varied from \(+10\) mV to \(-10\) mV. Our results show the feasibility of employing stimulated Raman scattering microscopy to probe the membrane potential without labeling. © 2015 AIP Publishing LLC.

The plasma membrane, which consists of a double layer of phospholipids with embedded proteins, is the outer boundary of a biological cell. The cell membrane functions as a physical barrier isolating the cytoplasm from the extracellular environment, thus protecting the cell from its surroundings. Since the cell membrane is selectively permeable, it regulates and organizes the cell by controlling the exchange of substances. In addition to these fundamental roles, the cell membrane also mediates the communication between different cells. Changes in the transmembrane potential can generate action potential and give cells the capability to convey messages and communicate with other partners. Therefore, methods of probing cell membrane properties are of extreme importance.

Fluorescence labeling is the most widely used method for cell membrane related study. Various fluorescent voltagesensitive indicators have been extensively used to sense the membrane potential.1 Immunofluorescence staining is a well-established method for the analysis of cell polarity.2 Cell membrane fluidity is frequently analyzed by monitoring the dynamics of fluorophores which tag either membrane proteins or lipids to mimic the behavior of cell membrane.3 For these fluorescence based approaches, perturbation of the membrane properties by chromophores is often a concern.

As a label-free analytical tool, Raman spectroscopy has been used to measure the responses of membranes to variations of temperature, pH, and transmembrane ion gradients.4,5 Because spontaneous Raman scattering is weak, extremely high laser power (400 mW at sample) was used in the previous study,7 which is close to the physical damage threshold. The weak signal also hinders the application of spontaneous Raman scattering as a dynamic imaging tool. To address this problem, coherent Raman scattering microscopy techniques6 were recently developed, which increased the acquisition speed by several orders of magnitude through coherent excitation of Raman bands in a spectral window of interest. With coherent anti-Stokes Raman scattering (CARS) microscopy,7 the orientation of lipid bilayers inside the erythrocyte ghost membrane was visualized by Potma and Xie.8 Compound Raman microscopy,9 which combines advantage of high-speed vibrational imaging offered by CARS and quantitative spectral analysis by spontaneous Raman scattering, offers an alternative way to study the membrane. Using a compound Raman microscope, Yue et al. demonstrated label-free analysis of apical polarity of live cell membranes in 3D culture.10

In this work, we present the use of stimulated Raman scattering (SRS) microscopy11 to detect the membrane potential in a label-free manner. SRS gives identical spectral profile as spontaneous Raman scattering and it provides several orders of magnitude stronger signal through stimulated scattering. By offering spectral information at each pixel, hyperspectral SRS microscopy has allowed for quantitative chemical imaging of cell12,13 tissue,14,15 and Caenorhabditis elegans.16 Herein, we demonstrate the employment of hyperspectral SRS microscopy for probing the transmembrane voltage with erythrocyte ghost as a model system.

Our hyperspectral SRS imaging system (Fig. 1) was built based on a previously reported spectral focusing scheme,13,17 where the Raman shift is controlled by the temporal delay of two chirped femtosecond pulses. An ultrafast laser system with dual output operating at 80 MHz (InSight DeepSee, Spectra-Physics) provided the excitation sources. The tunable output with a pulse duration of 120 fs was tuned to 800 nm, serving as pump beam. The other output centered at 1040 nm with a pulse duration of 220 fs was used as Stokes beam, modulated by an acousto-optic modulator (AOM, 1205-C, Isomet) at 2.3 MHz. An optical delay line was installed in the pump light path and a motorized

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**References**

translation stage (T-LS28E, Zaber) was employed to scan the delay between two beams. After combination by a dichroic mirror, both beams were chirped by two 12.7 cm long SF57 glass rods. In order to match the linear chirp parameter of two beams, the inbuilt prism compressor inside the laser system was used to adjust the dispersion of the pump beam. After glass rods, pulse durations of the pump and Stokes beams were stretched to ~1.3 and ~0.8 ps, respectively, measured with an autocorrelator (PulseScope, APE). The pump and Stokes beams were sent into a home-built laser-scanning microscope described elsewhere.\textsuperscript{15,18} Briefly, a 60\textdegree water immersion objective lens (NA = 1.2, UPlanApo/IR, Olympus) was utilized to focus the light into the sample, and an oil condenser (NA = 1.4, U-AAC, Olympus) was used to collect the signal. The stimulated Raman loss signal was detected by a Si photodiode (S3994–01, Hamamatsu) and extracted with a digital lock-in amplifier (HF2LI, Zurich Instrument).

We first calibrated the Raman shift with respect to the pump-Stokes temporal delay using the Raman peaks of known chemicals. Fig. 2(a) shows the spontaneous Raman spectra of dimethyl sulfoxide, methanol, acetic acid and cyclohexanone in the CH stretching region from 2800 to 3050 cm\textsuperscript{-1}, measured with a commercial spectrograph (Shamrock SR-303i-A, Andor Technology). Corresponding SRS spectra recorded by scanning the pump-Stokes pulse delay are shown in Fig. 2(b). The SRS intensity was normalized with the two-photon absorption signal of Rhodamine 6G which approximately reflects the cross-correlation between the pump and Stokes pulses. Using these distinguished Raman peaks shown in Fig. 2(a), the translation stage positions were then correlated to the Raman shifts by linear fitting, resulting in the calibration curve shown in Fig. 2(c). The correlation is almost linear (R\textsuperscript{2} = 0.995), which indicates that the laser pulses were nearly linearly chirped. The spectral resolution of our system is ~25 cm\textsuperscript{-1}, estimated from the SRS spectra of dimethyl sulfoxide and methanol.

Using the hyperspectral SRS imaging system, we recorded Raman spectrum from single erythrocyte membrane which is an established model system for membrane biochemistry.\textsuperscript{15,19} Fig. 3(a) shows a representative stack of hyperspectral SRS images of an erythrocyte ghost. Preparation of erythrocyte ghosts can be found in detail in supplementary material.\textsuperscript{20} A total of 50 frames were recorded from 2784.82 cm\textsuperscript{-1} to 3041.83 cm\textsuperscript{-1} with a step increment of ~5.25 cm\textsuperscript{-1}. Each frame was composed of 40 x 40 pixels with a pixel dwell time of 200 l\textsuperscript{s}. Acquisition of the entire stack took ~28 s. A representative SRS spectrum from single natural membrane, obtained through averaging the hyperspectral data from the pixels indicated by the rectangular box in Fig. 3(a), is shown in Fig. 3(b). Lorentzian fitting was performed to decompose the spectrum into 7 bands according to previous studies.\textsuperscript{10,21} The spectrum analysis and fitting parameters can be found in

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure1.png}
\caption{Schematic of the hyperspectral SRS microscope. $\lambda/2$, half wave plate; PBS, polarizing beam splitter; AOM, acousto-optic modulator; DM, dichroic mirror; SF57, SF57 glass; Obj, objective; Con, condenser; PD, photodiode; LIA, lock-in amplifier.}
\end{figure}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure2.png}
\caption{Calibration of Raman shifts in the spectral focusing system. (a) Normalized spontaneous Raman spectra of dimethyl sulfoxide (DMSO), methanol (MeOH), acetic acid (AcOH), and cyclohexanone (CYC). (b) Corresponding SRS spectra measured by scanning pump-Stokes delay. (c) Correlation of the Raman shift with respect to the pump-Stokes delay.}
\end{figure}
Supplementary material. The Raman band around 2850 cm⁻¹ is largely contributed by the CH₂ groups in lipid bilayers. The band at ~2930 cm⁻¹ is largely contributed by the CH₃ groups in membrane proteins. As the pump and Stokes beams were parallel-polarized, the SRS signal intensity [inset of Fig. 3(a)] showed a clear dependence on the orientation of lipid bilayers with respect to laser polarization.

To prove the concept of probing transmembrane potential via SRS microscopy, we changed the potential across the erythrocyte membrane in the CH₃ stretching region. Previous studies have shown that transmembrane ion gradients, temperature, and pH influence Raman spectrum of the erythrocyte membrane in the CH₃ stretching region. Mikkelsen et al. suggested that intensification of the ~2930 cm⁻¹ band reflected transfer of protein CH₃ groups from the apolar to more polar environments. Besides, external environments like temperature and pH may involve a rearrangement of protein residue orientation, which can be reflected by the I₂₉₃₀/I₂₈₅₀ ratio as well. SRS is a nonlinear enhancement process of spontaneous Raman scattering, and it exhibits identical spectrum as spontaneous Raman. Consistently, the observed spectral profile change herein could be assigned to the transmembrane potential induced conformation changes of the membrane proteins.

In summary, with erythrocyte ghost as a model system, we showed the feasibility of using hyperspectral SRS microscopy to probe membrane potential in a label-free manner. By further improving the temporal resolution, SRS dramatically from 2.91 ± 0.27 to 7.59 ± 0.84 with the membrane potential changing from +10 mV to −10 mV.

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microscopy can be potentially used to measure the membrane potential and even record the action potential propagation in living cells. This imaging technique can also be extended to detect mitochondrial membrane potential and study the apoptosis process.

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20See supplementary material at http://dx.doi.org/10.1063/1.4919104 for details of the preparation of erythrocyte ghosts, intravesicular and extravesicular ionic compositions, and SRS spectrum analysis.