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Three-dimensional imaging of biological cells with picosecond ultrasونics

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We use picosecond ultrasونics to image animal cells in vitro—a bovine aortic endothelial cell and a mouse adipose cell—fixed to Ti-coated sapphire. Tightly focused ultrashort laser pulses generate and detect GHz acoustic pulses, allowing three-dimensional imaging (x, y, and t) of the ultrasonic propagation in the cells with ~1 μm lateral and ~150 nm depth resolutions. Time-frequency representations of the continuous-wavelet-transform amplitude of the optical reflectivity variations inside and outside the cells show GHz Brillouin oscillations, allowing the average sound velocities of the cells and their ultrasonic attenuation to be obtained as well as the average bulk moduli.

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Characterization of the physical properties of cells is essential for understanding their behavior such as mitosis, apoptosis, adhesion, and mobility.1–3 In particular, various techniques have been used to investigate the mechanical properties of single cells,4 for example, embedded particle tracking,5 magnetic twist cytometry,6 micro-patterned substrates,7 micropipette aspiration,8 optical traps or laser tweezers,9 optical stretchers,10 magnetic traps,11 microneedles,12 atomic force spectroscopy,13 and biomembrane force probes.14 These techniques mainly provide information concerning cell adhesion and deformation directly related to the cell membrane and cytoskeleton. In contrast, conventional acoustic microscopy allows the non-invasive study of the mechanical properties of the intracellular medium, i.e., the cytoplasm and nucleus.15 This technique is sensitive to the sound velocity, density, acoustic impedance, and bulk modulus of cells. Although images and sections of cells have been obtained with this method, full three-dimensional (3D) scanning of a single cell has not been reported.16,17

Recently, picosecond ultrasونics has been introduced to characterize single cells after decades of being used to investigate the mechanical and thermal properties of metals and semiconductors on sub-micron scales.18,19 This technique monitors GHz Brillouin oscillations in the time domain, allowing acoustic properties—sound velocity and ultrasonic attenuation—for the nucleus and vacuole of vegetal cells to be measured.20–22 In addition, cell density, compressibility and adhesion, as well as mechanical properties of cell walls have been studied with this technique in both vegetal and mammalian cells.23–27 In spite of this work, the technique has never been applied to obtain spatiotemporal data that allow 3D imaging of cell properties.

In this letter, we demonstrate 3D cell imaging with picosecond ultrasونics. We investigate two types of mammalian biological tissues—a bovine aortic endothelial cell and a mouse adipose, i.e., fat cell. Endothelial cells play an important role in physiology of blood vessels and are useful in the study of biomechanics.28,29 Fat cells provide an interesting comparison because of their different geometry and contents.

Fluorescence micrographs superimposed on background differential-interference- or phase-contrast images of a typical collection of each type of cell are shown in Figs. 1(b) and 1(c). A bovine endothelial cell,28,33 of thickness ~2 μm, is prepared by use of a standard formalin fixation technique on a 150 nm polycrystalline Ti film deposited on a sapphire substrate of thickness 0.5 mm. A mouse fat cell, of thickness ~10 μm, is differentiated from a commercial 3T3-L1 cell line34 and fixed in a similar manner. Each (unstained) cell is covered by a single-cavity microscope slide filled with saline and buffer solutions for the endothelial and fat cells, respectively, and each slide is then placed on an x–y translation stage, allowing 2D raster scans. The experimental setup is shown in Fig. 1(a). An ultrashort-pulse mode-locked Ti:Sapphire laser with a 75.8 MHz repetition rate and a 0.2 ps pulse duration is used for pumping and probing. A 830-nm pump beam of pulse energy 0.1 nJ (ensuring linear response) is focused by a 50× objective lens at normal incidence to a ~3 μm diameter (full width at half maximum) spot on the film from the substrate side, heating the Ti film from underneath at a chopping frequency of 1 MHz. A signal at the chopping frequency is coupled to a lock-in amplifier for high signal-to-noise ratio measurement.19 A series of GHz picosecond ultrasonic pulses traverse the cell sample, and are monitored by a 415-nm (frequency-doubled) probe beam with a pulse energy of 0.03 nJ from the opposite side. This beam is focused to ~2 μm diameter spot by a 50× objective lens sharing the same central axis with the pump beam and temporally scanned up to 800–1000 ps (after each ultrasonic pulse arrives at the Ti-solution interface) by use of a mechanical delay line. The optical beams impose an overall steady-state temperature rise at the Ti-solution interface...
As the ultrasonic pulses, generated with strain amplitudes of \( \times \) any evident damage. (Time scans were executed for each and \( \times \), temporal scanning allow 3D imaging (STORM).\(^3\(^1\)\(^3\(^2\))\(^4\) For quantitative processing, we identify such movies\(^3\(^0\)) allow one to trace the ultrasonic propagation. The acquisition time for a single image (STORM).\(^3\(^1\)\(^3\(^2\))\(^4\) However, the optical probe pulses detect only a single frequency as a result of Brillouin scattering, as described later. Spatial and temporal scanning allow 3D imaging (\( x \), \( y \), and \( t \)) over 40 \( \mu m \times 40 \mu m \) areas with \( \sim 1 \mu m \) and 1 ps resolutions. Acquisition times for each complete 3D data set are typically 7 h for 800 images with 1 \( \mu m \) /pixel, but this did not lead to any evident damage. (Time scans were executed for each \( x \) and \( y \).) Such movies\(^3\(^0\)) allow one to trace the ultrasonic propagation. The acquisition time for a single image (\( \sim 30 s \)) is of the same order as that in super-resolution microscopy techniques such as photoactivated localization microscopy (PALM) and stochastic optical reconstruction microscopy (STORM).\(^3\(^1\)\(^3\(^2\))\(^4\) For quantitative processing, we identify regions inside and outside the cells from static probe-beam reflectivity (\( R \)) images of the cells, as shown in Figs. 1(b) and 1(c). Figures 2(a) and 2(b) show \( \delta R(t) \) averaged over the cells and surrounding regions (after subtracting a thermal background signal), whereas (c) and (d) show typical images of \( \delta R \) for the two cells. The GHz oscillations in \( \delta R \) arise from interference between probe-beam light reflected from the Ti film and that scattered from the ultrasonic wavefront travelling perpendicular to the film surface in the solution or cell. This so-called Brillouin scattering results in oscillations for normal probe incidence at frequency \( f_B = 2n\varepsilon_0\lambda_p \), where \( n \) and \( \varepsilon_0 \) are the local refractive index and longitudinal sound velocity (\( \sim 1500 \text{ m/s} \)) and \( \lambda_p \) is the vacuum optical wavelength of the probe. Measuring \( f_B \) for known (or negligibly varying) \( n \) allows one to map \( \varepsilon_0 \) within the samples. Ultrasonic attenuation can be accessed from the decay in oscillation amplitude.

Rather than work directly with \( \delta R(t) \), it is advantageous to follow \( f_B(t) \) in order to find \( \varepsilon_0(t) \). Here, we implement continuous-wavelet transforms (WTs), previously used to analyse \( \delta R \) in picosecond ultrasonics to reveal acoustic echoes hidden by Brillouin oscillations,\(^2\(^5\))\(^6\) as well as on sub-picosecond timescales to reveal carrier-phonon interactions,\(^7\) for example. \( \delta R(t) \) for the regions inside and outside the cells are averaged and wavelet transformed using the Morlet mother wavelet

\[
\begin{align*}
M(t) &= \left( \frac{1}{\pi} \right)^{1/4} \cos(5t) \exp\left( -\frac{t^2}{2} \right),
\end{align*}
\]

where in this equation, \( t \) is normalized. We plot the wavelet-transform amplitude \( A_{WT} \) in Figs. 3(a) and 3(b) for the saline solution and endothelial cell and in Figs. 3(c) and 3(d) for the buffer solution and fat cell, respectively. The

FIG. 1. (a) Experimental setup and sample. PBS: polarization beam splitter. ND filter: neutral density filter. (b) and (c) Fluorescence micrographs of the endothelial and fat cells. In (b), superimposed on a phase-contrast image, actin filaments are stained by rhodamine phalloidin and nuclei by DAPI. In (c), superimposed on a differential-interference-contrast image, neutral lipids are stained by BODIPY 493/503 and nuclei by Hoechst 33342. (d) and (e): images of the normalized probe-beam reflectivity \( R \). The white dashed areas are used for analysis. The solid (red) lines are cell boundaries. 1 and 2 indicate the cells and their surrounding solutions.

of \( \sim 13 \text{ K} \) above room temperature (25 °C).\(^3\(^0\)) Two photodiodes (A and B in Fig. 1(a)) operating in differential mode are used to measure the probe reflectivity variations \( \delta R \sim 10^{-5} \). As the ultrasonic pulses, generated with strain amplitudes \( \sim 10^{-5} \) at up to \( \sim 100 \text{ GHz} \) in the Ti film,\(^3\(^5\)) traverse the cells and the solutions, the pulse duration broadens.\(^3\(^5\)) However, the optical probe pulses detect only a single frequency as a result of Brillouin scattering, as described later. Spatial and temporal scanning allow 3D imaging (\( x \), \( y \), and \( t \)) over 40 \( \mu m \times 40 \mu m \) areas with \( \sim 1 \mu m \) and 1 ps resolutions. Acquisition times for each complete 3D data set are typically 7 h for 800 images with 1 \( \mu m \) /pixel, but this did not lead to any evident damage. (Time scans were executed for each \( x \) and \( y \).) Such movies\(^3\(^0\)) allow one to trace the ultrasonic propagation. The acquisition time for a single image (\( \sim 30 s \)) is of the same order as that in super-resolution microscopy techniques such as photoactivated localization microscopy (PALM) and stochastic optical reconstruction microscopy (STORM).\(^3\(^1\)\(^3\(^2\))\(^4\)
corresponding \( f_B \) are 9.6 ± 0.1, 10.3 ± 0.1, 9.8 ± 0.1, and 9.9 ± 0.1 GHz yielding ultrasonic wavelengths ~0.15 \( \mu \)m.

Spatially interpolated images of normalized \( A_{WT} \) at three separate times are shown in Figs. 4(a)–4(c) for the endothelial cell and in (d)–(f) for the fat cell. Corresponding vertical positions \( z \) calculated from the estimated \( v_l \) are also shown. The spatial resolution for \( z \) is limited to the order of the ultrasonic wavelength ~0.15 \( \mu \)m.) \( A_{WT}(t) \) reveals the ultrasonic attenuation. That of the endothelial cell is clearly greater than that of its surroundings, whereas that of the fat cell is similar. The spatial dependence of \( A_{WT}(t) \) shows evidence of inhomogeneity inside the cells, although our signal-to-noise ratio is not sufficient to identify specific internal cell structures. \(^{38} \) \( A_{WT}(x, y, \text{and } t) \) is available in movie form. \(^{30} \)

Given the frequency variation \( f_B(t) \) and refractive index \( n \) one can obtain \( v_l(t) = f_B(t)\sqrt{\frac{\rho}{\rho_n}}/2 \). Here, we assume \( n = 1.38 \) for the endothelial cell and 1.36 for the fat cell. \(^{39,46} \) For the surrounding solutions, we use the refractive index of water, \( n = 1.33,41,42 \) as a good approximation. Figures 5(a) and 5(b) show \( v_l(t) \) derived in this way for the endothelial and fat cells and for their surrounding solutions. For the range of 200–600 ps, \(^{43} \) \( v_l(t) \) for the endothelial cell shows a slight variation that is probably the result of the finite wavelet-transform frequency resolution, \(^{44} \) whereas for the saline solution \( v_l(t) \) is, as expected, constant to within the experimental error. The fat cell and its surrounding buffer solution both exhibit a relatively constant \( v_l(t) \). The average value of \( v_l \) for the endothelial cell is 1.55 ± 0.02 \( \mu \)m ns\(^{-1} \), while that for the fat cell is 1.51 ± 0.01 \( \mu \)m ns\(^{-1} \). On the other hand, \( v_l \) for the saline solution is 1.50 ± 0.02 \( \mu \)m ns\(^{-1} \) and that for the buffer solution is 1.53 ± 0.01 \( \mu \)m ns\(^{-1} \). For endothelial cells, \( v_l \) was reported to be 1.56 (Ref. 45), and for fat cells it was reported to be 1.48 \( \mu \)m ns\(^{-1} \), \(^{46} \) in agreement with our results to within 1% and 2%.

One can also calculate the corresponding average bulk moduli \( B = \rho v_l^2 \), where \( \rho \) is the density. Assuming constant average values \( \rho = 1080 \) kg/m\(^3 \) for the endothelial cell \(^{45} \) and \( \rho = 920 \) kg/m\(^3 \) for the fat cell, \(^{45} \) the corresponding moduli are 2.60 ± 0.05 and 2.09 ± 0.02 GPa, while that for the saline solution is 2.23 ± 0.04 GPa and for the buffer solution 2.32 ± 0.02 GPa, obtained using the value \( \rho = 992.9 \) kg/m\(^3 \).
for water at \( \approx 40^\circ \text{C} \).48 Values of \( B \) for a bovine aortic endothelial cell and for a fat cell were measured to be 2.63 (Ref. 45) and 2.03 GPa,46 in agreement with our results to within 2\% and 3\%. As expected, \( B \) derived for the surrounding solutions are very close to that of water (\( B = 2.2 \text{ GPa} \)).49

To access ultrasonic attenuation, we plot \( A_{WT} \) vs time at \( f_s \) for each medium in Figs. 5(c) and 5(d). We fit to exponential decays43 (dotted black curves), yielding average inverse attenuation times \( \tau^{-1} = 0.0054 \) and 0.0017 ps\(^{-1} \) for the endothelial and fat cells. The surrounding saline and buffer solutions yield \( \tau^{-1} = 0.0024 \) and 0.0016 ps\(^{-1} \). (The error in \( \tau^{-1} \approx 10^{-5} \text{ ps}^{-1} \).) The good fits are the result of \( \tau^{-1} \) being essentially constant (verified by extracting \( dA_{WT}/dt \) as a function of \( t \), and noting that the value was approximately constant). The average spatial ultrasonic attenuation coefficients \( \alpha = (v_t \tau^{-1})^{-1} \), calculated using averaged values of \( v_t \), are 3.47 \pm 0.04 mm\(^{-1} \) for the endothelial cell and 1.56 \pm 0.04 mm\(^{-1} \) for its surrounding solution, whereas for the fat cell the corresponding values are 1.10 \pm 0.01 and 1.05 \pm 0.01 mm\(^{-1} \). These values are typical for cells and solutions in the GHz range.20,22 Lipid droplets in fat cells are filled with neutral lipids, and do not contain proteins and DNA. In contrast, the cytoplasm of endothelial cells is filled with proteins and organelles. This accounts for the differences in average physical properties between fat cells and endothelial cells.

In conclusion, using picosecond ultrasونics, we have demonstrated 3D in vitro biological-cell imaging and shown how continuous-wavelet transforms can be used to process the data sets obtained. This technique allows one to derive important physical values of cell samples, namely, sound velocities, bulk moduli, and ultrasonic attenuation coefficients. The averaged values for these properties derived from experiment show good agreement with those previously documented. In future, by the use of longer data acquisition times or use of acoustic contrast agents it should be possible to extend this technique to locally study the features inside cells through images of \( A_{WT}(x, y) \), \( v_t(x, y) \), and \( \alpha(x, y) \) at different times. In addition, lowering the laser power or using substrates of higher thermal conductivity would open the way to in vivo imaging. Such studies should be invaluable for investigating the mechanical properties of cell organelles in both vegetal and animal cells.

30 See supplementary material at http://dx.doi.org/10.1063/1.4918275 for movies of $\delta R$ and $A_{WT}$ for the endothelial and fat cells and the calculation for the temperature distribution.
38 The slight horizontal banding is thought to be an artifact of the signal acquisition and interpolation.
43 The range of validity of the WT data was verified by synthetic sinusoidal signals: 200–600 ps and 200–800 ps for the endothelial and fat cells, respectively.