Vibrational imaging of newly synthesized proteins in live cells by stimulated Raman scattering microscopy

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Synthesis of new proteins, a key step in the central dogma of molecular biology, has been a major biological process by which cells respond rapidly to environmental cues in both physiological and pathological conditions. However, the selective visualization of a newly synthesized proteome in living systems with subcellular resolution has proven to be rather challenging, despite the extensive efforts along the lines of fluorescence staining, autoradiography, and mass spectrometry. Herein, we report an imaging technique to visualize nascent proteins by harnessing the emerging stimulated Raman scattering (SRS) microscopy coupled with metabolic incorporation of deuterium-labeled amino acids. As a first demonstration, we imaged newly synthesized proteins in live mammalian cells with high spatial-temporal resolution without fixation or staining. Subcellular compartments with fast protein turnover in HeLa and HEK293T cells, and newly grown neurites in differentiating neuron-like N2A cells, are clearly identified via this imaging technique. Technically, incorporation of deuterium-labeled amino acids is minimally perturbing to live cells, whereas SRS imaging of exogenous carbon-deuterium bonds (C-D) in the cell-silent Raman region is highly sensitive, specific, and compatible with living systems. Moreover, coupled with label-free SRS imaging of the total proteome, our method can readily generate spatial maps of the quantitative ratio between new and total proteomes. Thus, this technique of nonlinear vibrational imaging of stable isotope incorporation will be a valuable tool to advance our understanding of the complex spatial and temporal dynamics of newly synthesized proteome in vivo.

stable isotope labeling | stimulated Raman microscopy | protein synthesis

The proteome of a cell is highly dynamic in nature and tightly regulated by both protein synthesis and degradation to actively maintain homeostasis. Many intricate biological processes, such as cell growth, differentiation, diseases, and response to environmental stimuli, require protein synthesis and translational control (1). In particular, long-lasting forms of synaptic plasticity, such as those underlying long-term memory, require new protein synthesis and translational control to maintain homeostasis. Many intricate biological processes, such as cell growth, differentiation, diseases, and response to environmental stimuli, require protein synthesis and translational control (1). In particular, long-lasting forms of synaptic plasticity, such as those underlying long-term memory, require new protein synthesis and translational control to maintain homeostasis.

Extensive efforts have been devoted to probing protein synthesis via fluorescence contrast. The inherent fluorescence of green fluorescent protein (GFP) and its genetic encodability allow one to follow a given protein of interest inside living cells with high spatial and temporal resolution (5, 6). However, GFP tagging through genetic manipulation works only on individual proteins but not at the whole-proteome level. To probe newly synthesized proteins at the proteome level, a powerful technique named bioorthogonal noncanonical amino acid acid tagging (BONCAT) was developed by metabolic incorporation of unnatural amino acids containing reactive chemical groups such as azide or alkyne (7–13). A related labeling method was recently demonstrated using an alkyn analog of puromycin (14). Newly synthesized proteins can then be visualized through subsequent conjugation of the reactive amino acids to fluorescent tags via click chemistry (15). Unfortunately, these fluorescence-based methods generally require nonphysiological fixation and subsequent dye staining and washing.

In addition to fluorescence tagging, radioisotope or stable isotope labeling is another powerful tool to trace and quantify proteome dynamics. Classical radioisotope-labeled amino acids (e.g., [14C]methionine) provide rigorous analysis of global protein synthesis. However, samples must be fixed and then exposed to film for autoradiography. For stable isotopes, the discovery of deuterium by Urey in 1932 immediately led to the pioneer work of Schoenheimer and Rittenberg studying intermediary metabolism (16, 17). To study proteome changes between different cells or under different conditions, stable isotopic labeling by amino acids in cell culture (SILAC) coupled with mass spectrometry (MS) has matured into a popular method for quantitative proteomics (18–21). However, SILAC-MS does not usually provide spatial information down to subcellular level and its invasive nature also limits its application for live-cell imaging. The same limitation applies to the recent ribosome profiling study using deep sequencing technique (22).

Therefore, it is highly challenging and desirable to be able to quantitatively image proteome synthesis in live cells with high spatial-temporal resolution. Herein, we report using stimulated Raman scattering (SRS) microscopy, an emerging vibrational imaging technique, for the visualization of nascent proteins in live cells coupled through metabolic incorporation of deuterium-labeled amino acids (Fig. 1). Newly synthesized proteins are imaged via their unique vibrational signature of carbon–deuterium bonds (C–D). Vibrational imaging by Raman contrast is a rapidly growing field. Spontaneous Raman microscopy can offer spatially resolved chemical information based on the vibration frequencies of characteristic chemical bonds. However, spontaneous Raman scattering is an intrinsically weak process, hence not ideal for fast live-cell imaging (23). As a nonlinear technique, coherent anti-Stokes Raman scattering (CARS) offers much higher imaging speed by virtue of coherent amplification (24–28). Unfortunately, CARS suffers from spectral distortion, unwanted nonresonant background, nonstirred forward concentration dependence, and coherent image artifact (25). Most recently, SRS microscopy has emerged to supersede CARS microscopy in almost all aspects (29–38). Using Einstein’s stimulated emission principle (39, 40), SRS has achieved unprecedented sensitivity down to ~1,000 retinoic acid molecules and up to video rate imaging speed in vivo (30, 33). Unlike CARS, SRS microscopy exhibits straightforward image interpretation and quantification without complications from the nonresonant background and phase-matching conditions (41, 42). Consequently, not only is the signal-to-noise ratio improved over CARS, but the Raman spectral fidelity is higher.

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greatly amplified by about $10^7$ times when the energy difference of the two laser beams matches the particular chemical bond vibration, $\Omega_{\nu B}$ (41). Accompanying such stimulated activation of one vibrational mode, one photon is created into the Stokes beam and simultaneously another photon is annihilated from the Pump beam, a process called stimulated Raman gain and stimulated Raman loss, respectively. Essentially, the energy difference between the Pump photon and the Stokes photon is used to excite the vibrational mode, fulfilling energy conservation. As shown in Fig. 1B, a high-frequency modulation scheme, where the intensity of the Stokes beam is turned on and off at 10 MHz, is used to achieve shot noise-limited detection sensitivity by suppressing laser intensity fluctuations occurring at low frequencies. The transmitted Pump beam after the sample is detected by a large-area photodiode, and the corresponding stimulated Raman loss signal, which also occurs at 10 MHz, is demodulated by a lock-in amplifier. By scanning across the sample with a laser-scanning microscope, a quantitative map with chemical contrast can be produced from the targeted vibrating chemical bonds. As the SRS signal is dependent on both Pump and Stokes laser beams, the nonlinear nature herein provides a 3D optical sectioning ability.

Here, we detect the vibrational signal of C–D as an indicator for newly synthesized proteins that metabolically incorporate deuterium-labeled amino acids (Fig. 1B). When hydrogen atoms are replaced by deuterium, the chemical and biological activities of biomolecules remain largely unmodified. Intriguingly, the C–D stretching motion displays a distinct vibrational frequency from all of the other vibrations of biological molecules inside live cells. It is known from classical mechanics that the frequency of vibrational oscillation, $\Omega_{\nu B}$, inversely scales with the square root of the reduced mass of the oscillator $\mu = (1/2m_1 + 1/2m_2)/m$, where $m$ is the spring constant of the corresponding chemical bond, and $\mu$ denotes the reduced mass of the oscillator. The reduced mass of the C–D oscillator is increased by two folds when hydrogen is replaced by deuterium. Based on the above equation, $\Omega_{\nu B}$ would be reduced by $\sqrt{2}$. Indeed, the experimental C–D stretching vibrational frequency is shifted from ~2,950 cm$^{-1}$ of C–H to ~2,100 cm$^{-1}$ of C–D. Remarkably, the vibrational frequency of 2,100 cm$^{-1}$ is located in a cell-silent spectral window in which no other Raman peaks exist (Fig. S1), thus enabling detection of exogenous C–D with both high specificity and sensitivity.

SRS Imaging of Newly Synthesized Proteins by Metabolic Incorporation of Leucine-d$_{10}$ in Live HeLa Cells. Among the 20 natural amino acids, leucine is an essential one with both high abundance in protein (~9%) in mammalian cells and a large number of side-chains C–H that can be replaced by C–D (43). Hence, we first demonstrated the feasibility of our technique by detecting the metabolic incorporation of leucine-d$_{10}$ (1-leucine-2,3,4,5,5',5',5'-d$_{10}$ as shown in Fig. 2A) to nascent proteins in live HeLa cells. Fig. 2B shows the spontaneous Raman spectrum of HeLa cells incubated in the medium containing 0.8 mM free leucine-d$_{0}$ for 20 h (blue) overlaid with the spectrum of HeLa cells growing in the regular medium without leucine-d$_{10}$ (red) as well as the spectrum from a 10 mM free leucine-d$_{0}$ solution in PBS (black). As indicated by the comparison between the blue and the red spectra, the Raman peaks of leucine-d$_{10}$, exhibiting multiple peaks around 2,100 cm$^{-1}$ due to symmetric and asymmetrical C–D stretching, are indeed located in the cell-silent region. The comparison of the blue and the black spectra implies that leucine-d$_{10}$ incorporated into cellular proteome after 20 h is enriched to about 10 mM. Thus, a 10% incorporation yield of leucine-d$_{10}$ can be estimated at this condition based on the intrinsic leucine concentration of about 100 mM in proteins (calculated from protein concentration and leucine percentage in cells).

Based on the above spectra, we choose to target the central 2,133 cm$^{-1}$ vibrational peak of C–D to acquire SRS images of nascent proteins in live HeLa cells. As expected, HeLa cells growing in regular medium show no detectable SRS contrast at 2,133 cm$^{-1}$ (Fig. 2C), which is consistent with the flat spectral baseline (red in Fig. 2B) in the cell-silent region. In contrast, SRS image of HeLa cells growing in the medium containing 0.8 mM

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**Fig. 1.** Stimulated Raman scattering (SRS) microscopy principle and experimental scheme. (A) Principle of SRS microscopy. When the energy difference between the Pump beam photon and the Stokes beam photon matches the vibrational frequency ($\Omega_{\nu B}$) of a specific chemical bond, a molecule is efficiently driven from the vibrational ground state to its vibrational exited state, passing through a virtual state. A quanta of such vibrational activation results in a photon in the Pump beam being annihilated (stimulated Raman loss) and a photon in the Stokes beam being created (stimulated Raman gain), which serves as the contrast for SRS microscopy. (B) Experimental scheme of imaging proteins with metabolic incorporation of deuterium-labeled amino acids. By feeding live cells with deuterium-labeled amino acids, newly synthesized proteins can be specifically labeled with carbon–deuterium bonds (C–D). By tuning the energy difference between Pump and Stokes beams to match the vibrational frequency of C–D, the distribution of C–D carrying new proteins can be imaged in live cells by SRS with high sensitivity and resolution without additional fixation or staining.

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**Results and Discussion**

**Physical Principle of Isotope-Based SRS Imaging.** SRS microscopy is a molecular-contrast, highly sensitive imaging technique with intrinsic 3D sectioning capability. It selectively images the distribution of molecules that carry a given type of chemical bonds through resonating with the specific vibrational frequency of the targeted bonds (30, 33, 41). As Fig. L4 illustrates, by focusing both temporally and spatially overlapped Pump and Stokes laser pulse trains into samples, the rate of vibrational transition is also preserved and a linear concentration dependence is strictly followed (30).

First, we demonstrated the proof-of-concept of our technique on live HeLa cells using a single deuterium-labeled essential amino acid, leucine-d$_{10}$, Then we optimized the incorporation efficiency of the deuterium isotope into nascent proteins and showed broad applicability of the method on several mammalian cell lines, particularly, its unique advantage in generating spatial maps of the quantitative ratio between new and old proteomes. Furthermore, besides visualizing newly synthesized proteins in cell bodies, the ability to image nascent proteins in neurites of neuron-like mouse neuroblastoma Neuro-2A (N2A) cells upon differentiation was demonstrated (42). By scanning across the sample with a laser-scanning microscope, a quantitative map with chemical contrast can be produced from the targeted vibrating chemical bonds. As the SRS signal is dependent on both Pump and Stokes laser beams, the nonlinear nature herein provides a 3D optical sectioning ability.
leucine-d10 (Fig. 2D) shows a weak but clearly identifiable contrast outlining the cell shape. As a control, the off-resonant SRS image at 2,000 cm\(^{-1}\) of the same cells is background free (Fig. 2E). Such clean chemical contrast among Fig. 2C–E would be difficult for CARS microscopy due to the presence of its nonresonant background. As a protein reference, an image taken at 2,940 cm\(^{-1}\) shows both existing and newly synthesized proteins (Fig. 2F), the signal of which comes from the same regions but is much stronger than that in Fig. 2D. Thus, we have demonstrated the feasibility of using SRS imaging to detect newly synthesized proteins by specifically targeting the C–D vibrational signal of metabolically incorporated leucine-d10 in live HeLa cells. This opens up an imaging opportunity to capture nascent proteome dynamics in live cells under a myriad of cues.

**Imaging Optimization by Metabolic Incorporation of a Deuterium-Labeled Set of All Amino Acids in Live HeLa Cells with Multicolor SRS Imaging.** Although leucine is the most abundant essential amino acid, it only accounts for a small fraction of amino acids in proteins. Hence, we reasoned that deuterium labeling of all of the amino acids would lead to a substantial signal enhancement. Indeed, the spontaneous Raman spectrum (Fig. 3A) of HeLa cells incubated with a deuterium-labeled set of all 20 amino acids (prepared by supplying a uniformly deuterium-labeled whole set of amino acids to leucine-, lysine-, and arginine-deficient DMEM; for more details, refer to Materials and Methods) exhibits C–D vibrational peaks about five times higher than the blue spectrum in Fig. 2B under the same condition. The corresponding SRS image at 2,133 cm\(^{-1}\) (Fig. 3B) shows a significantly more pronounced signal than that in Fig. 2D under the same intensity scale. In particular, nucleoli (indicated by arrows in Fig. 3B and verified by differential interference contrast visualization) exhibit the highest signal, which is in accordance with previous reports using BONCAT and our own fluorescence staining results (Fig. S2). Nucleoli, the active sites for ribosomal biogenesis, have been reported to involve rapid nucleolar assembly and proteomic exchange (44–46). Such fast protein turnover is indeed reflected by the spatial enrichment of newly synthesized protein signals in those subcellular areas (Fig. 3B). Note that SRS imaging here is directly performed on live cells and hence free from potential complications due to fixation and dye conjugation. Again, the off-resonant image at 2,000 cm\(^{-1}\) is clean and dark (Fig. 3C), proving the specificity of SRS imaging of C–D at 2,133 cm\(^{-1}\). In addition to imaging newly synthesized proteins, SRS can readily image intrinsic biomolecules in a label-free manner. By simply adjusting the energy difference between the Pump and the Stokes beams to match the vibrational frequency of amide I, lipids, and total proteins, respectively, Fig. 3D–F shows the SRS images of amide I band at 1,655 cm\(^{-1}\) primarily attributed to proteins; CH\(_2\) stretching at 2,845 cm\(^{-1}\) predominantly for lipids; and CH\(_3\) stretching at 2,940 cm\(^{-1}\) mainly from proteins with minor contribution from lipids.

**Time-Dependent de Novo Protein Synthesis and Protein Synthesis Inhibition.** Being linearly dependent on analyte concentration, SRS contrast is well suited for quantification of de novo protein synthesis in live cells. Here, we show time-dependent protein synthesis images under the same intensity scale (Fig. 4A–C). As expected, the new protein signal (2,133 cm\(^{-1}\)) from 5-, 12-, and 20-h incubation increases substantially over time (Fig. 4A–C), whereas the amide I (1,655 cm\(^{-1}\)) signal remains at a steady state (Fig. 4D–F). Because protein distribution is often heterogeneous in biological systems, we presented a more quantitative representation by acquiring ratio images between the newly synthesized proteins and the total proteome (from either amide I or CH\(_3\)). Fig. 4G–I depicts the fraction of newly synthesized proteins (2,133 cm\(^{-1}\)) from the total proteome (1,655 cm\(^{-1}\)) and its spatial distribution. The fraction of newly synthesized proteins is growing with time from 5 to 20 h, gradually highlighting nucleoli as the subcellular compartments with fast protein turnover (44–46). Such quantitative ratio imaging of new versus old proteins would be very difficult to obtain using BONCAT or mass spectroscopy without the destruction of cells. More time-dependent cell images are shown in Fig. S3. Moreover, Fig. 4J shows time-lapse SRS images of a live dividing HeLa cell after...
20-h incubation in deuterium-labeled, all-amino acids medium, clearly proving the viability of cells under the imaging condition.

The effect of protein synthesis inhibition by chemical drugs is further tested to validate that the detected C–D signal indeed derives from newly synthesized proteins. HEK293T cells incubated with a deuterium-labeled set of all amino acids together with 5 μM anisomycin, which works as a protein synthesis inhibitor by inhibiting peptidyl transferase or the 80S ribosome system, for 12 h show the absence of the C–D signal in the spontaneous Raman spectrum (Fig. 4K). Furthermore, SRS imaging of the same samples (Fig. 4L) exhibits drastically weaker signal [Fig. S4 provides a more thorough analysis of the residual signal, which is possibly attributed to the intracellular free amino acid pool (47)] compared with Fig. 4B without the protein synthesis inhibitor. As a control, the corresponding 2,940 cm−1 image (Fig. 4M) of total proteome remains at a similar level as the non-drug-treated counterpart in Fig. 3F. Thus, the detected C–D SRS signal (Fig. 4 A–C) originates from deuterium-labeled nascent proteins, which vanishes upon adding the protein synthesis inhibitor.

Demonstration on HEK293T Cells and Neuron-Like Differentiable Neuroblastoma N2A Cells. To show the general applicability and potential of our method, we choose two additional mammalian cell lines for further demonstration: human embryonic kidney HEK293T cells, and neuron-like neuroblastoma N2A cells, which can be induced to differentiate with the growth of neurites (i.e., axons and dendrites). The spontaneous Raman spectrum (Fig. 5A) of HEK293T cells incubated with a deuterium-labeled set of all amino acids for 12 h exhibits a 2,133 cm−1 C–D channel signal nearly as high as the 1,655 cm−1 amide channel signal. The resulting SRS image shows a bright signal for new proteins with an intense pattern residing in nucleoli (Fig. 5B). As before, the off-resonant image (2,000 cm−1) displays vanishing background (Fig. 5C); the amide I channel (1,655 cm−1) image (Fig. 5D) exhibits consistent overall protein contribution similar to that in HeLa cells; CH2 (2,845 cm−1) image (Fig. 5E) depicts a more diffuse lipid distribution in cytoplasm compared with that in HeLa cells. Consistent with the results obtained in HeLa cells above, the ratio image (Fig. 5F) between the newly synthesized proteins (Fig. 5B) and the total proteins (Fig. 5D) highlights nucleoli for active protein turnover in HEK293T cells as well (44–46).

In addition to showing the ability to image newly synthesized proteins inside cell body, our technique can also be applied to tackle more complex problems, such as de novo protein synthesis in neuronal systems. As a model system to study neuronal differentiation, axonal growth, and signaling pathways. Under differentiation condition, N2A cells massively grow new neurites from cell bodies and form connections with other cells. Fig. 6A shows the image of newly synthesized proteins after induction for differentiation, by simultaneously differentiating the N2A cells and supplying with the deuterium-labeled set of all amino acids for 24 h. Similar to HeLa and HEK293T cells, N2A cell bodies are observed to display high-level protein synthesis. More interestingly, newly synthesized proteins are also observed in a subset of, but not all neurites (Fig. 6A and B), which implies that the observed neurites in Fig. 6A are newly grown under the differentiation condition. For a detailed visualization, Fig. 6C and D shows the zoomed-in regions in the dashed squares in Fig. 6A and B, respectively. A more comprehensive examination is illustrated by both the ratio image (Fig. 6E) between Fig. 6C and D and the merged image (Fig. 6F) with the red channel designating new protein signal from Fig. 6C and the green channel designating total protein signal from Fig. 6D. On one hand, both the ratio image and the merged image highlight the neurites with higher percentage of new proteins (indicated by stars), implying these neurites are newly grown. On the other hand, from the merged image, there are some neurites (indicated by arrows) showing obvious signals in the green channel (total proteins) only but with no detectable signal in the red channel (new proteins). Hence, the arrow indicated neurites are most likely older than their starred counterparts. In addition, the transition from green to red in the merged image (Fig. 6F) implies the growth direction by which new neurites form and grow. A second set of N2A images showing similar patterns as in Fig. 6 is also examined in Fig. S5. A more relevant system to study de novo protein synthesis and neuronal activities would be hippocampal neurons, which are known to be involved in long-term memory formation (2–4). SRS image (2,133 cm−1) of hippocampal neuron cells incubated with a deuterium-labeled set of all amino acids shows a newly synthesized protein pattern in the neurites (Fig. S6). The intricate relationship between protein synthesis and neuronal activities is currently under investigation.

Fig. 4. SRS imaging of time-dependent de novo protein synthesis and drug-induced protein synthesis inhibition effect in live HeLa cells incubated in a deuterium-labeled all-amino acid medium. (A–F) SRS image targeting the central 2,133 cm−1 vibrational peak of C–D displays a time-dependent signal increase [5 h (A), 12 h (B), 20 h (C)] of the newly synthesized proteins, with nucleoli being gradually highlighted. As a control, the amide I (1,655 cm−1) signal remains at a steady state over time [5 h (D), 12 h (E), 20 h (F)]. (G–I) Ratio images between the SRS image at 2,133 cm−1 (newly synthesized proteins) and the SRS image at 1,655 cm−1 (the amide I band from total proteins), representing the relative new protein fraction with subcellular resolution at each time point [5 h (G), 12 h (H), 20 h (I)]. The color bar ranging from black to red represents the ratio ranging from low to high. (J) Time-lapse SRS images of a live dividing HeLa cell during a 25-min time course after 20-h incubation with deuterated all-amino acids medium. (K) Spontaneous Raman spectrum of HeLa cells incubated with both deuterium-labeled all-amino acids and a protein synthesis inhibitor anisomycin (5 μM) for 12 h shows the drastic attenuation of the C–D Raman peak at 2,133 cm−1. (L) SRS image of the same sample displays near vanishing signal throughout the whole field of view. (M) As a control, the image of the same cells at 2,940 cm−1 confirms that anisomycin does not influence the total protein level.
mammalian cells under physiological conditions without any fixation or staining. From the perspective of biological applications, the bio-compatibility of both deuterium labeling and SRS imaging renders this technique the prospect of revealing spatial-temporal proteome dynamics in more complex systems such as live animals. From the perspective of imaging technology, nonlinear vibrational microscopy is well suited for visualizing the metabolic incorporation of isotope labeled precursors of macromolecules for its high sensitivity, specificity, and the non-invasive nature. We expect this strategy to be generalized and expanded to other stable isotopes such as $^{13}$C and $^{15}$N.

**Materials and Methods**

**SRS Microscopy.** The experimental setup is shown in Fig. 1B. Spatially and temporally overlapped pulsed Pump (tunable from 720 to 990 nm, 7 ps, 80-MHz repetition rate) and Stokes (1,064 nm, 5–6 ps, 80-MHz repetition rate, modulated at 10 MHz) beams, which are provided by picoEMERALD from Applied Physics & Electronics are coupled into an inverted laser-scanning microscope (FV1000 MPE, Olympus) optimized for near-IR throughput. A 60x water objective (UPlanAPO/IR; 1.2 N.A.; Olympus) is used for all cell imaging. After passing through the sample, the forward going Pump and Stokes beams are collected in transmission by a high N.A. condenser and imaged onto a large area Si photodiode. A high OD bandpass filter (890/220, Chroma) is used to block the Stokes beam completely and to transmit the Pump beam only for the detection of the stimulated Raman loss signal. The output current from the photodiode is terminated, filtered, and demodulated by a lock-in amplifier (SR844; Stanford Research Systems) at 10 MHz to ensure shot noise–limited detection sensitivity. For imaging, 512 × 512 pixels are acquired for one frame (26 s per frame) with a 100-μs pixel dwell time and 20-μs time constant from the lock-in amplifier. Powers after 60x IR objective used for imaging are as follows: 61 mW for modulated Stokes beam; 145 mW for the Pump beam of 2,133 cm$^{-1}$, 2,000 cm$^{-1}$.

**Conclusion**

The ability to visualize newly synthesized proteomes in biological systems will greatly advance our understanding of complex cellular functions occurring in space and time (1–4). Currently, this endeavor is mainly pursued by several distinct contrast mechanisms including fluorescence staining, autoradiography, and mass spectroscopy. Here, we report a new technique of SRS microscopy coupled with stable isotope labeling (deuterium labeling in this study) to address this challenge. The major advantages of our technique lie in the following aspects. First, our approach is essentially noninvasive and completely compatible with the live-cell physiology. This is in contrast with earlier methods of autoradiography and BONCAT. In terms of sample preparations, the deuterium isotope has a high degree of similarity with the cells’ endogenous counterpart (18–21). In terms of imaging conditions, SRS directly probes vibrational transitions in a stain-free manner using near-infrared lasers whose phototoxicity is low especially when using picosecond pulses. We note that a recent technique called multiisotope imaging mass spectrometry has also demonstrated a high-resolution isotope imaging ability (48, 49), but with a highly destructive nature due to the use of an ion microscope. Second, overcoming the major problems of CARS microscopy, SRS is an emerging nonlinear Raman microscopy with purely chemical contrast and high sensitivity, enabling fast imaging speed up to video rate in live animals and humans (41, 42). Our current electronics offers imaging speed of ~26 s per frame (512 × 512 pixels), which could be accelerated to video rate using a custom lock-in amplifier (33). As a comparison, spontaneous Raman microscopy relies on a feeble signal, which is easily overwhelmed by cell autofluorescence and needs long integration time (>hours) for imaging (23), and is thus undesirable for live-cell imaging. In fact, spontaneous Raman microscopy has been applied for detection of newly synthesized proteins, but was only possible with fixed cells (50). Third, SRS microscopy can readily offer the intrinsic total proteins distribution in a label-free manner. Such a valuable internal reference of total proteins is very helpful for obtain techniques such as BONCAT or mass spectroscopy without destruction of the cells.

Therefore, we have demonstrated SRS microscopy coupled with deuterium-labeled amino acids incorporation as an imaging technique for visualization of newly synthesized proteins in living mammalian cells.
and 1,655 cm$^{-1}$ channels, and 64 mW for Pump beam of 2,950 cm$^{-1}$ and 2,845 cm$^{-1}$ channels.

**Metabolical Labeling of the Newly Synthesized Proteins by Deuterium-Labeled Amino Acids.** Deuterium-labeled leucine-d$_{10}$ medium is made by adding leucine-d$_{10}$ (0.8 mM), lysine (0.8 mM), and arginine (0.4 mM) (Sigma) into leucine-, lysine-, and arginine-deficient DMEM (Sigma). Deuterium-labeled all-amino acids medium is made by adding uniformly deuterium-labeled amino acid mix (20 aa) (Cambridge isotope) into leucine-, lysine-, and arginine-deficient DMEM (Sigma). The final concentration of leucine-d$_{10}$ is adjusted to be 0.8 mM among the amino acid mix. (Because the starting medium is leucine-, lysine, and arginine deficient, by adding the deuterium-labeled 20-aa mix, we essentially deuteriate all of the leucine, lysine, and arginine as well as about one-half of the other amino acids.) Cells are seeded on a coverslip in a petri dish with 2 mL of regular DMEM with 10% (vol/vol) FBS and 1% penicillin/streptomycin (Invitrogen) for 20 h. The regular medium is then replaced with medium containing either leucine-d$_{10}$ or a deuterium-labeled set of all amino acids. After incubation for a certain amount of time, the coverslip is taken out to make an imaging chamber filled with PBS for SRS imaging. For N2A cells, in the process of induced cell differentiation with serum deprivation and 1 μM retinoic acid, the deuterium-labeled set of all amino acids is supplemented.

**Spontaneous Raman Spectroscopy.** The spontaneous Raman spectra were acquired using a laser Raman spectrometer (inVia Raman microscope; Renishaw) at room temperature. A 27-mW (after objective), 532-nm diode laser was used to excite the sample through a 50×, N.A. 0.75 objective (NPLAN EPI; Leica). The total data acquisition was performed during 80 s using the WIRE software.

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Supporting Information

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SI Materials and Methods

**Stimulated Raman Scattering Microscopy.** An integrated laser (picoEMERALD; Applied Physics & Electronics) was used as the light source for both Pump and Stokes beams. Briefly, picoEMERALD provides an output pulse train at 1,064 nm with 7-ps pulse width and 80-MHz repetition rate, which serves as the Stokes beam. The frequency-doubled beam at 532 nm is used to synchronously seed a picosecond optical parametric oscillator to produce a mode-locked pulse train [the idler beam of the optical parametric oscillator (OPO) is blocked with an interferometric filter] with 5- to ≈6-ps pulse width. The wavelength of the OPO is tunable from 720 to 990 nm, which serves as the Pump beam. The intensity of the 1,064 nm Stokes beam is modulated by a built-in acousto-optic modulator at 10 MHz driven by a square-wave function generator with a modulation depth of more than 70%. The Pump beam is spatially overlapped with the Stokes beam with a dichroic mirror inside picoEMERALD. The temporal overlap between Pump and Stokes pulse trains is ensured with a built-in delay stage and optimized by the stimulated Raman scattering (SRS) signal of pure dodecane liquid.

Pump and Stokes beams are coupled into an inverted laser-scanning microscope (FV1000MPE; Olympus) optimized for near-IR throughput. A 60× water objective (UPlanAPO/IR; 1.2 N.A.; Olympus) with high near-IR transmission is used for all cell imaging. The Pump/Stokes beam size is matched to fill the backaperture of the objective. The forward-going Pump and Stokes beams after passing through the sample are collected in transmission with a high N.A. condenser lens (oil immersion, 1.4 N.A.; Olympus), which is aligned following Kohler illumination. A telescope is then used to image the scanning mirrors onto a large area (10 × 10 mm) Si photodiode (FDS1010; Thorlabs) to descan beam motion during laser scanning. The photodiode is reversed bias by 64 V from a DC power supply to increase both the saturation threshold and response bandwidth. A high OD bandpass filter (890/220 CARS; Chroma Technology) is used to blocked the Stokes beam completely and transmit the Pump beam only. The output current of the photodiode is electronically prefILTERed by a bandpass filter (BBP-10.7; Mini Circuits) to suppress both the 80-MHz laser pulsing and the low-frequency contribution due to laser scanning cross the scattering sample. It is then fed into a radio frequency lock-in amplifier (SR844; Stanford Research Systems) terminated with 50 Ω to demodulate the stimulated Raman loss signal experienced by the Pump beam. The R-output of the lock-in amplifier is fed back into the analog interface box (FV10-ANALOG) of the microscope. The time constant is set for 20 μs (the shortest available with no additional filter applied). The current SRS imaging speed is limited by the shortest time constant available from the lock-in amplifier (SR844). For imaging, 512 × 512 pixels are acquired for one frame with a 100 μs of pixel dwell time and 20 μs of time constant from the lock-in amplifier. Laser powers after 60x IR objective used for cell imaging are as follows: 61 mW for modulated Stokes beam; 145 mW for the Pump beam of 2,133 cm⁻¹, 2,000 cm⁻¹, and 1,655 cm⁻¹ channels; and 64 mW for Pump beam of 2,950 cm⁻¹ and 2,845 cm⁻¹ channels.

**Sample Materials for SRS Microscopy.** Leucine-, lysine-, arginine-deficient DMEM (catalog no. D9443), t-lysine (catalog no. L8662), t-arginine (catalog no. A8094), t-leucine-2,3,4,5,5,5,5,5-d₁₀ (catalog no. 492949), and retinoic aid (catalog no. R2625) were obtained from Sigma. Uniformly deuterium-labeled cell free amino acid mix (20 aa) (DLM-6819) was from Cambridge Isotope Laboratories. Homopropargylglycine (Hpg) and Click-iT Hpg Alexa Fluor 488 Imaging Kit (catalog no. C10428) were purchased from Invitrogen.

Deuterium-labeled DMEM (containing either leucine-d₁₀ or deuterium-labeled all amino acids) was prepared by adding the appropriate amounts (same final concentration as in the regular DMEM) of leucine-d₁₀ together with non-deuterium-labeled regular lysine and arginine stock solutions; or of deuterium-labeled cell free amino acid mix solution, into leucine-, lysine-, arginine-deficient DMEM supplied with 10% (vol/vol) FCS, 100 U/mL penicillin, and 50 μg/mL streptomycin. The deuterium-labeled Neurobasal medium for culturing hippocampal neurons is made by supplying all of the essential nutrients with deuterium-labeled amino acids solution for neurons according to Neurobasal medium recipe (Invitrogen), which is then added with B-27 Serum-Free Supplement.

**Fluorescence Microscopy Using Bioorthogonal Noncanonical Amino Acid-Tagging Approach.** Cells were incubated with 1 mM Hpg, which is an alkyne-bearing analog of methionine, for 20 h. Then fluorescence labeling was conducted using a Click-iT Imaging Kit according to the manufacturer’s procedure. Briefly, cells were fixed with 3.7% formaldehyde for 15 min. Then, cells were washed twice with 3% BSA in PBS, followed by incubation with 0.5% Triton X-100 in PBS for 20 min. After washing, cells were incubated with Click-iT reaction mixture for 30 min. Then, after washing with 3% BSA in PBS four times, fluorescence images were obtained using a Leica TCS SP5 confocal microscope while the cells were immersed in PBS solution.
Fig. S1. Spontaneous Raman spectrum from 3,000 cm$^{-1}$ to 1,200 cm$^{-1}$ of HeLa cells growing in a regular medium clearly displays a cell-silent spectral region as highlighted by the blue dashed box. The 2,940 cm$^{-1}$ peak shows the signal of CH$_3$ stretching mainly from cellular proteins. In addition, the 1,655 cm$^{-1}$ peak shows the amide I stretching signal also primarily from cellular proteins.

Fig. S2. Fluorescence image of newly synthesized proteins in HeLa and HEK 293T cells using bioorthogonal noncanonical amino acid tagging (BONCAT). By metabolic incorporation of homopropargylglycine (Hpg) followed with fluorescence staining after fixation, permeabilization, and click chemistry using Click-iT Hpg Alexa Fluor 488 Imaging Kit, the newly synthesized proteins are shown in green. The fluorescence images of HeLa cells (A) and HEK293T cells (B) show the maps of newly synthesized proteins in the whole-cell level with nucleoli being highlighted.
Fig. S3. Multicolor time-dependent SRS imaging of live HeLa cells incubated in deuterium-labeled all-amino acid medium. (A–C) 2,133 cm\(^{-1}\) (C–D channel) images of new proteins for cells incubated for 5, 12, and 20 h displaying increasing signals over time. (D–F) The corresponding 1,655 cm\(^{-1}\) (amide I channel) images of primarily total proteins show the signals at a steady state. (G–I) 2,845 cm\(^{-1}\) (CH\(_2\) channel) images of primarily lipids. (J–L) 2,940 cm\(^{-1}\) (CH\(_3\) channel) images of mainly total proteins with minor contribution from lipids. (M–O) Ratio maps between the new proteins (2,133 cm\(^{-1}\)) and total proteins (1,655 cm\(^{-1}\)) over time, gradually highlighting the nucleoli.
A shows the SRS image targeting the $2,133$ cm$^{-1}$ of C–D vibration peak for newly synthesized proteins during 5 h of incubation with a deuterium-labeled set of all amino acids. Note that the color scale of this image adapts the same intensity scale as shown in Fig. 4 A–C to illustrate the time-dependent signal growth. In contrast, B displays the “vanishing” signal when $5 \mu$M anisomycin was coincubated in the medium to block protein synthesis. When the image color scale of both A and B are amplified by five times, C and D are the resulting amplified images. C shows the same cells but with five times higher brightness (even with partial saturation) than A. D now starts to exhibit some faint but identifiable image contrast, which is possibly due to the intracellular free amino acids pool (1).

Fig. S5. SRS imaging of newly synthesized proteins in both cell bodies and newly grown neurites of neuron-like differentiable mouse neuroblastoma (N2A) cells. During the cell differentiation process by serum deprivation and 1 μM retinoic acid, a deuterium-labeled set of all amino acids is also supplied for 24 h. (A) SRS images targeting the central 2,133 cm⁻¹ vibrational peak of C–D bond show the newly synthesized proteins. (B) SRS images targeting the 2,940 cm⁻¹ \( \text{CH}_3 \) show the total proteins. (C and D) Zoomed-in images as indicated in the white dashed squares in A and B. (E) Ratio image between new proteins (C) and total proteins (D). Although the starred neurites show relatively high percentage of new protein, the arrows indicated neurites displaying low new proteins percentage. (F) Merged image between new protein (C) (red channel) and total proteins (D) (green channel). Similarly, the starred regions show obvious new proteins, whereas the arrows indicate regions that have low-level new protein signal.

Fig. S6. SRS imaging of newly synthesized proteins in both cell bodies and neurites of live hippocampal neurons growing in Neurobasal medium supplied with deuterium-labeled amino acids for 24 h. (A) SRS image targeting 2,133 cm⁻¹ vibrational peak of C–D bond shows the newly synthesized proteins in both cell bodies and part of the neurites. SRS images at 2,940 cm⁻¹ (B) and 1,655 cm⁻¹ (C) are both attributed mainly to total proteins.