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COMMUNICATION

Direct organelle thermometry with fluorescence lifetime imaging microscopy in single myotubes †

Hideki Itoh,ab Satoshi Arai,cd Thankiah Sudhaharan, b Sung-Chan Lee, e Young-Tae Chang,fg Shin’ichi Ishiwata,c,h Madoka Suzuki,cd† E. Birgitte Laneb†

We describe organelle thermometry using an endoplasmic reticulum-targeting small molecule dye and cytosolic mCherry, whose fluorescence lifetimes reduce with increasing temperature and can be monitored by fluorescence lifetime imaging microscopy. The results show that heat production in single myotubes is highly localized and is coupled to a Ca2+ burst.

Optical sensors that allow the read-out of temperature with fluorescence signals have begun to emerge recently.1 We have developed microscopic thermometry methods with glass capillaries,2 nano-particles,3-5 and sheet-coated cover glass.6 For intracellular thermometry, the precise subcellular localization of these thermosensors is all-important. Because the measured temperature is a function of the distance from the heat source, the location of the probe critically affects the accuracy of the temperature measurements. To the best of our knowledge, there have been only two sensors reported that meet the spatial requirements for targeting endoplasmic reticulum (ER) or sarcoplasmic reticulum (SR), which produces heat in skeletal muscle during non-shivering thermogenesis.1-7

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One is the use of genetically-encoded protein-based ratiometric fluorescent thermosensors which can be expressed in the organelle of interest.8 Another is a small molecule-based fluorescent thermosensor that specifically localizes to ER or SR: this is easier to use by virtue of its small molecular size, leading to high cell permeability, and its photostability. In this study, fluorescence lifetime imaging microscopy (FLIM) was used because of its advantages over intensity-based methods, in that the fluorescence lifetime of the dye is independent of sample displacement, dye concentration and/or light source intensity.9 Currently two types of FLIM are available. Time-domain FLIM is the most frequently used, but requires several minutes to acquire one image with relatively higher spatial resolution. Frequency-domain FLIM takes about 10 sec to create a single image but with lower spatial resolution. By using both types of FLIM, we demonstrated subcellular-level quantitative imaging of Ca2+-induced heat production in myotubes, through the use of the ER-targeted fluorescent thermometer, ER thermo yellow (Fig. 1a), as described by us previously10, and cytosolic mCherry.

As the fluorescence intensity of ER thermo yellow is temperature sensitive, we first confirmed that ER thermo yellow would also work as a temperature sensor with FLIM in HeLa cells. The fluorescence lifetime of ER thermo yellow measured by time-domain FLIM shortened with temperature increase (Fig. S1, ESI†). The intensity-weighted average fluorescence lifetime (τint) of ER thermo yellow was 2.50 ns (3.31 ns, 35%; 1.64 ns, 65%) at 25 °C and 2.28 ns (3.19 ns, 29%; 1.49 ns, 71%) at 37 °C. The relationship between temperature and the average fluorescence lifetime of ER thermo yellow measured by frequency-domain FLIM (Fig. S2, ESI†) was linear (R2 = 0.97) in living HeLa cells between 23 °C and 40 °C (Table S1, Fig. S3a, ESI†). The temperature sensitivity was ~26 ps/°C and the accuracy of temperature measurement was 1.6 ± 0.4 °C (average ± standard deviation (SD)) in these experimental conditions. In ideal situations, the absolute temperature could be determined by means of FLIM. However, in our setup with frequency-domain FLIM, it was observed that the accuracy of absolute temperature measurements in this study was almost the same as the magnitude of temperature changes in response to chemical stimulations, as mentioned below. It was therefore decided to limit the analysis to the temperature difference in this study. Similar thermosensitivity with better...
accuracy was recorded in parallel experiments using fixed HeLa cells (Table S1, Fig. S3b, ESI †). Effects of the chemical environment on the fluorescence lifetime of ER thermo yellow were also measured by time-domain FLIM. In a physiological pH range (5.0–8.0), the fluorescence lifetime of ER thermo yellow fluctuated, but it was relatively stable at the normal pH of ER (pH 7.4 ± 0.2)11; τint = 2.18 ± 0.07 ns at pH 7 and τint = 2.16 ± 0.08 ns at pH 8 (Fig. S4a, ESI †). The emission spectra show a similar trend (Fig S4b, ESI †). τint was independent of ionic strength (0–500 mM KCl) (Fig. S4c, ESI †). We and others have measured a temperature increase of about 1 °C that is induced by Ca2+ shock with ionomycin, a Ca2+ ionophore5,9,10,12. Hence we validated the thermometry with the fluorescence lifetime of ER thermo yellow in living HeLa cells by means of ionomycin stimulation. The temperature increase after 2 μM ionomycin addition was 0.93 ± 0.2 °C (vehicle: 0.03 ± 0.42 °C), which was consistent with previous studies (Fig. S5, ESI †).

This thermometry can be used on C2C12 myotubes (Fig. 1b, Table S1, Fig. S6, ESI †). The NST of skeletal muscle is mediated by sarco/endoplasmic reticulum Ca2+-ATPase (SERCA), which is a Ca2+ pump located in the SR membrane that actively transfers Ca2+ from the cytosol to the SR lumen by means of ATP hydrolysis. The Ca2+ release through the ryanodine receptor (RyR), which is a Ca2+ channel located at SR7,8, triggers the NST. Using frequency-domain FLIM, we evaluated the efficacy of ER thermo yellow in monitoring the heat production in C2C12 myotubes. First, we stimulated RyR with 1 mM caffeine, which increases the open probability of RyR, so that Ca2+ is released from the SR lumen13,14. The myotubes were loaded with Ca2+ indicator (fluo-4) to visualize intracellular Ca2+ concentration. Immediately after 1 mM caffeine addition, a Ca2+ burst was observed, followed by a return to basal Ca2+ concentration at about 120 s (Fig. 2a, Fig. S7, ESI †). Thapsigargin, an inhibitor of SERCA15, also induced a Ca2+ burst when applied at 1 μM due to Ca2+ leakage through RyR and store-operated calcium entry16, which peaked at 60 s, and the Ca2+ signal remained high for 150 s (Fig. 2b, Fig. S7, ESI †). The fluorescence lifetime of ER thermo yellow shortened after 1 mM caffeine addition, and returned within 10 min (Fig. 2c, Fig. S8, ESI †), but no obvious change was observed upon 1 μM thapsigargin addition (Fig. 2d). As mentioned above (Fig. S4a, Fig. S4b, ESI †), the fluorescence lifetime of ER thermo yellow was changed between pH 6 and pH 7, but was constant below pH 6 and over pH 7. We observed the shortening of the fluorescence lifetime of ER thermo yellow by 1 mM caffeine stimulation, however, this cannot be explained by the decrease of pH from pH 7.4 ± 0.211. Myotubes were displaced due to the Ca2+ bursts (Fig. S9, ESI †). In this situation, single-colour intensity-based methods are problematic. In contrast, Fig. 2d shows the fluorescence lifetime of ER thermo yellow is independent of the displacement of myotubes, which is one of the advantages of FLIM. Upon 1 mM caffeine stimulation, the temperature increased after a Ca2+ burst (Fig. S10, ESI †), which was consistent with the gap between the Ca2+ burst and temperature increase of HeLa cells in response to ionomycin in our previous study7. The average trend showed a peak at 105 s (Fig. 2c), and the temperature changes at 105 s before (ΔTbefore) and after (ΔTafter) 1 mM caffeine addition were 0.07 ± 0.25 °C and 1.6 ± 0.6 °C (p = 1.7 × 10−3) respectively, whereas ΔTbefore and ΔTafter for the vehicle (DMEM) only addition were −0.14 ± 0.26 °C and −0.23 ± 0.28 °C (p = 0.36) respectively (Fig. 2e). Upon 1 μM thapsigargin stimulation, ΔTbefore (at −45 s) and ΔTafter (at 45 s) were very similar, 0.10 ± 0.28 °C and 0.38 ± 0.34 °C (p = 1.9 × 10−3) respectively; 0.01 ± 0.22 °C and −0.15 ± 0.28 °C (p = 7.5 × 10−3) respectively in the control (DMEM) measurements (Fig. 2f). The control time course data are shown in Fig. S11, ESI †. In addition, the changes of the fluorescence lifetime of ER thermo yellow were suppressed upon 1 mM caffeine stimulation in the presence of 1 μM thapsigargin (Fig. S12, ESI †), which is consistent with other results in this study.

Fig. 1 Properties of fluorescence lifetime of ER thermo yellow. (a) Chemical structure of ER thermo yellow. (b) Fluorescence lifetime images of living C2C12 myotubes loaded with ER thermo yellow at 25 °C and 37 °C. Scale bar, 50 μm; calibration bar, 2.3—3.3 ns.

Fig. 2 A key player of heat production, SERCA, in C2C12 myotubes. (a, b) Time course of the fluorescence intensity of fluo-4 in C2C12 myotubes stimulated by 1 mM caffeine or 1 μM thapsigargin. The thin lines, individual myotubes; the thick lines, the means (a, N = 4, n = 36; b, N = 4, n = 20). (c, d) Time course of the fluorescence lifetime changes of ER thermo yellow in C2C12 myotubes stimulated by 1 mM caffeine or 1 μM thapsigargin. The thin lines, individual myotubes; the thick lines, the mean (c, N = 3, n = 16; d, N = 3, n = 16).
Comparison of the ΔT of SR in C2C12 myotubes between 1 mM caffeine (red circles) and DMEM (blue circles) stimulations, or between 1 μM thapsigargin (red circles) and DMSO (blue circles) stimulations. The red and blue diamonds, the mean values in each condition, respectively.

Expression levels of RyR and SERCA increase with the differentiation process of C2C12 cells. Consistently, 1 mM caffeine induced no detectable Ca²⁺ burst in C2C12 myoblasts (Fig. 3a, Fig. S13, ESI†), nor the fluorescence lifetime change of ER thermo yellow (Fig. 3b). These results also confirm that caffeine itself has no significant effect on the fluorescence lifetime of ER thermo yellow. The temperature sensitivity of ER thermo yellow in myoblasts was ~25 ps/°C (Table S1, Fig. S14a, ESI†). The ΔTbefore (at −105 s) and ΔTafter (at 105 s) were 0.10 ± 0.28 °C and 0.09 ± 0.36 °C (p = 0.86) respectively (Fig. 3c). The control measurements showed essentially the same trend (Fig. S14b, ESI†); ΔTbefore (at −105 s) and ΔTafter (at 105 s) were 0.05 ± 0.35 °C and −0.10 ± 0.40 °C (p = 1.5 × 10⁻²⁴) respectively (Fig. 3c). We therefore concluded that there was no substantial temperature increase of ER in single C2C12 myoblasts upon 1 mM caffeine stimulation, that could be detected using these methods.

![Fig. 3 No substantial events in C2C12 myoblasts (undifferentiated cells) stimulated by 1 mM caffeine.](image)

We observed Ca²⁺-induced temperature increase (~1.6 °C) at the SR, a target organelle that we expected to be a major heat source, in C2C12 myotubes. The heat should diffuse out from the SR to the surrounding environment within the cell. Thus, the temperature around the SR should vary inversely with distance from the SR. To examine the temperature rise in the cytosol, we expressed mCherry (fluorescent protein) in the cytosol of C2C12 myotubes. Since the fluorescence quantum yield decreases with temperature increase, we expected that the fluorescence lifetime of mCherry would also be temperature dependent. In fact, the τint of cytosolic mCherry was 1.42 ns (1.85 ns, 35%; 1.01 ns, 65%) at 25 °C and 1.30 ns (1.77 ns, 29%; 0.93 ns, 71%) at 37 °C (Fig. 4a). These τint values were consistent with a literature value (~1.46 ns at 20 °C).²⁹

The relationship between temperature and the fluorescence lifetime of cytosolic mCherry was linear (R² = 0.98) between 24°C and 40 °C, and the temperature sensitivity was ~14 ps/°C (Fig. S15, ESI†). We then measured temperature changes, by means of the fluorescence lifetime of mCherry, in the cytosol of C2C12 myotubes exposed to 1 mM caffeine (Fig. S16a, ESI†). The temperature difference at 105 s before (ΔTbefore) and 105 s after (ΔTafter) the addition of 1 mM caffeine were −0.03 ± 0.12 °C and −0.07 ± 0.18 °C (p = 0.74) respectively (Fig. 4b). In the control measurement, the ΔTbefore and ΔTafter were −0.20 ± 0.14 °C and −0.15 ± 0.20 °C (p = 0.68) respectively (Fig. 4b, Fig. S16b, ESI†). We could detect no significant changes of the fluorescence lifetime of mCherry by 1 mM caffeine in the cytosol of C2C12 myotubes.

![Fig. 4 Temperature change measurements of cytosol in C2C12 myotubes (a) Fluorescence lifetime imaging of living C2C12 myotubes expressing cytosolic mCherry captured at 25 °C and 37 °C. Scale bar, 50 μm; calibration bar, 1.6—2.1 ns. (b) Comparison of the ΔT of cytosol in myotubes between 1 mM caffeine (red circles) and DMEM (blue circles) stimulations. The red and blue diamonds, the mean values in each condition, respectively.](image)
other types of cells (e.g. H35 cells: 15 pW/cell)\(^{28}\), and cells which naturally have a thermogenic role in the body would be expected to produce more heat. Another aspect considered is heterogeneous temperature increase in cells. No detectable temperature increase was observed in the fluorescence lifetime measurement of cytosolic mCherry of C2C12 myotubes stimulated by 1mM caffeine (Fig 4b). This result also shows an analogy with previous studies\(^{5,12}\), where heterogeneous thermogenesis was detected in HeLa and NIH3T3 cells in response to ionomycin stimulation, although their nanothermometers could not demonstrate location specificity of the thermogenesis.

This study demonstrates, for the first time to the best of our knowledge, Ca\(^{2+}\)-induced heat production at the single live cell level in a C2C12 myotube model, which may partially represent NST of skeletal muscle\(^{7,8}\). Contrary to our results showing no detectable temperature changes with 1 \(\mu\)M thapsigargin treatment (Fig 2d, Fig 2f), Kiyonaka et al. reported that cyclopiazonic acid (CPA), a SERCA inhibitor, induced temperature drop of SR\(^{2}\), suggesting that heat production is continuously occurring in untreated cells. Although the reason for these differences are not immediately clear, our results suggest that the substantial heat production at SR happens only when Ca\(^{2+}\) bursts occur in the cytosol.

In conclusion, we have monitored heat production coupled with Ca\(^{2+}\) burst in myotubes by means of frequency domain FLIM. This SERCA-mediated heat production induced by Ca\(^{2+}\) release through RyR matches the physiological mechanism of NST\(^{7,8}\), thus potentially leading to deeper understanding of its mechanism and applications.

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Notes and references