Dual-FRET imaging of IP₃ and Ca²⁺ revealed Ca²⁺-induced IP₃ production maintains long lasting Ca²⁺ oscillations in fertilized mouse eggs

Toru Matsu-ura¹, Hideki Shirakawa², Kenichi G. N. Suzuki³, Akitoshi Miyamoto¹,⁴, Kotomi Sugiuara⁵, Takayuki Michikawa⁶, Akihiro Kusumi⁷ & Katsuhiko Mikoshiba¹,⁸,⁹

In most species, fertilization induces Ca²⁺ transients in the egg. In mammals, the Ca²⁺ rises are triggered by phospholipase Cζ (pLCζ) released from the sperm; IP₃ generated by pLCζ induces Ca²⁺ release from the intracellular Ca²⁺ store through IP₃ receptor, termed IP₃-induced Ca²⁺ release. Here, we developed new fluorescent IP₃ sensors (IRIS-2s) with the wider dynamic range and higher sensitivity (Kd = 0.047–1.7 μM) than that we developed previously. IRIS-2s employed green fluorescent protein and Halo-protein conjugated with the tetramethylrhodamine ligand as fluorescence resonance energy transfer (FRET) donor and acceptor, respectively. For simultaneous imaging of Ca²⁺ and IP₃ using IRIS-2s as the IP₃ sensor, we developed a new single fluorophore Ca²⁺ sensor protein, DYC3.60. With IRIS-2s and DYC3.60, we found that, right after fertilization, IP₃ concentration ([IP₃]) starts to increase before the onset of the first Ca²⁺ wave. ([IP₃] stayed at the elevated level with small peaks followed after Ca²⁺ spikes through Ca²⁺ oscillations. We detected delays in the peak of [IP₃] compared to the peak of each Ca²⁺ spike, suggesting that Ca²⁺-induced regenerative IP₃ production through PLC produces small [IP₃] rises to maintain [IP₃] over the basal level, which results in long lasting Ca²⁺ oscillations in fertilized eggs.

In most species, rises in cytosolic Ca²⁺ concentration ([Ca²⁺]) trigger the egg-embryo transition. Unfertilized eggs, which are arrested at different stages of meiosis in different species, are “activated” and released from the arrest by fertilization¹². In mammals, egg activation is triggered by a periodic series of Ca²⁺ transients, known as Ca²⁺ oscillations³⁴. The response in mammalian eggs lasts for several hours and involves relatively low frequency, large amplitude Ca²⁺ increases⁵. The multiple increases in [Ca²⁺] are essential for completion of all the events of egg activation in mammals⁵,⁶. The first Ca²⁺ transient occurs some minutes after sperm-egg fusion⁷. The Ca²⁺ oscillations in mammalian eggs appear to be a result of Ca²⁺ release via the inositol 1,4,5-trisphosphate (IP₃) receptor/Ca²⁺ release channel (IP₃R) located on the intracellular Ca²⁺ stores⁸. A sperm-specific phospholipase Cζ (PLCζ)⁹, which produces IP₃ via hydrolysis of phosphatidyl 4,5-bisphosphate (PIP₂), is reported as an egg-activating sperm factor¹⁰ in

¹Laboratory for Developmental Neurobiology, Center for Brain Sciences, RIKEN, 2-1 Hirosawa, Wako, Saitama, 351-0198, Japan. ²Department of Applied Physics and Chemistry, The University of Electro-Communications, Tokyo, 182-8585, Japan. ³Center for Highly Advanced Integration of Nano and Life Sciences (G-CHAIN), Gifu University, 1-1 Yanagido, Gifu, 501-1193, Japan. ⁴Laboratory of Single-Molecule Cell Biology, Kyoto University Graduate School of Biostudies, Konoe-cho, Sakyoku, Kyoto, 606-8501, Japan. ⁵Laboratory of Biotechnological Optics Research, Center for Advanced Photonics, RIKEN, 2-1 Hirosawa, Wako, Saitama, 351-0198, Japan. ⁶Okinawa Institute of Science and Technology Graduate University, 1919-1 Tancha, Onna, Okinawa, 904-0495, Japan. ⁷Department of Pharmacology, Keio University School of Medicine, 35 Shinanomachi, Shinkujuku, Tokyo, 160-8582, Japan. ⁸Laboratory for Advanced Immunochemical Studies, ShanghaiTech University, Shanghai, 201210, China.
mammalian species. The microinjection of complementary RNA (cRNA) encoding PLCζ [1] or recombinant PLCζ proteins [1] into unfertilized mouse eggs triggers characteristic Ca²⁺ oscillations like those observed at fertilization. Sperm from transgenic mice with significantly reduced expression of PLCζ display a premature termination of Ca²⁺ oscillations following in vitro fertilization [1]. PLCζ shows extremely high Ca²⁺ sensitivity for its enzymatic activity compared with other PLC isoforms, with 70% maximal activity at 100 nM Ca²⁺ [12]. Therefore, it has been considered that basal cytosolic Ca²⁺ in the fertilized egg can stimulate PLCζ to produce an amount of IP₃ sufficient to trigger the initial release of Ca²⁺, which has not been confirmed experimentally, since single cell imaging using fluorescent IP₃ indicators, such as green fluorescent protein (GFP)-fused to pleckstrin homology domain (GFP-PHD) [14] and fretino-2 [15], failed to clearly detect IP₃ concentration ([IP₃]) changes evoked in fertilized mouse eggs [16,17].

Because all PLC isoforms including PLCζ are activated by Ca²⁺ [12,18-20], there will be further increase in IP₃ production when [Ca²⁺] starts to increase. This positive feedback has been proposed to play a central role for the generation of the upstroke of Ca²⁺ transients [21,22]. Except for PLCζ, members of each of the PLC families are expressed in eggs, and PLC3L1 is reported to contribute generation of Ca²⁺ transients [23] and in theory any of these could be involved in modulating Ca²⁺ oscillations. On the other hand, the positive feedback regulation of Ca²⁺ acting directly on the IP₃R has been proposed to drive regenerative Ca²⁺ increases [24]. Simultaneous detection of [Ca²⁺] and [IP₃] is necessary to figure out the contributions of Ca²⁺-induced IP₃ production from PLCs and Ca²⁺ release form IP₃R for the generation of fertilization-induced Ca²⁺ transients.

In the present study, we developed novel fluorescent resonant energy transfer (FRET)-based IP₃ sensor proteins, designated as IRIS-2s, to visualize IP₃ dynamics in fertilized mouse eggs. The novel IP₃ sensors possess an improved dynamic range compared with the previous sensor, IRIS-1 [25]. A high IP₃ binding affinity variant, IRIS-2.3, can be successfully used to monitor [IP₃] changes naturally induced in fertilized mouse eggs. IRIS-2s contain enhanced green fluorescent protein (EGFP) and Halo-protein with tetramethylrhodamine (TMR) ligand as FRET donor and acceptor, respectively. To monitor [Ca²⁺] and [IP₃] changes simultaneously, we also developed a new Ca²⁺ sensor protein, designated as DYC3.60, which has enhancedcyan fluorescent protein (ECFP) as a solo fluorophore. The pair of IRIS-2s and DYC3.60 contains a new set of fluorophores for dual-FRET imaging, and real-time monitoring with IRIS-2s and DYC3.60 provide us insights into the mechanism underlying the generation of Ca²⁺ oscillations in mouse fertilized eggs.

**Results**

**Construction of IRIS-2s.** We constructed novel IP₃ sensors composed of HaloTag protein (Promega), IP₃ binding domain (IP₃-BD) of mouse IP₃R1 [26], and mEGFP (upper panel in Fig. 1a). HaloTag protein is an engineered, catalytically inactive derivative of a hydrolase that forms a covalent bond with commercially available HaloTag ligands. We used HaloTag® tetramethylrhodamine (TMR) ligand (Promega) as a FRET acceptor for mEGFP. Amino acid residues 224–575 and 224–579 of mouse IP₃R1 were used for IRIS-2 and IRIS-2.3, respectively (upper panel in Fig. 1b), to manipulate the IP₃ binding affinity of the sensors. We constructed IRIS-2-Dmut, in which two critical amino acid residues (Thr267 and Lys508) for IP₃ binding have been replaced in IRIS-2, as a negative control [25] (upper panel in Fig. 1b). The upper panel in Fig. 1c shows emission spectrum of IRIS-2 when excited at 480 nm. Purified IRIS-2 with addition of HaloTag TMR ligand (IRIS-2-TMR) showed greater TMR emission (565 nm) and lesser EGFP emission (510 nm) (green line in Fig. 1c) compared with those of untreated IRIS-2 (red line in Fig. 1c), indicating that FRET between EGFP and TMR occurs in IRIS-2-TMR. The addition of 100 μM IP₃ increased the EGFP emission and decreased the TMR emission (blue line in Fig. 1c), indicating that the FRET efficiency of IRIS-2-TMR decreases upon IP₃ binding (Fig. 1a). The relative change in the EGFP/TMR emission ratio of IRIS-2 was measured with zero and 100 μM IP₃ (Fig. 1c). The ratio of IRIS-2 TMR monitored with zero and 100 μM IP₃ was 71 ± 3% (n = 3); Fig. 1d) was three times larger than that of IRIS-2 (1.7 ± 0.2 μM; n = 3; Fig. 1d). The Kd value of IRIS-2-TMR (0.047 ± 0.006 μM; n = 3; Fig. 1d) was 36-times smaller than that of IRIS-2 (1.7 ± 0.2 μM; n = 3; Fig. 1d). The Kd value of IRIS-2-TMR was 3-times larger than that of IRIS-1 (0.55 ± 0.06 μM) [25].

As a partner of IRIS-2s, we developed a FRET based Ca²⁺ indicator with single fluorophore to avoid fluorescent overloading with IRIS-2s. We introduced a non-fluorescent mutation (Y145W) into a yellow fluorescent protein, cp173Venus, of YC3.60 [27] (lower panels in Fig. 1a,b). The resultant protein have a fluorescent spectrum as same as ECFP, and addition of 100 μM Ca²⁺ decreased its emission by FRET quenching. The peak fluorescent amplitude was 71 ± 3% (n = 3) reduced after addition of Ca²⁺ in DYC3.60 (lower panel in Fig. 1c). Fluorescence from the three fluorophores used in IRIS-2s and DYC3.60 can be easily separated (Fig. 1e). Figure 1f shows time course changes of fluorescence from DYC3.60 and IRIS-2 in glutamate stimulated mGlur5-expressing HeLa cells. Less overlaps of excitation and emission spectra of IRIS-2 and DYC3.60 allowed dual-FRET imaging of Ca²⁺ and IP₃ even without spectral unmixing [28] (Fig. 1f).

**Characterization of IRIS-2s and DYC3.60 expressed in cultured mammalian cells.** IRIS-2-TMR and DYC3.60 were uniformly distributed within the cytosol when expressed in HeLa cells (Fig. 2a–e). Halo-TMR staining increased fluorescent signal detected by a 573–613-nm emission filter (Fig. 2b,d). The frequency of Ca²⁺ oscillations monitored with Indo-5F in mGlur5-expressing HeLa cells were not significantly different among IRIS-2s, IRIS-2-Dmut-, and DYC3.60-expressing cells (50 ± 16 Hz for IRIS-2, n = 9; 55 ± 6 Hz for IRIS-2-Dmut, n = 4; 47 ± 10 Hz, n = 6 for DYC3.60) (Fig. 2f-h). IRIS-2-TMR signals did not return to its basal level during the intervals between Ca²⁺ transients, and its fluctuation was synchronous with Ca²⁺ oscillations (Fig. 2f). These characteristic IP₃ dynamics monitored with IRIS-2-TMR in HeLa cells are almost same as those recorded with other FRET-based IP₃ sensors [15,25,29,30].
Initiation, maintenance, and termination of Ca\(^{2+}\) oscillations in HeLa cells. Figure 2i–k show imaging data of IRIS-2 and DYC3.60 in mGluR5-expressing HeLa cells. The dual FRET imaging clearly showed [IP\(_3\)] increase precedes [Ca\(^{2+}\)] rise as same as our previous report with IRIS-125 (Fig. 2i,j). Figure 2k shows a phase plane trajectory of [IP\(_3\)] and [Ca\(^{2+}\)] imaging data. [IP\(_3\)] gradually increased from 1st to 4th Ca\(^{2+}\) spikes, and then,
repeated \( \text{Ca}^{2+} \) spikes occurred in the certain range of \([\text{IP}_3]\) (Fig. 2k). In the range of \([\text{IP}_3]\), the trajectory cycled at almost the same orbit, suggesting that the trajectory is in a limit cycle (Fig. 2k). After termination of agonist stimulation, \([\text{IP}_3]\) decreased below the range of limit cycle maintenance, which resulted in the termination of \( \text{Ca}^{2+} \) oscillations (Fig. 2i,k). In the initial phase of \( \text{Ca}^{2+} \) oscillations, \([\text{IP}_3]\) increase precedes \( \text{Ca}^{2+} \) spikes (Fig. 2k), suggesting that \([\text{IP}_3]\) increases induce \( \text{Ca}^{2+} \) spikes. In the limit cycle phase, \( \text{Ca}^{2+} \) spikes occur without marked \([\text{IP}_3]\) increases (Fig. 2k), suggesting that \( \text{Ca}^{2+} \) induced positive and negative feedbacks to IP3R autonomously induce \( \text{Ca}^{2+} \) spikes24. \( \text{Ca}^{2+} \) oscillations last as long as \([\text{IP}_3]\) maintained in the range of limit cycle. Termination of agonist stimulation induces \([\text{IP}_3]\) decrease below to the range maintaining the limit cycle. \([\text{IP}_3]\) necessary to induce \( \text{Ca}^{2+} \) spike should be different at the initial state and later state of \( \text{Ca}^{2+} \) oscillations because \( \text{Ca}^{2+} \) directly or indirectly inactivates IP3R28,31. Thus, \( \text{Ca}^{2+} \) disappears even \([\text{IP}_3]\) above the basal level at the termination of \( \text{Ca}^{2+} \) oscillations (Fig. 2i).

**Characterization of IRIS-2 in UV-uncaging experiments.** Next, we checked the compatibility of \( \text{IP}_3 \) sensors with UV-uncaging. Caged-compounds are light-sensitive probes that functionally encapsulated biomolecules in an inactive form. The active compounds can be released from caged-compounds with UV light in most of caged-compounds. IRIS-1 or IRIS-2 were expressed in HeLa cells and irradiated by UV pulses (Supplementary Fig. 2a,b). We found UV irradiation caused temporal reduction of fluorescence of both EGFP and Venus in IRIS-1 expressing cells (Supplementary Fig. 2a). Because of the difference of the signal reduction between those fluorescent proteins, the fluorescent ratio of IRIS-1 was significantly reduced (\(-1.9 \pm 0.7\%\), \(n = 22\)). In contrast, the fluorescent signals from EGFP and HaloTag-TMR were stable after the UV irradiation (Supplementary Fig. 2b), which resulted in successful detection of \([\text{IP}_3]\) changes after UV-uncaging of caged-\( \text{IP}_3 \) (Supplementary Fig. 2c).

**Detection of \( \text{IP}_3 \) concentration changes in fertilized mouse eggs.** To detect \( \text{IP}_3 \) dynamics in fertilized mouse eggs, IRIS-1, IRIS-2, or IRIS-2.3 was expressed in eggs by cRNA injection. For the simultaneous monitoring of \( [\text{Ca}^{2+}] \) changes, we first used Indo-5F as a \( \text{Ca}^{2+} \) indicator according to the method described previously25. As shown in Supplementary Figure 3, we did not detect any changes of IRIS-1 signals in fertilized eggs. Not only the fails of the detection of \( \text{IP}_3 \) changes, it was difficult to detect \([\text{Ca}^{2+}] \) changes after addition of...
suggesting that IRIS-1 works as a significant IP3 buffer. We also tested IRIS-2 expressing eggs for 
Ca2+
 via Ca2+
arrow. The time point of the change in the rate of rise in the IRIS-2.3TMR signal is shown by a vertical broken 
line. (d) Left panel shows the rates of [IP3] increases before and after the shoulder point of the first Ca2+
 spike. Right panel shows peak [IP3] change before [Ca2+] rise and first [IP3] peak after [Ca2+] rise. n = 7. *p < 0.05, Student’s t-test.

Figure 3. IRIS-2 TMR and IRIS-2.3 TMR Signals in fertilized mouse eggs. (a,b) [IP3] and [Ca2+] dynamics detected 
by IRIS-2s and Indo-5F. Normalized emission ratio changes (ΔR/Rbase) of IRIS-2_TMR (a) and IRIS-2.3_TMR (b) 
are plotted with red lines. [Ca2+] changes detected with Indo-5F are shown with black lines (a,b). Sperm was 
added at time zero (a,b). Fluorescent images were acquired each 4 sec in (a) and each 10 sec in (b). (c) Time 
courses of emission ratio changes of Fura-2 (black) and IRIS-2.3_TMR (red) at a first Ca2+
 transient observed approximately 30 min after the onset of the first Ca2+
 transient, [IP3] continues to increase, and all the following Ca2+
 transients were less compared to IRIS-2-Dmut (number of Ca2+
 spikes: IRIS-1: 1.91 ± 0.13 (n = 3); IRIS-1-Dmut: 3.75 ± 0.5 (n = 4); p = 0.008, Student’s t-test), 
suggesting that IRIS-1 works as a significant IP3 buffer. We also tested IRIS-2 expressing eggs for in-vitro fertilization 
assay and found IRIS-2 expressing eggs had normal Ca2+
 spikes after fertilization (Fig. 3a). However, it was 
also hard to detect clear increases in FRET signals in IRIS-2-expressing eggs during the first Ca2+
 transient evoked after fertilization, while small repetitive transients of IRIS-2_TMR signals synchronous with Ca2+
 oscillations were observed approximately 30 min after the onset of the first Ca2+
 transient (Fig. 3a). On the other hand, we clearly 
detected IP3 increases during the all Ca2+
 transients, including the first Ca2+
 transient, in IRIS-2.3-expressing eggs (Fig. 3b). During the first large Ca2+
 transient, [IP3] continues to increase, and all the following Ca2+
 transients accompanied with a rapid increase and a following slow decline on the elevated level of [IP3] (Fig. 3b). Three 
independent experimental results of [IP3] and [Ca2+] imaging with IRIS-2.3 and Indo-5F at the onset of first Ca2+
 spike were shown in Supplementary Figure 4. We did not find significant difference of numbers of Ca2+
 spikes during 30 min after 1st Ca2+
 spike between IRIS-2 and IRIS-2.3 expressing eggs (IRIS-2: 5.17 ± 1.72, n = 6; IRIS-2.3: 6.33 ± 5.72, n = 9; p = 0.58, student’s t-test).

Initiation of [IP3] and [Ca2+] changes. Next, we investigated the temporal order of the onset of increase 
between [IP3] changes and [Ca2+] changes during the first Ca2+
 transient evoked after fertilization. To detect the initial [Ca2+] changes experimentally, we used Fura-2, whose affinity is higher than that of Indo-5F (Fura-
2: Kd = 135 nM; Indo-5F: Kd = 470 nM), as a Ca2+
 indicator to detect the timing of the onset of the first Ca2+
 transient as precise as possible. As shown in Fig. 3c, [IP3] rise preceded the onset of the initial step of the first Ca2+
 transient for 2.7 ± 2.4 min in 11 of 13 eggs. The initial [IP3] increase should initiate Ca2+
 release from IP3, IP3, and Ca2+
 are the co-agonist of IP3, and open probability of IP3 markedly increase with Ca2+
 in the presence of IP3. Thus, Ca2+-induced Ca2+- release (CICR) from IP3 should have major role to produce initial Ca2+-
spike. The first Ca2+
 transient observed in IRIS-2.3-expressing eggs was composed of two steps separated by a 
shoulder point (dashed line in Fig. 3c) as reported previously35. The peak amplitude and the rising speed of [IP3] 
increased after the shoulder point of the first Ca2+-
 spike (Fig. 3c,d), suggesting acceleration of IP3 production via Ca2+-induced activation of PLC isoforms.

Positive feedback loop to produce rising phase of Ca2+
 spikes. Each Ca2+-
spike of Ca2+-
 oscillations usually form as a result of an initial slow pacemaker rise in [Ca2+] followed by a rapid rise in [Ca2+] to achieve a peak. The accelerated rise of [Ca2+] is suggested that a regenerative process is involved in the generation of the abrupt upstroke35. Such regenerative processes require a positive-feedback element22, and CICR from IP3 and Ca2+-induced IP3 production through PLC have been proposed as candidates of the positive feedback element. In the previous
the amplitudes of IP3 fluctuations were relatively small (Fig. 3b), and the rate of [IP3] rise did not increase during the rising phase of the Ca2+ spikes and IRIS-2.3TMR were differentiated and aligned at the time when the rate of [Ca2+] rise reached its maximum (Fig. 4). In the early phase (from first to 5th transients) of fertilization-induced Ca2+ oscillations, the amplitudes of IP3 fluctuations were relatively small (Fig. 3b), and the rate of [IP3] rise did not increase during the rising phase of Ca2+ spikes, as found in cultured HeLa cells (Fig. 4a,b). The amplitudes of IP3 fluctuations were gradually increased during the later phase of Ca2+ oscillations (Fig. 3a,b), and contrary to the early phase, the onset of the rate of [IP3] rise precedes that of [Ca2+] (Fig. 4c). The result suggests that Ca2+-induced IP3 production through PLC may work as a part of the positive feedback loop to produce abrupt [Ca2+] spikes in later phase of Ca2+ oscillations. However, the peak of the rate of [IP3] rise always delayed from that of [Ca2+] (Fig. 4c), suggesting that CICR from IP3R has major role to produce the rising phase of Ca2+ spikes and [IP3] rises.

[IP3] stayed at the elevated level and did not return to the basal level through Ca2+ oscillations (Fig. 3b).

Dual-FRET imaging of [IP3] and [Ca2+] in fertilized mouse eggs. We also test our dual-FRET pair for [IP3] and [Ca2+] at fertilization of mouse eggs. We microinjected cRNAs of DYC3.60 and IRIS-2.3 into the eggs and stained the eggs with TMR. As shown in Fig. 5a, these fluorophores were distributed evenly in the egg. Well separation of excitation and emission spectra of these fluorophores enabled simultaneous detection of these fluorescence (Figs 1e, 5a,b). As same as the results we obtained with the pair of Indo-5F and IRIS-2.3TMR, we successfully detected fertilization-induced [Ca2+] and [IP3] changes at each Ca2+ spikes and [IP3] changes with DYC3.60 and IRIS-2.3TMR (Fig. 5b and Supplementary video 1). As same as HeLa cells, [IP3] at the termination was higher than that at the initiation of Ca2+ oscillations in fertilized mouse eggs (Fig. 2i and Supplementary Fig. 5).

Ca2+-induced regenerative IP3 production. We also detected delays in the peak of [IP3] compared to the peak of each Ca2+ spike (17 ± 11 sec, n = 63, Fig. 5c,d), suggesting that Ca2+-induced regenerative IP3 production through PLC produces small [IP3] rises at each Ca2+ spike to maintain [IP3] over the basal level, which results in long lasting Ca2+ oscillations in fertilized eggs. Which PLC isoforms contribute to this regenerative process? Eight PLC isoforms are known to express in the mouse egg: PLCβ1, PLCβ2, PLCγ1, PLCγ2, PLCδ1, PLCδ2, PLCε, and PLCζ. From these isoforms, knockout mice of PLCβ1, PLCβ2, PLCγ1, PLCγ2, PLCδ1, PLCδ2, PLCε, and PLCζ are born normally. On the other hand, knockout mice of PLCβ3, PLCδ1, PLCδ2, and PLCζ have problems on development of the embryo. However, dominant-negative experiments employing recombinant SH2 domain to inhibit PLCγ1 and -ε did not inhibit the Ca2+ oscillatory pattern during fertilization. Based on these findings, Igarashi et al. found that reduced expression of PLCβ1 by RNAi resulted in a significant decrease in Ca2+ transients and overexpression of PLCβ1 by cRNA injection resulted in perturbed duration and frequency of Ca2+ oscillations. Thus, Ca2+-induced activation of PLCβ isozymes are the strong candidates which play a pivotal role to the accelerated production of IP3 during Ca2+ spikes in fertilized mouse eggs. To determine the role of PLCβ in

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**Figure 4.** Rate of [Ca2+] and [IP3] changes at each Ca2+ spike. Rate of [Ca2+] and [IP3] changes are shown as differentiated signals of Indo-5F (Em. 460–510 nm) (upper panel) and IRIS-2.3TMR (lower panel) aligned by the time when the differentiated Indo-5F signal was at its maximum (frame 0, broken line) during first Ca2+ transients after fertilization (n = 9) (a), from 2nd to 5th Ca2+ transients (n = 20) (b) and from 10th to 15th Ca2+ transients (n = 20) (c). Error bars correspond to the SD. Arrowheads indicate the peak of the rate of [IP3] rise. Broken vertical lines indicate the peaks of differentiated Indo-5F signals.
the mouse egg, we stimulate unfertilized mouse eggs with 100 μM carbachole (Fig. 5f,g). The stimulation caused Ca²⁺ spikes and a monotonic [IP₃] rise (Fig. 5f). At fertilization, [IP₃] changes always follow after Ca²⁺ spikes. On the other hand, Ca²⁺ spikes did not accompany with delayed [IP₃] rises in carbachole stimulated unfertilized eggs (Fig. 5f). Particularly, IP₃ peak at the first Ca²⁺ spike preceded Ca²⁺ peak (Fig. 5g). These data showed that Ca²⁺-induced IP₃ producing activity is not strong in unfertilized eggs, suggesting that sperm derived PLCζ should participate Ca²⁺-induced IP₃ production. As we showed in Figs 3a,b and 4, Ca²⁺-induced [IP₃] rises increased later phase of fertilization-induced Ca²⁺ oscillations, suggesting that fertilization induces quantitative or qualitative changes of PLC in later phase of Ca²⁺ oscillations.

Discussion

In this study, we developed a dual-FRET pair of biosensors for the detection of [IP₃] and [Ca²⁺] in mammalian cells. The uniqueness of our dual-FRET pair is using single fluorophore for one of the pairs. Replacement of Y145 to W in EYFP is known to produce a non-fluorescent chromoprotein that retains its absorption of emission light. Introduction of the Y145W mutation into cp173Venus of YC3.60 resulted to produce single fluorophore with fluorescent quencher in the Ca²⁺ FRET sensor, DYC3.60. Usually, four fluorophores are necessary for dual-FRET imaging. Most of FRET sensors have cyan and yellow fluorescent proteins, and these fluorescent proteins cover a broad spectral profile. Thus, using FRET sensor with cyan and yellow proteins, it is difficult to find a partner FRET sensor for dual-FRET imaging without using spectral unmixing to distinguish each fluorescent signal mathematically from significantly overlapped fluorescent signals. We offer a dual-FRET imaging with three fluorophores, which gives easier detection and separation of fluorescent signals.

Figure 5. Delayed IP₃ pulses during Ca²⁺ oscillations in fertilized mouse eggs visualized by dual-FRET sensors. (a) Fluorescence images of DYC3.60 (ECFP) and IRIS-2.3TMR (EGFP and TMR) in a single mouse egg. DYC3.60 was illuminated with 425–445 nm light, and IRIS-2.3TMR was illuminated with 460–490 nm light. Scale bar, 10 μm. (b) Dual-FRET imaging of [IP₃] and [Ca²⁺] in a fertilized mouse egg. Signals from DYC3.60 (ECFP) and IRIS-2.3TMR (EGFP and TMR) from single fertilized mouse egg are shown in the left and right panels, respectively. (c) Emission changes in DYC3.60 (black line) and ratio changes of IRIS-2.3TMR (red line) are shown. Sperm was added at time zero. (d) Data shown in (c) on an enlarged time scale. The arrowheads indicate the time of peaks in DYC3.60 signal (black) and IRIS-2.3TMR signal (red). (e) A histogram of the peak time difference between DYC3.60 signals and IRIS-2.3TMR signals (n = 61). The positive value indicates that the peak of DYC3.60 signals precedes that of IRIS-2.3TMR signals (17 ± 11 sec). (f) Emission ratio changes of DYC3.60 (black line) and IRIS-2.3TMR (red line) in an unfertilized egg stimulated with 100 μM carbachole. Asterisks show the peak of each Ca²⁺ spike. Carbachole was added during the time indicated by the horizontal bar. (g) Data shown in (f) on an enlarged time scale around the rise of first Ca²⁺ spike. The dashed lines with red and black triangles indicate the time of peaks in IRIS-2.3TMR signal and DYC3.60 signal, respectively.
Ca\(^{2+}\) transient. The result is consistent with the expectation that highly Ca\(^{2+}\) sensitive PLC\(\zeta\) produces IP\(_3\) at the basal level of [Ca\(^{2+}\)] in the egg cytosol after sperm-egg fusion\(^{12}\). Mehlmann and Kline reported microinjection of small amount of IP\(_3\) (8.6 nM) is able to induce Ca\(^{2+}\) spike in unfertilized mouse eggs\(^{48}\). Our measurements showed the same results that the amount of IP\(_3\) produced in mouse eggs is small even after the fertilization because only IRIS-2.3, which shows the highest IP\(_3\) sensitivity (Kd = 47 nM) among the IP\(_3\) sensors developed, could detect [IP\(_3\)] increases at the onset of the first Ca\(^{2+}\) transients.

IP\(_3\)R has a bell-shaped calcium response curve: the open probability of IP\(_3\)R is activated by low [Ca\(^{2+}\)] and inhibited by high [Ca\(^{2+}\)]\(^{32}\). Based on this finding, De Young and Keizer reported a mathematical model to reproduce Ca\(^{2+}\) oscillations with constant [IP\(_3\)]\(^{24}\). In this previous reports, we showed sustained [IP\(_3\)] increase during Ca\(^{2+}\) oscillations in HeLa cells and fertilized mouse eggs\(^{27}\) (Figs 3 and 4), and the same results were obtained with other IP\(_3\) sensor proteins\(^{29,30}\). Consistently with our results, Mehlmann and Kline reported single microinjection of IP\(_3\) induces Ca\(^{2+}\) oscillations in unfertilized mouse eggs\(^{48}\). Jones et al. also reported Ca\(^{2+}\) oscillations with continuous low level caged-IP\(_3\) photolysis in unfertilized mouse eggs\(^{49}\). PLC\(\zeta\) is a smallest and simplest PLC isoform\(^9\). The activity of PLC\(\zeta\) is regulated by Ca\(^{2+}\) and localization into nucleus after pronuclear formation, and other regulations are not known\(^{50}\). PLC\(\zeta\) has highest Ca\(^{2+}\) sensitivity compared to the other PLC isoforms and is 70% active at the basal level [Ca\(^{2+}\)] in cells\(^{42}\). Thus, PLC\(\zeta\) should be continuously active after fertilization until pronuclear formation\(^{42}\), which should sustain continuous [IP\(_3\)] increase during fertilization-induced Ca\(^{2+}\) oscillations (Figs 3 and 4). We previously found that CICR dominantly work as a positive feedback loop to produce the rising phase of Ca\(^{2+}\) spikes in HeLa cells\(^{37}\). Our data suggest that the mechanism elicits the rising phase of Ca\(^{2+}\) spikes in fertilized mouse eggs is more complex. Initially, CICR dominantly works as the positive feedback loop, and Ca\(^{2+}\)-induced IP\(_3\) production gradually participates to produce Ca\(^{2+}\) spikes cooperatively with CICR in the later phase of Ca\(^{2+}\) oscillations. Ca\(^{2+}\)-induced IP\(_3\) production through PLC\(_{\zeta}\) produces [IP\(_3\)] at each Ca\(^{2+}\) spike to help keeping [IP\(_3\)] over the basal level, which results in long lasting Ca\(^{2+}\) oscillations in fertilized eggs.

In conclusion, we produced FRET sensors with new choices of fluorophores for dual-FRET imaging of [IP\(_3\)] and [Ca\(^{2+}\)]. Less overlaps of excitation and emission spectrum of IRIS-2s and DYC3.60 allowed dual-FRET imaging of Ca\(^{2+}\) and IP\(_3\) even without spectral unmixing. Because of the smaller number of fluorophores, our dual-FRET approach can reduce the effort to detect each fluorescent signal separately. The wider dynamic range and higher sensitivity achieved by IRIS-2.3 will enable the detection of subtle [IP\(_3\)] changes associated with [Ca\(^{2+}\)] changes at egg fertilization to local [Ca\(^{2+}\)] increase events.

Materials and Methods

Animals. Experiments used ddY mice for preparation of oocytes and sperm. All animal experiments were performed in accordance with the guidelines approved by the Animal Experiments Committee of RIKEN Brain Science Institute. All experiments were carried out in accordance with the approved ethical guidelines and regulations.

Gene construction. The FRET donor and acceptor of IRIS-1 were replaced with mEGFP and Halo-protein (Promega), respectively, to produce IRIS-2. Amino acid residues 224–575 of mouse IP3R1 in IRIS-2 were replaced with amino acid residues 224–579 of mouse IP3R1 to produce IRIS-2.3. The Y145W mutant of circular permuted Venus (cp173V-Y145W) was generated using the site-directed mutagenesis. The FRET acceptor of YC3.60 was replaced with cp173Venus-Y145W to produce DYC3.60. IRIS-2, IRIS-2.3 and DYC3.60 cDNAs were cloned into the XbaI site of pFastBac (Invitrogen). The recombinant baculovirus was used for the large-scale expression of IRIS-2 in Sf9 cells as described previously\(^{52}\). The expressed proteins were purified on a HiTrap heparin HP column (GE Healthcare Life Sciences) as described previously\(^{52}\). The full-length cDNA of IRIS-2 was isolated from pcDNA3.1 zeo (+) (Invitrogen) for the expression in HeLa cells. The cDNAs were cloned into the XbaI site of pTNTTM (Promega) with extended poly(A) tail (57 residues) and synthesized cRNAs were injected into mouse oocytes.

Protein expression and purification. The full-length cDNA of IRIS-2 was isolated from pcDNA3.1 zeo (+) by using NheI and XbaI sites and was cloned into the XbaI site of baculovirus transfer vector pFastBac1 (Invitrogen). The recombinant baculovirus was used for the large-scale expression of IRIS-2 in Sf9 cells as described previously\(^{52}\). The expressed proteins were purified on a HiTrap heparin HP column (GE Healthcare Life Sciences) as described previously\(^{52}\). HeLa cells were cultured in Dulbecco’s modified Eagle medium supplemented with 10% heat-inactivated fetal bovine serum. HeLa cells were transfected with expression vectors by transfection reagent (Mirus TransIT). One day after the transfection, cells were used for imaging experiments.

Preparation of RNA. Plasmids carrying IRIs or DYC3.60 were digested by NdeI, and linearized DNA fragments were purified with Wizard SV Gel and PCR clean-up Kit (Promega). They were used as the templates for RNA transcription by T7 polynuclease using T7 mMESSAGEmACHINE Kit (Ambion). RNA was purified using RNeasy MinElute Cleanup Kit (Qiagen) and stored at -80°C until use.

Preparation of gametes. Full grown immature oocytes were collected from the follicles in the ovaries of female mice 47–49 h after the injection of pregnant mare serum gonadotropin. Isolated oocytes were freed from cumulus cells mechanically by pipetting in M2 medium, and then cRNAs of IRIS-1, IRIS-2, IRIS-2.3, DYC3.60 or dKeima570 were injected as described below. Sperm was collected from the caudal epididymides and were incubated in M16 medium for 16 h at 37°C with 5% CO\(_2\). Only eggs maturated normally to metaphase II with the
first polar body were used in the following experiments. After loaded with 2 μM of Indo-5F or Fura-2 for 30 min in the M2 medium, eggs were freed from the zona pellucida by brief treatment with acidic Tyrode’s solution (pH 2.5) for insemination. Sperm was added during imaging experiments.

Imaging. After loading HeLa cells with 10 μM Indo-5F-AM (AnaSpec), imaging was performed under the constant flow (2 ml/min) of the balanced salt solution containing 20 mM Hepes, pH 7.4, 115 mM NaCl, 5.4 mM KCl, 1 mM MgCl2, 1.3 mM CaCl2, and 10 mM glucose as an imaging media at 37 °C through an inverted microscope (IX71 or IX81; Olympus) with a cooled charge-coupled device (CCD) camera (ORCA-ER; Hamamatsu Photonics) and a 40×, 1.35 NA, oil-immersion objective (Olympus). For the fluorescent images of IRIS-1 and Indo-5F, an emission splitter (W-view; Hamamatsu Photonics) was used with a light source exchanger (DG-4; Sutter Instrument Co.) on the IX71 inverted microscope. Sequential excitation of IRIS-1 and Indo-5F was performed by using a 450-nm dichroic mirror and two excitation filters (a 425–445 nm filter for IRIS-1 and a 360-nm filter for Indo-5F). Emissions from IRIS-1 and Indo-5F were split with a 460–510-nm filter (for IRIS-1 and Indo-5F), a long-path 520-nm (for IRIS-1) barrier filter, and two 505-nm dichroic mirrors equipped in W-view.

Eggs were incubated with M2 buffer at 37 °C on IX81 inverted microscope. Ca2+ and IP3 were visualized with sets of Indo-5F and IRIS-1, Indo-5F and IRIS-2, Indo-5F and IRIS-2.3, Fura-2 and IRIS-2.3, or DYC3.60 and IRIS-2.3, respectively. Sequential excitation of Ca2+ and IP3 indicators was performed by using dichroic mirrors (a 400-nm mirror for Indo-5F and a 450-nm mirror for IRIS-1 and DYC3.60 and a 505-nm mirror for IRIS-2 and IRIS-2.3) and excitation filters (a set of 340 and 380-nm filters for Fura-2 and a 380-nm filter for Indo-5F and a 425–445 nm filter for IRIS-1 and DYC3.60 and a 460–490-nm filter for IRIS-2 and IRIS-2.3). Emissions from Ca2+ and IP3 indicators were split with emission filters (a set of 400–420 and 460–510-nm filters for Indo-5F and a set of 460–510 and 525–565 filters for IRIS-1 and DYC3.60 and a 510–550 filter for Fura-2 and a set of 510–550 and 573–613-nm filters for IRIS-2 and IRIS-2.3, respectively, and three filter exchangers (Lamda 10; Sutter Instruments, IX2-RFACA; Olympus).

Image acquisition was performed with MetaFluor ( Molecular Devices). Data analysis was performed with MetaFluor and Igor Pro (WaveMetrics) softwares. The EGFPP/TMR emission ratio (IRIS-2s), the ECFP/Venus emission ratio (Indo-1) and the ratio of 510–550 nm emission excited at 340 nm and 510–550 nm emission excited at 380 nm (Fura-2) were defined as R. ∆R was defined as R - Rbase, where Rbase is the basal level of R. Baseline drift in each experiment was corrected with subtracting the trend line which is calculated with the line around the beginning of each experiment.

Uncaging of caged-IP3. HeLa cells transfected with IRIS-2 and DYC3.60 were loaded with 10 μM membrane permeable caged-IP3 (iso-Ins(1,4,5)P3/PM (caged), Enzo Life Science). The uncaging stimulation was done with extra light source (mercury lamp) equipped in IX81, filtered by a 333–348-nm filter and a 400-nm dichroic mirror, illuminated the cells through 20×, 0.50 NA, water-immersion objective (Olympus).

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Author Contributions
T. Matsu-ura, K. Suzuki, K. Sugiura and A.K. invented IRIS-2 variants. T. Matsu-ura and A.M. invented DYC3.60. T. Matsu-ura and H.S. established the method of the expression of IP3 sensors in mouse eggs. T. Matsu-ura performed other experiments. T. Matsu-ura and H.S. analyzed data. T. Matsu-ura, H.S., T. Michikawa, K. Suzuki, A.K. and K.M. wrote the manuscript. K.M. supervised the study.

Additional Information
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