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Dynamics of male and female chromatin during karyogamy in rice zygotes

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Chromatin from the rice egg nucleus initially moves into the sperm nucleus during karyogamy, which enlarges the latter, and then the male chromatin decondenses as nuclear fusion progresses.
Footnotes

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ABSTRACT

In angiosperms, the conversion of an egg cell into a zygote involves two sequential gametic processes: plasmogamy, the fusion of the plasma membranes of male and female gametes, and karyogamy, the fusion of the gametic nuclei. In this study, the nuclei and nuclear membranes of rice (*Oryza sativa* L.) gametes were fluorescently labeled using histone H2B-GFP/RFP and SUN2-GFP, respectively, which were heterologously expressed. These gametes were fused *in vitro* to produce zygotes, and the nuclei and nuclear membranes in the zygotes were observed during karyogamy. The results indicated that the sperm nucleus migrates adjacent to the egg nucleus 5 to 10 min after plasmogamy via an actin cytoskeleton, and the egg chromatin then appears to move unidirectionally into the sperm nucleus through a possible nuclear connection. The enlargement of sperm nucleus accompanies this possible chromatin remodeling. Then, 30 to 70 min after fusion, the sperm chromatin begins to decondense with the completion of karyogamy. Based on the present observations, the development of early rice zygotes from plasmogamy to karyogamy was divided into eight stages, and, using RT-PCR analyses, paternal and *de novo* synthesized transcripts were separately detected in zygotes at early and late karyogamy stages, respectively.
INTRODUCTION

In angiosperms, the sporophytic generation is initiated by double fertilization to form seeds (reviewed in Raghavan, 2003). In double fertilization, one sperm cell from the pollen grain fuses with the egg cell, and the resultant zygote develops into an embryo that can transmit genetic material from the parents to the next generation. The central cell fuses with the second sperm cell to form a triploid primary endosperm cell, which develops into the endosperm that nourishes the developing embryo/seedling (Nawaschin, 1898; Guignard, 1899; Russell, 1992). The conversion of the egg cell into the zygote is completed by two sequential gametic processes, plasmogamy, the fusion of the male and female gametes’ plasma membranes, and karyogamy, the fusion of the male and female gametes’ nuclei.

GENERATIVE CELL SPECIFIC 1/HAPLESS 2 (GCS1/HAP2) and EGG CELL 1 (EC1) have been identified as male and female gamete factors for plasmogamy, respectively. GCS1/HAP2 was identified as a key male membrane protein with a single transmembrane domain and a histidine-rich domain in the extracellular region (Mori et al., 2006; von Besser et al., 2006). Recently, Sprunck et al. (2012) indicated that small cysteine-rich EC1 proteins, which accumulated in the storage vesicles of Arabidopsis egg cells, were secreted via exocytosis upon sperm cell attachment to the egg cell, and the secreted EC1 proteins functioned in redistributing GCS1/HAP2 proteins to the sperm cell surface, resulting in successful gamete fusion.

Karyogamy involves the approach and fusion of male and female nuclei in a zygote. An intensively investigated karyogamic event is the fusion of two haploid nuclei during yeast
mating, in which cytoskeleton-dependent nuclear migration, termed nuclear congression, and chaperon/endoplasmic reticulum (ER)-protein-dependent nuclear fusion have been reported (Kurihara et al., 1994; Melloy et al., 2009; Tartakoff and Jaiswal, 2009; Gibeaux et al., 2013). Additionally, the cell cycle arrest and resumption induced by fertilization have been well studied using animal eggs (reviewed in Kishimoto, 2003). Interestingly, in starfish eggs, Tachibana et al. (2008) indicated that cyclin B-Cdk1 was required to form the sperm astral microtubules on which female and male pronuclei migrate during karyogamy. This suggested alternative cyclin B-Cdk1 function in addition to its well-known role in mitotic spindle assembly during M phase (Karsenti and Vernos, 2002). In animals, the pronuclear migration in the fertilized gamete during karyogamy is dependent on the microtubule system. However, changes in the actin distribution by cytocharasin D, an inhibitor of actin polymerization, in mouse eggs is also reported to prevent pronuclear migration (Maro et al., 1984).

In angiosperms, the mechanisms responsible for the migration of the sperm nucleus toward the egg nucleus in the fertilized egg cell still remain undetermined, although the actin corona/patch is organized around or in female gametes and supposed to function in gamete fusion (Huang and Russell, 1994; Fu et al., 2000). Using transmission electron microscopy, nuclear fusion in angiosperm zygotes was first observed in the pollinated ovaries of cotton (Jensen, 1964), and later, karyogamy in several kinds of plants, including grasses, was reported (Van Went, 1970; Wilms, 1981; Mogensen, 1982; You and Jensen, 1985; Hause and Schröder, 1987; Tian and Shen, 1992). From these observations, three main steps for nuclear fusion have been described: (I) the external nuclear membranes become closely apposed, or in contact, via the ER and then fuse; (II) the inner nuclear membranes fuse, and bridges are formed between
nuclei; and (III) the bridges enlarge and can entrap some cytoplasm. The male chromatin then begins to decondense.

To address the dynamics of karyogamy, Faure et al. (1993) obtained a precise time-table of the karyogamic progression by observing the nuclear approach and fusion in zygotes produced by the electro-fusion of isolated maize gametes. They indicated that karyogamy in the zygotes occurred within 2 to 3 h after gamete fusion. DAPI staining of barley zygotes isolated from pollinated ovaries also indicated the integration of male chromatin into the egg nucleus and the decondensation of the male chromatin in the fused nucleus occurs a few hours after pollination (Mogensen and Holm, 1995). As for the membrane fusion of gamete nuclei, Maruyama et al. (2010) revealed that Bip, an ER-localized HSP70 chaperon, is essential for the nuclear fusion of polar nuclei in the central cell. In addition, NFD1, a component of mitochondrial ribosomes, is also reported to be involved in the nuclear fusion of Arabidopsis karyogamy (Portereiko et al., 2006). After nuclear fusion, chromatin that was tightly packed in the sperm nucleus decondenses in the fused nucleus. Interestingly, nascent synthesis of the paternal mRNA and proteins in fused gametes are reported to coincide with male chromatin decondensation (Scholten et al., 2002), and the zygotic genome appears to switch on only hours after fertilization (Meyer and Scholten, 2007; Ingouff et al., 2007; Zhao et al., 2011; Nodine and Bartel, 2012). Moreover, it has been indicated that Histone 3 (H3) variants contributed by paternal or maternal chromatin are diminished from the Arabidopsis zygotic nucleus within a few hours after fertilization and these H3 variants are then de novo synthesized in the zygote, suggesting that the zygotic resetting of the H3 variants is part of the epigenetic reprogramming for zygotic development and embryogenesis (Ingouff et al., 2010). Although the mechanisms
behind karyogamy and zygotic development have gradually become evident, how the male nucleus approaches and fuses with the female nucleus still remains unclear. This may be because of their location in zygote within the embryo sac, which is deeply embedded in the ovaries, making them difficult to trace.

To investigate the mechanisms of gametic and/or early zygotic development in angiosperms, we previously established a procedure to isolate rice gametes and an *in vitro* fertilization (IVF) system to produce zygotes that can develop into fertile plants (Uchiumi et al., 2006; 2007). The rice IVF system has been used for analyzing gene/protein expression profiles in gametes/zygotes and polarity changes during zygotic development (Nakajima et al., 2010; Sato et al., 2010; Ohnishi et al., 2011; Abiko et al., 2013a; 2013b). In the present study, zygotes were prepared by IVF of rice gametes heterologously expressing histone H2B-GFP/RFP and SUN2-GFP, which labeled the nuclei and nuclear membranes, respectively, and the dynamics of karyogamy in the rice zygotes were monitored. The results indicated that the sperm nucleus migrated adjacent to the egg nucleus in an actin filament-dependent manner within 5 to 10 min after gamete fusion, and then the egg chromatin became detectable in the sperm nucleus. Upon female chromatin transfer to inside the sperm nucleus via a possible connection between nuclei, the size of the nucleus was enlarged. Thereafter, according to the progression of nuclear fusion, sperm chromatin began to decondense 30 to 70 min after fusion. *De novo* gene expression was detectable when male chromatin had almost completely decondensed, and then karyogamy was completed. In addition, based on the karyogamic progression, the precise staging of the early zygote was also presented for subsequent analyses on the early development of the fertilized egg/zygote.
RESULTS

Fluorescent labeling of chromatin and nuclear membrane in rice gametes

In egg cells prepared from transformed rice expressing H2B-GFP under the ubiquitin promoter, the nucleus was clearly visible (Fig. 1, A and B; Abiko et al. 2013b). In addition, observations for divisional profile of the zygotes, which were isolated from pollinated ovaries of the transformant, resulted in the possible identification of fluorescently labeled chromosomes during mitosis (Supplemental Fig. S1). These data are consistent with the report that H2B-GFP is incorporated into the chromatin and chromosome structures (Howe et al., 2012). In addition, nucleoli were strongly fluorescently labeled, although why such a strong signal was detected cannot be explained. A histone H2B-RFP fusion protein was heterologously expressed in rice plants under the DD45 promoter (Steffen et al., 2007), which is known to be egg-cell specific in Arabidopsis. In egg cells isolated from the transformants, the nuclei were labeled by the fusion protein (Fig. 1, C and D), suggesting that the DD45 promoter could function as an active promoter in rice egg cells as well. In pollen grains from transformants expressing H2B-GFP, two putative sperm nuclei and a vegetative nucleus were visible (Fig. 1, E and F). When pollen grains were immersed in mannitol adjusted to 370 mOsmol kg⁻¹ H₂O, the pollen grains burst, releasing their contents, and the nuclei of the two released sperm cells and the vegetative nucleus were visible (Fig. 1, G and H).

Arabidopsis SUN2 protein, a nuclear membrane protein (Graumann et al., 2010), was tagged with GFP, and the fusion protein was heterologously expressed in transformed rice under the
ubiquitin promoter. The putative nuclear membrane was clearly visible in isolated egg cells (Fig. 1, I-K) and sperm cells (Fig. 1, L-N).

**Male chromatin decondensation**

A sperm cell expressing histone H2B-GFP was electro-fused with a wild-type egg cell, and fluorescence in the resulting zygote was observed (Fig. 2, A and B). The sperm nucleus migrated adjacent to the egg nucleus approximately 5 to 10 min after fusion (Fig. 2, C–F; Supplemental Fig. S2, A and B). Then, the sperm nucleus appeared to enlarge (Fig. 2, E–H; Supplemental Fig. S2, A–D), and the signal from the male chromatin began to distribute within the fused nucleus (Fig. 2, I–L; Supplemental Fig. S2, E–J). These observations suggested that nuclear fusion, and the subsequent decondensation of sperm chromatin, had occurred. After or during decondensation of the male chromatin, two nucleoli were detectable (Fig. 2, M and N; Supplemental Fig. S2, K and L), and then the nuclear signal intensity became progressively stronger (Fig. 2, M–R; Supplemental Fig. S3). This gradual increase in the H2B-GFP protein signal suggested that the H2B-GFP transcript and protein were *de novo* synthesized in the zygotes during or immediately after nuclear fusion, consistent with the report that nascent mRNA and protein synthesis coincided with the decondensation of male chromatin in fused nuclei of maize zygotes (Scholten et al., 2002).

In this study, 50 independent zygotes produced by the fusion of a wild-type egg cell with a sperm cell expressing H2B-GFP were observed every 15 to 20 min after fusion to estimate the time-table from gamete fusion to the onset of sperm chromatin decondensation. Male chromatin decondensation was most prominent 31 to 70 min after fusion (Fig. 3), although the time ranged
from 31 to 170 min after fusion. These results indicated that the time course from gamete fusion to decondensation in rice zygotes was almost equivalent to that in zygotes of maize (2 h after gamete fusion; Scholten et al., 2002), barley (80 to 100 min after pollination; Mogensen and Holmb, 1995) and Arabidopsis (2 h after fertilization; Ingouff et al., 2010).

**Cytoskelton and karyogamy**

The effects of latrunculin B, an inhibitor of actin polymerization (Coue et al., 1987; Spector et al., 1989), and oryzalin, an inhibitor for tubulin assembly (Hugdahl and Morejohn 1993), on karyogamy in rice zygotes were observed. In zygotes, which were prepared from the fusion of a sperm cell expressing H2B-GFP with an egg cell treated with 200 nM latrunculin B, karyogamy progressed normally (Fig. 4, A-D), and the zygotic nucleus was clearly visible after the completion of karyogamy (Fig. 4, E and F). When an egg cell treated with 2 μM latrunculin B was used for the production of a zygote, the sperm nucleus did not approach the egg nucleus (Fig. 4, G-J). In zygotes at 12 h after introduction, the sperm nucleus was still unfused with the egg nucleus, and interestingly, the fluorescent signal was detected in both the sperm and egg nuclei (Fig. 4, K and L). The effect of latrunculin B on actin organization in egg cells was also observed (Supplemental Fig. S4). In the egg cell, a mesh-like structure labeled by Lifeact-tagRFP was clearly visible, and the structure remained evident after treatment with 200 nM latrunculin B for 120 min. However, the structure was not visible in egg cells treated with 2 μM latrunculin B for 90 min. The difference in the effects of latrunculin B on karyogamy between the 200 nM and 2 μM concentrations (Fig. 4) may be caused by differences in the actin filament structure at the respective inhibitor concentrations. In contrast to latrunculin B,
oryzalin showed no effect on karyogamy in rice zygotes at 10 µM (data not shown) and 100 µM (Fig. 4 M-R) concentrations.

A sperm cell and an egg cell expressing H2B-GFP and Lifeact-tagRFP, respectively, were fused to trace the migration of sperm nucleus and organization of actin filaments in a zygote. Actin filaments existed around or near sperm nucleus, and the sperm nucleus appeared to migrate along actin filaments (Fig. 5). In contrast, sperm nucleus hardly migrated in zygote when actin-depolymerization was induced by latrunculin B treatment.

Movement of female chromatin and enlargement of the sperm nucleus

A wild-type sperm cell and an egg cell expressing H2B-GFP were fused. Although the sperm nucleus was not visible during intracellular migration or shortly after contacting the egg nucleus (Supplemental Fig. S2, M and N), interestingly, the fluorescent signal was detected in the sperm nucleus as well as the egg nucleus 20 min after fusion (Supplemental Fig. S2, O and P). Thereafter, the nuclear fusion progressed and karyogamy was completed (Supplemental Fig. S1, Q–X). In addition, the fusion process of the nuclear membranes was observed when male and female gametes, whose nuclear membranes were labeled with SUN2-GFP, were fused (Supplemental Fig. S5).

To visualize karyogamy in detail, zygotes produced by electro-fusion of a sperm cell expressing H2B-GFP and an egg cell expressing H2B-RFP were observed using a confocal laser scanning (CLS) microscope. In zygotes, both the GFP- and RFP-derived signals from the sperm and egg nuclei, respectively, were observed, with the signal from the egg nucleus localizing inside the sperm nucleus at 20 min after fusion (Fig. 6, A–C). This suggests that female
chromatins labeled by H2B-RFP or free H2B-RFP proteins in egg nucleus were distributed into the sperm nucleus through possible connection between male and female nuclear membranes. Alternatively, the possibility cannot be excluded that H2B-RFP proteins translated in fused egg cell were transported into sperm nucleus and the H2B-RFP-derived signal became detectable. Therefore, an egg cell expressing H2B-RFP was treated with 2 μM latrunculin B and fused with an sperm cell expressing H2B-GFP to see whether H2B-RFP signal becomes detectable in sperm nucleus which does not contact with egg nucleus. The results indicated that the H2B-RFP-derived signal was not detected in sperm nucleus at least 150 min after fusion (Supplemental Fig. S6), suggesting that detection of the H2B-RFP in sperm nucleus attaching with egg nucleus within 20 min after fusion (Fig. 6, A-C) was not due to transport of translational products in fused egg but to migration of egg nucleus contents through possible connection between male and female nuclei. In addition, since most H2B-GFP proteins heterologously expressed in the rice egg cells and zygotes appears to be incorporated in chromatins and chromosomes (Supplemental Fig. S1), detected H2B-RFP signals would be from egg chromatin rather than free H2B-RFP proteins in egg nucleus. However, a high time-resolution observation should be conducted for judging which possibility is the plausible.

The putative egg chromatin signal became more condensed in the sperm nucleus (Fig. 6, D–I), but sperm chromatin was not detected in the egg nucleus. This indicated that sperm chromatin did not migrate into the egg nucleus at this karyogamic stage. Thereafter, nuclear fusion and the decondensation of male chromatin progressed (Fig. 6, J–R), and finally karyogamy was completed (Fig. 6, S–U).

The possible one-way migration of egg chromatin into the sperm nucleus explains the
enlargement of the sperm nucleus during or after contact with the egg nucleus (Fig. 2, E–H; Supplemental Fig. S2, A–D). Thus, a sperm cell expressing SUN2-GFP was fused with an egg cell expressing H2B-RFP, the serial sectional images of the nuclei in the fused cells were obtained with a CLS microscope, and the volume of the sperm nucleus before and after the incursion of female chromatin was measured. Female chromatin was not detectable in the sperm nucleus at 15 min after fusion, but the signal was observed at 20 min after fusion (Fig. 7, A and C). Although the detection of the egg nucleus contents in sperm nucleus suggests that the connection between male and female nuclear membranes was formed until 20 min after fusion, the SUN2-GFP signal was uniformly detected on the sperm nuclear membrane, and such possible connection could not be detected using fluorescent labeling of nuclear membrane by SUN2-GFP (Fig. 7, C). To observe the initial connection between nuclear membranes, termed inter-nuclear bridge, electron microscopic analysis is essential as reported (Jensen, 1964; Van Went, 1970; Wilms, 1981; Mogensen, 1982; You and Jensen, 1985; Hause and Schröder, 1987; Tian and Shen, 1992). When egg chromatin became detectable in the sperm nucleus, its volume was slightly larger than prior to the egg chromatin’s distribution (Fig. 7, B and D). The volume of the sperm nucleus continues to increase (Fig. 7, C-H) and a possible fusion point between the sperm and egg nuclei was observed (Fig. 7G). The fusing area enlarged as nuclear fusion progressed (Fig. 7, I and J). The volume of the sperm nucleus after the incursion of egg chromatin was approximately 1.75 fold larger compared with prior to egg chromatin incursion (Fig. 7K: before egg chromatin incursion, 129.34 ± 25.99 μm³; after egg chromatin incursion, 225.58 ± 24.08 μm³, n=3). Statistical tests indicated that this difference was highly significant (P<0.01).
DISCUSSION

Based on the present observations of karyogamy in rice zygotes, a schematic diagram of the progression of karyogamy is presented in Figure 8. After gamete fusion (stage I), the sperm nucleus migrates inside of the fused cell to appose the egg nucleus via an actin filament-dependent mechanism (stages II and III). In contrast to actin cytoskeleton-dependent nuclear migration, in many animal species, it has been known that pronuclear congression in fused egg depends on a microtubule aster, which is built from paternally inherited centrioles and maternally provided pericentriolar materials (Chambers, 1939; Schatten, 1994; Reinsch and Gönczy, 1998). The actin filament-dependent sperm nucleus migration in plant zygotes may be consistent with the characteristics of sperm cell, the male gamete in angiosperms, which does not possess the centrioles. After apposing of nuclei (stage III), female chromatin possibly moves into the sperm nucleus possible inter-nuclear connection, resulting in sperm nucleus enlargement, but the sperm chromatin remains tightly packed (stage IV). Male chromatin begins to decondense (stage V), and male chromatin further decondenses according to the progression of membrane fusion (stages VI and VII). The karyogamic event is finally completed at stage VIII.

The nascent synthesis of mRNAs and proteins from the parental genomes in zygotes was reported to be initiated either during karyogamy or within hours after fertilization in maize, Arabidopsis, and tobacco (Meyer and Scholten, 2007; Ingouff et al., 2010; Zhao et al., 2011; Nodine and Bartel, 2012). However, a relationship between a precise karyogamic stage and the start of de novo gene expression in the zygote was not clearly presented. We recently conducted
transcriptome analyses using isolated rice gametes and zygotes prepared from ovaries 2 to 4 h after pollination, and globally identified genes that were up- or down-regulated after fertilization (Abiko et al., 2013b). Therefore, in the study, the precise timing of the expression of several genes, which were known to be up-regulated after fertilization in rice zygotes, was monitored during karyogamy.

Zygotes produced by the fusion of sperm cells expressing H2B-GFP and wild-type egg cells were prepared and incubated. Then, zygotes that were in karyogamic stages III to IV, V to VI, and VII to VIII were selected for subsequent RT-PCR analyses. For the Os08g0562800 and Os01g0300000 genes, detectable expression levels occurred only at stages VII and VIII (Fig. 9). This indicates that the \textit{de novo} expression of these genes in rice zygotes starts between stage VII and VIII, at which time the male chromatin in the fused nucleus has mostly decondensed, but there was not \textit{de novo} gene expression during stages V and VI, at which time the initial decondensation of male chromatin occurs. For the Os07g0182900 gene, the transcript was detected from stages III/IV and V/VI, and the expression level increased during stages VII and VIII. The initial transcript detected at stages III to IV, approximate 10-20 min after fusion, will be derived from the paternal transcript, since the gene is not expressed in egg cell before fusion (Fig. 9) and it was reported that Os07g0182900 gene is highly expressed in sperm cells (Abiko et al., 2013b). Its increased expression level detected at stages VII and VIII will be a result of \textit{de novo} zygotic gene expression. As for the Os01g0840300 and Os10g0580900 genes, no expression was detected, suggesting that the expression of these genes occurs in zygotes after the completion of karyogamy (stage VIII).

The present study indicated that \textit{de novo} zygotic gene expression starts in rice zygotes at
stage VII to VIII of karyogamy, during which sperm chromatin has mostly decondensed in the fusing nucleus, and that paternally supplied transcripts were also detected in the zygote at stages III and IV of karyogamy before de novo expression started. In Arabidopsis, the contribution of such paternal mRNA for zygotic development is well known. After fertilization, vacuoles in Arabidopsis zygotes fragment, and the zygote elongates two- to three-fold before a large vacuole is re-assembled (Mansfield and Briarty, 1991; Faure et al., 2002). The polarized zygote divides asymmetrically into a two-celled proembryo consisting of an apical and a basal cell, which develop into the embryo proper and the suspensor/hypophysis, respectively, suggesting that cellular polarity in the zygote is tightly linked to the establishment of the initial apical-basal axis in plants (Jeong et al., 2011, Zhang and Laux 2011; Ueda and Laux 2012). Bayer et al. (2009) indicated that transcripts of SHORT SUSPENSOR (SSP), encoding a member of the Pelle/IRAK/receptor-like superfamily, are produced in sperm cells and delivered to the zygotes. Then, SSP proteins accumulate and activate the YODA-dependent MAPKKK signaling pathway that is thought to be crucial for the cell elongation of the zygote and the specified cell fate of the basal cell of Arabidopsis two-celled proembryo (Lukowitz et al., 2004). In addition, it has been reported that WRKY2 transcription factor expressing in sperm cell as well as egg cell functions in establishment of early body plan in Arabidopsis zygote (Ueda et al. 2011). In the study, transcripts from the Os07g0182900 gene encoding cytosine-5 DNA methyltransferase 1 (MET1) detected at early karyogamy stage were considered as paternally supplied mRNA. Interestingly, it has been reported that the establishment of polarity or asymmetric cell division of rice zygotes is partly affected by treatment of zygotes with a specific inhibitor of MET1, RG108 (Abiko et al., 2013b). This could be another example of sperm mRNA involvement in
zygotic development in angiosperms. Transcriptome analyses of rice zygotes at stages III to IV of karyogamy will result in the identification of genes whose transcripts are delivered from sperm cells and function in early zygotic development.

In rice zygote produced in vitro, de novo zygotic gene expression occurs during or after nuclear fusion in accordance with male chromatin decondensation in fused nucleus. However, in latrunculin B-treated zygotes, which is produced by fusion of a wild-type egg cell with a sperm cell expressing H2B-GFP, H2B-GFP signal became detectable in an egg nucleus as well as a sperm nucleus even if sperm nucleus did not fuse with egg nucleus (Fig. 4, K and L). This may indicate that egg activation and subsequent de novo gene expression progress in such zygote without completion of karyogamy. Using a Ca$^{2+}$-mediated IVF system with maize gametes, Antoine et al. (2000, 2001) indicated that gamete fusion induced egg cell activation, including rapid cell wall formation (Kranz et al., 1995), via a transitory increase of the Ca$^{2+}$ levels in the fertilized egg cells. H2B-GFP expression in karyogamy-defected zygotes may suggest that the egg activation can be triggered by fusion stimuli and subsequent deliver of sperm cell cytoplasm into fertilized egg, and that initial zygotic events, including transcriptional/translational initiation, histone H3 replacement, and chromatin remodeling, toward de novo gene expression occur in each unfused male and female nucleus without karyogamy.

MATERIALS AND METHODS

Plant materials, isolation of gametes and zygotes, and the electro-fusion of gametes
Oryza sativa L. cv. Nipponbare was grown in environmental chambers (K30-7248; Koito Industries, Yokohama, Japan) at 26°C with a 13/11-h light/dark cycle. The isolation of egg cells and sperm cells from rice flowers, and the electro-fusion of isolated gametes for zygote production, were conducted as described by Uchiumi et al. (2006, 2007).

Vector construction and preparation of transformants

Transformed rice expressing the histone H2B-GFP fusion protein was prepared as previously described (Abiko et al., 2013b). The plasmid vector, pENTRTM3C (Life Technologies, Carlsbad, CA, USA), harboring Arabidopsis DD45 promoter::H2B-TagRFP was generously gifted from Dr. T. Igawa, Chiba University, Japan (Igawa et al., 2013). After the expression cassette was transferred to the destination vector pGWB1 (Nakagawa et al., 2007) by the LR reaction using the Gateway LR Clonase enzyme mix (Invitrogen, Carlsbad, CA, USA), the vector was used as the DD45 promoter::H2B-RFP vector to transform rice plants. For actin labeling using Lifeact (Era et al., 2009), pGWB1 binary vector, harboring Arabidopsis DD45 promoter::Lifeact-TagRFP was generously gifted from Dr. T. Igawa, Chiba University, Japan, and used for rice transformation as below.

The promoter region of an ubiquitin gene (Os02g0161900) was subcloned into the HindIII and XbaI sites of pIG121-Hm, a binary plasmid vector harboring a GUS coding sequence, and the subsequent plasmid was used as Ubi promoter::GUS vector according to Abiko et al. (2013b). The pGWB405 vector harboring a DNA fragment encoding the GFP-tagged Arabidopsis SUN2 (At5g22880) was generously gifted from Dr. K. Tamura, Kyoto University, Japan (Tamura et al., 2013). This vector was used as the template for PCR amplification of the
DNA region encoding the H2B-GFP fusion protein using the specific primers 5′-GCTAGCATGCGGCGTCAACGCGT-3′ and 5′-GAGCTCTTACTTGTACAGCCTGCTCCA-3′. The amplified PCR product was subcloned into the pGEM-T Easy vector (Promega, Fitchburg, WI, USA), and the plasmid possessing the SUN2-GFP sequence was cut with *Nhe*I and *Sac*I. Then, the excised DNA fragment was subcloned into the *Xba*I–*Sac*I site of *Ubi* promoter:*GUS* vector, replacing the GUS sequence with SUN2-GFP. The vector was used as the *Ubi* promoter:*SUN2-GFP* vector to transform rice plants.

*Agrobacterium tumefaciens* LBA4404 was transformed with the above mentioned vectors, and transgenic rice plants were prepared by co-cultivation of scutellum tissue with *A. tumefaciens* according to Toki et al. (2006).

**Microscopic observation, 3D reconstruction and volume measurements of sperm nuclei**

Zygotes produced by IVF of isolated rice gametes were transferred into droplets of mannitol adjusted to 450 mOsmol kg⁻¹ H₂O. Gametes/zygotes expressing H2B-GFP or SUN2-GFP were observed with a BX-71 inverted fluorescence microscope (Olympus, Tokyo, Japan) with 460–490-nm excitation and 510–550-nm emission wavelengths (*U*-MWIBA2 mirror unit; Olympus). For H2B-TagRFP proteins in gametes/zygotes, observations were conducted with 520–550-nm excitation and >580-nm emission wavelengths (*U*-MWIG mirror unit; Olympus). Digital images of gametes, zygotes, and their resulting embryos were obtained through a cooled charge-coupled device camera (Penguin 600CL; Pixcera, Los Gatos, CA, USA) and InStudio software (Pixcera). In addition to BX-71 inverted fluorescence microscope, zygotes were
observed with BZ9000 microscope (Keyence, Osaka, Japan) with 440-470-nm excitation and 535-550-nm emission wavelengths (OP-66836 GFP-BP filter set; Keyence) for H2B-GFP proteins and 525–540-nm excitation and 605-655-nm emission wavelengths (OP-66837 TRITC filter set; Keyence) for H2B-RFP and Lifeact-tagRFP proteins. IVF-produced zygotes were observed also with an LSM 710 CLS microscope (Carl Zeiss, Jena, Germany) with 488-nm excitation and 505–530-nm emission wavelengths for H2B-GFP and SUN2-GFP, and with 543-nm excitation and >560- nm emission wavelengths for H2B-TagRFP.

The 3D structure of the sperm nucleus in the zygote was constructed using IMARIS software (Bitplane, Zürich, Switzerland) from a z-series of CLS images in which sperm and egg nuclei were labeled with SUN2-GFP and H2B-TagRFP, respectively. The volume of the sperm nucleus was measured by surface rendering using the Surface function of IMARIS.

**Inhibitor treatment of egg cells and zygotes**

Latrunculin B (Wako, Osaka, Japan) and oryzalin (Wako) were dissolved with methanol into stock solutions of 2 and 100 mM, respectively. Egg cells were incubated in droplets of mannitol solution adjusted to 370 mOsmol kg\(^{-1}\) H\(_2\)O containing latrunculin B (200 nM and 2 μM) or oryzalin (10 and 100 μM) for 1.5 h. The egg cells were used for IVF with sperm cells, and the resultant zygotes were transferred into droplets of mannitol adjusted to 450 mOsmol kg\(^{-1}\) H\(_2\)O containing the inhibitors for microscopic observation.

**RT-PCR**

Zygotes were produced by electro-fusion of sperm cells expressing H2B-GFP with wild-type
egg cells and incubated in droplets of mannitol solution adjusted to 450 mOsmol kg⁻¹ H₂O. The progression of karyogamy was observed by fluorescence microscopy as mentioned above, and the zygotes, at appropriate karyogamic stages, were selected and washed three times by transferring the cells into fresh droplets of mannitol solution on coverslips. The washed cells were submerged in 5 μl of the extraction buffer supplied in a PicoPure RNA Isolation Kit (Life Technologies) and stored at −80°C until use. cDNAs were synthesized from total RNAs of 10 to 12 zygotes using the High Capacity RNA-to-cDNA™ Kit (Life Technologies) according to the manufacturer’s instructions. For RT-PCR, 0.3 μl of first-strand cDNA was used as the template in a 20 μl PCR reaction with 0.3 μM of primers using KOD-FX DNA polymerase (Toyobo, Osaka, Japan) as follows: 45 cycles of 94°C for 1 min, 55°C for 30 s, and 72°C for 1 min. The expression of the ubiquitin gene (Os02g0161900) was monitored as an internal control. Primer sequences used for PCR analyses are listed in Table S1.

Supplemental Data

Supplemental Table S1. Nucleotide sequences of PCR primers.

Supplemental Figure S1. Mitotic division of an isolated rice zygote expressing histone H2B-GFP.

Supplemental Figure S2. Karyogamy in rice zygotes produced by in vitro fertilization.

Supplemental Figure S3. Karyogamy progression and de novo gene expression.

Supplemental Figure S4. Membrane fusion between rice sperm and egg nuclei.

Supplemental Figure S5. Effect of latrunculin B on the actin structure in rice egg cells.
**Supplemental Figure S6.** Male and female nuclei in zygote without karyogamy.

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We thank Ms. H. Maeda for help in preparing transformed plants and Ms. T. Mochizuki (both of Tokyo Metropolitan University) for isolating rice egg cells, Dr. T. Igawa (Chiba University) for the *DD45* promoter::*H2B-TagRFP* in pENTR™3C and the *DD45* promoter::*Lifeact-TagRFP* in pGWB1, Dr. K. Tamura (Kyoto University) for the *GFP–SUN2* in pGWB405, Dr. T. Nakagawa (Shimane University) for the pGWB1 and pGWB405 vectors, and the RIKEN Bio Resource Center (Tsukuba, Japan) for providing cultured rice cells (Oc line).
LITERATURE CITED


Zhao J, Xin H, Qu L, Ning J, Peng X, Yan T, Ma L, Li S, Sun MX (2011) Dynamic changes of transcript profiles after fertilization are associated with de novo transcription and maternal elimination in tobacco zygote, and mark the onset of the maternal-to-zygotic transition. Plant J 65: 131–145
FIGURE LEGENDS

Figure 1. Rice gametes expressing the histone H2B-GFP, H2B-RFP or SUN2-GFP fusion protein. A and B, An egg cell isolated from transgenic rice expressing histone H2B-GFP under control of the ubiquitin promoter visualized with bright-field (A) and fluorescence (B) microscopy. The arrowheads indicate nucleoli. C and D, An egg cell isolated from transgenic rice expressing histone H2B-RFP under control of the DD45 promoter visualized with bright-field (C) and fluorescence (D) microscopy. The arrowheads indicate nucleoli. E and F, A pollen grain expressing histone H2B-GFP under control of the ubiquitin promoter visualized with bright-field (E) and fluorescence (F) microscopy. Arrowheads and the arrow indicate sperm and vegetative nuclei, respectively. G and H, A pollen grain expressing H2B-GFP releasing its content in mannitol solution visualized with bright-field (G) and fluorescence (H) microscopy. Arrowheads indicate sperm cells and the arrow indicates a possible vegetative nucleus. I–K, An egg cell expressing SUN2-GFP under control of the ubiquitin promoter visualized with fluorescence (I) and bright-field (J) microscopy. Panel K contains the merged images of I and J. L–N, A pollen grain expressing SUN2-GFP releasing its content in mannitol solution (L). Two released sperm cells enclosed within the square in panel L were visualized with fluorescence microscopy (M). Panel N contains the merged bright-field and fluorescent images. Bars = 20 μm.

Figure 2. Karyogamy in rice zygotes. A and B, Alignment of a sperm cell expressing H2B-GFP with a wild-type egg cell on one of the electrodes under an alternating current field in a fusion
droplet. C–R, Gametes in panel B were electrically fused and the resulting fused cell (zygote) was observed at the time points indicated. The upper panels are fluorescent images, and the lower panels are merged bright-field and fluorescent images. Black and white arrowheads indicate sperm cells expressing histone H2B-GFP and nucleoli, respectively. Bar = 20 μm.

**Figure 3.** Time table of sperm chromatin decondensation in the fused nuclei of 50 independent rice zygotes. Zygotes were produced by *in vitro* fusion of a wild-type egg cell with a sperm cell expressing a histone H2B-GFP fusion protein, and the fluorescence signal was observed every 15 to 20 min after fusion. Y-axis indicates the number of zygote in which initial decondensation of sperm chromatin was observed.

**Figure 4.** Effects of latrunculin B and oryzalin on karyogamy in fused rice gametes. Egg cells were treated with inhibitors and used for electro-fusion with sperm cells expressing H2B-GFP. A–F, Egg cells and fused cells that were treated with 200 nM latrunculin B. G–L, Egg cells and fused cells that were treated with 2 μM latrunculin B. M–R, Egg cells and fused cells that were treated with 100 μM oryzalin. After *in vitro* fertilization, the fused cell were observed at 10 to 15 min (A, B, G, H, M, N), 80 to 120 min (C, D, I, J, O, P) and 10 to 12 h (E, F, K, L, Q, R) after fusion. In zygotes treated with 2 μM latrunculin B, the decondensation of sperm chromatin was not observed for 2 h after fusion (I, J), and the signal from H2B-GFP was detected both in sperm and egg nuclei at 12 h after fusion. Bar = 20 μm.

**Figure 5.** Actin organization and sperm nucleus migration. Egg cells expressing Lifeact-tagRFP
treated with or without 2 μM latrunculin B were fused with sperm cells expressing H2B-GFP, and resultant zygotes were observed at the time point indicated. An arrow indicates nucleus surrounded by actin filaments. Bars = 20 μm.

**Figure 6.** Chromatin state during or after nuclear fusion in rice zygotes. Zygotes produced by electro-fusion of a sperm cell expressing histone H2B-GFP and an egg cell expressing H2B-RFP were observed using a confocal laser scanning (CLS) microscope. The fluorescent signals from histone H2B-GFP and H2B-RFP are presented in the upper and middle panels, respectively, while the lower panels are merged images. Bar = 10 μm.

**Figure 7.** Changes over time in the volume of a rice sperm nucleus during contact with an egg nucleus. Zygotes produced by electro-fusion of sperm cells expressing SUN2-GFP and an egg cell expressing H2B-RFP were observed using a CLS microscope. Panels A, C, E, G and I are CLS images, and panels B, D, F, H and J are 3D structure images. An arrowhead indicates the possible fusion point between the nuclear membranes. Arrows indicate the fusing area between sperm and egg nuclear membranes. Numbers in panels of 3D images represent the volume of the sperm nucleus. Bars = 10 μm. K, Changes in the volume of the sperm nucleus before (panels A and B) or after egg chromatin incursion (panels G and H). The data are mean ±SD of three sperm nuclei.

**Figure 8.** A schematic diagram of the progression of karyogamy in rice zygotes. After gamete fusion, the sperm nucleus migrates inside of the fused cell to appose the egg nucleus via
possible actin filament-dependent machinery (stages I–III). Through a possible inter-nuclear connection, female chromatin moves into the sperm nucleus, but sperm chromatin remains tightly packed (stage IV). Male chromatin begins decondensing (Stage V), and male chromatin further uniformly distributes in accordance with the progression of nuclear fusion (Stages VI and VII). The karyogamic event is finally completed at stage VIII, and thereafter, the zygote develops into an embryo. Light blue and pink circles indicate sperm and egg nuclei/chromatin, respectively. Yellow indicates the merged sperm and egg chromatin.

**Figure 9.** Expression patterns of five genes putatively induced after fertilization in rice zygotes. Semiquantitative RT-PCR was performed on total RNAs isolated from 10-12 zygotes at the appropriate karyogamic stages. Ubiquitin mRNA was used as an internal control. See Table S1 for primer sequences.

**Figure S1.** Mitotic division of an isolated rice zygote expressing H2B-GFP. Zygotes were isolated from pollinated flower of transformed rice expressing H2B-GFP as described in Abiko et al. (2013b). The isolated zygotes were cultured and observed as described in Materials and Methods.

**Figure S2.** Karyogamy in rice zygotes produced by in vitro fertilization. A–L, Karyogamy in rice zygotes produced by fusion of wild-type egg cells with sperm cells expressing histone H2B-GFP. M–X, Karyogamy in rice zygotes produced by fusion of egg cells expressing histone
H2B-GFP with wild-type sperm cells. The upper panels are fluorescence images, and the lower panels are merged bright-field and fluorescence images. Arrowheads indicate sperm nuclei fusing with egg nuclei. Bars = 20 μm.

**Figure S3.** Karyogamy progression and *de novo* gene expression in zygotes. Rice zygotes produced by fusion of wild-type egg cells with sperm cells expressing H2B-GFP were observed using a BX-71 inverted fluorescence microscope with 460–490-nm excitation and 510–550-nm emission wavelengths (U-MWIBA2 mirror unit; Olympus). All fluorescent images were captured with same exposure time (1/1.8 sec), and the images were processed with same methods and parameters for comparison of fluorescence intensity. The upper panels are fluorescence images, and the lower panels are merged bright-field and fluorescence images. Although sperm nucleus was visible before nuclear fusion (A-D), fluorescent signal was hardly detected during or just after sperm chromatin decondensation by observation using same exposure time (E, F). Thereafter, the nuclear signal intensity became progressively strong according to the progression of karyogamy (G-L). Bar = 20 μm.

**Figure S4.** Effect of latrunculin B on the actin structure in rice egg cells. Egg cells were isolated from the transformed rice plants expressing Lifeact-TagRFP under the DD45 promoter, and incubated in droplets of mannitol solution adjusted to 370 mOsmol kg⁻¹ H₂O containing latrunculin B (200 nM and 2 μM). At the indicated time, cells were observed with a BX-71 inverted fluorescence microscope with 520–550-nm excitation and >580-nm emission wavelengths (U-MWIG mirror unit; Olympus).
Figure S5. Membrane fusion between rice sperm and egg nuclei. An egg cell and a sperm cell expressing SUN2-GFP were fused. Then, the zygotes was observed with a CLS microscope, and z-series serial images were obtained. A, 3D reconstructed image of nuclei in zygote at 98 min after fusion. Panels A1–A5 represent z-series serial images. B. 3D reconstructed image of the nuclei in a zygote at 103 min after fusion. Panels B1–B5 represent z-series serial images. At the possible fused area, the SUN2-GFP signal was not detected. C. 3D reconstructed image of the nuclei in a zygote at 110 min after fusion. Panels C1–C5 represent z-series serial images. Progression of membrane fusion was observed. Bars = 10 μm.

Figure S6. Co-localization of sperm and egg nuclei in a zygote without karyogamy. Egg cells were isolated from the transformed rice plants expressing H2B-RFP and treated with 2 μM latrunculin B. Then, the egg cells were fused with sperm cell expressing H2B-GFP, and the fused cells were observed at the time point indicated with a BZ9000 microscope with 440-470-nm excitation and 535-550-nm emission wavelengths (OP-66836 GFP-BP filter set, Keyence) for H2B-GFP, and 525–540-nm excitation and 605-655-nm emission wavelengths (OP-66837 TRITC filter set, Keyence) for H2B-RFP. Panels A, E, I and M; H2B-GFP signals, Panels B, F, J and N; H2B-RFP signals, Panels C, G, K and N; merged signal of H2B-GFP and H2B-RFP, Panels D, H, L and P; bright-field images. Bar = 20 μm.
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