

Title	Spermatozoa lacking Fertilization Influencing Membrane Protein (FIMP) fail to fuse with oocytes in mice
Author(s)	Fujihara, Yoshitaka; Lu, Yonggang; Noda, Taichi et al.
Citation	Proceedings of the National Academy of Sciences of the United States of America. 2020, 117(17), p. 9393-9400
Version Type	VoR
URL	https://hdl.handle.net/11094/78547
rights	© 2020 the Author(s). Published by PNAS. This article is licensed under a Creative Commons Attribution 4.0 International License.
Note	

## The University of Osaka Institutional Knowledge Archive : OUKA

https://ir.library.osaka-u.ac.jp/

The University of Osaka

# Spermatozoa lacking Fertilization Influencing Membrane Protein (FIMP) fail to fuse with oocytes in mice

Yoshitaka Fujihara<sup>a,b,c</sup>, Yonggang Lu<sup>a</sup>, Taichi Noda<sup>a</sup>, Asami Oji<sup>a,d</sup>, Tamara Larasati<sup>a,e</sup>, Kanako Kojima-Kita<sup>a,e</sup>, Zhifeng Yu<sup>b</sup>, Ryan M. Matzuk<sup>b</sup>, Martin M. Matzuk<sup>b,1</sup>, and Masahito Ikawa<sup>a,e,f,1</sup>

<sup>a</sup>Research Institute for Microbial Diseases, Osaka University, Suita, 565-0871 Osaka, Japan; <sup>b</sup>Center for Drug Discovery and Department of Pathology & Immunology, Baylor College of Medicine, Houston, TX 77030; <sup>c</sup>Department of Bioscience and Genetics, National Cerebral and Cardiovascular Center, Suita, 564-8565 Osaka, Japan; <sup>d</sup>Laboratory for Developmental Epigenetics, RIKEN Center for Biosystems Dynamics Research, Kobe, 650-0047 Hyogo, Japan; <sup>e</sup>Graduate School of Medicine, Osaka University, Suita, 565-0871 Osaka, Japan; and <sup>f</sup>The Institute of Medical Science, The University of Tokyo, Minato-ku, 108-8639 Tokyo, Japan

Contributed by Martin M. Matzuk, February 18, 2020 (sent for review October 1, 2019; reviewed by Jean-Ju L. Chung and Janice Evans)

Sperm-oocyte fusion is a critical event in mammalian fertilization, categorized by three indispensable proteins. Sperm membrane protein IZUMO1 and its counterpart oocyte membrane protein JUNO make a protein complex allowing sperm to interact with the oocyte, and subsequent sperm-oocyte fusion. Oocyte tetraspanin protein CD9 also contributes to sperm-oocyte fusion. However, the fusion process cannot be explained solely by these three essential factors. In this study, we focused on analyzing a testisspecific gene 4930451111Rik and generated mutant mice using the CRISPR/Cas9 system. Although IZUMO1 remained in 4930451111Rik knockout (KO) spermatozoa, the KO spermatozoa were unable to fuse with oocytes and the KO males were severely subfertile. 4930451111Rik encodes two isoforms: a transmembrane (TM) form and a secreted form. Both CRISPR/Cas9-mediated TM deletion and transgenic (Tg) rescue with the TM form revealed that only the TM form plays a critical role in sperm-oocyte fusion. Thus, we renamed this TM form Fertilization Influencing Membrane Protein (FIMP). The mCherry-tagged FIMP TM form was localized to the sperm equatorial segment where the sperm-oocyte fusion event occurs. Thus, FIMP is a sperm-specific transmembrane protein that is necessary for the sperm-oocyte fusion process.

CRISPR/Cas9 | fertilization | infertility | IZUMO1 | transgenic

ammalian fertilization comprises various steps including sperm migration through the female reproductive tract, changes in sperm physiology and morphology, sperm-egg interaction, and fusion (1, 2). Recent gene KO mouse studies revealed essential factors for fertilization (3-5). However, to date, only three proteins have been shown to play an essential role in sperm-oocyte fusion, with all three expressed on the cell surface. CD9 and JUNO modulate female fertility, while IZUMO1 regulates fusion in spermatozoa. CD9 consists of a group of tetraspanin membrane proteins and is expressed ubiquitously. However, while Cd9 KO females appeared healthy, their fertility was significantly reduced (6-8). Cd9 KO oocytes show impaired development of microvilli on the oocyte membrane and are unable to fuse with spermatozoa (9). IZUMO1 is a type I transmembrane protein localized on the sperm acrosomal membrane. One of the physiological changes of spermatozoa during the acrosome reaction is the presentation of IZUMO1 onto sperm surface. IZUMO1 translocates from the acrosomal membrane to the plasma membrane and spreads over the equatorial and postacrosomal regions. As such, IZUMO1 is used as a marker of the acrosome reaction (10, 11). Izumo1 KO male mice are sterile because spermatozoa lack the ability to fuse with the oocyte plasma membrane (12). Although the cytoplasmic tail of IZUMO1 is dispensable for fertility, the N-terminal region (5th through 113th amino acids) is found to be essential for sperm-oocyte fusion (13, 14). GPI-anchored protein JUNO

(officially named IZUMO1R) was identified as the oocyte receptor of IZUMO1, and *Juno* KO female mice were also infertile due to impaired sperm-oocyte fusion (15). Further structural analyses found that several amino acid residues of each protein are critical for the direct binding of JUNO and IZUMO1 in mice and humans (16–20). However, IZUMO1–JUNO interaction is not enough to trigger membrane fusion (21). Therefore, additional factors must contribute to the molecular mechanisms of sperm-oocyte fusion.

KO mouse studies have identified many essential genes for fertility by conventional gene-targeting approaches (1, 5). However, the ES cell-mediated gene targeting and subsequent chimeric mice production are laborious and costly. In 2013, the CRISPR/Cas9-mediated genome editing system emerged, enabling large-scale screening using KO mice (22, 23). In fact, while we have reported 96 reproductive tract enriched genes that are individually dispensable (24–27), we have also reported 18 genes and 2 gene clusters that are essential for male fertility (12, 28–43) CRISPR/Cas9 technologies have allowed us not only to disrupt a gene but also to analyze important regions of a gene of interest. One can easily delete protein domains or introduce a point mutation at the nucleotide level (31, 44).

### **Significance**

As the human body is composed of 60 trillion cells that originate from a fertilized egg, sperm—oocyte fusion is the initial event of our life. Few sperm—oocyte fusion factors have been unveiled to date, and only IZUMO1 has been identified as a sperm-specific fusion-mediating protein. Here, we identified the testis-specific 4930451111Rik gene important for male fertility, playing a role in sperm—oocyte fusion during fertilization. Based on its functional role, we renamed this gene fertilization influencing membrane protein (Fimp). We discovered a factor responsible for sperm—oocyte fusion in mammals, and this knowledge could be used to develop in vitro and in vivo infertility treatments as well as male contraceptives.

Author contributions: Y.F., M.M.M., and M.I. designed research; Y.F., Y.L., T.N., A.O., T.L., K.K.-K., Z.Y., and R.M.M. performed research; Y.F., Y.L., Z.Y., M.M.M., and M.I. analyzed data; and Y.F., M.M.M., and M.I. wrote the paper.

Reviewers: J.-J.L.C., Yale University; and J.E., Purdue University.

The authors declare no competing interest.

This open access article is distributed under Creative Commons Attribution License 4.0 (CC RY)

<sup>1</sup>To whom correspondence may be addressed. Email: mmatzuk@bcm.edu or ikawa@biken.osaka-u.ac.jp.

This article contains supporting information online at https://www.pnas.org/lookup/suppl/doi:10.1073/pnas.1917060117/-/DCSupplemental.

First published April 15, 2020.

Here, we focused on analyzing a testis-specific gene 4930451111Rik that is localized on mouse chromosome 7 and that is conserved in humans and other mammals. To examine the physiological role of this gene, we generated CRISPR/Cas9-mediated KO mice. The 4930451111Rik KO male mice were severely subfertile as a result of impaired sperm-oocyte fusion, even though IZUMO1 remained intact. Moreover, we found that 4930451I11Rik is expressed as two isoforms (transmembrane [TM] and secreted forms) in the testis due to alternative splicing. We generated 4930451I11Rik TM-deleted mice using the CRISPR/Cas9 system. The 4930451I11Rik TM-deleted mice demonstrated that the TM form is critical for spermatozoa to fuse with oocytes. Thus, we renamed 4930451I11Rik as fertilization influencing membrane protein (FIMP). In addition, the TM form of FimpmCherry Tg mice restored male infertility in Fimp KO mice and was detected on the sperm equatorial segment where the sperm-oocyte fusion event occurs. Thus, we have discovered FIMP as the second sperm factor that is necessary for spermatozoa to fuse with oocytes.

#### Results

Testis-Specific Expression of 4930451111Rik in Mice and Humans. Mouse 4930451111Rik gene is localized to chromosome 7. The expression of this gene in various mouse organs was examined by RT-PCR, and it showed testis-specific expression (Fig. 14). Next, the onset of 4930451111Rik expression in testes was examined by RT-PCR. This gene was expressed 20 d after birth (Fig. 1B). Furthermore, the human ortholog C16ORF92 showed testisspecific expression as well (Fig. 1C). Since we detected two bands (312 bp and 400 bp) of 4930451111Rik from the mouse testes (Fig. 1 A and B), we confirmed their identities using direct sequencing analysis (SI Appendix, Fig. S1A). The 4930451111Rik consists of four exons and is expressed in two variants from three or four exons, respectively. The difference between these two variants is whether the third exon encoding the transmembrane domain is present or not (Fig. 1D). We found that mouse 4930451111Rik has two isoforms, TM (+) and secreted (TM [-]; Fig. 1E). Protein structure analysis predicted that the TM form of 4930451I11Rik is a type I single-pass transmembrane protein containing a cleavable N-terminal signal peptide and a transmembrane-encoding sequence (SI Appendix, Fig. S1 B and C). Moreover, the protein sequence of 4930451I11Rik is highly conserved in mammals; the gene has been observed in more than 70 eutherian mammals including marsupials (wallaby; SI Appendix, Fig. S1D; http://www.informatics.jax.org/homology/ 52331). These data indicate that mouse 4930451111Rik and human ortholog C16ORF92 are both testis-enriched genes. While mouse 4930451I11Rik is expressed in two forms, TM and secreted, human C16ORF92 is only expressed as the TM form.

Male Fertility of 4930451111Rik KO Mice. To analyze the physiological role of 4930451I11Rik, we generated 4930451I11Rik KO mice by CRISPR/Cas9. A 1-bp deletion in the second exon of 4930451111Rik was confirmed by PCR and direct sequencing analysis (SI Appendix, Fig. S2A). The 1-bp deletion caused a frameshift mutation that resulted in a premature termination codon at the 43rd amino acid of 4930451I11Rik (TM [+]: 111 amino acids and TM [-]: 92 amino acids in wild-type mice; SI Appendix, Fig. S2B). Thus, both isoforms were disrupted by the 1-bp deletion. The KO (-1/-1) mice had no overt developmental abnormalities. Furthermore, no deleterious effects on testicular histology and sperm morphology were observed (SI Appendix, Fig. S2 C and D). To examine male fertility, adult KO males were mated with wild-type females for several months. The KO males were severely subfertile, showing normal mating behavior with successful ejaculation and vaginal plug formation. The average number of pups per plug was  $8.5 \pm 1.7$  in wild-type males (15 plugs) and  $0.5 \pm 0.8$  in the KO males (10 plugs;

Fig. 1F). These data indicated that 4930451111Rik is required for normal male fertility in mice.

In Vitro Fertilizing Ability in 4930451111Rik KO Spermatozoa. To analyze the cause of infertility in 4930451111Rik KO males, we performed in vitro fertilization (IVF) assays using cauda epididymal spermatozoa from 4930451111Rik KO males with cumulus-intact oocytes. 4930451111Rik KO spermatozoa could penetrate the zona pellucida (ZP), but rarely fertilize oocytes;  $91.5 \pm 9.3\%$  (397/434) fertilized eggs were observed when oocytes were treated with wild-type spermatozoa, whereas  $1.8 \pm$ 2.2% (7/384) fertilized eggs were observed with KO spermatozoa (Fig. 2A). While we were able to observe the pronuclei of control eggs 7 h after insemination, 4930451111Rik KO spermatozoa remained in the perivitelline space (Fig. 2B). Next, we measured the fusion ability of the spermatozoa mixed with ZP-free oocytes in vitro. 4930451111Rik KO spermatozoa had a significantly reduced ability to fuse with the oocyte membrane compared with wild-type (3.3  $\pm$  1.6 and 0.03  $\pm$  0.1 spermatozoa per egg using wild-type and KO spermatozoa, respectively; P < 0.01; Fig. 2 C and D). The KO spermatozoa showed impaired ability to fuse with the oocytes.

To examine the effects of 4930451I11Rik disruption on IZUMO1, we performed an immunoblot analysis of IZUMO1 in 4930451111Rik KO spermatozoa. There were no significant differences in the size or amount of IZUMO1 between wild-type and 4930451111Rik KO spermatozoa (Fig. 2E). Next, we observed IZUMO1 localization by immunostaining analysis. IZUMO1 changes its localization from the acrosomal cap to the entire head during the acrosome reaction, which is considered one of the most essential steps for fertilization. When we observed the cauda epididymal spermatozoa 4 h after incubation, there was no difference in the IZUMO1 staining pattern in 4930451111Rik KO spermatozoa and in wild-type (Fig. 2F). These results indicate that IZUMO1 remains and correctly translocates in 4930451111Rik KO spermatozoa.

Analysis of 4930451111Rik TM-Deleted Mice. Our data indicate that the 4930451I11Rik is an important factor for the sperm-oocyte fusion process. As shown in Fig. 1, 4930451I11Rik is expressed as two isoforms in the mouse testis: the TM form and the secreted form. Thus, to determine which isoform is essential, we deleted the third exon that encodes a TM region by the CRISPR/Cas9 system (Fig. 3A). A 246-bp deletion around the third exon of 4930451111Rik was confirmed by PCR, direct sequencing (Fig. 3B and SI Appendix, Fig. S3A), and RT-PCR analysis (Fig. 3C and SI Appendix, Fig. S3B). The 246-bp deletion (including 88 bp of the third exon and 12 bp of fourth exon) caused an in-frame mutation that appeared from a premature terminal codon at the 85th amino acid of 4930451I11Rik (Fig. 3D). We verified that only the TM form of 4930451I11Rik disappeared from the testis in 4930451I11Rik TM-deleted mice (Fig. 3C). We performed IVF assays to check the fertilizing ability of 4930451I11Rik TM-deleted mouse spermatozoa. Similar to the KO (-1/-1) mice in which both secreted and TM forms were disrupted, 4930451I11Rik TM-deleted spermatozoa could rarely fertilize oocytes (0.6  $\pm$  1.3%; 1 fertilized egg detected out of 176 oocytes/embryos analyzed) in contrast to wild-type spermatozoa, in which nearly all oocytes were fertilized (97.1  $\pm$  2.5%; 168 eggs/ 173 oocytes/embryos analyzed; Fig. 3E). 4930451I11Rik TMdeleted spermatozoa also had a significantly reduced ability to fuse with the oocyte membrane compared with the control (2.0  $\pm$ 1.0 and  $0.1 \pm 0.2$  spermatozoa per egg in the wild-type and TMdeleted spermatozoa, respectively; P < 0.01; Fig. 3F). Moreover, IZUMO1 was present and localized properly in the TM-deleted mice, as well as wild-type mice (SI Appendix, Fig. S3 C and D). To signify this newly discovered sperm-oocyte fusion-mediating factor, we renamed 4930451I11Rik to fertilization influencing

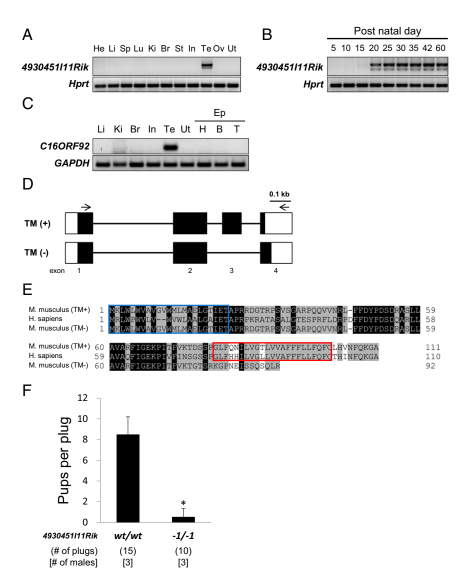


Fig. 1. Characterization of 4930451111Rik in mice and humans and male fertility in 4930451111Rik KO mice. (A) Testis-specific expression of mouse 4930451111Rik by multitissue RT-PCR analysis. The expression of each gene was examined by RT-PCR, using RNA isolated from various organs. 4930451111Rik was detected only in the mouse testis. The Hprt gene was used as an expression control. He, heart; Li, liver; Sp, spleen; Lu, lung; Ki, kidney; Br, brain; St, stomach; In, intestine; Te, testis; Ov, ovary; Ut, uterus. (B) RT-PCR analysis of mouse 4930451111Rik in mouse testis. 4930451111Rik was expressed 20 d after birth. (C) RT-PCR analysis of human C160RF92 (ortholog of mouse 4930451111Rik) in human tissues. Human C160RF92 was also detected only in the testis. The GAPDH gene was used as an expression control. Ep, epididymis; H, head (caput epididymis); B, body (corpus epididymis); T, tail (cauda epididymis). (D) Two isoforms, TM form and secreted form, of mouse 4930451111Rik gene. The 4930451111Rik mRNA consists of four exons. Although the TM (+) form contains all four exons, the TM (-) form expresses only three exons, as the third exon is spliced out. Exons are indicated by boxes. Protein-coding and noncoding regions are indicated in black and white, respectively. The sequences of the two isoforms are shown in SI Appendix, Fig. S1A. Arrows indicate primers used for RT-PCR analysis. (E) Amino acid sequence similarity of mouse 4930451111Rik and human C160RF92. The length of TM (+) form and TM (-) form of the mouse 4930451111Rik protein is 111 and 92 amino acids, respectively. The amino acids 1 to 77 are identical between TM (+) and (-) forms. Human C160RF92 has 110 amino acids match between mice (TM [+] form) and humans. The signal peptide region and putative transmembrane domain are indicated by the blue and red boxes, respectively. Black indicates a match in all sequences, whereas gray indicates a match in two sequences. (F) Average litter size of 4930451111Rik KO males. The average litter size was measured by the n

membrane protein (FIMP). Thus, the TM form of FIMP is important for the sperm-oocyte fusion process and functions independent of IZUMO1 in spermatozoa.

Analysis of Fimp-mCherry Tg Rescue Mice. To observe the localization of FIMP, we generated Fimp-mCherry Tg mice because no anti-FIMP antibodies are available for immunostaining and immunoblot analyses. The transgene expressed the mCherry-tagged TM form of FIMP (111 aa) under the testicular germ cell-specific Clgn promoter (Fig. 4A). To confirm that the Fimp-

mCherry transgene is functional, we obtained Tg mice on a Fimp KO (-1/-1) background (Fig. 4B). The infertile phenotype of Fimp KO males was fully rescued by the transgene  $(8.6 \pm 3.0, 0.4 \pm 0.9, \text{ and } 7.1 \pm 2.2 \text{ pups per plug in the wild-type [12 plugs], Fimp KO [15 plugs], and Tg rescued Fimp KO [23 plugs] males, respectively; <math>P < 0.01$ ; Fig. 4C). These results confirm that the TM form of FIMP regulates the sperm–oocyte fusion process. Next, we observed testicular sections and cauda epididymal spermatozoa in Fimp-mCherry Tg mice. Although spermatogenesis looked normal in testicular sections, the mCherry protein

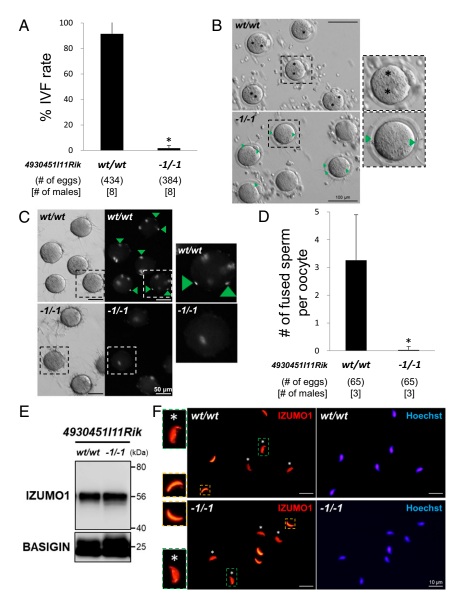


Fig. 2. In vitro fertilizing ability of 4930451111Rik KO spermatozoa. (A) IVF rates using wild-type and 4930451111Rik KO spermatozoa. Average fertilization rates of wild-type (wt) and 4930451111Rik KO (-1/-1) spermatozoa were 91.5  $\pm$  9.3% (397/434 eggs) and 1.8  $\pm$  2.2% (7/384 eggs), respectively. \*P < 0.01, Student's t test. (B) Observation of spermatozoa and oocytes 7 h after insemination. Although oocytes inseminated with wild-type (wt) spermatozoa formed pronuclei (indicated by asterisks), oocytes inseminated with 4930451111Rik KO (-1/-1) spermatozoa formed no visible pronuclei. Rather, spermatozoa were localized to the perivitelline space. 4930451111Rik KO spermatozoa could not properly fuse with the oocyte membrane. Pronuclei are indicated by asterisks. Green arrowheads indicate spermatozoa that remain in the perivitelline space after penetrating the zona pellucida. Enlarged images are indicated by dotted squares. (Scale bars: 100 µm.) (C) Observation of sperm-oocyte fusion in wild-type and 4930451111Rik KO spermatozoa. 4930451111Rik KO spermatozoa have impaired oocyte membrane fusion ability in vitro. Fused spermatozoa are indicated by green arrowheads. Enlarged images are indicated by dotted squares. (Scale bars: 50 µm.) (D) Average number of spermatozoa fused with the oocyte membrane in vitro. The number of fused spermatozoa in 4930451111Rik KO (-1/-1) mice  $(0.03 \pm 0.1$  spermatozoa; mean  $\pm$  SD) was significantly reduced compared with that of wild-type (wt) mice  $(3.3 \pm 1.6$  spermatozoa). \*P < 0.01, Student's t test. (E) Immunoblot analysis of IZUMO1 using sperm lysates collected from wild-type and 4930451111Rik KO mice. IZUMO1 is a sperm membrane protein essential for sperm-oocyte fusion. Although 4930451111Rik KO (-1/-1) spermatozoa showed impaired ability to fuse with the oocyte membrane, IZUMO1 was still present in 4930451111Rik KO spermatozoa. BASIGIN was used as a loading control. (F) Immunostaining of IZUMO1 in 4930451111Rik KO spermatozoa. IZUMO1 is used as a marker of the acrosome reaction. The acrosome reaction occurred in 4930451111Rik KO (-1/-1) spermatozoa. Acrosome reaction is determined by IZUMO1 localization (acrosome-intact spermatozoa: acrosomal cap pattern [orange dotted squares], acrosome-reacted spermatozoa: entire head pattern [green dotted squares]). Acrosome-reacted spermatozoa are indicated by asterisks. Enlarged images are indicated by dotted squares. (Scale bars: 10 μm.)

accumulated in the lumen of seminiferous tubules (SI Appendix, Fig. S44). Because most of the mCherry protein was concentrated in a cytoplasmic droplet on the sperm tail, it was difficult to check the localization of FIMP-mCherry on the sperm head by conventional fluorescence microscopic observation (SI Appendix, Fig. S4B). By confocal microscopic observation, we confirmed that FIMP-mCherry was present on all sperm heads, specifically on the equatorial segment, before the acrosome reaction (SI Appendix, Fig. S4C). However, both FIMP-mCherry fluorescent and nonfluorescent (39.0  $\pm$  3.5%) acrosome-reacted spermatozoa were observed (Fig. 4 D and E). These data indicated that the TM form of the FIMP protein is localized to the equatorial segment of the sperm head and can restore the infertility of Fimp KO males.

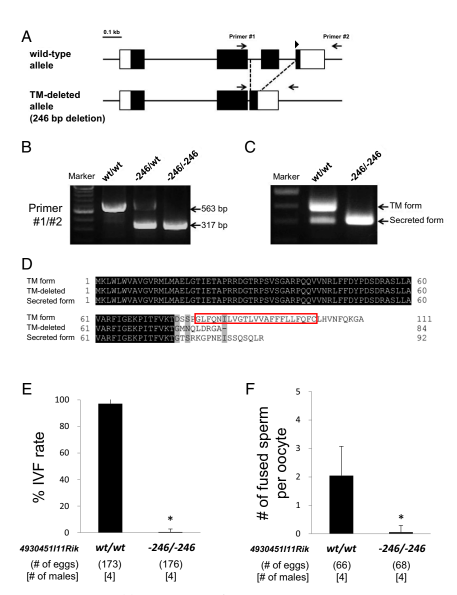


Fig. 3. Analysis of 4930451111Rik TM-deleted mice. (A) Targeting scheme of the 246-bp deletion in the 4930451111Rik locus. The third exon that encodes TM domain of 4930451111Rik was deleted by the CRISPR/Cas9 system. Arrowhead indicates gRNA targeting in the fourth exon. (Scale bar: 0.1 kb.) (B) Genotyping using PCR in 4930451111Rik TM-deleted mice. Both a 563-bp band representing the wild-type (wt/wt) allele and a 317-bp band representing the TM-deleted (-246/-246) allele were amplified by PCR using primers #1 and #2. (C) RT-PCR analysis of 4930451111Rik TM-deleted mouse testis. The coding region of 4930451111Rik was amplified by PCR in the testis from TM-deleted mice. The TM band (upper band in wild-type testis) disappeared from 4930451111Rik TM-deleted (-246/-246) mouse testis as expected. The lower band consisted of the secreted form and the in-frame mutation due to the 246-bp deletion as confirmed by direct sequencing (SI Appendix, Fig. S3B). (D) Amino acid sequence alignments of 4930451111Rik from wild-type and 4930451111Rik TM-deleted mice. A total of 90.5% of amino acids (76/84 amino acids) are identical with the secreted form of 4930451111Rik. Black indicates a match in all sequences, whereas gray indicates a match in two sequences. (E) IVF rates using wild-type and 4930451111Rik TM-deleted spermatozoa. Average fertilization rates of wild-type (wt) and 4930451111Rik TM-deleted (-246/-246) spermatozoa were 97.1 ± 2.5% (168/173 eggs) and 0.6 ± 1.3% (1/176 eggs), respectively. \*P < 0.01, Student's t test. (F) Average number of fused spermatozoa with oocyte membrane in vitro. The number of fused spermatozoa in 4930451111Rik TM-deleted (-246/-246) mice (0.1 ± 0.2 spermatozoa). \*P < 0.01, Student's t test.

In Vitro Cell-Oocyte Binding Assay. Cultured cells transiently expressing mouse IZUMO1 can tightly bind to the plasma membrane of ZP-free mouse oocytes (14). To confirm the relationship of FIMP and the oocyte membrane in sperm-oocyte interactions, we performed cell-oocyte binding assays using COS-7 and HEK293T cells expressing IZUMO1 and FIMP (SI Appendix, Figs. S4 D and E and S5). As shown in SI Appendix, Figs. S4E and S5B, FIMP-expressing cells could not bind to the oocyte surface, whereas IZUMO1-expressing cells could (4.0  $\pm$  0.4 cells per oocyte in IZUMO1-expressing cells). Moreover, coexpression of IZUMO1 and FIMP in transfected HEK293T cells

also did not enhance the binding ability to the oocyte membrane  $(4.5 \pm 0.5 \text{ and } 4.1 \pm 0.1 \text{ cells per oocyte in IZUMO1-expressing cells and IZUMO1 and FIMP coexpressing cells, respectively;$ *SI Appendix*, Fig. S5C). Thus, these results suggest that FIMP does not bind directly to the oocyte surface in sperm–oocyte fusion and does not directly promote IZUMO1 binding to JUNO on the oocyte plasma membrane.

#### Discussion

Before the advances in gene manipulation technology, oocyte and sperm proteins important for fertilization were discovered

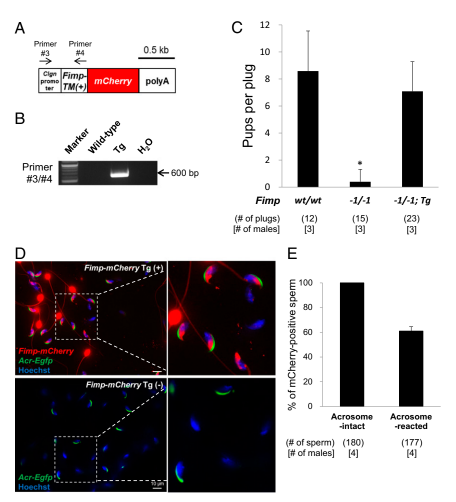


Fig. 4. Analysis of Fimp-mCherry Tg rescue mice. (A) Design of Fimp-mCherry transgene. Transgene designed in which the transmembrane isoform of FIMP and mCherry are expressed as a fused protein under the testicular germ cell-specific Clgn promoter (total, 1.9 kb). Arrows indicate primers used for genotyping. (Scale bar: 0.5 kb.) (B) Genotyping with PCR in Fimp-mCherry Tg mice. A 600-bp band representing the Tg allele was amplified by PCR using primers #3 and #4. (C) Average litter size of Fimp-mCherry Tg rescued Fimp KO male mice. The Fimp-mCherry transgene restored male infertility in Fimp KO mice. The average litter size was measured by the number of pups in each copulatory plug. The mean (± SD) litter size was 8.6 ± 3.0 in wild-type (wt) males, 0.4 ± 0.9 in Fimp KO (-1/-1) males, and 7.1 ± 2.2 in Fimp-mCherry Tg rescued Fimp KO (-1/-1; Tg) male mice. \*P < 0.01, Student's t test. (D) Confocal microscopic observation of Fimp-mCherry Tg mouse spermatozoa. The TM form of FIMP and mCherry fused protein (red signals) were localized to the equatorial segment of the sperm head in acrosome-intact spermatozoa (Acr-EGFP [green signals] positive). However, after the acrosome reaction (Acr-EGFP [green signals] negative), both fluorescent and nonfluorescent spermatozoa were observed. Red signals could not be detected in Acr-Egfp Tg mouse spermatozoa (without Fimp-mCherry transgene). (Scale bars: 10 μm.) (E) Fate of FIMP-mCherry protein on spermatozoa after the acrosome reaction. Although all of the acrosomeintact spermatozoa express FIMP-mCherry,  $39.0 \pm 3.5\%$  of acrosome-reacted spermatozoa lacked FIMP-mCherry in the sperm head (mean  $\pm$  SD). Acrosomereacted spermatozoa were observed after 3 h incubation.

via IVF experiments using biochemical approaches. However, in the last 2 decades, KO strategies have revealed that many "key" sperm proteins uncovered in vitro using antibody blocking or peptide competition experiments are not essential for in vivo fertilization (1, 5). In 2005, we found that the sperm membrane protein IZUMO1 is essential for spermatozoa to fuse with oocytes using the KO approach (12). Almost 10 years later, GPIanchored protein JUNO, the oocyte receptor of IZUMO1, was reported by Bianchi et al. (15). They identified JUNO by an avidity-based extracellular interaction screen (AVEXIS) assay that combines polymerized IZUMO1 and an iterative expression cloning approach using a mouse egg cDNA library. Recently, crystal structure analysis of the IZUMO1 and JUNO complex provided data on the amino acids critical in forming the intercellular bridge necessary for initiating sperm-oocyte fusion (45).

In this article, we focused on a testis-enriched gene 4930451111Rik and generated genetically modified mice that have a single gene disruption by the CRISPR/Cas9 system. We found that 4930451111Rik is necessary for the fertilizing ability of spermatozoa (Figs. 1F and 2A). Although 4930451111Rik KO mice produced normal-appearing epididymal spermatozoa, they showed impaired sperm-oocyte fusion (Fig. 2B and SI Appendix, Fig. S2D). Unexpectedly, although IZUMO1 is still intact, 4930451111Rik KO spermatozoa lacked the ability to fuse with oocytes (Fig. 2E). Thus, using CRISPR/Cas9, we have uncovered a KO mouse that produces sperm that cannot fuse with oocytes even though IZUMO1 remains. IZUMO1 changes its localization on the sperm head (from the acrosomal cap to the entire head) during the acrosome reaction, one of the most critical steps of fertilization (10, 11). This IZUMO1 translocation step is required for the ability of spermatozoa to fertilize eggs as evaluated by Spesp1 (sperm equatorial segment protein 1) and *Tssk6* (testis specific serine kinase 6) KO mice, which both showed reduced sperm fusion ability (46, 47). Since SPESP1 and TSSK6 are localized to the sperm head, abnormal sperm head formation may cause impaired translocation of IZUMO1 in these KO spermatozoa. However, IZUMO1 translocation was normal in 4930451111Rik KO spermatozoa (Fig. 2F). Our results indicate that 4930451I11Rik is a transmembrane protein that

regulates the sperm-oocyte fusion process via an IZUMO1-unrelated step.

Based on these results, we renamed 4930451I11Rik to fertilization influencing membrane protein (FIMP). Fimp is expressed as two variants in the mouse testis (Fig. 1 A and B). Direct sequencing and RT-PCR analysis confirmed the two forms: either localized to the transmembrane or secreted (Fig. 1D and SI *Appendix*, Fig. S1 A-C). In this study, we took advantage of the CRISPR/Cas9 system for quick and efficient genetic modification (48-51) of the Fimp allele in mice. To determine which isoform is essential for male fertility, we generated FIMP TMdeleted mice which introduced an in-frame mutation in the TM form (91.3% similarity of the secreted form; Fig. 3D). The TMdeleted mice showed the same phenotypes of Fimp KO mice in which both forms were disrupted (Fig. 3 E and F and SI Appendix, Fig. S3 C and D). Protein sequence alignment also shows that there is no secreted form among other mammalian species including humans (SI Appendix, Fig. S1D). Moreover, a transgene expressing the FIMP TM form restored male infertility in Fimp KO mice (Fig. 4C). Tg mouse analysis showed that FIMP is localized to the sperm equatorial segment membrane, which is attached to the oocyte membrane for fusion (SI Appendix, Fig. S4C). However, there were  $\sim 40\%$  of acrosome-reacted spermatozoa in which the FIMP-mCherry signals disappeared, in contrast to acrosome-intact spermatozoa, where their signals were localized to the equatorial segment (Fig. 4 D and E). The in vitro cell-oocyte binding assay showed that FIMP-expressing cells do not bind to the oocyte plasma membrane, in direct contrast to IZUMO1-expressing cells that directly bind to the oocyte surface (SI Appendix, Figs. S4E and S5B). Further studies are required to determine which population of acrosome-reacted spermatozoa (those that still express FIMP and/or those that do not) are required for the sperm-oocyte fusion process and how the TM form of FIMP (111 aa) regulates sperm fertilization. Although FIMP-expressing cells do not bind the oocyte in vitro, our results indicate that the TM form of FIMP is localized to the sperm equatorial segment to function during fusion with oocytes. FIMP may play a role after the initiation of the IZUMO1–JUNO complex for sperm-oocyte membrane binding, and/or FIMP could be part of a larger IZUMO1-independent sperm complex required for sperm-oocyte interactions.

In conclusion, our Fimp KO mice may prove useful in elucidating the physiological function of human FIMP (C16ORF92). In this study, however, we could not clarify the localization of endogenous FIMP in Izumo1 KO spermatozoa because anti-FIMP antibodies are not available for immunostaining and immunoblot analyses. Male infertility in Fimp KO mice may be caused by abnormalities in sperm protein or proteins other than IZUMO1. SPACA6 is a testis-specific transmembrane protein and was reported in a screening of male-infertile Tg mice that display sperm-oocyte fusion defects (52). However, no further phenotypic studies of Spaca6 KO mice have been reported. Thus, a targeted mutagenesis experiment that produces Spaca6 KO mice is required to reconfirm the reproductive phenotype and analyze whether it has a relationship to the Fimp KO phenotype. FIMP, IZUMO1, and SPACA6 are similar type I transmembrane proteins localized on the sperm head. FIMP and SPACA6 may also have an oocyte receptor counterpart, similar to IZUMO1. These proteins may function after the initiation of the IZUMO1-JUNO complex during sperm-oocyte fusion and/ or as part of a larger IZUMO1-independent fusion-required complex. As mentioned previously, further studies are needed to examine the cause of male infertility in Fimp KO mice and to clarify the interaction of IZUMO1, SPACA6, and FIMP. Our findings support a potential role of human FIMP in sperm function and could be used to develop infertility treatments as well as male-specific contraceptives.

#### Methods

Animals. All animal experiments were approved by the Animal Care and Use Committee of the Research Institute for Microbial Diseases, Osaka University. Human tissues were collected as nonhuman subject research by the Human Tissue Acquisition & Pathology Core at Baylor College of Medicine under the institutional review board-approved Protocol H-14435. Mice were maintain under a 12-h light/dark cycle (lights on from 8:00 to 20:00). Wild-type mice were purchased from CLEA Japan (Tokyo, Japan) and Japan SLC (Shizuoka, Japan). In this study, we generated genetically modified mouse lines, Fimp KO mice (B6D2-4930451111Rik<em10sb>); RBRC10114, and Fimp-mCherry and CAG/Acr-EGFP double Tg mice (B6D2-Tg(Clgn-4930451111Rik/mCherry)10sbTg(CAG/Acr-Egfp)C3-N01-FJ0020sb); RBRC10110. These were deposited to the RIKEN BioResource Research Center (https://mus.brc.riken.jp/en/) and the Center for Animal Resources and Development, Kumamoto University (http://card.medic.kumamoto-u.ac.jp/card/english/).

Generation of 4930451111Rik Mutant Mice with CRISPR/Cas9. 4930451111Rik KO mice were produced by microinjection of the pX330 plasmid (https:// www.addgene.org/42230/) into mouse embryos, as described previously (50, 51). A search for sgRNA and off-target sequences was performed using CRISPRdirect software (https://crispr.dbcls.jp/) (53). The sgRNA sequence used for microinjection were: 5'-GGAACCCGGCCCTCGGTTTC-3' for the second exon of 4930451111Rik (targeted for the common exon of two variants) and 5'-CCTCAGGAACTTCCAGAAAG-3' for the fourth exon of 4930451111Rik. Each sgRNA was injected into the pronuclei of fertilized eggs. The 2-cell stage embryos were transferred into the oviducts of pseudopregnant ICR females the next day. 4930451111Rik KO mice had a 1-bp deletion (5'-T-3') in the second exon. 4930451111Rik TM-deleted mice had a 246-bp deletion in the fourth exon. Both a 563-bp band as the wild-type allele and a 317-bp band as the TM-deleted allele were amplified by PCR. The primers used are listed in SI Appendix, Table S1. Detailed genotype information of mutant mouse lines is shown in the SI Appendix, Figs. S2A and S3A.

**Sperm–Oocyte Fusion Assay.** Sperm–oocyte fusion assay using mouse spermatozoa was performed as described previously (12). Briefly, ZP removal was performed by incubating the cumulus oocyte complex with collagenase (Sigma Aldrich, MO) at 100  $\mu$ g/mL for 10 min. Hoechst 33342 preloaded ZP-free oocytes were inseminated with 2  $\times$  10<sup>5</sup> cells/mL capacitated spermatozoa for 30 min. After fixing with 0.25% glutaraldehyde and washing with TYH medium, the number of fused spermatozoa was counted under a fluorescence microscope.

Generation of Fimp-mCherry Tg Rescue Mice. To make the Tg rescue construct, the TM form of mouse Fimp cDNA was amplified by PCR, using wild-type testis cDNA, fused with the amplified mCherry cDNA at the C terminus, and introduced downstream of the mouse Clgn promoter in pClgn1.1 vector (54). The Kpnl- and Sacl-digested transgene fragment (1.9 kb) was purified by gel electrophoresis and microinjected into the pronuclei of fertilized eggs from Fimp heterozygous KO mice using standard methods to create the Fimp-mCherry Tg rescue mouse line [B6D2-Tg(Clgn-4930451111RiklmCherry) 10sb]. For genotyping, a 600-bp band corresponding to the Tg allele was amplified by PCR, using primers as listed in SI Appendix, Table S1.

**Statistical Analysis.** Statistical analyses were performed using Student's t test inserted into Microsoft Excel after the data were tested for normality of distribution. Differences were considered significant at P < 0.01.

**Data Availability Statement.** All data are included in the paper or its supporting appendix.

ACKNOWLEDGMENTS. We thank Naoko Nagasawa and the Biotechnology Research and Development (nonprofit organization) for technical assistance and Ferheen Abbasi for critical reading of the manuscript. This work was supported by Ministry of Education, Culture, Sports, Science and Technology/Japan Society for the Promotion of Science KAKENHI grants (JP15H05573 and JP16KK0180 to Y.F.; JP18K16735 to Y.L.; JP18K14612 to T.N.; and JP25112007, JP17H01394, and JP19H05750 to M.I.); Japan Agency for Medical Research and Development Grant JP18gm5010001 to M.I.; Takeda Science Foundation grants to Y.F. and M.I.; the Chubei Itoh Foundation grant to Y.F.; Senri Life Science Foundation grant to Y.F.; Intramural Research Fund (30-2-5 and 31-6-3) for Cardiovascular Diseases of National Cerebral and Cardiovascular Center to Y.F.; the Eunice Kennedy Shriver National Institute of Child Health and Human Development grants (R01HD088412 and P01HD087157 to M.M.M. and M.I.); and the Bill & Melinda Gates Foundation grant (INV-001902 to M.M.M. and M.I.). The Human Tissue Acquisition & Pathology Core is funded through the P30 Cancer Center Support Grant (NCI CA125123).

- 2. R. Yanagimachi, "Mammalian fertilization." The Physiology of Reproduction, E. Knobil, J. D. Neill, Eds. (Raven Press, LTD., New York, ed. 2, 1994), pp. 189-317.
- 3. M. Okabe, Sperm-egg interaction and fertilization: Past, present, and future. Biol. Reprod. 99, 134-146 (2018).
- 4. Y. Fujihara, H. Miyata, M. Ikawa, Factors controlling sperm migration through the oviduct revealed by gene-modified mouse models. Exp. Anim. 67, 91-104 (2018).
- 5. M. M. Matzuk, D. J. Lamb, The biology of infertility: Research advances and clinical challenges. Nat. Med. 14, 1197-1213 (2008).
- 6. K. Miyado et al., Requirement of CD9 on the egg plasma membrane for fertilization. Science 287, 321-324 (2000).
- 7. F. Le Naour, E. Rubinstein, C. Jasmin, M. Prenant, C. Boucheix, Severely reduced female fertility in CD9-deficient mice. Science 287, 319-321 (2000).
- 8. K. Kaji et al., The gamete fusion process is defective in eggs of Cd9-deficient mice. Nat. Genet. 24, 279-282 (2000).
- 9. K. E. Runge et al., Oocyte CD9 is enriched on the microvillar membrane and required for normal microvillar shape and distribution. Dev. Biol. 304, 317-325 (2007).
- 10. Y. Satouh, N. Inoue, M. Ikawa, M. Okabe, Visualization of the moment of mouse sperm-egg fusion and dynamic localization of IZUMO1. J. Cell Sci. 125, 4985–4990 (2012).
- 11. M. Okabe, K. Takada, T. Adachi, Y. Kohama, T. Mimura, Inconsistent reactivity of an anti-sperm monoclonal antibody and its relationship to sperm capacitation. J. Reprod. Immunol. 9, 67-70 (1986).
- 12. N. Inoue, M. Ikawa, A. Isotani, M. Okabe, The immunoglobulin superfamily protein Izumo is required for sperm to fuse with eggs. Nature 434, 234-238 (2005).
- 13. S. A. Young et al., CRISPR/Cas9-mediated mutation revealed cytoplasmic tail is dispensable for IZUMO1 function and male fertility. Reproduction 152, 665-672 (2016).
- 14. N. Inoue et al., Molecular dissection of IZUMO1, a sperm protein essential for spermegg fusion. Development 140, 3221-3229 (2013).
- 15. E. Bianchi, B. Doe, D. Goulding, G. J. Wright, Juno is the egg Izumo receptor and is essential for mammalian fertilization. Nature 508, 483-487 (2014).
- U. Ohto et al., Structure of IZUMO1-JUNO reveals sperm-oocyte recognition during mammalian fertilization. Nature 534, 566-569 (2016).
- 17. K. Nishimura et al., The structure of sperm Izumo1 reveals unexpected similarities with Plasmodium invasion proteins. Curr. Biol. 26, R661-R662 (2016).
- 18. K. Kato et al., Structural and functional insights into IZUMO1 recognition by JUNO in mammalian fertilization. Nat. Commun. 7, 12198 (2016).
- L. Han et al., Divergent evolution of vitamin B9 binding underlies Juno-mediated adhesion of mammalian gametes. Curr. Biol. 26, R100-R101 (2016).
- 20. H. Aydin, A. Sultana, S. Li, A. Thavalingam, J. E. Lee, Molecular architecture of the human sperm IZUMO1 and egg JUNO fertilization complex. Nature 534, 562-565 (2016)
- 21. N. Inoue, Y. Hagihara, D. Wright, T. Suzuki, I. Wada, Oocyte-triggered dimerization of sperm IZUMO1 promotes sperm-egg fusion in mice. Nat. Commun. 6, 8858 (2015).
- D. Mashiko et al., Feasibility for a large scale mouse mutagenesis by injecting CRISPR/ Cas plasmid into zygotes. Dev. Growth Differ. 56, 122-129 (2014).
- 23. H. Wang et al., One-step generation of mice carrying mutations in multiple genes by CRISPR/Cas-mediated genome engineering. Cell 153, 910-918 (2013).
- 24. T. Noda et al., Nine genes abundantly expressed in the epididymis are not essential for male fecundity in mice. Andrology 7, 644–653 (2019).
- 25. Y. Lu et al., CRISPR/Cas9-mediated genome editing reveals 30 testis-enriched genes dispensable for male fertility in mice. Biol. Reprod. 101, 501-511 (2019).
- 26. R. J. Holcomb et al., The testis-specific serine proteases Prss44, Prss46, and Prss54 are dispensable for mouse fertility. Biol. Reprod. 102, 84-91 (2020).
- 27. H. Miyata et al., Genome engineering uncovers 54 evolutionarily conserved and testisenriched genes that are not required for male fertility in mice. Proc. Natl. Acad. Sci. U.S.A. 113, 7704-7710 (2016).
- 28. Y. Fujihara et al., Identification of multiple male reproductive tract-specific proteins that regulate sperm migration through the oviduct in mice. Proc. Natl. Acad. Sci. U.S.A. 116, 18498-18506 (2019).

- 29. Y. Fujihara, A. Oji, K. Kojima-Kita, T. Larasati, M. Ikawa, Co-expression of sperm membrane proteins CMTM2A and CMTM2B is essential for ADAM3 localization and male fertility in mice. J. Cell Sci. 131, jcs221481 (2018).
- 30. F. Abbasi et al., RSPH6A is required for sperm flagellum formation and male fertility in mice. J. Cell Sci. 131, jcs221648 (2018).
- 31. Y. Fujihara, A. Oji, T. Larasati, K. Kojima-Kita, M. Ikawa, Human globozoospermiarelated gene Spata16 is required for sperm formation revealed by CRISPR/Cas9-Mediated mouse models. Int. J. Mol. Sci. 18, E2208 (2017).
- 32. J. M. Castaneda et al., TCTE1 is a conserved component of the dynein regulatory complex and is required for motility and metabolism in mouse spermatozoa. Proc. Natl. Acad. Sci. U.S.A. 114, E5370-E5378 (2017).
- 33. S. A. Young et al., CABYR is essential for fibrous sheath integrity and progressive motility in mouse spermatozoa. J. Cell Sci. 129, 4379-4387 (2016).
- 34. H. Miyata et al., Sperm calcineurin inhibition prevents mouse fertility with implications for male contraceptive. Science 350, 442-445 (2015).
- 35. Y. Fujihara, M. Okabe, M. Ikawa, GPI-anchored protein complex, LY6K/TEX101, is required for sperm migration into the oviduct and male fertility in mice. Biol. Reprod. 90, 60 (2014).
- 36. Y. Fujihara et al., Expression of TEX101, regulated by ACE, is essential for the production of fertile mouse spermatozoa. Proc. Natl. Acad. Sci. U.S.A. 110, 8111-8116 (2013).
- 37. R. Yamaguchi, Y. Fujihara, M. Ikawa, M. Okabe, Mice expressing aberrant spermspecific protein PMIS2 produce normal-looking but fertilization-incompetent spermatozoa, Mol. Biol. Cell 23, 2671-2679 (2012).
- K. Tokuhiro, M. Ikawa, A. M. Benham, M. Okabe, Protein disulfide isomerase homolog PDILT is required for quality control of sperm membrane protein ADAM3 and male fertility. Proc. Natl. Acad. Sci. U.S.A. 109, 3850-3855 (2012).
- 39. Y. Fujihara et al., SPACA1-deficient male mice are infertile with abnormally shaped sperm heads reminiscent of globozoospermia. Development 139, 3583-3589 (2012).
- 40. M. Ikawa et al., Calsperin is a testis-specific chaperone required for sperm fertility. J. Biol. Chem. 286, 5639-5646 (2011).
- 41. Y. N. Lin, A. Roy, W. Yan, K. H. Burns, M. M. Matzuk, Loss of zona pellucida binding proteins in the acrosomal matrix disrupts acrosome biogenesis and sperm morphogenesis. Mol. Cell. Biol. 27, 6794-6805 (2007).
- 42. M. P. Greenbaum et al., TEX14 is essential for intercellular bridges and fertility in male mice. Proc. Natl. Acad. Sci. U.S.A. 103, 4982-4987 (2006).
- 43. M. Ikawa et al., The putative chaperone calmegin is required for sperm fertility. Nature 387, 607-611 (1997).
- 44. K. Nozawa, Y. Satouh, T. Fujimoto, A. Oji, M. Ikawa, Sperm-borne phospholipase C zeta-1 ensures monospermic fertilization in mice. Sci. Rep. 8, 1315 (2018)
- 45. Y. Satouh, M. Ikawa, New insights into the molecular events of mammalian fertilization. Trends Biochem. Sci. 43, 818-828 (2018).
- 46. Y. Fujihara et al., Sperm equatorial segment protein 1, SPESP1, is required for fully fertile sperm in mouse. J. Cell Sci. 123, 1531-1536 (2010).
- 47. J. Sosnik et al., Tssk6 is required for Izumo relocalization and gamete fusion in the mouse. J. Cell Sci. 122, 2741-2749 (2009).
- T. Noda, A. Oji, M. Ikawa, Genome editing in mouse zygotes and embryonic stem cells by introducing SgRNA/Cas9 expressing plasmids. Methods Mol. Biol. 1630, 67-80 (2017)
- 49. A. Oji et al., CRISPR/Cas9 mediated genome editing in ES cells and its application for chimeric analysis in mice. Sci. Rep. 6, 31666 (2016).
- 50. Y. Fujihara, M. Ikawa, CRISPR/Cas9-based genome editing in mice by single plasmid injection. Methods Enzymol. 546, 319-336 (2014).
- 51. D. Mashiko et al., Generation of mutant mice by pronuclear injection of circular plasmid expressing Cas9 and single guided RNA. Sci. Rep. 3, 3355 (2013).
- 52. D. Lorenzetti et al., A transgenic insertion on mouse chromosome 17 inactivates a novel immunoglobulin superfamily gene potentially involved in sperm-egg fusion. Mamm. Genome 25, 141-148 (2014).
- 53. Y. Naito, K. Hino, H. Bono, K. Ui-Tei, CRISPRdirect: Software for designing CRISPR/Cas guide RNA with reduced off-target sites. Bioinformatics 31, 1120-1123 (2015).
- 54. M. Ikawa et al., Calmegin is required for fertilin alpha/beta heterodimerization and sperm fertility. Dev. Biol. 240, 254-261 (2001).